Calcium-Dependent Calcium Decay Explains STDP in a Dynamic Model of Hippocampal Synapses

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Abstract

It is widely accepted that the direction and magnitude of synaptic plasticity depends on post-synaptic calcium flux, where high levels of calcium lead to long-term potentiation and moderate levels lead to long-term depression. At synapses onto neurons in region CA1 of the hippocampus (and many other synapses), NMDA receptors provide the relevant source of calcium. In this regard, post-synaptic calcium captures the coincidence of pre- and post-synaptic activity, due to the blockage of these receptors at low voltage. Previous studies show that under spike timing dependent plasticity (STDP) protocols, potentiation at CA1 synapses requires post-synaptic bursting and an inter-pairing frequency in the range of the hippocampal theta rhythm. We hypothesize that these requirements reflect the saturation of the mechanisms of calcium extrusion from the post-synaptic spine. We test this hypothesis with a minimal model of NMDA receptor-dependent plasticity, simulating slow extrusion with a calcium-dependent calcium time constant. In simulations of STDP experiments, the model accounts for latency-dependent depression with either post-synaptic bursting or theta-frequency pairing (or neither) and accounts for latency-dependent potentiation when both of these requirements are met. The model makes testable predictions for STDP experiments and our simple implementation is tractable at the network level, demonstrating associative learning in a biophysical network model with realistic synaptic dynamics.

Introduction

Activity-dependent change in synaptic strength is widely believed to underlie learning and memory, as originally proposed by Hebb [1]. To have behavioural significance, synapses must be equipped to extract statistical regularities in the spiking behaviour of their pre- and post-synaptic neurons. Numerous algorithms have implemented this principle in artificial neural networks, where the strength of connections between processing units changes as a function of correlations in their output rates [2–6]. These algorithms readily learn to associate patterns of network activity with one another, that is, they implement Hebbian associative learning [see 7]). Similarly, activity-dependent change in the strength of synapses, or synaptic plasticity, is well established in the forms of long term potentiation (LTP) [8] and long term depression (LTD) [9]. Notably, the direction of plasticity (LTP or LTD) has been shown to depend on pre- and post-synaptic activation [10,11], including pre- and post-synaptic spike timing correlations [see 12]).

Spike timing-dependent plasticity (STDP) refers to experimental data showing that the direction and magnitude of synaptic change can be controlled by the precise timing of repeated pre- and post-synaptic spike patterns [13–19]. STDP has played an important role in revealing the mechanisms underlying plasticity [see 20,21]), but ‘learning rules’ fit to these data do not easily generalize to deviations from the strict protocols under which the data were recorded [see 22]). In contrast, by simulating the underlying neurophysiology, synaptic models have provided mechanistic explanations for data recorded under different STDP protocols and at different synapses [23–30]. The fundamental premise of these models is that a biological variable captures the statistics of pre- and post-synaptic spiking and is used by intracellular signalling pathways to modify synaptic strength accordingly.

Here, we focus on the synapse that has been most intensively studied in plasticity experiments to date, that between pyramidal neurons in regions CA3 and CA1 of the hippocampus [22]. As with virtually all synapses, plasticity at CA3-CA1 synapses depends on post-synaptic calcium (Ca2+) in dendritic spines, where high levels of Ca2+ lead to LTP, and moderate, above-baseline levels lead to LTD [see 22,31,32]). We follow the approach of Shouval and colleagues [23] by further assuming that the relevant supply of Ca2+ is mediated by NMDA receptors (NMDARs), known to be required for LTP and LTD at synapses onto CA1 neurons [32]. In this regard, NMDARs provide a mechanism for pre- and post-synaptic coincidence detection: they bind glutamate, providing a marker for pre-synaptic spiking, but they are blocked by magnesium until sufficiently depolarized by back-propagating action potentials (BAPs), which provide markers for post-synaptic spiking. Ca2+ enters the post-synaptic spine when these two markers overlap.

Under STDP protocols, LTP at CA3-CA1 synapses not only depends on Ca2+ flux and NMDAR activation, but also requires
post-synaptic bursting and an inter-pairing frequency in the range of the hippocampal theta rhythm [19] (∼4−12 Hz). We build on earlier modelling work [23–30,33] by proposing a novel mechanistic explanation for these two requirements for LTP. We hypothesize that these requirements reflect the saturation of the mechanisms underlying Ca$^{2+}$ extrusion from the post-synaptic spine [34,35], supporting the buildup of Ca$^{2+}$ by preventing its decay with sufficiently-vigorous spiking activity. We test this hypothesis with a dynamic model of post-synaptic protocols, as well as experiments from two earlier, related studies.

A baseline time constant and a sigmoidal function of such, the time constant of decay was modelled as the summation of protocols at CA3-CA1 synapses [19,36]; we demonstrate that under biophysically plausible parameter values, the model cannot reproduce these findings without the dynamic Ca$^{2+}$ time constant; and we make predictions for experimental verification. We further demonstrate that under the same parameters that reproduce these data, our model supports Hebbian associative learning in a biophysical network model of hippocampal circuitry with realistic synaptic dynamics.

**Methods**

Our model is based on the premise that high levels of post-synaptic Ca$^{2+}$ trigger kinase-activated intracellular pathways, leading to LTP, and moderate, above-baseline levels trigger phosphatase-activated pathways, leading to LTD [31,37,38]. Consistent with evidence that plasticity at CA1 synapses is NMDAR-dependent [32], we assume that NMDARs provide a coincidence detector of pre- and post-synaptic activity, permitting the influx of Ca$^{2+}$ when their activation coincides with BAPs [23]. We take a simple approach in this regard, simulating the activation of NMDARs as the proportion of glutamate-bound channels at the synapse, simulating BAPs as the proportion of maximum dendritic depolarization, and simulating Ca$^{2+}$ flux as the decaying property of these two terms, scaled by a soft cap. Note that our model addresses homosynaptic plasticity, implicit in the assumption of coincidence detection by NMDARs. Like earlier authors [23,28,30,33], we assume that the tail of the BAP has a residual effect on the level of post-synaptic depolarization, so that the overlap between NMDARs and BAPs supports Ca$^{2+}$ flux when post-synaptic spikes precede pre-synaptic spikes during STDP protocols (see below).

To test the hypothesis that the post-synaptic burst-dependence and inter-pairing frequency dependence of LTP at CA3-CA1 synapses can be explained by the saturation of the mechanisms of Ca$^{2+}$ extrusion from the spine, we modelled the saturation of these mechanisms with a Ca$^{2+}$-dependent Ca$^{2+}$ time constant. As such, the time constant of decay was modelled as the summation of a baseline time constant and a sigmoidal function of Ca$^{2+}$. We directly simulated the STDP experiments of [19] by generating pre- and post-synaptic impulses according to their experimental protocols, as well as experiments from two earlier, related studies of CA3-CA1 synapses [15,36].

To demonstrate that our Ca$^{2+}$-based plasticity model can support Hebbian associative learning under biologically plausible conditions, we embedded the model in a network of leaky integrate-and-fire neurons with realistic synaptic dynamics. Our chosen task was auto-associative recall, a classic task for models of hippocampal circuitry [39,40].
transients can be measured. However, the binding of learning rates, we set $H_w$ by data from CA1 slices [46]. For the potentiation and depression of synaptic activity, but did not qualitatively effect our results.

LTD window as a function of the latency between pre- and post-

$Ca^{2+}$ burst requirements of LTP were not satisfied (shown in the Results). For the $Ca^{2+}$-dependent Ca$^{2+}$ time constant $\tau_{Ca}(t)$, we chose a saturating function because our model captures a saturating process (extrusion of Ca$^{2+}$ from the spine). The baseline Ca$^{2+}$ time constant was $\tau_{0,Ca} = 25$ ms [34] and the maximum Ca$^{2+}$ time constant was $T = 500$ ms, previously reported as the time constant of recovery from saturated extrusion in the spine at CA1 synapses [35]. We set the slope parameter $\vartheta = 15$ to a moderate value (see Figure 1C). The only remaining parameters were the LTD threshold $\Theta_d$ and the scale factor $\psi$, which we adjusted to produce a good approximation of the LTP data by [19] for 5 Hz pairings of a single pre-synaptic spike with two post-synaptic spikes (shown in the Results). The chosen values were $\Theta_d = 0.1$ and $\vartheta = 0.135$, which we used in our simulations of all other protocols. Note that within a reasonable range ($\sim 0.05 - 0.2$), the LTD threshold controlled the width of the LTD window as a function of the latency between pre- and post-synaptic activity, but did not qualitatively effect our results.

It is worth noting that the baseline time constant of calcium decay ($\tau_{0,Ca} = 25$ ms) refers to Ca$^{2+}$ extrusion from the spine, which dominates Ca$^{2+}$ clearance at synapses onto pyramidal neurons (see [47]). Measurement of $\tau_{0,Ca}$ is achieved by visualizing calcium transients with Ca$^{2+}$-sensitive fluorescent molecules (calcium indicators), i.e. the indicators bind calcium so the transients can be measured. However, the binding of Ca$^{2+}$ by indicators lengthens the duration of decay by providing an ‘exogenous buffer’ for Ca$^{2+}$, which slows its clearance. As such, numerous studies have reported Ca$^{2+}$ time constants on the order of hundreds of milliseconds, but these measurements included the effects of indicators. The time constant of Ca$^{2+}$ decay can be estimated by using multiple concentrations of indicator, measuring the time constant of decay for each concentration, and regressing to the case of 0-indicator. As such, the baseline time constant of Ca$^{2+}$ decay in dendritic spines is estimated to be several tens of milliseconds [34,48]. At high concentrations of spine Ca$^{2+}$, the mechanisms of extrusion saturate [35], providing the basis for our model. See [47] for a thorough review of these and related issues.

The Network Model

The local circuit model is a fully recurrent network of leaky integrate-and-fire neurons [49], comprised of $N_p = 200$ simulated pyramidal neurons and $N_i = 50$ fast-spiking inhibitory interneurons. Excitatory currents from pyramidal neurons were mediated by AMPA receptor (AMPA) and NMDAR conductances, and inhibitory currents from interneurons were mediated by GABA...
receptor (GABAR) conductances. Synaptic connections from pyramidal neurons to interneurons, from interneurons to pyramidal neurons, and from interneurons to interneurons were uniform. Synaptic connections from pyramidal neurons to pyramidal neurons were scaled by weight $w$, set to 0.5 at the start of each trial.

Each model neuron is described by

$$C_m \frac{dV}{dt} = -g_L(V-E_L) - I,$$

where $C_m$ is the membrane capacitance of the neuron, $g_L$ is the leak conductance, $V$ is the membrane potential, $E_L$ is the equilibrium potential, and $I$ is the total input current. When $V$ reaches a threshold $\delta$, it is reset to $V_{\text{reset}}$, after which it is unresponsive to its input for an absolute refractory period of $\tau_{\text{ref}}$.

For pyramidal neurons, $C_m = 0.5 \, \text{nF}$, $g_L = 25 \, \text{nS}$, $E_L = -70 \, \text{mV}$, $\delta = -50 \, \text{mV}$, $V_{\text{reset}} = -60 \, \text{mV}$ and $\tau_{\text{ref}} = 2 \, \text{ms}$. For interneurons, $C_m = 0.2 \, \text{nF}$, $g_L = 20 \, \text{nS}$, $E_L = -70 \, \text{mV}$, $\delta = -50 \, \text{mV}$, $V_{\text{reset}} = -60 \, \text{mV}$ and $\tau_{\text{ref}} = 1 \, \text{ms}$ [44].

The total input current to each neuron $i$ is given by

$$I_i = I_{\text{AMPA},i} + I_{\text{NMDA},i} + I_{\text{GABA},i} + I_{\text{bg},i} + I_{\text{el},i},$$

where $I_{\text{AMPA},i}$, $I_{\text{NMDA},i}$ and $I_{\text{GABA},i}$ are the summed AMPAR, NMDAR and GABAR currents from intrinsic (recurrent collateral) synapses, $I_{\text{bg},i}$ is background activity and $I_{\text{el},i}$ is selective input. The intrinsic currents are defined by

$$I_{\text{AMPA},i} = \sum_j G_{\text{AMPA}} \cdot g_{\text{AMPA},j} (V_i - V_E) \cdot w_{ij},$$

$$I_{\text{NMDA},i} = \sum_j G_{\text{NMDA}} \cdot g_{\text{NMDA},j} (V_i - V_E) \cdot w_{ij} \cdot \eta_i,$$

$$I_{\text{GABA},i} = \sum_j G_{\text{GABA}} \cdot g_{\text{GABA},j} (V_i - V_I),$$

where $G_{\text{AMPA}}$, $G_{\text{NMDA}}$ and $G_{\text{GABA}}$ are the respective strengths of AMPAR, NMDAR and GABAR conductance, $V_E = 0 \, \text{mV}$ is the reversal potential for AMPARs and NMDARs, $V_I = -70 \, \text{mV}$ is the reversal potential for GABARs, and index $i$ and $j$ indicate the synaptic connection to pyramidal neuron $i$ from pyramidal neuron $j$. AMPAR and GABAR activation (proportion of open channels) are described by

$$\frac{dg_{\text{AMPA}}}{dt} = - \frac{g_{\text{AMPA}}}{\tau_{\text{AMPA}}} + \delta(t-t_f),$$

$$\frac{dg_{\text{GABA}}}{dt} = - \frac{g_{\text{GABA}}}{\tau_{\text{GABA}}} + \delta(t-t_f),$$

where $t_f$ is the time of firing of a pre-synaptic neuron. NMDAR activation is described above in Equation 1. The voltage-dependence of NMDARs is captured by $\eta = 1/(1 + Mg \cdot \exp(-0.062 - V/3.57))$, where $M_g = 1 \, \text{mM}$ describes the extracellular magnesium concentration and $V$ is measured in millivolts [50].

At intrinsic synapses onto pyramidal neurons, $G_{\text{AMPA}} = 0.25 \, \text{nS}$, $G_{\text{NMDA}} = 1.25 \, \text{nS}$ and $G_{\text{GABA}} = 20 \, \text{nS}$. At synapses onto interneurons, these conductances were scaled by a factor of 0.75, a common approach in this class of network [44,45,51,32]. At extrinsic (feedforward) synapses onto pyramidal neurons, $G_{\text{AMPA}} = 2.5 \, \text{nS}$, supporting strong selective input to the network. Consistent with recordings in hippocampal slices, the synaptic time constants used were $\tau_{\text{AMPA}} = 3 \, \text{ms}$ [53] and $\tau_{\text{GABA}} = 10 \, \text{ms}$ [54] ($\tau_{\text{NMDA}}$ is provided above).

**Background activity.** Theta-rhythmic background activity was simulated by sinusoidal current injection. The current injected at each neuron was the product of a 5 Hz sine wave and the point conductance model of [55]. For each neuron, current $I_{\text{bg}}$ is defined by

$$I_{\text{bg}} = g_x(t)(V - V_E) + g_y(t)(V - V_I),$$

where reversal potentials $V_E$ and $V_I$ are given above (Equation 11). The time-dependent excitatory and inhibitory conductances $g_x(t)$ and $g_y(t)$ are updated at each timestep $\delta t$ according to

$$g_x(t + \delta t) = g_0 + [g_x(t) - g_0] \cdot e^{-t/\tau_x} + A_x \cdot Y,$$

and

$$g_y(t + \delta t) = g_0 + [g_y(t) - g_0] \cdot e^{-t/\tau_y} + A_y \cdot Y,$$

where $g_0$ and $g_0$ are average conductances, $\tau_x$ and $\tau_y$ are time constants, $Y$ is normally distributed random noise with 0 mean and unit standard deviation. Amplitude coefficients $A_x$ and $A_y$ are defined by

$$A_x = \frac{D_x \tau_x}{2} \left[1 - \exp \left(-\frac{2\delta t}{\tau_x}\right)\right],$$

and

$$A_y = \frac{D_y \tau_y}{2} \left[1 - \exp \left(-\frac{2\delta t}{\tau_y}\right)\right],$$

where $D_x = 2\sigma^2_x/\tau_x$ and $D_y = 2\sigma^2_y/\tau_y$ are noise ‘diffusion’ coefficients. See [55] for the derivation of these equations. We followed Table 1 of [55] for parameter values $\tau_x = 5 \, \text{ms}$, $\tau_y = 7.5 \, \text{ms}$, $\sigma_x = 5 \, \text{nS}$ and $\sigma_y = 7.5 \, \text{nS}$ for pyramidal neurons and interneurons and $\sigma_0 = 10 \, \mu\text{S}$ for pyramidal neurons. We gave nominal values to $g_0$ for interneurons and to $g_0$ for pyramidal neurons and interneurons, setting these conductances to 2.5 $\mu\text{S}$, i.e. the network’s intrinsic connectivity was sufficient to mediate realistic levels of inhibitory background activity onto pyramidal neurons and excitatory and inhibitory background activity onto interneurons. All simulations were run with timestep $\delta t = 0.1 \, \text{ms}$ and the standard implementation of Euler’s forward method. Our results were verified with the standard fourth order Runge-Kutta method.

**Results.** Before showing our results, it is useful to summarise the data being addressed. [19] recorded at CA3-CA1 slices in slice preparations, where repetitive pairings of a single pre-synaptic spike with a single post-synaptic spike (the doublet protocol) produced LTD, the magnitude of which was inversely related to the time between spikes $\Delta t$. This was the case regardless of the order of pre- and post-synaptic spiking (the sign of $\Delta t$) or the interpairing frequency (0.1 Hz, 0.5 Hz and 5 Hz). LTD not only required repetitive low-latency pairings of a pre-synaptic spike with at least two post-synaptic spikes (the triplet protocol), but also...
required an inter-pairing frequency of 5 Hz. As with the doublet protocol, negative latencies produced LTD under the triplet protocol, the magnitude of which was inversely related to latency. Notably, these conditions also produced an LTD window at intermediate positive latencies, that is, LTD occurred when a post-synaptic burst followed a pre-synaptic spike by several tens of milliseconds. These data suggest temporally symmetric learning during the hippocampal theta rhythm, where tight coincidence of burst firing yields LTP, loose coincidence yields LTD, and a lack of coincidence yields no synaptic change (see [22]). Note that these constraints fit naturally with auto-associative models of hippocampal learning [39,40,56], demonstrated in Results section Hebbian associative learning in the network model. We refer to this form of STDP as temporally symmetric, bidirectional STDP. We refer to STDP data in which pre-before-post pairings lead only to LTD and post-before-pre pairings lead only to LTD as temporally asymmetric, bidirectional STDP. In summary, [19] ran experiments with four combinations of spike pairings and inter-pairing frequencies: (1) the doublet protocol at 0.1 Hz and 0.5 Hz produced LTD; (2) the doublet protocol at 5 Hz produced LTD; (3) the triplet protocol at 0.5 Hz produced LTD; and (4) the triplet protocol at 5 Hz produced temporally symmetric, bidirectional STDP. Thus, LTP required short-latency pairings in the range of theta and post-synaptic bursting. LTD was produced if either of these requirements was not met.

In addition to the data by [19], we also addressed data from two earlier plasticity studies with inter-pairing frequencies in the theta range, one demonstrating the requirement of post-synaptic bursting for LTD at CA1 synapses under protocols different than those described above [36], and one showing temporally symmetric, bidirectional STDP at CA1 synapses without the requirement of post-synaptic bursting [15]. In the latter case, we demonstrate that the physiological explanation provided by [19] for the lack of burst-dependence found by [15] accounts for these data in our model. All these results were produced under the same parameters.

### Temporally Symmetric STDP at CA3-CA1 Synapses

To determine whether our Ca\(^{2+}\) plasticity model can account for the data, we followed three STDP protocols used by [19]. First, we conducted simulations of the doublet protocol at latencies \(-100 \leq \Delta t \leq 100\) ms, using inter-pairing frequencies of 0.5 Hz and 5 Hz. We refer to these pairings as pre-post for \(\Delta t \geq 0\) ms and post-pre for \(\Delta t < 0\) ms. Next, we simulated the triplet protocol (a single pre-synaptic spike and two post-synaptic spikes) over the same range of latencies \(\Delta t\), using an inter-pairing frequency of 0.5 Hz. In this case, \(\Delta t\) refers to the time between the pre-synaptic spike and the second post-synaptic spike. The post-synaptic spikes were separated by 10 ms. We refer to these pairings as pre-post for \(\Delta t \geq 10\) ms, post-pre for \(0 < \Delta t < 10\) ms, and post-post for \(\Delta t < 0\) ms. Finally, we repeated the triplet protocol with an inter-pairing frequency of 5 Hz. For all three sets of simulations, we used 75 pairings, consistent with the corresponding experiments ([19] used 70–100 pairings in different conditions). These protocols and corresponding terminology are depicted in Figure 2.

The doublet protocol at 0.5 Hz and 5 Hz produced LTD-only curves, where the magnitude of LTD was greatest at low latency, decreasing with increasing latency in the causal (\(\Delta t > 0\)) and anti-causal (\(\Delta t < 0\)) directions (Figure 3B). Very similar curves were produced for all frequencies less than or equal to 9 Hz. Quantitatively, these LTD-only curves are narrower than the Gaussian fit to the data shown in Figure 3A, but their qualitative resemblance is clear. The triplet protocol at 5 Hz produced an LTP window for right temporal coincidence (\(-1 \leq \Delta t \leq 25\) ms), surrounded by LTD windows in which the magnitude of LTD decreased with increasing latency on either side of the LTP window (Figure 4B, solid black). The corresponding data are reproduced from [19] in Figure 4A. Note that a graded transition between maximal LTP and LTD can be produced by tuning the LTD threshold. The triplet protocol at 0.5 Hz produced an LTD-only curve (Figure 4B, dotted grey). The mean value of the Ca\(^{2+}\)-dependent Ca\(^{2+}\) time constant \(\tau(\text{Ca}(t))\) is plotted in Figures 3C and 4C for each condition shown in panel B in each figure.

Our simulations thus support the hypothesis that the post-synaptic burst-dependence and inter-pairing frequency dependence of LTD at CA3-CA1 synapses can explained by the saturation of the mechanisms of Ca\(^{2+}\) extrusion from the post-synaptic density. The mechanistic basis of these results is clear in Figure 5, where the dynamic time constant \(\tau(\text{Ca}(t))\) and the corresponding values of the Ca\(^{2+}\)-like variable \(\text{Ca}(t)\) are shown for the above protocols. The dynamic time constant supports high levels of \(\text{Ca}(t)\) for the triplet protocol at 5 Hz, but under the other protocols, it decays to baseline between pairings (see figure caption for details).

### Table 1. A comparison of Ca\(^{2+}\)-based plasticity models.

| Model                  | NM | CT | IP | SS | CE | TD | LM |
|------------------------|----|----|----|----|----|----|----|
| Shouval et al. (2002)  | yes| int| no | yes| no | sym| no |
| Karmarkar and Buonomano (2002) | no | peak| no | yes| no | sym| no |
| Model 1                | no | peak| no | yes| no | sym| no |
| Model 2                | no | peak| no | no | no | asym| no |
| Abarabanel et al. (2003) | no | time| no | yes| no | sym| no |
| Rubin et al. (2005)    | no | time| no | yes| no | both| no |
| Hartley et al. (2006)  | yes| int| no | no | no | both| yes|
| Rackham et al. (2010)  | yes| peak| no | yes| no | sym| no |
| Kumar and Mehta (2011) | yes| int| no | yes| no | sym| no |
| Graupner and Brunel (2012) | no | int | yes| yes| no | asym| no |
| Linear model           | no | int | yes| yes| no | both| no |
| Nonlinear model        | no | int | yes| yes| no | asym| no |
| Bush and Jin (2012)    | yes| int| yes| yes| no | sym| no |
| Our model              | yes| int| no | yes| sym| yes| yes |

A comparison of Ca\(^{2+}\)-based plasticity models. In relation to our model, earlier models can be distinguished along dimensions including: whether NMDARs provide the only source of calcium for plasticity (NM: yes/no); whether plasticity outcomes are determined by integrated Ca\(^{2+}\), peak Ca\(^{2+}\) or the timecourse of Ca\(^{2+}\) transients (CT: int/peak/time); whether an intermediate process transforms calcium levels to plasticity outcomes (IP: yes/no); whether LTD and LTP depend on the same source of Ca\(^{2+}\) (SS: yes/no); whether saturation of the mechanisms of Ca\(^{2+}\) extrusion was simulated (CE: yes/no); whether the model accounts for symmetric and/or asymmetric STDP data (TD: sym/asym/both); and whether the model demonstrated learning and memory (LM: yes/no). doi:10.1371/journal.pone.0086248.t001
Simulations without the Dynamic Ca^{2+} Time Constant

To determine whether the dynamic time constant $\tau(Ca(t))$ is required for the model to reproduce the data, we repeated our simulations without it, i.e., with $\tau(Ca(t))$ set to the baseline Ca^{2+} time constant $\tau_{0,Ca}$. Because the parameter values given above were chosen for the intact model, it was necessary to choose parameter values without the dynamic time constant before running these simulations. To this end, we searched the space of values for three parameters: the baseline Ca^{2+} time constant $\tau_{0,Ca}$, the time constant of decay of the BAP tail $\tau_{p}$, and the LTP threshold $\Theta_{p}$. Both versions of the model (with and without the dynamic time constant) readily account for the LTD data under the doublet protocol, so we searched for parameter values that produce LTP under the triplet protocol at 5 Hz, but produce LTD under the triplet protocol at 0.5 Hz. We did not search the time constants of rise and decay of NMDAR activation because the LTP window shown by [19] was approximately $0 \leq \Delta t \leq 25$ ms, whereas the decay constant of NMDARs is $\tau_{NMDA} \geq 50$ ms, i.e., reasonable values of $\tau_{NMDA}$ govern the width of causal (pre-before-post) LTD, but do not effect the occurrence of LTP. We did not search the scale factor $\psi$ (Equation 5) because without the non-linearity of the dynamic time constant, this factor changes the scale of the simulations, not the phenomenology, i.e., any changes to $\psi$ can be overcome by changes to $\Theta_{p}$. We did not search the proportion of the BAP given to the peak and the tail because these parameters change the scale of calcium transients for a given value of $\tau_{0,Ca}$, but not their decay time. As long as Ca(t) does not have the opportunity to buildup between pairings, LTP will not occur at 0.5 Hz. Note that the available data support our chosen values, justified in Methods section Parameter values. Finally, we ignored the learning rates $k_p$ and $k_d$ because they govern the magnitude of LTP and LTD respectively, not their occurrence.

The results of our parameter search for the model without the dynamic time constant are shown in Figure 6, where the LTP window (expressed as $\Delta t^\text{LTP}$) is shown over the LTP threshold $\Theta_p$ for different values of the baseline Ca^{2+} time constant $\tau_{0,Ca}$ (panels A, B and C) and the BAP tail $\tau_{p}$ (panel D). The lower and upper curves show the lowest and highest values of $\Delta t^\text{LTP}$ under the triplet protocol for a given set of parameters. The black and grey curves show results for 5 Hz and 0.5 Hz respectively, where $\Theta_p$ was incremented by 1 $\times$ 10^{-3} in the search. For a given set of parameters to be consistent with the data, there should be an LTP window between approximately 0 and 25 ms for 5 Hz, but not for 0.5 Hz, i.e., there should be black curves, but not grey curves in this region. For $\tau_p = 40$ ms and $\tau_{0,Ca} = 25$ ms (justified in Methods section Parameter values), fine tuning of the LTP threshold permits a non-overlapping region for $11 \leq \Delta t^\text{LTP} \leq 12$ ms (Figure 6A). Doubling $\tau_{0,Ca}$ has little effect ($\sim 10 \leq \Delta t^\text{LTP} \leq 14$ ms, Figure 6B). The data can be approximately reproduced for $\tau_{0,Ca} = 150$ ms (Figure 6C), but this value is biologically implausible (see Methods section 0). Alternatively, with the plausible baseline Ca^{2+} time constant $\tau_{0,Ca} = 25$ ms, the data can be approximated with $\tau_p = 125$ ms (Figure 6D), but we are unaware of any data to support a BAP tail longer than ~40 ms. Thus, fine-tuning the model’s parameters allows the reproduction of the data without the dynamic time constant, but not under realistic parameter values. Further to this, we
considered the robustness of the model by including noise in simulations of the triplet protocol at 0.5 Hz. Without the dynamic time constant, the model was given $H_p \sim 0.6$ and the implausible baseline $Ca_{z} \sim 150 \text{ ms}$ (see Figure 6C). We added normally distributed noise with mean $\mu_x \sim 0$ and standard deviation $\sigma_x \sim 0.05$ to each parameter $x$ in each version of the model (with and without the dynamic time constant), and we ran 100 trials for each value of $\Delta t$ (the time between the pre-synaptic spike and the second post-synaptic spike) for $10 \leq \Delta t \leq 20 \text{ ms}$ (increments of 1 ms). This range of $\Delta t$ was used by [19] in their experiments with spike triplets at 0.5 Hz, producing LTD. The noise had little impact on the dynamic model, where only 6 out of 1100 trials produced LTP. Without the dynamic time constant, these simulations produced LTP on 256 trials out of 1100.

Simulations of Further Spike-timing Protocols at CA1 Synapses

The study by [19] is not the only one to demonstrate the requirement of post-synaptic bursting for LTP at CA1 synapses under spike-timing protocols with an inter-pairing frequency in the theta range. Earlier work by [36] also showed this requirement, where four protocols were followed: one pre-synaptic spike followed by one post-synaptic spike (1-pre-1-post), three pre-synaptic spikes followed by one post-synaptic spike (3-pre-1-post), one pre-synaptic spike followed by three post-synaptic spikes (1-pre-3-post), and three pre-synaptic spikes followed by three post-synaptic spikes (3-pre-3-post). For the 3-pre-1-post and 3-pre-3-post protocols, the time between pre-synaptic spikes was $D_{tpre} \sim 5 \text{ ms}$. For the 1-pre-3-post and 3-pre-3-post protocols, post-synaptic spikes were elicited by current pulses of 20 ms, during which time the post-synaptic neuron emitted three spikes. In all cases, pre-synaptic activity preceded post-synaptic activity by $10 \leq \Delta t \leq 20 \text{ ms}$. Ten pairings were provided with an inter-pairing frequency of 5 Hz, repeated four times at 10 s intervals. We simulated all four protocols for each integer value of $\Delta t$. We set the
spikes were separated by window for normally-occurring potassium. Cesium blocks potassium channels, the doublet protocol, also at 5 Hz. A possible explanation for this 5 Hz, earlier authors found LTP at these synapses by pairing a post-synaptic spike and an inter-pairing frequency of higher Ca concentration. Comparing data from different studies, especially given the much higher Ca concentration, the model requires biologically implausible parameter values to reproduce the data (panels C and D, see text). (A) The LTP window for parameter values justified in the Methods (t prer = 25 ms, t post = 40 ms). (B) The LTP window for t prer = 50 ms and t post = 40 ms. (C) An implausibly long baseline time constant of Ca2+ decay approximates the data (t prer = 150 ms, t post = 40 ms). (D) An implausibly long time constant of decay of the BAP tail approximates the data, though Dtp post expands slightly in the anti-causal direction (t prer = 25 ms, t post = 125 ms).

doi:10.1371/journal.pone.0086248.g007

Figure 7. The Ca2+-like variable Ca(t) in simulations of the experiments by [36]. Results are shown for simulations with Δt = 10 ms, Δt prer = 5 ms and Δt post = 10 ms, where Δt refers to the time between the last pre-synaptic spike and the first post-synaptic spike, Δt prer refers to the time between pre-synaptic spikes and Δt post refers to the time between post-synaptic spikes. The LTP threshold (horizontal line) is only exceeded under conditions with post-synaptic bursting (1-pre-3-post and 3-pre-3-post), while pre-synaptic bursting has little effect on Ca(t) (3-pre-1-post). The inter-pairing frequency was 5 Hz.
doi:10.1371/journal.pone.0086248.g008

Predictions for Experimental Enquiry

Having shown that the model accounts for the inter-pairing frequency dependence and post-synaptic burst dependence of LTP at CA3-CA1 synapses under the STDP protocols of [19], we ran further simulations to investigate more subtle variations of these two independent variables. First, we ran simulations of the doublet protocol at inter-pairing frequencies above 5 Hz. For frequencies lower than 9 Hz, the model produced LTD-only curves (Figure 3B), similar to those shown in Figure 3B. A narrow, causal LTP window emerged at 10 Hz, expanding in both directions as frequency was increased up to 14 Hz, shown in Figure 3B. Note that the decreasing range of Δt with increasing frequency in the figure reflects the shortening of the period. At 15 Hz, LTD was saturated for all Δt (Figure 9B). The model thus predicts that temporally symmetric, bidirectional STDP can be produced by the doublet protocol at higher frequencies within the theta band, including LTP under low-latency post-pre pairings, and that the spike which can broaden action potentials and enhance BAPs in the apical dendrite, where CA3 neurons synapse onto CA1 neurons [57]. [19] therefore made further recordings under the doublet protocol (Δt = 5 ms) with a cesium-based intracellular solution, finding that action potentials were broadened by a factor of 5. This manipulation was sufficient to rescue LTP. We simulated the effect of their cesium solution by broadening the decay constant of the BAP peak by a factor of 5 (from τ pre = 3 ms to τ pre = 15 ms) and re-running our simulations of the doublet protocol for the full range of latencies Δt. This small change to the model's parameters was sufficient to qualitatively reproduce the data by [15] (Figure 6).
timing-dependence of LTP is dominated by spike rate at frequencies higher than theta.

Next, we ran simulations of the triplet protocol with a range of inter-pairing frequencies. For frequencies \( \leq 3 \text{ Hz} \), the model produced LTD-only curves (Figure 10A), identical to the dotted grey curve in Figure 4B. At 4 Hz, a narrow, causal LTP window emerged. Similar to the higher-frequency doublet simulations, this LTP window expanded in both directions as frequency was increased up to 10 Hz (Figure 10B). At 11 Hz, LTP was saturated for all \( \Delta t \) (Figure 10B). The model thus predicts that under the triplet protocol, the LTP window of temporally symmetric, bidirectional STDP expands in both directions at higher frequencies within the theta band, including LTP under post-post-pre pairings, and that the spike timing-dependence of LTP is dominated by spike rate at sufficiently high frequency.

Having made predictions for a gradient of frequencies under the doublet and triplet protocols, we investigated the post-synaptic burst-dependence of LTP under low-frequency protocols with more than 2 post-synaptic spikes. We ran simulations at 2 Hz and 3 Hz with a quadruplet protocol, where one pre-synaptic spike was paired with three post-synaptic spikes. As with the doublet and triplet protocols, \( \Delta t \) was defined as the temporal difference between the pre-synaptic spike and the last post-synaptic spike (\( \Delta t \) is also defined this way for quintuplet and sextuplet pairings below). Under the quadruplet protocol, an LTP window emerged at 3 Hz, shifted in the causal direction (LTP onset at \( \Delta t = 9 \text{ ms} \), Figure 11A). At 2 Hz, a graded, LTD-only curve was produced by this protocol. Similarly, we ran simulations with a quintuplet protocol, where one pre-synaptic spike was paired with four post-synaptic spikes at frequencies of 1 Hz and 2 Hz. Temporally symmetric, bidirectional STDP was produced under these low inter-pairing frequencies (identical curves in Figure 11B). These predictions are important to the interpretation of the data by [19]. As shown in Figures 11C and D, there is a difference between the mechanism underlying LTP under the triplet protocol and under the quintuplet and sextuplet protocols (5 post-synaptic spikes). Under the triplet protocol, an inter-pairing frequency in the theta range prevents \( \text{Ca}(t) \) from decaying to baseline between pairings, so \( \text{Ca}(t) \) builds up over several pairings and eventually exceeds the LTP threshold (Figure 5). Under the quintuplet and sextuplet protocols, \( \text{Ca}(t) \) exceeds the LTP threshold during each post-synaptic burst, but decays to baseline between pairings. This mechanistic difference explains why the STDP curves under the quintuplet protocol were identical at 1 Hz and 2 Hz, i.e. the inter-pairing frequency made no difference (Figure 11B). The model thus predicts that with bursts of more than two or three post-synaptic spikes, LTP does not require theta-frequency pairings.

**Hebbian Associative Learning in the Network Model**

Having shown that our plasticity model qualitatively reproduces STDP data from CA3-CA1 synapses [15,19,36] and makes predictions for experimental enquiry, we embedded the model in a network of integrate-and-fire neurons with realistic synaptic dynamics to determine whether it supports Hebbian associative learning under biophysically plausible conditions. Our chosen task was auto-associative recall, a classic task for hippocampal models [39,40,56], in which a spatial pattern of neural activity is associated with itself by the modification of excitatory recurrent synapses (see Discussion section *Synaptic plasticity, learning and memory*). All neurons in the network received noisy, theta-rhythmic current injection (see Methods section *Background activity*), while the neurons in a selective population (neurons 50–100 in Figure 12A) each received independent, homogeneous Poisson spike trains at 500 Hz, mediated by AMPAR conductance synapses. The input to the selective population was provided for 2 s while the plasticity
model was implemented at all recurrent synapses. We then provided the 500 Hz spike trains to half the selective population (neurons 75–100 in Figure 12A) for 2s without the plasticity model. As shown in Figure 12, the model supported associative learning, i.e. the network recalled the full spatial activity pattern from partial input. The value of \( Ca^2+ \) at a randomly-chosen synapse between two neurons in the selective population is shown in Figure 12B. The final strength of all synapses in the network is shown in Figure 12C.

Discussion

Under STDP protocols, LTP at CA3-CA1 synapses depends on post-synaptic bursting and an inter-pairing frequency in the range of the hippocampal theta rhythm [19]. We hypothesize that these dependencies reflect the saturation of the mechanisms of \( Ca^2+ \) extrusion from the post-synaptic spine. We have tested this hypothesis with a minimal model of \( Ca^{2+} \)-dependent plasticity, where the saturation of the mechanisms of \( Ca^{2+} \) extrusion is simulated by the \( Ca^{2+} \)-dependence of the time constant of \( Ca^{2+} \) decay (Figure 1). Our model qualitatively reproduces the STDP data of [19], where latency-dependent LTD is elicited by pre- and post-synaptic spike doublets at frequencies of 5 Hz or less (Figure 3), and by spike triplets (one pre-synaptic spike and two post-synaptic spikes) at frequencies of 0.5 Hz or less (Figure 4). At a frequency of 5 Hz, triplet pairings produce temporally symmetric, bidirectional STDP (Figure 4). Without the \( Ca^{2+} \)-dependent \( Ca^{2+} \) time constant, the minimal model cannot reproduce these data under biologically plausible parameter values (Figure 6). The model further accounts for the dependence of LTP on post-synaptic bursting at CA1 synapses under other spike timing protocols, including those with multiple pre-synaptic spikes [36] (Figure 7); and accounts for temporally symmetric, bidirectional...
STDP under spike doublets with intracellular solutions that broaden BAPs at CA3-CA1 synapses [15,19] (Figure 8).

The model makes testable predictions about the burst-dependence and frequency-dependence of STDP at CA3-CA1 synapses. For example, our simulations predict that temporally symmetric, bidirectional STDP will emerge under the doublet protocol at higher frequencies within the theta band (~10–14 Hz), with the LTP window expanding in the causal and anti-causal directions with increasing frequency (Figure 9). The model predicts that this expansion will also occur under the triplet protocol at lower theta frequencies (~4–10 Hz, Figure 10) and that pairing pre-synaptic spikes with more than two or three post-synaptic spikes will elicit LTP at any sub-theta frequency (Figure 11). Our Ca\(^{2+}\) plasticity model is simple enough to be implemented at the network level and we have demonstrated associative learning and recall under theta oscillations in a biophysical network model with realistic synaptic dynamics (Figure 12).

Our model is based on the Ca\(^{2+}\) control hypothesis [23,30], where high levels of post-synaptic Ca\(^{2+}\) lead to LTP and moderate levels lead to LTD. Consistent with the NMDAR-depence of plasticity at CA3-CA1 synapses, NMDARs provide the relevant source of Ca\(^{2+}\). We refer to our model as 'minimal' because the only biophysical mechanisms included in the model are those required to reproduce the data being addressed: a marker for pre-synaptic spikes, a marker for post-synaptic spikes, and a variable to quantify the coincidence of these markers, the time constant of which is monotonically dependent on the magnitude of the variable. The only biophysical correlates of these computational requirements are NMDARs, long-tailed BAPs, Ca\(^{2+}\), and the Ca\(^{2+}\)-dependent Ca\(^{2+}\) time constant \(\tau(Ca(t))\). This parsimonious approach not only limits the assumptions of the model, but affords a simple implementation at the network level. Indeed, the chosen network model is endowed with the minimum biophysical detail to account for a range of data from perceptual and cognitive tasks (e.g. [45,51,32,58]). The only biophysical mechanism that we added to this generic network is the BAP, driving Ca\(^{2+}\) dynamics in conjunction with NMDARs.

**STDP and the Need for Physiologically-motivated Models**

Since the discovery of graded, temporally asymmetric STDP in cultured hippocampal cells [14], numerous computational models have sought to characterize synaptic plasticity according to these data. In much of this work, 'STDP rules' have been defined by curves fit to the data, which are assumed to capture the phenomenology of synaptic change with ongoing pre- and post-synaptic spikes [59–64]. This assumption is perfectly reasonable, indeed unavoidable, for the spike trains under which the data were recorded. Given the implicit Hebbian form of temporally asymmetric STDP, this approach has an intuitive appeal, i.e. pre-post spike doublets imply that a pre-synaptic cell 'repeatedly and persistently takes part in firing' [1] a post-synaptic cell and should therefore lead to LTP, whereas post-pre doublets imply a lack of causality and therefore LTD. Further appeal stems from competitive interactions between the LTP and LTD portions of asymmetric STDP curves, known to support synaptic scaling with uncorrelated [65] and correlated [64] pre- and post-synaptic spiking.

The application of asymmetric STDP rules to spike trains different from those that produced the underlying data has not been straightforward. In neural simulations, the sampling of STDP curves with each pre- and post-synaptic spike has not accurately predicted the outcome of various triplet, quadruplet, and quintuplet protocols (see [22]). Augmenting STDP rules with so-called spike interaction rules has been productive in this regard, limiting the contribution of individual spikes to plasticity [17,62,66]. Biological correlates have been proposed for the time-dependent suppression of interactions between spikes [67], but ultimately, this approach requires the continual addition of rules to accommodate data recorded under different spiking conditions. See [22] for a thorough treatment of the limitations of this phenomenological approach. Note that temporally symmetric, bidirectional STDP [13] was discovered near-concurrently to its asymmetric homologue, but its phenomenology has received little attention.

**Ca\(^{2+}\)-based plasticity models.** An alternative to the phenomenological approach is to model synaptic physiology, providing a mechanistic explanation for plasticity data recorded under any protocol. STDP data have motivated several such models, including ours. As described above, all such models are based on the Ca\(^{2+}\) control hypothesis, where high and moderate levels of Ca\(^{2+}\) lead to LTP and LTD via kinase- and phosphatase-activated pathways respectively. Our minimal model simulates Ca\(^{2+}\) transients by the product of NMDAR activation and the BAP, where the Ca\(^{2+}\)-dependent Ca\(^{2+}\) time constant \(\tau(Ca(t))\) captures the saturation of the mechanisms of post-synaptic Ca\(^{2+}\) extrusion. Earlier Ca\(^{2+}\) models differ along a number of dimensions, including their underlying assumptions, their levels of biophysical detail, and the data they address. All of these models assume that NMDARs provide Ca\(^{2+}\) for plasticity, but some assume that voltage-gated Ca\(^{2+}\) channels (VGCCs) are also required (e.g. [24–26,29]), an assumption supported by experimental data under some conditions [68,69]. Assumptions differ about the relevant aspect of supra-threshold Ca\(^{2+}\), such as integrated Ca\(^{2+}\) [23,27,29,30,33] and the peak [24,26] and timecourse [25,26] of Ca\(^{2+}\) transients.

Ca\(^{2+}\)-based models have been used to address temporally symmetric and asymmetric STDP data. Any model based only on post-synaptic Ca\(^{2+}\) levels will demonstrate anti-causal LTD [23,26] and will therefore account for symmetric STDP. Additional mechanisms are required to explain asymmetric STDP, for example, the inclusion of separate sources of Ca\(^{2+}\) for LTP and LTD processes [24,27] or competitive interactions between these processes and an LTD veto process for appropriate Ca\(^{2+}\) transients [26]. The former is supported by data from neocortical slices [70] and the latter is consistent with competition between kinases and phosphatases (e.g. [71]). Since BAPs are known to attenuate with dendritic distance from the soma, several studies have considered plasticity outcomes as a function of dendritic location [29,30,33], accounting for experimental data showing a switch from LTP to LTD at longer distances [72].

Like our model, some existing Ca\(^{2+}\)-based models have simulated LTP and LTD directly from supra-threshold levels of Ca\(^{2+}\) [23,24,28,33], but others have included an intermediate processes for each of LTP and LTD [26,27,29,30]. These processes correspond to intracellular signalling cascades that mediate plasticity outcomes from Ca\(^{2+}\) levels, which may be loosely equated with kinase- and phosphatase-activated pathways respectively. Two recent studies augmented this approach with a mechanism for synaptic bistability, addressing STDP data. In the model by [29], post-synaptic Ca\(^{2+}\) is the sum of independent pre- and post-synaptic Ca\(^{2+}\) components, assumed to be mediated by NMDARs and VGCCs respectively. Their Ca\(^{2+}\) variable is thus incremented with every pre- or post-synaptic spike, increasing or
decreasing the efficacy of bistable synapses when it exceeds the respective thresholds for LTP and LTD. While their Ca\(^{2+}\) variable has a time constant of a few milliseconds (like \(\kappa_{Ca}\) here), their synaptic changes occur on a timescale on the order of minutes, driving all-or-none transitions to a down-state or an up-state, to which the efficacy variable relaxes in the absence of synaptic activity. Their model reproduces a broad range of data by varying the magnitude of the independent pre- and post-synaptic Ca\(^{2+}\) components, the LTP and LTD thresholds, and the learning rates for LTP and LTD. In the model by [30], supra-threshold Ca\(^{2+}\) transients change the probability of switching from the down-state to the up-state or vice versa, where these changes tend to occur following spiking activity.

As a model of CA1 plasticity, our model addresses temporally symmetric STDP, where the Ca\(^{2+}\)-dependence of Ca\(^{2+}\) decay [35] accounts for the frequency and burst-dependence of LTP [19]. The coincidence of NMDAR activation and the BAP provides the only source [19]. The coincidence of NMDAR activation and the BAP [35] accounts for the frequency and burst-dependence of LTP, but we have shown that under our minimalist assumptions, the model requires implausible parameter values to account for these dependencies without the proposed mechanism. A concise comparison of the above models and ours is provided by Table 1.

It is worth noting that several authors have developed plasticity models that occupy the middle ground between phenomenological and physiological models. Under this approach, abstract variables are responsive to pre- and post-synaptic activity and their values serve as parameters to learning rules that determine changes to synaptic weights. These models have successfully generalized across timing- and rate-based plasticity protocols [73,74].

The timecourse of supra-threshold calcium transients for LTP and LTD. It has been suggested that high levels of Ca\(^{2+}\) leading to LTP can be brief, but that moderate levels leading to LTD must be prolonged (e.g. [26,75]). The technical challenges of controlling and measuring Ca\(^{2+}\) flux at individual synapses are considerable and direct evidence for or against this suggestion is consequently sparse. Perhaps most convincingly, [76] used Ca\(^{2+}\) ‘uncaging’ to control post-synaptic Ca\(^{2+}\) elevation over tens of seconds, where high elevations over \(\sim 10\) s produced LTP and moderate elevations over \(\sim 60\) s produced LTD. These authors noted that the magnitude and duration of Ca\(^{2+}\) elevation comprise a two-dimensional space that is largely unexplored. For instance, it is unclear whether a high-magnitude elevation over a long timescale (e.g. 60 s) would lead to LTP. To the best of our knowledge, this possibility has not been addressed. The flip side of this issue is whether LTD requires Ca\(^{2+}\) to remain above the LTD threshold for an unbroken period of time, as suggested by [26], or simply requires more repetitions of a given protocol than LTP, i.e. a higher number of discrete supra-threshold Ca\(^{2+}\) transients. If the latter, then the slow onset of LTD under several protocols (e.g. [9,19,77]) can be explained by a low LTD learning rate (\(\kappa_d < \kappa_p\) in our model). Notably, at least two studies have shown Ca\(^{2+}\) transients that reach higher amplitudes and decay more slowly with LTP-inducing stimuli than with LTD-inducing stimuli [78,79]. Our model is very robust to parametric variation and our results can be obtained qualitatively for parameters under which Ca\(^{2+}\) exceeds the LTP threshold either in discrete jumps or for an unbroken interval. For Ca\(^{2+}\) to remain above the LTD threshold for an unbroken interval, an additional mechanism is required, e.g. VGCCs. This characterization is apparent in Figure 5, where the Ca\(^{2+}\)-dependence of the Ca\(^{2+}\) time constant prevents high levels of Ca\(^{2+}\) from undergoing significant decay, but moderate levels of Ca\(^{2+}\) drop to baseline between pairings. Methods for controlling and measuring post-synaptic Ca\(^{2+}\) are continually improving (see [47]), so we expect these considerations will be addressed in due course.

Addressing Data from Different Plasticity Protocols

We have stated that Ca\(^{2+}\)-based models capture the underlying physiology of plasticity and are therefore able to address data recorded under different protocols. Several studies have taken this approach (e.g. [25,29]), though it is unclear that a given model should account for data recorded from different tissue preparations and brain regions, i.e. ‘caution is warranted when generalizing from one synapse to another’ [32]. We have also stated that our model contains the minimal biophysical detail to explain the post-synaptic burst-dependence and inter-pairing frequency-dependence of LTP at CA3-CA1 synapses under STDP protocols [19]. Notably, no additional mechanisms or parameter changes were required for the model to reproduce the major findings of studies in which axonal tracts onto CA1 neurons were subject to tetanic stimulation. These studies showed that a brief period of high-rate stimulation (e.g. 100 Hz stimulation for 1 s) produces LTP and a long period of low-rate stimulation (e.g. 1 Hz for 1000 s) produces LTD [9,19]. It is not surprising that we were able to reproduce these findings with an integrate-and-fire neuron. If pre-synaptic stimulation is strong enough to drive the neuron, then low-rate stimulation will lead to LTD in our model (Figures 3B, 9A and 10A). Under the same parameters, high-rate stimulation effectively produces multiple low-latency, pre- and post-synaptic spikes, leading to LTP (not shown).

An aspect of the data by [19] that we did not address above is the dependence of STDP on the number of triplet pairings. Specifically, these authors showed that with 20 – 30 triplet pairings (instead of 70 – 100), their data were fit with a latency-dependent LTP-only curve. These data can be explained by several mechanisms. For example, [29] allowed each of their independent pre- and post-synaptic Ca\(^{2+}\) transients to exceed the LTP threshold. Another possibility is short term fatigue of the mechanisms initiating LTP, readily simulated in our model by increasing the LTP threshold with ongoing LTP. To demonstrate this mechanism, we ran 25 pairings under the triplet protocol with an LTP-dependent LTD threshold. The resulting LTD-only curve is shown in Figure 13 (see the caption for details).

Synaptic Plasticity, Learning and Memory

The use of biophysical plasticity models in network simulations is an important direction in neuroscience, allowing the investigation of the network- and systems-level consequences of plasticity processes. Network- and systems-level modelling with synaptic resolution has a fruitful recent history. For example, our understanding of the respective roles of AMPARs and NMDARs in cortical processing has been greatly advanced by simulations of persistent mnemonic activity in working memory tasks [44,51,80], the integration of evidence in decision tasks [45,81], and the role of oscillations in the transfer of information between cortical networks [58,82]. Several recent papers have discussed the need for this level of biophysical resolution in network models with plastic synapses [22,29,30]. Our simulations of associative memory show that our plasticity model is tractable in this regard. We are aware of only one model to previously bridge this gap, doing so

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under different assumptions than ours and performing a different learning task [27].

Our Ca\textsuperscript{2+} plasticity model is based on data from CA3-CA1 synapses [19,36,83], but memory formation in our network model is perhaps more easily equated with plasticity at CA3-CA3 synapses. Hippocampal region CA3 has long been hypothesized to support episodic memory by auto-association, owing to the dense lateral synaptic connectivity among CA3 neurons and to the well-established role of the hippocampus in episodic memory. For review of this hypothesis, see [84]. The use of our plasticity model in this task seems reasonable, as there do not appear to be any substantial physiological or plasticity differences between CA3-CA3 and CA3-CA1 synapses (Alan Fine, personal correspondence). It is worth noting that temporally symmetric STDP is strikingly consistent with the hypothesis that CA3 supports auto-associative recall. For example, in the hippocampal models by Lisman, Jensen and colleagues, theta/gamma oscillations synchronize coupled neurons and memories are encoded when tight temporal coincidence of pre- and post-synaptic spiking leads to LTP and loose coincidence leads to LTD [56,85]. These conditions are captured by symmetric STDP data from hippocampal preparations [15,19]. In this regard, it is perhaps surprising that the phenomenological implications of symmetric STDP have not received more attention.

Conclusions and Future Work

Our plasticity model builds on earlier work that identified the minimal mechanisms for NMDAR-dependent synaptic plasticity: markers for pre- and post-synaptic activation and a variable (Ca\textsuperscript{2+}) to capture their timing correlations [23]. We have augmented this well-established approach with a previously-unexplored computational mechanism, namely the Ca\textsuperscript{2+} dependence of Ca\textsuperscript{2+} decay. This dynamic Ca\textsuperscript{2+} time constant captures the saturation of the mechanisms of Ca\textsuperscript{2+} extrusion from the postsynaptic spine [34,35]. Our simulations of STDP experiments demonstrate that this mechanism is sufficient to explain the burst-and-frequency-dependence of LTP at CA3-CA1 synapses [19]. Our simulations of untried STDP protocols make testable predictions for the implications of the proposed mechanism and our simulations of associative learning demonstrate that our plasticity model is tractable at the network level.

A limitation of our minimalist approach is the requirement of the BAP for plasticity. As such, the present model cannot reproduce plasticity data obtained under protocols that hold post-synaptic neurons at fixed levels of depolarization for prolonged periods (see [86]). Furthermore, BAPs are neither necessary nor sufficient for plasticity under some conditions [21]. More generally, our Ca\textsuperscript{2+}-like variable is the product of the post-synaptic BAP and pre-synaptic activation, so our model is limited to homo-synaptic plasticity. These considerations point to the need for studies addressing the functional consequences of different plasticity phenomena. The suitability of symmetric STDP to auto-associative learning provides an example of such a functional consequence.

Our study points to several lines of enquiry for future work. Our implementation of the Ca\textsuperscript{2+}-dependence of Ca\textsuperscript{2+} decay is purposefully simplified and a detailed implementation of the mechanisms of Ca\textsuperscript{2+} extrusion may reveal the limitations of our more abstract approach. If so, it may also reveal the computational properties of unexplored calcium dynamics and determine the conditions under which our present, efficient implementation is justifiable in network simulations.

Finally, our study is consistent with the hypothesis that synaptic plasticity is regulated by the modulation of Ca\textsuperscript{2+} extrusion from dendritic spines. In support of this hypothesis, a recent study demonstrated that LTP is not expressed by pyramidal neurons in hippocampal region CA2 largely because rates of extrusion are significantly higher than in regions CA1 and CA3, which express LTP [48]. In these experiments, LTP was blocked in CA1 and induced in CA2 by interventions that speed up and ‘overwhelm’ Ca\textsuperscript{2+} extrusion respectively. We hypothesize that LTP is only produced if the mechanisms of extrusion are overwhelmed. Biophysical mechanisms that may control the saturation of Ca\textsuperscript{2+} extrusion constitute an exciting direction for future research on synaptic plasticity and its functional outcomes.

Acknowledgments

We thank Gayle Wittenberg for thorough explanation of the STDP protocols used by [19]; Stefan Krueger, Alan Fine, Neil Magoski and Colin Lever for helpful discussions; and Michael Shepherd, Srinivas Sampalli and Evangelos Millos for financial support of this work during its early stages.

Author Contributions

Conceived and designed the experiments: DS TT. Performed the experiments: DS. Analyzed the data: DS. Contributed reagents/materials/analysis tools: DS TT GB. Wrote the paper: DS TT GB.

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Figure 13. LTP-only curves for a small number of pairings under the triplet protocol at 5 Hz. (A) Data reproduced from [19], fit with a Gaussian function (least squares fit). (B) LTP ‘fatigue’ produces an LTP-only curve for 25 pairings under the triplet protocol. Fatigue of the processes initiating LTP was simulated with an LTP-dependent LTP threshold \( \Theta_p \), where \( \Theta_p \) was given an initial value equal to the LTD threshold \( \Theta_L \), to which it decayed with a time constant of 1 s. For all positive values of \( \Delta w \), \( \Theta_p \) was incremented by (\( \Theta_p - \Theta_p^0 \))/100. All other parameters are identical to the simulations above. \( \Delta w/w \) refers to the change in synaptic strength relative to initial strength (\( \Delta t \) defined as in Figure 4).

doi:10.1371/journal.pone.0086248.g013
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