Cellular and Circuit Mechanisms Shaping the Perceptual Properties of the Primate Fovea

Graphical Abstract

Highlights
- Cone photoreceptor kinetics dictate temporal sensitivity of our visual perception
- Foveal midget ganglion cells (MGCs) receive little or no synaptic inhibition
- Foveal but not peripheral MGCs express few inhibitory postsynaptic receptors
- Postsynaptic inhibition controls gain but not kinetics of MGC responses

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In Brief
The unique perceptual features of the primate fovea are shaped by the properties of cone photoreceptors, rather than retinal circuit computations.

Sinha et al., 2017, Cell 168, 413–426
January 26, 2017 © 2017 Elsevier Inc.
http://dx.doi.org/10.1016/j.cell.2017.01.005
Cellular and Circuit Mechanisms Shaping the Perceptual Properties of the Primate Fovea

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SUMMARY

The fovea is a specialized region of the retina that dominates the visual perception of primates by providing high chromatic and spatial acuity. While the foveal and peripheral retina share a similar core circuit architecture, they exhibit profound functional differences whose mechanisms are unknown. Using intracellular recordings and structure-function analyses, we examined the cellular and synaptic underpinnings of the primate fovea. Compared to peripheral vision, the fovea displays decreased sensitivity to rapid variations in light inputs; this difference is reflected in the responses of ganglion cells, the output cells of the retina. Surprisingly, and unlike in the periphery, synaptic inhibition minimally shaped the responses of foveal midget ganglion cells. This difference in inhibition cannot however, explain the differences in the temporal sensitivity of foveal and peripheral midget ganglion cells. Instead, foveal cone photoreceptors themselves exhibited slower light responses than peripheral cones, unexpectedly linking cone signals to perceptual sensitivity.

INTRODUCTION

The fovea accounts for <1% of the surface area of the primate retina but accounts for ~50% of the retinal output neurons and provides input to ~50% of the cells in primary visual cortex (Wässle et al., 1989). Signals originating in the fovea enable the high chromatic and spatial acuity that dominate our everyday visual experience—including your ability to read this page. But foveal signaling limits sensitivity to other important aspects of the visual world. Specifically, perceptual sensitivity to rapidly-varying light inputs is lower for foveal vision than peripheral vision (Hecht and Verrijp, 1933), and this difference is present in the retinal outputs (Solomon et al., 2002). The low temporal sensitivity of foveal vision is reflected in the refresh rates of computer monitors and movies and will be an important factor in the design of visual prosthetics. The absence of a fovea in most mammals and technical challenges associated with intracellular recordings from the primate fovea mean that we know little about the cellular and synaptic basis of the functional differences between the fovea and peripheral retina.

The foveal and peripheral retina share a similar core circuit architecture: photoreceptors convey information to bipartite neurons that then convey information to output ganglion cells (reviewed by Hoon et al., 2014). Inhibitory interneurons can modulate this core excitatory pathway. Given this common circuit architecture, differences in the temporal sensitivity of foveal and peripheral retinal outputs could originate in the cone photoreceptors themselves or in the neural circuits that read out the cone signals. A likely circuit mechanism is synaptic inhibition, which shapes the kinetics of signals in many neural circuits (reviewed by Isaacson and Scanziani, 2011), including the retina (reviewed by Jadhinsky and Baccus, 2013). Inhibition can speed or slow responses depending on where it operates and how it interacts with other synaptic properties such as depression (Asari and Meister, 2012). The potential role of inhibition in shaping response kinetics is one example of a broader emergent theme in retinal research showing the importance of inner retinal inhibitory circuits in unexpected and sophisticated computations (reviewed by Jadhinsky and Baccus, 2013) such as direction selectivity (Wei et al., 2011; Yonehara et al., 2011) and sensitivity to local but not global motion (Baccus et al., 2008). These computations are distinct from the outer retinal horizontal cell-mediated surround inhibition that shapes ganglion cell receptive fields across the retina (Crook et al., 2011; McMahon et al., 2004; Wu, 1992).

Differences in temporal sensitivity could also originate in the cone photoreceptor responses. The importance of the kinetics of photoreceptor inputs to the retina is not without precedent: striking differences in the kinetics of the responses of rod and cone photoreceptors (Schneeweis and Schnapf, 1995) clearly contribute to night versus day differences in the temporal sensitivity of visual perception (Hecht and Verrijp, 1933). Cones themselves are often assumed functionally homogeneous across the retina except for differences in wavelength sensitivity, but this assumption has not been tested.

RESULTS

The results are organized around three main findings about the origin of differences in temporal sensitivity between foveal and peripheral midget ganglion cells (MGCs): (1) the kinetics of responses of foveal MGCs, unlike peripheral MGCs, are minimally
shaped by synaptic inhibition; (2) consistent with this functional difference, foveal MGCs express fewer inhibitory postsynaptic receptors than their peripheral counterparts; and, (3) the kinetics of the responses of foveal and peripheral cones differ dramatically.

**Midget Ganglion Cells Show Slower Response Kinetics in the Fovea than in the Periphery**

In vivo recordings of responses to sinusoidal gratings show that responses of foveal MGCs decline more quickly with temporal frequency than those of peripheral MGCs (Solomon et al., 2002). To confirm that these differences are present in a preparation that allows access to the underlying cellular and synaptic mechanisms, we recorded spike responses in whole mount retina in an in vitro preparation. We compared MGC spike output in foveal (1 cone/MGC, 0–1 mm eccentricity), central MGC spike output in foveal (1 cone/MGC, 0–1 mm eccentricity) and peripheral retina in an in vitro preparation. We compared MGC spike output in foveal (1 cone/MGC, 0–1 mm eccentricity), central (3–4 cones/MGC, 2–4 mm eccentricity, 8–16 degrees of visual angle, Figure 1A middle and peripheral retina (10–20 cones/MGC, >6 mm eccentricity, >24 degrees of visual angle, Figure 1A right). These regions differ dramatically in retinal circuitry and in the visual behavior that they support; notably foveal MGCs receive (indirectly via a midget bipolar cell) input from a single cone photoreceptor, forming a “private line” of signaling and the basis for the high spatial acuity of foveal vision, while central and peripheral MGCs receive (indirect) input from ~2–6 cones and ~10–30 cones, respectively (Calkins et al., 1994; Kolb and Marshall, 2003; Merigan et al., 1991; Schiller et al., 1990).

Full-field contrast steps elicited a robust increment in spike responses at the onset of the stimulus in ON MGCs—i.e., MGCs that depolarize to increases in light intensity (Figure 1B). The spike responses of peripheral MGCs to 100% contrast steps had a more pronounced transient peak compared to responses of foveal MGCs (Figure 1B). To further characterize MGC response kinetics, we presented a time-varying random stimulus and computed the average stimulus preceding a spike (the spike-triggered average or STA; Figures 1C–1E). The STA kinetics differed substantially, with foveal MGCs exhibiting the slowest kinetics and peripheral MGCs exhibiting the fastest kinetics (Figures 1C and 1D). Kinetics of central MGCs fell between those of foveal and peripheral MGCs, indicating a smooth dependence of response kinetics on retinal eccentricity (Figures 1C and 1D).

**Foveal MGCs Receive Little or No Light-Driven Inhibitory Synaptic Input**

The gradient in kinetics of MGC spike output could be created upstream of the MGC in the retinal circuitry or by integration of excitatory and inhibitory synaptic inputs in the MGC. The balance of inhibitory and excitatory synapses (the I/E balance) shapes synaptic integration and signaling in many neural circuits (reviewed by Isaacson and Scanziani, 2011). This general importance of I/E balance underscores the importance of comparing MGC inhibitory and excitatory synaptic inputs across retinal locations (foveal, central and peripheral).

Full-field contrast steps elicited robust excitatory synaptic inputs to ON MGCs across retinal locations (Figure 2A). Excitatory synaptic inputs, like spike outputs, were more transient in peripheral MGCs compared to foveal and central MGCs (Figure 2A). Furthermore, MGC inhibitory synaptic inputs elicited by these (and other) stimuli varied dramatically across retinal regions, with the smallest inhibitory inputs in the fovea and largest in the periphery (Figures 2A, 2C–2F). OFF MGCs in the fovea also exhibited robust light-driven excitatory inputs and minimal inhibitory inputs (Figure 2B). Previous work in central retina (2–4 mm eccentricity) found, similarly, that drifting gratings elicit small inhibitory inputs to MGCs (Crook et al., 2011). The ratio of inhibitory to excitatory synaptic input (the I/E ratio) changed more than 10-fold between foveal and peripheral cells (Figures 2C and 2D).

Uncaging GABA and glutamate provided a further test of the generality of these results. GABA elicited a much smaller response relative to glutamate in foveal MGCs compared to peripheral MGCs (Figure 2G; see STAR Methods). The lack of light-driven inhibitory input was not an artifact of our recordings, as foveal MGCs received spontaneous inhibitory input that was not correlated with the onset of the light stimulus (Figure 2G; see also Figure S5).

**Non-midget Ganglion Cells in the Fovea Exhibit Robust Postsynaptic Inhibition**

Do other ganglion cells in the fovea share the near-complete lack of light-driven synaptic inhibition exhibited by MGCs? MGCs account for ~90% of the ganglion cells in the fovea; the remaining ~10% are ganglion cell types with wider dendritic fields, which correspondingly receive (indirect) input from multiple cones (Calkins and Sterling, 2007; Grünewald et al., 1993). Non-MGC foveal cells are difficult to target for recording due to their sparsity and the similarity of their soma morphology to that of the much more numerous MGCs. Hence, we have compared the wide-field cells collectively to the MGCs. Figures 3A–3C show excitatory and inhibitory synaptic inputs to three foveal wide-field cells, and Figure 3D collects I/E ratios for foveal MGCs, foveal wide-field cells and peripheral MGCs. Among these cell types the foveal MGCs are unique in receiving minimal light-driven inhibitory synaptic input.

**Foveal MGCs Express Low Levels of Inhibitory Relative to Excitatory Postsynaptic Receptors**

The lack of light-driven inhibitory input to foveal MGCs was surprising given the importance of inhibition in other neural circuits (reviewed by Isaacson and Scanziani, 2011) and the high density of presumed inhibitory (non-ribbon) synapses made onto foveal MGC dendrites as revealed by electron microscopy (Calkins et al., 1994; Calkins and Sterling, 1996; Kolb and Dekorver, 1991). These findings could be reconciled if the stimuli used in our physiological recordings failed to engage inhibitory circuits. Alternatively, postsynaptic expression of excitatory and inhibitory receptors could differ independent of synapse number. As described below, assessment of receptor expression in MGC dendrites across eccentricity supports the latter explanation.

Excitatory and inhibitory postsynaptic receptors on the dendrites of individual GCs were visualized by biolistic transfection with plasmids encoding several fluorescently labeled
synaptic proteins: (1) postsynaptic density protein-95 (PSD95-CFP), a scaffolding protein at most retinal excitatory synapses (Koulen et al., 1998) and a “marker” for ionotropic glutamate receptors in retinal ganglion cells (Kerschensteiner et al., 2009; Morgan et al., 2008); (2) GABA_A receptor subunits (α1-YFP, β2/3 and γ2); and, (3) tdTomato to visualize dendrites (Figures 4A and 4B). We labeled the GABA_A α1 subunit because it is predominantly localized on primate MGCs (Abbott et al., 2012; Grünert et al., 1993). Fluorescently-tagged postsynaptic proteins were localized appropriately at their respective synapses (Figure S1).

Figure 1. Midget Ganglion Cells Show Slower Response Kinetics in the Fovea than in the Periphery
(A) Schematic depicting the convergence of the midget circuit in primate retina at three different eccentricities. MGCs were classified as foveal (<1 mm), central (2–4 mm), and peripheral (>6 mm) based on their distance from the foveal pit. Top panel shows a composite fluorescence micrograph of AMPA receptor subunit, GluA3, at the outer plexiform layer indicating the cone density and the bottom panel is the corresponding image of a neurobiotin-filled MGC at each eccentricity. (B) Spike trains from six trials of an exemplar MGC and the average peri-stimulus time histogram (binwidth of 20ms) across several MGCs at each retinal eccentricity in response to a full-field 100% contrast light step (background light intensity of ~4,000 R*/cone/sec). Note the transient spike response from peripheral MGCs. (C) Temporal receptive field probed with a full-field Gaussian white noise stimuli. A short excerpt of the stimulus and spike response (top) with exemplar time-reversed spike-triggered average (STA) of individual MGCs at each retinal eccentricity (bottom). (D) Quantification of the time to peak i.e., latency of the peaks in STAs yields a mean of 67 ± 6 ms for foveal MGCs, 53 ± 6 ms for central MGCs and 37 ± 4 ms for peripheral MGCs. The latency of the STAs is significantly longer for foveal MGCs than central or peripheral MGCs. (E) Quantification of the biphasic index of STAs i.e., the ratio of the peak amplitudes from the baseline yields a mean value of 0.33 ± 0.15, 0.30 ± 0.06 and 0.51 ± 0.35 for MGCs in foveal, central and peripheral retina, respectively. In all figures, error bars indicate s.d. and ‘n’ refers to the number of cells analyzed.
Figure 2. Foveal Midget Ganglion Cells Receive Little or No Light-Evoked Inhibition

(A) Average excitatory (Exc, in green) and inhibitory (Inh, in red) synaptic currents produced by a 100% contrast light step from voltage-clamped MGCs in foveal, central and peripheral retina.

(B) Mean excitatory (green) and inhibitory (red) synaptic current in response to a negative 100% contrast step measured from a voltage-clamped foveal OFF MGC.

(C) Ratio of the inhibitory to excitatory charge transfer analyzed from individual MGC response to a contrast light step as shown in A for each of the retinal locations. The mean I/E ratio is 0.06 ± 0.05, 0.33 ± 0.14 and 1.18 ± 0.35 for foveal, central and peripheral MGCs, respectively.

(D) 2D plot of absolute excitatory versus inhibitory charge transfer across eccentricity (raw values used to calculate I/E ratio in Figure 2C). The average value for peripheral MGCs lies on the unity line whereas foveal and central MGCs are below the unity line and skewed toward excitation.

(E) Exemplar average trace of inhibitory and excitatory synaptic currents of foveal ON MGCs in response to a time varying stimulus. Bottom panel shows a pairwise comparison of stimulus-driven variance in excitatory and inhibitory responses across 4 foveal ON MGCs.

(F) Mean excitatory (green) and inhibitory (red) synaptic current in response to a positive 200% contrast step measured from voltage-clamped foveal ON MGCs (n = 5).

(G) (i), Spontaneous inhibitory synaptic currents (Control Inh) are often observed in foveal MGCs and are blocked after bath application of GABA and Glycine receptor blockers (Inh block).
To quantify expression of excitatory and inhibitory receptors, we estimated the percent of the dendritic volume occupied by PSD95-CFP and by GABA(A)-YFP (volume occupancy, see STAR Methods). Two results emerged. First, the ratio of GABA(A)-YFP to PSD95-CFP expression (I/E ratio) was much smaller in foveal compared to peripheral MGCs (Figures 4B and 4D), consistent with the physiological observations. Second, wide-field ganglion cells in the fovea, such as parasol cells, exhibited considerably greater expression of inhibitory GABA(A)-YFP receptors than nearby foveal MGCs; with a GABA(A)-YFP/PSD95-CFP ratio near 1.0 (Figure 4D), also consistent with physiology (Figure 3).

Immunolabeling experiments provided additional evidence for the low expression of inhibitory receptors by foveal MGCs. Individual MGCs were filled with neurobiotin during physiological recordings and the retina was then immunostained with antibodies against the GluA3 subunit of the AMPA receptor (Figure 4C, raw images of immunostaining in Figure S2A, additional control labelings in Figure S6) and gephyrin (Abbott et al., 2012) and consistent with our physiological measurements (Figure 2). Wide-field ganglion cells in the fovea displayed high GABA(A)α1 levels relative to MGCs (Figure S4).

Together, both biolistic and immunolabeling experiments show that foveal MGCs express relatively few inhibitory receptors, and that foveal and peripheral wide-field GCs that receive input from multiple cones express a much higher ratio of inhibitory to excitatory receptors. We compare these findings with electron microscopic measures of presynaptic structures in the Discussion.

Inhibitory Synaptic Input Regulates the Gain but Not Kinetics of the Peripheral MGC Spike Output

Does the strong gradient in the ratio of inhibitory to excitatory synaptic input received by MGCs across retinal locations account for the difference in MGC response kinetics? We used the dynamic clamp technique to determine how integration of excitatory and inhibitory inputs shapes the MGC spike output. These experiments measure the voltage changes elicited by injecting currents corresponding to light-evoked conductances. We used three sets of conductances measured in response to randomly-varying stimuli (Figure 5A): (1) the excitatory conductance measured from a foveal MGC; (2) excitatory and inhibitory conductances measured simultaneously (Cafaro and Rieke, 2010) from a peripheral MGC; and (3) the peripheral MGC excitatory conductance alone. Three conclusions emerged from comparing the spike responses and STAs of foveal and peripheral MGCs (Figures 5B–5D). First, the conductances rather than the cell type they were injected into dominated the kinetics of the STAs—i.e., a foveal and a peripheral MGC injected with the same conductances responded similarly (Figure 5B). Second, STA kinetics were similar with and without the inhibitory conductance, providing direct evidence that inhibitory

excitatory postsynaptic receptors (Figures 4C and 4E), as previously shown (Abbott et al., 2012) and consistent with our physiological measurements (Figure 2). Wide-field ganglion cells in the fovea displayed high GABA(A)α1 levels relative to MGCs (Figure S4).

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synaptic input to the MGC did not regulate response kinetics (Figure 5B). Third, the spike rate and STA amplitude were both larger in the absence of the inhibitory conductance (Figures 5C and 5D).

These results indicate that inhibitory input acts to regulate gain but not kinetics of the responses of peripheral MGCs. Peripheral MGCs, unlike those in the fovea, receive input from multiple cones and hence may require gain regulation via inhibitory input to extend the range of contrasts they can encode (see Discussion).

Presynaptic Inhibition Does Not Shape the Excitatory Inputs of Foveal MGCs

The gradient in MGC response kinetics across retinal regions could, alternatively, be created by differences in kinetics of excitatory synaptic inputs onto MGCs. Consistent with this possibility, the kinetics of the linear filters (the filters that transform visual inputs into changes in excitatory synaptic currents; analogous to the STAs) characterizing the MGC excitatory synaptic inputs depended strongly on eccentricity (Figures 6A and 6B).

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<th>Presynaptic Inhibition Does Not Shape the Excitatory Inputs of Foveal MGCs</th>
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| (A) Dynamic clamp experiment illustrating the three sets of conductances injected into foveal or peripheral MGCs. (i) Synaptic conductances were measured in response to white noise stimuli as in Figure 1C: 1. Foveal $g_{\text{exc}}$, 2. Peripheral $g_{\text{exc}} + g_{\text{inh}}$, and 3. Peripheral $g_{\text{inh}}$. (B) Exemplar current-clamp recording from a foveal MGC where each of the three sets of conductances evokes corresponding spike responses. Exemplar STA from a foveal MGC for each of the three injected conductance sets. (B) Bar graph comparing the time to peak of the STAs for each of the three conductance sets for foveal and peripheral MGCs. MGCs show faster kinetics with shorter time to peak when injected with peripheral synaptic conductances at the fovea (mean of $36 \pm 3$ ms for $g_{\text{exc}}$ and $36 \pm 1$ ms for $g_{\text{inh}}$) and in the periphery (mean of $34 \pm 2$ ms for $g_{\text{exc}}$ and $34 \pm 2$ ms for $g_{\text{inh}}$). However, when MGCs are injected with peripheral conductances their response kinetics gets appreciably slower (mean of $57 \pm 4$ ms for foveal MGCs and mean of $55 \pm 4$ ms for peripheral MGCs). (C) Ratio of the STA amplitude measured from MGCs without peripheral inhibitory conductances (set 3 in A) to those measured with peripheral inhibitory conductances (set 2 in A) yields a mean value of $1.5 \pm 0.4$ for both foveal and peripheral MGCs. (D) The ratio of the mean spike rate yields a mean value of $2.2 \pm 1.4$ and $1.9 \pm 0.5$ for foveal and peripheral MGCs, respectively.

Figure 5. Inhibitory Synaptic Input Affects Gain but Not Kinetics of Peripheral Midget Ganglion Cell Outputs

Figure 4. Foveal Midget Ganglion Cells Express Low Levels of Inhibitory Postsynaptic Receptors Relative to Excitatory Postsynaptic Receptors

(A) Whole-mount view of retinal ganglion cells (RGCs) biolistically labeled by expression of tdTomato (Tom) in foveal (top) and peripheral (bottom) macaque retina. Inset shows a single MGC at each eccentricity. (B) Maximum intensity projection of a co-transfected MGC and parasol cell in the macaque fovea and a MGC in peripheral retina. Arrows point to the axon. PSD95-CFP (green) and GABA_A_{x1}-YFP (red) expression in tdTomato (Tom, blue) labeled cells. (C) Maximum intensity projections showing neurobiotin-filled (NB, blue) dendrites of MGCs colabeled with GluA3 (green) and gephyrin (Gep, red) or with GluA3 and GABA_A_{x1}-YFP (red) at each of the three different retinal eccentricities (foveal, central, peripheral). Top panel shows the combined raw immunolabeling signal (see Figure S2 for individual labeling panels). Middle panel shows the receptor signal within the masked MGC dendritic process. Bottom panel shows the receptor signal along a small dendritic stretch. (D) Quantification of the % volume occupancy of GABA_A_{x1}-YFP signal relative to PSD95-CFP signal within tdTomato filled dendrites in the fovea revealed a much higher $I_{\text{GABA_A}}/I_{\text{PSD-95}}$ ratio in parasol cells (mean of $1.2 \pm 0.17$) relative to MGCs (mean of $0.12 \pm 0.05$). MGCs also revealed a low-fovea to high-periphery $I_{\text{GABA_A}}/I_{\text{PSD-95}}$ ratio across eccentricity (periphery, mean of $0.93 \pm 0.12$). The overlapping data points have been shifted and color-coded for better clarity. (E) Quantification of the $I_{\text{GABA_A}}/I_{\text{GluA3}}$ ratio by estimation of the % volume (dendrite) occupancy of gephyrin signal as compared to the % occupancy of the GluA3 signal. The $I_{\text{GABA_A}}/I_{\text{GluA3}}$ ratio for neurobiotin filled MGCs was significantly lower in foveal (mean of $0.21 \pm 0.06$) compared to central (mean of $0.50 \pm 0.06$) and peripheral (mean of $0.60 \pm 0.05$) retina. Similar results were obtained upon quantification of the $I_{\text{GABA_A}}/I_{\text{GluA3}}$ ratio for neurobiotin filled MGCs (foveal, mean of $0.17 \pm 0.04$; central, mean of $0.49 \pm 0.04$; peripheral, mean of $0.76 \pm 0.03$). The overlapping data points have been shifted and color-coded for better clarity. See also Figures S1, S2, S3, S4, and S6.
6B). The kinetics of linear filters for peripheral MGCs were similar to the corresponding STAs (Figures 6A–6C), which reaffirms the conclusion that postsynaptic inhibition is unlikely to play a significant role in temporal filtering of MGC responses. These results suggest that a mechanism upstream of the MGCs creates the kinetic differences in retinal output across eccentricity.

Several mechanisms can shape the excitatory input to MGCs. Presynaptic inhibition, for example, could act on the axon terminals of midget bipolar cells that provide the dominant excitatory input to MGCs (Calkins et al., 1994; Kolb and Marshak, 2003). To assess the contribution of presynaptic inhibition, we measured the effect of suppressing GABA and glycine receptor activity on excitatory synaptic inputs to foveal and peripheral MGCs. Suppressing inhibitory receptors substantially altered excitatory inputs and spike outputs of peripheral MGCs, including revealing a large response at light offset in addition to the normal response at light onset (Figures 6F and 6H). The extent of these changes precluded an estimate of the impact of presynaptic inhibition on response kinetics. The same manipulation had little or no effect on amplitude and kinetics of the foveal MGC excitatory input and spike output (Figures 6D, 6E and 6G). Thus, neither pre- nor post-synaptic inhibition substantially shapes the kinetics of responses of foveal MGCs.

**Cone Photoreceptors Exhibit Slower Kinetics in the Fovea than the Periphery**

Differences in the kinetics of responses of foveal and peripheral MGCs could originate in the cones themselves (Seiple and Holopigian, 1996; Tyler, 1985). Foveal and peripheral cones differ dramatically in density and morphology (Wikler et al., 1990). The cone density decreases with distance from the fovea and is 100-fold less in the peripheral retina (Wikler et al., 1990). Cones in the fovea have axons that extend ~400 μm in length, whereas central and peripheral cones have much shorter axons (Hsu et al., 1998). Foveal cones also have longer inner and outer segments compared to central and peripheral cones (Polyak, 1941, 1957). However, the possibility that cones could differ functionally across retinal location has not been tested.

To compare the kinetics of responses of foveal and peripheral cones, we recorded cone inner segment voltages in response to Gaussian noise stimuli and computed the linear filters—i.e., the filters that transform visual inputs into changes in cone voltage (Figure 7A; stimuli and analysis were identical to those used for the MGC responses in Figures 1 and 2 and 6). We focused on long (L) and middle (M) wavelength sensitive cones, identified based on their distinct spectral sensitivities. Surprisingly, the light-dependent voltage changes of foveal L and M cones were considerably slower than those of peripheral cones (Figure 7B). Voltage changes elicited by...
brief light flashes exhibited similar differences (Figure 7C). These differences are comparable in magnitude to the differences observed in the MGC responses—i.e., the time to peak of both the cone and MGC responses are ~2-fold slower in the fovea than periphery.

The differences in kinetics of light-dependent voltage changes of foveal and peripheral cones could originate within the phototransduction process, from electrical properties of the inner segment, or from network interactions mediated by horizontal cells or gap-junctional coupling (Barrow and Wu, 2009; Endeman et al., 2012; Hornstein et al., 2005a; Hornstein et al., 2005b; Verweij et al., 2003). To test for contributions of inner segment conductances, we measured cone currents using voltage-clamp recordings. Voltage-clamp recordings suppress contributions of inner segment conductances, which are activated by voltage changes, and the contributions of membrane capacitance, which can slow changes in voltage. Cone capacitance, for example, could differ across eccentricity due to the longer axons of foveal cones compared to peripheral cones (Hsu et al., 1998). The kinetics of responses of foveal and peripheral cones differed in voltage-clamp recordings both for filters derived from responses to Gaussian noise (Figure 7D; polarity opposite to current clamp recordings in Figures 7B and C) and flash responses (not shown). Response differences were quantitatively very similar to those measured for voltage responses (i.e., in current-clamp recordings; Figures 7B and 7C). Current-clamp responses (Figures 7B and 7C) were slightly slowed compared to voltage-clamp responses (Figure 7D), likely due to membrane capacitance. The persistence of the differences in the kinetics of foveal and peripheral cone responses under voltage clamp (Figure 7D) indicates that they are not due to inner segment conductances.

Network interactions, as mentioned above, could also vary across retinal regions and hence contribute to differences in the kinetics of cone responses across eccentricity. To test for a contribution of horizontal cell feedback to cones, we suppressed horizontal cell responses with the AMPA-receptor antagonist NBQX (Dunn et al., 2007). Voltage responses of foveal and peripheral cones changed little following suppression of horizontal cell activity (Figures 7E and 7F). This suppression is consistent with work indicating that horizontal cell feedback modulates a conductance with a reversal potential close to the cone photoreceptor resting potential, and hence impacts transmitter release more noticeably than the cone light response (Endeman et al., 2012; Verweij et al., 2003). To test for a role of gap junctions, we performed voltage clamp recordings from foveal cones after omitting ATP and GTP from the intracellular solution. Omission of ATP and GTP suppresses the phototransduction cascade, causing the cGMP-gated channels to close in the recorded cone while retaining contributions from neighboring (coupled) photoreceptors. This manipulation strongly suppresses light responses in peripheral cones (Angueyra and Rieke, 2013). To further attenuate phototransduction, we included a poorly hydrolyzed GTP analog, GTPγS (Sampath and Rieke, 2004) in the intracellular solution during voltage clamp recordings from foveal cones. We also eliminated horizontal cell feedback by bath application of NBQX (Figure S7). These manipulations strongly suppressed light responses of foveal cones (Figure S7). The sensitivity of the light response to suppression of phototransduction suggests a small contribution of gap junctions between cone photoreceptors.

These experiments collectively suggest that the dramatic kinetic differences between foveal and peripheral cones originate from differences in the phototransduction process in the cone outer segment.

DISCUSSION

Despite the importance of the fovea for how we see, we know very little about the cellular and synaptic basis of its operation. Here, we reveal two important features that distinguish signaling in the fovea from that in the peripheral retina. First, a combination of intracellular recordings and structure-function analyses show that foveal MGCs receive minimal inhibitory synaptic input. This is unlike peripheral MGCs, which receive abundant inhibitory input. Second, we find that the responses of the very first retinal neurons—the cone photoreceptors—differ dramatically between the foveal and peripheral retina. These results, detailed below, provide a glimpse into the mechanistic specializations of the fovea, and provide a simple explanation for foveal/peripheral differences in sensitivity to rapidly-varying light inputs.

A Unique Mode of Synaptic Integration by Foveal MGCs

Understanding how excitatory and inhibitory inputs are integrated to shape neural responses is an extremely active research area (reviewed by Isaacson and Scanziani, 2011). Most circuits operate in a mode in which inhibition and excitation are balanced (I/E balance), and signaling relies on a momentary imbalance of the two.

Given the general importance of I/E balance, we were surprised to find that responses of MGCs in the primate fovea were minimally shaped by either pre- or post-synaptic inhibition. Foveal MGCs are unique in receiving input from single cone photoreceptors, and integration without inhibition ensures that cone signals are minimally altered before transmission to cortical neurons. The lack of inhibitory input is not shared by all foveal ganglion cell types. Thus, parallel and intertwined circuits in the fovea exhibit very different modes of synaptic integration. This physiological observation had a direct anatomical correlate (see below) and has important implications for the sophistication of foveal computations.

Integration of excitatory and inhibitory signals shapes computations performed by most sensory and cortical circuits—including feature selectivity, gain control, and coincidence detection (reviewed by Isaacson and Scanziani, 2011). A theme of recent retinal work has been the surprising computational specializations of non-foveal retinal circuits (Baccus et al., 2008; Buldyrev and Taylor, 2013; Farrow et al., 2013; Manookin et al., 2008; Münch et al., 2009; Sivyer et al., 2010; Venkataramani and Taylor, 2010; Venkataramani et al., 2014; Wei et al., 2011; Yonehara et al., 2011). These computations rely on synaptic inhibition generated by the diverse population of inner retinal amacrine cells (Farrow et al., 2013; Münch et al., 2009; Protti et al., 2014; Venkataramani et al., 2014; Wei et al., 2011; Yonehara et al., 2011). Our work here shows that this picture does not apply to computations by foveal MGCs.
Figure 7. Cone Signals Are Slower in the Fovea than Periphery

(A) Schematic illustrating the morphology of a cone photoreceptor. Whole cell recordings were performed at the inner segment as depicted in the schematic. A cell-fill image of a foveal (Fov) and peripheral (Peri) cone. Please note the dramatic difference in cone morphology, particularly the long axon of the foveal cone. A short excerpt of a foveal cone voltage response to a time varying stimulus. Cone voltage responses (B, C and E) are depicted with an opposite polarity compared to current responses (D).

(B) Average time-reversed linear filters for foveal and peripheral M and L cone voltage responses. Shaded regions represent SEM. Quantification of the time to peak for the linear filters yields a mean of 31.5 ± 2 ms and 58.4 ± 9 ms for peripheral and foveal M cones, respectively. The mean time to peak for peripheral and foveal L cones was 33.2 ± 3 ms and 59.3 ± 4 ms, respectively. Error bars represent s.d.

(legend continued on next page)
Instead, the reduced mechanistic repertoire shaping responses of foveal MGCs suggests that they lack the computational sophistication of their peripheral or wide-field counterparts. This distinction will be an important consideration in how visual computations are partitioned between retinal and cortical circuits in the fovea and periphery and between different foveal ganglion cell types.

**Low Level of Expression of Inhibitory Receptors by Foveal MGCs Supports Unique Mode of Synaptic Integration**
Foveal MGCs not only received minimal light-driven inhibitory input, but also expressed very few inhibitory receptors compared to excitatory receptors (low I/E ratio, see Figure 4). This was surprising given the higher I/E synapse ratios estimated previously from counts of ribbon and non-ribbon synapses onto foveal MGC dendrites using electron microscopy (Calkins et al., 1994; Calkins and Sterling, 1996; Kolb and Dekorver, 1991). These two types of experiments, however, measure different anatomical aspects of the synapses and hence are not necessarily contradictory.

Three possible issues could contribute to this difference, including (1) non-ribbon excitatory synapses, (2) differences in synaptic structure, and (3) postsynaptic receptors.

**Non-ribbon Excitatory Synapses**
Amacrine synapses were identified in electron microscopy studies based on the absence of ribbons (Calkins et al., 1994). MGCs could receive non-ribbon excitatory contacts, e.g., from glutamatergic amacrine cells (Lee et al., 2014; Marshak et al., 2015), and bipolar cells could make some conventional synapses on the dendrites of MGCs (Kolb and Dekorver, 1991).

**Differences in Synaptic Structure**
The MGC is contacted by a single midget bipolar cell with a claw-like synaptic terminal containing 25–50 ribbons (Calkins et al., 1994) packed into a small volume (~70 μm³). This is akin to the giant mossy fiber bouton contacting the CA3 pyramidal neuron (Bischofberger et al., 2006), a very robust excitatory synapse. The unusual anatomy of the bipolar-MGC synapse may produce strong excitatory synaptic input to the MGC that cannot be inferred by counting synapses.

**Postsynaptic Receptors**
Postsynaptic receptor densities at individual amacrine and bipolar synaptic contacts could differ. The differences we observe in dendritic volume occupancy of inhibitory and excitatory receptors support this possibility. However, we were not able to measure the size of the receptor clusters with confidence due to the limited resolution of light microscopy. Development of techniques that allow the simultaneous labeling of excitatory and inhibitory receptors on MGC dendrites at electron microscopy resolution could provide the needed estimates of receptor cluster size. Independent of the reason, the functional I/E ratios correlated more strongly with I/E ratios based on receptor expression than counts of presynaptic structures. This is an important result given the vast amount of electron microscopy data available now and in the near future and the desire to draw functional inferences from these data.

In many neural circuits, including the peripheral retina, inhibitory interneurons establish synapses before primary excitatory neurons (Ben-Ari, 2014; Hendrickson, 1994, 1996; Hoon et al., 2014). Interestingly, foveal synapses develop in the opposite sequence: excitatory ribbon synapses between bipolar cells and ganglion cells form prior to amacrine inhibitory synapses (Hendrickson, 1994, 1996; Hoon et al., 2014). This sequence suggests that the foveal midget circuitry from its inception is wired such that the excitatory input from single cones dominates synaptic integration.

**A Simple yet Surprising Mechanistic Link to Human Perception**
A wealth of human psychophysical experiments have shown that sensitivity to rapidly varying light inputs is higher in the periphery than in the fovea (Rovamo and Raninen, 1988; Tyler, 1985; Waugh and Hess, 1994). This difference is not subtle: the estimated time constant based on human perception differs ~2-fold for foveal and peripheral vision (Tyler, 1985). The reduced temporal sensitivity of foveal vision is an important consideration in the design of devices such as visual displays and will be an important consideration for visual prosthetics.

We were surprised to find that differences in inhibitory synaptic input to foveal and peripheral MGCs contributed minimally to differences in temporal sensitivity. Instead, post-synaptic inhibition controls the gain of peripheral MGC responses. This form of gain control may be needed in the periphery, where visual signals are pooled from several cone photoreceptors, but not in the fovea where MGCs receive (indirect) input from single cones.

Differences in temporal sensitivity could instead be explained by differences in the kinetics of the responses of the cone photoreceptors themselves. Indeed, responses of foveal cones were slower than those of peripheral cones. Importantly, cone photoreceptor responses (Figure 7), ganglion cell responses (Figure 1 and Solomon et al., 2002) and perception (Tyler, 1985) all show a near-identical 2-fold difference in kinetics between the fovea and periphery. The similarity of these measures suggests that the cones themselves account for the well-
established foveal/peripheral differences in sensitivity to rapid variations in light inputs. Consequently, foveal cones integrate their inputs over longer times—reducing sensitivity to rapidly-varying inputs while improving sensitivity to slowly-varying inputs. Peripheral cone circuits, in which MGCs receive input from multiple cones, improve sensitivity by averaging across multiple converging cones (Goodchild et al., 1996), and this spatial integration may obviate the need for temporal integration in the cones themselves.

Differences in the kinetics of foveal and peripheral cone responses appear to originate within the phototransduction cascade rather than from inner segment or network effects. These differences may complement anatomical differences between the cones (Tyler, 1985). For instance, the axons of foveal cones are ~10-fold longer than those of peripheral cones; the slower kinetics of foveal cone responses may be matched to the filtering associated with propagation of signals down their long axons (Hsu et al., 1998).

In summary, our study builds on previous studies in retina and human perception in the following ways: (1) Functional recordings from MGCs in the fovea allowed us to directly measure light-driven excitatory and inhibitory synaptic inputs to the MGCs and reveal a unique mode of synaptic integration without inhibition. (2) Anatomical analyses show that foveal MGCs express few inhibitory receptors compared to their peripheral counterparts, corroborating the physiological findings. (3) Comparison of responses of foveal and peripheral cone photoreceptors revealed a surprising 2-fold difference in kinetics and a simple explanation for the lower temporal sensitivity of foveal vision.

**STAR METHODS**

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

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**SUPPLEMENTAL INFORMATION**

Supplemental Information contains seven figures and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2017.01.005.

**AUTHOR CONTRIBUTIONS**

R.S., F.R. and J.B. performed the electrophysiological recordings and analysis. M.H. performed the immunohistochemistry, biolistics and all the anatomical analysis. H.O. helped with the biolistic experiments and image analysis. R.S., M.H. and F.R. conceived of experiments and analyses. R.S., M.H., R.O.W. and F.R. wrote the paper.

**ACKNOWLEDGMENTS**

We thank E.J. Chichilnisky, G. Horwitz, G.D. Field and J.M. Anguerya for helpful comments. We thank C. Puller for helpful discussions and suggestions. We thank R. Enz and H. Wässle for the GABAA antibody, T. Jacob for the GABAa1-fluorescent protein construct and T. Yoshimatsu for generating the GABAa1-YFP fusion construct. We thank M. Cafaro, S. Cunningham, and P. Newman for excellent technical assistance. This work is supported by the Human Frontier Science Foundation (long-term fellowship to R.S.), Knights Templar Eye Foundation (career starter grant to M.H.), Simons Foundation (324759 to F.R.), ARCS foundation (J.B.), HHMI (F.R.) and NIH grants (EY026070 to R.S., EY11850 to F.R. and EY10699 to R.O.W.).

Received: August 31, 2016
Revised: November 21, 2016
Accepted: January 5, 2017
Published: January 26, 2017

**REFERENCES**


## STAR★METHODS

### KEY RESOURCES TABLE

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to the Lead Contact Fred Rieke (rieke@uw.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Retinas from both male and female Macaques (M. mulatta, M. nemestrina, and M. fascicularis) aged between 4–21 years old were obtained through the Tissue Distribution Program of the Regional Primate Research Center at the University of Washington following procedures approved by the Institutional Animal Care and Use Committee. Animals were housed in primate center facilities, cared for by their veterinary staff and were in good health. For each dataset, we collected data from at least 3 retinas from at least 3 animals.

METHOD DETAILS

Tissue Preparation and Cell Identification

Electrical recordings from foveal (<1 mm from foveal pit), central (2–4 mm from foveal pit) and peripheral (>6 mm from foveal pit) primate retina followed approaches used previously for peripheral retinal recordings (Cafaro and Rieke, 2013). For all recordings, a piece of retina was isolated from the choroid and pigment epithelium and mounted flat in a recording chamber. The retina was mounted ganglion cell side up for ganglion cell recordings and photoreceptor side up for cone recordings. The retina was superfused with warmed (∼34°C) and oxygenated (5% CO2/95% O2) Ames solution (Sigma). The foveal pit was clearly identifiable when focusing on the cone array (Figure 1).

All functional recordings were performed at a background light level of ∼24,000 absorbed photons per cone per sec (R*/cone/s). Responses in the rods themselves were saturated at 10-fold lower light levels (data not shown), and hence the rods should contribute minimally to the reported ganglion cell or cone responses. Of note, the density of rod photoreceptors, which are abundant in central and peripheral retina, drops precipitously near the fovea—in fact, the central fovea (<0.3 mm from the foveal center) is devoid of rods altogether and has the highest cone density (Kolb et al., 2002; Wässle et al., 1995).

MGCs were initially identified based on their relatively sustained responses to light steps compared to the more transient responses of wide-field ganglion cells (Cafaro and Rieke, 2013; De Monasterio and Gouras, 1975). In central and peripheral retina identification was aided by the morphology and size of the MGC somas (Crook et al., 2011; Dacey, 1993; Kolb and Marshak, 2003). Cell identification was confirmed by fluorescence imaging (using Neurobiotin or Alexa 594 dye) as shown in Figure 1. The foveal MGCs have a characteristic small dendritic arbor of 5–10 μm (Figure 1) compared to 20 μm arbors of foveal wide-field ganglion cells (Figure 4).

The baseline spike rate of MGCs at the three retinal locations at a mean luminance of 4,000 R*/cone/s ranged from ∼5–30 Hz. Before initiating data collection, the sensitivity of the retina was determined from the spike responses of MGCs to a full field 100% contrast light step from a background of ∼4,000 R*/cone/s. Data were collected from cells in which this stimulus produced a peak increase in firing rate of >40 spikes/s in foveal and central retina and >100 spikes/s in the peripheral retina. This contrast gain—i.e., 0.4 spikes%/contrast—is within a factor of 2 of contrast gains from in vivo recordings of foveal MGCs at similar mean light levels (Benardete and Kaplan, 1999a, b; Kaplan and Benardete, 2001; Kaplan and Shapley, 1986; Lee et al., 1990; Purpura et al., 1988; Solomon et al., 2002). Recordings from the fovea were pooled across both L and M-cone driven ON-type MGCs. OFF-type MGCs occurred far less frequently near the top of the ganglion cell layer in the fovea (Yin et al., 2014); this restricted our ability to collect extensive data from foveal OFF MGCs and hence we report results predominantly from ON MGCs.

Light Stimulation

Light stimuli were delivered from LEDs with peak spectral outputs at 470 nm or 640 nm. The stimuli were delivered through the microscope condenser and glass bottom of the recording chamber. Light intensities are reported as isomerizations per cone per second (R*/cone/s), calculated using the calibrated LED output and spectra, the photoreceptor spectral sensitivity (Baylor et al., 1987), and a cone collecting area of 0.37 μm² (Schnapf et al., 1990). Full-field stimuli covered a 500-μm-diameter spot on the retina centered on the cell of interest. The spot size covered both the center and surround of the MGC receptive field at each of the retinal locations. Gaussian noise stimuli had 50% contrast (SD/mean) and 0–60 Hz bandwidth. To analyze responses to these stimuli, we computed...
the average stimulus preceding an action potential (the spike-triggered average or STA) or the corresponding linear filter for ganglion cell excitatory synaptic currents and cone photovoltages/photocurrents.

**Patch-Clamp Recordings**

Whole-cell patch-clamp recordings were made using a Multiclamp 700B amplifier (Molecular Devices). Pipettes for voltage-clamp recordings were filled with a Cs-based internal solution containing (in mM): 133 potassium aspartate, 10 KCl, 10 EGTA, 5 Mg-ATP, 0.5 Tris-GTP, and 2 QX-314, pH ~7.3, ~280 mOsm. An inhibitory cocktail (20 µM GABAzine, 50 µM TPMPA and 2 µM strychnine; Tocris) was added to the perfusion solution to block GABAA, GABAC and glycine receptors as indicated in Figures 2 and 6 and Figure S5. Voltage-clamp recordings were made using pipettes of 5 MΩ resistance for ganglion cells and of 10–14 MΩ resistance for cones. Whole-cell patch clamp recordings were performed from the inner segments of cone photoreceptors as described previously (Angueyra and Rieke, 2013). The internal solution contained (mM): 133 potassium aspartate, 10 KCl, 10 HEPES, 1 MgCl2, 4 ATP, 0.5 GTP, pH ~7.3, ~280 mOsm (Angueyra and Rieke, 2013). For cone recordings in Figure 7E and 7F, 10 µM NBQX (Tocris) was added to the bath perfusion solution. GTPγS (Sigma) was added at a concentration of 50 µM to the intracellular solution (Figure S7). Unless specified otherwise, reported voltages have been corrected for a ~10 mV liquid junction potential.

The resting membrane potential of foveal MGCs was ~−55 mV, similar to that of peripheral MGCs. For measurements of excitatory and inhibitory synaptic currents, cells were voltage-clamped at several holding potentials between −80 and +30 mV. Light steps were delivered while excitatory and inhibitory reversal potentials were assessed empirically by focusing on time points dominated by either excitatory or inhibitory input, and identifying holding potentials that minimized each (near −60 and +10 mV). The cell was held at the estimated excitatory reversal potential when inhibitory currents were recorded and at the estimated inhibitory reversal potential when excitatory currents were recorded. For measurement of current-voltage curves (see Figure S5), light stimuli were delivered starting 1–2 s after a voltage step to allow any transient change in current to subside.

Spontaneous IPSCs were apparent immediately after depolarizing a MGC to the reversal potential for excitatory input, and spontaneous IPSCs were stable in amplitude throughout the time the cell remained depolarized (as much as 20 min in some cases; Figure S5). Thus changes in intracellular chloride following depolarization did not appear to attenuate inhibitory currents. Spontaneous IPSCs in foveal and peripheral MGCs had similar amplitudes and kinetics (Figure S5C).

**Caged Compounds**

MNI-glutamate (2 mM, Tocris Biosciences) and RuBi-GABA (5–10 µM, R&D Systems) were puffed onto the tissue from a glass pipette. Photolysis of MNI-glutamate and RuBi-GABA was performed with light from a mercury arc lamp (X-cite, Excelitas Technologies). Light was delivered to the entire tissue in 50 ms pulses through the data-acquisition software (Symphony, Symphony-DAS, Seattle, USA). MGCs were identified based on their spike responses as shown in Figure 1 for fovea and periphery. Light-evoked responses were blocked by perfusing with a solution containing cadmium chloride (200 µM). Responses to uncaged GABA and glutamate were measured at the excitatory and inhibitory reversal potentials. A limitation in quantitative interpretation of these experiments is that previous studies have shown that MNI-Glutamate can inhibit GABA receptors (Finoc et al., 2009; Maier et al., 2005).

This could explain why the amplitude of GABA-mediated current is smaller than expected from the immunohistochemical analysis for both foveal and peripheral MGCs. Nonetheless, the relative amplitude of inhibitory to excitatory current is larger for peripheral MGCs compared to their foveal counterparts, consistent with the rest of our experiments.

**Dynamic Clamp**

Synaptic input was mimicked and manipulated during dynamic clamp experiments (Cafaro and Rieke, 2013). Cells were current-clamped using pipettes filled with a K-based internal solution (containing, in mM: 122 K-Aspartate, 10 KCl, 1 CaCl2, 1 MgCl2, 2 EGTA, 10 HEPES, 4 Mg-ATP, and 0.5 Tris-GTP, pH 7.2 with KOH, 280 mOsm). Current injected into a cell (I) during dynamic clamp experiments was calculated as

\[ I(t) = G_{\text{exc}}(t)(V(t) - E_{\text{exc}}) + G_{\text{inh}}(t)(V(t) - E_{\text{inh}}) \]

where \( G_{\text{exc}} \) and \( G_{\text{inh}} \) are the conductances recorded during light stimulation, \( V \) is the membrane potential, and \( E_{\text{exc}} \) (\( E_{\text{inh}} \)) is the reversal potential for excitatory (inhibitory) synaptic input. Current injected at time \( t \) was calculated from the measured voltage at time \( t - \Delta t \), with \( \Delta t = 100 \) µs. \( E_{\text{exc}} \) was 0 mV and \( E_{\text{inh}} \) was −90 mV. Excitatory and inhibitory synaptic conductances in response to Gaussian noise stimuli were measured near-simultaneously from peripheral MGCs using an alternating voltage approach (Cafaro and Rieke, 2010). The excitatory conductances from a foveal MGC to similar light stimuli were used for the dynamic clamp experiments. Upon initiating a dynamic clamp experiment, the amplitudes of the excitatory and inhibitory conductances were scaled by a common factor, while keeping their ratio constant, to achieve spike numbers near those recorded during light stimulation. As needed, a modest constant current (<100 pA) was injected to maintain a spontaneous firing rate of 5–20 spikes/s.

The dynamic clamp approach described in Equation 1 neglects the NMDA conductances that are present in the inputs to MGCs. Previous studies have shown that NMDA conductances, however, have little impact on peripheral MGCs’ spike responses under the conditions of our experiments (Cafaro and Rieke, 2013). Block of NMDA receptors with L-AP4 similarly had little effect on the spike responses of foveal MGCs to Gaussian noise stimuli (data not shown).
The STAs in Figure 5C are one component of a linear-nonlinear (LN) model. A change in the amplitude of the STA corresponds to a change in the x axis scaling of the static nonlinearity (Chichilnisky, 2001; Kim and Rieke, 2001). To compare STA amplitudes, the x-axes of the nonlinearities were scaled to produce maximal overlap and the STA amplitudes correspondingly adjusted. This associates any change in response with a change in the STA.

Immunohistochemistry

MGCs were filled intracellularly with a solution containing 5 mM neurobiotin and retinas were immersion fixed for either 15 min (for GluA3, GABA\(_a\)-z1 and gephyrin labeling) or 30 min (for GABA\(_c\)-labeling) with 4% paraformaldehyde in 0.1M phosphate buffered saline (PBS, pH 7.4). Post-fixation, the tissue was washed in PBS and then freeze-cracked in 30% sucrose (made in PBS) to facilitate antibody penetration. Antibody incubation was carried out in a blocking solution containing 5% donkey serum and 1% triton (Sigma) in PBS for 1 week at 4°C. Antibodies used were directed against the GluA3 subunit of AMPA receptor (goat polyclonal, 1:500, Santa Cruz Biotechnology), Gephyrin (mouse monoclonal mab3b11, 1:1000, Synaptic Systems), GABA\(_a\) receptor z1 subunit (mouse monoclonal mab339, 1:500, Millipore). As a control for non-specific association of the receptor signal within MGC dendrites, we labeled for GABA\(_a\) receptors (Enz et al., 1996) using an antibody against GABA\(_c\) receptor b subunit (rabbit polyclonal, 1:500, kindly provided by R. Enz and H. Wässle), known to localize at bipolar cell terminals, and found in negligible amounts on MGC dendrites (Figure S6). Secondary antibodies were anti-isotypic Alexa Fluor conjugates (1:1000, Invitrogen or Jackson ImmunoResearch). Secondary antibody incubation was carried out in blocking solution for 3–4 hr at room temperature, and after washes in PBS the retinas were subsequently mounted in Vectashield (Vector labs). To amplify the neurobiotin signal, streptavidin conjugated to Alexa Fluor 568 (1:1000, Invitrogen) was included with the secondary antibody.

Biolistic Transfection

Gold particles (1.6 μm diameter; 12.5 mg; Bio-Rad) were coated with DNA plasmids encoding tdTomato (24 μg), PSD95-CFP or PSD95-YFP (12 μg), GABA\(_a\)-z1-YFP (modified version of GABA\(_a\)-z1 tagged fusion constructs (Connolly et al., 1996; Jacob et al., 2008)—a generous gift from T. Jacobs) (19 μg), GABA\(_a\)-z2 (20 μg) and GABA\(_b\)-z2/3 (19 μg) under the control of the cytomegalovirus promoter. The particles were delivered to retinal ganglion cells in whole-mount retinas using a Helios gene gun (Bio-Rad). Ongnotypic culturing and transfection of retinas was performed as described previously (Lye et al., 2007; Percival et al., 2014). In brief, pieces of retina were mounted ganglion cell side up onto a 0.4 mm Millicell tissue culture inset (Millipore). Transfected retinas were incubated in a 95% O\(_2\)/5% CO\(_2\) environment for at least two days in sterile Ames medium at 32°C. The retinas were then immersion fixed in 4% paraformaldehyde and mounted on glass slides directly or used for immunostaining.

Imaging and Analysis

Image stacks of whole mounted retinas were acquired on an Olympus FV1000 laser scanning confocal microscope. The tissue was imaged using a 1.35 NA 60X oil immersion objective at a xyz voxel size of 0.069, 0.069, 0.3 μm. For acquiring overview images, a 0.85 NA 20X oil immersion objective was used. Each optical plane was averaged three to four times (Kalman filter). Raw image stacks were processed using MetaMorph (Molecular Devices) and Amira (FEI Visualization Sciences Group). All acquired images were median filtered in MetaMorph before their 3D visualization in Amira.

To quantify the percentage occupancy of receptor signal with the MGC dendritic mask, the dendrites of the MGC were at first masked in 3-dimensions using the Labelfield function in Amira such that pixels outside the mask were set to a value of zero. The MGC dendritic mask was multiplied by the receptor (GluA3, PSD95, Gephyrin or GABA\(_a\) tagged fusion construct) channel to isolate the receptor signal exclusively within the dendritic mask. A constant threshold at 4 standard deviations above the noise peak (see Figure S3) was selected to exclude background fluorescence and retain only pixels representing the signal (e.g., receptors). The threshold was chosen by fitting the distribution of pixel values for each immunolabeling with a gamma distribution to determine the peak and standard deviation of the noise (background) pixels (Figure S3). The volume of the receptor pixels above the threshold was expressed as a percentage of the total volume occupied by the pixels within the MGC dendritic mask.

To assess if the low I/E ratio in foveal MGCs is due to a lower overall amount of inhibitory receptor protein expression in foveal MGCs rather than size differences between foveal and peripheral MGC dendritic arbors, we compared the volume occupancy of the receptor signal of peripheral MGCs in the entire dendritic arbor with the receptor occupancy estimated in smaller stretches of peripheral MGC dendritic volume (70 μm\(^3\)) comparable to the arbor of foveal MGCs (Figure S1). The anatomical I/E ratio for peripheral MGCs was similar in both cases (full arbor versus smaller volume), indicating that the low I/E ratio of inhibitory receptors in foveal MGCs is not merely due to the foveal MGC dendritic arbor being smaller than the dendritic arbor for peripheral MGCs.

To assess the % colocalization between PSD95-YFP and GluA3 immunolabeling or GABA\(_a\)-YFP clusters and Gephyrin immunolabeling (see Figure S1), PSD95-YFP or GABA\(_a\)-YFP signal within the MGC dendritic mask was first isolated using the Amira software as described above. Thereafter, PSD95-YFP or GABA\(_a\)-YFP clusters were assessed for colocalization with GluA3 or Gephyrin immunolabeling respectively using a custom written MATLAB (Mathworks, USA) routine. GluA3 and Gephyrin signals were thresholded (similar to thresholds set in Figure S3) to exclude background fluorescence. For each PSD95-YFP punctum within the MGC dendrite the proportion of PSD95 pixels that also contain GluA3 signal was determined. Similarly, for each GABA\(_a\)-YFP punctum the proportion of GABA\(_a\) pixels that also contain Gephyrin signal was determined. A PSD95 punctum was considered as “colocalized” with the GluA3 signal, if the GluA3 signal occupied more than 20% of all the pixels representing the PSD95 punctum. Scoring for all the PSD95
puncta yielded a % colocalization (Figure S1). The % of GABA$\alpha_{1}$-YFP clusters colocalized with Gephyrin was calculated the same way (Figure S1). As a control, the GluA3 or Gephyrin channel were flipped 90° and their % random colocalization with PSD95-YFP and GABA$\alpha_{1}$-YFP clusters determined (Figure S1).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

We used the nonparametric Wilcoxon-Mann-Whitney rank sum test for all the statistical analysis. Error bars indicate SD. Statistical parameters including the exact value of n, precision measures (mean ± SD) and statistical significance are reported in the figures and in the figure legends (see individual sections). The significance threshold was placed at $\alpha = 0.05$ (n.s., $p > 0.05$; *, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.001$).
Figure S1. Transient Expression of Fluorescently-Tagged PSD95 and GABA_A Receptors on MGC Dendrites Largely Colocalizes with Endogenous Synaptic Components, Related to Figure 4

(A) Maximum intensity projection showing PSD95-YFP puncta (green) on a labeled (CMV-tomato) peripheral MGC dendrite. Immunostaining with an antibody against GluA3 subunit of AMPA receptors (red) shows that GluA3 puncta colocalize with PSD95-YFP.

(B) Maximum intensity projection showing GABA_A-YFP receptor puncta (green) on a labeled (CMV-tomato) peripheral MGC dendrite. Immunostaining with antibody against gephyrin (red) shows that the gephyrin signal colocalize with GABA_A-YFP receptors.

(C) Percent colocalization of the GluA3 signal within PSD95-YFP punctas (see STAR Methods) yields a mean value of 68% ± 11% given that ~70% of AMPA receptors are labeled by the GluA3 antibody in primate retina (Abbott et al., 2012). Flipping the GluA3 channel by 90° yields a significantly reduced (random) colocalization value of 5% ± 3%. For GABA_A-YFP puncta, 71% ± 2.0% colocalized with gephyrin, which is consistent with previous observations that gephyrin is not at all GABA_A receptor synapses (Abbott et al., 2012). Flipping the gephyrin channel by 90° yielded a significantly reduced colocalization value of 32% ± 7%. Together, these results indicate that most of the PSD95-YFP and GABA_A-YFP are properly targeted to their appropriate synaptic locations. Error bars indicate s.d.

(D) Quantification of the % volume occupancy of GABA_A-α1YFP receptors relative to PSD95-CFP signal for peripheral MGCs either within the entire tdTomato filled dendritic volume (full arbor) or a smaller dendritic volume (70 μm³) that is comparable to the arbor of foveal MGCs. Pairwise comparison shows no significant difference in the IGABA_A/EPSD-95 ratio (Full arbor - mean = 0.93 ± 0.11; 70 μm³ - mean = 1.04 ± 0.22).
Figure S2. GluA3, Gephyrin, and GABA\textsubscript{\alpha}1 Antibody Labeling across Eccentricities, Related to Figure 4

(A) Raw images of neurobiotin (NB) filled MGCs (white) immunostained with antibodies against GluA3 (green) and Gephyrin or GABA\textsubscript{\alpha}1 receptors (red) across eccentricities.

(B) Figure 2C replotted to show cells (color coded) where the physiological I/E was measured before the morphological analysis. Cells depicted by red circles were immunostained for Gephyrin and GluA3 and cells depicted by purple circles were immunolabeled for GABA\textsubscript{\alpha}1 and GluA3.
Figure S3. Threshold Selection of Immunoreactive Signal for Analysis of Percentage Volume Occupancy, Related to Figure 4

(A) Raw GluA3, gephyrin and GABAA xr1 signal within the foveal MGC dendritic processes illustrated in Figure 4. Panels below show the corresponding 3-dimensional surface plot of the immunoreactive signal within each mask. The red dotted line indicates the threshold set to quantify pixel volume for each label. (B) Top panel displays the raw GluA3 and gephyrin immunolabeling in fovea and periphery. (C) Middle panel shows the distribution of all the fluorescent pixels and the noise peak (black arrow). The red curve is a fit with a Gamma function and the threshold is set at 4 s.d. above the noise peak (dotted line). Bottom panel shows the GluA3 and gephyrin pixels detected after using this threshold.
Figure S4. Immunostained GABA<sub>α</sub>1 Receptors within Biolistically Labeled Foveal Ganglion Cells, Related to Figure 4

(A) Maximum intensity projections of GABA<sub>α</sub>1 (red) labeling together with CMV-tomato (blue) expressing foveal MGC and foveal wide-field ganglion cell. The GABA<sub>α</sub>1 immunoreactive signal at the soma of the MGC is negligible (top) but shows a ring-like pattern around the soma of wide-field ganglion cells (bottom, arrows) in the fovea. This is consistent with previous studies that show a selective labeling of GABA<sub>α</sub>1 around the soma of foveal wide-field ganglion cells, such as Parasol cells and a fraction of other wide-field ganglion cells (Grunert et al., 1993).

(B) Comparison of the GABA<sub>α</sub>1 receptor (red) signal within the dendrites of a foveal MGC and foveal wide-field ganglion cells (CMV-tomato expression, blue). Quantification of % volume occupancy of GABA<sub>α</sub>1 signal within the dendritic mask yields a mean value of 1.03% ± 0.65% for foveal MGCs and a mean value of 7.89% ± 2.0% for wide-field ganglion cells indicating relatively less GABA<sub>α</sub>1 expression by foveal MGCs compared to foveal wide-field ganglion cells.
Figure S5. Current-Voltage Relationship and Time-Dependent Analysis of Inhibitory Synaptic Currents of ON Foveal MGCs, Related to Figure 2

(A) Average current-voltage (IV) curves from voltage clamped ON foveal MGCs (n = 2 cells; holding potential not corrected for liquid junction potential) before (black trace) and after (red trace) bath application of GABA and Glycine receptor blockers (GABAzine, TPMPA and Strychnine). Peak amplitudes in response to a 10 ms light flash are plotted in the IV curves.

(B) Comparison of first two trials soon after depolarizing to the excitatory reversal potential (within 5–10 s) versus later trials. Average of eight foveal MGCs.

(C) Analysis of spontaneous IPSCs in foveal and peripheral MGCs. Average spontaneous IPSCs from foveal and peripheral MGCs (n > 100 IPSCs). Inset shows comparison of normalized trace.

(D) Time-dependent analysis of average spontaneous IPSCs from a foveal MGC calculated from trials recorded 20 min apart (n > 100 IPSCs).
Figure S6. Antibody Labeling against GABAC Receptors in MGCs, Related to Figure 4
Maximum intensity projections of a foveal and central MGC dendritic arbor filled with neurobiotin (NB, blue) showing negligible immunoreactive GABAC signal (red) within the dendritic mask.
Figure S7. Gap-Junctional Coupling between Photoreceptors Probed during Voltage Clamp Recordings, Related to Figure 7
Average responses of foveal cones (n = 4) under voltage clamp in absence of ATP and GTP and presence of 50 μM GTPγS from the intracellular pipette solution. Response to a 300% contrast light flash (10 ms), from a background of 4000 R'/cone/sec soon after breaking into the cell (black, initial) and 5 min after (red). 10μM NBQX was bath applied after recording initial responses.