Dynamics of spatial frequency tuning in mouse visual cortex

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THE STRUCTURAL AND FUNCTIONAL layout of the rodent visual system is similar in many aspects to that of carnivores or primates. However, there are some significant differences. Besides the general absence of a gyrated cerebral surface and laterally placed eyes, the rodent primary visual cortex lacks ocular dominance columns and is subdivided into small binocular and larger monocular regions (Hubener 2003; Tagawa et al. 2005; van Brussel et al. 2009). Moreover, orientation-tuned cells in mouse and rat V1 are not clustered in columns of similar orientation preference, as in cats and monkeys, but are randomly dispersed throughout V1 in a “salt-and-pepper pattern” (Mrsic-Flogel et al. 2007; Ohki et al. 2005). The lack of a columnar organization in V1 is not confined to nocturnal rodents such as rats and mice that do not rely on vision as much as primates for gaining information about their environment. Squirrels intensely rely on vision for their arboreal lifestyle yet equally lack orientation columns (Van Hooser et al. 2006; Van Hooser et al. 2005).

Despite these differences, neurons in rodent V1, and mouse V1 in particular, are highly tuned to a narrow range of stimulus parameters. For example, mouse V1 neurons show relatively sharp tuning for orientation, spatial frequency, temporal frequency, contrast, as well as stimulus length or size (Gao et al. 2010; Niell and Stryker 2008; Van den Bergh et al. 2010). While some aspects of neuronal responses, e.g., much larger receptive fields (RFs) and lower visual acuity, and a substantially lower level of surround suppression in mouse V1, can be different from those in primates, other tuning parameters are remarkably similar. The bandwidth for spatial frequency of neurons in mouse V1 when measured in octaves is similar to the spatial frequency bandwidth recorded in macaque or cat V1 (Gao et al. 2010; Van den Bergh et al. 2010), suggesting that similar circuitry in the visual system of rodents could be responsible for generating neuronal selectivity for spatial frequency.

Previous studies in cats and monkeys have shown that the spatial frequency tuning of V1 neurons changes over short time spans. V1 neurons are at first selective to lower spatial frequencies but gradually develop selectivity for higher spatial frequencies (Bredfeldt and Ringach 2002; Mazer et al. 2002; Nishimoto et al. 2005). This temporal shift in preferred spatial frequency can be attributed to an increase in the spatial frequency selectivity over time, generated by the suppression of lower spatial frequencies (Bredfeldt and Ringach 2002). While these spatial frequency tuning dynamics are clearly observable in the primary visual cortex, they might partly originate in the lateral geniculate nucleus (LGN) (Allen and Freeman 2006). In the wulst of the burrowing owl, a V1 analog in birds, similar temporal changes in spatial frequency tuning were observed without a parallel shift in preferred spatial frequency (Pinto and Baron 2010).

In an attempt to determine whether the spatial frequency tuning of individual neurons in mouse visual cortex depends on similar neuronal connectivity to that in macaque monkeys or cats, we analyzed the dynamics of spatial frequency tuning in mouse V1 using a similar reverse-correlation method to that used in monkey V1 (Bredfeldt and Ringach 2002). We also compared the dynamics of spatial frequency tuning between V1 and the extrastriate lateromedial (LM) field, which is a part of the lateral V2 complex (V2L) (Andermann et al. 2011; Marshel et al. 2011; Wang and Burkhhalter 2007; Wang et al. 2011). Our data show that, as in cats and monkeys, neurons in mouse visual cortex showed a clear shift in the preferred spatial frequency from low to high although this is not accompanied by an increase in tuning selectivity. There were no significant differences in the temporal spatial frequency tuning characteristics between V1 and LM in mice except for the strength of suppression.

MATERIALS AND METHODS

Subjects

Microelectrode recording experiments were conducted in anesthetized mice of the C57Bl/6 strain ranging in age from 3 to 5 mo (n = 9), housed under a conventional 13:11-h light-dark rhythm. All experiments were approved by the Ethical Committee of the univer-

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sity and were in strict accordance with the European Communities Council Directive of November 24th, 1986 (86/609/EEC).

Preparation

Mice were first given an intraperitoneal injection of the antipsychotic Chlorpromazine (5 μg/g body wt) to reduce the dose of urethane required to fully anesthetize the animals. Five minutes later, mice were anesthetized with urethane (1.2 mg/g body wt ip), the most common anesthesia used to study cortical physiology in mice, and, if necessary, additional doses of 0.2–0.3 mg/g body wt urethane were given to induce surgical anesthesia. A tracheotomy was performed, and a short plastic tube was inserted in the open end of the trachea, just below the larynx. A larger plastic tube blowing 95% oxygen-5% carbon dioxide was placed in front of the opening of the trachea tube enriching the inhaled room air with oxygen. The animal’s core temperature was kept at 37.5°C. The animal was mounted in a custom-built stereotaxic frame with the head held in place by two ear bars with fine tips (1.5 mm) and a mouth bar that fixed the upper incisors. Two electrocardiograph leads were inserted subcutaneously at the left and right side of the thorax to continuously monitor the heart rate, enabling us to assess the depth of anesthesia and the physiological condition of the animal, which is crucial for obtaining good visual responsiveness. The animal’s corneas were protected from drying out by regular application of silicon oil. With this procedure, we could maintain the reasonably good optical quality of the eyes. Because the nictitating membrane in mice does not block the cornea under anesthesia and the residual eye drift following urethane anesthesia (without paralysis) is negligible as expected from the large RF sizes in mice (Drager 1975; Gordon and Stryker 1996; Wagers et al. 1980; Wang and Burkhalter 2007), there was no need to employ special procedures to control these potential issues of their physiological optics. The ipsilateral eye was masked by a black cap over the eyeball, and responses were only recorded from the contralateral, right eye. A small (0.5 mm) craniotomy was performed over the central, binocular region of V1 of the left hemisphere. After the opening was made, the dura mater was locally removed using a sharp scalpel. The exposed brain was covered with a drop of mineral oil to prevent drying. Cardiac or respiratory pulsations of the brain surface could not be observed when making these small craniotomies.

Recording and Visual Stimulation

Tungsten-in-glass electrodes were used for isolating the activity of individual cortical neurons. The impedance of our electrode (tip impedance of ~1 or 9 MΩ) was higher than that of the silicon electrodes employed by Niell and Stryker (2008). We found that, with our electrodes, we could obtain both good isolation and excellent stability in mouse primary and extrastriate visual areas (as reported by many investigators in the visual cortex of higher species). The angle of the penetration was about 45° to the surface because the vertically oriented electrode entered the surface that had a slope of about 45° around the border between V1 and the lateral visual areas (Van den Bergh et al. 2010). This location of electrode penetration with respect to visual areas was determined according to the published map of V1 and its adjacent visual areas, response properties of neurons, and histological examination of electrode tracks (Kalatsky and Stryker 2003; Wagers et al. 1980; Wang and Burkhalter 2007).

Action potentials were extracellularly recorded and amplified using conventional methods. We searched for well-isolated units, and, if we encountered multiple units, we discriminated responses from these units using appropriately placed time-voltage windows. Our recordings started from the brain surface and ended either when the electrode entered the white matter or the animal’s deteriorated physiological situation precluded further recordings. Thus we typically sampled through all layers of cortex with intervals of at least 50 μm between subsequent cells. We determined RF positions as described earlier (Van den Bergh et al. 2010) by manually moving small circular optimized drifting gratings (10°) across the screen and locating the position where these gratings produced the strongest response. For mice, the center of the RF in the great majority of our units was located within 20° of the center of the visual field, with only a few units as far out as 60°. We did not dilate the pupil and did not determine the position of the optic disc to prevent a reduction in visual acuity. Because we could not use the optic disc as a reference point, we used the vertical plane through the midline of the animal as the vertical meridian, whereas the horizontal plane through the center of the eye was defined as the horizontal meridian. Recorded action potential sizes were digitized at 25 kHz, sampled at a rate of 60 Hz (16.7-ms bin widths for drifting grating experiments), and compiled into peristimulus time histograms that were equal in duration to and synchronized with the temporal cycle of the grating, by using data acquisition systems (System III; Tucker-Davis Technologies, Alachua, FL).

Visual stimuli were generated using custom-developed stimulation software using Matlab (The MathWorks, Natick, MA) and the Psychophysics Toolbox (Brainard 1997; Pelli 1997). Stimuli were presented on an LCD monitor (frame rate = 60 Hz, mean luminance = 27 cd/m², 1,280 × 800 pixels, 87° × 66°) at a distance of 25 cm from the eyes either perpendicular to the rostrocaudal axis of the animal for RFs at <30° from this axis, or at an angle of 45° for RFs at >30°. This limited distortion of the visual stimuli for RFs at higher azimuths.

RFs were first mapped on the tangent screen using moving dark edges. Optimal orientation and spatial frequency were quantitatively analyzed by showing drifting sinusoidal gratings that covered the entire monitor screen with a temporal frequency of 3 Hz and a contrast of 99%. Stimulus presentations were interleaved with blank gray screens of mean luminance to determine the spontaneous activity of the units. Spontaneous activity was subtracted from visually evoked responses before tuning curves were analyzed. In all tests, stimulus conditions were shown two times for 10 cycles (3 s at 3 Hz) in random order and interleaved with ≥3 s of gray background. For orientation tuning, 12 equally spaced orientations, 30° apart, were shown at near-optimal spatial frequency [generally around 0.02–0.06 cycles/degree (cpd)]. Spatial frequency was then analyzed at the optimal orientation by presenting drifting sinusoidal gratings with 11 different spatial frequencies ranging from 0.002 to 1.6 cpd. Stimuli were always presented monocularly to the eye contralateral of the hemisphere being sampled. Orientation and spatial frequency tuning were analyzed as described earlier (Van den Bergh et al. 2010).

Dynamics of Spatial Frequency Tuning

Dynamics of spatial frequency tuning were measured and analyzed using the reverse-correlation technique described by Bredfeldt and Ringach (2002). It is important to keep in mind that we employed the identical reverse-correlation method developed by Ringach et al. (1997) and extensively used in Bredfeld and Ringach (2002) to optimize the comparisons between macaque monkeys and mice with respect to dynamics of spatial frequency tuning in V1. Responses were recorded from individual cells of mice that were shown a rapid sequence (60 Hz) of luminance-modulated sinusoidal gratings with an orientation fixed at the optimal orientation of the cell, as measured using drifting sinusoidal gratings, and varying spatial frequencies and spatial phases. The luminance contrast (Michaelson contrast) was 99%. Spatial frequencies ranged from 0.0025 to 0.32 cpd in half-octave steps, encompassing most of the response ranges for spatial frequencies for all the cells we analyzed. Each of the spatial frequencies was shown for eight spatial phases, equally spaced and spanning 360°. Interleaved were blank gray images of uniform mean luminance that were shown to assess the cell’s baseline response. The probability of showing a blank image was equal to that of showing any one spatial frequency, irrespective of spatial phase. Stimuli were shown across a full screen to enable the stimulation of both the RF center and
surround. Each sequence consisted of 512 images drawn randomly from the image set of all spatial frequencies, spatial phases, and blanks and lasted ~9 s. Ninety sequences were shown with an interval of about 1.5 s, for a total experiment time of 15 min.

For each action potential the cell fired during visual stimulation, we recorded its arrival time. For each time lag \( \tau \), with 1 ms resolution, we determined the probability that a spike was preceded by a particular grating with a spatial frequency \( f \) at a particular time delay \( \tau \), independent of spatial phase: \( Pr(f, \tau) \). The probability that a blank image with uniform luminance preceded a spike at a specific delay time \( \tau \) was called the baseline \( B(\tau) \). Because the blank stimuli we used might drive the cells to a steady-state response that is above their spontaneous firing rates, this baseline response probability should more accurately reflect a maintained response level. The relative strength of the response to a given stimulus grating compared with the baseline (or maintained) response probability was then calculated:

\[
R(f, \tau) = \log_{10} \left( \frac{Pr(f, \tau)}{B(\tau)} \right)
\]

We analyzed the spatial frequency tuning dynamics by calculating \( R(f, \tau) \) at \( \tau \) values ranging from 0 to 200 ms. \( R(f, \tau) \) values of zero indicate that the cell’s response to a test stimulus was equal to that of the blank stimulus. Positive values indicate response enhancement while negative values correspond with suppression of the response (Fig. 1). Before \( \tau = 30 \) ms and after \( \tau = 150 \) ms, \( R(f, \tau) \) should be at zero, indicating that there is no increased response probability at these lag times, while between 30 and 150 ms increased or decreased response probabilities can occur.

To determine the time of response onset (\( \tau_{\text{onset}} \)) and decay (\( \tau_{\text{final}} \)), we looked at the variance of the response over time (Fig. 2A). For short time delays, before the signal has reached the visual cortex, the deviation of the response from the zero baseline is due to measure-

A Unit 1 (101 07 02)

B Unit 2 (090 11 01)

C Unit 3 (089 06 00)

Fig. 1. Examples of temporal dynamics of spatial frequency tuning for three representative neurons (A–C). The response tuning curves at different time delays are shown. The dashed lines indicate the baseline response level. Responses above this line are excitatory; responses below this level are suppressive.
defined was used to estimate the variance of the noise. For time delays with frequency octaves: the variance of the response \( V \) was defined as the peak of the smoothed tuning curve.

Because these maximum and minimum amplitudes vary over time, \( f_{\text{pk}} \) and \( f_{\text{min}} \) indicate the time lag on which the maximum amplitude of the response at the first 20 ms above the mean variance of noise (Bredfeldt and Ringach 2002).

The maximum and minimum amplitudes of the response were defined as

\[
M(x(\tau)) = \max_{\tau} \{R(f(\tau))\} \quad \text{and} \quad Mn(\tau) = \min_{\tau} \{R(f(\tau))\}
\]

respectively (Fig. 2B). Because these maximum and minimum amplitudes vary over time, \( F_{\text{pk}} \) and \( F_{\text{min}} \) indicate the time lag that showed strongest response enhancement or response suppression, respectively. We also defined \( F_{\text{pk}} \) and \( F_{\text{min}} \) as the time lag where \( M(x(\tau)) \) reached or decayed back to half its maximum amplitude \( R(f_{pk},/max) / 2 \).

For each time lag \( \tau \) we analyzed the spatial frequency tuning curves. To reduce the effects of noise, we smoothed the raw data by interpolating to 300 log-spaced data points and convolving with a Gaussian curve with \( \sigma \) of 0.4 log units. The peak of the spatial frequency tuning curve, the preferred spatial frequency \( f_{pk} \) at delay \( \tau \), was defined as the peak of the smoothed tuning curve.

To analyze the change of \( f_{pk} \) over time, we compared the peak spatial frequency \( f_{pk} \) over time by calculating the overall change in location of the preferred spatial frequency at \( F_{\text{pk}} \) and \( F_{\text{min}} \), in spatial frequency octaves:

\[
\Delta f_{pk} = \log 2 \left[ \frac{f_{pk}(\tau_{\text{decay}})}{f_{pk}(\tau_{\text{dev}})} \right]
\]

We also estimated the preferred spatial frequency of the cell for the time-averaged tuning curve:

\[
\bar{R}(f) = \frac{1}{(\tau_{\text{dev}} - \tau_{\text{onset}})} \int_{\tau_{\text{onset}}}^{\tau_{\text{dev}}} R(f, \tau) d\tau
\]

This time-averaged peak spatial frequency will be referred to as \( f_{pk} \).

To analyze spatial frequency tuning selectivity, we used the "quality factor" of the tuning curve (Bredfeldt and Ringach 2002), given by

\[
Q(\tau) = \frac{f_{pk}(\tau)}{f_{pk}(\tau - f_{low}(\tau) - f_{high}(\tau))}
\]

with \( f_{\text{high}}(\tau) \) and \( f_{\text{low}}(\tau) \) the high and low spatial frequency cut-offs, the spatial frequencies where the tuning curve drops below \( R(f_{pk},/sigma) / sqrt(2) \). \( f_{pk}(\tau) \) is the preferred spatial frequency at time \( \tau \) (Fig. 2C).

Cells with sharp spatial frequency tuning have a larger Q-factor, whereas cells with broad tuning have a Q-factor close to zero. Using the Q-factor to determine selectivity is advantageous over traditional methods such as spatial frequency bandwidth, since it allows for the analysis of low-pass-tuned responses, whereas spatial frequency bandwidth is undefined in this case (Bredfeldt and Ringach 2002).

To analyze changes in tuning selectivity over time, we define the change in selectivity as

\[
\Delta Q = Q_{\text{decay}} - Q_{\text{dev}}
\]

Positive values of \( \Delta Q \) indicate increased selectivity over time, whereas negative values indicate the cell became less selective over time.

To examine the steepness of the high and low spatial frequency limbs in our data, we use two indexes that allow us to separately analyze the low- and high-frequency flanks of the tuning curves:

\[
M_{L} = \frac{f_{low}}{f_{pk}} \quad \text{and} \quad M_{H} = \frac{f_{pk}}{f_{high}}
\]

An index close to zero indicates the spatial frequency limb has a shallow curve, whereas a steep curve is indicated by an index close to one.

In addition to the above selectivity measures that were used by Bredfeldt and Ringach (2002), we also computed an alternative and more global measure for tuning curve selectivity: low spatial frequency suppression (LSFS) (Xing et al. 2004) and analogously high spatial frequency suppression (HSFS). This corresponds, respectively, to the ratio between the response at the lowest spatial frequency tested and the response at the optimal spatial frequency (LSFS), and the ratio of the response at the highest spatial frequency tested and the response at the preferred spatial frequency (HSFS). The time-dependent changes in LSFS and HSFS were defined as follows:

\[
\Delta \text{LSFS} = \text{LSFS}(\tau_{\text{decay}}) - \text{LSFS}(\tau_{\text{dev}}) \quad \text{and} \quad \Delta \text{HSFS} = \text{HSFS}(\tau_{\text{decay}}) - \text{HSFS}(\tau_{\text{dev}})
\]
Strength of response suppression was measured as the ratio of the area of suppression vs. the total area beneath the curve, including enhancement and suppression over $\tau$ delay values ranging from $\tau_{\text{dev}}$ and $\tau_{\text{decay}}$, as well as from $\tau_{\text{decay}}$ to 150 ms:

$$\text{SupIndex} = \frac{\sum \tau A_S(\tau)}{\sum \tau A_S(\tau) + A_E(\tau)}$$

$A_S(\tau)$ and $A_E(\tau)$ are approximations of the excitatory and suppressive area under the curve.

The analysis of the phase dependence of the response enabled us to make a distinction between simple and complex cells, by calculating the FI-to-F0 ratio from the phase modulation at $\tau_{\text{max}}$. We found only six cells with simple cell-like characteristics (FI/F0 $\leq$ 1), which was not sufficient to make any reliable conclusions about possible differences in the spatial frequency tuning dynamics between simple and complex cells. Moreover, although we did not determine the optimal phase for simple cells but analyzed our data independent of spatial phase, this should not have affected our results because only six units were simple cells and our population thus primarily consisted of complex cells.

For neuronal response characteristics that showed normal distribution, we used standard parametric tests; otherwise, we used median values to evaluate these cellular properties and a Kruskal-Wallis rank test to test the significance of differences.

**Histology**

At the end of each penetration, two small electrolytic lesions (5 s, 10 $\mu$A, electrode negative) were made along the electrode track at intervals of $\sim$500 $\mu$m. The animals were given an overdose of pentobarbital sodium (Nembutal), intraperitoneally, and transcardially perfused with 1% paraformaldehyde in 0.15 M PBS (pH 7.4) followed by 4% paraformaldehyde in 0.15 M PBS (pH 7.4). Vibratome sections were cut and Nissl stained with thionine stain following standard procedures. Photographs of the histological patterns were obtained with a Zeiss Axio-Imager equipped with a Zeiss Axiocam. Brightness and contrast were adjusted using Adobe Photoshop CS2. Areal location and depth of the recording sites were reconstructed based on the positions of the electrolytic lesions and the recording depths along the electrode track as previously described (Van den Bergh et al. 2010). Because there appeared to be little layer-dependent differences in spatial frequency tuning dynamics and the variability of spatial frequency tuning properties within layers was generally as large as in the total cell population, we only placed our cells in subgroups based on areal location, but not based on depth or layers. Because the sample size for each layer was small, a larger sample would be required to make conclusive statements about layer differences.

All recording sites lateral of V1 were located in the lateral complex of visual areas, which is called lateral V2 (V2L) by Franklin and Paxinos (2008), recently described to comprise two different areas (Van der Gucht et al. 2007; Wang and Burkhelter 2007; Wang et al. 2011). The use of coronal Nissl-stained sections did not allow to determine the anterolateral (AL) and LM areas relative to V1 in cortical flat maps (Wang and Burkhelter 2007; Wang et al. 2011), the position of our electrode tracks along the anterior-posterior axis (between 2.0 and 3.6 mm posterior from bregma) and the fact that we were generally recording from the upper visual field in V1 before we crossed the V1/V2L complex border, we are confident that the recording sites outside V1 were all in LM and not in AL.

**Data Selection**

We recorded responses from 181 units from 9 mice. All cells in our analysis were stimulated with gratings oriented to within 30° of their preferred values. We excluded 66 units that did not show a significant response, i.e., where $V(\tau > 20)$ was less than 4 SD above the average variance of the noise ($[V(\tau \leq 20)]$. Twelve cells were excluded because their responses for the highest spatial frequencies were not at baseline. For two more cells, we could not get histological verification of cortical layer and visual area. A total of 101 cells passed these criteria and were thus included in the dataset comprising this study. The percentage of cells that passed these criteria (56%) was only slightly lower than the percentage of cells passing similar criteria in monkeys (61%) (Bredfeldt and Ringach 2002).

**RESULTS**

The goal of this study was to determine whether the development of spatial frequency tuning of mouse visual cortex neurons at a fine time scale was comparable to that of monkeys and cats. To this end, we quantitatively analyzed the temporal dynamics of spatial frequency tuning of 181 neurons in nine mice using a reverse-correlation method in the spatial frequency domain. For 101 cells, we obtained a significant response to the stimulus, 54 cells from V1 and 47 from LM. All units came from the binocular region of these areas, with most neurons located within 30° of the vertical meridian. The stimuli used for reverse correlation consisted of short movies of sine wave gratings of optimal orientation with spatial frequency and phase randomly changing at 60 Hz. We showed 15 different spatial frequencies, each at 8 different spatial phases. We averaged the reverse-correlated responses across all spatial phases.

Figure 1 shows the dynamic response of three representative cells. The response rate relative to the maintained discharge level, $R(f, \tau)$, is plotted for delays ranging from 23 to 114 ms. A dashed line at $R(f, \tau)$ equal to 0 indicates the response level to a blank stimulus of mean luminance. Positive values signify response enhancement to a specific spatial frequency, whereas negative values point to suppression below the response level of the blank stimulus. All three units showed a first observable response $R(f, \tau)$ at a delay time of around 30–35 ms. This response gradually increased in strength, reaching a maximum after 48–66 ms, followed by a gradual decrease in response, returning to the baseline level after about 80–90 ms.

All neurons showed spatial frequency selectivity, responding selectively to either very low spatial frequencies, with preferred spatial frequencies below 0.01 cpd (e.g., Fig. 1B, unit 2) or, as for most units, to preferred spatial frequencies between 0.01 and 0.1 cpd (e.g., Fig. 1, A and C, units 1 and 3).

Nonetheless, these tuning curves changed substantially over time. For two of the representative cells [units 1 and 2 (Figs. 1A, 1B, 1C, 3A, and 3D)], the preferred spatial frequency was not constant, but increased over time. Figure 1B (unit 2) shows that the unit initially was tuned at $\tau = 34$ ms to a very low spatial frequency (maximal response at an $f_{pk}$ of 0.0025 cpd). Over time, the spatial frequency with maximal response strength $f_{pk}$ shifted to 0.0085 cpd at $\tau = 82$ ms.

This shift in $f_{pk}$ was more clearly illustrated in Fig. 3, D and E, where we plotted $f_{pk}$ as a function of time delay $\tau$ between $\tau_{\text{dev}}$ and $\tau_{\text{decay}}$ (see Fig. 2 for definitions of these time points). The total shift of $f_{pk}$ during this period was 2.87 octaves. Another unit (unit 1, Figs. 1A, 3A, and 3B) was tuned to a higher preferred spatial frequency, and the preferred frequency also shifted over time. However, the shift was less pronounced, with $f_{pk}$ changing from 0.0278 to 0.0378 cpd between delay...
times \( \tau \) of 24 and 64 ms, resulting in a shift of 0.56 octaves between \( \tau_{\text{dev}} \) and \( \tau_{\text{decay}} \).

For both of these units, selectivity did not clearly increase over time (Fig. 3, A and D). The responses for spatial frequencies below \( f_{\text{pk}} \) visibly decreased at later delay times, but these responses were not obviously suppressed below baseline unlike in macaque monkeys (Bredfeldt and Ringach 2002). Some response suppression below baseline was present in these units, but, in general, this amounted to an overall decrease in response suppression at all or most spatial frequencies. In addition, it only occurred after the excitatory response again had reached baseline levels for all spatial frequencies (Figs. 1 and 3). Another unit (unit 3, Figs. 1C, 3G, and 3H) was tuned to a similar preferred spatial frequency as unit 1, but the tuning curve for this cell remained stable at all delays, with no observable shift in \( f_{\text{pk}} \) over time. Also here, selectivity did not change over time, neither was there increased response suppression at low spatial frequencies, except at very late \( \tau \).

**Population Data of Spatial Frequency Tuning**

**Preferred spatial frequency.** We measured the preferred spatial frequency \( f_{\text{pk}} \) from \( R(\tau, \tau_{\text{max}}) \) for both V1 and LM neurons. The distribution of the \( f_{\text{pk}} \) at \( \tau_{\text{max}} \) is shown in Fig. 4A. For most V1 and LM neurons, the \( f_{\text{pk}} \) ranged from 0.0025 to 0.06 cpd with some LM neurons reaching an \( f_{\text{pk}} \) at or above 0.1 cpd. The median \( f_{\text{pk}} \) at \( \tau_{\text{max}} \) for V1 (0.031 cpd) was not significantly different from LM (0.033 cpd) (Wilcoxon sign rank test; \( P = 0.96 \)).

To characterize the temporal dynamics of \( f_{\text{pk}} \), we measured the \( \Delta f_{\text{pk}} \), the difference in \( f_{\text{pk}} \) at \( \tau_{\text{dev}} \) and \( \tau_{\text{decay}} \). Figure 4, B–D, shows the distribution of \( \Delta f_{\text{pk}} \) for the V1 and LM cell populations. The scatter plot illustrates that the majority of the V1 and LM cells had higher preferred spatial frequencies at \( \tau_{\text{dev}} \) than at \( \tau_{\text{decay}} \); they lie above the unity line. The histogram shows the distribution of \( \Delta f_{\text{pk}} \) across the population of neurons. Positive values of \( \Delta f_{\text{pk}} \) indicate that \( f_{\text{pk}} \) shifted from lower to higher spatial frequencies between \( \tau_{\text{dev}} \) and \( \tau_{\text{decay}} \). Negative values correspond with decreased preferred spatial frequencies over the response delay. On average, \( f_{\text{pk}} \) changed positively (with average \( \Delta f_{\text{pk}} = 0.85 \) octaves in V1 and 0.61 octaves in LM), which is in the same range as the shifts observed in monkey V1 (Bredfeldt and Ringach 2002). The median values showed positive \( \Delta f_{\text{pk}} \) values as well (median \( \Delta f_{\text{pk}} = 0.52 \) octaves in V1, 0.47 octaves in LM), and most units had a higher preferred spatial frequency at \( \tau_{\text{decay}} \) than at \( \tau_{\text{dev}} \) (Paired Wilcoxon sign rank test, V1: \( P < 0.0001 \), LM: \( P = 0.02019 \)). We did not observe statistically significant differences in the changes of \( f_{\text{pk}} \) over time between V1 and LM neurons (Wilcoxon sign rank test, \( P = 0.57 \)).

Because the observed changes in \( f_{\text{pk}} \) were subtle and based on only two time points, we also determined the rate of change of the preferred spatial frequency. For each neuron, we plotted \( f_{\text{pk}} \) vs. \( \tau \) and fit a line to the \( f_{\text{pk}} \) values lying between \( \tau_{\text{dev}} \) and \( \tau_{\text{decay}} \). The distribution of the slopes, which provides the rate and direction of change of \( f_{\text{pk}} \) in V1 and LM, is shown in Fig. 5. Eighty-three percent (45/54) of the cells in V1 and 62% (27/48) of those in LM displayed earlier response latencies to low spatial frequencies while response latencies for higher spatial frequencies occurred later. The median slope was significantly different from zero in V1 (Wilcoxon sign rank test, \( P < 0.0001 \)) while, although the same trend was visible, it was only close to significance in LM (Wilcoxon sign rank test, \( P = \ldots \))
Spatial frequency selectivity. In macaque monkey, the shift in preferred spatial frequency over time is accompanied by an increase in spatial frequency tuning selectivity (Bredfeldt and Ringach 2002). To analyze whether the origin of the temporal increase in $f_{pk}$ of mouse visual cortex neurons is similar to that in macaque monkeys, we next analyzed the selectivity of spatial frequency tuning over time. To directly compare with the previously published data in macaque monkeys by Bredfeldt and Ringach, we used the Q-factor, the ratio of the peak spatial frequency over the difference between low and high cut-off frequencies, as a measure to estimate the spatial frequency tuning selectivity.

We measured the Q-factor for both $R(f, \tau_{max})$ and the time-averaged tuning curve $R(f)$. The selectivity was significantly higher for the mean tuning curve than at $\tau_{max}$, both in V1 and LM (Wilcoxon sign rank test, V1: $P = 0.0130$; LM: $P = 0.0009$). This was observed in monkey V1 as well (Bredfeldt and Ringach 2002). Figure 6 shows the distribution of the Q-factor of the time-averaged tuning curves and the tuning curve at $\tau_{max}$ for V1 and LM. For the mean tuning curve, the median selectivity was 0.73 for V1 neurons and 0.67 for LM neurons (Wilcoxon sign rank test: $P = 0.3302$), ranging from barely tuned (Q-factor of mean tuning curve = 0.11) to highly tuned [Q-factor $R(f) = 2.03$]. At $\tau_{max}$, selectivity was significantly higher in V1 than in LM (Wilcoxon sign rank test: $P = 0.0379$, V1: median Q = 0.64; LM: median Q = 0.55).

The representative cells shown in Figs. 1 and 3 demonstrated that, in these units, the Q-factor did not change much over time (Fig. 3, C, F, and I). Also, $\Delta Q$ (the difference between the Q-factor at $\tau_{dev}$ and $\tau_{decay}$) were very close to zero. To determine if the selectivity was unchanged for the cell population, we plotted the Q-factor at $\tau_{dev}$ and $\tau_{decay}$ (Fig. 7, A–D) for each unit. There was no trend toward an increase in spatial frequency selectivity over time for these neurons unlike in monkey V1. To the contrary, in the scatter plot, cells were located both above and below the unity line, and $\Delta Q$ values were distributed around zero. Most of the units did not show a large time-dependent change in selectivity, and, for cells whose selectivity changed, this change could go both toward a decrease or an increase in selectivity, resulting in the absence of significant net changes in selectivity (Wilcoxon paired sign rank test: V1: $P = 0.6115$; LM: $P = 0.6036$). We also did not...
observe differences in the distribution of $\Delta Q$ between V1 and LM neurons.

This apparent lack of a clear increase in tuning selectivity in mouse visual cortex neurons is contrary to what has been observed in macaque monkeys (e.g., Bredfeldt and Ringach 2002). In macaque monkeys, the increased preferred spatial frequency over time was shown to be related to an increase in tuning selectivity. Because there is also a minority of mouse units that showed a time-dependent decrease in $f_{pk}$ (see Fig. 4C) and thus have negative $\Delta f_{pk}$ values, we asked whether there is a correlation between the direction of changes in $f_{pk}$ and the changes in selectivity. We separately plotted the distribution of $\Delta Q$ for units with a negative $\Delta f_{pk}$ and for those with a positive $\Delta f_{pk}$ (Fig. 7, E and F). We would expect that, if the increase in $f_{pk}$ depended on increased selectivity, the $\Delta Q$ for units with a positive $\Delta f_{pk}$ would be positive as well, whereas the $\Delta Q$ for units with a time-dependent decrease of $f_{pk}$ would be negative. However, we found that both for units with negative as well as positive $\Delta f_{pk}$, the $\Delta Q$ was distributed around zero with an equal distribution to positive and negative $\Delta Q$ values. This was the case in both V1 and LM (Wilcoxon paired sign rank test: V1, $\Delta f_{pk} \leq 0$: $P = 0.8603$; V1, $\Delta f_{pk} > 0$: $P = 0.7416$; LM, $\Delta f_{pk} \leq 0$: $P = 0.8457$; LM, $\Delta f_{pk} > 0$: $P = 0.5911$).

An alternative way to look at changes in selectivity is to analyze time-dependent changes in the steepness of the flanks of the tuning curves. Figure 8 illustrates the distribution of the changes in steepness between $\tau_{dev}$ and $\tau_{decay}$ for the limbs of the tuning curve below ($\Delta M_{L}$) and above ($\Delta M_{H}$) the $f_{pk}$. Both $\Delta M_{L}$ and $\Delta M_{H}$ were distributed around zero, and the shape of their distribution histogram was similar as well, indicating that there is no bias to changed tuning curve steepness at either side of $f_{pk}$. This was in contrast to the increased curve steepness below $f_{pk}$ and the unchanged steepness above $f_{pk}$ in monkey (Bredfeldt and Ringach 2002). This was not surprising given the lack of changes in selectivity in mouse neurons that would occur when the steepness of the tuning curve would change over time (e.g., Bredfeldt and Ringach 2002 and Fig. 4D). There were no statistical differences between V1 and LM for these parameters.

Both the Q-factor and the steepness of the tuning curve flanks provide a selectivity measure that is limited to the central part of the curve. To obtain a more global selectivity measure that also could act as a time-dependent measure for the suppression of the response to spatial frequencies below $f_{pk}$, we calculated the difference in low spatial frequency suppression between $\tau_{decay}$ and $\tau_{dev}$ ($\Delta LSFS$). As a control, we also calculated the difference in high spatial frequency suppression ($\Delta HSFS$) to analyze possible suppression of the response at spatial frequencies higher than $f_{pk}$. Figure 9, A and B, shows the distribution for both $\Delta LSFS$ and $\Delta HSFS$ for V1 and LM neurons. From the distribution of $\Delta LSFS$, there was a marked reduction in the response to the lowest spatial frequency tested in this study compared with the response at $f_{pk}$. For V1, the median $\Delta LSFS$ was $-0.08$, with the LSFS at $\tau_{decay}$ and $\tau_{dev}$ significantly different (Wilcoxon paired sign rank test: $P = 0.0036$). In LM, $\Delta LSFS$ did not differ significantly from zero (median $\Delta LSFS = -0.03$, Wilcoxon paired sign rank test for LSFS at $\tau_{decay}$ and $\tau_{dev}$: $P = 0.7491$). This showed that, at least in V1, the response to the lowest extreme of tested spatial frequencies was markedly suppressed between $\tau_{dev}$ and $\tau_{decay}$. In contrast, high spatial frequency responses were markedly less suppressed at $\tau_{decay}$ than at $\tau_{dev}$ in V1 (median $\Delta HSFS = 0.13$, Wilcoxon paired sign rank test for HSFS at $\tau_{decay}$ and $\tau_{dev}$: $P < 0.0001$). Again, in LM, there was no difference between HSFS at $\tau_{decay}$ and $\tau_{dev}$ (Wilcoxon paired sign rank test: $P = 0.7973$; median $\Delta HSFS = 0.002$).

We separately plotted the $\Delta LSFS$ values for neurons with a decrease (Fig. 9C) and an increase (Fig. 9D) in the preferred spatial frequency over time. In V1, only for neurons with a positive $\Delta f_{pk}$ value, we could detect a significant increase in response suppression to the lowest spatial frequency (median $\Delta LSFS = -0.11$, Wilcoxon paired sign rank test for LSFS at $\tau_{decay}$ and $\tau_{dev}$: $P = 0.0030$). For units with a negative $\Delta f_{pk}$ or all units in LM, there were no significant deviations of $\Delta LSFS$ from zero.

Suppression. In monkeys, the increase in selectivity of the spatial frequency tuning curve over time was often accompanied by significant response suppression below the baseline for the spatial frequencies below the optimal (Bredfeldt and Ringach 2002). Indeed, Malone and Ringach (2008) demonstrated that inseparability was highest when enhancement and suppression were of roughly equivalent magnitude and that, when responses were dominated by enhancement without much suppression, the separability index was highest.

Response suppression below the baseline was therefore quantified by calculating the suppression index, the relative contribution of the suppression to the overall response. A value of 1.0 indicates that the response is purely suppressive; a value of 0 indicates complete absence of suppression; and a suppression index of 0.5 signifies that enhancement and suppression equally contribute to the response. While in monkey 69% of
the units with significant suppression had a mean suppression index of 0.27 (Bredfeldt and Ringach, 2002), we did not detect significant suppression during the response development between $\tau_{\text{dev}}$ and $\tau_{\text{decay}}$. The mean suppression index in V1 was 0.02 ± 0.05 and amounted to 0.01 ± 0.06 in LM (Fig. 10A).

Nevertheless, as shown by the representative cells in Fig. 1 as well as the minimum and maximum responses in Fig. 2B, about one-third of the units displayed suppression after the response had returned to baseline levels. Often, this suppressive phase encompassed all spatial frequencies tested. To quantify this general suppressive effect, we calculated the suppression index from $\tau_{\text{decay}}$ up to a delay of 150 ms (Fig. 10B). We observed strong suppression, with a mean suppression index for V1 of 0.46 ± 0.25 and 0.35 ± 28 in LM. There was a significant difference in suppression index between V1 and LM (Wilcoxon sign rank test: $P = 0.0491$).

**DISCUSSION**

The main finding of this study was that the spatial frequency tuning in mouse visual cortex was dynamic in the temporal domain. For the majority of neurons, the preferred spatial frequency was shifted toward higher spatial frequencies over the duration of their response. This shift in the preferred spatial frequency was not accompanied by an increase in tuning curve selectivity over time. Nevertheless, the response to the lowest spatial frequencies decreased over a short period of time relative to the response of the preferred spatial frequency. Also, there was no suppression below baseline levels. Instead, suppression only emerged after the tuning curve had returned to baseline levels. Another key finding was that there were no clear differences in inseparability of spatial frequency and time for the tuning curves for neurons in V1 and extrastriate region LM.

**Temporal Shift of Preferred Spatial Frequency**

The preferred spatial frequency measured in mice using the dynamic gratings and reverse-correlation method was rather low (around 0.031 cpd at $\tau_{\text{max}}$) compared with that of cat and monkey. However, it is very much in line with the preferred spatial frequency of mouse V1 neurons that was determined with drifting sinusoidal gratings. Recent studies established the median optimal spatial frequency in mouse V1 as being in the range from 0.03 cpd (Gao et al. 2010) and 0.032 cpd (Niell and Stryker 2008) to 0.04 cpd (Van den Bergh et al. 2010). Also in LM this corresponds with the data found in the literature [0.03 cpd vs. 0.04 cpd in V2L (Van den Bergh et al. 2010) and 0.028 cpd in LM (Marshel et al. 2011)].

As in cats and monkeys, the majority of V1 and LM neurons developed a time-dependent increase in the optimal spatial frequency. This shift in the peak spatial frequency of the tuning curves was qualitatively similar to that observed in earlier studies in monkeys (Bredfeldt and Ringach 2002; Frazor et al. 2004; Mazer et al. 2002) and cats (Frazor et al. 2004; Nishimoto et al. 2005). Importantly, the time-dependent shift in mice was comparable when quantified in octaves, which compensates for the much lower quality of vision in mice.

We were able to compare the magnitude of the shift in preferred spatial frequency with that estimated in Bredfeldt and Ringach (2002), since we employed essentially the same experimental paradigm and analysis methods as in their study. The only difference between the two studies was that we defined the delays of beginning and end of the response as $\tau_{\text{dev}}$ and $\tau_{\text{decay}}$, respectively, instead of $\tau_{\text{onset}}$ and $\tau_{\text{offset}}$. We decided on these delay times since $\tau_{\text{onset}}$ and $\tau_{\text{offset}}$, which are based on temporal variation in the tuning curve, were not clearly defined for several of the cells in our database, which had somewhat more variable responses for $\tau < 30$ ms. Because $\tau_{\text{dev}}$ and $\tau_{\text{decay}}$ are by definition the delays where the response reaches half-maximal values in contrast to $\tau_{\text{onset}}$ and $\tau_{\text{offset}}$ that delimit the beginning and end of the visual response, the total response time over which $\Delta f_{\text{pk}}$ was calculated was somewhat smaller than reported by Bredfeldt and Ringach (2002). Therefore, our procedure could result in a modest underestimation of the magnitude of change in $f_{\text{pk}}$ compared with that reported in...
monkeys. Nevertheless, the average shift in $f_{pk}$ of 0.85 octaves we observed in mouse V1 was similar but slightly higher than the value reported for monkeys (0.62 octaves).

Using a similar method, Mazer et al. (2002) detected only a small but significant shift in $f_{pk}$ in monkeys, which they did not quantify. They found that most cells had a positive rate of change of preferred spatial frequency over the response. Because they reported this rate of change as absolute values, we cannot make direct comparisons between mouse and monkey partly because of much lower acuity in mice. Nevertheless, the relative rate of the spatial frequency shift was similar in mouse V1, but less so compared with LM. Frazor et al. (2004), on the other hand, found a similar shift in $f_{pk}$ in cats (0.88 octaves) and monkeys (0.97 octaves), although they employed a markedly distinct method; 200-ms static gratings were shown with short interstimulus intervals, and then they analyzed the peristimulus time histograms across different spatial frequencies. This procedure eliminated potential overlap in the response across different stimuli. In another study with a reverse-correlation method to perform subspace mapping of the RFs in cats, the spatial frequency shift was visible (Nishimoto et al. 2005). The shift in $f_{pk}$ was analyzed both in area 17 and area 18, and the magnitude of the shift was somewhat smaller than in mouse: 0.22 and 0.24 octaves, respectively.

As in monkeys and cats, neurons in mouse visual cortex prefer coarser stimuli early during their response while finer stimuli preferentially elicit responses at a later time (Marr and Poggio 1979; Menz and Freeman 2004). As such, this temporal shift in spatial frequency preference (coarse-to-fine processing) might be a common property of visual cortex neurons of many mammals, even in those lacking orientation columns or having “low-quality” vision.

**Putative Mechanisms for Spatial Frequency Dynamics**

One of the most obvious differences between our results and those from monkeys was that we did not observe a time-dependent increase in the selectivity of the tuning curves in mouse. In monkeys, the change in the local selectivity measure over the response time (as estimated by $\Delta Q$) was positive for nearly all units, indicating a temporal increase in the sharpness of the tuning curves (Bredfeldt and Ringach 2002). Moreover, these investigators observed a rising steepness of the tuning curve over time below the peak spatial frequency and a time-dependent increase in suppression of the responses to lower spatial frequencies below baseline levels. These investigators concluded that the lagged suppression to low spatial frequency first sharpens the tuning curve and that, when the magnitude of the suppression increases relative to the magnitude of excitation, the increasing overlap of excitation and inhibition pushes the peak of the spatial frequency tuning curve toward higher frequencies (Bredfeldt and Ringach 2002). The absence of a local selectivity increase and a rise in the steepens...
ness of the tuning curve for the low spatial frequencies close to the preferred frequency in mice suggest that different mechanisms could be operating in mice for the observed dynamics of spatial frequency tuning.

An alternative explanation for the observed shift in peak spatial frequency in V1 considers the differences in the response properties of magnocellular (M) and parvocellular (P) neurons in the LGN. This feedforward model states that convergent input from M cells that fire faster than P cells but have RFs with twice the size than those of P cells could account for the latency differences as a function of spatial frequency in V1 (Frazor et al. 2004; Mazer et al. 2002). Because slow-conducting X and fast-conducting Y cells within cat LGN also provide inputs to area 17 differing in response latency that covaries with spatial frequency, this model could explain the observed dynamics of spatial frequency tuning in cat as well (Nishimoto et al. 2005). Although in mice no distinction between M and P, or X and Y LGN neurons has yet been established (Grubb and Thompson 2003; Huberman and Niell 2011), there are V1 neurons in mice with low spatial resolutions that prefer higher temporal frequencies while cells with high spatial resolutions prefer slowly moving stimuli, supporting the notion of “parallel” input channels into V1 (Gao et al. 2010). In one highly visual rodent, the squirrel, such parallel processing in LGN has been observed, with short-latency neurons having lower spatial resolution than longer-latency neurons (Van Hooser et al. 2003). This suggests that such parallel input into V1 might be a general feature of the visual system of rodents, and that of mice.

The key observation in our study that supports the above feedforward model comes from our analysis of tuning curve selectivity with a global selectivity measure, i.e., the LSFS (Xing et al. 2004). Here, we found, at least in V1, a small but significant increase in tuning curve selectivity due to a substantial reduction in response rates to the lowest spatial frequencies (Fig. 9). In the burrowing owl wulst, this shift in LSFS was also observed although it was accompanied by a decrease in tuning bandwidth but not by a change in the preferred spatial frequency (Pinto and Baron 2010). We did not analyze tuning bandwidth, since we knew from earlier studies that up to 30% of mouse units showed low pass tuning for spatial frequency (Van den Bergh et al. 2010) and would thus provide no useful metric for this property.

Another explanation for the shift in optimal spatial frequency in V1 was suggested by Allen and Freeman (2006). They demonstrated that spatial frequency tuning dynamics are present in the LGN and that a feedforward model could explain the observed dynamics at the cortical level. Because similar dynamics of spatial frequency tunings have not been studied in the mouse LGN, it is not possible to determine how much of what we observed in mouse V1 and LM can be explained by similar retinogeniculate mechanisms in the mouse visual system. However, despite similar shifts in optimal spatial frequency, the rate of change was lower in LM than in V1, suggesting that, apart from direct LGN input, at least some additional processing within cortex is required to observe these differences between cortical areas.

Suppression and Differences Between V1 and LM

It is unclear why we did not observe strong suppression for the lower spatial frequencies as reported in monkeys (Bredfeldt and Ringach 2002). We used a nearly identical reverse-correlation method where interleaved blank gray frames of average luminance were used to assess baseline responses. This allowed us to observe suppression of responses below the baseline after the excitatory responses had disappeared (Fig. 10B). However, we could not demonstrate strong suppressive effects while the excitatory response was still present (Fig. 10A) although we sporadically found neurons with weak suppression of spatial frequencies below the peak, concomitant with the excitatory response.

One possible explanation for the apparent lack of the “tuned suppression” in mouse V1 could be the use of urethane as an anesthetic/analgesic. Urethane tends to depress spontaneous and visually evoked cortical responses in rodents (e.g., Girman et al. 1999; Sceniak and Maciver 2006) although it is known to have little or no effect on the neurons’ excitatory or inhibitory synaptic inputs (Sceniak and Maciver 2006). Because the general loss of cell’s excitability occurs to the same degree in GABAergic and glutamatergic neurons, the use of urethane is unlikely to influence the excitation/inhibition balance. On the other hand, one of the significant consequences of reduced excitability, whether urethane induced or not, is an impoverished signal-to-noise ratio (SNR) of responses in mouse V1 neurons. If SNR in mouse V1 is substantially lower than that in macaque V1, the suppression for low spatial frequency could be “masked” or not readily “visible.” The visually evoked responses in mice V1 tend to be lower than those in macaque V1 while the maintained discharge is higher in mice (Niell and Stryker 2008; Van den Bergh et al. 2010). This means that “suppression” could be more easily demonstrated in mice, and
the variability of visually evoked spiking (fano factor) should be less in mice. Therefore, a potentially lower SNR in mouse V1 may not adequately explain the apparent absence of tuned suppression for low-spatial-frequency stimuli.

In this regard, the question remains as to why there is a global suppressive effect in mouse V1 only beyond the decay of the excitatory response with no local suppression of low spatial frequencies. One possibility is that, in mouse visual cortex, the response level for low spatial frequencies can still decrease due to weaker suppression, potentially giving rise to a small increase in preferred spatial frequency, but without reducing responses below baseline levels. Moreover, suppression of responses below the baseline might be “unmasked” only when the excitatory responses have already returned close to baseline levels. As a result, more global suppression at the end of the excitatory response emerges (Fig. 2). The broad tuning of the suppression at late time can arise from the broad tuning for spatial frequency found in mouse GABAergic neurons (Kerlin et al. 2010).

Related to these observations, we found differences in the strength of the late suppressive effect following the disappearance of the excitatory response between V1 and LM. In fact, this was one of the few clear-cut differences between the two visual areas in the diverse temporal properties of spatial frequency tuning we investigated. Our results did not show differences in the magnitude of the shift in preferred spatial frequency between V1 and LM. Similarly, Nishimoto et al. (2005) did not detect differences in the dynamics of the preferred spatial frequency between cat area 17 and 18.

An important question here would be why the delayed suppression we observed is stronger in V1 than in LM. We have previously found that RF surround suppression is stronger in V1 than in V2L. Moreover, the surround suppression in mouse V1 is significantly weaker than that observed in macaque V1 (Van den Bergh et al. 2010). Because surround suppression and the suppressive effects observed in the dynamics of spatial frequency tuning are likely to be supported by inhibitory cortical neurons (Angelucci et al. 2002; Bair et al. 2003; Bredfeldt and Ringach 2002), our data suggest that the general level of inhibition, either intracortical or as feedback from higher visual areas, is likely to be lower in LM than in V1. A relatively lower level of inhibition in mice vs. monkeys might be closely associated with the observed differences in the dynamics of spatial frequency tuning between the two species. Also, the difference between the “delayed” suppression in mouse V1 and LM suggests that the mechanism for this suppression occurs within these respective visual areas and does not originate from brain regions earlier in the visual pathway, such as the geniculate nucleus (e.g., Allen and Freeman 2006).

In this study, we demonstrated that, in mouse visual cortex, neurons display a time-dependent increase in preferred spatial frequency. The observed dynamics of spatial frequency tuning did not result from a significant increase of selectivity due to strong suppression of responses for low spatial frequencies as in macaque monkeys (Bredfeldt and Ringach 2002) but rather by a more subtle decrease in the response for low spatial frequencies. This temporal spatial frequency change in mice appears to be closely associated with weak suppression tuned to lower spatial frequency and/or the parallel feedforward input streams from the LGN that have shorter latencies for cells tuned toward lower spatial frequencies and longer latencies for neurons tuned to higher spatial frequencies. Taken together, our results suggest that there are subtle but significant differences between mice and macaque monkeys with respect to the cortical mechanisms underlying the dynamics of spatial frequency tuning of individual neurons.

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DISCLOSURES

No conflicts of interest, financial or otherwise, are declared by the authors.

AUTHOR CONTRIBUTIONS


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