Sensory and Motor Systems

Orientation Tuning Depends on Spatial Frequency in Mouse Visual Cortex

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DOI: http://dx.doi.org/10.1523/ENEURO.0217-16.2016

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Abstract

The response properties of neurons to sensory stimuli have been used to identify their receptive fields and to functionally map sensory systems. In primary visual cortex, most neurons are selective to a particular orientation and spatial frequency of the visual stimulus. Using two-photon calcium imaging of neuronal populations from the primary visual cortex of mice, we have characterized the response properties of neurons to various orientations and spatial frequencies. Surprisingly, we found that the orientation selectivity of neurons actually depends on the spatial frequency of the stimulus. This dependence can be easily explained if one assumed spatially asymmetric Gabor-type receptive fields. We propose that receptive fields of neurons in layer 2/3 of visual cortex are indeed spatially asymmetric, and that this asymmetry could be used effectively by the visual system to encode natural scenes.

Key words: visual cortex; cortical tuning; orientation selectivity; spatial frequency; two-photon; calcium imaging; mouse; asymmetry; Gabor model

Significance Statement

In this manuscript, we show that the orientation selectivity of neurons in primary visual cortex of the mouse is highly dependent on the stimulus spatial frequency (SF). This dependence is demonstrated quantitatively by a decrease in the selectivity strength of cells in nonoptimum SF, and, more importantly, it is also demonstrated qualitatively by a shift in the preferred orientation of cells in nonoptimum SF. We show that a receptive-field model of a 2D asymmetric Gabor, rather than a symmetric one, can explain this surprising observation. Therefore, we propose that the receptive fields of neurons in layer 2/3 of mouse visual cortex are spatially asymmetric, and this asymmetry could be used effectively by the visual system to encode natural scenes.

Introduction

Neurons in the primary visual cortex (V1) have been traditionally described by their receptive field (RF) structure and their response characteristics (Hubel and Wiesel, 1962; Skottun et al., 1991). They are classified into two major groups, simple and complex, and exhibit spatially localized receptive fields that consist of distinct elongated...
ON and OFF subfields. In mouse visual cortex, ~75% of the orientation-selective neurons in layer 2/3 are classified as simple cells, showing response characteristics similar to simple cells in visual cortex of monkeys or cats (Niell and Stryker, 2008).

To capture the selectivity of a neuron to a certain feature of a visual stimulus (e.g., orientation, spatial frequency (SF), size, position, and speed), it is convenient to measure one-dimensional (1D) tuning curves that show the average response of a neuron to a specific feature value. Customary tuning models propose that the response strength of a neuron can be predicted based on the similarity between the optimal stimulus of a neuron and the given stimulus. From the 1D tuning curves, which describe the response behavior to one stimulus feature, one can calculate several parameters such as the preferred stimulus, the strength of the selectivity, or the width of the selectivity that quantifies the specificity level of the neuron. In visual cortex, for example, since neurons are highly responsive to lines and edges, such curves, which commonly characterize simple cells in V1, are the orientation–tuning and spatial frequency–tuning curves. However, reducing the complexity of the RF spatial structure to 1D tuning curves in order to study individual features, comes at a cost of losing information that might be critical for understanding the neuronal representation of sensory information. Here, we measured the population responses of neurons in L2/3 of mice V1 to drifting gratings that varied in both orientation and SF. We used in vivo two-photon Ca2+ imaging to measure evoked responses from hundreds of V1 neurons. Accordingly, we calculated a two-dimensional (2D) tuning matrix, and studied the relationship between orientation and SF selectivity. Then, we compared orientation–tuning curves in various SFs. First, we found that the orientation selectivity of a neuron depends strongly on the stimulus SF, such that when we presented gratings with higher or lower SFs than the optimum, the orientation selectivity was significantly reduced. In addition to a quantitative change in the selectivity strength of the neurons, we also observed a qualitative change in the preferred stimulus of the neuron. As we moved away from the optimal SF, to either lower or higher SFs, there was a significant shift in the preferred orientation (Pref) of the neurons. Dependence between orientation and SF selectivity of cells has been previously suggested in the visual cortex of primates and cats (Andrews and Pollen, 1979; Vidyasagar and Sigüenza, 1985; Webster and De Valois, 1985; Jones et al., 1987; Zhu et al., 2010).

In order to explain this dependence between orientation selectivity and SF, we used the common Gabor model (Gabor, 1946) to predict the neuronal response to various stimuli. A Gabor filter is a Gaussian modulated sinusoid, which well describes the receptive fields of simple cells and successfully models their responses (Marcelja, 1980; Field and Tolhurst, 1986; Jones and Palmer, 1987b). However, the classic Gabor model, even though it succeeds in predicting multiple neuronal responses, cannot capture the full variety and complexity of the visual system. And, indeed, we found that the classic mathematical description of a 2D symmetric Gabor model (with either odd or even amplitude symmetry) cannot account for our experimental findings. However, spatially modifying the classic model to introduce a 2D asymmetry by way of tilting the Gabor against its elongated axis generates a fundamental change in the response predictions, which qualitatively explains our experimental observations.

The modified Gabor model presented in this article can explain the response characteristics of a population of neurons and suggests that the receptive field of many cells in layer 2/3 of visual cortex of mice demonstrates a central asymmetry in its 2D spatial organization.

Materials and Methods

Animals

Animal handling and experimentation were performed in accordance with the National Institutes of Health and Columbia University institutional animal care guidelines. Animals of both sexes were used and were housed in a temperature-controlled environment on a 12 h light/dark cycle. We used a total of five mice, either WT or VIP-Cre crossed with LSL-tdTomato [postnatal day 40 (P40) to P80; The Jackson Laboratory].

Surgery

The mice were placed on a warming plate (37°C) and anesthetized with isoflurane (initially 2%, and reduced to 1–1.5% during surgery) administered via nose cone. A custom-made titanium head plate was attached to the skull using dental cement. Subsequently, a craniotomy (~1 × 1 mm) was made over the primary visual cortex (3.5–4.5 mm posterior to bregma, 2.3–2.7 mm lateral to midline; putative monocular region) using a dental drill (Osada). An ophthalmic ointment was applied on both eyes to protect the eyes from dehydration during surgery, and was removed during visual stimulation.

Dye loading

For bulk loading of cortical neurons, Oregon Green Bapta-1 (OGB-1) AM (Molecular Probes) was first mixed with 4 μl of pluronic acid (20% in DMSO) and further diluted in 35 μl of dye buffer (150 mM NaCl, 2.5 mM KCl, and 10 mM HEPES, pH 7.4). Sulforhodamine-101 (SR101; Molecular Probes) at a concentration of 50 μM was added to the solution to label astrocytes (Nimmerjahn et al., 2004). Animals were head fixed, and the dye was slowly pressure injected into the left visual cortex at a depth of ~130–200 μm below the dura surface (layer 2/3) at an angle of 30° through a patch pipette (outer diameter, ~1–2 μm) using a Picospritzer II. Two to four injections were carried out at 10 psi for 8 min, each under visual control of a two-photon imaging microscope [850 nm excitation; 10× water-immersion objective; 0.5 numerical aperture (NA); Olympus]. After dye injections, the exposed cortex was covered with agarose (1.5–2%; Sigma-Aldrich) and a cover glass (World Precision Instruments) to reduce brain motion. Data collection began 60–90 min after injections to ensure dye uptake across a large number of cells. During data collection, light anesthesia was maintained by isoflurane (0.8–0.9%) administered via nose
cone (KOPF Instruments). Heart rate, respiration, and oxygen saturation were monitored throughout the experiments using MouseOx (STARR Life Sciences Corp), and the respiration rate was used to monitor and control anesthesia levels.

Two-photon Ca²⁺ imaging

Imaging was performed with a two-photon moveable objective microscope (Sutter Instrument) and a mode-locked dispersion-precompensated Ti:sapphire laser (Chameleon Vision II, Coherent). Frames were scanned through a 20× (0.95 NA; Olympus) or 25× (1.05 NA, Olympus) water immersion objective. Laser intensity was controlled via Pockels cell (Conoptics) and ranged between 20 and 70 mW. Scanning and image acquisition were controlled using Mscan (4.07 frames/s for 512 × 512 pixels; Sutter Instrument). OGB-1 fluorescence was excited at 950 nm. Fluorescence changes collected with a 20× objective typically varied between 5% and 50%, and between 10% and 70% with a 25× objective. Emission was collected using green (535/50 nm) and red (610/75 nm) filters (Chroma) simultaneously on two photomultiplier tubes.

Visual stimulation

Visual stimuli were generated in MATLAB (RRID: SCR_001622) using Psychophysics toolbox (Brainard, 1997; Pelli, 1997; Kleiner et al., 2007) and displayed on a gamma-corrected LCD monitor (Dell; 19 inches, 60 Hz refresh rate) positioned 15 cm from the contralateral eye, at ~45° to the long axis of the animal (spanning ~114° vertical by ~140° horizontal of visual space). The presentation of visual stimuli was synchronized with image acquisition using Mscan (Sutter Instrument) and a routine written in MATLAB, such that each stimulus presentation was triggered on the beginning of frame acquisition. The actual time of stimulus presentation was detected with a silicon photodiode (Hamamatsu) attached to the bottom right corner of the screen.

We presented square-wave drifting gratings (100% contrast) for 670 ms, followed by 3–5 s of uniform gray background (the mean luminance of the gratings) plus a blank condition. The gratings orientation was perpendicular to the drift direction. Gratings were presented at 12 directions of motion (in 30° steps), three sets of four or five spatial frequencies: [0.02, 0.03, 0.04, 0.06 cycles per degree (cpd)], [0.01 0.02 0.04 0.06 cpd], or [0.01 0.02 0.04 0.08 0.16 cpd], and a temporal frequency of 1.5 Hz. All stimuli were block randomized and repeated 5–10 times, and the initial phase of the drifting gratings was kept constant across all trials. Stimuli were presented in a pseudorandom order, but time courses are shown after sorting (Fig. 1C).

Data analysis

Image analysis

All data analyses were performed using built-in and custom software written in MATLAB (MathWorks). Images were first converted to TIFF format and registered to correct for x-y motion using Turboreg plug-in in ImageJ (Thévenaz et al., 1998; RRID:SCR_014308). Regions of interest (ROIs) were drawn around each cell using a semi-automated algorithm based on fluorescence intensity (mean projection), fluorescence change (SD projections), and cell size and shape, and were adjusted by visual inspection. Glia cells were excluded from further analysis using SR101 staining, which selectively labels astrocytes in rodent neocortex (Nimmerjahn et al., 2004). Pixels were averaged within each ROI for each image frame. Baseline Ca²⁺ fluorescence was computed for each trial as the mean over 2 s prestimulus. Then, fluorescence values were converted to the percentage change above baseline [response amplitude (ΔF/F)] according to the following: ΔF/F = (F – F₀)/F, where F is the baseline fluorescence and F₀ is the instantaneous fluorescence signal averaged over two frames (~500 ms) following stimulus onset (F₀+2, F₀+3), where F₀ is the stimulus onset frame).

Responsiveness and reliability criteria were defined as previously described (Marshel et al., 2011). Briefly, neurons were considered responsive if their mean ΔF/F to any stimulus exceeds 6%. Reliability (δ) was determined according to the following:

$$\delta = \frac{\mu_{\text{max}} - \mu_{\text{blank}}}{\sigma_{\text{max}} - \sigma_{\text{blank}}}$$

where μ_max and σ_max are the mean and SD of the response to the preferred stimulus, respectively, and μ_blank and σ_blank are the mean and SD of the response to the blank stimulus, respectively. Neurons were considered reliable if δ > 1. Only cells that demonstrated visual responsiveness and reliability were chosen for further analysis, which excluded between 38% and 43% of the total number of cells we observed per field of view (FOV). Therefore, we analyzed 85.4 ± 3.6 reliably responsive cells of 156.2 ± 7.67 cells in total (mean ± SEM; n = 5 mice).

Visual tuning

To calculate tuning curves, we averaged the evoked responses (ΔF/F) over two frames following stimulus presentation. Then we averaged the response over the number of repetitions (5–10) per stimulus direction (12 directions). Direction tuning curves generated from OGB-1 fluorescence are comparable to those recorded with electrophysiological techniques (Kerlin et al., 2010; Marshel et al., 2011).

Orientation selectivity index (OSI) was computed as follows:

$$\text{OSI} = \frac{\mu_{\text{max}} - \mu_{\text{orth}}}{\mu_{\text{max}} + \mu_{\text{orth}}}$$

where μ_max is the mean response to the Pref and μ_orth is the mean response to the orthogonal orientation (Orth; average of both directions). Only cells that demonstrated an OSI ≥0.3 were chosen for tuning comparisons.

Orientation–tuning curves were fitted with the sum of two Gaussians of identical width, as follows:
Figure 1. *In vivo* imaging of visual-evoked responses of layer 2/3 neurons. **A**, Top, A two-photon image (maximum projection) from L2/3 neurons in mouse primary visual cortex, loaded with OGB-1. Scale bar, 100 μm. **B**, Example traces of Ca2+ signals from 10 cells during the presentation of drifting gratings with 4 different spatial frequencies and 12 directions. **C**, Ca2+ responses of two neurons, displayed as a matrix of all stimulus conditions. Columns indicate the direction of the gratings, and rows indicate their SF. Each trial is shown in gray (n = 7); average response across trials of a given stimulus is shown in black. **D**, Tuning matrix of eight cells (cells 1 and 2 are shown in C). Left, Response matrices evoked by each stimulus. Pixels intensity corresponds to the average ΔF/F over two frames poststimulus presentation and over seven repetitions. Right, Direction–tuning curves fitted with a double Gaussian, measured in various SFs. Colors correspond to SFs marked with arrows next to the matrices on the left.

\[
R(\theta) = \alpha_0 + \alpha_1 e^{-\frac{(\theta-\theta_0)^2}{2\sigma_1^2}} + \alpha_2 e^{-\frac{(\theta-\theta_1)^2}{2\sigma_2^2}},
\]

where \(R(\theta)\) is the averaged response to gratings with direction \(\theta\), and \(\alpha_0\) is the mean ΔF/F of the four lowest points in the curve. \(\alpha_1\) and \(\alpha_2\) are the amplitudes of the two Gaussians, \(\theta_0\) is the preferred direction, \(\theta_1\) is the null direction, and \(\sigma\) is the SD of the Gaussian function. The sum of two Gaussians fitting was constrained, according to Mazurek et al. (2014), by using several initial conditions, considering the fit with the lowest least square error as the best fit of the data. Peak responses were the maximum ΔF/F values of the 2D Gaussian fit curve. Half-width at half-height (HWHH) was computed as follows:

\[
HWHH = \sqrt{2\ln2}\sigma,
\]

where \(\sigma\) is the SD of the Gaussian function.

To further assess the statistical robustness of the tuning curve fitting, we applied a bootstrap method where we randomly resampled the data with replacement for each cell and obtained a distribution of response values. This procedure was repeated 100 times, where each repetition was fitted with a double Gaussian (Mazurek et al., 2014). This yielded a distribution of values for each of the tuning curve parameters. The mean values of the resampled data were then used for comparing population parameters showing the same statistical significance as achieved by comparing parameters of tuning curves obtained by fitting the data that included all the trials.

Additionally, orientation selectivity was assessed using a metric based on 1– circular variance (CirVar; Ringach et al., 2002), as follows:

\[
1 - \text{CirVar} = \frac{\sum R(\theta)\exp(2i\theta)}{\sum R(\theta)}
\]

where \(R(\theta)\) is the averaged response to gratings with orientation \(\theta\). This metric takes into account both tuning width and depth of modulation, and was found to be more reliable than extracting parameters following curve fitting, and more sensitive for detecting differences in selectivity between two populations (Piscopo et al., 2013; Mazurek et al., 2014).

2D Gabor model

The two-dimensional Gabor function is the product of a 2D sinusoid wave with a circular Gaussian envelope (see Fig. 3A), which was defined as follows:

\[
g(x, y) = \exp\left(-\frac{x^2 + y^2}{2\sigma_2^2}\right)\cos\left(2\pi\frac{x'}{\lambda} + \psi\right),
\]

where \(\sigma\) is the SD of the Gaussian envelope, which determines the size of the receptive field; \(\gamma\) is the spatial aspect ratio of the Gaussian, which determines the ellipticity of the Gabor (in a circular Gaussian \(\gamma = 1\); \(\lambda\) is the wavelength of the sinusoid; \(\theta\) is the orientation; and \(\psi\) is the phase offset.

In the tilted Gabor model, we introduced another parameter, which is the angle of the Gaussian tilt \(\phi\) with respect to the sinusoid wave. To generate a tilted Gabor filter, we first generated a symmetric 2D Gaussian with a spatial aspect ratio of \(\gamma = 0.5\) (see also Fig. 4B) and then applied a tilt by multiplying it with a 2D rotation matrix \(A\) with \(\phi = 30^\circ\), as follows:

\[
A = \begin{bmatrix}
\cos\phi & -\sin\phi \\
\sin\phi & \cos\phi
\end{bmatrix}.
\]

Calculating the predicted response

To calculate the predicted response, \(R(\theta)\), of a neuron, we computed the inner product between the RF \(l(x,y)\)—either a symmetric or a tilted Gabor filter, and the stimulus \(l(x,y)\)—square-wave gratings drifting in 180 different orientations with \(1^\circ\) interval (see Fig. 3B, examples). Since we measured neuronal responses by recording Ca2+ signals, we obtained tuning matrices by maximizing the averaged evoked response per stimulus. Therefore, to compute the predicted response, we calculated the inner product per stimulus orientation for each phase of the stimulus between 0 and \(2\pi\) (21 samples, \(\pi/10\) apart; see Fig. 5) and then maximized the response across phase.

Results

Layer 2/3 cells in mouse V1 show dependence of orientation selectivity on spatial frequency

In lightly anesthetized mice, we identified cells in layer 2/3 of visual cortex using two-photon Ca2+ imaging and monitored the activity of neuronal populations (Marshel et al., 2011; Miller et al., 2014) evoked by a brief visual stimulus presented to the contralateral visual field. We characterized the response dynamics by optically recording Ca2+ signals of OGB-1 from cells in upper layer 2/3 (depth, \(\sim 130–200\) μm; \(n = 5\) animals) in a typical FOV (Fig. 1A,B). Specifically, we measured spatial frequency–tuning and direction–tuning curves of each neuron in our FOV by averaging across trials the response to each stimulus (Fig. 1C,D). Stimuli were square-wave drifting gratings presented at 1.5 Hz for 670 ms (approximately one cycle) and varied across 12 directions and four or five SFs (see Materials and Methods).
Initial characterization of the stimulus-evoked responses of the local network showed dependence of the orientation selectivity of the cells on SF. We characterized this dependence by calculating orientation–tuning curves for each SF and then estimating six parameters derived from these curves: (1) response amplitude of the Pref; (2) response amplitude of the Orth; (3) OSI; (4) 1 − CirVar; (5) HWHH; and (6) the preferred orientation (see Materials and Methods). For each cell (85.4 ± 3.6 cells/animal, mean ± SEM), we compared its tuning curve at the preferred SF, with the tuning curves at SFs below (Fig. 2A) and above (Fig. 2B) the preferred SF.

Tuning curves at SFs below the preferred SF had by definition reduced responses to the preferred orientation (\(p < 10^{-10}, n = 67\) cells; Wilcoxon signed-rank test), but also showed increased responses to the orthogonal-to-preferred orientation (\(p < 0.001\), Wilcoxon signed-rank test), a decrease in OSI (\(p < 10^{-7}\), Wilcoxon signed-rank test), a decrease in 1 − CirVar (\(p < 10^{-6}\), Wilcoxon signed-rank test), and an increase in HWHH (\(p < 10^{-9}\), Wilcoxon signed-rank test; Fig. 2A). In addition, we observed that some cells had a significant shift in their preferred orientation (Fig 2A, right; since we sampled 12 directions between 0º and 360º, for this analysis we subtracted 180º from the preferred direction of cells that had preferred direction >180º). To quantify this shift, regardless of its sign, we calculated the absolute value of the \(\Delta\) in preferred orientation between the preferred SF and SF below the preferred. We observed a mean shift of a 22.9º ± 2.8º (mean ± SEM; Fig. 2C, left).

Tuning curves measured with SFs above the preferred SF had a decreased response to the preferred orientation (\(p < 10^{-11}, n = 73\) cells; Wilcoxon signed-rank test); and showed a slight but significant decrease in the orthogonal-to-preferred responses (\(p < 0.01\), a decrease
in OSI ($p < 0.01$, Wilcoxon signed-rank test), a decrease in $1 - \text{CirVar}$ ($p < 10^{-4}$, Wilcoxon signed-rank test), and a slight decrease in HWHH ($p < 0.05$, Wilcoxon signed-rank test). Here too, we found cells with a significant shift in their preferred orientation (Fig. 2A, right), and, indeed, the $\Delta$ in preferred orientation was $18.5 \pm 2.5^\circ$ (mean $\pm$ SEM; Fig. 2C, right). Further analysis of the change in the preferred orientation in nonoptimal SFs revealed a monotonic averaged shift from SF below the preferred orientation to SF above the preferred orientation, which was quantified as the averaged shift across cells shown in Figure 2, C and D, and exemplified in the tuning curves of single cells shown in Figure 1D. The population results shown in Figure 2, A and B, were consistent across animals. The preferred response was decreased, as expected, at lower and higher SFs ($\Delta 4.73 \pm 0.74\%$ and $\Delta 5.39 \pm 0.66\%$, respectively; $n = 5$ animals; 85 $\pm$ 3.6 cells; $t$ test, $p < 0.005$); the Orth response was increased and decreased at lower and higher SFs, respectively ($\Delta 0.94 \pm 0.31\%$ and $\Delta 1.32 \pm 0.67\%$, respectively; $t$ test, $p < 0.05$); the mean OSI was significantly decreased at lower and higher SFs ($\Delta 0.26 \pm 0.05\%$ and $\Delta 0.12 \pm 0.02\%$, respectively; $t$ test, $p < 0.01$); the mean ($1 - \text{ CirVar}$) was significantly decreased at lower and higher SFs ($\Delta 0.084 \pm 0.010\%$ and $\Delta 0.087 \pm 0.009\%$, respectively; $t$ test, $p < 0.01$); the HWHH was increased and decreased at lower and higher SFs, respectively ($\Delta 6.7 \pm 1.7\%$ and $\Delta 3.1 \pm 1.1\%$, respectively; $t$ test, $p < 0.05$), and the preferred orientation had a mean shift of $22.1 \pm 0.9^\circ$ at a lower SF and a $18.3 \pm 0.6^\circ$ shift at a higher SF (mean $\pm$ SEM; Fig. 2E).

A tilted Gabor model explains a shift in the preferred orientation at different SFs

To our knowledge, the observed shift in the preferred orientation as a function of SF has not been previously reported in mice, although it has been demonstrated in cats (Jones et al., 1987). Therefore, we further investi-
gated whether a known RF model of simple cells in V1 may explain this finding. To do so, we used a Gabor filter model with odd or even symmetry ($\theta = 0$, or $\theta = \pi/2$ respectively, where $\theta$ denotes the sine-wave phase; see Materials and Methods; Fig. 3A) to compute the predicted response of a neuron. A Gabor filter with odd symmetry consists of two side-by-side antagonistic regions of equal strength, whereas a Gabor filter with even symmetry demonstrates a central region flanked by two antagonistic regions of equal strength (Fig. 4A). The neuronal response is predicted based on the dot product between the Gabor filter and the stimulus (square-wave drifting gratings). Since we imaged $Ca^{2+}$ signals that have slow dynamics, we could not deduce the optimal phase that each neuron preferred. Therefore, as the experimental tuning curves were calculated based on the maximum evoked responses, we used in our predictions the maximal response across phase (Fig. 5).

We calculated the predicted response for stimuli with various orientations and SFs, and, accordingly, computed the orientation–tuning curves at different SFs (Fig. 3C) and examined the behavior of their parameters (Fig. 3D). First, we observed a significant difference between a Gabor filter with odd symmetry and a Gabor filter with even symmetry in the predicted responses to the orthogonal-to-preferred orientation. This difference also led to a difference in OSI by its definition and in 1/CirVar. Second, and more importantly, we noticed that a symmetric Gabor filter, regardless of its phase, could not explain a shift in the preferred orientation of a neuron with a change in SF (Fig. 3D). Therefore, we also simulated the predicted response by altering different symmetry components of the Gabor filter [e.g., the phase of the sine wave ($\theta$) and the spatial aspect ratio of the Gaussian envelope; $\gamma$; see Materials and Methods; also see Fig. 4A,B], but these alterations alone could not qualitatively explain a shift in the preferred orientation of the cells at different SFs.

Figure 4. Altering the Gabor model parameters for predicting neural responses. A, A predicted response tuning matrix computed as the inner product between the Gabor RF model shown in Figure 3 and stimuli of square-wave gratings with various orientations (1° interval) and seven SFs. Each pixel represents the inner product between the RF and the stimulus (maximized across phase). The row marked with a blue arrow on the right denotes the preferred SF and was taken as a reference for comparing other SFs. Right, Orientation–tuning curves at various SFs, color coded according to the arrows shown next to the predicted tuning matrix. Each panel corresponds to RF with $\theta = 0$, $\theta = \pi/6$, $\theta = \pi/3$, and $\theta = \pi/2$. a.u., Arbitrary units. B, Same as in A only for an RF with $\theta = \pi/2$ and various spatial aspect ratios ($\gamma$). Each panel corresponds to RF with $\gamma = 1$, $\gamma = 1.33$, $\gamma = 1.5$, and $\gamma = 2$. Note that these parameter alterations alone could not qualitatively explain a shift in the preferred orientation of the cells at different SFs.
traditional model could explain the unexpected dependence of orientation on SF. We introduced an asymmetry by way of tilting the Gaussian envelope along the elongated axis of the 2D sine wave (Fig. 6A) and generated a filter that demonstrates a displacement of one subfield relative to the other along the RF orientation axis. We refer to this model as a tilted Gabor. Again, we generated two types of tilted Gabors according to the phase of the sine wave (\(\pi/2\), or \(3\pi/2\)) and then predicted the neuronal responses accordingly (Fig. 6B). We examined the dependence of orientation–tuning curves on SF (Fig. 6C) and found that using this modified RF we can predict the behavior of all parameters: a change in the Orth response, a change in OSI, a change in CirVar, a change in WHWH, and most importantly, a shift in the preferred orientation of the neuron. Here too we observed a difference in the preferred orientation shift between RF phases, such that RF with \(\pi/2\) showed a monotonic decrease with the increase in SF, whereas a tilted Gabor with \(3\pi/2\) showed a nonmonotonic behavior (Fig. 6D).

Collectively, the modified tilted Gabor model better explains our experimental data and suggests an asymmetric 2D organization of the RF ON and OFF subfields.

**Discussion**

In this study, we measured neuronal responses from layer 2/3 of primary visual cortex of mice presented with drifting gratings of various orientations and SFs. We observed a unique dependence of orientation–tuning curves on SF, which suggests that the receptive field of some cells in mouse V1 present spatial asymmetry in their RF structural organization.

**Investigating tuning curves in populations of neurons using calcium imaging**

Both optical and electrical recording techniques enable the monitoring of activity from hundreds of neurons simultaneously. Therefore, to conduct feasible experiments, one cannot tailor the repertoire of stimuli to the optimal stimulus of each and every neuron. Therefore, the number and the complexity of the experimental stimuli are reduced. In the visual cortex, one common way to reduce the stimulus dimensionality is to measure 1D tuning curves, which are composed of neuronal responses evoked by presenting drifting gratings of various orientations (typically, 8 or 12). This is based on the assumption that the 2D spatial structure of the receptive field can account for a large fraction of the orientation selectivity of simple cells (Lampl et al., 2001). However, orientation–tuning curves measured with a small number of points are just a 1D reduction of the receptive field of a neuron, which has a more complex structure in a high-dimensional space. As a result of this dimensionality reduction, we not only "pay" a quantitative price of reduced
neural responses due to nonoptimal stimulus, but we also pay a qualitative price, manifested as a shift in the preferred orientation of the cells in nonoptimal SFs that likely arises from an asymmetric RF structure.

Cortical neurons have an asymmetric RF spatial structure

The RF structure of a neuron in visual cortex describes the organization of ON and OFF subfields in visual space, and accordingly explains which visual features a neuron is sensitive to. Simple cells in primate and cat cortex have been found to present mainly 1D symmetric organization (i.e., odd-phase or even-phase symmetry along the x-axis, orthogonal to the preferred orientation), although there are studies showing that cells do not demonstrate only odd or even symmetry but also exhibit, to a smaller extent, various phases, which breaks the 1D symmetry that exists between the relative strength of each subfield (Kulikowski and Bishop, 1981; Jones and Palmer, 1987a; Ringach, 2002).

However, breaking the amplitude symmetry alone cannot explain the behavior of the observed tuning parameters (Fig. 4). Only a 2D spatial asymmetry that introduces a shift in the relative location of the subfields can predict our experimental findings. In fact, the critical characteristic in predicting a shift in the preferred orientation is the displacement of one RF-subfield relative to the other along the orientation axis, such that it breaks the symmetry in the orthogonal axis. This 2D displacement can be easily modeled by introducing a tilt in the 2D Gaussian envelope that generates an asymmetric Gabor filter.

Traditionally, the Gabor filter has been proposed as a model to describe the receptive field of simple cells, and successfully predicts the responses of cortical neurons of monkeys and cats (Marcelja, 1980; Kulikowski et al., 1982; Daugman, 1985; Field and Tolhurst, 1986; Jones and Palmer, 1987b). The Gabor model is optimal in terms of minimizing the uncertainty associated with localizing a...
signal in both space and SF (Marcelja, 1980; Daugman, 1985), and therefore can create a sparse representation of natural images. The modified model we present here keeps the basic characteristics of the classic model and introduces one additional parameter that accounts for spatial asymmetry.

Although the tilted model succeeds in predicting the observed responses, there is still no experimental study in mice that mapped the RF of cells directly, and accordingly characterized asymmetry based on observations. However, a study in cats (Jones and Palmer, 1987a) showed that the 2D response profile of simple cells is not necessarily Cartesian separable due to a relative displacement of the subfields, and that within a given receptive field, subfields need not be the same length. In addition, visual inspection of examples of cells from studies conducted in mice also revealed some displacement between the ON and OFF subfields of the RF of cells (Bonin et al., 2011; Ko et al., 2013; Cossell et al., 2015). However, Bonin et al. (2011) used wavelet stimuli to map the RF of cells (Selesnick et al., 2005), which are spatially symmetric by nature, while Ko et al. (2013) and Cossell et al. (2015) presented natural images but used a regularized pseudo-inverse method to estimate the RF of cells (Smyth et al., 2003), which introduces a two-dimensional smoothness constraint on the RF and might bias the RF structure toward more symmetric organization.

**Functional role of asymmetry in neuronal representations**

Breaking the symmetry of the receptive field of neurons has been previously reported in the retina and the hippocampus. In the retina, asymmetry is observed between the sizes of the circular receptive fields of ON and OFF ganglion cells (Chichilnisky and Kalmar, 2002), or between the relative dendritic field size of parasol and midget cells (Dacey and Petersen, 1992), or in the asymmetric adaptation of ON and OFF ganglion cells to photopic (day) and scotopic (night) conditions (Pandarinath et al., 2010).

Hippocampal place cells exhibit an asymmetry in the spatial arrangement of their place fields. That is, as the animal moves through the receptive field of a cell, the firing rate modulates asymmetrically, where at the start of the place field, the firing rate is low, and at the end of the field, the firing rate is higher. The asymmetry increases as a function of familiarity (Mehta et al., 2000). This asymmetry also exists at the level of subthreshold excitatory inputs to place cells, and has been proposed to arise from a change in the balance between inhibitory inputs at the pyramidal cell soma and increases in dendritic excitation (Harvey et al., 2009). This place cell asymmetry represents a prospective coding scheme, where the increased firing skewness early in the receptive field signals upcoming place field center. The network mechanisms of this asymmetry however, are not completely known.

Furthermore, asymmetry in the visual cortex might play an important role in contour integration, which is in line with the gestalt principles of perceptual grouping (Metzger, 2006), specifically the law of “good continuation” (Field et al., 1993). Since neighboring neurons share RF subfields (Smith and Häusser, 2010), then the integration of local populations of neurons, some of which exhibit spatial RF asymmetry, may be the basis for cortical representation of spatial continuity in curved shapes.

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