Widespread and Highly Correlated Somato-dendritic Activity in Cortical Layer 5 Neurons

**Highlights**
- Layer 5 neuron apical dendrites exhibit frequent GCaMP6f signals
- Calcium electrogenesis underlies dendritic GCaMP6f signals
- Dendritic activity is highly correlated with somatic activity
- Visual stimuli and running do not change somato-dendritic GCaMP6f correlation

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**In Brief**
Beaulieu-Laroche et al. perform near-simultaneous calcium imaging of somatic and dendritic activity to reveal that active dendritic integration is an integral feature of information processing in cortical pyramidal neurons.
Widespread and Highly Correlated Somato-dendritic Activity in Cortical Layer 5 Neurons

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SUMMARY

Dendritic integration can expand the information-processing capabilities of neurons. However, the recruitment of active dendritic processing in vivo and its relationship to somatic activity remain poorly understood. Here, we use two-photon GCaMP6f imaging to simultaneously monitor dendritic and somatic compartments in the awake primary visual cortex. Activity in layer 5 pyramidal neuron somata and distal apical trunk dendrites shows surprisingly high functional correlation. This strong coupling persists across neural activity levels and is unchanged by visual stimuli and locomotion. Ex vivo combined somato-dendritic patch-clamp and GCaMP6f recordings indicate that dendritic signals specifically reflect local electrogenesis triggered by dendritic inputs or high-frequency bursts of somatic action potentials. In contrast to the view that dendrites are only sparsely recruited under highly specific conditions in vivo, our results provide evidence that active dendritic integration is a widespread and intrinsic feature of cortical computation.

INTRODUCTION

Dendrites shape how synaptic inputs are integrated into behaviorally relevant outputs at the level of individual neurons (London and Häusser, 2005; Magee, 2000; Major et al., 2013; Stuart and Spruston, 2015). The amplification of synaptic inputs by voltage-gated ion channels can produce nonlinear dendritic spikes, which, in turn, trigger or modulate somatic action potential output. Layer 5 (L5) pyramidal neurons possess long apical dendrites with powerful spikes mediated by calcium electrogenesis (Harnett et al., 2013; Larkum et al., 1999; Schiller et al., 1997; Williams, 2004). Recordings in anesthetized animals have demonstrated that dendritic calcium electrogenesis occurs in vivo and usually coincides with somatic burst firing (Helmchen et al., 1999). Calcium imaging has revealed previously that apical dendritic activity can represent behaviorally relevant features in trained animals (Peters et al., 2017; Ranganathan et al., 2018; Takahashi et al., 2016; Xu et al., 2012). However, because somatic activity has not been concurrently assayed in awake animals, the relationship between compartments remains unclear. The prevalence of active dendritic integration in layer 5 is also unknown: prior experiments sparsely sampled neurons because of the GCaMP expression approach used and the neural dynamics during task performance. Is dendritic activity infrequent and independent of somatic activity, or is it a widespread and integral aspect of cellular computation?

To determine how dendritic processing relates to cortical neuron output, we simultaneously measured dendritic and somatic GCaMP6f signals from L5 neurons in the primary visual cortex (V1) of awake mice. Because there are no existing ground truth calibrations of GCaMP6f signals in L5 somata or in dendrites, we combined multi-site whole-cell patch-clamp electrophysiology with GCaMP6f imaging in acute mouse brain slices to interpret in vivo results.

RESULTS

We first imaged V1 neurons in awake head-fixed mice that were free to run on a cylindrical treadmill. We quasi-simultaneously recorded L5 somata (527.8 ± 9.1 μm deep, n = 12 fields of view in 5 mice) and their distal apical trunk dendrites in layer 2 (167.1 ± 6.2 μm deep, 360.8 ± 6.7 μm from the soma) at 15.46 Hz using an electrically tunable lens coupled to a two-photon microscope (Figures 1A–1C). Dendritic activity was detectable in all soma-dendrite pairs that exhibited somatic transients (Figure 1E; 156 of 157 pairs from 5 mice). This widespread dendritic engagement was associated with synchronous activity patterns between corresponding somata and dendrites (Figures 1C and S1), resulting in dendritic and somatic signals that were strongly correlated (Figure 1D; GCaMP6f signal correlation coefficient, 0.542 ± 0.008; n = 156 from 5 mice). Although corresponding somatic and dendritic GCaMP6f traces were similar, their amplitudes and kinetics were not closely matched (Figure 1C). To determine whether these differences reflect divergent features of dendritic and somatic compartments as opposed to noisy calcium dynamics or optical measurements, we analyzed soma-dendrite pairs and pairs of dendrites originating from the same soma that branched below the dendrite imaging plane (GCaMP6f signal correlation coefficient, 0.754 ± 0.033, n = 29 from 5 mice). Dendritic signals consistently had faster rise and decay kinetics than somatic signals (Figures 1H–1J). Thus, corresponding somatic and dendritic GCaMP6f signals are strongly...
In vivo multi-plane imaging

Figure 1. Dendritic GCaMP6f Signals Are Widespread across the Population and Highly Correlated with Somatic Signals
(A) Left: experimental approach to simultaneously record somatic and dendritic signals from L5 pyramidal neurons. Right: two-photon mean image at the two imaging planes. The yellow regions highlight a soma and its corresponding dendrite.

To dissect the relationship between somatic and dendritic activity, we isolated transients (Figure 2A; STAR Methods). We found that the majority of transients were paired with a transient in the other compartment (Figure 2B; paired dendritic transient; median [Q1-Q3]: 82.3% [69.1% 92.4%]; paired somatic transient; median [Q1-Q3]: 84.8% [65.8% 93.3%]; n = 156 from 5 mice). Furthermore, paired transients were much larger for the soma (somatic transient integral; paired, median [Q1-Q3]: 84.8% [65.8% 93.3%]; n = 156 from 5 mice). Furthermore, paired transients were much larger for the soma (somatic transient integral; paired, median [Q1-Q3]: 82.3% [69.1% 92.4%]; paired somatic transient; median [Q1-Q3]: 83.9% [71.1% 91.1%]; n = 156 from 5 mice).

Having established that most transients were paired, we next assessed the spatial coupling of these events. To do so, we correlated the first-order derivative of the GCaMP6f transients across all compartments, representing segments with the gray dashed box. We then computed a coupling index (Figure 2D), which reflects the time integral of the paired and unpaired transients, accounting for the different magnitudes of each dendrite and soma (***p < 10^{-22}, Wilcoxon rank-sum test, n = 156 pairs from 5 mice).

To estimate how much of the activity is coupled while accounting for the different magnitudes of the paired and unpaired transients, we computed a coupling index (Figure 2D), which reflects the time integral of paired transients over the time integral of all transients. This analysis revealed that ~97% of the activity is coupled (dendrite coupling index: median [Q1-Q3]: 0.965 [0.916 0.987]; soma coupling index: median [Q1-Q3]: 0.972 [0.897 0.992], n = 156 from 5 mice). GCaMP6f transients lasted up to minutes because of the temporal summation of multiple calcium signals. To assess coupling on a finer timescale, we separated rise and decay segments in the transients based on the first-order derivative of the GCaMP6f signal. We isolated rise events that correspond to segments with positive derivatives that likely reflect calcium entry and underlying electrical activity (Peters et al., 2017; Figure 2E; STAR Methods). Similar to the transients, most rise events were paired with a rise event in the other compartment (Figure 2F; paired dendritic rise event; median [Q1-Q3]: 83.9% [71.1% 91.1%]; paired somatic rise event; median [Q1-Q3]: 73.4% [56.8% 85.9%], n = 156 from 5 mice).

Furthermore, the amplitude of rise events was highly correlated with somatic signals (Figure 2F; paired somatic rise event; median [Q1-Q3]: 66.4% [36.9 133.2%]; paired dendritic rise event; median [Q1-Q3]: 83.9% [71.1% 91.1%]; paired somatic rise event; median [Q1-Q3]: 73.4% [56.8% 85.9%], n = 156 from 5 mice). To estimate how much of the activity is coupled while accounting for the different magnitudes of the paired and unpaired transients, we computed a coupling index (Figure 2D), which reflects the time integral of paired transients over the time integral of all transients. This analysis revealed that ~97% of the activity is coupled (dendrite coupling index: median [Q1-Q3]: 0.965 [0.916 0.987]; soma coupling index: median [Q1-Q3]: 0.972 [0.897 0.992], n = 156 from 5 mice). GCaMP6f transients lasted up to minutes because of the temporal summation of multiple calcium signals. To assess coupling on a finer timescale, we separated rise and decay segments in the transients based on the first-order derivative of the GCaMP6f signal. We isolated rise events that correspond to segments with positive derivatives that likely reflect calcium entry and underlying electrical activity (Peters et al., 2017; Figure 2E; STAR Methods). Similar to the transients, most rise events were paired with a rise event in the other compartment (Figure 2F; paired dendritic rise event; median [Q1-Q3]: 83.9% [71.1% 91.1%]; paired somatic rise event; median [Q1-Q3]: 73.4% [56.8% 85.9%], n = 156 from 5 mice). Furthermore, the amplitude of
Figure 2. The Majority of GCaMP6f Transients Are Paired between Corresponding Somata and Dendrites

(A–C) Somatic and dendritic transients were categorized as paired or unpaired based on their temporal overlap with transients in the other compartment. (A) GCaMP6f signals from the soma and dendrite, with detected transients shown in solid colors. The asterisk highlights an unpaired dendritic transient. (B) Percentage of paired somatic and dendritic events (n = 156 pairs from 5 mice). (C) Events from the neuron shown in (A), categorized as paired and unpaired. Gray and light red lines represent individual transients (750 ms before the onset and up to 2,600 ms thereafter), whereas averages are shown in black and red. The number of paired or unpaired transients over the total number of transients is indicated above each panel. (D) Coupling index for all soma-dendrite pairs (n = 156 pairs from 5 mice).

Figure 3A). Dendritic whole-cell recordings (348.0 ± 12.8 µm from the soma, n = 10 dendrites from 7 mice) revealed a nonlinear relationship between voltage and GCaMP6f fluorescence (Figures 3B–3D). Subthreshold depolarization produced negligible signals, whereas suprathreshold dendritic electrogenesis produced significant GCaMP6f signals (Figures 3B–3D). The rise events (maximum – minimum ΔF/F within an event) was correlated between the two compartments (Figures 2G and 2H; rise event amplitude correlation coefficient: median [Q1-Q3]: 0.554 [0.473 0.645], n = 156 from 5 mice). This strong somato-dendritic coupling resulted in similar tuning between the two compartments (Figure S1). These analyses reveal that most GCaMP6f events occur concurrently in corresponding dendritic and somatic compartments.

To determine the physiological events that underlie the correlated somato-dendritic GCaMP6f transients we observed in vivo, we performed whole-cell recordings from GCaMP6f-expressing L5 neurons in acute brain slices from mouse V1 (Figure 3A). Dendritic whole-cell recordings (348.0 ± 12.8 µm from the soma, n = 10 dendrites from 7 mice) revealed a nonlinear relationship between voltage and GCaMP6f fluorescence (Figures 3B–3D). Subthreshold depolarization produced negligible signals, whereas suprathreshold dendritic electrogenesis produced significant GCaMP6f signals (Figures 3B–3D). The amplitude of the GCaMP6f transients reflected the duration of the suprathreshold dendritic spike (Figure S2). Importantly, dendritic spikes of all durations produced significant ΔF/F changes (Figure S2). Backpropagating action potentials (bAPs) can evoke calcium influx into dendrites (Grienberger and Konnerth, 2012; Helmchen et al., 1996; Hill et al., 2013; Maravall et al., 2000; Markram et al., 1995; Ranganathan et al., 2018; Schiller et al., 1995; Spruston et al., 1995). Indeed, a critical frequency of bAPs (~100 Hz) has been shown to trigger dendritic electrogensis (Larkum et al., 1999; Shai et al., 2015; Williams and Stuart, 2000). We therefore investigated whether somatically evoked APs could elicit widespread somato-dendritic GCaMP6f signals. We performed dual whole-cell recordings (Figure 3F) and found that high-frequency bAPs (200 Hz), but not low-frequency bAPs (50 Hz), engaged dendritic electrogensis to produce dendritic GCaMP6f signals (Figure 3E). Somatic GCaMP6f signals also displayed a strong frequency dependence (Figures 3E and 3H; n = 21 cells from 9 mice) so that APs produced similar signals in the soma and dendrite (Figure 3I). Unlike L2/3 somata (Chen et al., 2013),
L5 somata exhibited negligible signals for low-frequency or single APs (Figures S3). Thus, low-frequency bAPs do not underlie dendritic GCaMP6f signals, but high-frequency bursts can trigger dendritic electrogenesis to produce dendritic GCaMP6f signals. Conversely, directly triggering dendritic input can trigger dendritic electrogenesis to produce dendritic GCaMP6f signals, but high-frequency bursts of somatic APs (Figures S3). Thus, low-frequency bAPs do not un-

Figure 3. Ex Vivo Calibration of Dendritic GCaMP6f Signals
(A) Left: two-photon z stack montage image of a GCaMP6f-expressing mouse L5 neuron. The dashed line indicates the location of calcium imaging next to the dendritic patch-clamp electrode 373 μm from the soma. Right: dendritic voltage (top) in response to step current injection (bottom). (B) Dendritic GCaMP6f signal (top) associated with dendritic voltage (bottom). (C) GCaMP6f peak as a function of peak voltage for increasing 100-ms step current injections for the dendrite shown in (A)—(C). (D) Dendritic GCaMP6f peak as a function of peak voltage for increasing 100-ms step current injections (n = 10 dendrites from 7 mice). (E) Membrane potential and GCaMP6f signals at the dendrite (red, top) and soma (black, bottom) during slow (50 Hz, left) and fast (200 Hz, right) APs evoked by somatic current injection (gray) in the neuron shown in (F). (F) Two-photon image of a GCaMP6f-expressing L5 neuron with somatic and dendritic (320 μm from the soma) patch-clamp electrodes. (G) Membrane potential and GCaMP6f signals at the dendrite (red, top) and soma (black, bottom) during short (20 ms, left) and long (100 ms, right) dendritic spikes evoked by dendritic current injection (light red) in the neuron shown in (F). (H) GCaMP6f signals at the soma (left) and dendrite (right) for 10 somatic APs at the indicated frequencies in the neuron shown in (F). (I) Somatic (black) and dendritic (red) GCaMP6f peak for 10 APs as a function of AP frequency (n = 21 neurons from 9 mice). Pooled data represent mean ± SEM. (J) Somatic (black) and dendritic (red) GCaMP6f peak as a function of dendritic spike duration (n = 10 neurons from 7 mice). Pooled data represent mean ± SEM. See also Figures S2 and S3.
demonstrate that visual inputs and locomotion do not alter strong somato-dendritic GCaMP6f coupling despite pronounced activity level changes.

**DISCUSSION**

By simultaneously imaging GCaMP6f in L5 distal apical dendrites and somata in awake animals, we reveal an unexpectedly broad engagement of active dendritic processing across the population of V1 L5 neurons as well as strong coupling between dendritic and somatic activity. Our ex vivo calibration experiments demonstrate that dendritic GCaMP6f signals specifically reflect dendritic electrogenesis. These dendritic spikes can trigger or be triggered by high-frequency action potentials, altering the pattern of axo-somatic output (Beaulieu-Laroche et al., 2018; Hamett et al., 2013; Shai et al., 2015; Williams, 2005; Williams and Stuart, 1999). Strong correlations between somatic and dendritic compartments indicate that apical dendritic spikes are an essential feature of single-cell computation in awake cortical circuits as opposed to rare coincidence detection events. Furthermore, widespread and frequent dendritic calcium influx could have important implications for plasticity induction and learning (Bittner et al., 2017; Golding et al., 2002; Guerguiev et al., 2017; Kampa et al., 2006; Magee and Johnston, 1997; Remy and Spruston, 2007; Sjöström and Häusser, 2006).

Although GCaMP6f signals show robust functional correlation, somatic and dendritic compartments exhibit electrical compartmentalization, especially at the subthreshold level (Beaulieu-Laroche et al., 2018; Fletcher and Williams, 2019; Spruston, 2008; Stuart and Spruston, 1998, 2015; Williams, 2004; Williams and Stuart, 2002). Previous studies indicate that electrical interactions between somata and dendrites can be variable (Helmchen et al., 1999) and modulated by inhibition (Larkum et al., 1999; Silberberg and Markram, 2007; Takahashi et al., 2016) or neuromodulators (Brombas et al., 2014; Labarrera et al., 2018; Williams and Fletcher, 2019). We show here that the coordination of suprathreshold activity is constant in Figure 4. Somato-dendritic GCaMP6f Correlations Are Conserved across Activity Levels and Unchanged by Visual Stimuli and Locomotion

(A) GCaMP6f signals from a soma and its corresponding dendrite during a dark screen (left) or presentation of natural movies (right).

(B) Comparison of somatic activity level (**p < 10^{-10}, Wilcoxon paired test, n = 156 pairs from 5 mice) during the dark and natural movie epochs.

(C) Comparison of dendritic activity level (**p < 10^{-19}, Wilcoxon paired test, n = 156 pairs from 5 mice) during the dark and natural movie epochs.

(D) Comparison of correlation coefficients (p = 0.81, Wilcoxon paired test, n = 156 pairs from 5 mice) in soma-dendrite pairs during the dark and natural movie epochs. The leftward point cluster represents neurons that were silent in the dark.

(E) GCaMP6f signals from the soma and dendrite (bottom) with running speed trace (top).

(F) Comparison of somatic activity level (**p < 10^{-8}, Wilcoxon paired test, n = 156 pairs from 5 mice) during the stationary and running epochs.

(G) Comparison of dendritic activity level (**p < 10^{-9}, Wilcoxon paired test, n = 156 pairs from 5 mice) during the stationary and running epochs.

(H) Comparison of correlation (p = 0.12, Wilcoxon paired test, n = 156 pairs from 5 mice) in paired soma-dendrites during the stationary and running epochs.

Pooled data represent median and interquartile range for (B)–(D) and (F)–(H). See also Figure S4.
the face of visual stimuli, locomotion, and activity level changes, at least within the limits of GCaMP6f imaging. However, flexible somato-dendritic communication may occur on a shorter timescale or may not act along simple human-interpretable categories (e.g., running versus not running). We conclude that active dendritic integration is a widespread and integral feature of information processing in L5 pyramidal neurons.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.neuron.2019.05.014.

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AUTHOR CONTRIBUTIONS

L.B.-L. designed the experiments, performed and analyzed slice experiments, acquired imaging data, analyzed in vivo experiments, prepared the figures, and wrote the manuscript. E.H.S.T performed surgeries, acquired imaging data, built analysis tools, and contributed to data analysis. N.J.B. performed surgeries and acquired imaging data. M.T.H conceived of and supervised the project.

DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES


## STAR METHODS

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METHOD DETAILS

Surgery
Mice were anesthetized with isoflurane and secured in a stereotaxic apparatus. A heating pad was used to maintain body temperature; additional heating was provided until fully recovered. The scalp was shaved, wiped with hair-removal cream, and cleaned with iodine solution and ethanol. After intraperitoneal injection of Dexamethasone (4 mg/kg) and Carprofen (5 mg/kg), and subcutaneous injection of slow-release Buprenorphine (0.5 mg/kg), the skull was exposed.

For in vivo imaging, using Rbp4-Cre+/+ mice aged 8-15 weeks, a 3 mm craniotomy was drilled centered at −3.8 mm anterior-posterior, +3.0 mm lateral from bregma. 50-100 nL of AAV1-syn-flex-GCaMP6f-WPRE-SV40 were injected in 3-4 sites, 650-700 μm from the surface of the brain. Cranial windows consisting of two stacked 3 mm coverslips (~0.1 mm) under a 5 mm coverslip were inserted into the craniotomy, and the edges of the larger glass were sealed with cyanoacrylate glue and dental cement. A head-post was implanted to allow head fixation. Imaging was performed 4-12 weeks post-surgery.

For slice experiments, using mice aged 7-12 weeks, 1-3 small craniotomies (burr holes) were performed in each hemisphere centered at −3.8 mm anterior-posterior, ± 3.0 mm lateral from bregma. 50-100 nL of the GCaMP6f virus described above were injected at each site at 650-750 μm from the surface of the brain in Rbp4-Cre+/− mice. For C57BL/6 wild-type, the GCaMP6f virus was mixed in equal part with a diluted AAV1-syn-Cre virus at 650-750 μm from the surface of the brain in Rbp4-Cre+/− mice. For C57BL/6 wild-type, the GCaMP6f virus was mixed in equal part with a diluted AAV1-syn-Cre virus at 650-750 μm or 150 μm (for L2/3; Figure S3) from the surface of the brain. Injected animals, including those some of those used for in vivo imaging, were used for slice physiology 4-12 weeks post-surgery.

Two-photon imaging
Imaging from behaving animals was performed with a Neurolabware standard microscope (http://neurolabware.com/microscope) equipped with GaAsP photomultiplier tubes. GCaMP6f was excited with a 980 nm ultrafast pulsed laser beam from a dispersion-compensated Insight DeepSee coupled to a 4x passive pulse splitter to reduce photodamage and bleaching (Ji et al., 2008). A water immersion objective (Nikon 16 x, 0.8 NA) was used for excitation and emission collection. Bi-directional frames (512 x 796 pixels) were simultaneously acquired at two planes with an electrically-tunable lens (Optotune EL-10-30-NIR-LD) at 30.92 Hz (15.46 Hz for each plane). Z stacks (1000-1500 frames every 5-7 μm) were acquired at the end of the experiments. Laser beam intensity was independently controlled with electro-optical modulators.

Visual stimuli were presented on the contralateral side via a monitor 20 cm from the mouse’s eye. Stimuli were luminance-normalized and spherically warped to compensate for the wide-angle field-of-view of the mice (http://help.brain-map.org/display/observatory/Documentation). A dark, opaque polymer cylinder was attached to the head-plate and surrounded the objective to limit light contamination from the monitor. Before imaging, animals were acclimatized over the course of several days to head-fixation on the behavior rig equipped with a cylindrical treadmill. For experiments in Figures 1, 2, 4, S1A, and S4, mice were presented with natural movies (black-and-white clips lasting 20-30 s). They were repeated ten times in a random order, separated by a ten-second gray screen. A black screen was presented before and after the visual stimuli series for 10-20 minutes each time. For experiments in Figures S1B–S1G, we presented only drifting gratings in separate imaging sessions. Drifting gratings (square waveform) were displayed for two seconds, followed by a five-second gray screen. Ten repetitions of each grating type were presented in a random order each session, sampled from eight orientations (0 to 315° in 45° increments) and four temporal frequencies (1, 2, 4, and 8 Hz), with a spatial frequency of 0.05 cycles/degree.

An in vitro galvanometer-based multiphoton microscope system (Bruker Ultima) with a water-immersion lens (60 x, 0.9 NA) was used to image brain slices. A dispersion-compensated Mai-Tai DeepSee laser was used to excite Alexa 594 at 880 nm and GCaMP6f at 920 nm (separated via dichroic mirrors to independent sets of GaAsP photosensor modules). Another photosensor module was used to collect transmitted-light Dodt gradient images for patch-clamp targeting. Laser beam intensity was independently controlled with electro-optical modulators. Line scan imaging was performed at somatic and dendritic sites at 400-1000 Hz with dwell times of 8 μs. Line scan and frame scan imaging produced comparable signals (Figure S3).
Cortical slice preparation
Cortical brain slices were prepared from adult (> 11 weeks old) mice using methods previously described (Beaulieu-Laroche and Hamnett, 2018; Beaulieu-Laroche et al., 2018). Mice were deeply anesthetized with isoflurane in balanced oxygen prior to immediate decapitation. 300 μm slices were prepared from the visual cortex. Slicing was performed with a vibrating blade microtome in ice-cold slicing artificial cerebrospinal fluid (aCSF) containing (in mM): sucrose 160, sodium bicarbonate 28, potassium chloride 2.5, sodium phosphate monobasic monohydrate 1.25, calcium chloride 1, magnesium chloride 7.5, glucose 7.25, HEPES 20, sodium pyruvate 3, and sodium ascorbate 3, 295-305 mOsm, saturated with 95% O₂ and 5% CO₂. Slices were then incubated for ~30 minutes at 35.5 °C in recovery aCSF containing (in mM): sodium chloride 92, sodium bicarbonate 28.5, potassium chloride 2.5, sodium phosphate monobasic monohydrate 1.2, magnesium chloride 2, magnesium chloride 4, glucose 25, HEPES 20, sodium pyruvate 3, and sodium ascorbate 5, 300-310 mOsm, saturated with 95% O₂ and 5% CO₂. Slices were kept in recovery aCSF at 18 °C until use.

Patch-clamp recording
Patch-clamp recordings were performed from the soma and apical dendrites of pyramidal neurons at 34–36 °C in recording aCSF containing (in mM): sodium chloride 120, potassium chloride 3, sodium bicarbonate 25, sodium phosphate monobasic monohydrate 1.25, calcium chloride 1.2, magnesium chloride 1.2, glucose 11, sodium pyruvate 3, and sodium ascorbate 1, 300-305 mOsm, saturated with 95% O₂ and 5% CO₂. Except for Figures S3G and S3H, recordings were restricted to GCaMP6f+ neurons with cytosolic baseline fluorescence and nuclear exclusion. Whole-cell dialysis did not impact GCaMP6f signals (Figure S3). Current-clamp recordings were performed in bridge mode with an Axopatch 200B or a Dagan BVC-700A amplifier with bridge fully balanced. Patch pipettes were made from thick-wall glass (1.5 O.D., 0.75 I.D.) or thin-wall glass (1.5 O.D., 1.1 I.D.). Pipettes had resistances ranging from 5 to 15 MΩ. The intracellular solution (in mM): potassium gluconate 134, potassium chloride 6, HEPES 10, sodium chloride 4, adenosine 5, sodium phosphate magnesium 4, guanosine 5'-triphosphate sodium 3, phosphocreatine di (tris) 14, and Alexa 594 0.05. For experiments in Figures S3G and S3H, 0.1 mM Oregon Bapta Green-1 (OGB-1) was added to the internal solution. Liquid junction potential was not corrected for. Current and voltage signals were filtered at 10 kHz and acquired at 20 kHz.

To elicit somatic action potentials, step current injection (2 ms) of constant amplitudes (1-3 nA) were employed. Dendritic spikes were elicited through suprathreshold step current injections of constant amplitude (300-1000 pA) for various durations (Figure S2).

QUANTIFICATION AND STATISTICAL ANALYSIS
All analyses were performed using custom-written MATLAB codes. In vivo imaging data was rigidly motion-corrected. Regions-of-interest (ROIs) were manually drawn on all the detectable somas. The use of Z stacks, corresponding dendritic ROIs were then drawn. Dendritic ROIs were excluded when spatially overlapping with other dendritic segments or when the path from the soma to the dendrite intersected with other dendrites. One soma-dendrite pair showed no transients for the whole recording session (correlation coefficient = 0.02) and was excluded from the analyses. Soma-dendrite pairs from different neurons (Figure 1D) correspond to all possible combinations of unpaired somas and dendrites from the same imaging sessions. Light contamination from the monitor was subtracted following robust regression of the baselined contamination fluorescence against each ROI’s raw fluorescence. To compute ΔF/F, the baseline F was estimated as the 10th percentile of the fluorescence using a 160 s rolling-window. For figure display, signals were low-pass filtered at 4 Hz with zero-phase filtering using the MATLAB function filtfit.

To detect transients, we first computed the mean and standard deviation of the raw signal without transients (< 0.2 ΔF/F) and defined threshold A as 3 standard deviations above the mean and threshold B as 1 standard deviation above the mean. We then low-pass filtered (filtfilt) the raw signal at 2 Hz and identified transients as points that were above threshold A. The start and end of each transient were defined as the points in the filtered signal that crossed threshold B. Overlapping transient were combined. To compare the paired and unpaired transient integrals, only cells with both paired and unpaired transients were used.

To detect events and decays, we computed the derivative of the filtered signal and low-pass filtered (filtfilt) it at 0.5 Hz. Rise events were identified as points in the filtered signals that were above threshold A and where the derivative was positive. The start and end of each event were defined as the points in the filtered signal that crossed threshold B or where the derivative crossed zero. Decay events were identified as points in the filtered signals that were above threshold A and where the derivative was negative. The start and end of each event were defined as the points in the filtered signal that crossed threshold B or where the derivative crossed zero. Overlapping events were combined.

For analyses in Figures 1 and 2, the whole recording sessions with the movies and the dark periods were used. The recording sessions were separated as dark versus movie and stationary versus running for Figures 4 and S4. Running speed was low-pass filtered (filtfilt) at 0.05 Hz, and 2.5 cm/s was used as the threshold to detect moving epochs. The percentage of paired events was computed only for pairs with events in both conditions (Figures S4C and S4E). The slope of rise events was computed only for pairs with at least 20 paired events in both conditions (Figures S4D and S4F).

For grating responses, we computed ΔF/F for each trial using the mean of the two seconds preceding the onset of the stimulus. We then averaged the ten trials for each condition and computed the mean responses during the two-second stimulus presentation. We determined whether the soma and dendrite of each pair were visually-responsive by computing a one-way ANOVA (p < 0.05) across

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the 32 conditions. Using nine fields of views from three mice, we found 34 out of 66 soma-dendrite pairs with at least one visually-responsive compartment. We then averaged the somatic and dendritic tuning curves to determine the preferred temporal frequency. The tuning curves and preferred orientation of corresponding somas and dendrites were then computed at that preferred temporal frequency.

Voltage signals were low-pass filtered at 5 kHz with zero-phase filtering using the MATLAB function filtfilt for some experiments. Line scan signals were low-pass filtered at 75 Hz with zero-phase filtering using the MATLAB function filtfilt. 1-10 trials were acquired per condition and averaged for both electrophysiological and optical traces. To compute $\Delta F/F$, the baseline $F$ was computed using 50-100 ms of baseline signals before eliciting spikes. Morphological and distance measurements for slice experiments were performed using ImageJ/FIJI (National Institutes of Health) on two-dimensional maximal intensity projections of 1.5–2 $\mu$m Z series collected at the end of the experiment.

Statistical analysis was performed in MATLAB. D’Agostino-Pearson tests were used to assess normality. For normal data, results are presented as mean ± SEM. For skewed data, the median and the lower and upper quartiles (Q1-Q3) are reported, and a Wilcoxon paired test or Wilcoxon rank sum test was used for statistical comparisons using MATLAB. Statistical details can be found in the figure legends and in the main text. Reported n values can be found in the figure legends and in the results. They include the number of pairs for in vivo results or recordings for ex vivo results, and the number of mice from which they were obtained.