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A pixel-encoder retinal ganglion cell with spatially offset excitatory and inhibitory receptive fields

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Graphical Abstract

Highlights
- A pixel-encoder retinal ganglion cell type (PixON-RGC) is labeled in Grik4-Cre mice
- PixON-RGCs have spatially offset excitatory and inhibitory receptive fields
- PixON-RGCs encode local image contrast approximately linearly
- PixON-RGCs project to the dLGN and likely contribute to visual perception

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In Brief
Johnson et al. genetically identify a pixel-encoder retinal ganglion cell type in mice (PixON-RGCs). PixON-RGCs have spatially offset excitatory and inhibitory receptive fields and encode local image contrast approximately linearly. Their axons project to the dorsolateral geniculate nucleus of the thalamus indicating that PixON-RGCs likely contribute to visual perception.
A Pixel-Encoder Retinal Ganglion Cell with Spatially Offset Excitatory and Inhibitory Receptive Fields

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SUMMARY

The spike trains of retinal ganglion cells (RGCs) are the only source of visual information to the brain. Here, we genetically identify an RGC type in mice that functions as a pixel encoder and increases firing to light increments (PixON-RGC). PixON-RGCs have medium-sized dendritic arbors and non-canonical center-surround receptive fields. From their receptive field center, PixON-RGCs receive only excitatory input, which encodes contrast and spatial information linearly. From their receptive field surround, PixON-RGCs receive only inhibitory input, which is temporally matched to the excitatory center input. As a result, the firing rate of PixON-RGCs linearly encodes local image contrast. Spatially offset (i.e., truly lateral) inhibition of PixON-RGCs arises from spiking GABAergic amacrine cells. The receptive field organization of PixON-RGCs is independent of stimulus wavelength (i.e., achromatic). PixON-RGCs project predominantly to the dorsal lateral geniculate nucleus (dLGN) of the thalamus and likely contribute to visual perception.

INTRODUCTION

The output of retinal computations is conveyed to the brain through the spike trains of >30 retinal ganglion cell (RGC) types. Recent surveys in mice have highlighted the morphological and functional diversity of RGCs (Sanes and Masland, 2015; Baden et al., 2016; Helmstaedter et al., 2013), but few RGC types have been genetically identified and studied in detail. Therefore, what specific information RGCs encode, how excitatory and inhibitory circuits give rise to their characteristic light responses, and where in the brain RGCs send their information remains, for most cell types, unknown.

RGCs serve at least three broad functions: (1) to encode local contrast in the retinal image and support visual perception (Sinha et al., 2017; Schwartz et al., 2012; Pang et al., 2003; van Wyk et al., 2009); (2) to detect specific forms of motion and drive reflexive behaviors (Huang et al., 2017; Münch et al., 2009; Sabbah et al., 2017; Zhang et al., 2012); and (3) to measure overall luminance and regulate non-visual light-dependent processes (Chen et al., 2011; Lazzerini Ospri et al., 2017). Individual RGC types can contribute to more than one of these functions, which involve different subcortical targets (Dhande et al., 2015). Recent studies in mice have identified numerous RGC types that detect specific forms of motion (Zhang et al., 2012; Jacoby and Schwartz, 2017; Münch et al., 2009; Borst and Euler, 2011) and intrinsically photosensitive RGC (ipRGC) types that regulate non-visual light-dependent processes (Chen et al., 2011; Lazzerini Ospri et al., 2017). What murine RGC types contribute to visual perception, which relies on signal propagation from the retina to dorsal lateral geniculate nucleus (dLGN) and primary visual cortex, is less well understood (Kerschensteiner and Guido, 2017). In primates, midget and parasol RGCs dominate input to primary visual cortex (Wässle et al., 1989; Field and Chichilnisky, 2007). Midget RGCs encode contrast and spatial information approximately linearly (i.e., pixel encoders) and mediate high acuity pattern vision, whereas parasol RGCs integrate spatial information nonlinearly and are sensitive to motion irrespective of the precise image patterns involved (Field and Chichilnisky, 2007; Petrusca et al., 2007; Cafaro and Rieke, 2013). Alpha RGCs in mice share response properties with parasol RGCs in primates and provide motion-sensitive input to dLGN (Cruz-Martín et al., 2014; Pang et al., 2003; Rompani et al., 2017; Schwartz et al., 2012). Whether the mouse retina contains a pixel-encoder RGC type that projects to dLGN, and, if so, what circuit mechanisms give rise to its responses remains unknown.

Throughout the nervous system, excitatory and inhibitory synaptic inputs shape the computations of neurons (Isaacson and Scanziani, 2011). Many RGCs receive excitation and inhibition via one of two circuit motifs (Cafaro and Rieke, 2013; Demb and Singer, 2015; Roska et al., 2006; Roska and Werblin, 2001). In feedforward inhibition, the same bipolar cells that excite an RGC activate amacrine cells that inhibit the RGC. In crossover inhibition, bipolar cells with opposite contrast preferences (ON versus OFF) excite and inhibit—via amacrine cells—an
RGC (Cafaro and Rieke, 2013; Demb and Singer, 2015; Roska et al., 2006; Roska and Werblin, 2001). In both motifs, excitatory and inhibitory receptive fields have their highest sensitivity over the dendritic field of the RGC (i.e., the receptive field center), with inhibition extending laterally beyond excitation (i.e., the receptive field surround). By contrast, inhibitory receptive fields of direction-selective ganglion cells (DSGCs) are shifted side-ways relative to their excitatory receptive fields and dendrites (Fried et al., 2002; Wei et al., 2011; Yonehara et al., 2011), indicating that non-canonical arrangements of excitatory and inhibitory receptive fields help diversify RGC light responses.

Here, we genetically identify a pixel-encoder RGC type in mice (PixON-RGCs). In two-photon-guided patch-clamp recordings, we find that PixON-RGCs encode local image contrast approximately linearly and identify a novel circuit motif (i.e., truly lateral inhibition), which gives rise to their responses. Combining genetic labeling and retrograde tracing, we show that PixON-RGCs project predominantly to the dLGN and likely contribute to visual perception.

RESULTS

Genetic, Morphological, and Functional Identification of PixON-RGCs

Characterizing the light responses, circuit mechanisms, and projection patterns of the >30 RGC types, which relay the output of retinal computations to the brain, is a prerequisite to understanding vision. To genetically identify unknown RGC types, we crossed a variety of Cre driver lines, including Grik4-Cre (Nakazawa et al., 2002), to the tdTomato reporter strain Ai9 (Madisen et al., 2010), and targeted fluorescent cells under two-photon guidance for patch-clamp recordings. Except where noted otherwise, we recorded cells in the ventral retina where cones express predominantly S-opsin (Baden et al., 2013; Szél et al., 1992) and presented stimuli in the UV spectrum (peak: 385 nm). In Grik4-Cre:Ai9 retinas, ON and ON-OFF DSGCs were labeled as previously reported (Ivanova et al., 2010; Martersteck et al., 2017). In addition, we frequently encountered RGCs with medium-sized, densely branched dendritic arbors stratifying close to the ganglion cell layer (Figures 1A–1D). These cells had high baseline firing rates, increased spiking in a sustained manner in response to light increments (i.e., ON stimuli), and decreased spiking during light decrements (i.e., OFF stimuli, Figures 1E–1G). Voltage-clamp recordings revealed that excitatory synaptic inputs matched the spike responses of these cells, which, remarkably, received no inhibitory input for stimuli restricted to their dendritic field (spot diameter: 300 μm, Figures 1E, 1H, 1I, and S1), and, which were not intrinsically photosensitive at light levels ranging from twilight to bright daylight (Figure S2)(Cronin et al., 2014; Johnsen et al., 2006; Milner and Do, 2017). Based on their genetic labeling, monomorphic dendrites, and consistent physiological properties, we identify these cells as a single RGC type. We refer to them as PixON-RGCs, because they appear to function as pixel encoders and increase firing to ON stimuli.

Linear Encoding of Contrast and Spatial Information by PixON-RGCs

To allow the brain to infer patterns accurately, pixel-encoder RGCs change their firing rates approximately linearly as...
a function of the light intensity over their dendritic fields, and do not respond to stimuli, including second-order motion (Demb et al., 2001), for which the overall light intensity in this area does not change (Sinha et al., 2017; Field and Chichilnisky, 2007).

Recording spike responses to spots (diameter: 300 \( \mu \text{m} \)) of varying intensity, we found that PixON-RGCs increased and decreased their firing rates approximately linearly to positive and negative contrast steps, respectively (Figures 2A and 2B). These responses matched excitatory synaptic inputs to PixON-RGCs, which increased and decreased linearly from high tonic levels (1.3 ± 0.2 nano-Siemens [nS], mean ± SEM, n = 9) to light increments and decrements, respectively (Figures 2C and 2D).

We next recorded the spike trains of PixON-RGCs during presentation and motion of textures of varying spatial scales. We balanced textures to keep the average light intensity above the dendritic field of each RGC constant (Figure 2E). PixON-RGCs responded neither to presentation nor motion of these textures (Figures 2F–2H), and neither elicited synaptic excitation (Figures 2I and 3K). By contrast, the ON DSGCs and ON-OFF DSGCs labeled in Grik4-Cre:Ai9 retinas responded robustly to both the presentation and motion of texture stimuli (Figure S3). Thus, PixON-RGCs encode contrast and spatial information linearly. They inherit these properties from their excitatory input and receive no synaptic inhibition from their receptive field center.

To characterize the temporal response properties of PixON-RGCs, we recorded spike trains during presentation of Gaussian white noise stimuli. Reverse correlation of responses with the stimulus revealed monophasic temporal filter kernels (Figure S4), with opposite polarity in the receptive field center and surround, in keeping with the sustained responses observed during contrast steps.

**Surround Suppression of PixON-RGCs and Its Mechanisms**

Many RGCs show attenuated responses to large stimuli (Kuffler, 1953; Demb and Singer, 2015). This enhances edges in their representation of the retinal image, which in turn facilitates pattern and object recognition in the brain (Egan et al., 2016; Biederman and Ju, 1988). To explore the stimulus size tuning of PixON-RGCs, we presented spots of varying diameter (20–1,200 \( \mu \text{m} \)) in pseudorandom sequences. ON responses increased up to a stimulus size of 300 \( \mu \text{m} \) and were strongly suppressed by larger...
stimuli (Figures 3A and 3B). Thus, PixON-RGCs fired at only 17.2% ± 8.8% (n = 22) of their maximal response rates when stimulated with 1,200-μm spots. Excitatory synaptic inputs similarly increased up to a stimulus size of 300 μm but were suppressed to a lesser degree than spike responses by larger stimuli (Figures 3C and 3D). Thus, 1,200-μm spots still activated 49.0% ± 9.4% (n = 9) of the maximal excitatory conductance of PixON-RGCs. Although spots restricted to their dendritic fields did not evoke inhibition, we found that larger stimuli gradually activated inhibitory synaptic inputs to PixON-RGCs (Figures 3E and 3F). Because PixON-RGCs received little or no tonic inhibition, only ON stimuli modulated these inputs (Figures 3E, 3F, and S5), and suppression of tonic excitation underlay the spike suppression at light OFF. Thus, PixON-RGCs encode stimulus contrast and spatial information linearly, reflecting properties of excitatory inputs from their receptive field center, and exhibit strong size selectivity (i.e., a preference for local contrast or edges) due to a combination of pre- and postsynaptic inhibition. The latter, unusually, is recruited only by stimuli that extend beyond the dendrites of PixON-RGCs into their receptive field surround.

Kinetics and Mechanisms of Inhibitory Synaptic Inputs to PixON-RGCs

Comparing the timing of synaptic inputs elicited by large stimuli (diameter: 1,200 μm), we found that inhibition and excitation were temporally matched (Figure 4A). Thus, neither the time to peak (Figure 5B) nor the percentages of sustained input (excitation: 47.8% ± 7.0%, inhibition: 46.1% ± 2.6%, n = 15, p > 0.8) were significantly different between excitation and inhibition. Because inhibition, in spite of originating in the receptive field surround, reached PixON-RGCs simultaneously with excitation from the receptive field center, we hypothesized that spiking GABAergic wide-field amacrine cells may be its source. We tested this hypothesis pharmacologically. Both gabazine and tetrodotoxin (TTX) blocked inhibition to PixON-RGCs (Figures 4C–4F), indicating that inhibition is mediated by GABAA receptors and provided by spiking GABAergic wide-field amacrine cells. In addition to using spikes to relay signals from the surround, at least one retinal circuit accelerates inhibition by driving amacrine cells via gap-junctional rather than glutamatergic input from bipolar cells (Farrow et al., 2013). Inhibition to PixON-RGCs was blocked completely by D-AP5 and NBQX (Figures 4G and 4H), suggesting that the spiking GABAergic wide-field amacrine cells in this circuit are activated by conventional glutamatergic input from bipolar cells.

Spatially Offset Inhibitory and Excitatory Receptive Fields of PixON-RGCs

For centered spots, we found that inhibition was restricted to large stimuli, which extended beyond PixON-RGCs’ dendritic fields (Figure 3). This could either be because the wide-field amacrine cells that provide this input need to integrate bipolar cell inputs over large areas to pass spike threshold (i.e., size-thresholded inhibition) or because the wide-field amacrine cells that inhibit a given PixON-RGC receive their excitatory input outside its dendritic field (i.e., spatially offset inhibition). We designed two sets of stimuli to distinguish between these possibilities. First, we gradually eroded a large centered spot (diameter: 1,200 μm) from the inside (Figure 5A). Inhibition elicited by this stimulus declined only after the inner stimulus boundary had receded beyond the dendritic territory of PixON-RGCs (Figures 5A–5C). Second, we presented a stimulus square (side length:
300 μm) in a 3 × 3 grid centered on a PixON-RGC (Figure 5D). Consistently, the center square failed to elicit inhibitory synaptic inputs, whereas all other squares evoked robust inhibition (Figures 5D–5F). The responses of PixON-RGC inhibition differed drastically from those of ONa-RGCs (Figure S6), which receive canonical feedforward inhibition. We conclude that the inhibitory receptive fields of PixON-RGCs are spatially offset from their dendrites and their excitatory receptive fields (Figure S7) and that PixON-RGCs receive inputs from a previously unknown circuit motif, which we refer to as truly lateral inhibition (Figure 5G).

Wavelength Independence of PixON-RGC’s Receptive Field Organization
To test whether the organization of PixON-RGC receptive fields depends on stimulus wavelength and whether these cells encode chromatic information, we recorded PixON-RGCs at different points along the dorsoventral gradient of cone opsin expression (Baden et al., 2013; Szél et al., 1992) noting their position in the retina (Figures 6A and 6B). In addition, we switched from stimulating in the UV spectrum (peak: 385 nm), which favors S-opsin activation, to stimulating with green light (peak: 532 nm), favoring M-opsin activation. Excitatory and inhibitory receptive fields of PixON-RGCs probed with green stimuli in the dorsal (Figures 6C and 6D; n = 4) and ventral (Figures 6G and 6H; n = 3) retina were activated exclusively by light increments and were spatially offset, identical to our observations for UV stimuli in the ventral retina (Figure 3). In addition, with the exception of one cell in the opsin transitional zone (OTZ) (Figures 6E and 6F), the ratio of center excitation and surround inhibition was similar across the retina and stimulus wavelengths. Thus, outside of the OTZ, where color-opponent responses arise in several RGC types without cone-type selective connectivity (Chang et al., 2013), PixON-RGCs appear not to encode chromatic information and their receptive field organization is wavelength independent.

PixON-RGC Axons Project to dLGN
Visual perception relies on retinal input to dLGN (Saalmann and Kastner, 2011; Kerschensteiner and Guido, 2017). To determine whether PixON-RGCs could contribute to visual perception, we retrogradely traced their axonal projection patterns. We injected either cholera toxin B conjugated to Alexa 488 (CTB488) or an adeno-associated virus expressing GCaMP6f in a Cre-dependent manner (AAV9-Syn-FLEX-GCaMP6f) into subcortical visual targets labeled in Grik4-Cre:Ai9 mice (Figure 7A). We then targeted RGCs co-labeled with tdTomato and CTB488 or GCaMP6f for patch-clamp recordings and identified PixON-RGCs by their characteristic morphology (Figure 7B) and light responses. Of the 74 PixON-RGCs included in this study, 23 were targeted in this way. PixON-RGCs accounted for approximately 1/3 of the cells co-labeled after dLGN injections. By comparison, PixON-RGCs made up smaller fractions of co-labeled cells after injections into ventral lateral geniculate nucleus (vLGN) or superior colliculus (SC), and no PixON-RGCs were labeled by injections into the medial terminal nucleus (MTN, Figure 7C). This projection pattern of PixON-RGC axons supports the notion that they contribute to visual perception.

DISCUSSION
In this study, we discover and characterize a pixel-encoder RGC type in the mouse retina (PixON-RGCs). The dendrites of PixON-RGCs, which morphologically resemble cluster 9n cells in the Eyewire museum (Bae et al., 2017), stratify near the boundary between the inner plexiform and ganglion cell layer, and receive...
Figure 5. Spatially Offset Inhibition of PixON-RGCs

(A) Annuli of constant 1,200-µm outer diameter and varying inner diameter were presented as 100% contrast steps (2 s ON) from a gray background (1,500 R*/rod/s).

(B) Representative IPSCs in response to annuli shown in (A).

(C) Summary data plotting inhibitory conductance as a function of the inner diameter of stimulus annuli. Circles (error bars) indicate mean (±SEM) of the recorded population (n = 7).

(D) The display was divided into a 3 × 3 grid of 300- × 300-µm squares and centered on the soma of the recorded cell. Between stimulus presentations all squares had the same luminance (1,500 R*/rod/s). During stimulus presentation, one of the nine squares increased in luminance (100% contrast, 2 s ON).

(E) Representative IPSCs in response to appearance of squares in the positions shown in (A). (F) Map of the normalized inhibitory conductance evoked by the appearance of a square at each position (n = 4). Coloration indicates the strength of inhibition. (G) Sensitivity profiles of excitatory and inhibitory receptive fields in feedforward inhibition, in crossover inhibition, and in truly lateral inhibition.

See also Figures S6 and S7.
responses are coupled with high gain to excitatory input elicited by small stimuli (diameter 300 μm: 17.5 ± 2.8 spikes sp/s/nS). For larger stimuli, spatially offset inhibition reduces the gain of the excitation-spike coupling (diameter 1,200 μm: 8.5 ± 3.6 sp/s/nS), which, together with presynaptic inhibition, accounts for the nearly complete surround suppression. Thus, truly lateral inhibition allows for a combination of high gain in the receptive field center and strong surround suppression. This enables PixON-RGCs to encode relatively faithfully the retinal image, deemphasizing regions of uniform intensity. Because excitation and inhibition are temporally matched (Figure 4), stimulation of the surround suppresses the amplitude of the center response without changing its dynamics. This further simplifies the apparent image encoding of PixON-RGCs.

We find that spiking GABAergic amacrine cells provide the inhibitory input to PixON-RGCs. The most parsimonious explanation for the spatial offset of inhibitory receptive fields is that these amacrine cells receive input and provide output in separate arbor (dendrite and axon, respectively) or separate regions of activity S- or M-opsin, indicating that outside the OTZ where color-opponency can arise in several RGC types without cone-type selective wiring (Chang et al., 2013), PixON-RGCs do not encode chromatic information. A recent paper described an RGC type, referred to as M5 (Stablo et al., 2018), which morphologically resembles PixON-RGCs and which likewise receives sustained excitatory inputs. However, M5 RGCs were suggested to consistently encode chromatic information and to exhibit intrinsic photocurrents. PixON-RGCs did not stain for melanopsin and did not exhibit intrinsic photocurrents at light levels equivalent to bright daylight (Figure S2) (Cronin et al., 2014; Johnsen et al., 2006; Milner and Do, 2017), which elicited large intrinsic photocurrents in M1 RGCs (data not shown). Thus, is seems that despite their morphologic similarity PixON-RGCs and M5 RGCs may be distinct cell types.

Combining retrograde tracing and genetic labeling, we found that PixON-RGC axons project predominantly to dLGN (Figure 7). To what extent information from PixON-RGCs is preserved in dLGN or whether it is recombined with other retinal inputs before
being sent to primary visual cortex remains to be determined. Interestingly, sustained ON responses that linearly encode contrast and spatial information have been recorded in dLGN (Piscopo et al., 2013), and RGCs with morphologies similar to PixON-RGCs have been shown to contribute to relay mode innervation of dLGN neurons (Rompani et al., 2017). These findings suggest that information of PixON-RGC may be preserved in dLGN and relayed faithfully to primary visual cortex. Future studies will have to test this experimentally and probe the contribution of PixON-RGCs to visual perception.

**EXPERIMENTAL PROCEDURES**

**Animals**

Throughout this study, we used Grik4-Cre mice (Nakazawa et al., 2002) (Jackson Laboratories stock #006474) crossed to the tdTomato reporter strain Ai9 (Madien et al., 2010) (Jackson Laboratories stock #007909) to label PixON-RGCs. We isolated retinas from young adult mice (postnatal day P21–P40) of both sexes. All experiments in this study were approved by the Institutional Animal Care and Use Committee of Washington University School of Medicine and were performed in compliance with the NIH Guide for the Care and Use of Laboratory Animals.

**Tissue Preparation**

Mice were deeply anesthetized with CO2, killed by cervical dislocation, and encéphalised. For patch-clamp recordings, mice were dark adapted overnight before their retinas were isolated under infrared illumination (>900 nm) in oxygenated mACSFHEPES containing (in mM) 119 NaCl, 2.5 KCl, 1 NaH2PO4, 2.5 CaCl2, 1.3 MgCl2, 20 HEPES, and 11 glucose (pH adjusted to 7.37 using NaOH), mounted flat on filter paper and fixed for 30 min in 10% sucrose in PBS at room temperature (RT). Flat mounts were washed in PBS (3 × 1 h) at RT, incubated with secondary antibodies for 1 day at 4°C, washed and thawed three times, and blocked with 10% normal donkey serum in PBS for 2 hr before incubation with primary antibodies for 5 days at 4°C. Flat mounts were washed in PBS (3 × 1 h) at RT, incubated with secondary antibodies for 1 day at 4°C, and washed in PBS (3 × 1 h) at RT. The following primary antibodies were used in this study: rabbit anti-Mela
dopsin (1:5,000, Abcam, RRID:AB_444842) and mouse anti-RFP (1:1,000, Abcam, RRID:AB_945213). Secondary antibodies were Alexa 488 and Alexa 568 conjugates (1:1,000, Invitrogen, RRID:AB_2556546 and RRID:AB_2534013).

**Immunohistochemistry**

Flat-mount preparations were cryoprotected (1 hr 10% sucrose in PBS at room temperature [RT]), 1 hr 20% sucrose in PBS at RT, and overnight 30% sucrose in PBS at 4°C. Frozen and thawed three times, and blocked with 10% normal donkey serum in PBS for 2 hr before incubation with primary antibodies for 5 days at 4°C. Flat mounts were washed in PBS (3 × 1 h) at RT, incubated with secondary antibodies for 1 day at 4°C, and washed in PBS (3 × 1 h) at RT. The following primary antibodies were used in this study: rabbit anti-Melan
dopsin (1:5,000, Abcam, RRID:AB_444842) and mouse anti-RFP (1:1,000, Abcam, RRID:AB_945213). Secondary antibodies were Alexa 488 and Alexa 568 conjugates (1:1,000, Invitrogen, RRID:AB_2556546 and RRID:AB_2534013).

**Confocal Imaging**

Image stacks of fixed tissue were acquired through a 20 × 0.85 numerical aperture (NA) oil immersion objective (Olympus) on an upright laser scanning confocal microscope (FV1000, Olympus) and processed with Fiji (Schindelin et al., 2012).

**Electrophysiology**

Cell-attached and whole-cell patch-clamp recordings were obtained primarily from the ventral halves of dark-adapted retinas, flat-mounted on transparent membrane discs (13 mm Whatman Anodisc) superfused (~7 mL/min) with warm (30°C–33°C) mACSFNaHCO3. Fluorescent RGCs were targeted under two-photon guidance (excitation wavelength: 940 nm) in Grik4-Cre:Ai9 retinas. The intracellular solution for current-clamp recordings contained (in mM) 125 K-glucuonate, 10 NaCl, 1 MgCl2, 10 EGTA, 5 HEPES, 5 ATP-Na2, and 0.1 GTP-Na (pH adjusted to 7.2 with KOH). The intracellular solution for voltage-clamp recordings contained (in mM) 120 Cs-glucuonate, 1 CaCl2, 1 MgCl2, 10 Na-HEPES, 11 EGTA, 10 TEA-Cl, 2 Qx314, ATP-Na2, and 0.1 GTP-Na (pH adjusted to 7.2 with CsOH). Patch pipettes had resistances of 3–6 MΩ (borosilicate glass). Signals were amplified with a Multiclamp 700B amplifier (Molecular Devices), filtered at 3 kHz (8-pole Bessel low-pass), and sampled at 10 kHz (Digitata 1500, Molecular Devices). For voltage-clamp recordings, series resistance (<15 MΩ) was compensated electronically by ~80%. All reported voltages were corrected for a liquid junction potential of ~15 mV. Excitatory postsynaptic currents (EPSCs) were measured near the reversal potential of inhibitory conductances (~70 mV) and inhibitory postsynaptic currents (IPSCs) were measured near the reversal potential of excitatory conductances (0 mV). In some experiments (Figure 5), the following pharmacological agents were individually added to mACSFNaHCO3 and bath-applied: gabazine (5 μM, Tocris), D-AP5 (30 μM, Tocris), NBQX (10 μM, Tocris), or TTX (1 μM, Sigma). To measure intrinsic photosensitivity (Figure S2), the following agents were applied simultaneously: L-AP4 (100 μM, Tocris), NBQX (10 μM, Tocris), gabazine (5 μM, Tocris), strychnine (2 μM, Sigma-Aldrich), and D-AP5 (30 μM, Tocris).

**Retrograde Labeling**

Mice were anesthetized with ketamine (1 mg/10 g) and Cholec
toxin B conjugated to Alexa 488 (1 g/L; 150 nL, Thermo Fisher Scientific) or AA
g-Syn-FLEX-GCaMP6f (University of Pennsylvania Vector Core) were injected into different brain regions using a Nanoject II (Drummond). Approximately 72 hr after injection, retinas were prepared for physiological recordings as described above, while brains were removed and placed in 4% paraformalde
hyde overnight. The following day, brains were sectioned, stained with Neuro
trace (Thermo Fisher Scientific), and mounted for confocal imaging. Brain slices were imaged on a confocal microscope to verify injection accuracy.

**Visual Stimulation**

All visual stimuli were written using the Cogent Graphics toolbox (John Romaya, Laboratory of Neurobiology at the Wellcome Department of Imaging Neuroscience, University College London) in MATLAB (MathWorks). A UV E4500 MKII PLUS II projector illuminated by a 385-nm light emitting diode (LED) (EKB Technologies) was used for stimulus presentation, except for Figures 6 and S2 in which the green (peak: 532 nm) and blue (peak: 452 nm) LEDs, respectively, of a DLP LightCrafter 4500 (Texas Instruments) were used. Stimuli were focused onto the photo receptors via a substage condenser of
an upright two-photon microscope (Scientifica). All stimuli were centered on the soma of the recorded cell. A background luminance of 1,500 rhodopsin isomerizations/rod/s (R*) was used for all visual stimuli unless otherwise noted. In Figures 1 and 4, a circular region (Figure 1 diameter: 300 μm, Figure 4 diameter: 1,200 μm) was square-wave modulated at a frequency of 0.25 Hz (Michelson contrast: 100%). In Figures 2 and S5, contrast sensitivity was tested by 1-s luminance steps within a circular region (Figure 2 diameter: 300 μm, Figure S5 diameter: 1,200 μm) every 3 s. To test spatiotemporal filtering (Figure S4), the receptive field center was stimulated by a 300-μm circle, and the surround was stimulated by a series of annuli of equal area to the center circle, but of varying inner and outer diameters. The intensities of these regions were chosen randomly from a normal distribution (root-mean-square [RMS] contrast: 40%) every 16.7 ms (refresh rate: 60 Hz) for 18–24 min. In another series of experiments, only the center was stimulated. In Figures 2, S3, and S5, textures of varying scale (2, 12, 20, 32, 48, 60 μm) were generated by convolving random binary maps with 2D Gaussian filters. Textures were masked within a 300-μm (Figures 2 and S3) or 1,200-μm (Figure S5) circle. The mean luminance of the textured region was equal to that of the background. In Figure 3, circles of varying diameters were presented in a pseudo random sequence and square-wave modulated at 0.25 Hz (Michelson contrast: 100%). In Figure S2, a full-field light pulse from a black background was presented for 20 s. The intensity of these light pulses (peak: 452 nm) ranged from ~1,000 R* (~2 × 10^11 photons/cm^2/s) to ~7 R* (~2 × 10^10 photons/cm^2/s), the equivalent of twilight and bright daylight, respectively (Lindell et al., 2014; Johnson et al., 2013). The display was divided into a 3 × 3 grid of squares (300-μm side lengths), with the center square centered on the recorded RGC soma. With each stimulus presentation, 1 of the 9 squares increased in luminance (+100% contrast) for 2 s. The visual stimulus in Figure 6 was analogous to the visual stimulus in Figure 3, but green light (peak: 532 nm) was used instead of UV (peak: 385 nm). In Figure S7, annuli of a constant width (60 μm) and varying inner and outer diameters were presented in a pseudorandom sequences as +100% contrast steps from background for 2 s.

Electrophysiology Analysis

Average spike rates or baseline-subtracted average conductances were measured during 100- to 200-ms time windows. The percentages of responses that were sustained were calculated by dividing the response 1.5 s after stimulus onset by the peak response. Excitation preference (Figure S1) was calculated as the difference between the peak excitatory and inhibitory conductances evoked by presentation of a 300-μm circle divided by the sum of these conductances. A value of 1 indicates pure excitation, and a value of −1 indicates pure inhibition. Following white noise stimulation (Figure S4), spike-triggered stimulus averages (STAs) were calculated by reverse correlation and used to map spatiotemporal receptive fields (Johnson et al., 2014). All analyses were performed using custom scripts written in MATLAB.

Morphological Analysis

To calculate dendritic field diameters, maximum intensity projections of Alexa 488 filled cells were made in Fiji. Using custom software written in MATLAB, a polygon was drawn around the edges of a cell’s dendrites, and the dendritic field diameter was calculated as the longest distance across the polygon. The area of the polygon was also calculated, and the equivalent diameter was calculated as the diameter of a circle with them same area as the polygon (2π*A/2). Neurite length was calculated by tracing 2 stack images of Alexa 488 filled cells using the Simple Neurite Tracer plugin (Longair et al., 2011) in Fiji. Dendritic branching patterns were analyzed using the Sholl Analysis plugin (Ferreira et al., 2014) in Fiji. To calculate inner plexiform layer (IPL) depth, IPL borders were detected from transmitted light images. Z stack images of filled cells were registered by their relative position within the inner plexiform layer (0%–100% from its border with inner nuclear layer to its border with the ganglion cell layer). Fluorescence intensity at each depth was measured using custom scripts written in MATLAB.

Statistics

Paired t tests and ANOVA were used to assess the statistical significance of observed differences. Unless otherwise noted, population data are reported as mean ± SEM, and n represents the numbers of cells analyzed.

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures and can be found with this article online at https://doi.org/10.1101/182758.

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AUTHOR CONTRIBUTIONS

The study was conceived and designed by K.P.J. and D.K.; data were acquired by K.P.J. and L.Z.; data were analyzed and interpreted by K.P.J. and D.K.; and the manuscript was written by K.P.J. and D.K.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

A Pixel-Encoder Retinal Ganglion Cell with Spatially Offset Excitatory and Inhibitory Receptive Fields

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Figure S1. Current-voltage relationship of synaptic inputs to PixON-RGCs elicited by small stimuli (related to Figure 1)

(A) Currents evoked by presentation of a 300 µm circle (2 s ON, 2 s OFF, background: 1500 R*/rod/s) at different holding potentials. (B) Summary peak current versus voltage curve (n = 5). Line is fit of data by a second order polynomial. (C) Histogram showing excitation preferences of all cells (gray) in Grik4-Cre: Ai9 for which both excitatory and inhibitory currents were recorded during presentation of a 300-µm circle. The histogram of PixON-RGCs’ excitation preference is overlaid in red.
Figure S2. PiXON-RGCs do not stain for melanopsin and are not intrinsically photosensitive (related to Figure 1)

(A) Melanopsin staining in a Grik4-Cre:Ai9 retina. (B) Representative EPSCs of a PiXON-RGC to full-field light stimulation in control conditions (red) and under blockade of synaptic inputs to PiXON-RGCs (light red). Dashed line shows baseline in the absence of stimulus. (C) Summary of excitatory conductance evoked by full-field light stimulation in the presence of synaptic blockers (n = 8). Stimuli were presented at ~1000 R* (~2 * 10^{11} photons / cm^2 / s) and ~10^7 R* (~2 * 10^{15} photons / cm^2 / s).
Figure S3. Responses of direction-selective ganglion cells in Grik4-Cre:Ai9 to texture stimuli (related to Figure 2)

(A) Representative texture stimuli of different spatial scales masked in a 300 µm circle. Textures appeared from a grey background (onset) and were then translated 33 µm in either the dorsal, ventral, nasal, or temporal direction (motion). Textured regions had the same mean luminance as the background. (B) Representative spike responses of an ON-OFF DSGC (ooDSGC) to presentation of textures of the spatial scales shown in (A). (C, D) Summary of ON-OFF DSGC spike responses to the onset (C, n = 4) and motion (D, n = 4) of texture stimuli of different spatial scales. ON-OFF DSGCs exhibited directional preference during texture motion. Firing rates shown are for motion in the preferred direction. (E) Representative spike responses of an ON DSGC (oDSGC) to presentation of textures of the spatial scales shown in (A). (F, G) Summary of ON DSGC spike responses to the onset (F, n = 4) and motion (G, n = 4) of texture stimuli of different spatial scales. ON DSGC exhibited directional preference during texture motion. Firing rates shown are for motion in preferred direction.
Figure S4. Spatiotemporal receptive fields of PixON-RGCs (related to Figures 1, 2 and 3)

(A) Representative spatiotemporal receptive field map of a PixON-RGC measured from its spike response to a circular white noise stimulus with rings of constant area. (B) Spike-triggered average (STA) responses of the center (300 µm circle) and surrounding rings (n = 3). An additional 11 cells were recorded in which the white noise stimulus was only presented within the center 300 µm circle. Lines (shaded areas) indicate the mean (± SEM) temporal kernels of the center (black) and surround (magenta) regions. (C) Summary of STA biphasic indices (STA peak/|STA trough|) (n = 14 center, n = 3 surround). (D) Summary of times to the STA peak (n = 14 center, n = 3 surround).
Figure S5. Contrast response function and texture responses of inhibition to PiX_{ON}-RGCs (related to Figure 2)

(A) Representative IPSC responses to contrast steps presented in a 1200-µm circle centered on the soma of the recorded cell. Dashed line shows baseline in the absence of stimulus. (B) Contrast response function of normalized inhibitory conductance (B, n = 4). (C) Texture stimulus as described in Figures 3 and S3, but here textures are masked in a 1200-µm circle. (D) Representative inhibitory responses of a PiX_{ON}-RGC to the presentation of textures of the spatial scales shown in (C). (E, F) Summary of Inhibitory responses to the onset (E, n = 3) and motion (F, n = 3) of texture stimuli of different spatial scales.
Figure S6. Inhibitory receptive fields of ONα-RGCs (related to Figure 5)

(A) Annuli of constant outer diameter (1200 µm) and varying inner diameter were presented as 100% contrast steps (2 s ON) from a grey background (1500 R*/rod/s). (B) Representative IPSCs in an ONα-RGC in response to annuli shown in (A). (C) Normalized inhibitory conductance versus annulus inner diameter (n = 3). (D) The display was divided into a 3 x 3 grid of 300 µm x 300 µm squares and centered on the soma of the recorded cell. Between stimulus presentations all squares had the same luminance (1500 R*/rod/s). During stimulus presentation, one of the nine squares increased in luminance (100% contrast, 2 s ON). (E) Representative IPSCs in an ONα-RGC in response to appearance of squares in the positions shown in (D). (F) Map of the normalized inhibitory conductance evoked by the appearance of a square at each position (n = 2).
Figure S7. Spatial separation of excitatory and inhibitory receptive fields of PixON-RGCs (related to Figure 5)

(A) Annuli of constant width (60 µm) and varying inner diameters were square-wave modulated (2 s ON, 2 s OFF, background: 1500 R*/rod/s). (B, D) Representative excitatory (B, red) and inhibitory (D, cyan) currents evoked by annuli shown in (A). Dashed lines show baselines in the absence of stimulus. (C) Excitatory conductance (inward only) versus annulus inner diameter (n = 4). (E) Inhibitory conductance (outward only) versus annulus inner diameter (n = 8).