Excitation creates a distributed pattern of cortical suppression due to varied recurrent input

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Abstract

Sensory cortex transforms its inputs, generating neural responses that represent features of the sensory world. Dense local, recurrent connections are a major feature of cortical circuits, yet how they affect these transformations is unclear, with some studies reporting that recurrent circuits amplify certain excitatory inputs, and other studies showing instead local suppression. Here we find that input to mouse V1 excitatory neurons generates salt-and-pepper patterns of excitation and suppression. We show that neurons’ responses are not strongly predicted by their optogenetic input alone, suggesting that recurrent inputs nonlinearly transform the population response. A balanced-state network model explains the observed dynamics, suppressed responses, and long tail of excited responses, but only with substantial variance in cell-to-cell connectivity. In sum, we find recurrent cortical connectivity has high variability and high average strength, which can decouple individual cells’ responses from the direct input each cell receives, leading to substantial suppression generated by excitatory input.
Introduction

The cerebral cortex of mammals is specialized into areas that perform different functions\(^1\). Animals from rodents to primates have several different visual cortical areas, each containing neurons with different types of selectivity\(^2\)–\(^4\). In principle, these different representations in different visual areas could be created purely by feedforward mechanisms, where transformations happen via projections from one area or layer to the next, without outputs of a neuron feeding back (directly or indirectly) to influence that neuron’s activity. In fact, in a variety of artificial neural networks, much or all computation is provided by feedforward mechanisms\(^5\).

Yet in the brains of animals and humans, cortical recurrent connectivity is extensive. Most excitatory connections that a cortical neuron receives originate within a few hundred microns of their cell bodies\(^6\)–\(^8\). Such recurrent connections can in principle have large effects on neural computation\(^9\), dramatically changing how cortical neurons respond to input.

How recurrent connections affect cortical computation is not fully understood, but important aspects of the structure of cortical recurrent connectivity have been determined. Some features of cortical network activity, such as irregular firing, are well-described by balanced-state models which assume strong recurrent coupling between excitatory and inhibitory neurons (either moderately strong, yielding ‘loose balance’, or very strong, yielding ‘tight balance’\(^10\)). Work using inhibitory perturbations has shown that not just excitatory-inhibitory connectivity is strong, but the average excitatory-excitatory connectivity is strong as well. More precisely, cortical recurrent excitatory coupling is strong enough that the excitatory network is unstable and self-amplifying, a phenomenon described by inhibition-stabilized network models (ISNs)\(^11\)–\(^14\).

While some consensus has developed on these average cortical connectivity properties (but see\(^15\)), the effect recurrent connections have on transforming sensory or input signals has been less clear. For example, some recent studies have shown that certain patterns of excitatory input can be amplified by the cortical network\(^16\),\(^17\), consistent with some theoretical predictions\(^16\),\(^19\). On the other hand, however, other studies have shown that nearby neurons can be substantially suppressed by stimulation that excites a single or a small ensemble of excitatory cortical neurons\(^20\),\(^21\). How excitatory and inhibitory neurons might interact through recurrent connections to create such suppression has not been determined.

Here, to understand how cortical neurons’ responses are shaped by the cortical recurrent network, we stimulate excitatory cells in the visual cortex optogenetically and record responses of local neurons with electrophysiology and two-photon imaging. First, we find that stimulation of excitatory cells leads to a salt-and-pepper pattern of local suppression, a pattern that is consistent with the pattern of excited and suppressed cells produced when animals see a strong visual stimulus. To understand how this suppression effect might arise from cortical recurrent circuitry, we examine both the patterns of firing rate changes and the dynamics of responses and compare them to cortical models. Recent theoretical work has shown that cortical visual responses can be “reshuffled” by additional excitatory input\(^22\) — that is, strong average recurrent coupling allows individual neurons’ firing to change significantly in response to input while the distribution of population activity is little-changed\(^23\),\(^24\). We implement this scenario in a
conductance-based simulation and find that it can explain the suppression we observe, and additionally we find our data is consistent with substantial variability in local recurrent connectivity, with some neurons receiving large net recurrent excitation and others smaller or net suppressive recurrent input. Our results go beyond prior work that found strong average recurrent connectivity, showing that variance in excitatory-excitatory connectivity must also be substantial, and further show that this variance in recurrent connectivity can decouple neurons’ firing rate responses from the direct input they receive.

The suppression we observe during excitatory cell stimulation occurs in individual cells, but the mean response is elevated. This increase in mean, however, seems at odds with the prior finding that single-cell stimulation leads to inhibition on average. To resolve this, we simulate the effect of single cell stimulation and find that the difference in the two results can be explained by the activation state of the cortical network. Increasing activity in the network with visual stimulation results in a slight decrease in mean responses to stimulation, showing the prior results and our current results can be described in the same model framework. Thus, we find that a balanced-state, excitatory-inhibitory model, with strong average coupling and variability in recurrent connectivity, explains many features of our data, including dynamics and neural response distributions. These results show how cortical neural suppression can be generated from excitatory input: variability in recurrent input means that firing rate responses are decoupled from (are only weakly affected by) the level of excitatory input we provide to that cell, as much of the input a cell receives comes from recurrent sources. Because recurrent input varies from cell to cell, what results is many excited cells, but also a substantial fraction of suppressed neurons.
Results

Strong visual input leads to salt-and-pepper distributed suppression in primary visual cortex

We first measured local patterns of suppression in visual cortex in response to visual stimuli. We presented small high-contrast visual stimuli to head-fixed mice while measuring activity in V1 layer 2/3 neurons via two-photon imaging (Fig. 1A). We expressed GCaMP7s in all neurons via viral injection (AAV-hSyn-GCaMP). Animals were kept awake and in an alert state with occasional drops of water reward, and to ensure animals did not drift into a quiet wakefulness or quiescent state, we monitored animals during data collection to verify they continued to drink the delivered reward.

We imaged responses to two types of high-contrast visual stimuli, a fast-changing stimulus designed to minimize adaptation (“oriented noise”, Fig. 1A) and a drifting grating (data...

Figure 1: V1 neurons show salt-and-pepper suppression to strong visual stimuli. (A) Experimental setup. Awake mice viewed a small (15 degree diameter) visual stimulus with rapidly changing frames of white noise, filtered to have a peak spatial frequency and orientation though pixel values changed from frame to frame (0° peak orientation, 90% contrast, frames presented at 60 Hz, total stimulus duration 3-5s, more details of stimulus in Methods). (B) Example 2-photon imaging data from layer 2/3 of mouse V1 in response to the oriented noise stimulus, during the transient and (C) steady-state periods. Time intervals used for averaging in (B-D) displayed in green and purple in (G). Note the emergence of intermixed (salt-and-pepper) elevated and suppressed responses during the steady-state period. (D) Deconvolved responses from (C), projected onto segmented cell masks (see Methods). (E) Example df/F trace for one elevated and one suppressed cell. Shaded regions: SEM across trials. Shaded red: optogenetic stimulation duration. (F) Deconvolution of the traces in (D) reveals an initial transient period and then a steady-state response. (G) Average response for all elevated and suppressed cells in (B). (H) Schematic of optogenetic and imaging experiments, to determine if suppression arises from recurrent sources. We expressed stChrimsonR selectively in excitatory neurons in V1 (AAV-FLEX-stChrimsonR; Emx1-Cre mouse line) and used two-photon imaging, extracellular physiology, and widefield imaging to measure cortical responses to excitatory cell stimulation.
shown in Fig. 1 Suppl. 1D). We found a salt-and-pepper mix of suppressed and excited cells (Fig. 1B,C), with suppression especially visible during the later part of responses to long stimulus presentations of the oriented noise stimuli (Fig. 1C). In other words, in response to both types of visual stimuli, we found some cells that responded with strongly elevated steady-state responses, and other cells that showed suppressed steady-state responses (Fig. 1C–F, Fig. 1 Suppl. 1).

Deconvolving fluorescence responses to yield a proxy measure of spike rate confirmed this salt-and-pepper pattern with substantial numbers of suppressed neurons intermingled with excited neurons (Fig. 1D,F-G).

Neurons showed sustained responses during the oriented noise stimuli (Fig. 1E), and deconvolving the timecourses revealed an initial transient in excited cells (Fig. 1F, population responses shown in Fig. 1G), followed by a steady-state period.

The viral expression strategy we used for these experiments results in both excitatory and inhibitory neurons that express GCaMP. However, the large fraction of suppressed neurons (Fig. 1C, 40% of responding neurons, Wald CI 29–51%; population: 44%, Wald CI 35–53%) implies that many excitatory neurons were suppressed, and it was not just that a group of inhibitory neurons was suppressed by stimulation. Below, we confirm with electrophysiology and imaging that optogenetic excitatory input produces suppression in many excitatory cells.

Optogenetic excitatory drive also results in sparse and distributed suppression

To probe the influence of recurrent excitatory-inhibitory circuits on local response properties, we next measured V1 responses while optogenetically stimulating excitatory cortical cells. Direct stimulation allows us to exclude some feedforward mechanisms for suppression – for example, to argue against the possibility that cortical suppression is generated principally by suppression of thalamic inputs29.

We injected a Cre-dependent excitatory opsin (soma-targeted ChrimsonR, or stChrimsonR) in layer 2/3 of a mouse line expressing Cre in excitatory neurons only (Emx1-Cre30), and expressed GCaMP7s in all neurons with a second virus (AAV-hSyn-GCaMP7s).

With optogenetic stimulation we also found a clear salt-and-pepper distribution of elevated and suppressed responses (Fig. 2C,E,G). We saw this pattern in imaging experiments with two different stimulation pulse lengths. First, we used short (200 ms) stimulation pulses and measured cells’ responses after stimulation onset to avoid optical artifacts (Fig. 2B,C). We also saw the same interspersed pattern in other imaging experiments where we used long (4 sec) stimulation pulses and imaged during stimulation (Fig. 2D,E).
Figure 2: Salt-and-pepper distribution of elevation and suppression to optogenetic excitation shown using 2-photon imaging. (A) Experimental setup. Animals expressing both GCaMP7s and stChrimsonR (both via AAV) were imaged using 920 nm 2-photon scanning laser, while stChrimsonR was stimulated with 595 nm light. (B) Example field of view, showing GCaMP fluorescence. Emx1-Cre animal, injected with AAV-DIO/FLEX-stChrimsonR and non-Cre dependent AAV-GCaMP7s. (C) Steady-state response to 1-photon stimulation (200 ms duration) Red: elevation of firing rate relative to baseline, blue: suppression. Data is dF/F measured after stimulus offset, deconvolved, with peak response for each cell normalized to peak dF/F% for that cell. Same field as in B, stChrimsonR expression not shown. (D) Field of view from a second animal stimulated with long (4 sec) optogenetic pulses; stimulation during imaging flyback (gray region at edges) to avoid contaminating imaging responses and allow stimulation during imaging. Data analysis same as for C but responses collected during stimulation (stimulation delivered on horizontal i photon scanning laser, while stChrimsonR was stimulated with 595 nm light. (E) Deconvolved response of cells in (D) during the steady-state period, colored similarly to (C). Salt-and-pepper suppression is evident. (F) Increasing power leads to stronger elevation and suppression (steady-state response) in their respective populations, as expected. Shaded region: SEM across cells. (G) Spatial distribution of elevated and suppressed cells, N = 3 animals, elevated cells n = 246, suppressed n = 139 cells (one sample t-test, p < 0.05). (H) Population timecourses for all cells in (E), Red shaded region: optogenetic stimulation period. Steady-state response calculated over 200 ms to 3750 ms post stimulus onset. Light lines: individual cell traces, heavy lines: population averages. Shaded region (largely obscured by thick lines): SEM across cells. (I) Selective expression of opsin in excitatory cells only, as expected for the double-inverted lox-site AAV vector and excitatory Cre mouse line (Emx1-Cre). The stChrimsonR opsin was fused to mRuby2, so we measured mRuby2 mRNA (red) and VGLUT1 mRNA (white), a marker of excitatory cells, via fluorescent in situ hybridization (RNAscope; Methods). Cell counting showed 76% of neurons are VGLUT1 positive (N = 195/257). Arrows highlight a few example neurons. As expected, all cells that express the opsin are excitatory, but not all excitatory neurons express the opsin (59% of VGLUT1 cells are mRuby2 positive: N = 115/195). (J) mRuby2 mRNA (red) and VGAT mRNA (green), a marker of inhibitory neurons. 24% of neurons are VGAT positive (N = 62/257), and zero (N=0/62, Cipoper-Pearson 95% CI 0-0.472), express the opsin.

Deconvolving the imaging responses revealed that elevated populations showed a transient response, then a steady-state response that lasted for the duration of the optogenetic stimulation (Fig. 2H).

These data suggest that the network is being driven to a new steady state or fixed point by input. While there was a slight decay in the excited population’s response at high power (perhaps due to network effects, spike rate adaptation, or opsin desensitization), at moderate stimulation power (1 mW, Fig. 2H), deconvolved firing rates are largely constant while stimulation is on.

We next verified that the opsin we used was expressed only in excitatory cells. We used a mouse line that expresses Cre in excitatory neurons, and a genetic strategy (FLIP/DIO31) that blocks expression in non-Cre-expressing cells. Still, to confirm expression in excitatory neurons, we labeled excitatory, inhibitory, and stChrimsonR-expressing neurons via fluorescence in-situ hybridization (RNAscope, ACD Inc; Fig. 2I,J). Excitatory neurons expressed the opsin (Fig. 2I), but as expected for AAV expression, not all excitatory neurons were positive (59%, N = 115/195, Fig. 2I). None of the inhibitory neurons (24% of neurons in the sample, N = 62/257) showed expression of the opsin (Fig. 2J), demonstrating, as expected from the Cre mouse and floxed virus strategy, that the opsin was expressed in excitatory neurons.

Next, we examined whether this salt-and-pepper pattern exists at larger distances from the stimulation site. Using the same opsin preparation, in awake mouse V1, we recorded neural responses to excitatory cell stChrimsonR activation using a silicon electrode array (Fig. 3A,E).

We found both elevation and suppression across all distances (Fig. 3B–D) and depths (Fig. 3F–H) from the stimulation site, suggesting a similar local salt-and-pepper organization of elevated and suppressed cells that we observed directly in the 2-photon response data.
Across the population of recorded neurons, 56.6% (77 of 136) showed an elevated steady-state response to the optogenetic stimulation, and 36.0% (49 of 136) showed a suppressed steady-state response. Both elevated and suppressed cells, on average, showed an initial (positive) transient followed by a (positive or negative) steady-state response (Fig. 3I,J). The electrophysiological recordings show similar dynamics as the deconvolved imaging timecourses (Fig. 2H), except for one feature: the recordings show an initial brief positive transient in the suppressed cells (Fig. 3B–D, F–H, blue lines; Fig. 3J) not just the elevated cells as in the imaging data. The imaging data likely conceal the initial transient in suppressed cells via the slower timescale of imaging. That is, due to the imaging frame rate (30 Hz; 33 ms frames), the transient would often be contained within one frame where it would be averaged with suppression.
163 **Global spatial patterns arise from trends in local salt-and-pepper suppression**

The neurophysiology data showed some evidence of a larger-scale organization on top of the local salt-and-pepper distribution of elevation and suppression. Over distances of more than a millimeter from the stimulation site, we found that the number of elevated units gradually decreased (Fig. 3K,L; Pearson’s chi-squared test, $\chi^2 = 51.31$, df = 3, $p < 0.001$) and the number of suppressed units gradually increased (Fig. 3M,N; $\chi^2 = 44.83$, df = 3, $p < 0.001$). There was also a similar trend in neurons’ firing rates (Fig. 3 Suppl. 2). Elevated single units showed less elevated firing rate with distance from the stimulation site, and suppressed single units showed...
more suppression with distance from the stimulation site, though the linear trend between
distance and population response was stronger in unit counts than in average population firing
rates (Pearson’s r = -0.11, df = 29, p = 0.56, Pearson’s r = 0.32, df = 46, p < 0.05; Fig. 3 Suppl.
2). Notably, however, the number of elevated neurons did not go to zero even at 1.2 mm from
the stimulation site: only the relative numbers of elevated and suppressed neurons changed.
This suggests that the salt-and-pepper organization we saw with imaging persists across the
cortex.

The trends over distance we saw with physiology, however, gave only a limited view of how
mean population responses varied with distance from the stimulation site. To characterize
population average responses more fully, we turned to widefield, mesoscale calcium imaging. In
this approach, each pixel in the imaging integrates signals from many neurons. For these
experiments, we expressed GCaMP in all excitatory cells using a genetic mouse line to
maximize consistency of GCaMP expression across cortical distance (Fig. 4A; Ai148::Cux2-
CreERT2, or Ai162::Cux2-CreERT2, see Methods). We expressed stChrimsonR also in
excitatory cells (AAV-CamKIIa-stChrimsonR) and stimulated while simultaneously imaging
responses.

We saw clear spatial patterns in widefield imaging, broadly consistent with the spatial trends we
saw in the electrophysiology data. During the initial frame of stimulation (~7 Hz imaging, 140 ms
frame period), we saw an increase in activity both at the center of the stimulation light and
extending some distance outside the center of expression (Fig. 4B,E,H). This first-frame
increase is consistent with the transient response in both elevated and suppressed cells we saw
in electrophysiology (Fig. 3I,J). Transient responses like this, before the network reaches a new
steady-state, are also consistent with neural responses being shaped by a strongly-coupled
recurrent network.

Later in the stimulation pulse, consistent with the large-scale patterns in the electrophysiological
recordings, a center-surround pattern emerged (Fig. 4C,F,I.) The area with maximum
stChrimsonR expression continued to show an elevated response, while a donut-shaped region
around it was suppressed. The activated area in the center reflected the area of expression,
measured with fluorescence imaging of the cortical surface (Fig. 4 Suppl. 2). This late
suppression also seemed consistent with the electrophysiology data, which showed
suppression not in transient responses but in the steady-state response. To examine these
timecourses (Fig. 4D, Fig 4. Suppl. 3), we deconvolved imaging responses to better reflect spike
rate changes. The deconvolution was essential to reveal the suppression because fluorescence
(dF/F) measurements, without deconvolution, integrate spiking-related calcium signals over
time. For responses with this timescale, dF/F measures without deconvolution mix initial
transients with later suppression, potentially hiding one or the other. Indeed, the dF/F signals
showed less suppression (Fig. 4 Suppl. 1). We compared several different deconvolution
Methods and found suppression in all cases (Fig. 4 Suppl 1). The suppressed area of the cortex was largest about 500 µm from the center of our laser stimulus, and extended over 1 mm from the stimulation center (Fig. 4 G–I).

In summary, the physiology and imaging data together support the idea that suppressed and elevated neurons are locally organized in a salt-and-pepper pattern, and that the proportion of suppressed to elevated neurons increases with distance from the stimulation site. This change in the proportion of suppressed cells results in a center-surround pattern that can be seen with population-level imaging, with net suppression in excitatory cells emerging, after an initial positive transient, about 500 µm away from the stimulation site.

Response dynamics support a balanced-state excitatory-inhibitory network that is driven to a new steady state by input

If suppression is due to local recurrent network effects, we would expect excitatory cells to be recruited first by stimulation, and then inhibitory cells should receive inputs from excitatory cells and respond slightly later. After this first few milliseconds, balanced-state models predict that excitatory and inhibitory cells should later show similar response distributions. This is in contrast to weakly-coupled models, or a feedforward inhibition framework, where excitatory and inhibitory populations change firing rates in opposite directions: input drives inhibitory cells to increase their rates, inhibiting excitatory cells, which then decrease their rates.

In the data we saw support for the balanced-state recurrent model (Fig. 5A) — differences in excitatory and inhibitory responses in the first few milliseconds, but later similar distributions of excitatory and inhibitory rates.

We classified cells into putative excitatory and inhibitory classes by waveform (Fig. 5B). We have previously confirmed with in vivo pharmacology that narrow-waveform cells are inhibitory interneurons, likely PV-positive fast-spiking cells, while wide-waveform cells are primarily excitatory neurons. We saw that wide-waveform (excitatory) neurons have a slightly faster onset latency than narrow-waveform inhibitory cells, by approximately 2.5 ms (Fig. 5A, inset,
Figure 5: Response dynamics are consistent with steady-state suppression shaped by a recurrent network mechanism. Units with wide waveforms (excitatory units) have slightly faster onset latencies than narrow (inhibitory) units, as expected when stimulating excitatory cells, but other quantities do not differ across neural populations. (A) Average traces for narrow- (inhibitory) and wide-waveform (excitatory) units. Narrow-waveform units are likely inhibitory, fast-spiking neurons, while wide-waveform units include mainly excitatory cells with a minority of wider-waveform inhibitory neurons. Inset: enlarged view of laser onset, highlighting latency difference between narrow and wide units (green markers: time to half peak.) (B) Wide/narrow sorting approach. The distribution of spike waveforms is bimodal (classification threshold: 0.445 ms.) (C) Onset latencies are slightly shorter for wide (excitatory) units, as expected when stimulating excitatory cells. See Results for test statistics and p-values. (D) No significant difference in initial transient firing rate (peak firing rate) between wide-waveform and narrow-waveform units (medians, wide: 24.5 spk/s, narrow: 31.2 spk/s.) (E) No significant difference in steady-state firing rate between wide and narrow units. For both putative inhibitory and excitatory cells, some units show suppressed rates and some elevated rates (medians, wide: 4.0 spk/s, narrow: 5.1 spk/s.) (F) No difference in onset latency for elevated and suppressed populations. Conventions as in C. (G) No difference in time to steady state for elevated and suppressed populations.

5C; narrow latency 7.9 ms, wide latency 5.4 ms, difference 2.5 ms, Mann-Whitney U = 1256.0, p < 0.01; onset latencies computed via curve-fitting to rising phases, see Fig. 5 Suppl. 1A for details).

An alternative to excitatory and inhibitory cells acting as a balanced-state network (with excitatory and inhibitory cells responding similarly but with slightly different latency) is that suppression was due to a subtype or subpopulation of inhibitory cells whose firing rate went up as excitatory rates went down. If such separate subnetworks existed, we might first expect to see differences in the dynamics of elevated and suppressed cells. But we found no significant differences in onset time or time to steady state for elevated and suppressed cells (Fig. 5F,G, onset time Mann-Whitney U = 1741.0, p = 0.17, time to steady state, Mann-Whitney U = 801.0, p = 0.32.)
Moreover, if there were suppressed and elevated excitatory cells that were part of different subnetworks, instead of acting as part of an overall coupled network, we might expect to see differences in the dynamics of suppressed and elevated sets of wide-waveform (excitatory) cells only. But there were no significant differences in onset time or latency for suppressed and elevated cells when we restricted these analyses to wide-waveform cells (Fig. 5 Suppl 1C,D).

Beyond the differences in onset latency, other response dynamics were not different between excitatory and inhibitory cells. Consistent with a recurrent network with strongly coupled excitation and inhibition, we found that both excitatory and inhibitory cell populations increase their average firing rate when excitatory cells are stimulated (wide mean Δ: 14.54 spk/s, t = 5.52, df = 93, p < 0.001, narrow mean Δ: 13.53 spk/s, t = 3.07, df = 41, p < 0.01). That is, both excitatory and inhibitory populations contain elevated and suppressed neurons, though elevated cells dominate both averages. Further, the initial transient and steady-state firing rate medians were not detectably different between inhibitory and excitatory cells (transient: Mann-Whitney U = 1816.0, p = 0.23, steady-state Mann-Whitney U = 1866.0, p = 0.31, Fig. 5D,E). Also, the time it took for wide-waveform and narrow-waveform cells to settle into their steady states did not differ (Fig. 5 Suppl 1B), consistent with the idea that the steady-state dynamics emerge from integration of both inhibitory and excitatory inputs. Overall, in a strongly-coupled network operating as an ISN, excitatory and inhibitory mean firing rates tend to vary together\(^{11,34}\), consistent with our observation that excitatory and inhibitory populations have similar dynamics and firing rate distributions.

**A balanced-state model with strong coupling explains many features of the data**

Thus far, a moderately- or strongly-coupled balanced-state network seems consistent with both the response distributions and dynamics we observe. Such network models, with excitatory and inhibitory neuron populations that have strong coupling across populations, explain experimental data in many, if not all, cortical areas\(^{10}\). In balanced networks, individual neurons each receive large excitatory and inhibitory inputs, each strong enough that by themselves they would control the neuron — either to fire at high rates or to be completely suppressed. But the two currents sum to a small net value, and the neurons fire based on fluctuations caused by transient shifts in excitatory/inhibitory balance.

The large excitatory and inhibitory inputs in balanced-state cortical models could underlie the suppression we see. Intuitively, stimulation would excite excitatory cells, the excited cells would excite their inhibitory targets, and this would lead to some inhibitory cells excited more than others. In turn, this would cause varying amounts of inhibition across network neurons, which in some cases would lead to suppression. Indeed, recent theoretical work\(^{22}\) has shown that this kind of heterogeneous network response occurs in strongly-coupled cortical networks. To understand if our experimental data could be explained by this mechanism, we examined recurrent network models with features reflecting our data, and determined which features of the recurrent network models were important to explain the suppression.
First, we used the imaging data to determine if the suppression we observed was tightly linked to the variability in opsin from cell to cell. In principle, it could be the case that only the cells that are excited by the opsin would increase their firing rates, while non-opsin cells would be suppressed. This might happen if recurrent excitatory input were small compared to optogenetic input. That is, a tight relationship between opsin and firing rate could mean network effects were not important.

Instead of a tight link, we found a very weak relationship between opsin expression and cells’ firing rate changes. To approximate the optogenetic drive to individual neurons, we measured fluorescence of mRuby2 (fused to stChrimsonR) in donut-shaped regions around each cell’s membrane using two-photon imaging (Fig. 6). The amount of opsin-related fluorescence explained little of the variance in steady-state responses (i.e. the responses were spread out along the y axis, and did not form a tight line; Fig. 6A-C shows single animal example at 2 mW stimulation, Pearson’s $r = 0.19$, df $= 111$, slope of linear fit $> 0$, $p < 0.05$). We found that the weak but positive relationship between opsin-related fluorescence and steady-state response is consistent across the population (Fig. 6D), though the slope trends back towards zero at high...
fluorescence, driven by a few cells that have high fluorescence but show little response, perhaps due to overexpression of opsin in a small number of neurons (excluding cells with high fluorescence, 1 mW: Pearson’s $r = 0.10$, df = 193, $p = 0.17$, 2 mW: Pearson’s $r = 0.17$, df = 193, $p < 0.05$; Fig 6E).

Because the amount of suppression was not well-explained by stChrimsonR expression, we next examined how network effects could lead to suppression. We first sought to confirm that our results are consistent with a balanced-state model (as in22) and that we could replicate these effects within a conductance-based network. We constructed and simulated conductance-based spiking neural networks, varying network connectivity and opsin drive across neurons in these models, and measured network responses to excitatory cell stimulation (Fig. 7A,B).

Each network consisted of two sparsely connected populations of conductance-based spiking neurons, one excitatory (8000 neurons; 80%) and one inhibitory population (2000 neurons; 20%). We used a model with a minimal assumption of only one inhibitory population. The inhibitory population may map onto fast-spiking parvalbumin-positive (PV) cells, which seem likely to be able to play a balancing role in the cortex given their strong input to excitatory cells13,35, consistent with the dynamics and responses we observed in narrow-waveform, likely PV, neurons (Fig. 5). Excitatory and inhibitory synaptic mean strengths were adjusted so the total recurrent excitatory and inhibitory inputs were similar: for every network, we adjusted a background input current to either excitatory or inhibitory neurons to hold the spontaneous firing rate of the neurons at a value (4–6 spk/s) consistent with the data. We drew the opsin input strength to each neuron from a distribution fit to the imaging data (log-normal distribution, Fig. 7C; Methods).

We first manipulated the mean connectivity strength of recurrent connections. We constructed three different models, varying the average strength of recurrent coupling in each (schematic, Fig. 7B). The “tightly balanced” network had the strongest recurrent coupling, and we scaled down the synaptic weights by a factor of 2 or 4 to create more weakly-coupled “50%” and “25% strength” networks.

We checked that the networks showed paradoxical suppression of inhibitory cells, a sign of strong recurrent coupling within the excitatory network and the ISN regime, as observed in visual cortex11,13,36. We stimulated the inhibitory cells in each network and found, as expected, paradoxical suppression (Fig. 7 Suppl. 1).
A schematic: data features

D

high proportion of suppressed cells

E

long-tailed positive response

F

data: ephys

G

data: 2-photon

H

sim: tight balance

I

simulations

J

simulations

K

L

simulations

M

simulations

N

simulations

\( \Delta \) steady-state firing rate

\( \Delta \) steady-state firing rate

\( \Delta \) steady-state firing rate

\( \Delta \) steady-state firing rate

firing rate (spks/s)

\( \Delta \) steady-state firing rate

\( \Delta \) steady-state firing rate

\( \Delta \) steady-state firing rate

\( \Delta \) steady-state firing rate
Figure 7: Strongly-coupled balanced-state recurrent neural network model with heterogeneous connectivity describes experimental data. (A) Simulation design: conductance-based spiking network model with 8000 excitatory cells and 2000 inhibitory cells (2% connection probability between cells, synaptic strength variability chosen to be of same order as mean (Chapeton et al., 2012).) (B) We varied the overall strength of recurrent connectivity by scaling the recurrent synaptic strengths. (C) Optogenetic input strengths (effects of opsin on each cell) were chosen based on the fluorescence values observed in 2-photon imaging data. A distribution function (log-normal) was fit to the fluorescent data and each weight was drawn independently from this distribution. (D) Schematic depicting the characteristic features of the response distribution we have observed. (E) Electrophysiological data and simulated network response to excitatory cell optogenetic stimulation, using input weights chosen to match the data (C). The more tightly balanced networks show a long tail of cell responses similar to the long tail seen in the data. On the right we display the same data but zoomed in on the suppressed (negative) values of the distribution. The network with strongest recurrent coupling shows the most suppression. (F) Cumulative distribution of responses to increasingly strong 1-photon stimulation during electrophysiology experiments. (G) 2-photon experiments, and (H) simulations. (I, left) Mean time course for elevated cells in the strongest recurrent network. (I, right) Mean time course for suppressed cells in the strongest recurrent network. (J) Balance index (Ahmadian and Miller, 2021) of the 3 networks represented in (B). Values between 0–0.1 correspond to tight balance, between 0.1 and 1 loose balance, and above 1 unbalanced, where there is little recurrent contribution to neurons’ responses. The strongest-coupled network (purple) has a median index corresponding to the tight balance regime. (K) Schematic of types of input variability. Number of recurrent connections across cells can vary (left) or the strength of those connections can vary (middle). Variability in number of input (here optogenetic input) can also contribute to response variability. (L) Simulated neural responses to optogenetic input, with (left, same as Fig. 7F) and without (center) variability in number of recurrent inputs. The relationship between optogenetic input and steady-state response strengthens with the variance in number of recurrent inputs held constant ($R^2$ original = 0.29, $R^2$ reduced conn. var. = 0.58, $\Delta R^2 = 0.29$). Further, when recurrent strength variability is reduced, the relationship gets even tighter (R right) ($R^2$ original = 0.29, $R^2$ reduced conn + strength var. = 0.70, $\Delta R^2 = 0.41$) Therefore, when recurrent variability is reduced, each cell’s steady-state response is more strongly dependent on opsin expression. (M) Cumulative distributions of baseline-subtracted steady-state firing rates. Variability from either recurrent or optogenetic sources is sufficient to approximate the long tail of elevated firing rates, however having both best approximates the amount of suppression we observe. “All var.:” network with parameters as in Fig. 7, including variability in input number, variability in optogenetic input weight (fit from data, Fig. 7C), and variability in synaptic strength. Pink, network with variability in number of recurrent connections removed. Orange, network with optogenetic input variability removed (all optogenetic input weights set to a constant). Yellow: both types of variability removed. (Right) Zoom of (M) to the suppressed portion of the distribution. Some suppression still exists if either source of variance is removed, but suppression is effectively extinguished when both sources of variance are removed. (N) Reducing the variance of the synaptic weights by a factor of 10, shows the cumulative distribution becomes strictly positive with little variance. Yellow: network with no connection number variability and no optogenetic input variability. Blue: same as yellow network but with variance in synaptic strength distributions reduced by a factor of 10.

Recurrent excitatory-inhibitory networks can be tightly or loosely balanced, depending on the total amount of recurrent excitatory and inhibitory input to network neurons. To classify the networks, we calculated the balance index, a ratio that measures how completely inhibitory input cancels out the excitatory input for each neuron in the networks (see Methods). We found that all three networks we constructed are balanced as expected due to their irregular spontaneous activity (balance index << 1), and the networks span a range from loose to tight balance (Fig. 7J).

The model replicates the long tail of positive responses, suppressed responses, and aspects of the dynamics

Two characteristic features of the response data we observed are the long-tailed positive response and the substantial proportion of suppressed cells (Fig. 7D). To match the input strength to the model, we scaled the input distribution until the 75th percentile of the model response matched the electrophysiological response to the strongest optogenetic input. All three simulated networks showed a long tail of elevated responses as in the data, with many neurons showing increases in firing rate to stimulation, and a few showing large increases (Fig. 7H). However, the amount of suppression did depend on recurrent coupling strength. Scaling the total excitatory and inhibitory input leads to more suppressed neurons when other network parameters are held constant (Fig. 7E). The network that best fit the fraction of suppressed cells

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we observed was the most strongly-coupled network, which was just inside the tight balance regime (Fig. 7E-J). Excitatory and inhibitory cells’ response distributions were similar in the model, as also seen as in the data (Fig. 7 Suppl 2). Finally, to further demonstrate how well the tightly-coupled model could reproduce the features of our response distributions, we simulated responses to parametrically-increasing strengths of the input distribution, and compared them to the electrophysiological and 2-photon responses to increasingly strong experimental optogenetic stimulation. The shapes of the response distributions in the most tightly coupled model and the data were similar (Fig. 7F–H).

We next examined response dynamics to determine if the dynamics we observed were also consistent with a balanced E-I model. We found that model responses were qualitatively similar (Fig. 7I) to the timecourses of responses seen in the data (Figs. 2–3). Excitatory cells first showed a brief, positive transient response before the network settled into a new steady state, with some cells excited and some suppressed. The biphasic response in suppressed cells (positive transient followed by suppression) is a key data point, as this biphasic response suggests a network mechanism where input initially excites many excitatory cells, but later recurrent inputs lead to suppression of some excitatory cells. A second matching feature of the dynamics is the offset dynamics in both elevated and suppressed cells: after stimulation ends, both show a slight suppression before returning to baseline. A third matching feature is that excitatory cells have earlier onset times, indicating that E cells were directly stimulated and I cells were recruited just a few milliseconds later (Fig. 7 Suppl. 2), before both populations then evolved to a new steady state.

One feature of the dynamics seen in the data but not the model is that for high stimulation powers there is a slight decay during the tonic or steady-state period (Fig. 3I). However, this decay effect is not seen at lower stimulation intensities, suggesting it arises from known opsin dynamics (inactivation at high light power, e.g.37) or other known biophysical, non-network, effects like spike-rate adaptation.

**Neurons’ responses are not tightly related to optogenetic input due to recurrent inputs**

Thus, a balanced excitatory-inhibitory spiking model with one inhibitory population shows the main features of both our two-photon and electrophysiological data. The observations reflected in the model include the distribution of responses as a function of optogenetic input, the dynamics, the long tail of excited neurons, and the suppression of a fraction of excitatory neurons.

In addition to these effects, we also saw another characteristic in the data that can be informative about the model: strikingly little correlation between opsin expression and neural response to stimulation (Fig. 6). This implies that the amount of optogenetic input a neuron receives is not a good predictor of a cell’s response, and further that recurrent input strongly governs neurons’ responses. In fact, the tightly-coupled model showed the same pattern of responses (Fig. 7L, left).

We therefore asked which sources of variability were important to explain why neural responses were weakly related to optogenetic input. To do this we varied sources of input variability in the
model (Fig. 7K). First, we reduced variability in either the number or strength of recurrent inputs and found that this created a tighter correlation between optogenetic input strength and response (Fig. 7L, middle and right) which made the model a worse fit for the data (Fig. 6). The original relationship was not recovered by increasing the recurrent strength of the network (Fig. 7 Supp 3), implying recurrent connection variability was required to produce this effect and higher recurrent strength could not substitute for it. Next, we asked whether variability in opsin input across cells was also essential to explain the suppression we observed. Removing the variance in optogenetic input across cells (so that each network E cell with optogenetic input received the same strength optogenetic input) significantly reduced the number of suppressed neurons and also produced a worse fit to the data (Fig. 7M; right inset highlights suppressed neurons).

If input variability was the primary source of variability in neural responses, then removing variability from both kinds of input — both optogenetic input variability and recurrent input variability — should substantially reduce the amount of suppression observed. This is what we found. Removing or reducing both types of variability produced a set of neural responses clustered tightly around the mean response (Fig. 7N) with no suppressed neurons. Thus, both variability in recurrent input and in optogenetic input are required to explain the data.

In these simulations, we set several parameters as fixed based on data. Those include: the variability of optogenetic input (estimated from Fig. 6) and the spontaneous rate of the networks (setting the spontaneous rates to ~ 5 spk/s, Fig. 3I–J). In each network, we also adjusted the optogenetic input strength to match the data (Fig. 7F–H, matching the 75th percentile of the data rather than the median or mean due to the long positive tail of responses in the data). Once those parameters were fixed, we varied only the mean strength of excitatory and inhibitory connections (Fig. 7E,J), and the variability of recurrent input (Fig. 7K–N) to determine how each affected network responses, finding that variability in recurrent connectivity was needed to describe the data.

Together these results show that both optogenetic input variability and recurrent connection variability help create the variability in the response of different neurons that receive similar optogenetic input. Why do both contribute? Each neuron’s firing rate is affected not just by the optogenetic input that particular cell receives, but also by recurrent input received from other neurons, which themselves receive different amounts of optogenetic and recurrent input. That means the firing rates of the input neurons to a given cell are affected by both the variability in their recurrent connections to the network, and the distribution of optogenetic input they receive. The resulting recurrent input to a given neuron varies from neuron to neuron, and this is what uncouples neurons’ responses from the amount of direct optogenetic input they receive, producing the weak relationship between optogenetic input and firing rate that we observed (Fig. 6).

A balanced-state network model with connection variability also explains expected responses to single cell stimulation

Past work has found that stimulating a single cell in visual cortex leads to average suppression in the surrounding population (averages across space and across orientation yield a negative
mean response, e.g. their Fig. 2D). Our results seem initially to contradict this finding, because our data and simulations both find a net positive response across the population when we stimulate many excitatory cells.

To determine if the effects of single-cell stimulation could also be explained by the balanced-state simulation that describes our data, we performed simulations of single cell stimulation in the same tightly-coupled network we examined previously, measuring the response of the non-stimulated population (Fig 8A,B). We found that, while single cell stimulation produced a range of individual cell responses (i.e. reshuffling, Fig. 8C), the mean response was not negative, but instead close to 0 (Fig. 8C,D; mean firing rate 95% CI [-0.006, 0.021], t (9998) for nonzero mean = 1.13, p = 0.26).

Different neurons in the network showed different responses based on recurrent connectivity, but this was insufficient to explain the negative mean response that results from single-cell stimulation. The excitatory cells that received a direct connection from the stimulated cell (feedforward, FF, cells, n=107 neurons) had an elevated response. Those that received a connection from the inhibitory cells which received a monosynaptic input from the 107 directly connected E cells had a very slightly suppressed response (i.e. E→I connections, or feedback (FB) cells, n=4380 neurons; Fig. 8E). The small set of strongly excited cells compensate for the large number of weakly suppressed cells, leading to a mean response of 0 (Fig. 8F,G). Single cell stimulation alone in our balanced-state model, therefore, could not account for the mean suppression observed in previous studies.

We next hypothesized that this difference could be due to difference in the activation state of the network. Chettih and Harvey (2019) stimulated during visual input, while here we delivered optogenetic input during spontaneous activity. One explanation for this effect could be explained by a balanced-state model: Sanzeni et al. (2022) found that increasing the firing rate of a similar network to our model reduced the mean response of the network. Further, in models of visual cortex with subnetwork connectivity (e.g. higher connectivity between neurons with similar orientation tuning38, it has also been shown that visual input can shift the network response to be negative39. Therefore, to test whether additional network drive could reproduce mean suppressive responses, we simulated single-cell stimulation on top of an input that mimics visual drive (Fig. 8G; Methods). This indeed shifted the mean population response negative (Fig. 8G; firing rate change: mean: -0.05; 95% CI: [-0.07, -0.04]). Further, we show that for single-cell activation, as seen by Sanzeni et al. (2022) for optogenetic activation of many neurons, that the relationship between baseline firing rate before input and network response to input is systematic. The more strongly the network is activated with visual-like input, the more negative the response to optogenetic input (Fig. 8H).

Therefore, we find that a strongly-coupled balanced-state model can explain both our results with stimulation of many neurons (a long tail of positive responses, positive mean, and a substantial fraction of suppressed cells) and the negative mean that results from single-cell stimulation. In both cases, strong mean connectivity, as well as variability in recurrent connectivity shape the responses of the network.
Figure 8: **Single cell stimulation produces elevated firing rates in a small subset of cells, but widespread weak suppression across the population.**

(A) A single cell was chosen and repeatedly stimulated in the network with the strongest coupling (Fig. 8B top). (B) Firing rate distribution during single cell stimulation shows that single cell stimulation only strongly modulates the stimulated cell and perhaps a few others. (C) Cumulative density functions during single cell stimulation and without stimulation (black and green, respectively). While single cell stimulation widens the distribution of responses, the mean remains around 0. (D) Same as (C), but with probability density representations. (E) Two subsets of the E cell population were drawn: cells that receive direct connections from the stimulated cell (n = 107; brown outlined cells), and cells that receive feedback inhibition as a direct result of the stimulated cell (n = 4380; dark blue outlined cells). While cells that receive direction connections from the stimulated cells show a positively shifted distribution with a heavy positive tail, the cells that receive feedback inhibition are biased towards suppression. (F) Bootstrapped means of the distributions mirror the effects described in earlier panels. The whole distribution (excluding the stimulated cell) shows a mean suppressed response (black). The small subset of E cells that receive direction connections from the stimulated E cell show a significantly elevated mean response (brown). The subset of cells that receive feedback inhibition as a direct result of the stimulated cell show an even greater bias towards suppression (dark blue), while the whole population (black) is mean 0. (G) To simulate Chettih and Harvey (2019) conditions more closely, where single cell stimulation was carried out during visual drive, we performed our simulations with various forms of synthetic visual drive by providing the network with large scale input during single cell stimulation. Either the previously modeled 1-photon stimulus was added during single cell stimulation, or all excitatory and inhibitory cells were provided with a uniform input. Adding single cell stimulation on top of our previously modeling 1-photon stimulus (red) leads to a negative bias in comparison to the single cell stimulation condition (black). (H) Parametric manipulation of a uniform input to both E and I cells shows that stronger network drive leads to the effect of mean suppression during single cell stimulation.
Discussion

In this work, we see robust suppression in visual cortex due to direct optogenetic drive to excitatory neurons. Using electrophysiology and 2-photon imaging, we found intermixed elevated and suppressed neurons distributed in a salt-and-pepper pattern. This salt-and-pepper distribution of responses resembles what is observed during visual input and arises without input to inhibitory neurons.

The salt-and-pepper local pattern shows global trends over larger areas of the cortex. With widefield calcium imaging, we saw mean elevated rates at the center of the stimulation site, and mean suppression in a concentric circle a few hundred microns. The suppressed cortical surround region is consistent with past data on surround suppression in vision (e.g.\textsuperscript{40}). Our observations show that cortical suppression can be driven by direct excitatory input to the cortex, suggesting that surround suppression also arises from recurrent interactions. Our data suggest recurrent connections are organized to produce a local salt-and-pepper input pattern of activation and suppression not just at the stimulation location (Fig. 2), but in the cortical surround (Fig. 3). The salt-and-pepper distribution of responses we observe is therefore overlaid on top of the global trends we observed in widefield imaging (Fig. 4) and electrophysiology (Fig. 3).

The response distributions and dynamics of the response to excitatory cell stimulation suggest a network mechanism for the observed suppression. Previous work with balanced-state models with strong coupling shows reshuffling: relatively large responses in individual neurons to excitatory cell stimulation, while the population response distribution remains relatively invariant, seen in both mice and monkeys\textsuperscript{42}. We confirm this mechanism applies in a conductance-based simulation (Fig. 7). Further, Sanzeni et al. (2022) shows that variability in optogenetic (external, non-recurrent) input across cells is required for this reshuffling. By measuring opsins expression (Fig. 6) we further show that variability in recurrent connectivity in the cortex is required to explain our data. Recurrent input variability, combined with external input variability, largely decouples external input strength and firing rate response in individual cells (Fig. 6). This is because the input activates many different neurons, which each provide recurrent input to other cells, and these network inputs have significant effects on neurons' responses.

Intuitively, the network mechanism that creates the salt-and-pepper excitation and suppression is that external inputs first elevate the firing rates of excitatory cells (Fig. 5A-C), some more than others. That activation excites inhibitory cells, also some more than others. The result is the network settles into a new steady state (Fig. 5E–G) with a very broad distribution of excitatory cell firing rate changes (Figs. 6,7). Each individual excitatory neuron receives a different amount of net input (excitatory minus inhibitory) and thus many cells' rates are elevated and at the same time a substantial fraction of neurons are suppressed by stimulation. This mechanism seems likely to underlie variability in visual responses as well (Fig. 1), because recurrent connection variability is present in the cortical network for all kinds of input, and thus is present both for visual input and for the optogenetic input we provide.
Variability in recurrent connectivity in the cortex: experimental evidence

We find that variability in connection strengths between L2/3 neurons creates heterogeneous responses to input. Several observations suggest that the brain has recurrent variability at least as large, and possibly larger, than we assumed in the simulations. First, electrophysiological studies often find a long tail of synaptic strengths between pairs of neurons with a few very large connections, though the variance of individual synaptic weights may be lower, with the larger connection strengths due to multiple synaptic contacts between neurons (though see for evidence of long-tailed synaptic bouton sizes). If there is a long tail in synaptic connection strengths, this would still support our finding of high recurrent variance, as it would increase the recurrent variability even beyond the weight distributions we used (truncated Gaussian with mean and variance equal). Second, we used a connection sparsity of 2%, and generally set the number of inputs a cell received from the recurrent network according to a binomial sum, with fixed connection probability between neurons. Connection probability in the brain may be higher than we used, as for example paired recording studies have found connection probabilities of 10% or higher. And higher connection probability will produce greater variance in net input into different cells, as binomially-distributed sums have a larger variance as connection probability increases (in the 0–50% range). Finally, patterned or subnetwork-specific connections, which we did not include, would also only increase variance, though specific connections seem to have just a moderate effect on connection probability — shared tuning changes the connection probability from 10–20% on average to 30–50% for like-tuned neurons, in some cases. Taken together, the substantial recurrent variability that explains our data is consistent with experimental measurements of recurrent connection variability.

Strong balance, loose balance: implications for models that describe cortical networks

In addition to supporting the idea that recurrent connections between neurons have substantial variability, our results also confirm that the mean V1 recurrent connectivity is strong — i.e. V1 operates an inhibitory-stabilized network, meaning that the excitatory network is unstable if inhibition could be frozen. Within the class of balanced networks, two sorts of balance have been distinguished: “loose” and “tight” balance. The best network in our results is on the border of the tight- and loose-balance regimes, with individual cells falling in either the tight or loose-balance regimes. A network near the transition from loose to tight balance is broadly consistent with past experimental data (reviewed in ref. 10) which do not suggest a very tightly-balanced regime for the cortex.

We find that a two-population excitatory-inhibitory model is sufficient to explain the data we observe. A priori, it could have been that a model with multiple inhibitory subtypes would be needed to reproduce the dynamics and population statistics we saw. Recent work has argued for particular roles for cortical inhibitory subtypes: that parvalbumin-positive (PV) neurons are the primary class providing inhibition stabilization, while somatostatin-positive (SOM) cells are involved in gain control. These separate roles are still consistent with our findings. PV cells are likely to be the primary inhibitory cell class in our data and model, as PV cells are the narrow-waveform cells that we identify in electrophysiology (see Fig. 5). Those cells show dynamic and response firing rate changes expected for the inhibitory population in an E-I model (slightly delayed onset latency, similar distribution of firing rate change as E cells). It is also
plausible that stimulating cortical excitatory cells as we did does not cause gain to vary, so that
a separate gain role of SOM neurons was not evident in our experiments. Thus, a two-
population inhibitory model (with PV cells likely making up a large part of the I population in the
model) seems to be sufficient to explain the suppression we observe.

The mechanism we find for suppression is strikingly different than paradoxical suppression in an
ISN when inhibitory cells are stimulated\textsuperscript{12–14}. In both cases, suppression is paradoxical: here we
excite excitatory cells and see suppression of excitatory cells, and in an ISN, exciting inhibitory
cells causes suppression in inhibitory cells. But in paradoxical inhibitory suppression, the \textit{mean}
firing rate of the inhibitory population decreases\textsuperscript{14}. Here with excitatory cell stimulation, the
mean firing rate change is non-paradoxical: excitatory cell average rates \textit{increase}. It is the
substantial variability or heterogeneity of recurrent connections in combination with variability of
input that causes many cells to be suppressed as others increase their firing. However, both
types of paradoxical suppression, when excitatory or inhibitory cells are stimulated, are only
present when the network operates as an ISN — that is, both effects happen in a network with
strong average recurrent coupling\textsuperscript{13,22}. The observed paradoxical suppression of excitatory cells,
however, requires variability around that strong average recurrent coupling.

\textbf{Recurrent computations: what potential roles in visual cortical function are there for
recurrent connections?}

Our data shows that recurrent connections in V1 affect the steady-state responses of neurons to
input. This is one possible role for recurrent connectivity in visual cortex: amplifying or
suppressing certain input patterns.

Artificial networks, such as deep network models, can perform very complex visual tasks without
recurrent connectivity. In advanced object recognition models, layers and areas connect to other
layers and areas, and the synaptic strengths of these feedforward connections control the
computation. This feedforward computation alone is extremely powerful — in principle, layered
feedforward networks can approximate any function, and can perform arbitrary classification
and transformation.

Why then, do recurrent connections seem to be able to have large effects on responses to input
in visual cortex? The suppression we see may be a signature of a common cortical operation,
normalization\textsuperscript{47}. Normalization suppresses activity of other neurons within a layer when some
neurons increase their firing rates. From a statistical or informational standpoint, normalization
within a cortical layer results in improved coding — that is, units whose responses are less
correlated — in models trained to reflect natural scene statistics\textsuperscript{48–50} (see also\textsuperscript{51} for biological
data). And in artificial networks, normalization is used to keep the response mean or response
distribution similar when some cells are activated by input, which often improves learning\textsuperscript{52}.

Thus, normalization in biological and artificial networks tends to result in small changes in the
mean of population firing rates even as individual neurons show large changes in firing rates,
consistent with what we see (e.g. Fig. 7F-H).

If the suppression we see is a consequence of normalization that improves learning or other
functions, there is still a question of why it is provided by recurrent connections. Normalization in
deep networks for vision is often provided without using recurrent connections, by adding extra feedforward layers (as in ResNet\textsuperscript{53}). Our data imply biological brains, on the other hand, collapse the normalization function into recurrent connections within a layer. Thus, brains seem likely to perform normalization via recurrence, while retaining the ability to perform feedforward computation via the pattern of connectivity between layers and areas. Models of visual cortex with more biological features, like the stabilized supralinear network (SSN)\textsuperscript{54} demonstrate this mechanism — they perform a normalization-like function via recurrent connectivity within a network that also transforms feedforward input.

**Future: subnetworks, computation, and interactions between areas**

These results could be extended in a few ways. First, here we did not consider how subnetwork connectivity between excitatory neurons in the cortex might influence the effects. Ko and colleagues (2011) showed approximately a 2–3 fold increase in probability of connection between V1 excitatory neurons that had similar tuning (orientation or direction) vs those with dissimilar tuning. Adding subnetwork connectivity would not qualitatively change our conclusions: that suppression results from recurrent influences, and that it depends on variability of connectivity within the network. However, future work stimulating within or across subnetworks might change the fraction of cells suppressed, given that input patterns would drive neuron populations with somewhat more or less connectivity with each other and the rest of the network. Cell-specific two-photon holographic stimulation\textsuperscript{16,55–57} seems well-placed to study how patterned activity in one subnetwork affects activity in another subnetwork.

While local collaterals probably contribute the majority of recurrent cortical input, it is possible that long-range, inter-areal, connections could contribute to the experimental results we observe. We simulate a population of excitatory neurons and another population of inhibitory neurons which are coupled to each other. One might assume that all these neurons are within a few hundred microns of each other, as estimates of connectivity falloff show most connections to a given neuron come from neurons within that distance\textsuperscript{6,7}. But in principle, cells in other areas could form part of the recurrent population. This could happen for example if projections from V1 to the thalamus recruited neurons there which connect back to the cortex. Understanding how long-range connections contribute to the local pool of recurrent processing could be achieved with stimulation studies with large scale silencing of other regions. However, our widefield imaging data (Fig. 4) shows that the influence of stimulation extends only a few hundred microns away from the stimulation site, suggesting relatively local influence. Therefore, it seems likely that the recurrent connections in the simulations primarily reflect local connections within V1 to nearby neurons.

**Conclusion**

Here we find paradoxical suppression of excitatory cells in the cortex when excitatory cells are stimulated. A balanced-state model with strong mean recurrent connectivity, and variability in drive across neurons, accounts for the data. These results suggest that a primary purpose of recurrent connectivity in visual cortex is to change the steady-state firing rate of network neurons, beyond just how inputs are transformed by feedforward connections. One purpose of this recurrent cortical transformation may be to perform normalization, which confers
Computational advantages in artificial neural networks and is found throughout many different biological networks. These results are a step forward in explaining how cortical networks fire in response to different patterns of input — a fundamental building block of neuronal computation.

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Author contributions

Electrophysiology data was collected by ZZ. 2-photon data was collected by PKL, AJL, and JFO. Widefield data was collected by JFO. Histological data collection and analysis was performed by ZZ and HCG. JFO and MHH designed and implemented simulation experiments. JFO, MHH, and ZZ curated and analyzed data. JFO and MHH wrote and edited the manuscript.
References


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Methods

Reagent Table

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Animals

All procedures were approved by the NIMH Institutional Animal Care and Use Committee (IACUC). Emx1-cre animals\(^{30}\) (\(N = 10\); https://www.jax.org/strain/005628) were used for 2-photon and electrophysiology experiments. For widefield imaging experiments, Ai162 (\(N = 2\); https://www.jax.org/strain/031562) and Ai148 (\(N = 1\); https://www.jax.org/strain/030328) animals\(^{58}\) were crossed with the Cux2-CreERT2 line\(^{59}\) (https://www.mmrrc.org/catalog/sds.php?mmrrc_id=32779), and GCaMP6f or GCaMP6s was induced via tamoxifen injection during adulthood (P22 or later, tamoxifen 2 mg intraperitoneally daily for 3 days). All animals were singly housed on a reversed light cycle.

Implants and injections

Details of the headplate and window procedures are described in previous studies\(^{13,60}\). Optical glass windows (3 mm diameter) were placed over V1 (center: -3 mm ML, +1.5 mm AP, relative to lambda) for 2p and widefield imaging. Windows were also used before electrophysiology for imaging to localize V1.

For Emx1-cre animals, 300 nL of AAV9-syn-jGCaMP7s-WPRE (RRID:Addgene_104487) and/or AAV9-Syn-DIO-stChrimsonR-mRuby (RRID:Addgene_105448) were injected 250 μm below the dura (200 nL/min) prior to cementing the cranial window. For Ai148 and 162 animals, AAV9-CamKIIa-stChrimsonR-mRuby\(^{2}\) was generated by cloning the CaMKIIa promoter (RRID:Addgene_120219) into a pAAV backbone containing stChrimsonR-mRuby\(^{2}\) (RRID:Addgene_105447) and packaged into an AAV (Vigene, Inc.). This was injected at the same depth as the hSyn-DIO-stChrimsonR virus, but with 100 nL volume at 100 nL/min.
Electrophysiology

Electrophysiological methods are described in detail in previous studies, and are summarized here. Animals were headfixed during recording. Before the first session of electrophysiology, the animal’s cranial window was removed and the craniotomy was flushed with saline. Between recording sessions, the craniotomy was covered using Kwik-Sil polymer (WPI, Inc.). A fiber optic cannula (400 μm diameter, 0.39 NA, Thorlabs) was placed to center light output at the center of stChR2mR expression. For light intensity calculations, spot area was defined as the area inside the 50% contour of light spot intensity on the cortex, measured with a camera by imaging the spot on the brain surface. 1–2% agarose (Type IIIA, Sigma) was placed over the dura at the start of each session, and an array of four electrodes (4 probes, 32 sites in total, part #A4x8-5mm-100-400-177-A32, NeuroNexus, Inc.) were lowered into the cortex using a micromanipulator (Sutter MPC-200). One probe was placed at the center of the light spot. Probes were advanced 600–1000 μm below the point in which the first probe touched the dura. Probes were not moved for 1 hour prior to recording, as we found this improved recording stability. Recording data was sampled at 30 kHz (Cerebus, Blackrock Microsystems).

Optogenetic stimulation was performed with randomly interleaved stimulation light pulses with several intensities over the range 0.2 mW/mm² to 15 mW/mm². Stimulation pulses were 600 ms long and delivered with a 4 s period.

For spike recordings, waveforms (bandpass filtered, 750 Hz – 7.5 kHz) were digitized and saved by storing a short data section around points where amplitude exceeded 3 times the RMS noise on that channel. Single units were identified (OfflineSorter, Plexon, Inc) based on clusters in the waveform PCA that were separate from noise and other clusters, had unimodal spike width distributions, and inter-spike intervals consistent with cortical neuron absolute and relative refractory periods. A single-unit score was assigned to each unit manually based on these factors. To compare these populations quantitatively, we calculated SNR for both single and multiunits. Median SNR for single units was larger than median SNR for multiunits (SU: 3.32, MU: 2.26; Fig. 3 Suppl. 1A), consistent with prior reports.

Visual stimulation

Visual stimuli were presented using MWorks (https://mworks.github.io/). Grating stimuli (sinusoidal contrast variation, 0.1 cyc/deg, orientation = 0 deg) were masked with a circular raised-cosine envelope (15 deg FWHM.) Visual stimuli were displayed on an LCD display, with center positioned 0–10 degrees of visual angle temporal to the central meridian. The oriented noise stimuli were generated by filtering white noise pixel arrays (each pixel drawn independently from a uniform distribution) with a spatial band-pass filter (peak orientation = 0 deg, orientation bandwidth = 10 deg, peak spatial frequency = 0.05, frequency bandwidth = 0.05). Frames were generated at 60 Hz and the noise pattern was independent from frame to frame. Visual stimuli were presented for 3 or 5 seconds, depending on the experiment.

2-photon imaging

During 2-photon experiments, animals were awake, water-scheduled, and given periodic water rewards (20% probability per trial, reward once every 30 s on average). If animals stopped...
licking in response to the rewards, data collection was ended. We imaged GCaMP7s responses (920 nm excitation) with either a galvo-galvo (5 Hz) or resonant scanning (30 Hz) two-photon microscope. stCrimsonR-mRuby2 expression was imaged at 1000 nm. The microscopes used for imaging were built using MIMMS components (https://www.janelia.org/open-science/mimms-21-2020) and other custom components, built in-house or provided by Sutter Inc. A second light path, combined with the 2p stimulation light path before the tube lens using a dichroic, was used to stimulate stCrimsonR using 530nm light (CoolLED, pE-4000). For 200 ms long optogenetic pulses, we measured responses in the first frame after stimulation. For 4 s long optogenetic light pulses, we imaged while stimulation was ongoing. To do this, we avoided stimulation artifacts by stimulating only during horizontal flyback (approximate pulse duration 19 µs, off time 44 µs, duty cycle 30%, line rate 8kHz). Optogenetic pulse period was 6 s.

Widefield imaging
For widefield imaging experiments, we used Ai162;Cux2-creERT2 or Ai148;Cux2-creERT2 animals, expressing GCaMP6f or 6s in L2/3 excitatory cells. Animals were headfixed and awake during widefield imaging experiments. Prior to imaging, a fiber optic cannula was aimed at the center of the focal stCrimsonR expression. Images were collected using a Zeiss microscope (Discovery V12) with a 1.0x objective using excitation light with wavelength centered at 475 nm (Xylis X-Cite XT720L). A Zyla 4.2 sCMOS camera (Oxford Instruments) collected images (100 ms exposure time, approximately 140 ms frame period) with 4-pixel binning. Laser powers were randomly interleaved, with 50 repetitions per laser power. Laser pulses were 600 ms long, and presented with 6 s period.

Analysis of electrophysiology data
For spike rate plots, spike counts were binned (1 ms bins), and smoothed via LOWESS 84. To classify units as having elevated or suppressed responses, we measured spike rate over 145–400 ms after stim onset, relative to baseline (-1020 ms–0 ms relative to stim onset) for 6 mW/mm² stimulation intensity. To classify cells as wide- or narrow-waveform, we used a spike width threshold of 0.445 ms based on the bimodal distribution of waveform widths (Fig. 5B). This threshold is consistent with pharmacological segregation of inhibitory and excitatory cells 13.

For analysis of onset times, we fit a sigmoid (logistic function) to each cell’s response from 100 ms before to 100ms after laser pulse onset:

\[ f(x) = \frac{L}{1 + e^{-k(x-x_0)}} + b \]

L: upper asymptote, b: lower asymptote, k: slope, \( x_0 \): onset latency. \( x_0 \) was constrained to the range [onset+0.5 ms, onset + 30 ms]. We defined onset latency as \( x_0 \), the time to half-max. To estimate the time to steady-state, the same function was fit to data from 500 ms before and after the laser onset, with the spike rates within a 50 ms window around the initial transient blanked by setting to the baseline firing rate. Each cell’s time to steady-state was computed as the difference between the steady-state onset and the initial onset (difference between the \( x_0 \) parameters of the two fits).
**Analysis of 2-photon data**

For short optogenetic stimulation (200 ms pulses) during two-photon imaging, we avoided stimulation light influencing imaging responses by measuring responses in the frame after the stimulus offset. For long pulses (4 s), we stimulated during imaging frames by restricting stimulation to imaging line flyback and intensities we give are the average intensity, corrected for the 30% stimulation duty cycle. Because we found that the LED device we used for stimulation (pE-4000, CoolLED Ltd; specified bandwidth 100 kHz) had some variability in onset/offset for each line, we removed pixels (~40% of frame) at left and right edges of field of view to ensure no stimulation light could affect images. Image frames were motion corrected using NoRMCorre through the CalmAn processing suite\(^66\). Deconvolution was done with OASIS\(^66\) via CalmAn. To ease interpretability of the deconvolution signals, each neuron’s deconvolved signal was normalized to have the same maximum value as the dF/F of the corresponding fluorescence trace. To segregate populations into elevated and suppressed cells, we performed a one sample t-test (\(\alpha = 0.05\); testing if cell response mean was larger or smaller than 0) on the deconvolved dF/F during the stimulation period (long pulses) or the frame just after the stimulation period (short pulses). For the short pulses, we used the frame just after stimulation to estimate responses for each neuron per trial. For the long pulses, we averaged data within the period 750 ms after stimulus onset to the stimulus offset in order to capture the steady-state response.

For visual response data, data were preprocessed in the same manner as the short optogenetic stimulation experiments. In order to characterize steady-state responses, the averaging period started 750 ms after stimulus onset and ended at stimulus offset.

**Analysis of widefield imaging data**

Widefield fluorescence images were motion corrected for rigid translation, and any linear trend across the full imaging session was estimated via regression and subtracted. Deconvolution was done via Widefield Deconvolution\(^67\), which differs from single-neuron deconvolution algorithms like OASIS by dropping the sparsity assumption that is useful for deconvolving spike trains of single neurons. This algorithm produces better results for aggregated signals, such as that from a single pixel during widefield imaging\(^67\). We rescaled the deconvolved signals to the maximum dF/F of the imaging data, as with the two-photon deconvolution. Comparison of Widefield Deconvolution, OASIS, and first-differencing is given in Fig. 4 Suppl. 1. To average images across animals, images were aligned on the basis of their maximum response during the first frame of the stimulation period. For quantification of spatial falloff (Fig 4G–I), we averaged pixels in concentric circles around the peak response and fit a LOWESS line to these radial distributions. Crossing points are the minimum distance at which the 95% confidence interval contains zero.

**Spiking network model**

We simulated a conductance-based neural network model with 10000 neurons (8000 excitatory, 2000 inhibitory) to understand the recurrent features that contribute to the response properties we observe during excitatory cell stimulation. Simulations were performed using Brian2\(^68\).
Membrane and synaptic dynamics evolve according to the following equations:

\[ C \frac{dv}{dt} = g_L(E_L - V) + g_E(E_E - V) + g_I(E_I - V) + g_{background} + c \cdot I_{opto}(t) \]

\[ \frac{dg_E}{dt} = -g_E/\tau_E \]

\[ \frac{dg_I}{dt} = -g_I/\tau_I \]

Each synapse was stepped by its corresponding connection weight for every presynaptic spike.

Connections between neurons were made with 2% probability, independently for each potential connection. This sparsity is lower than that observed in vivo (2% vs ~10%)\(^ {42,43} \), though this value has been used before in studies of spiking neural networks\(^ {69} \). Probability of connection can vary dramatically across distance\(^ {42} \) (though see\(^ {70} \)), and unconditional estimates vary across different studies\(^ {38,42} \). We chose a low probability to balance what has been observed in cortex with producing networks that show important physiological characteristics, such as asynchronous firing and low signal correlations.

Synaptic weights were drawn from truncated (rectified) Gaussian distributions. Allowing synaptic weights to vary allows us to further assess how variability in recurrent input may impact the variance in response to stimulation of excitatory cells. Mean connectivity parameters were selected to produce irregular and asynchronous firing. Background currents were chosen for inhibitory and excitatory cell populations to fix baseline firing-rates for each constructed network to be between 4–6 Hz. Network parameters are shown in Table 1.

Table 1: Spiking neural network model parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \tau_E )</td>
<td>5 ms</td>
<td>Mean ( W_{EE} )</td>
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<td>( \tau_I )</td>
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</tr>
<tr>
<td>( g_L )</td>
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<td>I cell ( I_{background} ) Full Network</td>
<td>150 pA</td>
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<tr>
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<td>E cell ( I_{background} ) 50% Network</td>
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<tr>
<td>Variance ( W_{EE} )</td>
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<td>I cell ( I_{background} ) 50% Network</td>
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<tr>
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<tr>
<td>Variance ( W_{EI} )</td>
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<td>I cell ( I_{background} ) 25% Network</td>
<td>193 pA</td>
</tr>
</tbody>
</table>

1-photon stimulation simulations
Optogenetic stimulation was provided as an additional constant current for the length of the stimulation period, with onset and offset ramped linearly over 3 ms. The strength of the optogenetic stimulation (c in Eq. 1) could vary across cells, and was initially chosen from a lognormal distribution fit, scaled between 0 and 1 (Fig. 7C) to our in-vivo fluorescence estimates of stChrimsonR expression, and later held constant (Fig. 7M,N). For each network, this stimulation distribution was scaled by a constant in order to reproduce the same response rate at the 75th percentile across excitatory cells.

Steady-state response was measured for each cell as their firing rate during the 1 s baseline period subtracted from the firing rate during the last 500 ms of the stimulation period.

To reduce connection strength variability (Fig. 7N), we reduced the variability of the truncated Gaussians that define connection strength by a factor of 100 (setting both the synaptic strength variability and connection number variability to zero produced a network that was less stable).

**Single cell stimulation simulations**

Single cell stimulation simulation experiments were performed in the network we constructed with the strongest recurrent coupling (described above). A single cell was chosen and stimulated at the maximum value of the stimulation distribution previously used to push the network to a matched 75th percentile point. This was done for 10,000 trials. A set of control experiments were performed with the same parameters but with no stimulation. We then compared response distributions across the stimulated distribution and the control distribution.

To further examine the effects of single cell stimulation, we segregated the E cell population into two subsets: cells that receive a projection stimulated cell, that is: cells that have a non-zero connection weight from the stimulated cell, and cells that receive projections from the inhibitory cells that the stimulated E cell projects to.

To simulate single cell stimulation under similar conditions to Chettih and Harvey (2019) with visual input, we provided single cell stimulation during either the log-normal 1-photon stimulation, as previously described, or during uniform input of both excitatory and inhibitory cells.

**Balance index**

We computed the balance index as described by Ahmadian and Miller (2021). For each neuron, we computed this index as the net current (excitatory + inhibitory) divided by the excitatory current. The index becomes smaller as balance becomes tighter, with component currents becoming larger, and the index becomes larger as inhibitory input from the network shrinks.
**Supplemental Figures**

**Fig. 1 Supplement 1**

- **A**
  - Oriented noise (stimulus: small, FWHM = 15 deg, stimulus approximately aligned to cells' receptive fields measured outside this experiment, same animal as in Fig. 1), corresponding to the deconvolved cell responses shown in Fig. 1B. Here and in Fig. 1B, responses are measured beginning 750 ms after stimulus onset to focus on steady-state response (Methods.) Evidence of suppression is seen here but is more evident when data is deconvolved (compare this panel to Fig. 1B), as expected for sustained suppression preceded by a transient, as the initial transient seen in Figs. 2, 3, 7.

- **B**
  - Drifting gabor patch (spatial frequency 0.1 cpd, FWHM 15 deg), from the same animal, showing cells that are elevated and suppressed in response to drifting gratings. Overall pattern of responses to noise stimulus and grating is similar.

- **C**
  - Deconvolved population response to oriented noise stimulus (replicated from Fig. 1F for comparison.) Stimulus on during time indicated by light red shaded box.

- **D**
  - Deconvolved cell responses to Gabor patch, same data as in (B). Gabor patches drive both steady-state elevation and suppression, though show signs of stronger off responses and potentially a larger onset transient.

- **E**
  - Population deconvolved response to oriented noise stimulus in two additional animals, consistent with effects from example animal.
Fig. 3 Suppl. 1: Sorting and quality of electrophysiology data (A) Single units demonstrate higher SNR (N=136, median = 3.32) than multi-units (N=184, median = 2.26). (B) Mean spike-waveform of putative excitatory units (wide) in red (N = 94), mean spike-waveform of putative inhibitory units (narrow) in blue (N = 42.) Bimodal histogram of spike widths is shown in Fig. 5B.
Fig. 3 Supplement 2

Steady-state firing rates of neurons in layer 2/3 follow a weak spatial gradient with similar trends as the spatial distribution observed in cell counts. (A) Elevated cell steady-state rates, with the highest and lowest powers for comparison. Rate is the difference in firing rate during stimulation relative to baseline. (B) Suppressed cell steady-state rates, with the highest and lowest powers for comparison, measured in relation to decreases from baseline.
Fig. 4 Suppl. 1: **Center-surround organization is present regardless of deconvolution method.**

(A) Mean whole-frame dF/F GCaMP response in an example animal. (B) We tested 3 different methods of deconvolution, OASIS, first-differences (i.e. subtracting one frame from the previous), and Widefield Deconvolution. Widefield Deconvolution is expected to be the best method, as it is designed for data like this and does not incorporate the sparse-event constraints of OASIS, which is designed for single neurons. We found similar time-series results for each of the methods. The first-differences method (i.e. deconvolution with an kernel that decays immediately) seems to overestimate decreases in firing rate, as might be expected. All panels use the same dataset. (E, F, G) Spatial distribution of responses during the early laser period. All deconvolution methods produce a qualitatively similar excitatory response during this early period. (H, I, J) Spatial distributions of responses during late laser period demonstrates slight differences in size of surround, but overall a qualitatively similar center-surround organization with all methods. (K, L, M) Spatial distributions of response during the late laser period, but with dashed contours depicting the manually-drawn regions of interest (ROIs) that we used to produce the time-series data in (B, C, D), with red dashed contours representing the center ROIs, and blue dashed contours representing the surround ROIs.
Fig. 4 Supplement 2

(A) Example animal's response during the late stim period. (B) Example animal's stChrimsonR expression pattern (gray: fluorescence) with overlaid contours of fluorescence intensity. (C) Example animal's response during the late stim period overlaid with their stChrimsonR contours.

Stimulation response correlates to the pattern of stChrimsonR expression.
Fig. 4 Supplement 3

**Figure 4 Suppl. 3:** Population time course of normalized center and surround deconvolved GCaMP response to laser stimulation (N = 3). Error represents the standard error to the mean across the 3 animals.
Fig. 5 Supplement 1

**Fig. 5 Suppl. 1:** Differences in dynamics are restricted to those seen between the onset of wide- and narrow-waveform cells. The excitatory and inhibitory (wide- and narrow-) onset latency difference is shown in Fig. 7C. Other quantities shown here do not differ: wide- vs narrow (excitatory vs inhibitory) time to steady state (B), and onset time and time to steady state (C,D) for elevated and suppressed groups of wide-waveform excitatory cells. (A) Example single neuron firing rate with fits. To obtain the onsets for individual cells, each cell’s mean timecourse was smoothed with width dependent on the detectability of the transient signal (SNR; Methods), then a logistic function was fit to data from time range [-100ms, 100ms]. The onset time (latency) was defined as the time to half-max of the logistic function. (B) No difference in median time to steady state was found across narrow-spiking and wide-spiking cells. (C) No difference in median onset time for elevated and suppressed groups of wide-waveform (excitatory) cells. (D) Same as B, but difference in median time to steady-state.
Fig. 7 Supplement 1

(A) To examine paradoxical suppression, we record the steady-state responses of the I cells in response to different levels of stimulation. We performed this experiment on all networks presented in Figure 8. (B) Steady-state responses of the I cell population to 5 levels of stimulation. Error bars are standard error to the mean. Graph has been zoomed into the region which clearly shows paradoxical suppression in all 3 networks. This paradoxical suppression is predicted for both loosely and tightly balanced networks. Our simulations used three recurrent strength values, one in the tight-balance regime and two in the loose-balance regime, and we confirmed that all three showed paradoxical effects of suppression when I cells are stimulated (C) Same as (B) but input normalized by the input value calculated in Fig. 8 to drive each network to the same firing rate (input level that achieves same value of the 75th percentile of evoked rates; see Fig. 8.)
Fig. 7 Supplement 2

**Inhibitory neurons in balanced state model show similar responses to excitatory neurons but are recruited after initial stimulation.** (A) Mean timecourses for elevated and suppressed inhibitory cells (left and right, respectively) show the same characteristic transient response followed by steady-state responses. (B) Cross-correlation analysis of E- and I-cell response. Network has no synaptic delays built into the model. E-cells respond to direct stimulation, and then I-cells are recruited after. (C) Population distribution of steady-state responses is similar across E- and I-cells, though excitatory cells show a slightly longer-tailed positive response (true in the data as well; Fig. 5E), as seen through the distribution of responses or their corresponding CDFs (D).
Fig. 7 Suppl. 3: Increasing strength of recurrent connections does not substitute for recurrent connection variability. (A) Cumulative distribution of responses to optogenetic stimulation in model with 1x recurrent strength, matched to the 75th percentile of the response measured using electrophysiology. Negligible recurrent variability in this simulation (same number of recurrent connections to each neuron, variability in recurrent strength ~1% of mean, see Methods), and so spread in responses as a function of input is due to optogenetic input variability. Distribution of input is inferred from data in Fig. 6 (lognormal fit; Methods.) (B) Same as (A) but in model with 2.5x recurrent strength. (C) Relationship between input and steady-state response in the model with 1x recurrent strength. Marginal distribution of response show on the right. (D) Same as (C), but in model with 2.5x recurrent strength. Note that both stimulations produce similar variability between input strength and firing rates. This variability is seen as spread in the red cloud of points around an imagined curve that could be fit through the points. (E) Estimated mean absolute error of the relationship between the input and output as measured by a LOWESS fit across all recurrent strength manipulations.