Early-Onset, Slow Progression of Cone Photoreceptor Dysfunction and Degeneration in CNG Channel Subunit CNGB3 Deficiency

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PURPOSE. To investigate the progression of cone dysfunction and degeneration in CNG channel subunit CNGB3 deficiency.

METHODS. Retinal structure and function in CNGB3−/− and wild-type (WT) mice were evaluated by electroretinography (ERG), lectin cytochemistry, and correlative Western blot analysis of cone-specific proteins. Cone and rod terminal integrity was assessed by electron microscopy and synaptic protein immunohistochemical distribution.

RESULTS. Cone ERG amplitudes (photopic b-wave) in CNGB3−/− mice were reduced to approximately 50% of WT levels by postnatal day 15, decreasing further to approximately 30% of WT levels by 1 month and to approximately 20% by 12 months of age. Rod ERG responses (scotopic a-wave) were not affected in CNGB3−/− mice. Average CNGB3−/− cone densities were approximately 80% of WT levels at 1 month and declined slowly thereafter to approximately 50% of WT levels by 12 months. Expression levels of M-opsin, cone transducin α-subunit, and cone arrestin in CNGB3−/− mice were reduced by 50% to 60% by 1 month and declined to 35% to 45% of WT levels by 9 months. In addition, cone opsin mislocalized to the outer nuclear layer and the outer plexiform layer in the CNGB3−/− retina. Cone and rod synaptic marker expression and terminal ultrastructure were normal in the CNGB3−/− retina.

CONCLUSIONS. These findings are consistent with an early-onset, slow progression of cone functional defects and cone loss in CNGB3−/− mice, with the cone signaling deficits arising from disrupted phototransduction and cone loss rather than from synaptic defects. (Invest Ophthalmol Vis Sci. 2011;52: 3557–3566) DOI:10.1167/iovs.10-6358

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The CNGB3 mouse line (on a C57BL/6N background) was generated by targeted deletion (Deltagen Inc., San Mateo, CA), as described previously. WT mice (C57BL/6) were purchased from Charles River Laboratories (Wilmington, MA). All mice were maintained under cyclic light (12-hour light/12-hour dark) conditions. Cage illumination was approximately 7 foot-candles during the light cycle. All experiments were approved by the local Institutional Animal Care and Use Committees (University of Oklahoma Health Sciences Center, Oklahoma City, OK) and conformed to the guidelines on the care and use of animals adopted by the Society for Neuroscience and the Association for Research in Vision and Ophthalmology (Rockville, MD).

Affinity-purified rabbit polyclonal antibodies against mouse M-opsin and cone arrestin (CAR) were provided by Cheryl Craft (University of Southern California, Keck School of Medicine). Rabbit polyclonal antibody against mouse S-opsin was provided by Muna Naash (University of Oklahoma Health Sciences Center). Rabbit polyclonal antibody against cone transducin α-subunit (Gnat2) was obtained from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). Monoclonal anti-actin antibody was purchased from Abcam, Inc. (Cambridge, MA). Mouse monoclonal antibody directed against C-terminal binding protein 2 (Grip2) was purchased from BD Transduction Laboratories (San Jose, CA). Rabbit polyclonal anti–complexin 3 was from Synaptic Systems (Göttingen, Germany). Biotinylated peanut agglutinin (PNA) and streptavidin-FITC were purchased from Vector Laboratories (Burlingame, CA) and Sigma-Aldrich (St. Louis, MO), respectively. Wheat germ agglutinin (WGA) and PNA conjugated to AlexaFlour-488 or -568, and fluorescent goat anti–mouse and goat anti–rabbit secondary antibodies conjugated to AlexaFlour-488, -568, or -647, were from Invitrogen-Molecular Probes (Carlsbad, CA). Horseradish peroxidase (HRP)-conjugated anti-rabbit or anti-mouse secondary antibodies were purchased from Kirkegaard & Perry Laboratories Inc. (Gaithersburg, MD).

Recording of Electroretinograms

Full-field ERG testing was carried out as described previously. Briefly, after overnight dark adaptation, animals were anesthetized by intra-peritoneal injection of 85 mg/kg ketamine and 14 mg/kg xylazine. ERGs were recorded using an LKC Technologies (Gaithersburg, MD) system. Potentials were recorded using a platinum wire contacting the corneal surface through a layer of 2.5% methylcellulose. For assessment of scotopic responses, a stimulus intensity of 1.89 log cd s/m² was presented to dark-adapted dilated mouse eyes in a Ganzfeld (GS-2000; Nicolet Instruments, Inc., Madison, WI). To evaluate photopic responses, mice were adapted to a 1.46 log cd s/m² light for 5 minutes, and then a light intensity of 1.89 log cd s/m² was given. Responses were differentially amplified, averaged, and stored using a signal averaging system (Compact-i, Nicolet Instruments, Inc.).

Eye Preparation, Immunohistochemistry, and Confocal Microscopy

Mouse retinal whole mounts or cross-sections were prepared for immunohistochemical analysis, as described previously. For whole mount preparations, eyes were enucleated, marked at the superior pole with green dye, and fixed in 4% formaldehyde (Polysciences, Inc., Warrington, PA) for 5 to 10 minutes. The cornea and lens were removed, and the eyes were fixed in 4% formaldehyde at 4°C overnight. The retina was then marked for orientation with a small cut left of the superior portion, and the retina was isolated. For retinal sections, mouse eyes were enucleated and fixed in 4% formaldehyde in 0.1 M sodium phosphate buffer, pH 7.4, at 4°C overnight. The superior portion of the cornea was marked for orientation before enucleation. Fixed eyes were then stored in 70% ethanol until processed and embedded in paraffin. Paraffin sections (5-μm thickness) passing vertically through the retina were prepared using a Leica (Wetzlar, Germany) microtome.

Immunohistochemical labeling of cones using anti-opsin antibodies and PNA was performed as described previously. Briefly, retinal whole mounts or sections were blocked in PBS containing 5% BSA and 0.5% Triton X-100 for 1 hour at room temperature. Primary antibody incubation (rabbit polyclonal anti–M-opsin, 1:500; rabbit polyclonal anti-S-opsin, 1:500) was performed at room temperature for 2 hours. After AlexaFluor-488 or -568 or FITC-conjugated secondary antibody incubation and rinses, slides were mounted and coverslipped. PNA immunohistochemistry was performed using biotinylated PNA (1:50) and streptavidin-FITC (1:200). Fluorescent signals were imaged using a fluorescence microscope (AX70; Olympus Corp., Center Valley, PA) with imaging software (QCapture; QIMaging Corp., Surrey, BC, Canada) or a confocal laser scanning microscope (IX81-FV500; Olympus, Melville, NY) (using excitation wavelengths of 543 nm for AlexaFluor-568 and 488 nm for FITC) with imaging software (FluoView; Olympus, Melville, NY). Fluorescence labeling intensities of different regions of the retinal sections (outer segment [OS], outer nuclear layer [ONL], and outer plexiform layer [OPL]) were analyzed using the intensity mapping feature of the software as described previously. Evaluation of cone density in retinal whole mounts was performed as described by Komeima et al. Briefly, images were taken using a 40× objective on an Olympus microscope approximately 1 mm from the optic nerve in the center of each quadrant. Image scale was calibrated, and cones were counted in four regions, each with dimensions of 125 μm × 125 μm (1.56 × 10⁴ μm²; 500 × 500 pixels), using imaging software (Image Pro 6; Media Cybernetics, Inc., Bethesda, MD). Averages of the counts from four regions of the quadrants were analyzed and graphed using graphing and statistics software (Prism; GraphPad Software, San Diego, CA).

To assess the integrity of photoreceptor synaptic terminals, paraffin sections prepared from WT and CNGB3 mouse eyes were triple labeled using methods similar to those described previously. Briefly, sections were deparaffinized and antigen was retrieved in citrate buffer and incubated in primary antibody overnight at 4°C. Mouse monoclonal antibody directed against CtBP2 (diluted 1:1000) was used to label synaptic ribbons. Rabbit polyclonal antibody against complexin 3 (diluted 1:1000) was used to selectively label cone terminals. Primary antibody was removed, and binding of primary antibodies was visualized using an appropriate combination of fluorescent goat anti–mouse and goat anti–rabbit secondary antibodies (diluted 1:200) and fluorescent conjugates of WGA or PNA (diluted 1:20–1:40) to visualize rod and cone terminal placement in the OPL and flat contacts onto cone terminals. After secondary antibody treatment, sections were rinsed and mounted using mounting medium plus DAPI (Prolong Gold; Invitrogen-Molecular Probes, Carlsbad, CA) to retard bleaching and visualize nuclei. Sections were imaged using a epifluorescence microscope (BX61-Wi; Olympus America, Center Valley, PA), a camera (ORCA-ER; Hamamatsu, Bridgewater, NJ), and software (Slidebook; Intelligent Imaging Innovations, Denver, CO). Figures were prepared by calibrating image scale, exporting images to image editing software (Photoshop; Adobe, Mountain View, CA), and adjusting brightness, contrast, and threshold to highlight specific labeling.

Retinal Membrane Preparation, SDS-PAGE, and Western Blot Analysis

Protein SDS-PAGE and Western blot analysis were performed as described previously. Briefly, retinas were homogenized in homogenization buffer (10 mM Tris-HCl, pH 7.4, 1 mM EDTA, 200 mM sucrose, 0.5% Triton X-100 for 1 hour at room temperature. Primary antibody incubation (rabbit polyclonal anti–opsin, 1:500; rabbit polyclonal anti–sorin, 1:500) was performed at room temperature for 2 hours. After AlexaFluor-488 or -568 or FITC-conjugated secondary antibody incubation and rinses, slides were mounted and coverslipped. PNA immunohistochemistry was performed using biotinylated PNA (1:50) and streptavidin-FITC (1:200). Fluorescent signals were imaged using a fluorescence microscope (AX70; Olympus Corp., Center Valley, PA) with imaging software (QCapture; QIMaging Corp., Surrey, BC, Canada) or a confocal laser scanning microscope (IX81-FV500; Olympus, Melville, NY) (using excitation wavelengths of 543 nm for AlexaFluor-568 and 488 nm for FITC) with imaging software (FluoView; Olympus, Melville, NY). Fluorescence labeling intensities of different regions of the retinal sections (outer segment [OS], outer nuclear layer [ONL], and outer plexiform layer [OPL]) were analyzed using the intensity mapping feature of the software as described previously. Evaluation of cone density in retinal whole mounts was performed as described by Komeima et al. Briefly, images were taken using a 40× objective on an Olympus microscope approximately 1 mm from the optic nerve in the center of each quadrant. Image scale was calibrated, and cones were counted in four regions, each with dimensions of 125 μm × 125 μm (1.56 × 10⁴ μm²; 500 × 500 pixels), using imaging software (Image Pro 6; Media Cybernetics, Inc., Bethesda, MD). Averages of the counts from four regions of the quadrants were analyzed and graphed using graphing and statistics software (Prism; GraphPad Software, San Diego, CA).

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Transmission Electron Microscopy

Mouse eye samples were prepared for transmission electron microscopy, as described previously. Briefly, mouse eyes were fixed in 4% formaldehyde in 0.1 M sodium phosphate buffer, pH 7.4, for 16 hours at 4°C and were transferred to PBS or 0.1 M sodium phosphate buffer, pH 7.4, containing 0.02% sodium azide, for storage until processing and embedding in plastic resin. Tissue sections were obtained from that in CNGB3 mice was approximately 40% of the level detected in WT mice at 1 month, whereas the response in CNGB3 mice was approximately 50% of the age-matched WT level (Fig. 1A). The difference between CNGB3 and WT mice was further increased at 1 month, with the response in CNGB3 mice at 3 months of age was not statistically different from that in CNGB3 mice at 1 month. However, a further progressive decrease in ERG b-wave amplitude was observed in CNGB3 mice after 6 months of age. Compared with the photopic b-wave amplitude of CNGB3 mice at 1 month, the b-wave amplitude was reduced by approximately 30%, 45%, and 55% at 12, 15, and 18 months, respectively. In contrast, no...
significant difference in b-wave amplitude was observed in WT mice up to 12 months of age (Fig. 1A). Figure 1B shows representative traces of photopic ERG recordings in CNGB3−/− and WT mice at 1, 6, and 12 months. Hence, cone dysfunction in CNGB3−/− mice showed early onset and was slowly progressive.

We also evaluated rod function in CNGB3−/− mice at different ages by scotopic ERG recordings (Figs. 1C–E). Although there was a progressive reduction in the scotopic a- and b-waves of both WT and CNGB3−/− mice with age, there was no significant difference between the amplitudes of the a-waves or b-waves of age-matched CNGB3−/− and WT mice (Figs. 1C, 1D). Figure 1E shows representative traces of scotopic ERG recordings in CNGB3−/− and WT mice at 1, 6, and 12 months.

**Early-Onset, Slow Progression of Cone Degeneration in CNGB3−/− Mice**

We previously showed cone loss in CNGB3−/− mice at 1 month of age.18 This included reduced cone density and increased photoreceptor apoptosis, as indicated by enhanced TUNEL labeling.18 We extended those findings by examining the progression of cone loss in CNGB3−/− mice. Cone density in CNGB3−/− and WT mice at different ages was examined by cone opsin staining and PNA labeling of retinal whole mounts and retinal cross-sections. We found that although cone density starts to decrease as early as 1 month, cone degeneration in CNGB3−/− mice progresses only slowly. Figure 2 shows immunofluorescence labeling of S-opsin on the retinal whole mounts prepared from WT and CNGB3−/− mice at 1, 4, 8, and 12 months (Figs. 2Aa and 2Bb, respectively) and the corresponding quantification results (Fig. 2B; images of inferior quadrant shown). Labeling of other quadrants was similar; see Supplementary Fig. S1, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.10-6358/-/DCSupplemental). Cone density in CNGB3−/− mice at 1 month was approximately 80% of the WT level. This number was reduced to approximately 50% at 12 months. M-opsin staining and PNA labeling on the retinal whole mounts showed a similar pattern (see Supplementary Fig. S2, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.10-6358/-/DCSupplemental). The early-onset, slow progression of cone degeneration was also shown by labeling of cones in retinal cross-sections. Figure 3 shows M-opsin labeling on retinal sections prepared from WT (Fig. 3a) and CNGB3−/− (Fig. 3b) mice at P15 and at 1, 3, 6, and 12 months.

Cone degeneration in CNGB3−/− mice was also investigated by examining the expression of cone-specific proteins. Western blot analysis was performed to examine the expression of M-opsin, Gnat2, and CAR using retinal protein extracts prepared from CNGB3−/− and WT mice. As shown in Figure 4A, expression of these cone-specific proteins was significantly reduced in CNGB3−/− mice compared with WT mice, even at 1 month of age. Densitometric analysis showed that M-opsin, Gnat2, and CAR levels in CNGB3−/− retinas were significantly reduced at 1 month and were further reduced at 9 months compared with the age-matched WT mice (Fig. 4B). These results also are consistent with an early-onset, slow progression of cone degeneration in CNGB3−/− mice.

Cone degeneration in CNGB3−/− mice shows an asymmetric pattern, with cones in the inferior portion of the retina degenerating faster than in the superior retina.29 We examined cone densities in the four different quadrants of retinal whole mounts prepared from CNGB3−/− mice to determine whether there was a topographic pattern of cone degeneration. Figure 5 shows M-opsin staining (Figs. 5Aa, WT; 5Ab, CNGB3−/−) and PNA labeling (Figs. 5Ba, WT; 5Bb, CNGB3−/−) on retinal whole mounts prepared from WT and CNGB3−/− mice at 1, 4, 8, and 12 months. Unpaired Student’s t-test was used to determine significance between age-matched CNGB3−/− and WT mice (P < 0.05).

**Figure 2.** Early-onset, slow progression of cone degeneration in CNGB3−/− mice evaluated by immunofluorescence labeling of S-opsin on retinal whole mounts. Immunofluorescence labeling was performed on retinal whole mounts prepared from CNGB3−/− and WT mice at 1, 4, 8, and 12 months. (A) Representative confocal images of S-opsin labeling in the inferior quadrants of the retinal whole mounts prepared from WT (a) and CNGB3−/− (b) mice. Scale bar, 10 μm. (B) Quantitative analysis of S-opsin labeling in the inferior quadrants of the retinal whole mounts. Data represent mean ± SD (n = 4–9 mice for each group). Unpaired Student’s t-test was used to determine significance between age-matched CNGB3−/− and WT mice (*P < 0.05).
mounts prepared from CNGB3−/− and WT mice at 8 months and the corresponding quantitative analysis (Fig. 5C). Cone density in CNGB3−/−/H11002 mice was approximately 72%, 70%, 78%, and 68% of the WT level for the superior, inferior, nasal, and temporal quadrants, respectively, as measured by M-opsin labeling (Fig. 5C, left), and was approximately 83%, 70%, 83%, and 78% of the WT level for the superior, inferior, nasal, and temporal quadrants, respectively, as measured by PNA labeling (Fig. 5C, right). Hence, the rate of cone degeneration in CNGB3−/−/H11002 mice did not differ significantly among the four quadrants.

Mislocalization of Cone Opsin

Disturbed trafficking of cone opsin was shown in CNGA3−/−/H11002 mice. We examined the localization of cone opsin in CNGB3−/−/H11002 mice. Similar to reported results in CNGA3−/−/H11002 mice, we detected substantial amounts of opsin in the inner segments, ONL, and OPL of CNGB3−/−/H11002 retina. This was particularly significant in young mice (P15). Figure 6A shows labeling for M-opsin and S-opsin on retinal cross-sections prepared from CNGB3−/−/H11002 (Fig. 6Aa) and WT (Fig. 6Ab) mice at different ages. Figure 6B shows the quantitative results of the fluorescence intensity of M-opsin labeling in the OS, ONL, and OPL of the CNGB3−/−/H11002 retina at P15 and P30, compared with that in the age-matched WT mice. Hence, CNGB3 deficiency, like CNGA3 deficiency, interferes with cone opsin trafficking to the outer segment.

Photoreceptor Terminals Are Unaffected by the Absence of CNGB3

To determine whether disruption of the synaptic terminals of the surviving cones might contribute to the decline in cone function noted in the ERG of the CNGB3−/−/H11002 retina, we performed triple labeling of retinas from WT and CNGB3−/−/H11002 mice for WGA, complexin III, and CtBP2 to visualize photoreceptor terminal distribution in the OPL, cone terminals, and synaptic ribbons, respectively (Fig. 7). The terminals of surviving cones and rods in the CNGB3−/−/H11002 retina showed normal placement, continued to express appropriate synaptic proteins, and retained their synaptic ribbons. Similarly, surviving cone terminals also retained their flat synaptic contacts, as identified by PNA labeling (not shown). Rod and cone terminals showed normal terminal ultrastructure in the terminals of rods and cones (Fig. 8). Both rod and cone terminals in the WT and CNGB3−/−/H11002 retina showed ribbon synaptic complexes with the triadic organization and ultrastructural features. As appropriate, cone terminals in the WT and CNGB3−/−/H11002 retina had multiple ribbon complexes and formed flat contacts with OFF-cone bipolar cell dendrites. Together, these results suggest that decreased cone signals in the ERG arose primarily from dis-
ruptured cone phototransduction and cone degeneration rather than from synaptic disruption.

**DISCUSSION**

The high correlation of mutations in CNGB3 with human cone diseases implies a critical role for this protein in human cone function. We have previously shown impaired cone function and cone degeneration in CNGB3/−/− mice and provided the first experimental evidence supporting the link between mutations in CNGB3 and human cone diseases. This study demonstrates an early-onset (by P15, when the mouse eye opens), slow progression of cone dysfunction and degeneration arising from CNGB3 deficiency. The phenotype in CNGB3/−/− mice is similar to the clinical symptoms of patients with CNGB3 mutations. It is important to note a phenotypic difference between human patients with achromatopsia, in whom there is an almost total lack of cone function from birth, and CNGB3/−/− mice, in which some residual cone function persists as indicated by the ERG. The reason for this difference remains to be identified but may be related to the heterogenetic causes of achromatopsia (three other genes have been identified in achromatopsia patients in addition to CNGB3) and species differences. A more thorough clinical phenotype/genetic disorder relationship study may help to better address this question. It is also worth noting that the residual cone function in CNGB3/−/− mice was not comparable to the percentage of cones that remained. The cone ERG response at 1 month was only approximately 30% of the WT level, whereas approximately 80% of cones remained at this age. Similarly, the cone response at 12 months was only approximately 20% of the WT level, whereas more than 50% of cones remained at this age in the CNG3B/−/− retina. In contrast, the reduction of cone response (approximately 70%–80%) did approximate the reduction of cone proteins (i.e., M-opsin, Gnat2, and CAR, which were reduced by approximately 50%–70%; see Fig. 4). Thus, although ample cones remained in the CNGB3-deficient retina as late as 12 months of age, the functional response and expression of phototransduction proteins of the surviving cones were seriously impaired. Indeed, we cannot exclude the possibility that an altered protein expression in cones also could contribute to the functional deficits. Furthermore, our studies showed that the terminals of cones and rods in the CNGB3/−/− retina had normal placement in the OPL and retained their appropriate synaptic components and ultrastructural organization. These findings suggest that the cone-signaling deficits observed arose from phototransduction deficits and progressive cone loss rather than from synaptic defects. Consistent with this finding, most of the surviving cones in the CNGA3/−/− retina have morphologically normal synapses.
and mice lacking both functional cones and rods (CNGA3−/−/Rh11002−/−/Rh11002−/−) have structurally normal synaptic contacts until the time of complete loss of photoreceptors.30

Cone degeneration has been shown in a variety of mouse lines that are deficient in cone-specific or cone-dominant proteins, such as cpfl1 (with cone PDE6C deficiency),31 RetGC1−/−/RetGC1−/−,32–34 and GCAP−/− (and GCAP L151F, Y99C mutant) mice.35 Mutations in these genes in humans are associated with cone defects36,3738 However, the retinal pathogenesis and disease progression in these mouse lines differ substantially from one another. Cone degeneration in cpfl1 mice has an early-onset, rapid progression; by 2 to 3 months, most of the cones (90%) are lost.31 In contrast, cone degeneration in RetGC1−/− mice has a relatively late onset, with normal numbers of cone cells at 4 and 5 weeks of age and then gradually decreasing by 6 months.34 This work and the study by Michalakis et al.29 showed that cone degeneration in CNG channel deficiency is of early-onset, slow progression. Of note, there is no clear cone degeneration in the mouse line with Gnat2 deficiency,39 although cone transduction is completely blocked in these mice. Thus, although deficiency of these molecules—which are involved directly or indirectly in the phototransduction cascade—leads to a common consequence (i.e., blockade or impairment of cone phototransduction), the progression of cone degeneration varies. The underlying mechanisms, therefore, might be expected to be distinct. It is also worth noting that even with deficiencies of the same gene, phenotypes can vary among species. A slow progression of cone degeneration with late onset is observed in canines with CNGB3 mutation or CNGB3 deficiency. In contrast to what we have found in the present study in mice, cone density does not start to decrease in CNGB3-deficient dogs until nearly 2 years of age (G. Aguirre, University of Pennsylvania, personal communication).

Little is known about the mechanism of cone degeneration in CNG channel deficiency. Cone degeneration in CNGB3−/− and CNGA3−/− mice is likely attributable to a decrease or loss of the functional channels and subsequent impairment or loss of cone phototransduction. We18 and others29 have shown that cone death occurs in CNG channel deficiency primarily by way of an apoptotic mechanism, but the molecular pathway connecting CNG channel deficiency to cone apoptotic death...
remains to be elucidated. It is recognized that studying the mechanism of cone degeneration in a rod-dominant mammalian retina is challenging because cones account for only 3% to 5% of the photoreceptor population. We have recently shown that the cone-dominant Nrl⁻/⁻ mouse line (deficiency of retinal leucine zipper transcription factor) is a valuable model for the study of cone CNG channel function and structure.17 The Nrl⁻/⁻ retina expresses abundant cone CNG channels and lacks expression of the rod CNG channel. Hence, the mouse lines with CNG channel deficiency in a cone-dominant retina (i.e., CNGA3⁻/⁻/Nrl⁻/⁻ and CNGB3⁻/⁻/Nrl⁻/⁻ mice) could be valuable models in which to study the mechanism of cone degeneration. One way to explore the triggering factors involved in cone degeneration is to examine the direct/immediate cellular consequences of CNGB3 deficiency. One strong possibility is that the cellular consequences of CNG channel deficiency (i.e., a lowered intracellular Ca²⁺ concentration and the subsequent accumulation of cGMP) may trigger cone death. Indeed, we observed a dramatically increased level of cGMP in CNGB3⁻/⁻/Nrl⁻/⁻ mice at a young age (P10-P30 days) that correlated with cone apoptosis (our unpublished observation). This observation suggests a potential role for cGMP accumulation in the early-onset cone degeneration in CNG channel deficiency. Cone degeneration in CNGB3⁻/⁻ mice might also be associated with a role of CNGB3 in outer segment morphogenesis and integrity, similar to its rod counterpart CNGB1, which is known to play a role in rod outer segment disc morphogenesis.40 Our work showing outer segment disorganization41 in some cones of CNGB3⁻/⁻ mice is consistent with this finding. The early onset of cone degeneration in CNG channel deficiency may be related to the mislocalization of cone opsin. Mislocalization of cone opsin in CNGB3⁻/⁻ mice (this study) and CNGA3⁻/⁻/Nrl⁻/⁻ mice was more profound in developing and young animals and correlated with cone apoptosis at the same age. Mislocalization of cone opsin in young and old animals could be useful models for these studies. One potentially useful application of these models would be to identify gene/protein expression profiles using microarray/proteomic analyses with retinal preparations from young and old mice compared with age-matched Nrl⁻/⁻ controls.

Mislocalization of cone opsin is observed in both CNGA3⁻/⁻ and CNGB3⁻/⁻ mice, suggesting that it is the functional channel complex, rather than the individual subunit, that is critical for the normal trafficking and outer segment targeting of cone opsin. The molecular details of cone opsin trafficking, however, are not yet known. In rod photoreceptors, rhodopsin is synthesized in the inner segment and...
then is actively transported through the connecting ciliost into the outer segment. An equivalent mechanism may exist in cones and could be impaired in CNG channel deficiency. Nevertheless, although the channel-deficient cones are able to synthesize opsins, they failed to transport or retain the protein properly into the outer segments. Mislocalization of cone opsin could have two potential consequences. First, it might be associated (or partially associated) with cellular stress and apoptosis. Second, the mislocalized opsin might be degraded more easily than the normal localized opsin. Indeed, the expression level of M-opsin in the CNGB3 knockout mouse is significantly reduced (see Fig. 4). We have shown previously that the mRNA level for cone opsin in CNGB3 mice is comparable to that of WT mice, suggesting that the reduction in the opsin protein level is likely due to enhanced degradation rather than decreased synthesis.

In summary, this work provides experimental evidence showing the nature of early-onset, slow progression of cone dysfunction and degeneration in CNGB3 deficiency. The phenotype of CNGB3 mice is similar in many respects to the symptoms in human patients with mutations of CNGB3. Thus, the CNGB3-deficient mouse line could serve as a valuable model in which to study the retinal pathogenesis of CNG channel deficiency and the mechanism of cone degeneration.

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