The role of mechano-electric feedbacks and hemodynamic coupling in scar-related ventricular tachycardia

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Abstract
Mechano-electric feedbacks (MEFs), which model how mechanical stimuli are transduced into electrical signals, have received sparse investigation by considering electromechanical simulations in simplified scenarios. In this paper, we study the effects of different MEFs modeling choices for myocardial deformation and nonselective stretch-activated channels (SACs) in the monodomain equation. We perform numerical simulations during ventricular tachycardia (VT) by employing a biophysically detailed and anatomically accurate 3D electromechanical model for the left ventricle (LV) coupled with a 0D closed-loop model of the cardiocirculatory system. We model the electromechanical substrate responsible for scar-related VT with a distribution of infarct and peri-infarct zones. Our mathematical framework takes into account the hemodynamic effects of VT due to myocardial impairment and allows for the classification of their hemodynamic nature, which can be either stable or unstable. By combining electrophysiological, mechanical and hemodynamic models, we observe that all MEFs may alter the propagation of the action potential and the morphology of the VT. In particular, we notice that the presence of myocardial deformation in the monodomain equation may change the VT basis cycle length and the conduction velocity but do not affect the hemodynamic nature of the VT. Finally, nonselective SACs may affect wavefront stability, by possibly turning a hemodynamically stable VT into a hemodynamically unstable one and vice versa.

Keywords: Cardiac electromechanics, Numerical simulations, Mechano-electric feedback, Stretch-activated channels, Ventricular tachycardia

1 Introduction
Cardiac arrhythmias result from an irregular electrical activity of the human heart. Among them, ventricular tachycardia (VT), which manifests with an accelerated heart rate, is one of the most life-threatening rhythm disorders. VT may be classified as either hemodynamically stable or unstable, depending on the capability of the heart to effectively pump blood in the circulatory system. In the former case antiarrhythmic drugs are generally employed, while in the latter case cardioversion is needed [21]. According to the specific pathogenesis, the stability of the VT remains the same or changes over time. Moreover, it may also degenerate towards ventricular fibrillation (VF), a life-threatening condition in which the ventricular activity is fully disorganized and chaotic, leading to heart failure [62].

In the clinical framework, these pathological scenarios can be hardly ever fully investigated and predicted for all patients. For this reason, biophysically detailed computational heart models could be used to provide a deeper understanding of the hemodynamic response to VT and to characterize the electromechanical substrate leading to dangerous arrhythmias.
While electrophysiological simulations are well-established for scar-related VT identification and treatment on human ventricles [1, 10, 19, 46], patient-specific electromechanical models coupled with closed-loop cardiovascular circulation have been just recently used for the first time to enhance our knowledge on VT [61]. Indeed, on one side, the physiological processes by which the mechanical behavior alters the electrical activity of the human heart, known as mechano-electric feedbacks (MEFs), are relevant and not fully elucidated [13, 31, 33, 64, 65, 68, 70]. On the other hand, the identification of the hemodynamic nature of the VT has significant clinical implications [61].

The goal of this study is threefold: (1) to combine electrophysiology, activation, mechanics and hemodynamics in several numerical simulations of scar-related VT, with the aim of uncovering the roles of myocardial deformation and the recruitment of SACs on VT stability; (2) to show that our computational model effectively reproduces both hemodynamically stable and hemodynamically unstable VT; (3) to capture relevant microscopic mechanisms during VT, such as the incomplete relaxation of sarcomeres, given the biophysical detail of our electromechanical model.

2 Mathematical models

We provide an overview of the 3D cardiac electromechanical model coupled with a 0D closed-loop model of the cardiovascular system (see [55, 61]).

We depict in Figure 1 the computational domain $\Omega_0$, which represents a human left ventricle (LV) taken from the Zygote Solid 3D heart model [28]. Its boundary $\partial \Omega_0$ is partitioned into epicardium $\Gamma_{\text{epi}}^0$, endocardium $\Gamma_{\text{endo}}^0$ and base $\Gamma_{\text{base}}^0$. We recall that $\partial \Omega_0 = \Gamma_{\text{epi}}^0 \cup \Gamma_{\text{endo}}^0 \cup \Gamma_{\text{base}}^0$.

2.1 3D-0D closed-loop electromechanical model

We consider a multiphysics and multiscale 3D-0D mathematical framework comprised of four different core models, namely cardiac electrophysiology ($E$) [12, 14, 36, 40], mechanical activation ($A$) [34, 39, 51, 58, 59], passive mechanics ($M$) [23, 24, 27, 41] and blood circulation ($C$) [3, 4, 8, 26, 32, 50, 55]. The volume conservation constraint ($V$) defines the coupling condition between the LV cardiac electromechanics and the remaining part of cardiovascular system. [55]. All these models represent several physiological processes, ranging from the cellular level to the organ scale.

Given the computational domain $\Omega_0$ and the time interval $t \in (0, T]$, the mathematical model reads:
by triggering the action potential at specific locations of the myocardium. The reaction terms
the transmembrane capacitance per unit area. The applied current \( I \) (specified by the ionic model at hand) couple the action potential propagation to the cellular dynamics. Specifically,

\begin{equation}
(1a) \quad \frac{\partial u}{\partial t} + I_{\text{ion}}(u, w) + I_{\text{SAC}}(u, F) = -\nabla \cdot (JF^{-1}DF^{-T}\nabla u) = J_{\chi_m}I_{\text{app}}(t) \quad \text{in } \Omega_0 \times (0, T], \quad \text{in } \Omega_0 \times (0, T], \quad \text{on } \partial \Omega_0 \times (0, T],
\end{equation}

with a suitable ionic model for the human ventricular action potential \([9, 14, 37, 69]\).

We model the electrical activity of the myocardium by means of Eq. (1a), that is the monodomain equation coupled
2.1.1 Electrophysiology

\begin{equation}
(1b) \quad \frac{\partial s}{\partial t} = K \left( s, [\text{Ca}^{2+}], SL, \frac{\partial SL}{\partial t} \right) \quad \text{in } \Omega_0 \times (0, T],
\end{equation}

\begin{equation}
(1c) \quad \rho_s \frac{\partial^2 d}{\partial t^2} - \nabla \cdot P(d, T_a(s, SL)) = 0 \quad \text{in } \Omega_0 \times (0, T], \quad \text{on } \Gamma_0^{\text{epi}} \times (0, T], \quad \text{on } \Gamma_0^{\text{endo}} \times (0, T], \quad \text{on } \Gamma_0^{\text{base}} \times (0, T],
\end{equation}

\begin{equation}
(1d) \quad \frac{dc(t)}{dt} = Z(t, c(t), p_{LV}(t)) \quad \text{for } t \in (0, T], \quad \text{for } t \in (0, T],
\end{equation}

\begin{equation}
(1e) \quad V_{LV}(c(t)) = V_{LV}^{3D}(d(t)) \quad \text{for } t \in (0, T],
\end{equation}

in which the model unknowns are:

\begin{align}
\begin{aligned}
& u: \Omega_0 \times (0, T] \to \mathbb{R}, \quad w: \Omega_0 \times (0, T] \to \mathbb{R}^n_w, \\
& s: \Omega_0 \times (0, T] \to \mathbb{R}^{n_s}, \quad d: \Omega_0 \times (0, T] \to \mathbb{R}^3, \quad c: (0, T] \to \mathbb{R}^{n_c},
\end{aligned}
\end{align}

(2)

where \( u \) is the transmembrane potential, \( w \) is the vector containing both gating and ionic variables, \( s \) are the
states variables of the active force generation model, \( d \) represents the tissue mechanical displacement, \( c \) defines the state vector of the circulation model (including pressures, volumes and fluxes of the different compartments of the cardiocirculatory system) and \( p_{LV} \) and \( p_{RV} \) are the left and right ventricular pressure, respectively. Finally, all
variables are endowed with suitable initial conditions in \( \Omega_0 \times \{0\} \):

\begin{align}
\begin{aligned}
& u = u_0, \quad w = w_0, \quad s = s_0, \quad d = d_0, \quad \frac{\partial d}{\partial t} = \dot{d}_0, \quad c = c_0.
\end{aligned}
\end{align}

(3)

2.1.1 Electrophysiology \((\mathcal{E})\)

We model the electrical activity of the myocardium by means of Eq. (1a), that is the monodomain equation coupled
with a suitable ionic model for the human ventricular action potential \([9, 14, 37, 69]\).

In the electrophysiological model \((\mathcal{E})\), \( \chi_m \) is the surface area-to-volume ratio of cardiomyocytes, \( C_m \) represents the transmembrane capacitance per unit area. The applied current \( I_{\text{app}} \) mimics the effect of the Purkinje network \([18, 35, 71]\) by triggering the action potential at specific locations of the myocardium. The reaction terms \( I_{\text{ion}} \) and \( H \) (specified by the ionic model at hand) couple the action potential propagation to the cellular dynamics. Specifically,
we use the ten Tusscher-Panfilov ionic model (TTP06), which is able to accurately describe ions dynamics across the cell membrane [69]. Furthermore, Eq. (1a) is equipped with homogeneous Neumann boundary conditions for \( u \) at the boundary \( \partial \Omega_0 \), which defines the condition of electrically isolated domain for \( \Omega_0 \). \( D \) represents the conductivity tensor, that reads:

\[
D = \eta \sigma_1 \frac{\mathbf{F}_0 \otimes \mathbf{F}_0}{\| \mathbf{F}_0 \|^2} + \eta \sigma_\tau \frac{\mathbf{F}_{\mathbf{S}0} \otimes \mathbf{F}_{\mathbf{S}0}}{\| \mathbf{F}_{\mathbf{S}0} \|^2} + \eta \sigma_n \frac{\mathbf{F}_{\mathbf{n}} \otimes \mathbf{F}_{\mathbf{n}}}{\| \mathbf{F}_{\mathbf{n}} \|^2},
\]

where \( \sigma_1, \sigma_\tau, \sigma_n \) are the longitudinal, transversal and normal conductivities, respectively. The parameter \( \eta = \eta(\mathbf{x}) \in [0, 1] \) takes into account the effect of scars, grey zones and non-remodeled regions. This parameter is also incorporated inside the formulation of the TTP06 model, as described in [61].

The deformation tensor \( \mathbf{F} = \mathbf{I} + \nabla \mathbf{d} \) and its determinant \( J = \det(\mathbf{F}) \) are needed to perform the mechano-electric coupling [67]. Indeed, we model the so called mechano-electric feedbacks (MEFs) [17, 25]. The geometry-mediated MEFs incorporate the effects of displacement \( \mathbf{d} \) on the cardiac tissue, while other physiological processes act at the level of single cardiomyocytes [36, 68]. Among them, some examples are selective (e.g. \( K^+ \)-permeable) or nonselective SACs and intracellular calcium [\( Ca^{2+} \)], binding to sarcolemmal buffers, the latter requiring more sophisticated ventricular ionic models [5, 66]. In this paper we model nonselective SACs by means of the following formulation [36]:

\[
I_{\text{SAC}}(u, \mathbf{F}) = G_s(\| \mathbf{F}_0 \|) - 1_+ (u - u_{\text{rev}}),
\]

where \( G_s \) and \( u_{\text{rev}} \) represent the conductance of the channels and the reversal potential, respectively. SACs may alter the shape of the action potential (AP) by lengthening or shortening its duration (APD) and by generating higher resting potentials. This may induce early or delayed afterdepolarizations (EADs or DADs) and premature excitation.

Both geometry-mediated MEFs and nonselective SACs are generally known to be pro-arrhythmic in pathological scenarios, because they increase the likelihood of having extra stimuli during cardiac contraction and relaxation [68].

In this work we consider Eq. (1a) with several degrees of complexity to assess similarities and differences in the outcomes of the electromechanical simulations during VT. We report in Tab. 1 the mathematical models and the parametrizations that we consider. In particular, the choice of three different mathematical models for the geometry-mediated MEFs, namely \((\mathcal{E}_g\text{-MEF-minimal}), (\mathcal{E}_g\text{-MEF-enhanced}) \) and \((\mathcal{E}_g\text{-MEF-full})\), is motivated by the numerous formulations of this type of feedback that can be found in the literature [11, 15, 36, 49]. Indeed, we range from minimal to complete inclusion of geometry-mediated MEFs in the monodomain equation.

### 2.1.2 Activation \((\alpha')\)

To model how the calcium wave following the AP triggers a series of chemo-mechanical reactions within sarcomeres, resulting in the generation of an active force in the muscle, we use the mean-field version of the model proposed in [51], henceforth denoted by RDQ20-MF. This mathematical model is based on a biophysically accurate description of regulatory and contractile proteins and their dynamics. Thanks to suitable dimensionality reduction techniques, the dynamics of stochastic processes underlying both chemical and mechanical microscale transitions are described in only 20 ODEs (that is \( s(t) \in \mathbb{R}^{20} \)). The computational cost of this model is thus comparable to that of phenomenological models [34, 38, 57], while providing a biophysically detailed description that is consistent with the level of detail and mechanistic understanding of the ionic model needed for the present study.

The inputs of the RDQ20-MF model are the intracellular calcium concentration [\( Ca^{2+} \)] coming from \((\mathcal{E})\), the sarcomere length (denoted by \( SL \)) and its time derivative (the latter allows to account for the so-called force-velocity relationship [30]). The variable \( SL \) is obtained as \( SL = SL_0 \sqrt{I_{AF}} \), where \( SL_0 \) is the sarcomere slack length and the fourth invariant \( I_{AF} = \mathbf{F}_{0} \cdot \mathbf{F}_{0} \) measures the tissue stretch in the fiber direction. Finally, the output of the RDQ20-MF model is the active tension generated at the microscale, that can be obtained as \( T_A(s, SL) \).

Differently from [61], here we consider a biophysically detailed active stress model. For this reason, we do not need to put the parameter \( \eta \) inside its parametrization. Indeed, the active force generation mechanisms are properly
Table 1: Modeling choices for the monodomain equation that have been used in this paper. We consider different parametrizations for $\mathcal{I}_{\text{SAC}}(u, F)$ in terms of $G_s$ and $u_{\text{rev}}$. $D_I$ indicates the conductivity tensor in Eq. (4) with $F = I$.

| Model name | Equation |
|------------|----------|
| ($\varepsilon$) | $\chi_m \left[ C_m \frac{\partial u}{\partial t} + \mathcal{I}_{\text{ion}}(u, w) \right] - \nabla \cdot (D_1 \nabla u) = \chi_m \mathcal{I}_{\text{app}}(t)$ |
| ($\varepsilon_{\text{gMEF-minimal}}$) | $\chi_m \left[ C_m \frac{\partial u}{\partial t} + \mathcal{I}_{\text{ion}}(u, w) \right] - \nabla \cdot (J F^{-1} D_1 F^{-T} \nabla u) = \chi_m \mathcal{I}_{\text{app}}(t)$ |
| ($\varepsilon_{\text{gMEF-enhanced}}$) | $\chi_m \left[ C_m \frac{\partial u}{\partial t} + \mathcal{I}_{\text{ion}}(u, w) \right] - \nabla \cdot (J F^{-1} D F^{-T} \nabla u) = \chi_m \mathcal{I}_{\text{app}}(t)$ |
| ($\varepsilon_{\text{gMEF-full}}$) | $J \chi_m \left[ C_m \frac{\partial u}{\partial t} + \mathcal{I}_{\text{ion}}(u, w) \right] - \nabla \cdot (J F^{-1} D F^{-T} \nabla u) = J \chi_m \mathcal{I}_{\text{app}}(t)$ |
| ($\varepsilon_{\text{SAC}}$) | $\chi_m \left[ C_m \frac{\partial u}{\partial t} + \mathcal{I}_{\text{ion}}(u, w) + \mathcal{I}_{\text{SAC}}(u, F) \right] - \nabla \cdot (D_1 \nabla u) = \chi_m \mathcal{I}_{\text{app}}(t)$ |
| ($\varepsilon_{\text{gMEF-full, SAC}}$) | $J \chi_m \left[ C_m \frac{\partial u}{\partial t} + \mathcal{I}_{\text{ion}}(u, w) + \mathcal{I}_{\text{SAC}}(u, F) \right] - \nabla \cdot (J F^{-1} D F^{-T} \nabla u) = J \chi_m \mathcal{I}_{\text{app}}(t)$ |

handled by the active stress model, which receives different intracellular calcium waves from ($\varepsilon$) according to the specific area of the myocardium (scar, grey zone or healthy) and provides physiological values of active tension in all cases.

2.1.3 Mechanics ($\varepsilon$)

We employ the momentum conservation equation reported in Eq. (1c) to model the dynamics of the displacement $\mathbf{d}$ of the myocardium. We also consider the hyperelasticity assumption, so that the strain energy function can be differentiated with respect to the deformation tensor $\mathbf{F}$ to obtain $\mathbf{P}$ [23, 41].

In the mechanical model ($\varepsilon$), $\rho_s$ represents the density of the myocardium. The Piola-Kirchhoff stress tensor $\mathbf{P} = \mathbf{P}(\mathbf{d}, T_s)$ is additively decomposed according to:

$$
\mathbf{P}(\mathbf{d}, T_s) = \frac{\partial \mathcal{W}(\mathbf{F})}{\partial \mathbf{F}} + T_s(s, SL) \frac{\mathbf{F} f_0 \otimes f_0}{\sqrt{J_{tf}}},
$$

where the first term stands as the passive part of the tensor $\mathbf{P}$, while the latter as the active one; $\mathcal{W} : \text{Lin}^+ \rightarrow \mathbb{R}$ is the strain energy density function, $T_s(s, SL)$ is the active tension, provided by the activation model ($\varepsilon$).

To model the passive behaviour of cardiac tissue, we employ the orthotropic Guccione constitutive law [24], whose strain energy function is defined as:

$$
\mathcal{W} = \frac{\kappa}{2} (J - 1) \log(J) + \frac{a}{2} (e^Q - 1),
$$

where the first term is the volumetric energy with the bulk modulus $\kappa$, which penalizes large variation of volume to enforce a weakly incompressible behavior [20, 42], and the latter is the deviatoric energy where $a$ is the stiffness scaling parameter and the exponent $Q$ reads:

$$
Q = b_0 E_{in}^2 + b_a E_{ns}^2 + b_m E_{nm}^2 + b_n \left( E_{in}^2 + E_{ns}^2 \right) + b_0 \left( E_{in}^2 + E_{nm}^2 \right) + b_n \left( E_{in}^2 + E_{ns}^2 \right),
$$

5
in which $E_{ij} = E_{ij0} \cdot j_0$ for $i, j \in \{f, s, n\}$ are the entries of $E = \frac{1}{2} (C - I)$, i.e. the Green-Lagrange strain energy tensor, being $C = F^T F$ the right Cauchy-Green deformation tensor.

To model the mechanical effects of the pericardial sac [22, 43, 63], we impose at the epicardial boundary $\Gamma_0^{\text{epi}}$ a generalized Robin boundary condition $P(d, T_0)N = K^{\text{epi}}d + C^{\text{epi}} \frac{\partial d}{\partial n}$ by defining the tensors $K^{\text{epi}} = K^{\text{epi}}(N \otimes N - I) - K^{\text{epi}}(N \otimes N - I) - K^{\text{epi}}(N \otimes N)$, where $K^{\text{epi}}$, $K^{\text{epi}}$, $C^{\text{epi}}$, $C^{\text{epi}} \in \mathbb{R}^+$ are the stiffness and viscosity parameters of the epicardial tissue in the normal and tangential directions, respectively. Normal stress boundary conditions were imposed at the endocardium of the LV $\Gamma_0^{\text{endo}}$, in which $p_{LV}(t)$ is the pressure exerted by the blood in the LV, modeled by means of the 0D closed-loop circulation model [55]. To take into account the effect of the neglected part, over the basal plane, on the ventricular domain, we set on $\Gamma_0^{\text{base}}$ the energy consistent boundary condition $P(d, T_0)N = |F^{-T}N|v_{LV}^0(t)$ [52], where:

$$v_{LV}^0(t) = \frac{\int_{\Gamma_0^{\text{endo}}} p_{LV}(t) JF^{-T}N d\Gamma_0}{\int_{\Gamma_0^{\text{base}}} |F^{-T}N| d\Gamma_0}.$$  \hfill (9)

2.1.4 Blood circulation ($\mathcal{C}$) and coupling conditions ($\mathcal{Y}$)

We model the blood circulation through the entire cardiovascular system by means of a closed-loop model recently proposed in [55]. In the 0D closed-loop model, systemic and pulmonary circulations are modeled with RLC circuits, heart chambers are described by time-varying elastance elements and non-ideal diodes stand for the heart valves [55].

We represent the circulation core model ($\mathcal{C}$) with a system of ODEs, which is expressed by Eq. (1d), where $Z$ is a proper function (defined in [55]) and $c(t)$ includes pressures, volumes and fluxes of the different compartments composing the vascular network:

$$c(t) = (V_{LA}(t), V_{LV}(t), V_{RA}(t), V_{RV}(t), p_{LV}^{\text{SYS}}(t), p_{LV}^{\text{VEN}}(t), p_{AR}^{\text{SYS}}(t), p_{AR}^{\text{VEN}}(t), p_{PUL}^{\text{SYS}}(t), p_{PUL}^{\text{VEN}}(t))^T.$$  

Here $V_{LA}$, $V_{RA}$, $V_{LV}$ and $V_{RV}$ refer to the volumes of left atrium, right atrium, left ventricle and right ventricle, respectively; $p_{AR}^{\text{SYS}}$, $p_{AR}^{\text{VEN}}$, $p_{LV}^{\text{SYS}}$, $p_{LV}^{\text{VEN}}$, $p_{LV}^{\text{PUL}}$, $p_{AR}^{\text{PUL}}$, $p_{PUL}^{\text{SYS}}$ and $p_{PUL}^{\text{VEN}}$ express pressures and flow rates of the systemic and pulmonary circulation (arterial and venous). For the complete mathematical description of the 0D circulation lumped model we refer to [55]. To couple the 0D circulation model ($\mathcal{C}$) with the 3D LV model, given by ($\mathcal{C}$)–($\mathcal{M}$)–($\mathcal{Y}$), we follow the same strategy proposed in [55]. In particular, we replace the time-varying elastance elements representing the LV in ($\mathcal{C}$) with its corresponding 3D electromechanical description and we introduce the coupling condition ($\mathcal{Y}$) where:

$$V_{LV}^{\text{3D}}(d(t)) = \int_{\Gamma_0^{\text{endo}}} J(t) ((h \otimes h) (x + d(t) - b)) \cdot F^{-T}(t)N d\Gamma_0$$

in which $h$ is a vector orthogonal to the LV centerline (i.e. lying on the ventricular base) and $b$ lays inside the LV [55].

Due to ($\mathcal{Y}$), in the 3D-0D coupled model (1), $p_{LV}(t)$ is not determined by the 0D circulation model (1d), but rather acts as Lagrange multipliers and enforces the constraint ($\mathcal{Y}$).

2.2 Reference configuration

Cardiac geometries are acquired from in vivo medical images through imaging techniques. These geometries are in principle not stress free, mainly because there is always a pressure acting on the endocardium. Therefore, one needs to estimate the unloaded (i.e. stress-free) configuration (also named reference configuration) to which the 3D-0D cardiac electromechanical model (1) refers. To recover the reference configuration $\Omega_0$, starting from a geometry acquired from medical images $\Omega$, we apply the same procedure proposed for the left ventricle in [55].
Specifically, we solve an inverse problem: find the solution $d = d_{\Omega}$ of the following differential problem

$$
\begin{align*}
\nabla \cdot P(d, T_a) &= 0 & \text{in } \Omega_0, \\
P(d, T_a)N + K^{cpi}d &= 0 & \text{on } \Gamma_0^{epi}, \\
P(d, T_a)N &= -p_{LV}(t) J F^{-T} N & \text{on } \Gamma_0^{endo}, \\
P(d, T_a)N &= |J F^{-T} N| v_{base}^{LV}(t) & \text{on } \Gamma_0^{base},
\end{align*}
$$

(10)

obtained for $p_{LV} = \tilde{p}_{LV}$ and $T_a = \tilde{T}_a$, such that we get the domain $\Omega = \{ x : x = x_0 + d \ \forall x_0 \in \Omega_0 \}$.

Finally, to properly initialize the numerical simulation, we inflate the ventricular reference configuration $\Omega_0$ by solving problem (10), where we set the pressures $p_{LV} = p_{ED}^{LV}$. The value $p_{ED}^{LV}$ is chosen to bring the left ventricle to a defined volume $V_{ED}^{LV}$, as explained in [45]. Then, the solution $d$ of the problem (10) is set as initial condition $d_0$ for $d$ (with $d_0 = 0$) in (\M).
We employ the Finite Element Method (FEM) for the space discretization of electrophysiological, activation and mechanical models [47, 48]. We use hexahedral meshes and $Q_1$ finite element spaces for the space discretization of all the core models. This multiphysics problem presents different space resolutions according to the specific model at hand [55, 56]. In the framework of cardiac electrophysiology, we consider a fine geometrical description (1'987'285 DOFs, 1'926'912 cells, $h_{\text{mean}} \approx 0.86 \text{ mm}$) to accurately capture the electric propagation due to fine-scale phenomena arising from the continuum modeling of the cellular level, especially with the aim of reproducing and properly address arrhythmias [60]. On the other hand, cardiac mechanics allows a lower space resolution (35'725 DOFs, 30'108 cells, $h_{\text{mean}} \approx 3.3 \text{ mm}$). This eases its numerical solution, which is computationally demanding, especially for the assembling phase [60]. Differently from [61], in this paper we consider a significant space scales separation between electrophysiology and mechanics because we are dealing with a simple distribution of ischemic regions.

For time discretization, we use backward differentiation formula (BDF) schemes [47]. In particular, we employ a third order BDF scheme for electrophysiology and a first order BDF scheme for activation, mechanics and circulation. These choices allow to accurately capture the fast time dynamics of the model variables without unbearable restrictions on the time step, while not introducing numerical instabilities. We treat the nonlinear terms coming from both electrophysiological and activation models in a semi-implicit fashion. Cardiac mechanics is numerically advanced in time with a fully implicit scheme. The cardiocirculatory model is solved with an explicit method [45]. To stabilize the numerical oscillations arising from the feedback of the tissue mechanics on the activation model, without having to resort to a monolithic strategy, we rely on the scheme proposed in [54]. We use a smaller time step for electrophysiology ($\tau = 50 \mu s$) than for activation, mechanics and circulation ($\Delta t = 500 \mu s$). We set the final time $T = 4 \text{ s}$ for all the numerical simulations.

The mathematical models of Sec. 2 and the numerical methods presented in this section have been implemented in lifeX (https://lifex.gitlab.io/lifex), a high-performance C++ library developed within the iHEART project and based on the deal.II (https://www.dealii.org) Finite Element core [2].

The numerical simulations were performed on a HPC facility available at MOX for the iHEART project. The entire cluster is endowed with 8 Intel Xeon Platinum 8160 processors, for a total of 192 computational cores and a total amount of 1.5TB of available RAM.

4 Numerical results

We present electromechanical simulations to evaluate the effects of MEFs. We consider different modeling choices for the monodomain equation, as reported in Tab. 1. We start from a baseline simulation with model ($\mathcal{E}$), in which we obtain a stable VT. Then, we compare the effects of different geometry-mediated MEFs, i.e. models ($\mathcal{E}$), ($\mathcal{E}_{\text{gMEF-minimal}}$), ($\mathcal{E}_{\text{gMEF-enhanced}}$) and ($\mathcal{E}_{\text{gMEF-full}}$). We also study the impact of different parametrizations for SACs, i.e. ($\mathcal{E}_{\text{SAC}}$), with respect to ($\mathcal{E}$). Finally, we evaluate the combined effects of geometry-mediated MEFs and nonselective SACs. We report in Appendix A the values of the parameters that we use to get the numerical results that will be discussed in this paper.

4.1 Baseline simulation

We depict in Fig. 3 the evolution of the transmembrane potential and the displacement magnitude for the baseline electromechanical simulation, where all MEFs are fully neglected (model ($\mathcal{E}$) in Tab. 1). We induce a sustained VT with a figure-of-eight pattern around the isthmus, which is laterally bordered by scars, that act as conduction blocks.

In Fig. 4, we compare the Pressure-Volume (PV) loop over different heartbeats for the baseline simulation under VT with a reference healthy PV loop in sinus rhythm obtained by removing scars and grey zones. We observe that the contractility increases while the stroke volume (SV) decreases. The ejection fraction (EF) remains approximately the same and we approach a steady state in which the electromechanical function is not impaired. For these reasons, we conclude that the VT is stable.
Figure 3: Time evolution of the transmembrane potential $V$ (left) and displacement magnitude $|d|$ (right) for the Zygote LV with an idealized distribution of ischemic regions. Each picture on the right side is warped by the displacement vector $d$. MEFs are neglected, i.e. we use model ($\mathcal{E}$).
Figure 4: Comparison between a reference healthy PV loop in sinus rhythm (red, Appendix A, heartbeat period equal to 0.8 s) and the one obtained in the baseline simulation under VT for $t \in [0,4]$ s (light blue). We underline that, differently from [61], here we induce a hemodynamically tolerated VT.

Table 2: BCL for different modeling choices in geometry-mediated MEFs. Model $(\mathcal{E}_{\text{gMEF-minimal}})$ significantly changes BCL with respect to $(\mathcal{E})$, $(\mathcal{E}_{\text{gMEF-enhanced}})$, $(\mathcal{E}_{\text{gMEF-full}})$.

| Model                  | $(\mathcal{E})$ | $(\mathcal{E}_{\text{gMEF-minimal}})$ | $(\mathcal{E}_{\text{gMEF-enhanced}})$ | $(\mathcal{E}_{\text{gMEF-full}})$ |
|------------------------|-----------------|----------------------------------------|-----------------------------------------|----------------------------------|
| BCL                    | 0.60 s          | 0.65 s                                 | 0.61 s                                  | 0.60 s                           |

VT associated with ischemia are known to disturb the normal isovolumetric processes and to influence the end systolic/diastolic pressure volume relationship (ESPVR/EDPVR). Simultaneously, a phenomenon called incomplete relaxation may occur, especially when the VT does not leave enough time for the uncoupling of all the actin-myosin bonds between two consecutive contraction phases [6]. The occurrence of this phenomenon is illustrated in Fig. 5, where we depict the time evolution of the minimum, maximum and average active stress in the computational domain $\Omega_0$ for a reference healthy case in sinus rhythm and the baseline simulation under VT. Specifically, in the healthy case there is always a time interval between two consecutive heartbeats (precisely, during ventricular diastole) in which the active stress is virtually zero in the LV. In the VT case, instead, the cardiac muscle is never fully relaxed, thus not allowing the LV to complete its emptying. All these details are properly captured by our electromechanical model thanks to its biophysical accuracy.

4.2 Effects of geometry-mediated MEFs

We consider four different modeling choices for the geometry-mediated MEFs, namely $(\mathcal{E})$, $(\mathcal{E}_{\text{gMEF-minimal}})$, $(\mathcal{E}_{\text{gMEF-enhanced}})$ and $(\mathcal{E}_{\text{gMEF-full}})$ in Tab. 1, while for the moment we completely neglect the impact of SACs. Then, we perform four different electromechanical simulations by employing these four different formulations for the monodomain equation.

We illustrate in Fig. 6 the development of transmembrane potential $V$ over time. We observe minor differences
Figure 5: Minimum, average and maximum active tension $T_a$ over time for a reference healthy case in sinus rhythm (left, Appendix A, heartbeat period equal to 0.8 s) and the baseline simulation under VT (right). We see that incomplete relaxation occurs during VT.

in action potential propagation among ($\epsilon$), ($\delta_{gMEF\text{-}\text{enhanced}}$) and ($\delta_{gMEF\text{-}\text{full}}$). These differences, as we can see for $t = 4 \text{s}$, are mainly focused on the depolarization wave and occur during VT. Moreover, by looking at Tab. 2, we notice that the VT BCL is very similar among these three models. Indeed, the BCL is approximately equal to 0.60 s, which is long if compared to more dangerous VT and justifies a stable ventricular excitation.

On the other hand, ($\delta_{gMEF\text{-}\text{minimal}}$) entails major changes in VT BCL, which increases from 0.60 s (for model ($\epsilon$)) to 0.65 s, and conduction velocity, that significantly decreases. These observations are in agreement with Fig. 7, where the electrophysiological, mechanical and hemodynamic variables retrieved in a random point of the computational domain are shifted forward for ($\delta_{gMEF\text{-}\text{minimal}}$), while ($\epsilon$), ($\delta_{gMEF\text{-}\text{enhanced}}$) and ($\delta_{gMEF\text{-}\text{full}}$) show a very similar pattern. This is also motivated by the change in the VT exit site, as we can see from Fig. 6 for $t = 2.4 \text{s}$. This phenomenon is particularly evident from the plot of sarcomere length over time (Fig. 7), which also presents different peak values for ($\delta_{gMEF\text{-}\text{minimal}}$) and $t \gtrsim 2.4 \text{s}$. Finally, we see that wave stability is not affected by geometry-mediated MEFs. Indeed, the VT always remains hemodynamically stable in the four different cases.

4.3 Effects of SACs

In this section, we fully neglect the effects of geometry-mediated MEFs and we focus on different parametrizations for SACs in terms of $G_s$ and $V_{\text{rev}}$. We also compare the outcomes of ($\delta_{\text{SAC}}$) with the ones of ($\epsilon$) to outline similarities and differences.

We notice from Fig. 8 that both APD and wave stability are affected by SACs parametrizations. Indeed, by combining the 3D information with the pointwise evaluations of Fig. 9, we discover that there is one choice of the parameters, namely $G_s = 100 \text{s}^{-1}$ and $V_{\text{rev}} = 0 \text{mV}$, that converts the VT from stable to unstable. The instability derives from an extra stimulus that is completely driven by contraction, which occurs in the superior-right part of the ventricle. This extra stimulus changes the VT morphology, along with its BCL, which is not the same over time.

By considering data of Tab. 3, we observe that the VT BCL remains the same when there is no stability transition.
Figure 6: Comparison among different models for geometry-mediated MEFs in terms of transmembrane potential $V$. 

$t = 0.20\text{s}$

$t = 0.45\text{s}$

$t = 2.4\text{s}$

$t = 4.0\text{s}$
Figure 7: Pointwise values of transmembrane potential $V$, intracellular calcium concentration $[\text{Ca}^{2+}]_i$, sarcomere length $SL$, active tension $T_a$, pressure $p_{LV}$ and volume $V_{LV}$ over time for $(\delta)$, $(\delta_{\text{gMEF-minimal}})$, $(\delta_{\text{gMEF-enhanced}})$ and $(\delta_{\text{gMEF-full}})$. 
Table 3: VT classification for different SACs parametrizations. The unstable VT has a BCL that ranges from 0.43 s to 0.58 s.

| Model     | VT type               |
|-----------|-----------------------|
| (Δ)       | Stable (BCL = 0.60 s) |
| (Δ_{SAC}), G_s = 100 s^{-1}, V_rev = -70 mV | Stable (BCL = 0.60 s) |
| (Δ_{SAC}), G_s = 100 s^{-1}, V_rev = -35 mV | Stable (BCL = 0.60 s) |
| (Δ_{SAC}), G_s = 100 s^{-1}, V_rev = 0 mV | Unstable (BCL_{avg} = 0.50 s) |
| (Δ_{SAC}), G_s = 50 s^{-1}, V_rev = 0 mV | Stable (BCL = 0.60 s) |

Table 4: VT classification for combinations of geometry-mediated MEFs and nonselective SACs. The unstable VT related to (Δ_{SAC}) has a BCL that ranges from 0.43 s to 0.58 s. The unstable VT related to (Δ_{SACMEF-full, SAC}) has a BCL that ranges from 0.44 s to 0.50 s.

| Model     | VT type               |
|-----------|-----------------------|
| (Δ)       | Stable (0.60 s)       |
| (Δ_{SAC}), G_s = 100 s^{-1}, V_rev = 0 mV | Unstable (BCL_{avg} = 0.50 s) |
| (Δ_{SACMEF-full, SAC}), G_s = 100 s^{-1}, V_rev = 0 mV | Unstable (BCL_{avg} = 0.47 s) |

Moreover, from the hemodynamic perspective, the onset of different types of arrhythmias is driven by the combined effects of G_s and V_rev. Indeed, in Fig. 9 we see that given G_s, different V_rev may change wave stability. On the other hand, in Fig. 10 we show that different G_s may affect wave stability, with V_rev fixed a priori.

4.4 Combined effects of geometry-mediated MEFs and SACs

We briefly evaluate the combined effects of geometry-mediated MEFs and nonselective SACs. Once SACs parametrization is fixed, we notice that switching between no formulation (i.e. (Δ_{SAC})) to full formulation (i.e. (Δ_{SACMEF-full, SAC})) of geometry-mediated MEFs entails significant differences. In particular, from Fig. 11 we see that (Δ_{SACMEF-full, SAC}) triggers the extra stimuli faster than (Δ_{SAC}). As we can notice from Fig. 12, the VT remains unstable but both pressure and volume traces over time are very different from each other for (Δ_{SAC}) and (Δ_{SACMEF-full, SAC}). From Tab. 4, we infer that the VT BCL for (Δ_{SACMEF-full, SAC}) is lower than the one of (Δ_{SAC}). This potentially defines a more dangerous VT. Finally, in Fig. 13 we highlight the joint contributions of electrophysiology, mechanics and hemodynamics in the (Δ_{SACMEF-full, SAC}) coupled model.

5 Discussion

We investigate how several types of geometry-mediated MEFs and different parametrizations for nonselective SACs affect the electric and hemodynamic stability of VT. In particular, we focus on the sustainment and the morphology
Figure 8: Comparison between model $\langle E \rangle$ and model $(E_{SAC})$, for different values of $V_{rev}$ ($G_s = 100 \text{s}^{-1}$).
Figure 9: Pointwise values of transmembrane potential $V$, intracellular calcium concentration $[\text{Ca}^{2+}]_i$, sarcomere length $SL$, active tension $T_a$, pressure $p_{LV}$ and volume $V_{LV}$ over time for $(\delta)$ and $(\delta_{\text{SAC}})$ with different choices of $V_{rev}$ ($G_s = 100 \text{s}^{-1}$).
of VT macro-reentrant circuits and blood supply, which is analyzed by means of PV loops.

Differently from previous studies [13, 31, 65], we keep into account tissue heterogeneity by introducing an idealized distribution of scars, grey zones and non-remodeled regions over the myocardium. We also consider a more sophisticated coupled mathematical model that embraces electrophysiology, activation, mechanics and cardiovascular fluid dynamics to analyze these mechano-electric couplings. To the best of our knowledge, this framework enables to study for the first time the hemodynamic effects of both geometry-mediated MEFs and nonselective SACs on VT by means of electromechanical simulations, showing strengths and weaknesses of some simplifying modeling choices that are commonly made when simulating this type of phenomenon [11, 15, 36, 49]. The coupling between the 3D electromechanical model and the 0D circulation model permits to identify the hemodynamic nature of the VT. With our approach, we discriminate between stable and unstable VT, which might result in hemodynamically tolerated or not tolerated VT [61].

We see that if a VT is triggered by a certain stimulation protocol and by neglecting all MEFs, the very same pacing protocol induces a VT for all possible combinations of MEFs [61].

In our numerical simulations, we do not observe a significant impact of geometry-mediated MEFs on the induction and sustainment of the VT. Most of the modeling choices for geometry-mediated MEFs, namely \((\mathcal{E})\), \((\mathcal{E}_{\text{MEF-enhanced}})\) and \((\mathcal{E}_{\text{MEF-full}})\), present very similar VT BCLs and conduction velocities, while showing a few differences in the depolarization wave [13]. On the other hand, \((\mathcal{E}_{\text{MEF-minimal}})\) manifests major differences in the VT BCL, which increases, and its exit site with respect to \((\mathcal{E})\), \((\mathcal{E}_{\text{MEF-enhanced}})\) and \((\mathcal{E}_{\text{MEF-full}})\), as shown in [61] for a patient-specific unstable VT. This can be justified by the simplifications introduced in \((\mathcal{E}_{\text{MEF-minimal}})\), while moving towards \((\mathcal{E}_{\text{MEF-full}})\) we almost totally recover the behavior observed in \((\mathcal{E})\). Therefore, the minimal MEFs modeling choice might lead to biased results which are in favor of less severe VT.

We observe that nonselective SACs may affect the hemodynamic nature of the VT, as they may induce EADs or DADs, which lead to ectopic foci that reactivate the LV [25]. These extra stimuli are generally located in the regions of the myocardium in which there is a transition between scar and border zone or between border zone and non-remodeled areas, where high stretches are likely to be present [29]. According to the specific combination of \(G_s\) and \(V_{\text{rev}}\), nonselective SACs affects both APD and AP resting values [68]. We remark that such spontaneous
Figure 11: Comparison among models (E), (E_{SAC}) and (E_{gMEF-full, SAC}).
Figure 12: Pointwise values of pressure $p_{LV}$ and volume $V_{LV}$ over time for $(\mathcal{E}^{E})$, $(\mathcal{E}^{E_SAC})$ ($G_s = 100 \text{s}^{-1}$, $V_{rev} = 0 \text{mV}$) and $(\mathcal{E}^{E_{gMEF-full, SAC}})$ ($G_s = 100 \text{s}^{-1}$, $V_{rev} = 0 \text{mV}$).

Arrhythmias triggered by myocardial stretches cannot be assessed in electrophysiological simulations, where the mechanical behavior is neglected.

Finally, in our study we also investigate the combined effects of geometry-mediated MEFs and nonselective SACs. Significant differences between $(\mathcal{E}^{E_{gMEF-full, SAC}})$ and $(\mathcal{E}^{E_{SAC}})$ are observed when geometry-mediated MEFs are combined with a parametrization of SACs that entails extra stimuli. Specifically, the VT BCL with $(\mathcal{E}^{E_{gMEF-full, SAC}})$ is lower than the one of $(\mathcal{E}^{E_{SAC}})$ because the extra stimuli driven by SACs is triggered more often in the former case. This completely changes the pressure-volume dynamics of the VT, whose stability is however still not affected by the geometry-mediated MEFs.

6 Limitations

There are other types of MEFs that could be investigated in future works. Among them, an important role is certainly played by the mechanical effects mediated by fibroblasts in the extracellular matrix, $[\text{Ca}^{2+}]$, buffers handling, alterations in transmembrane capacitance $C_m$ due to local stretch and ions selective SACs [33]. Indeed, modeling different cellular processes in cardiac electrophysiology might be of interest to further shed light on the underlying mechanisms of arrhythmias. Nevertheless, we have provided a broad study of geometry-mediated MEFs and nonselective SACs by combining electrophysiological, mechanical and hemodynamic observations.

Furthermore, MEFs should be properly quantified in patient-specific cases. Currently, our model for nonselective SACs permits to put one single value of both $G_s$ and $u_{rev}$ for the whole geometry, which is unlikely to happen in realistic scenarios. A precise space assessment of SACs combined with our mathematical framework could have significant clinical implications.
Figure 13: Coupled effects of electrophysiology, mechanics and hemodynamics for the numerical simulation with model ($E_{\text{MEF-full, SAC}}$). The extra stimuli in the upper right part of the LV, which is driven by SACs, activate the LV electrophysiologically and mechanically. This has a direct impact on both pressure and volume transients, which in turn have an effect on the electromechanical behavior of the LV.
7 Conclusions

We studied the effects of geometry-mediated MEFs and nonselective SACs on a realistic LV geometry endowed with an idealized distribution of infarct and peri-infarct zones. We performed numerical simulations of cardiac electromechanics coupled with closed-loop cardiovascular circulation under VT.

Our electromechanical framework allows for the hemodynamic classification of the VT, which can be either stable or unstable, and permits to capture mechanically relevant indications under VT, such as the incomplete relaxation of sarcomeres. Furthermore, by combining electrophysiology, activation, mechanics and hemodynamics, we observed several differences on the morphology of the VT with respect to electrophysiological simulations. In particular, geometry-mediated MEFs do not affect VT stability but may alter the VT BCL, along with its exit site [31]. On the other hand, the recruitment of SACs may generate extra stimuli, which may change VT stability. These extra stimuli are driven by myocardial contraction and are induced by changes in the APD or in the resting value of the transmembrane potential [29]. We conclude that both geometry-mediated MEFs and nonselective SACs define important contributions in electromechanical models with hemodynamic coupling, especially when numerical simulations under arrhythmia are carried out.

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Appendices

A Model parameters

We provide the full list of parameters adopted for the numerical simulations of this paper. Specifically, Tab. 5 contains the parameters related to the electrophysiological model, Tab. 6 those related to the sarcomere model RDQ20-MF, Tab. 7 the parameters of the mechanical model and, finally, Tab. 8 contains the parameters associated with the circulation model. For the TTP06 model, we adopt the parameters reported in the original paper (for endocardial cells) [69], with the only difference that we rescale the intracellular calcium concentration $[Ca^{2+}]_i$ by a factor of $\omega_{Ca}$ to get more physiological values [16].
### Table 5: Parameters of the electrophysiological model.

| Variable                          | Value      | Unit     |
|-----------------------------------|------------|----------|
| Conductivity tensor               |            |          |
| $\sigma_l$                        | 0.7643 · 10^{-4} | m^2 s^{-1} |
| $\sigma_t$                        | 0.3494 · 10^{-4} | m^2 s^{-1} |
| $\sigma_n$                        | 0.1125 · 10^{-4} | m^2 s^{-1} |
| Applied current                   |            |          |
| $T_{\text{app}}$                  | 17         | V s^{-1} |
| $t_{\text{app}}$                  | 3 · 10^{-3} | s        |
| Calcium rescaling                 |            |          |
| $\omega_{\text{Ca}}$              | 0.48       | -        |

### Table 6: Parameters of the sarcomere model RDQ20-MF (for the definition of the parameters, see [51]).

| Variable                          | Value      | Unit     |
|-----------------------------------|------------|----------|
| Regulatory units steady-state     |            |          |
| $\mu$                             | 10         | -        |
| $\gamma$                          | 30         | -        |
| $Q$                               | 2          | -        |
| $k_d$                             | 0.4        | µM       |
| $\alpha_k$                        | -0.2083    | µM µm^{-1} |
| Regulatory units kinetics         |            |          |
| $k_{\text{off}}$                   | 40         | s^{-1}   |
| $k_{\text{basic}}$                | 8          | s^{-1}   |
| Crossbridge cycling               |            |          |
| $\rho_{\mu}$                      | 32.255     | s^{-1}   |
| $\rho_{\gamma}$                   | 0.768      | s^{-1}   |
| $\rho_Q$                          | 134.31     | s^{-1}   |
| Upscaling                         |            |          |
| $a_{\text{XB}}$                    | 160        | MPa      |
| $SL_0$                            | 1.9        | µm       |

### Table 7: Parameters of the mechanical model.

| Variable                          | Value      | Unit     |
|-----------------------------------|------------|----------|
| Constitutive law                  |            |          |
| $B$                               | 50 · 10^3  | Pa       |
| $C$                               | 0.88 · 10^3 | Pa       |
| $b_{ff}$                          | 8          | -        |
| $b_{ss}$                          | 6          | -        |
| $b_{nn}$                          | 3          | -        |
| $b_{fs}$                          | 12         | -        |
| $b_{fn}$                          | 3          | -        |
| $b_{sn}$                          | 3          | -        |
| Boundary conditions               |            |          |
| $K_{\text{epi}}^\parallel$       | 2 · 10^4   | Pa m^{-1} |
| $K_{\text{epi}}^\perp$            | 2 · 10^3   | Pa m^{-1} |
| $C_{\text{epi}}^\parallel$       | 2 · 10^4   | Pa s m^{-1} |
| $C_{\text{epi}}^\perp$            | 2 · 10^3   | Pa s m^{-1} |
| Tissue density                    |            |          |
| $\rho_s$                          | 10^3       | kg m^{-3} |

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| Variable | Value | Unit |
|----------|-------|------|
| \(R_{SYS}^{R}\) | 0.64 | mmHg s mL\(^{-1}\) |
| \(R_{PUL}^{R}\) | 0.032116 | mmHg s mL\(^{-1}\) |
| \(R_{SYS}^{R}\) | 0.32 | mmHg s mL\(^{-1}\) |
| \(R_{PUL}^{R}\) | 0.035684 | mmHg s mL\(^{-1}\) |
| \(C_{SYS}^{E}\) | 1.2 | mL mmHg\(^{-1}\) |
| \(C_{PUL}^{E}\) | 10.0 | mL mmHg\(^{-1}\) |
| \(C_{SYS}^{C}\) | 60.0 | mL mmHg\(^{-1}\) |
| \(C_{PUL}^{C}\) | 16.0 | mL mmHg\(^{-1}\) |
| \(L_{SYS}^{L}\) | 5 \cdot 10^{-3} | mmHg s\(^2\) mL\(^{-1}\) |
| \(L_{PUL}^{L}\) | 5 \cdot 10^{-4} | mmHg s\(^2\) mL\(^{-1}\) |
| \(L_{SYS}^{L}\) | 5 \cdot 10^{-4} | mmHg s\(^2\) mL\(^{-1}\) |
| \(L_{PUL}^{L}\) | 5 \cdot 10^{-4} | mmHg s\(^2\) mL\(^{-1}\) |

| Variable | Value | Unit |
|----------|-------|------|
| \(E_{PUL}^{LA}\) | 0.18 | mmHg mL\(^{-1}\) |
| \(E_{PUL}^{RA}\) | 0.07 | mmHg mL\(^{-1}\) |
| \(E_{PUL}^{RV}\) | 0.05 | mmHg mL\(^{-1}\) |
| \(E_{PUL}^{max, LA}\) | 0.07 | mmHg mL\(^{-1}\) |
| \(E_{PUL}^{max, RA}\) | 0.06 | mmHg mL\(^{-1}\) |
| \(E_{PUL}^{max, RV}\) | 0.55 | mmHg mL\(^{-1}\) |
| \(V_{0, LA}\) | 4.0 | mL |
| \(V_{0, RA}\) | 4.0 | mL |
| \(V_{0, RV}\) | 16.0 | mL |

| Cardiac valves |
|----------------|
| \(R_{min}\) | 0.0075 | mmHg s mL\(^{-1}\) |
| \(R_{max}\) | 75006.2 | mmHg s mL\(^{-1}\) |

Table 8: Parameters of the circulation model.
References

[1] H. Arevalo, F. Vadakkumpadan, E. Guallar, and et al. “Arrhythmia risk stratification of patients after myocardial infarction using personalized heart models”. In: *Nature Communications* 7 (2016), pp. 113–128.

[2] D. Arndt, W. Bangerth, B. Blais, and et al. “The deal.II Library, Version 9.2”. In: *Journal of Numerical Mathematics* 28.3 (2020), pp. 131–146.

[3] T. Arts, T. Delhaas, P. Bovendeerd, and et al. “Adaptation to mechanical load determines shape and properties of heart and circulation: the CircAdapt model”. In: *American Journal of Physiology-Heart and Circulatory Physiology* 288 (2005), H1943–H1954.

[4] C. M. Augustin, M. A. F. Gsell, E. Karabelas, and et al. “A computationally efficient physiologically comprehensive 3D–0D closed-loop model of the heart and circulation”. In: *Computer Methods in Applied Mechanics and Engineering* 386 (2021), p. 114092.

[5] C. Bartolucci, E. Passini, J. Hyttinen, and et al. “Simulation of the Effects of Extracellular Calcium Changes Leads to a Novel Computational Model of Human Ventricular Action Potential With a Revised Calcium Handling”. In: *Frontiers in Physiology* 11 (2020), p. 314.

[6] M. B. Bastos, D. Burkhoff, J. Maly, and et al. “Invasive left ventricle pressure–volume analysis: overview and practical clinical implications”. In: *European Heart Journal* 41.12 (2019), pp. 1286–1297.

[7] J. D. Bayer, R. C. Blake, G. Plank, and N. Trayanova. “A novel rule-based algorithm for assigning myocardial fiber orientation to computational heart models”. In: *Annals of Biomedical Engineering* 40.10 (2012), pp. 2243–2254.

[8] P. J. Blanco and R. A. Feijóo. “A 3D-1D-0D computational model for the entire cardiovascular system”. In: *Computational Mechanics, eds. E. Dvorking, M. Goldschmit, M. Storti* 29 (2010), pp. 5887–5911.

[9] A. Bueno-Orovio, E. M. Cherry, and F. H. Fenton. “Minimal model for human ventricular action potentials in tissue”. In: *Journal of Theoretical Biology* 253 (2008), pp. 544–560.

[10] N. Cedilnik, J. Duchateau, R. Dubois, and et al. “Fast personalized electrophysiological models from computed tomography images for ventricular tachycardia ablation planning”. In: *EP Europace* 20.suppl 3 (2018), pp. iii94–iii101.

[11] D. Chapelle, M. A. Fernández, J. F. Gerbeau, and et al. “Numerical simulation of the electromechanical activity of the heart”. In: *International Conference on Functional Imaging and Modeling of Heart* 5528 (2009), pp. 357–365.

[12] P. Colli Franzone, L. F. Pavarino, and S. Scacchi. “A numerical study of scalable cardiac electro-mechanical solvers on HPC architectures”. In: *Frontiers in Physiology* 9 (2018), p. 268.

[13] P. Colli Franzone, L. F. Pavarino, and S. Scacchi. “Effects of mechanical feedback on the stability of cardiac scroll waves: A bidomain electro-mechanical simulation study”. In: *Chaos* 27 (2017), p. 093905.

[14] P. Colli Franzone, L. F. Pavarino, and S. Scacchi. *Mathematical Cardiac Electrophysiology*. Vol. 13. Springer, 2014.

[15] A. Collin, S. Imperiale, P. Moireau, and et al. “Apprehending the effects of mechanical deformations in cardiac electrophysiology: A homogenization approach”. In: *Mathematical Models and Methods in Applied Sciences* 29.13 (2019), pp. 2377–2417.

[16] R. Coppini, C. Ferrantini, L. Yao, and et al. “Late sodium current inhibition reverses electromechanical dysfunction in human hypertrophic cardiomyopathy”. In: *Circulation* 127.5 (2013), pp. 575–584.

[17] F. S. Costabal, F. A. Concha, D. E. Hurtado, and et al. “The importance of mechano-electrical feedback and inertia in cardiac electromechanics”. In: *Computer Methods in Applied Mechanics and Engineering* 320 (2017), pp. 352–368.
[18] F. S. Costabal, D. Hurtado, and E. Kuhl. “Generating Purkinje networks in the human heart”. In: Journal of Biomechanics 49 (2016), pp. 2455–2465.

[19] D. Deng, A. Prakosa, J. Shade, and et al. “Sensitivity of Ablation Targets Prediction to Electrophysiological Parameter Variability in Image-Based Computational Models of Ventricular Tachycardia in Post-infarction Patients”. In: Frontiers in Physiology 10 (2019), p. 628.

[20] S. Doll and K. Schweizerhof. “On the development of volumetric strain energy functions”. In: Journal of Applied Mechanics 67 (2000), pp. 17–21.

[21] A. E. Epstein and et al. “ACC/AHA/HRS 2008 Guidelines for Device-Based Therapy of Cardiac Rhythm Abnormalities: a report of the American College of Cardiology/American Heart Association Task Force on Practice Guidelines (Writing Committee to Revise the ACC/AHA/NASPE 2002 Guideline Update for Implantation of Cardiac Pacemakers and Antiarrhythmia Devices): developed in collaboration with the American Association for Thoracic Surgery and Society of Thoracic Surgeons”. In: Circulation 117 (2008), e350–e408.

[22] A. Gerbi, L. Dedè, and A. Quarteroni. “A monolithic algorithm for the simulation of cardiac electromechanics in the human left ventricle”. In: Mathematics in Engineering 1 (2018), pp. 1–37.

[23] J. M. Guccione and A. D. McCulloch. “Finite element modeling of ventricular mechanics”. In: Theory of Heart. Springer, 1991, pp. 121–144.

[24] J. M. Guccione, A. D. McCulloch, and L. K. Waldman. “Passive material properties of intact ventricular myocardium determined from a cylindrical model”. In: Journal of Biomechanical Engineering 113 (1991), pp. 42–55.

[25] A. Hazim, Y. Belhamadia, and S. Dubljevic. “A Simulation Study of the Role of Mechanical Stretch in Arrhythmogenesis during Cardiac Alternans”. In: Biophysical Journal 120.1 (2021), pp. 109–121.

[26] M. Hirschvogel, M. Bassilios, L. Jagschies, and et al. “A monolithic 3D-0D coupled closed-loop model of the heart and the vascular system: experiment-based parameter estimation for patient-specific cardiac mechanics”. In: International Journal for Numerical Methods in Biomedical Engineering 33 (2017), e2842.

[27] G. A. Holzapfel and R. W. Ogden. “Constitutive modelling of passive myocardium: a structurally based framework for material characterization”. In: Philosophical Transactions of the Royal Society A: Mathematical, Physical and Engineering Sciences 367 (2009), pp. 3445–3475.

[28] Z. M. G. Inc. Zygote Solid 3D heart Generation II Development Report. Technical Report. 2014.

[29] X. Jie, V. Gurev, and N. Trayanova. “Mechanisms of Mechanically Induced Spontaneous Arrhythmias in Acute Regional Ischemia”. In: Circulation Research 106.1 (2010), pp. 185–192.

[30] J. P. Keener and J. Sneyd. Mathematical Physiology. Vol. 1. Springer, 2009.

[31] R. H. Keldermann, M. P. Nash, H. Gelderblom, and et al. “Electromechanical wavebreak in a model of the human left ventricle”. In: American Journal of Physiology-Heart and Circulatory Physiology 299.1 (2010), H134–H143.

[32] R. C. P. Kerkhoffs, M. L. Neal, Q. Gu, and et al. “Coupling of a 3D finite element model of cardiac ventricular mechanics to lumped system models of the systemic and pulmonary circulation”. In: Annals of biomedical engineering 35 (2007), pp. 1–18.

[33] P. Kohl, F. Sachs, and M. R. Franz. Cardiac Mechano-Electric Coupling and Arrhythmias. Oxford University Press, 2013.

[34] S. Land, S. Park-Holohan, N. P. Smith, and et al. “A model of cardiac contraction based on novel measurements of tension development in human cardiomyocytes”. In: Journal of Molecular and Cellular Cardiology 106 (2017), pp. 108–109.
[35] M. Landajuela, C. Vergara, A. Gerbi, and et al. “Numerical approximation of the electromechanical coupling in the left ventricle with inclusion of the Purkinje network”. In: *International journal for numerical methods in biomedical engineering* 34 (2018), e2984.

[36] F. Levrero-Florencio, F. Margara, E. Zacur, and et al. “Sensitivity analysis of a strongly-coupled human-based electromechanical cardiac model: Effect of mechanical parameters on physiologically relevant biomarkers”. In: *Computer Methods in Applied Mechanics and Engineering* 361 (2020), p. 112762.

[37] C. Luo and Y. Rudy. “A dynamic model of the cardiac ventricular action potential. I. Simulations of ionic currents and concentration changes”. In: *Circulation Research* 74 (1994), pp. 1071–1096.

[38] S. A. Niederer, P. J. Hunter, and N. P. Smith. “A quantitative analysis of cardiac myocyte relaxation: a simulation study”. In: *Biophysical Journal* 90.5 (2006), pp. 1697–1722.

[39] S. A. Niederer, G. Plank, P. Chinchapatnam, and et al. “Length-dependent tension in the failing heart and the efficacy of cardiac resynchronization therapy”. In: *Cardiovascular research* 89 (2011), pp. 336–343.

[40] F. Nobile, A. Quarteroni, and R. Ruiz-Baier. “An active strain electromechanical model for cardiac tissue”. In: *International journal for numerical methods in biomedical engineering* 28 (2012), pp. 52–71.

[41] R. W. Ogden. *Non-linear elastic deformations*. Courier Corporation, 1997.

[42] S. H. Peng and W. V. Chang. “A compressible approach in finite element analysis of rubber-elastic materials”. In: *Computers & Structures* 62 (1997), pp. 573–593.

[43] M. R. Pfaffer, J. M. Hörmann, M. Weigl, and et al. “The importance of the pericardium for cardiac biomechanics: from physiology to computational modeling”. In: *Biomechanics and Modeling in Mechanobiology* 18 (2019), pp. 503–529.

[44] R. Piersanti, P. C. Africa, M. Fedele, and et al. “Modeling cardiac muscle fibers in ventricular and atrial electrophysiology simulations”. In: *Computer Methods in Applied Mechanics and Engineering* 373 (2021), p. 113468.

[45] R. Piersanti, F. Regazzoni, M. Salvador, and et al. “3D-0D closed-loop model for the simulation of cardiac electromechanics”. In: *MOX Report 57* (Politecnico di Milano, 2021).

[46] A. Prakosa, H. Arevalo, D. Dongdong, and et al. “Personalized virtual-heart technology for guiding the ablation of infarct-related ventricular tachycardia”. In: *Nature Biomedical Engineering* 2 (2018), pp. 732–740.

[47] A. Quarteroni. *Numerical models for differential problems*. Vol. 2. Springer, 2009.

[48] A. Quarteroni, L. Dedè, A. Manzoni, and C. Vergara. *Mathematical modelling of the human cardiovascular system: data, numerical approximation, clinical applications*. Cambridge Monographs on Applied and Computational Mathematics. Cambridge University Press, 2019.

[49] A. Quarteroni, T. Lassila, S. Rossi, and R. Ruiz-Baier. “Integrated Heart—Coupling multiscale and multiphysics models for the simulation of the cardiac function”. In: *Computer Methods in Applied Mechanics and Engineering* 314 (2017), pp. 345–407.

[50] A. Quarteroni, A. Veneziani, and C. Vergara. “Geometric multiscale modeling of the cardiovascular system, between theory and practice”. In: *Computer Methods in Applied Mechanics and Engineering* 302 (2016), pp. 193–252.

[51] F. Regazzoni, L. Dedè, and A. Quarteroni. “Biophysically detailed mathematical models of multiscale cardiac active mechanics”. In: *PLoS computational biology* 16 (2020), e1008294.

[52] F. Regazzoni, L. Dedè, and A. Quarteroni. “Machine learning of multiscale active force generation models for the efficient simulation of cardiac electromechanics”. In: *Computer Methods in Applied Mechanics and Engineering* 370 (2020), p. 113268.

[53] F. Regazzoni and A. Quarteroni. “Accelerating the convergence to a limit cycle in 3D cardiac electromechanical simulations through a data-driven 0D emulator”. In: *Computers in Biology and Medicine* 135 (2021), p. 104641.
[71] C. Vergara, S. Palamara, D. Catanzariti, and et al. “Patient-specific generation of the Purkinje network driven by clinical measurements of a normal propagation”. In: Medical & Biological Engineering & Computing 52 (2014), pp. 813–826.