HETEROGENEOUS FIRING RESPONSES PREDICT DIVERSE COUPLINGS TO PRESYNAPTIC ACTIVITY IN MICE LAYER V PYRAMIDAL NEURONS

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ABSTRACT

In the present study, we present a theoretical framework combining experimental characterizations and analytical calculus to capture the firing rate input-output properties of single neurons in the fluctuation-driven regime. We use this framework to investigate the functional impact of the heterogeneity in firing responses found experimentally in young mice layer V pyramidal cells. We first design and calibrate in vitro a simplified morphological model of layer V pyramidal neurons with a dendritic tree following Rall's branching rule. Then, we propose an analytical derivation for the membrane potential fluctuations at the soma as a function of the properties of the synaptic bombardment in dendrites. This mathematical description allows us to easily emulate various forms of presynaptic activities: either balanced, unbalanced, synchronized, purely proximal or purely distal synaptic activity. We found that those different forms of activity led to various comodulations of the membrane potential fluctuation properties, thus raising the question whether individual neurons might differentially couple to specific forms of activity because of their different firing responses. We indeed found such a heterogeneous response for all types of presynaptic activity. This heterogeneity was explained by different levels of cellular excitability in the case of the balanced, unbalanced, synchronized and purely distal activity. A notable exception appeared for proximal activity: increasing activity could either promote firing response in some cells or suppress it in some other cells whatever their individual excitability. This behavior could only be explained by various sensitivities to the speed of the fluctuations, which was previously associated to heterogeneous levels of sodium channel inactivation and density. Because local network connectivity targets rather proximal region, our results suggest that this biophysical heterogeneity might be relevant to neocortical processing by controlling how individual neurons couple to local network activity.

AUTHOR SUMMARY

Neocortical processing of sensory input relies on the specific activation of subpopulations within the cortical network. Though specific circuitry is thought to be the primary mechanism underlying this functional principle, we explore here a putative complementary mechanism: whether diverse biophysical features in single neurons contribute to such differential activation. In a previous study, we reported that, in young mice visual cortex, individual neurons differ not only in their excitability but also in their sensitivities to the properties of the membrane potential fluctuations. In the present work, we analyze how this heterogeneity is translated into diverse input-output properties in the context of low synchrony population dynamics. As expected, we found that individual neurons couple differentially to specific form of presynaptic activity (emulating afferent stimuli of various properties) mostly because of their differences in excitability. However, we also found that the response to proximal activity was controlled by the sensitivity to the speed of the fluctuations (which can be linked to various levels of density of sodium channels and sodium inactivation). Our study thus proposes a novel functional impact of biophysical heterogeneity: because of their various firing responses, individual neurons will differentially couple to local network activity.

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INTRODUCTION

The specific activation of subpopulations within neocortical networks appears to be the core mechanism for the cortical representation of sensory features. The details of how such specific activations happen are therefore key questions in systems neuroscience. As a primary source for specific activation, the neocortex is characterized by some degree of specific circuitry: neurons differ in their afferent connectivity. A classic example can be found in the primary visual cortex, layer IV simple cells specifically sample their input from ON and OFF cells in the thalamic nucleus [1,2]. Additionally, neocortical neurons also vary in their electrophysiological properties: for example, heterogeneous levels in the action potential threshold are routinely measured in vivo [3–5]. Thus, an emerging refinement is that the sensitivity of a neuron to a given feature do not only results from its stimulus specificity (e.g. orientation selectivity as a result of a specific afferent circuitry), but from the combination of its stimulus specificity and its biophysical specificity. Two somato-sensory cortex studies illustrates this point precisely. In Crochet et al. [3], during active touch, the spiking probability of a neuron (its sensitivity to whisker touch) follows from the combination of the reached level of synaptically-driven membrane potential deflection (its stimulus-specificity resulting from afferent circuitry, as quantified by post-synaptic reversal potentials) and its threshold for action potential triggering (its biophysical specificity). A similar result was found in the study of Yang et al. [5] for texture recognition, where the combination of those two quantities was shown to predict choicerelated spiking. Those results therefore suggest that heterogeneity in the biophysical properties of neocortical neurons might have an impact on their functional role during sensory processing.

In the present work, we further investigate the interaction between *stimulus specificity* and *biophysical specificity* in the light of the variability in the biophysical features reported in our previous study [6], namely that single neurons in juvenile mice cortex not only vary in their excitability (linked to the action potential threshold) but also in their sensitivity to the properties of the membrane potential fluctuations. Our previous communication introduced those new dimensions in the *biophysical specificity*, we here aim at understanding their functional impact. To this purpose, we implemented various stimuli onto layer V pyramidal cells (we varied the properties of presynaptic activity in the fluctuation-driven regime), and

we investigated whether individual neurons would differentially respond to those inputs because of their various firing rate response [6] (their *biophysical specificity*).

RESULTS

The results are organized as follows. We first formulate our theoretical framework for cellular computation which arbitrarily separate the question of dendritic integration and spiking probability. Then we present our simplified model of dendritic morphology which we calibrated on *in vitro* measurements. We derive an analytical approximation for the membrane potential fluctuations at the soma that we compare to numerical simulations using standard compartmental modeling. We implement various types of presynaptic activity and we analyze the individual responses of the layer V pyramidal cells (characterized in our previous study [6]).

A THEORETICAL FRAMEWORK FOR SINGLE CELL COMPUTATION IN THE FLUCTUATION-DRIVEN REGIME

Our study investigates the properties of single cell computation in the regime of low synchrony population dynamics [7,8] (the analogous, at the network level, of the fluctuation-driven regime at the cellular level) and aims at describing effects mediated by slow population dynamics $(T \ge 20-50 \text{ms})$. In this context, the cellular input-output function of a neocortical neuron corresponds to the function that maps the presynaptic variables to the spiking probability of the neuron. Our cellular model has five presynaptic variables (depicted in Figure 1A). Four of them are presynaptic firing rates (stationary release probabilities at the synapses) as those constitute the primary variables in this rate-based paradigm. To investigate their differential contribution, the proximal and distal parts of the dendritic trees have been separated and each of them has two presynaptic rates corresponding to the excitatory and inhibitory populations (hence four rate variables: v_e^p , v_i^p , v_e^d and v_i^d). The main motivation for this separation is to distinguish between two types of projections onto neocortical pyramidal neurons: synaptic inputs from the local network are thought to be more proximal while the distal apical tuft receives input from more distant cortical areas and thalamic locations [9]. Additionally, a global synchrony variable has been introduced for presynaptic events. This reproduces the effect of multi-innervation of a cell by its presynaptic afferent and, more importantly, the effect of pairwise

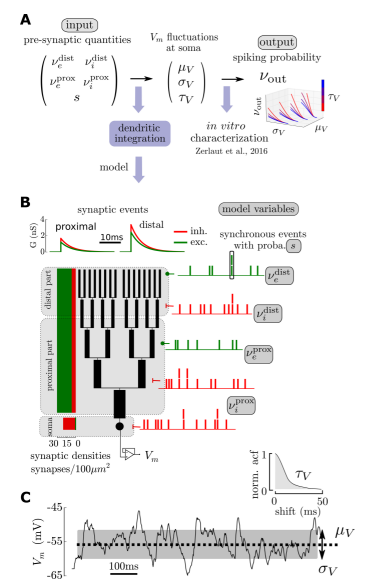


Figure 1. A theoretical framework for single cell computation in the fluctuation-driven regime. (A) Theoretical paradigm: to get the inputoutput function of a single cell, we split the relation from presynaptic quantities (the input) to the spiking probability (the output) into two steps. 1) passive dendritic integration shapes the membrane potential at the soma and 2) how those fluctuations are translated into spikes is captured by a firing response function determined in vitro. (B) Theoretical model for dendritic integration. A single cell is made of a lumped impedance somatic compartment and a dendritic tree. The dendritic tree is composed of B branches (here B=5), the branching is symmetric and follow Rall's 3/2 rule for the branch diameters. Synapses are then spread all over the membrane according to physiological synaptic densities. We define 3 domains: a somatic and proximal domain as well as a distal domain, excitatory and inhibitory synaptic input can vary independently in those domains. An additional variable: synaptic synchrony controls the degree of coincident synaptic inputs. (C) A given presynaptic stimulation (here $v_e^p = v_e^d = 0.2 Hz$, $v_i^p =$ $v_i^d = 1.2 \, Hz$ and s = 0.05) creates membrane potential fluctuations at the soma characterized by their mean μ_V , their amplitude σ_V and their autocorrelation time τ_V .

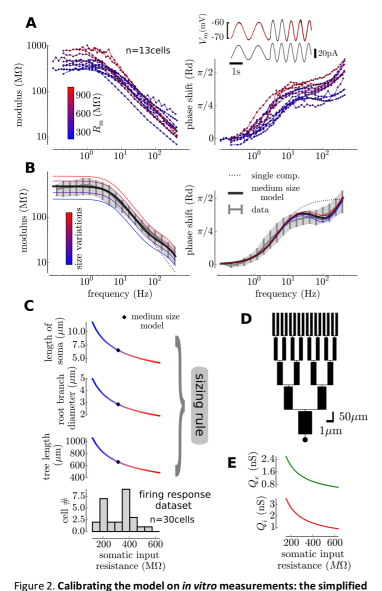
correlations associated to neocortical dynamics [10]. The synchrony degree in the presynaptic activity has been suggested to vary with stimulus statistics in the primary visual cortex [11,12] what motivates its introduction as a variable in our model.

To get the input-output function of a single cell, we split the relation from presynaptic quantities (the input) to the spiking probability (the output) into the two steps illustrated on Figure 1A: 1) passive dendritic integration shapes the membrane potential at the soma and 2) how those fluctuations are translated into spikes is captured by a firing response function determined *in vitro* [6]. The idea behind this approach rely on the fact that the action potentials are initiated at the axon initial segment [13] (i.e. electrotonically close to the soma) so that the fluctuations at the soma could determine the firing probability uniquely. We emphasize here that, given the complexity of neocortical cells, this approach only constitutes a first approximation of the cellular input-output function (see Discussion).

A SIMPLIFIED MORPHOLOGICAL MODEL FOR DENDRITIC INTEGRATION

The morphology of our theoretical model is a lumped impedance somatic compartment in parallel with a dendritic arborescence of symmetric branching following Rall's 3/2 branching rule (see Figure 1B and Methods). This morphology is of course a very reductive description of pyramidal cells: it does not discriminate between the distinct apical trunk and the very dense arborescence. Also, branching in pyramidal cell morphologies have been shown to deviate from Rall's 3/2 branching rule. Nonetheless this simplified model contains the important ingredient for our study: the fact that the transfer impedance to the soma of a synaptic input will strongly depend on its location on the dendritic tree. Indeed, as observed experimentally [14], distal events will be more low-pass filtered than proximal events in this model.

We spread synapses onto this morphology according to physiological densities [15] and describe synaptic events as transient permeability changes of ion-selective channels (see Methods). We arbitrarily separate the dendritic tree into two domains: a proximal and a distal domain (delimited by their distance to the soma, see Figure 1B). The distal part was taken as the last eighth of the dendritic tree to reproduce the large electronic distance to the soma characterizing distal synapses [14].



model and its size variations provides an approximation for the somatic input impedance of pyramidal cells and its heterogeneity over the recorded population. (A) Input impedance (left: modulus and right: phase shift) measured at the soma in intracellular recordings with sinewave protocols in current-clamp (inset). The color code indicates the input resistance and is likely to result from size variations of individual cells. (B) A medium size model accounts for the average data and varying the size of the dendritic tree and soma (according to the sizing rule shown in C) partially reproduces the variability in the individual measurements. Large cells (blue) have a lower modulus and a lower phase shift while small cells (red) have both a higher modulus and phase shift. (C) We obtain a map between input resistance and size of the morphological model. (D) Representation of the medium-size model. (E) Additionally the synaptic weights are rescaled with respect to the cell's somatic input resistance. Because the mean transfer resistance to soma is linked to the input resistance, this rescaling insures that the mean synaptic efficacy at soma is the same for all cells

Following experimental evidences [14], we set a higher synaptic efficacy for distal synapses. The synaptic parameters take physiological values [16] and can be found on Table 1. The passive and morphological parameters will be estimated from *in vitro* measurements.

MODEL CALIBRATION ON IN VITRO MEASUREMENTS

We will use the firing response function of our previous study, we therefore wanted a characterization of the passive and morphological properties on the same experimental system: layer V pyramidal neurons in the primary visual cortex of young mice. To this purpose, we performed measurements of the input impedance at the soma with intracellular recordings in vitro in n=13 cells. The key property on which this characterization relies is the fact that the input impedance at the soma cannot be accounted for only by the isopotential somatic compartment (i.e. a RC circuit). The input impedance also shows the contribution of the dendritic tree in parallel to the soma [17]. Indeed, both the modulus and the phase of the input impedance show deviations from the RC circuit impedance (see the comparison in Figure 22B): see for example the exponent of the power law scaling of the modulus (-1 exponent for the single compartment and ~-0.7 for pyramidal cells) or the decreased phase shift around 100Hz.

We first average all data (shown on Figure 2A) to obtain a mean input impedance (shown on Figure representative of a mean cellular behavior. We then performed a minimization procedure to obtain both the passive properties and the morphology corresponding to this average behavior (see Methods). The obtained passive properties were compatible with standard values, e.g. the resulting specific capacitance was 1.05 μ F/cm², close to the commonly accepted 1 μ F/cm² value, thus suggesting that the procedure could capture the physiological parameters of pyramidal cells, see Table 1 for the other parameters. Most importantly, the surface area was physiologically realistic, so that when using synaptic densities, we obtain an accurate number of synapses (see below). A representation of this mean morphology can be seen on Figure 2D.

Pyramidal cells show a great variability in input impedance, for example their input resistance almost spans one order of magnitude (both in the present n=13 cells, see the low frequency modulus values in Figure 2A, as well as in the firing response dataset, see bottom in Figure 2C). We found that varying the size of the

morphological model within a given range around the mean morphological model could partially reproduce the observed variability in the input impedance profiles (see Figure 2B). Size variations corresponds to a linear comodulation of the 1) tree length L_t , 2) the diameter of the root branch D_t and 3) the length of the somatic compartment L_{S} (see Figure $\,$ 2C for the range of their variations). On Figure 2A, the cells have been colored as a function of their input resistance while on Figure 2B, we vary the size of the size of the morphological model. Large cells (blue, low input resistance) tend to have a lower input resistance and phase shift than the small cells (red, high input resistance). Note that this simplistic account of morphological variations only very partially describes the observed behavior in pyramidal cells. In particular, 1) it strongly underestimates the variations of phase shifts at medium and high frequencies (f>20Hz) and 2) the relationship between size and impedance modulus at high frequencies (f>100Hz) is poorly captured. Those discrepancies are likely to be due to the details of dendritic arborescence that are not captured by the strong constraints of our dendritic model (symmetric branching, diameter rules, number of branches, etc...). Despite those discrepancies, size variations in our morphological model constitute a reasonable first approximation to account for cellular variety within the layer V pyramidal cell population.

This characterization, combined with the analytical tractability of the model (see Methods) allow us to construct a map between input resistance at the soma and size of the morphological model (the passive properties are set as identical, the one fitted on the mean impedance behavior). Thus, for each neuron in our previous "firing response dataset", because we have its input resistance at the soma, we can associate a given morphology. The association rule is shown in Figure 2C.

We now check what is the number of synapses obtained from the combination of our fitted morphologies with the physiological synaptic densities. We found a number of synapses of 3953 \pm 1748 (mean and standard deviation across the n=30 cells) with a ratio of excitatory to inhibitory numbers of synapses of 4.5 \pm 0.1. The fact that those numbers fall within the physiological range constitutes a validation of our approach (the morphology estimate through input impedance profile characterization).

AN ANALYTICAL APPROXIMATION FOR THE PROPERTIES OF THE MEMBRANE POTENTIAL FLUCTUATIONS AT THE SOMA

For each cell, we now want to translate the five variables of the model in terms of membrane potential fluctuations properties at the soma $(\mu_V, \sigma_V, \tau_V)$ on which we will apply the cell's firing response function. This constitutes the first step to obtain the final input-output function of individual cells (see Figure 1A).

Investigating dendritic integration detailed morphological structures is made difficult by the fact that this has to be done numerically with a relatively high spatial and temporal discretization. In the fluctuationdriven regime, one also needs to sample over long times $(T \gg \tau_V \sim 20 \text{ms})$ to obtain the statistical properties of the somatic V_m resulting from dendritic integration. addition, we have n=30 different morphologies in this study and we will explore a five dimensional parameter space (the five variable of our model). Under those conditions, if performed numerically, the computational cost of such a study is clearly prohibitive. We briefly describe here, why, in our simplified model, an analytical treatment is nonetheless possible and thus render this investigation feasible (see details in the Methods and in the Supplementary Material). The key ingredient is the ability to reduce the dendritic tree to an equivalent cylinder [17], we only adapted this reduction to the changes in membrane permeability associated to the high conductance state [18]. Two approximations underlie our estimation: 1) the driving force during an individual synaptic event is fixed to the level resulting from the mean bombardment [19] and 2) the effect at the soma of an synaptic event at a distance x in a branch of generation b, corresponds to the $\frac{1}{2^{b-1}}$ fraction of the post-synaptic response to the stimulation made of synchronous events at distance x in all the 2^{b-1} branches of the generation b. Luckily, the combination of those approximation is a favorable situation. Indeed, hypothesis 1) overestimates the size of post-synaptic events (because the driving force is not fixed, it diminishes during the PSP time course) while hypothesis 2) underestimates the size of post-synaptic events (because of the $2^{b-1} - 1$ synchronous events in neighboring branches, the membrane conductance is higher than in the case of a single event, consequently neighboring events have a shunting effect that artificially decreases the response). In addition, both of those approximation are likely to hold when single events are of

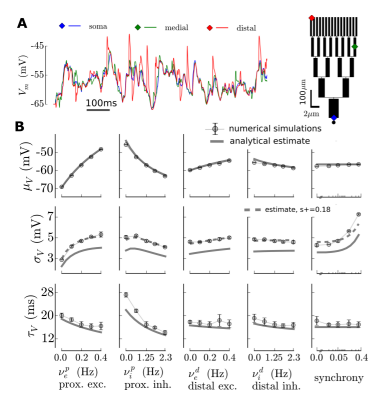


Figure 3. Accuracy of the analytical estimate for the properties of the membrane potential fluctuations at the soma: comparison between numerical simulations and the analytical approximation. Shown for the medium size model of Figure 2D. (A) In the numerical simulation, we explicitly simulate the whole dendritic arborescence, we show the membrane potential variations for the three locations shown on the left. (B) Properties of the membrane potential fluctuations (mean μ_V , standard deviation σ_V and autocorrelation time τ_V) for different configuration of presynaptic activity: analytical predictions and output from numerical simulations in *NEURON*. In each column, one variable is varied while the other variables are fixed to the mean configuration value corresponding to $v_e^p = v_e^d = 0.2$ Hz, $v_i^p = v_i^d = 1.2$ Hz and s=0.05. In the σ_V plots (middle panels, dashed gray lines), we added the prediction of the analytical estimate after a +0.18 correction for the synchrony (found with a Newton method).

low amplitude compared to the amplitude of the massive synaptic bombardment (see e.g. Kuhn et al. [19] for the validity of the first hypothesis).

In Figure 3, we compare the analytical approximation to the output of numerical simulations performed with the NEURON software [20]. We varied the five variables of the model around a mean synaptic bombardment configuration (see next section). Some discrepancies between the approximation and the simulations appeared, in particular one can see a $\sim\!1\text{mV}$ shift in the standard deviation σ_V of the fluctuation (meaning that single events are underestimated in the analytical treatment, so that hypothesis 2 is the most problematic

one). Because the synchrony controls the amplitude of the fluctuations (Figure 3B and next section), the analytical estimate could therefore be seen as an accurate estimate, modulo a shift in the synchrony (see Figure 3B, an increase of 0.18 in the synchrony corrects for the $\sim 1 \text{mV}$ shift in σ_V). Importantly, the trend in the variations of the fluctuations as a function of the model variables is globally kept between the analytical estimate and the numerical simulations. This relatively good agreement therefore shows that our analytical estimate is a valid tool to study dendritic integration in the fluctuation-driven regime.

PROPERTIES OF THE FLUCTUATIONS FOR DIFFERENT TYPES OF PRESYNAPTIC ACTIVITY

We now implement various types of presynaptic activity and investigate the properties of the resulting membrane potential fluctuations at the soma. In addition, we represent the variations of the somatic input conductance (relative to the leak input conductance) because, as it is routinely measured in intracellular studies in vivo, this quantity allows a comparison between the model and experimentally observed activity levels. On Figure 4, we present those different protocols, on the left (panel A), one can see how the five variables of the model are comodulated for each protocol (color coded, see bottom legend) and on the right (panel B), one can see the resulting properties of the membrane potential fluctuations. We present those results only for the medium-size model, but it was calculated for the morphologies associated to all cells. The variability introduced by the various morphologies is shown in Figure S2 and we found that the qualitative behavior discussed in this section was preserved in all cells.

We first introduced a baseline of presynaptic input corresponding to a low level of network activity: $\frac{g_{tot}^{soma}}{g_L} \sim 1.7, \text{ compared to } \sim 3\text{-}4 \text{ in activated states, reviewed in } [18]. This baseline activity is a mix of proximal and distal activity with a low degree of synchrony (<math>s$ =0.05). Similarly to Kuhn et al. [19], the inhibitory activity is adjusted to obtain a balance of the V_m fluctuations at -55mV. The firing values of this baseline level are very low $v_e^d = v_e^p = 0.2$ Hz for the excitation and $v_i^d = v_i^p = 1.2$ Hz for the inhibition) in accordance with the sparse activity characterizing mammalian neocortical dynamics [3,10]. On top of this non-specific background activity, we will now add a specific stimulation.

We start with unbalanced activity. We define it as a stimulation that brings the mean membrane potential above -55mV corresponding to the previously defined balance. The stimulation corresponds to an increase of the excitatory synaptic activity (still running within a very sparse range of activity, $v_e^d = v_e^p \in [0.05, 0.5]$ Hz) with an increasing inhibitory activity adjusted to linearly disrupt the balance between -55mV and -52mV (see Figure 4). The synchrony is kept constant and the activity indifferentially raises in the proximal and distal part. This increase of total activity raises the input conductance ratio close to four. In this moderate range, the variations of the amplitude of the fluctuations σ_V remains a monotonic increase (unlike the non-monotonic variations found in the singlecompartment study of Kuhn et al. [19] and the case of a proximal stimulation, see below), the fluctuations gets approximately twice faster (the normalized autocorrelation time $\frac{\tau_V}{\tau_m^0}$ decays from 100% to 50%) and, of course (by design), the mean depolarization has a linear increase of 3mV.

We now emulate purely proximal activity. To this purpose, we fix the distal presynaptic firing frequencies (v_e^d and v_i^d) as well as the synchrony to their baseline levels. To remain in a sparse activity level, we increase the proximal excitatory activity from the baseline level to 1.7Hz and we adjust proximal and somatic inhibitory activity to keep the balance at the soma. This would nonetheless correspond to large network activity level, as can be seen from the input conductance ratio (that raises up to 8). This situation gives results comparable to the single-compartment study of Kuhn et al. [19]. The amplitude of the fluctuations has a non-monotonic profile and the autocorrelation time strongly decreases. A notable difference is that, even if we investigated high activity levels, the autocorrelation time does not go to zero and the amplitude of the fluctuations has only a moderate decrease. This discrepancy is due to 1) the choice of non-negligible synaptic time constants compared to the membrane time constants (here $au_{syn} = 5 \text{ms}$ and $au_m^0 \sim 25 \text{ms}$, then au_V / au_m^0 would saturate at au_{syn}/ au_m^0 = 20% and 2) the fact that the synaptic input is distributed attenuates the strong shunting effects observed in the single compartment case.

For distal activity, we keep the proximal presynaptic frequencies (v_e^p and v_i^p) as well as the synchrony to their baseline levels. We increase the distal excitatory activity v_e^d from the baseline level to a moderate level: 0.7Hz. The distal inhibitory frequency v_i^d is again adjusted to keep

the balance at the soma. Here, we get a different picture than in the proximal case, the increase in activity leads to negligible increase of the somatic input conductance as expected from electrotonically distant input [21]. Also, the decrease of the speed of the fluctuations is much attenuated. The reason for this phenomenon is that only the distal part has a high conductance, consequently postsynaptic events are strongly low-pass filtered by the proximal part of the arborescence before reaching the soma. Here, the amplitude of the fluctuations strongly increases as a function of the input and do not show the non-monotonic relation found for proximal input. This is explained by the combination of the fact that 1) distal events are of higher amplitude and 2) the shunting of post-synaptic events is much reduced due to the relatively narrow localization of the synaptic conductances.

Finally, we emulate an increase in the presynaptic synchrony. Here, all synaptic frequencies are kept constant with respect to the baseline level and we simply increase the probability of coincident events for each synaptic spike train. Because there is no change in synaptic activity, this stimulation does not affect the input conductance ratio, neither the mean membrane potential or the speed of the fluctuations. However, presynaptic synchrony strongly affects the amplitude of the fluctuations in a near linear manner.

Note, that in addition to the sparse activity constraints or the balance constraints, the criteria for the ranges of the model variables was manually chosen to have the fluctuations in the same domain. For example, we investigated a lower activity range for the distal part (variations of v^{dist}) than for the proximal part (variations of v^{prox}) to avoid an explosion of σ_V , the range for the synchrony increase followed the same criteria.

HETEROGENEOUS FIRING RESPONSES INDUCE DIVERSE COUPLING TO PRESYNAPTIC ACTIVITY

For each one of the n=30 cells of our previous study [6], we now have 1) a morphological model and 2) a firing response function $v_{out} = \mathcal{F}(\mu_V, \sigma_V, \tau_V)$. Thanks to the analytical approximation, we can translate the five model variables $(v_e^p, v_i^p, v_e^d, v_i^d, s)$ into the stationary fluctuations properties $(\mu_V, \sigma_V, \tau_V)$ that, in turn, the firing response translate into a spiking probability. Thus, we finally get the full *input-output* function (within our theoretical framework) as illustrated on Figure 1A.

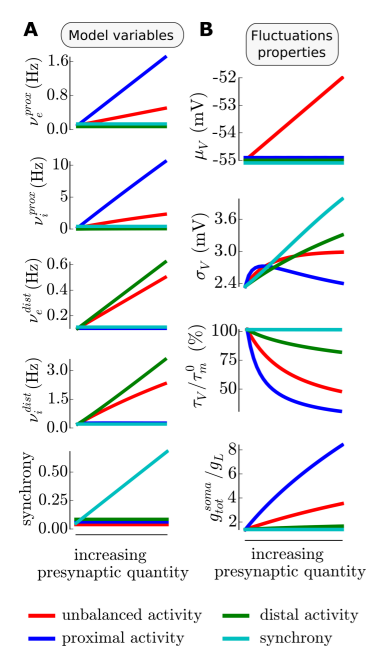


Figure 4. Properties of the membrane potential fluctuations for various types of presynaptic activity: either unbalanced (red), purely proximal (blue), purely distal (green), synchronized (cyan). A common baseline configuration of balanced proximal and distal activity at low rate gives rise to baseline fluctuations properties, on top of this, the increase of a given type of presynaptic activity corresponds to a given comodulations of the 5 model variables. (A) Comodulations of the model variables to achieve varying levels of the different types of activity. (B) Membrane potential fluctuations properties (mean μ_V , standard deviation σ_V and autocorrelation time τ_V) and somatic input conductance at the soma for the different protocols. Shown for the medium-size model, see Supplementary Material for the variability introduced by variations in cell morphologies.

We show on Figure 5 the response of four cells to the different types of presynaptic activity described in the previous section (those four cells were chosen as they were representative of different firing response behaviors, see Figure 5 and 6B in [6]). The input-output relationships show qualitative and quantitative differences, we briefly discuss them here and we perform a more rigorous analysis on the full dataset in the next section.

First, we can see that individual cells have a very different level of response to the baseline level of synaptic activity (initial response in Figure 5). Cell 1 has a baseline at $\sim 10^{-2}$ Hz while Cell2 or Cell3 have response above 1Hz, i.e. two orders of magnitude above.

Importantly, those cells have different preferences for particular types of stimulations. Cell 1 responds more to unbalanced activity whereas Cell 2 and Cell 4 respond more to an increase in synchrony and Cell 3 responds preferentially to proximal activity (within this range). This is what we mean by *preferential coupling*: individual neurons will respond preferentially to a particular type of synaptic activity. An even more pronounced discrepancy appears for proximal activity: the response can be either

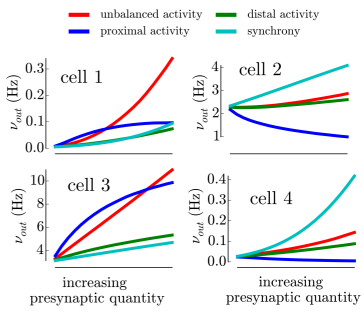


Figure 5. Examples of the firing response of 4 different cells for the various types of presynaptic activity (color-coded) shown in Figure 4. The abscissa "increasing synaptic quantity" corresponds to the comodulations of the model variables shown in Figure 4A (same color code). As an example, the response to a "proximal activity increase" (blue curve) corresponds to a linear increase of v_e^{prox} and v_i^{prox} while keeping v_e^{dist} , v_i^{dist} and s to their baseline level. See Figure 4A for the comodulations of the other types of increasing activity.

increased (Cell 1 and Cell 3) or decreased (Cell 2 and Cell 4) with respect to the baseline level.

Given the relative invariance of the fluctuations properties for each cell (see previous section and Figure S2, despite the various morphologies, the same input creates the same fluctuations), those differences are the consequences of the heterogeneity in the firing responses, we conclude that heterogeneous firing responses induce diverse coupling to presynaptic activity.

BIOPHYSICAL ORIGIN OF THE HETEROGENEOUS COUPLINGS TO PRESYNAPTIC ACTIVITIES

We now make this analysis more quantitative by computing the responses for all n=30 cells. We get their response to the baseline level ν_{bsl} and their mean response change for each stimulation type (the mean over the range of scanned presynaptic input): $\delta\nu_{ubl}$ for the unbalanced activity, $\delta\nu_{prox}$ for the proximal activity, $\delta\nu_{dist}$ for the distal activity and $\delta\nu_{synch}$ for an increased synchrony. We show the histogram of those values in the left column of Figure 6. We also look for the origin of the individual couplings by correlating them with the characteristics of the neuronal firing responses [6].

We first analyze the response to baseline activity v_{bsl} . When log-scaled (Figure 6A), the distribution is approximately normal and spans 2-3 orders of magnitude. This log-normal distribution of pyramidal cell firing rates during spontaneous activity seems to be a hallmark of mammalian neocortical dynamics (see e.g. [10] in human neocortex). We investigated what properties of the firing responses could explain this behavior, we therefore looked for correlations between our measures of the firing responses in the fluctuation-driven regime [6] and the baseline responses. Not surprisingly, we found a very strong linear correlation between the excitability $\langle V_{thre}^{eff} \rangle_{D}$ and the baseline response level, the other characteristics do not have an impact (Pearson correlations, see values in Figure 6A). Again, it should be stressed that presynaptic connections are non-specific, those results therefore show that the typical log-normal distribution of firing rates could very naturally emerge as a result of the normal distribution observed in pyramidal cell's excitabilities [6], thus suggesting that no specific circuitry might be needed to explain this neocortical property.

Despite the important differences in the fluctuations they create (see Figure 6B), the responses over cells to unbalanced activity, distal activity and an increased synchrony share a very similar behavior. First, those stimuli produce systematically (over all cells) an increase in firing rate. The firing increase again show a strong heterogeneity over cells, covering two orders of magnitude (see log y-axis on Figure 6B,D,E). This variability in responses was again highly correlated with the excitability. Surprisingly, the response was not dependent on any other of the characteristics of the firing response. For example, because synchrony controls the standard deviation σ_V , the variability observed during an increase in synchrony could have been linked to the to the sensitivity to the standard deviation $\langle \frac{\partial v}{\partial \sigma_V} \rangle_D$, but this effect was not significant (see Pearson correlation in Figure 6E). For those three protocols, none of the sensitivities to the fluctuations properties had a strong impact on the individual cellular responses (c<0.4 and p>0.01, Pearson correlations, see Figure 6). This analysis therefore revealed that, for those type of synaptic activities, those properties of the firing response have negligible impact compared to the very strong effect of the variability in excitabilities (see Discussion).

The response to proximal activity also showed a great variability but with a qualitatively different behavior (Figure 6C). Notably, firing could be suppressed or increased. This variability was independent of the excitability of the cells but was correlated with the sensitivity to the speed of the fluctuations $\langle \frac{\partial v}{\partial \tau_V^N} \rangle_D$. Indeed, the proximal stimulation implies a strong variations of the fluctuations speed (i.e. decreasing τ_V , while keeping moderate variations of σ_V and, by design, a constant μ_V) thus rendering the sensitivity to the fluctuation speed the critical quantity for this stimulation type. Those results therefore show that the response to proximal activity of an individual cell is controlled by its level of sensitivity to the speed of the fluctuations (see Discussion).

Discussion

In the present work, we introduced an analytically tractable description of single cell computation in the fluctuation-driven regime for neurons with a dendritic structure. We used it to investigate how the heterogeneity found experimentally in firing responses to fluctuating input shape the diverse input-output functions of neocortical pyramidal cells. Focusing on the regime of

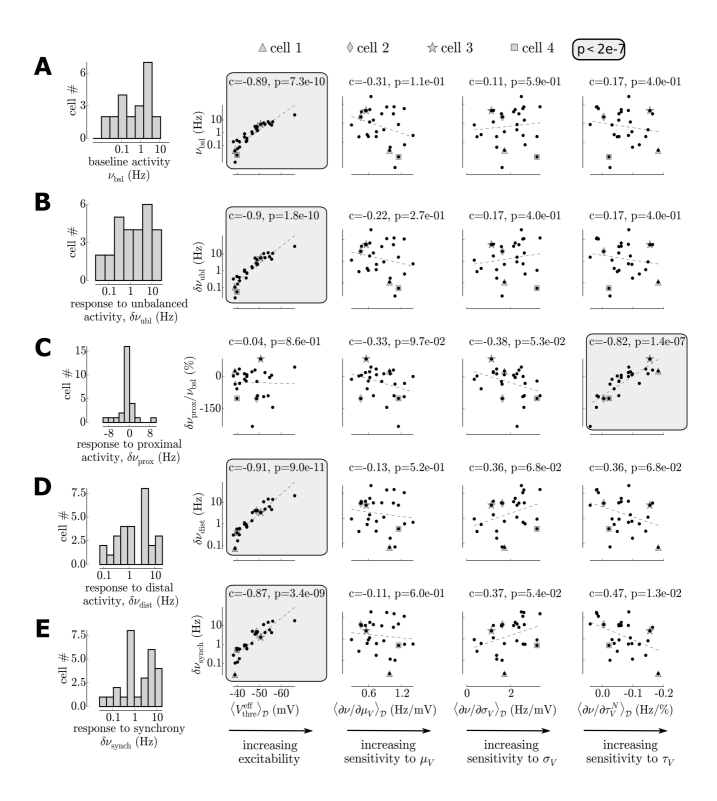


Figure 6. Diverse cellular responses to the various types of presynaptic activity and their link to the characteristics of their firing response function. Note the logarithmic scale for the firing responses in B,C,D. (A) Diverse response to baseline stimulation. (B) Diverse response to unbalanced activity. (C) Diverse response to proximal activity. Note that because the response also show negative changes of firing rate, the data cannot be log-scaled. Instead, they have been rescaled by the baseline response (i.e. we show $\frac{\delta v_{prox}}{v_{bsl}}$). (D) Diverse response to distal activity. (E) Diverse response to a synchrony increase.

near asynchronous population dynamics, we emulated various types of presynaptic activity in a theoretical model of dendritic integration. We showed that those different types of synaptic stimulation correspond to various comodulations of the fluctuations properties. This property is what motivated to fully scan the three dimensional space $(\mu_V, \sigma_V, \tau_V)$ in our previous study [6] instead of the response to a given presynaptic input type that corresponds to an arbitrary comodulation of $(\mu_V, \sigma_V, \tau_V)$. Importantly, we found that, because of their different response to the same fluctuations, individual neurons would respond differently to those stimulations.

Very naturally, a key quantity to explain the various neuronal responses was the cellular excitability. Indeed, the response to unbalanced activity, distal activity or an increase in synchrony was strongly correlated with cellular excitability. Also the sensitivity to the speed of the fluctuations had a crucial impact on the response to proximal activity. In our previous communication [6], theoretical modeling suggested that a high sensitivity to the speed of the fluctuations was enable by a high level of sodium inactivation (as only fast fluctuations allow to deinactivate sodium channels) and a high density of sodium channels (as it corresponds to a sharp spike initiation mechanism that enables to extract fast fluctuating input, reviewed in [22]). The present analysis thus proposes an important functional role for those biophysical properties: controlling the coupling to proximally targeting activity.

The present analysis also sheds light on the physiological relevance of the properties that we introduced in our previous study. In all four measures characterizing the cellular firing rate response function in the fluctuationdriven regime (shown at the bottom of Figure 6), only two of them seem to be physiologically relevant: the excitability $\langle \, \textit{V}^{eff}_{thre} \rangle_{D} \,$ and the sensitivity to the speed of the fluctuations $\langle rac{\partial
u}{\partial au_V^N}
angle_D$. The reason for the lack of functional impact of the sensitivity to depolarizations $\langle \frac{\partial v}{\partial u_D} \rangle_D$ and to the sensitivity to amplitude of the fluctuations $\langle \frac{\partial v}{\partial \sigma_V} \rangle_D$ was that they were estimated independently of the excitability (after rescaling by the different excitability levels). While this separation was useful to isolate and evidence the contribution of different biophysical mechanisms to the firing rate response function [6], the effect of a change in depolarization or an increase in amplitude of the fluctuations on the firing response of a neuron will actually be mainly led by the excitability level whereas those more subtle variations in the firing rate response function were shown to have a negligible impact.

The proposed theoretical framework for single-cell computation nonetheless suffers from several weaknesses. First, even if our description captures the various electrotonic distances associated to various synaptic locations (the crucial ingredient here to discriminate between proximal and distal inputs), the morphological model appears as a very poor description of layer V pyramidal cell. Deviations from the symmetric branching hypothesis and Rall's branching rule will have a significant impact on dendritic integration in the fluctuation-driven regime. To investigate those effects within the framework proposed in our study, one could benefit from the large body of theoretical work on the derivation of Green's function for arbitrary branched passive dendritic trees [23-26]. Another important limitation of our description lies in the absence active mechanisms in dendrites [27]. It is therefore a question how much those mechanisms could affect the picture provided in our study. Preliminary numerical analysis performed in presence of NMDA and Ca²⁺ currents [28] (see Figure S4 and Figure S5), showed that, provided excitation balances inhibition (a situation where NMDA channels keeps a relatively low level of stationary activation) and provided synchrony do not reach a too high level (for s≥0.4 excitatory events almost systematic lead to NMDA spikes), the qualitative behavior of the cellular input-output function remains unaffected. Our preliminary analysis thus suggest the hypothesis that active dendritic mechanisms may not be crucial for effects mediated by slow ($T \ge 20-50$ ms) variations of weakly synchronous population dynamics while being critical for operations performed on synchronous synaptic activations (T≤5-10ms), consistent with the current view on dendritic processing [29]. Future work should address this hypothesis in details.

Despite the current weaknesses of our description, we believe that having an analytical model for dendritic integration in the fluctuation-driven regime is a useful tool for many problems in theoretical neuroscience. The main advantage of this model is that you can very naturally plug in physiological parameters (because surface area as well as transfer resistance to soma can take physiological

values) and while still allowing an analytical treatment (though see deviations of the approximations in Figure 3). In the theoretical analysis of neural network dynamics, the literature is almost exclusively based on the reduction to the single-compartment (reviewed in [30]). Though being approximate, our framework thus opens the path toward a detailed mathematical analysis of recurrent network dynamics containing neurons with extended dendritic structures.

Additionally, it must be stressed that formulating the cell response with the somatic fluctuations properties $(\mu_V, \sigma_V, \tau_V)$ as an intermediate variable is very powerful because it allows one to apply the same measurements to various models. For example, with a single-compartment model it is easy to translate these variables into excitatory and inhibitory activities [19]. We showed here that it is also possible to obtain relations with synaptic inputs occurring in dendrites in a simplified morphological neuron model. The latter model uses the same measurements, so no experiments need to be redone. We could in principle also apply the same approach to more complex models and obtain more realistic transfer functions. This phenomenological two-step procedure thus offers a flexible complementary approach to the analytical approaches tackling the problem of the spiking behavior in presence of an extended dendritic structure [31-33].

Finally, we speculate about a putative link between a recent observation and the present findings. The in vivo study in mice visual cortex of Okun and colleagues [4] reported a strong heterogeneity in the coupling between individual cell's responses and the locally recorded population activity. The authors explained those observations by a variability in the local recurrent connectivity and found that this diverse coupling did not seem to be explained by biophysical heterogeneity (e.g. the coupling was independent from the action potential threshold, somehow a measure of the excitability). However, as local connectivity is thought to target more proximal regions such as the basal dendrites, our study proposes a biophysical mechanism that could also contribute to their observation. The diverse coupling to proximal activity was here explained by the sensitivity to the speed of the fluctuations, and similarly to their results, this coupling was found to be independent on the cellular

excitability (see Figure 6B in Zerlaut et al. [6]). Part of their results could therefore be explained by this mechanism. Also, even if this electrophysiological heterogeneity disappears in mature phenotypes, the preferential coupling present in young animals could be amplified by long term plasticity to form this strongly coupled local network. Future work could therefore address this hypothesis by combining recordings of population activity with a subtle and functionally-relevant analysis of single cell properties.

MFTHODS

ETHICS STATEMENT

Experiments were performed at Unité de Neurosciences, Information et Complexité, Gif sur Yvette, France. Experimental procedures with animals were performed following the instructions of the European Council Directive 2010 86/609/EEC and its French transposition (Décret 2013/118).

MORPHOLOGICAL MODEL

The morphology of our theoretical model is the following (depicted in Figure 1B): it is made of an isopotential somatic compartment (i.e. a leaky RC circuit) in parallel with a dendritic structure. The dendritic tree is an arborization of total length l_t containing B generation of branches. For simplicity all branches of a generation $b \in [1,B]$ have a length $\frac{l_t}{B}$. From one generation to the other, a branch divides into two branches where the diameter of the daughter branches follows Rall's 3/2 branching rule [17] $(d_{b+1})^{\frac{3}{2}} = \frac{(d_b)^{\frac{3}{2}}}{2}$, i.e. $d_b = 2^{-\frac{2}{3}}\,d_t$ where d_t

branching rule [17] $(d_{b+1})^2 = \frac{e^{-b^2}}{2}$, i.e. $d_b = 2$ 3 d_t where d_t is the diameter of the root branch of the dendritic tree. Excitatory and inhibitory input are then spread homogeneously over the soma and dendritic tree according to the densities of synapses \mathcal{D}_i^{soma} , \mathcal{D}_e^{soma} , \mathcal{D}_e^{tree} , \mathcal{D}_i^{tree} . The parameters of the model are presented on Table 1.

MODEL EQUATIONS: SYNAPTIC INPUT AND PASSIVE PROPERTIES

The cable equation describes the temporal evolution and spatial spread of the membrane potential along the branches of the dendritic tree [17]:

$$\frac{1}{r_i} \frac{\partial^2 v}{\partial x^2} = i_m(v, x, t)$$

$$= c_m \frac{\partial v}{\partial t} + \frac{v - E_L}{r_m}$$

$$- i_{syn}(v, x, t)$$
(1)

the membrane current $i_m(v,x,t)$ is a linear density of current (the presented cable equation already includes the radial symmetry, i.e. $r_m = \frac{1}{G_L\pi D}, r_i = \frac{4\,R_i}{\pi\,D^2}, c_m = C_m\pi\,D$, where G_L,R_i,C_m are the passive membrane parameters, see Table 1). Though the modeled system has several branches, the equation can be written as a single spatial dependency x because the symmetry of the model across branches imply that the properties of the input are identical at a given distance to the soma.

Synaptic input is modeled by local (infinitely small) and transient changes of membrane permeability to selective ionic channels. Both excitatory (accounting for AMPA synapses) and inhibitory synapses (accounting for GABAa synapses) are considered, their reversal potential is E_e =0mV and E_i =-80mV respectively. Each synaptic event is generated by a point process and its effect on the conductance is an increase of a quantity $Q_{s\in\{e,i\}}$ followed by an exponential decay $\tau_{s\in\{e,i\}}$. The form of the synaptic current is therefore:

$$\begin{cases} i_{syn}(v,x,t) = g_e(x,t)(E_e - v) + g_i(x,t)(E_i - v) \\ g_e(x,t) = \sum_{\{x_e,\{t_e\}\}} \delta(x - x_e) \sum_{\{t_e\}} \mathcal{H}(t - t_e) Q_e(x) e^{-\frac{t - t_e}{\tau_e}} \\ g_i(x,t) = \sum_{\{x_i,\{t_i\}\}} \delta(x - x_i) \sum_{\{t_i\}} \mathcal{H}(t - t_i) Q_i(x) e^{-\frac{t - t_i}{\tau_i}} \end{cases}$$
 (2)

where g_e and g_i are linear densities of conductances. Each synapse, indexed by s, has a position x_s and a set of presynaptic events $\{t_s\}$, hence the iteration over $\{x_s, \{t_s\}\}$ for the sum over synapses for each synaptic type. $\mathcal H$ is the Heaviside step function. The presynaptic events $\{t_s\}$ are generated by point processes at fixed frequencies $\{v_s\}$ with a given degree of synchrony, see details in the next section.

The model distinguishes two domains : a proximal domain $(x \in [0, l_p])$ with the upper index p and a distal domain $(x \in [l_p, l])$ with the upper index d (see Figure 1), where l_p is the length of the somatic compartment and l is the total length of the dendritic tree. The space-dependent quantities (presynaptic frequencies, synaptic quantal and synaptic decay time constant) can be written as:

$$\begin{cases} v_{e}(x) = v_{e}^{P} + (v_{e}^{d} - v_{e}^{p}) \mathcal{H}(x - l_{p}) \\ v_{i}(x) = v_{i}^{P} + (v_{i}^{d} - v_{i}^{p}) \mathcal{H}(x - l_{p}) \\ Q_{e}(x) = Q_{e}^{P} + (Q_{e}^{d} - Q_{e}^{p}) \mathcal{H}(x - l_{p}) \\ Q_{i}(x) = Q_{i}^{P} + (Q_{i}^{d} - Q_{i}^{p}) \mathcal{H}(x - l_{p}) \end{cases}$$
(3)

The continuity of the membrane potential and of the current at the boundaries between the proximal and distal part imply:

$$\begin{cases} v(l_p^-, t) = v(l_p^+, t) \\ \frac{1}{r_i} \left(\frac{\partial v}{\partial x}\right)_{l_p^-} = \frac{1}{r_i} \left(\frac{\partial v}{\partial x}\right)_{l_p^+} \end{cases}$$
(4)

where the limit with upper index \pm indicate the limit taken from the left or the right respectively.

At the soma, x=0, we have a lumped impedance compartment. It has leaky RC circuit properties and also receives synaptic inhibition, the somatic membrane potential therefore follows:

$$\begin{cases} C_{M} \frac{dV}{dt} + \frac{V - E_{L}}{R_{M}} + G_{I}(t)(V - E_{i}) + I(t) = 0 \\ \\ G_{i}(t) = \sum_{N_{i}} \sum_{\{t_{i}\}} Q_{i}^{p} e^{-\frac{(t - t_{i})}{\tau_{i}^{p}}} \mathcal{H}(t - t_{i}) \end{cases}$$

where I(t) is the time-dependent input current from the soma into the dendrite. R_M and C_M are the RC properties of the lumped compartment (capital letters will indicate the somatic properties throughout the calculus). N_i is the number of somatic synapses, each of them generates a point process $\{t_i\}$ of inhibitory synaptic events. The properties of the somatic synapses v_i^S , Q_i^S are equivalent to the proximal ones.

This equation with the membrane potential continuity will determine the boundary condition at the soma (x=0). We identify V(t)=v(0,t), then I(t) is the current input into the dendritic tree at x=0 so it verifies:

$$\left(\frac{\partial \mathbf{v}}{\partial \mathbf{x}}\right)_{\mathbf{x}=\mathbf{0}} = -r_i I(t)$$

So:

$$\left(\frac{\partial v}{\partial x}\right)_{x=0} = r_i \left(C_M \left(\frac{\partial v}{\partial t}\right)_{x=0} + \frac{v(0,t) - E_L}{R_m} + G_I(t) \left(v(0,t) - E_i\right)\right)$$
(5)

Finally, the last boundary condition is that all branches terminate with an infinite resistance that impede current flow (sealed-end boundary conditions):

$$\left(\frac{\partial \mathbf{v}}{\partial \mathbf{x}}\right)_{\mathbf{x}=\mathbf{1}} = 0 \tag{6}$$

Together with the biased Poisson process for event generation (see the next section), the final set of equations that describes the model, is therefore the combination of Equations 1, 2, 3, 4, 5, 6.

MODEL OF PRESYNAPTIC ACTIVITY

Presynaptic activity is modelled as a discrete set of presynaptic events. Because of the apparent random spiking activity in the fluctuation-driven regime, the basis for the generation of those discrete events is the Poisson process. Nonetheless, we want to reproduce the additional degree of presynaptic synchrony found in neocortical assemblies that result mainly for two phenomena: 1) pairwise correlation between neurons and 2) multiinnervation of a cell by a presynaptic neuron. We therefore introduce a variable s that will bias the event generation of the Poisson process ($s \in [0,1]$). For simplicity in the analytical treatment, synchrony in presynaptic activity is not shared across different synapses. We arbitrarily limit the number of coincident events to four events, therefore for a degree of synchrony s: single events have a probability 1-s, double events have a probability $s - s^2$, triple events have a probability $s^2 - s^3$ and quadruple events have a probability s^3 . To generate a biased Poisson process of frequency ν with a degree of synchrony s, we therefore generate a Poisson process of frequency:

$$v_{\text{synch}} = \frac{v}{1 + s + s^2 + s^3 3} \tag{7}$$

and we duplicate (from up to four events) each event according to their probabilities of occurrence.

This is a very simplistic and limited model of presynaptic synchrony but it is sufficient for reproducing the impact of synchrony on the quantities investigated in this paper (only the variance of the membrane potential fluctuations).

NUMERICAL IMPLEMENTATION

The full model has been implemented numerically using the *NEURON* software [20]. The branched morphology was created and passive cable properties were introduced (see Table $_1$). The spatial discretization was nseg=30 segments per branch. On each

segment, one excitatory and one inhibitory synapse were created, the shotnoise frequency was then scaled according to the segment area and the synaptic density to account for the number of synapses on this segment (using the properties of the Poisson process, N synapses at frequency ν is a synapse at frequency $N\nu$). Custom event generation was implemented to introduce correlations (instead of classical NetStim) and fed NetCon objects attached to each synapses (ExpSyn synapses). Each simulation had a time step dt=0.01ms and a length of 10s, the simulation was repeated over 4 seeds to yield a mean and a standard deviation in the estimate of the membrane potential fluctuations at the soma (see Figure 3B).

ANALYTICAL DERIVATION OF THE FLUCTUATION PROPERTIES: STRATEGY

We present here a derivation that provides an analytical approximation for the properties of the fluctuations of the membrane potential at the soma for our model. Summing up its properties, we get: 1) a morphology with a lumped somatic compartment and a dendritic tree of symmetric branching following Rall's rule 2) conductance-based synapses 3) independent excitatory and inhibitory shotnoise input spread all over the morphology 4) asymmetric properties between a proximal part and a distal part and 5) a certain degree of synchrony in the pre-synaptic spikes.

The properties of the membrane potential fluctuations at the soma correspond to three stationary statistical properties of the fluctuations: their mean μ_V , their standard deviation σ_V and their global autocorrelation time τ_V . Following [6], we emphasize that the global autocorrelation time is a partial description of the autocorrelation function (as the autocorrelation function is not exponential) but it constitutes the first order description of the temporal dynamics of the fluctuations.

A commonly adopted strategy in the *fluctuation-driven* regime to obtain statistical properties is to use stochastic calculus after having performed the *diffusion approximation*, i.e. approximating the synaptic conductance time course by a stochastic process [34]. This approach is nonetheless not easily generalizable to conductance input in an extended structure and render the inclusion of asymmetric properties (proximal vs distal) complicated. We rather propose here an approach that combines simplifying assumptions and analytical results from point process theory, it extends the approach proposed in Kuhn et al. [19] to dendritic structures following Rall's branching rule.

For each set of synaptic stimulation $\{v_p^e, v_i^p, v_e^d, v_i^d, s\}$, the derivation corresponds to the following steps:

I. We transform the dendritic structure to its equivalent cylinder. The reduction to the equivalent cylinder is "activity-dependent" and captures the changes in

membrane properties that results from the mean synaptic conductance levels.

- II. We derive a mean membrane potential $\mu_V(x)$ corresponding to the stationary response to constant densities of conductances given by the means of the synaptic stimulation. We use this space-dependent membrane potential $\mu_V(x)$ to fix the driving force all along the membrane for all synapses. The relation between synaptic events and the membrane potential now becomes linear.
- III. We derive a new cable equation that describes the variations of the membrane potential around this $\mu_V(x)$ solution.
- IV. We calculate the effect of one synaptic event on a branch b, $b \in [1,B]$ at a distance x. We calculate the post-synaptic membrane potential event $PSP_b(x,t)$ at the soma resulting from b synchronous synaptic events occurring at the distance x from the soma. We approximate the effect of only one event by rescaling the response by the number of input $\frac{PSP_b(x,t)}{b}$.
- V. We use shotnoise theory to compute the power spectrum density of the membrane potential fluctuations resulting from all excitatory and inhibitory synaptic events (including the synchrony between events).

The full derivation has been conducted with the help of the python modulus for symbolic computation: *sympy*. The resulting expression were then exported to *numpy* functions for numerical evaluation. Details are provided in Text S1

FROM FLUCTUATION PROPERTIES TO SPIKING PROBABILITY

How layer V pyramidal neurons translate membrane potential fluctuations into a firing rate response was the focus of our previous communication [6]. We re-use here the same dataset, i.e. the individual characterizations over n=30 single neurons of the firing responses $v_{out} = \mathcal{F}(\mu_V, \sigma_V, \tau_V)$.

Importantly, our study introduced four quantities to describe the mean properties of a single neuron response in the fluctuation-driven regime: (i) a measure of the excitability given by the mean phenomenological threshold $\langle V_{thre}^{eff} \rangle_{\mathcal{D}}$ of the firing response function, (ii) a sensitivity to the mean of the fluctuations $\langle \frac{\partial v_{out}}{\partial \mu_{V}} \rangle_{\mathcal{D}}$, quantifying how much a mean depolarization is translated into a change in firing rate, (iii) a sensitivity to the amplitude of the fluctuations $\langle \frac{\partial v_{out}}{\partial \sigma_{V}} \rangle_{\mathcal{D}}$, quantifying how much an increase of the standard deviation of the fluctuations is translated into a change in firing rate, and (iv) a sensitivity to the speed of the fluctuations

 $\langle \frac{\partial v_{out}}{\partial \tau_V^N} \rangle_D$, quantifying how much a change in the autocorrelation time of the fluctuations is translated into a change in firing rate.

Note that in the main text, we removed the n=3/30 cells that had too low excitabilities, as it is hard to conduct an analysis on 6 orders of magnitudes, $\nu \in [10^{-5}, 10^1]$ Hz (for data visualization). This reduction limited the output variations to $\nu \in [10^{-2}, 10^1]$ Hz. In the supplementary figure S 3, we reintroduce the discarded cells and we show that they do not affect the results presented in the main text.

EXPERIMENTAL PREPARATION AND ELECTROPHYSIOLOGICAL RECORDINGS

Experimental methods were identical to those presented in [6]. Briefly, we performed intracellular recordings in the current-clamp mode using the perforated patch technique on layer V pyramidal neurons of coronal slices of juvenile mice primary visual cortex. For the n=13 cells presented in this study, the access resistance R_S was $13.3 \mathrm{M}\Omega \pm 5.4$, the leak current at -75mV was -25.7pA ± 17.3 , cells had an input resistance R_m of $387.3 \mathrm{M}\Omega \pm 197.2$ and a membrane time constant at rest of $32.4 \mathrm{ms} \pm 23.1$.

INPUT IMPEDANCE CHARACTERIZATION

To determine the input impedance at the soma, we injected sinusoidal currents in the current-clamp mode of the amplifier (Multiclamp 700B, Molecular Devices), we recorded the membrane potential response to a current input of the form $I(t) = I \sin(2\pi f t)$, we varied the frequencies f and amplitudes I over 40 episodes per cell. The frequency range scanned was [0.1, 500] Hz. For each cell, we determined manually the current amplitude I_0 that gave a \sim 5mV amplitude in a current step protocol, from this value, the value of I was scaled exponentially between I_0 at 0.1 Hz and 50 I_0 at 500Hz. The reason for varying the current amplitude (and not only the oscillation frequency) in those input impedance protocols is to anticipate for the low pass filtering of the membrane and insure that the membrane potential response at high frequencies is far above the electronic noise level \sim 0.1 mV.

After removing the first 3 periods of the oscillations (to avoid transient effects), we fitted the membrane potential response to the form: $V(t) = E_L + R \, I \, sin(2\pi f t - \phi)$, where E_L, R, ϕ were fitted with a least-square minimization procedure. The frequency dependent values of R and ϕ give the modulus and phase shift of the input impedance presented in Figure 2A.

FITTING PASSIVE PROPERTIES AND A MEAN MORPHOLOGY

Because the variables combined discrete (the branch number) and continuous variables, the minimization consisted in taking the minimum over a grid of parameters. The parameter space had 7 dimensions: the branch number ($B \in [2,7]$), the somatic length ($l_S \in [5,20]\mu$ m), the total length of the tree ($l_t \in [1,20]\mu$)

 $[300,800]\mu$ m), the diameter of the root branch ($d_t\in[0.5,4]\mu$ m), the leak specific resistance ($r_m\in100,1000]\mu$ S/cm²), the intracellular resistivity ($r_i\in[10,90]\,\Omega$.cm), the specific capacitance ($c_m\in[0.8,1.8]\,\mu\text{F/cm}^2$). Each dimension was discretized in 5 points, the scan of the 7 dimensional space then consisted in finding the least square residual of the product of the modulus and phase of the impedance over this 5^7 points. The resulting parameters are shown on Table 1.

REFERENCES

- Hubel DH, Wiesel TN. Receptive fields, binocular interaction and functional architecture in the cat's visual cortex. J Physiol. 1962;160: 106–154. doi:10.1113/jphysiol.1962.sp006837
- Ferster D, Chung S, Wheat H. Orientation selectivity of thalamic input to simple cells of cat visual cortex. Nature. 1996;380: 249–252. doi:10.1038/380249a0
- Crochet S, Poulet JFA, Kremer Y, Petersen CCH. Synaptic mechanisms underlying sparse coding of active touch. Neuron. Elsevier Inc.; 2011;69: 1160–1175. doi:10.1016/j.neuron.2011.02.022
- Okun M, Steinmetz N a., Cossell L, lacaruso MF, Ko H, Barthó P, et al. Diverse coupling of neurons to populations in sensory cortex. Nature. 2015;521: 511– 515. doi:10.1038/nature14273
- Yang H, Kwon SE, Severson KS, O'Connor DH. Origins of choice-related activity in mouse somatosensory cortex. Nat Neurosci. 2015;19: 127–134. doi:10.1038/nn.4183
- Zerlaut Y, Teleńczuk B, Deleuze C, Bal T, Ouanounou G, Destexhe A. Heterogeneous firing rate response of mice layer V pyramidal neurons in the fluctuation-driven regime. J Physiol. 2016;13: 3791–3808. doi:10.1113/JP272317
- Ecker AS, Berens P, Keliris GA, Bethge M, Logothetis NK, Tolias AS. Decorrelated neuronal firing in cortical microcircuits. Science. American Association for the Advancement of Science; 2010;327: 584–7. doi:10.1126/science.1179867
- Renart A, Rocha J De, Bartho P, Hollender L, Parga N, Reyes A, et al. The asynchronous state in cortical circuits. Science. 2010;327: 587–90. doi:10.1126/science.1179850
- 9. Spruston N. Pyramidal neurons: dendritic structure and synaptic integration. Nat Rev Neurosci. 2008;9: 206–21. doi:10.1038/nrn2286
- 10. Peyrache A, Dehghani N, Eskandar EN, Madsen JR,

- Anderson WS, Donoghue J a, et al. Spatiotemporal dynamics of neocortical excitation and inhibition during human sleep. Proc Natl Acad Sci U S A. 2012;109: 1731–6. doi:10.1073/pnas.1109895109
- Baudot P, Levy M, Marre O, Monier C, Pananceau M, Frégnac Y. Animation of natural scene by virtual eyemovements evokes high precision and low noise in V1 neurons. Front Neural Circuits. 2013;7: 206. doi:10.3389/fncir.2013.00206
- 12. El Boustani S, Marre O, Béhuret S, Baudot P, Yger P, Bal T, et al. Network-state modulation of power-law frequency-scaling in visual cortical neurons. PLoS Comput Biol. 2009;5: e1000519. doi:10.1371/journal.pcbi.1000519
- Debanne D, Campanac E, Bialowas a., Carlier E, AlcarazG. Axon Physiology. Physiol Rev. 2011;91: 555–602.doi:10.1152/physrev.00048.2009
- 14. Magee JC. Dendritic integration of excitatory synaptic input. Nat Rev Neurosci. 2000;1: 181–190. doi:10.1038/35044552
- DeFelipe J, Fariñas I. The pyramidal neuron of the cerebral cortex: morphological and chemical characteristics of the synaptic inputs. Prog Neurobiol. 1992;
- Destexhe A, Mainen ZF, Sejnowski TJ. Kinetic Models of Synaptic Transmission. 1998.
- 17. Rall W. Electrophysiology of a dendritic neuron model. Biophys J. The Biophysical Society; 1962;2: 145.
- Destexhe A, Rudolph M, Paré D. The high-conductance state of neocortical neurons in vivo. Nat Rev Neurosci. Nature Publishing Group; 2003;4: 739–751. doi:10.1038/nrn1289
- Kuhn A, Aertsen A, Rotter S. Neuronal integration of synaptic input in the fluctuation-driven regime. J Neurosci. Soc Neuroscience; 2004;24: 2345–56. doi:10.1523/JNEUROSCI.3349-03.2004
- 20. Hines ML, Carnevale N. The NEURON simulation environment. Neural Comput. 1997;9: 1179–209.
- Koch C, Douglas R, Wehmeier U. Visibility of synaptically induced conductance changes: theory and simulations of anatomically characterized cortical pyramidal cells. J Neurosci. Soc Neuroscience; 1990;10: 1728–44.
- 22. Brette R. What Is the Most Realistic Single-Compartment Model of Spike Initiation? PLoS Comput Biol. 2015;11: e1004114.

doi:10.1371/journal.pcbi.1004114

- 23. Abbott LF, Farhi E, Gutmann S. The path integral for dendritic trees. Biol Cybern. 1991;66: 49–60.
- 24. Cao BJ, Abbott LF. A new computational method for cable theory problems. Biophys J. 1993;64: 303–313. doi:10.1016/S0006-3495(93)81370-5
- Bressloff PC, Coombes S. Physics of the Extended Neuron. Int J Mod Phys B. 1997;11: 2343–2392. doi:10.1142/S0217979297001209
- 26. Coombes S, Timofeeva Y, Svensson C-M, Lord GJ, Josić K, Cox SJ, et al. Branching dendrites with resonant membrane: a "sum-over-trips" approach. Biol Cybern. 2007;97: 137–149. doi:10.1007/s00422-007-0161-5
- 27. London M, Häusser M. Dendritic computation. Annu Rev Neurosci. 2005;28: 503–532. doi:10.1146/annurev.neuro.28.061604.135703
- 28. Larkum ME, Nevian T, Sandler M, Polsky A, Schiller J. Synaptic integration in tuft dendrites of layer 5 pyramidal neurons: a new unifying principle. Science (80-). 2009;325: 756–760. doi:10.1126/science.1171958
- Stuart GJ, Spruston N. Dendritic integration: 60 years of progress. Nat Neurosci. 2015;18: 1713–1721.

doi:10.1038/nn.4157

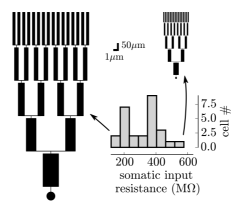
- Renart A, Brunel N, Wang X-J. Mean-field theory of irregularly spiking neuronal populations and working memory in recurrent cortical networks. Comput Neurosci A Compr approach. Boca Raton, CRC Press; 2004; 431–490.
- 31. Svensson C-M, Coombes S. Mode Locking in a Spatially Extended Neuron Model: Active Soma and Compartmental Tree. Int J Bifurc Chaos. 2009;19: 2597–2607. doi:10.1142/S0218127409024347
- 32. Saparov A, Schwemmer MA. Effects of passive dendritic tree properties on the firing dynamics of a leaky-integrate-and-fire neuron. Math Biosci. 2015;269: 61–75. doi:10.1016/j.mbs.2015.08.014
- Schwemmer MA, Lewis TJ. Bistability in a Leaky Integrate-and-Fire Neuron with a Passive Dendrite.
 SIAM J Appl Dyn Syst. 2012;11: 507–539. doi:10.1137/110847354
- 34. Tuckwell HC, Wan FYM, Rospars J-P. A spatial stochastic neuronal model with Ornstein--Uhlenbeck input current. Biol Cybern. Springer; 2002;86: 137–145.

SUPPORTING INFORMATION

TEXT S1

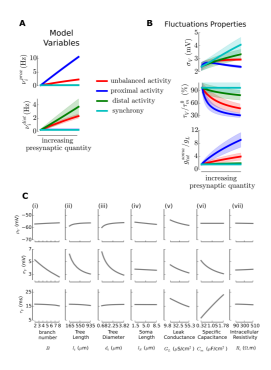
Supplementary material contains the detailed mathematical derivation of the membrane potential fluctuations at the soma as a function of the model variables (see PDF added at the end of the manuscript).

FIGURE S1



Heterogeneity in morphologies. Graphical representation of the estimated morphologies of the largest (the lowest input resistance) and the smallest (the highest input resistance) cells present in the dataset.

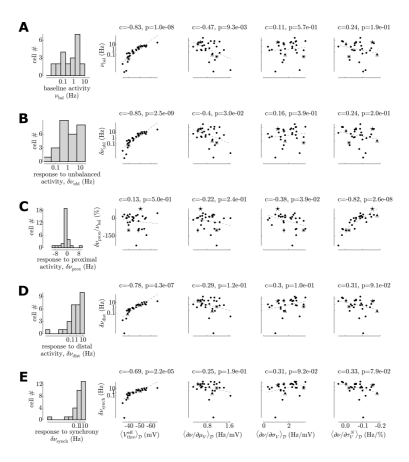
FIGURE S2



Variability in the fluctuations properties introduced by different morphologies. (A) Variability in the model variables across protocols introduced by the different morphologies (sizes). We show only the quantities that vary across cells, all other variables (v_e^p , v_e^d , s) are fixed across cells. The balance μ_V is adjusted for each cells and the cells have different surfaces, so different number of synapses (and especially different ratio of excitatory to inhibitory numbers) hence the need to adjust

inhibitory activity slightly differently for each cell. (B) Variability in the properties of the membrane potential fluctuations (standard deviation σ_V and autocorrelation time τ_V) across protocols introduced by the different morphologies (sizes). We show only the quantities that vary across cells, μ_V is fixed across cells by design. (C) Dependency of the fluctuations properties as a function of the morphology parameters (all parameters are varied of -70% and +70% around those of the mean model, one parameter is varied while all others are fixed to those of the mean model, see Table 1). We fixed the presynaptic stimulation to a level of $v_e^p = v_e^d = 0.3 Hz$, $v_i^p = v_i^d = 1.7 Hz$ and s = 0.05. Globally, the mean polarization μ_V is poorly affected by the morphology, given the balanced nature of the input and the homogenous spread of synapses on the membrane. On the other hand, the morphology parameters strongly affect the properties of the amplitude and speed of the fluctuations. (i) Dependency on the branching number. Because the branching number has an impact on the area, the number of synapses increases with the number of branches, then the amplitude σ_V of the fluctuations strongly decreases with the number of branches because of the law of large numbers. (ii) Dependency on the tree length. The same argument as in (i) holds. The length of the tree increases the number of synapses so that it reduces the amplitude of the fluctuations. (iii) Dependency on the tree diameter. The same argument as in (i) holds again. The diameter of the tree increases the number of synapses so that it reduces the amplitude of the fluctuations. (iv) Dependency on the size of the soma. Here, this increases only the number of inhibitory synapses at the soma. Within this range this has very weak effects on the fluctuations. (v) Dependency on the leak conductance. The leak conductance sets the amplitude and low pass filtering of individual postsynaptic events (because $\tau_m \sim C_m/G_L$, see correction introduced by the ongoing activity in the main text). An increasing leak conductance will "shunt" post-synaptic events. This lowers the amplitude of the fluctuations σ_V and lowers the autocorrelation time of the fluctuations au_V . (vi) Dependency on the specific capacitance. Again because $au_m \sim C_m/G_L$, an increasing capacitance renders the fluctuations much slower (increasing τ_V). Also because PSP events have a much lower frequency content, they lead to a reduced amplitude of the fluctuations σ_V . (vii) Dependency on the intracellular resistivity. This has a moderate effect within this range.

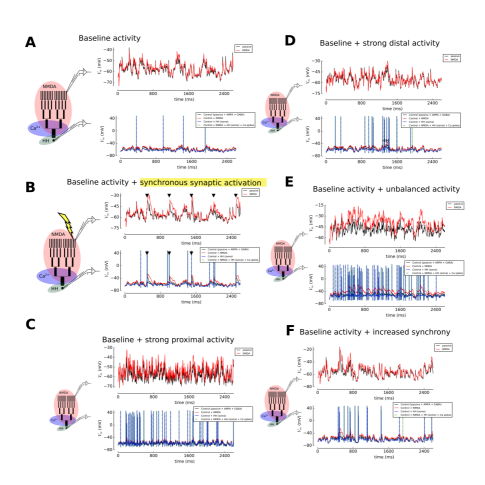
FIGURE S3



Re-including all cells in the study of the correlation between the firing rate responses and the firing response function characteristics. We re-introduce here the recorded cells of very low excitability that produce very low firing rate responses

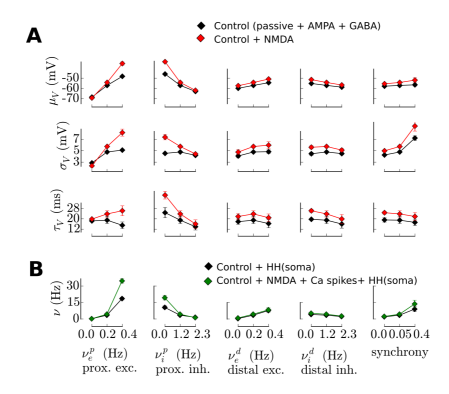
 $(\nu_{out} < 10^{-3})$ and partially impede data visualization (because they shift extend the axis limits far from the majority of the data). Note that the strong correlations discussed in the result are not affected.

FIGURE \$4



Including dendritic non-linearities (numerically) within our framework for single cell computation. Active mechanisms are taken from Larkum et al. [28]. On top of each panel, we present the recording of the membrane potential in the far distal dendrite (depicted with the upper electrode on the drawing) in the Control and Control+NMDA case. On the bottom, we present the membrane potential recorded at the soma (depicted with the lower electrode on the drawing) in four cases, the Control, the Control+NMDA case, the Control case with a HH model inserted in the soma and the Control+NMDA+Ca²⁺spikes+HH(soma). (A) Response to baseline activity. $v_e^p = v_e^d = 0.2$ Hz, $v_i^p = v_i^d = 1.2$ Hz and s = 0.05. (B) Response to baseline activity with the synchronous activation of 20 synapses every 500ms (marked by a triangle), note the impact of the NMDA mechanism under this stimulation type. (C) Response to an increase of proximal activity with respect to the baseline level ($v_e^p = 1.6$ Hz, $v_i^p = 10$ Hz). (D) Response to an increase in distally targeting presynaptic activity ($v_e^d = 0.6$ Hz, $v_i^d = 3$ Hz). (E) Response to an increase in unbalanced activity ($v_e^p = 0.4$ Hz, $v_i^p = 2$ Hz, $v_e^p = 0.5$ Hz, $v_i^p = 1.7$ Hz). (F) Response to an increase in presynaptic synchrony (s = 0.4).

FIGURE S5



Active dendritic mechanisms do not qualitatively affect the relationship between presynaptic activity and firing rate provided that excitation does not unbalance inhibition ($\mu_V \lesssim -50$ mV), and provided that synchrony remains in a relatively low range ($s \lesssim 0.4$). Because our framework for single cell computation consists in (i) evaluating analytically the somatic subthreshold membrane potential fluctuations at the soma as a function of the presynaptic quantities and (ii) convert those fluctuations into a spiking probability thanks to a firing response function determined in vitro in individual neuron thanks to somatic recordings, it needs to be tested against two phenomena: (I) whether active dendritic mechanisms qualitatively affect the relationship between presynaptic quantities and somatic membrane potential fluctuations, i.e. checking the validity of step (i), and (II) whether dendritic non-linearities introduce a coupling between presynaptic activity and spike emission that renders the decomposition into the successive steps i & ii problematic. Those phenomena will naturally appear to some extent and will produce quantitative deviations with respect to our predictions (that do not include dendritic non-linearities, e.g. trivially, all cells should be more excitable). As our results do not rely on the absolute values of the input-output functions but rather on the relative behaviors from cell to cell (that emerge from differences across individual firing response functions), we only look for putative "qualitative deviations". (A) We first investigate whether active dendritic mechanisms qualitatively affect the relationship between presynaptic quantities and somatic membrane potential fluctuations. We observe that NMDA channels exert an additional small depolarization (increased μ_V), they also increase the standard deviation of the fluctuations because they increase the amplitude of depolarizing events (increased σ_V) and they slow down the fluctuations because they increase the low frequency content of post-synaptic depolarization (increased τ_{ν}). We also observe, that, despite this small shift in all quantities, the qualitative trend of all curves remains identical. Only for high excitation, a qualitative difference can be seen (see bottom left plot): increasing excitation makes V_m fluctuations faster in the passive case, whereas increasing excitation makes fluctuations slower in the presence of NMDA currents (decreasing vs increasing τ_V curve respectively). (B) We then investigate whether dendritic non-linearities introduce a coupling between presynaptic activity and spike emission that would complicate our picture. This was tested by including both NMDA and Ca²⁺ currents. We first included numerically a spike emission model at the soma (Hodgkin-Huxley like, Na+ and K+ currents) in the morphological model, to allow the model to produce the full input-output function (from presynaptic quantities to spiking probability). Trivially, we observe an increased spiking probability because cells are more excitable (NMDA and Ca²⁺ currents are depolarizing currents). Notably, when excitation does not balance inhibition ($v_e^p = 0.4$ Hz or $v_i^p = 0$ Hz), the spiking probability is greatly enhanced (almost doubled). For the other variables, those effects are much more moderate.

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FIGURES

FIGURE 1

Figure 1. A theoretical framework for single cell computation in the fluctuation-driven regime. (A) Theoretical paradigm: to get the input-output function of a single cell, we split the relation from presynaptic quantities (the input) to the spiking probability (the output) into two steps. 1) passive dendritic integration shapes the membrane potential at the soma and 2) how those fluctuations are translated into spikes is captured by a firing response function determined *in vitro*. (B) Theoretical model for dendritic integration. A single cell is made of a lumped impedance somatic compartment and a dendritic tree. The dendritic tree is composed of B branches (here B=5), the branching is symmetric and follow Rall's 3/2 rule for the branch diameters. Synapses are then spread all over the membrane according to physiological synaptic densities. We define 3 domains: a somatic and proximal domain as well as a distal domain, excitatory and inhibitory synaptic input can vary independently in those domains. An additional variable: synaptic synchrony controls the degree of coincident synaptic inputs. (C) A given presynaptic stimulation (here $v_e^p = v_e^d = 0.2Hz$, $v_i^p = v_i^d = 1.2~Hz$ and s = 0.05) creates membrane potential fluctuations at the soma characterized by their mean μ_V , their amplitude σ_V and their autocorrelation time τ_V .

FIGURE 2

Figure 2. Calibrating the model on *in vitro* measurements: the simplified model and its size variations provides an approximation for the somatic input impedance of pyramidal cells and its heterogeneity over the recorded population. (A) Input impedance (left: modulus and right: phase shift) measured at the soma in intracellular recordings with sine-wave protocols in current-clamp (inset). The color code indicates the input resistance and is likely to result from size variations of individual cells. (B) A medium size model accounts for the average data and varying the size of the dendritic tree and soma (according to the sizing rule shown in C) partially reproduces the variability in the individual measurements. Large cells (blue) have a lower modulus and a lower phase shift while small cells (red) have both a higher modulus and phase shift. (C) We obtain a map between input resistance and size of the morphological model. (D) Representation of the medium-size model. (E) Additionally the synaptic weights are rescaled with respect to the cell's somatic input resistance. Because the mean transfer resistance to soma is linked to the input resistance, this rescaling insures that the mean synaptic efficacy at soma is the same for all cells.

FIGURE 3

Figure 3. Accuracy of the analytical estimate for the properties of the membrane potential fluctuations at the soma: comparison between numerical simulations and the analytical approximation. Shown for the medium size model of Figure 2D. (A) In the numerical simulation, we explicitly simulate the whole dendritic arborescence, we show the membrane potential variations for the three locations shown on the left. (B) Properties of the membrane potential fluctuations (mean μ_V , standard deviation σ_V and autocorrelation time τ_V) for different configuration of presynaptic activity: analytical predictions and output from numerical simulations in *NEURON*. In each column, one variable is varied while the other variables are fixed to the mean configuration value corresponding to $v_e^p = v_e^d$ =0.2Hz, $v_i^p = v_i^d$ =1.2Hz and s=0.05. In the σ_V plots (middle panels, dashed gray lines), we added the prediction of the analytical estimate after a +0.18 correction for the synchrony (found with a Newton method).

FIGURE 4

Figure 4. Properties of the membrane potential fluctuations for various types of presynaptic activity: either unbalanced (red), purely proximal (blue), purely distal (green), synchronized (cyan). A common baseline configuration of balanced proximal and distal activity at low rate gives rise to baseline fluctuations properties, on top of this, the increase of a given type of presynaptic activity corresponds to a given comodulations of the 5 model variables. (A) Comodulations of the model variables to achieve varying levels of the different types of activity. (B) Membrane potential fluctuations properties (mean μ_V , standard deviation σ_V and autocorrelation time τ_V) and somatic input conductance at the soma for the different protocols. Shown for the medium-size model, see Supplementary Material for the variability introduced by variations in cell morphologies.

FIGURE 5

Figure 5. Examples of the firing response of 4 different cells for the various types of presynaptic activity (color-coded) shown in Figure 4. The abscissa "increasing synaptic quantity" corresponds to the comodulations of the model variables shown in Figure 4A (same color code). As an example, the response to a "proximal activity increase" (blue curve) corresponds to a linear increase of v_e^{prox} and v_i^{prox} while keeping v_e^{dist} , v_i^{dist} and s to their baseline level. See Figure 4A for the comodulations of the other types of increasing activity.

FIGURE 6

Figure 6. Diverse cellular responses to the various types of presynaptic activity and their link to the characteristics of their firing response function. Note the logarithmic scale for the firing responses in B,C,D. (A) Diverse response to baseline stimulation. (B) Diverse response to unbalanced activity. (C) Diverse response to proximal activity. Note that because the response also show negative changes of firing rate, the data cannot be log-scaled. Instead, they have been rescaled by the baseline response (i.e. we show $\frac{\delta v_{prox}}{v_{bsl}}$). (D) Diverse response to distal activity. (E) Diverse response to a synchrony increase.

TABLES

TABLE 1

Parameters	Name	Symbol	Value	Unit
Passive				
	Leak conductance density	G_L	325	μS/cm²
	Intracellular resistivity	R_i	30	$\Omega.cm$
	Specific capacitance	C_m	1.05	μF/cm²
	Leak reversal potential	E_L	-65	mV
Synaptic				
	Inhibitory reversal potential	E_i	-80	mV
	Excitatory reversal potential	E_e	0	mV
	Somatic inhibitory density	\mathcal{D}_i^{soma}	20	Synapses/(100μm²)
	Somatic excitatory density	\mathcal{D}_e^{soma}	0	Synapses/(100μm²)
	Dendritic inhibitory density	\mathcal{D}_i^{tree}	6	Synapses/(100μm²)
	Dendritic excitatory density	\mathcal{D}_e^{tree}	30	Synapses/(100μm²)
	Proximal inhibitory weight	$Q_i^{0,p}$	1.0	nS
	Proximal excitatory weight	$Q_e^{0,p}$	0.7	nS
	Distal inhibitory weight	$Q_i^{0,d}$	1.5	nS
	Distal excitatory weight	$Q_e^{0,d}$	1.05	nS
	Inhibitory decay time	$ au_i$	5	ms
	Excitatory decay time	$ au_e$	5	ms
Mean Morphology				
	Soma length	l_{S}	5.0	μm
	Soma diameter	d_S	15.0	μm
	Root branch diameter	d_t	2.25	μm
	Tree length	l_t	550.0	μm
	Branch number	В	5	
	Proximal tree fraction	f_{prox}	7/8	

TABLE 1. MODEL PARAMETERS

Supplementary Material Heterogenous firing responses leads to diverse coupling to presynaptic activity in mice layer V pyramidal neurons

Y. Zerlaut & A. Destexhe

June 20, 2016

Contents

1	Reduction to the equivalent cylinder	2
2	Mean membrane potential	2
3	Membrane potential response to a synaptic event	4
4	Deriving the power spectrum of the membrane potential fluctuations	g
5	Deriving the fluctuations properties $(\mu_V, \sigma_V, \tau_V)$	10
6	References	11

1 Reduction to the equivalent cylinder

The key of the derivation relies on having the possibility to reduce the complex morphology to an equivalent cylinder (Rall, 1962). We adapted this procedure to capture the change in integrative properties of the membrane that results from the mean synaptic bombardment during active cortical states, reviewed in Destexhe et al. (2003).

For a set of synaptic stimulation $\{\nu_p^e, \nu_i^p, \nu_e^d, \nu_i^d, s\}$, let's introduce the following stationary densities of conductances:

$$\begin{cases} g_{e0}^{p} = \pi \, d \, \mathcal{D}_{e} \, \nu_{e}^{p} \, \tau_{e}^{p} \, Q_{e}^{p} & ; \quad g_{i0}^{p} = \pi \, d \, \mathcal{D}_{i} \, \nu_{i}^{p} \, \tau_{i}^{p} \, Q_{i}^{p} \\ g_{e0}^{d} = \pi \, d \, \mathcal{D}_{e} \, \nu_{e}^{d} \, \tau_{e}^{d} \, Q_{e}^{d} & ; \quad g_{i0}^{d} = \pi \, d \, \mathcal{D}_{i} \, \nu_{i}^{d} \, \tau_{i}^{d} \, Q_{i}^{d} \end{cases}$$
(1)

where \mathcal{D}_e and \mathcal{D}_i are the excitatory and inhibitory synaptic densities.

We introduce two activity-dependent electrotonic constants relative to the proximal and distal part respectively:

$$\lambda^{p} = \sqrt{\frac{r_{m}}{r_{i}(1 + r_{m}g_{e0}^{p} + r_{m}g_{i0}^{p})}} \quad \lambda^{d} = \sqrt{\frac{r_{m}}{r_{i}(1 + r_{m}g_{e0}^{d} + r_{m}g_{i0}^{d})}}$$
 (2)

For a dendritic tree of total length l, whose proximal part ends at l_p and with B evenly spaced generations of branches, we define the space-dependent electrotonic constant:

$$\lambda(x) = (\lambda^p + \mathcal{H}(x - l_p)(\lambda^d - \lambda^p))2^{-\frac{1}{3} \lfloor \frac{B \cdot x}{l} \rfloor}$$
(3)

where $\lfloor . \rfloor$ is the floor function. Note that $\lambda(x)$ is constant on a given generation, but it decreases from generation to generation because of the decreasing diameter along the dendritic tree. It also depends on the synaptic activity and therefore has a discontinuity at $x = l_p$.

Following Rall (1962), we now define a dimensionless length X:

$$X(x) = \int_0^x \frac{dx}{\lambda(x)} \tag{4}$$

We define L = X(l) and $L_p = X(l_p)$, the total length and proximal part length respectively (capital letters design rescaled quantities).

2 Mean membrane potential

We derive the mean membrane potential $\mu_V(x)$ corresponding to the stationary response to constant densities of conductances given by the means of the synaptic stimulation. We obtain the stationary equations by removing temporal derivatives in Equation, the set of equation governing this mean membrane potential in all branches is therefore:

$$\begin{cases}
\frac{1}{r_{i}} \frac{\partial^{2} \mu_{v}}{\partial x^{2}} = \frac{\mu_{v}(x) - E_{L}}{r_{m}} \\
- g_{e0}^{p} (\mu_{v}(x) - E_{e}) - g_{0i}^{p} (\mu_{v}(x) - E_{i}) \quad \forall x \in [0, l_{p}] \\
\frac{1}{r_{i}} \frac{\partial^{2} \mu_{v}}{\partial x^{2}} = \frac{\mu_{v}(x) - E_{L}}{r_{m}} \\
- g_{e0}^{d} (\mu_{v}(x) - E_{e}) - g_{0i}^{d} (\mu_{v}(x) - E_{i}) \quad \forall x \in [l_{p}, l] \\
\frac{\partial \mu_{v}}{\partial x|_{x=0}} = r_{i} \left(\frac{\mu_{v}(0) - E_{L}}{R_{m}} + G_{i0}^{S} (\mu_{v}(0) - E_{i}) \right) \\
\mu_{v}(l_{p}^{-}, t) = \mu_{v}(l_{p}^{+}, t) \\
\frac{\partial \mu_{v}}{\partial x|_{r_{p}^{-}}} = \frac{\partial \mu_{v}}{\partial x|_{l_{p}^{+}}} \\
\frac{\partial \mu_{v}}{\partial x|_{x=l}} = 0
\end{cases}$$
(5)

Because the reduction to the equivalent cylinder conserves the membrane area and the previous equation only depends on density of currents, the equation governing $\mu_v(x)$ in all branches can be transformed into an equation on an equivalent cylinder of length L. We rescale x by $\lambda(x)$ (see Equation 4) and we obtain the equation verified by $\mu_V(X)$:

$$\begin{cases}
\frac{\partial^{2} \mu_{v}}{\partial X^{2}} = \mu_{v}(X) - v_{0}^{p} & \forall X \in [0, L_{p}] \\
\frac{\partial^{2} \mu_{v}}{\partial X^{2}} = \mu_{v}(X) - v_{0}^{d} & \forall X \in [L_{p}, L] \\
\frac{\partial \mu_{v}}{\partial X}|_{X=0} = \gamma^{p} \left(\mu_{v}(0) - V_{0}\right) \\
\mu_{v}(L_{p}^{-}) = \mu_{v}(L_{p}^{+}) \\
\frac{\partial \mu_{v}}{\partial X}|_{L_{p}^{-}} = \frac{\lambda^{p}}{\lambda^{d}} \frac{\partial \mu_{v}}{\partial X}|_{L_{p}^{+}} \\
\frac{\partial \mu_{v}}{\partial X}|_{X=L} = 0
\end{cases}$$
(6)

where:

$$v_{0}^{p} = \frac{E_{L} + r_{m} g_{e0}^{p} E_{e} + r_{m} g_{i0}^{p} E_{i}}{1 + r_{m} g_{e0}^{p} + r_{m} g_{i0}^{p}}$$

$$v_{0}^{d} = \frac{E_{L} + r_{m} g_{e0}^{d} E_{e} + r_{m} g_{i0}^{d} E_{i}}{1 + r_{m} g_{e0}^{d} + r_{m} g_{i0}^{d}}$$

$$\gamma^{p} = \frac{r_{i} \lambda^{p} (1 + G_{i}^{0} R_{m})}{R_{m}}$$

$$V_{0} = \frac{E_{L} + G_{i}^{0} R_{m} E_{i}}{1 + G_{i}^{0} R_{m}}$$
(7)

We write the solution on the form:

$$\begin{cases} \mu_v(X) = v_0^p + A \cosh(X) + C \sinh(X) \quad \forall X \in [0, L_p] \\ \mu_v(X) = v_0^d + B \cosh(X - L) + D \sinh(X - L) \quad \forall X \in [L_p L] \end{cases}$$
(8)

- Sealed-end boundary condition at cable end implies D=0
- Somatic boundary condition imply: $C = \gamma^p (v_0^p V_0 + A)$
- Then v continuity imply : $v_0^p + A \cosh(L_p) + \gamma^p (v_0^p V_0 + A) \sinh(L_p) = v_0^d + B \cosh(L_p L)$
- Then current conservation imply: $A \sinh(L_p) + \gamma^p (v_0^p V_0 + A) \cosh(L_p) = \frac{\lambda^p}{\lambda^d} B \sinh(L_p L)$

We rewrite those condition on a matrix form:

$$\begin{pmatrix}
\cosh(L_p) + \gamma^p \sinh(L_p) & -\cosh(L_p - L) \\
\sinh(L_p) + \gamma^p \cosh(L_p) & -\frac{\lambda^p}{\lambda^d} \sinh(L_p - L)
\end{pmatrix} \cdot \begin{pmatrix} A \\ B \end{pmatrix} = \begin{pmatrix} v_0^d - v_0^p - \gamma^p (v_0^p - V_0) \sinh(L_p) \\
-\gamma^p (v_0^p - V_0) \cosh(L_p)
\end{pmatrix}$$

And we solved this equation with the solve_linear_system_LU method of sympy

The coefficients A and B are given by:

$$A = \frac{\alpha}{\beta} \qquad B = \frac{\gamma}{\delta} \tag{10}$$

where:

$$\alpha = V_0 \gamma^P \lambda^D \cosh(L_p) \cosh(L - L_p) + V_0 \gamma^P \lambda^P \sinh(L_p) \sinh(L - L_p)$$

$$- \gamma^P \lambda^D v_0^d \cosh(L_p) \cosh(L - L_p) - \gamma^P \lambda^P v_0^d \sinh(L_p) \sinh(L - L_p)$$

$$- \lambda^P v_0^d \sinh(L - L_p) + \lambda^P v_0^p \sinh(L - L_p)$$

$$\beta = \gamma^P \lambda^D \cosh(L_p) \cosh(L - L_p) + \gamma^P \lambda^P \sinh(L_p) \sinh(L - L_p) + \lambda^D \sinh(L_p) \cosh(L - L_p) + \lambda^P \sinh(L - L_p) \cosh(L_p)$$

$$\gamma = \lambda^D (V_0 \gamma^P + \gamma^P v_0^d \cosh(L_p) - \gamma^P v_0^d$$

$$- \gamma^P v_0^p \cosh(L_p) + v_0^d \sinh(L_p) - v_0^p \sinh(L_p))$$

$$\delta = \gamma^P \lambda^D \cosh(L_p) \cosh(L - L_p) + \gamma^P \lambda^P \sinh(L_p) \sinh(L - L_p)$$

$$+ \lambda^D \sinh(L_p) \cosh(L - L_p) + \lambda^P \sinh(L - L_p) \cosh(L_p)$$

3 Membrane potential response to a synaptic event

We now look for the response to $n_{src} = \lfloor \frac{B x_{src}}{l} \rfloor$ synaptic events at position x_{src} on all branches of the generation of x_{src} , those events have a conductance

 $g(t)/n_{src}$ and reversal potential E_{rev} . We make the hypothesis that the initial condition correspond to the stationary mean membrane potential $\mu_V(x)$. This potential will also be used to fix the driving force at the synapse to $\mu_v(x_{src}) - E_{rev}$, this linearizes the equation and will allow an analytical treatment. To derive the equation for the response around the mean $\mu_v(x)$, we rewrite Equation 9 in main text with $v(x,t) = \delta v(x,t) + \mu_v(x)$, we obtain the equation for $\delta v(x,t)$:

$$\begin{cases}
\frac{1}{r_{i}} \frac{\partial^{2} \delta v}{\partial x^{2}} = c_{m} \frac{\partial \delta v}{\partial t} + \frac{\delta v}{r_{m}} (1 + r_{m} g_{e0}^{p} + r_{m} g_{i0}^{p}) \\
-\delta(x - x_{src}) \left(\mu_{v}(x_{src}) - E_{rev}\right) \frac{g(t)}{n_{src}}, \quad \forall x \in [0, l_{p}]
\end{cases}$$

$$\begin{cases}
\frac{1}{r_{i}} \frac{\partial^{2} \delta v}{\partial x^{2}} = c_{m} \frac{\partial \delta v}{\partial t} + \frac{\delta v}{r_{m}} (1 + r_{m} g_{e0}^{d} + r_{m} g_{i0}^{d}) \\
-\delta(x - x_{src}) \left(\mu_{v}(x_{src}) - E_{rev}\right) \frac{g(t)}{n_{src}}, \quad \forall x \in [l_{p}, l]
\end{cases}$$

$$\begin{cases}
\frac{1}{r_{i}} \frac{\partial \delta v}{\partial x^{2}} = C_{M} \frac{\partial \delta v}{\partial t} + \frac{\delta v(0, t)}{R_{m}} (1 + R_{m} G_{i0}^{S}) \\
\delta v(l_{p}^{-}, t) = \delta v(l_{p}^{+}, t)
\end{cases}$$

$$\begin{cases}
\frac{\partial \delta v}{\partial x} = \frac{\partial \delta v}{l_{p}^{-}} = \frac{\partial \delta v}{\partial x} + \frac{\partial \delta v}{l_{p}^{+}} \\
\frac{\partial \delta v}{\partial x} = 0
\end{cases}$$

$$\begin{cases}
\frac{\partial \delta v}{\partial x} = 0
\end{cases}$$

Because this synaptic event is concomitant in all branches at distance x_{src} , we can use again the reduction to the equivalent cylinder (note that the event has now a weight multiplied by n_{src} so that its conductance becomes g(t)), we obtain:

eight multiplied by
$$n_{src}$$
 so that its conductance becomes $g(t)$), we
$$\begin{cases} \frac{\partial^2 \delta v}{\partial X^2} = \left(\tau_m^p + (\tau_m^d - \tau_m^p)\mathcal{H}(X - L_p)\right) \frac{\partial \delta v}{\partial t} + \delta v \\ - \left(\mu_v(X_{src}) - E_{rev}\right) \delta(X - X_{src}) \times \\ \frac{g(t)}{c_m} \left(\frac{\tau_m^p}{\lambda^p} + (\frac{\tau_m^d}{\lambda^d} - \frac{\tau_m^p}{\lambda^p})\mathcal{H}(X_{src} - L_p)\right) \end{cases} \\ \begin{cases} \frac{\partial \delta v}{\partial X}|_{X=0} = \gamma^p \left(\tau_m^S \frac{\partial \delta v}{\partial t}|_{X=0} + \delta v(0,t)\right) \\ \delta v(L_p^-, t) = \delta v(L_p^+, t) \\ \frac{\partial \delta v}{\partial X}|_{L_p^-} = \frac{\lambda^p}{\lambda^d} \frac{\partial \delta v}{\partial X}|_{L_p^+} \\ \frac{\partial \delta v}{\partial X}|_{X=L} = 0 \end{cases}$$

$$(13)$$

where we have introduced the following time constants:

$$\tau_m^D = \frac{r_m c_m}{1 + r_m g_{e0}^d + r_m g_{i0}^d}
\tau_m^P = \frac{r_m c_m}{1 + r_m g_{e0}^p + r_m g_{i0}^p}
\tau_m^S = \frac{R_m C_m}{1 + R_m G_{i0}^S}$$
(14)

We now use distribution theory (see Appel (2008) for a comprehensive textbook) to translate the synaptic input into boundary conditions at X_{src} , physically this corresponds to: 1) the continuity of the membrane potential and 2) the discontinuity of the current resulting from the synaptic input.

$$\begin{cases}
\delta v(X_{src}^{-}, f) = \delta v(X_{src}^{+}, f) \\
\frac{\partial \delta v}{\partial X_{src}^{+}} - \frac{\partial \delta v}{\partial X_{src}^{-}} = -(\mu_{v}(X_{src}) - E_{rev}) \times \\
(\frac{\tau_{m}^{p}}{\lambda^{p}} + (\frac{\tau_{m}^{d}}{\lambda^{d}} - \frac{\tau_{m}^{p}}{\lambda^{p}}) \mathcal{H}(X_{src} - L_{p})) \frac{g(t)}{c_{m}}
\end{cases}$$
(15)

We will solve Equation 13 by using Fourier analysis. We take the following convention for the Fourier transform:

$$\hat{F}(f) = \int_{\mathbb{R}} F(t) e^{-2i\pi ft} dt \tag{16}$$

We Fourier transform the set of Equations 13, we obtain:

urier transform the set of Equations 13, we obtain:
$$\begin{cases}
\frac{\partial^{2} \hat{\delta v}}{\partial X^{2}} = \left(\alpha_{f}^{p} + (\alpha_{f}^{d} - \alpha_{f}^{p})\mathcal{H}(X - L_{p})\right)^{2} \hat{\delta v} \\
\frac{\partial \hat{\delta v}}{\partial X_{|X=0}} = \gamma_{f}^{p} \hat{\delta v}(0, f) \\
\hat{\delta v}(X_{src}^{-}, f) = \hat{\delta v}(X_{src}^{+}, f) \\
\frac{\partial \hat{\delta v}}{\partial X_{X_{src}^{-}}} = \frac{\partial \hat{\delta v}}{\partial X_{X_{src}^{+}}} - \left(\mu_{v}(X_{src}) - E_{rev}\right) \times \\
\left(r_{f}^{p} + \left(r_{f}^{d} - r_{f}^{p}\right)\mathcal{H}(X_{src} - L_{p})\right) g(\hat{f}) \\
\hat{\delta v}(L_{p}^{-}, f) = \hat{\delta v}(L_{p}^{+}, f) \\
\frac{\partial \hat{\delta v}}{\partial X_{L_{p}^{-}}} = \frac{\lambda^{p}}{\lambda^{d}} \frac{\partial \hat{\delta v}}{\partial X_{L_{p}^{+}}} \\
\frac{\partial \hat{\delta v}}{\partial X_{X=L}} = 0
\end{cases} \tag{17}$$

where

$$\alpha_f^p = \sqrt{1 + 2i\pi f \tau_m^p} \qquad r_f^p = \frac{\tau_m^p}{c_m \lambda^p}$$

$$\alpha_f^d = \sqrt{1 + 2i\pi f \tau_m^d} \qquad r_f^d = \frac{\tau_m^d}{c_m \lambda^d}$$

$$\gamma_f^p = \gamma^p \left(1 + 2i\pi f \tau_m^S\right)$$
(18)

To obtain the solution, we need to split the solution into two cases:

1. $X_{src} \leq L_p$

Let's write the solution to this equation as the form (already including the boundary conditions at X = 0 and X = L):

$$\hat{\delta v}(X, X_{src}, f) = \begin{cases}
A_f(X_{src}) \left(\cosh(\alpha_f^p X) + \gamma^p \sinh(\alpha_f^p X) \right) \\
\text{if } : 0 \le X \le X_{src} \le L_p \le L \\
B_f(X_{src}) \cosh(\alpha_f^p (X - L_p)) + C_f(X_{src}) \sinh(\alpha_f^p (X - L_p)) \\
\text{if } : 0 \le X_{src} \le X \le L_p \le L \\
D_f(X_{src}) \cosh(\alpha_f^d (X - L)) \\
\text{if } : 0 \le X_{src} \le L_p \le X \le L
\end{cases} \tag{19}$$

We write the 4 conditions corresponding to the conditions in X_{src} and L_p to get A_f, B_f, C_f, D_f . On a matrix form, this gives:

$$M = \begin{pmatrix} \cosh(\alpha_f^p X_{src}) + \gamma_f^p \sinh(\alpha_f^p X_{src}) & -\cosh(\alpha_f^p (X_{src} - L_p)) & -\sinh(\alpha_f^p (X_{src} - L_p)) & 0 \\ \alpha_f^p \left(\sinh(\alpha_f^p X_{src}) + \gamma_f^p \cosh(\alpha_f^p X_{src}) \right) & -\alpha_f^p \sinh(\alpha_f^p (X_{src} - L_p)) & -\alpha_f^p \cosh(\alpha_f^p (X_{src} - L_p)) & 0 \\ 0 & 1 & 0 & -\cosh(\alpha_f^d (L_p - L)) \\ 0 & 0 & \alpha_f^p & -\alpha_f^d \frac{\lambda^p}{\lambda^d} \sinh(\alpha_f^d (L_p - L)) \end{pmatrix}$$

$$M \cdot \begin{pmatrix} A_f \\ B_f \\ C_f \\ D_f \end{pmatrix} = \begin{pmatrix} 0 \\ -r_f^p I_f \\ 0 \\ 0 \end{pmatrix} \tag{21}$$

And we will solve it with the solve_linear_system_LU method of sympy. For the $A_f(X_{src})$ coefficient, we obtain:

$$A_f(X_{src}) = \frac{a_f^1(X_{src})}{a_f^2(X_{src})}$$
 (22)

with:

$$a_{f}^{1}(X_{src}) = I_{f}r_{f}^{P} \left(-\alpha_{f}^{D}\lambda^{P} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} - L_{p}\alpha_{f}^{P} + X_{s}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{D}\lambda^{P} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} - X_{s}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} - L_{p}\alpha_{f}^{P} + X_{s}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} - X_{s}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{D}\lambda^{D} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{D}\lambda_{f}^{P}\lambda^{P} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) - \alpha_{f}^{D}\lambda^{P} \sinh\left(-L\alpha_{f}^{D} + L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{D}\lambda^{P} \sinh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda_{f}^{P}\lambda^{D} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda_{f}^{P}\lambda^{D} \cosh\left(L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \sinh\left(-L\alpha_{f}^{D} + L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \sinh\left(-L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \sinh\left(-L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \sinh\left(-L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

$$\left. + \alpha_{f}^{P}\lambda^{D} \sinh\left(-L\alpha_{f}^{D} - L_{p}\alpha_{f}^{D} + L_{p}\alpha_{f}^{P} \right) \right.$$

2. $L_p \leq X_{\rm src}$

Let's write the solution to this equation as the form (already including the boundary conditions at X = 0 and X = L:

$$\hat{\delta v}(X, X_{src}, f) =
\begin{cases}
E_f(X_{src}) \left(\cosh(\alpha_f^p X) + \gamma^p \sinh(\alpha_f^p X) \right) \\
\text{if } : 0 \le X \le L_p \le X_{src} \le L \\
F_f(X_{src}) \cosh(\alpha_f^d (X - L_p)) + G_f(X_{src}) \sinh(\alpha_f^d (X - L_p)) \\
\text{if } : 0 \le L_p \le X \le X_{src} \le L \\
H_f(X_{src}) \cosh(\alpha_f^d (X - L)) \\
\text{if } : 0 \le L_p \le X_{src} \le X \le L
\end{cases}$$
(24)

We write the 4 conditions corresponding to the conditions in X_{src} and L_p to get A_f, B_f, C_f, D_f . On a matrix form, this gives:

We rewrite this condition on a matrix form:

$$M_{2} = \begin{pmatrix} \cosh(\alpha_{f}^{p}L_{p}) + \gamma_{f}^{p}\sinh(\alpha_{f}^{p}L_{p}) & -1 & 0 & 0 & 0\\ \alpha_{f}^{p}\left(\sinh(\alpha_{f}^{p}L_{p}) + \gamma_{f}^{p}\cosh(\alpha_{f}^{p}L_{p})\right) & 0 & -\alpha_{f}^{d}\frac{\lambda^{p}}{\lambda^{d}} & 0\\ 0 & \cosh(\alpha_{f}^{d}\left(X_{src} - L_{p}\right)) & \sinh(\alpha_{f}^{d}\left(X_{src} - L_{p}\right)) & -\cosh(\alpha_{f}^{d}\left(X_{src} - L\right))\\ 0 & \alpha_{f}^{d}\sinh(\alpha_{f}^{d}\left(X_{src} - L_{p}\right)) & \alpha_{f}^{d}\cosh(\alpha_{f}^{d}\left(X_{src} - L_{p}\right)) & -\alpha_{f}^{d}\sinh(\alpha_{f}^{d}\left(X_{src} - L\right)) \end{pmatrix}$$
(25)

$$M \cdot \begin{pmatrix} E_f \\ F_f \\ G_f \\ H_f \end{pmatrix} = \begin{pmatrix} 0 \\ 0 \\ 0 \\ -r_f^d I_f \end{pmatrix} \tag{26}$$

And we will solve it with the solve_linear_system_LU method of sympy. For the $E_f(X_{src})$ coefficient, we obtain:

$$E_f(X_{src}) = \frac{e_f^1(X_{src})}{e_f^2(X_{src})}$$
 (27)

with:

$$e_f^1(X_{src}) = 2I_f \lambda^P r_f^D \cosh\left(\alpha_f^D (L - X_s)\right)$$

$$e_f^2(X_{src}) = -\alpha_f^D \gamma_f^P \lambda^P \cosh\left(-L\alpha_f^D + L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^D \gamma_f^P \lambda^P \cosh\left(L\alpha_f^D - L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$- \alpha_f^D \lambda^P \sinh\left(-L\alpha_f^D + L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^D \lambda^P \sinh\left(L\alpha_f^D - L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^D \gamma_f^P \lambda^D \cosh\left(-L\alpha_f^D + L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^P \gamma_f^P \lambda^D \cosh\left(L\alpha_f^D - L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^P \gamma_f^P \lambda^D \sinh\left(-L\alpha_f^D + L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^P \lambda^D \sinh\left(-L\alpha_f^D + L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^P \lambda^D \sinh\left(L\alpha_f^D - L_p \alpha_f^D + L_p \alpha_f^P\right)$$

$$+ \alpha_f^P \lambda^D \sinh\left(L\alpha_f^D - L_p \alpha_f^D + L_p \alpha_f^P\right)$$

From this calculus, we can write the PSP at the soma on the form:

$$\hat{\delta v}(X=0, X_{src}, f) = K_f(X_{src}) \left(\mu_v(X_{src}) - E_{rev}\right) g(\hat{f})$$
 (29)

where $K_f(X_{src})$ given by:

$$K_f(X_{src}) = \begin{cases} A_f(X_{src}) & \forall X_{src} \in [0, L_p] \\ E_f(X_{src}) & \forall X_{src} \in [L_p, L] \end{cases}$$
(30)

This is obtained by taking a unitary current $I_f = 1$ in the previous calculus.

4 Deriving the power spectrum of the membrane potential fluctuations

The calculus rely on the ability to obtain the power spectrum of the membrane potential fluctuations at the soma $P_V(f)$.

This can be obtained from shotnoise theory (Daley and Vere-Jones, 2007) (see also El Boustani et al. (2009) for an application similar to ours), the general form of the power spectrum density can be expressed as:

$$P_V(f) = \sum_{\{syn\}} N_{syn} F_{\text{synch}} \nu_{syn} \| PS\hat{P}_{syn}(f) \|^2$$

(31)

where $\{syn\}$ is the set of identical synapses, each having N_{syn} synapses, a Poisson release probability: ν_{syn} and creating a post-synaptic event $PSP_{syn}(t)$. In addition, F_{synch} is a synchrony factor (depending on the variable s in the model), it accounts for the effects of the synchronous arrivals of presynaptic events. Given the synchrony generator considered in the main text, the synchrony factor takes the form:

$$F_{\rm synch} = (1-s) + (s-s^2)2^2 + (s^2-s^3)3^2 + s^34^2 \end{(32)}$$

because single events arise with a probability 1 - s, double events with a probability $s - s^2$ (and the PSP are squared in Eq. 4, hence the 2^2 factor), etc...

Now obtaining the power spectrum density $P_V(f)$ in our situation requires to explicit the sum over synapses: $\sum_{\{syn\}}$, In our cases, we need to sum over 1) their type (excitatory/inhibitory, $\sum_{s\in\{e,i\}}$), 2) their location (we will integrate over the dendritic length $\int_0^L dx$) 3) branches.

$$P_{V}(f) = \sum_{s \in \{e,i\}} \int_{0}^{L} dx \, \pi \, \mathcal{D}_{s} \left(d_{t} \, 2^{-\frac{2}{3} \lfloor \frac{B \, x}{l} \rfloor} \right) \, 2^{\lfloor \frac{B \, x}{l} \rfloor} \, F_{\text{synch}} \, \nu_{s}(x) \, \| \hat{\delta v}_{s}(0, x, f) \|^{2}$$

$$+ \pi \, \mathcal{D}_{i} \, l_{S} \, d_{S} \, F_{\text{synch}} \, \nu_{i}(0) \, \| \hat{\delta v}_{i}(0, 0, f) \|^{2}$$

$$(33)$$

where $\delta v_s(0, x, f)$ is given by Eq. 29 (note that the dependency on synaptic type s comes from the reversal potential term E_{rev} in Eq. 29). The factor $2^{\lfloor \frac{Bx}{l} \rfloor}$ corresponds to the sum of the synapses over the different branches at the distance x. The term $\left(d_t \, 2^{-\frac{2}{3} \, \lfloor \frac{Bx}{l} \rfloor}\right)$ is the diameter of the branches at the distance x.

The last term in Eq. 33 corresponds to the contribution of somatic inhibitory synapses (number of somatic inhibitory synapses: $\pi \mathcal{D}_i l_S d_S$.

5 Deriving the fluctuations properties $(\mu_V, \sigma_V, \tau_V)$

The final expressions for the fluctuation propeorties as a function of $(\nu_e^p, \nu_i^p, \nu_e^d, \nu_i^d, s)$ are thus given by:

- μ_V : we obtain the mean of the fluctuations at the soma by taking $\mu_V(0)$ in Equation 8.
- σ_V : we obtain the standard deviation of the fluctuations from the power spectrum density in Equation 33 and the expression:

$$\sigma_V^2 = \int_{\mathbb{R}} P_V(f) \, df \tag{34}$$

This integral expression was discretized and evaluated numerically

• τ_V : we obtain the autocorrelation time of the fluctuations from the power spectrum density in Equation 33 and the expression (Zerlaut et al., 2016):

$$\tau_V = \frac{1}{2} \left(\frac{\int_{\mathbb{R}} P_V(f) \, df}{P_V(0)} \right)^{-1} \tag{35}$$

This integral expression was discretized and evaluated numerically

6 References

Appel W (2008) Mathématiques pour la physique et les physiciens H. et K. Editions.

Daley DJ, Vere-Jones D (2007) An introduction to the theory of point processes: volume II: general theory and structure, Vol. 2 Springer Science & Business Media.

Destexhe A, Rudolph M, Paré D (2003) The high-conductance state of neocortical neurons in vivo. *Nature reviews. Neuroscience* 4:739–751.

El Boustani S, Marre O, Béhuret S, Baudot P, Yger P, Bal T, Destexhe A, Frégnac Y (2009) Network-state modulation of power-law frequency-scaling in visual cortical neurons. *PLoS computational biology* 5:e1000519.

Rall W (1962) Electrophysiology of a dendritic neuron model. *Biophysical* journal 2:145.

Zerlaut Y, Telenczuk B, Deleuze C, Bal T, Ouanounou G, Destexhe A (2016) Heterogeneous firing response of mice layer V pyramidal neurons in the fluctuation-driven regime. *The Journal of Physiology* in press.