A Novel Achromatopsia Mouse Model Resulting from a Naturally Occurring Missense Change in Cngb3

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PURPOSE. A local colony of inbred mice (129S6/SvEvTac origin), in isolation for over a decade, were found to have absent light-adapted electroretinogram (ERG) responses. We investigated the inheritance and genetic basis of this phenotype of cone photoreceptor function loss.

METHODS. An affected 129S6/SvEvTac colony animal was outcrossed to a C57BL/6j mouse and intercrossed to investigate inheritance in the F2 generation. We performed ERG testing and targeted resequencing on genes of interest (Gnat2, Cnga3, Cngb3, Pde6c, Hcn1, Syn2). The eyes of a subset of animals underwent histologic immunostaining.

RESULTS. All 129S6/SvEvTac colony animals tested lacked cone pathway function by ERG testing (n = 12), although rod pathway–based ERG responses remained unaffected. Outcross-intercross breeding showed a recessive inheritance pattern. A novel missense mutation was identified in the Cngb3 gene, which causes an amino acid substitution at a conserved residue (NM_013927:c.692G>A; p.(R231H)). The recessive phenotype only affected homozygotes (χ² = 39, P = 3.2e -10). Cones had normal morphology at postnatal day (PND) 70, but cone cell counts declined from PND 30 to PND 335 (P = 0.058), indicating progressive cone photoreceptor death.

CONCLUSIONS. We identified the spontaneous occurrence of a 10th model of cone photoreceptor function loss (cpfl10) in an isolated line of inbred mice. Our results indicate that this is caused by a novel missense mutation in the Cngb3 gene, with a fully recessive inheritance pattern. This mouse may provide a more appropriate background against which to assess CNGB3 achromatopsia gene therapy for missense mutations.

Keywords: retina, achromatopsia, Cngb3, cones

We noticed that a particular colony of inbred mice (of 129S6/SvEvTac origin) had poor light-adapted visual responses and hypothesized that a novel genetic mutation affecting cone photoreceptors might be responsible. Cone photoreceptors facilitate daytime vision, color perception, and homeostatic functioning of cones has multiple linked steps and a single cone gene mutation can cause loss of cone function.1

Mice are valuable for the identification and investigation of genes and mutations that may affect normal cone photoreceptor function, particularly in achromatopsia. There have been nine mouse models of cone photoreceptor function loss (cpfl) reported since cpfl1 was first described (Chang B, et al. IOVS 2001;42:ARVO Abstract 527). The cpfl retinal phenotype varies across the models and the genetic basis is different for each. Yet, all nine models display the common trait of partial or complete loss of cone function from birth, sometimes with progressive cone cell loss (Chang B, et al. IOVS 2001;42:ARVO Abstract 527) (Hawes NL, et al. IOVS 2003;44:ARVO E-Abstract 4531) (Chang B, et al. IOVS 2006;47:ARVO E-Abstract 2294) (Hawes NL, et al. IOVS 2004;45:ARVO E-Abstract 3590) (Hawes NL, et al. IOVS 2006;47:ARVO E-Abstract 4579) (Hawes NL, et al. IOVS 2007;48:ARVO E-Abstract 1350) (Chang B, et al. IOVS 2015;ARVO E-Abstract B0250).1,2 Mouse models also facilitate the development of retinal gene and molecular therapies; two cpfl models have been central in developing adeno-associated virus (AAV) gene therapies before trials in humans.3,4 The cpfl phenotypes model human achromatopsia, a rare (1 in 30,000-80,000), congenital visual condition characterized by diminished or absent cone photoreceptor function.5 Patients have severely reduced visual acuity (~20/200), nyctagmus, and photophobia.6 Causative autosomal recessive mutations largely affect genes of the cone phototransduction cascade and are summarized elsewhere.7

CNGB3 forms a heterotetramer cyclic nucleotide–gated (CNG) transmembrane channel with CNGA3 and is the most commonly affected gene in human achromatopsia.8 CNGB3 is the minor component in direct channel conductance and plays a modulatory role.9 In murine retinal sections, CNGB3 and CNGA3 colocalize and are expressed exclusively in cone photoreceptors.10 CNGB1 and CNGA1 form an analogous CNG channel specific to rod photoreceptors.11 Here we described the cpfl10 mouse phenotype in detail and identified the causative missense mutation in exon 6 of Cngb3, producing a substitution of histidine for arginine. We identified a slow degeneration of cones in this model and demonstrate that heterozygotes do not suffer any intermediate phenotype.
METHODS

Mice and Breeding

All animals used in this study were treated humanely in accordance with the UK Home Office Regulations and the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Animals were housed in a 12:12 hour light-dark cycle. An affected 129S6/SvEvTac colony animal was outcrossed to a wild-type C57BL/6J mouse to produce first filial generation hybrid offspring (129SB6F1). The 129SB6F1 animals were intercrossed to investigate inheritance in the F2 generation. A further intercross of affected homozygous F2 offspring was performed to produce an F3 generation to confirm penetrance. Complementation breeding of an 129S6/SvEvTac mouse with a Cnga3/0 colony produced offspring to exclude Cnga3 as a possible affected gene. In confocal scanning laser ophthalmoscopy (cSLO), optical coherence tomography (OCT), and qPCR experiments, mice from a commercially obtained 129S2/SvHsd colony (Envigo, Cambridgeshire, UK) were used as approximate strain-matched controls.

Electroretinography

Before the ERG procedure, mice were dark adapted (>1 hour) and the procedure was performed under dim red illumination. The mouse was anaesthetized with a single intraperitoneal injection (80 mg/kg ketamine and 10 mg/kg xylazine) and the pupils were dilated with tropicamide 1% and phenylephrine 2.5% eye drops. The mouse was positioned on a heated mat in front of the testing console (Colordome Electroretinography machine; Diagnosys LLC, Vision Park, Cambridge, UK), which also generated and controlled the light stimulus. Corneal electrodes (Diagnosys LLC) coupled with custom lenses produced from achromatic Aclar embedding (Honeywell International, Morris Plains, NJ, USA) were positioned on the cornea by using micromanipulators and viscous coupling media (Hypermellose 1%; Alcon, Fort Worth, TX, USA). Subcutaneous ground and reference electrodes (platinum needles) were placed in the flank and scalp.

The test protocol consisted of three dark-adapted and seven light-adapted steps. Dark-adapted recordings commenced with a dim single flash −4 log cd.s/m² stimuli, with an interstimulus interval (ISI) of 5 seconds, and averaged over 9 trials. A brighter single flash of 1 log cd.s/m² stimuli was also used, with an ISI of 30 seconds, and averaged over 4 trials. Continuous stimuli of 7 Hz were delivered at −2 log cd.s/m² and averaged over 30 responses. After dark-adapted recordings were completed, animals were exposed to a full-field 30 cd/m² white background for 10 minutes; subsequent steps were delivered on top of this continuous background. Single-flash stimuli after light adaptation consisted of −0.5, 0, 0.5, 1, and 1.5 log cd.s/m²; 20 responses were averaged, with an ISI of 1 second. Continuous stimuli of 7 Hz and 20 Hz were delivered at 1 log cd.s/m² and averaged over 30 responses.

ERG a-wave and b-wave amplitudes were measured (Espion v6; Diagnosys LLC) by using manual and automated methods. Technically inadequate replicates of traces were excluded. A-wave and b-wave cursors were placed manually. Full b-wave amplitudes in rod-isolating test steps were measured from the trough of the a-wave to the peak of the b-wave. If the a-wave was undetectable then b-wave amplitude was measured from the baseline at stimulus onset.

cSLO and OCT

The retinal phenotype was assessed in vivo by using a combined confocal scanning laser ophthalmoscope and spectral-domain optical coherence tomograph (Spectralis HRA; Heidelberg Engineering, Heidelberg, Germany). Fundus autofluorescence imaging was performed by following a previously described protocol using 488- and 790-nm excitation light, and near-infrared reflectance imaging using a 820-nm laser. OCT captures cross-sectional images of the retina by using low-coherence interferometry of infrared light. The section density, orientation, and technical replicates of scan patterns were customized to include a high-definition linear scan and a radial scan. Total and outer retinal thickness measurements were made manually by using calipers on Heidelberg software.
Tail tissue was collected and genomic DNA (gDNA) was extracted with a commercial DNA mini kit (QIAGEN, Manchester, UK). Custom primer pairs were designed to the reference coding sequence for *Gnat2*, *Cngb3*, and *Pde6c*.

Initially, gDNA from six pilot mice was amplified with PCR and the target amplicon confirmed with agarose gel electrophoresis. The PCR product was extracted with a commercial kit according to the manufacturer’s instructions. Samples underwent commercial fluorescent dideoxynucleic acid sequencing and the results were analyzed with Geneious v8.0 software (Biomatter, Auckland, New Zealand). Single nucleotide polymorphisms (SNPs) were compared to the National Center for Biotechnology Information (NCBI) dbSNP database; established SNPs were assessed as nonpathologic and not investigated further. The *Cngb3* base pair variation not documented on NCBI dbSNP was sequenced further in the full 77 mouse cohort of *cpfl10* (n = 12), F2 intercross (n = 59), and F3 intercross mice (n = 6) to confirm cosegregation with phenotype.

Sanger sequencing results were validated by a custom RFLP assay. Genomic DNA was amplified by using custom primers targeting *Cngb3* exon 6 (FW-5’-TTCTCTCCACACCGTGCAG-3’ and RC-5’-ACACTGACAAATGTCTATCCACAGA-3’) and amplicons underwent restriction digest with *Apa*LI enzyme (NEB, Ipswich, MA, USA). On an agarose gel, wild-type mice showed bands at 454 bp and 15 bp, whereas homozygous mutants lacked the larger band and showed 267-bp, 187-bp, and 15-bp bands.
FIGURE 3. (A) Representative en face confocal scanning laser ophthalmoscopy images show no retinal changes indicating degeneration across all imaging modalities (PND 56). (B) Representative optical coherence tomography cross-sections show equal retinal thickness (PND 56) that is consistent across all mice measured ($n = 5$ cpfl10 and $n = 5$ 126SvEvTac wild types). (C) Immunohistochemical staining for Arr3 in the cpfl10 retina shows grossly normal cone morphology at PND 70 ($n = 4$ cpfl10 eyes compared to $n = 2$ wild-type eyes). Scale bar: 50 μM. (D) The mean retinal cone count in cpfl10 mice declines significantly from PND 30 to PND 335. (E) Representative retinal flat mounts with Arr3 staining showing the difference in cone number and morphology between cpfl10 and controls. Scale bar: 10 μM.
fragments—the G>A missense mutation introduced a new *ApaI* cut site in the middle of the exon 6 amplicon (see Figs. 4B, 4C).

### mRNA Extraction and qPCR

Retinal tissue was homogenized in a lysis solution of B-mercaptoethanol and chaotropic buffer. Passing the solution 10 times each through a 25-gauge and 30-gauge needle optimized homogenization. The solution was then bound to a silica-gel-membrane spin column by centrifugation. DNase was added to degrade contaminant DNA, before further wash steps with buffer. The RNA was eluted in water.

Eluted retinal RNA was reverse transcribed to cDNA by using the Superscript III synthesis system (Life Technologies, Paisley, UK) as per the manufacturer's instructions. cDNA was purified for PCR by using the QIAquick PCR purification kit (QIAGEN, Manchester, UK). Standard curve analysis was performed to ensure efficiencies were close to 100% and the optimal primer pairs for each gene were used (Table).

All qPCR reactions used a commercial kit (SYBR Green PCR Master Mix; Applied Biosystems, Foster City, CA, USA) and 2 μM final concentration of each forward and reverse primer. All qPCR experiments were performed in triplicate with a commercial real-time PCR machine (CFX Connect Optics Module; BioRad, Hercules, CA, USA). Reactions were performed with the following settings: an initial denaturation step of 95°C for 10 minutes, followed by 40 cycles of 95°C for 10 seconds, an annealing temperature of 55°C, and extension at 72°C for 30 seconds.

![Figure 4](image-url)

**Figure 4.** (A) The chromatogram for the c.692G>A mutation is shown for heterozygotes (top), homozygotes (middle), and wild-type littermates (bottom). (B) The c.692G>A mutation introduces a novel *ApaI* restriction site in the *Cngb3* gene sequence. (C) An RFLP assay can thus identify homozygous, heterozygous, or wild-type mice; wt band = 469 bp, mutant band = 250 bp + 190 bp + 15 bp fragments. Sanger sequencing results for wt (+) or mutant (−) displayed below each lane show 100% concordance across all samples tested with RFLP (n = 43 animals assayed). NTC is negative template control, DNA ladder in kilobases. (D) A representation of the complementation and outcross breeding undertaken to identify the inheritance pattern of the *cpfl10* phenotype. The panel also lists the numbers of mice genotyped as heterozygous (het), homozygous (hom), or wild type for the c.692G>A missense mutation.
**RESULTS**

**Functional and Structural Retinal Phenotype of the Mouse cpf10 Mutant**

The cpf10 phenotype can be easily identified by ERG. For all mice tested in the original colony (n = 12), dark-adapted rod responses were equivalent to C57BL/6j wild-types (linear regression, P = 0.4), and light-adapted cone responses were completely absent or substantially reduced to 15% to 20% of littermate b-waves (n = 12 mice; Fig. 1). A further 114 mice derived from outcrosses and intercrosses of cpf10 were phenotyped. All of the original cpf10 mice (n = 12), the entire F2 cohort (n = 59), and the F3 intercross cohort (n = 6) were genotyped. Figure 2 gives detailed analysis of the F2 and F3 cohort ERG phenotype and the relationship to genotype. The cpf10 retinal fundus appeared normal on all applied cSLO imaging modalities when compared with a commercially obtained 129S2/SvHsd colony (Envigo) wild-type (Fig. 3A, investigated up to postnatal day [PND] 60). OCT assessment of the retinal layers showed normal architecture and thickness measurements that were not different when compared to wild type (194 ± 0.03 μM, 193 ± 0.09 μM; P = 0.43; Fig. 3B). Immunohistochemistry (IHC) staining for cone arrestin at PND 70 showed normal retinal appearance and cone morphology (Fig. 3C). However, retinal flat mounts stained for cone arrestin (Figs. 3D, 3E) in cpf10 mice at PND 30 (n = 3) and PND 335 (n = 3) showed a significant reduction in retinal cone counts over time (P = 0.039; t-test). The cone loss observed in cpf10 mice at PND 335 was far more severe than what might be expected for normal aging (145 ± 13 nuclei/field of view for cpf10 versus 302 ± 17.8 nuclei/field for approximately age- and strain-matched controls). Yet, surviving cones in cpf10 mice were slightly more numerous than in age-matched mice from the Cnga3<sup>−/−</sup> line (87 ± 3 cone nuclei/field of view at PND 355), which is known to experience progressive cone photoreceptor loss.

**Inheritance Pattern**

Figure 4D shows that complementation breeding of the cpf10 mouse with a Cnga3<sup>−/−</sup> colony produced offspring (n = 21) all with normal cone function on light-adapted ERG testing. The lack of complementation showed the cpf10 mutation is not allelic to Cnga3 (χ² = 3.87, df = 1, P < 0.001).

In outcrosses with C57BL/6j mice, the cpf10 phenotype occurred in an autosomal recessive pattern. None of the C57BL/6j outcross F1 progeny (129860F1) displayed a deficit in cone ERG function (n = 28), excluding a dominant inheritance pattern (χ² = 57, P < 0.001). The F1 progeny were then intercrossed to each other to produce an F2 generation to further explore inheritance. Approximately one quarter in the F2 generation (n = 15/59) showed the original cpf10 phenotype, indicating an autosomal recessive inheritance pattern (χ² = 13, P < 0.001). Intercross of two affected F2 mice showed full penetrance of the phenotypes in the subsequent F3 generation (n = 6). The cpf10 ERG phenotype cosegregated with the c.692G>A mutation in 100% of mice sequenced (χ² = 39, P = 3.2e−10).

**Mutation Analysis**

Targeted resequencing revealed that affected mice did not carry any of the spontaneous cpf10 mutations previously described. No mutations were detected in affected mice during sequencing of the entire coding sequence of Gnat2—the known Gnat2<sup>Exor3</sup> exon 6 mutation was excluded (Chang B, et al. IOVS 2001;42:ARVO Abstract 327). No mutations were
found in those exons of Hcn1 gDNA that were sequenced (approximately 70% of exons), which had been previously linked to achromatopsia (Chang B, et al. IOVS 2006;47:ARVO E-Abstract 2294). Further sequencing of the remaining Hcn1 exonic gDNA was stopped once the causative mutation was found in the Cngb3 gene. The Synem2 mutation (c.13978C>T; p.Q4660*) was excluded. Cngb3 was not sequenced, as it was excluded by complementation testing.

A novel missense mutation, c.692G>A; (p.Arg231His), was identified in exon 6 of the Cngb3 gene. The mutation introduces a unique Apal restriction site that can be exploited in a genotyping RFLP assay to confirm the Sanger sequencing results (Fig. 4). There are no other known cpfl10 genes located on chromosome 4 along with Cngb3 that could cosegregate.

The arginine at position 231 is the first extracellular amino acid immediately adjacent to the first transmembrane domain of the CNGB3 protein and is highly conserved across species (Fig. 5A). The arginine residue in the same location in CNGA3 is also conserved across species (Fig. 5B). Fewer than 25% of residues are conserved between both CNGA3 and CNGB3 across multiple species. The CNG channels are heterotetramers and thus important residues and protein domains share similarities between both component proteins. The location of the affected residue within the larger structure of CNGB3 and the heterotetramer is displayed in Figures 5C and 5D.

In silico analysis of the mouse mutation predicted a “deleterious” effect, with a PROVEAN score of −4.985, well beyond the cutoff of −2.5. Based on cross-species conservation, the predicted human mutation (NM_0190998.3:746G>A; p:(R259H)) was explored by using peer-reviewed PolyPhen II in silico tools and a score of 1.00 predicted this equivalent mutation to be “probably damaging” in humans.

**Effect of cpfl10 Mutation on mRNA and Protein Expression**

Regarding mRNA, retinal samples from cpfl10 mice (n = 5 cpfl10 mice; PND 84) and controls (n = 3 12952/SvHsd mice [Envigo]; PND 80) expressed both Cngb3 and Cnga3 transcripts on qPCR. After normalization to the ActB reference gene, Cngb3 expression levels in cpfl10 mice were similar to controls (ΔΔCt = 0.83 ± 0.85; P = 0.262). Expressed as a ratio, Cngb3 expression in control mice was 1.7 ± 0.95 times greater than Cnga3 (P = 0.003, Fig. 5E). In cpfl10 mice, the Cngb3 transcript expression levels were 3.5 ± 0.94 times higher than Cnga3 levels (P < 0.001), a ratio that was significantly higher than controls (P = 0.037).

However, the Rbo levels of cpfl10 mice were significantly lower than controls (ΔΔCt = 0.56 ± 0.85, P = 0.011), which suggests more extensive strain-related differences in the retinal transcriptome beyond the effects of the cpfl10 mutation.

Regarding protein, the previously described Cngb3 knockout mouse does not produce CNGB3 protein.12 The CNGB3 IHC performed after successful AAV/CNGB3 gene therapy rescue of this model was performed by using a custom-generated antibody that is not commercially available. All three commercially available antibodies to human CNGB3 (ABIN571515, Antibodies Online, Atlanta, GA, USA; CPBT 31596RH, Creative Diagnostics, Shirley, NY, USA; Orb156415, Biorbyt, Cambridge, UK) were tested on wild-type murine retinal sections at a range of concentrations. A nonspecific binding pattern was observed that could not be improved with optimization steps. Therefore, the protein expression of CNGB3 in the cpfl10 mouse is currently unknown.

**DISCUSSION**

This is the first naturally occurring mouse model of achromatopsia resulting from a mutation in Cngb3. A spontaneously occurring G>A nucleotide substitution in exon 6 of Cngb3 mutates a conserved arginine residue. The missense mutation is located at the beginning of the first extracellular loop of the B-subunit of the cone-specific cyclic nucleotide-gated ion channel. In silico analysis on multiple platforms confirms a deleterious effect across species for mutations at this residue.

Normal cone function is lost in Cngb3<sup>cpfl10</sup> mice. The extent of cone function loss is potentially more severe than the incomplete achromatopsia observed in the previously published Cngb3<sup>−/−</sup> (Cngb3<sup>knockout</sup>) mouse. This model contains a targeted deletion in exon 6, resulting in a premature stop codon c.735_749del; (p.A247*), and yet still reliably retains 20% to 30% of wild-type ERG function.25 This may arise because the CNGA3 subunits form a channel on their own. It is also possible that truncated CNGB3 is still translated and contributing to function—the authors have used a far C-terminal antibody that would be unable to detect truncated products.25 Notably, the cpfl10 model involves a slow progressive cone degeneration over the first year, which has been observed in other models of achromatopsia such as the cpfl10 mouse.24 Adaptive optic retinal imaging in human CNGB3 and CNGA3 achromatopsia cases has shown a slow decline in foveal cone density and changes in cone reflectivity throughout lifetime.25 In both the cpfl10 mouse and human cases, cones survive long enough to provide a therapeutic window for treatment.

CNGB3 forms a heterotetramer cyclic nucleotide-gated transmembrane channel with CNGA3.13 The subunit configuration of cone CNG channels is thought to be either 2A:2B or an equal 2A:2A configuration, although homomeric 4A channels can form in vitro.26,27 A mutant CNGB3 could therefore impair function by altering normal subunit ratios, interfering with subunit assembly, preventing heterotetramer insertion in the cell membrane, or affecting heterotetramer function after assembly. The transcriptome results showing increased Cngb3 expression in the cpfl10 mouse could be consistent with increased promoter induction in response to reduced CNGB3 protein.

Theoretically, a diminished ability of the mutant membrane channel to fully open or remain close could therefore impair hyperpolarization in the cone phototransduction cascade. The affected arginine residue is 1 of only 48 residues in human CNGB3 conserved across all vertebrates. Missense mutations account for 4% of all human CNGB3 achromatopsia mutations, but there are currently no reported human cases of mutation at this residue.11 The incidence of achromatopsia in the Pinegelp islander population is approximately 10% due to a CNGB3 missense mutation at a highly conserved residue: c.1304C>T; (p.S433F). The Cngb3<sup>cpfl10</sup> mouse model may provide a more appropriate preclinical model for developing genetic treatments for missense mutations in humans than the existing Cngb3<sup>−/−</sup> (Cngb3<sup>knockout</sup>) mouse model.

Furthermore, this model may be beneficial in further exploring and understanding any differences in treatment effect observed in subgroups of patients (missense versus truncation mutations) in the current CNGB3 phase I clinical trials in humans (NCT03001310 and NCT02599922). For example, CNGB3 gene therapy vectors might be more effective in cases of premature stop codons rather than missense mutations. Such a difference could be explored through a direct comparison of AAV in the different missense (cpfl10) and truncation mouse models of Cngb3 achromatopsia.
**FIGURE 5.** (A) The p.(R231H) missense mutation occurs at an arginine residue conserved across all vertebrates (within black box). (B) The equivalent arginine residue in Cnga3 is also highly conserved across vertebrates and other more distant species. (C) Structural modelling of Cngb3 shows the affected first residue of the first extracellular loop (red residue) and (D) in the context of the CNG heterotetramer. (E) The ratio of Cngb3 transcript expression normalized to Cnga3 transcript expression (dotted line) in cpfl10 (n = 5; PND 84) and 129S2/SvHsd control mice (n = 3; PND 80) as measured with qPCR. Values are mean ± SEM.
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