Hierarchic Stochastic Modelling Applied to Intracellular Ca\textsuperscript{2+} Signals

Gregor Moenke\textsuperscript{1}, Martin Falcke\textsuperscript{1}, Keven Thurley\textsuperscript{1,2}*

\textsuperscript{1}Mathematical Cell Physiology, Max Delbrück Center for Molecular Medicine, Berlin, Germany, \textsuperscript{2}Institute for Theoretical Biology, Charité Universitätsmedizin, Berlin, Germany

Abstract

Important biological processes like cell signalling and gene expression have noisy components and are very complex at the same time. Mathematical analysis of such systems has often been limited to the study of isolated subsystems, or approximations are used that are difficult to justify. Here we extend a recently published method (Thurley and Falcke, PNAS 2011) which is formulated in observable system configurations instead of molecular transitions. This reduces the number of system states by several orders of magnitude and avoids fitting of kinetic parameters. The method is applied to Ca\textsuperscript{2+} signalling. Ca\textsuperscript{2+} is a ubiquitous second messenger transmitting information by stochastic sequences of concentration spikes, which arise by coupling of subcellular Ca\textsuperscript{2+} release events (puffs). We derive analytical expressions for a mechanistic Ca\textsuperscript{2+} model, based on recent data from live cell imaging, and calculate Ca\textsuperscript{2+} spike statistics in dependence on cellular parameters like stimulus strength or number of Ca\textsuperscript{2+} channels. The new approach substantiates a generic Ca\textsuperscript{2+} model, which is a very convenient way to simulate Ca\textsuperscript{2+} spike sequences with correct spiking statistics.

Introduction

The molecular state transitions and interactions inside cells forming pathways and functional units are inherently random [1–3]. Some cellular subsystems involve sufficiently large molecule numbers to be well described by deterministic mean field dynamics, but many are best described as stochastic processes. Additionally, cells show considerable heterogeneity even within clonal populations. Biological noise and cell-to-cell variability have been identified and studied in chemotaxis [4], gene expression [5], cell signalling [6–8] and cell differentiation [9,10]. A variety of mathematical strategies can be used to analyse these stochastic dynamical systems, including approximation techniques such as Langevin and Fokker-Planck Equations [11,12], and exact methods like the chemical master equation or its simulation by the Gillespie algorithm [13]. Approximation techniques are often based on the assumption of Gaussian white noise, which is only valid for large copy-numbers of identical components and fluctuations which are small compared to the mean value. However, in biological processes like gene expression [1] or Ca\textsuperscript{2+} dynamics [14,15], the change of state of a single molecule may have a huge impact on systems dynamics [see also [16]]. To describe such systems without rough approximations, we recently developed a new modelling framework based on emergent behaviour of biomolecules [17–19].

Ca\textsuperscript{2+} is a ubiquitous second messenger transmitting information in many cases by repetitive cytotoxic concentration spikes [14,15,18]. An important class of Ca\textsuperscript{2+} signals is mediated by Inositol-1,4,5-trisphosphate (IP\textsubscript{3}), which is produced in response to hormonal activation of cell surface receptors [14,15]. IP\textsubscript{3} then binds IP\textsubscript{3} receptors (IP\textsubscript{3}R) in the endoplasmic reticulum (ER) and thus sensitisises them for activation by Ca\textsuperscript{2+}. IP\textsubscript{3}Rs are organised as clusters of about 1 to 20 IP\textsubscript{3}R molecules [20–22]. Active IP\textsubscript{3}Rs act as Ca\textsuperscript{2+} channels, releasing Ca\textsuperscript{2+} ions from the ER lumen into the cytosol. Upon sensitisation by IP\textsubscript{3} they are successively activated by Ca\textsuperscript{2+} -induced Ca\textsuperscript{2+} release (CICR). This mechanism is based on the opening probability of IP\textsubscript{3}Rs, which increases with the local Ca\textsuperscript{2+} concentration, up to a threshold value where further increase of the Ca\textsuperscript{2+} concentration becomes inhibitory [23,24]. The outflux of Ca\textsuperscript{2+} eventually stops either because of depletion of the ER Ca\textsuperscript{2+} stores or the threshold [Ca\textsuperscript{2+}] for channel inhibition is reached. Sarco-endoplasmic reticulum Ca\textsuperscript{2+} ATPases pump Ca\textsuperscript{2+} back into the ER after release. The transient increases in cytosolic [Ca\textsuperscript{2+}] trigger downstream effects like activation of protein kinase C [15,25]. The clustering of IP\textsubscript{3}Rs implies that global cell-wide Ca\textsuperscript{2+} signals result from a hierarchic cascade of single channel opening (‘blips’) over cluster opening (‘puffs’) to opening of several clusters (‘wave’ or ‘spike’), which arise when open clusters open neighbouring clusters due to Ca\textsuperscript{2+} diffusion and CICR. This cascade is well characterised by live cell imaging and mathematical modelling studies [14,15]. We and others have recently demonstrated that cellular Ca\textsuperscript{2+} signals are repetitive stochastic events [6,18,26,27], although at certain conditions they appear regular and have long been analysed with deterministic mathematical models [28,29]. Thus, the Ca\textsuperscript{2+} spikes arise by a multiscale stochastic process emerging by clustering of IP\textsubscript{3}Rs. If we seek for a theory to...
in intracellular Ca\textsuperscript{2+} signalling, what are the experimental observations such a theory should explain? Ca\textsuperscript{2+} spike sequences are random with average interspike intervals (ISI) from a few tens of seconds to hundreds of seconds. Due to this random spiking, simulation of individual time series of the Ca\textsuperscript{2+} concentration is less meaningful than with deterministic limit cycle oscillators. There are, however, well-defined properties of the stochastic sequences characterising the system. The moment relation between the standard deviation $\sigma$ and the average $\tau_{\text{av}}$ of ISI is linear with a positive slope between 0.2 and 1 [6], the slope of the moment relation of individual cells agrees with the population relation [50,31], i.e. it is robust against cell-to-cell variability and cell type specific [32]. These results were established by single cell experiments with astrocytes, microglia, PLA stem cells and HEK cells [6]. The ISI obey simple two-parameter distributions [6]. Weakening spatial coupling between IP\textsubscript{3}R clusters by Ca\textsuperscript{2+} buffers like EGTA destroys cellular spikes and any variation of Ca\textsuperscript{2+} signalling on their time scale [27]. Therefore, a mechanistic theory should be able to explain the emergence of cellular spiking on the basis of cluster properties and cluster coupling only. Moreover, the theory should link the statistical properties to biological parameters like stimulation strength and buffer concentrations.

We meet all those requirements by an approach called hierarchic stochastic modelling [17]. What is measured in vivo can be correctly described only by waiting-time probabilities since it represents transitions between observable states like ‘open’ or ‘closed’ of a specific IP\textsubscript{3}R cluster. Hierarchic stochastic modelling deals with waiting-time probabilities for state-changes in a reduced state-space. The reduction of the state space focuses on functional or observational aspects of the system. In contrast the full state space contains all biochemically possible states of the involved molecules. Their dynamics can then be described by rate laws yielding the usual chemical master equation. To avoid computational overhead by computing non observable state transitions and facilitate analytic solutions, all microscopic states corresponding to one function (e.g. for an ion channel ‘open’) are subsumed into one observable state. The set of all observable states forms the reduced state-space. This entails non-exponential waiting-time probabilities instead of rate-laws for the description of transitions between the observable states.

Here, we develop a convenient solution technique for hierarchic stochastic systems by the Laplace transform, which directly leads to analytical expressions for statistical properties of important system variables. Moreover, we present a minimal mechanistic model for stochastic Ca\textsuperscript{2+} spikes based on recent data from live cell imaging. We provide analytic approximations for the dependence of system statistics on biological parameters. Finally, based on the observation of a very robust time scale separation in the mechanistic model, we find a surprisingly convenient description of Ca\textsuperscript{2+} spike statistics, which is in excellent agreement with mechanistic models and with experimental data.

**Results**

Cellular Ca\textsuperscript{2+} dynamics is a multiscale stochastic process driven by huge gradients in the cytosolic Ca\textsuperscript{2+} concentration, and detailed computer simulations are difficult to implement and time-consuming [14,32]. A concise, analytically treatable mechanistic model was published only recently by two of us [17]. In the following, we start by explaining the method that made this possible and presenting an extension to the method, which is solution by Laplace transform of generalised exponential distributions. After that, we construct the input data required for the modelling approach in terms of analytical functions of cellular parameters. The third section analyzes a minimal mechanistic Ca\textsuperscript{2+} model, which allows for analytical solutions and elucidates the role of a global negative feedback. Finally, based on that analysis, we present a proper derivation of a very simple description of stochastic Ca\textsuperscript{2+} dynamics called generic model, which very nicely describes the experimental Ca\textsuperscript{2+} spike statistics.

**Hierarchic Stochastic Modelling Analysed by the Laplace Transform**

The traditional formulation of a stochastic model starts from the state transitions of the IP\textsubscript{3}R subunits [14]. In most single channel models [33], three binding sites are assumed for each of the four subunits, one activating and one inhibiting site for Ca\textsuperscript{2+} and one site for IP\textsubscript{3}. A complete stochastic description determines the time course of the probability for each system state. Their dynamics are described by the master equation [11].

$$\frac{d}{dt} P_i(t) = \sum_{j=1}^{N} [q_{ij} P_j(t) - q_{ji} P_i(t)]. \tag{1}$$

$N$ is the number of system states ($N=\infty$ is possible) and $P_i(t) = P(i,t|j,0)$ is the conditional probability that the system is in state $i$ at time $t$ conditioned by being in state $j$ at time $t=0$. The $q_{ij}$ are the rates for state transitions $i \rightarrow j$ (see Table 1 for a summary of the notation). Figure 1A illustrates the state scheme of a hypothetical receptor channel molecule $x$ with only four states $x_1, \ldots, x_4$ (3 closed states, 1 open state). This case can easily be treated directly by Eq. 1. However, if we consider several interacting molecules with more than 100 possible states each, like those in typical ion channels [33–35], Eq. 1 becomes intractable due to the large number of system states.

When describing a biological system with a mathematical model, we are often not interested in all the molecular details, but rather in few states that are accessible to experimental quantification or are important for system function. In the case of Ca\textsuperscript{2+} dynamics the main question is whether the channel is closed or not, and not which of the many possible closed states it is in. Nevertheless, the channel state depends on the dynamics between all its closed states, such that a rigorous treatment demands the inclusion of all possible microscopic state changes into Eq. 1. We circumvent state space explosion by lumping microscopic states belonging to the same observable state (Fig. 1B). Consider stochastic transitions between observable system states $S_l$ each representing a set of microscopic system states. Because a sequence of microscopic transitions occurs between two consecutive transitions, each transition $S_l \rightarrow S_k$ is not governed by a simple rate-law with exponentially distributed waiting-time, as in Eq. 1. The sequence of microscopic transitions causes a non-exponential waiting time distribution (Fig. 1C). These waiting time distributions introduce memory into the process such that the Markov property is lost (see Text S1). The probability densities of the waiting times until specific state transitions $i \rightarrow l$ occur, $P_l(t)$, are considered as input data for our Ca\textsuperscript{2+} model (see Section Materials and Methods).

For analytical analysis, we need to solve a non-Markovian master equation determined by the $P_l(t)$. As an extension to the previously published theory [17], we find that analytical solutions for the stationary occupancy probabilities in the observable system states (see Fig. 1) can be obtained by Laplace transform followed by solution of linear algebraic systems (see Section Materials and Methods and Text S1). Importantly, we do not need to calculate...
the often awkward inverse Laplace transform. We can obtain the average and standard deviation of interevent times which are key characteristics of biological processes like Ca\textsuperscript{2+} signalling or chemotaxis [17,36]. The expressions for the moments \( \langle t^n \rangle \) of interevent times are particularly convenient if the system can be represented as a linear chain with \( N \) distinct states \( X \in \{0,1,\ldots,N-1\} \) and state transitions \( 0 \rightarrow 1 \rightarrow \ldots \rightarrow N-1 \) (see Section Materials and Methods). For instance, in the case of \( N = 3 \) we obtain,

\[
\langle t^n \rangle = (-1)^n \sigma^n \sum_{i=0}^{n} \left[ \frac{\Psi_{0i}(s)\Psi_{12}(s)}{1 - \Psi_{0i}(s)\Psi_{10}(s)} \right] s^{-n},
\]

where \( \Psi_{ij}(s) \) denotes the Laplace transformed conditioned waiting times. From Eq. 2, average \( \langle t \rangle \) and standard deviation \( \sigma = \sqrt{\langle t^2 \rangle - \langle t \rangle^2} \) of interevent times are readily computable.

A limiting step in this solution strategy is the availability of the Laplace transform \( \Psi_{10}(s) \). Multistep processes like the ones considered here (see Fig. 1) are often well described by \( \gamma \)-distributions [11], which are also frequently observed in biological dynamical systems [27,37,38]. However, expressions like Eq. 2 are impractical for \( \gamma \)-distributed conditioned waiting times. Instead of that, we found that generalised exponential (GE) distributions [39].

\[
\psi(t) = 2\lambda(1 - e^{-\lambda t})^{a-1}e^{-\lambda t}
\]

are very convenient for operations in Laplace space and approximate the multistep processes leading to Ca\textsuperscript{2+} spikes equally well as \( \gamma \)-distributions (see below). In Eq. 3, \( a \geq 1 \) is the shape parameter and \( \lambda > 0 \) is the scale parameter. Note that for \( a = 1 \), Eq. 3 is an exponential distribution with parameter \( \lambda \).

With the use of Laplace transformed GE distributions, we have found a straightforward strategy to solve for dynamic properties of non-Markovian systems. Importantly, computational effort is essentially reduced to solution of a linear algebraic system, and thus the strategy can easily be extended to larger systems, in contrast to the Volterra integral equations discussed in Ref. [17].

**Waiting Times of Ca\textsuperscript{2+} Dynamics and their Dependence on Cellular Parameters**

Intracellular Ca\textsuperscript{2+} dynamics can be described in terms of an observable open and an observable closed state of clusters of IP\textsubscript{3}R [14,17,18]. All microscopic states in which all channels of the cluster are closed belong to the observable closed state. All other states belong to the observable open state. The subunits of individual IP\textsubscript{3}R continuously change states by binding and unbinding of regulatory molecules (microscopic dynamics), but only the closing of the last open channel and the first opening when all channels are closed represent transitions between the observable states of a cluster of IP\textsubscript{3}R (see Fig. 1). The waiting times until a cluster opens or closes are determined by true probability densities, which we denote by \( \psi_o(t) \) and \( \psi_c(t) \), respectively. Cellular Ca\textsuperscript{2+} dynamics consist of the action of several interacting clusters, constituting a state space \( S_1, \ldots, S_N \) of possible configurations of open and closed clusters. The conditioned waiting times \( \Psi_{ij}(t) \) are products of the basic transition probability \( \psi_o(t) \) or \( \psi_c(t) \) of a cluster and the probabilities that all other clusters remain in their states, \( \tilde{\psi}_o(t) \) can be derived from experimental data (see Ref. [17] and Text S1 for details). \( \tilde{\psi}_o(t) \) depends on the number of IP\textsubscript{3}Rs per cluster \( N_0 \) and on the IP\textsubscript{3}Rs

---

**Figure 1. Schematic illustration of hierarchic stochastic modelling.** (A) Markovian models consider all possible changes of a molecule \( x \), e.g. binding of regulatory factors, production, degradation. Thus, microscopic state transitions \( x_i \rightarrow x_j \) are governed by rate parameters \( q_{ij} \). In the case of IP\textsubscript{3}R, the \( q_{ij} \) refer to binding (unbinding) of IP\textsubscript{3} and Ca\textsuperscript{2+} to (of) their respective binding sites on certain IP\textsubscript{3}R subunits. (B) A typical path of state changes according to (A) includes repetitive switches between individual states before a specific state of interest (here, \( x_{IP3 Rs} \)) is reached. The observable states \( S_i \) are subsets of all microscopic states (here e.g. \( S_i = \{x_1,x_2,x_3\} \)). In Ca\textsuperscript{2+} dynamics, observable states may represent open or closed clusters of IP\textsubscript{3}Rs. (C) Microscopic transitions \( x_i \rightarrow x_j \), with parameters \( q_{ij} \) induce exponential waiting time probabilities \( q_{ij} \exp(-\sum q_{ij}) \). Observable state transitions \( S_i \rightarrow S_j \) follow waiting time probabilities \( \Psi_{ij}(t) \), which can have one or several maxima due to their internal dynamics.
concentration, which represents the strength of a hormonal stimulus in our model. Intracellular IP$_3$ dynamics are still an open question and beyond our scope here. Because of CICR, the individual opening probabilities depend on the local Ca$^{2+}$ concentration, which is a function of the number of open clusters. The value of the Ca$^{2+}$ concentration rise caused by an open cluster at the locations of the other clusters gives the strength of spatial coupling between clusters.

All parameters that modify properties of Ca$^{2+}$ diffusion in the cell and describe the spatial configuration of clusters affect the coupling [14]. Therefore, we seek for a valid analytical approximation of $\psi_o(t)$ and its dependence on [Ca$^{2+}$], $N_{ch}$ and [IP$_3$], which was not available in our previous article [17]. Recent data from TIRF microscopy revealed key properties of $\psi_o(t)$ [27,40], but are not sufficient for a closed mathematical model, because the dependencies of the Ca$^{2+}$ and IP$_3$ concentration have not yet been established. A good approximation was achieved by using two-parametric probability distributions like the waiting time density of the inhomogeneous Poisson process or the $\gamma$-distribution, indicating recovery times between successive opening events [27]. However, in some cases, simple exponential distributions without any recovery process described the data equally well [27,40]. This coincides with our observation that single channel models generate approximately exponential distributions at low (basal) local Ca$^{2+}$ concentration [17]. This is probably because intermediate inhibitory states are rarely reached at low [Ca$^{2+}$] in these models [14]. Based on these findings, we assume for the purposes of this study that the transition from 0 to 1 open clusters (puff probability) is governed by a pure exponential distribution with average interpuff interval as recorded for SH-SY5Y cells for typical values of cellular parameters. For subsequent cluster openings (1$\rightarrow$2$\rightarrow$3, ... open clusters), we use GE distributions (see Eq. 3), which fit the data equally well as $\gamma$-distributions (Fig. 2A) but are more convenient for operations in Laplace space. That means, for the puff probability distribution, we set $\gamma=1$ and identify $\lambda=\lambda_0$ with the puff rate.

As we have determined the functional form of the $\psi_o(t)$, it remains to establish the dependencies of the distribution parameters $\lambda$ and $\gamma$ on the cellular parameters [Ca$^{2+}$], $N_{ch}$ and [IP$_3$]. Unfortunately, for such dependencies no directly usable experimental data are available. However, earlier work has shown that $\psi_o(t)$ can be computed from single channel models by solution of a Markovian master equation [41]. The single channel models were partly based on data from in vitro studies, and the new in vitro experimental data [27] confirmed the type of distributions resulting from those models. In particular, we adopt an implementation of the De Young Keizer model [34,41], one of the most accepted single channel models in the field (see Table S1 and Text S1 for details). To obtain an analytical description of the parameters $\lambda$ and $\gamma$, we performed simulations of the De Young Keizer model with varying cellular parameters and inspected functional forms matching the simulations. The base-level Ca$^{2+}$
concentration, an important (and experimentally undetermined) parameter needed for the simulations, was chosen by comparison with the measured puff rate [27].

We found (Fig. 2B–C) that the parameter dependencies of the puff rate \( \lambda_0 \) can be approximated by a Hill function.

\[
\lambda(x) = \frac{U_x x^{n_x}}{(L_x)^{n_x} + x^{n_x}}, \tag{4}
\]

where \( x = [\text{IP}_3] \) or \( N_{ch} \). Note that \( N_{ch} \) is a discrete variable, and Eq. 4 should only be evaluated at integer values in this case. This is also true for the dependencies of the parameter \( \lambda \) of the GE distribution on \( x = \text{Ca}^{2+} [\text{IP}_3]\) or \( N_{ch} \) (Fig. 2D–F). Dependencies of \( \lambda_0 \) on \( N_{ch} \) and of \( \lambda_0 \) on \( [\text{Ca}^{2+}] \) are close to the linear regime (Fig. 2G,F). The dependence on all parameters (Fig. 2G–I) can be fitted by another Hill equation,

\[
\alpha(x) = \frac{V_x x^{n_x}}{(K_x)^{n_x} + x^{n_x}} + 1, \tag{5}
\]

where the ‘+1’ in Eq. 5 reflects the fact that GE-distributions have a different shape at \( x < 1 \). The fits of analytical functions to computed opening probabilities \( \psi_0 \) at varying cellular parameters provide all the information needed to perform calculations of the hierarchic stochastic model (see Text S1). Fit parameters for the examples shown in Fig. 2 are given in Table S2.

The analytical parameter dependencies of waiting times presented here are a huge advantage over the approach followed in Ref. [17], where the single channel models were evaluated de novo for each of model simulation at a certain set of cellular parameters. In particular, Eqs. 5–4 allow for integration of the hierarchic stochastic model into models of signal transduction networks in future research.

**Analysis of a Minimal Mechanistic Ca\(^{2+}\) Model**

Solution of the hierarchic stochastic model by the Laplace transform is most convenient in the case of a linear chain (see Section Materials and Methods). As applied to stochastic \( \text{Ca}^{2+} \) dynamics, cluster arrangements with equal distances between all clusters and identical clusters lead to linear chain models. That can be achieved only up to 4 clusters in a tetrahedral arrangement, but this number of clusters leads to physiologically meaningful results concerning spike statistics already [17], in particular in conjunction with the generic model introduced below. Further details needed to derive the \( \text{Ca}^{2+} \) model are given in Text S1. By using the analytic approximations, Eqs. 3–4, we can relate cellular parameters like the cytosolic \( \text{IP}_3 \) concentration to the parameters of the opening and closing waiting time densities.

Figure 3 shows a typical model calculation with realistic parameters and spiking statistics. The opening probabilities (Fig. 3A–B) are split into two temporal regimes for opening of the first cluster with the constant rate \( \lambda_0 \) (puff probability, inset) and successive cluster openings. If the first cluster has opened, the probability for the opening of successive clusters is greatly enhanced. This reflects CICR in our stochastic model. The stationary occupancy probabilities (Fig. 3C) for the different states can be computed from Eq. 14. We performed delayed stochastic simulations (Fig. 3D) by methods similar to the Gillespie algorithm [13,17] to provide a picture of the actual dynamics.

We define a spike in the tetrahedron model by opening of all four clusters. The average ISI \( \langle t \rangle \equiv T_{av} [\text{IP}_3],N_{ch} \) can then be calculated analytically by Eq. 15. Figure 4A shows that \( T_{av} \) changes by about one order of magnitude with varying \( N_{ch} \) and \([\text{IP}_3]\), respectively. If we suppose a realistic coverage of the cellular parameter range, a strong cell to cell variability in terms of \( T_{av} \) should be expected in experiments, and is indeed reported [6,26]. Even though we cannot calculate the ISI density directly, apart from very simple cases (see Text S1) where the inverse Laplace transform is feasible, we can characterize it with higher moments, which are directly available from our theory (see Methods, Eq. 15).

A Generic Model Based on Local Puff Dynamics

We use the simple model for the global puff process introduced in Ref. [6] to introduce a global feedback into the hierarchic stochastic process. The global feedback is included as a slow recovery of the spike rate after a spike occurred, which leads to an inhomogeneous Poisson process with time dependent spike rate.

\[
\kappa(t) = \kappa_0 (1 - e^{-\xi t}), \tag{6}
\]

with the Poisson spike rate \( \kappa_0 \) and the recovery rate \( \xi \). With this formulation the probability for a spike at \( t = 0 \) equals zero. This represents an inhibitory or negative global feedback, at high \( \text{Ca}^{2+} \) concentrations, which naturally occurs after a global \( \text{Ca}^{2+} \) spike.

The probability for a spike is the probability for a puff times the probability that the puff opens the other clusters. The latter is in mathematical terms a splitting probability that a single puff triggers a consecutively opening of all clusters. For the tetrahedron
This means to go from one open cluster to four open clusters before reaching the ground state with all clusters closed again. It is analytically given by \[17\].

\[ C_{14} = \frac{C_{12}C_{23}C_{34}}{1 + C_{12}(C_{23} - 1) + C_{23}(C_{34} - 1)}, \]

with the one-step splitting probabilities \( C_{i,i+1} \).

Figure 5 shows the dependence of \( C_{14} \) on cellular parameters. As in Ref. [17], we formulate puff generation as inhomogeneous Poisson process with time-dependent rate \( \lambda_0(1 - e^{-\xi t}) \), where \( \xi \) is the recovery rate. This effectively inhibits the occurrence of a local puff event right after a spike and hence incorporates global negative feedback. The puff rate asymptotically approaches the stationary value \( \lambda_0 \) as time increases. We now resample the puff process into the global spike process with probability \( C_{14} \) and its complementary process with probability \( (1 - C_{14}) \). The complementary process is the Poisson process describing the failed puffs, the ones that did not lead to a global Ca\(^{2+}\) spike.

In general, a cellular spike will require \( N \) open clusters, the splitting probability should therefore be denoted by \( C_{i,N} \). Clusters are heterogeneous with respect to the number of channels per cluster as well as to their distance to neighboring clusters. Hence, the puff probability as well as the splitting probability are cluster specific. We introduce the individual puff rate \( \lambda_{0,i} \) and splitting probability \( C_{1,N,i} \) for every cluster \( i \). Some failed puffs occur between two spikes. The probability for the interspike interval \( t \) is then given by the sum over the probabilities for all possible numbers of puffs in between two spikes from 0 to \( \infty \). This finally leads to the expressions constituting the inhomogeneous Poisson splitting (see also Text S1).

\[ \kappa(t) = \sum_{i=1}^{N} \lambda_{0,i} C_{1,N,i}(1 - e^{-\xi t}) \]

\[ p(t) = \kappa(t)e^{-\lambda_0 t}, \]

Figure 2. Parameter dependencies of the opening probabilities \( \psi \). (A) Comparison of fits of Gamma and GE distributions to the De Young-Keizer model (see Ref. [34] and Table S1) at \([\text{Ca}^{2+}] = 0.1 \mu\text{M}\). (B–I) Circles are fits of results from the De Young-Keizer model to GE distributions (Eq. 3) with parameters \( x \) and \( \lambda \), as in A. Lines are (B,D,E) nonlinear fits to Eq. 4, (C,F) linear fits, (G–I) nonlinear fits to Eq. 5. (B–C) show the dependencies for the puff-rate, i.e. at base-level \([\text{Ca}^{2+}] = 0.1 \mu\text{M}\) (see Text S1). (D–H) show parameter dependencies in the regime of a puff, i.e. with one nearby cluster already open, yielding \([\text{Ca}^{2+}] = 0.6 \mu\text{M}\) (see Text S1). Other parameter values (if not varied on the x-axis) are \( N_{ch} = 5 \), \([\text{IP}_3]\) = 1 \mu\text{M}\).

doi:10.1371/journal.pone.0051178.g002
where \( A(t) = \sum_{i=1}^{N} \lambda_i c_{i,N} \int_0^t (1 - e^{-C}) dt \) is the intensity and \( p(t) \) is the density function of the probability that a global Ca\(^{2+}\) spike occurs at time \( t \) after the previous spike.

Comparison with Eq. 6 shows that we found an expression of the Poisson spike rate \( k_0 \) in terms of cluster properties,

\[
k_0 = \sum_{i=1}^{N} \lambda_i c_{i,N,i}.
\]

Thus, \( k_0 \) can be taken as total spike rate, summing up the individual contributions of clusters \( i \) to the spike rate. That is, the individual puff rates \( \lambda_i \) of cluster \( i \) are rescaled with the coupling strength \( C_{i,N,i} \). The average interspike interval \( T_{av} \) and the standard deviation \( \sigma \) can then be calculated analytically with the results given in Ref. [6].

To confirm the validity of the generic model in the presence of global feedback, we performed stochastic simulations of the hierarchical stochastic model as described in Ref. [17] and compared them with the analytic results of the generic model. For the homogeneous tetrahedron cluster arrangement, Eq. 10 gives \( k_0 = 4 \lambda_C C_{14} \). This new generic model approximates the average ISI very well (Fig. 6A). The experimental moment relations are in very good agreement with the moment relations obtained for \( \xi = const \) (Fig. 6B). For a single cell the specific value of \( k_0 \) positions the cell on that moment relation as indicated by the symbols in Fig. 6. Note that very small slopes of the moment relation below 0.3, which have been reported for HEK293 cells [6], can be generated by adding cooperativity to the generic model [18]. The saturation for both \( \lambda_0 \) and \( C_{14} \) with respect to the IP\(_3\) concentration (see Figs. 2 and 5) gives rise to an intrinsic minimal ISI in the model (Fig. 6A–B). The relation of this minimal ISI to the physiological minimal ISI reported from single cell measurements [6] should be elucidated by further research.

By applying Poisson splitting (see Text S1 for details), we implicitly neglect the possible dynamics between the puff and the spike events. Most notably we completely miss the duration of the failed puffs. The number of those failed puffs is given on average by \( 1/C_{14} \), so we expect the error we make by using the splitting scales exactly with \( 1/C_{14} \). By comparing our exact analytic results for the tetrahedron model with our generic model without recovery we show that this is indeed the case (Fig. S1). Moreover we have computed the relative error and state that it is bounded and smaller than 10% (see Text S1). The great advantage of the generic model is its capability of providing a closed expression for the ISI distribution reproducing the moment relation between the standard deviation \( \sigma \) and the average \( T_{av} \) of ISIs found experimentally, incorporating the local puff dynamics. The key parameter for the local dynamics is the coupling strength \( C_{14} \) obtained by hierarchical stochastic modelling.

**Discussion**

We present an extension to a description of complex stochastic networks in terms of observable states and non-exponential transition time distributions. The approach can be used for very efficient stochastic simulations and in many cases analytical solutions can be obtained. We apply hierarchical stochastic modelling to spatially resolved stochastic Ca\(^{2+}\) dynamics. Apart from using a new solution method, we made further progress in the Ca\(^{2+}\) model published in [17] by developing analytic approximations of model dependencies on cellular parameters and taking into account recent experimental data for model.
calibration. This permitted us to rationalise time scale separation between puffs and waves in a physiological setting, and subsequently to find a derivation of a generic description of the stochastic spike generating process.

The theory presented here is able to calculate the statistical properties of cellular Ca\(^{2+}\) spiking from cluster properties and spatial coupling. We have derived the moments of the ISI distribution by Laplace transform of the non-Markovian master equation, and we approximated the complete ISI distribution with the generic model. This simplistic description was postulated earlier in the Falcke group [6] but only now we can give a mechanistic explanations for its validity. The generic model is only applicable for a large time scale separation between the interpuff intervals and the ISIs. However, this separation was shown experimentally for many cell types [27,44]. As is often the case with approximate solutions, the generic model allows for deeper insight than the direct solution of the integro-differential equations. The theory explains the role of spatial arrangement of clusters and strength of spatial coupling in the spike probability in a simple and intuitive way. We introduced a phenomenological factor \(k_0\) which rescales the local puff dynamics to the global spike dynamics. This factor also explains the observed large cell-to-cell variability as differences in cluster properties and cluster arrangement. While we started with the assumption that the puff probability slowly recovers from negative feedback, Eqs. 8 also allows for a time dependency of the splitting probability \(C_{14}\). Hence, more complicated recovery processes arising from a variety of feedbacks could replace the simple relaxation chosen here. The model exhibits almost linear moment relations between standard deviation and average in agreement with experimental results. Changes in \(k_0\) of an individual cell by experimental manipulations or between cells due to cell-to-cell variability do not alter the slope of the moment relation. That explains its robustness against changes of local properties and coupling strength, i.e. cell-to-cell variability within a given cell type, which has been observed [17,32]. The rate of recovery from global feedback strength \(\xi\) is cell type specific, which explains the cell type specificity of the moment relation.
Our theory is based on recent experimental data from live cell imaging for opening and closing transition times of Ca\textsuperscript{2+} channel clusters. A description of cellular Ca\textsuperscript{2+} dynamics further requires information about the coupling of individual Ca\textsuperscript{2+} channel clusters and the dependence on cellular parameters like strength of the upstream signal (IP\textsubscript{3} concentration) and cluster size. These functional relations could, up to now, only be obtained from single channel models, which are well established and have successfully been used in earlier modelling studies [14,17,32]. Based on this, we suggest analytical approximations for model dependencies on cellular parameters. This approach can be used for integration of the model into mathematical descriptions of larger signalling cascades.

We calculated the splitting probability $C_{iK}$ for $N=4$, where $N$ is the number of open clusters forming a cellular release spike. Typically, a cellular spike will consist of more than 4 open clusters. However, the calculation provides a good description of cellular spiking, if a cell has a preferred nucleation area initiating most of the cellular spikes. The stronger the release in such an area is, the more likely is the nucleation of a wave due to Ca\textsuperscript{2+} diffusion and CICR. At a sufficient value, this probability will be very close to one. The tetrahedron model is a good description of cellular spiking with nucleation areas, if that value is reached by 4 open clusters. Preferred nucleation areas have indeed been observed, e.g. in hepatocytes [45]. In the spirit of hierarchical modelling, the results with one nucleation area can then easily be generalised to such areas.

We find that the minimal tetrahedral system considered here stays in the regime of a Poisson process for Ca\textsuperscript{2+} spikes, characterized by a coefficient of variation equal to one, even for relatively large deviations from the physiological parameters. Only a drastic reduction of the time scale separation (or increase of the coupling strength between clusters), mediated by increased puff rate $\lambda_0$ and reduction of the channel closing rate $\gamma$, can reduce the noise level by means of array enhanced coherence resonance [42,43]. Therefore, we suppose that the lower noise levels observed in many cell types [6] indeed stems from feedback processes, as suggested by earlier results [17]. However, the experimental proof for the importance of such feedback processes for Ca\textsuperscript{2+} spike statistics is still lacking. Given an inhomogeneous Poisson process for the formation of a puff, the subsequent nucleation process (one to many open clusters) is very fast compared to a typical ISI. This time scale separation is the key assumption behind the generic model. By incorporation of a negative feedback it leads to ISI distributions close to experimental data.

Hierarchic stochastic modelling uses non-exponential transition times to reflect microscopic state changes occurring during sojourns in observable states. Remarkably, such non-exponential transition times have already been measured or computed for other cellular systems like gene expression [37,38], autocatalytic reactions [46,47] or signal transduction cascades [48,49]. Using non-exponentially distributed transition times instead of kinetic constants can also be interpreted as dynamic modelling with stochastic delay – the delay arises from the underlying microscopic dynamics (see Fig. 1). Biological systems with stochastic delay were previously investigated [50–53], but these studies lack the possibility to implement molecular interactions on the lumped state level, and have not been generalised to hierarchical networks. Hierarchic stochastic modelling can also be related to published methods which exploit the hierarchic structure of networks [54–56]. Similar to our approach, such methods reduce the state space based on emergent behaviour of biomolecules arising by the dynamic hierarchy from molecular interactions to cellular properties [19,57,58]. However, to our knowledge, these studies have been limited to a reduction of the state space or inference of system parameters in deterministic systems so far. Since complex stochastic networks are frequently observed in cell biology [1,2], we expect that our approach has a wide range of potential applications. On the one hand, pure Markov modelling means that one has to take into account all microscopic variables to make an exact description of the system. This is impossible for most, if not all, processes analysed in cell biology due to the large numbers of system states and because microscopic state transitions can rarely be measured in vivo. On the other hand, the assumption of Gaussian white noise, which is often used in approximation techniques, is questionable for many biological systems because the number of independent identical processes is too small for application of the law of large numbers. Therefore, hierarchic stochastic modelling brings theoretical concepts closer to experimental data.

The application to intracellular Ca\textsuperscript{2+} dynamics presented here makes an important step forward in the development of a mechanistic but comprehensive mathematical description of the cellular release. Experimental data is the nucleation of a wave due to the cellular spikes. The stronger the release in such an area is, the spiking, if a cell has a preferred nucleation area initiating most of the cellular release spike. Typically, a cellular spike will consist of more than 4 open clusters.
process. We are now in a position to characterise the stochastic process in the physiologic parameter regime. A key finding is that the time scale separation between single cluster pulls and subsequent openings is very strong. And that despite the non-Markovian nucleation process, a coefficient of variation smaller than one can not be realised without a negative feedback. These results pave the road to the generic description in the form of a time-dependent Poisson process, which can be sampled by standard methods. Our derivation shows how the phenomenological parameters of the generic model depend on cellular parameters. The generic model can therefore be used to generate Ca^{2+} spikes with correct statistics in models of larger signalling cascades in future research.

Materials and Methods

The Conditioned Waiting Times \( \Psi_d(t) \)

In the mechanistic stochastic Ca^{2+} model recently published by two of us [17], we not only lumped the microscopic states of individual \( \text{IP}_{3}: \text{Rs} \) but the whole ensemble of Ca^{2+} channels forming a cluster. Therefore, our effective reduced state space just contains the time dependent cluster configuration, i.e. the number of open and closed Ca^{2+} clusters at a specific time point. In the theoretical framework of semi-Markov processes (see Text S1 for details) transitions within that state space can be described by the probability given by \( p_{ij}\Psi_d(t) \). The first factor describes the transition probability of an ordinary discrete time Markov chain for the transition \( i \rightarrow j \), the latter gives the possibly non-exponential waiting time distribution. The \( \Psi_d(t) \) are not true probability densities for more than two possible state transitions, as they are not normalised individually, but rather by the condition \( \sum_0^\infty \Psi_d(t)dt = \sum_{i,j} p_{ij}=1 \). Therefore the term conditioned waiting times (or conditioned dwell times) is appropriate. The waiting times can be calculated from a mathematical model or measured directly (see Section Results). Exact definitions of the \( \Psi_d(t) \) and for the considered semi-Markovian stochastic process are given in Text S1. Apart from the analytical approach given below, the \( \Psi_d(t) \) can also be used very efficiently in stochastic simulations [17].

Solution of the Non-Markovian Master Equation

State-transition with non-exponential waiting times are not covered by Eq. 1, but rather by a generalised (non-Markovian) master equation [17,59].

\[
\frac{\partial}{\partial t} P_{ij}(t) = \sum_{k=1}^{N} \left[ R_{ik}(t) - P_{ik}(t) \right], \tag{11}
\]

with the probability fluxes given by [59]

\[
P_{ij}(t) = \int_0^t \Psi_d(t-x) \sum_{k=1}^{N} I_{ik}(x)dx + f_{ij}(t). \tag{12}
\]

Note that the probability fluxes \( f_{ij}(t) \) can no longer be expressed in terms of the \( P_{ij}(t) \) like in Eq. 1, but are solutions of a Voltera integral equation determined by the conditioned waiting times \( \Psi_d(t) \) and appropriate initial functions \( f_{ij}(0) \). The initial functions are related to the initial probabilities \( P_{ij}(0) \) in Eq. 11, e.g. \( P_{i1}(0)=1, P_{i0}(0)=0 \) sets \( f_{i1}(0) = \Psi_d(t) \) and \( f_{i0}(0)=0 \) for all \( i \neq j \) and all \( I \).

The form of Eq. 11 suggests a solution based on the Laplace transform \( \mathcal{L}\left[f(t)\right] = \int_0^\infty e^{-st}f(t)dt = F(s) \) (see also [60]), because the convolution theorem implies.

\[
\mathcal{L}\left[P_{ij}(s)\right] = \Psi_d(s) \sum_{k=1}^{N} \left[ R_{ik}(s) + f_{ij}(s) \right]. \tag{13}
\]

This constitutes a linear system of equations for the probability fluxes in Laplace space. We obtain the stationary occupancy probabilities as.

\[
\lim_{t \to \infty} P_{ij}(t) = \lim_{s \to 0} \left\{ \sum_{k=1}^{N} \left[ R_{ik}(s) - \tilde{P}_{ij}(s) \right] \right\}, \tag{14}
\]

where the limits in Eq. 14 are equal due to the final value theorem of the Laplace transform.

Certain dynamical system properties can be characterised by statistical moments of the density function of the first passage time (FPT) \( \tilde{F}(t) \) giving the probability for the first visit of state \( i \) when starting at state \( j \). The moments of \( \tilde{F}(t) \) obey [11]:

\[
\langle t^n \rangle = (1)^n \sum_{i,j=0}^{\infty} \mathcal{L}\left[ F_i(s)\right] \delta_{ij} , \tag{15}
\]

where the Laplace transformed FPT density can be found by the formula \( \mathcal{L}\left[ F_i(s)\right] = \mathcal{L}\left[ F_i(s)/F_j(s)\right] \) [11]. By using the property \( \mathcal{L}\left[ f(t)\right] = f(s) - f(0) \), the next step is to apply the Laplace transform to Eq. 11. This yields

\[
s \tilde{P}_{ij}(s) - \delta_{ij} = \sum_{l=1}^{N} \left[ R_{il}(s) - \tilde{P}_{il}(s) \right], \tag{16}
\]

stating that \( P(i,0|j,0)=1 \) for \( i=j \) and zero otherwise. From this, we find

\[
\tilde{F}_i(s) = \frac{\sum_{l=1}^{N} \left[ R_{il}(s) - \tilde{P}_{il}(s) \right]}{\sum_{l=1}^{N} \left[ R_{ij}(s) - \tilde{P}_{ij}(s) \right] + 1}. \tag{17}
\]

This expression is particularly simple in the case of a linear chain with \( N=K+1 \) distinct states \( X=\{0,1,\ldots,K\} \), where we have to compute the \( 2K \) probability fluxes \( I_{0,1},I_{1,0},\ldots, I_{K-1,K},I_{K,K-1}. \) Suppose the system is in state \( 0 \) at \( t=0 \) and we want to calculate the moments of the FPT distribution to reach the highest state \( K \) using Eq. 15 derived above. In the case of a linear chain, we arrive at

\[
\tilde{F}_K(s) = \frac{I_{0,K-1}}{I_{0,K-1} + I_{K-1,K}}, \tag{18}
\]

The conditioning on either state \( 0 \) or state \( K \) at \( t=0 \) leads effectively to two distinct non-homogeneous linear systems of equations for the fluxes. They differ only with respect to the initial functions, namely \( f_{01} = \Psi_{01} \) and \( f_{00} = \Psi_{00} \). Therefore, in the case of \( N=3 \) we obtain Eq. 2 (see Text S1 for details).
Model Calculations
Details of the mechanistic Ca\(^{2+}\) model are given in Text S1. We used Wolfram’s Mathematica 8 for linear algebra and a homemade simulation tool [17] for stochastic simulations.

Supporting Information

Figure S1 Comparison of the generic model with the exact analytic results provided by the hierarchic stochastic model. Shown are the results for the realistic channel closing rate \(\gamma\) (black lines) and two examples with lower time scale separation. (A) The approximation error grows linear with the average number of failed puffs, given by \(1/C_{\text{A}}\). (B) The relative approximation error is smaller than 10% for the realistic closing rate.

Table S1 Parameter values for the De Young-Keizer model. The De Young-Keizer model with the parameters in this table is used to compute the opening transition times \(\psi_{\text{o}}\).

Table S2 Analytical approximation of the GE distribution parameters. The values in this table result from nonlinear fitting to Eqs. 4–5 (see Fig. 2) and are used for the dependencies of the GE distribution parameters \(z\) and \(\lambda\) on cellular parameters \(N_{\text{A}}\) and \([\text{IP}_3]\) and Ca\(^{2+}\) (see Text S1 for details). In each row, only one parameter is varied, and other parameters are kept constant at \(N_{\text{A}} = 5\), \([\text{IP}_3] = 1 \mu\text{M}\), and \([\text{Ca}^{2+}]\) given by Eq. 41 in Text S1. Every system state is described by the number of open clusters (o. cl) and has its own set of parameters.

References

1. Raj A, van Oudenaarden A (2008) Nature, nurture, or chance: stochastic gene expression and its consequences. Cell 135: 216–26.
2. Balazsi G, van Oudenaarden A, Collins JJ (2011) Cellular decision making and biological noise: from microbes to mammals. Cell 144: 910–23.
3. Abschuleier JS, Wu LF (2010) Cellular heterogeneity: do differences make a difference? Cell 141: 559–63.
4. Alon U, Surette MG, Barkai N, Leibler S (1999) Robustness in bacterial chemotaxis. Nature 397: 168–71.
5. Elowitz MB, Levine AJ, Siggia ED, Swain PS (2002) Stochastic gene expression in a single cell. Science 297: 1183–6.
6. Skupin A, Kettemann H, Winkler U, Wartenberg M, Sauer H, et al. (2008) How does intracellular Ca\(^{2+}\) oscillate: by chance or by the clock? Biophys J 94: 2404–11.
7. Feinerman O, Veiga J, Dorfman JR, Germain RN, Altan-Bonnet G (2008) Variability and robustness in T cell activation from regulated heterogeneity in protein levels. Science 321: 1081–4.
8. Frische-Geuehter R, Wizel F, Sieber A, Herr R, Schmid N, et al. (2011) Strong negative feedback from Erk to Raf confers robustness to MAPK signalling. Mol Syst Biol 7: 489.
9. Cagatay T, Turcotte M, Elowitz MB, Garcia-Ojado J, Suel GM (2009) Architecture-dependent noise discriminates functionally analogous differentiation circuits. Cell 139: 512–22.
10. Mariani L, Schutz EG, Leshchuk MH, Helmstetter C, Raabeuch A, et al. (2010) Short-term memory in gene induction reveals the regulatory principle behind stochastic IL-4 expression. Mol Syst Biol 6: 359.
11. Van Kampen NG (2002) Stochastic Processes in Physics and Chemistry. Amsterdam: Elsevier Science B.V.
12. Tanase-Nicola S, ten Wolde P (2008) Regulatory control and the costs and benefits of biochemical noise. PLoS Comput Biol 4: e1000125.
13. Gillespie DT (1977) Exact stochastic simulation of coupled chemical reactions. J Phys Chem 81: 2340–2361.
14. Falcke M (2004) Reading the patterns in living cells - the physics of Ca\(^{2+}\) dynamics in larvalcocytes. Biophys J 95: 2193–202.
15. Berridge MJ, Lipp P, Bootman MD (2000) The versatility and universality of calcium signalling. Nat Rev Mol Cell Biol 1: 11–21.
16. Thul R, Falcke M (2008) Stability of membrane bound reactions. Phys Rev Lett 93: 180603.
17. Thurley K, Falcke M (2011) Derivation of Ca\(^{2+}\) signals from puff properties reveals that pathway function is robust against cell variability but sensitive for control. Proc Natl Acad Sci U S A 108: 427–32.
18. Thurley K, Skupin A, Thul R, Falcke M (2012) Fundamental properties of Ca\(^{2+}\) signals. Biochim Biophys Acta 1820: 1185–94.
19. Skupin A, Thurley K (2012) Calcium Signaling: From Single Channels to Pathways. Adv Exp Med Biol 740: 531–51.
20. Marchant JS, Callamaras N, Parker I (1999) Initiation of IP\(_3\)-mediated Ca\(^{2+}\) waves in Xenopus oocytes. EMBO J 18: 5285–5299.
21. Tafur Ur R, Skupin A, Falcke M, Taylor CW (2009) Clustering of InsP\(_3\) receptors by InsP\(_3\) returns their regulation by InsP\(_3\) and Ca\(^{2+}\). Nature 458: 455–9.
22. Smith IF, Parker I (2009) Imaging the quantal substructure of single IP\(_3\)R channel activity during Ca\(^{2+}\) puffs in intact mammalian cells. Proc Natl Acad Sci U S A 106: 6404–9.
23. Bogorovavni I, Wiatra J, Ehrlich BE (1991) Bell-shaped calcium-response curves of Ins(1,4,5)P\(_3\) and calcium-gated channels from endoplasmic reticulum of cerebellum. Nature 351: 751–4.
24. Taylor CW, Tovey SC (2010) IP\(_3\) Receptors: Toward Understanding Their Activation. Cold Spring Harb Perspect Biol 2: a004010.
25. Onoza E, Meyer T (1998) Protein kinase C as a molecular machine for decoding calcium and diacylglycerol signals. Cell 95: 307–18.
26. Dupont G, Abou-Lovergne A, Combettes L (2008) Stochastic aspects of oscillatory Ca\(^{2+}\) dynamics in larvalocytes. Biophys J 95: 2193–202.
27. Thurley K, Smith IF, Tovey SC, Taylor CW, Parker I, et al. (2011) Timescales of IP\(_3\)-Evoked Ca\(^{2+}\) Spikes Emerge from Ca\(^{2+}\) Puffs Only at the Cellular Level. Biophys J 101: 2630–44.
28. Meyer T, Stryer L (1989) Molecular model for receptor-stimulated calcium spiking. Proc Natl Acad Sci U S A 86: 5051–5.
29. Schuster S, Marhl M, Hofer T (2002) Modelling of simple and complex calcium oscillations. From single-cell responses to intercellular signalling. Eur J Biochem 260: 1333–55.
30. Skupin A, Falcke M (2007) Statistical properties and information content of calcium oscillations. Genome Inform 18: 44–53.
31. Skupin A, Falcke M (2009) From puffs to global Ca\(^{2+}\) signals: how molecular properties shape global signals. Chaos 19: 037111.
32. Skupin A, Kettemann H, Falcke M (2010) Calcium signals driven by single channel noise. PLoS Comput Biol 6.
33. Sneyd J, Falcke M (2005) Models of the inositol trisphosphate receptor. Prog Biophys Mol Biol 89: 207–45.
34. De Young GW, Keizer J (1992) A single-pool inositol 1,4,5-trisphosphate-receptor-based model for agonist-stimulated oscillations in Ca\(^{2+}\) concentration. Proc Natl Acad Sci U S A 89: 9895–9.
35. Sneyd J, Falcke M, Dubour JF, Fox C (2004) A comparison of three models of the inositol trisphosphate receptor. Prog Biophys Mol Biol 85: 121–140.
36. Ueda M, Shihata T (2007) Stochastic signal processing and transduction in chemotactic response of eukaryotic cells. Biophys J 93: 11–20.
37. Harper CV, Finkenstadt B, Woodcock DJ, Friedrichsen S, Sengrini S, et al. (2011) Dynamic analysis of stochastic transcription cycles. PLoS Biol 9: e1000607.
38. Suter DM, Molina N, Gatfield D, Schneider K, Schüller U, et al. (2011) Mammalian genes are transcribed with widely different bursting kinetics. Science 332: 472–4.
39. Gupta RD, Kundu D (2007) Generalized exponential distribution: Existing results and some recent developments. J Statist Plann Inference 137.
40. Pal N, Bandyopadhyay S, Paria A (2007) Stochastic mathematical models for calcium dynamics in a single cell. J Theor Biol 243: 343–351.
41. Skupin A, Kettemann H, Falcke M (2010) Calcium signals driven by single channel noise. PLoS Comput Biol 6.
42.钩 McDonnell MB, Abbott D (2009) What is Stochastic Resonance? Definitions, Misconceptions, Debates, and Its Relevance to Biology. PLoS Comput Biol 5: e1000381.
43. Zhou, C., Kurths, J., Hu, B. (2001) Array-enhanced coherence resonance: nontrivial effects of heterogeneity and spatial independence of noise. Phys Rev Lett 87: 098101.
44. Marchant, J.S., Parker, I. (2001) Role of elementary Ca^{2+} puffs in generating repetitive Ca^{2+} oscillations. EMBO J 20: 65-76.
45. Dupont, G., Swijnen, S., Clair, G., Tordjmann, T., Combettes, L. (2000) Hierarchical organisation of calcium signals in hepatocytes: from experiments to models. Biochim Biophys Acta 1490: 134–132.
46. Delbrück, M. (1945) The burst size distribution in the growth of bacterial viruses (bacteriophages). J Bacteriol 50: 131-5.
47. Delbrück, M. (1940) Statistical Fluctuations in Autocatalytic Reactions. J Chem Phys 8: 120-124.
48. Bel, G., Minsky, B., Nemenman, I. (2010) The simplicity of completion time distributions for common complex biochemical processes. Phys Biol 7: 016003.
49. Lu, T., Shen, T., Zong, C., Hasty, J., Wolynes, P.G. (2006) Statistics of cellular signal transduction as a race to the nucleus by multiple random walkers in compartment/phosphorylation space. Proc Natl Acad Sci USA 103: 16752-7.
50. Schlicht, R., Winkler, G. (2008) A delay stochastic process with applications in molecular biology. J Math Biol 57: 613-648.
51. Barrio, M., Burrage, K., Leier, A., Tian, T. (2006) Oscillatory Regulation of Hes1: Discrete Stochastic Delay Modelling and Simulation. PLoS Comput Biol 2: 1017–1030.
52. Bratsun, D., Volfson, D., Tsimring, L.S., Hasty, J. (2005) Delay-induced stochastic oscillations in gene regulation. Proc Natl Acad Sci USA 102: 14593-8.
53. Josic, K., Lopez, J.M., Ott, W., Shiau, L., Bennett, M.R. (2011) Stochastic delay accelerates signaling in gene networks. PLoS Comput Biol 7: e1002264.
54. Borisov, N.M., Markovich, N.I., Hock, J.B., Khodolenko, B.N. (2003) Signaling through receptors and scaffolds: independent interactions reduce combinatorial complexity. Biophys J 89: 951-66.
55. Lade, S.J., Gross, T. (2012) Early warning signals for critical transitions: a generalized modelling approach. PLoS Comput Biol 8: e1002360.
56. Blochl, F., Wittmann, D.M., Tendel, E. (2011) Effective parameters determining the information flow in hierarchical biological systems. Bull Math Biol 73: 706-25.
57. Laughlin, R.B., Pines, D. (2000) The theory of everything. Proc Natl Acad Sci USA 97: 28-31.
58. Perelson, A. (2002) The riddle of “life,” a biologist’s critical view. Naturwissenschaften 96: 1-23.
59. Prager, T., Fälcke, M., Schimansky-Geier, L., Zaks, M.A. (2007) Non-Markovian approach to globally coupled excitable systems. Phys Rev E 76: 011118.
60. Thal, R., Fälcke, M. (2007) Waiting time distributions for clusters of complex molecules. Eur Phys Lett 79: 30003.