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All-optical electrophysiology reveals the role of lateral inhibition in sensory processing in cortical layer 1

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Abstract

Cortical layer 1 (L1) interneurons have been proposed as a hub for attentional modulation of underlying cortex, but the transformations that this circuit implements are not known. We combined genetically targeted voltage imaging with optogenetic activation and silencing to study the mechanisms underlying sensory processing in mouse barrel cortex L1. Whisker stimuli evoked precisely timed single spikes in L1 interneurons, followed by strong lateral inhibition. A mild aversive stimulus activated cholinergic inputs and evoked a bimodal distribution of spiking responses in L1. A simple conductance-based model that only contained lateral inhibition within L1 recapitulated the sensory responses and the winner-takes-all cholinergic responses, and correctly predicted that the network would function as a spatial and temporal high-pass filter for

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Author contributions: L.Z.F., A.E.T. and A.E.C. conceived and designed the study. L.Z.F. designed and conducted all the experiments except for patch clamp, built the optical system, programmed the software, analyzed the data, constructed conductance-based model, performed network simulations and co-wrote the manuscript. S.K. and H.W. built the optical system in early stage. U.B. performed simulations of passive membrane model and data fitting. K.D.P. performed patch clamp for SomArchon. K.D.P. and E.S.B. shared SomArchon. M.E.X. helped analyze the data and performed simulations of composite movies. V.P. helped with slice experiments. Y.H. performed patch clamp for GtACR2. K.E.E. helped clone GtACR2. A.E.C. performed network simulations, supervised the research and co-wrote the manuscript with input from A.E.T..

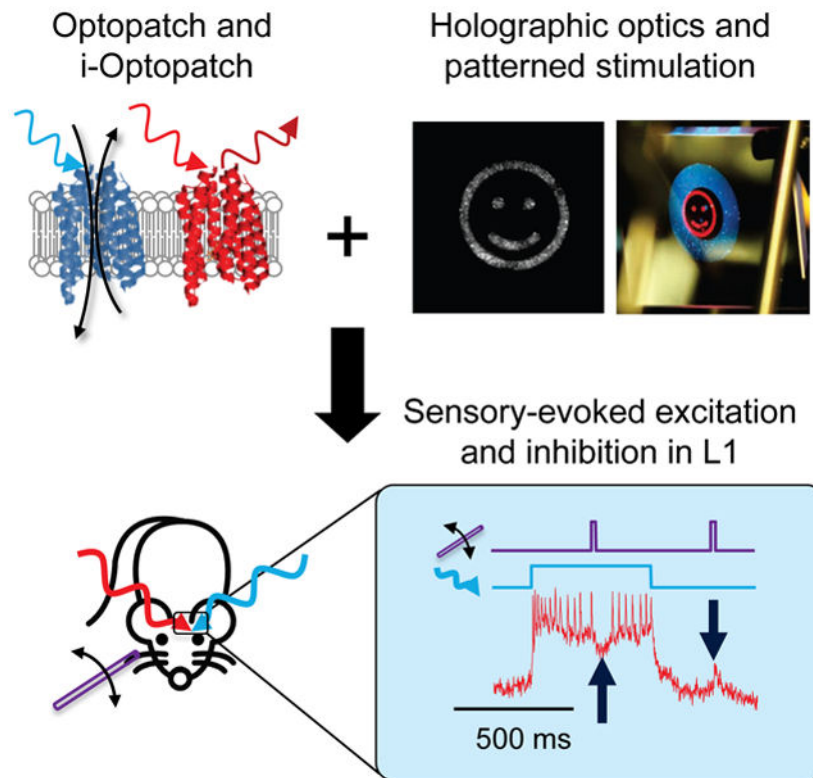
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excitatory inputs. Our results demonstrate that all-optical electrophysiology can reveal basic principles of neural circuit function *in vivo* and suggest an intuitive picture for how L1 transforms sensory and modulatory inputs.

Graphical Abstract



In brief

By simultaneously combining genetically targeted voltage imaging with optogenetic modulation of neuronal activity, Fan et al. demonstrate that all-optical electrophysiology in awake animals can be a powerful tool for revealing hidden principles of neural circuit function.

Keywords

Layer 1; barrel cortex; lateral inhibition; acetylcholine; optical electrophysiology; E/I balance

Introduction

The brain receives myriad sensory inputs. It must distinguish the relevant from the irrelevant. An input can merit attention either through its intrinsic properties (novelty, salience) or through learned associations. The sparse interneurons of neocortical Layer 1 (L1) have been hypothesized as a hub for integrating these factors and modulating the underlying cortex to gate sensory processing (Jiang et al., 2013; Letzkus et al., 2011; Takesian et al., 2018). L1 interneurons receive thalamic (Cruikshank et al., 2012; Takesian et

al., 2018; Zhu, 2004), cortico-cortical (Lee et al., 2013; Palmer et al., 2012), and neuromodulatory (cholinergic (Letzkus et al., 2011; Poorthuis et al., 2018; Takesian et al., 2018), serotonergic (Lee et al., 2010; Poorthuis et al., 2018) and adrenergic (Lam and Sherman, 2019)) inputs. The primary outputs of L1 appear to be via inhibition of deeper-layering interneurons and thereby dis-inhibition of pyramidal neurons (Jiang et al., 2013; Lee et al., 2013; Letzkus et al., 2011; Takesian et al., 2018), and via inhibition of apical dendrites of deeper layer pyramidal cells (Abs et al., 2018; Lee et al., 2015; Takesian et al., 2018).

Electrophysiological, genetic, and anatomical studies have identified four sub-types of interneurons within L1 (Schuman et al., 2018). These cells fall into two broad classes (Jiang et al., 2015). Laterally projecting neurogliaform cells primarily synapse within L1 and show late-spiking non-adapting firing pattern in acute slices (Cadwell et al., 2016; Chu et al., 2003; Cruikshank et al., 2012; Schuman et al., 2018). Downward projecting single bouquet-like cells primarily dis-inhibit underlying pyramidal neurons and show a non-late spiking adapting firing pattern (Jiang et al., 2013; Lee et al., 2013; Letzkus et al., 2011; Takesian et al., 2018). It is not known how these properties combine in awake animals to transform L1 inputs to outputs. Indeed, the function of L1 has been called a “crowning mystery” (Schuman et al., 2018).

L1 is unique among cortical layers for having purely inhibitory short-range connectivity. In mutually inhibitory networks, competition between concurrently activated units can lead to winner-takes-all or point-attractor dynamics (Koyama et al., 2016). This motif has been proposed as a general means to implement multiple forced-choice calculations, i.e. in which small disparities in excitatory inputs are amplified to select one among multiple possible firing modes (Machens et al., 2005). Considering the proposed role of L1 in gating sensory processing—where multiple inputs compete for attention—it is interesting to ask what kinds of dynamics arise in this circuit *in vivo*.

Electrophysiological studies in L1 *in vivo* have been challenging due to the sparseness of neuronal cell bodies. While a few whole-cell patch clamp recordings have been performed in anesthetized rats (Egger et al., 2015; Jiang et al., 2013; Zhu, 2004), technical difficulties prevented similar acquisitions in awake animals. Recent advances in genetically encoded voltage indicators (GEVIs) enabled voltage imaging with single-neuron, single-spike resolution *in vivo* in near-surface neurons (Abdelfattah et al., 2019; Adam et al., 2019; Piatkevich et al., 2019; Villette et al., 2019). Paired expression of near infrared GEVIs based on Archaeorhodopsin 3 (Arch) (Kralj et al., 2012) with blue-excited channelrhodopsin actuators enabled optical measurements of excitability *in vivo* (Optopatch) (Adam et al., 2019; Lou et al., 2016) and of synaptic transmission in primary culture and acute slices (Fan et al., 2018). These advances open the possibility for optical explorations of L1 circuit function in awake animals.

Using a holographic structured illumination imaging system and a recently developed somalocalized GEVI derived from Arch, SomArchon (Piatkevich et al., 2019), we probed the circuit function of L1 interneurons in barrel cortex of awake mice. We developed an all-optical technique to resolve the separate contributions of excitatory and inhibitory synaptic inputs to membrane potential. Experiments with patterned optogenetic activation and

silencing, paired with voltage imaging, revealed that strong lateral inhibition within the L1 microcircuit plays a dominant role in setting the network responses to sensory and neuromodulatory inputs. Mutual inhibition led to high-pass temporal and spatial filtering of excitatory inputs, while under strong cholinergic drive the circuit showed winner-takes-all attractor dynamics.

A simple conductance-based model that incorporates the known electrophysiology and connectivity of L1 interneurons captured the main features of our sensory and neuromodulatory data and predicted responses to several types of inputs which we tested experimentally. The model also predicted that the sensitivity of the L1 microcircuit to thalamic inputs should show an inverted-U dependence on cholinergic drive, a prediction which can be tested in future experiments.

Results

***In vivo* voltage imaging with holographic patterned illumination**

We analyzed the sources of signal and noise in optical voltage recordings with the goal to achieve millivolt-sensitivity recordings of subthreshold dynamics *in vivo*. Voltage signals in tissue arise solely from the neuronal membrane. In epifluorescence images, the soma perimeter appeared brighter than the center, a geometrical projection effect from viewing membranes edgeon. We thus reasoned that incident photons would most efficiently produce signal if targeted to the soma perimeter. Confocal-like excitation combined with spatially filtered emission also minimized optical crosstalk from out-of-focus cells.

We built a holographic structured illumination system, similar to Ref. (Lutz et al., 2008), to achieve this precisely targeted illumination with red ($\lambda = 639$ nm) light for excitation of SomArchon (Fig. 1A, Table S1, Methods and Methods S1). SomArchon fluorescence from all holographically targeted spots was recorded simultaneously on a scientific CMOS camera. All recordings were performed at a 1 kHz frame rate. Spatial filters were applied digitally in postprocessing to separate signal from background (Methods). A digital micromirror device (DMD) patterned blue illumination for targeted optogenetic stimulation (Fig. 1A, Table S1, Methods S1).

We characterized the performance of the imaging system in cortical L1 and superficial L2/3 *in vivo* (Methods). Under wide-field red illumination, SomArchon-expressing neurons were not visible due to high background from scattered and out-of-focus light (Fig. 1B). Illumination targeted to the somas revealed individual cells (Fig. 1B, Methods S1). Holographic membrane focal illumination significantly improved the signal-to-noise ratio of optically detected spikes (spike amplitude:baseline noise in a 1 kHz bandwidth 11.4 ± 1.4 membrane focal vs. 7.5 ± 1.3 whole soma, mean \pm s.e.m., paired measurements in $n = 10$ cells, matched laser power per cell between illumination patterns; Fig. 1C).

A challenge with 1-photon voltage imaging is to ensure that the fluorescence ascribed to each cell is not contaminated by crosstalk from other simultaneously imaged cells or from out-of-focus background. To characterize crosstalk between in-focus cells, we quantified the fluorescence signal (spike amplitude) in concentric rings centered on individual

holographically illuminated cells (Fig. S1). At 6 μm from the cell boundary, signal amplitude decayed to $7 \pm 6\%$ of the on-cell signal (mean \pm s.d., $n = 4$ cells), and was undetectable ($< 1\%$ of on-cell signal) at 20 μm from the boundary.

To characterize out-of-focus crosstalk, we recorded from individual cells as a function of optical defocus. With membrane focal illumination, fluorescence amplitude decayed by 50% over 15 μm deeper defocus, but more slowly with shallower defocus, whereas with whole-soma illumination this fluorescence decay was \sim two-fold slower (Fig. S1).

We used activity-based image segmentation to remove residual out-of-focus crosstalk. A penalized matrix decomposition algorithm separated true voltage signals from crosstalk based on the different spatial profiles of these two signal sources (Buchanan et al., 2018). Application of the algorithm to simulated data with realistic noise (Adam et al., 2019), and to composite movies where optical crosstalk was inserted 'by hand' confirmed that the extracted signals were highly robust to crosstalk (Fig. S1, Methods).

Finally, we used simultaneous patch clamp and fluorescence measurements in acute slices to assess the precision of the optical voltage measurements. Optical and electrical recordings showed close correspondence, both for spikes and subthreshold events (Fig. S1).

The holographic optical system enabled high-resolution recordings in several genetically defined cell types across L1 and superficial L2/3 in awake head-fixed mice (Fig. 1D). 5HT_{3A}R-Cre transgenic mice expressed primarily in L1 interneurons (Lee et al., 2010; Takesian et al., 2018); somatostatin (SST)-Cre transgenic mice expressed in interneurons in deeper layers (Neske et al., 2015); and CKII(0.4)-Cre virus drove dense pan-neuronal expression in wild-type mice. In all three populations, two-photon fluorescence images of an appended eGFP tag showed membrane-localized and somatically restricted expression (Fig. S1).

Action potentials were recorded with SNR 12 ± 4 (mean \pm s.d., $n = 16$ cells) throughout L1 at depths between 20 and 150 μm at a mean laser power of 3 mW/cell. In SST-Cre mice optical recordings were made in superficial L2/3 (100 – 230 μm depth) with SNR 9 ± 4 (mean \pm s.d., $n = 6$ cells). Fig. 1D shows a recording at depth of \sim 230 μm with SNR 7. SNR was approximately two-fold greater for near-surface neurons (< 100 μm depth, 14 ± 4 , mean \pm s.d., $n = 22$ neurons) than at 150 – 200 μm (8 ± 3 , mean \pm s.d., $n = 7$ neurons, Fig. 1E). We imaged multiple neurons simultaneously to a depth of 150 μm (Fig. 1D).

For an SNR of 7 and a spike-detection threshold set at 4σ above the baseline noise, the expected false-positive rate is < 0.3 spikes per 10 s recording, and the expected false-negative rate (missed spikes) is $< 0.14\%$ of true spikes (Methods). We used the refractory period after true spikes to test the false-positive rate. A spike-triggered autocorrelogram showed a probability $< 10^{-3}$ of two spikes occurring within 5 ms of each other ($n = 27$ cells, Fig. 1F), confirming the physiological origin of the detected spikes.

Simultaneous optogenetic perturbations and voltage imaging: Optopatch and i-Optopatch

Next, we sought to combine optogenetic perturbations (activation and silencing) with simultaneous voltage imaging. For optogenetic activation, we paired SomArchon with a blue

light-activated soma-localized channelrhodopsin, SomCheRiff (Adam et al., 2019; Fan et al., 2018; Hochbaum et al., 2014). This channelrhodopsin has previously been shown to have negligible activation at the 639 nm wavelength used to excite Arch-derived GEVIs (Hochbaum et al., 2014). For co-expression *in vivo*, we made a Cre-dependent bicistronic AAV construct which we called Optopatch4.

In awake head-fixed 5-HT_{3A}R-Cre mice expressing Optopatch4, single-cell targeted optogenetic stimulation through a cortical window ($I = 1.8 - 21 \text{ mW/mm}^2$) evoked sustained spiking which was clearly resolved via holographically targeted voltage imaging (Fig. 1G, H). Measurements using laterally offset spots showed little crosstalk of stimulation to surrounding cells, which had a mean spacing of $\sim 60 \mu\text{m}$ (Meyer et al., 2013) (Fig. S2).

Optogenetic silencing can be a powerful tool to determine the roles of specific neural populations in network dynamics, but optogenetic silencing has not previously been paired with voltage imaging. The action spectrum of a soma-localized blue-shifted chloride channel, stGtACR2 (Mahn et al., 2018) suggested it could be spectrally orthogonal to SomArchon. In human embryonic kidney (HEK293) cells expressing stGtACR2, blue light ($I = 0.2 - 1 \text{ mW/mm}^2$) evoked large inhibitory photocurrents ($\sim 0.5 \text{ nA}$), but red light at intensities used for voltage imaging (635 nm, $0.1 - 10 \text{ W/mm}^2$) did not induce observable photocurrents, nor did it impair blue light-mediated channel gating (Fig. S3). Thus SomArchon and stGtACR2 constitute a spectrally orthogonal pair appropriate for inhibitory Optopatch (i-Optopatch).

In head-fixed awake mice expressing i-Optopatch in L1 (Methods), targeted optogenetic stimuli ($I = 1.8 - 21 \text{ mW/mm}^2$) inhibited spontaneous spiking (spontaneous spike rate $3.2 \pm 0.7 \text{ Hz}$ vs. $0.4 \pm 0.3 \text{ Hz}$ with blue light at $I = 1.8 \text{ mW/mm}^2$, $p = 3 \times 10^{-3}$, two-sided paired-sample *t*-test, Fig. 1I, K). Optogenetic stimulation also hyperpolarized resting membrane potential by $-13.5 \pm 4\%$ of spike height (mean \pm s.e.m., $n = 14$ neurons, $I = 1.8 \text{ mW/mm}^2$, 2 mice, Fig. 1J, K). These results establish i-Optopatch as a tool for assessing how targeted optogenetic silencing affects network dynamics.

Whisker stimulation evoked excitation followed by inhibition in L1

We used this suite of tools to dissect the circuit function of L1 during sensory processing (Fig. 2A, B). We first characterized the intrinsic excitability properties of L1 neurons. In acute slices from 5HT_{3A}R-Cre mice expressing Optopatch4, single-cell L1-targeted optogenetic stimuli evoked characteristic firing patterns, including bursting adapting and late-spiking non-adapting phenotypes, as previously reported (Chu et al., 2003), though not all neurons had a clear classification (Fig. S4). Neurons recorded *in vivo* sequentially under isoflurane anesthesia and then wakefulness showed characteristic firing patterns preserved between the two brain states (Fig. S4). The *in vivo* recordings did not clearly resolve into distinct sub-classes based on firing patterns, consistent with prior results (Cadwell et al., 2016; Chu et al., 2003). We therefore treated all 5-HT_{3A}R-positive neurons measured *in vivo* as a single population.

Next, we used voltage imaging to characterize the sensory-evoked responses in L1 interneurons. Barrel fields corresponding to individual whiskers (B2, C2, D2) were

identified by intrinsic imaging (Fig. S5, Methods). In mice anesthetized with isoflurane and in awake mice, brief stimuli to individual whiskers (~1 mm deflection, ~8 mm from the base, 20 ms duration, repeated at 0.5 Hz, Methods) elicited excitatory post-synaptic potentials (EPSPs) and often single spikes in L1 neurons in the corresponding barrel fields (Fig. 2C, D). Whisker stimuli were substantially more effective in eliciting spikes in anesthetized mice than in awake mice (anesthetized: 135 spikes from 153 trials, $n = 24$ cells vs. awake: 73 spikes from 126 trials, $n = 24$ cells), consistent with prior studies in other cortical layers (Haider et al., 2013).

The delay from stimulus onset to spike peak was 16 ± 2 ms in anesthetized mice (mean \pm s.d., $n = 135$ events, 24 neurons, 3 mice, Fig. 2E) and 16 ± 3 ms in awake mice (mean \pm s.d., $n = 73$ events, 21 neurons, 3 mice, Fig. S5). Similar delay and jitter were previously reported in L4 pyramidal and fast-spiking neurons, both of which receive direct thalamic inputs (Gabernet et al., 2005). To assess how much of the jitter reflected variation in input timing vs. variation in the intrinsic responses of individual L1 neurons, we then compared the relative delay in spike time between simultaneously recorded pairs of neurons in anesthetized mice. The mean relative delay was 0.9 ms ($n = 14$ pairs, 66 trials in which both cells spiked, Fig. 2E). Remarkably, despite the small delay, the order of firing was preserved in all pairs in all trials, to within our 1 ms time resolution, i.e. if cell A fired before cell B in a single trial, then cell A fired before cell B in all trials. These observations indicate that abrupt sensory inputs drove millisecond-precision spike sequences in L1.

We observed striking differences in the subthreshold dynamics between spontaneous vs. whisker-evoked spikes, in both anesthetized and awake animals. Spike-triggered average (STA) waveforms of spontaneous spikes rode atop a baseline depolarization that both preceded and followed the spike, whereas whisker-evoked spikes arose abruptly and were followed by a period of hyperpolarization (Fig. 2F, G). Stimulus-triggered average waveforms of whisker deflection trials that did not induce spikes also showed a depolarization followed by a hyperpolarization (Fig. S5). Together, these results implied that whisker stimulus evoked synaptic excitation followed by synaptic inhibition.

Consistent with this hypothesis, we found that in awake animals, whisker stimulation evoked single spikes and then transiently suppressed spontaneous L1 activity (before stimulus firing rate: 3.6 ± 0.85 Hz; 30 to 105 ms post-stimulus: 1.2 ± 0.65 Hz, mean \pm s.e.m., $n = 24$ cells $p = 0.004$, Kruskal-Wallis test, Fig. 2H). Sensory-evoked network suppression was particularly apparent when the sensory stimulus fortuitously arrived during a burst of spontaneous activity (Fig. 2H).

Sensory-evoked network suppression also affected subsequent sensory-evoked responses. When a whisker was subjected to a series of 20 ms deflections at 10 Hz in an anesthetized mouse, the response to the second stimulus was undetectable, while the mean responses to the third and fourth stimuli were ~75% of the response to the first stimulus (Fig. S5). While this observation might be explained by effects upstream of L1, it is also consistent with rapid sensory-evoked inhibition onto L1.

Optical dissection of sensory-evoked excitation and inhibition

We developed an all-optical technique to resolve the distinct contributions of excitatory and inhibitory synaptic inputs to sensory-evoked responses. Rapid inhibition is mediated by GABA_A receptors, ligand-gated chloride channels with a reversal potential of ~ -70 mV. L1 interneurons in anesthetized rats have been reported to rest at -65 to -70 mV (Zhu, 2004), suggesting that inhibitory inputs should have only small effects on membrane potential at rest. Borrowing from well-established patch clamp protocols (Segal and Barker, 1984), we reasoned that optogenetic depolarization would increase the driving force for inward chloride current, and thereby amplify the impact of GABA_A receptor activation on the inhibitory postsynaptic potential (IPSP) (Fig. 3A,B).

In both awake and anesthetized mice, whisker stimuli in the absence of optogenetic stimulation evoked clear spikes or EPSPs in L1 interneurons, as in prior experiments (Fig. 3C, S5). Optogenetic stimuli targeted to single cells (500 ms duration, 1.8 to 21 mW/mm², repeated at 1 Hz) reliably evoked stimulus intensity-dependent spiking. Remarkably, whisker stimuli applied during single-cell optogenetic stimulation hyperpolarized membrane potential and suppressed spiking (Fig. 3C, D, E, S5). We quantified the sensory-evoked subthreshold waveforms by digitally removing spikes (Methods) and calculating a stimulus-triggered average at different optogenetic stimulus strengths (Fig. 3E, S5). In both awake and anesthetized brain states, whisker stimuli had opposite effects in the absence vs. presence of baseline optogenetic depolarization, illustrating dramatic non-additivity of sensory and optogenetic inputs to the same neuron. This observation mimicked the prior observation that sensory stimuli evoked single spikes in quiescent neurons but suppressed ongoing bursts in active neurons (Fig. 2H).

We developed a simple biophysical model to test our interpretation that single cell-targeted optogenetic stimulation amplified the effect of network inhibition. We assumed a transient excitatory synaptic input followed shortly by a transient inhibitory input. With only passive conductances (leak, channelrhodopsin, AMPA receptor and GABA_A receptor), the model captured the main features of the subthreshold dynamics (Fig. 3F, G, S5, Methods), including several subtleties. In the anesthetized state, as the strength of the optogenetic drive increased, the sensory-evoked IPSP amplitude first increased—as explained above—but then decreased (IPSP amplitude $29 \pm 5\%$ of spike height at $I_{\text{blue}} = 5.8$ mW/mm², vs. $16 \pm 4\%$ of spike height at $I_{\text{blue}} = 21$ mW/mm², mean \pm s.e.m., $n = 15$ neurons, 3 mice, $p = 0.001$, two-sided paired-sample t -test, Fig. S5). The model revealed that this decrease was due to shunting of the membrane potential toward the CheRiff reversal potential (~ 0 mV) at high CheRiff conductance (Methods). This simple model thus connected the complex context-dependent whisker-evoked responses in L1 interneurons to basic membrane biophysics.

To constrain the possible sources of network inhibition, we next studied the relative timing of excitatory and inhibitory inputs. We delivered whisker stimuli alternately with and without baseline weak optogenetic stimulation targeted to single neurons (5.8 mW/mm², Fig. 3H). In trials that evoked spikes, stimulus-triggered average waveforms \pm blue light overlapped for the first 2 ms after onset of whisker-evoked depolarization. Thereafter, the waveform in the presence of optogenetic stimulation fell below the waveform in the absence, signaling the onset of inhibition (Fig. 3I, inset). This finding implied a ~ 2 ms delay between

onset of excitation and inhibition, suggesting at most a difference of one synapse in the respective paths (Gabernet et al., 2005). Slower inhibitory signals (e.g. from GABA_B receptors or polysynaptic mechanisms) may also have contributed to inhibition at later times.

Center/surround optogenetic stimulation reveals lateral inhibition in L1

We sought to identify the source of the sensory-evoked inhibition. Patch clamp measurements in acute slices identified inhibitory connections between L1 interneurons (Chu et al., 2003; Cruikshank et al., 2012; Schuman et al., 2018). We hypothesized that whisker stimulation evoked near-synchronous spiking in L1, which then led to network inhibition via lateral connections within the L1 population. To test whether L1 activation was sufficient to evoke lateral network inhibition we performed an all-optical measurement of functional connectivity *in vivo* (Fig. 4A).

We expressed Optopatch4 in 5-HT_{3A}R-Cre mice and targeted voltage imaging to 1-3 L1 interneurons in the center of the field of view. We then defined two optogenetic stimulus patterns. The first pattern comprised small disks targeted individually to the central neurons, with sustained optogenetic depolarization (500 ms, 25 mW/mm²) to increase the driving force for inhibitory currents. The second pattern comprised an annulus (inner diameter ~200 μm, outer diameter of ~400 μm, Fig. 4B, C, Methods), surrounding the central neurons. Midway through the stimulation of the central neurons, a brief flash (20 ms, 25 mW/mm²) was applied to the annulus to evoke synchronized spiking of the surrounding cells.

Optogenetic stimulation of the central neurons evoked robust spiking (41 ± 6 Hz, $n = 25$ neurons, 3 mice, mean \pm s.e.m.). Stimulation of the surrounding neurons transiently suppressed this spiking (spike rate 12 ± 4 Hz in the 25 ms following the annular flash, $p = 4 \times 10^{-4}$, two-sided paired-sample *t*-test, Fig. 4D, E, F). The mean fluorescence waveform following the annular flash showed robust hyperpolarization of the central neurons ($-27 \pm 3\%$ of spike height, Fig. 4G). (Control experiments without the central optogenetic stimulus revealed that the initial depolarization after the annular flash was an artifact from light scatter, Fig. S2). The spike patterns and subthreshold hyperpolarization dynamics in these experiments closely resembled the corresponding data for a sensory stimulus (Fig. 3E, F, I). We observed similar results in neuroderived neurotrophic factor (Ndnf)-Cre mice (Poorthuis et al., 2018; Tasic et al., 2016) which drove Optopatch4 expression selectively in neurogliaform and canopy cells whose axons project laterally within L1 (Schuman et al., 2018) (Fig. S6). Together, these results established that synchronous activation of laterally projecting L1 neurons was sufficient to elicit rapid lateral inhibition within the L1 microcircuit.

Patterned optogenetic inhibition reveals that sensory stimulation evokes lateral inhibition

The existence of a lateral inhibitory connection within L1 did not prove that this connection was responsible for the observed sensory-evoked inhibition. To address this question, we tested how optogenetic inhibition of neurons in an annular region modulated sensory-evoked responses in a centrally located neuron (Fig. 5A, B).

We expressed Cre-on-SomArchon-EGFP and Cre-on-flp-off-stGtACR2-CFP in 5-HT_{3A}R-Cre mice (Fig. 5C, Methods). To prevent scattered blue light from inducing spurious

hyperpolarization of the measured cells, we used low-titer Flpo virus to turn off expression of stGtACR2 in a sparse subset of L1 neurons. We used 2-photon imaging of the appended fluorescent tags (GFP and CFP) to identify neurons that expressed SomArchon and not stGtACR2 (Fig. 5D). These cells were targeted for voltage imaging in awake mice. We then illuminated a surrounding annulus with blue light (inner diameter ~200 μm , outer diameter ~400 μm) to inhibit the surrounding cells while delivering whisker stimuli.

Whisker stimuli in the absence of surround optogenetic inhibition evoked excitation followed by inhibition ($-12 \pm 4\%$ of spike height, mean \pm s.e.m., $n = 13$ neurons, 2 mice, Fig. 5E,F), recapitulating earlier experiments performed with a different reporter construct (Fig. 2G). The same cells were then recorded during whisker stimulation with surround optogenetic inhibition (500 ms, 5.8 mW/mm^2 , Fig. 5E,F right). The post-stimulus hyperpolarization was replaced by a post-stimulus depolarization ($+16 \pm 5\%$ of spike height, Fig. 5F). Some cells that showed prompt single spikes in response to whisker stimulation alone showed delayed and sustained spiking when lateral inhibition was suppressed. Together, these results established that sensory stimulation evoked lateral inhibition between L1 interneurons and that this lateral inhibition contributed to precise timing of whisker stimulus-evoked spikes.

Cholinergic activation induces diverse neuronal responses

Cholinergic modulation of L1 interneurons can enhance cortical L2/3 responses to sensory inputs (Letzkus et al., 2011). As a first step toward understanding the integration of modulatory and sensory inputs in L1, we studied the response of barrel cortex L1 neurons to a stimulus designed to activate cholinergic inputs (Fig. 6A, B). An air puff to the face excites cholinergic neurons in basal forebrain (Hangya et al., 2015), and these neurons are known to innervate cortical L1 (Eggermann et al., 2014; Letzkus et al., 2011; Mechawar et al., 2000).

We imaged L1 neurons in awake mice while delivering a mild air puff (100 ms duration, ~5 psi) to the ipsilateral eye (to avoid incidental whisker stimulation, Fig. 6B). The air puff induced a significant increase in mean spike rate (from 2 ± 1 Hz to 8 ± 4 Hz, 50 to 200 ms after stimulus onset, $p = 0.048$, $n = 15$ neurons, 3 mice, paired two-sample t -test) and evoked a mean depolarization that grew over ~100 ms and lasted for ~1600 ms (Fig. 6C).

The $\alpha 4$ nicotinic acetylcholine receptor is highly expressed in L1 interneurons (Arroyo et al., 2012; Takesian et al., 2018), so we hypothesized that this receptor mediated the neuromodulatory response. We made paired recordings of the same L1 interneurons before and after systemic administration of the $\alpha 4$ nAChR blocker dihydro- β -erythroidine hydrobromide (DH β E, 1.5 mg/kg i.p., Fig. 6C). The AD_{50} for this drug in mice has been reported to range from 0.45 to 1.7 mg/kg (Damaj et al., 1995), implying partial block at the dose used (higher doses are toxic (Damaj et al., 1995)). DH β E did not significantly affect the spontaneous spike rate (before drug: 2 ± 1 Hz, after drug 3 ± 3 Hz, $p = 0.53$, two-sided paired-sample t -test, $n = 15$ neurons, 3 mice, Fig. 6D), but the drug abolished the air puff-induced increase in spike rate (3 ± 3 Hz before air puff to 3 ± 2 Hz after air-puff, $p = 0.79$; Fig. 6D), and significantly decreased the air puff-induced subthreshold depolarization from $22 \pm 4\%$ of spike height to $10 \pm 3\%$ ($p = 0.02$, two-sided paired-sample t -test, Fig. 6E,F).

Thus $\alpha 4$ -mediated cholinergic excitation was largely responsible for the air puff-induced responses, though we cannot rule out other contributions to these responses.

The single-cell traces showed striking variability, so we explored in detail the responses of individual cells. In 15 of 21 L1 interneurons, the air puff evoked a clear depolarization. In 6 of these neurons the air puff evoked one or more spikes and in 3 of these neurons, the air puff evoked a barrage of firing that lasted ~ 1 s, strikingly different from the precisely timed single spikes evoked by whisker stimulation (Fig. 6G). In pairs of simultaneously recorded neurons, we often observed distinct air puff-induced responses in the two cells, indicating that the variability in response was intrinsic to the L1 circuit and was not due to trial-to-trial variation (Fig. S7).

To explore how cholinergic activation affected synaptic inputs, we paired the air puff with single cell-targeted optogenetic depolarization. When the air puff was applied in the middle of an optogenetic stimulus (500 ms, 5.8 mW/mm²), there was no significant change in mean spike rate (Fig. 6H), though some individual neurons showed dramatic increases or decreases in spike rate (see e.g. Fig. 6G, second and fourth traces). During baseline optogenetic stimulation, the air puff on average depolarized the membrane potential (Fig. 6I), opposite to the hyperpolarizing transient evoked by a whisker stimulus under comparable conditions (e.g. Fig. 3C). However, individual neurons sometimes showed strong air puff-induced hyperpolarization (Fig. S7). There are many possible mechanisms underlying the heterogeneous air puff responses. Below we suggest a simple mechanism by which this heterogeneity could arise in a mutually inhibitory network.

Network model of L1 dynamics

Due to the strong mutual inhibition between L1 neurons, one cannot intuitively predict the network responses to excitatory inputs. We developed a simple numerical model of the L1 microcircuit that explained much of our data and predicted network responses to some new inputs (Fig. 7A, Methods S2). The model contained elongated neurogliaform cells (eNGCs) with mutually inhibitory connections, and downward-projecting single-bouquet cells (SBCs) that received inhibitory inputs from eNGC neurons but did not feed back into the simulated network. Both cell types received thalamic and neuromodulatory inputs (Cruikshank et al., 2012).

Conductance-based point-neuron simulations approximately recapitulated the firing properties of the two major cell classes in L1 (Fig. 7B, Methods, Table S2) (Wang and Buzsáki, 1996). We used published anatomical and patch clamp data to set the soma positions, synaptic weights, and thalamocortical input strength (Chu et al., 2003; Jiang et al., 2015; Schuman et al., 2018) (Methods S2, Table S2). Our simulations comprised 51 neurons (34 eNGCs, 17 SBCs), approximately matching the number of interneurons above a single whisker barrel (Meyer et al., 2013).

We simulated the L1 network with thalamic excitation and patterned optogenetic stimulation. Simulated thalamic inputs alone evoked precisely timed single spikes followed by a period of hyperpolarization and suppressed spontaneous activity (Fig. S7), recapitulating the results of Fig. 2. Simulations containing single-cell targeted optogenetic

stimulation amplified the effect of network inhibition in the targeted cell, closely matching the experimental results in Fig. 3 (Fig. 7C). We also reproduced the center/surround excitation experiments from Fig. 4 (Fig. S7).

Simulations predicted that localized excitation would evoke sustained spiking, while wide-area excitation would lead to strong firing rate adaptation due to lateral inhibition (Fig. 7D, E, F). Experiments in anesthetized mice confirmed that optogenetic stimulation of individual cells led to sustained spiking (Fig. 7G, H, I). Optogenetic stimulation of a 400 μm wide spot elicited a prompt spike in nearly all neurons (17 of 20 cells), followed by a complete suppression of spiking in nearly all neurons (18 of 20 cells) during the subsequent 20 ms (Fig. 7G, H). Most neurons remained silent during the subsequent ~ 100 ms, though a few (4 of 20 cells) resumed tonic spiking (Fig. 7G). The sub-threshold potentials also showed a clear difference between localized and distributed optogenetic excitation (Fig. 7I). Localized excitation led to a sustained depolarization, whereas distributed excitation led to a peak in potential followed by a ~ 100 ms inhibition (Fig. 7I).

These results demonstrate the input-output transformations of L1. The circuit acts as a temporal high-pass filter for spatially distributed inputs. This filtering manifests in the precisely timed single spikes from whisker stimulation. The circuit also acts as a spatial high-pass filter for temporally sustained inputs, i.e. spatially localized inputs can evoke sustained local firing, but spatially distributed inputs cannot.

We modeled cholinergic input as activation of an excitatory conductance in all L1 interneurons, with rise time of 300 ms and duration ~ 1 s. To our surprise, sustained spatially distributed excitation of all cells led to winner-takes-all spiking patterns above a critical excitation strength. Some cells spiked a little faster than their neighbors, suppressing spiking of the neighbors and amplifying the disparity in spike rates. For cholinergic inputs above the critical strength, the simulated single-cell firing traces (Fig. 7J) fell into two classes (sustained spiking vs. transient depolarization or single spikes), closely resembling the bimodal single-cell cholinergic responses we observed in response to an air puff (Fig. 7K). A similar bimodal distribution of cholinergic responses had previously been noted in L1 interneurons in auditory cortex in response to a tail shock (Letzkus et al., 2011), but had not been explained. We emphasize that the bimodal responses in the model are an emergent property of a homogeneous mutually inhibitory network; not the result of hard-wired single-cell properties or network connections. These simulation results suggest that strong cholinergic input to L1 can temporarily lock the circuit into a winner-takes-all firing pattern.

Discussion

By combining soma-localized voltage indicators with holographic structured illumination microscopy we achieved high-sensitivity genetically targeted recordings of membrane voltage in cortical L1 and superficial L2/3 in awake mice. We introduced optogenetic techniques for resolving excitatory from inhibitory contributions to subthreshold voltage and for mapping functional connectivity *in vivo*. We developed tools for simultaneous optogenetic inhibition and voltage imaging (i-Optopatch) to reveal the roles of specific neural populations in network function.

We applied these tools to study the input-output properties of L1. Whisker deflections evoked precisely timed single spikes in L1 interneurons, followed by periods of hyperpolarization, reduced spontaneous activity and reduced response to sensory inputs. Optogenetic depolarization of individual cells revealed that whisker stimuli evoked excitation, followed by strong network inhibition. Optogenetic stimulation of neighboring L1 neurons evoked similar patterns of network inhibition. Optogenetic silencing of neighboring L1 neurons abolished sensory-evoked network inhibition. Together these experiments showed that lateral inhibition within L1 plays a critical role in determining the network response to sensory inputs. Our results in L1 are reminiscent of similar results showing sensory-evoked concurrent excitation and inhibition in pyramidal cells in L4 (Gabernet et al., 2005; Wehr and Zador, 2003) and in L2/3 (Petersen et al., 2003; Sachdev et al., 2004).

Our results further showed that neuromodulatory excitation drove tonic spiking in a subset of cells, and drove single spikes or purely subthreshold depolarizations in other cells. While the origin of this heterogeneity is not known, simulations suggested that this heterogeneity could arise in a homogeneous mutually inhibitory network via winner-takes-all dynamics. However, alternate explanations are not ruled out. The heterogeneous cholinergic responses may reflect variations in acetylcholine receptor expression between L1 neuronal subtypes (Schuman et al., 2018), or specific features of the wiring. A definitive test will require population-level voltage imaging in defined subtypes of L1 interneurons, combined with precisely controlled cholinergic activation. This type of experiment is likely to be technically feasible in the near future.

The convergence of sensory and cholinergic inputs onto L1 interneurons raises the question of how these modalities interact. Our model predicts that weak neuromodulatory excitation sensitizes the network to thalamic inputs by bringing all cells closer to threshold; but strong neuromodulatory excitation transiently locks the network into a symmetry-broken winner-takes-all state comprising strongly excited and strongly inhibited neurons. In this state, responses to thalamic inputs are suppressed. Thus the sensitivity of the L1 network to sensory inputs is predicted to show an inverted-U dependence on cholinergic drive.

Considering the importance of L1 in modulating sensory processing, we speculate that the L1 microcircuit may contribute to the inverted-U dependence of performance on arousal observed in many sensory processing tasks (McGinley et al., 2015), often called the Yerkes-Dodson law (Yerkes and Dodson, 1908). We further speculate that rapid firing in a subset of L1 interneurons under strong cholinergic drive may be important for L1-mediated plasticity (Abs et al., 2018; Doron et al., 2019). A clear goal for future work will be to test these predictions by studying the interaction of sensory and modulatory inputs in each subclass of L1 interneurons.

Future applications of the tools described here will benefit from voltage imaging deeper in the brain. Due to light scattering, the present 1-photon imaging approach faces a tradeoff between the density of signal sources and the imaging depth, with a maximum cortical depth for sparse cells of ~230 μ m. Several approaches to circumventing this tradeoff are conceivable, such as adaptive optics (Ji, 2017), graded-index (GRIN) lens endoscopes (Jung

et al., 2004) or prism-based imaging approaches (Andermann et al., 2013). Efforts are also underway to develop two-photon voltage imaging *in vivo* (Villette et al., 2019), though improvements in molecular indicators and optics are likely necessary. Two-photon targeted optogenetic stimulation has recently been paired with calcium imaging *in vivo* (Emiliani et al., 2015; Forli et al., 2018; Mardinly et al., 2018; Marshel et al., 2019; Packer et al., 2015; Rickgauer et al., 2014). Combination of two-photon stimulation with voltage imaging may enable fine-grained circuit dissection *in vivo*.

The all-optical tools introduced here could be used to dissect circuit mechanisms in many contexts. For instance, interactions between excitation and inhibition are critical in network models of reinforcement learning (Schultz et al., 1997) and predictive coding (Rao and Ballard, 1999) but many aspects of these models have not been tested. The ability to silence or activate defined sub-populations while imaging voltage can reveal otherwise hidden inhibition, and elucidate the necessity and sufficiency of specific connections for network function.

STAR Methods

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Adam Cohen (cohen@chemistry.harvard.edu). Plasmids generated in this study and their sequences are available from Addgene.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell lines—HEK293T cell line were bought from ATCC and incubated in DMEM supplemented with 10% FBS and penicillin/streptomycin in a 37 °C incubator under 5% CO₂.

Mice—All procedures involving animals were in accordance with the National Institutes of Health Guide for the care and use of laboratory animals and were approved by the Institutional Animal Care and Use Committee at Harvard University.

For slice experiments, P16-P28 5HT_{3A}R-Cre^{+/-} mice were used.

For *in vivo* experiments, 35-60-day-old wild-type C57BL/6, 5HT_{3A}R-Cre^{+/-}, SST-Cre^{+/-}, and NDNF-Cre^{+/-} mice were used. All mice were housed in standard conditions (reverse 12-hour light/dark cycles, with water and food *ad libitum*). Mice of both sexes were used.

METHOD DETAILS

Design of Optopatch4—Optopatch4 construct (SomArchon-eGFP-P2A-somCheRiff) was cloned into an AAV vector with Cre-dependent expression driven by the hSyn promoter.

LZF1735 pAAV_hSyn-DiO-SomArchon-eGFP-P2A-somCheRiff (Optopatch4, Addgene #126512)

For expression of SomArchon alone we used: pAAV_CAG-FLEX-SomArchon-eGFP (Addgene #126943)

High-titer AAV2/9 virus with Optopatch4 (1.74×10^{13} GC/mL, LZF1735) was obtained from the Janelia Farm Vector Core. High-titer AAV2 virus with SomArchon (6.30×10^{12} GC/mL) was obtained from the UNC Vector Core. High-titer AAV9 virus with CKII(0.4)-Cre (2.8×10^{13} GC/mL) was obtained from UPenn Vector Core.

Design of i-Optopatch—i-Optopatch constructs include:

LZF1826 pAAV_hSyn-DiO-SomArchon-eGFP-P2A-stGTACR2 (Addgene #135412)

LZF1827 pAAV_hSyn-Flp-off-Cre-on-stGtACR2-CFP (Addgene #135413)

pAAV_hSyn1-SIO-stGtACR2-FusionRed was a gift from Ofer Yizhar (Addgene plasmid #105677). stGtACR2 cDNA segments were generated from the template of pAAV_hSyn1-SIO-stGtACR2-FusionRed (Addgene #105677).

High-titer AAV2/9 virus with LZF1826 (2.30×10^{13} GC/mL) and LZF1827 (2.50×10^{13} GC/mL) were obtained from the Janelia Farm Vector Core. pAAV-EF1a-Flpo was a gift from Karl Deisseroth (Addgene plasmid #55637). High-titer AAV5 virus with EF1a-Flpo (2.6×10^{12} GC/mL) was obtained from the UNC Vector Core.

Optical system for holographically targeted voltage imaging and patterned optogenetic stimulation—The optical system combined a red laser ($\lambda = 639$ nm) path for holographic targeted illumination voltage imaging, a blue laser ($\lambda = 488$ nm) path for micromirror-patterned optogenetic stimulation, a two-photon (2P) path for structural imaging, and a wide-field epifluorescence imaging path.

Red laser path.: A red laser (CNI Inc., MRL-FN-639, $\lambda = 639$ nm, 700 mW single transverse mode) was coupled into the setup via a photonic crystal polarization maintaining fiber (NKT Photonics, LMA-PM-15). The fiber output was collimated with an $f = 100$ mm focal length lens (Thorlabs, AC254-100-A-ML) to form a beam with approximately ~ 10 mm diameter. The polarization of the beam was set with a zero-order half-wave plate. The beam was directed onto a holographic reflection-mode liquid crystal spatial light modulator (SLM, Meadowlark 1920SLM VIS) with a resolution of 1920×1152 pixels. Zero-order diffraction was blocked by a home-made anti-pinhole comprised of a dot of solder on a glass slide, placed in a plane conjugate to the sample image plane. The SLM was re-imaged onto the back-focal plane of the objective via a series of relay optics. The objective lens was a $25\times$ water immersion objective, numerical aperture 1.05 (Olympus XLPLN25XWMP2). A mechanical shutter blocked the red laser between data acquisitions. A series of OD filters were placed after the red laser for modulating intensity.

In the first generation of the setup, we used a variable focal length camera lens (Sigma macro 18-200 mm) to control the magnification of the SLM at the back focal plane of the objective. Demagnifying the SLM decreased the effective numerical aperture of the illumination at the sample, leading to bigger spots in the sample, but also to a larger region

that could be targeted with red light. In the second-generation system, we used a fixed lens after the SLM to minimize aberrations. All relay lenses are specified in Table S1.

The SLM device was controlled by custom software. A user specified a set of lines for the SLM to target by drawing on a wide-field epifluorescence image or a 2P fluorescence image. These lines were discretized into a set of spots. The SLM phased pattern was calculated using the Gerchberg-Saxton algorithm.

Red laser intensity was ~ 3 mW per cell for *in vivo* imaging, ~ 1 mW per cell for acute slice imaging.

Blue laser path.: A blue laser (Cobolt, 06-01 series, $\lambda = 488$ nm, 60 mW) was modulated in intensity via an acoustooptic tunable filter (AOTF; Gooch and Housego TF525-250-6-3-GH18A). The beam was focused into a singlemode optical fiber. The output was collimated with an $f = 60$ mm focal length lens (Thorlabs, AC254-060-A-ML) to form a beam with approximately a ~ 17 mm diameter. The beam was then sent to a digital micromirror device with a resolution of 1024×768 pixels (DMD, Vialux, V-7001 VIS). The patterned blue beam was combined with the patterned red beam via a dichroic mirror. The DMD was re-imaged onto the sample at a magnification such that one DMD pixel corresponded to $0.62 \mu\text{m}$ in the sample plane. The DMD optical system enabled patterned blue light stimulation across a field of view of $\sim 450 \times \sim 520 \mu\text{m}$.

The DMD was controlled by custom software. For excitability measurement, a pixel bitmap was preloaded onto and projected from the DMD. For lateral inhibition experiments, pixel bitmaps were loaded into the on-board RAM and digital clock pulses triggered the DMD to sequence through the pre-defined set of exposure patterns.

Wide-field fluorescence imaging path.: The image was relayed from the objective to the camera via a series of three lenses. The final image formation step was performed by a 4x objective (Olympus XLFLUOR 4X/340) serving the role of the tube lens. Fluorescence was collected on a scientific CMOS camera (Hamamatsu ORCA-Flash 4.0). The final magnification of the optical system was 16.7, corresponding to $0.39 \mu\text{m}$ in the sample plane per camera pixel.

Fluorescence from the sample was separated from the blue and red excitation beams via a dichroic mirror (Di03-R405/488/561/635-t3-40x55). An emission filter (Semrock 635 nm long-pass, BLP01-635R-25) further separated SomArchon fluorescence from scattered excitation light. An IR-blocking emission filter (Semrock, FF01-842/SP-25) was placed for blocking scattered infrared excitation light.

All movies are acquired at 1 kHz. To image at 1 kHz, the camera region of interest (ROI) was restricted to typically 200 rows, centered on the image-sensor midline.

The imaging system was designed for a magnification lower than the nominal 25x of the objective for two reasons. First, lower magnification increased the number of neurons that could be imaged simultaneously onto the limited detector area accessible at 1 kHz. Second, by concentrating sample photons onto as few camera pixels as possible, we sought to

minimize the contribution from camera electronic noise, so that all signals would be in the shot noise-limited regime.

Two-photon imaging path.: Light from a femtosecond tunable pulsed infrared laser (Spectra Physics DeepSee) was sent to a pair of galvo mirrors (Cambridge Technologies 6215H). The galvos were re-imaged onto the back focal plane of the objective via an optimized scan lens (Thorlabs, SL50-CLS2) and tube lens (Thorlabs, TL200-CLS2). The visible (blue and red) and near-infrared beams were combined using a 785 nm long-pass dichroic mirror (Semrock, Di03-R785-t3-40x55). GFP fluorescence was directed to the 2P detection path via a removable 550 nm long-pass dichroic. Scattered excitation light was blocked by a 550 nm long-pass emission filter, an IR-blocking emission filter (BSP01-785R-25) and a band-pass emission filter (FF03-525/50-25). A pair of lenses (focal lengths 75 mm and 16 mm) re-imaged the back-aperture of the objective onto a photomultiplier tube (Hamamatsu, H11706P-40). The output of the photomultiplier was amplified and low pass filtered through an amplifier unit (Hamamatsu C7319) and then digitized.

For i-Optopatch experiments, GFP fluorescence was excited by 950 nm and selected by a band-pass emission filter (Chroma, ET525/36m). CFP fluorescence was excited by 900 nm and selected by a band-pass emission filter (FF01-475/28-25).

Control software.: The entire setup was controlled by custom software written in LabView. Interfacing was via a National Instruments DAQ (NI PCIe-6363).

The software contained routines for registration of the DMD, SLM, 2P microscope coordinates to the camera via affine transformations. The camera served as the global reference coordinate system.

Experimental protocols were specified by a set of images (to the SLM and the DMD), output waveforms (to the galvos, the AOTF, the shutters, the update clock on the DMD, the piezo whisker stimulator, and the air puff controller), and analog input streams (from the PMT, the camera exposure clock, and a patch clamp electrophysiology setup not used in the present work).

The Hamamatsu camera uses an internal 100 kHz clock to synchronize image row readout. We found that when the camera exposure times were triggered by the DAQ in synchronous mode, the camera rounded the exposure time to the nearest 10 μ s, leading to 1% jitter in exposure time for 1 kHz imaging. To address this noise source, we used a custom firmware upgrade to access the 100 kHz camera clock. This clock became the master clock for the DAQ system, and all analog and digital input/output functions were synchronized to the camera clock.

Imaging in acute slices—All procedures involving animals were in accordance with the National Institutes of Health Guide for the care and use of laboratory animals and were approved by the Institutional Animal Care and Use Committee at Harvard University.

Virus injection for acute slice measurements. Virus comprising AAV2/9 hSyn-Dio-SomArchon-eGFP-P2A-somCheRiff (1.74×10^{13} GC/mL) was diluted in PBS and injected at a final titer of $\sim 2 \times 10^{12}$ GC/mL.

5HT_{3A}R-Cre^{+/-} mice were crossed with wild-type C57BL/6 mice. Pups were cryo-anesthetized at P0-P2 and immobilized dorsal side up under a stereotaxic microscope. Injections were made using home-pulled micropipettes (Sutter P1000 pipette puller), mounted in a microinjection pump (World Precision Instruments Nanoliter 2010) controlled by a microsyringe pump controller (World Precision Instruments Micro4). The micropipette was positioned using a stereotaxic instrument (Stoelting Digital Mouse Stereotaxic Instrument). Pups were injected in the left hemisphere, 1 mm lateral and 1.2 mm anterior to lambda. Starting at a depth of 0.3 mm beneath the surface of the skull, virus injections (40 nL, 1 nL/s) were performed at 0.1 mm increments as the pipette was withdrawn. Pups were placed back in their home cage once they were awake.

Genotyping. Genotyping for 5HT_{3A}R-Cre, SST-Cre and NDNF-Cre was performed with the PCR primer pairs: Cre 5': 5' TAT CTC ACG TAC TGA CGG TG 3' and Cre 3': 5' AGA CTA ATC GCC ATC TTC CAG C 3' to yield a 500 bp band from Cre.

Acute slice preparation. Acute brain slices were prepared from P16–P28 5HT_{3A}R-Cre^{+/-} mice. The mice were anesthetized by isoflurane and then perfused with carbogen (95% O₂, 5% CO₂)-saturated ice-cold slicing solution with the following composition (in mM): 110 choline chloride, 2.5 KCl, 1.25 NaH₂PO₄, 25 NaHCO₃, 25 glucose, 0.5 CaCl₂, 7 MgCl₂, 11.6 Na-ascorbate, and 3.1 Na-pyruvate. Mice were then decapitated and the brains were rapidly coronally sliced with 300 μm thickness on a vibratome (Leica VT 1200S).

Slices were incubated for 45 min at 34 °C in a carbogenated artificial CSF (ACSF) with the following composition (in mM): 127 NaCl, 2.5 KCl, 1.25 NaH₂PO₄, 25 NaHCO₃, 25 glucose, 2 CaCl₂, and 1 MgCl₂. The osmolarity of all solutions was adjusted to 300–310 mOsm and the pH was maintained at 7.3 under constant bubbling with carbogen.

Imaging acute slices. Measurements were conducted in ACSF at 23 °C under ambient atmosphere. The slice was immobilized in a slice recording chamber using a slice anchor (Warner Instruments, SHD-40/2). ACSF, perfused with carbogen, was flowed through the chamber at a rate of 2 mL/minute.

Cranial windows and virus injections

Cranial window surgery and virus injection for imaging barrel cortex L1. For Optopatch experiments, virus comprising AAV2/9 hSyn-Dio-SomArchon-eGFP-P2A-somCheRiff (1.74×10^{13} GC/mL) was diluted in PBS and injected at a final titer of $\sim 3 \times 10^{12}$ GC/mL.

For i-Optopatch experiments (Fig. 1I-K and Fig. 5), virus comprised AAV2/9 hSyn-Flp-off-Cre-on-stGtACR2-CFP (final concentration $\sim 1 \times 10^{12}$ GC/mL) mixed with AAV2 CAG-FLEX-SomArchon-eGFP (final concentration $\sim 6.3 \times 10^{11}$ GC/mL) and AAV5 EF1a-Flpo (final concentration $\sim 2 \times 10^{11}$ GC/mL).

The procedure for surgery and imaging in barrel cortex L1 followed the protocol from Andermann (Goldey et al., 2014). 35-60-day-old heterozygous 5HT_{3A}R-Cre mice (male and female) were deeply anesthetized with 2% isoflurane and maintained with ~1% isoflurane throughout the surgery. Eyes were kept moist using ophthalmic eye ointment. Body temperature was continuously monitored and maintained at 37 °C using a heating pad (WPI, ATC2000). The skull was exposed and thoroughly dried and a 3 mm round craniotomy (3.3 – 3.4 mm lateral, 1.6 mm caudal of bregma) was opened using a biopsy punch (Miltex). Virus was then injected in 4 - 8 locations in the center of the craniotomy. Starting at a depth of 0.2 mm beneath the surface of the dura, virus injections (60 nL, 30 - 60 nL/min) were performed at 0.1 mm increments as the pipette was withdrawn. Brain surface was kept moist with saline throughout the injection.

A window was prepared prior to the surgery and comprised two 3 mm round #1 cover glasses and one 5 mm round #1 cover glass (Harvard apparatus) cured together with UV curable adhesive (Norland Products, NOA 81). Following the virus injection, the window was then placed covering the barrel cortex and cemented to the skull with dental cement (C&B metabond, Parkell, No. 242-3200). After the window cured, a titanium headplate (similar to the design in Ref. (Goldey et al., 2014)) was glued around the window and any exposed skull was covered with dental cement. Animals were returned to their home cage for recovery and treated for 3 days with Carprofen (5 mg/kg) and Buprenorphine (0.1 mg/kg) twice a day. To avoid damage to the implant, mice were housed in separate cages.

Cranial window surgery and virus injection for L1-3 imaging (Fig. 1D): The procedure for surgery and imaging in visual cortical L1-3 followed the same protocol as above, except that 35-60-day-old wild-type C57BL/6 mice or heterozygous SST-Cre mice (male and female) were used. The coordinates for the 3 mm round craniotomy were 2.4 mm lateral and 2.7 mm caudal of bregma. Virus was then injected in 4 - 8 locations in the center of the craniotomy. Starting at a depth of 0.3 mm beneath the surface of the dura, virus injections (60 nL, 30 - 60 nL/min) were performed at 0.2 mm increments as the pipette was withdrawn.

For wild-type mice, injected virus comprised AAV2 CAG-FLEX-SomArchon-eGFP (final titer $\sim 0.5 \times 10^{12}$ GC/mL) mixed with CKII(0.4)-Cre virus (UPenn vector core, final titer $\sim 1 \times 10^9$ GC/mL).

For SST-Cre mice, injected virus comprised AAV2 CAG-FLEX-SomArchon-eGFP (final titer $\sim 0.5 \times 10^{12}$ GC/mL).

Tracking, whisker stimulation, intrinsic imaging and *in vivo* voltage imaging—

For general behavior monitoring during acquisitions we recorded video of the mouse's face. An IR LED light (850 nm) was placed in front of the animal. A PointGrey camera (GS3-U3-51S5M-C, Mono Grasshopper3 USB 3.0 Camera) with a Fuji lens (Fuji Photo Optical 1:1.4/25 Fujinon-TV Camera Lens) and an IR-passing optical filter (Thorlabs, FB850-40) was placed on the side to track the animal's face (whisker motion and eye blinks) during data acquisitions.

Whisker stimulation.: An individual whisker was selected for stimulation (typically B2, C2 or D2). Other whiskers were trimmed to prevent direct contact with the glass pipette. The whisker was inserted into a glass pipette glued to a piezoelectric actuator. The actuator was connected to an amplifier (Krohn-Hite 7602M) and controlled by the DAQ. A spike was classified as ‘evoked’ if it occurred within 30 ms of onset of whisker stimulus.

Intrinsic imaging.: Intrinsic imaging was performed on the same setup as described above. A 4x objective (Olympus XLFLUOR 4X/340) was used to image the entire 3 mm cranial window. The whisker stimulation was 10 Hz for 4 s with a 16 s interstimulus interval. A red LED (625 nm) illuminated the window surface from the side. Reflected light was imaged onto the camera at 10 Hz frame rate. A decrease in reflectance from the brain indicated the barrel, which could be localized relative to the blood vessel pattern as visualized with 488 nm illumination.

Imaging anesthetized animals.: Imaging started 3 weeks post-surgery. Mice were lightly anesthetized (0.7–1% isoflurane), head-fixed under the upright microscope using the titanium head plate and held in a body tube. Eyes were kept moist using ophthalmic eye ointment. Body temperature was continuously monitored and maintained at 37 °C using a heating pad (WPI, ATC2000). A typical imaging session lasted 1–2 hours, and then animals quickly recovered and returned to their home cage. Recordings targeting L1 neurons *in vivo* were performed at a depth < 150 μm in both anesthetized and awake animals. Recordings typically targeted 1 – 4 neurons simultaneously.

Habituation and imaging awake animals.: Habituation started 2 weeks post-surgery. Each animal was acclimated to the head restraint in a body tube for at least 3 days before starting the imaging sessions. For imaging in awake animals, a 3D-printed paw blocker was placed in front of the forepaws to prevent them from pushing away the glass pipette for whisker stimulation.

Air puff.: The timing of the air puff was controlled by a solenoid valve (WPI). The strength of the air puff was set ~ 5 psi and controlled by a pressure regulator (Festo, pressure regulator LRP-1/4-4). The air puff was delivered through a blunt needle at ~ 5 mm from the eye ipsilateral to the brain hemisphere used for voltage imaging (to avoid spurious whisker stimulation arriving in the imaged barrels). Air pressure and needle position were adjusted to achieve an air puff strength just strong enough to evoke an eye blink response and increase the pupil diameter.

DHβE administration.: To modulate cholinergic signaling, dihydro-β-erythroidine hydrobromide (DHβE, Tocris; 2349) was diluted in saline. The drug was administered systemically (1.5 mg/kg i.p.). Optopatch and air puff measurements were performed before, and then 30 min. after drug administration, on the same sets of cells. The experiment was performed on each animal twice, on successive days.

Center/surround optogenetic stimulation—For lateral inhibition experiments, we defined two optogenetic stimulus patterns. “Central masks” covered individually the cell bodies of 1 – 3 neurons at the center of the field of view. We calculated the geometrical

centers of these masks individually. The mean of these centers was set as the coordinates for the center for the surrounding “annulus mask”.

The annulus inner radius was set to be ~100 μm from the most non-centered central mask. This distance was selected to minimize the impact of scattered light from the annulus mask. The outer radius of the annulus mask was set as the largest value at which the annulus would be contained within the FOV. Typical outer radii were ~200 μm .

The image sequence was composed of three composite masks: (1) central masks only; (2) central mask and annulus mask together; (3) annulus mask only. These masks were preloaded into the on-board RAM and digital clock pulses triggered the DMD to sequence through the pre-defined set of exposure patterns.

QUANTIFICATION AND STATISTICAL ANALYSIS

Data were analyzed with homemade code written in MATLAB.

Corrections for photobleaching and motion artifacts.—Movies were first corrected for motion using the NoRMCorre algorithm (Pnevmatikakis and Giovannucci, 2017). Movies were then corrected for photobleaching by dividing the movie by an exponential fit of the mean fluorescence.

Image segmentation and waveform extraction.—We divided the movie into sub-movies comprising single cells and performed activity-based image segmentation separately in each sub-movie. Whereas correlations often arose between subthreshold voltages and out-of-focus background, we assumed that spiking was not correlated with background, and furthermore that the spatial footprint associated with spiking would be the same as for true subthreshold dynamics. For segmentation purposes, we first removed subthreshold signals via a 100 Hz high-pass filter. Movies were then segmented semi-automatically using one of two activity-based segmentation algorithms. In one approach, principal components analysis was followed by time-domain independent components analysis (PCA/ICA) (Mukamel et al., 2009). The spatial masks from PCA/ICA were then applied to the original movies without high-pass filtering to extract fluorescence traces.

In a second approach, high-pass filtered movies were segmented via penalized matrix decomposition nonnegative matrix factorization algorithm (PMD-NMF) (Buchanan et al., 2018). The spatial masks were applied to the original movies without high-pass filtering to extract initial fluorescence traces. An estimate of the background signal was calculated from pixels not on the targeted cells. Signals from the cellular footprints were calculated with the constraint that the background spatial profile should be maximally smooth. Both approaches to image segmentation gave similar results, a consequence of the sparsity of L1 interneurons and the optical sectioning from tightly focused holographic excitation.

Validation of the PMD-NMF signal extraction algorithm.—The algorithm has been extensively tested on simulated data comprising realistic shot-noise limited signals from individual cells, correlations in sub-threshold voltages between cells, and correlated out-of-focus background (See Extended Data Fig. 6 of Ref (Adam et al., 2019)). To test the

algorithm with data from cortical L1 interneurons, we took independent single-cell recordings and extracted the fluorescence dynamics from each. We made composite movies by adding two movies together, with ~50% overlap between cell bodies. We extracted the signals from the composite movies, and compared to the ground truth data extracted from the individual movies. This test is more stringent than any real-world data because in-focus cells never physically overlapped in real data.

Removing scattering background for lateral inhibition measurements.—Flashes of blue light delivered in an annular pattern induced tissue autofluorescence which bled into the voltage recordings from centrally located neurons. To correct for this crosstalk, background fluorescence from a region surrounding the central imaged cells was subtracted from the fluorescence of the recorded cells.

Spike finding and scaling of fluorescence recordings.—A simple threshold-and-maximum procedure was applied for spike detection. Fluorescence traces were first high-pass filtered, and initial threshold was set at 4 times the noise level. This threshold was then manually adjusted.

All fluorescence signals were normalized to spike height for spike-triggered average or stimulation-triggered average.

Spike detection: false-positive and false-negative rates.—For an SNR of 7 and a spike-detection threshold set at 4σ about the baseline noise, the false-positive rate is calculated as the probability that samples from a Gaussian distribution lie more than 4σ above the mean. This probability is $(1-p_4)/2 * 1 \text{ kHz} * 10 \text{ s}$, where $p_4 = 0.999937$. The false-negative rate is calculated as the probability that spike height values fall more than 3σ below the mean. This probability is $(1-p_3)/2$, where $p_3 = 0.9973$.

Spike removal for calculation of subthreshold waveforms.—Spikes were digitally removed and replaced with linear interpolations of the surrounding data. Spike width was estimated by viewing individual fluorescence recordings. Linear interpolations were performed between data-points 1 ms beyond the edges of the spike.

Statistics: All error ranges represent standard error of the mean, unless otherwise specified. For the same neurons before and after drug administration, in anesthetized and awake states, with and without surrounding inhibition, and under local and global optogenetic stimulation, paired sample t-test was used. For two-sample comparisons of a single variable, student's t-test was used. In cases where the underlying distributions were non-Gaussian, the Kruskal-Wallis test was used. Probabilities of the null hypothesis $p < 0.05$ were judged to be statistically significant.

Biophysical model of subthreshold membrane potential: The evolution of subthreshold membrane potential in the presence of synaptic inputs and optogenetic stimulation (Fig. 3) was simulated with a passive single compartment model using the following equation:

$$C_m \frac{dV}{dt} = g_e(E_e - V) + g_i(E_i - V) + g_l(E_l - V) + g_{ChR}(E_{ChR} - V), \quad (1)$$

where C_m is the membrane capacitance, g_e , g_i , g_l and g_{ChR} are the conductance of excitatory, inhibitory, leak and channelrhodopsin channels, and E_e , E_i , E_l and E_{ChR} are the respective reversal potentials. The time course of conductance upon excitatory or inhibitory synaptic input was simulated using an alpha function:

$$g(t) = \begin{cases} g_{syn} \frac{t - t_0}{\tau} e^{1 - \frac{t - t_0}{\tau}} + g_{baseline} & \text{for } t \geq t_0 \\ g_{baseline} & \text{for } t < t_0 \end{cases}. \quad (2)$$

Here g_{syn} is the strength of the synaptic input, t_0 is the time of the synaptic input, τ is a time-constant of synaptic input, and $g_{baseline}$ reflects the tonic level of synaptic input excluding the event of interest.

In principle, the values of $g_{baseline}$ for the inhibitory and excitatory synaptic inputs could be wrapped into the definitions of g_l and E_l . Doing so would not affect the solutions to Eq. 1. We chose to keep the baseline synaptic conductances as separate parameters to facilitate explorations of the model under different brain states (e.g. anesthesia vs. wakefulness). In this approach, g_l and E_l reflect cell-autonomous leak conductances (e.g. K_{ir} channels), assumed to be independent of brain state, while $g_{baseline}$ captures the effect of network-dependent inputs.

To simulate lateral inhibition in L1, we assumed inhibition lagged excitation by 2 ms. Other parameters are listed below:

Parameter	Value
$g_{e\ syn}$	1.5 nS
$g_{e\ baseline}$	0.1 nS
$g_{i\ syn}$	5 nS
$g_{i\ baseline}$	0.1 nS
g_l	3.33 nS
g_{ChR}	0 – 10 nS
E_e	–5 mV
E_i	–70 mV
E_l	–70 mV
E_{ChR}	0 mV
C_m	150 pF
τ	1 ms

Equation 1 was numerically integrated using Euler's method.

The model above can be solved analytically for the steady-state voltage by setting $\frac{dV}{dt} = 0$, which yields:

$$V = \frac{E_e g_e + E_i g_i + E_l g_l + E_{ChR} g_{ChR}}{g_e + g_i + g_l + g_{ChR}}. \quad (3)$$

The PSP amplitudes are obtained by calculating the difference in steady-state voltage,

V_{PSP} , after vs. before the sensory perturbations to g_e and g_i assuming that all other parameters do not vary during the synaptic event. If one assumes that g_e and g_i are both zero before the synaptic event (i.e. by absorbing the pre-stimulus values of g_e and g_i into the definition of g_j and E_j), then one finds:

$$\Delta V_{IPSP} \approx \frac{g_{ChR}[g_e(E_e - E_{ChR}) + g_i(E_i - E_{ChR})] + g_l g_e(E_e - E_l) + g_l g_i(E_i - E_l)}{(g_l + g_{ChR})(g_i + g_l + g_{ChR})}. \quad (4)$$

Since only differences in voltage appear in Eq. 4, one can arbitrarily choose to set one of the voltages to zero, and measure all other voltages relative to this reference. For convenience, we set E_l to zero for the fitting. Eq. 4 is linear in g_{ChR} in the numerator, and quadratic in g_{ChR} in the denominator, suggesting 5 fitting parameters to specify the function

$V_{IPSP}(g_{ChR})$. However, the proportionality between V and F is not *a priori* known. Also the proportionality between g_{ChR} and I_{488} is not *a priori* known.

To estimate the waveforms of the post-synaptic potentials, we first normalized all fluorescence traces by spike height. We then calculated the mean fluorescence over 10 ms before the whisker stimulus, and the mean fluorescence over 10 ms starting 30 ms after the whisker stimulus. The difference between these values was taken as the amplitude of the post-synaptic potential. Equation 4 was fitted to the data using the nonlinear least-squares method in Matlab.

Analytical approximation.—To gain an intuition for the responses, one can make a simple estimate of the amplitude of the IPSP by assuming that $g_e = 0$ at all times, that $g_i = 0$ before the synaptic input, and that the resting potential is the same as the inhibitory reversal potential, i.e. $E_l \approx E_i$. One then obtains:

$$\Delta V_{IPSP} \approx \frac{g_i g_{ChR}(E_l - E_{ChR})}{(g_l + g_{ChR})(g_i + g_l + g_{ChR})}. \quad (5)$$

Within the blue light intensity range used in our experiments, the CheRiff conductance is well approximated by a linear function of the blue intensity, though the proportionality factor depends on the (unknown) CheRiff expression level and attenuation of the blue light by scattering.

Eq. 5 shows that for small g_{ChR} the amplitude V_{IPSP} is linear in g_{ChR} , while for large g_{ChR} the quadratic term in the denominator dominates and V_{IPSP} decreases inversely with g_{ChR} .

The value of g_{ChR} that gives the largest amplitude IPSP is $g_{ChR}^{max} = \sqrt{g_l(g_l + g_i)}$, or for weak inhibition, $g_{ChR}^{max} \approx g_l$. If g_l is large, then the shunting from g_{ChR} is suppressed. The membrane time constant is approximately $\tau \approx \frac{C}{g_l + g_{ChR}}$. The inverse relation between τ and g_{ChR} is consistent with our observation of faster recovery at stronger stimulus strength.

Numerical model of L1 dynamics

Single-cell properties.: We considered two broad classes of L1 interneurons: laterally projecting neurogliaform cells (eNGC) and downward projecting single bouquet-like cells (SBC). To capture the firing properties of SBC, we used a modified form of the conductance-based model for hippocampal interneurons in Ref (Wang and Buzsáki, 1996) (Table S2). The voltage gated conductances responsible for the late-spiking pattern of eNGC have not been identified (Overstreet-Wadiche and McBain, 2015). We included an A-type inactivating potassium conductance to capture the firing pattern (Bard Ermentrout and Terman, 2010). We then connected the two neuronal subtypes with distance dependent inhibitory synapses. ENGc cells synapsed onto each other and onto SBC cells. SBC cells did not synapse within L1. Simulations comprised 51 L1 interneurons (34 eNGC, 17 SBC) corresponding approximately to a single barrel (Table S2).

We judged more detailed channel-based biophysical models to have too many unknown parameters. Simpler spike rate-based models did not capture the details of the spike timing or subthreshold potentials, which we judged important for L1 circuit function.

Each SBC neuron was described by a single compartment and simulated with the following equation:

$$-C_m \frac{dV}{dt} = -I(t) + g_k n^4 (V - E_k) + g_{Na} m_{inf}^3 h (V - E_{Na}) + g_{leak} (V - E_{leak}) + g_{ChR} (V - E_{ChR}) + g_e (V - E_e) + g_i (V - E_i) + \epsilon(t) \quad (6)$$

$$\frac{dn}{dt} = \varphi(\alpha_n(V))(1 - n) + \beta_n(V)n; \quad (7)$$

$$\frac{dm}{dt} = \alpha_m(V)(1 - m) + \beta_m(V)m; \quad (8)$$

$$\frac{dh}{dt} = \varphi(\alpha_h(V))(1 - h) + \beta_h(V)h \quad (9)$$

$$\alpha_n = \frac{-0.01(V + 24)}{e^{-0.1(V + 24)} - 1}; \beta_n = 0.125 * e^{-\frac{V + 34}{80}};$$

$$\alpha_m = \frac{-0.1(V+25)}{e^{-0.1(V+25)} - 1}; \beta_m = 4 * e^{-\frac{V+50}{18}};$$

$$\alpha_h = 0.07 * e^{-\frac{V+48}{20}}; \beta_h = \frac{1}{e^{-0.1(V+18)} + 1};$$

where $C_m = 1 \mu\text{F}/\text{cm}^2$; and $I(t)$ is the injected current; the conductance of leak current, $g_{leak} = 0.23 \text{ mS}/\text{cm}^2$ so that the passive time constant $\tau = 4.3 \text{ ms}$; $E_{leak} = -66.8 \text{ mV}$; $g_{Na} = 35 \text{ mS}/\text{cm}^2$; $E_{Na} = 55 \text{ mV}$; $\varphi = 5$; $g_k = 9 \text{ mS}/\text{cm}^2$; $E_k = -90 \text{ mV}$. The quantity $\epsilon(t) = N(0, \sigma^2)$ represents a Gaussian distributed white noise current with $\sigma = A (\delta t)^{1/2}$, where $A \sim 1 \text{ pA cm}^{-2} \text{ ms}^{-1/2}$ and $\delta t = 0.02 \text{ ms}$ (or in some cases 0.05 ms) is the integration time-step. The equations were integrated using the Euler method.

The dynamics of eNGC neurons followed similar equations with the following difference: we included an A-type inactivating potassium conductance to capture the firing pattern. This channel contributed a current:

$$I_A = g_A(0.6 * ha_1 * ma_1^4 + 0.4 * ha_2 * ma_2^4)(V - E_A)$$

$$\frac{dma_1}{dt} = \frac{ma_{1,inf} - ma_1}{t_{ma}}; \frac{dma_2}{dt} = \frac{ma_{2,inf} - ma_2}{t_{ma}};$$

$$\frac{dha_1}{dt} = \frac{ha_{1,inf} - ha_1}{t_{ha,1}}; \frac{dha_2}{dt} = \frac{ha_{2,inf} - ha_2}{t_{ha,2}};$$

$$t_{ma} = \frac{1}{e^{(V+35.82)/19.69} + e^{-(V+79.69)/12.7}} + 0.37;$$

$$ma_{1,inf} = \frac{1}{1 + e^{-\frac{(V+60)}{8.5}}}; ma_{2,inf} = \frac{1}{1 + e^{-\frac{(V+36)}{20}}}; ha_{inf} = \frac{1}{1 + e^{(V+78)/6}};$$

$$t_{ha,1} = \begin{cases} \frac{1}{e^{(V+46.05)/5} + e^{-(V+238.4)/37.45}} & \text{for } V < -63 \\ 19 & \text{for } V \geq -63 \end{cases}$$

$$t_{ha,2} = \begin{cases} \frac{1}{e^{(V+46.05)/5} + e^{-(V+238.4)/37.45}} & \text{for } V < -73 \\ 60 & \text{for } V \geq -73 \end{cases}$$

where $g_A = 10 \text{ mS/cm}^2$; $E_A = -75 \text{ mV}$.

To introduce firing rate adaptation in SBC cells, we included a slow outward potassium current. This channel contributed a current:

$$I_{sk2} = g_{sk2} * m_k * h_k * (V - E_{sk2})$$

$$\frac{dm_k}{dt} = \frac{mk_{inf} - m_k}{t_{mk}}; \frac{dh_k}{dt} = \frac{hk_{inf} - h_k}{t_{hk}};$$

$$t_{mk} = \frac{1}{e^{(V-80.98)/25.64} + e^{-(V+132)/17.953}} + 9.9;$$

$$t_{hk} = \frac{1}{e^{(V-1329)/200} + e^{-(V+129.7)/7.143}} + 120;$$

$$mk_{inf} = \frac{1}{\left(1 + e^{-\frac{(V+43)}{17}}\right)^4}; \quad hk_{inf} = \frac{1}{1 + e^{-\frac{(V+58)}{10.6}}};$$

where $g_{sk2} = 10 \text{ mS/cm}^2$; $E_{sk2} = -70 \text{ mV}$.

For the figure in Fig. S7, in order to produce high spontaneous firing, we set the resting potential higher at -55 mV for both eNGCs and SBCs.

Channelrhodopsin activation.: The channelrhodopsin CheRiff was modeled as an excitatory conductance with reversal potential 0 mV and conductance proportional to blue light illumination intensity. Channel gating kinetics were assumed to be instantaneous. Patterns of blue light were targeted to one or more cells and modulated in time during the simulation.

Synaptic properties: Inhibition.: Examination of patch clamp recordings of inhibitory post-synaptic potentials (IPSPs) in acute slices showed a rapid onset followed by a slow recovery. The recovery was not well captured by a single exponential. To approximate these dynamics we used the following function:

$$g_{inh}(t) = g_{inh}^0 \left(\frac{t}{\tau_1} e^{1 - \frac{t}{\tau_1}} + 0.6 \frac{t}{\tau_2} e^{1 - \frac{t}{\tau_2}} \right)$$

$\tau_1 = 5$ ms, $\tau_2 = 30$ ms. We found that the qualitative network dynamics were insensitive to variations in the functional form or time constants. One may think of the two terms as representing GABA_A and GABA_B receptors respectively, though a more accurate implementation of a GABA_B-mediated hyperpolarization would use the K⁺ reversal potential (−90 mV) rather than the Cl[−] reversal potential (−70 mV).

To calibrate the value of g_{inh}^0 for eNGC → eNGC and eNGC → SBC synapses, we set up a model circuit with one eNGC cell synapsing onto an eNGC cell and an SBC cell. We adjusted a simulated channelrhodopsin conductance ($E_{ChR} = 0$ mV) to depolarize the downstream cells to −55 mV. We then triggered the upstream cell to spike. We adjusted g_{inh}^0 to induce IPSP amplitudes of −1.5 to −2 mV, to match the patch clamp data. The model did not contain short term inhibitory synaptic plasticity.

The IPSP amplitudes recorded via patch clamp were assumed to represent the strongest possible IPSPs. The IPSP strength between each pair of neurons in the circuit was modulated by a Gaussian function of separation, with a length-scale set by the sum of sizes of the presynaptic axonal arbor and the postsynaptic dendritic arbor (Jiang et al., 2015).

Synaptic properties: Excitation.: We modeled thalamocortical excitation in L1 using the function:

$$g_{exc}(t) = g_{exc}^0 \frac{t}{\tau} e^{1 - \frac{t}{\tau}}$$

We calibrated the amplitude of g_{exc}^0 and τ by simulating thalamic inputs to eNGC cells and SBC-like cells and matching to literature data which showed that whisker-evoked EPSP amplitudes *in vivo* were 3 – 7 mV. We assumed that the timecourse of thalamic excitation to all neurons was identical. Excitatory synaptic strengths were randomized by 20-30% between cells to prevent numerical degeneracies. The model did not contain short-term excitatory synaptic plasticity.

Synaptic properties: Neuromodulation.: The greatest uncertainty in the model surrounded the time-course and strength of neuromodulatory inputs. We assumed that neuromodulatory inputs activated an excitatory conductance with reversal potential 0 mV. We assumed that the strength and timecourse of neuromodulatory action was the same on the eNGC and SBC-like neurons. If the coupling of neuromodulatory input to the SBC-like neurons was sufficiently strong, then these neurons could be driven to spike by neuromodulatory inputs, even in the absence of thalamic inputs.

Omissions from the model.: Many possibly relevant features were omitted from the model. These include: gap junction connections between eNGC neurons (Chu et al., 2003),

activation of GABA_B receptors or of muscarinic acetylcholine receptors (Brombas et al., 2014), and possible feedback inhibition from deeper layer Martinotti cells (Abs et al., 2018). We did not consider a finer classification of L1 interneurons into sub-types. Our model also did not include cortico-cortical inputs. We did not study or simulate the effects of L1 interneuron activation on apical dendrites of deeper layer pyramidal cells. Activation of 5HT_{3A} ionotropic serotonin receptors is expected to have similar electrophysiological effects to activation of nicotinic acetylcholine receptors, though the distribution of these two receptor-types in the different L1 interneuron sub-classes may be different.

DATA AND CODE AVAILABILITY

Data, code and custom software are available upon request.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- All-optical electrophysiology reveals synaptic excitation and inhibition, *in vivo*.
- Whisker stimuli evoke concurrent excitation and inhibition in L1 interneurons.
- Cholinergic inputs evoke winner-takes-all spiking in L1 interneurons.
- Lateral inhibition within L1 governs whisker and cholinergic responses.

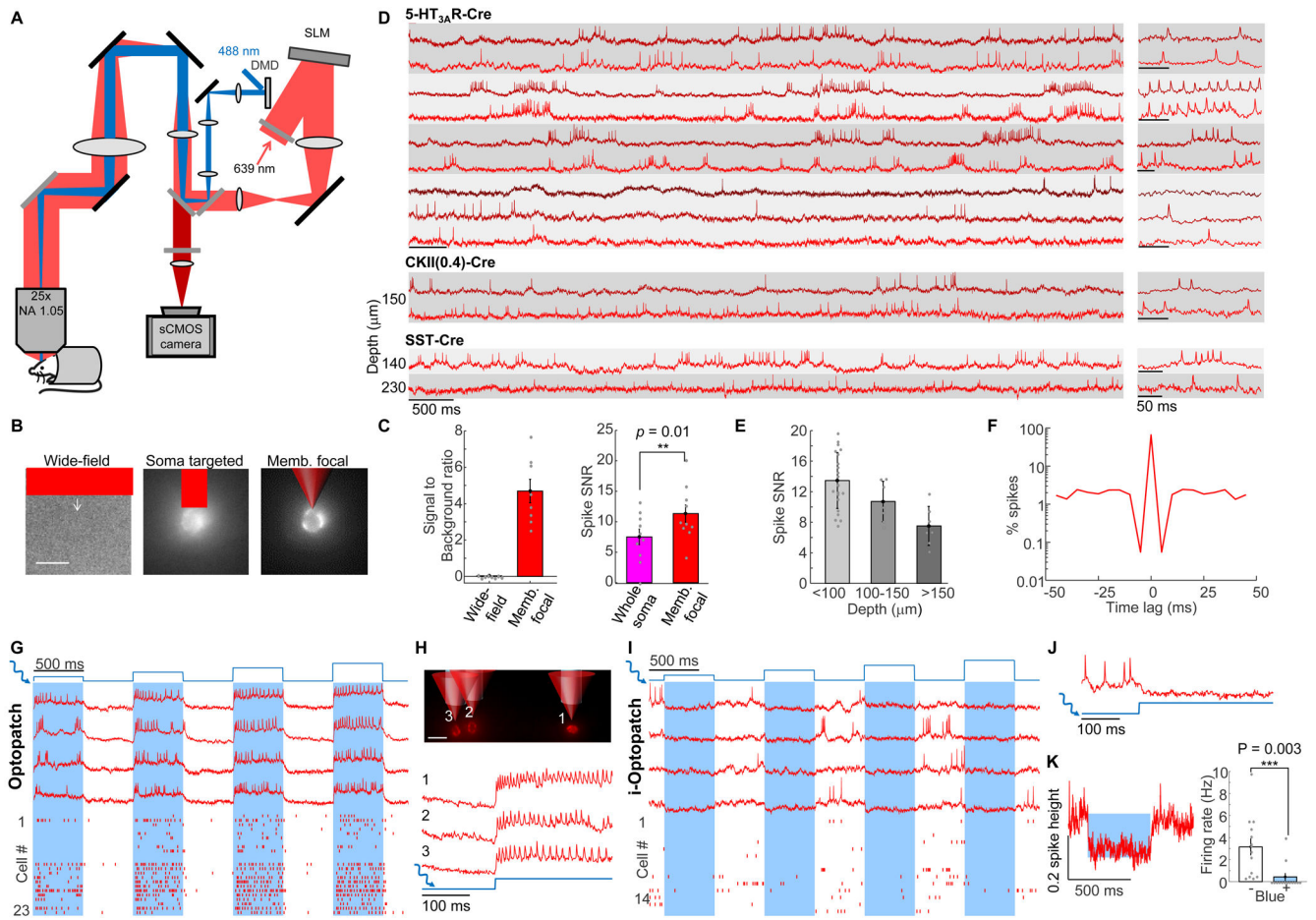


Figure 1. All-optical electrophysiology *in vivo*.

(A) Optical system for holographic structured illumination voltage imaging (red) and micromirror-patterned optogenetic stimulation (blue). Details in Methods. (B) Comparison of wide-field epifluorescence, soma-targeted, and membrane-targeted holographic illumination of the same field of view containing a SomArchon-expressing cortical neuron. Arrow indicates location of cell in wide-field image. Scale bar 20 μm . (C) Left: Signal-to-background ratio for wide-field and membrane-targeted imaging modalities ($n = 8$ cells). Right: SNR of action potentials with soma-wide and membrane-targeted holographic illumination ($n = 10$ cells). (D) Voltage imaging in cortical neurons at different depths with different promoters in awake mice. Grey bands indicate simultaneously recorded cells. Right: magnified views. Traces corrected for photobleaching but not otherwise filtered. (E) SNR of action potentials at different depths (error bars: mean \pm s.d.; $n = 7 - 22$ cells per depth). (F) Spike-triggered autocorrelogram showing refractory period ($n = 27$ cells). (G) Simultaneous optogenetic stimulation and voltage imaging in L1 interneurons in awake mice expressing Optopatch4. Groups of 1 – 3 cells were stimulated with patterned blue light (1.8 to 21 mW/mm^2). Bottom: Spike raster ($n = 23$ cells, 3 mice). (H) Top: Three targeted cells, with red and blue illumination overlaid. Scale bar 20 μm . Bottom: Fluorescence traces from the three cells in response to blue illumination. (I) Simultaneous optogenetic inhibition and voltage imaging (i-Optopatch). Cells co-expressed stGtACR2 and SomArchon. Groups of 1

– 3 cells were inhibited with patterned blue light (1.8 to 21 mW/mm²). Bottom: Spike raster ($n = 14$ cells, 2 mice). (J) Magnified view showing hyperpolarization and silencing upon blue light onset. (K) Left: Mean fluorescence response to blue illumination (1.8 mW/mm²). Right: Blue light stimulation significantly decreased the spontaneous firing rate ($n = 14$ cells, 2 mice, 1.8 mW/mm²).

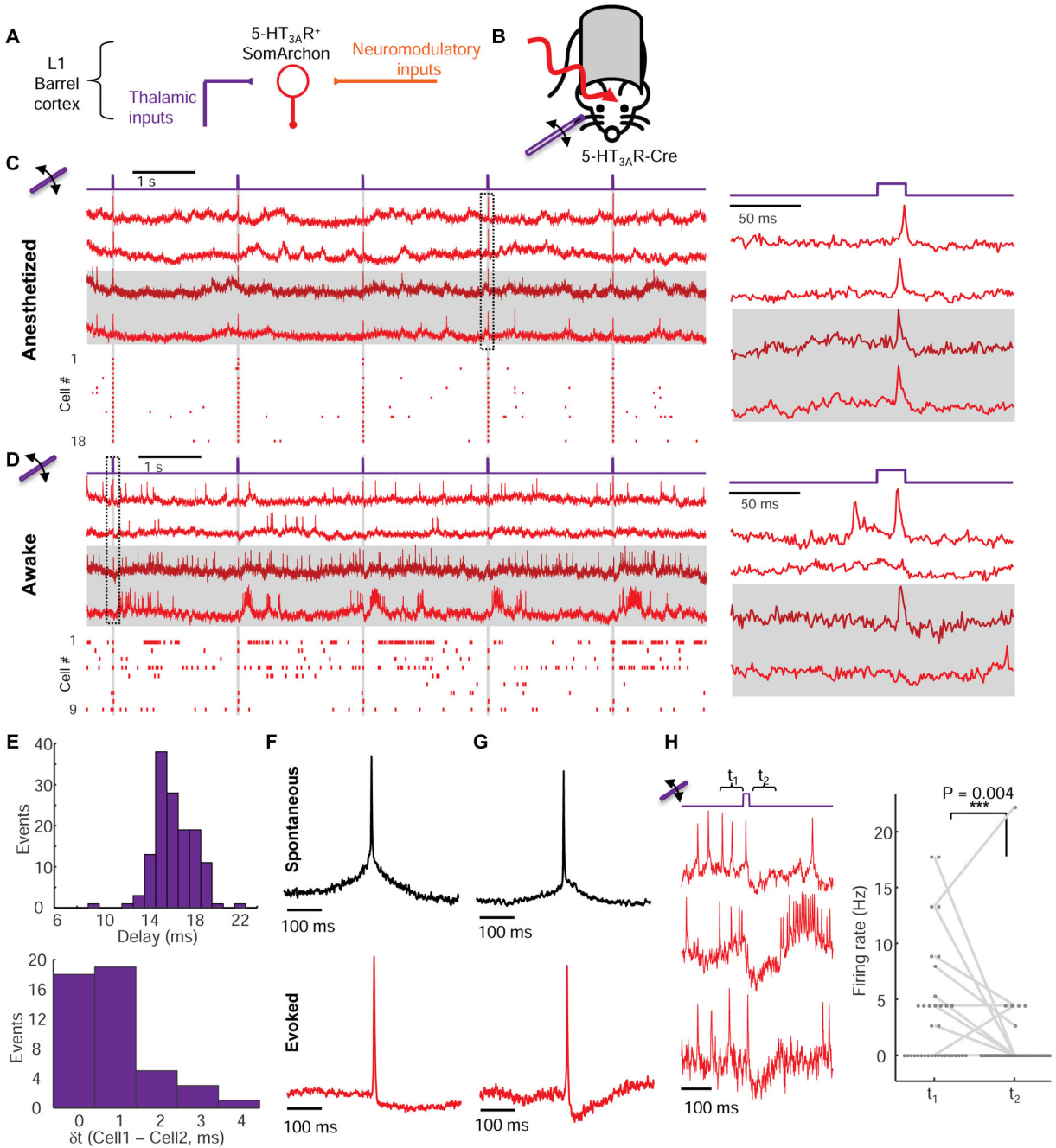


Figure 2. Sensory-evoked responses in L1 neurons *in vivo*.

(A) 5-HT_{3A}R-positive interneurons in barrel cortex L1 receive thalamocortical (sensory) and neuromodulatory inputs. (B) Simultaneous sensory stimulation and voltage imaging in L1. (C) Fluorescence transients in single L1 interneurons evoked by whisker stimuli (20 ms deflections at 0.5 Hz) in isoflurane anesthetized mice. Left, top: example fluorescence traces. Horizontal stripes indicate simultaneously recorded cells. Bottom: spike raster ($n = 18$ neurons, 3 mice). Right: Fluorescence waveforms from the boxed region at left. (D) Same as C but in awake mice. (E) Top: Distribution of delays between stimulus onset and peak of evoked spike ($n = 24$ cells, 135 trials). Bottom: Distribution of relative delays in sensory-

evoked action potential peaks between simultaneously recorded pairs of cells ($n = 14$ pairs, 66 trials). **(F)** Spike-triggered average waveform of spontaneous (top, $n = 17$ neurons) and whisker stimulus-evoked (bottom, $n = 24$ neurons) action potentials. **(G)** Same as **F** but in an awake mouse. Spontaneous: $n = 22$ neurons. Evoked: $n = 21$ neurons, 3 mice. Stimulus-evoked spikes had smaller after-spike hyperpolarization under anesthesia than under wakefulness (anesthetized: $11 \pm 2\%$ spike height, $n = 24$ neurons, 3 mice vs. awake: $17 \pm 1\%$ of spike height, $n = 21$ neurons, 3 mice, $p = 0.02$, two-tailed t -test), consistent with a more depolarized resting potential under wakefulness (Constantinople and Bruno, 2011). **(H)** Sensory stimulation induced a period of reduced spontaneous activity in awake mice.

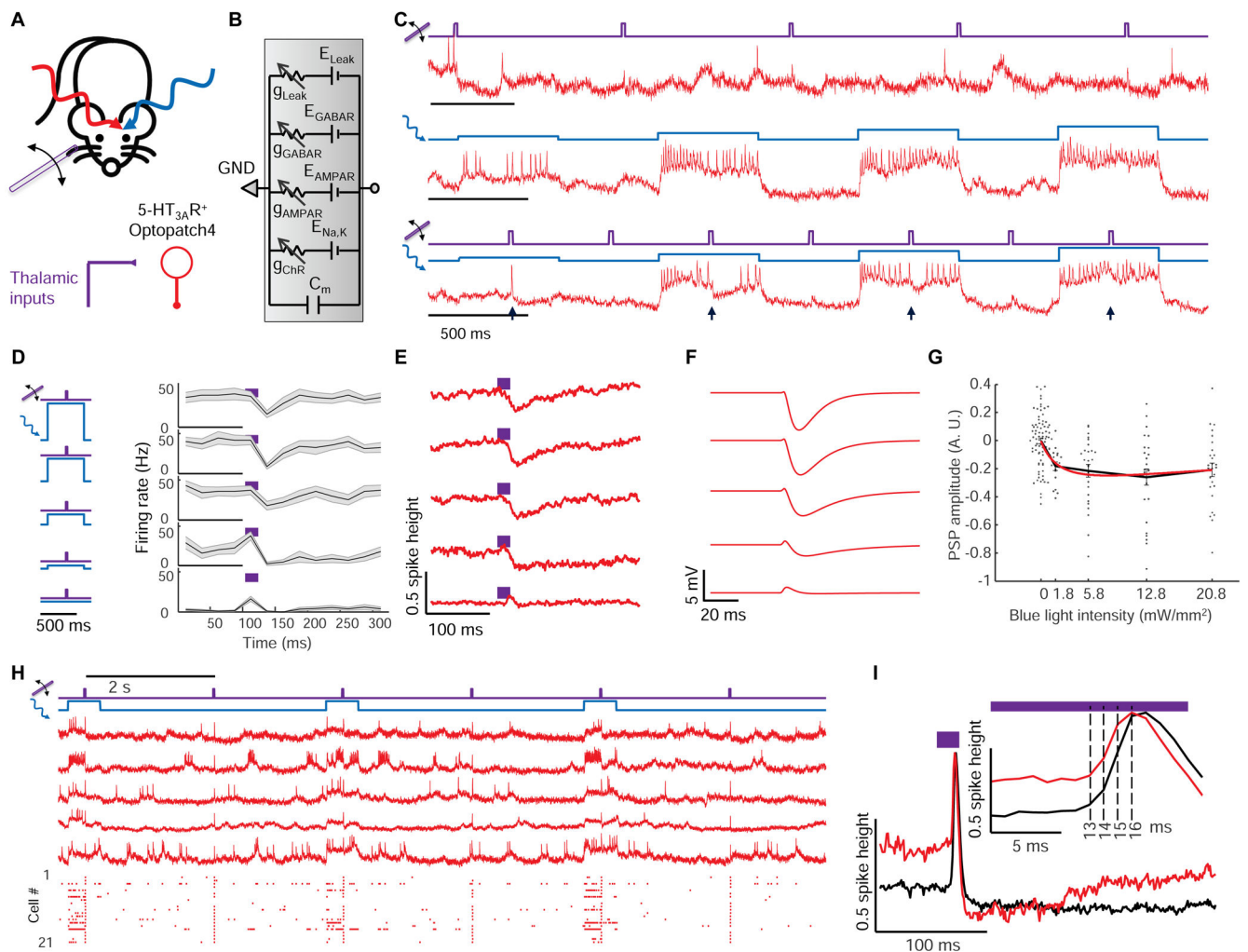


Figure 3. Optical dissection of excitation and inhibition in L1 interneurons in awake mice. (A) Whisker stimuli and single cell-targeted optogenetic stimuli were paired in 5HT_{3A}R-Cre mice expressing Optopatch4. (B) Conductance-based model of subthreshold membrane potential. This simple model only contained passive conductances, with gating by light (Channelrhodopsin, ChR), glutamate (AMPA), and GABA (GABA). A leak conductance set the resting potential. (C) Three recordings from a single neuron showing response to (top) whisker stimulus, (middle) targeted optogenetic stimulus, and (bottom) simultaneous optogenetic and whisker stimuli. Arrows show whisker stimulus-evoked inhibition. (D) Mean spike rate evoked by whisker stimuli at different levels of targeted optogenetic depolarization. In the absence of optogenetic stimulation, whisker stimuli evoked precisely timed single spikes. In the presence of optogenetic stimulation, whisker stimuli suppressed spiking. The suppression decreased in amplitude and duration as the strength of the optogenetic stimulus increased, a consequence of ChR shunting. Shading represents s.e.m. from $n = 27$ neurons, 4 mice. (E) Mean whisker stimulus-evoked subthreshold waveforms at different levels of optogenetic drive. Spikes were digitally removed prior to averaging (Methods). (F) Simulated membrane voltage waveforms under different levels of optogenetic drive, using the model shown in B. Excitation was assumed to lead inhibition by

2 ms (Methods). **(G)** Comparison of simulated (red) and measured (points) PSP amplitude as a function of optogenetic stimulus strength. **(H)** Repetitive measurements of whisker stimulus-evoked responses in anesthetized mice, with and without baseline optogenetic stimulation. Top: example recordings. Bottom: spike raster ($n = 21$ neurons, 3 mice). **(I)** Stimulus-triggered mean fluorescence responses without (black) and with (red) baseline optogenetic stimulation. Traces were aligned vertically to their peak. Inset: Magnified view showing dynamics near the peak.

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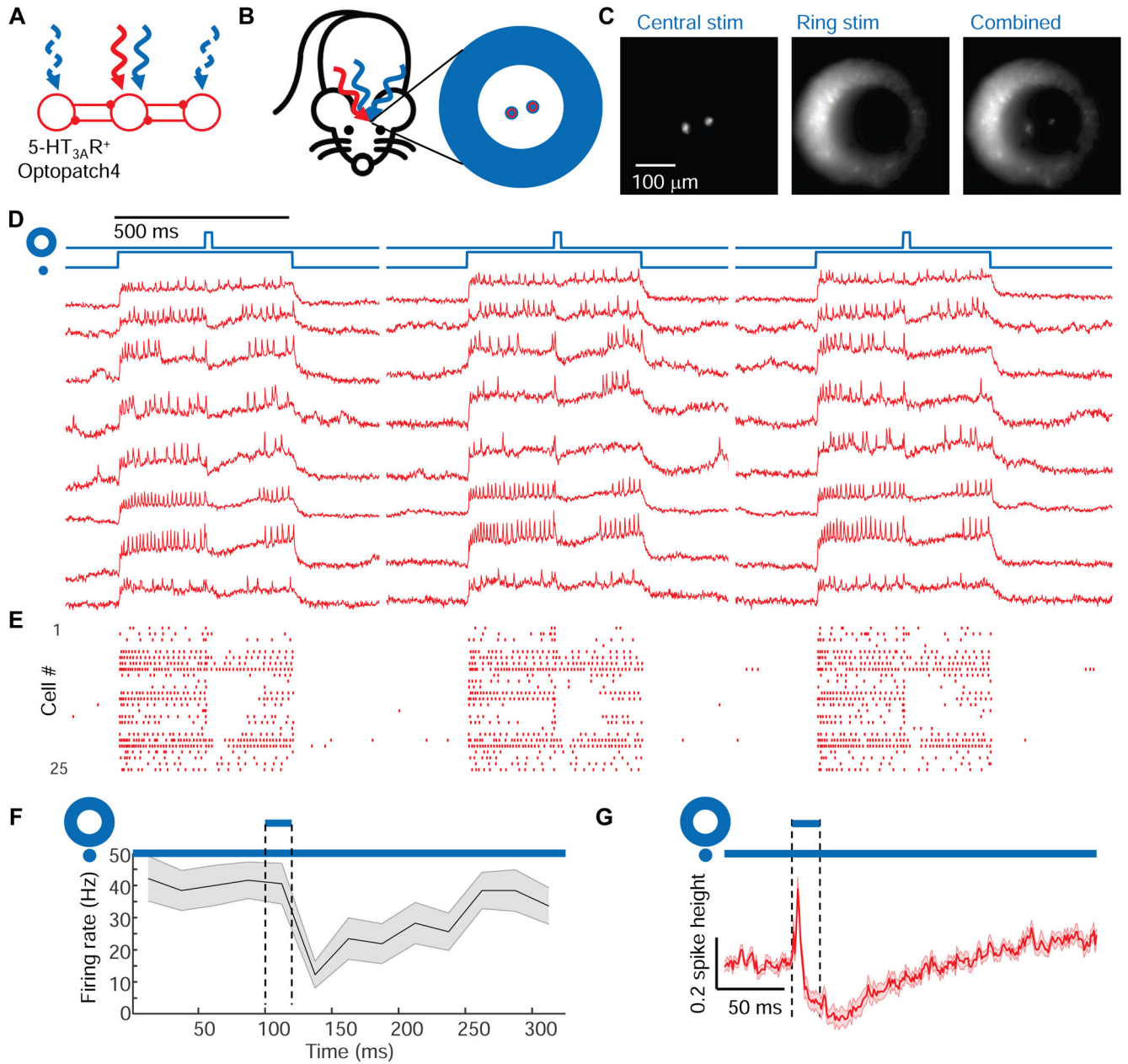


Figure 4. Center/surround optogenetic stimulation reveals lateral inhibition in barrel cortex L1. (A) Simple model of L1 circuit with lateral inhibition. Tonic optogenetic depolarization increased the driving force for inhibitory currents in the central neuron. Pulsed optogenetic stimulation of the surrounding neurons (blue) evoked lateral inhibition, revealed by voltage imaging (red). (B) Configuration of stimulation and imaging spots to probe lateral inhibition in L1 of 5-HT_{3A}R-Cre mice. (C) Epifluorescence images showing the illumination patterns *in vivo*. (D) Fluorescence waveforms from the central neurons under center/surround optogenetic stimulation. (E) Spike raster ($n = 25$ neurons, 3 mice). (F) Mean spike rate during central stimulation, before and after surround stimulation, $n = 25$ neurons, 3 mice. Shading represents s.e.m. (G) Mean subthreshold voltage in an optogenetically depolarized

central neuron during, before and after surround stimulation. The initial voltage increase was due to scattered light from the surround stimulus (Fig. S2). Shading represents s.e.m.

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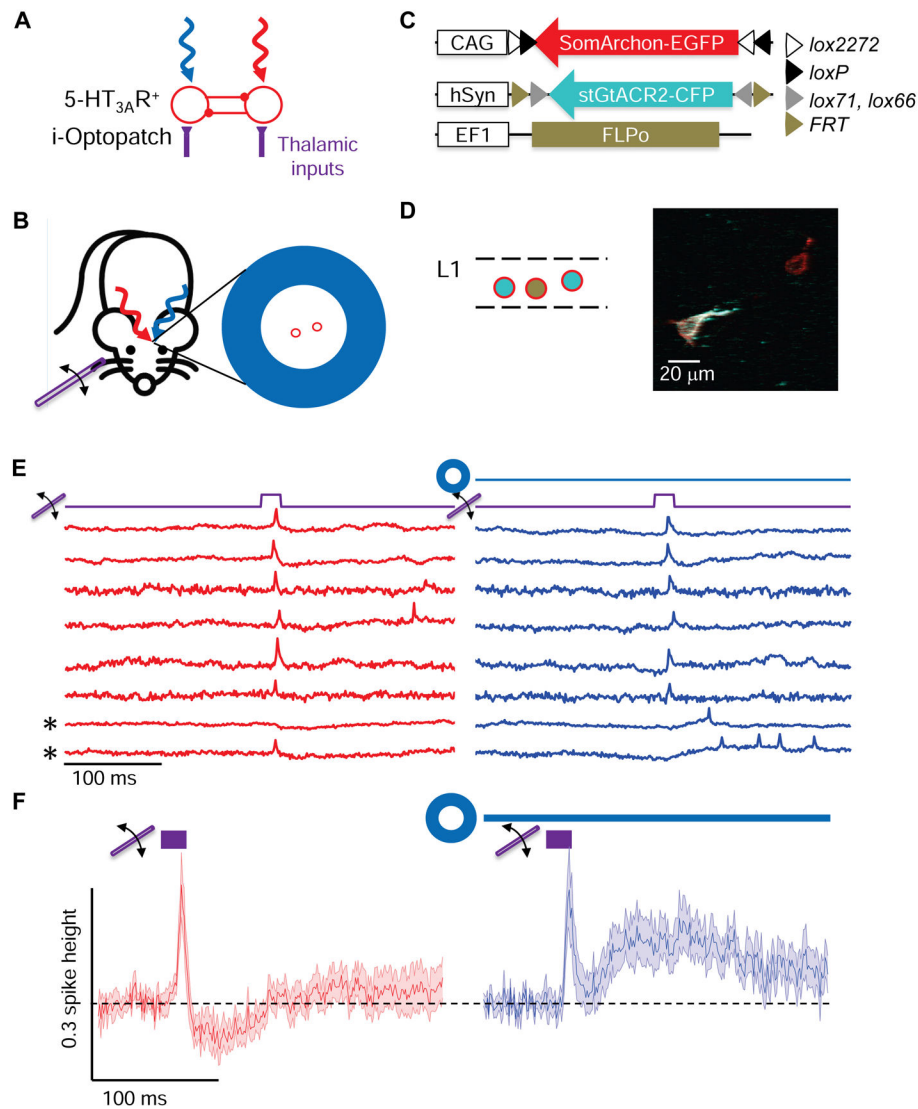
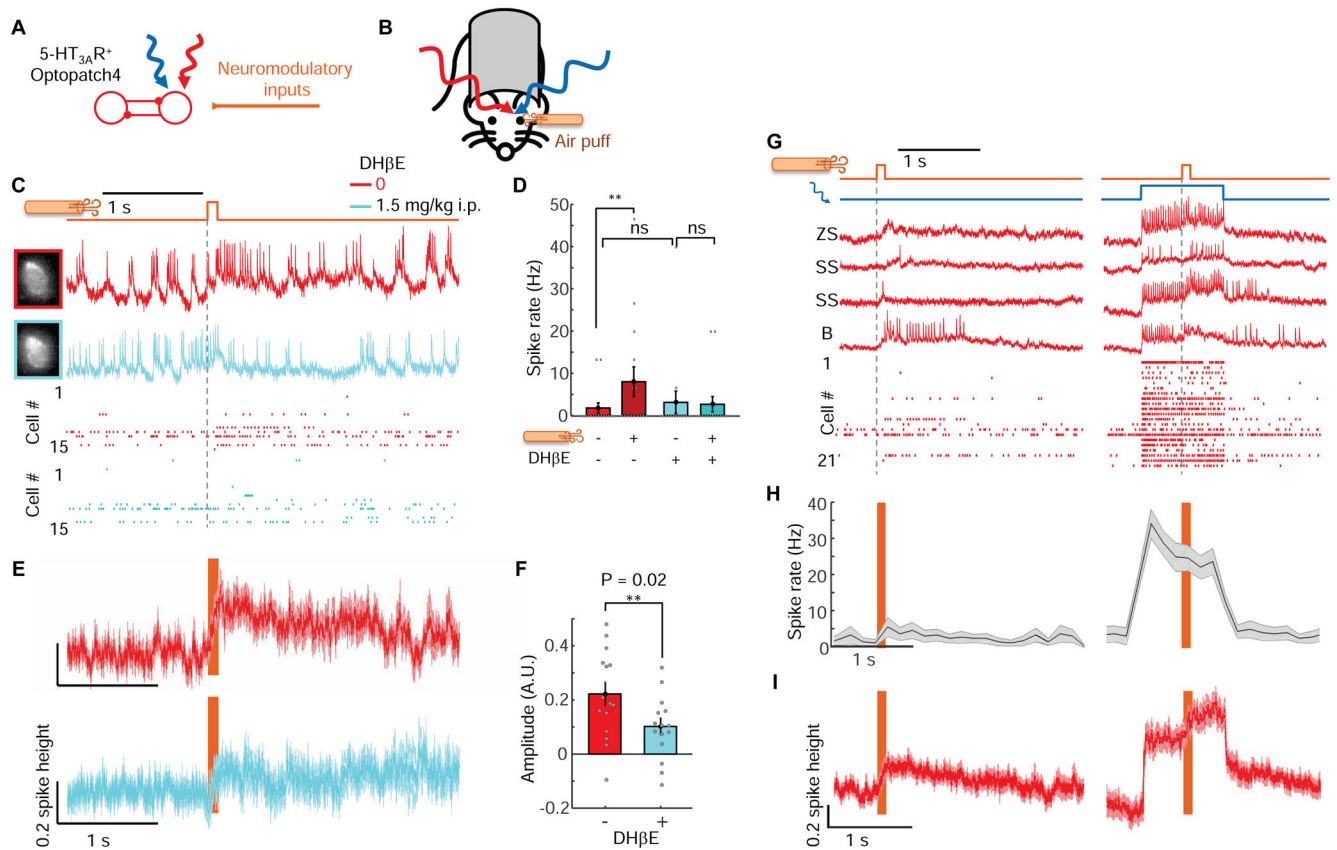


Figure 5. Optogenetic silencing reveals that L1 network activity is necessary for sensory-evoked lateral inhibition.

(A,B) Sensory-evoked responses in centrally located neurons were recorded (red) either without or with optogenetic silencing of surrounding L1 neurons (blue). (C, D) Targeted gene expression. Cre-on SomArchon-EGFP expressed in all 5-HT_{3A}R-Cre⁺ neurons. Cre-on-flp-off stGtACR2-CFP and low-titer Flpo virus were combined, so stGtACR2-CFP expressed in Cre⁺/Flp⁻ neurons. (D) Left: schematic of gene expression patterns. Colors indicate constructs in (C). Right: Composite two-photon fluorescence image of GFP fluorescence from SomArchon-EGFP and CFP fluorescence from stGtACR2-CFP. Voltage imaging was performed only in neurons that did not express stGtACR2. (E) Paired recordings in awake mice of sensory-evoked responses from individual centrally located neurons, either without (red) or with (blue) inhibition of surrounding neurons. Asterisks indicate recordings where sensory stimuli evoked delayed spiking when lateral inhibition was suppressed. (F) Whisker stimulus-triggered average membrane potential without (red) or with (blue) surround inhibition ($n = 13$ neurons, 2 mice, shading represents s.e.m.).



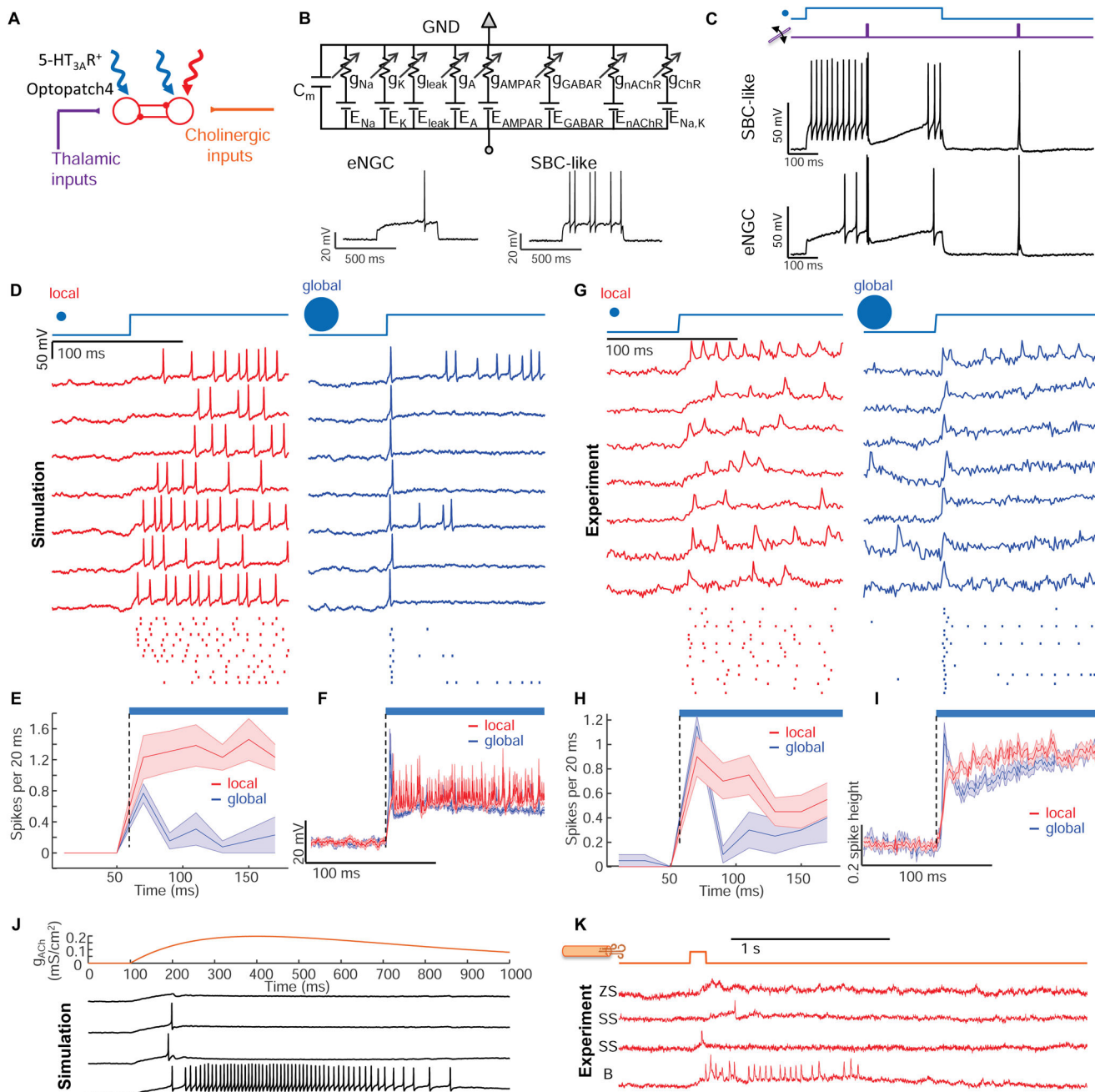


Figure 7. Numerical simulations capture input-output properties of L1 microcircuit. (A) Model of L1 function. Thalamic, cholinergic, and optogenetic inputs are modulated by lateral inhibition. (B) Conductance-based models for two subtypes of L1 interneurons mimic the intrinsic firing patterns. (C) Simulation of L1 network with network-wide thalamic excitation and single-cell targeted optogenetic stimulation. The traces show two simulations in which the optogenetic stimulation and voltage measurement were targeted either to an eNGC cell or to an SBC-like cell. Compare to Figs. 3C, S5. (D) Simulation of L1 microcircuit response to patterned optogenetic stimulation. Localized optogenetic stimulation induced sustained firing while global optogenetic stimulation induced firing in

all neurons only at the onset of the stimulation. **(E)** Population-average firing rates in response to local or global stimulation. **(F)** Population-average membrane potential in response to local or global stimulation. **(G-I)** Experimental test of the predictions in **D – F** with patterned optogenetic stimulation. For the local stimulation, spot sizes were 30 μm diameter. For global stimulation, spot sizes were 400 μm in diameter. **(J)** Simulated responses to cholinergic inputs showing winner-takes-all dynamics. **(K)** Experimental responses to air puffs (same data as in Fig. 6G). The air puff responses closely matched the simulated winner-takes-all dynamics.

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KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and Virus Strains		
LZF1735 pAAV_hSyn-DiO-SomArchon-eGFP-P2A-somCheRiff (Optopatch4)	Janelia Farm Vector Core	N/A
pAAV_CAG-FLEX-SomArchon-eGFP	UNC Vector Core	N/A
CKII(0.4)-Cre	UNC Vector Core	N/A
LZF1826 pAAV hSyn-DiO-SomArchon-eGFP-P2A-stGTACR2	Janelia Farm Vector Core	N/A
LZF 1827 pAAV_hSyn-Flp-off-Cre-on-stGtACR2-CFP	Janelia Farm Vector Core	N/A
pAAV-EF1a-Flpo	UNC Vector Core	N/A
Chemicals, Peptides, and Recombinant Proteins		
dihydro- β -erythroidine hydrobromide (DH β E)	Tocris	cat# 2349
TransIT-293	Mirus	cat# MIR2704
glass capillaries	World precision Instrument	cat# TW120-3
Experimental Models: Cell Lines		
Hek293T	ATCC	CRL-11268
Experimental Models: Organisms/Strains		
5HT _{3A} R-Cre transgenic mice	Takao Hensch Lab	N/A
C57BL/6 wild-type mice	Charles River	N/A
SST-Cre transgenic mice	Jackson Lab	Stock #013044
NDNF-Cre transgenic mice	Jackson Lab	Stock #028536
Recombinant DNA		
LZF1735 pAAV_hSyn-DiO-SomArchon-eGFP-P2A-somCheRiff (Optopatch4)	This work	Addgene 126512
LZF1826 pAAV hSyn-DiO-SomArchon-eGFP-P2A-stGTACR2	This work	Addgene 135412
LZF1827 pAAV_hSyn-Flp-off-Cre-on-stGtACR2-CFP	This work	Addgene 135413
pAAV_hSyn1-SIO-stGtACR2-FusionRed	Ofer Yizhar Lab	Addgene 105677
UB061 FCK-stGtACR2-fusionRed (Lenti-vector)	This Work	Addgene 135765
Software and Algorithms		
Matlab R2015b	Mathworks	https://www.mathworks.com/products/matlab.html
Labview	National Instrument	N/A
Other		
Holographic structured illumination microscope	This work	N/A