Probing Neurochemical Structure and Function of Retinal ON Bipolar Cells With a Transgenic Mouse

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ABSTRACT

Retinal ON bipolar cells make up about 70% of all bipolar cells. Glutamate hyperpolarizes these cells by binding to the metabotropic glutamate receptor mGluR6, activating the G-protein G<sub>α1</sub>, and closing an unidentified cation channel. To facilitate investigation of ON bipolar cells, we here report on the production of a transgenic mouse (Grm6-GFP) in which enhanced green fluorescent protein (EGFP), under control of mGluR6 promoter, was expressed in all and only ON bipolar cells. We used the mouse to determine density of ON bipolar cells, which in central retina was 29,600 cells/mm<sup>2</sup>. We further sorted the fluorescent cells and created a pure ON bipolar cDNA library that was negative for photoreceptor unique genes. With this library, we determined expression of 27 genes of interest. We obtained positive transcripts for G<sub>α</sub> interactors: regulators of G-protein signaling (RGS), Ret-RGS1 (a variant of RGS20), RGS16, RGS7, purkinje cell protein 2 (PCP2, also called L7 or GPSM4), synembryn (RIC-8), LGN (GPSM2), RAP1GAP, and Cβ5; cGMP modulators: guanylyl cyclase (GC) 1 and 2, phosphodiesterase (PDE) 1C, and PDE9A; and channels: inwardly rectifying potassium channel Kir2.4, transient receptor potential TRPC2, and sperm-specific cation channels CatSper 2–4. The following transcripts were not found in our library: AGS3 (GPSM1), RGS10, RGS19 (GAIP), calbindin, GC1 and 2, PDE5, PDE2A, amiloride-sensitive sodium channel ACCN4, and CatSper1. We then localized Kir2.4 to several cell types and showed that, in ON bipolar cells, the channel concentrates in their dendritic tips. The channels and modulators found in ON bipolar cells likely shape their light response. Additional uses of the Grm6-GFP mouse are also discussed. J. Comp. Neurol. 510:484 – 496, 2008.

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Indexing terms: GluR6; CatSper; Kir2.4; RGS; G-protein; guanylyl cyclase; phosphodiesterase; EGFP

About 70% of all retinal bipolar cells are ON bipolar cells; these include approximately five types of ON cone bipolar cells that mediate daylight vision as well as rod bipolar cells that mediate night vision (Sterling and Demb, 2004). The high proportion of ON bipolar cells and their key position as linkers between photoreceptors and ganglion cells justifies the significant efforts that have been made to understand their physiological responses and their underlying molecular mechanisms. ON bipolar cells receive their input from photoreceptors, which transmit their signal to both ON and OFF bipolar cells by modulating glutamate release (Ayoub et al., 1989). Glutamate has two modes of action: it depolarizes OFF bipolar cells via conventional ionotropic receptors and hyperpolarizes ON bipolar cells via a metabotropic glutamate receptor, mGluR6 (Nomura et al., 1994; Vardi et al., 1998, 2000, 2002; Haverkamp et al., 2000). Under scotopic conditions when photoreceptors are continuously releasing glutamate, bound mGluR6 activates the heterotrimeric G-protein G<sub>α1</sub> (Nawy, 1999; Dhingra et al., 2000, 2002),...
which eventually closes an unidentified nonspecific cation channel (Nawy and Jahr, 1990; de la Villa et al., 1995). This maintains the cell in a hyperpolarized state. A light increment reduces glutamate release, unbinds mGluR6, and inactivates Go\textsubscript{m}, and the channels are permitted to open. This depolarizes the ON bipolar cells, whose summed responses give rise to the b-wave of the noninvasive electroretinogram (ERG; Robson and Frishman, 1999).

Progress in solving the transduction cascade that mediates the light response in ON bipolar cells has been slow because, unlike photoreceptors, which can be purified and easily studied biochemically, the ON bipolar cells cannot be purified. Moreover, because they are imbedded in an inner layer of the retina, recording from them is difficult. Thus, we still cannot identify the effector for Go\textsubscript{m}, what forms the transduction channel, what modulates the response, and what is the function of cGMP. Such information would be difficult or impossible to achieve by physiological recordings if the key proteins were novel or if the known drugs were useless. To alleviate some of these difficulties in studying ON bipolar cells, especially their mGluR6 transduction cascade, we produced several mouse lines in which enhanced green fluorescent protein (EGFP) expression was driven under the control of mGluR6 promoter. Here we use this transgenic mouse to create an ON bipolar cDNA library and to examine the expression profile of these cells.

**MATERIALS AND METHODS**

**Generation of the transgenic mouse**

mGluR6 promoter sequence was extracted by digesting pN2N (a generous gift of Y. Nakajima and S. Nakanishi) with NotI, blunt filling with Klenow fragment polymerase, and then digesting with Xhol. pEGFP1 (Clontech, Cambridge, United Kingdom) was digested with SmaI and Xhol and ligated with mGluR6 promoter. Background was removed by digesting these plasmids with SacII. After transformation, clone sequence was confirmed by digestion and sequencing. This was then digested with Xhol and SphI and run on a gel. The 10.5-kb band consisting of mGluR6 promoter fused to EGFP coding sequence was extracted by digesting these plasmids with SacII. After transformation, clone sequence was confirmed by digestion and sequencing. This was then digested with Xhol and SphI and run on a gel. The 10.5-kb band consisting of mGluR6 promoter fused to EGFP coding sequence was eluted for microinjection into fertilized eggs (performed by the University of Pennsylvania Transgenic and Chimeric Mouse facility). The founder mice (in B6SJLF1/J background) were screened by genomic PCR for the presence of mGluR6 promoter. Here we use this transgenic mouse to create an ON bipolar cDNA library and to examine the expression profile of these cells.

**Cell count**

Mice were deeply anesthetized by intraperitoneal injection (85 µg/g ketamine plus 13 µg/g xylazine), and the eye orientation was noted by an incision at the nasal pole. Eyes were enucleated, and the mice were then killed by anesthetic overdose (3× initial doses). Animals were treated in compliance with federal regulations and University of Pennsylvania policy. The eye was fixed and the retina was flat mounted, ganglion cell up. A filter paper was placed around the retina to provide support for the coverslip and reduce compression. The inner nuclear layer was scanned with a confocal microscope with 1 µm steps in the Z-axis. Two retinas were scanned under a ×100 oil immersion lens (NA 1.4; Leica microscope), where the scanned field was 100 µm × 100 µm, and two retinas were scanned under ×60 oil (NA 1.42; Olympus), where the scanned field was 140 µm × 140 µm. Cells were identified through a stack of focal planes taken at 1 µm intervals and counted manually using Openlab software (Improvision, Lexington, MA). For the first two retinas, three counts were made from each field, and the counts were averaged; for the other two, only one count per field was made, because the variance between mice was greater than between counts.

**Immunocytochemistry**

After anesthesia, the eye was incised at the ora serrata and fixed by immersion in phosphate buffer (pH 7.4) containing 4% (or 2%) paraformaldehyde for 1 hour (or 10 minutes). It was then rinsed in buffer, soaked overnight in 30% buffered sucrose, and embedded in a mixture of two parts 20% sucrose in phosphate buffer and one part Tissue Freezing Medium. Retina was cryosectioned radially at 10–15 µm thickness and immunostained according to a standard protocol. Sections were soaked in diluent containing 10% normal goat serum, 5% sucrose, and 0.3% Triton X-100 in phosphate buffer. They were then incubated in primary antibodies overnight at 4°C, washed and incubated (3 hours) in anti-rabbit P(ab\textsubscript{g}) fragment conjugated to a fluorescent marker, and then rinsed and mounted in Vectashield.

HEK 293T cells were cultured in minimal essential medium supplemented with Penstrept (Invitrogen, Carlsbad, CA) and 10% heat-inactivated fetal bovine serum at 37°C in a 5% CO\textsubscript{2} incubator. Cells were transiently transfected with Kir2.4 (in pcDNA3.1) using Fugene transfection reagent (Invitrogen) and harvested 48 hours later. Retinas and HEK cells were imaged with either a Leica or an Olympus confocal microscope. All images were taken with the default pinhole (of one airy disk) calculated by the confocal microscopes. These give the following approximate z-resolutions: for ×100 NA 1.4 (Leica) and ×60 NA 1.42 (Olympus) objectives 500–600 nm and for ×40 NA 1.25 (Leica) and ×40 NA 1.3 (Olympus) 650–750 nm. For display, images were contrast adjusted with Adobe Photoshop 7.0 (Adobe Systems, San Jose, CA).

**Western blotting**

Fresh or frozen mouse retinas or HEK cells were homogenized in 0.2–1 ml ice-cold homogenization buffer (5 ml per g wet weight; in mM: 320 sucrose, 5 Tris-HCl, and 2 EDTA, 2.5 β-mercaptoethanol, pH adjusted to 7.4 at room temperature) containing protease inhibitor cocktails (P8340; Sigma Aldrich, St. Louis, MO) and centrifuged at 6,000g for 10 minutes, and the supernatant was collected. Protein assay was carried out by using BCA protein reagent (Bio-Rad, Hercules, CA). Twenty micrograms of protein was mixed with NuPAGE LDS sample buffer (Invitrogen), incubated for 10 minutes at 95°C, then loaded onto 10% Tris-glycine gel (Bio-Rad). Electrophoresis was performed with a Mini-Protein II electrophoresis cell (Bio-
Rad) under reduced denaturing conditions. The resolved proteins were transferred to a nitrocellulose membrane (Bio-Rad) in transfer buffer (192 mM glycine, 25 mM Tris-HCl, pH 8.3, and 20% methanol) for 40–50 minutes at 80 V using a Bio-Rad Trans-Blot semi-dry transfer cell. The membrane was blocked with 7% skim milk in TBST (10 mM Tris-HCl, 150 mM NaCl, 0.1% Tween 20, pH 7.4) for 1 hour and incubated with anti-Kir2.4 (1:5,000) overnight at 4°C in TBST-milk. After extensive washes in TBST, the membrane was incubated with the secondary antibody conjugated to peroxidase (Jackson Immunoresearch, West Grove, PA; 1:3,000) in TBST-milk for 2 hours at room temperature. After extensive washes in TBST, bound antibody was detected with SuperSignal West Femto Maximum Sensitivity Substrate (Pierce, Rockford, IL).

Antibodies

Three antibodies were used for marking cells. Rod bipolar cells were stained with a mouse monoclonal antibody against PKCs purified from bovine brain (clone MC5; Sigma-Aldrich; catalog No. P 4334) and shown to recognize specifically a 79-kDa protein, which can be blocked with peptides 296–317. They were also stained with a polyclonal antibody that was raised in rabbit against a synthetic peptide corresponding to rat PKCs amino acids 659–672 (Sigma-Aldrich; catalog No. P5704) and gives a staining pattern identical to that of the monoclonal. ON bipolar cells were marked with a polyclonal antibody that was raised in rabbit against peptide aa 47–59 of mouse G-α13 and whose specificity in retina was established by correlating transcript (RT-PCR) and immunostaining (Huang et al., 1999, 2003). This antibody was a gift of Robert Margolskee, Mount Sinai School of Medicine. Other antibodies were used for localizing proteins of interest. The only antibody that could be shown to be specific in retina was a polyclonal antibody against Kir2.4, which was raised in rabbit against the less well conserved C-terminus amino acids 395–434 of rat Kir2.4 and used at 1:500 dilution for immunocytochemistry. Its specificity in rat has been confirmed by transfecting COS cells and by correlating results with in situ hybridization (Liu et al., 2001; Pruss et al., 2003); its specificity in mouse retina is described in Results. Several other antibodies used in this study either did not reveal staining or their staining specificity could not be established in retina. We chose to report on their performance here in order to provide preliminary information that may benefit future studies. A polyclonal antibody against CatSper1 raised in rabbit against the first 150 amino acids of the amino terminus (a gift of Dejian Ren, University of Pennsylvania; Ren et al., 2001) gave similar Western blots and immunostaining in wild-type and CatSper1-null retina. Two polyclonal antibodies against CatSper2, one against the N-terminus and the other against the C-terminus (gifts from Timothy Quill, University of Texas; Quill et al., 2001), gave similar Western blots and immunostaining in wild-type and CatSper2-null mouse. A polyclonal antibody against TRPC2 raised in guinea pig against an N-terminal peptide (CSSDASGAGPGGPRNNE; gift from Richard Axel, Columbia University; Leybold et al., 2002) did not detect any band in Western blotting and did not stain retina. Monoclonal antibody to RGS16 raised in mouse against a partial recombinant human protein aa 90–180 (Abnova, Taipei City, Taiwan; catalog No. H00006004-A01) revealed several bands by Western blotting, but the major bands were not at the correct molecular weight. Immunocytochemistry for RGS16 revealed strong staining in the outer plexiform layer. Similarly, a polyclonal antibody against RIC-8 (synembrin) raised in goat against a synthetic protein corresponding to a human sequence (Abcam, Cambridge, MA; catalog No. ab 24383) gave several bands at unexpected molecular weights, and immunostaining revealed strongest staining in the photoreceptor outer segment. Polyclonal antibody to PDE9A raised in rabbit against a synthetic peptide corresponding to human C-terminus peptide CDISNEVRPMEVAEPWVDCLLEE (Genetex, San Antonio, TX; catalog No. GTX14625) recognized PDE9A in transfected HEK cells but gave five major bands in Western blotting. Immunostaining with this antibody gave strong punctate staining in the outer plexiform layer.

Cell isolation and FACS

Retinas were isolated and incubated in Ca²⁺- and Mg²⁺-free HBSS with 20 mM HEPES, 3 mg/ml BSA, 3 mg/ml glucose, 1.5 mM MgSO₄, and 0.0025% trypsin (Sigma-Aldrich) at 37°C for 25 minutes, with gentle shaking. Enzymatic reaction was stopped by adding 0.25 mg/ml trypsin inhibitor and 40 μg/ml DNaseI. Tissue was triturated with a 1-ml pipette tip, followed by a 100-μl tip until the suspension was homogenous. Cell viability and quality of dissociation were checked with EthD-1 (1 μM; Invitrogen). The dissociated cells were sorted by fluorescent-assisted cell sorting (FACS) using FACSVantage (BD Biosciences, San Jose, CA). Briefly, about 100,000 dissociated cells were resuspended in Ca²⁺- and Mg²⁺-free HBSS. Debris and aggregates were gated out by forward and side scatter. The cells were then sorted based on size and fluorescence intensity. Only sizes between 6 and 8 μm and intensities higher than 100 units were selected. For RNA extraction, the sorted cells were collected in the lysis buffer. For viewing the cells, they were collected in phosphate buffer, spun down at 2,500 rpm, and settled on a glass slide.

Library production

Total RNA from isolated cells was extracted with Nucleospin RNA II kit and reverse transcribed with the aid of the “SMART cDNA Library Construction Kit” (both from Clontech, Logan, UT). About 1 μg of RNA was primed to CDSIII primer containing the SfiI restriction site on the 5’ end and a stretch of dT on the 3’ end. The newly transcribed cDNA was extended with an oligonucleotide containing the SfiI sequence (SMART IV primer). The cDNA was then amplified with the CDSIII and a 5` primer, and the double-stranded cDNA was fractionated. The first four fractions containing cDNA were pooled, digested with SfiI, and ligated with the SfiI-digested pEX1. The ligation mixture was transformed in DH10B cells, and the number of independent clones was calculated to be >10⁶. The library was amplified, and aliquots were stored at −80°C.

PCR

PCR on library DNA was carried out with gene specific or degenerate primers on a programmable thermocycler (Perkin Elmer, Branchburg, NJ). The PCR was performed for 35 cycles with the following conditions: denaturation at 94°C for 30 seconds, annealing at 52–62°C (depending on the primers used) for 30 seconds, followed by extension at 72°C for 30 seconds. All primer sets were applied to the
ON bipolar library and to reverse transcribed retinal cDNA. For most transcripts, amplification from retina was used as positive control, and, unless otherwise noted, retina was indeed positive. For each test transcript, we designed three PCR primers: one lower primer and two upper primers (Table 1). At least one product with a predicted size from each primer set was sequenced to confirm that the product was not only the correct size but also the correct sequence. When the product with a predicted size from each primer set was sequenced to confirm that the product was not only the correct size but also the correct sequence. When the correct product size was not obtained, we sequenced the corrected size but also the correct sequence. When the product with a predicted size from each primer set was sequenced to confirm that the product was not only the correct size but also the correct sequence. When the correct product size was not obtained, we sequenced the following. Each sequence of an upper primer is preceded by the number of the first nucleotide at the 5' end, whereas each lower primer is preceded by the number of the last nucleotide at the 3' end (reverse primer). Lower primers were used with either of the two upper primers. The expected PCR product size (in bp) for each primer pair is given in parentheses. NT, N-terminus; CNGC, cyclic nucleotide-gated channel; GRK, G-protein receptor kinase; R, results of amplification from ON bipolar library; –, no amplification; ?, results were ambiguous; for CatSper2, the obtained PCR products were shorter than expected; for CatSper3, first primer indicates “–” because the full-length transcript could not be amplified from the library, but was amplified from retinal cDNA; proteins that are positive in ON bipolar cells are in boldface.

Each sequence of an upper primer is preceded by the number of the first nucleotide at the 5' end, whereas each lower primer is preceded by the number of the last nucleotide at the 3' end (reverse primer). Lower primers were used with either of the two upper primers. The expected PCR product size (in bp) for each primer pair is given in parentheses. NT, N-terminus; CNGC, cyclic nucleotide-gated channel; GRK, G-protein receptor kinase; R, results of amplification from ON bipolar library; –, no amplification; +, positive amplification; +/-, mostly negative; ?, results were ambiguous; for CatSper2, the obtained PCR products were shorter than expected; for CatSper3 and CatSper1, first primer indicates “–” because the full-length transcript could not be amplified from the library, but was amplified from retinal cDNA; proteins that are positive in ON bipolar cells are in boldface.

1For Ret-RGSI N-terminus, the mouse sequence is not available in GenBank; the accession No. given is for an EST (Dhingra et al., 2004).

2Names of isoforms as they appear in the NCBI databank is incorrect; we adapted the names according to Zabel et al. (1998) and Koesling et al. (2004).

3TRPC degenerate primers are based on Liman et al. (1999).

TABLE 1. Primers Used for PCR Amplification and Results

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Each sequence of an upper primer is preceded by the number of the first nucleotide at the 5' end, whereas each lower primer is preceded by the number of the last nucleotide at the 3' end (reverse primer). Lower primers were used with either of the two upper primers. The expected PCR product size (in bp) for each primer pair is given in parentheses.
remaining bands to test whether they represented different splice variants.

Electroretinogram (ERG) recording

The experimental apparatus, methods of light stimulation and quantification, ERG recording, and cone signal isolation have been described in detail previously (Lyubarsky et al., 1999, 2000). Briefly, a mouse was dark adapted overnight. Then, under dim red light, it was deeply anesthetized by intraperitoneal injection of ketamine (20 mg/g), xylazine (8 mg/g), and urethane (800 mg/g) and placed onto a warmed (37°C) platform. The eyes were dilated with 1% Mydriacyl (Alcon, Fort Worth, TX). A platinum recording electrode contacted both corneas, and the reference electrode lay in the animal’s mouth. The animal was then placed inside a light-proof Faraday cage that also served as a Ganzfeld, with appropriate ports and baffles to ensure uniform illumination. The light stimuli were brief flashes (~10 msec) generated by xenon flash lamps and delivered in a multiport, customized Ganzfeld through calibrated filters. Stimulus intensity and spectral composition were controlled with neutral density and bandpass interference filters. Light intensities were calibrated and converted to estimated number of photoisomerizations per photoreceptor (R*) as described previously (Lyubarsky et al., 1999, 2000). ERGs were recorded from both eyes using differential amplifiers with a bandwidth of 0.1 Hz to 1 kHz. The filtered traces were digitized at 5 kHz. Data were analyzed with Clampfit software, and the responses of wild-type and CatSper-null mice or Grm6-GFP mice were compared with a Student’s t-test.

RESULTS

Description of the EGFP-positive lines

Our initial screen revealed six lines of transgenic mice with EGFP expression in the retina. To determine which retinal cell types expressed EGFP, radial cryosections of mature retinas were performed, and the location of the fluorescing cells was examined. In all lines, EGFP was expressed only in cells of the inner nuclear layer that were restricted to the upper tiers. None of the lines showed fluorescence in the ganglion cell layer or the photoreceptor layer.

To verify that the fluorescing cells were only ON bipolar cells, we immunostained the retina with an antibody against G-γ13, the gamma subunit of gustducin. This subunit was shown to be expressed exclusively by ON bipolar cells (Huang et al., 2003). In two lines (lines 3 and 5), EGFP was expressed by all ON bipolar cells and nowhere else in the retina (Fig. 1). Tangential views of retina from line 5 mice illustrate the dense array of ON bipolar somas in the INL with sparse “holes” that probably indicate OFF bipolar somas and possibly Müller cells (Fig. 2A). The axons of these bipolar cells traversed sublamina a without branching or terminating (Fig. 2B). They then terminated in all strata of sublamina b with a dense network of large

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terminals in strata 5 (left region in Fig. 2C) and fine arborizations in strata 3 and 4 (right region in Fig. 2C). Because the fields are slightly slanted, structures on the left are deeper in the retina. For example, in C, the left field shows sublamina 5, where large structures belonging to rod bipolar terminals are seen, whereas the right field shows sublaminas 3 and 4, with smaller terminals of ON cone bipolar cells. D: ON bipolar cell density vs. distance from the optic disk (OD); distance of 0 is defined as the edge of the disk. All points are temporal to the disc, except the X at 100 μm, which is nasal. Different symbols are from different mice. Scale bar = 30 μm.

Fig. 3. Grm6-GFP transgenic mouse (line 1) shows a mosaic expression of EGFP in ON bipolar cells. A: A low-magnification radial section (projection of two confocal planes) of line 1 retina showing that EGFP is restricted to cells whose somas lie at the upper tiers of the INL and whose axons arborize in sublamina b. B: A radial section stained for G-γ13 (red). All EGFP-positive cells are stained for G-γ13 (double arrowheads), indicating that they are all ON bipolar cells. However, many G-γ13-positive cells are EGFP negative (arrows), indicating that not all ON bipolar cells are marked by EGFP in this mouse line. C: A radial section stained for PKC (red). Many PKC-positive cells are EGFP negative (arrows). Double arrowheads point to EGFP-positive rod bipolar cells, and asterisks indicate EGFP-positive ON cone bipolar cells. Scale bars = 30 μm in A; 10 μm in B,C.

Fig. 2. Density of ON bipolar cells varies little across eccentricity. A–C: Tangential view of line 5 retina at the level of the INL (A), sublamina a of the IPL (B), and sublamina b of the IPL (C). Because the fields are slightly slanted, structures on the left are deeper in the retina. For example, in C, the left field shows sublamina 5, where large structures belonging to rod bipolar terminals are seen, whereas

Four retinas were analyzed: near the optic disk, the density was more variable than in periphery; in most retinas, it was highest at about 100–300 μm away from the disk edge, and it stayed relatively constant up to 2 mm away. Taking the highest density from each retina (regardless of eccentricity), the average was about 32,000 cells/mm² (Fig. 2D). The average at 200–600 μm was 29,600 ± 3,700 cells/mm², and the average across eccentricities was 25,250 ± 1,520 cells/mm².

In another line of transgenic mice (line 1), staining for G-γ13 revealed that EGFP was expressed only in ON bipolar cells, but not in all of them (Fig. 3A,B). To determine whether the EGFP-expressing cells were rod bipolar cells, we stained for PKC. We found that certain PKC-stained (rod bipolar) cells did not express EGFP and that certain EGFP-expressing cells were not rod bipolar cells (Fig. 3C). Thus, in this mouse line, the EGFP-expressing cells were mosaics of rod and cone bipolar cells. In the rest of our mouse lines, EGFP expression was weak or not restricted to ON bipolar cells, so these lines and line 3 (which was redundant with line 5) were terminated.

The line 5 transgenic mouse has a normal ERG

We next tested the viability of the EGFP-expressing ON bipolar cells and their usefulness for future studies. Thus we carried out ERG recordings under dark- and light-adapted conditions. Grm6-GFP and wild-type mice were dark adapted overnight, and their ERG responses were tested first with dim stimuli, then with increasing flash...
Next, we tested response recovery with a pair of flashes separated by either 350 or 500 msec (Fig. 4D). Finally, we light adapted the mice and tested their cone ERGs (Fig. 4C). The wild-type and Grm6-GFP mice exhibited similar ERG wave shape and amplitude as well as similar adaptation effects (Fig. 4C,F). Also, the times to peak of the a- and b-waves were similar for the two mouse groups (data not shown).

Expression profiling using an ON bipolar cDNA library

The fluorescence of dissociated ON bipolar cells was sufficiently strong to permit FACS (Fig. 5A,B). The ON bipolar cells were sorted according to fluorescence intensity and size. Only sizes between 6 and 8 μm with intensities higher than 100 units were selected. Combined Nomarsky and fluorescence optics showed that the purity of the collected cells was about 98% (Fig. 5C,D). The sorted cells were lysed, and their mRNA was used to create a mammalian expression library. For practice and comparison, we also made a library from the mouse rd/rd retina (in which all rods and most cones degenerate) and tested that as well.

We tested the purity of the ON bipolar cDNA library by testing whether RNA unique to photoreceptors and not known to be expressed by ON bipolar cells is present in the library. This is a stringent test because photoreceptors are the most likely source of contamination. Consequently, we used primers specific for opsin, RGS9, GRK1, and cGMP-gated channels. OpsiB, which always gave the expected band when amplified from cDNA obtained from whole mouse retina, never gave a band when amplified from the ON bipolar library (at least four amplifications; Fig. 5G). In the rd library, these primers gave a correct band infrequently (two of five trials), probably as a result of the fact that some cones survived. RGS9, GRK1, CNGA1, and CNGA3, cone cGMP-gated channel; CNGA3, cone cGMP-gated channel. E and G come from the same gel. Scale bars = 10 μm.
in the bipolar cell library. This may reflect the 2% contamination we estimated after FACS. In contrast, amplification of mGluR6, Goα1, Purkinje cell protein 2 (PCP2), or Ret-RGS1, proteins known to be expressed by ON bipolar cells, gave bands for all samples (Fig. 5E, shown only for Goα1). Control samples, samples that did not include DNA in the PCR mixture, never gave any bands. These results show that our experimental library is reasonably pure.

We next used our library to determine whether, for 27 proteins of interest, their transcript is expressed by ON bipolar cells. Proteins of interest were divided into three categories: potential interactors of Gα, modulators of cGMP, and channels. To test the expression of each molecule, we used two sets of primers, a common lower primer and two upper primers (Table 1, Fig. 6). For all transcripts we report here to be positive, the expected PCR products were sequenced at least once and found to be correct.

**Gα interactors.** Modulators of interest were based primarily on our yeast two-hybrid screen of a retinal library, in which we used a constitutively active mutant of Goα1 as bait (Dhingra et al., 2004). Among 10 potential Gα interactors, the following transcripts were positive in ON bipolar cells: regulator of G-protein signaling (RGS) Ret-RGS1 (a variant of RGS20), Purkinje cell protein 2 (PCP2, also called L7 or GPSM4), RGS16, synembryon (RIC-8), LGN (GPSM2), and RAP1GAP. Activator of G-protein signaling 3 (AGS3, also called GPSM1), RGS10, RGS19 (GAIP), and calbindin were positive when amplified from retina but negative when amplified from the ON bipolar library. Because new immunocytochemistry data suggest that RGST coupled to Gβ5 is localized to ON bipolar dendritic tips and thus may contribute to the light response (Morgans et al., 2007; Rao et al., 2007), we also tested the expression of these transcripts and found both to be positive.

**cGMP modulators.** In early studies, cGMP was reported to be the second messenger of the mGluR6 transcription cascade (Nawy and Jahr, 1990; Shiells and Falk, 1990), and, in later studies, it appeared to have a modulatory function (Snellman and Nawy, 2004). In either case, it is important to identify enzymes that synthesize or hydrolyze cGMP. Because physiological experiments suggest that the guanylyl cyclase in ON bipolar cells is soluble (Shiells and Falk, 1992), we tested for GC1α1, GC1α2, GC1β1, and GC1β2. We found that only GC1α1 and GC1β1 were expressed by ON bipolar cells. GC1α2 was positive in retina but negative in ON bipolar cells, and GC1β2 was negative in both retina and ON bipolar cells. For hydrolysis, we concentrated on phosphodiesterase PDE1C, PDE2A, PDE5, and PDE9A, all known to hydrolyze cGMP. We found that only PDE1C and PDE9A were expressed in ON bipolar cells. PDE5 was positive in retina and negative in bipolar cells; PDE2A was ambiguous.

**Channels.** Our yeast two-hybrid screen also fished two channels: amiloride-sensitive cation channel 4 (ACCN4) and inwardly rectifying potassium channel J14 (KCNJ14 or Kir2.4). The amiloride-sensitive channel was positive in retina but negative in ON bipolar cells. PCR amplification of the library showed that Kir2.4 transcript was expressed by ON bipolar cells, and its full-length transcript was identical to that cloned in mouse; thus it represents a known potassium channel. Next we tested whether the canonical TRP channels (TRPC) are expressed in ON bipolar cells, because these channels are cation channels and TRPC2 is expressed together with Gβ in the vomeronasal organ (Takami et al., 2001). By using degenerate primers for this subfamily, we amplified a single band that, upon sequencing, was found to be TRPC2. The expression of this transcript was confirmed with specific primers. However, when we tested TRPC2 protein expression using immunocytochemistry and West-
Fig. 7. CatSper-null mice show normal ERG responses to light stimuli. A: Response of CatSper4-null mouse to an intense flash (10⁵ R*/rod) with an interflash interval of 300 msec. Time scale bar = 100 msec.

We produced several transgenic mouse lines and maintained two lines (1 and 5) in which EGFP was expressed under the control of mGluR6 promoter. In line 5, EGFP was expressed in all ON bipolar cells and only in these cells. With this line, we estimated the density of ON bipolar cells near the optic disk on the horizontal meridian to be about 30,000 cells/mm². The density of all bipolar cells in a similar eccentricity can be estimated to be 42,000 cells/mm² (Strettoi and Pignatelli, 2000), thus accounting for only 43% of ON bipolar cells. Rod bipolar cells peak at about 1 mm (15,000 cells/mm²), where the average ON bipolar cell density is 22,800, so at this eccentricity rod bipolars make up 66% of ON bipolar cells. From line 5, we also created an ON bipolar cell library that we then used to determine expression of Kir2.4 in their dendritic tips, but possibly other structures in the outer plexiform layers also express this channel.

### DISCUSSION

For one protein, Kir2.4, we tested expression with a polyclonal antibody against the C-terminus of the rat channel (Liu et al., 2001; Pruss et al., 2003). Immunostaining of Kir2.4-transfected HEK293T cells with this antibody recognized the expressed protein and gave no background in untransfected cells (Fig. 8A,B). Western blots revealed two bands, one at ~52 kDa (corresponding to the monomer; unmodified protein is expected to be 48 kDa) and the second at ~110 kDa (corresponding to the dimer). Western blots of retinal protein revealed the same two bands and no other, showing that the antibody is specific (Fig. 8C). Immunostaining of retinal sections with this protein gave staining throughout the retina. Staining appeared strongest in ganglion cell somas, photoreceptor inner segments, and outer plexiform layer, where labeling appeared punctate (Fig. 8D). When fixation was weak (4% paraformaldehyde for 10 minutes), all somas in the inner nuclear layer were also lightly stained. The ubiquitous expression of Kir2.4 shown here by immunostaining is consistent with in situ hybridization, by which all nuclear layers were labeled (Hughes et al., 2000). To determine whether the punctate labeling in the outer plexiform layer represents ON bipolar dendritic tips, we stained sections of the transgenic mouse and found that indeed most (but not all) of these puncta colocalized with EGFP (Fig. 8E–G). Puncta in the INL localized mainly to the edge of the somas, suggesting that the channel clustered at several membrane regions. Thus ON bipolar cells express Kir2.4 in their dendritic tips, but possibly other structures in the outer plexiform layers also express this channel.
Potential uses for the Grm6-GFP transgenic mouse

The ON bipolar cells in transgenic mouse line 5 are highly fluorescent, and their physiological properties remain unchanged. Thus this mouse line can be used for multiple purposes, several of which have already been exploited. In a previous study, we followed the developmental course of ON bipolar cells in these mice. We found that fluorescent cells could already be detected at postnatal day 3, and they had an unusual mode of axogenesis and dendritogenesis (Morgan et al., 2006). In the current study, we used the transgenic mouse to determine the ON bipolar cell density across different eccentricities and to generate an ON bipolar cDNA library and quest for their gene expression profile.

In the future, this mouse could be used to facilitate several more tasks such as to 1) produce cDNA libraries from different developmental stages and examine differential expression patterns, 2) facilitate targeting of ON bipolar cells in physiological experiments either in a slice preparation or after cell dissociation, 3) facilitate target-
ing of OFF bipolar cells by choosing EGFP-negative cells located in the middle of the inner nuclear layer (a procedure that is especially important since OFF bipolar cell density is relatively low), and 4) determine the connectivity pattern of ON bipolar cells with ganglion cells. In addition, scanning of other organs of the mouse showed EGFP-positive structures throughout the body, including the brain (Vardi, unpublished data), and this will no doubt add to potential uses. The high density of fluorescent cells in line 5 makes it difficult to reconstruct single cells, but, because EGFP expression in line 1 is patchy, it is better suited for this purpose.

**G	extsubscript{o} interactors in ON bipolar cells**

Light hyperpolarizes a photoreceptor but depolarizes the ON bipolar cells. This response reversal is mediated by the metabotropic glutamate receptor mGluR6, which couples to the heterotrimeric G-protein G	extsubscript{o}, whose activation causes closure of nonspecific cation channels. Although the effector for G	extsubscript{o} and the channel remains unknown, it is clear that a major site for modulation is the activation and inactivation of G	extsubscript{o}. By using a constitutively active mutant of Go, as bait in a yeast two-hybrid system, we previously fished out a number of potential interactors for G	extsubscript{o}. We then localized the protein that was hit most often, Ret-RGS1, to bovine ON bipolar cells and found that it can facilitate the rising phase of the light response (Dhingra et al., 2004). Here we probed for the other potential modulators to determine whether they too are expressed by ON bipolar cells.

After Ret-RGS1, the molecule that was hit the most times by the yeast two-hybrid screen was PCP2. PCP2 is known to be expressed by ON bipolar cells (Berrebi et al., 1991), so PCR amplification of its transcript was expected to be positive. Nevertheless, the library provided new information in that RACE PCR showed that retinal PCP2 represents a novel splice variant (Ret-PCP2; deposited in GenBank, accession No. EU164759). Further studies of this protein showed that Ret-PCP2 accelerates the ON bipolar light response (Xu et al., submitted). Among the other “fished” molecules, four more, RGS16, synembryn, LGN, and RAP1GAP, were found to be positive and are transcribed in full length. Nonetheless, the library is not contaminated by amacrine cells.

**Modulators of cGMP concentration**

An important modulator of cGMP concentration is guanylyl cyclase, by stimulating with NO donors and immunostaining for cGMP, gave staining in ON cone bipolar cells but not in rod bipolar cells (Keistinaho et al., 1993; Blute et al., 1998; Gotzes et al., 1998; for review see Sitaramayya, 2002). There are four isoforms of soluble guanylyl cyclase: two GC1a1s and two GC1b.s. GC1b2 appears to be nonfunctional (Koesling et al., 2004), and indeed our PCR from retinal cDNA did not amplify its mRNA. Among the other three isoforms, only GC1a1 and GCb1 were expressed, suggesting that they form the functional heterodimer in ON bipolar cells (Koesling and Fribe, 1999).

Another modulator of cGMP is PDE. The ON bipolar transduction cascade was thought to be similar to phototransduction, where transducin activates PDE6, which then hydrolyzes cGMP and leads to the closing of cGMP-gated channels (Nawy and Jahr, 1990; Shells and Falk, 1990). Consequently, we tested antibodies against photoreceptor PDEs (PDE6A-C) and against PDE	extsubscript{y}. All gave nice staining of photoreceptor outer segments but no staining in bipolar cells (Dhingra et al., 2004; unpublished data). We then examined other PDEs that hydrolyze cGMP and found two to be positive: PDE1C and PDE9A. The positive transcript for PDE1C is consistent with immunolocalization of this isoform to cells located high in the inner nuclear layer (Santone et al., 2006). PDE1C is activated by Ca\textsuperscript{2+}/calmodulin and hydrolyzes both cGMP and cAMP (Bender and Beavo, 2006). This being the case, it can be involved in multiple signaling cascades, such as regulation of GABA response by dopamine via cAMP (Feigenspan and Bormann, 1994). Unlike PDE1C1, PDE9A hydrolyzes only cGMP. This protein is highly expressed in the brain, especially in the olfactory bulb and Purkinje cells (Andreeva et al., 2001; van Staveren et al., 2002; van Staveren and Markerink-van Ittersum, 2005). Among the 20 known PDE isoforms, PDE9A has the highest affinity for cGMP (Fishier et al., 1998; Soderling et al., 1998). Thus cGMP levels in ON bipolar cells might be low, a situation that could explain why cGMP in rod bipolar cells remained undetected after NO stimulation of the cyclase.

**Channels**

Very little is known about the molecular identity and physiological properties of the transduction cation channel in ON bipolar cells. We do know that this channel is nonselective for cations and is thought to permeate calcium. Moreover, the channel is modulated by cGMP, but it does not contain a cGMP-binding domain (Nawy, 1999; Snellman and Nawy, 2004). Guided by this knowledge, we examined the expression of members of the CatSper family that permeate calcium and other cations, are modulated by cyclic nucleotides, and do not have a cyclic nucleotide-binding domain (Ren et al., 2001; Carlson et al., 2003). Four isoforms of this channel are known, and here we found that at least three (CatSper2–4) are transcribed by ON bipolar cells (Qi et al., 2007; Jin et al., 2007). CatSper2 transcript skips exons 2–5 and thus cannot be translated correctly. However, both CatSper3 and -4 are positive and are transcribed in full length. Nonetheless, light responses recorded by ERG and whole-cell mode from CatSper3- and CatSper4-null mice showed that responses were normal, suggesting that, if these channels do contribute to the light response, the contribution is minor.

We also examined Kir2.4 because it was fished out by our yeast two-hybrid screen. This inwardly rectifying po-
tassium channel (Topert et al., 1998, 2000) is expressed by all retinal cell types, including ON bipolar cells, consistent with in situ hybridization results (Hughes et al., 2000). Interestingly, in ON bipolar cells, Kir2.4 is expressed in dendritic tips more strongly than in somas. Somatic expression is limited to a few clusters in three or four regions of the membrane. Because Kir2.4 is a potassium channel and interacts with Goδ1 (Sulaiman, unpublished data), it probably contributes to regulation of the resting potential, possibly regulated by Gα.

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LITERATURE CITED


