Enhanced neuronal response induced by fast inhibition

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We report a facilitatory role of inhibitory synaptic input that can enhance a neuron’s firing rate, in contrast to the conventional belief that inhibition suppresses firing. We study this phenomenon using the Hodgkin-Huxley model of spike generation with random Poisson trains of subthreshold excitatory and inhibitory inputs. Enhancement occurs when, by chance, brief inhibition leads excitation with a favorable timing and counter-intuitively induces a reduction of the spike threshold. The basic mechanism is also illustrated with the phase-plane analysis of a two variable model.

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Inhibitory synaptic inputs play an important role in generating collective neuronal mechanisms of synchronization [1–3], wave propagation [4], chaos [5,6], asynchronous behavior [7], persistent states [8,9], neural oscillations [10], and a formation of cluster states [11]. At the single neuron level, inhibition is also effective in gain control of neuronal signals [12] and temporal sensitivity to coincident inputs [13]. In the classical view, an inhibitory input hyperpolarizes the membrane away from its spike threshold resulting in a reduction of the spike probability. Thus inhibition has conventionally been viewed as a suppressor of neuronal response [14–17], and, in particular, causing either divisive or subtractive effect on the output firing rate [14]. But inhibition playing a facilitatory role was recognized about 50 years ago [18,19], to the best of our knowledge, in the form of postinhibitory rebound (PIR) and is thought to play a major role in central pattern generator networks [20]. In PIR a neuron fires after being released from a long-lasting hyperpolarizing input. Here we report a facilitatory mechanism by which brief inhibitory inputs can, in contrast to the conventional belief, enhance firing probability during ongoing stimulation by trains of brief excitatory inputs. Unlike PIR, this mechanism does not require that an inhibitory input by itself leads to a rebound spike. In our case both the excitatory and inhibitory single inputs are subthreshold in magnitude.

We study this phenomenon using a Hodgkin-Huxley model neuron [21] with external excitatory and inhibitory input conductances. The inputs are subthreshold $\alpha$ functions timed at random independent Poisson intervals. For pure excitatory driving, the neuron responds with a finite output rate due to temporal summation of nearly coincident inputs. When inhibitory inputs are included some spikes are lost but other ones are added. With respect to the onset time of these evoked spikes, inhibitory events form a temporally localized distribution leading ahead of a similar distribution of excitatory events. The leading inhibition can transiently reduce the spike threshold, and a well timed brief subthreshold excitation can utilize this to evoke a spike. We term this phenomenon as the postinhibitory facilitation (PIF) [22]. The enhanced output response could also consist of inhibitory events that are paired with another set of inhibitory events displaying PIF for temporally brief inputs. Our result stresses the importance of the timing of the prespike input events rather than postspike [23,24] or the diffusion process response [25] of the membrane. Experimentally, some neurons’ firing probability and precision in response to excitatory synaptic input is enhanced by a preceding fast inhibition [26].

The membrane potential of the Hodgkin-Huxley model evolves according to the following equation [21,26]:

$$\frac{dV}{dt} = -I_N - I_e - I_{syn}$$

where $I_N = G_N m^3 h (V - E_N)$, $I_e = G_e E_e \delta(t - t_e) \exp(1 - \tau_e/(V - E_e)) H(t)$, and $I_{inh} = G_{inh} (V - E_{inh})$ are, respectively, sodium, potassium, and leakage currents through the membrane. The gating variables, activation $m$, and inactivation $h$ of the sodium current and activation $n$ of the potassium current, evolve according to the following equations [21,26]:

$$\frac{dm}{dt} = \alpha_m (1 - m) - \beta_m m$$

and

$$\frac{dn}{dt} = \alpha_n (1 - n) - \beta_n n$$

are voltage dependent, and are expressed via standard formu-
of 28% (24.7 Hz). For further increments in $G_{\text{inh}}$, the mean firing rate plateaus (explained later). In contrast, a suppression of the spike rate with $G_{\text{inh}}$ would occur if the membrane received a suprathreshold excitatory signal [shown in Fig. 1(b)]. The inhibition-induced enhancement of a firing rate for subthreshold excitatory inputs is the major result of our paper. The voltage reverse correlations, in the presence of inhibition, many of the spikes are preceded by a leading and well timed inhibitory arrival ($i$). The voltage reverse correlations, in the presence of inhibition, many of the spikes are preceded by a leading and well timed inhibitory arrival ($i$).

We present in Fig. 2(a) the relative contributions of these spike-causing pairs to the total spike rate. The effect of the $ee$ contribution decreases as expected in the presence of inhibition, but a prominent contribution is coming from $ie$ pairs. As the strength of inhibition is increased, more and more $ie$ pairs that were previously too far apart now become capable of causing spikes and the resultant contribution to the total spike rate increases. However, the number of such pairs is limited by the input arrival rate. Thus an increasing $G_{\text{inh}}$ recruits these pairs up to the maximum available number resulting in a saturation of the spike rate.

FIG. 2. (a) Breakdown of the total spike rate (normalized to control, 19.3 Hz) into four contributors. The $ie$ pairs contribute to most of the enhancement overcoming the $ee$ losses. (b) A precisely timed $ie$ pair can evoke a spike in a resting membrane: an excitatory input is given at $t=30$ ms, and an inhibitory input preceding it by $\delta$ ms ($G_{\text{inh}}=1$). ($G_{\text{ex}}=0.05$.)

A precisely timed $ie$ pair can evoke a spike in a resting membrane: an excitatory input is given at $t=30$ ms, and an inhibitory input preceding it by $\delta$ ms ($G_{\text{inh}}=1$). ($G_{\text{ex}}=0.05$.)
feedback in the system) of the membrane voltage above the rest level, \(V_{\text{rest}}\). The time \(t'\) at which \(V\) crosses \(V_{\text{rest}}\) upwardly depends continuously on \(\tau_{\text{inh}}\) [solid line in Fig. 3(b)] and nearly coincides with the mean of the \(ie\) relative timing distributions. The finite spread of these distributions is reminiscent of the finite width of the excitabile region shown in Fig. 2(b), and this latter region is shown here again [gray in Fig. 3(b)] but now as a function of \(\tau_{\text{inh}}\). The width and extent of this gray PIF region (and thus the relative \(ie\) timing distribution widths) can be controlled by the level of \(G_{\text{ex}}\). The relative hyperexcitability (i.e., effective threshold reduction), which is also reflected by the relative heights of the distributions, varies across the PIF window; it is dynamic and also depends on \(\tau_{\text{inh}}\). For example, at the point indicated by “+” (\(\tau_{\text{inh}} = 1\) ms, \(\delta = 6.5\) ms) a spike can be elicited with a \(G_{\text{ex}}\) value that is only 30% of the amount needed from rest.

What actual biophysical mechanism is behind the enhancement phenomenon? To address this question we reduce the full model equations to a two-variable model [37] described by \(\frac{V}{C} = -I_{\text{fast}}(V) - G_{\text{ex}} n^4 (V - E_K) - I_{\text{syn}} \), \(n = n_0 (1 - n) - \beta_n n\), where \(I_{\text{fast}}(V) = G_{\text{Na}} m^3 (V) h (V - E_{\text{Na}}) + I_{\text{L}}\). And \(n\) is the negative feedback variable. The rest state at \(-60\) mV is a focus, as in the full system, and as visualized in this phase plane (Fig. 4). In this reduced model, excitability corresponds to a spike-upstroke trajectory which is generated by a transient input that drives the \(V-n\) trajectory across the stable manifold (SMF) of the saddle. The SMF is the unique pair of phase plane trajectories (marked with double arrows in the figure) that enter the saddle point. Note that a SMF crossing could be evoked from a leftward-driving or rightward-driving stimulus. During a transient input, the \(V\) nullcline (i.e., the curve on which \(V=0\)) and the SMF move dynamically; then, after the stimulus, the phase point moves along the flow lines of the resting system. An inhibitory input that is faster than the intrinsic relaxation time scale perturbs the phase point with nullclines and SMF virtually returned to their “resting” positions. For inputs that would be subthreshold from rest, there still are two mechanisms by which a spike upstroke can be achieved by temporally sequencing such inputs. For a subthreshold inhibitory input, the trajectory returns to the focus in a spiral (solid curve). If during this return toward rest the brief excitation (which would be subthreshold if applied at rest) is delivered (\(\delta\) ms later, say at the point labeled \(\delta\) in Fig. 4 inset) when the trajectory is close to the SMF the phase point is pushed rightward across the SMF and a spike results. For a given strength of inhibition, as before, a range of such favorable times (marked by the thick portion on the solid curve) can be found that define the \(\delta\)-window where the spike threshold is effectively reduced and where PIF may occur. (\(G_{\text{ex}} = 0.05\), \(G_{\text{inh}} = 1\).)

FIG. 3. (Color online) \(\delta\)-window distributions and comparison with response to isolated \(ie\) pairs. (a) Distribution of relative timing of \(i\) with respect to \(e\) immediately preceding a spike onset for different \(\tau_{\text{inh}}\). (b) The mean values of these distributions are plotted (as filled circles) as a function of \(\tau_{\text{inh}}\). The gray region shows the parametric dependence of the width of hypersensitive region shown in 2(b). The solid curve distinguishes two subregions where \(V > V_{\text{rest}}\) (gray portion with \(\delta > t'\), see text) and where \(V < V_{\text{rest}}\) (\(\delta < t'\) following an isolated inhibitory input. See text for other markings.

FIG. 4. (Color online) Demonstration of PIF and PIR in a 2D reduced model. Three trajectories emerging from the rest state in the direction of arrows are shown corresponding to a lone inhibition delivered at \(t=0\) (solid curve, subthreshold oscillation), an inhibition (at \(t=0\)) followed by an identical inhibition delivered \(d=3\) ms later (dot-dashed curve, PIR), and an inhibition (at \(t=0\)) followed by an excitation delivered \(\delta=8\) ms later (dashed curve, PIF). The thick portion of the solid curve indicates the reduced threshold region in which PIF may occur. (\(G_{\text{ex}}=0.05\), \(G_{\text{inh}}=1\).)
nonresonator, but does not eliminate the counterclockwise flow or PIF.

Using biophysical Hodgkin-Huxley model equations, we have shown that brief inhibitory synaptic inputs that are usually associated with spike-rate suppression can, in fact, enhance the spike rate. The enhancement of firing probability stems from favorable temporal pairings of inhibitory inputs with subthreshold excitatory inputs. Such pairings will occur in any neuronal system that is subjected to random inputs. Of course, some neurons or models will show less (or more) hyperexcitability. But surprisingly, previous studies have not considered these per chance timing effects between random excitatory and inhibitory inputs on the output response. This could partly be due to the fact that studies of response to stochastic input are often carried out with more analytically tractable leaky integrate-and-fire (LIF) type one-variable models. The LIF, for example, disallows the PIF mechanism since it has no negative feedback variable that could be transiently reduced from rest in response to an inhibitory input. The LIF formalism may be insufficient to capture the full implications of fast inhibition and may have to be modified appropriately. Our results carry implications for the role of fast inhibition both in recurrent networks and in feedforward contexts, say, in sensory pathways. Random brief inhibition could upregulate the spontaneous firing of sensory neurons, many of which have high spontaneous activity. We are reporting separately [28] experimental in vitro evidence for PIF behavior in auditory brain stem neurons, in circuitry where inhibition can be quite fast. The coincidence detecting sensitivity of such neurons, that carry out the neural computation for sound localization, is shaped by fast inhibition [13]. Recent in vitro studies have revealed that fast inhibition can promote synchronizations and rhythmicogenesis among neurons in hippocampal circuits [31] and in the subthalamic nucleus [26]. In the latter case, the timing of inhibition before excitation was shown to be especially effective. Suggestions that the transient hyperpolarizing current that preceded depolarizing input contributes to spike firing have been made based on reverse correlation analysis of random inputs and firings [32–34]. Our own theoretical extensions of the PIF phenomenon to the network level with noisy inputs showed that PIF mediated the onset of synchrony as well as an increase of the network’s frequency over control levels [35]. Our findings of the enhancement to firing probability (by per chance PIF events) should be considered in seeking to interpret the roles by brief inhibition in such networks.


$m$ is fast and is well approximated by its steady state voltage dependent value of $m_{\infty}(V)$. $h$ is the lowest, and we set it equal to its rest value $h_0=0.596$. 

\[ C_m(=1 \mu F/cm^2) \] is the membrane capacitance. 
\[ G_{Na}=120.0 \text{ mS/cm}^2, \qquad G_K=36.0 \text{ mS/cm}^2, \] and \[ G_L=0.3 \text{ mS/cm}^2. \] The reversal potentials are \[ E_{Na}=55.0 \text{ mV}, \qquad E_K=-72.0 \text{ mV}, \qquad E_{Na}=49.387 \text{ mV}, \qquad E_{L}=-10.0 \text{ mV}, \qquad E_{Na}=70.0 \text{ mV}. \]