Modeling and Analysis of Calcium Signaling Events Leading to Long-Term Depression in Cerebellar Purkinje Cells

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ABSTRACT Modeling and simulation of the calcium signaling events that precede long-term depression of synaptic activity in cerebellar Purkinje cells are performed using the Virtual Cell biological modeling framework. It is found that the unusually high density and low sensitivity of inositol-1,4,5-trisphosphate receptors (IP₃Rs) are critical to the ability of the cell to generate and localize a calcium spike in a single dendritic spine. The results also demonstrate the model's capability to simulate the supralinear calcium spike observed experimentally during coincident activation of the parallel and climbing fibers. The sensitivity of the calcium spikes to certain biological and geometrical effects is investigated as well as the mechanisms that underlie the cell's ability to generate the supralinear spike. The sensitivity of calcium release rates from the IP₃Rs to calcium concentrations, as well as IP₃ concentrations, allows the calcium spike to form. The diffusion barrier caused by the small radius of the spine neck is shown to be important, as a threshold radius is observed above which a spike cannot be formed. Additionally, the calcium buffer capacity and diffusion rates from the spine are found to be important parameters in shaping the calcium spike.

INTRODUCTION

One cellular basis for learning is the phenomenon of synaptic plasticity that has been observed experimentally in neurons. Synaptic plasticity refers to a temporary or sustained depression or potentiation of the activity level of a synapse. An important form of synaptic plasticity related to motor-learning tasks such as the vestibular-ocular reflex, eye-blink conditioning, posture and locomotion adaptation, motor coordination, and hand/arm movement adaptation is observed in cerebellar Purkinje cells (see Fig. 1 for a micrograph of a Purkinje cell). This particular form of synaptic plasticity, known as long-term depression (LTD), is a lasting decrease in the activity of the synapses between spines on the Purkinje cell dendrites and axons of neighboring granule cells, often referred to as parallel fibers (PF).

It has been shown experimentally that LTD is induced by the repeated association of the PF and climbing fiber (CF) inputs (2). A Purkinje cell is normally in contact with a single CF. Even though a single Purkinje cell may have as many as 175,000 PF synapses (3), a single dendritic spine, generally, has a synapse with only a single PF. Although activation of a lone PF, therefore, normally results in events occurring in a single spine of a Purkinje dendrite, activation of the CF produces a delocalized depolarization of the Purkinje cell membrane. Activation of either a PF or CF results in signaling events involving ionic calcium (Ca²⁺). In the case of the CF, the resulting depolarization of the Purkinje cell opens voltage-sensitive calcium channels, allowing for Ca²⁺ entry into the cytosol from the extracellular space. Activation of the PF results in release of glutamate across the synapse that is then detected by metabotropic glutamate receptors (mGluR) on the neighboring Purkinje spine. The signaling pathway depicted in Fig. 2 is then activated resulting in the release of Ca²⁺ from the endoplasmic reticulum (ER) mediated by the inositol-1,4,5-trisphosphate receptors (IP₃Rs). Alpha-aminohydroxy-5-methyl-4-isoxazolepropionate receptors on the membrane also detect glutamate and respond by opening channels that allow for entry of Ca²⁺ into the cytosol from the extracellular space.

Ca²⁺ elevation has been shown to be required for LTD induction in Purkinje cells. For example, experiments using Ca²⁺ chelators, such as EGTA (4,5) or BAPTA (2) resulted in blockage of LTD. Additionally, activation of Ca²⁺ channels by current step-elicited membrane depolarization induced LTD when combined with PF stimulation (2,6). Experiments in which the IP₃R Ca²⁺ channels were inactivated (7) or otherwise not present (8) resulted in a lack of LTD, indicating that Ca²⁺ release from these internal stores is necessary for induction of LTD.

It has been found experimentally that coincident activation of the PF and CF inputs results in a supralinear increase in [Ca²⁺]. In other words, the change in [Ca²⁺] that is observed is significantly more than the sum of the Ca²⁺ responses obtained by exciting the PF and CF separately (3). It is hypothesized that this supralinear calcium response is the mechanism by which the cell detects the coincident activation of the PF and CF and is the first step in the mechanism leading to LTD. Under normal coincident activation conditions, these supralinear spikes may be confined to single spines.

The properties of the IP₃R involved in this Ca²⁺ signaling pathway are likely to play a key role in the generation of the
calcium release (CICR) (this is also observed in the Ryanodine receptors found in the Purkinje dendritic shafts, but not the spines themselves). Therefore, there is a positive feedback present in the system in which Ca\(^{2+}\) release stimulates further Ca\(^{2+}\) release. Note that there is an apparent threshold Ca\(^{2+}\) concentration above which an IP\(_3\)R becomes increasingly deactivated but at a slower timescale (9–12).

The IP\(_3\)R in Purkinje cells have certain, unique properties as compared to other neuronal or peripheral cells. Namely, the receptor density is significantly higher (13–16) and the sensitivity to IP\(_3\) is extraordinarily lower in vivo (17,18). It has been shown that the sensitivity of the Purkinje cell IP\(_3\)R is similar to those of other cells when analyzed in isolation, leading to the hypothesis that the reduced sensitivity is due to interference by an inhibiting protein found in Purkinje cells (19,20). The physiological significance of these properties in terms of Ca\(^{2+}\) release before the onset of LTD is unclear at this time. For example, would similar Ca\(^{2+}\) release profiles be observed if the cell had a significantly lower IP\(_3\)R density but a much higher sensitivity, i.e., the density and sensitivity common in other cells? It is possible that factors beyond those involved in the LTD-induction mechanism mandate these properties. Additionally, as is supported by the results of this work, it is likely that the CICR phenomenon at the IP\(_3\)R is the basis of the coincidence detection mechanism that triggers LTD. Therefore, it is important that the unique properties and influence of the IP\(_3\)R be correctly modeled and analyzed as part of a study of the Ca\(^{2+}\) dynamics that precede LTD.

When considering the origins of Ca\(^{2+}\) release patterns in Purkinje spines, the unique geometry of the spines must also be taken into account. It is hypothesized that the thin neck connecting a spine to its dendrite aids in localizing LTD to a particular spine by creating a significant diffusional resistance (21). This could aid in biochemically decoupling the spine from the parent dendrite (22–25), which is desirable since the Ca\(^{2+}\) signaling events can thus be made specific to a single synapse. In summary, the available data in the literature implies that the Purkinje cell’s ability to produce the supralinear behavior may rely on a variety of biochemical and geometrical effects.

Given the importance of Ca\(^{2+}\) signaling to the induction of the mechanisms leading to LTD, the objective of this work is to use mathematical models of a Purkinje cell that focus on the relevant Ca\(^{2+}\) signaling networks to investigate the significance of certain unique characteristics of the Purkinje cell, such as the sensitivity and density of the IP\(_3\)R and the geometry of the spines, in terms of LTD induction. The results of this work will aid in identifying those features of the cell that are most critical to the onset of LTD, including consideration of both biochemical and geometrical effects. The wide availability of experimental data on calcium dynamics in Purkinje cells makes such a modeling study feasible.

A recent modeling study of Purkinje spine Ca\(^{2+}\) signaling (26) contained analysis of a compartmental model of a single
spine that did not consider diffusion out of the spine into the adjacent dendritic shaft or the unique sensitivity and density of the IP₃R. These are key differences between the models proposed here and those in the other work. Indeed, it is these features of Ca²⁺ signaling in Purkinje cell spines that set it apart from other systems. As will be shown, without taking into account the unique in vivo sensitivity of the IP₃R, significant Ca²⁺ release would be produced at low [IP₃] in direct disagreement with available experimental data (e.g., (17)). This combination of biochemical and geometric features also assures that the signal is confined to the activated spine. Additionally, as shown experimentally by Noguchi et al. (25) in pyramidal neurons, diffusion through the spine neck is an important factor in spine Ca²⁺ signaling, suggesting that dynamic spine morphology can be a powerful modulator of synaptic plasticity. The modeling results shown here demonstrate that the spine neck diameter can behave as a sensitive switch for the Ca²⁺ signal.

In the next section, the models used in this work are introduced and the sources for all parameter values identified. In the subsequent sections, simulation results are presented for a one-dimensional spatial model, a compartmental model, and a two-dimensional spatial model. These results demonstrate the models’ ability to rationalize the high density and low sensitivity of the spine IP₃R, to generate the supralinear Ca²⁺ spike, and to elucidate the mechanisms underlying the supralinear behavior. In the last section, conclusions are drawn and future research directions discussed.

MODEL DEVELOPMENT

The models developed in this work are based on an assembly of components from our own previous work and those available in the literature relating to cellular calcium dynamics. Additionally, many parameter values are taken from a variety of pertinent references and are summarized in Table 1. The modeling and simulation are performed using the Virtual Cell (http://vcell.org) biological modeling framework (27,28). All models used in this work are publicly available in the Virtual Cell (http://vcell.org) biological modeling framework (27,28). In what follows, the models described in this article, log on to the Virtual Cell (free registration is required), and go to File → Open → BioModel. In the BioModel Database dialog, the two-dimensional and compartmental models can be found in the Shared Models directory under “hernjak.” The one-dimensional model is similarly available as a MathModel. In what follows, the features shared by all of the models are introduced and the sources for the parameter values discussed. A concise summary of all of the models is provided in the Appendix.

The following species are present in the models: Ca²⁺, IP₃, calbindin (CD28k), parvalbumin (PV), and magnesium (Mg²⁺). CD28k and PV are buffer species that bind Ca²⁺. PV also has a strong affinity for Mg²⁺, thus Mg²⁺ is explicitly included in the model to effectively diminish the Ca²⁺-buffering capacity of PV. The general form of the expression used to model the Ca²⁺ dynamics is given as

\[
\frac{\partial [Ca^{2+}]}{\partial t} = R_{\text{diffusion}} + R_{\text{buffering}} + R_{\text{channel}} + R_{\text{entry}}. \tag{1}
\]

Each of the terms on the right-hand side of Eq. 1 will be discussed individually in what follows. Because the methodology for modeling diffusion differs among the models developed in this work, discussion of the different forms of \( R_{\text{diffusion}} \) can be found in the corresponding sections below.

The rate of binding of Ca²⁺ to a buffering species \( X \) is modeled as

\[
R_{\text{buffering,}X} = -k_{\text{on,}X}[Ca^{2+}][X] + k_{\text{off,}X}[XB], \tag{2}
\]

where \( XB \) denotes the Ca²⁺-bound form of the buffer. CD28k is found to contain both medium- and high-affinity binding sites (29), accounted for in this model under the assumption that the sites bind independently of each other. The overall \( R_{\text{buffering}} \) term in Eq. 1 is the summation of the buffering rates for all of the buffers, each modeled as in Eq. 2. Parvalbumin and calbindin are the only endogenous buffers considered in this work. According to the experimental data of Schmidt et al. (30), the intracellular concentrations of other buffers (e.g., calmodulin) are considerably lower than those of parvalbumin and calbindin and provide small contributions to the overall Ca²⁺ dynamics in the system. Similarly, the role of nonprotein buffering species is also likely to be negligible, given the conditions explored in this work.

The behavior of the IP₃R is modeled using the formulation of Li and Rinzel (31). This model, which is a simplified version of the DeYoung-Keizer (32) model, also accounts for the CICR behavior that the receptor demonstrates. Although a number of other models for calcium release from the ER have been proposed (e.g., the models of (33–40)), the Li-Rinzel model is adequate for this work’s analysis as it represents the critical dynamics of the calcium release via the IP₃R in a compact form. The Li-Rinzel model for Ca²⁺ flux across the ER membrane (including the behavior of SERCA pumps at the ER and an allowance for Ca²⁺ leakage from the ER) is defined as

\[
R_{\text{channel}} = a \left( 1 - \frac{[Ca^{2+}]}{[Ca^{2+}]_{\text{ER}}} \right) \left\{ \frac{h[Ca^{2+}][IP₃]}{\left( [Ca^{2+}] + d_{0} \right) [IP₃] + d_{0}} \right\}^{3} - V_{\text{max}} \left[ \frac{[Ca^{2+}]}{[Ca^{2+}]_{\text{ER}}} \right]^{2} \left( 1 - \frac{[Ca^{2+}]}{[Ca^{2+}]_{\text{ER}}} \right) + L \left( 1 - \frac{[Ca^{2+}]}{[Ca^{2+}]_{\text{ER}}} \right), \tag{3}
\]

\[
\frac{\partial h}{\partial t} = \left( K_{1} - [Ca^{2+}] + K_{1}h \right) K_{2}.
\]

The variable \( h \) is the probability of an inhibition site on the receptor being unoccupied. For the purposes of this work, the most important parameters in the model are \( a \) and \( d_{0} \). The \( a \) parameter corresponds to the density of IP₃R in the system, with large values of \( a \) indicating a high density of IP₃R. The


TABLE 1 Parameters and initial conditions used in the simulations

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Value</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>r_s (r_f)</td>
<td>Spine radius</td>
<td>0.29 μm</td>
<td>(49)</td>
</tr>
<tr>
<td>r_n</td>
<td>Spine neck radius</td>
<td>0.1 μm</td>
<td>(49)</td>
</tr>
<tr>
<td>r_d</td>
<td>Dendrite radius</td>
<td>1 μm</td>
<td>(30)</td>
</tr>
<tr>
<td>l_s (l_f)</td>
<td>Spine neck length</td>
<td>0.66 μm</td>
<td>(49)</td>
</tr>
<tr>
<td>s</td>
<td>Linear spine density</td>
<td>14 μm⁻¹</td>
<td>(49)</td>
</tr>
<tr>
<td>l_a</td>
<td>Length of adjacent dendrite compartment</td>
<td>27.98 μm</td>
<td></td>
</tr>
<tr>
<td>l_{c3}</td>
<td>Distance between adjacent and distal dendrite compartments</td>
<td>5.63 μm</td>
<td></td>
</tr>
</tbody>
</table>

| D_{Ca} | Ca²⁺ diffusion coefficient | 223 μm² s⁻¹ | (51) |
| D_{IP3} | IP₃ diffusion coefficient | 283 μm² s⁻¹ | (51) |
| D_{PV} | PV diffusion coefficient | 43 μm² s⁻¹ | (30) |
| D_{CD28k} | CD28k diffusion coefficient | 28 μm² s⁻¹ | (30) |
| D_{CG} | CG diffusion coefficient | 15 μm² s⁻¹ | (52) |
| k_{on,CD28k,high} | Forward rate coefficient | 5.5 μM⁻¹ s⁻¹ | (29) |
| k_{on,CD28k,med} | Forward rate coefficient | 43.5 μM⁻¹ s⁻¹ | (29) |
| k_{off,CD28k,high} | Reverse rate coefficient | 2.5 s⁻¹ | (29) |
| k_{off,CD28k,med} | Reverse rate coefficient | 35.8 s⁻¹ | (29) |
| k_{on,IP3,Ca} | Forward rate coefficient | 107.0 μM⁻¹ s⁻¹ | (30) |
| k_{on,IP3,Mg} | Forward rate coefficient | 0.8 μM⁻¹ s⁻¹ | (30) |
| k_{off,IP3,Ca} | Reverse rate coefficient | 0.95 s⁻¹ | (53) |
| k_{off,IP3,Mg} | Reverse rate coefficient | 25.0 s⁻¹ | (53) |
| k_{on,CG} | Forward rate coefficient | 430.0 μM⁻¹ s⁻¹ | (30) |
| k_{off,CG} | Reverse rate coefficient | 140.0 s⁻¹ | (54) |
| a | IP₃-R Ca²⁺ release amplitude (abundance) | 21,000.0 μM s⁻¹ | 10× the value in (41) |
| [Ca²⁺]_{ER} | ER [Ca²⁺] | 400 μM (constant) | (55) |
| d_{Ca} | IP₃-R Ca²⁺ binding constant | 0.3 μM | (41) |
| d_{IP3} | IP₃-R IP₃ binding constant | 20.0 μM | 10× the value in (41) |
| V_{max} | Amplitude of SERCA pump intake | 3.75 μM s⁻¹ | (41) |
| k_{er} | Pump binding constant | 0.27 μM | (56) |
| L | Leak constant | 0.12 μM s⁻¹ | (41) |
| K₁ | Dissociation constant for IP₃-R | 0.2 μM | (41) |
| K₂ | Forward rate coefficient, Ca²⁺ binding to inactivating IP₃-R site | 2.7 μM⁻¹ s⁻¹ | (41) |
| τ₁ | Start time of CF Ca²⁺ influx | 0.1 s | (30) |
| τ₂ | End time of CF Ca²⁺ influx | 0.105 s | (30) |
| τ_{cb,1} | Magnitude of CF Ca²⁺ entry signal in the spine | 13.25 s⁻¹ | (30) |
| τ_{cb,2} | Magnitude of CF Ca²⁺ entry signal in the adjacent dendrite compartment | 6.25 s⁻¹ | (30) |
| [Ca²⁺]_{ex} | Extracellular [Ca²⁺] | 1.0 mM (constant) | (57) |
| f_p | IP₃ pulse magnitude | 80.0 μM s⁻¹ | Calculated |
| τ_f | Time between IP₃ pulses | 0.012 s | (10) |
| K_f | IP₃ pulse decay factor | 1.188 s⁻¹ | (41) |
| K_{krg} | IP₃ degradation rate | 0.14 s⁻¹ | (41) |
| P | Ca²⁺ pumping rate | 8.0 μM⁻¹ s⁻¹ | (46) |
| [Ca²⁺]_{T} | Threshold [Ca²⁺] | 0.2 μM | (46) |
| [Ca²⁺]_{lo} | Initial [Ca²⁺] | 0.045 μM | (58) |
| [IP₃]₀ | Initial [IP₃] | 0.16 μM | (41) |
| [PV]₀ | Total [PV] | 40 μM | (30) |
| [CD28k]₀ | Total [CD28k] | 40 μM | (30) |
| [CG]₀ | Total [CG] | 160 μM | (30) |
| [Mg²⁺] | [Mg²⁺] | 590 μM (constant) | (30) |

value of d_{IP3} controls the sensitivity of the IP₃-R to [IP₃], with increasing values of d_{IP3} indicating decreased sensitivity of the IP₃-R to [IP₃] changes. Physically, d_{IP3} is the dissociation constant for IP₃ binding to the channel. The d_{Ca} parameter performs the same task in determining the sensitivity of the CICR behavior to changes in [Ca²⁺]. Although most of the Li-Rinzel model parameters are taken to be the values obtained from modeling studies of other neurons (such as neuroblastoma cells (41)), the values for a and d_{IP3} were both increased to correspond to the increased density and decreased sensitivity of the IP₃-R found in Purkinje cells. Implicit in the model described by Eq. 3 is the assumption that the concentration of Ca²⁺ in the ER is constant. Although researchers have adapted the Li-Rinzel model to

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account for varying [Ca\(^{2+}\)]\(_{ER}\) (e.g., (42–44)), the assumption of constant [Ca\(^{2+}\)]\(_{ER}\) is appropriate for this work, since the timescales of the Ca\(^{2+}\) dynamics investigated here are very fast relative to the proposed timescales for Ca\(^{2+}\) dynamics in the ER and the magnitudes of the Ca\(^{2+}\) changes that are observed in the cytosol are more than an order-of-magnitude less than [Ca\(^{2+}\)]\(_{ER}\). Also note that the magnitude of the leakage term in Eq. 3 is normally very small relative to the other terms, and likely plays a negligible role; however, it is left in the equation for completeness.

To simulate the effect of depolarization of the Purkinje cell by the CF, expressions of the following type are added to the dendrite and spine compartments:

\[
R_{\text{entry}} = J_{\text{ch}}(t > \tau_1)(t < \tau_2)([\text{Ca}^{2+}]_{\text{ex}} - [\text{Ca}^{2+}]) \tag{4}
\]

(where \(\tau_1 < \tau_2\)). [Ca\(^{2+}\)]\(_{ex}\) is the concentration of Ca\(^{2+}\) in the extracellular space and the \(J_{\text{ch}}\) parameter controls the magnitude of the Ca\(^{2+}\) entry rate (see Table 1). The two conditional statements in Eq. 4 are equal to \(I\) when true and \(0\) when false. Therefore, this expression allows for a pulse of Ca\(^{2+}\) to flow between the extracellular space and the cytosol during the time period \(\tau_1 < t < \tau_2\). Experimental data showing [Ca\(^{2+}\)]\(_{ex}\) changes during CF activation in Purkinje cells (30) were used to find realistic \(J_{\text{ch}}\) values for the spine and adjacent dendrite regions.

The production of IP\(_3\) due to activation of the PF is modeled using techniques developed in a previous modeling study of neuroblastoma cells (41). Based on the experimental observations of Fink et al. ((41), Fig. 4), activation of mGluR is modeled as an exponentially decaying flux of IP\(_3\) from the membrane of a spine. Multiple activations produce multiple pulses of IP\(_3\). Activation of a PF does not trigger IP\(_3\) production in the dendrite shaft under normal conditions, but generates IP\(_3\) only in the spine. IP\(_3\) can, however, diffuse into the dendritic shaft through the spine neck, but this geometrical feature will be described in the next section. The IP\(_3\) pulses are modeled as

\[
\frac{\partial [\text{IP}_3]}{\partial t} = J_p \sum_{i=0}^{n-1} (t > (i \tau_3)) e^{-(t-i\tau_3)K_3} - K_{\text{deg}}([\text{IP}_3] - [\text{IP}_3]_{0}) + R_{\text{diffusion}}, \tag{5}
\]

where \(n\) is the number of pulses, \(\tau_3\) is the time between pulses, and \(J_p\) and \(K_3\) control the pulse magnitudes and decay times, respectively. Based on commonly adopted experimental protocols for induction of LTD (e.g., Wang et al. (3)), \(n\) was chosen as 12 with a time between pulses (\(\tau_3\)) of 0.012 s, corresponding to a frequency of \(~80\) Hz (10). The second term in Eq. 5 accounts for the degradation of IP\(_3\) to IP\(_2\) and IP\(_4\) where \(K_{\text{deg}}\) is the degradation rate and [IP\(_3\)]\(_{0}\) is the nominal concentration of IP\(_3\). The value for \(K_{\text{deg}}\) was obtained from experimental data included in a modeling study of neuroblastoma cells (41). It is possible that the value for \(K_{\text{deg}}\) in Purkinje cells may differ significantly from the neuroblastoma values, but data does not exist to support the use of other values of this parameter.

In regard to IP\(_3\) dynamics, an experimentally validated basis does not exist for the usage of expressions more detailed than Eq. 5 in modeling the IP\(_3\) response to PF activation. However, we have examined whether our system might be sensitive to certain unknown effects. For example, it has been shown that the rate of IP\(_3\) conversion to IP\(_4\) via 3-kinase is sensitive to [Ca\(^{2+}\)] (45). Using Eq. 3 from Dupont and Erneux (45) and their parameters, an estimate of the rate of degradation of IP\(_3\) via 3-kinase at the peak concentrations of IP\(_3\) and Ca\(^{2+}\) observed in this work is 0.49 \(\mu \text{M/s}\). Because the rate of diffusion of IP\(_3\) from the spine is on the order of \(10^{-2}–10^{-3} \mu \text{M/s}\), any degradation effect will be minimal. We have performed sensitivity analysis to examine this issue further (see Supplementary Material) and have confirmed that [Ca\(^{2+}\)] is only weakly sensitive to \(K_{\text{deg}}\) given the conditions explored in this work. Similar arguments demonstrate that any Ca\(^{2+}\)-sensitivity of the PLC isoform in Purkinje cells would have a negligible effect on the IP\(_3\) profile, particularly since the IP\(_3\) pulses required to induce the supralinear Ca\(^{2+}\) behavior are very localized and occur over short time-windows. Again, sensitivity analysis (included in the Supplementary Material) provides quantitative support for this argument.

**ONE-DIMENSIONAL SPATIAL MODEL**

The first model considered in this work is a one-dimensional spatial model of a dendritic segment of length 500 \(\mu\text{m}\), as shown in Fig. 3. The results discussed in this section will demonstrate, via thorough modeling of spatial effects, that the relevant Ca\(^{2+}\) dynamics are localized to the activated spine under normal conditions. The governing equation for diffusion and reaction for each species in the dendrite has the form of the standard reaction-diffusion partial differential equation (PDE),

\[
\frac{\partial [X]}{\partial t} = D_x \frac{\partial^2 [X]}{\partial z^2} + \sum R_i, \tag{6}
\]

where [X] represents the concentration of species \(X\) in the dendrite, \(z\) represents the length dimension, and \(R_i\) refers to the rates of the buffering reactions and other pertinent reactions as discussed in the previous section. Species diffusion constants \((D_x)\) were obtained from experimental data (30). The validity of this one-dimensional model depends on the assumption that the dimensions of the spines and the radius of the dendrite are small enough such that no spatial gradients exist in those directions. Thus, only diffusion along the length of the dendrite needs to be explicitly considered. The large length of the geometry was selected to make sure that boundary conditions have no effect on the solution. A standard Fick’s law expression is used to account for diffusion from a spine to the adjacent dendritic shaft region.
FIGURE 3 Visualization of the geometry used to simulate calcium dynamics in a dendritic spine. For the one-dimensional spatial model (see article), the diffusion along the dendrite \( J_{23} \) is modeled explicitly using the PDE in Eq. 6 and the dynamics in the spines is modeled using Eq. 7. For the compartmental model (see article), the first two of the following three compartments are each modeled by a set of ordinary differential equations: 1. The spine under consideration; 2. The region of the parent dendrite directly adjacent to the spine; and 3. The distal regions of the dendrite. Species concentrations in compartment 3 are held constant so that this compartment acts as a sink for material that enters compartment 2. Compartment 1 is modeled as a sphere with volume \( V_1 = 4/3 \pi r_1^3 \) where \( r_1 \) is the radius of the spine. Compartment 2 is modeled as a cylinder with volume \( V_2 = \pi r_2 l \) where \( r_2 \) is the radius of the dendrite and \( l \) is the length of the segment of the dendrite encompassing compartment 2. The fluxes, \( J_{ip} \), between compartments are modeled using Fick’s law as shown in Eq. 7.

through the diffusion barrier posed by the spine neck. Specifically, the change in concentration of species \( X \) in a spine due to flux between the spine and a point along the dendrite adjacent to the spine and the various flux rates discussed in the previous section is modeled using an expression such as

\[
\frac{d[X]_k}{dt} = \frac{D X A_n}{l_n V_s} ([X]_s - [X]_k) + \sum R_i,
\]

where \( l_n \) is the length of the spine neck, \( A_n \) is the cross-sectional area of the spine neck \( (A_n = \pi r_n^2) \), \( V_s \) is the volume of the spine \( (V_s = 4/3 \pi r_s^3) \), \([X]_s \) is the concentration of the species in the proximal dendrite, and \([X]_k \) is the concentration of species \( X \) in the spine. Production of IP\(_3\) due to PF activation (as in Eq. 5) is only simulated in the spine at the center of the dendrite so that the Ca\(^{2+}\) dynamics resulting from stimulation of a single spine can be studied. This model also accounts for Ca\(^{2+}\) extrusion through the plasma membrane using

\[
R_{\text{extrusion}} = \sigma P ([\text{Ca}^{2+}] - [\text{Ca}^{2+}]_T)/([\text{Ca}^{2+}]_T + [\text{Ca}^{2+}]_T),
\]

where \( \sigma \) is the surface/volume ratio of either the spine or the dendritic shaft, \( P \) is the rate of pumping, and \([\text{Ca}^{2+}]_T \) is a threshold Ca\(^{2+}\) concentration below which the pumps are inactive. Equation 8 was obtained from the work of Fink et al. (41) and is based on the experimental observations of Herrington et al. (46), which indicate that the pumping mechanisms behave linearly at moderate \([\text{Ca}^{2+}] \) above a threshold value. Because high values of \([\text{Ca}^{2+}] \) are reached only for very short time-intervals in this work, this approximation is sufficient. This approximation may tend to overestimate pumping rates at some values of \([\text{Ca}^{2+}] \) since pumps are known to saturate. Likewise, the effect of other species’ concentrations (e.g., Mg\(^{2+}\)) on the capacity of the pumps is ignored for similar reasons. The overall system of equations is solved by the Virtual Cell through use of a finite volume algorithm with computation points evenly distributed at 0.5-\( \mu \)m intervals along the dendrite.

Fig. 4 shows the IP\(_3\) profile in the activated spine region (\(-0.5 \mu \text{m} < z < 0.5 \mu \text{m}\)) as a function of time as well as a kymograph (a plot of the spatial concentration profile as a function of time) showing [IP\(_3\)] in neighboring spines distributed along a 50-\( \mu \text{m} \) central dendrite segment as a function of time. During PF activation, the level of IP\(_3\) in the activated spine region reaches a maximum of nearly 70 \( \mu \text{M} \). The kymograph shows that the pulses of IP\(_3\) affect only the [IP\(_3\)] in the spine region that is activated by the PF. Spines beyond that region do not show an appreciable accumulation of IP\(_3\), indicating that IP\(_3\) accumulates much more rapidly in the activated region than it can diffuse out into the adjacent dendrite and spines. Fig. 5 contains a kymograph showing the spatial distribution of IP\(_3\) in the dendrite during PF stimulation. The results show that IP\(_3\) disperses along the dendrite and never reaches a concentration >3 \( \mu \text{M} \). For comparison, Fig. 5 also includes the kymograph from Fig. 4 (i.e., spine [IP\(_3\)]) plotted with a color scale corresponding to that used to plot the dendrite [IP\(_3\)] data as well as timecourses.
showing spine and dendrite [IP$_3$] at 5 µm to the right of the activated spine. The similarities between the data in the two kymographs and the timecourses show that the neighboring spines reach similar [IP$_3$] as the adjacent sections of the dendrite, meaning that the diffusion barrier caused by the spine neck is not sufficient to block IP$_3$ entry at such slow time scales. Data obtained from the one-dimensional model.

The results shown in the kymographs demonstrate that the particular geometry of the dendritic spines aids in compart-

FIGURE 5  

**Top** Kymograph showing the spatial distribution of IP$_3$ in the dendritic shaft during PF activation of the center spine. **(Middle)** The kymograph from Fig. 4 is shown again but with the color scaling set to correspond to that of the top figure (hence the color saturation for the center spine data) for comparison. **(Bottom)** Comparison of the [IP$_3$] time courses obtained in the spine and dendrite segment at 5 µm to the right of the center spine. The results show that the neighboring spines reach similar [IP$_3$] as the adjacent sections of the dendrite, meaning that the diffusion barrier caused by the spine neck is not sufficient to block IP$_3$ entry at such slow time scales. Data obtained from the one-dimensional model.

FIGURE 6  

**Top** Kymograph showing the spatial distribution of Ca$^{2+}$ in the spines during PF activation of the center spine. **(Middle)** Kymograph showing the spatial distribution of Ca$^{2+}$ in the spine during coincident PF and CF activation. Ca$^{2+}$ spikes rapidly in the center spine and then quickly returns to steady-state values, as shown in the timecourse of [Ca$^{2+}$] in the center spine obtained during coincident activation of the PF and CF **(bottom)**. The kymographs demonstrate that Ca$^{2+}$ does not appreciably spread to other spines during PF or coincident PF and CF activation. Data obtained from the one-dimensional model.

central activated spine only amounts to 10 nM. Also shown in Fig. 6 is a spine [Ca$^{2+}$] kymograph obtained during coincident activation of the PF and CF. Although the concentration in the activated spine reaches ~22 µM (the supralinear effect that will be discussed further in the following section), note that the neighboring spine regions are still generally unaffected.

The results shown in the kymographs demonstrate that the particular geometry of the dendritic spines aids in compart-
mentalizing signals due to the diffusional resistance created by the spine neck (21). By adjusting the neck radius parameter in the model, the magnitude of this effect can be quantified. Fig. 7 shows data that demonstrate the effect of the spine neck radius on the magnitude of the Ca\(^{2+}\) spike generated by coincident activation of the PF and CF. The data show that as the neck radius is increased, the magnitude of the Ca\(^{2+}\) spike decreases. This is in line with intuition, since increasing the neck radius should decrease the resistance to diffusion for material leaving the spine (21,47). Interestingly, when the spine neck is increased by >30%, an apparent threshold is reached, since the Ca\(^{2+}\) spike is quickly extinguished. This switch behavior demonstrates the strongly nonlinear behavior of the model in the parameter space necessary to observe the supralinear Ca\(^{2+}\) spike. These results suggest how the dynamic morphology that has been observed in two-photon recordings of spines (21,47) could serve as an important mediator of LTD.

Because this one-dimensional model demonstrates that the important events under consideration in this work are restricted to a single spine, we decided to develop a simpler compartmental model that would capture the key process of diffusion down the dendrite without solving PDEs. This model, described in the next section, facilitates a detailed analysis of the interplay of biochemical, electrophysiological, and geometrical parameters in shaping the spine Ca\(^{2+}\) signal.

**COMPARTMENTAL MODEL**

In this section, simulation results are presented that demonstrate the ability of a compartmental model to generate a supralinear Ca\(^{2+}\) spike given physically realistic Ca\(^{2+}\) signals originating from the PF and CF inputs. Additional simulation results are then presented that demonstrate the sensitivity of the system to certain biochemical features. Results are also shown that elucidate the mechanisms leading to the supralinear Ca\(^{2+}\) spike. In total, the results demonstrate that a wide range of biochemical and geometrical effects act in concert to allow the system to generate the supralinear Ca\(^{2+}\) signal that triggers the LTD process.

The model used in this section focuses on events that occur in a single spine and in the region of the parent dendrite directly adjacent to the spine. The modeling approach is summarized in Fig. 3. As seen in the figure, the model treats the region as consisting of three well-mixed compartments: the spine under consideration, the region of the parent dendrite directly adjacent to the spine, and a compartment representing the combined regions found at the distant ends of the dendrite. Material is allowed to diffuse from one region to the next as depicted in the diagram in Fig. 3. Species’ concentrations in the distal dendrite regions are held constant, implying that the distal regions are sufficiently far from the proximal dendrite that they act as a sink for material coming from the spine. This approximation is justified by the results of our model of a 500-μm length of dendrite, as described in the previous section.

Flux from one compartment to the next is modeled assuming diffusion occurs according to Fick’s law, as represented in Eq. 7. To determine the effective distance from the proximal dendrite to the distal dendrite (\(l_{23}\)), the one-dimensional PDE model of diffusion along a dendrite was used to identify the value of the length parameter that yields the best agreement between the compartmental model and the PDE model. A key benefit of the use of the ODE model is the significant reduction in model complexity and computational cost with only a small loss in fidelity. Further details about this model may be found in the Appendix.

In all simulations, 160 μM Calcium Green-1 (CG) is included in the model as an indicator dye to provide better comparison of the results to experimental data. In the analysis, the indicator is treated as a buffering species and, in most cases, free [Ca\(^{2+}\)] is reported. Using the Virtual Cell

**FIGURE 7** *(Top)* The percent change in Ca\(^{2+}\) spike magnitude during coincident PF and CF activation as a function of the percent change in the radius of the spine neck. *(Bottom)* Several Ca\(^{2+}\) timecourses obtained during coincident activation with varying neck radii. As seen in the figure on the left, neck radii increased by 30% or more prevent a substantial Ca\(^{2+}\) spike from being formed. Data obtained from the one-dimensional model.
modeling framework, the relative fluorescence change, $\Delta F/F$, can also be calculated based on the fluorescence properties of CG versus CG bound to Ca$^{2+}$. In any event, it is important to keep the indicator species in the model because, due to its buffering capability, it may have an effect on the magnitude of the Ca$^{2+}$ signals observed (41).

**Inducing the supralinear Ca$^{2+}$ signal**

Fig. 8a shows the Ca$^{2+}$ profiles in the spine and adjacent dendrite compartments during the simulated CF activation with no PF stimulation. These results were obtained by adjusting the $J_{ch}$ parameters to match the experimental data of Schmidt et al. (30). During the simulated opening of the voltage-sensitive calcium channels, [Ca$^{2+}$] rapidly spikes in both the spine and adjacent dendrite with a larger magnitude spike in the spine. Decay rates are comparable in both compartments. Note that the magnitudes of both spikes are in the submicromolar concentration range.

Fig. 8b shows the [IP$_3$] and [Ca$^{2+}$] profiles in the spine during the simulated PF stimulation with no activation of the CF. The cumulative effect of the 12 pulses of IP$_3$ that simulate the PF firing activity on [IP$_3$] in the spine cytosol yields an increase on the order of 70 μM, similar to what is observed for the one-dimensional model. Consistent with IP$_3$ flash photolysis experiments discussed in the literature (17), two orders-of-magnitude higher [IP$_3$] is required to stimulate these levels of Ca$^{2+}$ release from the IP$_3$R as compared to that which is observed in other cells. The model demonstrates that such high levels of IP$_3$ can be attained because of the high surface/volume ratio of the spine. A pulse train of 12 stimuli identical to those used in our earlier neuroblastoma cell model (41) is sufficient to produce the requisite IP$_3$ accumulation in the small spine volume. The rise in [Ca$^{2+}$] yielded by the model is on the order of 0.02 μM, followed by a decay to steady-state levels. The change in fluorescence, based on the fraction of CG sites occupied, is also shown. Data obtained from the compartmental model.

Fig. 9 is a plot of [Ca$^{2+}$] in the spine during coincident activation of the PF and CF inputs. A plot showing the relative timing of the individual Ca$^{2+}$ signals is also included in the figure. This relative timing was found by trial and error to be optimal in the sense that if the timing between inputs strayed too far from the selected timing, the large-magnitude supralinear spike was not observed. The effect of the relative

![FIGURE 8](image)

**FIGURE 8** (a) Simulated Ca$^{2+}$ signals in the spine (top) and proximal dendrite (bottom) cytosol during CF activation. Data obtained from the compartmental model. (b) Simulated IP$_3$ (top) and Ca$^{2+}$ (middle) signals in the spine cytosol during activation of the PF with 12 stimuli at 0.012 s intervals. The percent change in fluorescence (bottom) based on the fraction of bound CG sites is also shown. Data obtained from the compartmental model.

![FIGURE 9](image)

**FIGURE 9** Relative timings of the individual Ca$^{2+}$ signals in the spine cytosol (top) and the resulting supralinear Ca$^{2+}$ spike in the spine cytosol during coincident activation (bottom). Data obtained from the compartmental model.
timing of the inputs on the supralinear behavior will be discussed in more detail in the following subsection. The \([\text{Ca}^{2+}]\) results in Fig. 9 clearly show the ability of the model to demonstrate the critical supralinear behavior of the Purkinje cell \(\text{Ca}^{2+}\)-signaling network. Note that the peak magnitude is on the order of 10 \(\mu\text{M}\), as compared to the individual peak magnitudes in Fig. 8 that are orders-of-magnitude lower. This shows that the nonlinearity inherent in the model causes the calcium signal resulting from coincident activation of the two inputs to be greater than the linear sum of the calcium signals caused by the individual inputs. This is consistent with experimental evidence (e.g., Wang et al. (3)). Additionally, the results show an IP3-sensitive delay before the supralinear rise in \([\text{Ca}^{2+}]\), which is also consistent with experimental data (see Supplementary Material).

As will be discussed in more detail, \(\text{Ca}^{2+}\) buffers play an important role in the generation of the \(\text{Ca}^{2+}\) spike. Included in the list of buffers is the fluorescent indicator dye CG. This dye was added to the model to allow for an accurate comparison to experimental data on CF activation. Because CG is a high-affinity dye \(\left(K_D = 325 \text{ nM}\right)\), it is possible that its presence is reducing the magnitude of the supralinear spike by binding large amounts of free \(\text{Ca}^{2+}\) and reducing the CICR effect. In the experimental work of Wang et al. (3), the effect of coincident activation of the PF and CF inputs when the indicator dye is Magnesium Green \(\left(K_D = 19 \text{ \mu M}\right)\) is investigated yielding similar results to those seen in this simulation work.

**Sensitivity of the supralinear behavior**

The results in this section will show that the ability of the system to demonstrate the strongly nonlinear behavior evidenced by the supralinear \(\text{Ca}^{2+}\) spike is strongly dependent on many features of the system, including biochemical effects and geometrical effects (i.e., spine geometry, as discussed previously). Recall that the IP3R in Purkinje cells are more abundant and less sensitive to IP3 than those found in other neurons or peripheral cells. Our earlier neuroblastoma cell model, for example, produced robust \(\text{Ca}^{2+}\) release with \(a\) and \(d_{IP3}\) each an order-of-magnitude lower than the values used thus far in this work. By adjusting the parameters \(a\) (abundance of IP3R) and \(d_{IP3}\) (sensitivity of IP3R to IP3) found in Eq. 3, the effect of adjusting the IP3R abundance and sensitivity can be investigated in terms of the ability of the system to generate the \(\text{Ca}^{2+}\) spike that induces LTD. In Fig. 10 \(a\), the results of a simulation involving coincident activation of the PF and CF are shown for parameter values that represent an order-of-magnitude decrease in the IP3R abundance and an order-of-magnitude increase in the IP3 sensitivity. As can be seen in the figure, altering those parameters by an order of magnitude effectively extinguishes the supralinear \([\text{Ca}^{2+}]\) spike under these conditions. It is important to stress that \(a\) and \(d_{IP3}\) were increased in our model to conform to the experimental findings of extraordinarily high IP3R density (13) and low sensitivity to IP3 (17). They were not adjusted simply to fit the desired \(\text{Ca}^{2+}\) signaling behavior.

Another interesting set of results is obtained by returning the IP3R abundance parameter, \(a\), to its high value but leaving the \(d_{IP3}\) parameter at the value corresponding to high IP3 sensitivity. Simulation results given this parameter set during coincident PF and CF activation are shown in Fig. 10 \(b\). At first glance, it appears that these conditions generate two

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**FIGURE 10**

(a) Spine cytosol \(\text{Ca}^{2+}\) signal during coincident activation when the abundance of IP3R is decreased by a factor of 10 and the sensitivity \((d_{IP3})\) is increased by a factor of 10. The supralinear behavior is negligible under these conditions. Data obtained from the compartmental model. (b) \(\text{Ca}^{2+}\) signals in the spine and proximal dendrite cytosol during coincident activation. The top plot shows the responses given the normal Purkinje IP3R sensitivity, whereas the bottom plot shows the responses with the sensitivity to IP3 \((d_{IP3})\) increased by a factor of 10. When the sensitivity is increased, a secondary \(\text{Ca}^{2+}\) spike is observed in the dendrite that diffuses into the spine. This effect may prevent localization of LTD to the intended spine. Data obtained from the compartmental model.
Ca$^{2+}$ spikes in the spine: a large magnitude spike followed by a smaller spike ~0.5 s later. In actuality, a secondary Ca$^{2+}$ spike is being generated in the proximal dendrite compartment which then diffuses into the spine compartment. Thus, the second spine Ca$^{2+}$ spike is effectively an echo of the first and is, likewise, manifest in all the proximal spines emanating from this region of the dendrite. This is because spine density is $\sim 14 \mu m^{-1}$ (49). These results suggest that the low sensitivity of Purkinje cell IP$_3$R aids in the localization of the Ca$^{2+}$ spike to only the activated spine by preventing Ca$^{2+}$ spiking in the dendrite. These results attest to the importance of the unique IP$_3$R sensitivity and density. Without these features, the spike is either not formed (normal IP$_3$R density) or not localized (normal IP$_3$R sensitivity) to a single spine.

As alluded to in the previous subsection and shown by Wang et al. (3), the relative timing of the CF and PF inputs is critical to the formation of the supralinear Ca$^{2+}$ spike. Experimental results show that the largest magnitude Ca$^{2+}$ spikes are obtained by activating the CF briefly after the PF input train begins (3). As shown in Fig. 11, simulated results agree qualitatively with the experimental evidence in terms of both peak and integral fluorescence. To obtain results that could be correctly compared to the data of Wang et al. (3), the indicator dye was changed to Magnesium Green (K$_D$ = 19 $\mu$M at a concentration of 375 $\mu$M) and the IP$_3$ pulse train length was reduced to include only four pulses. The results show that the maximum [Ca$^{2+}$] is obtained when the CF is activated ~0.05 s after the PF input train begins. Given timings of this order of magnitude, the CF [Ca$^{2+}$] spike is occurring just before the time that [IP$_3$] is reaching its peak magnitude at the end of the PF pulse train, thus providing a large Ca$^{2+}$ influx when the maximum release of Ca$^{2+}$ from the ER is occurring. Given this relative timing, [Ca$^{2+}$] peaks at approximately the same time that [IP$_3$] peaks. Timing shifts in either direction result in a decrease in maximum [Ca$^{2+}$], with shifts in the CF activation time to earlier times resulting in a sharper decrease in maximum [Ca$^{2+}$]. These results show a strong agreement with the experimental results of Wang et al. (3), in terms of the time when the maximum Ca$^{2+}$ peak occurs. Also in qualitative agreement with the experimental data, the results in Fig. 11 show that the integrated Ca$^{2+}$ data trend is more symmetric than the peak Ca$^{2+}$ data trend. Analysis of the results suggests that proper

![Figure 11](image-url)

**FIGURE 11** (a) Maximum spine [Ca$^{2+}$] observed during coincident activation as a function of the relative timing of the PF and CF inputs reported in terms of fluorescence. Timing is calculated as time of CF activation − time of initial PF activation. For comparison to data of Wang et al. (3), an IP$_3$ pulse train consisting of four pulses was used with Magnesium Green as an indicator dye. The results show that the maximum fluorescence is reached when the CF is activated ~0.05 s after the IP$_3$ input train begins. These results are qualitatively similar to the fluorescence results reported by Wang et al. (3) in Fig. 6 b of their article (shown in b, copyright permission granted by Nature) in terms of the time at which the maximum calcium is observed. Data obtained from the compartmental model. (c) The total integrated Ca$^{2+}$ signal (computed over the full transient period) also reaches its maximum value at a relative timing near 0.05 s, in agreement with the experimental data of Wang et al. as shown in d (copyright permission granted by Nature).
timing of CF activation relative to PF activation may stimulate the CICR phenomenon at the IP₃R to a degree that is optimal in terms of yielding maximum Ca²⁺ release given the fixed IP₃ profile, as is necessary for induction of LTD.

**Physical mechanisms underlying suprilinearity**

A key issue to address in the study of the Ca²⁺ spike that leads to LTD is the determination of the actual physical mechanism that leads to the supralinear behavior. Although the results in the previous subsection provide an understanding of when and under what conditions the supralinear spike forms, the question of how the spike forms still needs to be addressed. In this subsection, two underlying mechanisms that act in concert to generate the supralinear behavior are explored using the compartmental model.

The influx (CF-activated) contribution to the total Ca²⁺ in the spine is a linear function of the density of channels, the open timing, and the current per channel. These are all held constant in our models. However, the IP₃R Ca²⁺ release is intrinsically nonlinear as it is sensitive to the level of Ca²⁺ in the system, with higher levels of Ca²⁺ inducing increased Ca²⁺ release. This positive feedback is part of the mechanism that leads to the Ca²⁺ spike. As evidence, simulations were performed under coincident activation conditions but with the model parameter that controls the sensitivity of the IP₃R to Ca²⁺ levels, $d_{IP₃}$, increased to correspond to a lessened sensitivity to Ca²⁺. Given an order-of-magnitude increase in this parameter, the Ca²⁺ spike is extinguished. This result shows that the positive feedback in the CICR mechanism is necessary to generate the supralinear spike. In agreement with experimental evidence, the IP₃R with its concerted IP₃ and Ca²⁺ dependencies is necessary for the formation of the supralinear Ca²⁺ spike.

A second important aspect of the mechanism leading to the Ca²⁺ spike is the role played by Ca²⁺ buffers in the system. The Ca²⁺ buffers act as significant stores of bound Ca²⁺. Fig. 12a is a plot of the fraction of buffer sites occupied for each of the buffers as a function of time during a normal coincident activation simulation. The model results shown in Fig. 12a indicate that the buffers approach but do not reach saturation during the Ca²⁺ spike. Of course, as the buffers approach saturation, there must be a concomitant increase in free Ca²⁺ (Figs. 6, 9, and 12b, bottom). The rate of buffer diffusion from the spine may also be important, since the bound form of the buffers are free to diffuse from the spine to the dendrite and to be replaced by buffers with free sites, thus providing additional Ca²⁺-buffering capacity. A useful estimate of the characteristic time for diffusion is given by $L^2/D_X$, where $L$ is a characteristic diffusion length and $D_X$ is a diffusion coefficient. If $L$ is taken to be the length of the spine neck and $D_X$ is taken to be the fastest buffer diffusion coefficient in the model ($D_{PV} = 43 \, \mu m^2/s$), the fastest characteristic diffusion time is 0.010 s. This timescale is the same order of magnitude as the timing of the Ca²⁺ spike formation observed in the data obtained in this work. This implies that, in the best case, the rate of buffer diffusion may not be fast enough to replenish the spine with unbound buffers, thus aiding in the saturation of the buffers present in the spine. The results in Fig. 12a also suggest that the finite rate at which Ca²⁺ binds to the buffers may be slower than the rate of formation of the Ca²⁺ spike. This is particularly true in the case of PV, where the results suggest that Mg²⁺ must unbind first to free sites for Ca²⁺ to occupy since Mg²⁺ is preferentially bound. All of these effects suggest that the buffers are unable to bind all of the Ca²⁺ that is released into the cytosol thus resulting in the large, nonlinear increase in Ca²⁺ observed during coincident activation.

Another method for observing the role of the buffers in formation of the Ca²⁺ spike is by calculating the total Ca²⁺ in the spine (bound and free, not counting that which is in the stores in the ER or the extracellular space) as well as the proportion of total Ca²⁺ in the system that is free versus that

**FIGURE 12** (a) Fraction of calbindin (top) and parvalbumin (bottom) buffer sites occupied during coincident activation in the spine cytosol. Data obtained from the compartmental model. (b) Total spine Ca²⁺ (bound and free) during coincident activation and independent activation of the PF and CF. Fraction of spine Ca²⁺ that is bound during coincident and independent activation (bottom). Data obtained from the compartmental model.
which is bound to the buffers, including the CG indicator. As shown in Fig. 12 b, the total Ca$^{2+}$ in the spine during coincident activation is approximately three times the amount that is observed during CF activation and is orders-of-magnitude more than that which is observed during PF activation. Compare this result to the data in Fig. 9, which show that the change in cytosolic [Ca$^{2+}$] during coincident activation is ~10 times higher than the CF Ca$^{2+}$ signal. The difference lies in the proportion of free Ca$^{2+}$ versus bound Ca$^{2+}$. As seen in the lower plot in Fig. 12 b, during the comparably slow release of Ca$^{2+}$ from the ER during PF activation, there is no significant change in the proportion of bound versus free Ca$^{2+}$. During the fast influx of Ca$^{2+}$ observed during CF activation, a small increase in the proportion of free Ca$^{2+}$ is observed. During coincident PF-CF activation, a much larger proportion of free Ca$^{2+}$ is observed, indicating that a significant amount of Ca$^{2+}$ is not being bound by the buffers. Therefore, as the buffers approach saturation, the fraction of free Ca$^{2+}$ increases disproportionately, causing the spike in cytosolic [Ca$^{2+}$].

**TWO-DIMENSIONAL SPATIAL MODEL**

Although the results above demonstrate that the one-dimensional and compartmental models are sufficient to model the localized effects considered in this work, future research may consider effects that occur over longer length scales. This type of study will require a model that represents the geometry of the dendrite more accurately than the somewhat simplified one-dimensional model. In this section, a two-dimensional model of IP$_3$ diffusion in a segment of the dendrite is introduced and used to demonstrate its ability to yield results similar to those seen in the previous sections.

In the two-dimensional model, the PDE in Eq. 6 is solved across the geometry shown in Fig. 13. No approximations are made that would allow for the use of ODE expressions. Diffusion in spines is treated explicitly as in the dendritic shaft. The geometry is based on the high-magnification micrograph in Fig. 1. The Virtual Cell system allows electronic images such as this to be used in defining system geometries. As can be seen in the figure, the geometry used in this model can account for spines on either side of the dendritic shaft as opposed to the one-dimensional model that can only model a single spine at each point along the shaft. Because the one-dimensional model represented a 500-μm segment of dendrite where the boundaries could be safely assumed to have reached basal IP$_3$ concentrations at all times, the one-dimensional model was used to establish the time-dependent boundary conditions at the open ends of the two-dimensional geometry.

Similar to the kymographs in Figs. 4 and 5, Fig. 13 shows the spread of IP$_3$ during PF activation of a single spine in the center of the bottom side of the dendrite as a function of time. In the activated spine, the [IP$_3$] reaches values of >40 μM, whereas the remainder of the dendrite never reaches [IP$_3$] >10–15 μM. Given the available experimental data and the results in the previous section, this is an insufficient level of IP$_3$ to activate Ca$^{2+}$ spikes that would lead to LTD in other regions of the dendrite. Note that the levels of [IP$_3$] observed in the dendrite and the other spines are slightly higher than those observed in the one-dimensional model results in Figs. 4 and 5. Similar to the conclusions drawn from the one-dimensional model results, the results again indicate that the spine is able to compartmentalize signals and prevent the spread of LTD. This further justifies the use of the reduced-order compartmental model for local analysis.

Although the two-dimensional model was found to produce the same types of results as the compartmental model for analysis of events caused by stimulation of a single spine, this type of modeling will become necessary when consideration is given to events occurring in multiple spines. In that case, the results of this work will be instrumental in identifying parameter values and simulation conditions to be used in extending the low-dimensional models to higher-dimensional cases. The two-dimensional model discussed and demonstrated in this section is an example of such a model.

**CONCLUSIONS**

The results obtained in this work demonstrate that a unique interplay of biochemical and morphological specializations...
contribute to the Purkinje cells’ ability to generate localized Ca^{2+} spikes in dendritic spines. These features range from the biochemical effects of buffer capacity and the unique properties of the IP_{3}R to the diffusional barrier imposed by the spine neck. The simulation results suggest that the system is designed in such a way that a lack of any of the features and mechanisms discussed in this work may prevent the onset of LTD. In addition, the results obtained in this work provide some understanding of the degree of sensitivity of the supralinear Ca^{2+} response to many of the important system features.

In the case of the unusual IP_{3}R sensitivity and abundance observed in Purkinje cells, the results indicate that these features are necessary for the system to induce and localize the supralinear Ca^{2+} spike that is a prerequisite for LTD. The system geometry is also a critical feature in the Ca^{2+} signaling network due to the role it plays in limiting diffusion from the spine for the majority of the system species. In the case of IP_{3}, its higher diffusion coefficient allows it to overcome the spine neck’s diffusion barrier during repeated PF activation. This is consistent with the lack of evidence of rapid degradation of IP_{3} after PF activation ends.

Although the modeling and simulation performed in this work focused on events occurring at the level of a single dendritic spine, the ability of the Virtual Cell to model spatially varying systems using PDE models will allow for further study of events occurring over larger length scales, such as branchlet-wide Ca^{2+} signaling, which is generally dominated by voltage-dependent Ca^{2+} entry. The capability to model and simulate PDE systems efficiently will permit the study of events occurring at multiple spines including identification of particular events and mechanisms that cause the spread of LTD effects from one spine to others. Additionally, the role of the unique geometry of the dendritic arbor could be investigated in this context. The parameters identified in this work using the simplified model and existing experimental data will be very useful in extending the model so that Ca^{2+} dynamics on the scale of branchlet segments, or even full branchlets, will be possible. For example, an obvious next step would be to consider simulated activation of multiple spines in a region of dendritic shaft. Neglecting to consider diffusion of material from the spines (i.e., consideration of only compartmental models) would have served to prevent the applicability of the models to problems such as these.

Additionally, proper spatial modeling is required as one begins to consider accurately the role of the mechanisms leading to the production of IP_{3}. For example, PIP_{2} is bound to the membrane and, therefore, its availability for reaction may be limited by lateral diffusion along the membrane. This would limit the Purkinje cell’s ability to produce the large amount of IP_{3} required to stimulate the IP_{3}R. Membrane diffusion capabilities will soon be available in the Virtual Cell, thus allowing this type of analysis to be performed in conjunction with a larger spatial modeling study including explicit consideration of PIP_{2} hydrolysis.

**APPENDIX**

This Appendix includes a summary of each of the models used in this work. All parameter values can be found in Table 1.

**One-dimensional model**

In this model, each species X (X = Ca^{2+}, IP_{3}, PV, PVB_c, PVB_m, CD28, CD28kB_{1}, CD28kB_{2}, and CD28kB_{1,2}) at time t and location z is represented by two concentrations: [X]_{d}(z,t) and [X]_{s}(z,t) in the dendrite and spine, respectively. These concentrations are governed by the equations

\[
\frac{\partial [X]_d}{\partial t} = D_X \left( \frac{\partial^2 [X]_d}{\partial z^2} + \frac{k_1}{l_{12}} ([X]_s - [X]_d) \right) + R_{X_d},
\]

\[
\frac{d[X]_s}{dt} = -\frac{D_X k_2}{l_{12}} ([X]_s - [X]_d) + R_{X_s},
\]

where \(D_X\) is the diffusion coefficient of the species X, \(l_{12}\) is the spine neck length, and the coefficients \(k_1\) and \(k_2\) are determined by the spine neck radius \(r_{ne}\), the spine radius \(r_s\), the dendrite radius \(r_d\), and the linear spine density \(\zeta_s\): \(k_1 = \zeta_s (r_d/r_s)^2\), \(k_2 = (3/r_s)(r_d/2r_s)^2\). The last terms in the governing equations (the source terms) are described below. They include rates of all processes, other than diffusion, that influence the corresponding concentration. \(R_{X_d}\) and \(R_{X_s}\) have identical structures: \(R_{X_d}\) can be obtained from \(R_{X_s}\) by simply replacing all dendrite parameters and concentrations by their spine counterparts (except in the case of IP_{3}, which is discussed below).

The source terms for calcium and IP_{3} are detailed in Eqs. 2–5 and Eq. 8 of the main text. The rate of calcium buffering includes the sum of terms of the type defined in Eq. 2 that correspond to binding to PV and CD28k. The surface/volume ratio \(\sigma\) in the rate of calcium extrusion, Eq. 8, is \(2r_d/3r_s\) for the dendrite and spine, respectively. To reflect activation of only one spine by PF, the first term in Eq. 5 must be zero everywhere except on the segment of length \(L = 1/s\) (note that this term is absent from the so-called dendrite equation),

\[
R_{IP_{3,d}} = \theta(L/2 - z)J_D \exp(-K_{df}) \sum_{i=0}^{n-1} \theta(t - i\tau_s) \exp(iK_{df}\tau_s)
\]

\[-K_{df}([IP_{3}]_d - [IP_{3}]_s) \cdot R_{IP_{3,s}} = -K_{df}([IP_{3}]_s - [IP_{3}]_d),\]

where \(\theta(x)\) is a step function,

\[\theta(x) = \begin{cases} 
1, & x \geq 0 \\
0, & x < 0 
\end{cases}\]

The “dendrite” source terms (the “spine” source terms are analogous) for calcium buffers are

\[
R_{PV_{,d}} = -(k_{a_{PV, Ca}}[Ca^{2+}]_d + k_{a_{PV, Mg}}[Mg^{2+}]_d)[PV]_d + k_{off_{PV, Ca}}[PVB_c]_d + k_{off_{PV, Mg}}[PVB_m]_d.
\]

\[
R_{PV_{,c,d}} = k_{a_{PV, Ca}}[Ca^{2+}]_d[PV]_d - k_{off_{PV, Ca}}[PVB_c]_d,
\]

\[
R_{PV_{,m,d}} = k_{a_{PV, Mg}}[Mg^{2+}]_d[PV]_d - k_{off_{PV, Mg}}[PVB_m]_d.
\]

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The definitions of the source terms, species \( R \) are modeled using ODEs of the form
\[
R_{CD2k,d} = -(k_{on,CD2k,high}[CD28k]_d + k_{on,CD2k,med}[CD28k]_d + k_{off,CD2k,high}[CD28kB2]_d + k_{off,CD2k,low}[CD28kB1]_d)[Ca^{2+}]_d
\]
\[
+ k_{off,CD2k,high}[CD28kB2]_d + k_{off,CD2k,low}[CD28kB1]_d + (k_{off,CD2k,high} + k_{off,CD2k,low})[CD28kB12]_d.
\]

The set of ODEs was solved numerically using the Runge-Kutta-Fehlberg method (50), a default ODE solver in the Virtual Cell.

### Two-dimensional model

The two-dimensional model simulates the dynamics of IP\(_3\) across the geometry depicted in Fig. 13. The governing equation,
\[
\frac{\partial [IP_3]}{\partial t} = D_{IP_3} \nabla^2 [IP_3] - K_{deg}([IP_3] - [IP_3]_0),
\]
is solved in the presence of the IP\(_3\) influx restricted to the membrane of the central spine. The corresponding flux density, \( J(t) \), reflects repetitive single spine excitations,
\[
J(t) = J_0 \exp(-K_i t) \sum_{i=0}^{n-1} \theta(t-i\tau_s) \exp(iK_i\tau_s),
\]
with the parameter values as in Fink et al. (41). The conditions at the domain boundaries were obtained from the corresponding one-dimensional model simulated with the identical parameter set. The simulations of the two-dimensional model were performed with the 10-\(\mu\)m time step on a 0.04-\(\mu\)m grid overlaying the 27.9 \(\times\) 6.5 \(\mu\)m geometry. The computational error is estimated to be <1%.

### SUPPLEMENTARY MATERIAL

An online supplement to this article can be found by visiting BJ Online at http://www.biophysj.org.

The Virtual Cell project is supported by the National Institutes of Health National Center for Research Resources grant No. RR13186.

### REFERENCES


