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Reducing runtime of an electrophysiological cancer cell model from X to Y and live simulation dashboard

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Abstract: Through a reimplementation in Rust and numerical optimization approaches we were able to reduce the runtime of the A549 electrophysiological cancer cell model [1] from X to Y.

Keywords: Please insert your keywords here, separated by commas.

1 Introduction

Lung cancer is one of the most widespread pathologies worldwide and its mechanisms, specifically those of individual A549 cells, are not well understood. Computational techniques can help with a better understanding of the behaviour of these cancer cells. We work with the A549 model introduced in [1], together with a calcium channel extension introduced in [2], reimplementing the model in the Rust programming language and performing a number of numerical optimizations such as adaptive timestepping. We also verify the model's performance in Floating mode, as compared to an individual simulation of the estimated number of channels. We introduce a visualisation approach of the entire model in the form of a live simulation dashboard¹. The entire source code behind this simulation is freely available on GitHub², and reusable through three different channels: the simulation interface (powered by WebAssembly), the Rust linkable library implementation and a Python package. These three interfaces all originate from the same source code and our aim behind a distribution in this way is to make the simulation as accessible as possible. In order to find appropriate parameters for the model, an optimization procedure is performed. Multiple opti-

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mization approaches for the solution of the corresponding inverse problem (fitting model parameters to measurement data) are put in comparison. The measurements are obtained using a *Patch-Clamp System*, where one records the current through the membrane given a voltage protocol.

2 Methods

A cell's membrane consists of multiple ion channels, categorized into $M \in \mathbb{N}$ different types. Each ion channel is represented in one of $N_{s,k} \in \mathbb{N}$ states, which, in physical terms, is related to a positional configuration of a protein within the ion channel. Only some states can be observed directly, and its development only depends on the one previous state. Hence, we are working with a Hidden Markov Model (HMM). For many ion channel categories, their transition probabilities are voltage or ion-concentration dependent.

The whole cell current $I: T \to \mathbb{R}$ over time $t \in T \subset \mathbb{R}^+$ is then obtained as the sum of all individual channel contributions $I_k, k \in \{1, ..., M\}$ over $M \in \mathbb{N}$ channel types

$$I(t) := \sum_{k=1}^{M} N_k I_k(t) = \sum_{k=1}^{M} N_k g_k p_{o,k} \left(V(t) - E_k \right) , \quad (1)$$

where N_k is the number of channels of type $k \in \{1,...,M\}$, g_k is the respective ion channel's conductivity, $p_{o,k} \in [0,1]$ is the probability of observing the channel in a state where an ion current can flow ("open states"), $V: T \to \mathbb{R}$ is the voltage across the membrane and $E_k \in \mathbb{R}$ the reversal potential.

Within the simulation, we sample the state and current at discrete time points $T_{\rm meas}\subset T$, for example

$$T_{\text{meas}} := \left\{ t_n := \sum_{i=0}^n (\Delta t)_i \mid n \in \mathbb{N}_0 \mid n < N_t \right\}$$

for N_t measurements with step size $(\Delta t)_n$, which may be chosen equally large for all $n \in \{0,...,N_t-1\}$. We adapt this time interval $(\Delta t)_n \in \mathbb{R}^+$ per simulation step based on a state change heuristic, cf. Section 2.1.

At each time step,

$$s_{k,n+1} = H_k(V(t_n), C(t_n), t_n) s_{k,n}$$
 (2)

¹ https://in-silico-cancer-cell.waldert.at/

 $^{{\}bf 2}\ https://github.com/MrP01/InSilicoCancerCell$

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where $s_{k,n} \in \mathbb{R}^{N_{s,k}}$ is the state vector of ion channel type k at the n-th time step, $H_k\left(V,C,t_n\right) \in \mathbb{R}_+^{N_{s,k} \times N_{s,k}}$ the transition matrix for type k with $\sum_{j=1}^{N_{s,k}} \{H_k\}_{i,j} = 1 \ \forall i, \ V(t_n)$ the voltage across the membrane at time t_n and $C(t_n) \in \mathbb{R}^4$ the concentrations of Kalium, Calcium, Sodium and Chlorine at time t_n . We initialize the simulation at $t_0 = 0$ with $s_{k,0} = (1,0,...,0)^T$ for all k.

Given the state s_k , current measurements are then simply

$$I := (I(t_0), I(t_1), ..., I(t_{N_t-1}))^T \in \mathbb{R}^{N_t},$$

with I as stated above in Equation (1) and

$$p_{o,k} = \sum_{n_s \in \mathcal{S}_{o,k}} \{ \boldsymbol{s}_{n,k} \}_{n_s} \,,$$

where $S_{o,k}$ is the set of all states contributing to the ion channel current, the "open states".

2.1 Adaptive Timestepping

In order to accelerate the simulation in areas where there is little change to the dynamics, we choose an adaptive step size based on

$$(\Delta t)_{n+1} = (\Delta t)_n \left(\frac{\Delta^{\text{tol}}}{\sum_{k=1}^{M} N_k \|\mathbf{s}_{k,n+1} - \mathbf{s}_{k,n}\|_2} \right)^{1/2},$$
(3)

for all n, where $\Delta^{\rm tol} \in \mathbb{R}^+$ is a measure for the allowed state change in between steps. When the state changes too quickly in between time steps, the above heuristic will decrease $(\Delta t)_{n+1}$ and vice-versa. In principle, it would be feasible to apply the adaptive timestepping to each ion channel type individually, however this would make current sampling and data synchronization between channels hard to realize, considering the ion concentration dependence of H_k . Within this paper, we set $\Delta^{\rm tol} = 2 \cdot 10^{-7}$.

2.2 Inverse Problem

When regarding the cell model as a whole, the number of ion channels N_k per type k may be put into a configuration vector $\mathbf{N} := (N_1,...,N_M)^T \in \mathbb{N}_0^M$ and the total simulated current I sampled at measurement points T_{meas} can be expressed as a matrix-vector product

$$I = \sum_{k=1}^{M} N_k I_k = AN, \qquad (4)$$

where $A \in \mathbb{R}^{N_t \times M}$ is the matrix of all current measurements per channel type.

Tab. 1: Ion Channel Types: Most fitting configuration and corresponding references.

Channel Type	Channel Count $N_{m k}$	Reference
Kv13	1	[3]
Kv31	1	[3]
Kv34	1	[3]
Kv71	1	[3]
KCa11	1	[3]
KCa31	1	[3]
Task1	1	[3]
CRAC1	1	[3]
TRPC6	1	[3]
TRPV3	1	[3]
CLC2	1	[3]

Given the individual ion channel type models' parameters, which we know from literature (cf. Table 1), the question that remains is how many channels there are of each type to fit the measurements. This problem can be solved using a number of optimization approaches.

However, the formulation in Equation (4) also gives rise to a least-squares formulation, by projecting the measured current into the space of all individual channel currents. More specifically, we want to find

$$N_{\text{opt}} = \underset{N \in \mathbb{N}_0^M}{\operatorname{argmin}} \|AN - I_{\text{meas}}\|_2^2,$$
 (5)

with $I_{\text{meas}} \in \mathbb{R}^{N_t}$ the experimentally measured current. The most important constraint here is that of integer non-negativity, $N_{\text{opt}} \in \mathbb{N}_0^M$, which makes this problem hard to solve directly.

2.3 Implementation and Usage

The core simulation is implemented in the Rust programming language [4]. Each channel is

2.4 Live Simulation

The live

3 Results

3.1 Adaptive Timestepping

With the standard forward iteration approach, the simulation takes 412 million steps to simulate over all voltage protocols and cell cycle phases, while the adaptive timestepping only requires 23 million steps for the same configuration. While

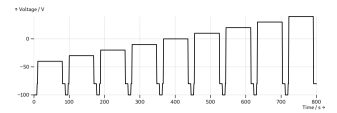


Fig. 1: The *activation* voltage pulse protocol V(t) used for the measurement and simulation. This is the voltage set across the A549 cell's membrane using a Patch-Clamp-System. The corresponding current is then recorded and depicted in Figure 2 along with its simulation result.

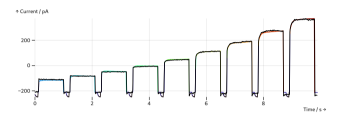


Fig. 2: The simulated and measured current response I(t) across the membrane of an A549 cancer cell, given the *activation* voltage protocol V(t) in Figure 1, recorded in the G0 phase of the cell cycle.

keeping the same level of accuracy when matching with the experimental data, our adaptive timestepping method is therefore 17 times faster than the standard approach, on average.

4 Outlook

Numerically, the stability of the simulation varies greatly with the time step state change tolerance $\Delta^{\rm tol}$, this could be improved using a higher-order integration scheme. Regarding the simulation dashboard, there are still many adjustments that could improve and enable further usage perspectives.

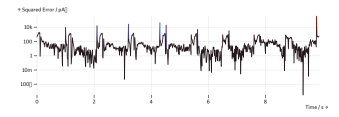


Fig. 3: Pointwise squared error between the measured current $I_{\rm meas}$ and the simulated current I, showing potential systematic problems within the computational model's representation of the ion channels as compared to the experimental results.

Tab. 2: Comparison of Optimization Approaches

Algorithm	Accuracy
Particle Swarm Optimization	1
Gradient Descent	2
LBFGS	3
QR-based LSQ	4
NNLS	5

Author Statement

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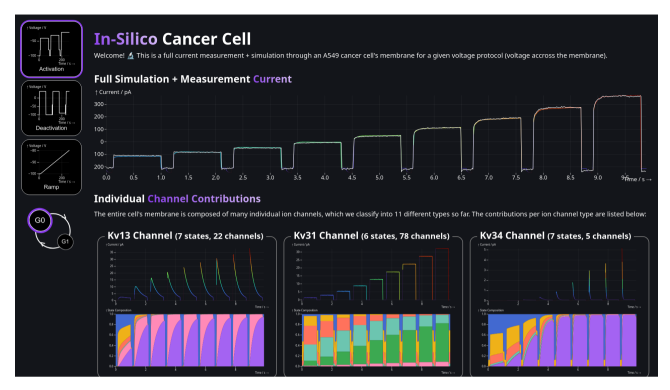


Fig. 4: Screenshot of the live simulation dashboard, available here: https://in-silico-cancer-cell.waldert.at/. This is the graphical user interface to the simulation written in Rust. The full simulation, completing within around 500ms, runs live in the browser using Rust's bindings to WebAssembly. One can choose the voltage protocol and cell cycle phase (G0/G1) in the menu on the left, while the full current and individual channel currents and internal states can be seen on the right. The remaining ion channel types have been cut off on this screenshot.