Spatiotemporal specificity of contrast adaptation in mouse primary visual cortex

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Prolonged viewing of high contrast gratings alters perceived stimulus contrast, and produces characteristic changes in the contrast response functions of neurons in the primary visual cortex (V1). This is referred to as contrast adaptation. Although contrast adaptation has been well-studied, its underlying neural mechanisms are not well-understood. Therefore, we investigated contrast adaptation in mouse V1 with the goal of establishing a quantitative description of this phenomenon in a genetically manipulable animal model. One interesting aspect of contrast adaptation that has been observed both perceptually and in single unit studies is its specificity for the spatial and temporal characteristics of the stimulus. Therefore, in the present work we determined if the magnitude of contrast adaptation in mouse V1 neurons was dependent on the spatial frequency and temporal frequency of the adapting grating. We used protocols that were readily comparable with previous studies in cats and primates, and also a novel contrast ramp stimulus that characterized the spatial and temporal specificity of contrast adaptation simultaneously. Similar to previous work in higher mammals, we found that contrast adaptation was strongest when the spatial frequency and temporal frequency of the adapting grating matched the test stimulus. This suggests similar mechanisms underlying contrast adaptation across animal models and indicates that the rapidly advancing genetic tools available in mice could be used to provide insights into this phenomenon.

Keywords: adaptation, mouse vision, primary visual cortex, sinusoidal gratings, pattern-specificity, electrophysiology, context

INTRODUCTION

Our perception of the world around us, and the neural activity underlying this experience, is strongly dependent on the recent stimulus history. In the visual system, there is evidence for a number of self-calibration mechanisms that rapidly adapt visual processing according to the prevailing attributes of the stimulus being viewed (Carandini, 2000). Contrast adaptation has been used extensively to study this form of short-term plasticity. In psychophysical studies, prolonged viewing of a high-contrast pattern can produce a perceived fading of the adapting stimulus and reduce sensitivity to low contrasts, but it can also improve sensitivity and discrimination around the adapting contrast (Blakemore and Campbell, 1969; Greenlee and Heitger, 1988; Foley and Chen, 1997; Abbonizio et al., 2002). Primary visual cortex (V1) neurons have sigmoidal contrast response functions when spike rate is plotted as a function of stimulus contrast, and contrast adaptation has been shown to shift the most sensitive part of the curve toward the adapting contrast (Movshon and Lennie, 1979; Ohzawa et al., 1982, 1985; Sclar et al., 1989; Bonds, 1991; Ibbotson, 2005). A case has also been made that contrast adaptation (and similar processes) must be incorporated into models of V1 to better predict the responses of real neurons to natural stimuli (Carandini et al., 2005). Thus, there is converging evidence that contrast adaptation is a fundamental process that the visual system uses to make moment-to-moment adjustments in its sensitivity to incoming input.

Both psychophysical observations and single unit recording studies in V1 indicate that contrast adaptation is pattern-specific such that its magnitude can depend on the spatial frequency (SF), temporal frequency (TF), or orientation of the adapting and test stimuli (Blakemore et al., 1973; Vautin and Berkley, 1977; Movshon and Lennie, 1979; Albrecht et al., 1984; Ohzawa et al., 1985; Saul and Cynader, 1989a,b; Snowden and Hammett, 1996; Müller et al., 1999). This pattern-specificity has been used to constrain possible mechanisms underlying contrast adaptation. For example, both psychophysical and V1 data indicate that contrast adaptation is strongest when the SF of the adapting stimulus matches the test stimulus (psychophysics: Blakemore and Campbell, 1969; Blakemore and Nachmias, 1971; Blakemore et al., 1973; Snowden and Hammett, 1996; neurophysiology: Movshon and Lennie, 1979; Ohzawa et al., 1985; Saul and Cynader, 1989a), but this SF specificity must develop in the cortex because contrast adaptation in the lateral geniculate nucleus (LGN) does not appear to be SF specific (Duong and Freeman, 2007).

Several cellular and circuit mechanisms have been proposed to play a role in contrast adaptation (for a review see Kohn, 2007), but understanding of the cause of contrast adaptation remains incomplete. Several useful genetic tools available in mice could provide another avenue to explore contrast adaptation, but baseline conditions must first be established in this species to make any genetic manipulation related to contrast coding interpretable.
Several recent studies of mouse $V_1$ have revealed similarities between mice and higher mammals, including tuning for spatial and temporal frequencies, selectivity for orientation and direction, and the presence of simple and complex cells (Niell and Stryker, 2008; Gao et al., 2010; Van den Bergh et al., 2010). However, contrast adaptation in mouse $V_1$ has been reported in only two studies (Niell and Stryker, 2008; Stroud et al., 2012). Stroud et al. (2012) investigated the orientation specificity of contrast adaptation, but the other two aspects of the pattern-sensitivity of contrast adaptation that have been so important for linking electrophysiological studies in higher mammals to human psychophysical observations, namely specificity for SF and TF, remain unexplored. Therefore, we examined the spatiotemporal specificity of contrast adaptation in mouse $V_1$ using a top-up adaptation protocol that was comparable with previous studies in cat and monkey (Movshon and Lennie, 1979; Duong and Freeman, 2007; Dhruv et al., 2011). We also used dynamic contrast ramp stimuli of varying SF and TF to obtain rapid measures of contrast adaptation with a wide variety of adaptors (Crowder et al., 2008; Stroud et al., 2012).

Mouse $V_1$ neurons showed robust contrast adaptation when the adapting grating matched the neuron's preferred stimulus, which confirms earlier findings (Stroud et al., 2012). Furthermore, in the top-up protocol contrast adaptation was diminished or absent when the SF or TF of the adaptor did not match the neuron's preference, indicating that mouse $V_1$ neurons show adaptation specificity similar to that observed in cats and primates. Adaptation observed in the contrast ramp experiments was also pattern-selective, but maximal adaptation often occurred at slightly higher-than-preferred SFs, indicating that the exact properties of the contrast adaptation observed depends on the nature of the testing protocol.

**MATERIALS AND METHODS**

**ANESTHESIA AND SURGICAL PROCEDURES**

The experimental procedures reported herein conform to the guidelines established by the Canadian Council on Animal Care, which were approved by the University Committee on Laboratory Animals at Dalhousie University. Electrophysiological recordings were made from 25 adult male C57 BL/6 J mice weighing between 20 and 30 g, which were purchased from Jackson Laboratories (Bar Harbor, Maine). In early experiments, mice ($n = 15$) were sedated with chlorprothixene (5 mg/kg ip; Sigma, St. Louis, MO), and then anesthetized with urethane (0.5–1.2 g/kg ip; Sigma). If needed, a small dose of ketamine (20 mg/kg ip; Wyeth) was given to accelerate descent to the surgical plane of anesthesia, and allow a tracheotomy to be performed quickly (see Moldestad et al., 2009 for details). Mice were left free-breathing throughout the experiment and a tube located in front of the mouse delivered oxygen (0.1 L/min) to supplement room air. In later experiments, mice ($n = 10$) were sedated with chlorprothixene (5 mg/kg ip) and anesthetized with isoflurane delivered through a customized nose cone (2.5% during induction, 1.5% during surgery, and 0.4–1% during recording), which decreased preparation time by eliminating the need for a tracheotomy. Gas anesthesia did not appear to affect the frequency of encountering responsive units, and produced no significant differences in the tuning strength or selectivity of recorded units (assessed with discrimination indices, see Initial data analysis below; two-sample $t$-tests, $p > 0.2$ for all; c.f. Kaneko et al., 2012). For all mice, body temperature was maintained at 37.5$°$C with a heating pad, and their corneas were protected by frequent application of a thin layer of optically neutral silicone oil (30000 cSt; Sigma). The skull was stabilized in a stereotax, and a craniotomy ($\sim 1$ mm$^2$) was made over the monocular retinotopic representation in primary visual cortex ($\sim 0.8$ mm anterior and 2.3 mm lateral to lambda; Paxinos and Franklin, 2001). Recordings were made using either glass micropipettes (2–5 μm tip diameter, filled with 2 M NaCl or carbon-fiber in glass microelectrodes (0.6–1.5 Ω impedance). Electrode depth was controlled using a micromanipulator (FHC, Bowdoin, ME). Extracellular signals from individual units were amplified (Xcell 3+, FHC) and filtered (bandpass: 50–2000 Hz) before being digitized (Cambridge Electronic Design Power1401 with Spike2, Cambridge, England). Acquired signals were sampled at 40 kHz, and online analysis was performed on triggered TTL pulses with Spike2, but subsequent analysis was done offline.

**VISUAL STIMULI**

Upon isolation of a visually responsive unit, the receptive field (RF) was mapped using hand-driven light bars and spots. Quantitative testing was then performed with custom computer generated visual stimuli programmed in MatLab (MathWorks, Natick, MA) using the Psychophysics Toolbox extension (Brainard, 1997; Pelli, 1997), and presented on a calibrated CRT monitor (LG Flatron 915FT plus 19” display, 100 Hz refresh, 1024 × 768 pixels, mean luminance $30$ cd/m$^2$) at a viewing distance of 10–25 cm. All stimuli were presented in a circular aperture surrounded by a gray field of mean luminance. Orientation selectivity and surround suppression were characterized online using drifting square wave gratings. Spatiotemporal tuning was then assessed with full contrast drifting sine wave gratings with 36 combinations of SFs [0.01, 0.02, 0.04, 0.08, 0.16 and 0.32 cycles per degree (cpd)] and TFs (0.25, 0.5, 1, 2, 4, 8 Hz). All spatiotemporal and adapting stimuli were presented at the optimal orientation and size for each unit, and drifted in the direction that elicited maximal excitation. Presentations of each combination of SF and TF were randomized with 8–10 repeats for each stimulus. The presentation time of the stimulus was 1.5 s, and a gray of mean luminance was shown between stimuli for 0.5 s. Grating start-phase was staggered on each repetition to average out periodic firing of phase-sensitive neurons. The spatiotemporal tuning of each unit was then examined online and appropriate adaptors were selected for the subsequently presented contrast adaptation protocols. Two stimulus protocols that have previously been used to investigate contrast adaptation in mice, cats, and primates were modified to study the spatiotemporal specificity of contrast adaptation in mouse $V_1$: top-up adaptation (Sclar et al., 1989; Duong and Freeman, 2007; Stroud et al., 2012), and contrast ramps (Crowder et al., 2008; Stroud et al., 2012).

**Top-up adaptation**

We chose the top-up contrast adaptation protocol because it has commonly been used to study the stimulus specificity of contrast adaptation in higher mammals (e.g., Movshon and Lennie, 1979;
Duong and Freeman, 2007; Dhruv et al., 2011), which facilitates cross-species comparisons. Sine-wave contrast is defined as:

$$\text{Michelson contrast} = \frac{(\text{Luminance}_{\text{max}} - \text{Luminance}_{\text{min}})}{(\text{Luminance}_{\text{max}} + \text{Luminance}_{\text{min}})}$$  (1)

where \(\text{Luminance}_{\text{max}}\) and \(\text{Luminance}_{\text{min}}\) are the maximum and minimum luminances, respectively. Non-adapted contrast response functions were obtained by recording responses to ten contrasts (0.04, 0.08, 0.12, 0.16, 0.24, 0.32, 0.48, 0.64, 0.82, 1) presented in random order for 0.5 s tests (8–12 repetitions) interleaved with 4 s of mean luminance. Adapted contrast response functions were collected in blocks where: (1) the adapting grating matched the cell’s spatiotemporal peak; (2) the SF of the adapting grating was 1–3 octaves higher or lower than the cell’s preferred SF; and (3) the TF of the adapting grating was set to 8 Hz. Adaptation blocks consisted of 60 s of the adapting grating at a contrast of 0.32 followed by 0.5 s tests (aforementioned contrasts for 8–12 repetitions) interleaved with 4 s adaptation top-ups. An adapting contrast of 0.32 was chosen because our previous study of contrast adaptation in mouse V1 (Stroud et al., 2012) indicated that this contrast produced reliable adaptation while still allowing the data to be easily fit with sigmoid curves (see Curve Fitting below).

**Contrast ramps**

One drawback of the top-up protocol described above is that it takes a long time to record even a single adapted contrast response function (Sclar et al., 1989; Crowder et al., 2006). Therefore, when exploring the SF or TF specificity of contrast adaptation, only a few conditions can be examined for any single cell. To more fully assess the nature of contrast adaptation in the spatiotemporal domain, we used contrast ramp stimuli. Contrast ramps are dynamic contrast stimuli where the contrast of the sine wave grating is changed linearly on each animation frame over the time-course of the presentation. Importantly, these ramps are able to measure several key markers of contrast adaptation with fairly short presentation times (Crowder et al., 2008; Stroud et al., 2012). Contrast ramp stimuli were first presented at a contrast of 0, and contrast was increased linearly over 2 s until it reached 1 (rising phase). The contrast of the grating was then ramped back down from 1 to 0 (falling phase) over the next 2 s. Thus, the neuron is presented with identical contrasts in the rising and falling phases, but the order of presentation (i.e., temporal context) is reversed. A full screen gray of mean luminance was shown between ramp stimuli for 2 s. In this protocol, the spatiotemporal specificity of contrast adaptation was tested by varying the SF and TF of the contrast ramps using the 36 combinations of SFs (0.01, 0.02, 0.04, 0.08, 0.16, and 0.32 cpd) and TFs (0.25, 0.5, 1, 2, 4, 8 Hz) that were directly comparable with the spatiotemporal profile obtained for each neuron. Contrast ramps with different spatiotemporal combinations were randomized and repeated 8–12 times for each combination.

We were interested in determining whether the spatiotemporal combination that caused maximal firing also caused maximum hysteresis between the rising and falling portions of the contrast ramp. In order to test this, we used a symmetrical contrast ramp procedure, which maintained the same spatiotemporal parameters for both the rising and falling phase of the ramp. We also collected a second type of contrast ramp from a subset of neurons referred to as peak-tested contrast ramps that were more directly comparable to the top-up protocol. In the peak-tested protocol, the rising phase of the contrast ramp was one of the 36 combinations of SF and TF, but the falling phase was always shown at the neuron’s preferred SF and TF (see Results), as chosen from the online tuning function.

**INITIAL DATA ANALYSIS**

Spike sorting was performed offline with Spike2 software, which first searched for and sorted spikes using a supervised template-matching algorithm, and then displayed candidate spikes with a principle components analysis for approval. Data was exported to MatLab and neuronal responses were represented as spike density functions (SDF) with 1 kHz resolution generated by convolving a delta function at each spike arrival time with a Gaussian window. For each unit, we calculated the magnitude of orientation, size, and spatiotemporal tuning using a discrimination index (DI) (DeAngelis and Uka, 2003):

$$\text{DI} = \frac{(\text{Resp}_{\text{Max}} - \text{Resp}_{\text{Min}})}{\left( (\text{Resp}_{\text{Max}} - \text{Resp}_{\text{Min}}) + 2\sqrt{\text{SSE}}/(N - M) \right)}$$  (2)

Resp\(_{\text{Max}}\) is the neuron’s max response, while Resp\(_{\text{Min}}\) is the neuron’s minimum response. SSE is the sum of squared error of the mean, \(N\) is the total number of presentations of the stimuli, and \(M\) is the number of different stimuli presented. In order to classify cells as simple or complex, we divided the first Fourier coefficient of a neuron’s response to a grating near the spatiotemporal peak (\(F_1\)) by the mean time-averaged response to this grating (\(F_0\)) (Movshon et al., 1978a,b; Skottun et al., 1991). Despite some recent controversy (Mechler and Ringach, 2002; Crowder et al., 2007; Henry and Hawken, 2013; Hietanen et al., 2013), the \(F_1/F_0\) ratio has been used to quantitatively classify simple and complex cells in numerous studies, and an \(F_1/F_0\) ratio less than 1 indicates a cell is complex.

**Curve fitting**

We used the least squares method to fit contrast response functions. Sigmoid curves (Albrecht and Hamilton, 1982) were fit to the mean responses from top-up contrast response functions and SDFs produced by contrast ramps:

$$R(\text{c}_i) = \frac{R_{\text{max}} \times c_i^n}{c_i^n + c_{50}^n} + M$$  (3)

where \(R(\text{c}_i)\) is the amplitude of the evoked response at contrast \(c_i\), \(M\) is the spontaneous rate, \(n\) is the exponent that determines the steepness of the curve, \(R_{\text{max}}\) is the maximum elevation in response above the spontaneous rate, and \(c_{50}\) is the contrast that generates a response elevation of half \(R_{\text{max}}\). Response saturation was evident for almost all non-adapted top-up contrast response functions and rising ramp responses allowing for well constrained fits. When fitting adapted curves where the response to maximal
contrast was similar to or less than the non-adapted response but saturation was not evident, we assigned an upper bound on the adapted R\text{max} of 15% above the non-adapted R\text{max} in order to obtain tractable fits.

**Neuronal latency**

To examine the amount of hysteresis for each contrast ramp, responses were latency-corrected as previously described (Crowder et al., 2008; Stroud et al., 2012). Briefly, for each unit a response threshold was established based on the 99% cut-off from a Poisson distribution fitted to the spontaneous firing rate. Each unit’s response latency was calculated as the first time the spiking rate in the response to gratings of optimal SF and TF (from the spatiotemporal tuning stimulus) exceeded the aforementioned Poisson threshold and stayed above the threshold for the subsequent 25 ms (Price et al., 2005). For each unit, responses to contrast ramps were shifted back in time by the neural latency then split into the rising and falling phases and re-plotted using units of contrast on the abscissa instead of time (which resulted in the falling phases of contrast ramps being flipped left-to-right).

**RESULTS**

Recordings were collected from 188 visually responsive units in the primary visual cortex of 25 C57BL/6J mice. We obtained contrast adaptation data from 65 units using the top-up protocol, and 125 units using the ramp protocol (n = 90 for symmetrical contrast ramps; n = 35 for peak-tested contrast ramps). The stimulus preferences of units in our sample were generally consistent with previous reports. Discrimination indices for orientation selectivity (0.49 ± 0.1; mean ± s.d.), size tuning (0.62 ± 0.1), and spatiotemporal selectivity (0.64 ± 0.08) were similar to those reported by Gao et al. (2010). Peak SFs and TFs were broadly distributed, with preferred SFs ranging from 0.01 to 0.18 cpd (mean = 0.03 cpd) and preferred TFs ranging from 0.25 to 8 Hz (mean = 1.77 Hz). Figure 1A shows the grid-like array of responses used to measure the spatiotemporal tuning of a sample neuron, and Figure 1B shows how these responses can be summarized as a contour plot to indicate the combination of SF and TF that produced the maximal response. Our range of peak SFs and TFs were similar to recent electrophysiological studies of mouse visual cortex (Niell and Stryker, 2008; Gao et al., 2010; LeDue et al., 2012), and within the ranges shown by recent multi-photon calcium imaging studies (Andermann et al., 2011; Marshel et al., 2011). Finally, 157 units were classified as complex (F1/F0 ratio < 1) and 35 units were classified as simple (F1/F0 ratio > 1). Since simple and complex cells showed similar trends for all measures of contrast adaptation, they were pooled into a single group.

**TOP-UP CONTRAST ADAPTATION**

Robust contrast adaptation following prolonged exposure to an adaptor of the preferred SF and TF has been shown previously in mouse V1 (Stroud et al., 2012). However, to our knowledge the spatial and temporal frequency specificity of contrast adaptation in mouse V1 have not been explored. Therefore, we compared the magnitude of contrast adaptation induced by an adaptor with preferred SF and TF with that induced by an adaptor with non-preferred SF or TF. We chose a non-preferred adapting TF of 8 Hz because high TFs rarely elicited strong responses. This high TF adaptor also permitted comparisons with primate work, which has shown that high TFs can reliably induce contrast adaptation in V1 without strongly driving the recorded neurons (Dhruv et al., 2011), presumably by inducing adaptation in the LGN (Solomon et al., 2004). We selected non-preferred SFs 1–3 octaves higher or lower than the peak SF depending on the breadth and location of the recorded unit’s spatiotemporal tuning. Care was taken to ensure non-preferred SFs elicited weak responses from the recorded unit but also were within the range of peak SFs of our sample population and below the mean SF cutoff reported in previous studies of LGN and V1 (Grubb and Thompson, 2003; Gao et al., 2010). Higher adapting SFs were selected more often than lower ones since SFs lower than 0.01 cpd can begin to appear as global changes in luminance within the stimulus aperture.

Figures 1C–F shows the SF and TF specificity of contrast adaptation for four example neurons. Contrast response functions are shown for non-adapted (black squares), preferred adapted (red circles), non-preferred SF adapted (green triangles), and non-preferred TF adapted (blue stars) conditions. For the cell in Figure 1F, contrast response functions from two different non-preferred SFs (low SF = pink diamonds; high SF = green triangles) are shown. The spatiotemporal tuning of each unit is shown inset with the SF and TF values of the adapting stimuli indicated with matching symbols. In each case the preferred adaptor induced the most contrast adaptation. Non-preferred adaptors either induced virtually no adaptation (Figures 1C,D), or less adaptation than the preferred stimulus (Figures 1E,F).

Sigmoid fits to each contrast response function are shown as thin lines in Figure 1, and we used the c50 and R\text{max} parameters extracted from these fits to quantitatively analyze changes in contrast response functions following top-up adaptation. For each adaptation condition we measured the change from the non-adapted curve as a difference-over-sum calculation (parameter\text{shift} = [(adapted − non-adapted)/(adapted + non-adapted)]), and plotted this metric as population histograms in Figure 2. For Figures 2A,C,E positive values of c50-shift indicate a rightward shift in the adapted contrast response function. Nearly all cells showed a rightward shift following preferred adaptation (Figure 2A, mean c50-shift = 0.26), but the population was centered closer to zero for both adaptors with non-preferred SF (Figure 2C, mean c50-shift = 0.09) and TF (Figure 2E, mean c50-shift = 0.05). A One-Way ANOVA followed by a Tukey-Kramer post-hoc indicated that the preferred adaptation produced significantly larger values of c50-shift than the other two adaptation conditions [F(2, 169) = 27.54, p < 0.001], while non-preferred SF and TF c50-shift did not differ. For Figures 2B,D,F negative values of R\text{max-shift} indicate a decrease in firing to maximal contrast following adaptation. Most cells showed a modest decrease in R\text{max} following preferred adaptation (Figure 2B, mean R\text{max-shift} = −0.17), but the population was centered near zero for both adaptors with non-preferred SF (Figure 2D, mean R\text{max-shift} = 0.02) and TF (Figure 2F, mean R\text{max-shift} = 0.05). A One-Way ANOVA followed by a Tukey-Kramer post-hoc showed similar results to the c50 data, with preferred adaptation producing significantly more negative values of R\text{max-shift} than the other 2 adaptation conditions [F(2, 169) = 7.72, p < 0.001], while
FIGURE 1 | Spatiotemporal specificity of top-up contrast adaptation.
The spatiotemporal selectivity of a sample neuron is presented in (A), with
a grid of SDFs showing the neuron’s response to gratings with different
combinations of SF (columns) and TF (rows). A scale bar depicting time
vs. impulses per second (ips) is shown in the bottom right SDF (0.32 cpd
and 0.25 Hz). Mean responses from SDFs are summarized as a grayscale
contour plot in (B) (maximal firing shown in white), with SF on the
absissa and TF ordinate. (C–F) show contrast response functions from
top preferred neurons with contrast on the abscissa and mean response
rate on the ordinate. Non-adapted, preferred adapted, non-preferred SF
adapted, and non-preferred TF adapted responses are shown as black
squares, red circles, green triangles, and blue pentagrams, respectively. (F)
shows responses following an additional non-preferred SF adaptor as pink
diamonds. Error bars represent SEM, and thin lines represent best fits to a
sigmoid function (see Materials and Methods). Spatiotemporal tuning of
each sample neuron is shown in the corresponding inset with adapting
stimulus SF and TF indicated with symbols matched to contrast response
functions.

non-preferred SF and TF $R_{\text{max-shift}}$ did not differ. Another way of
quantifying the spatiotemporal specificity of contrast adaptation
is simply to rank order the adapted curves for each cell. Preferred
adaptation $c_{50}$ values were larger than $c_{50}$ values measured fol-
lowing non-preferred SF adaptation for 90% of cells, and non-
preferred TF adaptation for 92% of cells. Preferred adaptation $R_{\text{max}}$
values were smaller than $R_{\text{max}}$ values measured following non-preferred SF adaptation for 75% of cells, and non-preferred
TF adaptation for 70% of cells.

CONTRAST RAMP ADAPTATION
The top-up adaptation data above clearly demonstrates that the
magnitude of contrast adaptation in mouse V1 depends on the
adapting SF and TF, however, as noted in the Methods section
only a few adaptation conditions can be studied for any sin-
gle cell due to the time constraints imposed by this protocol.
Therefore, we used symmetrical contrast ramp stimuli to more
extensively map the spatiotemporal selectivity of contrast adap-
tation. Figure 3A shows the response of a representative neuron
to a contrast ramp of optimal SF and TF. Even though the ris-
ing and falling phases of the ramp stimulus are symmetrical,
the spiking response shows clear hysteresis. If this spiking response is
latency-corrected and re-plotted with contrast on the abscissa (see
Materials and Methods), the difference between the responses to
the rising (red) and falling (blue) phases of the contrast ramp is
accentuated further (Figure 3B). As in previous studies (Crowder
et al., 2008; Stroud et al., 2012), the SDFs were fit to sigmoid
curves (thin lines). The most useful parameter extracted from
the sigmoid fits was $c_{50}$, since it captured the rightward shift
in the contrast response function by comparing semi-saturation
contrasts of the rising (upward pointing arrowhead) and falling
phases (downward pointing arrowhead) of the contrast ramp. For
our sample ($n = 90$), $c_{50}$ values from the rising phase were almost always smaller than $c_{50}$ from the falling phase (Figure 3C), and this difference was significant ($p < 0.01$, paired $t$-test). This replicates earlier findings in cats (Crowder et al., 2008), and mice (Stroud et al., 2012).

To map the spatiotemporal specificity of contrast adaptation we measured the hysteresis of ramp responses when the SF and TF of the ramp grating were varied (for easy comparison to the spatiotemporal tuning also obtained for each neuron we used the same 36 combinations of SF and TF). This stimulus protocol examined the spatiotemporal specificity of contrast adaptation from a slightly different perspective than the top-up protocol. The top-up protocol measured whether the magnitude of contrast adaptation was affected if the adapting grating did not match the test grating, which emphasized the importance of the adapting stimulus. Symmetrical contrast ramps measured the combination of SF and TF that produced the most hysteresis, which emphasized the importance of the cell’s own preferred stimulus in determining the strength and specificity of the adaptation effect (Saul and Cynader, 1989a). Figure 4A shows a grid of SDF ramp responses from a sample cell, each with the same format as Figure 3B. This neuron had strong ramp responses with substantial hysteresis around 0.02–0.04 cpd and 1–2 Hz. Responses to lower TFs (~0.25 Hz) showed little hysteresis despite monotonic increases in firing with contrast, and the entire ramp response flattened out at the highest SFs and TFs. The former effect was observed in 81/90 neurons, indicating that diminished adaptation was not solely due to lack of responding. We wanted to summarize the pattern of hysteresis for each neuron as a contour plot, but $c_{50}$ values taken from sigmoid fits to SDFs were unreliable for spatiotemporal combinations away from the peak, so we measured adaptation by calculating the mean difference between the responses to rising and falling phases of the contrast ramps. During adaptation, the semi-saturation contrast of the falling phase ramp response shifts to higher values, causing the falling ramp response to be lower than the rising phase response at most contrasts. This method of analysis has been shown by Stroud et al. (2012) to capture the major features of contrast adaptation without relying on fitting the ramp SDFs to sigmoid functions. Figure 4B shows the contour plot for this neuron summarizing the magnitude of hysteresis evoked by each combination of SF and TF. The first feature to note is the clear peak around 0.04 cpd and 1–2 Hz. Likewise, 85 out of 90 units produced contour plots with an easily identifiable single peak that was at least four times higher than the level of hysteresis produced by the least effective ramp. This supports our earlier finding of spatiotemporal specificity of contrast adaptation using a different method. The second feature of the hysteresis contour plot that we were interested in was whether the combination of SF and TF that produced maximum hysteresis for contrast ramps matched the
neuron’s peak in the spatiotemporal domain tested with regular grating blocks (Figure 4C). For this neuron, the two contour plots look similar (R = 0.78 from a 2D correlation analysis), but the gratings that produced maximum hysteresis had a slightly higher SF than the gratings that produced maximum firing. Figure 5 shows two more example cells, one where the spatiotemporal locations of maximum firing and maximum hysteresis match quite closely (Figures 5A, B; R = 0.87), and another where spatiotemporal location of maximum hysteresis is at a higher SF and lower TF (Figures 5C, D; R = 0.51). Figure 5E plots the difference in peak locations from the two types of contour plots in the spatiotemporal domain for each cell as “hatpins” (n = 85), with the empty dots indicating the location of maximum hysteresis. No pattern was apparent, indicating that there was not one specific combination of SF and TF that universally induced maximal hysteresis across cells. Figure 5F normalized the data from Figure 5E by calculating the octave difference in SF and TF between each pair of peaks to show the location of maximal contrast hysteresis (empty dots) relative to each cell’s peak in spatiotemporal tuning (all normalized to 0). Although the differences were small (45% of cells had peaks within 1 octave of each other, and the median R-value from 2D correlations was 0.71), at the population level the gratings that produced maximum hysteresis tended to have slightly higher SFs (mean: 0.47 octaves) and lower TFs (mean = −0.22 octaves) than the gratings that produced maximum firing. A 2-way repeated measures ANOVA (with peaks from spatiotemporal tuning vs. hysteresis contour plots and SF vs. TF as factors) showed a significant main effect of peak type $[F_{(1, 84)} = 3.95, p < 0.05]$, indicating that the spatiotemporal location of peak hysteresis and peak firing tended to be different. Furthermore, a significant interaction between factors indicated that the difference between peaks was larger for SF than for TF $[F_{(1, 84)} = 15.75, p < 0.001]$

Considering that we consistently observed the strongest adaptation when using the preferred SF and TF in the top-up protocol we were surprised by the results of the ramp protocol. However, as noted above there was one key difference between adapting procedures: in the top-up protocol the adapting grating varied but the test gratings were always set at the preferred SF and TF, whereas in the symmetrical ramp protocol the SF and TF of the grating remained constant throughout the rising and falling phases of the contrast ramp. To determine whether the different adaptation effects observed between the top-up and contrast ramp protocols were due to switching the SF/TF between adapting and test stimuli or the dynamic nature of the contrast ramp stimuli we presented a subset of cells with a modified ramp protocol referred to as peak-tested ramps. For these peak-tested ramps, the rising phase could have any one of the 36 combinations of spatiotemporal frequencies (a proxy for the adapting gratings in the top-up protocol), but the falling phase was always shown at the neuron’s preferred spatiotemporal frequency (a proxy for the test gratings). For these stimuli, we compared the response to the falling phase at the spatiotemporal peak with the falling phase responses at every other spatiotemporal combination since these stimuli were identical with only the preceding rising phase differing (Figure 6A). We expected the difference to be large if no contrast adaptation occurred, or small if contrast adaptation did occur. Responses from a representative neuron are shown in Figures 6A–C. We again represented the spatiotemporal specificity of adaptation for each neuron as a contour plot (Figure 6B), compared the adaptation contour plot to each neuron’s spatiotemporal profile (Figure 6C), and calculated the octave difference in SF and TF between peaks for the population (Figure 6D). For peak-tested ramps, there were clear peaks in the hysteresis contour plots of every cell (e.g., Figure 6B). Importantly, the contrast hysteresis and spatiotemporal profile contour plots were much more similar using this protocol. For the sample neuron shown in Figures 6A–C the 2D correlation between contour plots was 0.96, and the population median was 0.86. Furthermore, the 2D correlations between contour plots for the peak-tested protocol were significantly higher than for the symmetrical ramp protocol ($p < 0.0001$; t-test). Mean octave differences in peak location between contour plots were $-0.08$ and $-0.06$ for SF and TF, respectively (Figure 6D). A Two-Way repeated measures ANOVA (with peaks
from spatiotemporal tuning vs. hysteresis contour plots and SF vs. TF as factors) indicated that neither the main effect of peak type \( F(1, 34) = 0.31, p > 0.57 \), nor the interaction between factors were significant \( F(1, 34) = 0.01, p > 0.92 \). Thus, switching the SF/TF between adapting and test stimuli appear to be the important difference between the top-up and symmetrical ramp protocols because when the contrast ramp stimulus was altered to more closely resemble the top-up protocol the adaptation effects also matched the top-up results more closely. Overall, each of the three adaptation protocols along with their differing methods of analysis demonstrated the spatiotemporal specificity of contrast adaptation in mouse V1, even though differences between protocols produced some subtle variations in the nature of the adaptation.

**DISCUSSION**

This study demonstrated that contrast adaptation in mouse V1 is specific in the spatiotemporal domain. The magnitude of contrast adaptation observed in single units was found to depend on
both the SF and TF of the adapting grating, and the nature of the adaptation effect could also be affected by the SF and TF of the test stimuli. The properties of contrast adaptation we observed were broadly similar to single unit studies in higher mammals (monkeys: Sclar et al., 1989; Dhruv et al., 2011; cats: Movshon and Lennie, 1979; Ohzawa et al., 1982, 1985; Saul and Cynader, 1989a,b; Bonds, 1991) and psychophysical data (e.g., Blakemore and Campbell, 1969). This suggests that contrast adaptation can be thought of as a general feature of the mammalian geniculostriate pathway along with other classical response properties (e.g., Niell and Stryker, 2008; Gao et al., 2010; Van den Bergh et al., 2010). Despite marked differences between animal models (frontal eyes vs. lateral eyes, nocturnal vs. diurnal, acuity that varies over several orders of magnitude), adaptation in mouse visual cortex appears to follow similar rules and is of similar complexity to higher mammals. We believe that these findings uphold the viability of the mouse model for studying vision, and support the validity of a multi-species approach for investigating cortical visual processing.

COMPARISON WITH PREVIOUS STUDIES
To our knowledge only two previous studies have investigated contrast adaptation in mouse V1 (Niell and Stryker, 2008; Stroud et al., 2012). Stroud et al. (2012) were able to make direct comparisons between adaptation in mouse and cat V1 neurons, and reported that most key features of contrast adaptation were similar between species. When adapted and tested with an optimal grating, adaptation shifted contrast response functions down and to the right. Moreover, contrast ramps produced relatively robust contrast adaptation given their brief presentation times. The current study is in agreement with these previous findings, so the Discussion will focus on the spatiotemporal specificity of contrast adaptation.

Several previous papers have used some version of the top-up protocol to investigate either the SF or TF dependence of contrast adaptation (e.g., Movshon and Lennie, 1979; Duong and Freeman, 2007; Dhruv et al., 2011), and are therefore readily comparable to our own top-up data. Within our top-up protocol, the test stimuli were always at the optimal SF and TF for each neuron, which is most similar to the stimuli used by Dhruv et al. (2011) in their study of TF and orientation specificity of contrast adaptation in macaque V1. For these two studies, one central question was whether an adapting stimulus that itself does not strongly drive the recorded neuron could induce adaptation. Another study examining the SF specificity of contrast adaptation in cat V1 used a complementary design where the test gratings were not at each neuron’s optimal SF, but rather at an SF that evoked approximately the same firing rate as the adapting grating (Movshon and Lennie, 1979). Regardless of these design differences, the general findings of these studies indicate that contrast adaptation is most robust when the parameters of the adapting grating are similar to the test grating. The current study extends this finding into a genetically tractable animal model where there exists an expanded toolbox to investigate the mechanisms underlying this specificity.

Comparing orientation specificity and spatiotemporal specificity of contrast adaptation in mouse V1 is also worthwhile. Most neurons in V1 adapt to any orientation, even ones that elicit low firing rates (Stroud et al., 2012), which contrasts with our current results in the spatiotemporal domain. This pattern of results could be produced if cortical adaptation mechanisms pooled over orientation (or the sharpness of tuning was diluted by non-oriented cells), but were at least somewhat selective in the spatiotemporal domain (Andermann et al., 2011; LeDue et al., 2012).

It has been shown that adapting gratings with a high TF can induce modest but reliable contrast adaptation in macaque V1 without strongly driving the recorded neuron (Dhruv et al., 2011), presumably by inducing adaptation in magnocellular cells in the LGN (Solomon et al., 2004). Therefore, we were somewhat surprised to observe only occasional adaptation to higher TFs in our data set. In their study of macaque V1, Dhruv et al. (2011) used an adaptor with a TF of 30–50 Hz, which was 2–3 times higher than the peak TF of LGN neurons (10–16 Hz: Derrington and Lennie, 1984; Hawken et al., 1996). For our top-up protocol, the high TF adaptor (8 Hz) was also approximately double the peak TF of mouse LGN neurons (3.8 Hz: Grubb and Thompson, 2003), yet we observed little consistent adaptation to this stimulus. We had also predicted that the peaks in the contrast hysteresis contour plots obtained with our contrast ramp stimuli may be skewed toward higher TFs, but this was not the case. We are unsure what underlies this apparent species difference, but as outlined in a model of multiple sources of adaptation used by Dhruv et al. (2011), it suggests that less adaptation is occurring in (or being inherited from) the LGN in mice. This observation, in conjunction with the finding that contrast adaptation in cat LGN does not show SF specificity (Duong and Freeman, 2007), provide two good reasons for future work to investigate adaptation in mouse LGN.

Finally, the contrast ramp stimuli we used in this study were quite unique and therefore less comparable to previous papers (although the staircase-like stimuli used by Bonds (1991) also measured hysteresis when contrasts were presented in an ordered manner). However, this data is relevant to a longstanding issue in the contrast adaptation literature. Vautin and Berkley (1977) were the first to discuss how the adaptation measured in a recorded neuron could arise from processes occurring within the cell itself (intrinsic) or be inherited from other neurons in the circuit/network (extrinsic). Saul and Cynader (1989a) suggested that intrinsic mechanisms may be more narrowly tuned since they depend on the cell’s own tuning, while the aforementioned model by Dhruv et al. (2011) specified that some extrinsic sources of adaptation should be broadly tuned for certain stimulus attributes like orientation. The main strength of the contrast ramp stimulus is that it can measure features of contrast adaptation on a relatively short time-scale, and this allows for a large stimulus-space to be explored in a reasonable amount of time. This seems ideal for exploring the putative differences in tuning between intrinsic and extrinsic sources of adaptation. In the current study, both the symmetrical and peak-tested contrast ramp protocols support the spatiotemporal specificity of contrast adaptation initially described using the top-up protocol despite the fact ramp stimuli measured adaptation on a different time scale and used different metrics. It would be interesting to obtain comparative data from higher mammals for these stimuli.
USING MOUSE MODELS TO STUDY CONTRAST ADAPTATION

Electrophysiological studies in various animal models additively suggest that contrast adaptation is initiated at pre-cortical stages (retina: reviewed in Demb, 2008; LGN: Sanches-Vives et al., 2000a; Solomon et al., 2004; Duong and Freeman, 2007), and then refined and strengthened in the cortex (Ohzawa et al., 1985; Carandini, 2000; Dhruv et al., 2011). Furthermore, hyperpolarization of the membrane potential has been associated with contrast adaptation (Carandini and Ferster, 1997; Sanches-Vives et al., 2000a,b). However, questions about the specific mechanisms involved, their relative contributions, and the stage each one is implemented remain unanswered. It seems that some of the investigative tools currently most readily applied in the mouse could provide insights into these cellular and circuit mechanisms. The same biochemical and genetic flexibility that has allowed the use of optogenetic modulation to attribute particular functions to genetically defined inhibitory neurons within mouse V1 (e.g., Andermann et al., 2012; Atallah et al., 2012), or allowed genetically encoded calcium-indicator proteins to explore the response properties of hundreds of visually responsive neurons simultaneously (e.g., Andermann et al., 2011), could also be used to explore the mechanisms underlying contrast adaptation. Moreover, if specific mechanisms are isolated they could be knocked-out or modulated in real-time to probe the perceptual relevance of contrast adaptation using psychophysical tasks developed for the mouse (e.g., Busse et al., 2011). The possibility of causally linking neural processing to contrast perception is especially intriguing considering psychophysical studies of the performance enhancement conferred by contrast adaptation have been somewhat equivocal (Barlow et al., 1976; Määttänen and Koenderink, 1991; Abbonizio et al., 2002; Kohn, 2007). We are hopeful that insights gleaned from the mouse model will be relevant to higher mammals because the properties of cortical contrast adaptation that have already been explored appear quite similar between species.

AUTHOR CONTRIBUTIONS

Nathan A. Crowder and Emily E. LeDue designed the study; Emily E. LeDue, Jillian L. King, Kurt R. Stover, and Nathan A. Crowder collected data; Emily E. LeDue, Jillian L. King, and Nathan A. Crowder analyzed the data; Nathan A. Crowder and Emily E. LeDue wrote the manuscript.

ACKNOWLEDGMENTS

This work was supported by the Natural Sciences and Engineering Research Council of Canada and the Canada Foundation for Innovation.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 11 July 2013; accepted: 12 September 2013; published online: 03 October 2013.

Citation: LeDue EE, King JL, Stover KR and Crowder NA (2013) Spatiotemporal specificity of contrast adaptation in mouse primary visual cortex. Front. Neural Circuits 7:154. doi: 10.3389/fncir.2013.00154

This article was submitted to the journal Frontiers in Neural Circuits.

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