Estimation of Parameters for an Archetypal Model of Cardiomyocyte Membrane Potentials

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Abstract: Contemporary realistic mathematical models of single-cell cardiac electrical excitation are immensely detailed. Model complexity leads to parameter uncertainty, high computational cost and barriers to mechanistic understanding. There is a need for reduced models that are conceptually and mathematically simple but physiologically accurate. To this end, we consider an archetypal model of single-cell cardiac excitation that replicates the phase-space geometry of detailed cardiac models, but at the same time has a simple piecewise-linear form and a relatively low-dimensional configuration space. In order to make this archetypal model practically applicable, we develop and report a robust method for estimation of its parameter values from the morphology of single-stimulus action potentials derived from detailed ionic current models and from experimental myocyte measurements. The procedure is applied to five significant test cases and an excellent agreement with target biomarkers is achieved. Action potential duration restitution curves are also computed and compared to those of the target test models and data, demonstrating conservation of dynamical pacing behaviour by the fine-tuned archetypal model. An archetypal model that accurately reproduces a variety of wet-lab and synthetic electrophysiology data offers a number of specific advantages such as computational efficiency, as also demonstrated in the study. Open-source numerical code of the models and methods used is provided.

Keywords: Mathematical models, Cardiac action potential, Electrophysiology, Parameter estimation.

Introduction
Models of the action potential of cardiac cells are routinely used to interpret and integrate experimental findings, extrapolate animal data to human system context, and test novel hypotheses [21, 55]. It is frequently proposed that these models will soon make it possible to devise patient-specific precision therapies, and accelerate cardiac drug discovery and development [3, 16, 54]. Since the pioneering work of Noble [39], over 150 models have been published with the aim to capture in detail the electrophysiology of a wide variety of cardiac cell types under a broad range of experimental, physiological and pathological conditions [19]. Contemporary detailed models largely follow the Hodgkin-Huxley paradigm but have grown to a staggering complexity [47]. For example, a recent model of the human ventricular action potential [41] consists of 49 ordinary differential equations and includes 206 model parameters. Because models are typically developed by extending and re-using components from previous models, as advocated by large international initiatives like the Physiome Project [6] and the CellML Project [32],
many of these parameters and equations are poorly constrained and in many cases redundant. For instance, the meta-analysis of [38] demonstrates that the modern human ventricular models [52, 23] include parameters that have been inherited from studies in at least 9 different species over a range of 6 different temperatures; in other words, it is questionable whether they represent any human ventricular myocyte. While intense research effort is expended to estimate parameter uncertainty [12], calibrate models to identifiable and reliable experimental protocols [60, 13], increase reproducibility [14] and build trustworthy models [24], detailed cardiac cell models remain difficult to benchmark and to adapt to situations to which they have not been fitted [61] and are computationally expensive especially in tissue-scale simulations [11]. Most importantly, detailed cardiac models are becoming increasingly difficult for causal inference [9].

Thus, there is a certain need for simplified mathematical models that are accurate and flexible enough, computationally affordable, amenable to mathematical analysis and to mechanistic understanding. Starting with the early work of van der Pol [57] a number of conceptual models have been proposed to address this need, e.g. [20, 36, 2, 18, 33], and most of them have become popular and frequently used in the place of detailed ones. However, these conceptual models rely mostly on ad hoc assumptions and generally have a FitzHugh-Nagumo structure that leads to certain shortcomings [9]. In contrast, in [10, 9] we developed an asymptotic method that allows for a systematic and controlled reduction of arbitrary detailed cardiac ion current models. The method preserves the phase-space geometry of detailed models, different from the FitzHugh-Nagumo one, and reveals qualitatively new features of topological nature [49]. Following this approach in [9] we reduced Noble’s model of purkinje fibre electrophysiology [39]. We obtained a mathematically simple conceptual model that consists of three piece-wise linear differential equations and contains only 13 intrinsic model parameters and so it is rather inexpensive to integrate numerically. Further, the model admits closed-form analytic solutions when spacially-clamped [9], and closed-form travelling wave solutions when spacially-extended [49]. These exact solutions aid mechanistic understanding, extensive exploration of parameter space as well as benchmarking of numerical codes. More importantly the model is archetypal in the sense that it has the generic asymptotic structure of modern detailed cardiac ionic models and it is thus capable of reproducing slow repolarization, slow sub-threshold response, fast accommodation, front dissipation, variable peak voltage and other features of cardiac excitability crucial for understanding and controlling arrhythmogenesis [9] where most other ad hoc conceptual equations often fail.

In order to be practically useful beyond its utility as a conceptual tool, the parameter values of the archetypal model [9] must be determined so that it replicates the behaviour of state-of-the-art models of ventricular and atrial excitation and captures experimental measurements quantitatively. This is the goal of the present work. To this end we describe in the following the implementation of a standard parameter estimation procedure and use it to fit the archetypal model to a typical mammalian ventricular model [28], a typical human atrial model [15], as well as to experimental data for rabbit ventricular myocytes available from the literature [29]. In addition, we provide an open-source numerical code permanently available at [4] that can be used by the reader to apply the methodology to other detailed models and data of their own interest. Further, fitting data and detailed models to a common set of equations gives the opportunity to compare and contrast such models directly, which is otherwise impossible due to their different mathematical structure and components. The scale of the computational effort required to perform large-scale tissue and whole heart simulations is immense and simulations in real-time are beyond current computational capability. A number of strategies for reducing calculation
time are used at present including code parallelisation, lookup tables, exponential solutions for gating variables, operator splitting, adaptive time and space stepping, using graphics processing units and using simplified models [11]. In this connection, from a software engineering viewpoint it could be very beneficial to be able to replace the variety of different cell models, e.g. ventricular, atrial, sinoatrial, involved in large-scale tissue and whole-heart simulations, by a single model, as proposed here, with different parameter values at different spacial positions.

**Model formulation and methods of parameter estimation**

*The archetypal cardiac cell model*

Cardiac cell membranes are composed of a biphospholipid layer, impermeable to charged particles and maintaining a non-zero equilibrium voltage potential across the membrane [47]. The layer is protruded by voltage-gated ion channels – large proteins that open and close depending on the instantaneous value of the voltage and allow in/outflux of ion currents. When a cell is “excited”, these currents cause the formation of a large transmembrane voltage excursion known as an action potential. The action potential propagates within the myocardium and signals cardiac cell contraction thus controlling the heartbeat – the main function of a living heart. To a first approximation the physiology of cell membranes can be modelled as an electrical circuit consisting of a capacitor $C_m$ representing the biphospholipid layer, and an active resistor supporting ionic currents $I_{ion}(E)$ representing ionic channels, that are connected in parallel, giving rise to the ordinary differential equation $C_m \dot{E} = I_{ion}(E)$ for the transmembrane voltage potential $E(t)$ [19]. The models of the ionic current $I_{ion}(E)$ encapsulate the electrophysiological properties of the cardiac membrane mentioned above and over the last 70 years have grown to a staggering complexity as discussed in the Introduction.

Here, we consider the following archetypal model for the action potential of a single cardiac cell

\[
\frac{d}{dt} E = \frac{1}{\varepsilon_1 \varepsilon_2} G_{Na} (E_{Na} - E) \theta(E - E_\tau) h + \frac{1}{\varepsilon_2} \left( g_2(E)n + G(E) \right), \tag{1a}
\]

\[
\frac{d}{dt} h = \frac{1}{\varepsilon_1 \varepsilon_2} f_h \left( \theta(E_\tau - E) - h \right), \tag{1b}
\]

\[
\frac{d}{dt} n = F_n(E) \left( \theta(E - E_\tau) - n \right). \tag{1c}
\]

The model takes the form of a set of piecewise-linear ordinary differential equations for the evolution in time $t$ of three state variables – the voltage $E$, and the gating variables $h$ and $n$ describing the inactivation of a fast inward current and the activation of the time-dependent channel of a slow outward currents, respectively. Here $\theta(\cdot)$ is the Heaviside step function and

\[
g_2(E) = g_{21} \theta(E_\tau - E) + g_{23} \theta(E - E_\tau), \quad F_n(E) = f_n \left( r \theta(E_\tau - E) + \theta(E - E_\tau) \right), \tag{1d}
\]

\[
G(E) = \begin{cases} 
  k_1(E_1 - E), & E \in (-\infty, E_\tau), \\
  k_2(E - E_2), & E \in [E_\tau, E_3), \\
  k_3(E_3 - E), & E \in [E_3, +\infty), 
\end{cases} \tag{1e}
\]

\[
E_2 = \left( \frac{k_1}{k_2} + 1 \right) E_\tau - E_1 k_1/k_2, \quad E_3 = \left( \frac{k_2}{k_3} + 1 \right) E_3 - E_2 k_2/k_3. \tag{1f}
\]

The constants $f_h$, $f_n$ are time scales of approach of channel gating variables $h$ and $n$ to their respective steady states, with the parameter $r$ further modulating $f_n$. The steady states of $h$ and $n$ are assumed to have a “perfect switch” behaviour and the parameter $E_\tau$ is the voltage value at which switching occurs. An additional unnamed ultra-fast gating variable that switches
instantaneously at \( E_\ast \) is implicitly included in the model by the factor \( \theta(E - E_\ast) \). This is a conventional Hodgkin-Huxley description of channel gating kinetics [26] and the perfect switching assumption is often used in other simplified models [18, 22]. In addition to the time-dependent channel, the slow outward current has an instantaneous voltage-dependent channel modelled by assumption is often used in other simplified models [18, 22]. In addition to the time-dependent channel, the slow outward current has an instantaneous voltage-dependent channel modelled by assumption is often used in other simplified models [18, 22].

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The archetypal model (1) was first introduced in [9] as an asymptotic embedding of the original Noble purkinje fiber equations [39] using a set of verifiable transformations with simplification errors that can be measured and controlled accurately. For full details we refer to [9], and here we only note briefly that the asymptotic embedding procedure takes into account the following empirical properties that the original Noble model has in common with the vast majority of other detailed ionic current models: (a) the large differences in the time-scales for evolution of state variables, (b) the large maximal value of the sodium current \( I_{Na} \) compared with other currents and (c) the quasi-stationary permeability of the \( I_{Na} \) ionic gates in certain potential ranges. Equations (1) differ from the system introduced in [9] only in that \( n \) instead of \( n^4 \) is used in equation (1a), with the goal to obtain a fully linear system with even simpler closed-form solutions and because parameter values will be adjusted anyway as discussed further below. The archetypal model (1) has already been used to derive asymptotic expressions for the conduction velocity restitution in cardiac tissues [49], to understand the formation of excitation waves [8], and most recently its fast-time subsystem was employed to elucidate the conditions for arrhythmogenesis and refractoriness in atrial tissue with myocyte-fibroblast coupling [34, 35].

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Problem (1) has 17 free parameters. Four of them are not intrinsic: \( \varepsilon_1, \varepsilon_2, E_{stim} \) and \( B \). The positive constants \( \varepsilon_1, \varepsilon_2 \in [0, 1] \) are asymptotic parameters embedded in the model to enable formal asymptotic analysis but both will be kept fixed to unity in the present study. The stimulus voltage \( E_{stim} \) and the basic cycle length \( B \) are typically specified as a part of an external periodic cell stimulation protocol, also cf. equation (3) below. The remaining 13 parameters are intrinsic to equations (1a) to (1f) and we represent them as the components of a column vector

\[
p = [k_1, k_2, k_3, E_1, E_{Na}, E_{\ast}, f_h, f_{r}, r, G_{Na}, g_{21}, g_{22}]^T.
\]

The objective of the study is to find appropriate values for these parameters as discussed next.
Target models and experimental data

We seek to estimate the values of the protocol-independent parameters (2) of the archetypal model (1) so that the model outputs reproduce the behaviour and the biomarkers of target detailed ionic models or experimental measurements. Here, as target models/data we consider the models of Noble [39], Luo-Rudy [28], and Courtemanche [15], as well as the measurements of [29] for rabbit ventricular myocytes.

The Noble model [39] describes the action potential of Purkinje fibre cells. It incorporates a sodium current and two different types of potassium current. The model is based on the Hodgkin-Huxley formulation adjusted to the action potential of Purkinje cells, which is significantly different from that of the squid giant axon in terms of plateau duration. This model is the first ever mathematical model of the action potential of cardiac myocytes and is the ancestor of most current detailed ionic models and the basis of the archetypal model (1).

The Luo-Rudy (LR) model [28] captures the single-cell ventricular action potential of guinea pigs. It includes six ionic currents (sodium, slow inward, time-dependent, time independent and plateau potassium currents and a background current) controlled by seven gate variables and a description of the intracellular calcium concentration.

The Courtemanche et al. (CRN) model [15] describes the action potential of human atrial myocytes. It has 13 ionic currents including formulations of K+, Na+ and Ca2+ currents and representations of pump, exchange and background currents. The model is capable of responding to rate changes, calcium channel inhibition and sodium-calcium pump exchanger blockade.

Machine readable implementations of three mathematical models are available from the CellML model repository [32], and are also included with our code [4]. The models are supplemented by stimuli currents $I_{stim}(t)$ that take the form of periodic trains of rectangular impulses with amplitude $I_s$, duration $t_s$, and period (basic cycle length) $B$,

$$I_{stim}(t) = I_s \left[ 1 + \text{sgn} \left( \frac{\pi t}{B} \right) \text{sgn} \left( \sin \left( \frac{\pi (t - B - 2t_s)}{B} \right) \right) \right].$$

(3)

The inclusion of these currents is known as “stimulation by current”. The currents are used to excite action potentials in the same way as in an experiment and provide an equivalent alternative to the “stimulation by voltage” given by equations (1g) and (1h) for the archetypal model.

The experimental recordings of [29] consist of measurements of action potential and intracellular calcium transient characteristics in isolated myocytes from sub-epicardial, mid-myocardial and sub-endocardial regions of the rabbit left ventricles. These measurements were recorded under both healthy and heart failure conditions. Results showed that in the heart failure group, AP duration and calcium transient duration were prolonged in both sub-epicardial and mid-myocardial cells. These changes were significant at lower stimulus frequencies but the relative effect diminished at higher frequencies. Below we will only consider the measurements of mid-myocardium cells in both healthy and failing myocytes.

Numerical solution and biomarkers

The archetypal model (1) and each of the three target models are integrated in time with a relative tolerance of $10^{-6}$ employing an adaptive-step, adaptive-order method for systems of stiff ordinary differential equations based on the numerical differentiation formulas of [46] as implemented in Matlab(TM) [53] functions ode15s and ode23. The resting states of the three target
models were used as their respective initial conditions. While a closed-form analytic solution of the archetypal models is available [9], a numerical solution is used here for consistency with the target models.

The following biomarkers are computed for each model.

1. Discretized voltage trace during the $k$-th basic cycle length period $t \in (kB, (k+1)B]$. More precisely, for the archetypal model this takes the form of a set of ordered pairs $\{(tk_i, E_{ki})\}$, consisting of discrete values of time $tk_i = kB + i\Delta t$ and discrete values of the voltage $E_{ki} = E(tk_i)$ and $M = B/\Delta t$. A step of $\Delta t = 0.1$ ms is used which is sufficient to resolve the traces.

2. Action potential duration at 90% of the voltage peak amplitude in the $k$-th basic cycle length period $I = (kB, (k+1)B]$, defined as the solution $A_k$ of the equation

$$E(A_k + kB) = 0.9\left(\max_{t \in I} E(t) - \min_{t \in I} E(t)\right),$$

that satisfies the condition $\dot{E}(A_k + kB) < 0$. We refer to this biomarker as APD$_{90}$, an abbreviation often used in the experimental and the physiological literature.

The same type of biomarkers are also available from the experimental measurements of McIntosh et al [29]. Their values were extracted from the published manuscript using the online tool WebPlotDigitizer [44].

**Parameter estimation**

Having defined appropriate biomarkers, we now compare the archetypal model (1) to each of the target models and the experimental data using a residual (also known in the literature as “error” or “cost” or “objective”) function of the form

$$R(p) = \frac{1}{2}\left(\frac{|A(p) - A'|}{\alpha'} + \frac{1}{M+1} \sum_{i=0}^{M} \left|\frac{E_i(p) - E_i'}{\max_{j=0..M} \delta_j - \min_{j=0..M} \delta_j}\right|\right).$$

Here, $E_i$ are the discrete values of the voltage trace, $A$ is the APD$_{90}$ of the archetypal model while the calligraphic symbols $E_i'$ and $A'$ denote the corresponding values for the target models and data. In particular, the biomarkers in the initial basic cycle period $k = 0$ are used and, for brevity, the subscript $k$ is omitted. This form of the residual measures the discrepancy in the morphologies of the action potentials between the archetypal and the target models/data. The first term of the residual allows the fitting algorithm to assign extra weight to matching the value of APD$_{90}$ when optimising action potential morphology. We remark that, in general, for a complete comparison of the models, the residual needs to include differences between all dynamical variables of the models compared, including all gating variables and ion concentrations. This however, is not possible because of the difference in model formulations, in particular, because different models include different ionic currents and have different dynamical variables. Similarly, model quantities are not easily measured in experiments. The voltage is often the only quantity in common between different models and between models and data.

We now compute parameter values $\tilde{p}$ of the archetypal model (1) such that the residual (5) is minimised, in symbolic form,

$$\tilde{p} = \operatorname{arg\,min}_{p \in \Omega} R(p),$$
where $\Omega \subset \mathbb{R}^{13}$ is the ball $|p - p_0| < \tau$ centred at the default values $p_0$ of the archetypal model parameters given in the second column of Table 1 and radius $\tau = |p_0|$. More explicitly, for the parameter estimation function $\arg\min(\cdot)$, we use a MATLAB implementation [17] of the bounded gradient-free Nelder-Mead simplex method [37, 27] for minimisation of real-valued multivariate functions.

We provide an open-source numerical code including the models and methods described in this section. The code is permanently available at [4] and can be used by the readers to reproduce the results described below an/or to apply the methodology to other detailed models and data of their own interest.

**Results and analysis**

**Parameter sensitivity analysis**

Prior to estimating the parameter values (2) of the archetypal model (1), we perform a local sensitivity analysis in order to observe how each parameter affects the action potential morphology and to establish whether it is necessary to include all thirteen of them in the optimazation search (6). A formal local sensitivity analysis, see e.g. [50], involves computing a sensitivity matrix $S$ whose entries $S_{ij}(t)$ describe the normalised effect of perturbing the $j$-th parameter on the $i$-th state-variable, defined as

$$S_{ij}(t) \equiv \left. \frac{\partial x_i(t; p)}{\partial p_j} \right|_{p=p_0} = \left. \frac{\partial \log x_i(t; p)}{\partial \log p_j} \right|_{p=p_0}, \quad x = [E, h, n]^T. \quad (7)$$

However for clarity, instead of illustrating the elements of the sensitivity matrix, we employ a simpler and more intuitive approach. We vary the value of each of the thirteen model parameters by $\pm 20\%$ from their default values listed in the second column of Table 1 at all other parameter values fixed and observe how this variation affects the traces of the voltage $E$ and the slow gating variable $n$. Figure 1 illustrates the results from this experiment and also serves to illustrate clearly the effect each of the parameters has on the action potential morphology. For example, the peak membrane potential (PMP) is controlled by the value of $E_{Na}$ while the resting membrane potential (RMP) is influenced by $E_1$ only. Some of the parameter values affect the action potential morphology in a more complex way. For instance, $E_\ast, E_\dagger, g_{22}$ and $f_n$ control repolarisation, but $E_\ast$ and $E_\dagger$ also contribute to the duration of the plateau. The fast gating variable $h$ exhibits negligible variation from its quasi-stationary value $\bar{h} = 1$ apart from two very short time intervals during the front and the back of the action potential and for this reason is not included in Figure 1.

**Parameter estimation**

Table 1 lists the results of applying the minimisation procedure (6) to estimate the parameter values (2) of the archetypal model (1) so that it closely reproduces the action potential morphology of the target models [39, 28, 15] and the target data [29]. The agreement obtained in action potential morphology between the archetypal model and the targets is shown in Figure 2. Target models and the archetypal model were stimulated at a basic cycle length $B = 1500$ ms. The archetypal model is able to capture the action potential morphologies of all target models and data used well. The fits of the archetypal model to the CRN [15] and the LR [28] models show the biggest discrepancy as seen in Table 1(b) and Figures 2(b) and (c). The discrepancies are most significant in the neighbourhoods of the post overshoot drop and plateau regions. In particular, the archetypal model is somewhat inaccurate in capturing the deep notch produced by the CRN [15] model. This occurs because the archetypal model lacks transient outward currents $K^+$ and $Na^+ / Ca^{2+}$ exchanger currents [45, 42] that act to form notch phase in the CRN [15].
Figure 1: Sensitivity of the profiles of the voltage $E$ and the slow gating variable $n$ of archetypal model (1) to the variation of a single parameter as denoted by the corresponding symbol in each panel. In all panels, the model outputs generated from the default parameter values in Table 1 are shown by solid black line. The dashed blue line and dash-dotted red line are the model outputs obtained with $-20\%$ and $+20\%$ perturbation from the default, respectively.

The fit of the archetypal model to the Noble model [39] is rather satisfactory. The peak voltage is controlled by the fast sodium current. The peak voltage differs in different regions of the heart due to variation in magnitude of the sodium current. In the archetypal model, the magnitude of the fast inward current is modulated by parameter $E_{Na}$ only. The higher the peak voltage, the higher the value of $E_{Na}$. This is consistent with our finding (Figure 2 and Table 1), where the LR [28] model has the highest $E_{Na}$ because the model exhibits the largest action potential amplitude. Plateau phase is the long phase of the action potential during which the membrane potential remains depolarised and changes more slowly. This occurs due to balance...
Table 1: Estimates of the parameter values of the archetypal model (1) to selected target models and data as described in text. (a) Parameter values fitted by (6). (b) The residual error in action potential morphology (5) between the archetypal model and targeted models/data. The time is measured in ms, other units are given in the table if dimensional.

between some inward and outward currents [45]. In the archetypal model, $E^*$ is one of the parameters that controls the voltage value at the plateau region. The estimated value is consistent for each model, where $E^*$ in the LR [28] model is the biggest since the LR model has the largest value of voltage during this phase, while the smallest is shown by the Noble model.

To fit the archetypal model to isolated cardiac ventricular cell data we use a pacing rate of 0.3Hz ($B = 3333$ ms) at both healthy and heart-failure conditions as this is the basic cycle length employed in [29]. Figure 2(d) shows the action potential morphologies after the fitting process and the new estimated parameter values are shown in last two columns of Table 1. Overall, the archetypal model exhibits good correspondence with the targeted data, with minor discrepancies in the plateau region. The average relative error between the action potentials is relatively small as seen in Table 1(b). Figure 3(c) shows that after fitting the archetypal model to heart-failure data the parameter values most strongly affected are those related to the $n$ gating variable (the slow-gating potassium channels), namely $k_1$, $k_2$, $k_3$, $r$ and $g_{22}$. These parameters need to be adjusted in order to compensate the large APD$_{90}$ exhibited in the heart-failure group, which is commonly reported due to down-regulation of potassium current [7, 1]. Other archetypal parameter values like $E_{Na}$ and $E_1$ are similar in both heart-failure and healthy cells since the action potentials have identical amplitude and resting membrane potential.

Tests of APD restitution and computational speedup

In order to test the validity of our parameter estimation results beyond the conditions at which they are obtained, we compare the action potential duration (APD) restitution curves for each of the target models and the experimental data to corresponding curves computed using the parameter estimates reported in Table 1. Figure 3(a) demonstrates excellent agreement between the restitution curves of the authentic target models and the fitted archetypal models, while Fig-
Figure 2: Agreement in action potential morphology between the archetypal model at parameter values listed in Table 1 (broken lines) and corresponding target models (panels (a)-(c), solid lines) and experimental data (panel (d), solid lines). Model names are specified in the panel legends.

Figure 3(b) shows similarly good agreement with the restitution curves of the cells from healthy and heart-failure groups. In particular, at a given stimulation frequency failing myocytes exhibit larger APD_{90} than healthy myocytes. At high stimulation frequency the APD_{90} values for both groups show a less pronounced difference. The archetypal model is slightly better able to reproduce the accurate APD restitution curve for the HF myocyte, compared to healthy myocyte. For healthy myocyte, the discrepancy occurs at several stimulation frequencies, and it gets pronounced at stimulation frequency larger than 2 Hz, where the archetypal model produces smaller APD_{90} than the experimental data from healthy myocytes.

Because of the practical value of APD restitution curves, it is important that the fitted archetypal model is able to reproduce the restitution behaviour of the target models and data. These curves describe the dependence of the APD on the duration of the preceding diastolic interval (DI). Nolasco and Dahlen [40] noted that in a single-cell setting and with a fixed period of excitation, a slope of the APD(DI) curve greater than one indicates instability of the train of action potentials. For this reason, the restitution curves are considered an important tool in understanding instabilities of excitation waves leading to onset of cardiac arrhythmias [56] and is routinely measured experimentally, e.g. in the experimental work [29] that we compare with.

In order to quantify the computational speedup gained by using the archetypal model with fitted
Parameter values in comparison with authentic target models we measured the time taken using each model to compute numerically a train of 1000 action potentials at a fixed value of the basic cycle length $B = 1000$ ms. With the numerical methods described above in the paper, the archetypal model takes approximately about 180 sec to complete the numerical simulation with any set of parameter values. The LR [28] and CRN [15] model require 1246 sec and 1634 sec, respectively for the same simulation. Thus we conclude that using the archetypal model is 6 times faster than using the Luo-Rudy model [28] and 9 times faster than the using Courtemanche et al. model [15]. While the comparison was demonstrated at the single-cell level, we expect the improvement in the computational speed-up would also be seen in simulations of whole-heart or 3D tissue with realistic geometries. The numerical speedup advantage of using the archetypal model is likely even more pronounced in case of comparison with contemporary models more detailed than the LR [28] and the CRN [15] models. We also recall that the archetypal has closed form solutions that can be evaluated directly irrespective of parameter values used and this can be exploited to further reduce computational expenses or even eliminate the need of computation entirely.

**Discussion**

**Summary**

Contemporary mathematical models of single-cell cardiac electrical excitation have become immensely detailed. Along with increasing physiological realism such model complexity leads to parameter value uncertainty, high computational cost and barriers to mechanistic understanding. There is thus a need for conceptually and mathematically simple but physiologically accurate reduced models of the cardiac action potential. A single-cell cardiac excitation model that replicates the phase-space geometry of detailed cardiac models but is much simpler in both functional form and number of free parameters was derived in [9] and applied to a number of idealised problems [49, 8, 34, 35]. In order to render the archetypal model of [9] also practically applicable to the description of physiological measurements and to whole-heart and tissue numerical simulations, in this study we report a robust method for estimation of the parameter values of the model so as to approximate the action potential biomarkers of contemporary detailed ionic models as well as experimental data from direct wet-lab cell measurements. The parameter estimation procedure relies on the well-known and popular Nelder-Mead method.
for minimisation of multi-variable functions by direct simplex search. Here, the optimal parameter values of the archetype are determined by minimising the residual difference between the morphologies of single-stimulus action potentials of the archetypal model and the target model/data. The procedure is then applied to 5 test cases, namely (a) to the authentic Noble model [39], the precursor to all detailed cardiac ionic current models; (b) to the Luo-Rudy ventricular cell model [28], the original second generation model; (c) to the Courtemanche, Ramirez and Natel atrial cell model [15], a popular modern cardiac system, as well as (d,e) to wet lab experimental measurements of rabbit ventricular cells from both healthy and heart-failure samples [29]. In all cases, sets of values of the archetypal model parameters have been found so that the morphologies of stable single-stimulus action potential target transients are well reproduced. As this is an optimisation problem in high-dimensional parameter space that may have several local minima, it is difficult to ascertain if a given solution of the minimisation procedure using Nelder-Mead method is the best possible one. To further test and validate the method, action potential duration restitution curves are also computed and compared to those of the target models and data, again with excellent agreement. We conclude that compared to more sophisticated parameter estimation methods for cardiac models such as maximum-likelihood estimation [31], principal-axis fitting [58], genetic algorithms [51, 25], as well as the widely practised empirical “hand-tuning” of free parameters [5, 30], our rather straightforward approach provides comparable quality of approximation and performs remarkably well. This is likely due to the generic structure and the small number of parameters of the archetypal model we consider. An open-source Matlab(TM) implementation of the models and methods is made permanently available at [4] and can be used by the readers to fit the archetypal model to models and data of their own choice.

Many processes that occur in excitable cells, including cardiac cells are still not fully understood. None of the detailed models are themselves ultimate, rather they are continually improved and in some cases discarded in the light of new experimental measurements. The approach of parameter adjustment used in the present work, is a way to accurately model cellular electrical excitation even if fine details of cell physiology are not included.

Perspectives for future work
A major advantage of the archetypal model (1) is that it has three features that, to our knowledge, were not available in combination for any other model prior to this work.

(a) Model (1) admits both asymptotic and closed-form analytical solutions, see [9] and the extended discussion in the Introduction.
(b) Model (1) captures essential cardiac excitability characteristics such as slow repolarization, slow subthreshold response, fast accommodation, variable peak voltage, and front dissipation that other ad hoc simplifications do not, see [9].
(c) With the results of the present work, model (1) can now be fitted to reproduce accurately the electrophysiological responses of a variety cardiac cell types.

There is a large body of already developed theory for conceptual understanding of the dynamics of nonlinear wave processes in cardiac tissue that underlie arrhythmias, fibrillation and defibrillation [43]. The unique features of our model open the way to apply this theory to realistic experimental and clinical situations with a dramatic increase of quantitative accuracy. Particular examples include: applying known mathematical conditions of propagation block in terms of fitted myocyte parameters, making realistic analytical estimates of the vulnerability to extrastimuli, realistic prediction of the frequency and stability of functional re-entrant circuits and likelihood of recurrent fibrillation after a defibrillating shock. Similarly, robust relationships
between controllable cell parameters and outcome of experiments can now be obtained that will have the potential to allow more confident planning of experiments and facilitate development and improvement of antiarrhythmic strategies. These applications are planned for future work.

We wish to comment, in particular, on the perspectives that our works opens for the development of novel efficient numerical methods for excitation propagation and tissue simulations. The archetypal model (1) allows computational speed-up to be achieved in two essentially different ways.

(a) Firstly, a straightforward replacement of large detailed ion current models by model (1) with appropriately fitted parameters will lead to a several-fold speed-up as measured above for the LR and the CRN models. This is already a significant improvement as a speed-up of 6 to 9 times is comparable to the speed-up of using e.g. lookup tables [11].

(b) Secondly, more important benefits may be achieved by employing the asymptotic structure readily encoded in (1) to split the model to a fast-time subsystem describing only the front of the action potential coupled to a slow-time subsystem describing its plateau and recovery phases.

To explain the importance of (b), we note that physiological cell ionic models are stiff because the dynamics of the action potential front is orders of magnitude faster than the dynamics during the plateau and recovery. This requires very small time and spacial discretisation steps to be used in numerical schemes to adequately resolve propagating action potentials. After asymptotic splitting of the archetypal model (1), a numerical scheme can be used that will require a fine resolution only for solution of the fast-time front subsystem and allow a much coarser resolution to be used for solution of the non-stiff plateau and recovery subequations, e.g. the heterogeneous multiscale method [59]. However, hybrid asymptotic-numerical methods are not well developed in higher dimensions, which are needed for calculation of activation sequences. The equation of motion for the front is a partial differential equation of motion of a line (in 2D) or surface (in 3D). One immediate difference to the 1D case is that propagation of the front no longer depends only on the pre-front voltage and slow variables, but also on its own spatial configuration. Fortunately, unless the shape of the front deviates very strongly from plain, the effect of its shape can be taken into account via its mean curvature that can be easily incorporated [48]. This approach can be used to describe normal activation sequences in the heart, when the graph of the front solution in the space-time is a manifold without internal discontinuities. More serious challenges occur if there are propagation blocks and/or wave breaks, which introduce discontinuities of the front manifold in space-time. In such cases, a separate asymptotic description for the codimension-two areas, the wave break trajectories and the propagation block loci, are needed; obtaining such asymptotic description is another important direction for further research. These issues will be much easier to tackle using our simple (and now accurate) archetypal model rather than complex physiologically detailed cell ionic models.

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