Physiological Gain Leads to High ISI Variability in a Simple Model of a Cortical Regular Spiking Cell

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Abstract

To understand the interspike interval (ISI) variability displayed by visual cortical neurons (Softky and Koch, 1993), it is critical to examine the dynamics of their neuronal integration as well as the variability in their synaptic input current. Most previous models have focused on the latter factor. We match a simple integrate-and-fire model to the experimentally measured integrative properties of cortical regular spiking cells (McCormick et al., 1985). After setting RC parameters, the post-spike voltage reset is set to match experimental measurements of neuronal gain (obtained from in vitro plots of firing frequency vs. injected current). Examination of the resulting model leads to an intuitive picture of neuronal integration that unifies the seemingly contradictory " $1/\sqrt{N}$ " and "random walk" pictures that have previously been proposed. When ISI's are dominated by post-spike recovery, $1/\sqrt{N}$ arguments hold and spiking is regular; after the "memory" of the last spike becomes negligible, spike threshold crossing is caused by input variance around a steady state, and spiking is Poisson. In integrate-and-fire neurons matched to cortical cell physiology, steady state behavior is predominant and ISI's are highly variable at all physiological firing rates and for a wide range of inhibitory and excitatory inputs.

1 Introduction

Softky and Koch (1993) have shown that the spike trains of cortical cells in visual areas V1 and MT display a high degree of variability, as measured by their interspike interval (ISI) distributions at fixed mean firing rates. Does this result necessitate revision of the classical notion of information transmission in cortical neurons? By "classical notion", we mean the view that neurons integrate many synaptic inputs until reaching some voltage threshold, and at that point produce a spike. Spikes are then followed by a refractory period, during which the cell is less likely to produce another spike. If inputs are uncorrelated, the mean spike rate of such a model neuron depends on the average frequency of presynaptic events, and thus can embody the standard notion of "rate coding."

Softky and Koch (1993) argued that high ISI variability is inconsistent with a neuron acting as an EPSP integrator. The integration of many uncorrelated excitatory synaptic potentials should result in a time to threshold (or ISI) that is very regular, since the neuron averages over many random events. One measure of spike variability is the coefficient of variation (CV) of the ISI distribution, defined as the standard deviation divided by the mean. For an EPSP integrator, this should be approximately $1/\sqrt{N}$, where N is the number of events necessary to reach threshold. In contrast, cortical cells have CVs in the range .5 – 1, more closely resembling a random (i.e. Poisson) process for which CV = 1. As an alternative to simple integration, Softky and Koch (1993) proposed that active dendritic conductances may act to amplify submillisecond coincidences in synaptic input, leading to large random pulses of synaptic current. Thus, they suggested that high ISI variability may be more consistent with the notion (Abeles, 1982) of neurons acting as "coincidence detectors" rather than "rate encoders."

Shadlen and Newsome (1994), following Gerstein and Mandelbrot (1964), argued that if uncorrelated synaptic inputs to a cell consist of balanced excitation and inhibition, then the membrane voltage follows a "random walk", leading to ISIs that are quite variable. This balanced inhibition model is consistent with the classical notion of synaptic integration and rate coding, but requires that excitation and inhibition be tightly balanced in order to achieve ISI variability consistent with the data.

These arguments can be more clearly understood by considering spike production as a two-step process. First, synaptic inputs are integrated by an extensive and complex dendritic tree resulting in a total synaptic current I(t). Second, the cell

produces spikes in response to this synaptic current. Thus, we expect that spike variability should depend on two factors: the sensitivity of the spike mechanism to changes in the synaptic current, and the variability in this input current.

Previous discussions of ISI variability have largely focused on the second term, that is, under what conditions can sufficient input variability be achieved? Thus, Softky and Koch (1993) argued that input current variability was too low to account for output spiking variability, while Shadlen and Newsome (1994) argued that a balance of excitation and inhibition could yield high input variability. Other models demonstrated that network dynamics can lead to correlations among synaptic inputs sufficient to yield high variability (Usher et al., 1994; Hansel and Sompolinsky, 1996) (N.B., afferent inputs also have significant correlations (Alonso et al., 1996)). However, these investigations generally did not address the sensitivity of cortical neurons. Neuronal sensitivity has been addressed by Bell et al. (1995), who considered some of the same factors as the present work in the context of a Hodgkin-Huxley model. Our work differs in considering simple integrate-and-fire dynamics, in focusing on the roles of gain and refractoriness, and, most importantly, in matching model parameters to experimental data from cortical neurons.

For the purposes of this article, we define neuronal gain as the slope of a plot of firing frequency f=1/ISI vs. the (constant) level of injected current I. While there is no direct relationship between this measure and neuronal sensitivity under physiological conditions — neuronal gain is measured in the absence of input variability — we will argue that the two should be well correlated. Here we demonstrate that matching a simple integrate-and-fire model to experimentally measured values of neuronal gain can largely solve the "conundrum" of high ISI variability in visual cortical cells with many random synaptic inputs. A careful examination of this model leads to a relatively simple, intuitive picture of cortical neuronal integration that unifies the seemingly contradictory " $1/\sqrt{N}$ " and "random walk" pictures that have been previously proposed.

A preliminary report of this study has appeared as an abstract (Troyer and Miller, 1995).

¹Bell et al. (1995) and Wilbur and Rinzel (1983) also investigated bistability in neuronal dynamics as a mechanism that could contribute to ISI variability.

2 The High-Gain Model

In an integrate-and-fire model, the slower, integrative properties of neurons are modeled as passive changes in subthreshold membrane voltage. The fast spiking conductances are lumped into a stereotyped event that is "pasted onto" the voltage trace when the cell reaches threshold. After spiking, there is an absolute refractory period during which the cell cannot spike, followed by a relative refractory period during which the cell's ability to spike is reduced. We will use refractoriness to refer to the combined effect of all processes occurring after a spike that make a cell less likely to respond to a given pattern of input current with a subsequent spike. Refractoriness can be defined quantitatively as a function r of the time t after a given spike as follows. Suppose a just-threshold DC current is applied at or before the spike time, t=0. The refractoriness, r(t), is the magnitude of a brief current pulse that, superimposed upon this ongoing DC current at time t, is just sufficient to elicit a spike. Note that this definition includes all factors that contribute to the cell's refractoriness, including sodium channel inactivation and the effects of other active conductances, as well as spike after-hyperpolarization.³ In the simplest models, refractory effects are modeled by waiting for a fixed, absolute refractory period after spike onset, and then resetting the voltage to a value significantly below threshold. Thus, a single parameter, the depth of the after-spike voltage reset, serves to model all of the elements contributing to the cell's relative refractoriness.⁴ More realistic models of the complex events following a cortical spike may deepen our understanding of cortical integration, but at present such models are experimentally poorly constrained, and can lose the clarity motivating the use of simple models.

²Assume a spike occurs at t=0, and that after an absolute refractory period t_{refract} the voltage is reset to V_{reset} . Let τ be the membrane time constant, g_{leak} the leak conductance, and Δt the duration of the brief current pulse. Beginning from V_{reset} at time $t=t_{\text{refract}}$, let $I_{t'}$ be the DC current that leads to a spike at time t' (so I_{∞} is the just-threshold DC current). Let $V_{t'}(t)$ be the time course of the voltage given $I_{t'}$. Then $r(t) \equiv \frac{\tau}{\Delta t} g_{\text{leak}} \left(V_t(t) - V_{\infty}(t) \right) = (I_t - I_{\infty}) (1 - e^{-(t - t_{\text{refract}})/\tau}) \frac{\tau}{\Delta t}$.

³After-hyperpolarization – the voltage state of the cell – is often regarded as affecting integration rather than refractoriness, with "refractoriness" reserved for processes that alter the cell's spike responses at a given voltage, e.g. spike threshold elevation. When considering a cell's reduced responsiveness to input currents, however, it is most useful to combine all factors contributing to the reduction.

⁴Alternatively, one may consider a two (or three) parameter model of refractoriness in which, instead of (or in addition to) post-spike voltage reset, there is a post-spike elevation of the shold that decays at some rate (reviewed in Tuckwell, 1988, Table 3.1).

We use a simple, conductance-based integrate-and-fire model:

$$C \frac{\partial V}{\partial t} = g_{\text{leak}}(V_{\text{rest}} - V) + I \tag{1}$$

Parameters were matched to slice recordings of regular spiking cells⁵ (values from McCormick et al., 1985, noted in parentheses): $V_{\rm rest} = -74~(73.6\pm1.5)~mV$; $1/g_{\rm leak} = 40~(39.9\pm21.2)~M\Omega; C = \tau g_{\rm leak}$ where $\tau = 20~(20.2\pm14.6)~msec$. The absolute refractory period was taken to be the spike width, $t_{\rm spike} = 1.75~(1.74\pm.41)~msec$. Spike threshold $V_{\rm thresh}$ was set 20 mV above $V_{\rm rest}$: $V_{\rm thresh} = -54~mV$. This leaves the after-spike reset voltage, $V_{\rm reset}$, as the only free parameter. This parameter is commonly set equal to $V_{\rm rest}$, but this choice lacks physiological justification.

We set V_{reset} to match the model's f-I curve to that observed physiologically, as reported in McCormick et al. (1985) (see Fig. 1A). That is, we set V_{reset} to match the physiologically observed neuronal gain.⁶ This yields $V_{\text{reset}} = -60 \text{ mV}$, 6 mV below threshold. Using the f-I curve to set V_{reset} points to the strong connection between refractoriness and neuronal gain. Once τ and g_{leak} are determined from biophysical measurements, neuronal gain is determined by the cell's refractory behavior. A cell that is more refractory requires more input current to spike at any given frequency, resulting in an inverse relationship between refractoriness and neuronal gain.⁷ Similarly, greater refractoriness implies weaker sensitivity to physiological inputs. This explains why we connect high gain measured with constant currents to high sensitivity under physiological conditions: for a given membrane time constant and input resistance, weaker refractoriness implies both higher gain and higher physiological sensitivity.

Synaptic input took the form of Poisson distributed delta function conductance changes, $I_{\text{syn}}(t) = \sum_k g_{\text{syn}} \delta(t - t_k) (V_{\text{syn}} - V)$, where t_k denotes the arrival time of the kth presynaptic spike. g_{syn} represents the total conductance, integrated over the time course of the synaptic event, and thus has units nS msec. g_{syn} was taken from an exponential distribution with mean \bar{g}_{syn} ; values greater than $4\bar{g}_{\text{syn}}$ were then reset to

⁵The term "regular spiking" denotes the class of cortical cells that respond to constant current injection *in vitro* with single spike firing and spike-rate adaptation (McCormick et al., 1985). These constitute most cortical excitatory cells. The term does not imply regularity of firing *in vivo*.

⁶Matching the slope of the model f-I curve at 100 Hz to the average gain reported by McCormick et al. (1985) for cortical regular firing cells gives $V_{\text{reset}} = -59.75 \text{ mV}$.

⁷Using the definitions of footnote 2, $r(t) = (I_t - I_\infty)(1 - e^{-(t - t_{\text{refract}})/\tau}) \frac{\tau}{\Delta t}$. Writing gain G(f) as a function of firing frequency f(G(f)) is the slope of the f-I curve at f), this yields $r(t) = (1 - e^{-(t - t_{\text{refract}})/\tau}) \frac{\tau}{\Delta t} \int_{f=0}^{f=1/t} df/G(f)$. Thus, high gain corresponds to low refractoriness.

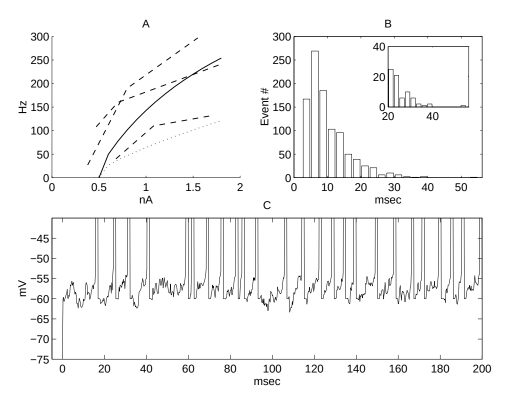


Figure 1:

A. f-I plots for our "high gain" model (solid) and experiment (dashed). For comparison, the dotted line represents the more common "low gain" model, in which $V_{\rm reset}$ is set to $V_{\rm reset}$. Experimental data shown are piecewise linear fits to data for first interspike interval from three cells (McCormick et al. 1985, fig 1). B. ISI histogram from 10 sec of model data: $\lambda_{\rm ex} = 7015~Hz$; $\lambda_{\rm in} = 3263~Hz~(R = .75)$; average firing rate = 98.4 Hz; CV = .6. Inset shows ISIs > 20 msec. C. Voltage trace of first 200 msec of simulation in B.

this maximum value.⁸ For excitatory synapses, $V_{\rm ex}=0mV$, and $\bar{g}_{\rm ex}=3.4~nS~msec$. This yields EPSPs at rest of mean .49 mV and maximum 2 mV. For inhibitory synapses, $V_{\rm in}=-70mV$, and $\bar{g}_{\rm in}=22.8~nS~msec$. This yields IPSPs that are twice as large as EPSPs at threshold.

The amount of inhibition was expressed as the ratio R of the mean inhibitory current to the mean excitatory current at threshold:

$$R = \frac{\lambda_{\rm in} \bar{g}_{\rm in} |V_{\rm in} - V_{\rm thresh}|}{\lambda_{\rm ex} \bar{g}_{\rm ex} |V_{\rm ex} - V_{\rm thresh}|}$$
(2)

Here λ expresses the mean rate of Poisson input. Note that R=1 implies that, when voltage is clamped at $V_{\rm thresh}$, the total synaptic current has mean zero; R<1 corresponds to a surplus of excitatory synaptic current at threshold, while R>1 corresponds to a surplus of synaptic inhibition. The leak current adds hyperpolarizing current. Estimates for $\lambda_{\rm ex}$ and $\lambda_{\rm in}$ rely on counting arguments only weakly constrained by experimental data (Shadlen and Newsome, 1994). Thus, to examine how model behavior depends on $\lambda_{\rm ex}$ and $\lambda_{\rm in}$, we fix R and then covary $\lambda_{\rm ex}$ and $\lambda_{\rm in}$ to yield varying output firing rates for that R. We have considered values of R ranging from 0 to 1.25; we will take R=.75 to be a reasonable guess for the ratio of inhibition to excitation in cortex.

Balanced inhibition models (Gerstein and Mandelbrot, 1964; Shadlen and Newsome, 1994) commonly assume equally likely positive and negative voltage steps, yielding a random walk in voltage. In a conductance-based model, this translates into a balance of inhibition and excitation sufficient to give a subthreshold mean current (so that there is not a steady drift up to near $V_{\rm thresh}$). We will roughly equate "balanced inhibition" with $R \geq 1$ (although smaller values are conceivable, since even for R = 1 the leak current maintains the mean voltage somewhat below threshold⁹).

3 Simulation Results

The results of a typical simulation (R = .75) are shown in figure 1. Postsynaptic average firing rate was 98 Hz for 10 sec of simulated data. We measured variability

⁸Thus the mean of the final distribution for $g_{\rm syn}$ is $(1 - e^{-4})\bar{g}_{\rm syn}$.

⁹The exact distance below threshold depends on the relative magnitudes of the leak and mean synaptic conductances. In the limit of high input rates, the leak is negligible and R=1 corresponds to a mean synaptic current just sufficient to elicit depolarization to threshold.

by taking the coefficient of variation (CV) of the ISI histogram, and found CV = .6, which falls in the physiological range of .5 to 1 reported in Softky and Koch (1993).

What expectations do we have for CV in these simple models? We expect CV to be bounded above by the CV of a Poisson process with dead time equal to the minimum recovery period. With mean ISI μ and dead time $t_{\rm dead}$, $CV_{\rm max} = (\mu - t_{\rm dead})/\mu$. For the simulation in Fig. 1, $t_{\rm dead} = t_{\rm spike}$ gives $CV_{\rm max} = .83$, while taking $t_{\rm dead}$ equal to the shortest interval observed (2.75 msec) yields $CV_{\rm max} = .73.^{10}$ A rough lower bound can be obtained from the case of pure EPSP integration. In our high gain model, it takes 16 average sized EPSPs to reach threshold from reset (recall that EPSPs are reduced by 3/4 near threshold). Again accounting for dead time of $t_{\rm dead} = t_{\rm spike}$, a $1/\sqrt{N}$ argument yields $CV = .83/\sqrt{16} = .2$. With variable sized PSPs, this estimate should be adjusted upward by a factor of roughly $\sqrt{2}$ yielding $CV \approx .28$ (Softky and Koch, 1993; Stein, 1967). The reason why $1/\sqrt{N}$ arguments fail to accurately estimate CV will be discussed shortly.

We measured the dependence of CV on postsynaptic firing rate by varying input rates while keeping the inhibition to excitation ratio fixed at R=.75 (Fig. 2A). Physiologically reasonable levels of CV are obtained across firing rates. Also, CV decreases at higher firing rates, as observed experimentally (Softky and Koch, 1993, Fig. 3). To test the notion that a balance of excitation and inhibition is required to achieve high variability (Shadlen and Newsome, 1994; Bell et al., 1995), we varied the inhibition ratio R, and observed ISI variability at input rates that yield postsynaptic firing rates between 95 and 105 Hz (Fig. 2B). To compare more closely with previous models that have used low gain, we also ran simulations with $V_{\rm reset} = V_{\rm rest}$. Balanced inhibition models are those with large R (e.g. R = 1.25), while the low gain model with excitation only (R = 0, marked "O") is similar to the simple EPSP integrator discussed in Softky and Koch (1993). The model with physiological (high) gain gives physiological CV over a broad range of R. In contrast, "balanced" inhibition (R > 1) is necessary to achieve CV > .5 in the low gain model.

¹⁰For a Poisson process with dead time, the shortest interval, the dead time, and the peak of the ISI distribution are the same.

 $^{^{11}\}mathrm{Shadlen}$ and Newsome (1994) used $V_{\mathrm{reset}}=V_{\mathrm{rest}}$ and hence considered a low gain balanced inhibition model.

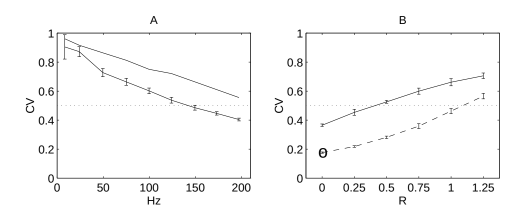


Figure 2:

A. CV vs. output firing rate, for fixed inhibition ratio R=0.75. Data shown are mean \pm standard deviation for 10 simulations (using different Poisson trains of inputs) of 10 sec duration. $\lambda_{\rm ex}$ ranged from 3396 Hz to 20,509 Hz, while $\lambda_{\rm in}$ correspondingly varied from 1274 Hz to 7691 Hz. Thin line shows CV for Poisson process with dead time equal to the shortest ISI observed. With increasing firing rate, the refractory period is a greater fraction of a typical ISI. Thus CV decreases with increasing rate. This effect is also seen in the biological data (Softky and Koch, 1993, Fig. 3). B. CV vs. inhibition ratio R for high gain (solid; $V_{\rm reset}=-60~mV$) and low gain (dashed; $V_{\rm reset}=V_{\rm rest}$) models. R=1.25 corresponds to example balanced inhibition models. Low gain with R=0 corresponds to EPSP integrator model (marked 'O'). At a given level of R, values of $\lambda_{\rm ex}$ and $\lambda_{\rm in}$ were found for which the 95% confidence interval for the mean spike rate was within the range from 95 to 105 Hz (10 simulations of 10 sec each). Data shown are from 10 subsequent trials at these input rates. $\lambda_{\rm ex}$ ranged from 4448 Hz to 19,584 Hz for high gain and from 7500 Hz to 23,261 Hz for low gain; $\lambda_{\rm in}$ varied from 0 Hz to 12,240 Hz (high gain) and from 0 Hz to 14,538 Hz (low gain). Dotted line shows CV=.5.

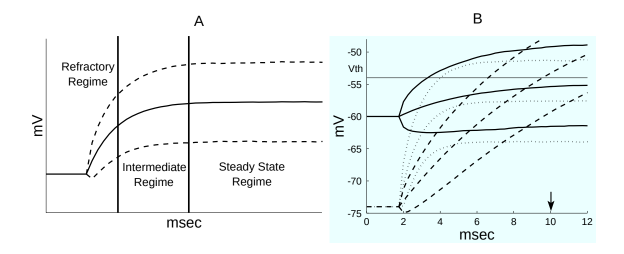


Figure 3:

Mean \pm standard deviation of the distribution of voltages. A. Schematic showing three regimes of behavior (see text). B. Data obtained from repeated (n=10,000) experiments starting the model (without spiking) from $V_{\rm reset}$. Rates set to achieve firing rates of 100 (± 5) Hz. Solid lines are from the high gain model ($V_{\rm reset}=-60~mV$; R=.75; $\lambda_{\rm ex}=8885~Hz$; $\lambda_{\rm in}=3332~Hz$). Broken lines are low gain models ($V_{\rm reset}=-74mV=V_{\rm rest}$). Dashed lines are from the EPSP integrator (R=0; $\lambda_{\rm ex}=7500~Hz$; $\lambda_{\rm in}=0~Hz$). Dotted lines are from a low gain, balanced inhibition model (R=1.25; $\lambda_{\rm ex}=23,261~Hz$; $\lambda_{\rm in}=14,538~Hz$). Arrow points to mean ISI.

4 An Intuitive Picture

An intuitive grasp of spike variability in simple integrate-and-fire models can be obtained by roughly dividing the ISI into three regimes (Fig. 3A; see also Abeles, 1991, Chap. 4; Smith, 1992). In the initial refractory regime, the state of the neuron is dominated by the recovery from the previous spike. In this regime, $1/\sqrt{N}$ arguments are valid (Smith, 1992) and spiking is regular. In the final, steady state regime, the memory of the last spike has decayed and random synaptic variation causes the cell to fluctuate about some steady state mean voltage (necessarily subthreshold). Thus, threshold crossings are equally likely to occur in any small interval and spike statistics are Poisson. In the intermediate regime, the mean voltage is still significantly rising, yet typical voltage fluctuations can be sufficient to cross threshold. Spike variability in this regime should be a mixture of $1/\sqrt{N}$ integration effects and Poisson random crossings. ISI variability depends on the relative amount of time spent in

these regimes.

As shown in Fig. 3B, integration in the EPSP integrator model is primarily in the refractory regime, and hence spiking is regular. In the low gain balanced inhibition model, large, transient changes in the balance of excitation and inhibition are large enough to dominate the cell's refractoriness rather quickly and the cell spends most of its time in the intermediate and steady state regimes. Note that this recovery is aided by a reduction of the membrane time constant due to the large synaptic conductance. The small reset of the high gain model causes the cell to spend most of the time in the intermediate or steady state regimes for virtually any input combinations that yield biologically reasonable firing rates.

Note that any mechanism that acts to reduce the importance of the transient, refractory regime contributes to variable spiking. In our model, $V_{\rm reset}$ is the dominant parameter determining the significance of the refractory regime. However, reducing the membrane time constant τ shortens transients and can emphasize steady state behavior. Increasing the variability of the synaptic current results in a more rapid transition to the intermediate regime and hence will also increase spike variability. Thus "balanced inhibition", which reduces τ and increases input variability, and "high gain" mechanisms are independent effects that both can contribute to variable spiking. It has been suggested (Bell et al., 1995) that spike variability depends on the cell "hovering" near threshold, so that the integration of only a few inputs can cause a spike. While this is similar to our explanation of steady state behavior, there is a significant difference. In the steady state regime, threshold crossings result from random fluctuations in the input current, so spiking is always Poisson, regardless of steady state voltage level. Thus, the nearness to threshold in the steady state regime affects spike rate, but does not directly affect spike variability.

5 Discussion

We have shown that matching simple integrate-and-fire neurons to experimental data results in ISI distributions with physiological CV. In additional studies, to be presented elsewhere, we have found that high CV is also produced by more realistic integrate-and-fire models incorporating the finite time course of synaptic conductances, realistic combinations of fast and slow excitatory synapses (AMPA and NMDA), and spike rate adaptation. Lengthening the synaptic time course can result

in input correlations longer than the shortest ISIs and lead to burst-like behavior that can actually increase CV; a realistic mixture of fast and slow synapses avoids such burstiness while achieving high CV.¹²

Does an understanding of the intermediate and steady-state regimes have implications for neural coding? In these regimes, both the mean synaptic current and the variance in this current contribute to neuronal spiking. A neuron operating in these regimes is certainly capable of rate coding: coding mean input rates with a mean output rate. When inputs are Poisson, the mean input rate of excitatory and inhibitory events determines both the mean and variance of the total synaptic current and consequently the mean output rate. However, such a neuron is also capable, in principle, of responding to input fluctuations with high temporal resolution. The possible contributions of these different forms of coding will depend on signal-to-noise considerations — how reliably can a given statistical attribute of the input be transmitted and be distinguished from random input fluctuations — that will depend on the details of the statistics of the neuronal input. Thus, the fact of high CV alone does not serve to distinguish between "rate coding" and "coincidence detection" or "spike timing" models of neural encoding.

One difficulty with high gain models is that they display low dynamic range, i.e. relatively small fluctuations in input current can lead to saturated outputs. In other studies (Troyer and Miller, 1995 and in preparation), we have found that spike rate adaptation helps to solve this problem by providing negative feedback that reduces gain on longer time scales, while preserving sensitivity to small input fluctuations on the time scale of typical interspike intervals. This fits well with the empirical observations that all excitatory cortical neurons show spike rate adaptation, while most inhibitory cortical neurons do not adapt but can sustain high rates of firing without saturating (McCormick et al., 1985).

Our high gain model results in ISI variability that falls in the lower half of the range .5 to 1 reported in Softky and Koch (1993). If one includes the refractory period, even a Poisson model can not account for the upper reaches of this data. The most likely explanation is that cortical ISI variability results from a combination of mechanisms (including high gain) that have generally been explored in isolation. However, ISI variability alone is inadequate to distinguish the relative contributions

¹²The burst-like behavior gives an unrealistically low CV2, a measure of variability that compares only adjacent interspike intervals (Holt et al., 1996). The mix of fast and slow synapses gives a realistically high CV2.

of these mechanisms to cortical integration. Stronger constraints may be found from experiments designed to probe the biophysical basis of integration and spiking in cortical cells, and from further statistical studies. The latter might examine, in vivo, the statistics of synaptically driven voltage and current fluctuations as well as spike train statistics beyond CV, and in vitro, the dependence of firing rate and CV on both mean and variance of white noise input current. One recent technique that may shed light on the contribution of cortical inhibition is the ability to pharmacologically block inhibitory input to a single cell (Nelson et al., 1994).

To understand cortical ISI variability, classical notions of neuronal integration must be modified, not by abandoning the integrate-and-fire neuron, but by deepening our understanding of the dynamics of such model neurons. In particular, the common intuition of dynamics dominated by recovery, with $1/\sqrt{N}$ integration behavior, must be supplemented by an understanding of a regime in which spike behavior is Poisson. Such a regime has been well studied in voltage-based integrate-and-fire models using a variety of statistical approaches (Gerstein and Mandelbrot, 1964; Stein, 1967; Smith, 1992; Tuckwell, 1988, chap. 9); in certain limits the regime can be described as an unbiased random walk of voltage with decay. The importance of such a random, steady-state regime for cortical processing has been discussed by several previous authors (Abeles, 1991; Shadlen and Newsome, 1994; Bell et al., 1995). Our contributions are to make specific links between refractoriness and neuronal gain, and to demonstrate that an integrate-and-fire model fitted to known cortical physiology will operate in the intermediate and steady-state regimes for physiologically reasonable firing rates and a wide range of ratios of inhibition to excitation. In these regimes, variable spike statistics dominate and hence the model produces physiological CV. Thus, in contrast to the intuition presented in Softky and Koch (1993), one should expect a high degree of ISI variability from cortical neurons. Of course, a full biophysical understanding of cortical integration may yet require abandonment of this simple view of cortical processing; but the fact of cortical ISI variability alone does not.

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References

Abeles M. (1982). Role of the cortical neuron: Integrator or coincidence detector? *Israel J. Med. Sci.*, 18:83–92.

Abeles M. (1991). Corticonics: Neural circuits of the cerebral cortex. Cambridge University Press.

Alonso J.M., Usrey W.M, and Reid R.C. (1996). Precisely correlated firing in cells of the lateral geniculate nucleus. *Nature*, 383:815–819.

Bell A.J., Mainen Z.F., Tsodyks M., and Sejnowski T.J. (1995). 'Balanced' conductances may explain irregular cortical spiking. Technical Report INC-9502, Institute for Neural Computation, Salk Institute.

Gerstein G.L. and Mandelbrot B. (1964). Random walk models for the spike activity of a single neuron. *Biophys. J.*, 4:41–68.

Hansel D. and Sompolinsky H. (1996). Chaos and synchrony in a model of a hypercolumn in visual cortex. J. Comput. Neurosci., 3:7–34.

Holt G.R., Softky W.R., Koch C., and Douglas R.J. (1996). A comparison of discharge variability in vitro and in vivo in cat visual cortex neurons. *J. Neurophysiol.*, 75:1806–14.

McCormick D.A., Connors B.W., Lighthall J.W., and Prince D.A. (1985). Comparative electrophysiology of pyramidal and sparsely spiny stellate neurons of the neocortex. *J. Neurophysiol.*, 54:782–805.

Nelson S., Toth L., Sheth B., and Sur M. (1994). Orientation selectivity of cortical neurons during intracellular blockade of inhibition. *Science*, 265:774–777.

Shadlen M.N. and Newsome W.T. (1994). Noise, neural codes and cortical organization. *Curr. Opin. Neurobiol.*, 4:569–579.

Smith C.E. (1992). A heuristic approach to stochastic models of single neurons. In McKenna T., Davis J., and Zornetzer S.F., editors, *Single Neuron Computation*, chapter 21, pages 561–588. Academic Press, Boston.

Softky W. and Koch C. (1993). The highly irregular firing of cortical-cells is inconsistent with temporal integration of random EPSPS. J. Neurosci., 13:334–350.

Stein R.B. (1967). The information capacity of nerve cells using a frequency code. *Biophys. J.*, 7:797–826.

Troyer T.W. and Miller K.D. (1995). A simple model of cortical excitatory cells linking NMDA-mediated currents, ISI variability, spike repolarization and slow AHPs. *Soc. Neuro. Abs.*, 21:1653.

Tuckwell H.C. (1988). *Introduction to Theoretical Neurobiology*. Cambridge University Press.

Usher M., Stemmler M., Koch C., and Olami Z. (1994). Network amplification of local fluctuations causes high spike rate variability, fractal firing patterns and oscillatory local field potentials. *Neural Comp.*, 6:795–836.

Wilbur W.J. and Rinzel J. (1983). A theoretical basis for large coefficient of variation and bimodality in neuronal interspike interval distributions. *J. Theor. Biol.*, 105:345–368.