Stochastic Representations of Ion Channel Kinetics and Exact Stochastic Simulation of Neuronal Dynamics

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Abstract

In this paper we provide two representations for stochastic ion channel kinetics, and compare the performance of exact simulation strategies with different, commonly used, approximate strategies. The first representation we present is a random time change representation, popularized by Thomas Kurtz, with the second being analogous to a “Gillespie” representation. Stochastic models of ion channel kinetics typically consist of an ordinary differential equation, governing the voltage, coupled with a stochastic jump process, governing the number of open ion channels. The processes are coupled because the parameters of the ODE for the voltage depend upon the number of open ion channels, and the propensity for the opening and closing of the channels depends explicitly upon the time-varying voltage. Exact stochastic algorithms are provided for the different representations, which are preferable to either (a) fixed time step or (b) piecewise constant propensity algorithms, which still appear in the literature. As examples, we provide versions of the exact algorithms for the Morris-Lecar conductance based model, and detail the error induced, both in a weak and a strong sense, by the use of approximate algorithms on this model. We include ready-to-use implementations of the random time change algorithm in both XPP and Matlab. Finally, through the consideration of parametric sensitivity analysis, we show how the representations presented here are useful in the development of further computational methods. The general representations and simulation strategies provided here are known in other parts of the sciences, but less so in the present setting.

1 Introduction

Fluctuations in membrane potential arise in part due to stochastic switching in voltage-gated ion channel populations [11, 28, 47]. We consider a stochastic modeling, i.e.
master equation, framework \[5, 10, 12, 29, 48\] for neural dynamics, with noise arising through the molecular fluctuations of ion channel states. We consider model nerve cells that may be represented by a single isopotential volume surrounded by a membrane with capacitance \(C > 0\). Mathematically, these are hybrid stochastic models which include components, for example the voltage, that are continuous and piecewise differentiable and components, for example the number of open potassium channels, that make discrete transitions or jumps.

These models are typically described in the neuroscience literature by providing an ODE governing the absolutely continuous portion of the system, which is valid between jumps of the discrete components, and a chemical master equation providing the dynamics of the probability distribution of the jump portion, which itself depends upon the solution to the ODE. In this paper, we take a different perspective. We will introduce here two pathwise stochastic representations for these models that are similar to Itô SDEs or Langevin models. The difference between the models presented here and Langevin models is that here the noise arisres via stochastic counting processes as opposed to Brownian motions. These representations give a different type of insight into the models than master equation representations do, and, in particular, they imply different exact simulation strategies. These strategies are well known in some parts of the sciences, but less well known in the current context \[5\].

From a computational standpoint the change in perspective from the master equation to pathwise representations is useful for a number of reasons. First, the different representations naturally imply different exact simulation strategies. Second, and perhaps more importantly, the different representations themselves can be utilized to develop new, highly efficient, computational methods such as finite difference methods for the approximation of parametric sensitivities, and multi-level Monte Carlo methods for the approximation of expectations \[2, 4\]. Third, the representations can be utilized for the rigorous analysis of different computational strategies and for the algorithmic reduction of models with multiple scales \[3, 7\].

We note that the types of representations and simulation strategies highlighted here are known in other branches of the sciences, especially operations research and queueing theory \[19, 22\], and stochastically modeled biochemical processes \[1, 5\]. See also \[36\] for a mathematical treatment of such stochastic hybrid systems and some corresponding approximation algorithms. However, the representations are not as well known in the context of computational neuroscience and as a result approximate methods for the simulation of sample paths including (a) fixed time step methods, or (b) piecewise constant propensity algorithms, are still utilized in the literature in situations where there is no need to make such approximations. Thus, the main contributions of this paper are: (i) the formulation of the two pathwise representations for the specific models under consideration, (ii) a presentation of the corresponding exact simulation strategies for the different representations, and (iii) a comparison of the error induced by utilizing an approximate simulation strategy on the Morris-Lecar model. Moreover, we show how to utilize the different representations in the development of methods for

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\[1\] For an early statement of an exact algorithm for the hybrid case in a neuroscience context see \[9\], Equations 2-3. Strassberg and DeFelice further investigated circumstances under which it is possible for random microscopic events (single ion channel state transitions) to generate random macroscopic events (action potentials) \[45\] using an exact simulation algorithm. Keener and Newby used an exact algorithm in a recent study of channel noise dependent action potential generation in the Morris-Lecar model \[24, 34\].
parametric sensitivity analysis.

The outline of the paper is as follows. In Section 2 we develop the different representations and provide the corresponding numerical methods for the two different representations. In Section 3 we present a stochastic representation for a particular conductance based model, the planar Morris-Lecar model, with a single ion channel treated stochastically. Here, we illustrate the numerical strategy on this example and provide in the appendix both XPP and Matlab code for this model. In Section 4 we present an example of a conductance based model with more than one stochastic gating variable, namely the Morris-Lecar model with both ion channel types (calcium channels and potassium channels) treated stochastically. To illustrate both the strong and weak divergence between the exact and approximate algorithms, in Section 5 we compare trajectories and histograms generated by the exact algorithms and the piecewise constant approximate algorithm. In Section 6 we show how to utilize the different representations presented here in the development of methods for parametric sensitivity analysis, which is a powerful tool for determining parameters to which a system output is most responsive. In Section 7 we provide conclusions and discuss avenues for future research.

2 Two stochastic representations

In Section 2.1 we heuristically present the random time change representation for the relevant models, whereas in Section 2.2 we present a “Gillespie” representation. In each section, we provide the corresponding numerical simulation strategies. See [5] for a development of the representations in the biochemical setting, and see [26, 27] for a more general rigorous mathematical development. For a more recent addition to the literature, see also [36] in which the authors, among other things, explicitly prove that the models satisfying the representations below are Markov processes satisfying the corresponding master equation.

2.1 Random time change representations

We begin with an example. Consider a model of a system that can be in one of two states, $A$ or $B$, which can represent “closed” and “open,” respectively. We model the dynamics of the system by assuming that the dwell times in states $A$ and $B$ are exponential random variables with parameters $\alpha > 0$ and $\beta > 0$, respectively. A graphical representation for this model is

$$A \xrightleftharpoons{\alpha}{\beta} B.$$  \hspace{1cm} (1)

The usual formulation of the stochastic version of this model is to note that the probability that a closed channel opens in the next increment of time $\Delta s$ is $\alpha \Delta s + o(\Delta s)$, whereas the probability that an open channel closes is $\beta \Delta s + o(\Delta s)$. This type of stochastic model is often described mathematically by providing the “chemical master
Thus, the propensity for seeing another jump is precisely $\lambda(s)$. Returning to the discussion directly following (1), and noting that $X_1(s)+X_2(s) = 1$ for all time, we note that the propensities of reactions 1 and 2 are

$$\lambda_1(X(s)) = \alpha X_1(s), \quad \lambda_2(X(s)) = \beta X_2(s).$$

Combining all of the above implies that we can represent $R_1$ and $R_2$ via

$$R_1(t) = Y_1 \left( \int_0^t \alpha X_1(s) ds \right), \quad R_2(t) = Y_2 \left( \int_0^t \beta X_2(s) ds \right),$$
and so a pathwise representation for the stochastic model (1) is

$$X(t) = X_0 + Y_1 \left( \int_0^t \alpha X_1(s) ds \right) \begin{pmatrix} -1 \\ 1 \end{pmatrix} + Y_2 \left( \int_0^t \beta X_2(s) ds \right) \begin{pmatrix} 1 \\ -1 \end{pmatrix},$$

(2)

where $Y_1$ and $Y_2$ are independent, unit-rate Poisson processes.

Slightly more generally, suppose now that $X_1(0) + X_2(0) = N \geq 1$. For example, perhaps we are now modeling the number of open and closed ion channels out of a total of $N$, as opposed to simply considering a single such channel. Suppose further that the propensity, or rate, at which ion channels are opening can be modeled as

$$\lambda_1(t, X(t)) = \alpha(t) X_1(t)$$

and the rate at which they are closing has propensity

$$\lambda_2(t, X(t)) = \beta(t) X_2(t),$$

where $\alpha(t), \beta(t)$ are non-negative functions of time, perhaps being voltage dependent.

That is, suppose that for each $i \in \{1, 2\}$, the conditional probability of seeing the counting process $R_i$ increase in the interval $[t, t+h)$ is $\lambda_i(t, X(t)) h + o(h)$. The expression analogous to (2) is now

$$X(t) = X_0 + Y_1 \left( \int_0^t \alpha(s) X_1(s) ds \right) \begin{pmatrix} -1 \\ 1 \end{pmatrix} + Y_2 \left( \int_0^t \beta(s) X_2(s) ds \right) \begin{pmatrix} 1 \\ -1 \end{pmatrix}. \quad (3)$$

Having motivated the time dependent representation for the simple model above, we turn to the more general context. We now assume a jump model consisting of $d$ chemical constituents (or ion channel states) undergoing transitions determined via $M > 0$ different reactions. For example, in the toy model above, the chemical constituents were $\{A, B\}$, and so $d = 2$, and the reactions were $A \rightarrow B$ and $B \rightarrow A$, giving $M = 2$. We suppose that $X_i(t)$ determines the value of the $i$th constituent at time $t$, so that $X(t) \in \mathbb{Z}^d$, and that the propensity function of the $k$th reaction is $\lambda_k(t, X(t))$. We further suppose that if the $k$th reaction channel takes place at time $t$, then the system is updated according to addition of the reaction vector $\zeta_k \in \mathbb{Z}^d$,

$$X(t) = X(t-) + \zeta_k.$$

The associated pathwise stochastic representation for this model is

$$X(t) = X_0 + \sum_k Y_k \left( \int_0^t \lambda_k(s, X(s)) ds \right) \zeta_k,$$

(4)

where the $Y_k$ are independent unit-rate Poisson processes. The chemical master equation for this general model is

$$\frac{d}{dt} P_{X_0}(x, t) = \sum_{k=1}^M P_{X_0}(x - \zeta_k, t) \lambda_k(t, x - \zeta_k) - P_{X_0}(x, t) \sum_{k=1}^M \lambda_k(t, x),$$

where $P_{X_0}(x, t)$ is the probability of being in state $x \in \mathbb{Z}_{\geq 0}^d$ at time $t \geq 0$ given an initial condition of $X_0$.  

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When the variable \( X \in \mathbb{Z}^d \) represents the randomly fluctuating state of an ion channel in a single compartment conductance based neuronal model, we include the membrane potential \( V \in \mathbb{R} \) as an additional dynamical variable. In contrast with neuronal models incorporating Gaussian noise processes, here we consider the voltage to evolve deterministically, conditional on the states of one or more ion channels. For illustration, suppose we have a single ion channel type with state variable \( X \). Then, we supplement the pathwise representation (4) with the solution of a differential equation obtained from Kirchoff’s current conservation law:

\[
C \frac{dV}{dt} = I_{\text{app}}(t) - I_V(V(t)) - \left( \sum_{i=1}^{d} g_i^o X_i(t) \right) (V(t) - V_X) \tag{5}
\]

Here, \( g_i^o \) is the conductance of an individual channel when it is the \( i \)-th state, for \( 1 \leq i \leq d \). The sum gives the total conductance associated with the channel represented by the vector \( X \); the reversal potential for this channel is the constant \( V_X \). The term \( I_V(V) \) captures any deterministic voltage-dependent currents due to other channels besides channel type \( X \), and \( I_{\text{app}} \) represents a time-varying, deterministic applied current. In this case the propensity function will explicitly be a function of the voltage and we may replace \( \lambda_k(s, X(s)) \) in (4) with \( \lambda_k(V(s), X(s)) \). If multiple ion channel types are included in the model then, provided there are a finite number of types each with a finite number of individual channels, the vector \( X \in \mathbb{Z}^d \) represents the aggregated channel state. For specific examples of handling a single or multiple ion channel types, see Sections 3 and 4, respectively.

2.1.1 Simulation of the representation (4)-(5)

The construction of the representation (4) provided above implies a simulation strategy in which each point of the Poisson processes \( Y_k \), denoted \( \tau_n \) above, is generated sequentially and as needed. The time until the next reaction that occurs past time \( T \) is simply

\[
\Delta = \min_k \left\{ \Delta_k : \int_0^{T+\Delta_k} \lambda_k(s, X(s)) ds = \tau_T^k \right\},
\]

where \( \tau_T^k \) is the first point associated with \( Y_k \) coming after \( \int_0^{T} \lambda_k(s, X(s)) ds \). The reaction that took place is indicated by the index at which the minimum is achieved. See [1] for more discussion on this topic, including the algorithm provided below in which \( T_k \) will denote the value of the integrated intensity function \( \int_0^{T} \lambda_k(s, X(s)) ds \) and \( \tau_k \) will denote the first point associated with \( Y_k \) located after \( T_k \).

All random numbers generated in the algorithm below are assumed to be independent.

**Algorithm 1** (For the simulation of the representation (4)-(5)).

1. Initialize: set the initial number of molecules of each species, \( X \). Set the initial voltage value \( V \). Set \( t = 0 \). For each \( k \), set \( \tau_k = 0 \) and \( T_k = 0 \).
2. Generate \( M \) independent, uniform(0,1) random numbers \( \{r_k\}_{k=1}^{M} \). For each \( k \in \{1, \ldots, M\} \) set \( \tau_k = \ln(1/r_k) \).
3. Numerically integrate \[5\] forward in time until one of the following equalities hold:
\[
\int_t^{t+\Delta_k} \lambda_k(V(s),X(s)) \, ds = \tau_k - T_k.
\]

4. Set \( \Delta = \min_k \{\Delta_k\} \) and let \( \mu \) be the index where the minimum was achieved.

5. For each \( k \), set
\[
T_k = T_k + \int_t^{t+\Delta} \lambda_k(V(s),X(s)) \, ds.
\]

6. Set \( t = t + \Delta \) and
\[ X \leftarrow X + \zeta_\mu. \]

7. Let \( r \) be uniform \((0,1)\) and set \( \tau_\mu = \tau_\mu + \ln(1/r) \).

8. Return to step 3 or quit.

Note that the above algorithm relies on us being able to calculate a hitting time for each of the \( T_k(t) = \int_0^t \lambda_k(s,X(s)) \, ds \) exactly. Of course, in general this is not possible. However, making use of any reliable integration software will almost always be sufficient.

2.2 Gillespie representation

There are multiple alternative representations for the general stochastic process constructed in the previous section, with a “Gillespie” representation probably being the most useful in the current context. Following [5], we let \( Y \) be a unit rate Poisson process and let \( \{\xi_i, i = 0, 1, 2 \ldots \} \) be independent, uniform \((0,1)\) random variables that are independent of \( Y \). Set
\[
\lambda_0(V(s),X(s)) \equiv \sum_{k=1}^M \lambda_k(V(s),X(s)),
\]
\[
q_0 = 0 \text{ and for } k \in \{1, \ldots, M\}
\]
\[
q_k(s) = \lambda_0(V(s),X(s))^{-1} \sum_{\ell=1}^k \lambda_\ell(V(s),X(s)),
\]
where \( X \) and \( V \) satisfy
\[
R_0(t) = Y \left( \int_0^t \lambda_0(V(s),X(s)) \, ds \right)
\]
\[
X(t) = X(0) + \sum_{k=1}^M \zeta_k \int_0^t 1 \{\xi_{R_0(s-)} \in [q_{k-1}(s-),q_k(s-)]\} \, dR_0(s)
\]
\[
C \frac{dV}{dt} = I_{\text{app}}(t) - I_V(V(t)) - \left( \sum_{i=1}^d g_i^o X_i(t) \right) (V(t) - V_X).
\]
Then the stochastic process \((X, V)\) defined via (6)-(8) is a Markov process that is equivalent to (4)-(5), see [5]. An intuitive way to understand the above is by noting that \(R_0(t)\) simply determines the holding time in each state, whereas (7) simulates the embedded discrete time Markov chain (sometimes referred to as the skeletal chain) in the usual manner. Thus, this is the representation for the “Gillespie algorithm” with time dependent propensity functions [1].

2.2.1 Simulation of the representation (6)-(8)

Simulation of (6)-(8) is the analog of using Gillespie’s algorithm in the time-homogeneous case. All random numbers generated in the algorithm below are assumed to be independent.

Algorithm 2 (For the simulation of the representation (6)-(8)).

1. Initialize: set the initial number of molecules of each species, \(X\). Set the initial voltage value \(V\). Set \(t = 0\).

2. Let \(r \) be uniform(0,1) and numerically integrate (8) forward in time until
\[
\int_t^{t+\Delta} \lambda_0(V(s),X(s))ds = \ln(1/r).
\]

3. Let \(\xi \) be uniform(0,1) and find \(k \in \{1,\ldots,M\} \) for which
\[
\xi \in [q_{k-1}((t + \Delta) - ), q_k((t + \Delta) - )].
\]

4. Set \(t = t + \Delta\) and
\[
X \leftarrow X + \zeta_k.
\]

5. Return to step 2 or quit.

3 Morris-Lecar

As a concrete illustration of the exact stochastic simulation algorithms, we will consider the well known Morris-Lecar system [37], developed as a model for oscillations observed in barnacle muscle fibers [32]. The deterministic equations, which correspond to an appropriate \(N \rightarrow \infty\) limit of the system, constitute a planar model for the evolution of the membrane potential \(v(t)\) and the fraction of potassium gates, \(n \in [0,1]\), that are in the open or conducting state. In addition to a hyperpolarizing current carried by the potassium gates, there is a depolarizing calcium current gated by a rapidly equilibrating variable \(m \in [0,1]\). While a fully stochastic treatment of the Morris-Lecar system would include fluctuations in this calcium conductance, for simplicity in this section we will treat \(m\) as a fast, deterministic variable in the same manner as in the standard fast/slow decomposition, which we will refer to here as the planar Morris-Lecar model [13]. See Section 4 for a treatment of the Morris-Lecar system with both the potassium and calcium gates represented as discrete stochastic processes.
The deterministic or mean field equations for the planar Morris-Lecar model are:

\[
\frac{dv}{dt} = f(v,n) = \frac{1}{C} (I_{\text{app}} - g_{Ca}m_\infty(v)(v - v_{Ca}) - g_L(v - v_L) - g_Kn(v - v_K)) \tag{9}
\]

\[
\frac{dn}{dt} = g(v,n) = \alpha(v)(1 - n) - \beta(v)n = (n_\infty(v) - n)/\tau(v) \tag{10}
\]

The kinetics of the potassium channel may be specified either by the instantaneous time constant and asymptotic target, \(\tau\) and \(n_\infty\), or equivalently by the \textit{per capita} transition rates \(\alpha\) and \(\beta\). The terms \(m_\infty, \alpha, \beta, n_\infty\) and \(\tau\) satisfy

\[
m_\infty = \frac{1}{2} \left( 1 + \tanh \left( \frac{v - v_a}{v_b} \right) \right) \tag{11}
\]

\[
\alpha(v) = \frac{\phi \cosh(\xi/2)}{1 + e^{2\xi}} \tag{12}
\]

\[
\beta(v) = \frac{\phi \cosh(\xi/2)}{1 + e^{-2\xi}} \tag{13}
\]

\[
n_\infty(v) = \alpha(v)/(\alpha(v) + \beta(v)) = (1 + \tanh \xi)/2 \tag{14}
\]

\[
\tau(v) = 1/(\alpha(v) + \beta(v)) = 1/ (\phi \cosh(\xi/2)) \tag{15}
\]

where for convenience we define \(\xi = (v - v_c)/v_d\). For definiteness, we adopt values of the parameters

\[
v_K = -84, v_L = -60, v_{Ca} = 120 \tag{16}
\]

\[
I_{\text{app}} = 100, g_K = 8, g_L = 2, C = 20 \tag{17}
\]

\[
v_a = -1.2, v_b = 18 \tag{18}
\]

\[
v_c = 2, v_d = 30, \phi = 0.04, g_{Ca} = 4.4 \tag{19}
\]

for which the deterministic system has a stable limit cycle. For smaller values of the applied current (e.g. \(I_{\text{app}} = 75\)) the system has a stable fixed point, that loses stability through a subcritical Hopf bifurcation as \(I_{\text{app}}\) increases \([13], \S3\).

In order to incorporate the effects of random ion channel gating, we will introduce a finite number of potassium channels, \(N_{\text{tot}}\), and treat the number of channels in the open state as a discrete random process, \(0 \leq N(t) \leq N_{\text{tot}}\). In this simple model, each potassium channel switches between two states – closed or open – independently of the others, with voltage-dependent \textit{per capita} transition rates \(\alpha\) and \(\beta\), respectively. The entire population conductance ranges from 0 to \(g_K^2 N_{\text{tot}}\), where \(g_K^2 = g_K/N_{\text{tot}}\) is the single channel conductance, and \(g_K\) is the maximal whole cell conductance. For purposes of illustration and simulation we will typically use \(N_{\text{tot}} = 40\) individual channels\(^2\).

Our random variables will therefore be the voltage, \(V \in (-\infty, +\infty)\), and the number of open potassium channels, \(N \in \{0, 1, 2, \ldots, N_{\text{tot}}\}\). The number \(N_{\text{tot}} \in \mathbb{N}\) is taken to be a fixed parameter. We follow the usual capital/lowercase convention in the

\(^2\)Morris and Lecar used a value of \(g_K = 8\) mmhos/cm\(^2\) for the specific potassium conductance, corresponding to 80 picoSiemens (pS) per square micron \([33]\). The single channel conductance is determined by the structure of the potassium channel, and varies somewhat from species to species. However, conductances around 20 pS are typical \([42]\), which would give a density estimate of roughly 40 channels for a 10 square micron patch of cell membrane.
stochastic processes literature: \( N(t) \) and \( V(t) \) are the random processes and \( n \) and \( v \) are values they might take. In the random time change representation of Section 2.1, the opening and closing of the potassium channels are driven by two independent, unit rate Poisson processes, \( Y_{\text{open}}(t) \) and \( Y_{\text{close}}(t) \).

The evolution of \( V \) and \( N \) is closely linked. Conditioned on \( N \) having a specific value, say \( N = n \), the evolution of \( V \) obeys a deterministic differential equation,

\[
\frac{dV}{dt} \bigg|_{N=n} = f(V, n).
\]

Although its conditional evolution is deterministic, \( V \) is nevertheless a random variable. Meanwhile, \( N \) evolves as a jump process, i.e. \( N(t) \) is piecewise-constant, with transitions \( N \rightarrow N \pm 1 \) occurring with intensities that depend on the voltage \( V \). Conditioned on \( V = v \), the transition rates for \( N \) are

\[
N \rightarrow N + 1 \text{ with per capita rate } \alpha(v) \tag{21}
\]

\( (i.e. \text{ with net rate } \alpha(v) \cdot (N_{\text{tot}} - N)) \),

\[
N \rightarrow N - 1 \text{ with per capita rate } \beta(v) \tag{22}
\]

\( (i.e. \text{ with net rate } \beta(v) \cdot N) \).

Graphically, we may visualize the state space for \( N \) in the following manner:

\[
\begin{align*}
0 \xrightarrow{\alpha \cdot N_{\text{tot}}} 1 \xrightarrow{\beta} 2 \cdots (k - 1) \xrightarrow{k \beta} (k + 1) \cdots (N_{\text{tot}} - 1) \xrightarrow{\alpha \cdot (N_{\text{tot}} - k)} N_{\text{tot}},
\end{align*}
\]

with the nodes of the above graph being the possible states for the process \( N \), and the transition intensities located above and below the transition arrows.

Adopting the random time change representation of Section 2.1, we write our stochastic Morris-Lecar system as follows (cf. equations (9-10)):

\[
\frac{dV}{dt} = f(V(t), N(t)) \tag{23}
\]

\[
= \frac{1}{C} (I_{\text{app}} - g_{Ca}m_\infty(V(t))(V(t) - V_{Ca}) - g_L(V - V_L) - g_K N(t)(V(t) - V_K))
\]

\[
N(t) = N(0) - Y_{\text{close}} \left( \int_0^t \beta(V(s))N(s)ds \right) + Y_{\text{open}} \left( \int_0^t \alpha(V(s))(N_{\text{tot}} - N(s))ds \right). \tag{24}
\]

Appendices [A.1.1 and A.1.2] provide sample implementations of Algorithm 1 for the planar Morris-Lecar equations in XPP and Matlab, respectively. Figure [I] illustrates the results of the Matlab implementation.

### 4 Models with more than one channel type

The algorithms presented above readily generalize to systems with more than one type of stochastic channel, such as the Hodgkin-Huxley system \([23, 43]\). As an illustration, we include code in Appendices [A.2.1 and A.2.2] for XPP and Matlab, respectively, implementing a stochastic version of the original Morris-Lecar model, in a three dimensional phase space with both the calcium and potassium channels taken to be discrete, using the random time change algorithm ((4)-(5)).
Figure 1: Trajectory generated by Algorithm \[1\] (the random time change algorithm, \[(4)-(5)\]) for the planar Morris-Lecar model. We set $N_{\text{tot}} = 40$ potassium channels and used a driving current $I_{\text{app}} = 100$, which is above the Hopf bifurcation threshold for the parameters given. **Top Panel:** Number of open potassium channels ($N$), as a function of time. **Second Panel:** Voltage ($V$), as a function of time. **Bottom Panel:** Trajectory plotted in the ($V, N$) plane. Voltage varies along a continuum while open channel number remains discrete. Red curve: $v$-nullclcline of the underlying deterministic system, obtained by setting the RHS of equation \[(9)\] equal to zero. Green curve: $n$-nullcline, obtained by setting the RHS of equation \[(10)\] equal to zero.
4.1 Random Time Change Representation for the Morris-Lecar Model with Two Stochastic Channel Types

In Morris and Lecar’s original treatment of voltage oscillations in barnacle muscle fiber [33] the calcium gating variable \( m \) is included as a dynamical variable. The full (deterministic) equations have the form:

\[
\frac{dv}{dt} = F(v, n, m) = \frac{1}{C} \left( I_{\text{app}} - g_L(v - v_L) - g_{Ca}m(v - v_{Ca}) - g_Kn(v - v_K) \right) \quad (25)
\]

\[
\frac{dn}{dt} = G(v, n, m) = \alpha_n(v)(1 - n) - \beta_n(v)n = (n_{\infty}(v) - n)/\tau_n(v) \quad (26)
\]

\[
\frac{dm}{dt} = H(v, n, m) = \alpha_m(v)(1 - m) - \beta_m(v)m = (m_{\infty}(v) - m)/\tau_m(v) \quad (27)
\]

Here, rather than setting \( m \) to its asymptotic value \( m_{\infty} = \alpha_m/(\alpha_m + \beta_m) \), we allow the number of calcium gates to evolve according to (27). The planar form (Equations 9–10) is obtained by observing that \( m \) approaches equilibrium significantly more quickly than \( n \) and \( v \). Using standard arguments from singular perturbation theory [37, 38], one may approximate certain aspects of the full system (25–27) by setting \( m \) to \( m_{\infty}(v) \), and replacing \( F(v, n, m) \) and \( G(v, n, m) \) with \( f(v, n) = F(v, n, m_{\infty}(v)) \) and \( g(v, n) = G(v, n, m_{\infty}(v)) \), respectively. This reduction to the slow dynamics leads to the planar model (9–10).

To specify the full 3D equations, we introduce \( \xi_m = (v - v_a)/v_b \) in addition to \( \xi_n = (v - v_c)/v_d \) already introduced for the 2D model. The variable \( \xi_x \) represents where the voltage falls along the activation curve for channel type \( x \), relative to its half-activation point \( v_a \) for calcium and \( v_c \) for potassium and its slope (reciprocals of \( v_b \) for calcium and \( v_d \) for potassium). The per capita opening rates \( \alpha_m, \alpha_n \) and closing rates \( \beta_m, \beta_n \) for each channel type are given by

\[
\alpha_m(v) = \frac{\phi_m \cosh(\xi_m/2)}{1 + e^{2\xi_m}}, \quad \beta_m(v) = \frac{\phi_m \cosh(\xi_m/2)}{1 + e^{-2\xi_m}} \quad (28)
\]

\[
\alpha_n(v) = \frac{\phi_n \cosh(\xi_n/2)}{1 + e^{2\xi_n}}, \quad \beta_n(v) = \frac{\phi_n \cosh(\xi_n/2)}{1 + e^{-2\xi_n}} \quad (29)
\]

with parameters \( v_a = -1.2, v_b = 18, v_c = 2, v_d = 30, \phi_m = 0.4, \phi_n = 0.04 \). The asymptotic open probabilities for calcium and potassium are given, respectively, by the terms \( m_{\infty}, n_{\infty} \), and the time constants by \( \tau_m \) and \( \tau_n \). These terms satisfy the relations

\[
m_{\infty}(v) = \alpha_m(v)/(\alpha_m(v) + \beta_m(v)) = (1 + \tanh \xi_m)/2 \quad (30)
\]

\[
n_{\infty}(v) = \alpha_n(v)/(\alpha_n(v) + \beta_n(v)) = (1 + \tanh \xi_n)/2 \quad (31)
\]

\[
\tau_m(v) = 1/\left( \phi \cosh(\xi_m/2) \right) \quad (32)
\]

\[
\tau_n(v) = 1/\left( \phi \cosh(\xi_n/2) \right). \quad (33)
\]

Assuming a population of \( M_{\text{tot}} \) calcium gates and \( N_{\text{tot}} \) potassium gates, the voltage evolves as according to the sum of the applied, leak, calcium, and potassium currents:

\[
\frac{dV}{dt} = F(V(t), N(t), M(t)) = \frac{1}{C} \left( I_{\text{app}} - g_L(V - v_L) - g_{Ca} M_{\text{tot}}(v - v_{Ca}) - g_K N_{\text{tot}}(V - v_K) \right), \quad (34)
\]
while the number of open $M$ and $N$ remain constant except for unit increments and
decrements.

Figure 2 shows the results of the Matlab implementation for the 3D Morris-Lecar
system, with both the potassium and calcium channel treated discretely, using Algo-



\[ M_{\text{tot}} = N_{\text{tot}} = 40 \]

and the applied current $I_{\text{app}} = 100$ puts the deterministic system at a stable limit cycle
close to a Hopf bifurcation.

Figure 2: Trajectory generated by Algorithm 1 (the random time change algorithm, (4)-(5))
for the full three-dimensional Morris-Lecar model (equations (25)-(27)). We set $N_{\text{tot}} = 40$
potassium channels and $M_{\text{tot}} = 40$ calcium channels, and used a driving current $I_{\text{app}} = 100$,
a value above the Hopf bifurcation threshold of the mean field equations for the parameters
given. **Top Left Panel:** Number of open calcium channels ($M$), as a function of time. **Second Left Panel:** Number of open potassium channels ($N$), as a function of time. **Third Left Panel:** Voltage ($V$), as a function of time. **Right Panel:** Trajectory plotted in the ($V, M, N$) phase space. Voltage varies along a continuum while the joint channel state
remains discrete. Note that the number of open calcium channels makes frequent excursions
between $M = 0$ and $M = 40$, which demonstrates that neither a Langevin approximation
nor an approximate algorithm such as $\tau$-leaping (Euler’s method) would provide a good
approximation to the dynamics of the system.

5 Comparison of the Exact Algorithm with a Piecewise Constant Propensity Approximation.

Exact versions of the stochastic simulation algorithm for hybrid ion channel models
have been known since at least the 1980s [9]. Nevertheless, the implementation one
finds most often used in the literature is an approximate method in which the per
capita reaction propensities are held fixed between channel events. That is, in step 3
of Algorithm 1 the integral
\[ \int_{t}^{t + \Delta_k} \lambda_k(V(s), X(s)) \, ds \]
is replaced with
\[ \Delta_k \lambda_k(V(t), X(t)) \]
leaving the remainder of the algorithm unchanged. Put another way, one generates
the sequence of channel state jumps using the propensity immediately following the
most recent jump, rather than taking into account the time dependence of the reaction
propensities due to the continuously changing voltage. This piecewise constant
propensity approximation is analogous, in a sense, to the forward Euler method for the
numerical solution of ordinary differential equations.

Figure 3 shows a direct comparison of pathwise numerical solutions obtained by
the exact method provided in Algorithm 1 and the approximate forward method
detailed above. In general, the solution of a stochastic differential equation with a given
initial condition is a map from the sample space \( \Omega \) to a space of trajectories. In
the present context, the underlying sample space consists of one independent unit rate
Poisson process per reaction. For the planar Morris-Lecar model a point in \( \Omega \) amounts to fixing two Poisson processes, \( Y_{\text{open}} \) and \( Y_{\text{closed}} \), to drive the transitions of
the potassium channel. For the full 3D Morris-Lecar model we have four processes,
\( Y_1 \equiv Y_{\text{Ca,open}}, Y_2 \equiv Y_{\text{Ca,closed}}, Y_3 \equiv Y_{\text{K,open}} \) and \( Y_4 \equiv Y_{\text{K,closed}} \). The “exact” algorithm
provides a numerical solution of the map from \( \{Y_k\}_{k=1}^4 \in \Omega \) and initial conditions
\( (M_0, N_0, V_0) \) to the trajectory \( (M(t), N(t), V(t)) \). The approximate forward algorithm
gives a map from the same domain to a different trajectory, \( (\tilde{M}(t), \tilde{N}(t), \tilde{V}(t)) \). To
make a pathwise comparison for the full Morris-Lecar model, therefore, we fix both
the initial conditions and the four sample processes, and compare the resulting trajectories. Several features are evident in Figure 3. Both algorithms produce a sequence
of noise-dependent voltage spikes, with similar firing rates. The trajectories \( (M, N, V) \)
and \( (\tilde{M}, \tilde{N}, \tilde{V}) \) initially remain close together, and the timing of the first spike (taken,
e.g., as an upcrossing of \( V \) from negative to positive voltage) is similar for both algorithms.
Over time, however, discrepancies between the trajectories accumulate. The timing of the second and third spikes is noticeably different, and before ten spikes have accumulated the spike trains have become effectively uncorrelated.

Even though trajectories generated by the exact and approximate algorithms diverge when driven by identical Poisson processes, the two processes could still generate
sample paths with similar time-dependent or stationary distributions. That is, even though the two algorithms show strong divergence, they could still be close in a weak sense.

Given \( M_{\text{tot}} \) calcium and \( N_{\text{tot}} \) potassium channels, the density for the hybrid Markov
process may be written
\[ \rho_{m,n}(v, t) = \frac{1}{dV} \Pr \{ M(t) = m, N(t) = n, V \in [v, v + dv) \}, \]
(35)
Figure 3: Comparison of the exact algorithm with the “forward” piecewise constant propensity approximation. Blue solid lines denote the solution \((M(t), N(t), V(t))\) obtained using the Algorithm 1. Red dashed lines denote the solution \((\tilde{M}(t), \tilde{N}(t), \tilde{V}(t))\) obtained using the forward approximation. Both algorithms were begun with identical initial conditions \((M_0, N_0, V_0)\), and driven by the same four Poisson process streams \(Y_1, \cdots, Y_4\). Note the gradual divergence of the trajectories as differences between the exact and the forward approximate algorithms accumulate, demonstrating “strong” (or pathwise) divergence of the two methods. The exact and approximate trajectories diverge as time increases, even though they are driven by identical noise sources.
and obeys a master equation

\[
\frac{\partial \rho_{m,n}(v,t)}{\partial t} = - \frac{\partial}{\partial v} \left( F(v,n,m) \rho_{m,n}(v,t) \right) - (\alpha_m(v) (M_{tot} - m) + \beta_m(v) m + \alpha_n(v) (N_{tot} - n) + \beta_n(v) n) \rho_{m,n}(v,t) + (M_{tot} - m + 1) \alpha_m(v) \rho_{m-1,n}(v,t) + (m + 1) \beta_m(v) \rho_{m+1,n}(v,t) + (N_{tot} - n + 1) \alpha_n(v) \rho_{m,n-1}(v,t) + (n + 1) \beta_n(v) \rho_{m,n+1}(v,t),
\]

with initial condition \( \rho_{m,n}(0,0) \geq 0 \) given by any (integrable) density such that \( \int_{v \in \mathbb{R}} \sum_{m,n} \rho_{m,n}(v,0) \, dv = 1 \), and boundary conditions \( \rho \to 0 \) as \( |v| \to \infty \) and \( \rho \equiv 0 \) for either \( m, n < 0 \) or \( m > M_{tot} \) or \( n > N_{tot} \).

In contrast, the approximate algorithm with piecewise constant propensities does not generate a Markov process, since the transition probabilities depend on the past rather than the present values of the voltage component. Consequently they do not satisfy a master equation. Nevertheless it is plausible that they may have a unique stationary distribution.

Figure 3 shows pseudocolor plots of the histograms viewed in the \((v, n)\) plane, i.e. with entries summed over \( m \), for Algorithm 1 (“Exact”) and the approximate forward (“Euler”) algorithm, with \( M_{tot} = N_{tot} = k \) channels for \( k = 1, 2, 5, 10, 20 \) and 40. The two algorithms were run with independent random number streams in the limit cycle regime (with \( I_{app} = 100 \)) for \( t_{max} \approx 200,000 \) time units, sampled every 10 time units, which generated \( \geq 17,000 \) data points per histogram. For \( k < 5 \) the difference in the histograms is obvious at a glance. For \( k \geq 10 \) the histograms appear increasingly similar.

Figure 5 shows bar plots of the histograms projected on the voltage axis, i.e. with entries summed over \( m \) and \( n \), for the same data as in Figure 4 with \( M_{tot} = N_{tot} = k \) channels ranging from \( k = 1 \) to \( k = 40 \). Again, for \( k \leq 5 \), the two algorithms generate histograms that are clearly distinct. For \( k \geq 20 \) they appear similar, while \( k = 10 \) appears to be a borderline case.

To quantify the similarity of the histograms we calculated the empirical \( L_1 \) difference between the histograms in two ways: first for the full \((v, n, m)\) histograms, and then for the histograms collapsed to the voltage axis. Let \( \rho_{m,n}(v) \) and \( \tilde{\rho}_{m,n}(v) \) denote the stationary distributions for the exact and the approximate algorithms, respectively (assuming these exist). To compare the two distributions we approximate the \( L_1 \) distance between them, i.e.

\[
d(\rho, \tilde{\rho}) = \int_{v_{\min}}^{v_{\max}} \left( \sum_{m=0}^{M_{tot}} \sum_{n=0}^{N_{tot}} |\rho_{m,n}(v) - \tilde{\rho}_{m,n}(v)| \right) \, dv
\]

where \( v_{\min} \) and \( v_{\max} \) are chosen so that for all \( m, n \) we have \( F(v_{\min}, n, m) > 0 \) and \( F(v_{\max}, n, m) < 0 \). It is easy to see that such values of \( v_{\min} \) and \( v_{\max} \) must exist, since \( F(v, n, m) \) is a linear and monotonically decreasing function of \( v \) for any fixed pair \((n, m)\), cf. equation (25). Therefore, for any exact simulation algorithm, once the voltage component of a trajectory falls in the interval \( v_{\min} \leq v \leq v_{\max} \), it remains in this interval for all time. We approximate the voltage integral by summing over 100 evenly spaced bins along the voltage axis. Figure 6, top panel, shows empirical estimates of \( d(\rho, \tilde{\rho}) \) for values of \( M_{tot} = N_{tot} \) ranging from 1 to 40. The bottom panel shows the histogram for voltage considered alone.

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Figure 4: Normalized histograms of voltage and $N$ for exact and approximate (“Euler”) algorithms. The $V$-axis was partitioned into 100 equal width bins for each pair of histograms. The $N$-axis takes on $N_{tot} + 1$ discrete values. Color scale indicates relative frequency with which a bin was occupied (dark = infrequent, lighter = more frequent).
Figure 5: Histograms of voltage for exact and approximate (“Euler”) algorithms. The $V$-axis was partitioned into 100 equal width bins for each pair of histograms.
Figure 6: $L_1$ differences between empirical histograms generated by exact and approximate algorithms. $H_j(v, n, m)$ is the number of samples for which $V \in [v, v + \Delta v), N = n$ and $M = m$, where $\Delta v$ is a discretization of the voltage axis into 100 equal width bins; $j = 1$ represents an empirical histogram obtained by an exact algorithm (Algorithm 1), and $j = 2$ represents an empirical histogram obtained by a piecewise-constant propensity approximation or “Euler” algorithm. Note the normalized $L_1$ difference of two histograms can range from 0 to 2. **Top:** Difference for the full histograms, $L_1(H_1 - H_2) \equiv \sum_{v,n,m} |H_1(v, n, m) - H_2(v, n, m)| / \# \text{samples}$. Here $v$ refers to binned voltage values. **Bottom:** Difference for the corresponding voltage histograms, $L_1(VH_1 - VH_2) \equiv \sum_v |VH_1(v) - VH_2(v)| / \# \text{samples}$, where $VH_j(v) = \sum_{n,m} H_j(v, n, m)$. 


6 Coupling, variance reduction, and parametric sensitivities

It is relatively straightforward to derive the exact simulation strategies from the different representations provided here. Less obvious, and perhaps even more useful, is the fact that the random time change formalism also lends itself to the development of new computational methods that potentially allow for significantly greater computational speed, with no loss in accuracy. They do so by providing ways to couple two processes in order to reduce the variance, and thereby increase the speed, of different natural estimators.

For an example of a common computational problem where such a benefit can be had, consider the computation of parametric sensitivities, which is a powerful tool in determining parameters to which a system output is most responsive. Specifically, suppose that the intensity/propensity functions are dependent on some vector of parameters $\theta$. For instance, $\theta$ may represent a subset of the system’s mass action kinetics constants, the cell capacitance, or the underlying number of channels of each type. We may wish to know how sensitively a quantity such as the average firing rate or interspike interval variance depends on the parameters $\theta$. We then consider a family of models $(V^\theta, X^\theta)$, parameterized by $\theta$, with stochastic equations

\[
\frac{d}{dt} V^\theta(t) = f \left( \theta, V^\theta(t), X^\theta(t) \right) \tag{37}
\]

\[
X^\theta(t) = X^\theta_0 + \sum_k Y_k \left( \int_0^t \lambda^\theta_k \left( V^\theta(s), X^\theta(s) \right) ds \right) \zeta_k,
\]

where $f$ is some function, and all other notation is as before. Some quantities arising in neural models depend on the entire sample path, such as the mean firing rate and the interspike interval variance. Let $g(\theta, V^\theta, X^\theta)$ be a path functional capturing the quantity of interest. In order to efficiently and accurately evaluate the relative shift in the expectation of $g$ due to a perturbation of the parameter vector, we would estimate

\[
\tilde{s} = \epsilon^{-1} \mathbb{E} \left[ g \left( \theta', V^\theta', X^\theta' \right) - g \left( \theta, V^\theta, X^\theta \right) \right]
\]

\[
\approx \frac{1}{\epsilon N} \sum_{i=1}^N \left[ g \left( \theta', V^\theta_{[i]}, X^\theta_{[i]} \right) - g \left( \theta, V^\theta_{[i]}, X^\theta_{[i]} \right) \right] \tag{38}
\]

where $\epsilon = ||\theta - \theta'||$ and where $(V^\theta_{[i]}, X^\theta_{[i]})$ represents the $i$th path generated with a parameter choice of $\theta$, and $N$ is the number of sample paths computed for the estimation. That is, we would use a finite difference and Monte Carlo sampling to approximate the change in the expectation.

If the paths $(V^\theta_{[i]}, X^\theta_{[i]})$ and $(V^{\theta'}_{[i]}, X^{\theta'}_{[i]})$ are generated independently, then the variance of the estimator (38) for $\tilde{s}$ is $O(N^{-1} \epsilon^{-2})$, and the standard deviation of the estimator is $O(N^{-1/2} \epsilon^{-1})$. This implies that in order to reduce the confidence interval of the estimator to a target level of $\rho > 0$, we require

\[
N^{-1/2} \epsilon^{-1} \lesssim \rho \implies N \gtrsim \epsilon^{-2} \rho^{-2},
\]

which can be prohibitive. Reducing the variance of the estimator can be achieved by coupling the processes $(V^\theta_{[i]}, X^\theta_{[i]})$ and $(V^{\theta'}_{[i]}, X^{\theta'}_{[i]})$, so that they are correlated, by
constructing them on the same probability space. This process is analogous to the
construction of simulation paths according to the random time change algorithm and a
piecewise constant propensity approximation algorithm used to illustrate strong diver-
gence (Figure 3). The representations we present here lead rather directly to schemes
for reducing the variance of the estimator. We discuss different possibilities.

1. **The common random number (CRN) method.** The CRN method simulates
both processes according to the Gillespie representation (6)-(8) with the same
Poisson process \(Y\) and the same stream of uniform random variables \(\{\xi_i\}\). In
terms of implementation, one simply reuses the same two streams of uniform
random variables in Algorithm 2 for both processes.

2. **The common reaction path method (CRP).** The CRP method simulates
both processes according to the random time change representation with the same
Poisson processes \(Y_k\). In terms of implementation, one simply makes one stream of
uniform random variables for each reaction channel, and then uses these streams
for the simulation of both processes. See [35], where this method was introduced
in the context of biochemical models.

3. **The coupled finite difference method (CFD).** The CFD method utilizes a
“split coupling” introduced in [2]. Specifically, it splits the counting process for
each of the reaction channels into three pieces: one counting process that is shared
by \(X^\theta\) and \(X^\theta'\) (and has propensity equal to the minimum of their respective
intensities for that reaction channel), one that only accounts for the jumps of
\(X^\theta\), and one that only accounts for the jumps of \(X^\theta'\). Letting \(a \wedge b = \min\{a, b\}\), the
precise coupling is

\[
X^\theta(t) = X^\theta_0 + \sum_k Y_{k,1} \left( \int_0^t m_k(\theta, \theta', s) ds \right) \zeta_k
+ \sum_k Y_{k,2} \left( \int_0^t \lambda_k^\theta(V^\theta(s), X^\theta(s)) - m_k(\theta, \theta', s) ds \right) \zeta_k
\]

\[
X^\theta'(t) = X^\theta(t) + \sum_k Y_{k,1} \left( \int_0^t m_k(\theta, \theta', s) ds \right) \zeta_k
+ \sum_k Y_{k,3} \left( \int_0^t \lambda_k^\theta(V^\theta(s), X^\theta(s)) - m_k(\theta, \theta', s) ds \right) \zeta_k,
\]

where

\[
m_k(\theta, \theta', s) = \lambda_k^\theta(V^\theta(s), X^\theta(s)) \wedge \lambda_k^\theta(V^\theta'(s), X^\theta'(s)),
\]

and where \(\{Y_{k,1}, Y_{k,2}, Y_{k,3}\}\) are independent unit-rate Poisson processes. Implementa-
tion is then carried out by Algorithm 1 in the obvious manner.

While the different representations provided in this paper imply different exact
simulation strategies (i.e. Algorithms 1 and 2), those strategies still produce statistically
equivalent paths. This is not the case for the methods for parametric sensi-
tivities provided in this section. To be precise, each method constructs a coupled pair of
processes \((V^\theta, X^\theta), (V^\theta', X^\theta')\), and the marginal processes \((V^\theta, X^\theta)\) and \((V^\theta', X^\theta')\)
are all statistically equivalent no matter the method used. However, the covariance
\( \text{Cov} \left( (V^\theta(t), X^\theta(t)), (V^{\theta'}(t), X^{\theta'}(t)) \right) \) can be drastically different. This is important, since it is variance reduction we are after, and for any component \( X_j \),

\[
\text{Var}(X_j^\theta(t) - X_j^{\theta'}(t)) = \text{Var}(X_j^\theta(t)) + \text{Var}(X_j^{\theta'}(t)) - 2 \text{Cov}(X_j^\theta(t), X_j^{\theta'}).
\]

Thus, minimizing variance is equivalent to maximizing covariance. Typically, the CRN method does the worst job of maximizing the covariance, even though it is the most widely used method \[44\]. The CFD method typically does the best job of maximizing the covariance, though examples exist in which the CRP method is the most efficient \[2, 44\].

7 Discussion

We have provided two general representations for stochastic ion channel kinetics, one based on the random time change formalism, and one extending Gillespie’s algorithm to the case of ion channels driven by time-varying membrane potential. We believe that the random time change representation (Algorithm \[1\]-(\[5\])) will be particularly useful to the computational neuroscience community as it allows for generalizations of computational methods developed in the context of biochemistry, in which the propensities depend upon the state of the jump process only. For example, variance reduction strategies for the efficient computation of first and second order sensitivities \[2, 6, 35\], as discussed in Section \[3\], and for the efficient computation of expectations using multilevel Monte Carlo \[4, 16\] now become feasible.

The random time change approach avoids several approximations that are commonly found in the literature. In simulation algorithms based on a fixed time step chemical Langevin approach, it is necessary to assume that the increments in channel state are approximately Gaussian distributed over an appropriate time interval \[15\] \[20\] \[21\] \[31\] \[46\]. However, in exact simulations with small membrane patches corresponding to \( M_{\text{tot}} = 40 \) calcium channels, the exact algorithm routinely visits the states \( M(t) = 0 \) and \( M(t) = 40 \), for which the Gaussian increment approximation is invalid regardless of time step size. The main alternative algorithm typically found in the literature is the piecewise constant propensity or approximate forward algorithm \[14\] \[25\] \[41\]. However, this algorithm ignores changes to membrane potential during the intervals between channel state changes. As the sensitivity of ion channel opening and closing to voltage is fundamental to the neurophysiology of cellular excitability, these algorithms are not appropriate unless the time between openings and closing is especially small. The exact algorithm \[9\] \[24\] \[34\] is straightforward to implement and avoids these approximations and pitfalls.

For any simulation algorithm, it is reasonable to ask about the growth of complexity of the algorithm as the underlying stochastic model is enriched. For example, the natural jump Markov interpretation of Hodgkin and Huxley’s model for the sodium channel comprises eight distinct states with twenty different state-to-state transitions, each driven by an independent Poisson process in the random time change representation. Recent investigations of sodium channel kinetics have led neurophysiologists to formulate models with as many as 26 distinct states connected by 72 transitions \[30\]. While the random time change representation extends naturally to such scenarios, it may also be fruitful to combine it with complexity reduction methods such as the
stochastic shielding algorithm introduced by Schmandt and Galán [39], and analyzed by Schmidt and Thomas [40]. For example, of the twenty independent Poisson processes driving a discrete Markov model of the Hodgkin-Huxley sodium channel, only four of the processes directly affect the conductance of the channel; fluctuations associated with the remaining sixteen Poisson processes may be ignored with negligible loss of accuracy in the first and second moments of the channel occupancy. Similarly, for the 26 state sodium channel model proposed in [30], all but 12 of the 72 Poisson processes representing state-to-state transitions can be replaced by their expected values. Analysis of algorithms combining stochastic shielding and the random time change framework are a promising direction for future research.

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A Sample Implementations of Random Time Change Algorithm (Algorithm 1)

A.1 Morris Lecar with Stochastic Potassium Channel

A.1.1 XPP: ml-rtc-konly.ode

```plaintext
# ML with stochastic potassium channels
# uses the exact random time change algorithm

# membrane potential
v'=(I-gca*minf*(V-Vca)-gk*wtot*(V-VK)-gl*(V-Vl))/c
# unit exponentials
t[3..4]'=0
# number of potassium (w) /calcium (m) channels open
w'=0
# int_0^t beta(V(s)) ds
# for the 4 reactions
awp'=aw
bwp'=bw
# initialize unit exponentials
t[3..4](0)=-log(ran(1))
# look for crossings, reset integrals, increment channels, choose next time
global 1 awp-t3 {t3=-log(ran(1));awp=0;w=w+1}
global 1 bwp-t4 {t4=-log(ran(1));bwp=0;w=w-1}
# parameters
par Nw=100
init v=-50
# fraction of open channels
wtot=w/Nw
# ML channel kinetic definitions
minf=.5*(1+tanh((v-va)/vb))
winf=.5*(1+tanh((v-vc)/vd))
tauw=1/cosh((v-vc)/(2*vd))
alw=winf/tauw
blw=1/tauw-alw
# independent, so rates are just multiples of number in each state
aw=alw*(Nw-w)*phi
bw=blw*w*phi
param vk=-84, vl=-60, vca=120
param i=75, gk=8, gl=2, c=20, phim=.4
param va=-1.2, vb=18
param vc=2, vd=30, phi=.04, gca=4.4
# set up some numerics and plotting stuff
@ dt=.01, nout=100, total=4000, bound=1000
@ maxstor=5000, meth=euler
@ xhi=4000, ylo=-65, yhi=50
done
```

function \([V,N,t,Ntot]=ml\_rtc\_exact\_konly(tmax,Ntot)\)

%function \([V,N,t,Ntot]=ml\_rtc\_exact\_konly(tmax,Ntot);\)

% Exact solution of Morris Lecar with stochastic potassium channel.
% Using the random time change algorithm. We track two reactions:
% % Rxn 1: closed -> open (per capita rate alpha)
% Rxn 2: open -> closed (per capita rate beta)
% % Default Ntot=40, tmax=4000.
% % Author: PJT July 2013, Case Western Reserve University.

% Applied current "Iapp" set on line 47 below.

%%% Use global variables to represent the channel state
% and random trigger for Poisson process
global Npotassium\_tot % total number of potassium channels
global Npotassium % number of open potassium channels
global tau1 T1 % time to next opening event (internal to reaction 1)
global tau2 T2 % time to next closing event (internal to reaction 2)

%%% Set defaults for input arguments
if nargin < 2, Ntot=40; end
Npotassium\_tot=Ntot;
if nargin < 1, tmax=4e3; end
%%% Parameters
% Standard Morris-Lecar parameters giving a globally attracting limit cycle
% (if applied current Iapp=100) or a stable fixed point (if Iapp=75).
va = -1.2;
vb=18;
vC = 2;
vD = 30;
phi = 0.04;
%%% Functions for Morris-Lecar
global Iapp; Iapp=@(t)100; % applied current.
global minf; minf=@(v)0.5*(1+tanh((v-vA)/vB)); % m-gate activation
global xi; xi=@(v)(v-vC)/vD; % scaled argument for n-gate input
global ninf; ninf=@(v)0.5*(1+tanh(xi(v))); % n-gate activation function
global tau\_n; tau\_n=@(v)1./(phi*cosh(xi(v)/2)); % n-gate activation t-const
% global alpha; alpha=@(v)(ninf(v)/tau\_n(v)); % per capita opening rate
% global beta; beta=@(v)((1-ninf(v))/tau\_n(v)); % per capita closing rate
%%% ODE options including reset
options=odeset('Events',@nextevent);
%%% Initialize
t0=0;
t=t0;  % global "external" time
tau1=-log(rand);  % time of next event on reaction stream 1 ("internal time")
tau2=-log(rand);  % time of next event on reaction stream 2 ("internal time")
T1=0;  % integrated intensity function for reaction 1
T2=0;  % integrated intensity function for reaction 2
V0=-50;  % start at an arbitrary middle voltage
V=V0;
N0=ceil(Ntot/2);  % start with half of channels open
Npotassium=N0;
N=N0;  % use N to record the time course

while t(end)<tmax

%% Loop over events
%% Identify which reaction occurred, adjust state, and
%% continue, and set trigger for next event.
mu=event_idx;  % next reaction index
if mu==1  % next reaction is a channel opening
    N0=N0+1;  % increment channel state
    tau1=tau1-log(rand);  % increment in tau1 is exp'l with mean 1
elseif mu==2  % next reaction is another channel closing
    N0=N0-1;  % decrement channel state
    tau2=tau2-log(rand);  % increment in tau2 likewise
end
Npotassium=N0;
if N0>Npotassium_tot, error('N>Ntot'), end
if N0<0, error('N<0'), end
T1=T1+Uout(end,2);
T2=T2+Uout(end,3);
V0=V(end);
end  % while t(end)<tmax

%% Plot output
figure
subplot(4,1,1),plot(t,N),xlabel('time'),ylabel('N')
subplot(4,1,2:3), plot(V,N,'.-'), xlabel('V'), ylabel('N')
subplot(4,1,4), plot(t,V), xlabel('time'), ylabel('V')
shg
end % End of function ml_rtc_exact_konly
%% Define the RHS for Morris-Lecar
% state vector for integration is as follows:
% u(1) = voltage
% u(2) = integral of activation hazard function
% u(3) = integral of inactivation hazard function
% u(4) = N (number of open potassium channels)
function dudt = dudtfunc(t,u)
global Iapp % Applied Current
global minf % asymptotic target for (deterministic) calcium channel
global Npotassium Npotassium_tot % num. open, total num. of channels
global alpha beta % per capita transition rates
%% Parameters
vK = -84; vL = -60; vCa = 120;
gK =8; gL =2; C=20; gCa = 4.4;
%% calculate the RHS;
v=u(1); % extract the voltage from the input vector
dudt=[(Iapp(t)-gCa*minf(v)*(v-vCa)-gL*(v-vL)-...
gK*(Npotassium/Npotassium_tot)*(v-vK))/C; % voltage
alpha(v)*(Npotassium_tot-Npotassium); % channel opening internal time
beta(v)*Npotassium;% channel closing internal time
0]; % N is constant between events
end
%% Define behavior at threshold crossing
function [value,isterminal,direction] = nextevent(~,u)
global tau1 T1 % timing trigger for reaction 1 (opening)
global tau2 T2 % timing trigger for reaction 2 (closing)
value=u(2)-(tau1-T1);u(3)-(tau2-T2)];
isterminal=[1;1]; % stop and restart integration at crossing
direction=[1;1]; % increasing value of the quantity at the trigger
end
A.2 Morris Lecar with Stochastic Potassium and Calcium Channels

A.2.1 XPP: ml-rtc-exact.ode

# ML with both potassium and calcium channels stochastic
# uses the exact random time change algorithm.

# membrane potential
v'=(I-gca*mtot*(V-Vca)-gk*wtot*(V-Vk)-gl*(V-Vl))/c

# unit exponentials
t[1..4]'=0

# number of potassium(w) /calcium (m) channels open
w'=0
m'=0

# int_0^t beta(V(s)) ds
# for the 4 reactions
amp'=am
bmp'=bm
awp'=aw
bwp'=bw

# initialize unit exponentials
t[1..4](0)=-log(ran(1))

# look for crossings, reset integrals, increment channels, choose next time
global 1 amp-t1 {t1=-log(ran(1));amp=0;m=m+1}
global 1 bmp-t2 {t2=-log(ran(1));bmp=0;m=m-1}
global 1 awp-t3 {t3=-log(ran(1));awp=0;w=w+1}
global 1 bwp-t4 {t4=-log(ran(1));bwp=0;w=w-1}

# parameters
par Nm=100,Nw=100
init v=-50
param vk=-84,vl=-60,vca=120
param i=75,gk=8,gl=2,c=20,phim=.4
param va=-1.2,vb=18
param vc=2,vd=30,phi=.04,gca=4.4

# fraction of open channels
wtot=w/Nw
mtot=m/Nm

# ML channel kinetic definitions
minf=.5*(1+tanh((v-va)/vb))
\[ \text{winf} = 0.5 \times (1 + \tanh((v - vc)/vd)) \]
\[ \text{tauw} = 1 / \cosh((v - vc)/(2*vd)) \]
\[ \text{taum} = 1 / \cosh((v - va)/(2*vb)) \]
\[ \text{alm} = \text{minf}/\text{taum} \]
\[ \text{blm} = 1 / \text{taum} - \text{alm} \]
\[ \text{alw} = \text{winf}/\text{tauw} \]
\[ \text{blw} = 1 / \text{tauw} - \text{alw} \]

# independent, so rates are just multiples of number in each state
\[ \text{am} = \text{alm} \times (N_m - m) \times \text{phim} \]
\[ \text{bm} = \text{blm} \times m \times \text{phim} \]
\[ \text{aw} = \text{alw} \times (N_w - w) \times \text{phi} \]
\[ \text{bw} = \text{blw} \times w \times \text{phi} \]

# set up some numerics and plotting stuff
\[ @ \text{dt} = 0.01, \text{nout} = 100, \text{total} = 4000, \text{bound} = 1000 \]
\[ @ \text{maxstor} = 5000, \text{meth} = \text{euler} \]
\[ @ \text{xhi} = 4000, \text{ylo} = -65, \text{yhi} = 50 \]
done
A.2.2 Matlab: mlexactboth.m

function [V,M,N,t,Mtot,Ntot]=mlexactboth(tmax,Mtot,Ntot)

%function [V,M,Mtot,N,Ntot]=mlexactboth(tmax,Mtot,Ntot);
%
% Exact solution of Morris Lecar with discrete stochastic potassium channel
% (0<=N<=Ntot) and discrete stochastic calcium channel (0<=M<=Mtot).
% Using the random time change representation, we track four reactions:
%
% Rxn 1: calcium (m-gate) closed -> open (per capita rate alpha_m)
% Rxn 2: calcium (m-gate) open -> closed (per capita rate beta_m)
% Rxn 3: potassium (n-gate) closed -> open (per capita rate alpha_n)
% Rxn 4: potassium (n-gate) open -> closed (per capita rate beta_n)
%
% Default Mtot=40, Ntot=40, tmax=4000.
%
% Applied current "Iapp" set internally.
%
% PJT June 2013, CWRU. Following Bard Ermentrout’s "ml-rtc-exact.ode".

%% Use global variables to represent the channel state
% and random trigger for Poisson process
%
% Calcium
global Mcalcium_tot % total number of calcium channels
global Mcalcium % number of calcium channels in conducting state
global tau1 T1 % dummy variables for time to next Ca-opening event
global tau2 T2 % dummy variables for time to next Ca-closing event
%
% Potassium
global Npotassium_tot % total number of potassium channels
global Npotassium % number of potassium channels in conducting state
global tau3 T3 % dummy variables for time to next K-opening event
global tau4 T4 % dummy variables for time to next K-closing event

%% Set defaults for input arguments
if nargin < 3, Ntot=40; end
Npotassium_tot=Ntot;
if nargin < 2, Mtot=40; end
Mcalcium_tot=Mtot;
if nargin < 1, tmax=4e3; end

%% Parameters
phi_m=0.4;
va = -1.2; vb=18;
vc = 2; vd = 30;
phi_n = 0.04;

%% Functions for Morris-Lecar
global Iapp; Iapp=@(t)100; % applied current

global xi_m; xi_m=@(v)(v-va)/vb; % scaled argument for m-gate input voltage
global minf; minf=@(v)0.5*(1+tanh(xi_m(v))); % m-gate activation function
global tau_m; tau_m=@(v)1./(phi_m*cosh(xi_m(v)/2)); % m-gate time constant
global alpha_m; alpha_m=@(v)(minf(v)/tau_m(v));
global beta_m; beta_m=@(v)((1-minf(v))/tau_m(v));

global xi_n; xi_n=@(v)(v-vc)/vd; % scaled argument for n-gate input voltage
global ninf; ninf=@(v)0.5*(1+tanh(xi_n(v))); % n-gate activation function
global tau_n; tau_n=@(v)1./(phi_n*cosh(xi_n(v)/2)); % n-gate time constant
global alpha_n; alpha_n=@(v)(ninf(v)/tau_n(v));
global beta_n; beta_n=@(v)((1-ninf(v))/tau_n(v));

%% ODE options including reset
options=odeset('Events',@nextevent);

%% initialize

t0=0;
t=t0; % global "external" time

tau1=-log(rand); % time of next event on reaction stream 1 ("internal time")
tau2=-log(rand); % time of next event on reaction stream 2 ("internal time")
tau3=-log(rand); % time of next event on reaction stream 3 ("internal time")
tau4=-log(rand); % time of next event on reaction stream 4 ("internal time")
T1=0; % integrated intensity function for reaction 1
T2=0; % integrated intensity function for reaction 2
T3=0; % integrated intensity function for reaction 3
T4=0; % integrated intensity function for reaction 4
V0=-50; % start at an arbitrary middle voltage
V=V0; % initial voltage
M0=0; % start with calcium channels all closed
Mcalcium=M0; % initial state of calcium channel
M=M0; % use M to record the time course

NO=ceil(Ntot/2); % start with half potassium channels open
Npotassium=NO; % initial state of potassium channel
N=N0; % use N to record the time course

%% Loop over events
while t(end)<tmax

%% integrate ODE for voltage, until the next event is triggered
% State vector U for integration contains the following components
% 1 [voltage;
integral of (# closed)*alpha_m(v(t));
integral of (# open)*beta_m(v(t));
integral of (# closed)*alpha_n(v(t));
integral of (# open)*beta_n(v(t));
# open calcium channels;
# open potassium channels].
U0=[V0;0;0;0;0;M0;N0];
tspan=[t(end),tmax];
[tout,Uout,~,~,event_idx]=ode23(@dudtfunc,tspan,U0,options);
Vout=Uout(:,1); % voltage at time of next event
Mout=Uout(:,6); % number of calcium channels open at end of next event
Nout=Uout(:,7); % number of potassium channels open at end of next event
t=[t,tout'];
V=[V,Vout'];
M=[M,Mout'];
N=[N,Nout'];

Identify which reaction occurred, adjust state, and continue;
% and set trigger for next event.
mu=event_idx; % next reaction index
if mu==1 % next reaction is a calcium channel opening
    M0=M0+1; % increment calcium channel state
    tau1=tau1-log(rand); % increment in tau1 is exponentially distributed with mean 1
elseif mu==2 % next reaction is a calcium channel closing
    M0=M0-1; % decrement calcium channel state
    tau2=tau2-log(rand); % increment in tau2 is exponentially distributed with mean 1
elseif mu==3 % next reaction is a potassium channel opening
    N0=N0+1; % increment potassium channel state
    tau3=tau3-log(rand); % increment in tau3 is exponentially distributed with mean 1
elseif mu==4 % next reaction is a potassium channel closing
    N0=N0-1; % decrement potassium channel state
    tau4=tau4-log(rand); % increment in tau4 likewise
end
Mcalcium=M0;
Npotassium=N0;
if M0>Mcalcium_tot, error('M>Mtot'), end
if M0<0, error('M<0'), end
if N0>Npotassium_tot, error('N>Ntot'), end
if N0<0, error('N<0'), end
T1=T1+Uout(end,2);
T2=T2+Uout(end,3);
T3=T3+Uout(end,4);
T4=T4+Uout(end,5);
V0=V(end);
end % while t(end)<tmax

%% Plot output

figure
subplot(6,1,1),plot(t,M),ylabel('M','FontSize',20),set(gca,'FontSize',20)
subplot(6,1,2),plot(t,N),ylabel('N','FontSize',20),set(gca,'FontSize',20)
subplot(6,1,6),plot(t,V),xlabel('Time','FontSize',20)
ylabel('V','FontSize',20),set(gca,'FontSize',20)
subplot(6,1,4:6),plot3(V,M,N,'.-'),xlabel('V','FontSize',20)
ylabel('M','FontSize',20),zlabel('N','FontSize',20),set(gca,'FontSize',20)
ggrid on, rotate3d, shg

end % End of function mlexactboth

%% Define the RHS for Morris-Lecar
% state vector for integration is as follows:
% u(1) = voltage
% u(2) = integral of Ca-opening hazard function (Rxn 1)
% u(3) = integral of Ca-closing hazard function (Rxn 2)
% u(4) = integral of K-opening hazard function (Rxn 3)
% u(5) = integral of K-closing hazard function (Rxn 4)
% u(6) = M (number of open calcium channels)
% u(7) = N (number of open potassium channels)

function dudt=dudtfunc(t,u)
global Iapp % applied current
global Mcalcium Mcalcium_tot
global Npotassium Npotassium_tot
global alpha_m beta_m

%% Parameters
vK = -84; vL = -60; vCa = 120;
gK = 8; gL = 2; C=20; gCa = 4.4;

%% calculate the RHS;
v=u(1); % extract the voltage from the input vector
dudt=[... % voltage
    (Iapp(t)-gCa*(Mcalcium/Mcalcium_tot)*(v-vCa)-gL*(v-vL)... 
    -gK*(Npotassium/Npotassium_tot)*(v-vK))/C;
% Calcium chan. opening, internal time elapsed
alpha_m(v)*(Mcalcium_tot-Mcalcium);
% Calcium chan. closing, internal time elapsed
beta_m(v)*Mcalcium;
% Potassium chan. opening, internal time elapsed
alpha_n(v)*(Npotassium_tot-Npotassium);

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% Potassium chan. closing, internal time elapsed
beta_n(v)*Npotassium;
0; % M is constant between events
0]; % N is constant between events
end

%% define behavior at threshold crossing

function [value,isterminal,direction] = nextevent(~,u)
    global tau1 T1 % timing trigger for reaction 1 (calcium opening)
global tau2 T2 % timing trigger for reaction 2 (calcium closing)
global tau3 T3 % timing trigger for reaction 3 (potassium opening)
global tau4 T4 % timing trigger for reaction 4 (potassium closing)
    value=[u(2)-(tau1-T1);u(3)-(tau2-T2);u(4)-(tau3-T3);u(5)-(tau4-T4)];
    isterminal=[1;1;1;1;1]; % stop and restart integration at crossing
direction=[1;1;1;1;1]; % increasing value of the quantity at the trigger
end
References

[1] David F Anderson. A modified next reaction method for simulating chemical systems with time dependent propensities and delays. J Chem Phys, 127(21):214107, Dec 2007.

[2] David F. Anderson. An efficient finite difference method for parameter sensitivities of continuous time Markov chains. SIAM Journal on Numerical Analysis, 50(5):2237 – 2258, 2012.

[3] David F. Anderson, Arnab Ganguly, and Thomas G. Kurtz. Error analysis of tau-leap simulation methods. Ann. Appl. Probab., 21(6):2226–2262, December 2011.

[4] David F. Anderson and Desmond J. Higham. Multi-level Monte Carlo for continuous time Markov chains, with applications in biochemical kinetics. SIAM: Multiscale Modeling and Simulation, 10(1):146 – 179, 2012.

[5] David F. Anderson and Thomas G. Kurtz. Design and Analysis of Biomolecular Circuits, chapter 1. Continuous Time Markov Chain Models for Chemical Reaction Networks. Springer, 2011.

[6] David F. Anderson and Elizabeth Skubak Wolf. A finite difference method for estimating second order parameter sensitivities of discrete stochastic chemical reaction networks. J. Chem. Phys., 137(22):224112, 2012.

[7] Karen Ball, Thomas G. Kurtz, Lea Popovic, and Greg Rempala. Asymptotic analysis of multiscale approximations to reaction networks. Ann. Appl. Prob., 16(4):1925–1961, 2006.

[8] Yang Cao, Daniel T Gillespie, and Linda R Petzold. Efficient step size selection for the tau-leaping simulation method. J Chem Phys, 124(4):044109, Jan 2006.

[9] J R Clay and L J DeFelice. Relationship between membrane excitability and single channel open-close kinetics. Biophys J, 42(2):151–7, May 1983.

[10] D. Colquhoun and A. G. Hawkes. Single-Channel Recording, chapter The Principles of the Stochastic Interpretation of Ion-Channel Mechanisms. Plenum Press, New York, 1983.

[11] Alan D. Dorval, Jr. and John A. White. Channel noise is essential for perithreshold oscillations in entorhinal stellate neurons. The Journal of Neuroscience, 25(43):10025–10028, Oct. 26 2005.

[12] Berton A. Earnshaw and James P. Keener. Invariant manifolds of binomial-like nonautonomous master equations. Siam J. Applied Dynamical Systems, 9(2):568–588, 3 June 2010.

[13] G. Bard Ermentrout and David H. Terman. Foundations Of Mathematical Neuroscience. Springer, 2010.

[14] Karin Fisch, Tilo Schwalger, Benjamin Lindner, Andreas V M Herz, and Jan Benda. Channel noise from both slow adaptation currents and fast currents is required to explain spike-response variability in a sensory neuron. J Neurosci, 32(48):17332–44, Nov 2012.
[15] Ronald F. Fox and Yan-nan Lu. Emergent collective behavior in large numbers of globally coupled independently stochastic ion channels. Phys Rev E Stat Phys Plasmas Fluids Relat Interdiscip Topics, 49(4):3421–3431, Apr 1994.

[16] Mike B. Giles. Multilevel Monte Carlo path simulation. Operations Research, 56:607–617, 2008.

[17] D. T. Gillespie. Exact stochastic simulation of coupled chemical reactions. J. Phys. Chem., 81:2340–2361, 1977.

[18] Daniel T Gillespie. Stochastic simulation of chemical kinetics. Annu. Rev. Phys. Chem., 58:35–55, 2007.

[19] Peter W. Glynn. A GSMP formalism for discrete event systems. Proc. of the IEEE, 77(1):14–23, 1989.

[20] Joshua H Goldwyn, Nikita S Imennov, Michael Famulare, and Eric Shea-Brown. Stochastic differential equation models for ion channel noise in Hodgkin-Huxley neurons. Phys Rev E Stat Nonlin Soft Matter Phys, 83(4 Pt 1):041908, Apr 2011.

[21] Joshua H Goldwyn and Eric Shea-Brown. The what and where of adding channel noise to the Hodgkin-Huxley equations. PLoS Comput Biol, 7(11):e1002247, Nov 2011.

[22] Peter J. Haas. Stochastic Petri Nets: Modelling Stability, Simulation. Springer, New York, first edition, 2002.

[23] A. L. Hodgkin and A. F. Huxley. A quantitative description of membrane current and its application to conduction and excitation in nerve. J Physiol, 117:500–544, 1952.

[24] James P Keener and Jay M Newby. Perturbation analysis of spontaneous action potential initiation by stochastic ion channels. Phys Rev E Stat Nonlin Soft Matter Phys, 84(1-1):011918, Jul 2011.

[25] T. Kipersky and J. A. White. Stochastic models of ion channel gating. Scholarpedia, 3(1):1327, 2008.

[26] Thomas G. Kurtz. Representations of Markov Processes as Multiparameter Time Changes. Ann. Prob., 8(4):682–715, 1980.

[27] Thomas G. Kurtz. Approximation of population processes, CBMS-NSF Reg. Conf. Series in Appl. Math.: 36, SIAM, 1981.

[28] Carlo Laing and Gabriel J. Lord, editors. Stochastic Methods in Neuroscience. Oxford University Press, 2010.

[29] Chang Lee and Hans Othmer. A multi-time-scale analysis of chemical reaction networks: I. deterministic systems. Journal of Mathematical Biology, 60:387–450, 2010. 10.1007/s00285-009-0269-4.

[30] Lorin S Milescu, Tadashi Yamanishi, Krzysztof Ptak, and Jeffrey C Smith. Kinetic properties and functional dynamics of sodium channels during repetitive spiking in a slow pacemaker neuron. J Neurosci, 30(36):12113–27, Sep 2010.

[31] Hiroyuki Mino, Jay T Rubinstein, and John A White. Comparison of algorithms for the simulation of action potentials with stochastic sodium channels. Ann Biomed Eng, 30(4):578–87, Apr 2002.
[32] C Morris and H Lecar. Voltage oscillations in the barnacle giant muscle fiber. Biophys J, 35(1):193–213, Jul 1981.

[33] C. Morris and H. Lecar. Voltage oscillations in the barnacle giant muscle fiber. Biophysical Journal, 35(1):193 – 213, 1981.

[34] Jay M Newby, Paul C Bressloff, and James P Keener. Breakdown of fast-slow analysis in an excitable system with channel noise. Phys Rev Lett, 111(12):128101, Sep 2013.

[35] Muruhan Rathinam, Patrick W. Sheppard, and Mustafa Khammash. Efficient computation of parameter sensitivities of discrete stochastic chemical reaction networks. Journal of Chemical Physics, 132:034103, 2010.

[36] Martin Riedler and Girolama Notarangelo. Strong Error Analysis for the Θ-Method for Stochastic Hybrid Systems. arXiv preprint arXiv:1310.0392, 2013.

[37] J. Rinzel and G.B. Ermentrout. Analysis of neural excitability and oscillations. In C. Koch and I. Segev, editors, Methods in Neuronal Modeling. MIT Press, second edition, 1989.

[38] D. Terman J. Rubin. Geometric singular perturbation analysis of neuronal dynamics. In B. Fiedler, editor, Handbook of Dynamical Systems, vol. 2: Towards Applications, pages 93–146. Elsevier, 2002.

[39] Nicolaus T Schmandt and Roberto F Galán. Stochastic-shielding approximation of Markov chains and its application to efficiently simulate random ion-channel gating. Phys Rev Lett, 109(11):118101, Sep 2012.

[40] Deena R Schmidt and Peter J Thomas. Measuring edge importance: a quantitative analysis of the stochastic shielding approximation for random processes on graphs. arXiv preprint arXiv:1308.5179, 2013.

[41] Tilo Schwalger, Karin Fisch, Jan Benda, and Benjamin Lindner. How noisy adaptation of neurons shapes interspike interval histograms and correlations. PLoS Comput Biol, 6(12):e1001026, 2010.

[42] R Shingai and FN Quandt. Single inward rectifier channels in horizontal cells. Brain Research, 369(1-2):65–74, Mar 26 1986.

[43] E Skauge and L Walløe. Firing behaviour in a stochastic nerve membrane model based upon the Hodgkin-Huxley equations. Acta Physiol Scand, 107(4):343–63, Dec 1979.

[44] Rishi Srivastava, David F. Anderson, and James B. Rawlings. Comparison of finite difference based methods to obtain sensitivities of stochastic chemical kinetic models. Journal of Chemical Physics, 138(7):074110, 2013.

[45] Adam F. Strassberg and Louis J. DeFelice. Limitations of the Hodgkin-Huxley formalism: Effects of single channel kinetics on transmembrane voltage dynamics. Neural Computation, 5:843–855, 1993.

[46] J.A. White, C.C. Chow, J. Ritt, C. Soto-Trevino, and N. Kopell. Synchronization and oscillatory dynamics in heterogeneous, mutually inhibited neurons. Journal of Computational Neuroscience, 5:5–16, 1998.

[47] J.A. White, J.T. Rubinstein, and A.R. Kay. Channel noise in neurons. Trends Neurosci., 23:131–137, 2000.
[48] Darren J. Wilkinson. *Stochastic Modelling for Systems Biology*. Chapman &
Hall/CRC, Nov 2011.