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Asymmetric Activation of ON and OFF Pathways in the Degenerated Retina

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Abstract

Retinal prosthetics are one of the leading therapeutic strategies to restore lost vision in patients with retinitis pigmentosa and age-related macular degeneration. Much work has described patterns of spiking in retinal ganglion cells (RGCs) in response to electrical stimulation, but less work has examined the underlying retinal circuitry that is activated by electrical stimulation to drive these responses. Surprisingly, little is known about the role of inhibition in generating electrical responses or how inhibition might be altered during degeneration. Using whole-cell voltage–clamp recordings during subretinal electrical stimulation in the rd10 and wild-type (wt) retina, we found electrically evoked synaptic inputs differed between ON and OFF RGC populations, with ON cells receiving mostly excitation and OFF cells receiving mostly inhibition and very little excitation. We found that the inhibition in the OFF pathway arises from glycinergic amacrine cells, and the stimulation of the ON pathway contributes to inhibitory inputs to the RGC. We also show that this presynaptic inhibition in the rd10 retina, compared with that in the wt retina.

Key words: circuit; electrical stimulation; retina; retinal degeneration; retinal prosthetics; synapse

Significance Statement

Changes in circuit processing may have deleterious effects on vision restoration for patients with retinitis pigmentosa. Prior research has focused on feedforward excitatory drive and not the interactions between excitation and inhibition that comprise normal retinal function. This study demonstrates that retinal ganglion cells respond to electrical stimulation in three broad functional groups that correspond to their anatomical structure. We show that while both degenerated and *wt* retinas display the same three groups, the degenerated retina has an increased amount of OFF pathway presynaptic inhibition limiting their excitatory output to OFF ganglion cells.

Introduction

Optogenetics and optoelectronic retinal prosthetics are both currently being pursued for vision restoration in patients with retinitis pigmentosa. Recently several new optogenetic therapies have entered clinical trials for retinitis pigmentosa (Drag et al., 2023); however, subretinal prosthetics are the only therapy in trials to restore lost vision in age-related macular degeneration (Pixium Vision, NCT03392324; Rubner et al., 2022), which is the most common form of retinal blindness (Bourne et al., 2020). This highlights the importance of continuing to pursue improvements in retinal prosthetics. Retinal prosthetics aim to recreate vision through implanted optoelectronic arrays to replace the function of photoreceptors and reactivate the remaining retinal circuitry with electrical stimulation. While implant technologies have been made available to patients, the quality of restored vision has been modest (Stingl et al., 2015, 2017; Palanker et al., 2020), and there are many questions remaining about the best strategies for vision restoration (Weiland et al., 2016; Damle et al., 2017).

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Continued on next page.

The vertebrate retina is subdivided into two parallel pathways, the ON and OFF pathway (Schiller, 1992; Masland, 2001; Wässle, 2004). These pathways work together to encode increments and decrements of light, respectively. When light activates photoreceptors, ON bipolar cells (BCs) are depolarized, and OFF BCs are hyperpolarized. These parallel processing channels are maintained throughout the retina and lateral geniculate nucleus and into primary visual area V1. Subretinal prosthesis aim to restore purposeful vision, but a major concern surrounding retinal prosthetics is the ability to restore meaningful ON and OFF pathway responses. Past work has shown that ON and OFF retinal ganglion cells (RGCs) maintain their distinctive ON- and OFF-specific morphology following retinal degeneration (Marc et al., 2003; Margolis et al., 2008). Intriguingly, a few studies have shown that ON and OFF responses can be evoked during electrical stimulation in a degenerated retina (Sekhar et al., 2017; Ho et al., 2018, 2020). However, it remains unknown if these electrical ON and OFF responses originate in "true" ON and OFF RGCs or reflect anomalous signaling from artificial stimulation. Past work has shown that healthy ON and OFF pathways have stereotypical patterns of excitation and inhibition (Pang et al., 2003; Murphy and Rieke, 2006), which are maintained following degeneration in the rd1 mouse (Margolis et al., 2008); however, the circuit mechanisms that generate electrical ON and OFF responses and the role of inhibition remain completely unknown.

Inhibition arises from a diversity of amacrine cells to shape the spatial and temporal response properties of both RGCs and BCs. Inhibition can be further characterized as feedback (onto BCs), feedforward (onto ganglion cells), crossover (activity in one pathway suppresses elements of the other), or serial (onto other amacrine cells); for reviews, see Masland (2012) and Diamond (2017). It is plausible that prosthetic vision signals could activate any combination of inhibitory signaling. Ideally prosthetic vision restoration would activate inhibitory circuitry as in normal vision. However, electrical stimulation may directly and indiscriminately activate amacrine cells. Thus, to generate prosthetic visual signals with appropriate spatiotemporal characteristics, we must understand how it interacts with inhibitory signaling. Unfortunately, few studies have examined how inhibition in the degenerated retina is recruited in response to electrical stimulation (Margalit and Thoreson, 2006; Tsai et al., 2011; Walston et al., 2018).

To address these questions, we used whole-cell voltage–clamp techniques to measure electrically evoked excitatory and inhibitory synaptic inputs to RGCs in the *rd10* retina. We found three functional response patterns that corresponded to morphologically identified ON, OFF, and ON/OFF ganglion cells, with the prominent difference being in the excitation to inhibition ratio (E/I), where ON RGCs had an E/I ratio >1 and OFF cells had an E/I ratio <1. Interestingly, this is consistent with previous work for light-evoked inputs for a subset of RGCs (Pang et al., 2003; Murphy and Rieke, 2006) and with spontaneous inputs in rd1 mice (Margolis et al., 2008). Pharmacological dissection revealed that presynaptic feedback inhibition in the OFF pathway prevented OFF excitation due to a glycinergic amacrine cell and that direct inhibition in OFF RGCs originated in the OFF pathway.

Materials and Methods

Retinal explant and voltage-clamp recording. All experimental methods and animal care procedures were conducted in accordance with NIH guidelines and were approved by the University of California Institutional Animal Care and Use Committee. Thirty-four adult [age range, postnatal day (P)62–P296] *rd10* and eight C57BL mice of either sex were anesthetized with isoflurane, killed by cervical dislocation, and enucleated, and their retinas were dissected free and maintained in Ames' medium oxygenated and equilibrated with 95% O₂, 5% CO₂. Retina pieces, ~2 × 2 mm, were transferred to a custom recording chamber and placed over stimulating electrodes on the bottom of the custom recording chamber, ganglion cell side up. The chamber was placed under an upright microscope and perfused with Ames' (Ames and Nesbett, 1981) media (United States Biological) at a rate of 4 ml/min at 35°C. RGCs were visualized and targeted using infrared (IR) differential interference contrast (DIC) video microscopy. Contact between the outer portion of the retina and the electrode array surface was also confirmed with microscopy. Stimulation was delivered on the nearest electrode to the target RGCs.

Sputtered iridium oxide film (SIROF) stimulating electrode arrays were fabricated as described previously on a borosilicate glass disk, which formed the bottom of the recording chamber (Damle et al., 2021). Briefly, the array consisted of a 4×8 grid of

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30-µm-diameter SIROF electrodes spaced 50 µm apart from center to center. SIROF electrodes were formed by reactive DC sputtering to a thickness of 600 nm over indium tin oxide (ITO) traces. ITO traces terminated in gold contact pads at the edge of the disk for connection to a 32-channel RHS2000 stim and recording system (Intan Technologies). Charge-balanced, anodic–first, square, biphasic current pulses were generated on the RHS2000, triggered by our acquisition software, and delivered to an individual stimulating electrode nearest to the cell of interest. Arrays of 30-µm-diameter iridium oxide electrodes were used to stimulate the inner nuclear layer with 1 ms, 20 µA biphasic anodic-first pulses (Fig. 1*A*). Distances were measured using Scientifica LinLab2 and IR-DIC microscopy. The cell was positioned in the center of the field of view and marked as the coordinate zero point. The stage was then moved to the center of the stimulating electrode, and the *x* and *y* coordinates from LinLab2 were recorded.

Recording electrodes were pulled from borosilicate capillary glass to have a final resistance of 4–5 MΩ and filled with internal solution comprised of 90 mM CsCH3SO3, 20 mM TEA-Cl, 10 mM HEPES, 10 mM EGTA, 10 mM phosphocreatine disodium salt, 4 mM Mg-ATP, and 0.4 mM Na-GTP. Whole-cell voltage–clamp recordings were made from ganglion cells, and currents were recorded in voltage-clamp mode using a MultiClamp 700b (Molecular Devices) patch-clamp amplifier. Signals were filtered at 4 kHz (four-pole Bessel), digitized at 20 kHz with an ITC-18 (HEKA Elektronik) data acquisition board, and saved to a personal computer for off-line analysis using a custom acquisition software in Igor Pro 8 (WaveMetrics).

Pharmacological agents when used were added to the superfusate. Inhibition was blocked with a cocktail of 5 μ M strychnine (Sigma-Aldrich), 5 μ M SR-95531, and 50 mM TPMPA (Tocris Bioscience), to block glycine, GABA_A, and GABA_C receptors, respectively. Gap junctions were blocked using 100 μ M meclofenamic acid (MFA; Sigma-Aldrich), and rod BCs were hyperpolarized using 5 μ M L-AP4 (Tocris Bioscience).

Cell identification. RGCs were targeted for morphological analysis using an electrode containing internal solution with the addition of 2% Lucifer yellow (LY; Invitrogen). Cells were whole-cell voltage-clamped to record the functional response properties; during the recording, LY was diffused into the cell. The retina was then removed from Ames' media and fixed in 4% paraformaldehyde in 0.1 M phosphate buffer (pH 7.4) for 45 min. Retinas were then washed in 0.1 M phosphate buffer for 25 min before being placed in 3% agarose (Sigma-Aldrich). Agarose blocks containing the retina were oriented vertically on a PELCO easiSlicer (Ted Pella) and sliced at 200 µm width. The sliced retina was plated and imaged on an epifluorescence microscope (Olympus, BX43). The inner and outer boundaries of the inner plexiform layer (IPL) were determined by the ganglion cell layer and the inner nuclear layer (Fig. 2). Cells were classified as ON, OFF, or ON/OFF based on dendrite depth within the IPL. The top boundary was denoted as 0% depth, and the bottom boundary was 100% depth; based on prior methods from Margolis and Detwiler (2007), we used 50% as the ON/OFF subliminal boundary. Dendritic arbors were traced using NeuronJ (ImageJ, National Institutes of Health).

Analysis. Data processing and statistical analyses were performed in Igor Pro 8. Conductance analysis was performed in a similar method to previous studies (Borg-Graham, 2001; Oesch and Taylor, 2010). Stimulus-evoked responses were recorded at 13 holding potentials (from -80 to +40 in 10 mV steps, Fig. 1B,C). I–V relationships were measured at 5 ms intervals for the duration of the voltage step, relative to the baseline in the preceding 500 ms. Voltage series were repeated two times and averaged. Synaptic currents were assumed to arise from a sum of linear excitatory and inhibitory synaptic inputs. Excitation is mediated by nonselective cation channels with a reversal potential of $V_e = 0$ mV, while inhibition is mediated by chloride channels with a reversal potential of $V_i = -52$ mV. Each I–V was fit with a line between -80 and -40 mV where the I–V relation was most linear. The slope and intercept were determined from the fit, thus producing a discrete measurement of conductance at every point.



Figure 1. Experimental setup. *A*, 4×8 , 30 µm iridium oxide stimulating array. *B*, Schematic of subretinal stimulation and current-clamp recording. *C*, Voltage-step paradigm. Voltage from -80 to +40 mV. A biphasic, anodic-first, 1 ms, 20 µA stimulation pulse occurred at time 1.5 s, halfway through the voltage step. *D*, Example currents recorded at each voltage step.



Figure 2. Analysis of cell morphology. *A*, Image of example filled cell in whole mount. *B*, The same retina vertically sliced at 200 µm to show dendritic stratification. *C*, Semiautomatic tracing performed in NeuronJ. *D*, Extracted skeleton from tracing. *E*, Boundaries of IPL defined at the edges of the ganglion cell layer and inner nuclear layer. *F*, The skeleton alighted to boundaries identified in (*E*).

Results

Inhibition performs many functions in the retina to process signals, such as center-surround organization (Cook and McReynolds, 1998; Taylor, 1999; Flores-Herr et al., 2001), gain modulation (Sagdullaev et al., 2006; Oesch and Diamond, 2019; Nagy et al., 2021), decorrelation (Franke et al., 2017), and mediating information transfer between ON to OFF pathways (Molnar et al., 2009; for review, see Diamond and Lukasiewicz, 2012). The role of inhibition in response to electrical stimulation or how inhibition might function in the rd10 retina is not yet clear (Cameron et al., 2013; Walston et al., 2018). To examine excitatory and inhibitory network responses to electrical stimulation in the rd10 retina, we used whole-cell voltage-clamp techniques to measure stimulation-evoked synaptic conductance. We recorded electrical stimulation-evoked currents from 60 cells over a range of holding potentials between -80 and +40 mV. We found that reversal potentials for stimulation-evoked currents were not normally distributed (Shapiro-Wilk test; p = 0.0006) and had three distinct modes (Fig. 3A). One mode was dominated by excitatory conductance, one dominated by inhibitory conductance, and one weaker intermediate mode with an equal mix of both excitation and inhibition. We divided the distribution into three groups centered around these modes by setting boundaries at the minima between the peaks of the multipeak Gaussian fit. Points where the fit decreased to zero were considered boundaries. Group 1 ranged from +40 to -10 mV (n = 23), Group 2 ranged from -80 to -35 mV (n = 27), and Group 3 ranged from -35 to -10 mV (n = 10). Plotting the average I-V and SEM for each mode showed distinct and nonoverlapping I-V relations for each group, which had roughly similar total conductance despite distinct reversal potentials (Fig. 3B). When we compared the excitatory and inhibitory conductance for the three groups identified by the distribution of I-Vs, we found that Group 1 excitation was significantly larger than inhibition and Group 2 cells had inhibition that was significantly larger than excitation $(p = 5.9 \times 10^{-7}, t \text{ test}; p = 3.8 \times 10^{-9}, t \text{ test})$. When we plot the difference between peak inhibitory and excitatory conductance, we find the SD of each group does not overlap with the neighboring groups (Fig. 3C).

Given the large sample of RGCs within each group, it is likely that each group is itself comprised of a heterogeneous population of RGC subtypes, but we reasoned that these broad E/I groups may share some properties within groups. Cells in Groups 2 and 3 displayed fast inhibition with a similar time to peak (p = 0.82, Tukey's post hoc test) and little variation in the population of waveforms. In contrast, Group 1 cells displayed a significantly slower time to peak for inhibitory conductance and significantly more variability in responses, possibly indicating a diversity of network-mediated inhibitory effects contributing to Group 1 inhibition ($p = 2.05 \times 10^{-5}$, ANOVA; p < 0.05, Bartlett's variance test; Fig. 3*D*,*E*). Although Group 2 cells had very little excitatory conductance, the kinetics of excitation were similar for all groups (Fig. 3*F*,*G*).

Morphological and functional classification

Previous work suggested that similar patterns of light-evoked E/I balance correspond to ON and OFF functional groups for certain subtypes of RGCs (Pang et al., 2003; Murphy and Rieke, 2006; Roska et al., 2006; Margolis and Detwiler, 2007), and we hypothesized that electrically evoked E/I Groups 1, 2, and 3 may correspond to ON, OFF, and ON/OFF RGCs, respectively. Margolis et al. (2008) found a similar pattern of E/I balance in the spontaneous synaptic inputs between



Figure 3. Functional grouping for 60 cells. *A*, Cells were divided into three groups based on the histogram of their reversal potential; dotted lines represent boundaries for individual groups. *B*, Average I–V curves for each cluster. Error bars represent 1 SD. Cluster A (red line, n = 23), Cluster B (blue line, n = 27), and Cluster C (black line, n = 10). *C*, Peak Gi–peak Ge for cells from *B*; shading represents 1 SD for each group. *D*, Average inhibitory conductance traces for each cluster. Shading represents SEM. *E*, Box plot for excitatory conductance time to peak values for Cluster A, Cluster B, and Cluster C. *F*, Average excitatory conductance traces for each cluster. *G*, Same as *E* but for inhibitory conductance. Error bars represent 1 SD; red diamond is the mean; * indicates significance p < 0.05.

morphologically identified ON and OFF cells in the rd1 mouse; however, the E/I ratio of stimulation-evoked synaptic inputs was not measured for light or electrical responses. Whether these patterns of stimulation-evoked excitation and inhibition hold true following degeneration has not yet been shown. We examined spontaneous excitatory and inhibitory inputs to RGCs in our groups to see if stimulation-evoked E/I groups were consistent with previous findings for spontaneous activity. We found an ~10-fold difference in the power of inhibitory conductance between Group 1 and Group 2 RGCs in the frequency range of 2–12 hz (Fig. 4*B*); consistent with stimulation-evoked E/I, groups corresponded to ON and OFF RGCs. This indicates that the E/I balance of electrically evoked synaptic inputs in Group 1 and 2 RGCs is consistent with previous findings of a unique E/I balance for degeneration-induced spontaneous synaptic inputs in ON and OFF RGCs, respectively.

To determine if there was a correlation between physiological groups and ON and OFF morphological structure, we filled cells with LY and recovered RGC morphology postrecording in a subset of recordings (n = 12, at least three from each

А



Figure 4. Oscillatory currents and conductance. *A*, Example current traces for a putative ON (top), OFF (middle), and ON/OFF (bottom) cell at holding potentials of 0 mV and –70 mV. *B*, Average excitatory and inhibitory conductance PSD for all ON (top) and OFF (bottom) cells. Gray represents excitatory conductance, and solid black represents inhibitory conductance; error bars are 1 SD.

group, Fig. 5). For all filled cells, the recovered cell morphology matched the predicted functional group: Cluster A corresponded to ON cells, Cluster B corresponded to OFF cells, and Cluster C corresponded to ON/OFF cells. Thus, both morphological and functional analyses of clusters indicate that the three electrical stimulation-evoked response patterns we observed are representative of the three main classes of RGCs, ON, OFF, and ON/OFF.

Our results demonstrate three separable groups based on E/I with ON/OFF RGCs occupying an intermediate E/I group. Interestingly, although past work only focused on a few subtypes of RGCs (Pang et al., 2003; Murphy and Rieke, 2006), our large sample of RGC recordings indicates that this stereotypical E/I grouping may apply to many subtypes of RGCs (Roska et al., 2006), although we did not confirm the individual subtype identity.

Mechanisms underlying E/I ON and OFF pathway differences

Given that subretinal electrical stimulation is expected to activate ON and OFF BCs to the same extent (Farzad et al., 2023; but see Twyford et al., 2014), we were surprised to find that OFF RGCs received little excitatory input. There are three primary possibilities to account for this: (1) stimulation-evoked inhibition suppresses OFF BCs, (2) OFF BCs may be hyperpolarized during degeneration and are unable to reach threshold, or (3) OFF BCs are not excited by electrical stimulation.



Figure 5. Morphological analysis of recorded RGCs. RGCs were filled with LY during the recording, fixed and sliced vertically to image dendritic termination in the IPL. *A*, Top: RGC dendrites terminating in the ON layer of the IPL. Left: example excitatory and inhibitory conductance trace from the filled cell. Right: I–V curve from the filled cell. *B*, Same as *A* for a different filled RGC with dendrites terminating in the ON layer. *C*, Top: RGC dendrites terminating in the OFF layer of the IPL. Left: example excitatory and inhibitory conductance trace from the filled cell. Right: I–V curve from the filled cell. *B*, Same as *A* for a different filled RGC with dendrites terminating in the OFF layer of the IPL. Left: example excitatory and inhibitory conductance trace from the filled cell. Right: I–V curve from the filled cell. *D*, Same as *C* for a different filled RGC with dendrites terminating in the OFF layer of the IPL. Left: example excitatory and inhibitory conductance trace from the filled cell. *B*, Same as *C* for a different filled RGC with dendrites terminating in the OFF layer. *E*, Top: RGC dendrites terminating in the OFF layer of the IPL. Left: example excitatory and inhibitory conductance trace from the filled cell. *B*, Same as *E* for a different filled RGC with dendrites terminating in the OFF layer. Inhibitory conductance, and gray dotted traces represent excitatory conductance.

Presynaptic inhibition of BC terminals is a well-described inhibitory motif in the retina. We therefore hypothesized that stimulation may activate some amacrine cells, either directly or indirectly, to limit the OFF BC output during stimulation. To test this possibility, we recorded from RGCs during the GABA_A, GABA_C, and glycine receptor block with SR-95531, TPMPA, and strychnine, respectively. Consistent with inhibitory suppression of the OFF BC output, OFF RGC excitatory conductance was significantly increased in the presence of inhibitory antagonists from 1.28 ± 5.96 to 4.88 ± 1.88 nS (p = 0.02; paired *t* test; n = 5; Fig. 6*A*,*C*). For ON RGCs, the average excitatory conductance did not change, although individual cell responses were quite heterogeneous with some showing an increase and some showing a decrease in excitation (Fig. 6*B*,*D*). This suggests that ON BCs may receive more diverse inhibitory signaling than OFF BCs, where some are suppressed by presynaptic inhibition and in others inhibition may function to increase gain (Sagdullaev et al., 2006; Eggers et al., 2007, p. 200; Oesch and Diamond, 2019; Nagy et al., 2021). In contrast, OFF cells showed a consistent increase in excitatory conductance under inhibitory blockade, indicating that presynaptic inhibition plays a uniform role in gating the excitatory output of OFF BCs to OFF RGCs, at least during electrical stimulation.

Given that the glycinergic All amacrine cell (All) is well known to inhibit the OFF bipolar output during ON pathway excitation, as in the primary night vision pathway (Murphy and Rieke, 2008; Münch et al., 2009; Tian et al., 2010), we hypothesized that Alls mediate OFF BC inhibition during stimulation. To examine this, we applied the glycine receptor antagonist strychnine. We found that on average OFF cell excitatory conductance increased, although to a smaller degree than for the complete inhibition block, from 2.61 ± 2.6 to 6.93 ± 4.98 nS (Fig. 7A,E; p = 0.018; paired *t* test; n = 10). Most OFF RGCs showed this effect (8 out of 10), indicating that most, but not all, presynaptic gating of OFF excitation is mediated by glycinergic amacrine cells. ON cells displayed a heterogeneous mix of responses during the glycine block but on average had the same amount of peak excitatory conductance, similar to results for the complete inhibition block (Fig. 6B,F).



Figure 6. Change in excitatory conductance with application of inhibitory antagonists SR-95531, TPMPA, and strychnine. *A*, A pairwise plot of peak excitatory conductance between the control and inhibition block groups in OFF cells (n = 5). Red diamonds indicate the mean. *B*, Same as *A* but for ON cells (n = 7). *C*, *D*, Average excitatory conductance trace for cells from the panels above; shading represents 1 SD. * indicates significance p < 0.05.

Many OFF RGCs receive direct inhibitory inputs from All amacrine cells (Manookin et al., 2008; Murphy and Rieke, 2008; Münch et al., 2009; van Wyk et al., 2009; Anderson et al., 2011) or other glycinergic amacrine cells (Buldyrev et al., 2012). For most OFF RGCs (6 out of 10), direct inhibitory conductance also decreased in strychnine compared with that in the control (Fig. 7*C*,*G*; 16.7 ± 9.03 to 6.98 ± 8.82 nS; p = 0.009; paired *t* test; n = 10). These six RGCs were also among the cells that showed glycine-mediated gating of excitation, leaving two cells which showed glycine-mediated presynaptic modulation of excitation, but no direct glycinergic inputs. This indicates that most OFF RGCs that receive direct glycinergic input also receive inputs from OFF BC subtypes that are modulated by glycinergic inputs. For ON RGCs, inhibition was not significantly changed (p = 0.33, *t* test) in the glycine receptor blockade with no distinct pattern between the direction of excitation and inhibition between the control and strychnine (Fig. 6*D*,*H*; 3.56 ± 9.03 nS to 1.85 ± 2.63 nC).

The ON pathway shapes OFF cell inhibition

Our results show that glycinergic inhibition mediates both direct inhibitory inputs to a majority of OFF RGCs and gates excitatory inputs to the same subset of OFF RGCs. Although a variety of glycinergic amacrine cell subtypes could be involved in both pre- and postsynaptic inhibition (Buldyrev et al., 2012), the All amacrine cell can potentially mediate both effects and has been heavily implicated in both functions (Manookin et al., 2008; Münch et al., 2009; van Wyk et al., 2009; Buldyrev and Taylor, 2013; McLaughlin et al., 2021); for review, see Demb and Singer (2012). Considering this, we hypothesized that crossover signals from the ON pathway may be mediating inhibitory signaling to both OFF BC terminals and OFF RGCs. To examine the contributions of the ON cone BCs and rod bipolar-mediated inputs that would drive All-mediated crossover inhibition, we first applied the gap junction blocker MFA, which should block signals that arise in the ON cone BCs. OFF RGC excitatory inputs remained unchanged in the MFA block (Fig. 8*A*,*E*). However, during the gap junction block, inhibitory inputs to OFF RGCs were decreased but were not significantly different from the control (Fig. 8*B*,*F*, paired *t* test, p = 0.1). Together this indicates that activation of OFF RGCs. Although MFA is often



Figure 7. Change in conductance in the glycine block. *A*, A pairwise plot of peak excitatory conductance between control and strychnine in OFF cells (n = 10). Red diamonds indicate the mean. *B*, Same as *A* but for ON cells (n = 10). *C*, A pairwise plot of peak inhibitory conductance between the control and strychnine in OFF cells (n = 10). *D*, Same as *C* but for ON cells (n = 10). *E*–*H*, Average conductance traces for cells from the subsequent panels above; shading represents 1 SD. * indicates significance p < 0.05.



Figure 8. Contributions of the ON pathway to OFF cell inhibition and excitation. *A*, A pairwise plot of peak excitatory conductance between control and MFA groups in OFF cells (n = 6). Red diamonds indicate the mean. *B*, Same as in *A* but for inhibitory conductance. *C*, A pairwise plot of peak excitatory conductance between control and MFA and AP4 groups in OFF cells (n = 6). *D*, Same as in *C* but for inhibitory conductance. *E*, Average excitatory conductance trace for cells from each group; shading represents 1 SD. *F*, Same as *E* but average inhibitory conductance. * indicates significance p < 0.05.

used to dissect this circuit (Liang and Freed, 2010), it is important to note that there are numerous gap junctions throughout the retina and the effect of MFA be mediated by effects at other gap junctions or even off-target effects at other receptor systems (Lee and Wang, 1999; Peretz et al., 2005).

Stimulation of rod bipolar cells (RBCs) could also be responsible for driving All activity. Since we cannot block the RBC-mediated glutamatergic drive to Alls without blocking excitatory inputs to OFF RGCs, we reasoned that we might reduce their contribution by hyperpolarizing RBCs with the mGluR6 agonist AP4 (Slaughter and Miller, 1983) in addition

to MFA. It is important to note that this may not completely eliminate the RBC to All input, because electrical stimulation may still be sufficient to evoke RBC release in the presence of AP4.

Despite this caveat, we observed a 50% reduction in inhibitory inputs to OFF RGCs when blocking both gap junctions and attenuating RBC input. This indicates that the RBC to the All pathway does contribute to direct inhibition of OFF RGCs (p = 0.03, paired *t* test, Fig. 8*D*), thus implicating the All. We did not, however, observe a significant increase in excitatory conductance in OFF RGCs (Fig. 8*A*,*C*). These results indicate that separate populations of glycinergic amacrine cells may contribute to pre- and postsynaptic OFF RGC inhibition and that postsynaptic inhibition involves a crossover inhibitory pathway. Alternatively, it is possible that one cell such as the Alls mediates both types of inhibition, but its output synapses must have different input–output functions for bipolar and ganglion cell synapses, which differentially process information routed to these two targets. While our data supports the All as contributing to crossover signals, we cannot rule out other amacrine cells or disinhibition of amacrine cells also playing a role in this phenomenon.

Remnant cone effect on E/I balance

We have interpreted these findings as resulting from direct stimulation of bipolar and/or amacrine cells; however, recent work has demonstrated that cone inner segments remain intact and light responsive in the rd10 mouse for a significant amount of time (around p67) after outer segment degeneration (Ellis et al., 2023). Given that electrical cone stimulation would increase glutamate release, increasing OFF pathway activity and decreasing ON activity, it is not obvious how cone activation could explain our results. Regardless, it may be possible that some residual cone inner segments remain and contribute to abnormal signaling in the ON or OFF pathways. Since cones eventually fully degenerate, the influence of the putative residual cone-mediated responses should decrease with age if they contribute. We plotted peak excitatory and inhibitory conductance against age for ON and OFF RGCs. For OFF RGCs, we found that age had no effect on peak excitatory or inhibitory conductance (Fig. 9A). In ON RGCs, there was a significant decrease in the excitatory conductance (Fig. 9B, Pearson's test, R = -0.475). This could indicate that residual cone inner segments provide some contribution to ON excitation; alternatively, this could be due to other disease-mediated changes in ON pathway synaptic transmission (Puthussery and Taylor, 2010). Given that Ellis et al. showed that residual cones should be almost completely gone by our youngest ages, we favor the latter hypothesis, although we cannot distinguish between these two possibilities here.

E/I groupings are present in both the rd10 and wt retina

Past work has demonstrated E/I ratios in the wild-type (*wt*) retina consistent with the E/I balance we observed in the *rd10* retina, but only in a few identified subtypes. It may be possible that the correlation between E/I ratios extends broadly across multiple subtypes in the *wt* retina, as our data for *rd10* suggest, or it may be that the general relationship between E/I ratio across ON and OFF RGC class is unique to retinal degeneration. To address this question, we measured E/I balance for electrical stimulation in the *wt* retina as we did for *rd10* RGCs. We found *wt* cells had a distribution of reversal potentials similar to those observed in *rd10* corresponding to three distinct groups (Fig. 10*A*–*C*). These groups once again showed distinct I–V relations, with reversal potentials consistent with *rd10* for Groups 1 and 3; however, *wt* Group 2 cells had a more positive reversal potential than *rd10* Group 2 cells (Fig. 10*B*). Consistent with this observation, when we compared the E/I ratio between *rd10* and *wt* OFF cells, there was a significant increase in the E/I ratio from *rd10* to *wt* (Fig. 10*D*); *p* = 0.0009; *t* test). ON cells showed no difference in E/I ratio between *rd10* and *wt* (Fig. 10*D*).

To examine the source of this difference in reversal potential, we compared the magnitude of excitatory and inhibitory conductance between rd10 and wt OFF RGCs. We found that excitatory conductance was significantly greater (p = 0.001, t test) in wt OFF RGCs compared with that in rd10 OFF RGCs, but there was no significant difference in peak inhibitory conductance (Fig. 10*E*, p = 0.18, t test). Therefore, greater excitation, not reduced inhibition, drives the more positive reversal potential and E/I ratio closer to 1 for OFF RGCs in wt compared with those in rd10. Given that we observed significant presynaptic inhibition of excitatory OFF inputs to RGCs in rd10 retina, we reasoned that this difference may be due to increased presynaptic inhibition in rd10 OFF circuit. To test this hypothesis, we compared the magnitude of excitatory conductance in wt OFF RGCs to the magnitude of OFF RGC conductance when we blocked glycine, to relieve presynaptic glycinergic inhibition to the OFF BCs. When glycinergic inhibition was blocked, the difference between wt and rd10 OFF RGCs was eliminated (Fig. 10*F*, p = 0.3, t test). For ONs, we found no significant differences between ON excitatory and inhibitory conductance (Fig. 10*G*, p = 0.9, t test) showing that the excitation—inhibition balance of the ON pathway is not disrupted in degeneration. Taken together, this indicates abnormal inhibitory signaling in the rd10 OFF RGCs.

Discussion

Here we provide evidence that electrical stimulation differentially modulates responses in the ON and OFF pathways. Similar to light stimulation, electrically evoked inputs for ON cells have an E/I ratio >1 with excitation dominating, while OFF cells had an E/I ratio <1 weighted toward inhibition. For ON cells, the time course of inhibitory conductance was variable suggesting multiple diverse sources of inhibitory inputs, while OFF cells had fast inhibitory conductance with little variance in kinetics, indicating more stereotyped inputs. We showed that reduced OFF excitation is due to the inhibition



Figure 9. Peak excitatory and inhibitory conductance as a function of animal age. *A*, OFF cell excitatory (open circle) and inhibitory (closed circle) peak conductance plotted against age. Linear regression plots for each. *B*, Same as *A* for ON cells.

of OFF BC terminals, which is largely mediated by glycinergic inhibition. We also showed that the ON pathway significantly contributes to the inhibition of OFF RGCs, but not OFF BC terminals, and that gap junctions are not required to drive inhibitory outputs. ON/OFF cells displayed a proportional excitatory to inhibitory balance indicating that the ON and OFF layer inputs are activated equally by electrical stimulation in contrast to the asymmetry seen for ON and OFF RGCs separately. Finally, we showed that inhibition to OFF BCs may be abnormally strong in the *rd10* retina, causing a lower E/I balance in OFF cells in the *rd10* retina compared with those in the *wt* retina. Taken together, these results show that the consideration of how inhibitory circuits are activated by prosthetic electrical stimulation is necessary to guide rational stimulus design for vision restoration and that changes resulting from degeneration also need to be considered.



Figure 10. Functional grouping for *rd10* versus *wt* ON and OFF cells. **A**, Cells clustered into three groups based on the histogram of their reversal potential; dotted lines represent cutoff points for individual groups. **B**, Average I–V curves for each cluster. Shading represents 1 SD. Cluster A (red line, n = 15), Cluster B (blue line, n = 12), and Cluster C (black line, n = 10); dotted lines represent the average I–Vs for *rd10* groups in Figure 2. **C**, Peak Gi–peak Ge for cells form **B**; triangles indicate *wt*, and dots indicate *rd10* cells; shading represents 1 SD for each group. **D**, E/I ratios for OFF cells (black) and ON cells (red); dotted line shows 1 on the right axis. **E**, Violin plots showing the peak excitatory and inhibitory conductance in OFF cells of the *wt* and *rd10* retina. **F**, A violin plot comparison of *wt* Ge versus *rd10* Ge in strychnine application. **G**, Same as **D** but for ON cells. Red diamond is the mean; * indicates significance p < 0.05.

Putative functional class grouping

Past work has shown that α ganglion cells show an asymmetry in E/I balance between some ON and OFF RGCs, similar to what we have shown for *rd10* RGCs here (Pang et al., 2003; Murphy and Rieke, 2006). Some evidence suggests this trend may extend to other RGC cell types (Roska et al., 2006), although E/I balance has yet to be systematically explored for all RGC subtypes. Our data for *wt* indicates that this general trend may hold true for a broad population of RGC subtypes. Our results also showed that ON/OFF RGCs have an equal E/I balance, consistent with other studies that have measured this, primarily in direction-selective ganglion cells (DSGCs; Poleg-Polsky and Diamond, 2016; Jain et al., 2020). In this work, instead of focusing on a subset of RGC types such as α ganglion cells, we attempted to sample across a large population of RGCs in an unbiased manner.

We performed whole-cell voltage–clamp recording and conductance analysis on 60 RGCs. As expected, this resulted in a population with heterogeneous response properties; however, the distribution of properties such as reversal potential was not unimodal. We identified two main modes of reversal potential consistent with conductance dominated by excitation and inhibition. In between these two modes was a weaker third mode. Although these three modes were not perfectly separable, we set boundary conditions at the troughs between modes (Fig. 3A) The individual groups resulting from this strategy were well separated, and the SDs of any one group did not overlap with the others (Figs. 3C, 10C). Although there were a small number of cells which lie close to the boundaries between modes and could potentially be misclassified, this only amounts to 3–6 cells out of 60 cells and excluding these cells does not change the conclusions of any subsequent analysis (data not shown).

Although we only filled a subset of cells (n = 12) to recover morphology, all the morphologies that we recovered matched the putative ON, OFF, and ON/OFF groups. We cannot be certain there are no deviations from this pattern in the entire sample; however, the majority of RGCs in each mode likely reflect the three distinct functional groups of RGCs. Importantly, the *wt* retina showed a similar multimodal distribution of reversal potential and E/I balance, albeit with a positive shifted E/I balance in the putative OFF group. Taken together, we believe that this work extends the basic finding of

asymmetric E/I balance between ON and OFF RGCs to be a trend among many RGC subtypes rather than a subset. It also suggests that equal E/I ratio may be a general property of ON/OFF RGCs extending beyond DSGCs.

The role of inhibition in prosthetic vision

Inhibition has many integral roles in vision formation (see above, Discussion) and has been shown to have a modulatory effect on degeneration-induced spontaneous activity (Margolis and Detwiler, 2011; Haq et al., 2014; Margolis et al., 2014; Carleton and Oesch, 2022), but the effect of electrical stimulation on inhibitory circuitry remains largely unexplored. Amacrine cells are the most diverse cell type of the retina with 60 identified cell types serving unique functions (Masland, 2012; Yan et al., 2020). These cells shape spatiotemporal responses to stimuli, create directional responses, and create center-surround responses, among a myriad of other integral mechanisms in creating vision; for reviews, see Diamond (2017) and Masland (2012). The field of retinal prosthetics has thus far focused on the generation of spikes with less attention on the underlying mechanisms that drive spiking during prosthetic stimulation (but see Walston et al., 2018). The inner nuclear layer is the main target of stimulation for subretinal prosthetics, and while BCs are depolarized in response to a stimulation of amacrine and BCs is important in understanding how the degenerated circuit will respond to prosthetic stimulation. Prior work assessing the electrical stimulation of starburst amacrine cells identified that they were activated by inputs from presynaptic cells and not direct electrical stimulation; however, these studies were performed in the healthy rabbit retina, where photoreceptors and retinal thickness may impact the direct stimulation of the INL (Tsai et al., 2011; Cameron et al., 2013).

Previous studies have shown that electrical stimulation preferentially activates ON responses, although this has only been confirmed in studies on the healthy retina (Im and Fried, 2016; Lee and Im, 2019). Moreover, retinal prosthetic user reports are also consistent with preferential stimulation of the ON pathway (Humayun et al., 2003, 2012; Fujikado et al., 2007; Naycheva et al., 2012; Luo et al., 2016). How the true OFF pathway is recruited during retinal prosthetic stimulation is less certain, although OFF responses in some RGCs can be formed (Sekhar et al., 2017; Ho et al., 2018, 2020). Our finding that the OFF pathway receives significant inhibition during electrical stimulation may provide insight to past reports of poor OFF RGC responses to prosthetic stimulation; however, an explanation has not been previously put forth. The observation of altered inhibition in the *rd10* retina is consistent with general compensatory changes in inhibitory signaling shown in response to acute photoreceptor ablation (Care et al., 2020; Lee et al., 2022), although the specificity for the OFF versus ON pathway and presynaptic versus postsynaptic inhibition was not reflected in these studies. With a more complete understanding of the mechanisms driving electrical responses in ON and OFF RGCs, we may be able to better target stimulation that allows for pathway-specific activation.

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