Vision and Locomotion Shape the Interactions between Neuron Types in Mouse Visual Cortex

Highlights

- Effects of locomotion on baseline activity and visual responses are not related

- Locomotion effects on visual responses are diverse across stimuli and cell types

- *Pvalb*, but not *Sst* or *Vip*, population activity linearly tracks pyramidal activity

- A network model predicts each cell type’s visual responses from the other types

Authors

Mario Dipoppa, Adam Ranson, Michael Krumin, Marius Pachitariu, Matteo Carandini, Kenneth D. Harris

Correspondence

m.dipoppa@ucl.ac.uk

In Brief

Dipoppa et al. record visual responses of four types of neurons in mouse visual cortex, revealing a complex and diverse interaction between stimulus size and locomotion. A recurrent neural field model in which locomotion modulates synapses predicts each cell type’s responses.
Vision and Locomotion Shape the Interactions between Neuron Types in Mouse Visual Cortex

Mario Dipoppa, Adam Ranson, Michael Krumin, Marius Pachitariu, and Kenneth D. Harris

SUMMARY

Cortical computation arises from the interaction of multiple neuronal types, including pyramidal (Pyr) cells and interneurons expressing Sst, Vip, or Pvalb. To study the circuit underlying such interactions, we imaged these four types of cells in mouse primary visual cortex (V1). Our recordings in darkness were consistent with a “disinhibitory” model in which locomotion activates Vip cells, thus inhibiting Sst cells and disinhibiting Pyr cells. However, the disinhibitory model failed when visual stimuli were present: locomotion increased Sst cell responses to large stimuli and Vip cell responses to small stimuli. A recurrent network model successfully predicted each cell type’s activity from the measured activity of other types. Capturing the effects of locomotion, however, required allowing it to increase feedforward synaptic weights and modulate recurrent weights. This network model summarizes interneuron interactions and suggests that locomotion may alter cortical computation by changing effective synaptic connectivity.

INTRODUCTION

Neocortical interneurons are divided into genetically distinct types and are arranged in stereotypic recurrent circuits (Jiang et al., 2015; Kepcs and Fishell, 2014; Markram et al., 2015; Pfeffer et al., 2013; Tasic et al., 2016; Tremblay et al., 2016; Zeisel et al., 2015). The behavior of recurrent circuits can be counterintuitive and cannot always be understood using intuitive arguments (Ozeki et al., 2009; Rubin et al., 2015; Tsodyks et al., 1997). To understand how different types of interneurons influence each other and shape the activity of excitatory neurons, one must therefore constrain quantitative circuit models with measurements from different neuronal classes during diverse neural computations.

In primary visual cortex (V1), two computations that are thought to arise from interneuron interactions are size tuning and locomotor modulation. Size tuning—the suppression of activity seen when visual stimuli increase beyond a preferred size—was suggested to depend on interneurons expressing somatostatin (Sst), which integrate inputs from wide regions of cortex (Adesnik et al., 2012; Zhang et al., 2014). Modulation of firing by locomotion (Ayaz et al., 2013; Erksen et al., 2014; Fu et al., 2014; Niell and Stryker, 2010) has been suggested to arise from a disinhibitory circuit, where interneurons expressing vasoactive intestinal peptide (Vip) inhibit Sst interneurons and thereby disinhibit pyramidal (Pyr) cells.

This disinhibitory circuit rests on substantial anatomical and functional evidence, but its role in the modulation of sensory cortex is debated. The connectivity is well established: Vip interneurons principally target Sst interneurons (Acsády et al., 1996a, 1996b; Fu et al., 2014; Garcia-Junco-Clemente et al., 2017; Karnani et al., 2016a; Pfeffer et al., 2013; Pi et al., 2013), and Sst neurons, in turn, inhibit most cortical neuronal classes except other Sst cells (Jiang et al., 2015; Karnani et al., 2016b; Pfeffer et al., 2013). In barrel cortex, disinhibition could explain the effects of whisking, which increases activity in Vip cells and Pyr dendrites and decreases it in Sst cells (Gentet et al., 2012; Lee et al., 2013). In visual cortex, locomotion increases activity in Vip cells (Fu et al., 2014; Reimer et al., 2014) and putative Pyr cells (Ayaz et al., 2013; Erksen et al., 2014; Fu et al., 2014; Niell and Stryker, 2010). However, it is not clear that it decreases the activity of Sst cells (Fu et al., 2014); some studies observed mixed or even opposite effects (Pakan et al., 2016; Polack et al., 2013; Reimer et al., 2014).

Here, we used two-photon microscopy to measure responses of Sst, Vip, and Pvalb interneurons and Pyr cells in V1. We found that locomotor modulation of each cell class depends critically on the stimulus size, with modulation of sensory responses following fundamentally different rules than modulation of spontaneous activity. We then used our data to constrain a model for the circuit connecting these neuronal classes. This model provided a quantitative account for all our measurements. It also captured the complexity of the interaction between locomotion, stimulus size, and cell class, thanks to a simple reweighting of feedforward versus recurrent synapses.

RESULTS

We used two-photon imaging to measure the activity of Pyr, Pvalb, Vip, and Sst neurons in mouse V1 (Figure 1;
Figure 1. Genetic Targeting and Activity Statistics Identify Pyr, Pvalb, Vip, and Sst Cells in the Awake Cortex

(A1) Experimental setup showing the air-suspended ball surrounded by the three screens for stimulus presentation.
(A2) Green fluorescence from an Emx1-Cre mouse expressing GCaMP6m via virus injections.
(A3) Normalized fluorescent trace from a representative Pyr neuron. Blue shading above axes represents periods of locomotion (>1 cm/s).
(A4) Histogram of fluorescence values for the example neuron in (A3). The number indicates the skewness of the distribution.
(A5) Distribution of skewness values over all Pyr neurons.
(B1) Green fluorescence from a mouse expressing GCaMP6 following virus injection. Scale bars, 100 μm.
(B2) Red fluorescence from the recordings in (B1), indicating tdTomato expression in Pvalb neurons.
(B3 and B4) Same as (A3) and (A4) for a representative Pvalb neuron.
(C1 and C2) Similar analysis for Vip cells.
(D1 and D2) Same as (A3) and (A4) for all Pvalb neurons.
(C3 and C4) Normalized fluorescent traces from an unlabeled neuron recorded simultaneously with the Vip example in (D3) and (D4).
(C5) Distribution of skewness values over all unlabeled neurons. Unlabeled cells above a skewness threshold of 2.7 (dashed vertical line) are classified as putative Pyr (E5).

Figure S1). Mice were head fixed and free to run on an air-suspended ball (Niell and Stryker, 2010) while viewing a grating in a circular window of variable diameter (Figure 1A1). The raw fluorescence traces were corrected for out-of-focus fluorescence (neuropil correction; Figure S2; Chen et al., 2013; Peron et al., 2015).
Genetic Targeting and Activity Statistics Identify Pyr, Pvalb, Vip, and Sst Cells in the Awake Cortex

To identify neurons belonging to a specific class, we used one of two genetic approaches (Figure 1, columns 1 and 2). In the first approach, we expressed GCaMP6m virally in all neurons in mice in which a class of interneurons was labeled with tdTomato (Figures 1B–1D, columns 1 and 2). This approach allowed us to record the activity of identified interneurons in the labeled class and of many unlabeled neurons, which will comprise mainly, but not exclusively, Pyr cells. In the second approach, we expressed the calcium indicator exclusively in a chosen cell class either by injecting a Cre-dependent GCaMP6m virus into an appropriate transgenic driver line (Figure 1A2, Figure S2B2) or via a triple-transgenic line that expressed GCaMP6s specifically in superficial layer Pyr cells.

Interneurons of all three classes fired much more frequently than pyramidal neurons (Figure 1, columns 3 and 4). As expected from the sparse firing of superficial-layer pyramidal cells (Niell and Stryker, 2010), identified pyramidal cells showed rare isolated calcium events (Figure 1A3), yielding a distribution of fluorescence that was highly skewed (Figure 1A4). By contrast, identified Pvalb, Vip, and Sst interneurons showed frequent calcium events (Figures 1B–1D, column 3), yielding distributions of fluorescence with little skewness (Figures 1B–1D, column 4).

These differences in skewness allowed us to use this measure to identify putative Pyr cells among the concurrently measured unlabeled neurons (Figure 1E). Similar to identified Pyr cells, most unlabeled neurons showed sparse activity and high skewness (e.g., Figures 1E3 and 1E4). To identify putative Pyr cells, we thus set a threshold on skewness. Its precise value made little difference to our results; we chose a conservative value of 2.7, as it provided a small false-positive rate (24/1,511 Pvalb neurons, 29/1,385 Vip cells, and 91/537 Sst cells exceeded this threshold; Figures 1B–1D, column 5) while correctly classifying most identified pyramidal cells (2,598/4,949; Figure 1A4), while unlabeled neurons exceeding this threshold were likely to be pyramidal, cells below the threshold could be of any type and were therefore excluded from further analysis (Figure 1E1). This procedure worked well for all methods of GCaMP expression (Figure S3).

The Effects of Locomotion on Baseline Activity Depend on Cell Type and Depth

We next asked how locomotion affected baseline (spontaneous) activity, measured when the screens were uniform gray (Figure 2). These measurements showed strong effects of locomotion on baseline activity of interneurons and unexpected dependences on cortical depth. They also revealed ways in which apparently conflicting reports on Sst cells could be reconciled.

Consistent with previous results (Fu et al., 2014; Niell and Stryker, 2010; Polack et al., 2013; Saleem et al., 2013), the effects of locomotion on the baseline activity of Pyr cells were weak and diverse (Figure 2A). The sparse baseline activity of a typical Pyr cell changed only weakly with running speed (Figure 2A3). Across Pyr cells, the average correlation between baseline activity and running speed was close to zero ($\rho_{gray} = 0.03 \pm 0.01$ SE, $n = 7,553$ identified Pyr cells; Figure 2A3). Nevertheless, 35% of identified Pyr cells showed a significant positive or negative correlation with speed ($p < 0.05$, shuffle test), significantly more than the 5% expected by chance ($p < 10^{-16}$, Fisher’s combined probability test). Similar results were seen in the putative Pyr neurons identified by skewness (Figures S4A and S4B).

The effects of locomotion on the baseline activity of Pvalb interneurons were stronger and more varied and depended on cortical depth (Figure 2B). For example, in two Pvalb cells imaged simultaneously, activity decreased with running speed ($\rho = -0.54$, $p < 0.01$, shuffle test) in the more superficial cell and increased ($\rho = 0.54$, $p < 0.01$, shuffle test) in the deeper cell (darker and lighter traces in Figure 2B1). These results were typical of the population, where correlations were strong and depended significantly on depth (robust regression, $p < 10^{-22}$, $n = 1,730$; Figure 2B3), with high consistency across experiments ($p < 0.018$, t test; Figure S5). Among Pvalb cells in superficial L2/3 (depth < 300 µm, $n = 833$), correlation with speed was significantly negative in 36% of the cells and significantly positive in only 24% of the cells ($p < 0.05$, shuffle test). The situation was reversed in deeper L2/3, with correlations significantly positive in 47% of cells and negative in only 18% of cells ($p < 0.05$, shuffle test). Therefore, when pooling across depth, a wide variety of effects was seen (Figure 2B3), echoing the wide and bimodal range of correlations observed previously (Fu et al., 2014).

Even larger effects were seen in Vip cells, where correlations were overwhelmingly positive (Figure 2C). Consistent with previous results (Fu et al., 2014; Pakan et al., 2016), the typical Vip cell increased baseline activity markedly with locomotion (Figure 2C1), and the overall population showed almost exclusively positive correlations with running speed, with a mean correlation of $\rho_{gray} = 0.27 \pm 0.03$ (SE, $n = 1,393$). The correlation increased significantly with cortical depth (robust regression, $p < 10^{-10}$), an effect that was robust across experiments ($p < 0.01$, t test; Figure S5).

Perhaps, surprisingly, locomotion also generally increased the baseline activity of Sst cells (Figure 2D). The typical Sst cell increased its baseline activity markedly with locomotion (Figure 2D1), and across the population the correlation of baseline activity with running speed was on average positive ($\rho_{gray} = 0.18 \pm 0.02$, SE, $n = 636$; Figure 2D3) regardless of depth (robust regression, $p = 0.39$; Figure 2D3). This did not reflect altered rate during running onset, as the correlation persisted after removing transition periods between locomotion and stationary periods from the analysis (Figure S6). These effects of locomotion on the baseline activity of Sst cells confirm some previous results (Pakan et al., 2016; Polack et al., 2013), but they appear to disagree with other measurements (Fu et al., 2014).

To confirm these observations in Sst cells, we first ensured that they were not due to background fluorescence that might originate from other cell classes. We repeated the measurements in mice expressing the calcium indicator only in Sst cells (Sst-ires-Cre mice injected when adult with a Cre-dependent GCaMP6m virus; Figure S2B3). These experiments confirmed our results: the average correlation of baseline activity with running speed was positive (Figures S2C–S2E) in all locations where GCaMP6m had strong expression, be it cell bodies or neuropil.
We next asked whether the disagreement on the effects of locomotion on the baseline activity of Sst cells could be due to differences in visual conditions (Pakan et al., 2016). Fu et al. (2014) made their measurements in darkness, whereas we (Figure 2D) and Polack et al. (2013) had the mouse face a gray screen. We thus turned off the screen and found that the effects of locomotion on baseline activity of Sst cells were now overall negative ($r_{\text{dark}} = -0.07 \pm 0.02$, SE; across experiments: $p = 0.019$, t test; Figures S7A, S7B, S7D, and S7E). The same cell could show different modulation by locomotion depending on screen illumination (Figure S7B): for example, of the cells showing significant modulation in both conditions, 26% showed $r_{\text{dark}} < 0$ and $r_{\text{gray}} > 0$. Not all cells, however, showed this diversity. On average, in fact, Sst cells showed a positive correlation between $r_{\text{gray}}$ and $r_{\text{dark}}$ (Pearson correlation 0.34; $p < 10^{-5}$).

These measurements, therefore, reconcile the apparent divergence of previous results (Fu et al., 2014; Pakan et al., 2016; Polack et al., 2013): the effect of locomotion on baseline activity of Sst cells is overall positive when mice view a gray screen and mildly negative when mice are in darkness.

Figure 2. The Effects of Locomotion on Baseline Activity Depend on Cell Type and Depth
(A1) Fluorescence of representative L2/3 pyramidal neuron (top) and simultaneous running speed trace (bottom).
(A2) Correlation coefficient of recorded pyramidal cells with running speed plotted versus cell depth. Circles represent cells with significant correlations at $p < 0.05$ (shuffle test); dots represent cells with insignificant correlations. For clearer visualization, we plotted only a random subsample of 800 of genetically identified Pyr cells. Dashed line represents fitted dependence of correlation versus depth. Black circle indicates example cell shown in (A1).
(A3) Histogram of correlation coefficients of all pyramidal cells. Solid bars indicate significant correlations at $p < 0.05$ (shuffle test). Values left and right of the histogram represent the percentage of cells with a negative or positive correlation, respectively.
(B) Similar analysis for Pvalb neurons. The two traces in (B1) (top) show fluorescence traces of representative Pvalb cells of upper and lower L2/3 (blue and cyan, respectively). The average correlation with speed was slightly negative ($r_{\text{gray}} = -0.05 \pm 0.03$, SE) among Pvalb cells in superficial L2/3 (depth < 300 \(\mu\)m, $n = 843$) and weakly positive ($r_{\text{gray}} = 0.11 \pm 0.04$, SE; $n = 831$) in deeper L2/3.
(C) Similar analysis for Vip cells.
(D) Similar analysis for Sst cells.
This observation is specific to Sst cells. In the other cell types, the effects of locomotion on baseline activity were similar whether the screen was gray or dark. In agreement with results obtained in darkness (Fu et al., 2014), when the monitors were switched off, locomotion continued to have little overall effect on baseline activity of Pyr cells ($\rho_{\text{dark}} = 0.00 \pm 0.01$, SE; Figures S7A1 and S7B1; Pearson correlation between $p_{\text{gray}}$ and $p_{\text{dark}}$ 0.33; $p < 10^{-7}$). Similar observations were made in Pvalb cells ($\rho_{\text{dark}} = − 0.14 \pm 0.01$, SE; Figures S7A2 and S7B2; Pearson correlation between $p_{\text{gray}}$ and $p_{\text{dark}}$ 0.49, $p < 10^{-22}$) and Vip cells ($\rho_{\text{dark}} = 0.30 \pm 0.06$, SE; Figures S7A3 and S7B3; Person correlation between $p_{\text{gray}}$ and $p_{\text{dark}}$ 0.54; $p < 10^{-48}$).

**Locomotion Modulates Sensory Responses Differently from Baseline Activity**

Having explored the effects of locomotion on baseline activity, we turned to its effect on visual responses (Figure 3). We presented drifting gratings of various diameters and focused first on the neurons whose receptive field centers were located within 10° of the stimulus center. We computed a modulation index, $M_B$, to measure the effect of locomotion on each cell’s visual responses and a corresponding modulation index, $M_0$, for baseline activity (see STAR Methods).

In contrast to its effects on baseline activity, locomotion tended to increase visual responses in all cell classes (Figure 3, columns 1 and 3). For example, a typical Pyr cell would markedly increase its visual responses while the animal ran (Figure 3A1, $M_0 = 0.31$), and this increase was common among Pyr cells ($p < 10^{-22}$, t test; Figure 3A3). Similar effects can be seen in the visual responses of the example Pvalb cells, both superficial and deep (Figure 3B1, $M_0 = 0.77$ and $M_0 = 0.34$, respectively), and was common across the Pvalb population ($p < 10^{-33}$, t test; Figure 3B3). For Vip cells, the effect of locomotion on visual responses was more mixed, with several cells having a weak modulation (e.g., Figure 3C1, $M_0 = 0.15$), but was still overall positive ($p < 10^{-3}$, t test; Figure 3C3). Finally, locomotion typically increased visual responses in Sst cells (Figure 3D1, $M_0 = 0.47$), an effect that was significant across the population ($p < 10^{-18}$, t test; Figure 3D3).

These effects of locomotion on visual responses were very different from those on baseline activity. Indeed, our previous analysis (Figure 2) had shown that locomotion decreased baseline activity in approximately half of Pyr and Pvalb neurons and in a significant fraction of Sst cells.

Indeed, we typically saw no correlation between the locomotor modulation of an individual cell’s baseline and its evoked activity (Figure 3, column 4). The correlation between the effects of locomotion on baseline activity $M_B$ and on visual responses $M_0$ was weak in Pyr cells ($\rho = −0.07$, $p = 0.041$; Figure 3A4) and not significant in the remaining cell types (Pvalb cells: $\rho = −0.12$, $p = 0.11$; Figure 3B4; Vip cells: $\rho = −0.10$, $p = 0.18$; Figure 3C4; Sst cells, $\rho = −0.05$, $p = 0.65$; Figure 3D4).

Finally, while in some cell classes the effect of locomotion on visual responses depended weakly on depth, the direction of this modulation often differed from that seen during baseline activity (Figure 3, column 2). The effect of locomotion did not vary significantly with cortical depth in Pyr cells ($p = 0.28$, robust regression; Figure 3A4), but it decreased with depth in Pvalb cells ($p < 10^{-3}$, robust regression; Figure 3B4), increased with depth in Vip cells ($p < 10^{-7}$, robust regression; Figure 3C4), and decreased with depth in Sst cells ($p < 10^{-2}$, robust regression; Figure 3D4).

**Locomotion Increases Sst Cell Responses to Large Stimuli and Vip Cell Responses to Small Stimuli**

We next asked how locomotion modulated responses to stimuli of different sizes. We focused on visually responsive neurons (significant effect of stimulus size, $p < 0.05$, one-way ANOVA) with receptive fields centered within 10° of the stimulus center. To discount possible effects of eye movements (whose occurrence might change during locomotion), we considered only trials in which pupil position was within 5° of its average. Loosening this criterion would make us underestimate the selectivity of neurons for stimulus size (Figure S8).

Pyr cells were selective for small stimuli and typically exhibited mild but diverse locomotor modulation (Figure 4A1). A typical Pyr neuron responded substantially more to a stimulus of diameter 5° than to a stimulus of diameter 60° regardless of locomotion (Figure 4A1), showing clear selectivity for smaller stimuli (Figure 4A2). Similar effects were seen in the overall population of identified Pyr cells ($n = 1,250$; Figure 4A3), in which cells preferring large (Figure S9A) or small (Figure S9B) stimuli were rare. On average, locomotion slightly increased responses to both a 5° diameter stimulus ($p < 10^{-16}$, paired t test across cells, $p < 0.01$, paired t test across experiments) and a 60° diameter stimulus ($p < 10^{-10}$ and $p = 0.02$). However, this effect was diverse among cells, with locomotion significantly increasing or decreasing responses in 17% and 3% of Pyr cells, respectively ($p < 0.05$; two-way ANOVA, main effect of locomotion over stimuli of diameter 5° and 60°; Figure 4A4). Many cells (17%) showed a significant interaction of locomotion and stimulus size ($p < 0.05$; two-way ANOVA over stimuli of diameter 5° and 60°; Figure 4A4; Figures S10A and S10B, column 1). In these cells, locomotion changed the relative response to large and small stimuli, as seen previously in deeper layers (Ayaz et al., 2013). Similar results were found in putative pyramidal cells identified by the sparseness of their calcium traces (Figures S4C and S4D).

Pvalb interneurons were similarly selective for smaller stimuli but showed a stronger and overwhelmingly positive effect of locomotion (Figure 4B1). A typical Pvalb interneuron responded strongly to small stimuli and more weakly to larger stimuli, and its responses markedly increased during locomotion (Figures 4B1 and 4B3). These effects were highly consistent across Pvalb interneurons ($n = 277$; Figure 4B4), with locomotion increasing firing rate in practically all cells (Figure 4B4). This increase was seen in responses to both large stimuli ($p < 10^{-11}$, paired t test across cells, $p < 0.01$, paired t test across experiments) and small stimuli ($p < 10^{-13}$ and $p = 0.02$), with no significant interaction between stimulus size and locomotion ($p = 0.23$, two-way ANOVA over stimuli of diameter 5° and 60°; Figures S10A and S10B, column 2).

The responses of Vip interneurons ($n = 233$) were selective for stimulus size and increased with locomotion, but this increase was generally restricted to responses to small stimuli (Figure 4C). A typical Vip interneuron responded most strongly to small stimuli during locomotion (Figures 4C1 and 4C2). Similar results were
seen across the population: Vip interneurons showed clear size tuning, and locomotion increased their responses to 5° stimuli ($p < 10^{-11}$, paired t test across cells, $p = 0.059$, paired t test across experiments), but not to 60° stimuli ($p = 0.95$ and $p = 0.77$; Figure 4C4), with a significant interaction of size and locomotion ($p < 10^{-10}$, two-way ANOVA over stimulus diameters 5° and 60°; Figure 4C4; Figures S10A and S10B, column 3).

Sst interneurons tended to prefer large stimuli, and their responses increased with locomotion (Figure 4D). As observed by Adesnik et al. (2012), a typical Sst interneuron responded better to 60° than 5° stimuli, especially while the animal was running (Figures 4D1 and 4D2). Similar results were seen across the population ($n = 191$, Figure 4D3), with overall activity peaking at diameter $\sim 15°$ during stationary conditions and $\sim 25°$ during

---

**Figure 3. Locomotion Modulates Sensory Responses Differently from Baseline Activity**

(A1) Visual responses of representative L2/3 pyramidal neuron (top) and simultaneous running speed trace (bottom) for the same cell as in Figure 2A1. In the top plot, dots represent the size of visual responses relative to the mean response to the presented stimulus, while the continuous line represents a smoothed interpolation of these points.

(A2) Average increased visual responses of recorded pyramidal cells by locomotion (modulation of visual responses by locomotion) plotted versus cell depth. Continuous line represents fitted dependence of correlation versus depth. Black circle indicates example cell shown in (A1).

(A3) Histogram of modulation of visual responses by locomotion for all pyramidal cells. Solid bars indicate significant correlations at $p < 0.05$ (t test). Values left and right of the histogram represent the percentage of cells with a negative or positive modulation, respectively.

(A4) Modulation of spontaneous activity by locomotion as a function of modulation of visual responses by locomotion. Each point corresponds to a different neuron. Black circle indicates example cell shown in (A1) and Figure 2A1.

(B) Similar analysis for Pvalb neurons. The two traces in (B1) (top) show fluorescence traces of the same representative Pvalb cells of upper and lower L2/3 (blue and cyan, respectively) as in Figure 2B1.

(C) Similar analysis for Vip cells with the example cell in (C1) being the same as in Figure 2C1.

(D) Similar analysis for Sst cells with the example cell in (D1) being the same as in Figure 2D1.
locomotion. Sst cells showed a significant interaction between stimulus size and locomotion (p < 10^{-7}, two-way ANOVA over diameters 5° and 60°), consistently across experiments (p < 0.01, t test; Figure S10A) and mice (p < 0.01, t test; Figure S10B). While locomotion did not significantly affect the responses to 5° stimuli (p = 0.051), paired t test across cells, p = 0.62, paired t test across experiments, Figure 4D), it strongly increased the responses to 60° stimuli (p < 10^{-8} and p = 0.02).

Some Sst cells, however, did show size tuning (Figure 4D; Figures S9C and S9D). This observation is consistent with observations in anesthetized mice (Pecka et al., 2014) but differs from those of Adesnik et al. (2012) in awake mice. We reasoned that this may reflect high sensitivity of these cells to stimulus centering and thus studied how size tuning varies with distance between receptive center and stimulus center. Size tuning emerged when stimulus distance was small (Figure 11A); when the two were distant, cells of all classes preferred larger stimulus sizes (Figure S11B). For Sst cells, in particular, size tuning appeared when stimuli were within a radius of 20° from the receptive field center (Figure S11, column 4).

For all cell types, we saw similar interactions of size tuning and locomotion when cells were recorded in the binocular and monocular regions of visual cortex (Figure S12). Furthermore, the results did not change if we deconvolved the calcium traces to estimate spike rates (Figure S13).

In summary, the effects of running on cortical cell classes depend on which stimuli are present. With small stimuli, locomotion boosts responses in Vip cells while having little effect on Sst cells, but with large stimuli, it has the opposite effect, doing little to Vip cells but boosting Sst cells.

The Relationship of Interneurons with Pyr Cells Is Linear in Pvalb Cells and Nonlinear in Vip and Sst Cells

We next examined the correlations between interneuron types and Pyr cells (Figure 5). According to the “balanced inhibition” theory, inhibitory activity should closely track the mean firing of the Pyr population, thereby stabilizing network function (Shu et al., 2003; van Vreeswijk and Sompolinsky, 1998; Wehr and Zador, 2003). To measure Pyr-interneuron correlation, we relied on our ability to simultaneously record identified interneurons and putative Pyr neurons (unlabeled cells with skewness > 2.7, Figure 1).

Consistent with the view that Pvalb interneurons track the activity of Pyr cells (Cruikshank et al., 2007; Isaacson and Scanziani, 2011; Okun and Lampl, 2008; Ozeki et al., 2009; Renart et al., 2010), we found strong positive correlations between the Pvalb and putative Pyr populations (Figure 5A). In each experiment, we compared the summed activity of the Pvalb population to that of the simultaneously recorded putative Pyr cells (using only cells with receptive fields within 10° of the stimulus center).
Spontaneous activity

Stationary Locomotion

Stimulus responses

Stationary Locomotion

Neuron 98, 1–14, May 2, 2018
In a typical experiment, spontaneous correlations ($r_0$) were strongly positive whether the mouse was stationary ($r_0 = 0.70$; Figure 5A) or running ($r_0 = 0.68$; Figure 5A). Similar results were seen across experiments ($r_0 = 0.60 \pm 0.04$, SE, during rest and 0.49 ± 0.03 during locomotion). Population signal correlations—i.e., the relationship of the mean summed responses of the Pvalb and Pyr cells across stimuli—also showed strong, positive correlations and a linear relationship in both stationary ($r_0 = 0.95$ for the example in Figure 5A, 0.86 ± 0.04 across all experiments) and locomotion ($r_0 = 0.97$ for the example in Figure 5A, 0.91 ± 0.02 across all experiments) conditions. Noise correlations (i.e., the relationship between trial-to-trial variability in summed activity of the Pvalb and Pyr populations) were also large (Figures S14A and S14D). These results were not biased by the exclusion of low-skewed Pyr cells: when we lowered the threshold (therefore including more Pyr cells but also some unlabeled inhibitory neurons), the correlations of putative Pyr cells with Pvalb cells remained high (Figures S14E–S14G).

Vip cells showed markedly different behavior (Figure 5B). Their correlations with the putative Pyr population differed from those of Pvalb cells in two respects. First, while spontaneous and noise ($r_0$) correlations tended to be positive ($r_0 = 0.43 \pm 0.03$, SE, during stationarity and 0.42 ± 0.06 during locomotion; $r_0 = 0.33 \pm 0.07$, SE, during stationarity and 0.37 ± 0.02 during locomotion), they were weaker than those of Pvalb cells, at least during stationarity (Figure 5D1; Figures S14B and S14D). Second, the relationship between the population mean responses of Vip and putative Pyr cells was nonlinear (Figures 5B2 and 5B3). This nonlinearity reflects the different size tuning of Vip and Pyr cells, with Vip responses peaking at smaller stimulus sizes than Pyr responses (compare Figures 4C2 and 4A2). It further suggests that, unlike for Pvalb cells, the sensory tuning of Vip cells during locomotion cannot be explained by a simple tracking of excitatory activity.

Sst interneurons showed yet a different sort of behavior, which depended on locomotion (Figure 5C). The correlation of the Sst and Pyr populations was positive in stationary conditions ($r_0 = 0.25$ for the example in Figure 5C1, 0.23 ± 0.10 SE across experiments) but weak during locomotion ($r_0 = -0.01$ for the example in Figure 5C2, 0.13 ± 0.13, SE, across experiments). Noise correlations were also positive (Figures S14C and S14D; $r_n = 0.39 \pm 0.11$, SE, stationary; $r_n = 0.30 \pm 0.04$, SE, locomotion). Signal correlations showed a nonlinear character, but this differed to that of Vip cells (Figures 5C3 and 5C4).

To quantify the linearity of the relationship between the mean responses of interneurons and of putative Pyr cells on a session-by-session basis, we parameterized their relationship using two angles, $\theta_1$ and $\theta_2$, that captured the signal correlation along the increasing and decreasing slopes of the size-tuning curve (Figure 5E; Figures S14H and S14I). This analysis confirmed the reliable linearity of Pyr–Vip correlations (indicated by $\theta_2$ close to 0) and also revealed a nonlinear relationship for Sst and Vip cells, indicated by a significant difference in $\theta_1$ (stationary: $p = 0.026$, locomotion: $p < 0.01$, Watson’s $U^2$ permutation test, $n = 1,000$ permutations) and $\theta_2$ between (stationary: $p = 0.039$, locomotion: $p = 0.036$, Watson’s $U^2$ permutation test, $n = 1,000$ permutations). Thus, while balanced inhibition is accurate for Pvalb neurons, the activity of Sst and Vip cells diverges substantially from balance with the Pyr population.

**A Recurrent Network Model Accurately Predicts the Visual Responses of Each Neuron Type**

Our data indicate that a disinhibitory circuit in which Sst neurons suppress Pyr responses to large stimuli cannot account for size tuning, as the Sst neurons themselves show size tuning; furthermore, the complex and size-dependent effects of locomotion on Vip and Sst cells do not match what one might intuitively expect from simple disinhibition.

We therefore asked whether a more complete recurrent network model could reproduce the different classes’ size tuning. We fit the model with a novel approach: we estimated the synaptic input parameters for each class of neurons to be those optimally predicting that class’s average population sensory responses, with the average population activity of all other cell classes clamped to their measured values.

We modeled the activity of each cell class by a “neural field”: a number that varied across the retinotopic cortical surface, representing the mean activity of all cells of that class at that location. In the model, the sensory response of class $\alpha$ was a function $f_\theta^\alpha(s, r)$ of stimulus size $s$ and retinotopic position $r$ (relative to stimulus center); dependence on $r$ was assumed to be circularly symmetric. The superscript $\nu$ indicates locomotion.

---

**Figure 5. The Relationship of Interneurons with Pyr Cells Is Linear in Pvalb Cells and Nonlinear in Vip and Sst Cells**

- $(A_1$ and $A_2$) Summed activity of Pvalb population versus Pyr population in the gray screen condition during stationary periods $(A_1)$ and during locomotion $(A_2)$. Each circle represents the simultaneous normalized value of the excitatory and the inhibitory populations at one time point. Dashed line indicates linear regression estimate of signal correlation.
- $(A_3$ and $A_4$) Average stimulus response of Pvalb population versus average response of Pyr population in a typical experiment during stationary periods $(A_3)$ and during locomotion $(A_4)$. Each point represents a response to a stimulus, with larger circles representing larger stimuli. Dashed line represents nonlinear interpolation of the Pvalb and Pyr size tuning obtained from their size-tuning curves.
- $(B$ and $C)$ Same as $(A)$, but for Vip $(B)$ and Sst $(C)$ interneurons. Note the nonlinear signal correlations.
- $(D_1$ and $D_2$) Summary plots of spontaneous correlations during stationary periods $(D_1)$ and during locomotion $(D_2)$. Error bars correspond to standard error.
- $(D_3$ and $D_4$) Linear signal correlation $r_n$ between the Pyr population and the three classes of interneurons averaged across all experiments during stationary periods $(D_3)$ and during locomotion $(D_4)$. Error bars correspond to standard error.
- $(E_1$) Same plot represented in $(C_4$) showing the characteristic angles used to illustrate the nonlinear relationship between each interneuron class and Pyr mean visual responses. Circles with black outline indicate the minimum size (diameter 5°, black filled), maximum sizes (diameter 60°, red filled), and the Pyr cells’ preferred size (white filled); $\theta_1$, is the angle relative to the horizontal axis of the line joining the response to stimuli of diameter 5° and the preferred stimulus for Pyr neurons; $\theta_2$ is the angle between the population line and the line joining the response at the Pyr cells’ preferred size to the response to stimuli of diameter 60°.
- $(E_2$ and $E_3$) Angle $\theta_1$ versus $\theta_2$ as defined in $(E_1$) for each experiment during stationary periods $(E_2)$ and during locomotion $(E_3)$. In $E_3$, arrows indicate examples in $(A_3$–$C_3)$; in $E_3$, arrows indicate examples in $(A_3$–$C_3)$. 

Neuron 98, 1–14, May 2, 2018 9
condition ($v = 0$: rest; $v = 1$: locomotion). We estimated these functions from the data using a smoothing method (Figures S11C and S11D). The connection strength to a cell of type $\alpha_1$ at location $r_1$ from a cell of type $\alpha_2$ at location $r_2$ was a two-dimensional Gaussian function, $G_{\alpha_{1}\alpha_{2}}(r_1 - r_2)$. We fit the strength and spatial spread of these connections by exhaustive search, minimizing the squared error between the predicted and actual rates. Synaptic integration followed a threshold-linear function, and we chose divisive or subtractive inhibition for each inhibitory synapse type to minimize errors. The firing of dLGN inputs $h^{V}(s, r)$ was modeled using a ratio of Gaussians (Ayaz et al., 2013), with parameters (for both rest and locomotion conditions) fit to the data of Erisken et al. (2014).

The model accurately predicted the size tuning of each class for both centered and off-centered stimuli (Figure 6), but this success was predicated on certain conditions. Specifically, to predict the strong response of Sst cells to large, off-center stimuli (Figures 6D$_3$ and 6D$_5$), the model required external excitatory input to these cells (e.g., from thalamus or from other cortical layers), because large stimuli elicited little response in Pyr cells (Figures 6A$_2$, 6A$_3$). A good fit was only obtained if this feedforward input to Sst cells had broad size tuning, as would be seen in thalamic neurons or perhaps in excitatory neurons of other cortical layers (Figure 6D$_1$). Moreover, to obtain similar tuning of Pvalb and Pyr cells (Figures 6A and 6B), the model required these classes to have similar inhibition from Sst cells. The model required Pyr neurons to lack Vip input. Finally, our parameter search only gave good results with divisive inhibition from Sst to Vip cells (Figure 6C$_1$): subtractive inhibition could not produce the observed sharp size tuning favoring small stimuli (Figures 6C$_2$, 6C$_3$).

We next asked what modifications of the model parameters could explain the effects of locomotion on the sensory responses of each cell type. Modeling the locomotor modulation of size tuning required that locomotion change synaptic strengths: for example, if we kept Pyr input synapses fixed between the stationary and locomotion conditions, we obtained a poor prediction of their size-tuning modulation (Figure S15A). To capture the effects of locomotion on each cell type, we searched for all possible ways that locomotion could modulate the thalamocortical and recurrent synaptic weights of each class (Figures S15A and S15B).

For Pyr and Pvalb cells, we could prove analytically that the fit quality depended only on the strength of their “effective connections” (gray connections in Figure 6; see STAR Methods), which take into account the amplification caused by recurrent excitation and Pvalb inhibition (Douglas et al., 1995). Thus, while the model fit identified unique values of the effective connections, these values could, in turn, be achieved through multiple possible strengths of the synaptic parameters (Prinz et al., 2004).

Capturing the locomotor modulation of Pyr tuning required an increase in effective connection from external inputs to Pyr cells and a decrease in effective connection of Sst to Pyr cells. Together, this produced the observed strong increase in responses of centered cells to medium-sized stimuli, along with a milder increase in response to larger stimuli (Figure 6A$_2$ versus 6A$_4$). However, this change in effective connection strength did not require a weakening of the physical Sst→Pyr connections: the same change in effective connection could also be achieved by an increase in external excitation together with a decrease in recurrent excitation. Intriguingly, these two effects are precisely those observed in vitro during cholinergic modulation of cortical synapses (Gol et al., 1997).

Consistent with the close correlations of Pvalb and Pyr cells, modeling the observed effects of locomotion on Pvalb size tuning required similar modulations to those required for Pyr cells: a decrease in effective inhibition from Sst cells. Again, however, this did not necessarily require a weakening of Sst→Pvalb synapses, as the same effect could be obtained via strengthening of Pvalb→Pvalb connections and of the external input. Producing the observed effects of locomotion on Vip cell tuning required no further changes in effective connection: locomotion only increased the responses of centered Vip cells to small stimuli (Figure 6C$_2$ versus 6C$_1$), and this increase could be readily provided by increased activity of local Pyr cells. Finally, correct modulation of Sst firing required boosting the external excitatory inputs responsible for their responses to large stimuli (Figure 6D$_4$) but did not require any change to their inhibitory inputs.

In summary, we were able to capture the effects of locomotion on all cell types through a reweighting of feedforward and recurrent connections: an increase in external excitatory input to all cell types, a decrease in recurrent excitation between Pyr cells, and an increase in recurrent inhibition between Pvalb cells.

**DISCUSSION**

We have shown that locomotion does not simply increase or decrease the activity of a particular cell class: its effects depend on the precise sensory conditions and even on cortical depth. The effect of locomotion on sensory responses was, on average, an increase in all cell classes, but the increase varied with cell type and stimulus, being largest in Sst responses to large stimuli and in Vip responses to small stimuli. The effects of locomotion on baseline neural activity (as assessed by gray screen viewing) were more complex: locomotion increased activity in most Sst and Vip neurons and had diverse effects on Pvalb and Pyr cells, suppressing most Pvalb cells in superficial L2/3 and increasing activity in deeper Pvalb cells.

Although studies on locomotor modulation of visual cortical responses have given apparently contradictory findings, our results are, in fact, consistent with most of their observations once differences in experimental methods are accounted for. While the fraction of cells showing locomotor modulation in our data might appear smaller than in previous electrophysiological recordings (Niell and Stryker, 2010), this may reflect the increased ability of two-photon microscopy to detect weakly responsive cells. Additionally, two studies on the spontaneous activity of Sst cells reported opposite effects of locomotion: increased (Polack et al., 2013) and decreased (Fu et al., 2014) activity. When we replicated their experimental conditions (gray screen for the first study and complete darkness for the second), we reproduced both observations in a common set of neurons. These results reconcile the apparent contradiction between these studies (see also Pakan et al., 2016). Our results also reinforce the importance of correcting for out-of-focus fluorescence in two-photon calcium imaging (neuronal correction). Indeed,
without correcting for this confound, one would observe an artificial negative correlation of fluorescence with running speed, particularly in image regions with weak GCaMP expression (Figure S2).

Our results are consistent with the “balanced inhibition” model, but only for Pvalb interneurons. In this model, the activity of inhibitory cells tracks that of excitatory cells, thereby stabilizing network activity over a wide dynamic range of inputs (Shu...
et al., 2003; van Vreeswijk and Sompolinsky, 1998; Wehr and Zador, 2003). We found that the mean activity of Pvalb, but not Vip or Sst, interneurons was in all cases linearly correlated with the summed local excitatory population despite the heterogeneity of individual Pyr cell responses. These data are therefore consistent with a primary role for Pvalb cells of stabilizing the activity of the local circuit via tracking summed excitatory firing (Cruikshank et al., 2007; Ozeki et al., 2009; Renart et al., 2010) rather than directly sculpting visual preferences. Our model was able to reproduce the similar tuning of Pvalb and Pyr cells only if they received similar inputs from other inhibitory classes: specifically, consistent with in vitro observations (Pfeffer et al., 2013), it required that both Pyr and Pvalb cells received inhibition from Sst, but not Vip, cells (data not shown). The linear relationship of Pvalb and Pyr cells allowed us to greatly simplify our network models by “integrating out” the activity of Pvalb cells and recurrent excitatory connections so that Pyr activity could be modeled using an “effective connection” from thalamic and Sst cells only.

Our results suggest that previous theories, which operated at the level of intuitive arguments, are not sufficient to explain the role of interneuron classes in V1 function. One such theory holds that size tuning is mediated by Sst neurons, which were reported to show negligible size tuning (Adesnik et al., 2012). Our experiments replicated this finding only when stimuli were poorly centered on the receptive field. Inhibition of Pyr cells by local Sst neurons therefore cannot be sufficient to explain size tuning, as for stimuli of diameter > 30°, the responses of Pyr cells continue to decrease with stimulus size, while Sst firing does not increase. Furthermore, although off-center Sst neurons respond to large stimuli, neither centered nor off-centered Pyr cells respond strongly enough to drive them. Our network model was able to replicate our experimental results but only under two conditions that we propose as experimental predictions: that Pyr cells integrate inhibitory input from spatially dispersed Sst neurons and that Sst cells receive a feedforward sensory input, i.e., an excitatory input conveying visual input other than from local Pyr cells. Whether Sst cells receive direct thalamic inputs is controversial (Cruikshank et al., 2010; Lee et al., 2013; Tan et al., 2008). Even without direct thalamic afferents, such an input could be conveyed to superficial Sst cells via other cortical layers. Interestingly, the optimal model parameters required that the external inputs that Sst cells receive be spatially diffuse as would be expected if this input had experienced an additional round of divergence through Pyr cells of other layers.

A second theory is the “disinhibition” hypothesis: that during locomotion, increased Vip activity would inhibit Sst cells, thus increasing Pyr activity (Fu et al., 2014). Although there is ample evidence for inhibitory synaptic connections between Vip and Sst cells, synaptic inhibition does not necessarily imply anticorrelation: for example, in “inhibitory stabilized network” models (Litwin-Kumar et al., 2016; Ozeki et al., 2009; Rubin et al., 2015; Tsodyks et al., 1997), hyperpolarization of inhibitory cells could cause a paradoxical increase in total inhibitory activity. Our results pointed to a more complex picture than either scenario, with the effects of locomotion depending on the precise visual stimulation conditions. Consistent with previous results (Fu et al., 2014; Pakan et al., 2016), we found that, in darkness, locomotion weakly decreased activity in Sst cells. However, locomotor modulation of stimulus responses was uncorrelated with modulation of spontaneous firing. Furthermore, the effects of locomotion on sensory responses depended on stimulus size, boosting Vip cells in the presence of small stimuli and Sst cells in the presence of large stimuli.

Our network model was able to reproduce these results but to do so required that locomotion change the effective synaptic connections, i.e., the effect of one class on another after taking into account amplification through recurrent excitation and inhibition. These changes in effective connections could, in turn, be instantiated through multiple possible modulations of physical synaptic strengths. The activity produced by a neural circuit is not always sufficient to constrain its underlying connectivity (Prinz et al., 2004); for the current model, we could mathematically prove that multiple underlying connectivity patterns yield identical sensory responses. Nevertheless, the parameter space consistent with our experimental observations favored one particularly attractive possibility, where locomotion would increase external excitatory input to all cell types, decrease recurrent excitation between Pyr cells, and increase recurrent inhibition between Pvalb cells. The first two of these are known effects of cholinergic modulation on cortical circuits (Gil et al., 1997).

In summary, our results suggest a set of simple rules for the interactions between pyramidal neurons and three classes of interneurons in the cerebral cortex and for how behavioral correlates such as locomotion may alter these interactions. We derived these rules from observations and made these observations in primary visual cortex. Future work will be needed to test these rules causally and to establish whether they describe a canonical circuit that is common to all of neocortex.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- **KEY RESOURCES TABLE**
- **CONTACT FOR REAGENT AND RESOURCE SHARING**
- **EXPERIMENTAL MODEL AND SUBJECT DETAILS**
  - Mice
  - Animal Preparation and Virus Injection
- **METHOD DETAILS**
  - Intrinsic Imaging
  - Retinotopic Mapping from Intrinsic Imaging
  - Visual Stimuli
  - Eye-Tracking Movie Acquisition and Analysis
  - In Vivo Calcium Imaging
- **QUANTIFICATION AND STATISTICAL ANALYSIS**
  - Calcium Data Processing
  - Pixel Maps of Calcium Data
  - Background Fluorescence Correction
  - Analysis of Neural Activity
  - Cell Selection
  - Phase of Locomotion
  - Correlation of Running Modulation with Depth
  - Modulation Index
DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes fifteen figures and one table and can be found with this article online at https://doi.org/10.1016/j.neuron.2018.03.037.

ACKNOWLEDGMENTS

We thank Charu B. Reddy, Chris Burgess, L. Federico Rossi, Miles Wells, Andrea Pisauro, and Sylvia Schröder for help with experiments or with software. We thank Boris S. Gutkin for constructive discussion and Laura Busse for sharing size-tuning curves of dLGN cells neurons. For the use of GCaMP6f we acknowledge Vivek Jayaraman, Rex A. Kerr, Douglas S. Kim, Loren L. Looger, and Karel Svoboda from the GENIE Project, Janelia Farm Research Campus, Howard Hughes Medical Institute. This work was supported by the Wellcome Trust (grants 095666, 095669, 108728, and 205093), M.D. was supported by a Marie Curie Intra-European Fellowship for Career Development (grant 627787), K.D.H. received support from ERC (grant 694401). M.C. holds a Marie Curie Intra-European Fellowship for Career Development (grant 694401), M.C. holds the GlaxoSmithKline/Fight for Sight Chair in Visual Neuroscience.

AUTHOR CONTRIBUTIONS

M.D. performed the experiments, analyzed the data, and implemented the model; M.D., A.R., and M.P. performed the surgeries; A.R. and M.K. built the eye-tracking software; M.K. implemented the eye-tracking software; K.D.H. and M.P. implemented the ROI detection algorithms; M.D., A.R., M.C., and K.D.H. designed the experiments; M.D. and K.D.H. designed the model; M.D., M.C., and K.D.H. wrote the paper with input from all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: June 9, 2016
Revised: July 26, 2017
Accepted: March 21, 2018
Published: April 12, 2018

REFERENCES


and high-throughput Cre reporting and characterization system for the whole mouse brain. Nat. Neurosci. 73, 133–140.


# STAR METHODS

## KEY RESOURCES TABLE

<table>
<thead>
<tr>
<th>REAGENT or RESOURCE</th>
<th>SOURCE</th>
<th>IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bacterial and Virus Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-Flex GCaMP6m</td>
<td>University of Pennsylvania Viral Vector Core</td>
<td>AAV1.Syn.GCaMP6m.WPRE.SV40</td>
</tr>
<tr>
<td>Flex GCaMP6m</td>
<td>University of Pennsylvania Viral Vector Core</td>
<td>AAV1.Syn.Flex.GCaMP6m.WPRE.SV40</td>
</tr>
<tr>
<td><strong>Experimental Models: Organisms/Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>tdTomato</td>
<td>The Jackson Laboratory; Madisen et al., 2010</td>
<td>Gt(ROSA)26Sor &lt; tm14(CAG-tdTomato)Hze &gt; ; RRID: IMSR_JAX:007909</td>
</tr>
<tr>
<td>Pvalb-Cre</td>
<td>The Jackson Laboratory; Hippenmeyer et al., 2005</td>
<td>Pvalb &lt; tm1(cre)Arbr &gt; ; RRID: IMSR_JAX:008069</td>
</tr>
<tr>
<td>Vip-Cre</td>
<td>The Jackson Laboratory; Taniguchi et al., 2011</td>
<td>Vip &lt; tm1(cre)Zjh &gt; ; RRID: IMSR_JAX:010908</td>
</tr>
<tr>
<td>Sst-Cre</td>
<td>The Jackson Laboratory; Taniguchi et al., 2011</td>
<td>Sst &lt; tm2.1(cre)Zjh &gt; ; RRID: IMSR_JAX:013044</td>
</tr>
<tr>
<td>Gad-Cre</td>
<td>The Jackson Laboratory; Taniguchi et al., 2011</td>
<td>Gad2 &lt; tm2(cre)Zjh &gt; ; RRID: IMSR_JAX:010802</td>
</tr>
<tr>
<td>Emx-Cre</td>
<td>The Jackson Laboratory; Gorski et al., 2002</td>
<td>Emx1 &lt; tm1(cre)Krj &gt; ; RRID: IMSR_JAX:005628</td>
</tr>
<tr>
<td>Rasgrf</td>
<td>Madisen et al., 2015</td>
<td>CamK2a-ITA; Ai94(TITL-GCaMP6s); Rasgrf2-2A-dCre</td>
</tr>
<tr>
<td><strong>Software and Algorithms</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MATLAB</td>
<td>MathWorks</td>
<td>N/A</td>
</tr>
<tr>
<td>Suite2P</td>
<td>Pachitariu et al., 2016</td>
<td>N/A</td>
</tr>
<tr>
<td>ScanImage</td>
<td>Pologruto et al., 2003</td>
<td>ScanImage 4.2</td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nano-injector</td>
<td>Drummond Scientific Company</td>
<td>Nanoject II injector</td>
</tr>
<tr>
<td>Pipette puller</td>
<td>Sutter</td>
<td>P-97</td>
</tr>
<tr>
<td>High-power LED (central wavelength: 560 nm)</td>
<td>Thorlabs</td>
<td>M565L3</td>
</tr>
<tr>
<td>CMOS camera (for intrinsic imaging)</td>
<td>Photonfocus</td>
<td>MV-D1024E-160</td>
</tr>
<tr>
<td>Microscope objective (for intrinsic imaging)</td>
<td>Olympus</td>
<td>UPLFLN, 4x, NA: 0.13, FN: 26.5</td>
</tr>
<tr>
<td>Collimated infrared LED (peak = 850 nm)</td>
<td>Mightex Systems</td>
<td>SLS-0208-B</td>
</tr>
<tr>
<td>Collimated infrared LED controller</td>
<td>Mightex Systems</td>
<td>SLC-AA02-US</td>
</tr>
<tr>
<td>Monochromatic camera</td>
<td>The Imaging Source</td>
<td>DMK 21BU04.H</td>
</tr>
<tr>
<td>Zoom lens (for eye tracking)</td>
<td>Navitar</td>
<td>MVL7000</td>
</tr>
<tr>
<td>Long-pass filter (for eye tracking)</td>
<td>The Imaging Source</td>
<td>092/52 × 0.75</td>
</tr>
<tr>
<td>Short-pass filter (for eye tracking)</td>
<td>Thorlabs</td>
<td>FES0900</td>
</tr>
<tr>
<td>Two-photon resonant-scanning microscope</td>
<td>ThorLabs</td>
<td>B-scope</td>
</tr>
<tr>
<td>Objective lens (for two-photon imaging)</td>
<td>Nikon</td>
<td>CFI75 LWD 16xW N.A.0.80, W.D.3.0mm</td>
</tr>
<tr>
<td>Piezo for z-scanning</td>
<td>PI</td>
<td>P-725.4CA (with E-665.CR controller)</td>
</tr>
<tr>
<td>Pockel’s cell</td>
<td>Conoptics</td>
<td>M350-80-LA-BK-02</td>
</tr>
<tr>
<td>Pockel’s driver</td>
<td>Conoptics</td>
<td>302 RM</td>
</tr>
<tr>
<td>Multifunction I/O</td>
<td>National Instruments</td>
<td>PCIe-6321</td>
</tr>
</tbody>
</table>

Please cite this article in press as: Dipoppa et al., Vision and Locomotion Shape the Interactions between Neuron Types in Mouse Visual Cortex, Neuron (2018), https://doi.org/10.1016/j.neuron.2018.03.037
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources should be directed to and will be fulfilled by the Lead Contact Mario Dipoppa (m.dipoppa@ucl.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

All experimental procedures were conducted according to the UK Animals Scientific Procedures Act (1986). Experiments were performed at University College London under personal and project licenses released by the Home Office following appropriate ethics review.

Mice

Experiments in which an interneuron class was labeled with tdTomato and recorded together with other cells were conducted in double-transgenic mice obtained by crossing Gt(Rosa)26 Sor < tm14(CAG-tdTomato)Hze > reporters (Madisen et al., 2010) with appropriate drivers: Pvalb<tm1(cre)Arbr> (Hippenmeyer et al., 2005) (2 males, 3 females), Vip<tm1(cre)Zjh> (Taniguchi et al., 2011) (3 males, 2 females), Sst<tm2.1(cre)Zjh> (Taniguchi et al., 2011) (2 males, 1 female), and Gad2<tm2(cre)Zjh> (Taniguchi et al., 2011) (2 females). Experiments in which indicator was expressed uniquely in one neuron class were conducted in single transgenic mice: Emx1-ires(cre) (n = 1), Pvalb<tm1(cre)Zjh> (n = 1), Vip<tm1(cre)Zjh> (n = 1), Sst<tm2.1(cre)Zjh> (n = 3), referred to as Vip-Cre and Sst-Cre respectively. Experiments in which pyramidal cells were labeled exclusively were conducted in CamK2a-tTA; Ai94(TITL-GCaMP6s); Rasgrf2-2A-dCre triple transgenic mice (n = 3) (Madisen et al., 2015). Mice were used for experiments at adult postnatal ages (P54-110).

Animal Preparation and Virus Injection

The surgeries were performed in adult mice (P35–P76) in a stereotaxic frame and under isoflurane anesthesia (5% for induction, 0.5%–3% during the surgery). During the surgery we implanted a head-plate for later head-fixation, made a craniotomy with a cranial window implant for optical access, and, on relevant experiments, performed virus injections, all during the same surgical procedure. In experiments where an interneuron class was recorded together with other cells, mice were injected with an unconditional GCaMP6m virus, AAV1.Syn.GCaMP6m.WPRE.SV40 (referred to as non-flex.GCaMP6m). In experiments where an interneuron class was labeled by unique expression, mice were injected with AAV1.Syn.Flex.GCaMP6m.WPRE.SV40 (flex.GCaMP6m) and AAV2/1.CAG.FLEX.tdTomato.WPRE.bGH (flex.tdTomato); all viruses were acquired from University of Pennsylvania Viral Vector Core. At the time of the injection, the mice were already adult, thus excluding the off-target expression that might occur in cells expressing Cre only transiently during development (Hu et al., 2013). Viruses were injected with a beveled micropipette using a Nanoject II injector (Drummond Scientific Company, Broomall, PA 1) attached to a stereotaxic micromanipulator. One to three boli of 100–200 nl virus (2.33×10¹² GC/ml for non-flex.GCaMP6m; 2.71×10¹² for flex.GCaMP6m) were slowly (23 nl/min) injected unilaterally into monocular V1 (Wagor et al., 1980), 2.1-3.3 mm laterally and 3.5-4.0mm posteriorly from Bregma and at a depth of L2/3 (200-400 μm).

METHOD DETAILS

Intrinsic Imaging

Prior to performing calcium imaging experiments, we performed intrinsic imaging of the optically accessible cortex to confirm the location of V1 within the cranial window (Figures S1A and S1B). The intrinsic imaging was performed in all mice (n = 22) about 7-14 days after the surgery. We illuminated the cortex through the epi-illumination path using a high-power LED (central wavelength: 560 nm, M565L3, Thorlabs, Ely, UK), and acquired images at 5 Hz at 1024 pixels using a CMOS camera (MV-D1024E-160; Photonfocus, Lachen, Switzerland) combined with a microscope objective (4x, NA: 0.13, FN: 26.5, UPLFLN, Olympus, Tokyo, Japan). To prevent the light contamination from the computer monitors we optically shielded the recording chamber with a custom black cone surrounding the objective.

Retinotopic Mapping from Intrinsic Imaging

To obtain retinotopic maps from intrinsic imaging we used the methods described in Pisauro et al. (2013). Briefly, we first removed global fluctuations from the signal, which are not stimulus driven. The residual signal reflects the retinotopic, stimulus-evoked responses. Visual stimuli were periodic drifting and flickering bars (Kalatsky and Stryker, 2003). Flickering bars (flicker frequency 2 Hz) drifted (speed = 0.8 deg/s) across –135° to 45° of the horizontal visual field (with bars oriented vertically) and –45° to 45° of the vertical visual field (with bars oriented horizontally) for 3 cycles. We calculated retinotopic maps using the method described in Kalatsky and Stryker (2003). Retinotopic contours (Figure S1B) where obtained after removal of artifactual extreme values (e.g., red regions in the top and bottom left corners of Figure S1A) and replacing the removed values by values interpolated using sum of normalized Gaussian functions with standard deviation of 20 μm centered on non-artifactual pixels. Consistent with a location in V1, the imaged regions (Figure S1C) were within an area of diameter at least 2 mm where the gradient of vertical retinotopy was
aligned from anterior to posterior (lower to higher values of elevation) and the gradient of the horizontal retinotopy was aligned from medial to lateral (temporal to central).

**Visual Stimuli**
Stimuli were horizontal gratings drifting downward, presented in a location adjusted to match the center of GCaMP expression, on one of two screens that together spanned $-45^\circ$ to $+135^\circ$ of the horizontal visual field and $\pm 42.5^\circ$ of the vertical visual field (left and central screens in Figure 1A). During gray screen presentation, the screens were set to a steady gray level equal to the background of all the stimuli presented for visual responses protocols. Gratings had a duration of 1-2 s temporal frequency of 2 Hz and spatial frequency of 0.15 cycles/deg. Note that during the presentation of all stimuli we switched off the red gun of the monitors in order to reduce an artifact of light from the monitors contaminating the red fluorescent channel. Hence what we defined as gray screen actually corresponds to the color cyan.

**Eye-Tracking Movie Acquisition and Analysis**
For eye tracking we used a collimated infrared LED (SLS-0208-B, $\lambda_{\text{peak}} = 850\text{nm}$; controller: SLC-AA02-US; Mightex Systems, Toronto, Canada) to illuminate the eye contralateral to the recording site. Videos of eye position were captured at 30 Hz with a monochromatic camera (DMK 21BU04.H, The Imaging Source, Bremen, Germany) equipped with a zoom lens (MVIL7000; Navitar, Rochester, NY), and positioned at approximately $50^\circ$ azimuth and $50^\circ$ elevation relative to the center of the mouse’ field of view. Contamination light from the monitors and the imaging laser was rejected using an optical band-pass filter (700-900nm) positioned in front of the camera objective (long-pass 092/52x0.75, The Imaging Source, Bremen, Germany; short-pass FES0900, Thorlabs, Ely UK).

To calibrate pupil displacement relative to the mouse visual field, we recorded additional movies at the end of each experiment while the mouse was still in exactly the same position as during the experiment. The eye was illuminated sequentially from a grid of known locations, the reflections were captured by the camera, and then this reflected grid was used to map the pupil displacement in pixels to pupil displacement in degrees of visual field.

Movie processing was performed offline using custom code written in MATLAB (Mathworks, Natick, MA) on a frame-by-frame basis. Briefly, each frame was mildly spatially low-pass filtered to reduce noise, then the pupil contour was detected by a level-crossing edge detector, and finally the position and the area of the pupil were calculated from the ellipse fit to the pupil contour. The output of the algorithm was visually inspected, and adjustments to the parameters (e.g., spatial filter strength, or level-crossing threshold) were made if necessary.

**In Vivo Calcium Imaging**
Experiments were performed 16-34 days after virus injection (P54-110). We used a commercial two-photon microscope with a resonant-galvo scanhead (B-scope, ThorLabs, Ely UK) controlled by ScanImage 4.2 (Pologruto et al., 2003), with an acquisition frame rate of about 30Hz (at 512 by 512 pixels, corresponding to a rate of 4-6 Hz per plane), which was later interpolated to a frequency of 10 Hz, common to all planes. Recordings were performed in the area where expression was strongest. In most recordings (n = 16) this location was in the monocular zone (MZ, horizontal visual field preference $> 30^\circ$) (Wagor et al., 1980). Other recordings (n = 11) were performed in the callosal binocular zone (CBZ, $n = 4$, 0-15°) (Wang and Burkhalter, 2007) and others (n = 7) in the acallosal binocular zone (ABZ, 15-30°). We observed no difference in results between recordings in monocular and binocular zones (Figure S12).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Calcium Data Processing**
Raw calcium movies were analyzed with Suite2p, which performs several processing stages (Pachitariu et al., 2016). First, Suite2p registers the movies to account for brain motion, then clusters neighboring pixels with similar time courses into regions of interest (ROIs). ROIs were manually curated in the Suite2p GUI, to distinguish somata from dendritic processes based on their morphology. Cells expressing tdTomato were identified semi-automatically using an algorithm based on their average fluorescence in the red channel. For spike deconvolution from the Calcium traces, we used the default method in Suite2p (Pachitariu et al., 2016). Whether we performed spike deconvolution or analyzed raw calcium signals made no difference to our results (Figure S13).

**Pixel Maps of Calcium Data**
To confirm the correlation of running speed and fluorescence independent of ROI detection, we computed correlation maps (Figure S2C), showing for each pixel the Pearson correlation between the activity of the pixel and the running speed (c.f. Freeman et al., 2014). Prior to correlation, the activity of each pixel was smoothed by convolving with a spatial Gaussian with standard deviation equal to 1.5 pixels, and a temporal Hamming window of 1 s width.

The correlation of baseline fluorescence with running speed varied across the field of view. In regions where GCaMP expression level was high, baseline fluorescence correlated positively with running speed, likely indicating an increase in axonal and dendritic activity in locomoting animals. However, in areas where GCaMP fluorescence was weak, the correlation of the background with running speed was negative, likely indicating that in absence of GCaMP the signal is dominated by increased hemodynamic filtering...
of the light due to stronger blood flow during running (Huo et al., 2015). To ensure this did not affect our results, we removed background fluorescence from the detected fluorescence of recorded neurons (see below).

**Background Fluorescence Correction**

With two-photon GCaMP imaging, an important concern is that out-of-focus fluorescence can contaminate the signal ascribed to particular neurons; this is of particular concern in situations where the surrounding GCaMP-labeled neuropil may itself show modulation by stimuli or behaviors such as locomotion. In order to correct out-of-focus contamination, we adopted the method of Peron et al. (2015). A “neuropil mask” was defined as the region up to 35 μm from the ROI border, excluding pixels corresponding to other detected cells (Figure S2A), and the fluorescence signal in this mask region was subtracted from that of the cell soma, weighted by a correction factor \( α_{exp} \) that was determined separately for each experiment.

To determine the correction factor, we estimated the linear relationship specifying the lowest possible somatic fluorescence compatible with any value of fluorescence in the neuropil mask (Figure S2A). To do so, for each cell \( i \) we binned the neuropil signals \( N_i(t) \) into 20 intervals, and for each one estimated the 5th percentile of the raw somatic fluorescence \( F_i(t) \). We computed \( α_i \) by linear regression, which accurately matched the lower envelope of the scatterplot of neuropil versus somatic fluorescence (Figure S2A). This method gave consistent results for sparse firing cells, but not always for densely firing cells for which a correlation of cellular activity with the neuropil signal could lead to misestimated slopes, as densely firing cells might only rarely exhibit baseline fluorescence. We therefore computed the correction factor \( α_{exp} \) for each experiment by averaging \( α_i \) over cells with high skewness (> 4). The corrected fluorescence was computed as \( F(t)−α_{exp}N(t) \) (Figure S2A). In experiments where only interneurons (and thus low skewed cells) expressed GCaMP6, we used as a correction factor an average from all the other experiments equal to \( \langle α_{exp} \rangle = 0.82 \).

**Analysis of Neural Activity**

The average fluorescence response to each stimulus was defined by \( ΔF/F_0 = (F-F_0)/F_0 \), where \( F \) is the average raw calcium signal during the first second of the stimulus presentation, and \( F_0 \) is the global minimum of the fluorescence trace filtered with a Hamming window of duration 0.5 s. The correlation of neural activity with locomotion speed during gray screen presentation was assessed by the Pearson correlation coefficient between the calcium signal and the locomotion speed trace, on an interpolated timescale of 10 Hz, smoothed (5 points) and decimated (1 Hz). To ascertain the significance of this correlation we used a shuffling method, in which the speed trace was randomly circularly shifted relative to the fluorescence trace 1,000 times – this was necessary because serial correlation in the time series of fluorescence and speed rendered successive samples statistically dependent.

The size of a cell’s response to a stimulus was defined by the difference of \( ΔF/F_0 \) between the first 1 s of the stimulus period, and the 1 s of baseline activity prior to stimulus presentation. We defined a neuron to have significant size tuning if it passed in at least one of the two locomotion conditions (rest or running) a one-way ANOVA test (\( p < 0.05 \)) comparing the mean visual responses to different stimuli.

To measure each cell’s retinotopic location, in the majority of datasets (\( n = 24 \)) receptive fields were obtained from responses to sparse, uncorrelated noise. The screen was divided into squares of 5 by 5 degrees, and each square was independently turned on and off randomly at a 5Hz overall rate. At any time, 5% of all squares were on. Each cell’s response to each square was obtained using stimulus-triggered averaging of the non-neuropil corrected trace. The RFs were smoothed in space and their peak was identified as the preferred spatial position. In a subset of early experiments (\( n = 3 \)), sparse noise was not presented, and RFs were assessed with flickering vertical or horizontal bars appearing in different locations; we verified in a further \( n = 4 \) datasets that the two measures of a cell’s preferred spatial position were compatible with any value of fluorescence in the neuropil mask (Figure S2).

When computing size tuning curves, we normalized the calcium activity (Figure 4, column 3 and Figure S12) or the spike rate (Figure S13) in the following way: for each cell, the response \( ΔF_0 \) to the “blank condition” (i.e., a stimulus of contrast 0) during stationary periods was subtracted from the raw cell response \( ΔF \) (1 s during stimulation minus 1 s of baseline activity): \( ΔF = ΔF_0 - ΔF_0 \). Then, for each recording we computed the average \( \langle ΔF \rangle_{all \ cells} \) over all selected cells whose distance of the receptive field from the stimulus center \( r \) was within a radius of 20°. Finally we divided the average response of either the centered cells (radius: \( r < 10° \)) or the off-centered cells (radius: \( r > 10° \) \( r < 20° \)) by the maximum value of all the cells combined: \( \langle ΔF \rangle_{cent} / \max (\langle ΔF \rangle_{all \ cells}) \) or \( \langle ΔF \rangle_{off-cent} / \max (\langle ΔF \rangle_{all \ cells}) \). We then averaged these values across all recording sessions.

When computing the interaction between locomotion and size (Figure 4, column 4 and Figures S10A and S10B) we normalized the responses of the individual cells for each experiment in the following way: after subtracting the average response to the blank during stationary periods \( ΔF = ΔF_0 - ΔF_0 \), we divided the responses by the average minimal calcium trace across cells \( \langle F_0 \rangle_{cells} \) within the same experiment or mouse (Figures S10A and S10B).

The relationship between calcium fluorescence and spiking is not completely characterized and might differ between cell types; nevertheless, we are confident that the specific measurements we perform here are unlikely to be affected by cell-type differences in calcium handling. First, we do not attempt to estimate firing rates or exact spike times in the main text: the deconvolution method of Figure S13 is not used for our main analyses (with the exception of Figures S5A–S5D, column 1,2 as detailed in the methods), but indicates that performing deconvolution analysis made no difference to our results. Second, even if the spike-to-GCaMP transform were several times larger in one cell type than another, this would not affect our conclusions: our analyses of run-speed modulation and size tuning always take place within a cell type, and correlations between cells (e.g., Figure 4) would not be affected by absolute firing rate changes. Third, differences in the kinetics of calcium handling between cell types (in particular their relaxation dynamics)
would not affect our results: we averaged calcium signals over the entire stimulus presentation, and used very long inter-stimulus intervals (2-5 s), to make sure that even a very long-lasting effect of the response to previous stimulus would only minimally interfere with the response to the current stimulus. Fourth, a difference in the nonlinearity of the spike-GCaMP coupling between cell types could not change the direction of locomotor tuning in individual cells, nor the existence of size tuning or preferred size, although it might exaggerate the “peakiness” of size tuning in some cell classes compared to others. While no systematic cell-type comparisons of spike-GCaMP coupling have been carried out to our knowledge, the data of Chen et al. (2013) suggests a linear relationship between the number of action potentials and peak ΔF/F₀, including at least Pvalb interneurons (their Figure S12F). In summary, we are confident that our findings are robust to most conceivable differences in spike-GCaMP coupling between cell types.

Cell Selection
For characterization of skewness (Figure 3, columns 3 and 4) and spontaneous activity (Figure 2; Figure 5, columns 1 and 2) we analyzed all detected cells.

For analysis of visual responses (Figure 3) and size tuning (Figure 4, columns 1–3; Figure 5, columns 3 and 4), we selected only centered cells: we chose cells whose RF was within 10° of the stimulus center and in which the effect of size on visual responses was significant during either stationary or locomotion periods (p < 0.05, significant effect of stimulus size, p < 0.05, 1-way ANOVA). For analysis of the effect of locomotion on different stimulus sizes (Figure 4, column 4) we used the same selection criterion of visual responses but the cells’ RF was allowed to be at a distance of 15° from the stimulus center.

To analyze how visual responses depended on both stimulus size and centering (Figure 6, columns 2–5) we used all detected cells. We did not use orientation tuning as a criterion for cell selection in any of our figures.

Phase of Locomotion
Previous studies have suggested that the modulation of spontaneous activity by locomotion can depend on the phase of the locomotion period, with stronger responses at locomotion onset (Vinck et al., 2015). However, for all cell types we found similar correlations between fluorescence and running speed after removing transition periods between locomotion and stationary periods from the analysis (Figure S6).

Correlation of Running Modulation with Depth
To determine whether running modulation of a given cell class varied significantly with cortical depth (Figure 2, column 2; Figure S7C), we computed ρ(gray) for each cell as the Pearson correlation of that cell’s neuropil-corrected fluorescence (without spike deconvolution) and running speed. We then assessed a significant relationship of ρ(gray) with depth using robust regression (bisquare-weighting).

Modulation Index
To measure how locomotion modulated baseline and evoked activity (regardless of stimulus size), we computed two indices M_B and M_R. In order to compute M_B we first computed, for each trial t, the baseline-subtracted responses ΔF_s = F_{post}^t - F_{pre}^t as the difference between the average activity during 1 s after the stimulus onset and the 1 s before the stimulus onset. We computed the average response to each stimulus size s as ΔF_s = mean{ΔF_i}_i(S(t)-s), where S(t) represents stimulus size on trial t. We then computed each trial’s residual response as d_i = ΔF_i - ΔF_{post}/S(D), and collected these residuals into a set for each locomotor condition (ν = 0: stationary, ν = 1: locomotion): d(ν) = {d_i}_i(ν)-ν. Finally, we computed the modulation index M_R for each cell’s visual responses as M_R = |⟨d(1)⟩ - ⟨d(0)⟩|/√σ^2[ΔF(1)] + σ^2[ΔF(0)].

To compute the baseline modulation M_B, we divided the period of uniform screen presentation into “virtual trials” of 1 s duration, and computed the average activity F_i for each of these. We separated the virtual trials into two sets according to locomotion condition V: F(ν) = {F_i}_i(V=ν). We finally computed the modulation index M_B on baseline (σ) for each cell as M_B = |⟨F(1)⟩ - ⟨F(0)⟩|/√σ^2[F(1)] + σ^2[F(0)]].

Curve Fitting
We fitted the size tuning curves of Figure 4 and Figures S4, S8, S9, S12, and S13 by least-squares with the following function family:

f(s) = R[erf(s/σ₁) - k erf(s/σ₂)] where erf(x) corresponds to the error function and s is the size of the stimulus. The free parameters of the function are R, k, σ₁ and σ₂. To estimate nonlinear signal correlation curves (Figure 5, columns 3 and 4), we first smoothed the responses to large sizes (diameter: s > 20°) for each population with a boxcar moving average method with span 25°. Then we smoothed again the responses for all sizes with a boxcar moving average method with span 20°. Finally we interpolated the values between the measured size with a shape-preserving piecewise cubic interpolation.

Size-Tuning Maps
To compute how size tuning depends on stimulus centering, we computed two-dimensional maps illustrating how each cell class’ average activity depends on stimulus diameter s, and the offset of the receptive field center from the stimulus center r. (Figure S11). To
do so, we first computed for each cell \(i\) a normalized tuning curve \(n_{ij}(s)\), where \(v\) represents locomotion condition, by using a shape-preserving piecewise cubic interpolation of \(\Delta F\). Dependence on \(r\) was estimated by smoothing: a two-dimensional map was made for each cell as an outer product: \(f_{i,j}(s,r) = n_{ij}(s)g_{ij}(r)\), where \(g_{ij}(r) = e^{-(r/r_0^2)/(2\sigma_r^2)}\) is a Gaussian centered at the offset value \(r\), of width \(\sigma_r = 5\). We chose \(\sigma_r = 5\) to match the size of the square pixels (5’ width) used in the sparse noise stimulus. Then we summed the maps belonging to one recording session \(j\) and divided by the sum of all the Gaussians centered at different offsets: \(m_{ij}(s,r) = \sum_j f_{ij}(s,r)/\sum_i g_{ij}(r)\) where the sum over \(i\) of \(g_{ij}(r)\) corresponds to the density distribution of the cells across \(r\); dividing by \(\sum_i g_{ij}(r)\) is necessary to avoid edge effects, ensuring that the average visual response for each \(s\) and \(r\) is normalized by the occurrence of cells having a particular value of \(r\). We then normalized this value for each experiment by the value at stationary, \(0^\circ\) offset and diameter \(10^\circ\) \(f_{ij}(s,r) = m_{ij}(s,r)/m_{ij,-1}(s = 10, r = 0)\). Finally, for each cell class we obtained the size-tuning offset maps by averaging across experiments: \(f_{ij}(s,r)\).

**Inter-population Correlation Analysis**

To compute spontaneous correlations (Figure 5, columns 1 and 2), we first normalized the deconvolved spike trace \(S\) of each cell \(i\) over time \(t\): \(n(i,t) = S(i,t)/\max(S)\). Then, for each experiment and each cell class \(c\) (interneurons or putative Pyr cells) we computed the average population rate across cells \(i\) belonging to class \(c\): \(R_c(t) = \langle n(i,t) \rangle_{i \in c}\). We then smoothed \(R_c(t)\) with a boxcar moving average method with span of 1 s and then decimated the sampling rate to 1 point every 1 s. To make the plots of different experiments visually comparable we normalized these responses: \(K_c(t) = (R_c(t) - R_c^0)/\sigma_i(R_c)\). Where \(R_c^0\) is the 1st percentile of \(R_c\) and \(\sigma_i(R_c)\) the standard deviation of \(R_c(t)\) across time.

To compute signal correlations (Figures 5, columns 3 and 4), for each experiment and each cell class \(c\) we first computed the average population response for each stimulus size \(s\) and locomotion condition \(v\) (\(v = 0\) stationary, \(v = 1\) running) by averaging over all cells \(i\) belonging to that class: \(\Delta R_c(s,v) = \langle \Delta f(i,s,v) \rangle_{i \in c}\). To make the plots of different experiments visually comparable we normalized the responses: \(\Delta R_c(s,v) = \Delta F_c(s,v)/\sigma_i(\Delta F_c)\). Finally, we subtracted the blank response during the resting condition: \(\Delta K_c(s,v) = \Delta R_c(s,v) - \Delta R_c(s = 0, v = 0)\).

To compute noise correlations (Figures S14A–S14D), for each experiment and each cell class \(c\) we first computed the average population response for each trial \(t\) by averaging over all cells \(i\) belonging to that class: \(\Delta F_c(t) = \langle \Delta f(i,t) \rangle_{i \in c}\). Then for each stimulus and locomotion condition we subtracted the mean response from the related trials: \(\Delta N_c(t; s,v) = \Delta F_c(t; s,v) - \langle \Delta F_c(t) \rangle_{s,v}\).

Finally, to make the plots of different experiments visually comparable, we normalized the responses by z-scoring over all trials: \(\Delta Z_c(t) = (\Delta N_c(t) - \langle \Delta N_c \rangle)/\sigma(\Delta N_c)\).

When measuring signal and noise correlations and for both interneurons and putative Pyr neurons, we selected cells whose receptive field center was within a radius of \(10\) from the stimulus center. For Figure 5 we selected putative Pyr cells as unlabeled (non tdTomato) neurons whose skewness was \(＞2.7\). In a control analysis (Figure S14) we show that the value of the skewness threshold makes little difference to these results and similar classification results have been obtained if using kurtosis instead of skewness (Ringach et al., 2016). A skewness value of 0 corresponds to the case where we selected all unlabeled cells as putative Pyr cells.

**Computational Model**

We asked whether we could predict the mean size tuning of each cell class using a neural field theory model. In this model, the mean firing rate of cells of type \(c\) is captured by a function \(f_c(s,r)\), where \(s\) represents the stimulus size, and \(r\) represents position on the cortical surface, measured in retinotopic coordinates. We model the external excitatory input arriving at point \(r\) (e.g., from thalamus or cortical layers) by a function \(h(s,r)\), again in retinotopic coordinates.

We denote the experimentally measured responses of cell class \(c\) by \(f^{\text{Exp}}(s,r)\), where \(v\) denotes locomotion condition (\(v = 0\): stationary, \(v = 1\): running). We assume that responses are circularly symmetrical, i.e., that responses depend on \(r\) only through the radial distance of the receptive field center from the stimulus center, \(r\). The response of each cell class \(f^{\text{Exp}}(s,r)\) is modeled by the following equations:

\[
\begin{align*}
    f_s &= R[\text{w}_{EV}(h + \text{w}_{sE} \text{[G}_{SE} + f_s]) - \text{w}_{sE} \text{[G}_{ES} + f_s])]; \\
    f_R &= R[\text{w}_{Ph}(h + \text{w}_{sE} \text{[G}_{RE} + f_s]) - \text{w}_{sE} \text{[G}_{RP} + f_s])]; \\
    f_v &= R[\text{w}_{sG} \text{[G}_{SH} + h]) + \text{w}_{sE} \text{[G}_{SV} + f_s])]; \\
    f_v &= R[\text{w}_{sG} \text{[G}_{SV} + f_s])]; \\
    f_v &= \text{w}_{sE} \text{[G}_{EV} + f_s])]; \\
    w_{sE} &= \text{w}_{sE} \text{[G}_{EV} + f_s])]; \\
    w_{sE} &= \text{w}_{sG} \text{[G}_{SV} + f_s])]; \\
    w_{sE} &= \text{w}_{sE} \text{[G}_{EV} + f_s])]; \\
    w_{sE} &= \text{w}_{sG} \text{[G}_{SV} + f_s])];
\end{align*}
\]

Only synaptic connections demonstrated in vitro (Pfeffer et al., 2013) are included in this equation; however, adding other potential synapses (Vip->Pyr) did not improve the fit (Figure S15).

For each postsynaptic cell class we tested different combination of subtractive and divisive inhibition from Sst and Vip cells:

\[
    R(x;y) = \begin{cases} 
    x - y, & \text{subtractive} \\
    x / (1 + y), & \text{divisive}
    \end{cases}
\]

As we discuss later in the text, the model predicts a subtractive inhibition from Sst to Pyr and Pvalb cells and from Vip to Sst cells, while it predicts a divisive inhibition from Vip to Sst cells.
Here, \( f_E, f_P, f_S, \) and \( f_V \) reflect the visual responses of the Pyr, Pvalb, Sst, and Vip cells respectively; \( z_+ \) is the positive part of \( z; w_{i,j} \) are the peak synaptic weights between the presynaptic cell class \( i \) and the postsynaptic cell class \( j \) (which can in principle depend on running condition \( v \)); \( G_{a,j} \) is a two-dimensional Gaussian function defined by \( G_{a,j}(z) = \exp[-(z^2/(2\sigma_{a,j}^2))]/(2\pi\sigma_{a,j}), \) with radius \( \sigma_{a,j} \) that depend on the pre- and post-synaptic cell type; and \( * \) represents convolution over retinotopic space: \( [G_{a,j} * f_j](s,r) = \int G_{a,j}(t-r)/f_j(s,r) d^2 r. \)

The equations describing the activity of the cell classes can be simplified with the following assumptions:

- \( \forall (v, s, r), f_E > 0, f_P > 0 \) (Pyr and Pvalb cells are not suppressed by stimuli, as seen in the data).
- The recurrent connections of Pyr and Pvalb neurons, and the connection from Pvalb to Pyr are local:
  \( \Gamma_{PE}(r) = \Gamma_{PP}(r) = 0 \) (where \( \delta(r) \) is the Dirac delta function).
- \( f_P = \mu_E \) (i.e., Pvalb activity closely tracks Pyr activity, as seen in the data).

We can then rewrite the equations as:

\[
\begin{align*}
  f_E &= R[w_E h; w_{ES}(G_{ES} * f_S)]/(1 - w_{EE} + \mu_{c,PE}) \\
  f_P &= R[w_{PH} h + w_{PE}(G_{PE} * f_E); w_{PS}(G_{PS} * f_S)]/(1 + w_{PP}) \\
  f_S &= R[w_{SH}(G_{SH} + h) + w_{SE}(G_{SE} * f_E); w_{SV}(G_{SV} * f_V)] \\
  f_V &= R[w_{VE}(G_{VE} * f_E); w_{VS}(G_{VS} * f_S)]
\end{align*}
\]

We can further simplify this equation as:

\[
\begin{align*}
  f_E &= R[w_{EH} h; w_{ES}(G_{ES} * f_S)] \\
  f_P &= R[w_{PH} h + w_{PE}(G_{PE} * f_E); w_{PS}(G_{PS} * f_S)] \\
  f_S &= R[w_{SH}(G_{SH} + h) + w_{SE}(G_{SE} * f_E); w_{SV}(G_{SV} * f_V)] \\
  f_V &= R[w_{VE}(G_{VE} * f_E); w_{VS}(G_{VS} * f_S)]
\end{align*}
\]

Estimation of Thalamic Input

The response to a stimulus of size \( s \) in thalamic cells with receptive fields located at position \( r \) in running condition \( v \) was modeled by a function \( h^{(i)}(s,r), \) estimated from the retinotopic recordings of Erskine et al. (2014). We first estimated the firing of centered cells as a function of stimulus size \( h^{(i)}(s,0) \), for stationary and locomotion periods separately. We fit the empirical size tuning curve of each cell \( (i = 1 \ldots 21) \) using the same method as Erskine et al. (2014): \( h^{(i)}(s,0) = b_i \text{erf}(s/m_i)^2 / (1 + b_i \text{erf}(s/m_i)^2), \) and estimated the mean thalamic response by averaging across all cells (after normalizing each \( h^{(i)}(s) \) by its maximum across \( s \) and \( v \)). The empirical data we had were only of LGN neurons. To extrapolate to off-center responses, we used a Ratio-of-Gaussians model (Ayaz et al., 2013) as a parametrized function: \( h^{(i)}(s,r) = a_i u(s,r,\sigma_i)/[1 + a_2 s(r,r,\sigma_i)] \) with \( u(s,r,\sigma) = \text{erf}(s/\sigma)/\sigma + \text{sign}(\sigma r) - \text{erf}(|s - r|)/\sigma \) fitted on the centered responses. The estimated parameters during the stationary periods were \( a_1 = 1.2, a_2 = 1.9, \sigma_1 = 36.7, \) and \( \sigma_2 = 33.9 \) while during locomotion \( a_1 = 0.5, a_2 = 0.4, \sigma_1 = 24.7, \) and \( \sigma_2 = 10.0. \)

Estimation of Presynaptic Inputs

To fit the model, we clamped the firing rate functions \( f^{(i)}_a(s,r) \) in Equation (1) to their experimentally measured values, and fit synaptic parameters to reduce the discrepancy between the right and left sides. To do so required extending our experimental data to continuous functions of \( s \) and \( r. \) For retinotopic positions \( r < r_0 = 33^\circ, f(s,r) \) was fitted from the data with a difference of Gaussians function: \( f(s,r) = R_i \text{erf}(s/\sigma_1) - k_r \text{erf}(s/\sigma_2a), \) because our data for cells off-center by a radius of more than \( 33^\circ \) were sparse, we extrapolated the values for \( r > r_0 \) with a decaying exponential approximation: \( f(s,r) = f(s,r_0) e^{-(r-r_0)/b}, \) to obtain the parameter \( b \) we first fit the values \( f(s,r) \) in the range \( r_m \leq r \leq r_0 \) (where \( r_m \) is the offset value that maximizes the response of that cell class) with an exponential decaying function with spatial coefficient \( c(s) \) for each stimulus size \( s. \) Then we estimated \( b \) as the average of \( c(s) \) between the values \( 0^\circ \leq s \leq 30^\circ \). The values of \( b \) that we obtained were 15.2 (stationary) and 13.5 (locomotion) for Pyr, 32.1 (stationary) and 20.1 (locomotion) for Pvalb, 22.3 (stationary) and 24.8 (locomotion) for Vip and 13.1 (stationary) and 21.5 (locomotion) for Sst cells.

Parameter Estimation

To estimate the parameters we minimized an objective function equal to the normalized mean-square error, plus additional penalty terms to favor simpler models:

\[
\text{Err} = \left( \frac{f^{(i)}_a(s,r) - f^{(i)}_a(s,r)}{\text{var}(f^{(i)}_a(s,r))} \right)^2 + \lambda_1 \sigma_{aw}^2 + \lambda_2 \sigma_{cw}^2 + \lambda_3 R^2
\]
Here, \( f^{(v)}_{nu} \) represents the measured firing rate, \( \hat{f}^{(v)}_{nu} \) represents the right hand side of Equation (1) and \( \text{var}_x(f^{(v)}_{nu}(s, r)) \) denotes the variance of normalized visual responses for each value of \( s, r \) and \( v \) over all experiments \( x \). Each experiment \( x \) was performed in a different field of view (and therefore different neurons), different days and group of experiments were performed in different mice. The normalization factor of \( 1/\text{var}_x(f^{(v)}_{nu}(s, r)) \) ensures that conditions with high inter-experiment variability do not overly influence the objective function; the normalized error can also be interpreted as the log-likelihood of the model fit under a Gaussian distribution estimated from all experiments \( x \). The averaging operator \( \langle \cdot \rangle_{s,v,r} \) runs over the space \( 0^\circ \leq s \leq 60^\circ \) and \( 0^\circ \leq r \leq 33^\circ \), in both stationary and locomotion conditions.

The last three terms represent regularization parameters. The first regularization term controls the number of synaptic strengths that are allowed to change with locomotion, \( n_{aw} \) (this L0 regularization method penalizes according to the number of non-zero weights, without regard to their magnitude), and for the current analysis we used a value of \( \lambda_1 = 0.1 \). The second regularization term controls the spatial distribution of synaptic weights; we used parameters \( \sigma_L = 40^\circ, \sigma_2 = 0.01 \). The final term \( R_a \), with \( \lambda_3 = 0.2 \), represents a factor to add biologically motivated constraints in the \( \text{Pvalb} \) and \( \text{Sst} \) equations.

We assume that \( \text{Sst} \) cells receive most of their input from a local area, therefore

\[
R_s = \sum_{v=1,2} \frac{\text{W}_{SH}^{(v)}}{\text{W}_{SE}^{(v)} + \text{W}_{SV}^{(v)}}.
\]

The activities of \( \text{Pyr} \) and \( \text{Pvalb} \) are very similar, so to avoid \( f_c \) dominating the \( f_P \) equation we have

\[
R_p = \sum_{v=1,2} \frac{\text{W}_{PE}^{(v)}}{\text{W}_{PD}^{(v)} + \text{W}_{PS}^{(v)}}.
\]

To determine the optimal parameters (reported in Table S1) of the model we first performed an exhaustive search over the extent of the spatial integration \( \sigma_{ab} \) parameters of all the presynaptic cell classes for each postsynaptic cell class. For inputs from the visual input and excitatory cells, we searched the ranges from 1° to 40° at 15 equally spaced steps. For inputs from \( \text{Sst} \) and \( \text{Vip} \) cells we searched a range from 1° to 100° at 12 equally spaced steps (Figure S15C). For each combination of \( \{\sigma_{ab}, \sigma_{d2}, \ldots\} \) we then found the optimal synaptic strength parameters \( \text{W}^{(v)}_{ab} \) (or effective strengths \( \tilde{w} \) for \( \text{Pyr} \) and \( \text{Pvalb} \) cells) using a combination of the trust region reflective and Levenberg-Marquardt algorithms (MATLAB), after 50 random initialization of the initial parameters \( \text{W}^{(v)}_{ab} \). We then chose the values of \( \sigma_{ab} \) and \( \text{W}^{(v)}_{ab} \) minimizing \( \text{Err} \).

To fit the way locomotion affects synaptic strengths, we sequentially evaluated models of increasing complexity, each of which was penalized by the L0 regularization penalty \( \lambda_1 n_{aw} \). We first evaluated equal weights in locomotion and stationary conditions, i.e., \( \text{W}^{(v)}_{ab} = \text{W}^{(v)}_{ab} \) and \( n_{aw} = 0 \); next, we fixed all but one of the synaptic weights \( \text{W}^{(v)}_{ab} \) (i.e., \( n_{aw} = 1 \)), and so on. We found the minimum error for each value of \( n_{aw} \), and selected between these using the penalized total error function \( \text{Err} \) (Figure S15A).

**DATA AND SOFTWARE AVAILABILITY**

The data that support the findings of this study are available from the corresponding authors upon request.
Supplemental Information

Vision and Locomotion Shape the Interactions between Neuron Types in Mouse Visual Cortex

Mario Dipoppa, Adam Ranson, Michael Krumin, Marius Pachitariu, Matteo Carandini, and Kenneth D. Harris
Supplementary Figure 1 (related to Figure 1). Identifying V1 with retinotopic mapping

A) Pseudocolor map of preferred horizontal (A_1) and vertical (A_2) position, ascertained by a periodic drifting and flickering bar (flicker frequency 2 Hz, speed = 0.8 deg/s) (Kalatsky and Stryker, 2003). See STAR Methods for further information.

B) Image of vasculature in the same region shown in A with iso-azimuth (B_1) and iso-elevation (B_2) contours after interpolation (see STAR Methods) of artefactual extreme values in A (e.g. blood vessels). Black square: field of view for 2-photon imaging experiments. Location in V1 is confirmed by visual inspecting the direction of retinotopic mapping (see STAR Methods).

C) 2-photon fluorescence mean image of GCaMP6m (C_1) and tdTomato (C_2) in this field of view. Scale bar: 100 μm.
Supplementary Figure 2 (related to Figures 1 and 2). Background fluorescence correction.

A1) For every cell, a “neuropil mask” was defined, extending 35 µm circularly from the cell’s center, and excluding any pixels belonging to detected neurons.

A2) Scatter plot of the raw signal vs. neuropil trace (1 point every 1s). Red line: fit of the lower 5% percentile of the scatter plot, showing a slope of $\alpha_i = 1.06$. For each experiment, we computed the mean correction factor $\alpha_i$ (excluding cells with low skewness), and used this to weight the neuropil signal subtracted from each cell’s fluorescence trace.

A3) Example raw signal trace (top), neuropil trace (center), and corrected signal trace (bottom).

B) Map showing mean GCaMP fluorescence level: (B1) GCaMP expressed in all neurons (same field of view as in Figure 1D1); (B2) GCaMP is expressed only in Sst neurons.

C) Maps showing correlation of running speed with fluorescence for each pixel, for the same fields of view shown in B1 and B2. Black contours represent outlines of Sst cells detected by tdTomato expression (c.f. B1 and Fig. 1A4, 1B4) or by GCaMP expression (c.f. B2). Note that while positive correlations are seen in regions where GCaMP is strongly expressed, negative correlations are seen in regions of low GCaMP expression, presumably reflecting hemodynamic filtering.

D, E) Correlation of fluorescence with running speed, as a function of baseline fluorescence level $F_0$, before (D) and after (E) neuropil subtraction for these two fields of view. Continuous black lines represent the average correlation in each plot. Note that prior to neuropil subtraction in the experiment with unconditional expression, a negative correlation is observed in cells of with very weak GCaMP expression, similarly to the surrounding neuropil.
Supplementary Figure 3 (related to Figure 1). Choosing the skewness threshold to identify putative excitatory cells.

A) Probability density distribution of skewness in mice where cells expressed GCaMP6m via virus injections (A1), GCaMP6s via virus injection (A2), or where GCaMP6s was expressed in a transgenic line (A3).

B) Receiver operating characteristic curve for Pyr classification in mice where cells expressed GCaMP6m via virus injections (B1), or GCaMP6s via virus injection (B2). Black circle represents values at skewness threshold = 2.7.

C) Skewness for each cell and experiment measured in mice where GCaMP6m was expressed by virus injection (see STAR Methods). Circles: Y-axis represents skewness for each cell with randomized position along the x-axis. Black triangles represent average skewness for each experiment.
Supplementary Figure 4 (related to Figure 2). Size tuning and locomotion modulation in putative Pyr cells identified by sparse firing.

A) Correlation of neural fluorescence with running speed as a function of cell depth. Circles indicate cells with significant speed correlations at p<0.05 (shuffle test), dots indicate cells of insignificant correlation. Dashed line corresponds to a linear fit of correlation vs. depth. Correlations were small on average ($\rho_{gray} = -0.01 \pm 0.01$; SE, n = 5,666).

B) Marginal histogram of correlation values for cells. Filled portions of bars indicate cells that are significantly modulated by speed. Correlations were significantly positive or negative in 19% and 30% of cells (p<.05, shuffle test). The number of cells significantly positively or negatively correlated with speed (p < 0.05) was significantly more than expected by chance (p < 10^{-16}, Fisher's combined probability test).

C) Mean size tuning curve averaged over all cells. Error bars correspond to standard error.

D) Scatter plots showing modulation by locomotion of responses to large stimuli (y-axis) and small stimuli (x-axis). Circles correspond to cells whose responses have a significant interaction between size and locomotion (p<0.05, two-way ANOVA over stimuli of diameter 5° and 60°); squares correspond to cells that did not have a significant interaction but did have a significant effect of locomotion; dots correspond to cells with no significant effect of locomotion.
Supplementary Figure 5 (related to Figure 2). Repeatability across experiments of dependence of $\rho_{\text{gray}}$ on depth.

A) Correlation coefficient of recorded cells with running speed, plotted vs. cell depth for different experiments. The scatter plots represent single experiment, each point representing a single cell, with correlation on the x-axis and depth between 0um and 400 $\mu$m on the y-axis. Circles represent cells with significant correlations at $p<0.05$ (shuffle test); dots represent cells with insignificant correlations. Dashed line represents fitted dependence of correlation vs. depth.

B) Histograms of slopes of fits computed in (A) for each experiment. Dark bars represent significant (robust regression, $p < 0.05$) dependence of $\rho_{\text{gray}}$ on depth; hollow bars indicate insignificant correlations.

C) Same as B) for each mouse, where cells of different experiments from the same mouse where grouped together.
Supplementary Figure 6 (related to Figure 2). Onset and offset of locomotion do not have an impact on correlation coefficients of each cell type’s activity with running speed, $\rho_{gray}$. Each plot shows a histogram of $\rho_{gray}$ for each cell type, after excluding transitions between locomotion and stationarity (0.5 s after locomotion onset; 1 s after locomotion offset), and also excluding periods of intermediate running speed (> 0 cm/s, < 2 cm/s). Solid bars indicate significant correlations at p<0.05 (shuffle test). Note the similarity to column 3 of Figure 2.
Supplementary Figure 7 (related to Figure 2). Correlation of neural activity with locomotion in darkness.
A) Histograms of correlations of neural activity with running speed in darkness. Filled portions of bars represent cells that are significantly modulated by speed (p < 0.05, shuffle test).
B) Correlation of neural activity with locomotion during gray screen presentation vs. in darkness. Black squares represent population averages. Filled black circle in B4 points to the example cell in (D,E).
C) Correlation coefficient of recorded cells in darkness with running speed, plotted vs. cell depth. Dashed line represents fitted dependence of correlation vs. depth. For Pvalb cells, we found a relationship between $\rho_{\text{dark}}$ and cortical depth in darkness (n = 361, p < 0.01, robust regression), similar to that with uniform gray screen stimulation. For other cell classes however the results differed to those with the uniform condition (Figure 2, column 2): we found a significant decrease of $\rho_{\text{dark}}$ as a function of cortical depth for Pyr (n = 3,693, p < 0.04, robust regression) and Sst (n = 278, p = 0.02, robust regression) cells while we found a non-significant change for Vip cells (n = 786, p = 0.44, robust regression). However when assessing the significance of the $c = \rho_{\text{dark}} / \text{depth}$ dependency, we did not find any significant effect across experiments (Pyr: p = 0.85, Pvalb: p = 0.32, Vip: 0.52, Sst: p = 0.39; t-test across experiments). Note that the cell number in (C) is smaller than that in (A,B) since for some experiment we had only an approximate information on cortical depth.
D) Fluorescence trace (D1) and running speed (D2) for an example recording in darkness.
E) Fluorescence trace (E3) and running speed (E4) for the same neuron in (D) when the screen was constantly gray.
Supplementary Figure 8 (related to Figure 4). Effect of eye movements on size tuning.
A) Image of the mouse’s eye. Yellow circles correspond to the position of the pupil center during retinotopy experiments.
B) Position of the pupil relative to the visual field. Symbols indicate pupil positions during trials in which the mouse was stationary (blue) and moving (red). Concentric circles indicate distances of 5 deg and 10 deg from average eye position (central dot). Scale bar: 10 deg.
C) Probability density for eye position relative to average position.
D) Visual responses to stimuli of 5 deg diameter, as a function of relative eye position.
E) Average tuning curves for centered (< 5 deg, black) and off-center (≥ 5 deg, red) eye position during locomotion (top) and stationary (bottom) conditions. Error bars correspond to standard error.
Supplementary Figure 9 (related to Figure 4). Diversity of size tuning between cells.

A1) Mean responses of a single Pyr neuron to grating stimuli of diameter 5° and 60°. Black curves show trial-averaged response in the stationary (dashed line) and locomotion (continuous line) conditions. Gray shaded region of each subpanel indicates the 1 s stimulus presentation period.

A2) Size tuning curves for this example cell. Solid line: locomotion; dashed line: stationary. Error bars correspond to standard error.

B) Same as (A) for another Pyr cell with different size tuning.

C,D) Same as (A) for two example Sst cells that showed diverse size tuning.
Supplementary Figure 10 (related to Figure 4). Repeatability of size-locomotion interaction. Every point represents a cell with different colors representing different experiments or mice.

A) Modulation of size tuning by speed. Columns represent experiments, with x-axis position within columns jittered for visibility. The Y-axis shows the interaction of locomotion with size tuning, defined as \( F(\text{loc.},\text{large}) - F(\text{stat.},\text{small}) - F(\text{small},\text{loc.}) + F(\text{small},\text{stat.}) \). Circles represent cells whose responses have a significant interaction between size and locomotion (multi-way ANOVA over stimuli of diameter 5° and 60° stimuli); dots represent cells with no significant effect of locomotion. Black squares represent average modulation for each experiment. Stars represent significant modulation across the population (\( p < 0.05 \), t-test).

B) Same as A) for each mouse, where cells of different experiments from the same mouse were grouped together.
Supplementary Figure 11 (related to Figure 4). Dependence of size tuning on stimulus centering.

A) Mean size tuning curve for centered cells (receptive field offset radius: 0°) during stationary (empty circles) and locomotion (filled circles) periods. Fitted model in Figure 6 is represented with dashed (stationary periods) and continuous (locomotion) lines. Panels 1 to 4: Pyr, Pvalb, Vip and Sst cells. Error bars correspond to standard error.

B) Same as (A) for off-center cells (receptive field offset radius: 20°).

C) Top: normalized firing rate maps as a function of the distance of the receptive field from the stimulus center (RF offset) and the stimulus size (stimulus diameter) during stationary periods. Bottom: fitted model to the stationary period data.

D) Same analysis as in (C) for locomotion periods.
Supplementary Figure 12 (related to Figure 4). Size tuning curves are similar in binocular and monocular zones in all cell classes. A) Size tuning curves averaged over experiments computed from calcium traces recorded in the monocular zone. Error bars correspond to standard error. B) Same as in (A) but calcium traces were recorded in the binocular zone.
Supplementary Figure 13 (related to Figure 4). Results are not substantially affected by spike deconvolution.  
A1) Calcium trace and A2) stimulus-triggered average of the calcium response of an example Pyr cell.  
B) Reconstructed firing rate (B1) and stimulus-triggered average of the spike response (B2) for the example Pyr cell in A.  
C,D) same as in A and B respectively for an example Sst neuron.  
E) Size tuning curves averaged over experiments computed from reconstructed spike traces. Error bars correspond to standard error.
Supplementary Figure 14 (related to Figure 5). Noise correlations are positive for most conditions and cell classes; correlation measures are robust to the skewness threshold used to identify putative Pyr cells.

A1) Summed visual responses of Pvalb population vs. Pyr population condition during stationary periods relative to the average visual response of the stimulus of each particular trials. Each circle represents the simultaneous normalized activity of the excitatory and Pvalb populations during one trial after removing the average response for each stimulus. Dashed line indicates linear regression estimate of signal correlation.

A2) Same as A1 during locomotion condition.

B) Same as A for Vip cells.

C) Same as A for Sst cells.

D1) Summary plots of noise correlations during stationary periods for all experiments. Error bars correspond to standard error.

D2) Same as D1 during locomotion.

E) Spontaneous, F) noise, and G) signal correlations computed as function of the skewness threshold used to identify putative Pyr neurons. Continuous lines represent mean across experiments, dashed lines represent 95% confidence bands. Colors represent correlations of Pyr population with Sst (red), Vip (green) and Pvalb (blue) populations. Skewness threshold = 0 corresponds to selecting all unlabeled cells as Pyr neurons (low false negative rate, high false positive), skewness threshold $\gg 1$ corresponds to selecting only (a subset) of Pyr neurons (low false positive rate but higher false negative). No major dependence on the threshold was observed.

H, I): The angles $\theta_1$ and $\theta_2$ summarizing the nature of nonlinear correlations $\theta_1$ and $\theta_2$ (as related to Figure 5), as functions of the skewness threshold. Again, no major dependence on threshold was observed.
Supplementary Figure 15 (related to Figure 6). Model selection.

A) Error of several competing models to fit the size tuning curves of Pyr cells. “H”, “E”, “V” and “S” represent models where the synaptic inputs from thalamic, Pyr, Vip or Sst cells respectively were modulated by locomotion; “Null” represents model where no weights were modulated. Models with subtractive (light gray bars) and divisive (dark gray bars) inhibition were tested. Star represents the optimal model. Models with divisive inhibition from Vip cells are not shown, but did not provide better performance.

B) Contours indicate regions of optimal model fit (0.1% percentile) in the 2d space of effective connection strengths (see STAR Methods), for stationary and locomotion conditions. Black points represent the optimal values chosen for the models in Figure 6.

C) Colormaps indicating regions of optimal model fit in the 2d space of spatial synaptic connections (see STAR Methods), for stationary and locomotion conditions. White circles represent the optimal values chosen for the models in Figure 6. For the synaptic input of Pvalb and Sst cells the error map was plotted in turn against two of the input spatial coefficients as axis $\sigma_{ab}$, $\sigma_{ac}$ while fixing the optimal values of the remaining input spatial coefficient $\sigma_{bc}$.
<table>
<thead>
<tr>
<th>Inputs to Pyr ($\alpha = E$)</th>
<th>Visual ($\beta = H$)</th>
<th>Sst ($\beta = S$) - subtractive</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\tilde{w}_{\alpha \beta}^{(0)}$ (stationary)</td>
<td>3.62</td>
<td>11.21</td>
</tr>
<tr>
<td>$\tilde{w}_{\alpha \beta}^{(1)}$ (locomotion)</td>
<td>8.52</td>
<td>7.14</td>
</tr>
<tr>
<td>$\sigma_{\alpha \beta}$ (Deg)</td>
<td>1.0</td>
<td>46.0</td>
</tr>
<tr>
<td>Inputs to Pvalb ($\alpha = P$)</td>
<td>Visual ($\beta = H$)</td>
<td>Pyr ($\beta = E$)</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>$\tilde{w}_{\alpha \beta}^{(0)}$ (stationary)</td>
<td>1.10</td>
<td>1.25</td>
</tr>
<tr>
<td>$\tilde{w}_{\alpha \beta}^{(1)}$ (locomotion)</td>
<td>1.86</td>
<td>1.25</td>
</tr>
<tr>
<td>$\sigma_{\alpha \beta}$ (Deg)</td>
<td>1.0</td>
<td>14.9</td>
</tr>
<tr>
<td>Inputs to Sst ($\alpha = S$)</td>
<td>Visual ($\beta = H$)</td>
<td>Pyr ($\beta = E$)</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>$w_{\alpha \beta}^{(0)}$ (stationary)</td>
<td>0.97</td>
<td>1.64</td>
</tr>
<tr>
<td>$w_{\alpha \beta}^{(1)}$ (locomotion)</td>
<td>1.77</td>
<td>1.64</td>
</tr>
<tr>
<td>$\sigma_{\alpha \beta}$ (Deg)</td>
<td>14.9</td>
<td>14.9</td>
</tr>
<tr>
<td>Inputs to Vip ($\alpha = V$)</td>
<td>Pyr ($\beta = E$)</td>
<td>Sst ($\beta = S$) - divisive</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>----------------------</td>
<td>----------------------</td>
</tr>
<tr>
<td>$w_{\alpha \beta}^{(0)}$ (stationary)</td>
<td>2.85</td>
<td>2.60</td>
</tr>
<tr>
<td>$w_{\alpha \beta}^{(1)}$ (locomotion)</td>
<td>2.85</td>
<td>2.60</td>
</tr>
<tr>
<td>$\sigma_{\alpha \beta}$ (Deg)</td>
<td>9.4</td>
<td>10.00</td>
</tr>
</tbody>
</table>

Supplementary Table 1 (related to Figure 6). Parameters of the model best fitting our experimental data. For the Pyr and Pvalb equations we estimated the effective connections strengths: $\tilde{w}_{EH}$ and $\tilde{w}_{ES}$ are given by $w_{EH}/(1 - w_{EE} + \mu w_{EP})$ and $w_{ES}/(1 - w_{EE} + \mu w_{EP})$, while $\tilde{w}_{PH}$, $\tilde{w}_{PE}$, and $\tilde{w}_{PS}$ are given by $w_{PH}/(1 + w_{PP})$, $w_{PE}/(1 + w_{PP})$, and $w_{PS}/(1 + w_{PP})$. Italics denote connection strengths that do not change between stationary and locomotion periods.