Ciliary genes \textit{arl13b}, \textit{ahi1} and \textit{cc2d2a} differentially modify expression of visual acuity phenotypes but do not enhance retinal degeneration due to mutation of \textit{cep290} in zebrafish

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Abstract

Mutations in the gene Centrosomal Protein 290 kDa (\textit{CEP290}) result in multiple ciliopathies ranging from the neonatal lethal disorder Meckel-Gruber Syndrome to multi-systemic disorders such as Joubert Syndrome and Bardet-Biedl Syndrome to nonsyndromic diseases like Leber Congenital Amaurosis (LCA) and retinitis pigmentosa. Results from model organisms and human genetics studies, have suggest that mutations in genes encoding protein components of the transition zone (TZ) and other cilia-associated proteins can function as genetic modifiers and be a source for \textit{CEP290} pleiotropy. We investigated the zebrafish \textit{cep290} \textit{fh297/fh297} mutant, which encodes a nonsense mutation (p.Q1217\textsuperscript{*}). This mutant is viable as adults, exhibits scoliosis, and undergoes a slow, progressive cone degeneration. The \textit{cep290} \textit{fh297/fh297} mutants showed partial mislocalization of the transmembrane protein rhodopsin but not of the prenylated proteins rhodopsin kinase (GRK1) or the rod transducin subunit GNB1. Surprisingly, photoreceptor degeneration did not trigger proliferation of Müller glia, but proliferation of rod progenitors in the outer nuclear layer was significantly increased. To determine if heterozygous mutations in other cilia genes could exacerbate retinal degeneration, we bred \textit{cep290} \textit{fh297/m297} mutants to \textit{arl13b}, \textit{ahi1}, and \textit{cc2d2a} mutant zebrafish lines. While \textit{cep290} \textit{fh297/m297} mutants lacking a single allele of these genes did not exhibit accelerated photoreceptor degeneration, loss of one alleles of \textit{arl13b} or \textit{ahi1} reduced visual performance in optokinetic response assays at 5 days post fertilization. Our results indicate that the \textit{cep290} \textit{fh297/m297} mutant is a useful model to study the role of genetic modifiers on photoreceptor degeneration in zebrafish and to explore how progressive photoreceptor degeneration influences regeneration in adult zebrafish.
Introduction

Ciliopathies refer to a group of recessive disorders stemming from defects in the biogenesis, structure or function of cilia [1]. These disorders exhibit both clinical and genetic heterogeneity [2], with mutations in dozens of genes resulting in a spectrum of diseases sharing overlapping symptoms. Clinically, ciliopathies can manifest as non-syndromic disorders, such as in Leber Congenital Amaurosis (LCA; OMIM: 204000), which is an inherited form of childhood retinal dystrophy, to more pleiotropic diseases, such as Joubert Syndrome (JTBS; OMIM 213300), Meckel Syndrome (MKS; OMIM 249000), Bardet-Biedl Syndrome (BBS; OMIM 209900), and Nephronophthisis (NPHP; OMIM 256100), each of which impact unique combinations of organ systems [3].

Mutations in the gene for Centrosomal Protein 290 (CEP290) cause JTBS, MKS, BBS, and NPHP [3], but also account for 15–25% of cases of isolated blindness in LCA with no associated systemic disease [4, 5]. Most CEP290 lesions in humans are stop codons that result from frameshift and nonsense mutations occurring throughout the gene, whereas pathogenic missense mutations are rare [6]. Despite the identification of more than 130 mutations in human CEP290, efforts to establish obvious genotype-phenotype correlations have been unsuccessful [6]. CEP290 mutations are strongly associated with retinal dystrophy and photoreceptor degeneration is one of the most common symptoms of ciliopathies [7]. The most frequent CEP290 mutation in LCA is a deep intronic mutation (c.2991+1655 A>G) that activates a cryptic splice site and creates a stop codon, resulting in early termination of the protein [4]. In a recent study of LCA patients with CEP290 mutations, visual acuity varied considerably although most patients had significant vision loss and undetectable electroretinograms (ERGs) regardless of genotype [8]. In spite of the severe loss of vision, multiple studies have reported that cone photoreceptors persist within the central retina of CEP290-LCA patients [8–11], suggesting that a window of opportunity may exist for therapeutic intervention.

In humans, CEP290 encodes a 2479 amino acid protein (~290 kDa) that can localize to the basal body [12] and/or the transition zone (TZ) of cilia [13, 14] in a tissue-specific manner. The TZ refers to the most proximal region of the ciliary axoneme, immediately distal to the basal body. The connecting cilium in vertebrate photoreceptors is homologous to the TZ of a prototypic primary cilium [15]. The TZ is believed to function as a ciliary gate that regulates protein entry and exit to the cilium. Defects in the ciliary gate may result in abnormal accumulation of non-outer segment proteins within the outer segment [16] and/or disrupt normal protein delivery to the outer segment. Work from C. elegans have proposed roles for Cep290 ranging from cell adhesion to TZ assembly [13, 17, 18], but in vivo studies in vertebrate models have not fully elucidated a role for Cep290 or explained the variability in photoreceptor phenotypes [14, 19–21]. Abyssinian cats exhibit a high degree of inherited retinal degeneration due to the rdAc allele in the feline Cep290 gene and this rdAc allele can also be found at elevated frequencies in several other cat breeds [22, 23]. In rodents, the rd16 allele reflects an in-frame deletion of Cep290 that leads to rapid photoreceptor degeneration [20]. Although a targeted gene knockout of Cep290 causes embryonic lethality in mice [14], truncating nonsense mutations in humans can often result in attenuated pathologies that range from multisystem dysfunction to mild retinal disease. Indeed, LCA patients with two truncating CEP290 mutations can sometimes maintain photoreceptor architecture and retain limited visual acuity [8, 9, 24]. These unexpectedly mild phenotypes have recently been attributed to basal exon skipping and nonsense-mediated alternative splicing of the CEP290 mRNA [24, 25]. Nevertheless, patients with identical genotypes can still exhibit very different retinal phenotypes [9].

One hypothesis to explain the variable phenotypic expression is the effects of mutations in second-site modifiers [26–29]. Genetic [26–28] and biochemical studies [18, 30] in C. elegans
and cultured mammalian cells have identified two molecular complexes within the TZ, termed the NPHP and MKS modules. The proteins Cc2d2a, Ah1, Mks1, and at least 5 other proteins form the MKS module [28], while the NPHP module consists of Nphp1 and Nphp4 [31], although proteomic studies suggest additional factors likely exist [30]. Homozygous mutations in genes from both an MKS and NPHP module severely disrupt cilia formation [27, 28, 31, 32]. These genetic interactions, however, reflect phenotypes resulting from double homozygous mutants. In humans, the frequency of pathogenic alleles in cilia genes is low and a more realistic scenario is that heterozygous mutant alleles in one cilia gene may influence phenotypic expression resulting from homozygous mutations in CEP290. Heterozygous missense alleles of the AHI1 gene were identified in LCA patients with severe neurological involvement, suggesting that alleles of AHI1 may influence phenotypic expression [33].

In this study, we evaluated the zebrafish cep290fh297/fh297 mutant in an effort to test ah1, cc2d2a, and arl13b as potential genetic modifiers of retinal degeneration. We report that the cep290fh297/fh297 mutant shows progressive and predominant cone degeneration. We found that the phenotype observed in these mutants was not the consequence of nonsense-associated alternative-splicing, a phenomenon hypothesized to explain phenotypic variation in humans [37]. We report that heterozygous mutations in ah1 and arl13b, were associated with decreased visual acuity, whereas the absence of one allele of cc2d2a had no effect on visual acuity. Retinal degeneration in the cep290fh297/fh297 mutant was not exacerbated by heterozygosity of any of these genes. Furthermore, the mild phenotypes observed in cep290fh297/fh297 mutants was not due to retinal regeneration These data demonstrate a role for Cep290 in cone survival in zebrafish and provide a foundation for future analysis of potential modifier genes of cep290-associated retinal degeneration.

Materials and methods

Zebrafish husbandry

Adult zebrafish were maintained and raised on an Aquatic Habitats recirculating water system (Pentair; Apopka, FL, USA) in a 14:10 hr light-dark cycle with ambient room lighting. Larvae used for optokinetic response measurements were kept in the laboratory in a 28.5˚C incubator with a clear glass door with ambient room lighting. The Cleveland Clinic Institutional Animal Care and Use Committee (IACUC) approved all experimental procedures (Protocol number: 2018–1980). The cep290fh297/fh297 mutant was identified by the zebrafish TILLING consortium and was a gift of Dr. Cecilia Moens (Fred Hutchinson Cancer Center, Seattle, WA. USA). The line was outcrossed to wild-type animals for at least five generations to reduce the presence of additional background ENU mutations. Animals were anesthetized with tricaine and euthanized by immersion in ice-water.

Sequencing

Using sequence data from Ensembl (http://useast.ensembl.org/Danio_rerio/Info/Index; transcript: cep290-202), primers were designed to span exon 29 (5′-GTCTGATGAAAGGCCCTGA-3′ and 5′-CCTCCAAGCCTTCAAGCCTTT-3′) for the cep290fh297 allele. Samples were
sequenced at the Genomics Core of the Cleveland Clinic Lerner Research Institute using the high-throughput, 96-capillary ABI 3730x/DNA Analyzer.

Genotyping

**cDNA extraction.** Tail-clips from embryos or adults were placed in 0.5 ml individual tubes and 25 μL (embryos) or 50 μL (adults) of lysis buffer (50 mM Tris pH 8.5, 5 mM EDTA, 100 mM NaCl, 0.4% SDS, 100 μg/mL proteinase K) was added to each tube and then incubated at 60°C for 4 hrs (embryos) to overnight (adults). Samples were then diluted 1:10 in nuclease-free water and heat inactivated at 95°C for 5 min.

**PCR and high resolution melt analysis (HRMA).** HRMA primers targeting *cep290* exon 29 for the *cep290*fh297 allele (5’ – ACAAACACAGTCTGCAGAAACTGGACGCG – 3’ and 5’ – CTGCTGTTGCTCATCCAG TT – 3’) were designed flanking the point mutation. PCR products were 95 bp. High-resolution melt curve analysis was performed using Bio-Rad Precision Melt reagent in 8μl reactions with a CFX96 Touch Real-Time PCR Detection System (Bio-Rad, Hercules, CA, USA) at standard cycling conditions. Melt curves were analyzed using the Precision Melt Analysis Software version 1.2 (Bio-Rad, Hercules, CA, USA).

**Micro-computed tomography (μCT)**

Adult zebrafish were euthanized and fixed in 4% paraformaldehyde (in 1X PBS) overnight at 4°C. Specimens were washed in 1X PBS and immersed in 70% ethanol in a 15 mL conical tube. Samples were scanned with an Explore Locus RS (GE Medical Systems, London, Ontario, Canada) at 45 μm. Images were analyzed and reconstructed using MicroView software version 2.5.0–3768 (Parallax Innovations Inc.; Ilderton, Ontario, Canada).

**Light and electron microscopy**

Light-adapted larvae were bisected through the swim bladder, and heads were prepared for transmission electron microscopy. Tails were used for genomic DNA extraction and genotyping as described above. For adult animals, enucleations were performed at the designated time points and samples prepared for transmission electron microscopy. Briefly, the eyes were enucleated from light-adapted animals and the anterior segment was dissected away in primary fixative (0.08M cacodylate buffer containing 2% paraformaldehyde and 2% glutaraldehyde). The tissue was fixed for 1 hr at room temperature in primary fixative and then washed with cacodylate buffer and post-fixed in 1% osmium tetroxide for 1 hr at 4°C. Samples were washed again and then dehydrated in a graded methanol series before embedding them in Embed-812/DER736 (Electron Microscopy Sciences; Hatfield, PA, USA), using acetonitrile as a transition solvent. Semi-thin sections were made with a Leica EM UC7 ultramicrotome (Leica Microsystems; GmbH Vienna, Austria), stained with Toluidine Blue, and imaged with a Zeiss Axio Imager.Z2 (Carl Zeiss Microscopy, Thornwood, NY, USA). Ultrathin sections were stained with uranyl acetate and lead citrate following standard procedures, and electron microscopy was performed on a Tecnai G2 Spirit BioTWIN 20–120 kV digital electron microscope (FEI Company; Hillsboro, OR, USA). Micrographs were acquired with a Gatan image filter and an Orius 832 CCD Camera (Gatan, Inc.; Pleasanton, CA, USA).

**Optokinetic response (OKR)**

OKR measurements on 5–6 dpf larvae were conducted between 12–6 pm using the Visio-Tracker system (VisioTracker 302060 Series, TSE Systems, GmbH Bad Homburg, Germany). Contrast sensitivity was assessed as described previously [38, 39]. For the spatial frequency...
response function [39, 40], the contrast was held constant at 70% and we tested stimuli of 0.02, 0.04, 0.06, 0.08, 0.12, and 0.16 cycles/degree by first increasing and then decreasing the frequency. Each spatial frequency stimulus was presented for 3 seconds before reversing direction for another 3 seconds to minimize saccade frequency. All OKR stimuli were presented with a constant angular velocity of 7.5 degrees per second. The genotypes of individual larvae were confirmed following OKR tests.

**Immunohistochemistry and fluorescence imaging**

Larvae were fixed for 2 hrs. at 4 °C. Adult eyes were fixed at the designated time points. Fixation protocols varied depending on the primary antibodies being used. For Zpr-1, Zpr3, GRK1, and GNB1, samples were fixed in 4% paraformaldehyde in 0.8X PBS at 4 °C overnight. For peanut agglutinin (PNA) and acetylated tubulin staining, heads were fixed in 4% paraformaldehyde in 0.8X PBS at 4 °C for a maximum of 2 hrs. All samples were cryoprotected in 30% sucrose overnight. Cryosections (10 μM) were cut and dried at room temperature overnight. Blocking solution (1% BSA, 5% normal goat serum, 0.2% Triton-X-100, 0.1% Tween-20 in 1X PBS) was applied for 2 hrs in a dark, humidified chamber. Primary antibodies were diluted in blocking solution as follow: Zpr-1 and Zpr-3 (1:200; Zebrafish International Resource Center, University of Oregon, Eugene, OR, USA), GNB1 (1:100; Abgent AP5036a), GRK1 (1:50; Abclonal A6497), and acetylated-α-tubulin (1:5000; Sigma 6-11-B1). Conjugated secondary antibodies were purchased from Invitrogen Life Technologies (Carlsbad, CA, USA) and used at 1:500 dilutions and 4,6-diamidino-2-phenylenedole (DAPI; 1:1000) was used to label nuclei. Optical sections were obtained with a Zeiss Axio Imager.Z2 fluorescent microscope fitted with the Apotome.2 for structured illumination (Carl Zeiss Microscopy, Thornhill, NY, USA). ImageJ was used to create image panels. Figures were assembled in Adobe Photoshop. To quantify cone outer segment density, the number of PNA-positive outer segments was determined from images of transverse sections of dorsal retinas. The distance of retina measured in each section was determined using ImageJ and density was calculated as the number of cone outer segments per 50 μm of retina. Each data point represented a minimum of one section from a distinct retina. The total number of fish used per experiment are noted in the figures. To quantify rhodopsin mislocalization, ImageJ was used to measure the integrated fluorescence density (IFD) across a region of interest (ROI) of defined area that was placed in the rod outer segments (ROS; proper localization), inner segment/outer nuclear layer (IS/ONL; mislocalized) or the inner nuclear layer (INL; background fluorescence). Corrected fluorescence intensities were calculated by subtracting the background fluorescence. The total rhodopsin fluorescence was calculated as the sum of the IFD from the ROS and IS/ONL. The percentage of mislocalized rhodopsin was calculated as the IFD from the IS/ONL (numerator) per total rhodopsin (denominator).

**RT-PCR**

For traditional RT-PCR, total RNA was extracted from pooled wild-type larvae at 5 dpf for positive control (tunicamycin), and from 4 isolated retinas from wild-type and cep290fh297/fh297 mutants at the designated time points using TRIzol according to standard protocols. cDNA was reverse transcribed using SuperScript II reverse transcriptase and random hexamers according to the manufacturer’s instructions (ThermoFisher Scientific, Waltham, MA, USA). RT-PCR was performed according to standard protocols and cycling conditions.

### Exon skipping

cDNA from wild-type and cep290fh297/fh297 mutants was obtained as described above. Primers were designed to encompass the mutated exon as follow: primers targeting exons 27–32 for cep290fh297/fh297 (5′–AGAATCACTGAAGTTGAGAAAACAG– 3′ and
TTCTTTTCTTTTAGCTTCTCTTCC– 3’ with products sizes– 1040 bp when mutated exon is included and 593 when mutated exon is skipped.

**Droplet digital PCR.** RNA was isolated from whole eyes of 6-month old cep290 mutant and sibling control animals (n = 9 per group) with Trizol (ThermoFisher 15596026). Reverse transcription using 1 microgram RNA was performed with the iScript cDNA Synthesis kit (Bio-Rad 1708891). Intron-spanning primers and probes for cep290 and ef1a1l1 were designed with Primer3Plus (http://primer3plus.com). Sequences are as follows: cep290F – ACACCGTCACTCAGCTGAAG; cep290R – CTGGCAAGACCTTCGTCAGT; cep290probe(FAM)–ACGTCCCTGTGGAAGCGACC; ef1a1l1F – CGTCTGCCACTTCAGGATGT; ef1a1l1R – CCCAGCCTTCAGATGTTCCAG; ef1a1l1probe(HEX)–ACTGTGCGCTGTGGGACGTGT. Multiplexed PCR reactions using 100 ng cDNA were prepared with the ddPCR supermix for probes (No dUTP, Bio-Rad 1863024) and fractionated into >20,000 droplets using the Bio-Rad QX200 droplet generator with droplet generation oil for probes (Bio-Rad 1863005). PCR cycling was performed using a 60 degree C annealing temperature, and droplet signal was detected with a QX200 droplet reader (Bio-Rad). Target copy number was determined with QuantaSoft Analysis Pro software (Bio-Rad) after droplets were manually thresholded relative to no-template control reactions.

**Statistics**

Graphs were generated using Prism6 (GraphPad Software; San Diego, USA). Statistical analyses were performed using a one-way ANOVA with a Multiple Comparisons test and Tukey’s correction or 2-way ANOVA with a Multiple Comparisons test and Sidak corrections. For all tests, P-values less than 0.05 were considered significant.

**Results**

**Identification of a nonsense mutation in zebrafish cep290 gene**

In zebrafish, the primary cep290 transcript (RefSeq: NM_001168267) encodes a 2439 amino acid protein. There is no known evidence of endogenous alternative splicing that would affect the coding region of the cep290 gene. The cep290fh297 allele was identified by the Zebrafish TILLING Consortium. This C>T transition mutation results in a stop codon (p.Gln1217X) downstream of the Cc2d2a binding domain [12] and upstream of the putative Rab8a binding domain. This mutation was predicted to truncate the protein by almost half (Fig 1A) and is near a similar mutation in humans (p.Gln1265X) that associated with LCA and JBTS (https://cep290base.cmgg.be/). We confirmed the mutation by direct sequencing (Fig 1B). To date, no parologue to cep290 has been reported in zebrafish. To assess the impact of the fh297 allele on gene expression, cep290 mRNA was quantified by digital droplet PCR (ddPCR). In adult animals, retinal expression of cep290 mRNA was reduced by 55% in mutants compared to expression in wild-type siblings (Fig 1C). Efforts to measure Cep290 protein by western blot were unsuccessful, despite multiple attempts with both commercial [41] and custom designed antibodies [42]. In fibroblasts derived from an LCA patient with the c.2991+1665A>G mutation, wild-type CEP290 transcripts were similarly reduced by ~60%, which resulted in a corresponding reduction in protein levels by ~80% [43].

At 5 days post fertilization (dpf) cep290fh297/fh297 mutants exhibited a sigmoidal tail curvature and did not yet have an inflated swim bladder (Fig 1D). These phenotypes were not fully penetrant with only 29% of mutant larvae exhibiting such characteristics (17 of 58 confirmed cep290fh297/fh297 mutants). Furthermore, these phenotypes were similar to, but distinct from phenotypes of other zebrafish mutants with defective cilia formation, such as ifi88 or cc2d2a mutants, which exhibit a ventral axis curvature [40, 46, 47]. All cep290fh297/fh297 mutants exhibited normal otolith numbers and only 6.8% (4 of 58 confirmed cep290fh297/fh297 mutants)
developed kidney cysts by 5 dpf. Although cep290fh297/fh297 mutants routinely survived to adulthood in Mendelian ratios, approximately 17% fewer cep290fh297/fh297 mutants were present at 12 months of age than would be expected (25 out of 120 total fish). The cep290fh297/fh297 mutants were unable to breed and 100% of the mutants exhibit a severe scoliosis (Fig 1E and 1F), a phenotype previously linked to defective cilia [48] and a pathology has been reported in a subset of Joubert Syndrome patients with CEP290 mutations [49]. The abnormal spinal curvature was also assessed by micro-computed tomography and revealed a significant

https://doi.org/10.1371/journal.pone.0213960.g001

Fig 1. Genetic mapping and identification of cep290 mutant allele. (A) Schematic structure of zebrafish Cep290 illustrating the location of predicted protein domains. Domain structure is based on prior results [12, 44, 45]. The cep290fh297 allele generates a premature stop codon at amino acid 1217. (B) Chromatograms of Sanger sequencing reactions of cDNAs from wild-type and homozygous cep290fh297/fh297 mutant confirming the C>T replacement. (C) Quantification of cep290 mRNA in 6 month old wild-type and mutant retinas by digital droplet PCR (ddPCR). (D) Lateral views of representative wild-type (top), heterozygous (middle), and homozygous (bottom) mutants at 5 dpf. At larval stage 30% of cep290fh297/fh297 mutant animals show ventral tail curvature. (E) Lateral view of 7 month old wild-type, heterozygous and a representative cep290fh297/fh297 mutant animals showing distorted vertebral column. At adult stage 100% of the homozygous mutant animals show scoliosis of the vertebral column. (F) Representative micro-CT-generated images of lateral (top) and dorsal (bottom) views of adult wild-type and cep290fh297/fh297 mutants. Images demonstrate that spinal curvature deviates within the dorsal/ventral plane as well as curves laterally (arrows). (G) Forward (F1) and reverse (R1) PCR primers were designed to bind sequences in exons 27 and 32 in order to flank exon 29 harboring the mutant cep290fh297 allele. Exon skipping would result in a truncated PCR product of 593 bp, while retention of exon 29 would result in a full-length 1040 bp product. (G, bottom) Results of PCR reactions from cDNAs generated from wild-type and samples of two individual mutants resulted in full-length products of 1040 bp. 100 bp ladder shown in lane 1. (H) Optokinetic response (OKR) contrast response function of 5 dpf wild-type larvae (n = 11; open circles) and cep290fh297/fh297 mutants (n = 26; closed circles). No significant differences were found. (I) OKR spatial frequency results for wild-type larvae (n = 13) and cep290fh297/fh297 mutants (n = 15). Error bars = s.e.m.
deviation in spinal curvature that was pronounced in both the dorsal-ventral axis as well as a lateral curvature (Fig 1F).

As the \textit{cep290}^{fh297} allele encodes a nonsense mutation, we were curious about the relatively mild phenotype compared to the \textit{Cep290} mouse knockout models [14]. A recent hypothesis proposed that if nonsense mutations occur in exons that begin and end in the same reading frame, those exons can be preferentially skipped in a process known as “nonsense-induced alternative splicing” [50]. These alternatively spliced transcripts can escape nonsense-mediated decay and produce near-full-length protein. Such phenomenon were reported to occur in Leber Congenital Amaurosis and Senor-Løken Syndrome patients with mutations in \textit{CEP290} [25, 37]. The \textit{cep290}^{fh297} allele is a nonsense mutation occurring in exon 29 and exon skipping would maintain the reading frame. We performed RT-PCR on cDNA from adult retinas from wild-type and \textit{cep290} mutant retinas and used primers that spanned the mutated exon. We showed that the mutant exon was present in all detectable transcripts, indicating that nonsense-mediated alternative splicing did not occur for this mutation in zebrafish (Fig 1G).

**Functional vision is preserved in \textit{cep290} mutant larvae**

As humans with \textit{CEP290} mutations report variable loss of visual function [9], we asked whether the visual performance of zebrafish \textit{cep290} mutants was also compromised. Visual acuity is a measure of the spatial resolving power of the visual system and is mainly driven by cones [51, 52]. Larval zebrafish visual function can be readily assessed using the optokinetic response (OKR) assay by presenting larvae with a moving grating stimuli that varies in either contrast or spatial frequency [39]. Detecting contrast differences of a stimulus presented at fixed spatial and temporal frequencies is a general method of testing function vision, while detecting the changes in spatial frequency of a stimulus at a fixed contrast under bright illumination assesses cone acuity. Because larval zebrafish rely exclusively on cone function at 5–6 dpf [53, 54], all recordings were done under photopic conditions [55]. We measured the OKR gain, which is defined by the ratio between stimulus velocity and eye velocity [38–40], of wild-type, and \textit{cep290}^{fh297/fh297} mutants between 5–6 dpf using established parameters [39, 56] and reproduced by our laboratory [38, 40]. Wild-type larvae (n = 12) showed a linear relationship between gain and the logarithm of contrast (Fig 1H). Interestingly, the \textit{cep290}^{fh297/fh297} mutants (n = 26) exhibited normal OKR responses to changes in stimulus contrast and spatial frequency (Fig 1H and 1I).

**Photoreceptor degeneration in \textit{cep290}^{fh297/fh297} mutants**

We next examined the retinal anatomy of wild-type and \textit{cep290}^{fh297/fh297} mutants by light microscopy at 5 dpf, 3 months post fertilization (mpf), 6 mpf, and 12 mpf (Fig 2). Normal retinal lamination and cellular differentiation was observed in \textit{cep290}^{fh297/fh297} mutants at 5 dpf (Fig 2A), indicating that retinal development did not require Cep290. At 3 mpf, we noticed fewer and more disorganized cone outer segments in \textit{cep290}^{fh297/fh297} mutants (Fig 2B, white arrow). Cone disorganization in \textit{cep290}^{fh297/fh297} mutants was progressive and by 6 mpf the loss of cone outer segments (COS) was more evident (Fig 2C). By 12 mpf, only a few discernible cones remained in the \textit{cep290}^{fh297/fh297} mutants (Fig 2D, arrows). Also noticeable was the continued thinning of the retinal outer nuclear layer (ONL) in \textit{cep290}^{fh297/fh297} mutant retinas when compared to wild-type animals (Fig 2E and 2F). The thickness of the ONL was reduced across the peripheral and central retina, although the difference was only statistically significant in the dorsal retina (Fig 2E). When the rows of nuclei in the ONL was quantified, a statistically significant difference was seen in the periphery of the dorsal retina (Fig 2F; 4.1±0.3 vs 3.1±0.3 rows of nuclei, \(P < 0.05\); n = 6).
We next used electron microscopy to examine retinal sections of wild-type and cep290<sup>fh297/fh297</sup> mutant adults (6 mpf) to determine how loss of Cep290 affected photoreceptor structure. In cep290<sup>fh297/fh297</sup> mutants, few cone outer segments were observed and the outer retina of cep290<sup>fh297/fh297</sup> mutants was more disorganized than that seen in wild-type retinas (Fig 3A–3C). At higher magnification, whereas the outer segments of wild-type animals contained highly organized stacks of disc membranes, the disc membranes were fragmented and the outer segments appear to be disintegrating in cep290<sup>fh297/fh297</sup> mutants (Fig 3D and 3E). We did not, however, observe any consistent accumulation of vesicular material adjacent to the ciliary base or other signs of disrupted ciliary trafficking (Fig 3D and 3E; white arrowheads).

Fig 2. Progressive cone loss in adult cep290<sup>fh297/fh297</sup> mutants. Methylene blue stained transverse histological sections of retinas from wild-type (top) and cep290<sup>fh297/fh297</sup> mutants (bottom) at 5 dpf (A); 3 months of age (B), 6 months of age (C), and 12 months of age (D). At 3 months, the cep290<sup>fh297/fh297</sup> mutants (bottom) had noticeably fewer cones (white arrows) and thinning of the cone outer segment (COS) layer. Few cones were observed at 12 months of age in cep290<sup>fh297/fh297</sup> mutants (black arrows). (E) Quantification of ONL thickness and (F) rows of nuclei in the ONL at different distances from the optic nerve in both the dorsal (negative numbers; left) and ventral (positive numbers; right) retina of cep290<sup>fh297/fh297</sup> mutants and wild-type sibling controls at 8 months of age. Data are shown as means ± SEM (n = 6, *P < 0.05). ROS, rod outer segments; COS, cone outer segments; ONL, outer nuclear layer; OPL, outer plexiform layer; INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer. Scale bar: 100 μm.

https://doi.org/10.1371/journal.pone.0213960.g002
These results suggest that loss of Cep290 disrupts cone outer segment structure and causes cell death.

**Progressive cone degeneration in cep290<sup>f297/fh297</sup> mutants**

To track the progression of photoreceptor degeneration, immunohistochemistry was performed on retinal cryosections of cep290<sup>f297/fh297</sup> mutants at 3, 6, and 12 months of age. Retinas were stained with peanut agglutinin lectin (PNA) to label the interphotoreceptor matrix surrounding cone outer segments [57] and Zpr-1, a monoclonal antibody that recognizes arrestin-3 like (Arr3L) on the cell bodies of red- and green-sensitive double cones [58, 59]. Similar to the results from plastic histology, the PNA-labeled cone outer segments were less...
organized and the inner segments appeared less densely packed in the cep290fh297/fh297 mutants at 3 months of age as compared to wild-type siblings (Fig 4A–4F). Cone degeneration in the cep290fh297/fh297 mutants was more apparent by 6 months of age (Fig 4G–4L) and by 12 months of age, very few cones remained (Fig 4M–4R). In older cep290fh297/fh297 mutants, some cones had Zpr-1 positive inner segments but lacked PNA-positive outer segments (Fig 4L and 4R; arrows), indicating that outer segment loss preceded cone death. This is consistent with a role for Cep290 in sensory cilia maintenance. The density of PNA-positive cone outer segments in wild-type and cep290fh297/fh297 mutants were quantified at each time point (Fig 5). The results indicated that cone degeneration initiated by 3 months of age in cep290fh297/fh297 mutants and progressed such that very few cone outer segments (1.6 COS / 50 μm) remained by 12 months of age.

**Distribution of rod outer segment proteins in cep290fh297/fh297 mutants**

Loss of Cep290 is associated with rapid loss of rods and rhodopsin mislocalization in the mouse cep290d1416/d1416 mutant [20]. Rhodopsin is a G-protein coupled receptor with seven transmembrane domains that passes along the ciliary plasma membrane before becoming incorporated into disc membranes within the outer segment. To evaluate the effects of Cep290 loss on rods in zebrafish, we stained retinal cryosections from cep290fh297/fh297 mutants and wild-type siblings with the monoclonal antibody Zpr3, which recognizes rhodopsin. At 3 mpf, when the first signs of cone degeneration were observed in cep290fh297/fh297 mutants, rhodopsin localized to the outer segments of both cep290fh297/fh297 mutants and wild-type control animals (Fig 6A–6D). By 6 mpf, when cone degeneration was pronounced, a significant amount...
of rhodopsin mislocalized to the inner segments and cell bodies (Fig 6E–6H; arrowheads). Rhodopsin continued to be mislocalized at 12 mpf (Fig 6l–6l; arrowheads), but significant loss of rod outer segment material was not observed. At 12 mpf, however, we noticed that the gap between the rod outer segments and the ONL, which typically is occupied by cone nuclei, was...
considerably smaller in 12 month-old animals as compared to wild-type siblings (Fig 6I and 6J; white brackets).

Active transport of cytosolic and transmembrane proteins (e.g. rhodopsin) through the ciliary TZ to the photoreceptor outer segments requires intraflagellar transport (IFT), while transport of lipidated protein cargo requires a distinct targeting system utilizing either PDE6D or UNC119 [60]. We therefore investigated the localization rhodopsin kinase (GRK1) and the β subunit of rod transducin (GNB1), to determine if loss of Cep290 also disrupts trafficking of lipidated ciliary proteins. GRK1 is a prenylated membrane protein that requires the function of Retinitis Pigmentosa 2 (RP2) for proper transport to rod outer segments [61]. Membrane association of GNB1 requires direct binding to the protein RP2 and loss of RP2 disrupts outer segment trafficking of both GNB1, GRK1, and other prenylated proteins [62–64]. In retinal cryosections from animals at both 6 months and 12 months of age, we found that GRK1 localized to the rod outer segments of \textit{cep290}^{fh297/fh297} mutants, similar to wild-type siblings (Fig 7A–7H). At both 6 and 12 months of age, the majority of GNB1 also localized to the rod outer segments of both wild-type and \textit{cep290}^{fh297/fh297} mutants (Fig 7I–7P). Taken together, these results suggest that loss of Cep290 specifically affects rhodopsin localization without broadly impairing transport of all outer segment proteins.

In response to retinal injury, zebrafish typically exhibit a robust capability of regenerating lost neurons, including photoreceptors [65, 66]. In the uninjured retina, Müller glia in the inner nuclear layer (INL) will periodically divide and produce unipotent rod progenitor cells that migrate to the ONL where they can proliferate as rod precursors and differentiate into rod photoreceptors [67–69]. In response to acute retinal damage, however, the Müller glia will dedifferentiate, undergo cellular reprogramming, and produce multipotent retinal progenitors that proliferate and are able to differentiate into all retinal cell types, including cones [66, 70]. Given this regenerative capacity, it was surprising to observe photoreceptor degeneration in \textit{cep290}^{fh297/fh297} mutants. To determine if \textit{cep290}^{fh297/fh297} mutants attempted regeneration, retinas from 3-month and 6-month old \textit{cep290}^{fh297/fh297} mutants and wild-type siblings were stained with antibodies against proliferating cell nuclear antigen (PCNA), which is a marker of cell proliferation, and quantified the number of PCNA+ cells in the ONL and inner nuclear layer (INL). Only a small number of individual proliferating cells were seen in the INL of 3 month old \textit{cep290}^{fh297/fh297} mutants (7.7±3.5) or wild-type siblings (10.9±1.5) and no statistical difference was found (Fig 8A and 8B; n = 6; P = 0.15). Compared to the INL, there were up to 10-fold more proliferating cells in the ONL of 3-month old mutant (82.7±17.2) and wild-type siblings (62.3±12.8), but still no statistical difference seen (Fig 8A and 8C; n = 6; P = 0.06). At 6 months of age, however, significantly more proliferating cells were found in both the INL and ONL of \textit{cep290}^{fh297/fh297} mutants (Fig 8B and 8C). Compared to wild-type siblings, there were 3-fold more proliferating cells in the INL and 10-fold more cells in the ONL of \textit{cep290}^{fh297/fh297} mutants. Of note, proliferating cells were 10-fold more abundant in the ONL than in the INL of \textit{cep290}^{fh297/fh297} mutants (note differences in Y-axes in Fig 8B and 8C). This suggests photoreceptor degeneration in \textit{cep290}^{fh297/fh297} mutants triggers robust proliferation of rod progenitor cells but only limited proliferation of Müller glia.

**Effects of the combined loss of \textit{cep290} and cilia genes \textit{ahi1}, \textit{cc2d2a}, or \textit{arl13b} differentially affects visual acuity but does not exacerbate photoreceptor degeneration**

A leading hypothesis to explain phenotypic variability in ciliopathies is the effects of mutations in second-site modifiers [26–29]. Typically, heterozygosity (i.e. loss of one allele) in cilia genes exacerbates the phenotypes observed in homozygous mutants of other cilia genes. Analysis of
animals with homozygous mutations in two distinct genes may reflect the additive effect of two independent phenotypes and not necessarily a role for second-site modifiers. Because the loss of cep290 leads to slow cone degeneration in zebrafish, we asked whether heterozygous mutations in genes encoding other cilia proteins would accelerate degeneration. The Cc2d2a and Ahi1 proteins are components of the MKS module that make up part of the TZ [28]. In humans, mutations in AHI1 and CC2D2A cause Joubert Syndrome and both genes have been proposed as potential modifiers of CEP290 [12, 33]. Cep290 directly binds Cc2d2a through an N-terminal domain of Cep290 [12] and Cep290 is predicted to bind Ahi1, suggesting that these connections are critical for TZ assembly or stability. Mutations in ARL13B also result in

Fig 7. Immunolocalization of GRK1 and GNB1 in cep290fh297/fh297 mutants at 6 and 12 months of age. Retinal cryosections of cep290fh297/fh297 mutants and wild-type siblings were stained with polyclonal antibodies against rhodopsin kinase (A–H; GRK1) or with GNB1 polyclonal antibodies against the β subunit of rod transducin (I–P) to label rod outer segments at both 6 and 12 months of age. ROS, rod outer segments; COS, cone outer segments; ONL, outer nuclear layer; OPL, outer plexiform layer; INL, inner nuclear layer; RGC, retinal ganglion cells. Scale bar: 50 μm.

https://doi.org/10.1371/journal.pone.0213960.g007
Joubert Syndrome. The Arl13b protein is required for axoneme extension and photoreceptor outer segment formation [34]. Importantly, the zebrafish cc2d2a-/- and ahi1-/- mutants show defects in photoreceptor OS structure during larval stages [40, 47] while the arl13b-/- zebrafish mutant undergoes a progressive photoreceptor degeneration [35]. The known and proposed biochemical and genetic interactions, as well as similar protein localization patterns in the transition zone and axoneme, suggested that these genes could potentially modulate phenotypes associated with Cep290 mutations.

As cep290fh297/fh297 adults were unable to breed naturally, heterozygous animals were mated to generate cep290+/fh297;ahi1+/-, cep290+/fh297;cc2d2a+/-, and cep290+/fh297;arl13b+/- adults. Pairwise crosses from these adults generated all possible genotypes in the expected Mendelian ratios. The double homozygous mutants (e.g. cep290fh297/fh297;ahi1+/-, cep290fh297/fh297;cc2d2a+/-; and cep290fh297/fh297;arl13b+/-) did not survive beyond 14 dpf. The cep290fh297/fh297;ahi1+/-, cep290fh297/fh297;cc2d2a+/-; and cep290fh297/fh297;arl13b+/- mutants were viable beyond 12 months and were indistinguishable from cep290fh297/fh297 mutants.

We next wanted to determine if loss of cep290 sensitizes photoreceptors to the additional loss of one allele of either ahi1, arl13b, or cc2d2a and would accelerate cone or rod degeneration. We first assessed cone degeneration by immunohistochemistry using the markers PNA and Zpr-1 on retinas from 6-month old cep290fh297/fh297;ahi1+/-, cep290fh297/fh297;cc2d2a+/-, and cep290fh297/fh297;arl13b+/- adults. No additional increase in cone loss was observed in the cep290fh297/fh297;ahi1+/-, cep290fh297/fh297;cc2d2a+/-; or cep290fh297/fh297;arl13b+/- mutants (Figs 9D, 10D and 11D). Retinal sections were also stained with antibodies against rhodopsin and rhodopsin kinase (GRK1) to assess trafficking of rod outer segment proteins (Figs 9B,
Rhodopsin mislocalization was quantified by measuring integrated fluorescence density in the rod inner and outer segments (see Methods). Whereas considerable rhodopsin mislocalization was observed in cep290fh297/fh297 mutants, there was no significant exacerbation of this phenotype by the additional loss of one allele of ahi1, cc2d2a, or arl13b (Figs 9E, 10E and 12E). GRK1 localized to the rod outer segments in wild-type animals and in all mutant genotypes (Figs 9C, 10C and 11C).

Finally, we asked if visual function of cep290fh297/fh297 mutant larvae could be diminished by the additional loss of one allele of ahi1, cc2d2a, or arl13b. We performed pairwise crosses of cep290fh297/fh297;ahi1+/−, cep290fh297/fh297;cc2d2a+/−, or cep290fh297/fh297;arl13b+/− adults and measured the OKR gain for both contrast sensitivity and visual acuity for all offspring at 5 dpf and subsequently determine the genotype for each animal (Fig 12). We previously reported that ahi1+/− mutants have disrupted photoreceptor outer segments but do exhibit normal visual behavior.
Although the cep290fh297/fh297;ahit1/+ mutants had no measurable defect in contrast sensitivity responses (Fig 12A), a significant reduction in spatial resolution discrimination was observed (Fig 12B). Interestingly, the cep290fh297/fh297;cc2d2a+/- mutants had no measurable defect in either contrast sensitivity or spatial resolution responses, although the cep290fh297/fh297;cc2d2a-/- mutants were more significantly affected (Fig 12C and 12D). The arl13b-/- single mutants showed significant impairment of both contrast sensitivity and spatial frequency discrimination (Fig 12E and 12F). Although the cep290fh297/fh297;arl13b+/- mutants were not statistically different from cep290fh297/fh297;arl13b-/- single mutants in contrast sensitivity, there was a statistically significant difference between cep290fh297/fh297 single mutants and cep290fh297/fh297;arl13b+/- mutants in spatial resolution (Fig 12F, purple bar). Interestingly, removing one allele of cep290 significantly enhanced the defects in both contrast sensitivity and spatial frequency of arl13b+/- mutants (Fig 12E and 12F; blue bars). Taken together, these results suggest that loss of cep290...
cep290 is differentially sensitive to the loss of one allele of ahi1, arl13b, and cc2d2a. The data also suggest that in zebrafish, arl13b may not function as a modifier of cep290, but cep290 may instead function as a modifier of arl13b in zebrafish. We did not include results from double homozygous mutants because this may reflect an additive effect from two independent phenotypes rather than a true modifier effect. Furthermore, the likelihood that both genes carry two mutant alleles is a highly unlikely to occur in humans with retina disease.

**Discussion**

Mutation of CEP290 is a major cause of ciliopathies and non-syndromic retinal degeneration. CEP290 mutations result in a variety of disorders with overlapping but clinically distinct phenotypes and significant phenotypic variation exists between patients diagnosed with the same syndrome. For example, the best corrected visual acuity of several LCA patients with CEP290 mutations varied from 20/50 to the absence of light perception [9]. The cause of this variation...
is often attributed to the presence of mutations in second-site modifier genes [33, 71]. Several reports have confirmed a role for modifier genes. For example, an allele of \textit{RPGRIP1L} enhances retinal degeneration across a spectrum of ciliopathies [29] and mutations in \textit{AHI1} were suggested to increase the severity of photoreceptor degeneration in nephronophthisis patients [72]. The effect of genetic modifiers on \textit{CEP290} phenotypes has been less clear. In humans, mutations in \textit{AHI1} and \textit{CC2D2A} cause JBTS and both genes have been proposed as potential modifiers of \textit{CEP290} [12, 33]. Missense alleles of \textit{AHI1} were associated with increased neurological involvement in a small number of \textit{CEP290}-LCA patients [33], while morpholino knockdown of \textit{cep290} increased the frequency of kidney cysts in \textit{cc2d2a} \textsuperscript{-/-} mutant zebrafish [12]. The absence of one \textit{Bbs4} allele enhanced photoreceptor degeneration in \textit{Cep290} \textit{rd16/rd16} mice [71]. However, mutation of a single allele of \textit{Bbs6} (\textit{Mkks}) rescued the photoreceptor degeneration of \textit{Cep290} \textit{rd16/rd16} mice [73]. Interestingly, a \textit{Cep290} gene-trap mouse model lacking almost all \textit{Cep290} protein was fertile and viable in Mendelian ratios when produced on the 129/Ola background, yet exhibited embryonic lethality when produced on the C57BL/6 background [74]. Such differences indicate that genetic modifiers within the C57BL/6 strain can

\textbf{Fig 12. Loss of \textit{ahi1} or \textit{arl13b} impairs visual function of \textit{cep290} \textit{fh297/fh297} mutants.} (A, C, E) OKR contrast response function of 5 dpf larvae (left) and bar graph (right) of corresponding data points at the 70\% contrast setting (hatched box). (B, D, F) OKR spatial resolution results of 5 dpf larvae (left) and bar graph of corresponding data points at the 0.039 spatial frequency (hatched box). Genotypes are indicated in the legend and in the X-axes. Inset values indicate the total number larvae tested for each genotype. \textdagger P < 0.05; \textdagger\dagger P < 0.01; \textdagger\dagger\dagger P < 0.001; \textdagger\dagger\dagger\dagger P < 0.0001 as determined by a 2-way ANOVA with a Multiples Comparisons test and Tukey corrections. Data are presented as means \pm s.e.m.

https://doi.org/10.1371/journal.pone.0213960.g012

https://doi.org/10.1371/journal.pone.0213960.g012
lead to significant phenotypic differences. Nevertheless, there remains some skepticism regarding the influence of genetic modifiers on human populations [75]. An analysis of 371 patients from 265 families with severe ciliopathies found that independent families harboring the same founder mutation had little phenotypic variation [75]. As the cohort under investigation was highly consanguineous, it is possible that modifiers were underrepresented in the population. It is also possible that the prevalence of certain phenotypes, such as retinal degeneration, may depend more on the influence of modifier genes when compared to the cardinal phenotypes required to establish a definitive clinical diagnosis.

In this study, we report that the zebrafish cep290<sup>fh297/fh297</sup> mutant retina undergoes progressive cone photoreceptor degeneration beginning at 3 months of age and is accompanied by rhodopsin mislocalization and thinning of the outer nuclear layer by 6 months of age. Retinal development occurs normally and the cep290<sup>fh297/fh297</sup> mutants have normal visual acuity as larvae. This is not inconsistent with clinical studies of LCA patients with point mutations in CEP290. Younger patients are more likely to have a normal fundus appearance, with older patients showing white flecks or pigmentary retinopathy [8, 33]. These observations indicate that in humans, photoreceptor development is preserved while the long-term photoreceptor survival is affected, similar to what is observed in the cep290<sup>fh297/fh297</sup> mutant.

We also investigated how heterozygous mutations in the ahi1, arl13b, or cc2d2a genes impacted retinal architecture and visual function of cep290<sup>fh297/fh297</sup> mutants. Compared to cep290<sup>fh297/fh297</sup> mutants, we found that the absence of one allele of these genes did not accelerate retinal degeneration or reduce viability on a cep290<sup>fh297/fh297</sup> mutant background. However, loss of one allele of ahi1 or arl13b did decrease the spatial frequency function of cep290<sup>fh297/fh297</sup> mutants while the cep290<sup>fh297/fh297; cc2d2a<sup>+/+</sup></sup> mutants were indistinguishable from cep290<sup>fh297/fh297</sup> mutants. We therefore conclude that the retinal phenotypes in zebrafish lacking cep290 are differentially sensitive to the loss of one allele of ahi1, arl13b, or cc2d2a. These experiments were prompted by a previous study of zebrafish cc2d2a<sup>−/−</sup> mutants that found that injection of morpholinos targeting cep290 significantly increased the frequency of pronephric cysts at 4 dpf [12], thereby indicating a potential genetic interaction between cc2d2a and cep290 in zebrafish. In a separate report, morpholino-induced knockdown of cep290 in zebrafish also prevented photoreceptor outer segment formation and other ciliopathy defects [76]. We rarely observed pronephric cysts in cep290<sup>fh297/fh297</sup> mutants and the frequency of cysts was not increased by the additional loss of cc2d2a, consistent with the lack of genetic interactions in the eye. It should also be noted that the cep290<sup>sa1383/sa1383</sup> mutant allele, which introduces a stop codon at amino acid 1844, was reported to have no overt phenotype [77]. Indeed, we did not detect any evidence of photoreceptor degeneration in cep290<sup>sa1383/sa1383</sup> mutants that were at least 15 months of age (data not shown). These contrasting results may reflect the observed differences observed between morpholino-induced phenotypes and mutant phenotypes. Such differences have been attributed to off-target effects of morpholinos or genetic compensation by mutants but not morphants [78, 79]. The slow photoreceptor degeneration observed in the zebrafish cep290<sup>fh297/fh297</sup> mutant differs from the phenotypes observed in mice lacking Cep290. The cep290<sup>rd16</sup> mouse undergoes almost complete loss of rods within 4 weeks of age [20], while a complete knockout of Cep290 is embryonic lethal [14]. Although the cep290<sup>fh297</sup> allele encodes a nonsense mutation, mutant cep290 transcripts were downregulated by 55% compared to wild-type levels. This is similar to what has been observed in human tissues. Fibroblasts derived from an LCA patient with the c.2991+1665A>G mutation had a 60% reduction in wild-type CEP290 transcripts that resulted in a corresponding ~80% reduction in protein levels [43]. A recent study determined that iPSC-derived RPE from a patient with biallelic truncating mutations in CEP290 maintained protein expression at levels at least 10% of wild-type expression [80]. Furthermore, CEP290-LCA patients carrying two nonsense alleles do not undergo the rapid photoreceptor degeneration observed in
Cep290<sup>rd16/rd16</sup> mice or the increased mortality of the Cep290<sup>ko/ko</sup> mice [8, 21]. Several possibilities exist that could explain these differences. It is possible that truncating mutations are subject to exon skipping in humans, thus leading to partial protein production [25]. Exon skipping was not detected in cep290 transcripts in the zebrafish retina, but perhaps retinal cells differ from other somatic cells in their sensitivity to mutations in the Cep290 gene. We acknowledge that the effect of the fh297 mutation on protein production in zebrafish remains unknown. Despite multiple attempts with both commercial antibodies [41] and custom-designed antibodies [42], we were unable to detect Cep290 protein in lysates of wild-type or cep290 mutants by immunoblotting or by immunohistochemistry on wild-type larvae, so the possibility that the mutated gene produces a truncated polypeptide with partial function cannot be excluded. Such partial and truncated polypeptides may also exist in humans with nonsense mutations.

Despite the loss of cone photoreceptors, the rod outer segments appear preserved. Zebrafish typically show a robust capacity to regenerate damaged photoreceptors following acute damage such as intense light exposure, mechanical injury, or chemical-induced toxicity [70, 81–83], but few studies have directly examined whether adult zebrafish have the capacity to regenerate photoreceptors in a model of inherited retinal degeneration. Following retina injury, the Müller glia within the INL undergo a reprogramming event and proliferate as retinal stem cells to regenerate lost neurons. Immunohistochemical analyses with proliferating cell nuclear antigen (PCNA) found a small increase in proliferating cells in the INL of cep290<sup>fh297/fh297</sup> mutants, but a significant increase in proliferating cells was seen in the ONL, which have been shown to be rod precursors [69, 84, 85]. It is possible that rods do undergo a slow degeneration in the cep290<sup>fh297/fh297</sup> mutants but the dying rods are being continually replaced from the rod progenitor population in the ONL. Because the Müller cells are not proliferative, cone regeneration does not occur. This obviously raises several intriguing questions about how the zebrafish retina differentially responds to acute injury versus a progressive hereditary degeneration.

Zebrafish cep290 mutants survive to adulthood and reinforce the important role of Cep290 in photoreceptor outer segment maintenance. Furthermore the cep290<sup>fh297/fh297</sup> mutant represents a model for slow retinal degeneration that mimics the ocular involvement of CEP290-dependent LCA and provides a unique platform to screen for genetic modifiers that accelerate or prevent photoreceptor degeneration. In addition, future work with this model can provide insight into the mechanisms required to trigger photoreceptor regeneration in zebrafish and the signaling pathways required to regenerate lost photoreceptors in CEP290 patients.

**Acknowledgments**

The authors would like to thank Dr. Iain Drummond from Massachusetts General Hospital and members of the Perkins lab for helpful discussions. This work was supported by the National Institutes of Health [NIH R01-EY017037 to B.D.P., NIH P30-EY025585 to Cole Eye Institute]; and the Research to Prevent Blindness [Doris and Jules Stein Professorship to B.D.P.].

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