

GPR179 Is Required for Depolarizing Bipolar Cell Function and Is Mutated in Autosomal-Recessive Complete Congenital Stationary Night Blindness

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Complete congenital stationary night blindness (cCSNB) is a clinically and genetically heterogeneous group of retinal disorders characterized by nonprogressive impairment of night vision, absence of the electroretinogram (ERG) b-wave, and variable degrees of involvement of other visual functions. We report here that mutations in *GPR179*, encoding an orphan G protein receptor, underlie a form of autosomal-recessive cCSNB. The *Gpr179^{nob5/nob5}* mouse model was initially discovered by the absence of the ERG b-wave, a component that reflects depolarizing bipolar cell (DBC) function. We performed genetic mapping, followed by next-generation sequencing of the critical region and detected a large transposon-like DNA insertion in *Gpr179*. The involvement of GPR179 in DBC function was confirmed in zebrafish and humans. Functional knockdown of *gpr179* in zebrafish led to a marked reduction in the amplitude of the ERG b-wave. Candidate gene analysis of *GPR179* in DNA extracted from patients with cCSNB identified *GPR179*-inactivating mutations in two patients. We developed an antibody against mouse GPR179, which robustly labeled DBC dendritic terminals in wild-type mice. This labeling colocalized with the expression of GRM6 and was absent in *Gpr179^{nob5/nob5}* mutant mice. Our results demonstrate that GPR179 plays a critical role in DBC signal transduction and expands our understanding of the mechanisms that mediate normal rod vision.

Congenital stationary night blindness (CSNB) is a severe disability that impairs night vision. Complete CSNB (cCSNB) is a genetically heterogeneous form of the disorder that is caused by mutations in genes that are required for signal transduction through retinal depolarizing bipolar cells (DBCs).^{1–8} The function of photoreceptors and DBCs can be assessed noninvasively with the electroretinogram (ERG), and their light-induced activities are reflected in the a-wave and b-wave, respectively.⁹ Individuals with cCSNB and animal models of the disorder have an ERG waveform that lacks the b-wave because of a failure to transmit the photoreceptor signal through the DBCs. Depolarization of the DBCs is initiated by a metabotropic glutamate receptor-mediated (GRM6)¹⁰ modulation of a transient receptor potential melastatin 1 cation channel (TRPM1).^{11–13} This G protein signal transduction cascade utilizes *Gαo1*,¹⁴ *Gβ5*,¹⁵ and depends on the auxiliary protein nyctalopin.^{16,17} Mutations in *GRM6* (MIM 604096), *TRPM1* (MIM 613216), or *NYX* (MIM 300278),

which encodes nyctalopin, all can cause cCSNB in humans.^{1–8} Mice with mutations in *Grm6*, *Trpm1*, *Gna0*, and *Gnb5* or *Nyx* also have a no b-wave (nob) ERG phenotype.^{10–16,18–20} In this report, we define a critical role for GPR179, a previously uncharacterized orphan G protein receptor, in the DBC signal transduction cascade and in human cCSNB. Specifically, mutations in *GPR179* in humans are responsible for a form of cCSNB. Consistent with this result, *nob5* mice have a mutation in *Gpr179* (*Gpr179^{nob5/nob5}*) and a nob ERG phenotype. Finally, zebrafish, whose *gpr179* expression is knocked down via morpholino injection, have a reduced ERG b-wave amplitude.

The *nob5* mouse arose as a spontaneous mutation in a colony of C3H mice and was identified via ERG when this line was crossed to a line of C3H mice lacking the *rd1* mutation (C3H-*f^{+/+}*). To identify the causative mutation, we crossed affected *nob5* mice to wild-type (WT) C57BL/6J mice and the resulting F1 mice were intercrossed to generate a segregating mapping cross. We identified F2

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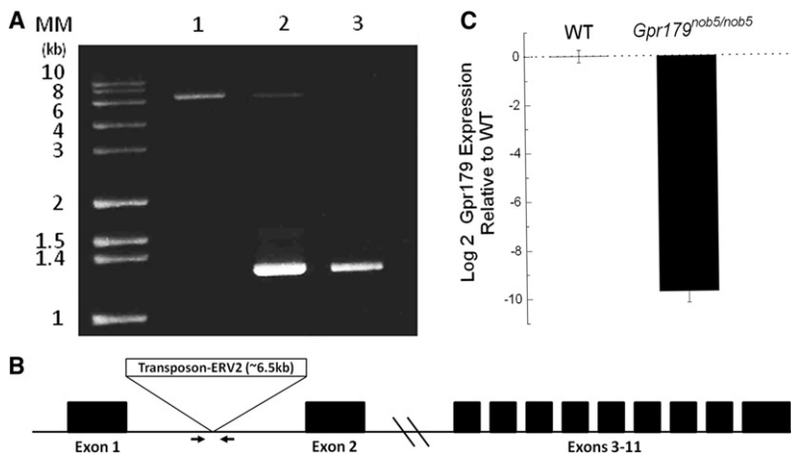


Figure 1. Transposable Element Disrupts *Gpr179* Expression in *nob5* Mouse

(A) PCR fragments from *Gpr179*^{nob5/nob5} (lane 1), *Gpr179*^{nob5/+} (lane 2), and WT C3H (lane 3). The insertion is ~6.5 kb.

(B) Schematic exon map of *Gpr179* indicating location of *nob5* insertion mutation. The arrows indicate location of PCR primers used in (A).

(C) Quantitative PCR of *Gpr179* cDNA generated from mRNA isolated from retinas of WT and *Gpr179*^{nob5/nob5} mice. Expression of *Gpr179* was normalized to that of *18S* RNA and is relative to the *Gpr179* expression in WT. The error bars indicate mean \pm standard deviation for three mice. *Gpr179* expression in *Gpr179*^{nob5/nob5} retina is significantly reduced ($p < 0.0005$). All animal studies were approved by the local institutional animal care and use committees and conformed to all regulatory standards.

progeny homozygous for the *nob5* locus by ERG and used them to map the phenotype by using a genome-wide screen with 103 simple sequence length polymorphic markers distributed throughout the genome.²¹ Initial mapping localized the gene to chromosome 11. Subsequently, > 600 additional informative meioses refined the map location of the *nob5* locus to between *D11Mit54* and *D11Mit67*. This 1.3 Mb region contains over 90 genes, none of which were known to be involved in DBC signal transduction or had been identified in molecular analyses of enriched pools of DBCs.^{22–24} To identify the mutation underlying the *nob5* phenotype we used genome capture and high-throughput sequencing. Comparison of the sequence encompassing the critical region in *nob5/nob5* and WT C3H mice revealed the presence of an insertion in intron 1 of *Gpr179* (Figure 1A). The next-generation sequence data provided only 10 bp of sequence on either side of the insertion, but these data suggested the insertion was a transposable element. To examine this directly, we used PCR to amplify the insertion and its flanking intronic DNA. This revealed the presence of the predicted 1.3 kb fragment in WT mice and a 7.8 kb fragment in homozygous affected littermates, indicating the insertion is ~6.5 kb (Figure 1A). Both bands were seen in heterozygotes. Henceforth, the mutant *nob5* allele will be referred to as *Gpr179*^{nob5}. Sequence analyses of the ends of the insertion indicated it was an endogenous retroviral element of the ERV2 class (Figure 1B). To evaluate the impact of this insertion on *Gpr179* expression, we used a quantitative intron-spanning Taqman RT-PCR assay to determine *Gpr179* mRNA levels of WT and *Gpr179*^{nob5/nob5} retinas (Figure 1C). The expression of mRNA representing *Gpr179* in the *Gpr179*^{nob5/nob5} retina was decreased more than 800-fold compared to the expression of mRNA in the WT retina. These data indicate that the *Gpr179*^{nob5/nob5} phenotype is caused by a large insertion mutation in intron 1 of *Gpr179*; this insertion dramatically reduces gene expression and probably represents a null allele.

A series of dark-adapted ERGs obtained from representative WT, *Gpr179*^{nob5/+}, and *Gpr179*^{nob5/nob5} mice are shown

in (Figure 2A). Throughout the stimulus range examined, WT ERGs are dominated by a positive polarity b-wave, which increases in amplitude with increasing flash luminance and reflects the light-induced activity of DBCs.²⁵ At higher flash luminance, the b-wave was preceded by a negative polarity a-wave, reflecting the light-induced closure of cation channels along rod photoreceptor outer segments.²⁶ ERG responses in heterozygous *Gpr179*^{nob5/+} mice resembled the responses of WT mice, consistent with autosomal-recessive inheritance. In contrast, whereas large a-waves are obtained from homozygous *Gpr179*^{nob5/nob5} mice, these responses lack the b-wave component, revealing slow PIII, an ERG component generated by the radial Müller glial cells.²⁷ Summary plots for the major components of the dark-adapted ERGs are shown in Figure 2B. ERG a-wave amplitudes were comparable across the three genotypes and the b-waves of *Gpr179*^{nob5/+} and WT mice were indistinguishable. The b-wave component is absent in *Gpr179*^{nob5/nob5} mice, and therefore, these data are not plotted. This ERG phenotype, in which the b-wave is absent while the a-wave is preserved, indicates that rod phototransduction is unaffected in *Gpr179*^{nob5/nob5} mice, whereas synaptic transmission between photoreceptors and DBCs, or DBC activity itself, is grossly abnormal.²⁸ Light-adapted ERGs obtained from representative WT, *Gpr179*^{nob5/+}, and *Gpr179*^{nob5/nob5} mice are shown in Figure 2C. In WT mice, the cone ERG was dominated by the positive polarity b-wave and higher frequency oscillatory potentials, which reflect activity through the DBC pathway.²⁹ In contrast, cone ERGs of *Gpr179*^{nob5/nob5} mice are electro-negative. Summary plots for cone ERGs recorded from all three genotypes are shown in Figure 2D. When cone ERG amplitude is measured from the negative trough to the following positive peak, the *Gpr179*^{nob5/nob5} response is reduced in amplitude, whereas those from WT and *Gpr179*^{nob5/+} heterozygotes are comparable. The *Gpr179*^{nob5/nob5} ERG phenotype is essentially indistinguishable from those of mouse mutants for other proteins involved in DBC signal transduction, including

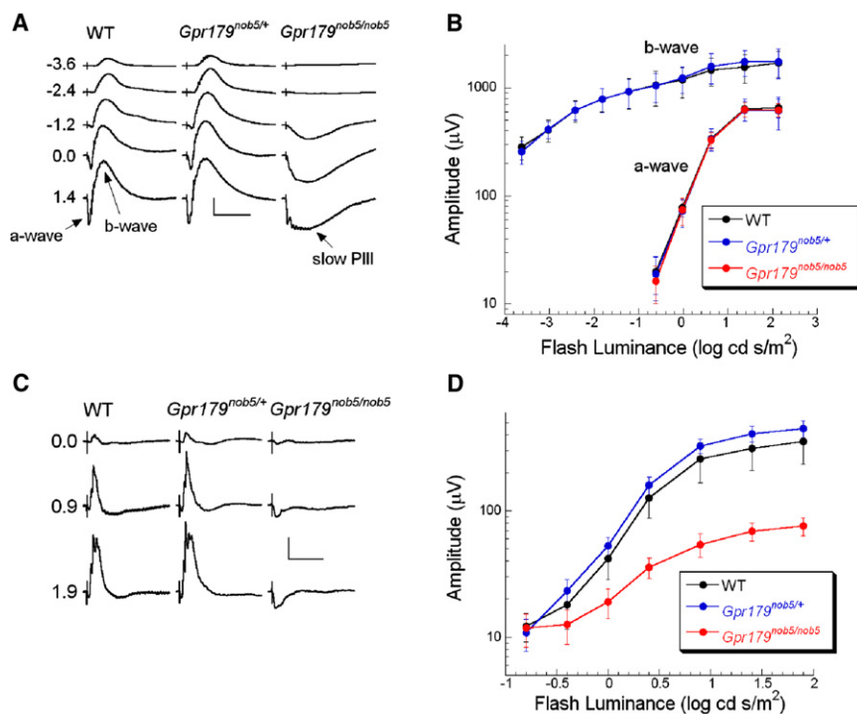


Figure 2. ERG Phenotype of *Gpr179^{nob5/nob5}* Mice

ERGs were recorded from mice anesthetized with ketamine (80 mg/kg) and xylazine (16 mg/kg) after overnight dark adaptation via a published procedure.¹⁶ (A) Dark-adapted ERG series obtained from representative WT (left), *Gpr179^{nob5/+}* (middle), and *Gpr179^{nob5/nob5}* (right) littermates at 6 months of age. The scale bars indicate 100 ms and 500 μ V. Values to the left of each row of waveforms indicate flash luminance in log cd s/m². (B) Luminance-response functions for the major components of the dark-adapted ERG. The b-wave component is absent in *Gpr179^{nob5/nob5}* mice and therefore these data are not plotted. (C) Cone-mediated ERG series obtained from WT (left), *Gpr179^{nob5/+}* (middle), and *Gpr179^{nob5/nob5}* (right) littermates at 6 months of age. Scale bar indicates 100 ms and 100 μ V. (D) Luminance-response functions for the cone ERG b-wave. Values to the left of each row of waveforms indicate flash luminance in log cd s/m².

GRM6,^{10,19,20} TRPM1,^{11–13} and NYX,^{16,18} a protein required for correct localization of TRPM1 channels in DBC dendrites.¹⁷

GPR179 encodes a predicted orphan G protein-linked receptor that has not been previously characterized in any cell or tissue. The *nob5* phenotype predicts that the *GPR179* gene product is required for DBC function and therefore should be present in DBCs. Because DBCs receive input from photoreceptors via ribbon synapses in the outer plexiform layer (OPL), we initially characterized gross retinal morphology and OPL ultrastructure of the *Gpr179^{nob5/nob5}* retina. However, all cellular and synaptic layers appeared normal in the *Gpr179^{nob5/nob5}* retina (Figure 3A). Furthermore, *Gpr179^{nob5/nob5}* ribbon synapses are indistinguishable from WT (Figure 3B). A normal retinal morphology is typical of all other mouse models of cCSNB.^{10,12,13,18,30}

To examine the cellular localization of GPR179 in the mouse retina, we developed a polyclonal sheep antibody to a peptide (KVQEETPGEDLDRPVLQKR) located within the amino terminal extracellular domain. Retinal cryosections from *Gpr179^{nob5/+}* and *Gpr179^{nob5/nob5}* littermates were reacted with our antibody to GPR179 and an antibody against PKC α to label rod DBCs. In *Gpr179^{nob5/+}* retina, PKC α labeled the entire rod DBC (Figures 3C and 3E), whereas the GPR179 antibody produced a punctate-labeling pattern (Figures 3D and 3E), which corresponds to the location of the OPL. In the *Gpr179^{nob5/nob5}* retina, labeling for PKC α (Figures 3F and 3H) is comparable to *Gpr179^{nob5/+}* and labeling for GPR179 is absent (Figures 3G and 3H), consistent with decreased *Gpr179* mRNA expression in *Gpr179^{nob5/nob5}* retinas (Figure 1C). The punctate-labeling pattern of GPR179 in the OPL is typical of

proteins that are localized to the dendritic tips of DBCs, including GRM6, TRPM1 and NYX.^{12,16} To confirm this localization, we double-labeled retinal sections with antibodies to GPR179 (Figure 3I) and GRM6 (Figure 3J). The punctate labeling for GPR179 colocalizes with GRM6 (Figure 3L), the glutamate receptor known to mediate signaling in DBCs. These data show that GPR179 is expressed on the dendritic terminals of DBCs. GPR179 is not expressed elsewhere in the retina (data not shown). The combined ERG, genetic and immunohistochemical data argue strongly that the *Gpr179^{nob5/nob5}* mouse phenotype is caused by the insertion and results in a functionally null allele of *GPR179*.

To directly determine whether reduced GPR179 expression could recapitulate the reduced b-wave phenotype, we used morpholino knockdown in zebrafish, which have a single copy of *gpr179* in their genome. We injected 1 cell stage zebrafish embryos with morpholinos (MOs) targeted against the *Gpr179* translation start site (MO-*Gpr179* 5'-GCCCATACTTTTAGCAACTGCTTCT-3'), and recorded ERGs at 4–6 days post fertilization. As comparisons, MOs against either the *nyx* translation start site (MO-*Nyx* 5'-GATGAAACACATCACTGGCTTC-3')³¹ or control (C-MO 5'-CCTCTTACCTCAGTTACAATTTATA-3') were injected. Embryos injected with MO-*Gpr179* had a significantly reduced ERG b-wave amplitude, similar to embryos injected with MO-*Nyx* (positive control) (Figures 4A and 4B). The b-wave was unaffected following injection of the control MO (Figure 4B). The b-wave/a-wave ratio decreased from 3.79 ± 0.46 ($n = 4$) in control to 0.75 ± 0.11 ($n = 7$) in *Gpr179*-MO ($p = 0.005$). These results indicate that *gpr179* expression is required for normal DBC function in zebrafish.

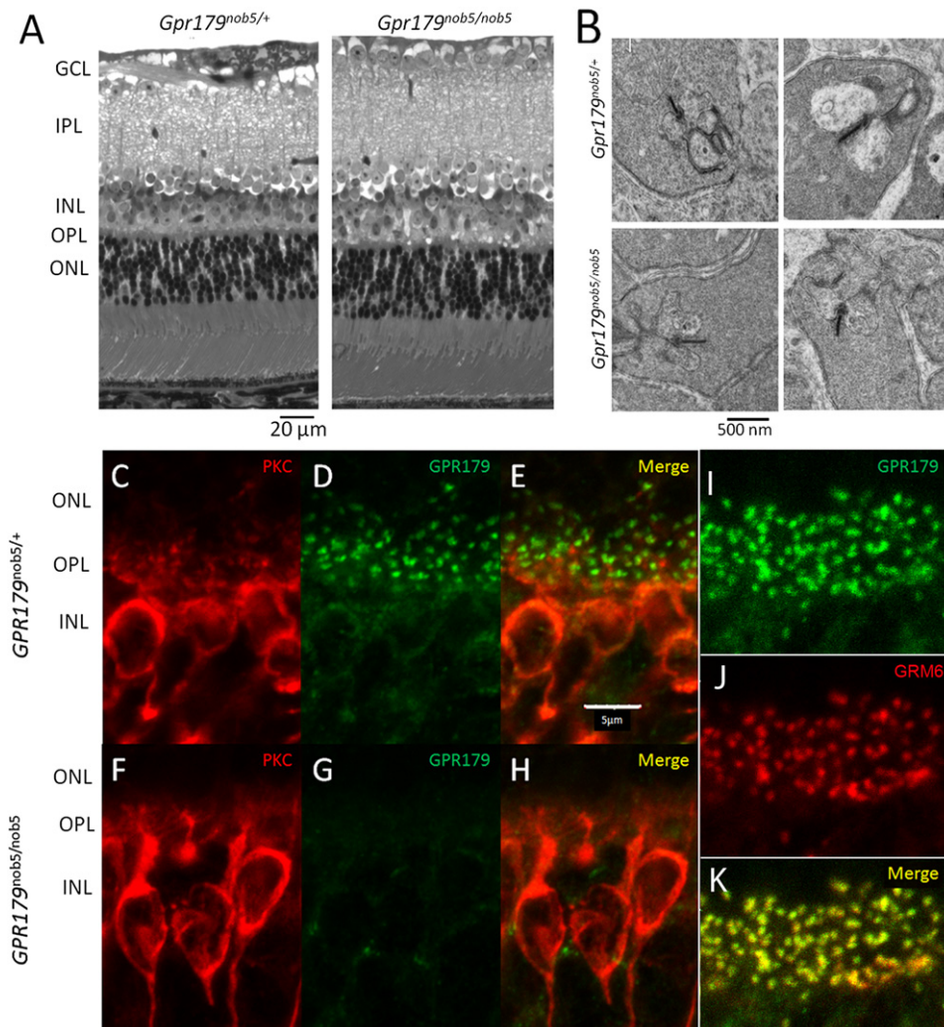


Figure 3. Anatomical Phenotype of *Gpr179^{nob5/nob5}* Retina

(A) Retinal cross-sections obtained from *Gpr179^{nob5/+}* and *Gpr179^{nob5/nob5}* mouse retinas fixed with 2.5% glutaraldehyde/2% paraformaldehyde and prepared according to published procedures.³⁵

(B) Electron micrographs of ribbon structures in *Gpr179^{nob5/+}* and *Gpr179^{nob5/nob5}* mouse retinas prepared according to published procedures.

(C–K) Confocal immunohistochemistry of retinas fixed with 4% paraformaldehyde for 15 min and prepared according to published procedures.¹⁶ The scale bar indicates 5 μ m. (C–E) *Gpr179^{nob5/+}* retina labeled with antibodies against (C) PKC α , (D) GPR179, and (E) merge of (C) and (D). (F–H) *Gpr179^{nob5/nob5}* retina labeled with antibodies against (F) PKC α , (G) GPR179, and (H) merge of (F) and (G). (I–K) *Gpr179^{nob5/+}* retina labeled with antibodies for (I) GPR179, (J) GRM6, and (K) merge of I–J. The following antibodies were used: GPR179; affinity purified polyclonal sheep anti-GPR179 peptide (KVQEETPGEDLDRPVLPQKR), 1:1,000; mouse monoclonal anti-ctbp2/Ribeye, 1:1,000 (BD Biosciences); guinea pig anti-GRM6, 1:1000 (see Koike et al.¹²); rabbit anti-PKC α (1:1,000, Sigma). The following secondary antibodies were used: Alexa-488 donkey anti-sheep, Alexa-555 goat anti-rabbit, and Alexa-633 goat anti-guinea pig (all at 1:1,000; Invitrogen).

Mutations in *NYX*, *GRM6* or *TRPM1* have been identified in patients with cCSNB.^{1–8} However, a small number of individuals with cCSNB did not bear mutations in any of these genes.⁸ To evaluate the potential involvement of *GPR179* in cCSNB patients lacking DBC function, we sequenced the 11 exons and flanking splice sites of the human gene in 44 patients (see Table 1 for primers). All human studies were undertaken with the approval of the appropriate institutional review board. We identified two probands with inactivating mutations in the *GPR179* gene. Proband 1 had no family history of night

blindness or consanguinity and was 10 years old at the time of diagnosis. He presented with 20/70 best corrected visual acuities, mild myopic refractive error, congenital nystagmus, a history of early onset nightblindness, a normal retinal appearance and full Goldmann visual fields. ERGs obtained under ISCEV standard conditions from proband 1 are shown in Figure 5A. Under dark-adapted conditions (upper traces), the ERG b-wave recorded to a low luminance stimulus was markedly reduced in amplitude, whereas the ERG obtained to a high flash luminance had a robust a-wave without the

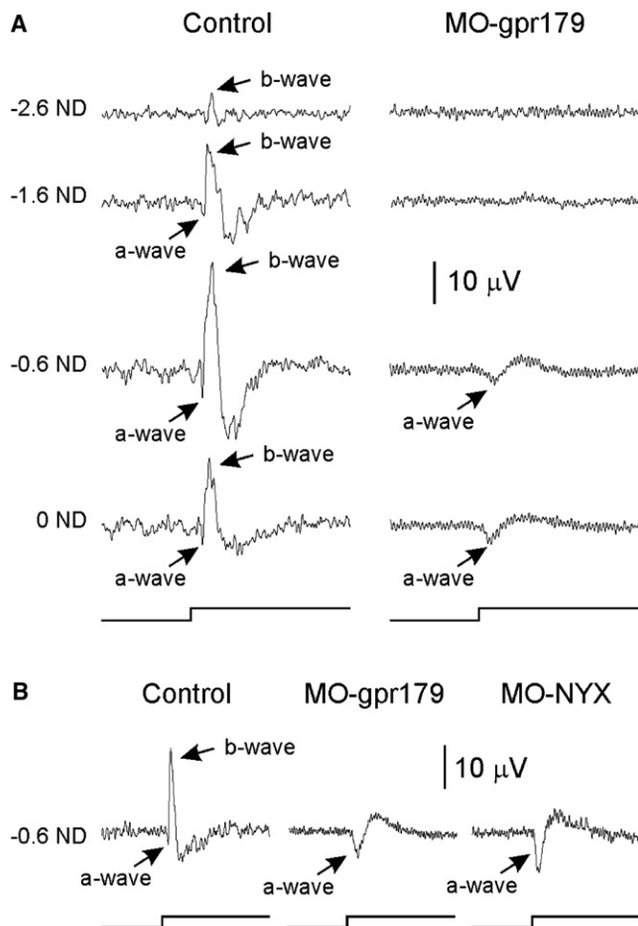


Figure 4. GPR179 Knockdown in Zebrafish Decreases ERG b-Wave

Injected MOs (Gene Tools) were designed against the *gpr179* translation site (MO 5'-GCCCATACTTTTAGCAACTGCTTCT-3'), which occurs as a single copy in the zebrafish genome, or the *nyx* translation site (MO 5'-GATGAAACACATCACTGGCTTC-3') or were a standard control (MO 5'-CCTCTTACCTCAGTTACAATTATA-3'). In each case, 30 ng of MO was injected into the chorions of one-cell-stage zebrafish embryos. ERGs were recorded from larvae at 4–6 days post fertilization with a 1 s stimulus after 30 min of dark adaptation, as previously described.³⁶

(A) ERGs from embryos injected with control (left column) or MO-*gpr179* (right column) MOs and tested at four flash intensities. Flash intensity at 0 log is 9.3 W/m². Neutral density (ND) filters reduced the intensity by the indicated amount in log units. The b-wave amplitudes obtained from MO-*gpr179* injected embryos were strongly reduced at all intensities. The b-wave/a-wave ratio decreased from 3.79 ± 0.46 ($n = 4$) in control to 0.75 ± 0.11 ($n = 7$) in MO-*gpr179* ($p = 0.005$), showing that a specific knockdown of *gpr179* reduces DBC function.

(B) The magnitude of b-wave reduction for MO-*gpr179* and MO-*nyx* were comparable. Injections of MO-control did not affect the ERG (data not shown).

subsequent b-wave seen in controls (middle traces). Under light-adapted conditions (lower pair of traces), the ERG waveform showed a square a-wave but retained a late positive ERG component. ERGs of proband 2 showed a similar selective absence of the dark-adapted b-wave and a square light-adapted ERG a-wave (data not shown). These ERG abnormalities have been uniquely associated

with human cases of DBC dysfunction^{32–34} and readily distinguish cCSNB from other retinal disorders with a reduced b-wave, such as incomplete CSNB (MIM 300071 and 610427).³³

Proband 1 was a compound heterozygote for two frameshift mutations in *GPR179*, c.187delC and c.984delC (NM_001004334.2) resulting in predicted protein truncations p.Leu63Serfs*12 and p.Ser329Leufs*4, respectively (Figure 5B). The premature chain termination is expected to result in functional null alleles. The probands' unaffected parents were each heterozygous for one of the mutations (Figure 5B).

Proband 2 was 20 years old at the time of diagnosis. She is of Norwegian descent and presented with rotatory nystagmus, a very unusual blond fundus, and congenital nightblindness and was able to see 20/30 with a -12.00 D prescription. Although not known to be related, proband 2 also carried the c.984delC frameshift mutation identified in proband 1, suggesting that this might be a founder mutation. Proband 2 carried a second mutation, c.659A>G, that would result in missense mutation, p.Tyr220Cys. (Figure 5C). The three variants in probands 1 and 2 were not present in 210 healthy control chromosomes. Two out of three Alamut analyses, which predict mutation impact on function, classified the p.Tyr220Cys as potentially pathogenic. Moreover, by introducing a new cysteine into the GRP179 protein, the mutation is likely to impact its structure. The functional importance of Tyr220 is supported by its conservation across species ranging from human to *Tetraodon* (Figure 5D). DNA of family members of proband 2 was not available for segregation analysis.

The combined data indicate that GPR179 is required for DBC signal transduction and that mutations that disrupt the function of GPR179 cause a recessive form of cCSNB. Although *Gpr179*^{nob5/nob5} mice lack expression of GPR179, we have not detected any anatomical defect in the retina, and the mice have no apparent health problem. Similarly, the two human patients we have identified with cCSNB and mutations in *GPR179* have no other known health problems. Given the ERG phenotype in mice, zebrafish, and humans, as well as the colocalization of GPR179 with GRM6, we postulate that GPR179 plays a critical role in DBC signal transduction, possibly by forming heterodimers with GRM6. Future studies will be required to define the specific role of GPR179 in this process. The availability of the *Gpr179*^{nob5/nob5} mouse model will be an important tool with which to address this question.

Acknowledgments

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Table 1. PCR Primers Used to Generate Products for Sequence Analysis

Name	Forward Primer (5' → 3')	Reverse Primer (5' → 3')	Amplicon Size (bp)
GPR179-EXON1_1	GACTGCAGCCAGCCTCTG	GCATGTTGAGAAAATTGGCG	402
GPR179-EXON1_2	AGATGCCAGCAGCTATCAC	TCAACTCGCTTCTTCAGG	410
GPR179-EXON1_3	ACCTTTAACCTCCACCAGG	GGACCTGACTTCTTCCCC	410
GPR179-EXON2	AAAGTGCCATAATTCCGAGC	GAGATGCTGGGGTAGCTTTG	330
GPR179-EXON3	GACCCAAGTCCCATGGATAG	GTACCCTCCAACCATCCTC	261
GPR179-EXON4	GGCTGTGGTGACTATCAGGG	TAGTCTCAGGCCTCCAGTTG	386
GPR179-EXON5	ACAGGGTGAAGCCTTAGCTC	GTGCCAGGTTTGGCTATG	241
GPR179-EXON6	TGGTGATGGCTCTATTGCAG	ATGAGTGAGCTGGGGTGG	284
GPR179-EXON7	CAGCAATGGACAGAGACCAG	CCAGAACAAGAGGAACCCTG	401
GPR179-EXON8	TCTTTCAGAGAAAGGGGTGG	GTAGCAGTGTGCCAGGATG	313
GPR179-EXON9	CTAAGCGTATCAGGGTTGGG	CTCCACAGTGGCCCTGAC	274
GPR179-EXON10	TGAGTGCAGGAGTGGACAAG	GACTTGCAGTGGCACATAGC	311
GPR179-EXON11-1	AAGGGTGGAGGAACAGACAG	CGCTGGCTGACCTGAAG	497
GPR179-EXON11-2	GACCCGCCTTCTTACTGACTC	TAGAAACCTGGTGCCTGATG	472
GPR179-EXON11-3	AGGCCATCAGCCAGGAG	GCCCTGCTTTTCTACTGTC	473
GPR179-EXON11-4	CAGCCAAGCAAGAAAATGTG	TCTGTTGGTTCTGGAGGGAC	471
GPR179-EXON11-5	CGTTCTCGGAGCACCTACAG	GCTGACGCGTTTCTGATTC	481
GPR179-EXON11-6	ATCAAAGAAACCCTGTTCG	TCAGGGGTATGAGCTTCTGG	480
GPR179-EXON11-7	CTGGGAGACGAGTGAAGGAG	AAGTCTCCTCCCGTTTCTG	474
GPR179-EXON11-8	AGTGAAGGAGGCGAGGATG	TCTCCCTGGGACAACTGAC	463
GPR179-EXON11-9	AATGCTGAGTCTGGGGACAG	GTCTTGAGGACGTGGTTGTG	529
GPR179-EXON11-10	TGGAATAATCGAAATCGAAG	ACCTGGATGCTGAGAACAGG	460
GPR179-EXON11-11	CCCAGGGAGAGAGCTGTTAC	GCTGCTGGGCAAGTCTGAG	474
GPR179-EXON11-12	GCCTGTTCTCAGCATCCAG	TGGGTTATGTGTCTGGGAAG	482
GPR179-EXON11-13	GAGTAGTGAAGTGGCAGAGGG	CTGAACTGGCACACCTTTTG	487
GPR179-EXON11-14	AATCTGACAGCAGTGCCAAG	AGGAACATCTCTCCTTGCCC	445
GPR179-EXON11-15	CTGTGGGAAGTGGTAGAGGC	TTTCAGGAGCTGTGGGG	473
GPR179-EXON11-16	AGGGTACCATGGCAGACATC	TTCAGCTCTTGGGAACACC	486

(R21EY021852 to N.S.P. and R.G.G.; R01EY12354 to R.G.G.; R01EY014701 to M.A.M.; RR017890 to A.F.X.G.; R01EY016501 and P30CA34196 to P.M.N.; R01EY004864 and P30EY006360 to P.M.I.; and R01NS43459 and R01EY020821 to G.T.); the Veterans Administration Medical Research Service; the Netherlands Organization for Scientific Research (Zon-MW NWO to M.K.); ODAS and European Commission FP7 Grant RETICIRC HEALTH-F2-2009-223156 (to M.K.); Algemene Nederlandse Vereniging ter Voorkoming van Blindheid (to A.A.B.B.); the Foundation Fighting Blindness Canada (to R.K.K.); the Canadian Institutes for Health Research (to R.K.K.); Reseau Vision (to R.K.K.); Fonds de recherche du Québec-Santé (to R.K.K.), a Foundation Fighting Blindness Center Grant to the Cole Eye Institute, Cleveland Clinic; and unrestricted awards from Research to Prevent Blindness to the Department of Ophthalmology, Cleveland Clinic Lerner College of Medicine; to the Department of Ophthalmology, Emory University; and to the Department of Ophthalmology and Visual Sciences, Univer-

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Web Resources

The URLs for data presented herein are as follows:

Human Genome Variation Society, <http://www.hgvs.org>

Online Mendelian Inheritance in Man (OMIM), <http://www.omim.org/>

Primer3, <http://frodo.wi.mit.edu/primer3/>

SNPCheck, <https://ngl.manchester.ac.uk/SNPCheckV2/>

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