Behavioral/Systems/Cognitive

The Suppressive Field of Neurons in Lateral Geniculate Nucleus

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The responses of neurons in lateral geniculate nucleus (LGN) exhibit powerful suppressive phenomena such as contrast saturation, size tuning, and masking. These phenomena cannot be explained by the classical center–surround receptive field and have been ascribed to a variety of mechanisms, including feedback from cortex. We asked whether these phenomena might all be explained by a single mechanism, contrast gain control, which is inherited from retina and possibly strengthened in thalamus. We formalized an intuitive model of retinal contrast gain control that explicitly predicts gain as a function of local contrast. In the model, the output of the receptive field is divided by the output of a suppressive field, which computes the local root-mean-square contrast. The model provides good fits to LGN responses to a variety of stimuli; with a single set of parameters, it captures saturation, size tuning, and masking. It also correctly predicts that responses to small stimuli grow proportionally with contrast: were it not for the suppressive field, LGN responses would be linear. We characterized the suppressive field and found that it is similar in size to the surround of the classical receptive field (which is eight times larger than commonly estimated), it is not selective for stimulus orientation, and it responds to a wide range of frequencies, including very low spatial frequencies and high temporal frequencies. The latter property is hardly consistent with feedback from cortex. These measurements thoroughly describe the visual properties of contrast gain control in LGN and provide a parsimonious explanation for disparate suppressive phenomena.

Key words: cat; gain control; receptive field; masking; adaptation; inhibition

Introduction

The visual responses of neurons in lateral geniculate nucleus (LGN) reveal suppressive phenomena that cannot be explained by the classical center–surround receptive field. First, responses saturate when stimulus contrast increases [contrast saturation (Derrington and Lennie, 1984; Chino and Kaplan, 1988; Sclar et al., 1990; Kremers et al., 2001; Alitto and Usrey, 2004)]. Second, responses to a stimulus are reduced by superimposition of another [masking (Cudeiro and Sillito, 1996; Felisberti and Derrington, 1999; Freeman et al., 2002; Solomon et al., 2002)]. Third, responses decrease when stimulus size is increased beyond an optimal value [size tuning (Cleland et al., 1983a; Murphy and Sillito, 1987; Jones and Sillito, 1991; Solomon et al., 2002; Ozeki et al., 2004)]. These three phenomena are suppressive, because responses are smaller than would be expected from the classical center–surround receptive field alone.

There is little agreement as to the physiological basis of these suppressive phenomena. Some have ascribed suppression to inhibitory signals within thalamus (Hubel and Wiesel, 1961; Levick et al., 1972; Cleland et al., 1983a; Sclar, 1987; Felisberti and Derrington, 1999). Others attributed it to cortical feedback (Murphy and Sillito, 1987; Przybyszewski et al., 2000; Webb et al., 2002; Alitto and Usrey, 2003).

Moreover, there is little agreement as to how the suppressive phenomena should be explained functionally: how should the receptive field model be extended to account for response suppression? Contrast saturation has been described by a compressive relationship between stimulus intensity and cell output (Derrington and Lennie, 1984; Sclar et al., 1990; Felisberti and Derrington, 1999; Przybyszewski et al., 2000; Kremers et al., 2001; Solomon et al., 2002; Webb et al., 2002; Alitto and Usrey, 2004). Masking has been described by a divisive gain control mechanism (Freeman et al., 2002; Solomon et al., 2002). Size tuning has been attributed to a large subtractive region surrounding the receptive field (Solomon et al., 2002; Nolt et al., 2004; Ozeki et al., 2004). These proposals aim to account for one suppressive phenomenon at a time and not for all suppressive phenomena at once. Moreover, they fail to explain known relationships between suppressive phenomena. For example, size tuning depends on contrast, being strong at high contrast and absent at low contrast (Solomon et al., 2002; Ozeki et al., 2004).

We suggest that the suppressive phenomena are parsimoniously explained by a single mechanism of contrast gain control, which operates in retina and is possibly strengthened in thalamus. Indeed, the stimulus manipulations that elicit the suppressive phenomena invariably increase local contrast. Increasing lo-
Model. As illustrated in Figure 1, the model contains two pathways, one linear and one nonlinear.

The linear pathway involves a classical center (ctr) – surround (srd) receptive field, a difference-of-Gaussians, $RF = G_{ctr} - k_{srd}G_{srd}$, where $G_{ctr}$ and $G_{srd}$ are two-dimensional Gaussian densities of width $\sigma_{ctr}$ and $\sigma_{srd}$ and $k_{srd}$ is the relative weight of the surround. The receptive field is convolved with the stimulus $S(x,y,t)$ to produce a linear response:

$$L(t) = [S \ast RF](0,0,t).$$

For our stimuli, which consist of the sum of a “test” and a “mask” drifting gratings, $L(t)$ is the sum of the responses to the individual gratings:

$$L(t) = L_{test}(t) + \sigma_{mask}L_{mask}(t),$$

where $\sigma_{mask}$ denotes the relative effectiveness of the temporal frequency of the mask in driving the receptive field.

The nonlinear pathway involves a suppressive field that computes the SD of local luminance $c_{local}$. Assuming that the stimulus $S(x,y,t)$ has a mean of zero, local contrast follows

$$c_{local} = \sqrt{\int \int S^2(x,y,t)G_{mask}(x,y)dx dy dt},$$

where $G_{mask}$ denotes the suppressive field, a Gaussian of width $\sigma_{SD}$, and $S(x,y,t)$ denotes the stimulus processed through a bank of filters, $\{S + H(x,y,t)\}$. We model each of these filters as a difference-of-Gaussians: $H = G_{ctr} - k_{srd}G_{srd}$.

The suppressive field controls neural gain by dividing the output of the receptive field. The result is

$$V(t) = \frac{V_{max}L(t)}{c_{local} + c_{local}},$$

where $c_{local}$ determines the strength of the suppressive field, and $V_{max}$ captures the overall responsiveness of the neuron.

Firing rate is a rectified version of $V$, with threshold $V_0$:

$$R(t) = [V(t) - V_0]^+.$$

Model characterization. We fitted the model onto the harmonic components of the spike trains at the temporal frequencies of the stimuli. Let $\{f_i\}$ denote the spike times of a response to a stimulus of temporal frequency $f_i$; the harmonic component follows $r = 1/N\sum_{x=0}^{N-1} \exp[-2\pi if_i x]$, where $[\|x\|]$ denotes the modulus of the complex number $x$, and $N$ is the number of spikes.

Fits minimize the square error between responses of neuron and model $\sum_{i=0}^{N-1} (r_i - m_i)^2$, where $r_i$ denotes the response of a neuron to trial $i$ of stimulus $j$, and $m_i$ is the response predicted by the model.

To estimate model parameters, we fitted Equations 1 – 5 to a sequence of four experiments, each constraining one or more parameters. The first experiment constrained the parameters of the receptive field $\{\sigma_{ctr}, \sigma_{srd}, k_{srd}, k_{srd}\}$ and involved drifting gratings varying in spatial frequency. Gratings had 50% contrast, optimal temporal frequency, >20° in diameter, and one of >14 logistically spaced spatial frequencies. The subsequent three experiments constrained the parameters of the suppressive field $\{c_{local}, \sigma_{SD}, \sigma_{SD}, \sigma_{SD}, k_{srd}, \sigma_{SD}\}$ and involved sums of a test grating and a mask grating, in which we varied mask contrast (see Fig. 3A, D), mask diameter (see Fig. 3B, E), and mask spatial frequency (see Fig. 3C, F). Test diameter, spatial frequency and temporal frequency were optimized cell by cell to elicit maximal response. Unless varied, the attributes of the mask were also optimal for the cell. Each experiment included test gratings presented alone at more than six contrasts and sums of a 50% contrast test and a mask of variable attributes. Parameter estimates obtained when fitting one experiment were held fixed in fits to subsequent experiments. We repeated this sequence of fits until parameters changed by <1% since the previous sequence iteration. The result was a single set of parameters used to predict responses to all experiments.

Parameters $V_{max}$ and $V_0$ were allowed to vary across experiments to account for slow changes in neural responsiveness and spontaneous ac-

cal contrast decreases the responsiveness (gain) of retinal ganglion cells (Shapley and Victor, 1978, 1981; Victor, 1987; Baccus and Meister, 2002). This effect is thought to be enhanced in LGN (Kaplan et al., 1987; Sclar, 1987; Cheng et al., 1995). It could potentially explain the suppressive phenomena.

Materials and Methods

We characterized the responses of 34 well isolated neurons recorded in LGN of eight anesthetized, paralyzed cats. These neurons were held long enough (>2 h, commonly 4 h) to perform a series of more than six experiments, which involved >180 stimuli.

Recording. Adult cats were anesthetized with ketamine (20 mg/kg) mixed with acepromazine (0.1 mg/kg) or xylazine (1 mg/kg). Anesthesia was maintained with a continuous intravenous infusion of pentothal mixed with acepromazine (0.1 mg/kg) or xylazine (1 mg/kg). Anesthesia experiments, which involved “test” and a “mask” drifting gratings, $L(t)$ is the sum of the responses to the individual gratings:

$$L(t) = L_{test}(t) + \sigma_{mask}L_{mask}(t),$$

where $\sigma_{mask}$ denotes the relative effectiveness of the temporal frequency of the mask in driving the receptive field.

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tivity. During the course of a recording session with a given neuron (which lasted on average 289 ± 72 min, SD; n = 34), the spontaneous firing rate varied over 18 ± 11 spikes/s, with variations >25 spikes/s in 9 of 34 neurons. Similarly, the overall responsiveness varied substantially over time. The harmonic response to a 50% contrast stimulus varied on average by 25 ± 16 spikes/s, with variations of >40 spikes/s in 4 of 34 neurons. These variations in spontaneous firing rate and responsiveness were not statistically correlated (p > 0.2). We tracked them by letting $V_{\text{max}}$ and $V_0$ vary from one experiment to another. In each experiment, we fixed $V_0$ to minus the mean spontaneous firing rate, which we obtained from the responses to blank stimuli, and we chose $V_{\text{max}}$ to minimize square error between model and data. Had we fixed $V_{\text{max}}$ and $V_0$ across experiments, the quality of predictions would have been compromised in many neurons.

Masking experiments. The logic of the masking experiments is that the test elicits strong responses, and the mask probes how the suppressive field influences these responses (Fig. 2). In general, both test (Fig. 2A, B) and mask (Fig. 2C, D) elicit responses when presented alone. These responses oscillate at the drift frequency and are well predicted by a simple model consisting of a receptive field followed by rectification (Fig. 2A–D, dotted curves). We chose the drift rates of test and mask to be incomensurate (e.g., 7.8 and 12.5 Hz) so that, even when test and mask are superimposed (Fig. 2E, F), we can distinguish a “test response” (the component of the response at the test frequency) and a “mask response” (the component at the mask frequency). The average temporal frequencies of test and mask were 7.3 ± 2.5 and 12.6 ± 2.9 Hz (n = 34).

In the presence of the mask, the test response is smaller than predicted by the receptive field and rectification (Fig. 2E, F, dotted curves). This masking effect is explained by the full model because adding the mask increases the output of the suppressive field. From the response suppression caused by the mask, we deduce the degree to which the mask drives the suppressive field.

We obtained one parameter of the receptive field, $\alpha_{\text{mask}}$, from the mask responses and all parameters of the suppressive field from the test responses. We estimated the strength of the suppressive field $\sigma_{\text{smax}}$ by varying mask contrast over more than six logarithmically spaced values (Fig. 3A, D). We estimated the size of the suppressive field $\sigma_{\text{sup}}$ by varying mask diameter over >10 logarithmically spaced values (Fig. 3B, E). We estimated the parameters that describe the filter bank ($\sigma_n$, $\sigma_m$, and $\delta_n$) by varying mask spatial frequency over >10 logarithmically spaced values (Fig. 3C, F). Finally, we used the mask responses to constrain $\alpha_{\text{mask}}$ (Eq. 2). We estimate $\alpha_{\text{mask}}$ from data in which mask contrast is varied (Fig. 3D) and use the same value for the remaining experiments (Fig. 3E, F).

Model evaluation. To judge the quality of model fits, we calculated the percentage of the variance in the averaged data that is explained by the model. This quantity is given by $100(1-n^{-1} \sum_{j=1}^{n}(\tilde{r}_j - m_j)/\sqrt{\text{var}(\tilde{r}_j)})$, where $\tilde{r}_j$ is the response to stimulus $j$ averaged across trials, and $n$ is the number of stimuli.

We tested model predictions with drifting gratings varying in contrast and diameter (see Fig. 5). This experiment included >50 stimuli covering a full matrix of more than six contrasts and more than eight diameters distributed on a logarithmic scale. The spatial frequency and temporal frequency of the gratings were set to the values eliciting maximal response. Parameters $V_{\text{max}}$ and $V_0$ were left free to account for changes in neural responsiveness. The remaining parameters were held fixed to their previously estimated values.

To compare data and model predictions, we used two independent methods. First, to investigate the ability of the model to predict the effects of contrast on the size tuning curves, we fitted a difference-of-Gaussians to the tuning curves at each contrast (Sceniak et al., 1999). This function provided excellent fits (Ozeki et al., 2004). From these fits, we calculated the preferred stimulus diameter and the degree of size tuning (expressed as a fraction of the peak response) and compared results for data and for model predictions. Second, to investigate the ability of the model to predict the effects of stimulus size on contrast saturation, we fitted a power law to the contrast–response curves at each diameter. The power law provided excellent fits, so the power-law exponent $\beta$ captures the linearity of the growth of responses with contrast: a value close to 0 indicates strong saturation, and a value close to 1 indicates linearity.

To measure the precision of model predictions for individual stimuli, we calculated z-scores of deviations between measured and predicted responses. The z-score of deviations is given by $z = (\tilde{r} - m)/\tilde{R}$, where $\tilde{r}$ is a robust estimate of the SD of responses calculated across trials. To estimate $\tilde{r}$, we fitted a power law relating the observed SD to the mean $\tilde{r}$ and used the resulting predictions as estimate of SD.

Properties of the suppressive field. We estimated the sizes of receptive field and suppressive field from model parameters. We defined size as the diameter of the stimulus eliciting 95% of the maximum response. The receptive field center and surround sizes are given by $\sigma_{\text{max}}$ and $\sigma_{\text{sup}}$, measured and predicted responses to sum of test and mask. Calibration bars in A and B indicate 50 spikes/s. Cell 44.4.2.
the suppressive field by subtracting the test responses measured with mask from the test response measured without mask, normalizing and averaging across cells. We estimated the preferences of the receptive field by normalizing and averaging mask responses.

Results
Our results are organized as follows. First, we devise a simple model of contrast gain control based on the established model of ganglion cell responses proposed by Shapley and Victor (Shapley and Victor, 1981; Victor, 1987). We extend this model to the domain of arbitrary spatial images, by specifying how local contrast should be integrated across space. Then we test the model on a large array of responses of cat LGN neurons and show that it captures response suppression. Finally, we measure the visual properties of contrast gain control and discuss how these properties constrain the possible sources of the suppressive signals.

Model
Our hypothesis is that LGN responses, and in particular the suppressive phenomena, can be explained by a model of contrast gain control such as the one proposed by Shapley and Victor for retinal ganglion cells (Shapley and Victor, 1981; Victor, 1987). In this model, the temporal responses of ganglion cells result from a cascade of low-pass and high-pass filters, whose time constants depend on contrast.

We reformulated the Shapley–Victor model to directly express the effect of contrast on response gain. In Appendix, we simplify the expressions of the model by concentrating on responses to stimuli that vary slowly, because these responses are most affected by gain control. We considered only the effects on response amplitude, ignoring the effects on response timing. The result is a single divisive stage, in which the output of the receptive field appears in the numerator and a measure of local contrast appears in the denominator:

\[
R = \frac{L}{c_{\text{global}} + c_{\text{local}}}.
\]  

(6)

where \( R \) is the response of the neuron, \( L \) is the response of the receptive field (which grows linearly with contrast), \( c_{\text{global}} \) is a constant, and \( c_{\text{local}} \) is local contrast. This expression resembles models of divisive normalization used in visual cortex (Albrecht and Geisler, 1991; Heeger, 1992; Carandini et al., 1997).

Shapley and Victor (1978) did not specify how to compute local contrast \( c_{\text{local}} \) but revealed important constraints. First, \( c_{\text{local}} \) is invariant to stimulus position within the receptive field. Second, \( c_{\text{local}} \) is an “even function” of stimulus intensity (Victor, 1987) (i.e., one that is the same for stimuli that are lighter or darker than the mean luminance). Third, \( c_{\text{local}} \) grows with stimulus area (Shapley and Victor, 1981).

Guided by these constraints, we assumed that local contrast \( c_{\text{local}} \) is the SD of the luminance falling under a Gaussian region concentric with the receptive field. We call this region "suppressive field," because its properties match the term coined by Levick et al. (1972) to describe a region that (1) is engaged equally by light or dark stimuli and (2) suppresses the responses of a cell. The SD is proportional to root-mean-square contrast. It fulfills the above constraints, and it can be generalized to arbitrary images.

The resulting model contains two image-processing pathways, the receptive field and the suppressive field (Fig. 1). The two pathways meet at a divisive stage, in which the receptive field provides the numerator and the suppressive field provides the denominator. The resulting output \( V \) is then rectified to obtain a positive firing rate \( R \).

The receptive field operates linearly: its output involves a simple weighted sum of the image intensities, with weights given by the receptive field profile. We model the receptive field with a difference-of-Gaussians (Rodieck, 1965; Enroth-Cugell and Robson, 1966), which has three free parameters: the width of the center Gaussian, the width of the surround Gaussian, and the strength of the surround relative to the center.

The operation of the suppressive field, instead, is nonlinear, because it involves square root and squaring operations. To compute local contrast, in particular, the suppressive field (1) processes the image through a bank of filters ["subunits" (Shapley and Victor, 1978)], (2) weights the filter outputs by a two-dimensional Gaussian, a window that represents the spatial extent of the suppressive field, and (3) computes the SD of the resulting signals.

We next show that, with a single set of parameters, the model predicts masking, size tuning, and contrast saturation. We obtain the parameters of the model by fitting a series of masking experiments and then hold those parameters fixed and show that they account for size tuning and contrast saturation.

Masking
The model predicts masking, the suppression observed when two stimuli are overlaid (Fig. 3). Masking occurs when either one grating extends in the region surrounding the other (Sillito et al., 1993; Felisberti and Derrington, 1999; Solomon et al., 2002) or two gratings are superimposed (Freeman et al., 2002). In our measurements, the two gratings are a fixed optimal test at 50%
contrast and a mask whose attributes are varied. Test and mask have incommensurate temporal frequencies, which elicit distinct response components (Bonds, 1989): a test response that oscillates at the temporal frequency of the test and a mask response that oscillates at the frequency of the mask. We first investigate how test responses are affected by mask attributes.

The model predicts that adding a mask should reduce test responses. The predicted test responses are given by Equation 6, in which the mask affects only the output of the suppressive field (the term $g_{local}$ in the denominator). Adding the mask, therefore, increases $c_{local}$ and thus reduces test responses.

We test the predictions of the model by varying mask contrast (Fig. 3A), diameter (Fig. 3B), and spatial frequency (Fig. 3C) and find that the model yields good fits (solid curves). (1) Increasing mask contrast progressively suppresses the test response (Fig. 3A). The model predicts this effect because increasing mask contrast increases the local contrast $g_{local}$. With masks of low contrast, there is little suppression because $g_{local}$ is dominated by the constant $c_{50}$. (2) Increasing mask diameter progressively suppresses test responses until responses reach a plateau (Fig. 3B). The model captures this decrease because increasing mask diameter increases local contrast $g_{local}$. The plateau is reached once the mask extends beyond the suppressive field. (3) Suppression depends on mask spatial frequency (Fig. 3C), being strong at low spatial frequencies and absent at high frequencies. The model captures this behavior because of the filters at the front end of the suppressive field, which endow the suppressive field with frequency preferences.

The suppressive field plays a critical role in explaining these observations. The fits performed without it (Fig. 3, dashed curves) are clearly inadequate, indicating that neither the receptive field nor the rectification stage can account for masking. The fits performed including the suppressive field, instead, have high quality: overall, the model explains 94.4% of the variance (median, $n = 34$).

The model can also predict the mask responses, the components of responses at the temporal frequency of the mask (Fig. 3D–F). In this case, varying mask attributes varies the outputs of both receptive field and suppressive field. Mask responses display strong saturation (Fig. 3D), they are tuned for stimulus size (Fig. 3E), and their tuning for spatial frequency is bandpass (Fig. 3F).

The resulting model predictions yield good fits to the mask responses (Fig. 3, solid curves). Fits of similar quality were obtained in the remaining cells, with the model explaining 87.9% of the variance (median, $n = 34$).

### Estimating model parameters

We use these masking experiments to constrain key model parameters, which we henceforth keep fixed. From the effects of mask contrast, we estimate the constant $c_{50}$ (Fig. 3A). From the effects of mask diameter, we estimate the width of the suppressive field $\sigma_{SF}$ (Fig. 3B). From the effects of mask spatial frequency, we estimate the parameters of the filters at the front end of the suppressive field (Fig. 3C). The result is one set of parameters for each cell (Table 1).

The fits yielded reliable estimates of the parameters (Fig. 4). For most parameters, the values yielding the best fits fell in a narrow range. This is the case, for example, for the width of the receptive field $\sigma_{rd}$ (Fig. 4A), for the strength of suppressive field $c_{50}$ (Fig. 4B), and for the width of the suppressive field $\sigma_{SF}$ (Fig. 4C). The exception to this rule is the strength of the surround of the subunits that compose the suppressive field, $k_d$ (Fig. 4D). For this parameter, any value $<0.8$ would be approximately equally good at fitting the responses.

### Size tuning and contrast saturation

Having characterized the model, we ask whether it predicts size tuning and contrast saturation. We test the model on responses to single drifting gratings whose diameter and contrast are varied independently. These data provide a stringent test, because they were not used to constrain the model parameters.

The model predicts size tuning and how this tuning depends on contrast (Fig. 5A). Size tuning is pronounced at high contrast (Fig. 5A, lighter symbols) but weak at low contrast (Fig. 5A, darker symbols).
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Figure 5. Size tuning and contrast saturation. Stimuli are gratings varying in diameter and contrast. Curves are predictions of model with parameters held fixed from previous measurements (Fig. 3; \( V_{\text{max}} = 128; V_0 = -2 \)). A, Responses as a function of diameter, for selected contrasts. B, Same data, plotted as a function of contrast, for selected diameters. Stimuli had optimal attributes: 0.24 cycles/deg and 7.8 Hz. Cell 33.1.3 (93.9% explained variance).

darker symbols) (Solomon et al., 2002; Ozeki et al., 2004). The model captures these effects (Fig. 5A, curves). It exhibits size tuning because extending the stimulus beyond the confines of the receptive field center increases the output of the suppressive field, \( c_{\text{local}} \), but not the output of the receptive field. At low contrasts, instead, \( c_{\text{local}} \) is much smaller than the constant \( c_{\text{opt}} \); the output of the suppressive field is ineffectual, and size tuning is not observed.

The dependence of size tuning on stimulus contrast is marked in our sample (Fig. 6A). In most cells, size tuning is strongest at high contrasts (Fig. 6A, open symbols) and weakest at low contrasts (Fig. 6A, filled symbols). To quantify the degree of size tuning, we estimate the amplitude of plateau responses, the responses to the largest stimulus tested, as a fraction of the peak responses. At 10% contrast, plateau responses are 27 ± 20% smaller than the peak responses (mean ± SD; \( n = 34 \)). The degree of size tuning increases with contrast and peaks at 100% contrast, at which plateau responses are 39 ± 18% below peak responses. The model predicts a similar trend, with values of 11 ± 9% at 10% contrast and 37 ± 15% at 100% contrast.

A consequence of these effects is that preferred stimulus size decreases with contrast (Kremers et al., 2001; Solomon et al., 2002; Nolt et al., 2004). To quantify this effect, we estimate the diameter of the stimulus eliciting maximal response. This preferred stimulus diameter decreases as contrast is increased, ranging from 3.8 ± 2.0° at 10% contrast to 2.4 ± 1.7° at 100% contrast. The model captures this effect with predicted preferred diameters of 4.1 ± 3.0° at 10% contrast and of 2.5 ± 1.6° at 100% contrast.

The model also predicts how responses saturate with increasing contrast (Fig. 5B). Responses to large stimuli show strong saturation (Fig. 5B, lighter symbols). For these stimuli, the outputs of both receptive field and suppressive field are simply proportional to grating contrast \( c \), so the model predicts that firing rate obeys \( R \propto c/(c_{\text{opt}} + c) \). This expression is known to capture the contrast responses of LGN neurons (usually contrast is elevated to a power of \( n \), with \( n \) close to 1) (Derrington and Lennie, 1984; Sclar et al., 1990; Felisberti and Derrington, 1999; Kremers et al., 2001).

The model predicts a novel phenomenon: that contrast saturation is pronounced only for large stimuli (Fig. 5B). Responses to small stimuli grow nearly linearly with contrast (Fig. 5B, darker symbols), whereas responses to large stimuli show clear saturation (Fig. 5B, lighter symbols). The model predicts linearity because small stimuli induce in the suppressive field a response that is negligible compared with the constant \( c_{\text{opt}} \). These data suggest that there is nothing intrinsically nonlinear about LGN responses: were it not for the suppressive field, responses would grow linearly with contrast.

This pronounced dependence of saturation on stimulus size is seen throughout our sample (Fig. 6B). In most cells, saturation is strongest for large stimuli (Fig. 6B, white symbols), weaker for optimal stimuli (Fig. 5B, gray symbols), and nearly absent for the smallest stimuli (Fig. 6B, black symbols). To summarize the degree of saturation across cells, we fit a power law to the contrast–response curve at each stimulus diameter. Power-law exponents close to 0 indicate strong saturation; exponents close to 1 indicate linear growth. With an average exponent of 0.78 ± 0.41, saturation is weakest at the smallest stimulus diameter eliciting a reliable response. Saturation monotonically increases with stimulus diameter, with an average exponent of 0.46 ± 0.18 at the largest diameters tested. The model predicts the trend yielding exponents of 0.61 ± 0.11 for small diameters and 0.40 ± 0.16 for large diameters.

As expected from previous results (Shapley and Victor, 1978; Sclar, 1987), the effects of gain control appear stronger in Y cells than in X cells. The strength of the suppressive field \( c_{\text{opt}} \) averages 0.21 ± 0.06 in X cells (\( n = 28 \), bootstrap estimates) and 0.14 ± 0.06 in Y cells (\( n = 6 \)). The power-law fits of the contrast–response curve (at the largest stimulus tested) yielded exponents of 0.46 ± 0.03 in X cells and 0.40 ± 0.10 in Y cells. These differences, however, did not reach statistical significance (ANOVA, \( p > 0.1 \)).

There are also indications that gain control is more pronounced in ON-center cells than in OFF-center cells, as has been demonstrated in retina (Chander and Chichilnisky, 2001; Zaghloul et al., 2003). The constant \( c_{\text{opt}} \) averages 0.15 ± 0.03 (\( n = 19 \)) in ON cells and 0.27 ± 0.08 in OFF cells (\( n = 15 \)). The exponents average 0.39 ± 0.05 in ON cells and 0.50 ± 0.05 in OFF cells. These differences are consistent with previous results but are not statistically significant (\( p > 0.1 \)).

Model performance
The model performs well in a vast majority of neurons, explaining overall 91.1% of the variance in the contrast-diameter experiments (median, \( n = 34 \)). This performance is remarkable, given that model parameters were estimated from a distinct data set (Fig. 3). To assess model performance in individual stimulus conditions, we computed z-scores of the deviations between measured and predicted responses. The amplitude of deviations is mostly below 1 (\( |z| = 0.65 \), median; \( n = 34 \)) indicating that, for most stimuli, model predictions approach the level of accuracy set by the trial-to-trial variability of responses.

The model predictions, however, showed small but significant biases. We estimated accuracy from the unsigned z-scores, conditioned on stimulus contrast or diameter. Predictions are most...
accurate for diameters slightly larger than optimal ($|z| = 0.52$, median) and least accurate for smaller and larger diameters ($|z| = 0.68$ and $0.74$). Similarly, predictions are most accurate at 40% contrast ($|z| = 0.52$) and least accurate at 5 and 100% contrast ($|z| = 0.86$ and $0.64$). We estimated bias from signed $z$-scores. The model overestimates the amplitudes of responses to large stimuli ($|z| / H11005 / H11002 = 0.71$), whereas predictions for small stimuli show little bias ($|z| / H11005 / H11002 = 0.16$). Likewise, the model overestimates the amplitudes of responses to low-contrast stimuli ($|z| / H11005 / H11002 = 0.89$), whereas predictions for high-contrast stimuli show little bias ($|z| / H11005 / H11002 = 0.06$).

Because fit quality is good and biases are small, we conclude that our simple model of contrast gain control accounts for several suppressive phenomena observed in LGN responses. The model predicts contrast saturation, size tuning, and masking and explains how these phenomena are related to one another.

Spatial extent

Studies in LGN suggested that the sources of suppression extend well beyond the classical receptive field (Levick et al., 1972; Murphy and Sillito, 1987; Cudeiro and Sillito, 1996; Felisberti and Derrington, 1999; Przybyszewski et al., 2000; Kaplan and Benedete, 2001; Sillito and Jones, 2002; Webb et al., 2002; Alitto and Usrey, 2003). How large is the suppressive field, and how does its spatial extent relate to that of the receptive field?

To our surprise, we found that suppressive field rarely extends beyond the surround of the classical receptive field (Fig. 7A, B). We obtained the extent of suppressive field and receptive field from model fits (Fig. 3). To compare the two, we concentrated on the 31 of 34 cells for which the receptive field surround was strong

![Figure 7. Spatial extent of receptive field and suppressive field. A, Comparison of extent of suppressive field and receptive field center obtained from model fits. Dashed line indicates linear regression. B, Same, for receptive field surround. C, Control measurements of spatial extent of receptive field surround and suppressive field. Filled symbols indicate responses to disks containing drifting gratings. Open symbols indicate responses to annuli containing a uniform field whose contrast is modulated in time. Arrows indicate estimated extents of receptive field surround (RF) and suppressive field (SF). Responses averaged across cells. We normalized the abscissa and the ordinate of each curve so that the size tuning curves peak at 1 with a value of 1. Error bars (invisible because smaller than symbols) indicate ± 1 SE ($n = 15$).](image)
(k_{cd} > 0.2; see Materials and Methods). The receptive fields of these cells had an average eccentricity of 13.9 ± 9.1° (SD). The suppressive field has an average extent of 10.3 ± 6.6°. This is ∼300% larger than the receptive field center (3.1 ± 2.2°) (Fig. 7A) and approximately the same size as the receptive field surround (9.1 ± 5.1°) (Fig. 7B).

The sizes of receptive field and suppressive field are strongly correlated across the population. The correlation between the sizes of receptive field center and suppressive field (r = 0.67 ± 0.12, bootstrap estimates) is comparable with the correlation between the sizes of receptive field center and surround (r = 0.53 ± 0.16). The sizes of suppressive field and receptive field surround are also significantly correlated (r = 0.38 ± 0.16).

These results seem discordant with previous studies, which describe suppression as originating beyond the classical receptive field (Levick et al., 1972; Murphy and Sillito, 1987; Cudeiro and Sillito, 1996; Felisberti and Derrington, 1999; Przybyszewska et al., 2000; Kaplan and Benardete, 2001; Sillito and Jones, 2002; Solomon et al., 2002; Webb et al., 2002; Alitto and Usrey, 2003). We therefore felt the need to confirm them with independent measurements (Fig. 7C). Indeed, the measures of receptive field extent presented above were obtained by fitting a difference-of-Gaussians to responses to moving gratings of different spatial frequencies. Although this method is standard and has been thoroughly validated, it is indirect because it involves Fourier analysis.

To obtain a more direct measure of receptive field size, we recorded responses to contrast-modulated circular annuli (n = 15, in two additional animals) and varied annulus inner diameter (Fig. 7C, open symbols). The harmonic responses to these stimuli yield an annulus tuning curve that reflects the center–surround organization of the receptive field (Kilavik et al., 2003; Kremers et al., 2004). The inner diameter at which this curve reaches the floor (within ±1 SD of the response to a blank stimulus) indicates the size of the receptive field surround (Fig. 7C, open arrow). From the same cells, we also recorded responses to optimal drifting gratings and varied the diameter of the disk window (Fig. 7C, filled symbols). The diameter for which these curves reach a plateau (within ±1 SD of the response to the largest stimulus) indicates the size of the suppressive field (Fig. 7C, filled arrow). The results of these analyses confirm that, just as seen earlier using model parameters, the suppressive field does not extend beyond the region of the receptive field. On average, the estimated receptive field and suppressive field sizes were indistinguishable (8.2 ± 0.9 and 7.9 ± 1.9 times larger than the optimal disk diameter, SE; n = 15). The normalized receptive field size had a median of 8.1 and a 50% confidence interval of 4.7–11.9 (Fig. 7C, open arrow and error bar). The normalized suppressive field size had a median of 4.6 and a confidence interval of 2.8–11.9. Although the suppressive field may appear smaller than the receptive field, this difference is not statistically significant (Wilcoxon’s rank sum test, p > 0.2).

These findings confirm that the suppressive field is typically not larger than the surround of the classical receptive field and suggest an explanation for the discrepancy with the literature. Previous studies might have underestimated the actual extent of the receptive field and, in particular, of the receptive field surround. The surround only provides a weak drive to the cell, which can easily go unnoticed. Indeed, our results reveal that the method commonly used to estimate receptive field size systematically underestimates true size, typically by a remarkable factor of 8. Previous studies of LGN responses (Jones et al., 2000; Sillito and Jones, 2002; Solomon et al., 2002; Alitto and Usrey, 2003; Sun et al., 2004; Webb et al., 2005) have estimated the extent of the classical receptive field by measuring responses to high-contrast moving gratings windowed by disks of various sizes and finding the disk size that results in the maximal response. As we have seen (Fig. 7C), this measure underestimates the size of the receptive field surround by a factor of 8.2 ± 0.9. Given that the receptive field surround is approximately three times larger than the center (Table 1), this measure also underestimates the size of the receptive field center by a factor of 2.4. Given that previous studies were most likely underestimating the size of the receptive field surround, it is not surprising that they would describe the suppressive influences as originating outside the receptive field.

Visual preferences

We complete our characterization of gain control in LGN by studying the preferences of the suppressive field for spatial frequency, temporal frequency, and orientation. A quantitative assessment of these preferences provides constraints on the possible sources of suppressive signals (retina, thalamus, or cortex). It is also a necessary step toward a future application of the model to complex stimuli such as natural scenes.

As expected from previous qualitative assessments in retina and in LGN (Shapley and Victor, 1979; Cudeiro and Sillito, 1996; Sun et al., 2004), the suppressive field responds to a broader range of spatial frequencies than the receptive field (Fig. 8A, B). To estimate the tuning of receptive field and suppressive field for spatial frequency, we relied on model parameters obtained as described above. The tuning of the suppressive field is low pass in spatial frequency (Fig. 8B), with low-frequency cutoffs mostly below 0.01 cycles/°, the lowest spatial frequency tested. By comparison, the tuning of the receptive field has strong bandpass characteristics (Fig. 8A) and low-frequency cutoffs in the range of 0.05–0.2 cycles/° (Maffei and Fiorentini, 1973; Dawis et al., 1984; Cheng et al., 1995). The low-frequency cutoff of the suppressive field was only 0.23 ± 0.12 times that of the receptive field (n = 34, bootstrap estimates). The suppressive field also responds to frequencies that often extend beyond the resolution limit of the receptive field: on average, its high-frequency cutoff is 1.27 ± 0.19 times that of the receptive field. Being broadly tuned, the suppressive field barely affects the spatial frequency selectivity of the neurons. Indeed, this selectivity depends only weakly on contrast. We measured spatial frequency tuning curves for stimuli at three contrast levels (∼15, 50, and 100%) and calculated the high-frequency cutoff of each curve. The cutoff frequencies changed little with increasing contrast, from 0.33 ± 0.04 to 0.37 ± 0.07 to 0.39 ± 0.07 cycles/° (mean ± SE; n = 7). These values were not significantly different from one another (ANOVA, p > 0.7). This contrast invariance of spatial frequency tuning seems consistent with the data, if not with the interpretation, of recent studies of spatial frequency tuning at different contrasts (Kremers et al., 2001; Nolt et al., 2004).

The suppressive field is broadly selective for temporal frequency and prefers frequencies of 10–20 Hz or higher (Fig. 8C, D). To estimate the preferences of suppressive field and receptive field, we recorded responses to sums of test and mask gratings and varied mask temporal frequency. By measuring the reduction in test response caused by the different mask temporal frequencies, we estimated the temporal frequency tuning of the suppressive field. By measuring the response to the mask itself, we estimated the temporal frequency tuning of the receptive field. We obtained these measurements in 40 neurons (which include the 34 of our main sample). The suppressive field and receptive field exhibit similar selectivity: both respond weakly at low frequencies and strongly at high frequencies (Fig. 8C, D). These selectivity...
profiles resemble those obtained from responses to single drifting gratings varying in temporal frequency (Fig. 8C, dashed line) (Dawis et al., 1984; Saul and Humphrey, 1990; Alitto and Usrey, 2004). We did not explore frequencies above 21 Hz because, in our stimuli, test and mask were interleaved, and the refresh rate of the monitor was 124 Hz. Therefore, we do not know how the suppressive field responds to very high frequencies.

This assessment of the selectivity of the suppressive field for temporal frequency complements previous studies of gain control in retina and LGN (Shapley and Victor, 1978, 1981; Benardete and Kaplan, 1999). These studies found that gain control affects responses to low frequencies and barely affects responses to high frequencies. Here, we concentrated on the responses to low frequencies, and we investigated the selectivity for temporal frequency of the suppressive field. Consistent with an early anecdotal report in retina (Shapley and Victor, 1979), we found this signal to be strong at high temporal frequencies but weak at low frequencies.

The suppressive field has no preferences for orientation: its bias for orientation is even weaker than that of the receptive field (Fig. 8E,F). To measure the orientation preferences of suppressive field and receptive field, we recorded responses to sums of test and mask gratings and varied mask orientation, following the same logic as the experiments in which we varied temporal frequency. We obtain these measurements in 30 of the 34 neurons in the main sample. The responses of the suppressive field are poorly tuned for orientation (Fig. 8F), reaching significance (ANOVA, p < 0.05) in only 7 of 30 neurons. By comparison, the receptive field often shows consistent biases for orientation (Fig. 8E) (Vidyasagar and Urban, 1982; Soodak et al., 1987; Shou and Leventhal, 1989; Sun et al., 2004), which reach significance (ANOVA, p < 0.05) in 20 of 30 neurons. This bias had no consistent relationship with that seen occasionally in the suppressive field: aligning responses with the orientation that gave the peak receptive field response (Fig. 8E) and averaging across neurons yields a curve that is essentially flat (Fig. 8F). The suppressive field, therefore, is essentially insensitive to stimulus orientation.

This finding agrees with some reports (Solomon et al., 2002; Webb et al., 2002; Sun et al., 2004) but not with others (Sillito et al., 1993; Cudeiro and Sillito, 1996). The latter studies used gratings enclosed in a disk and an annulus and found that the orientation of the grating in the annulus affected the responses to the disk. We suggest that this result might be attributable to interactions between responses elicited by the disk and the annulus in the receptive field. The annulus in these studies was placed just outside the diameter that yields the maximal response for high contrast disk stimuli. We have seen (Fig. 7C) that this diameter is approximately two times smaller than the receptive field center and eight times smaller than the receptive field surround. It is thus reasonable to assume that the annulus stimulated the receptive field surround and elicited some response of its own. This response would oscillate at the same frequency as the response to the disk. Depending on the details of the stimulus and of the receptive field, the two responses could interact destructively or constructively. The result could be an apparent effect of mask orientation.

In summary, we characterized the spatial extent and visual preferences of the suppressive field and found that it is generally (1) not larger than the receptive field surround, (2) broadly tuned for spatial frequency, (3) broadly selective for high temporal frequencies, and (4) not selective for orientation.

Discussion

We have formalized a simple model of gain control and have shown that it provides a parsimonious explanation for the diverse suppressive phenomena seen in LGN responses. The model predicts the phenomena of masking, size tuning, and contrast saturation and explains how these phenomena are interrelated. In particular, it explains how size tuning is pronounced at high contrast and weak at low contrast, and how increasing stimulus contrast reduces the size of the preferred stimulus. Furthermore, the model predicts a novel phenomenon: contrast saturation is present only for large stimuli, and responses to small stimuli grow almost linearly with contrast. This result indicates that there is no saturating nonlinearity at the output of LGN neurons; for stimuli that are small enough, LGN responses grow nearly linearly with contrast, with saturation being entirely explained in contrast gain control.

Origins of the suppressive field

Our measurements of the visual preferences of the suppressive field provide some constraints on the sources of suppression. The possible sources include the following: (1) mechanisms of suppression or gain control operating in retina (Cleland et al., 1983b;
Kaplan et al., 1987; Sclar, 1987; Cheng et al., 1995; Girardin et al., 2002; Nolt et al., 2004); (2) inhibitory circuitry in thalamus (Hubel and Wiesel, 1961; McIlwain and Creutzfeldt, 1967; Singer and Creutzfeldt, 1970; Levick et al., 1972; Cleland et al., 1983b; Cleland and Lee, 1985; Sherman and Koch, 1986; Kaplan et al., 1987; Funke and Eysel, 1998); and (3) negative feedback from visual cortex (Ahlén and Lindström, 1983; Sherman and Koch, 1986; Murphy and Sillito, 1987; Sillito et al., 1993; Cudeiro and Sillito, 1996; Sillito and Jones, 2002; Webb et al., 2002; Worgotter et al., 2002; Alitto and Usrey, 2003).

The visual preferences of the suppressive field are entirely consistent with an origin in retina: just as the suppressive field, retinal ganglion cells respond as well to low spatial frequencies, follow high temporal frequencies, and are barely tuned for stimulus orientation. The mechanisms of contrast gain control in retina have been described by the work of Shapley and Victor (1978, 1981) and are beginning to be understood in terms of anatomy and biophysics (Chander and Chichilnisky, 2001; Kim and Rieke, 2001; Baccus and Meister, 2002; Demb, 2002; Zaghoul et al., 2005). To test whether the suppressive field in LGN is entirely inherited from retinal afferents, one could perform experiments with a test grating in one eye and a mask grating in the other eye.

Our measurements, however, cannot exclude a contribution from thalamic inhibition. There is evidence that gain control is stronger in LGN than in retina (Kaplan et al., 1987; Cheng et al., 1995). LGN cells receive inhibition from two sources (Dubin and Cleland, 1977; Ahlén et al., 1982; Sherman and Koch, 1986): feedforward from local interneurons and feedback from the perigeniculate nucleus (PGN). Interneurons have visual properties that resemble the properties of relay cells (Dubin and Cleland, 1977). It might therefore not be trivial to obtain a broad tuning for spatial frequency by pooling the outputs of interneurons. A more likely source of suppression lies in PGN neurons, whose visual preferences resemble those of the suppressive field. PGN neurons (1) have large receptive fields (Sanderson, 1971; Uhlrich et al., 1991; Wrobel and Bekisz, 1994; Funke and Eysel, 1998), (2) respond well to low spatial frequencies (Price and Morgan, 1987; Xue et al., 1988; Murphy et al., 1994; Wrobel and Bekisz, 1994), (3) respond equally to light increments and decrements (ON/OFF responses) (Sanderson, 1971; Dubin and Cleland, 1977; So and Shapley, 1981; Xue et al., 1988; Uhlrich et al., 1991; Wrobel and Bekisz, 1994; Funke and Eysel, 1998), and (4) show poor selectivity for stimulus size (Jones and Sillito, 1994). The last property is consistent with the observation that suppression is maintained for large diameters (Fig. 3B).

Finally, the visual preferences of the suppressive field are hardly consistent with a cortical origin. The suppressive field has the same extent as the receptive field surround, whereas neurons in primary visual cortex (V1) have larger receptive fields (Jones et al., 2000). More importantly, the suppressive field responds strongly to low spatial frequencies (Fig. 8B) and high temporal frequencies (Fig. 8D), which elicit little response in V1 neurons (Ikeda and Wright, 1975; Movshon et al., 1978; Saul and Humphrey, 1992; DeAngelis et al., 1993; Freeman et al., 2002).

Cortical feedback, however, does seem to affect LGN responses (Alitto et al., 2002; Sillito and Jones, 2002; Worgotter et al., 2002). Cortical ablation or inactivation modifies the gain of the neurons (Przybyszewski et al., 2000; Webb et al., 2002) and reduces size tuning (Murphy and Sillito, 1987; Alitto and Usrey, 2003). These effects might possibly be explained if cortex modulated the strength of the suppressive field, for example by controlling the value of the parameter $\epsilon_{\text{SO}}$, which we have considered to be constant (Fig. 3A,B).

Limitations of our study

A limitation of our study is that we did not consider the nonlinear spatial summation seen in both retina and LGN. We have modeled the component of the responses oscillating at the stimulus frequency. This component accounts for most of the variance in the responses of X cells. Y cells, however, also respond to moving gratings with an elevated mean firing rate (Enroth-Cugell and Robson, 1966; Hochstein and Shapley, 1976; Derrington et al., 1979; Demb et al., 1999). We did not consider the mean rate of responses and therefore could not test the suggestion that the subunits underlying the responses of Y cells correspond to the bank of filters in the suppressive field (Shapley and Victor, 1978, 1981; Victor, 1987).

Another example of nonlinear summation that we have not considered is the “shift effect” (also known as the “periphery effect”), whereby the firing rate of a neuron can be increased by stimulus moving well outside the receptive field (McIlwain and Creutzfeldt, 1964; Levick et al., 1965; Ikeda and Wright, 1972; Kruger and Fischer, 1973; Derrington et al., 1979; Passaglia et al., 2001). Our model cannot explain this increase. Remote stimulation, however, can also induce suppressive effects that have been related to the shift effect (Felisberti and Derrington, 1999, 2001; Passaglia et al., 2001; Girardin et al., 2002). These effects resemble the effects of the suppressive field in many aspects. First, remote stimulation has a divisive effect on the responses to a central target (Felisberti and Derrington, 1999). Second, it affects the contrast gain of the neurons (Felisberti and Derrington, 1999; Girardin et al., 2002). Third, suppression can be induced by stimuli as distant as $\sim 10^\circ$ away from the receptive field center. This finding might be consistent with our finding that the suppressive field extends over a large region of visual field ($10.3 \pm 6.6^\circ$ in diameter at an average eccentricity of $13.9 \pm 9.1^\circ$).

A major limitation of our work is that we have not considered the time course of the responses. Contrast gain control not only affects the gain of the responses but also their temporal dynamics (Shapley and Victor, 1978, 1981). Increasing contrast reduces the integration time of the neurons affecting the phase of responses and their selectivity for temporal frequency. The model does not capture these effects; in fact, we intentionally simplified the Shapley–Victor model so that it would predict only the amplitude of the responses. Our model therefore only works for stimuli of fixed temporal frequencies.

Moreover, we have not considered the dynamics of contrast gain control (i.e., how a change in contrast at a given time can influence gain at a later time). We cannot study these dynamics because, in our stimuli, contrast was fixed through time. In particular, we cannot distinguish the fast mechanism of contrast gain control (Victor, 1987; Baccus and Meister, 2002; Mante et al., 2005; Zaghoul et al., 2005) from the slower mechanism of contrast adaptation (Chander and Chichilnisky, 2001; Kim and Rieke, 2001; Baccus and Meister, 2002; Solomon et al., 2004). We suspect that slow adaptation plays little role in our results, because it is pronounced only for high-frequency stimuli (Solomon et al., 2004) and is weak at lower frequencies (Ohzawa et al., 1985; Shou et al., 1996; Sanchez-Vives et al., 2000; Yang et al., 2003; Solomon et al., 2004). Moreover, our stimuli were brief (typically 1–2 s, always $<4$ s), presented in randomized order, and separated by $\sim 1$ s of blank screen; these properties would minimize the effects of slow adaptation, which take seconds to develop and are strongest when low contrast stimuli are preceded by long periods of strong stimulation (Solomon et al., 2004).
Implications for V1 responses

Our formulation of retinal contrast gain control in terms of division reveals close similarities with models of gain control or normalization in primary visual cortex (Albrecht and Geisler, 1991; Heeger, 1992; Carandini et al., 1997). These models predict expressions similar to Equation 6. In fact, neurons in primary visual cortex exhibit contrast saturation (Albrecht and Hamilton, 1982; Sclar et al., 1990), masking (Bonds, 1989; Freeman et al., 2002), and size tuning (Sceniak et al., 1999; Cavanaugh et al., 2002). The exact contribution of LGN suppression to these phenomena is not clear. It is evident that some cortical effects, such as the orientation selectivity of surround suppression (Cavanaugh et al., 2002), are not easily explained by suppression in LGN. Nonetheless, our results indicate that the suppressive phenomena seen in LGN would be expected to contribute to those seen in V1.

The fact that a model with similar structure can explain gain control in retina, in LGN, and in cortex is very good news. It paves the way for a long-due assessment of the relative roles of these stages of visual processing to phenomena of suppression, gain control, and normalization. Once these phenomena are fitted with the same model in the different stages, then the contribution of each stage can be described by the parameters of the model (Sclar, 1987; Sclar et al., 1990).

Function of the suppressive field

The suppressive field is distinct from the receptive field in many respects. First, it is responsible for suppressive phenomena that cannot be explained by the receptive field alone (Fig. 3, dashed lines). Second, it responds well to high and low spatial frequencies (Fig. 8B), frequencies that generally elicit poor responses in the receptive field (Fig. 8A). Third, it computes the SD of stimulus intensities (Fig. 1).

What use might the visual system make of divisive suppression? One possible use is to map the wide range of contrasts encountered in the natural environment onto the limited dynamic range of the neurons (Heeger, 1992). Suppression may also play a role in encoding of visual information by populations of neurons: a divisive gain control model applied to responses in primary visual cortex was shown to increase independence of the responses across neurons (Schwartz and Simoncelli, 2001; Valerio and Navarro, 2003).

We suggest that the suppressive field could also play a role in the joint encoding of spatial frequency and spatial position, which are subject to a stringent tradeoff (Daugman, 1985). In LGN, selectivity for spatial frequency results from the antagonistic effect of the receptive field center and surround. This selectivity comes, however, at the cost of lower spatial localization because of the large size of the receptive field surround. The suppressive field counteracts this effect by reducing the effective integration area of the neuron. This improvement in encoding of spatial position comes at no cost because the suppressive field barely affects the tuning of responses for spatial frequency (Fig. 3F).

Toward a functional model of LGN neurons

Our model is a parsimonious description of the processing performed by LGN neurons on spatial images. It features few free parameters, each of which has a clear functional interpretation. These parameters can be reliably estimated from a limited set of measurements and held fixed to predict data obtained in new experiments.

This research, thus, brings us closer to a central goal of visual neuroscience, the development of functional models of neural responses to visual stimuli. Functional models summarize the image processing performed by neural circuits, guide research in the underlying biophysics, and facilitate the study of subsequent visual stages. Functional models are particularly desirable for LGN, whose responses reflect the complex computations performed in retina, and constitute the principal input to the visual cortex. As the first inroads toward thalamic visual prostheses are being made (Pezaris and Reid, 2004), the need for such a model may become much more pressing.

Appendix: divisive effect of gain control

Here we show that the model of retinal contrast gain control by Shapley and Victor (1981) predicts that contrast has a divisive effect on the responses. The model consists of a fixed low-pass filter followed by a high-pass stage whose time constant varies with contrast.

For simplicity, we consider the version of their model described by Victor (1987), in which responses to a stimulus with contrast $c$ and temporal frequency $\omega/(2\pi)$ are given by the following:

$$R(c, \omega) = cL(\omega) \left(1 - \frac{h}{1 + i\omega\tau(c)}\right),$$

where $L(\omega)$ is a low-pass filter, and the expression in the large parentheses is a high-pass filter with gain $h$ and time constant $\tau(c)$. Following Victor (1987), we take the latter to depend on contrast $c$ as follows:

$$\tau(c) = \frac{\tau_0}{1 + (c/c_0)^n},$$

with $n$, $\tau_0$, $c_0$ constants.

Fixing $h = 1$ (Benardete and Kaplan, 1999), we substitute Equation 8 into Equation 7 and calculate the absolute value to obtain the following:

$$|R(c, \omega)| = |L(\omega)| \frac{c}{\sqrt{(c_0^2 + c^2)^{1/n} + \omega^2 \tau_0^2 c_0^{2n}}},$$

where $|L(\omega)|$ is the amplitude of the linear response, and the right-hand term captures the effect of gain control. The denominator in Equation 9 is responsible for the effects of gain control. Gain control is pronounced at low temporal frequencies ($\omega \ll 1/\tau_0$) for which the denominator grows with contrast. Gain control is absent at high frequencies ($\omega \gg 1/\tau_0$) for which the denominator becomes independent of contrast. With $\omega$ close to 0, Equation 9 reduces to a hyperbolic ratio:

$$|R(c)| = \frac{|L(0)| c}{(c_0^2 + c^2)} ,$$

which demonstrates that contrast gain control in retina is divisive, because contrast appears not only in the numerator but also in the denominator.

In cat X ganglion cells, $\tau_0$ has a median value of 0.23 s (Victor, 1987), so the approximation provided by Equation 9 is exact only for frequencies less than $1/0.23/(2\pi) = 0.69$ Hz. However, we found that the expression provides good fits to responses at frequencies as high as 12 Hz.

Our Equation 6 follows from Equation 10 once one sets $n = 1$. Indeed, this value seems more consistent with the data (Fig. 3) (Sclar et al., 1990) than the value $n = 2$ used by Victor (1987).
References


