# Neural sensitization improves encoding fidelity in the primate retina

<sup>2</sup> Todd R. Appleby <sup>1,2,3</sup> and Michael B. Manookin <sup>2,</sup>	3*
<sup>3</sup> <sup>1</sup> Graduate Program in Neuroscience, University of Washington, Seatt	le, WA 98195, USA
<sup>4</sup> <sup>2</sup> Department of Ophthalmology, University of Washington, Seattle	, WA 98195, USA
<sup>5</sup> <sup>3</sup> Vision Science Center, University of Washington, Seattle, WA	98195, USA
<sup>6</sup> *Correspondence: manookin@uw.edu	

# ABSTRACT

An animal's motion through the environment can induce large and frequent fluc-7 tuations in light intensity on the retina. These fluctuations pose a major challenge 8 to neural circuits tasked with encoding visual information, as they can cause cells to 9 adapt and lose sensitivity. Here, we report that sensitization, a short-term plasticity 10 mechanism, solves this difficult computational problem by maintaining neuronal sen-11 sitivity in the face of these fluctuations. The numerically dominant output pathway 12 in the macaque monkey retina, the midget (parvocellular-projecting) pathway, under-13 goes sensitization under specific conditions, including simulated eye movements. Sen-14 sitization is present in the excitatory synaptic inputs from midget bipolar cells and is 15 mediated by presynaptic disinhibition from wide-field amacrine cells. Direct physio-16 logical recordings and a computational model indicate that sensitization in the midget 17 pathway supports accurate sensory encoding and prevents a loss of responsiveness 18 during dynamic visual processing. 19

## INTRODUCTION

The fundamental constraints on sensory coding require that neural circuits adjust 20 their outputs based on the statistical properties of their recent inputs (Srinivasan et 21 al., 1982; Barlow, 1961; Laughlin, 1981). Neurons respond to dynamic inputs using 22 two distinct strategies—adaptation and sensitization. Adapting cells respond to strong 23 stimulation by decreasing their sensitivity and this decrease in responsiveness can 24 persist for several seconds after the stimulus intensity decreases (Baccus and Meister, 25 2002; Carandini and Ferster, 1997; Kim and Rieke, 2001; Laughlin, 1981; Manookin 26 and Demb, 2006; Smirnakis et al., 1997; Solomon et al., 2004). Thus, adapting cells are 27 relatively insensitive to weak stimuli occurring during these transition periods. Sensi-28 tizing cells show the opposite pattern-increasing their responsiveness at these transi-29 tions (Kastner and Baccus, 2011; Kastner and Baccus, 2013; Nikolaev et al., 2013). For 30 this reason, adaptation and sensitization are commonly thought to constitute opposing 31 and complementary forms of short-term neural plasticity (Kastner and Baccus, 2011; 32 Kastner and Baccus, 2013). 33

This hypothesis requires that a sensitizing cell type have an adapting counterpart that 34 encodes common information (Kastner and Baccus, 2011). However, this constraint 35 could potentially decrease the amount of information that can be encoded in an neu-36 ral ensemble and increase the metabolic demands on a sensory tissue (Laughlin, 1981; 37 Balasubramanian et al., 2001). Alternatively, adaptation and sensitization could be sig-38 natures of fundamentally distinct neural coding strategies (Młynarski and Hermund-39 stad, 2018). Further, these alternative hypotheses are not mutually exclusive—adapting 40 and sensitizing cells could mirror each other in some species and neural pathways and 41 not in others, depending on the particular coding and metabolic constraints in those 42 systems (Laughlin, 1981; Barlow, 1961; Levy and Baxter, 1996; Balasubramanian et al., 43 2001). However, given that neural sensitization was only recently discovered, relatively 44 little is known about its roles in neural information processing. 45

To address this issue, we recorded from five types of output neurons in the macaque monkey retina—broad thorny, On and Off parasol (magnocellular-projecting), and On and Off midget (parvocellular-projecting) ganglion cells. These cells have well described roles in visual processing and no known functional counterparts. We studied how these cells responded to global fluctuations in contrast and other stimulus statistics. We report that whereas broad thorny and parasol cells strongly adapted, midget cells sensitized—increasing their responsiveness to certain types of visual stimulation,

including high contrast and simulated eye movements. Synaptic current recordings re-53 vealed that this increased sensitivity was present in the excitatory input from midget 54 bipolar cells and was mediated by presynaptic disinhibition. A computational model 55 based on synaptic input recordings further indicated that this increase in sensitivity 56 greatly enhanced the fidelity of encoding natural scenes. Moreover, the lack of an 57 adapting counterpart to midget cells indicates that sensitizing circuits perform a dis-58 tinct role in primate retina relative to that observed in other vertebrate neural sys-59 tems (Kastner and Baccus, 2011; Kastner and Baccus, 2013; Nikolaev et al., 2013; Cohen-60 Kashi Malina et al., 2013). 61

## RESULTS

The midget pathway of the primate retina is commonly believed to lack short-term 62 plasticity mechanisms such as contrast gain control. This belief is based on reports that 63 midget cells did not exhibit noticeable changes in responsiveness following transitions 64 from high to low contrast regimes (Solomon et al., 2004; Benardete et al., 1992). The 65 assay used to measure adaptation was a sinusoidally modulated drifting grating with 66 bar widths tuned to the size of the midget cell receptive field center, which is narrower 67 than many other retinal cell types. Thus, if plasticity in the midget pathway depended 68 on mechanisms with broader spatial tuning, this assay would not engage such mecha-69 nisms. 70

To determine whether short-term plasticity in the midget pathway depended on 71 the spatial properties of the stimulus, we repeated this assay while varying the spa-72 tial tuning of the gratings. At the offset of high contrast, midget cells did not exhibit 73 a notable change in firing relative to the period that preceded high-contrast stimula-74 tion (Solomon et al., 2004; Benardete et al., 1992) (Figure 1C; spatial frequency, 3.5 75 cycles degree<sup>-1</sup>). To determine whether this lack of either adaptation or sensitiza-76 tion persisted across a range of stimulus conditions, we varied the spatial frequency 77 content of the drifting gratings. Following the offset of low spatial frequency grat-78 ings, most midget cells showed an increase in spiking relative to the period preced-79 ing grating onset (Figure 1D; spatial frequency, 0.35 cycles degree<sup>-1</sup>). This increase 80 in spiking following high contrast is characteristic of the contrast sensitization ob-81 served in other vertebrate retinas (Kastner and Baccus, 2011; Kastner and Baccus, 2013; 82 Nikolaev et al., 2013). The presence of sensitization at low spatial frequencies suggested 83 that sensitization depended on the ability to engage elements in the midget cell recep-84

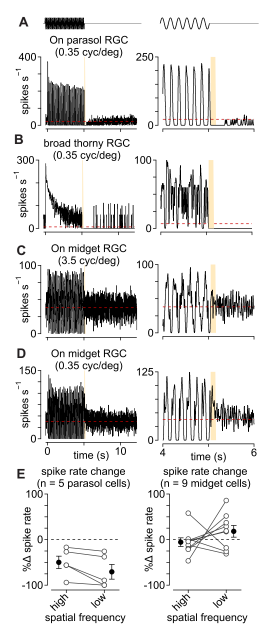


Figure 1. Parasol and midget cells exhibit opposing forms of plasticity. (A) Spike rate in an On parasol ganglion cell to a low spatial frequency drifting grating presented for five seconds (temporal frequency, 6 Hz; spatial frequency, 0.35 cycles degree<sup>-1</sup>). After the offset of high contrast, the spike rate declined below the level prior to grating onset (red dashed line). Right, zoom of transition period. (B) Same as (A) in a broad thorny (On-Off type) ganglion cell. (C) Same as (A) in an On midget ganglion cell to a high spatial frequency grating  $(3.5 \text{ cycles degree}^{-1})$ . (D) Spike responses from the same cell as in (C) to a low spatial frequency grating (0.35 cycles degree<sup>-1</sup>). (E) Change in spike rate for the period directly after grating offset relative to period prior to grating onset in parasol (left) and midget ganglion cells (right).

tive field with broad spatial tuning relative to the midget bipolar cell.

Parasol and broad thorny cells responded very differently than midgets. At the tran-86 sition from high to low contrast, these cells showed a pronounced decrease in spiking 87 relative to the period before the grating turned on and several seconds were required 88 for the spike rate to recover (Figure 1A, B; high contrast, 1.0; low contrast, 0.0; spatial 89 frequency, 0.18-3.5 cycles degree<sup>-1</sup>; grating size, 730  $\mu$ m  $\times$  730  $\mu$ m). This behavior is 90 characteristic of contrast adaptation-during periods of high contrast, circuit mecha-91 nisms reduce the gain to avoid saturation and the gain remains low for several seconds 92 following the transition to a low-contrast regime (Chander and Chichilnisky, 2001; 93

<sup>94</sup> Benardete and Kaplan, 1999; Solomon et al., 2004).

#### 95 Wide-field stimulation evokes contrast sensitization in midget ganglion cells

Our next goal was to determine how this putative wide-field component of the midget 96 cell receptive field contributed to contrast coding. To accomplish this goal, we sought a 97 more spatially and temporally precise assay of sensitivity following wide-field adapta-98 tion. Contrast tuning of parasol and midget cells was determined with spots centered 99 on the receptive field (duration, 0.1 s; parasol diameter, 80-200 µm; midget diameter, 100 40-80 µm). Contrast responses were measured in isolation (unadapted condition) or 101 50-100 ms following the offset of an adapting stimulus (adapted condition). The adapt-102 ing stimulus was a large, high-contrast spot modulated at 20-30 Hz (diameter, 730 µm; 103 contrast, 0.5-1.0, duration, 1.25 s). Presentations of the adapted and unadapted stimuli 104 were interleaved to account for any potential variability in cellular responses over time. 105

Example spike responses to this stimulus paradigm are shown in Figure 2. Parasol 106 cells increased their spike rate at the onset of the adapting stimulus and the spike rate 107 quickly decreased to a steady-state rate by ~0.25 s. Test flashes presented after the offset 108 of the adapting stimulus evoked fewer spikes relative to the unadapted control (Figure 109 2A). Both of these patterns—a transient increase in spike rate following the transition to 110 high contrast and a decrease in spiking after the transition to low contrast-are char-11 acteristic of cells undergoing contrast adaptation (Kim and Rieke, 2001; Baccus and 112 Meister, 2002; Brown and Masland, 2001). 113

We modeled the variation in the contrast-response function following the adapting 114 stimulus as a change in the slope (gain) and a horizontal shift relative to the control 115 condition (see Methods). Following the adapting stimulus, parasol cells showed a large 116 decrease in gain (-30.2  $\pm$  4.5%; n = 5 cells; p = 1.3  $\times$  10<sup>-3</sup>; Wilcoxon signed rank test, 117 here and below) and a small rightward horizontal shift (+3.6  $\pm$  1.5% contrast; p = 3.0  $\times$ 118  $10^{-2}$ ) relative to the unadapted control (Figure 2C). This result confirms previous re-119 ports that parasol cells readily adapt to contrast by continuously adjusting their sensi-120 tivity to match the statistics of incoming visual inputs (Chander and Chichilnisky, 2001; 121 Solomon et al., 2004; Benardete et al., 1992). 122

<sup>123</sup> Midget cells showed several striking differences relative to the pattern observed in <sup>124</sup> parasol cells. First, the decrease in gain was much smaller in midget cells ( $-5.4 \pm 4.3\%$ ; <sup>125</sup> n = 14 cells; p = 0.12). Second, an increase in spike rate was observed at the offset <sup>126</sup> of the adapting stimulus relative to the unadapted control (Figure 2B). This increase

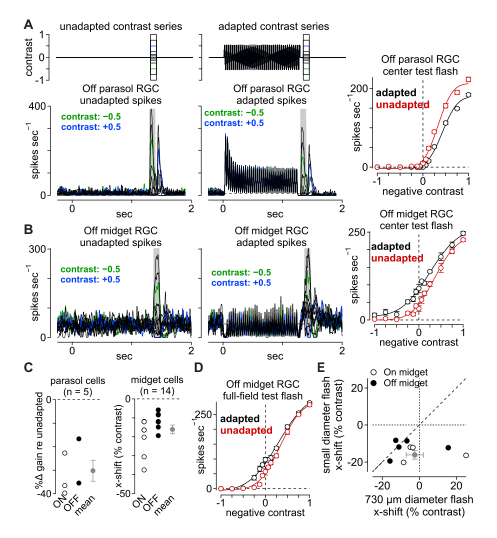


Figure 2. Midget ganglion cells display contrast sensitization. (A) Spike responses from an Off parasol ganglion cell to a series of spots centered over the receptive-field. Spots were either presented alone (left) or 50 ms following the offset of an adapting stimulus (right). Shaded regions indicate sampling windows. *Right*, Average spike rate across the shaded regions. The wide-field adaptation evoked a decrease in the slope (gain) of the contrast-response curve (black) relative to the unadapted control condition (red). (B) Same as (A) for an Off midget ganglion cell. *Right*, Average spike rate across the shaded regions. The wide-field adaptation evoked a leftward shift in the contrast-response curve (black) relative to the unadapted control condition (red). (C) Left, Population data showing the change in slope (gain) for the adapted condition relative to the unadapted condition in On (open circles) and Off (closed circles) parasol ganglion cells (n = 5). *Right*, Population data showing the x-axis shift for adapted relative to unadapted conditions for small-diameter test flashes in On (open circles) and Off (closed circles) midget cells (n = 14). Gray circle and bars indicate mean  $\pm$  SEM. (D) Average spike rate evoked by wide-field test flashes for the Off midget cell in (B). (E) Population data showing the x-axis shift for adapted relative to unadapted conditions for wide-field test flashes versus small-diameter test flashes in On (open circles) and Off (closed circles) midget cells. Gray circle and bars indicate mean  $\pm$  SEM.

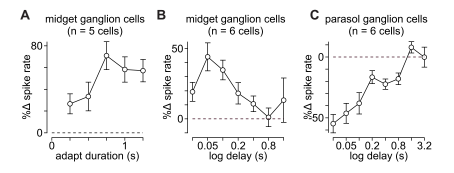
in spiking was evident at all contrasts tested including the zero-contrast condition in 127 which the spot intensity was equal to the average background intensity. The elevation 128 in spiking following the adapting stimulus produced a leftward shift in the contrast-129 response curve relative to control ( $-16.2 \pm 2.3\%$  contrast; p =  $5.2 \times 10^{-6}$ ). A negative 130 horizontal shift value occurred when the adapted curve was shifted to the left of the 131 control curve and this indicated that a weaker stimulus was required to elicit the same 132 spike response from a midget cell following the adapting stimulus. This observation 133 was consistent with previous reports demonstrating that a decreased spike threshold, 134 increased baseline response, and slight decrease in gain are characteristic of contrast 135 sensitization (Kastner and Baccus, 2011; Kastner and Baccus, 2013). 136

#### <sup>137</sup> Contrast sensitization is reduced for wide-field stimulation

Midget cells show narrow receptive-field centers with strong input from the receptive-138 field surround (Crook et al., 2011; De Monasterio and Gouras, 1975; Derrington et al., 139 1984). Thus, the effect of sensitization may be diminished following the adapting stim-140 ulus depending on the relative influences of the direct midget bipolar cell input and 141 wide-field mechanisms in contrast sensitization. To determine whether contrast sen-142 sitization varied with the size of the test flash, we repeated the adaptation experiment 143 but used wide-field test flashes to measure the contrast tuning of midget cells (diam-144 eter, 730 µm). The wide-field test flash evoked a slight leftward shift for the adapted 145 condition relative to control, but this shift was much smaller than was observed for the 146 small-diameter test flash in the same cell (compare Figure 2B, D). This trend held true 147 across midget cells-horizontal shifts were more negative for the small-diameter test 148 flash than for the wide-field test flash in the same cell and these shifts were not statis-149 tically significant for the wide-field test flashes (x-shift,  $-2.4 \pm 4.6\%$  contrast; p = 0.30; 150 gain change,  $-5.8 \pm 5.8\%$ ; n = 9 cells; p = 0.17; Figure 2E). These data indicated that the 15 relative activations of narrow-field and wide-field mechanisms during and following 152 the adapting stimulus were critical to contrast sensitization in midget ganglion cells. 153 Moreover, this result agrees well with previous findings that did not report contrast 154 sensitization to wide-field noise (Chander and Chichilnisky, 2001). 155

# 156 Time course for the onset and persistence of sensitization

<sup>157</sup> We next sought to determine the amount of stimulation needed to evoke sensitiza-<sup>158</sup> tion and also how long sensitization persisted after its onset. To determine the stim-



**Figure 3.** Time course of contrast sensitization and adaptation. (A) Change in spike rate for the adapted condition relative to unadapted control for adaptation periods (contrast,  $\pm 0.25$ -0.5; delay 0.05 s). Adaptation period was varied between 0.25-1.25 s (*x*-axis). (B) Duration of contrast sensitization in midget ganglion cells. Test flashes (contrast,  $\pm 0.25$ -0.5) were presented at different delays (*x*-axis) following the offset of an adapting stimulus. Percent change in spike rate for the adapted condition relative to the unadapted condition is shown on the *y*-axis. (C) Same as (B) for parasol ganglion cells. Error bars indicate mean  $\pm$  SEM.

<sup>159</sup> ulation period needed to initiate sensitization, we varied the presentation time of the <sup>160</sup> adapting stimulus and measured the change in spike rate relative to the unadapted con-<sup>161</sup> trol (adaptation duration, 0.25-1.25 s; contrast,  $\pm 0.5$ ; duration, 0.1 s). For each period <sup>162</sup> of adaptation, midget cells showed an elevation in spiking relative to unadapted con-<sup>163</sup> trols (Figure 3A). Thus, sensitization could be elicited even with fairly brief stimulus <sup>164</sup> presentations.

To determine the time course of sensitization in midget cells, we measured spot re-165 sponses at different times following the offset of the adapting stimulus (delay, 0.025-1.6 166 sec; contrast,  $\pm 0.5$ ; duration, 0.1 sec). Relative to the unadapted control, the adapting 167 stimulus elicited higher spike rates to the test flash in midget cells at delays of 0.025-168 0.4 seconds (Figure 3B). This elevation in spiking, characteristic of sensitization, was 169 greatest 0.05-0.1 seconds after the offset of the adapting stimulus. Parasol cells, on the 170 other hand, showed a reduction in spiking to the same stimulus that persisted for ap-171 proximately one second (Figure 3C). Together, these data indicated that sensitization 172 in midget cells could be elicited even with fairly brief stimulus presentations and that 173 it persisted for several hundred milliseconds. 174

# 175 Sensitization enhances chromatic processing in midget cells

Midget ganglion cells in the central retina exhibit strong chromatic opponency which
 is formed from differential input from long-wavelength cones (L cones) and middle-

wavelength cones (M cones) to the receptive-field center and surround (Crook et al.,
2011; De Monasterio and Gouras, 1975; Derrington et al., 1984). To determine whether
sensitization affected chromatic processing, we measured contrast responses in midget
cells with purely chromatic (isoluminant) test flashes following the adapting stimulus.

Isoluminant (equiluminant) stimuli are commonly employed to study color mecha-182 nisms in isolation. These stimuli are created by modulating L and M cones in oppos-183 ing phases to silence achromatic mechanisms that sum inputs from these cone types 184 (i.e., L+M). We measured contrast-responses to purely chromatic (isoluminant) flashes 185 (duration, 0.1 sec) in the presence or absence of an achromatic adapting stimulus, as 186 above. As with the achromatic stimuli, the adapting stimulus elicited a leftward shift 187 to chromatic test contrasts (Figure 4). This shift was reminiscent of that observed for 188 achromatic stimulation (-11.3  $\pm$  4.1% contrast; n = 8 cells; p = 1.5  $\times$  10<sup>-2</sup>). These data indi-189 cated that contrast sensitization enhanced both achromatic and chromatic processing 190 in midget cells. 191

While chromatic processing was affected by sensitization, the observation that an achromatic adapting stimulus was sufficient to evoke sensitization indicated that chromatic circuits were not necessary to elicit the phenomenon. These data did not, however, rule out contributions from purely chromatic mechanisms to contrast sensitization.

## <sup>197</sup> Sensitization does not arise from a chromatic mechanism

To determine whether such a chromatic mechanism contributed to the observed 198 contrast sensitization, we presented a chromatic adapting stimulus. This stimulus was 199 specifically designed to modulate chromatic mechanisms that differentiate L- and M-200 cone inputs (L-M; isoluminant) while silencing achromatic mechanisms that sum in-201 puts from the L- and M-cone pathways (L+M; isochromatic). Following the adapting 202 stimulus, an isoluminant contrast series was used to measure the input-output rela-203 tionship. In the same cell, we compared chromatic contrast-responses following a 204 chromatic (L-M) or achromatic (L+M) adapting stimulus. 205

The achromatic adapting stimulus produced a leftward shift in the chromatic contrastresponse relation. The chromatic adapting stimulus, however, produced no such shift (x-shift,  $-1.1 \pm 4.7\%$  contrast; n = 8 cells; p = 0.41). We interpreted this result as evidence that contrast sensitization arose from an achromatic mechanism in the midget cell receptive-field. Moreover, given the role of horizontal cells in forming the L-versus-M

<sup>211</sup> opponent receptive-field surround, these data excluded horizontal cells as the source

<sup>212</sup> of sensitization in the midget pathway (Crook et al., 2011).

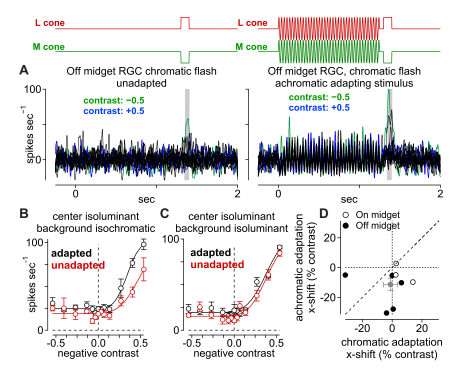


Figure 4. Sensitization arises from an achromatic mechanism. (A) Spike responses from an Off midget ganglion cell to a chromatic (isoluminant) contrast series. Spots were either presented alone (left) or 50 ms following the offset of an achromatic adapting stimulus (right). Shaded regions indicate sampling windows. (B) Average spike rate across the shaded regions indicated in (A). Achromatic adaptation evoked a leftward shift in the contrast-response curve (black) relative to the unadapted control condition (red) for the chromatic test flash. (C) Same as (B) for a chromatic adapting stimulus. The chromatic adapting stimulus did not evoke change in the contrast-response curve relative to control. (D) Population data showing the *x*-axis shift for adapted relative to unadapted conditions for a chromatic adapting stimulus (*x*-axis) relative to an achromatic adapting stimulus (*y*-axis) in On (open circles) and Off (closed circles) midget cells. Gray circle and bars indicate mean  $\pm$  SEM.

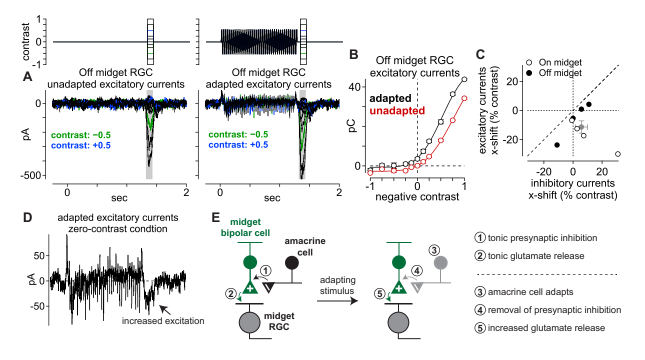
#### <sup>213</sup> Sensitization is present in excitatory synaptic input from midget bipolar cells

The experiments above found contrast sensitization in the spike output of midget ganglion cells. Our next goal was to understand the circuit mechanisms mediating sensitization. To accomplish this goal, we measured the direct excitatory and inhibitory synaptic inputs to midget ganglion cells with whole-cell, voltage-clamp recordings (see Methods). Excitatory currents were isolated by holding a cell's membrane voltage at

the reversal potential for inhibition (-70 mV), and likewise, inhibitory currents were 219 recorded at the excitatory reversal potential (0 mV). An increase in excitatory input to 220 a cell was indicated by a more negative (inward) current relative to the leak current. In-221 deed, the adapting stimulus evoked larger inward excitatory currents relative to the un-222 adapted control at all contrasts tested (Figure 5A). Plotting excitatory charge as a func-223 tion of contrast revealed a similar pattern to that observed in the spike recordings-the 224 adapting stimulus evoked a leftward shift in the contrast-response curve relative to the 225 unadapted control (Figure 5B). On average, the adapting stimulus elicited a horizontal 226 shift of -11% contrast ( $-11.3 \pm 4.2\%$  contrast; n = 8 cells; p =  $1.95 \times 10^{-2}$ ). These results 227 indicated that contrast sensitization was present in the excitatory synaptic input from 228 midget bipolar cells to midget ganglion cells. 229

We also tested for the presence of sensitization in the inhibitory synaptic inputs 230 to midget cells. Unlike the pattern observed in spiking and excitatory currents, the 231 adapting stimulus did not consistently elicit leftward shifts in the inhibitory contrast-232 response functions relative to control ( $+5.8 \pm 4.3\%$  contrast; n = 8 cells; p = 0.25; Figure 233 5C). These data indicated that contrast sensitization arose at or prior to the level of 234 glutamate release from midget bipolar cells. This finding was consistent with the cir-235 cuit model for contrast sensitization in bipolar cells in the retinas of fish, salamander, 236 mice, and rabbits (Kastner and Baccus, 2013; Kastner and Baccus, 2011; Nikolaev et 237 al., 2013). This model posited a mechanism in which a strongly adapting amacrine 238 cell drove sensitization by a mechanism of presynaptic inhibition at the bipolar cell 239 terminal (Kastner and Baccus, 2013). During the adapting stimulus, the amacrine cell 240 adapted such that it decreased release of inhibitory neurotransmitter to the bipolar cell 241 synaptic terminal relative to the tonic level following stimulus offset. This presynap-242 tic disinhibition, in turn, depolarized the bipolar cell synaptic terminal, allowing the 243 cell to utilize its full dynamic range in signaling via glutamate release to postsynaptic 244 ganglion cells. 245

Cleanly measuring the effects of presynaptic inhibition on circuit function has proven exceedingly difficult as use of inhibitory receptor antagonists typically cause many offtarget effects that make data interpretation highly tenuous (Cook et al., 1998). Indeed, adding inhibitory antagonists in primate retina evoked significant increases in tonic glutamate release from bipolar cells and changed the contrast polarity of On parasol cells (Manookin et al., 2018). Nonetheless, our spike and whole-cell recordings strongly supported the proposed model in which contrast sensitization arose from disinhibi-



**Figure 5.** Sensitization present in excitatory synaptic input from midget bipolar cells. (A) Excitatory currents from an Off midget ganglion cell to a series of spots (diameter, 40-80 µm) centered over the receptive field. Spots were either presented alone (*left*) or 50 ms following the offset of an adapting stimulus (*right*; diameter, 730 µm). Shaded regions indicate sampling windows. (B) Average spike rate across the shaded regions indicated in (A). The wide-field adaptation evoked a leftward shift in the contrast-response curve (black) relative to the unadapted control condition (red). (C) Population data showing the x-axis shift for adapted relative to unadapted conditions for excitatory versus inhibitory synaptic currents in On (open circles) and Off (closed circles) midget cells. Mean values are shown in gray. Error bars indicate mean  $\pm$  SEM. (D) Excitatory current recordings from the Off midget cell in (A) under the condition in which the stimulus intensity returned to the mean luminance after the offset of the adapting stimulus and an additional test flash was not presented (zero-contrast condition). A sustained increase in excitatory current was observed at the offset of that stimulus. (E) Proposed model for contrast sensitization in midget bipolar cells.

tion at the presynaptic bipolar cell terminal (Kastner and Baccus, 2013). First, the lack of sensitization to a purely chromatic (isoluminant) adapting stimulus indicated that sensitization did not arise in the outer retina at the level of horizontal cell feedback (Figure 4). Second, the effect of presynaptic disinhibition was seen in our excitatory current recordings (Figure 5D). In one of our stimulus conditions the test flash contrast was zero such that the stimulus intensity returned to the average background intensity at the offset of the adapting stimulus. Although this stimulus lacked a change in con-

trast following the adapting stimulus, we observed an increase in excitatory synaptic input (Figure 5D). This response pattern was consistent with a decrease in presynaptic inhibition following the offset of the adapting stimulus, resulting in an increase in glutamate release from midget bipolar cells. Thus, our recordings in midget pathway of primate retina were consistent with the circuit motif proposed in other vertebrate species (Figure 5E; (Kastner and Baccus, 2013)).

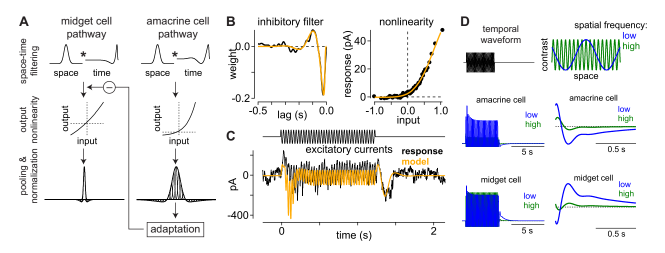
## <sup>266</sup> A contrast sensitization model reproduces midget cell responses

Having established the presence of contrast sensitization in midget bipolar cells, we 267 next sought to understand the relevance of this neural computation to visual process-268 ing in primates. To accomplish this goal, we developed a computational model of the 269 proposed circuit in which bipolar cell glutamate release was modulated through presy-270 naptic amacrine cell inhibition (Kastner and Baccus, 2013; Kastner and Baccus, 2011; 271 Nikolaev et al., 2013). Model parameters were determined by recording excitatory and 272 inhibitory synaptic current responses from midget ganglion cells to a Gaussian white 273 noise stimulus (see Methods). 274

We modeled the midget bipolar and presynaptic amacrine cell pathways using the 275 classical linear-nonlinear model with two modifications: 1) adaptation occurred at the 276 amacrine cell output and 2) the amacrine cell output was applied to the bipolar cell 277 model prior to the bipolar cell output nonlinearity (Figure 6A). The model parameters 278 controlling presynaptic sensitization were fit from direct excitatory current recordings. 279 In the same cell from which these parameters were determined, we measured excita-280 tory current responses to the wide-field adapting stimulus (see Figure 5), and the model 28 qualitatively reproduced the increase in excitatory currents following the offset of this 282 adapting stimulus (Figure 6C). 283

We further tested the model using the drifting grating stimuli presented in Figure 284 1. The model produced distinct outputs for the high and low spatial frequency grat-285 ings. The high frequency grating produced a relatively small response and, as a result, 286 little adaptation in the presynaptic amacrine cell (Figure 6D, middle row). This was 287 due to the broad receptive field center size of the amacrine cell relative to the bars of 288 the grating. Low frequency gratings, however, strongly modulated the amacrine cell 289 and produced significant adaptation; this adaptation, in turn, caused a removal of in-290 hibition at the level of the bipolar cell following the offset of the grating, resulting in 29 sensitization (Figure 6D, bottom row). The model predictions were qualitatively simi-292

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**Figure 6.** Sensitization model reproduces experimental results. (A) Sensitization model structure. Visual inputs were convolved with a spatiotemporal linear filter comprised of a Gaussian in space and a biphasic filter in time. Signals in the amacrine cell pathway were then passed through an output nonlinearity before passing to the adaptation stage of the model. The output of the amacrine cell model provided inhibitory input to the midget bipolar cell pathway upstream of the bipolar cell output nonlinearity. (B) Inhibitory temporal filter (*left*) and input-output nonlinearity (*right*) determined from noise recordings. These filters were then used as components of the computational model (A). (C) Excitatory current recording from an Off midget ganglion cell to the wide-field adapting stimulus (see Figure 5). Model prediction (orange) was generated from excitatory synaptic current recordings to the noise stimulus in the same cell. (D) Model output for drifting grating stimuli at high and low spatial frequencies.

lar to our direct recordings from midget cells, indicating that contrast sensitization in
 primate retina can be well explained via presynaptic disinhibition as in other species
 (Kastner and Baccus, 2013; Kastner and Baccus, 2011; Nikolaev et al., 2013).

# Sensitizing circuits more accurately reconstruct natural stimuli than adapting cir cuits

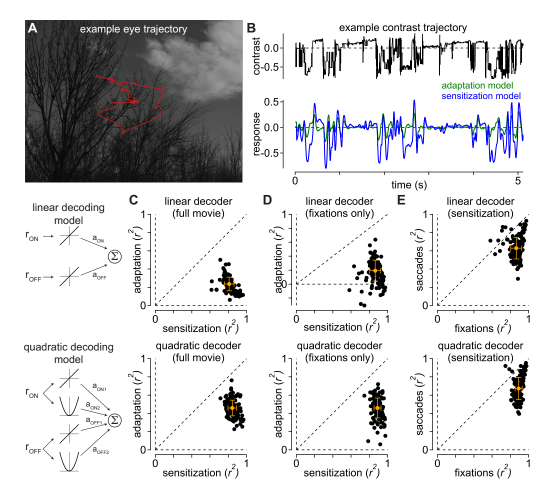
We next sought to understand how these differing strategies of adaptation and sensitization impacted encoding during naturalistic vision. This was done by testing the ability of adapting and sensitizing models to accurately encode natural scenes. We specifically wanted to determine how accurately downstream visual circuits could reconstruct naturalistic input stimuli based on the outputs of populations of model On and Off midget ganglion cells. The naturalistic stimuli used in the model were taken from the DOVES database—a dataset of eye movements in humans recorded while observing natural images (Van Der Linde et al., 2009). Reconstruction accuracy was determined by calculating the correlation between the stimulus and response of each model (see Methods). Periods of fixation between ballistic eye movements are critically important to visual coding in primates; thus, model performance was separately calculated for the complete movie or for periods of fixation only.

We considered two different decoding models for estimating the stimulus contrast 310 based on the outputs of On and Off midget ganglion cells. The first model utilized a 311 linear decoding scheme in which stimulus contrast was estimated by taking the scaled 312 difference between the On and Off cell outputs. We also tested a quadratic decoding 313 model that squared the On and Off outputs prior to differencing (see Methods). Us-314 ing these decoders, we compared the performance of the sensitization model with a 315 model in which the midget bipolar underwent contrast adaptation. Regardless of the 316 decoding scheme used, the sensitizing model showed higher accuracy for reconstruct-317 ing the entire stimulus trajectory than the adapting model (linear  $r^2$ : sensitization, 0.81 318  $\pm$  0.05; adaptation, 0.23  $\pm$  0.07; p = 2.7  $\times$  10<sup>-54</sup>; guadratic  $r^2$ : sensitization, 0.84  $\pm$  0.05; 319 adaptation,  $0.45 \pm 0.09$ ; p =  $2.9 \times 10^{-54}$ ; n = 161 movies; mean  $\pm$  SD; Figure 7C). The sen-320 sitizing model also outperformed the adapting model when the analysis was restricted 321 to periods of fixation (Figure 7D). 322

The sensitizing model showed increased encoding accuracy for periods of fixation relative to periods of ballistic eye movements (movement  $r^2$ ,  $0.63 \pm 0.12$ ;  $p = 2.9 \times 10^{-35}$ ; Figure 7E). This finding suggested that sensitization could play a particularly important role in vision during periods of fixation following the offset of global motion. We, thus, sought to determine whether background motion could evoke contrast sensitization with direct recordings from midget ganglion cells.

## 329 Background motion evokes contrast sensitization in midget cells

To determine whether background motion elicited sensitization, we measured contrast responses in midget cells following the offset of a full-field moving texture (speed, 5-11 degrees s<sup>-1</sup>; duration, 1 s). The goal was to simulate, as closely as possible, the brief periods of fixation following eye movements and to test sensitivity during these fixation periods. We interleaved these recordings with measurements when the texture was stationary throughout the trial. The moving textures elicited an increase in spiking and a leftward shift in the contrast-response functions relative to the control condition



**Figure 7.** Sensitization increases the fidelity of encoding natural movies. (A) Example image from the DOVES database. The observer's eye trajectory is shown in red. (B) *Top*, temporal contrast sequence from the eye movement data in (A). *Bottom*, responses of the adaptation and sensitization models to the example contrast sequence. (C) Performance of the sensitization (*x*-axis) and adaptation (*y*-axis) models at reconstructing 161 natural movies in the database. Performance was measured as the Pearson correlation between the stimulus and model predictions after adjusting for temporal lag. Performance for each movie is indicated by a black dot. Gray dot and bars indicate mean  $\pm$  SD. The sensitization model outperformed the adaptation model for each of the movies. (D) Model performances as in (C), but restricted to periods of fixation. The sensitization model outperformed the adaptation model in each case. (E) Sensitization model performance of the model was typically higher during periods of fixation.

- in which the texture was stationary (Figure 8). On average, the shift was -25% contrast
- for spike recordings ( $-25.4 \pm 4.4\%$  contrast; n = 10 cells; p =  $2.4 \times 10^{-2}$ ) and -12% contrast
- for excitatory current recordings ( $-12.5 \pm 5.1\%$  contrast; n = 4 cells; p =  $2.4 \times 10^{-2}$ ).

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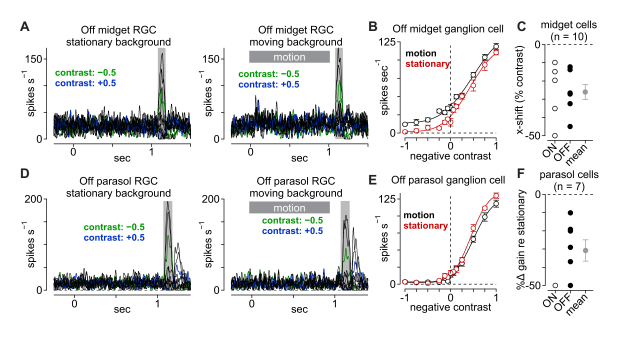


Figure 8. Background motion evokes contrast sensitization in midget cells. (A) Spike responses from an Off midget ganglion cell to a series of spots centered over the receptive field. Spots were either presented alone (left) or 50 ms following the offset of background motion (speed, 11 degrees  $s^{-1}$ ). Shaded regions indicate sampling windows. (B) Average spike rate across the shaded regions indicated in (A). The wide-field adaptation evoked a leftward shift in the contrast-response curve (black) relative to the unadapted control condition (red). (C) Horizontal shift (x-shift) in contrast-response function following background motion relative to control condition in which the background was stationary. Data are shown for On and Off midget ganglion cells (n = 10). Gray circle and bars indicate mean  $\pm$  SEM. (D) Same as (A) for an Off parasol ganglion cell. (E) Same as (B) for the Off parasol cell in (D). The cell showed a decrease in spike output following the offset of background motion-the opposite pattern to that observed in the Off midget cell. (F) Change in gain in the contrast-response function following background motion relative to the control condition. On average, background motion elicited a decrease in gain of ~30% relative to the control condition in which the background was stationary (n = 7 cells). Gray circle and bars indicate mean  $\pm$  SEM.

These data were consistent with our circuit model of contrast sensitization. The amacrine cell providing presynaptic inhibition to the midget bipolar cell adapted during background motion; at the offset of motion, the cell hyperpolarized and reduced presynaptic inhibition to the bipolar terminal. Thus, similar to circuits described in other vertebrates, the midget pathway could utilize presynaptic inhibition to account for self-motion (Olveczky et al., 2003; Baccus et al., 2008; Kastner and Baccus, 2013).

## DISCUSSION

Our results support a novel role for neural sensitization in primates relative to the 346 function proposed in other species. Sensitizing cells are commonly thought to coun-347 teract the loss of responsiveness experienced by adapting cells during transitions from 348 high to low variance environments (Kastner and Baccus, 2011). This hypothesis re-349 quires that sensitizing cells have an adapting counterpart that encodes similar infor-350 mation about the environment. Midget (parvocellular-projecting) ganglion cells are 35 well known for their roles in both chromatic and achromatic vision (Crook et al., 2011; 352 De Monasterio and Gouras, 1975; Derrington et al., 1984). Functional parallelism in 353 the midget pathway is achieved by splitting signals between different classes of cone 354 photoreceptor (L versus M) or bipolar cell (On versus Off) inputs to the midget cell 355 receptive-field. Further, we found that both On- and Off-type midget cells exhibited 356 sensitization (Figure 1-4, 8), and the primate retina lacks an adapting functional coun-357 terpart to midget cells with similar chromatic opponency or spatial acuity (Wässle, 358 2004); thus, sensitization does not counterbalance adaptation in another functionally 359 parallel pathway. 360

Instead, our findings indicate that sensitization maintains the responsiveness of the 361 midget pathway during dynamic visual processes, such as head or eye movements, 362 that cause rapid fluctuations in light intensity on the retina. We base this conclusion 363 on several key observations. First, sensitization was strongest following wide-field stim-364 ulation (Figure 1-4) or background motion (Figure 8). Second, sensitization persisted 365 for >0.2 s (Figure 3), a period that roughly corresponds to the durations of fixations fol-366 lowing eye movements in primates (reviewed in (Rayner, 1998)). Finally, sensitization 367 greatly improved the fidelity of encoding natural movies, particularly during periods 368 of fixation following ballistic eye motion (Figure 7). Thus, sensitization appears to play 369 a unique and crucial role in neural coding in primates. 370

A parallel study also found evidence supporting the link between the sensitization 371 mechanisms that we observed in midget ganglion cells and visual perception in hu-372 mans (Naecker and Baccus, 2018). Subjects showed a significant enhancement in con-373 trast sensitivity following the offset of wide-field motion; and this increase in sensitivity 374 was manifest as a leftward horizontal shift in the perceptual input-output relationship, 375 just as we observed in midget cells (compare Figure 2 in our study with Figure 5 of 376 (Naecker and Baccus, 2018)). Together, these findings provide a rare example of a be-377 havior that can be directly tied to a specific neural circuit motif. 378

#### <sup>379</sup> Distinct functions of adaptation and sensitization in primate retina

Our findings also speak to the roles of neural adaptation in the parasol and broad 380 thorny ganglion cell pathways. Previous work proposed that adapting cells could pro-38 duce a nearly optimal faithful encoding of sensory inputs (Fairhall et al., 2001). Our 382 computational model, however, indicates that sensitizing circuits outperform adapt-383 ing circuits in encoding natural movies (Figure 7). The improved reconstruction accu-384 racy of the sensitizing model was consistent with a recent theoretical report indicating 385 that sensitizing cells are better for encoding faithful representations of sensory input 386 than adapting cells (Młynarski and Hermundstad, 2018). According to this paradigm, 387 sensitizing cells such as midget ganglion cells would be useful for directly encoding 388 information about the properties of the input (e.g., contrast, color). Adapting cells, 389 on the other hand, are optimized for performing inference tasks (Wark et al., 2009; 390 Młynarski and Hermundstad, 2018). 391

Adapting cells dynamically adjust their input-output properties to align with the re-392 cent stimulus distribution (Baccus and Meister, 2002; Smirnakis et al., 1997). These 393 adjustments make the cells exquisitely sensitive to changes in stimulus statistics, allow-394 ing them to infer when salient properties of the environment change. For example, 395 quickly detecting object motion is an ethologically relevant and phylogenetically an-396 cient neural computation (Frost et al., 1990; Lettvin et al., 1959); by decreasing their 397 responsiveness during periods in which the background is either stationary or coher-398 ently moving, adapting neural circuits would be poised to report when an object moves 399 relative to the background (Olveczky et al., 2003; Puller et al., 2015). Interestingly, both 400 adapting parasol and broad thorny ganglion cells have been implicated in motion pro-401 cessing (Manookin et al., 2018; Puller et al., 2015) and project to retinorecipient brain re-402 gions in the lateral geniculate body, superior colliculus, and inferior pulvinar that con-403 tribute significantly to motion vision (Rodieck and Watanabe, 1993; Crook et al., 2008; 404 Kwan et al., 2018). 405

#### <sup>406</sup> Relationship to psychophysical measurements in humans

It has long been recognized that eye movements play important computational roles in visual processing (reviewed in (Martinez-Conde et al., 2004; Rucci and Victor, 2015)). Periods in which an image is stabilized on the retina cause that image to fade from perception (Troxler, 1804) and small fixational eye movements appear to counteract this fading (Rucci et al., 2007; Schütz et al., 2008). These eye movements can, how-

ever, produce large temporal fluctuations in contrast, particularly when viewing highcontrast objects. This would, in turn, produce fading phenomena in cells that strongly
adapt, such as parasol ganglion cells—a prediction that was confirmed with our computational model (Figure 7).

Neural mechanisms such as sensitization may serve to counteract adaptation by main-416 taining the sensitivity of certain visual pathways during eye movements. Indeed, our 417 computational model and direct measurements indicated that contrast sensitization in 418 the midget ganglion cell pathway was engaged well by background motion such as that 419 observed during eye movements (Figure 7, 8). Thus, contrast sensitization might act 420 to maintain sensitivity of image-forming visual pathways following eye movements 421 that are commonplace in primate vision. Indeed, psychophysical studies in humans 422 indicated that contrast sensitivity increases following both ballistic (saccade) and fixa-423 tional eye movements (Rucci et al., 2007; Schütz et al., 2008). Moreover, this increase 424 in sensitivity was limited to chromatic stimuli and high-spatial-frequency achromatic 425 stimuli, mirroring our results in midget ganglion cells. 426

## METHODS

Experiments were performed in an *in vitro*, pigment-epithelium attached preparation of the macaque monkey retina (Manookin et al., 2015). Eyes were dissected from terminally anesthetized macaque monkeys of either sex (Macaca *fascicularis, mulatta*, and *nemestrina*) obtained through the Tissue Distribution Program of the National Primate Research Center at the University of Washington. All procedures were approved by the University of Washington Institutional Animal Care and Use Committee.

# 433 Tissue Preparation and Electrophysiology

The retina was continuously superfused with warmed (32-35 °C) Ames' medium 434 (Sigma) at ~6-8 mL min<sup>-1</sup>. Recordings were performed from macular, mid-peripheral, 435 or peripheral retina (2-8 mm, 10-30° foveal eccentricity), but special emphasis was 436 placed on recording from more centrally located cells. Physiological data were ac-437 quired at 10 kHz using a Multiclamp 700B amplifier (Molecular Devices), Bessel filtered 438 at 3 kHz (900 CT, Frequency Devices), digitized using an ITC-18 analog-digital board 439 (HEKA Instruments), and acquired using the Symphony acquisition software package 440 developed in Fred Rieke's laboratory (http://symphony-das.github.io). 441

<sup>442</sup> Recordings were performed using borosilicate glass pipettes containing Ames medium

for extracellular spike recording or, for whole-cell recording, a cesium-based internal 443 solution containing (in mM): 105 CsCH<sub>3</sub>SO<sub>3</sub>, 10 TEA-Cl, 20 HEPES, 10 EGTA, 2 QX-444 314, 5 Mg-ATP, and 0.5 Tris-GTP, pH ~7.3 with CsOH, ~280 mOsm. Series resistance 445 (~3-9 M $\Omega$ ) was compensated online by 50%. The membrane potential was corrected 446 offline for the approximately -11 mV liquid junction potential between the intracel-447 lular solution and the extracellular medium. Excitatory and inhibitory synaptic cur-448 rents were isolated by holding midget ganglion cells at the reversal potentials for in-449 hibitory/chloride (E<sub>Cl</sub>, ~-70 mV) and excitatory currents (E<sub>cation</sub>, 0 mV), respectively. 450

## 451 Visual Stimuli and Data Analysis

Visual stimuli were generated using the Stage software package developed in the 452 Rieke lab (http://stage-vss.github.io) and displayed on a digital light projector (Lightcrafter 453 4500; Texas Instruments) modified with custom LEDs with peak wavelengths of 405, 454 505 (or 475), and 640 nm. Stimuli were focused on the photoreceptor outer segments 455 through a 10X microscope objective. Mean light levels were in the low to medium 456 photopic regimes ( $-3 \times 10^3 - 3.4 \times 10^4$  photoisomerizations [R\*] cone<sup>-1</sup> sec<sup>-1</sup>). Con-457 trast values for contrast-response flashes are given in Weber contrast and for periodic 458 stimuli in Michaelson contrast. All responses were analyzed in MATLAB (R2018a+, 459 Mathworks). 460

For extracellular recordings, currents were wavelet filtered to remove slow drift and amplify spikes relative to the noise (Wiltschko et al., 2008) and spikes were detected using either a custom k-means clustering algorithm or by choosing a manual threshold. Whole-cell recordings were leak subtracted and responses were measured relative to the median membrane currents immediately preceding stimulus onset (0.25-0.5 s window). Summary data are presented in terms of conductance (g), which is the ratio of the current response (I) to the driving force:

$$g = \frac{I}{V_m - E} \tag{1}$$

where  $V_m$  is the holding potential (in mV) and *E* is the reversal potential (in mV). Reversal potentials of 0 mV and -70 mV were used for excitatory and inhibitory inputs, respectively.

# 471 Sensitization and adaptation models

We modeled spatiotemporal integration in bipolar cells and amacrine cells as the 472 product of a Gaussian spatial filter and a biphasic temporal filter which was then passed 473 through an input-output nonlinearity. The output of this nonlinear stage of the amacrine 474 cell model was then passed through an adaptation stage; adaptation in the amacrine 475 cell provided inhibitory input to the bipolar cell model prior to the output nonlinearity 476 (Figure 6A). Following the subunit output, model midget ganglion cells and amacrine 477 cells pooled (summed) inputs from bipolar cell subunits and the weights of these inputs 478 were normalized by the subunit location relative to the receptive field center using a 479 Gaussian weighting. 480

To estimate the excitatory and inhibitory circuit components for the computational 481 model, we recorded excitatory and inhibitory synaptic currents from midget ganglion 482 cells in response to a full-field Gaussian flicker stimulus. The contrast of each frame 483 was drawn randomly from a Gaussian distribution and that value was multiplied by 484 the average contrast. Average contrast was updated every 0.5 s and drawn from a uni-485 form distribution (0.05-0.35 RMS contrast). The linear temporal filters (F) were calcu-486 lated by cross-correlating the stimulus sequence (S) and the leak-subtracted response 487 (R) (Baccus and Meister, 2002). 488

$$F(t) = \int R(\tau)S(t+\tau)d\tau$$
 (2)

where  $\tau$  is the temporal lag. These filters were then modeled as a damped oscillator with an S-shaped onset (Schnapf et al., 1990; Angueyra and Rieke, 2013):

$$F(t) = A \frac{(t/\tau_{rise})^n}{1 + (t/\tau_{rise})^n} e^{-(t/\tau_{decay})} \cos\left(\frac{2\pi t}{\tau_{period}} + \varphi\right)$$
(3)

<sup>491</sup> where *A* is a scaling factor,  $\tau_{rise}$  is the rising-phase time constant,  $\tau_{decay}$  is the damping <sup>492</sup> time constant,  $\tau_{period}$  is the oscillator period, and  $\varphi$  is the phase (in degrees).

The input-output nonlinearity was calculated by convolving the temporal filter (F) and stimulus (S) to generate the linear prediction (P).

$$F(t) = \int R(\tau)S(t-\tau)d\tau$$
(4)

495 The prediction (x-axis) and response (y-axis) were modeled as a cumulative Gaussian

<sup>496</sup> distribution (Chichilnisky, 2001).

$$N(x) = \varepsilon + \frac{\alpha}{\sqrt{2\pi}} \int_{-\infty}^{x} e^{\frac{-(\beta t + \gamma)^2}{2}} dt$$
(5)

<sup>497</sup> where  $\alpha$  indicates the maximal output value,  $\epsilon$  is the vertical offset,  $\beta$  is the sensitivity <sup>498</sup> of the output to the generator signal (input), and  $\gamma$  is the maintained input to the cell.

The spatial component of the bipolar and amacrine cell receptive fields was modeled as a Gaussian function with a 2-SD width of 18 µm and 90 µm, respectively. Each midget ganglion cell was modeled as receiving input from a single bipolar cell, as is typically the case in the central retina. Sensitization parameters were determined by fitting linear-nonlinear model predictions relative to the excitatory currents recorded to the Gaussian flicker stimulus.

The amacrine cell providing direct inhibition to the midget ganglion cells is likely 505 distinct from the cell providing presynaptic inhibition at the level of the midget bipo-506 lar cell (see Figure 5). Thus, our inhibitory synaptic recordings likely did not grant us 507 direct access to the properties of the amacrine cell responsible for contrast sensitiza-508 tion. These recordings do, however, provide an estimate of the time-course of signals 509 passing through the presynaptic amacrine cell to midget bipolar cells. Signals passing 510 through this amacrine cell proceed from cone photoreceptors to bipolar cells and then 511 to the amacrine cell in question before providing input to the midget bipolar cell. In 512 the same way, the amacrine cell providing direct inhibition to midget ganglion cells 513 must pass through an extra synapse. Thus, our recordings of direct synaptic inhibition 514 were useful in approximating the time course of presynaptic inhibition at the midget 515 bipolar terminal. 516

#### 517 Evaluating model performance to naturalistic movies

<sup>518</sup> We evaluated the performance of the adaptation and sensitization models in re-<sup>519</sup> constructing the naturalistic movie sequences using linear and quadratic decoding <sup>520</sup> paradigms. To estimate stimulus contrast, the linear decoder ( $f_{LINEAR}$ ) summed the <sup>521</sup> scaled outputs of the model On and Off midget ganglion cells:

$$f_{\text{linear}}(t) = a_{\text{on}}r_{\text{on}}(t) + a_{\text{off}}r_{\text{off}}(t) + k \tag{6}$$

where  $a_{ON}$  and  $a_{OFF}$  are scaling constants and k is an offset constant. The quadratic

model was similar in structure except that the response from each pathways was squared
prior to summation:

$$f_{\text{quadratic}}(t) = a_{\text{on1}}r_{\text{on}}(t) + a_{\text{on2}}r_{\text{on}}^2(t) + a_{\text{off1}}r_{\text{off}}(t) + a_{\text{off2}}r_{\text{off}}^2(t) + k$$
(7)

For each of the 161 movies in the database, the input stimulus was shifted to the peak of the midget temporal filter (~35 ms) and then scaling and offset coefficients were determined using least-squares curve fitting. The Pearson correlation was then calculated between the temporal trajectories of the model and the movie.

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#### **AUTHOR CONTRIBUTIONS**

<sup>541</sup> Conceptualization, M.B.M.; Methodology, M.B.M.; Software, M.B.M.; Formal Anal<sup>542</sup> ysis, M.B.M.; Investigation, M.B.M., T.R.A; Resources, M.B.M.; Data Curation, M.B.M.,
<sup>543</sup> T.R.A.; Writing – Original Draft, M.B.M.; Writing – Review & Editing, M.B.M., T.R.A.;
<sup>544</sup> Visualization, M.B.M.; Supervision, M.B.M.; Project Administration, M.B.M.; Funding
<sup>545</sup> Acquisition, M.B.M. The ORCID number for M.B.M. is 0000-0001-8116-7619.

#### **COMPETING INTERESTS**

<sup>546</sup> The authors declare no competing interests.

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