α-crystallin R49Cneo mutation influences the architecture of lens fiber cell membranes and causes posterior and nuclear cataracts in mice

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αA-crystallin R49Cneo mutation influences the architecture of lens fiber cell membranes and causes posterior and nuclear cataracts in mice

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Abstract

Background: αA-crystallin (CRYAA/HSPB4), a major component of all vertebrate eye lenses, is a small heat shock protein responsible for maintaining lens transparency. The R49C mutation in the αA-crystallin protein is linked with non-syndromic, hereditary human cataracts in a four-generation Caucasian family.

Methods: This study describes a mouse cataract model generated by insertion of a neomycin-resistant (neo') gene into an intron of the gene encoding mutant R49C αA-crystallin. Mice carrying the neo' gene and wild-type Crya were also generated as controls. Heterozygous knock-in mice containing one wild type gene and one mutated gene for αA-crystallin (WT/R49Cneo) and homozygous knock-in mice containing two mutated genes (R49Cneo/R49Cneo) were compared.

Results: By 3 weeks, WT/R49Cneo mice exhibited large vacuoles in the cortical region 100 μm from the lens surface, and by 3 months posterior and nuclear cataracts had developed. WT/R49Cneo mice demonstrated severe posterior cataracts at 9 months of age, with considerable posterior nuclear migration evident in histological sections. R49Cneo/R49Cneo mice demonstrated nearly complete lens opacities by 5 months of age. In contrast, R49C mice in which the neo' gene was deleted by breeding with CreEIIa mice developed lens abnormalities at birth, suggesting that the neo' gene may suppress expression of mutant R49C αA-crystallin protein.

Conclusion: It is apparent that modification of membrane and cell-cell interactions occurs in the presence of the αA-crystallin mutation and rapidly leads to lens cell pathology in vivo.

Background

Cataracts involving mutations in lens crystallin genes have received considerable attention in recent years [1-5]. Vertebrate lens crystallins are divided into two families, α and βγ. α-crystallin is essential for lens transparency and accounts for nearly 50% of the protein mass in human lenses. It is a large multimeric complex with an aggregate molecular mass of 500,000–1,200,000 Da, and is isolated from lens fiber cells as a complex of αA- and αB-crystallin in a 3:1 stoichiometry [6]. αA-crystallin/HSPB4 is a member of the small heat shock protein family, which also includes αB-crystallin/HSPB5 and Hsp27/HSPB1 [7]. The etiology of lens disease is diverse, but a common pathological endpoint is the formation of large protein aggregates that scatter light. The capacity of α-crystallins to efficiently trap aggregation-prone denatured proteins is thought to delay age-related cataract in humans. α-crystallin and homoaggregates of αA and αB-crystallin interact in...
vitro with non-native proteins and prevent their irreversible aggregation and insolubilization [8]. Using recombinant proteins, αA and αB-crystallin have been shown to interact with many partially denatured substrates. Their stability, dynamic properties, and ability to transition from large assemblies to smaller dimeric and monomeric species appear to be critical for this chaperone ability [9-13]. The binding of αA and αB-crystallin to misfolded proteins occurs with a high efficiency [14]; however, once all the α-crystallin in lens fiber cells has been depleted, the concentration of irreversibly denatured proteins could increase, resulting in cataract. αA-crystallin binds to lens cell membranes and this association is increased in cataracts. Indeed, many models of cataract involve alterations in lens cell membranes [15-17]. The interaction of α-crystallin with the cytoskeleton is also of major importance in maintaining lens transparency [18-22].

αA-crystallin transcripts are present in mouse lens epithelium at embryonic day 10.5 and continue to be expressed during embryonic development and postnatally [23]. Studies on α-crystallin knockout mice suggest that loss of αA-crystallin may increase the concentration of unstable proteins and affect the solubility of αB-crystallin and γ-crystallin [24,25]. In addition to its essential role in the optical and refractive properties of the eye lens, αA-crystallin performs other functions; αA-crystallin knockout mice exhibit increased lens epithelial cell death and reduced cell proliferation [26,27], and αA-crystallin is also expressed in the retina, brain, spleen, and thymus [28] although its role in these tissues is not fully understood.

Transfection studies show that α-crystallins protect cells from stress-induced apoptosis [27]. Both αA and αB-crystallin are negative regulators of apoptosis in lens cells [29,30], and αA-crystallin knockout lens epithelial cells have a higher level of cell death than wild type cells [31]. Thus, cellular protection by crystallins may delay the onset of age-related and hereditary cataracts.

Several single-point mutations in α-crystallins have been linked with hereditary human cataracts [1,32-37]. In αA-crystallin, R49C and W9X mutations in the N-terminal domain, and R116C, R116H, and G98R mutations in the C-terminal domain cause human cataracts, as do C-terminal R120G, 450delA, D140N mutations in αB-crystallin. Many of these mutations are inherited by autosomal dominant mechanisms. Study of these αA-crystallin gene mutations would enhance our understanding of the mechanisms of cataract formation [1,5,32,34,37-39]. Mutations in αA-crystallin increase the abundance of aggregation-prone proteins. In vitro studies suggest that mutant αA-crystallin may aggregate causing increased co-precipitation of substrate proteins [40-43]. The most common effect of single-point mutations and truncations is an increase in the size of the oligomeric complex [44-46]. The R49C mutant of αA-crystallin has a slightly higher mass and radius of gyration, but the main effect of the mutation is increased protein insolubility [42]. Arginine is a highly charged residue, and its replacement by the less polar cysteine might alter protein assembly and solubility. Moreover, the positive charge on arginine 49 has been highly conserved during evolution, and is important for in vitro chaperone activity of αA-crystallin [47,48].

Several model systems are available to study the effect of single-point mutations of crystallin genes. Transgenic mice have been used to investigate the effect of the R116C mutation in αA-crystallin in vivo [49], and several studies have used transfected cells expressing mutant proteins [1,38,50]. Naturally occurring and mutagenesis-induced mouse models have also been analyzed [51-54]. Naturally occurring αA-crystallin mutations in mice are CryaA lop1 (R54H), and CryaA R54C mutations in the N-terminal region, and Y118D and CryaA Aey7 (V124E) mutations in the C-terminal domain. While significant information has been obtained by studying mutant αA-crystallin in vitro, such studies have certain limitations. Mutations affecting protein interactions at low concentrations may have little relevance to how they associate in vivo at higher concentrations in the lens. Thus, an optimal model must investigate the mechanism of hereditary cataracts, this laboratory recently generated R49C knock-in mice and compared wild type lenses with heterozygous and homozygous mouse models which show milder lens effects in vivo, thus allowing examination of the effects of the mutant protein at potentially changed levels of expression. The aim of the present study was to characterize the lens abnormalities in WT/R49C
ewt and R49C
ewt/R49C
ewt mouse cataract models which show previous reports of WT/R49C and R49C/R49C lenses [42,55].

Methods

Animals
Mice were maintained at Washington University Division of Comparative Medicine by trained veterinary staff. All protocols and animal procedures were approved by the Washington University Animal Studies Committee. Lens opacity was monitored by slit lamp biomicroscopy. The left eye of the animals was examined, and pupils were dilated with a mixture of 10% phenylephrine hydrochloride and 1% tropicamide (Alcon, Fort Worth, Texas). Mice carrying the R49C
ewt locus were generated from one embryonic stem cell clone (129 background) as previ-
ously described [55]. This locus carries a neomycin cassette flanked by lox P sites inserted into an XhoI site in intron 1 of Cryaa. One ES clone with the neo insertion but no Cryaa mutation was used to generate WT/WT<sub>neo</sub> and WT<sup>neo</sup>/WT<sub>neo</sub> mice. Two lines of mice expressing the mutation in Cryaa, R49C<sub>neo</sub>KI3 and R49C<sub>neo</sub>KI4, were bred. Another mouse line, WT<sub>neo</sub>KI2, expressing wild type Cryaa was also generated on 129 background. Mice containing one copy of the targeted knock-in allele (homozygous mice) were interbred to generate homozygous mice. Genotyping primers were used to identify the knock-in construct containing the neomycin cassette as described previously [55]. Transgenic mice expressing Cre-ELAs on a C57BL/6 background were bred with homozygous Cryaa knock-in mice to delete the neo<sup>e</sup> gene. These mice were described in previous studies [42,55].

**Slit lamp examination**

Dilated mouse eyes were examined in unanesthetized mice. Stages of cataracts were defined as follows: Stage 0 – clear lens; Stage 1 – loss of normal appearance of posterior lens and prominence of γ-suture line; Stage 2 – discrete posterior changes accompanied by light nuclear opacity; Stage 3 – nearly mature cataract, involving approximately three-fourths of the lens with bubbles and opacity; Stage 4 – completely mature cataract involving the cortex with bubbles and vacuoles.

**Histology**

For conventional histology, eyes were fixed overnight at 4°C in formalin (Sigma, St. Louis, MO). After a thorough wash in phosphate-buffered saline, lenses were dehydrated through graded acetone and infiltrated in metacrylate resin (H-8100; Technovit, Kulzer, Germany) according to the following schedule: 1:2 resin:acetone, 1 day; 1:1 resin:acetone, 1 day; 100% resin, 4 days. Blocks were polymerized for 1 hour at 4°C. Sections (3 μm) were cut and stained with haematoxylin and eosin.

**Immunofluorescence**

Immunofluorescence was performed as previously described [31]. Primary antibody to lens membrane intrinsic protein MIP was purchased from Alpha Diagnostic International. Alexa<sup>568</sup> labeled secondary antibody (Molecular Probes) was used at 1:500 dilution.

**Immunoblotting**

Lens extracts were separated into water-soluble and insoluble fractions, and examined by SDS-PAGE and immunoblotting with an antibody to α-crystallin [56]. Protein concentration of the water-soluble fractions was determined by the bicinchoninic acid (BCA) protein assay according to the manufacturer’s instructions (Thermo Scientific-Pierce Chemical Co, Rockford IL). 40–50 μg of extract was loaded on the gel. The primary antibody was either a polyclonal antibody (used at 1:500 dilution) to total α-crystallin or a monoclonal antibody (a gift from Dr. Paul FitzGerald; used at 1:100 dilution) to αA-crystallin, and horseradish peroxidase-conjugated secondary antibodies as described previously [27,42,56]. The relative intensity of the mouse αA-crystallin band at approximately 20 kDa was analyzed by Image J software. The decrease in αA-crystallin soluble protein with increasing numbers of R49C<sub>neo</sub>-containing genes was calculated as the mean of three independent experiments.

**Cryoimmuno electron microscopic analysis**

Lens sections were examined by cryoimmuno electron microscopy. Sections (50–80 nm) were treated with a primary monoclonal antibody to αA-crystallin [27] and antimouse IgG conjugated with 18 nm gold particles as secondary antibody (Sigma). Specimens were stained with uranyl acetate and examined in a 1200EX transmission electron microscope as described previously [57].

**Analytical chromatography and analysis**

WT/WT<sub>neo</sub> and WT/R49C<sub>neo</sub> mouse lenses (1.5 to 2.5 month old) were used to isolate water-soluble proteins [42]. Six lenses of each genotype were homogenized in 500 μl of phosphate-buffered saline (Sigma) and the water-soluble protein fraction was separated by centrifugation at 15,000 g. FPLC chromatography was performed on a chromatography system containing an inline detector for UV absorption at 280 nm. A 16 cm × 60 cm Superdex 200 column (GE Healthcare Biosciences) equilibrated with 10 mM Tris.Cl; pH 6.8 containing 100 mM NaCl, 1 mM DTT and 0.5 mM EDTA was used. Proteins were injected at 1 mg/ml for different chromatographic runs. One ml fractions were collected, and proteins mass was determined by calibrated molecular mass standards. The area under each peak was used to determine changes in abundance of α<sub>α</sub>, β and γ-crystallin fractions. The results are representative of 6 lenses per genotype. The analysis was repeated with almost the same results.

**Results**

While constructing a gene knock-in mouse model of R49C αA-crystallin, we generated mice carrying a floxed neo<sup>e</sup> gene in an intron of the Cryaa gene (Figure 1). Mating these to a mouse expressing Cre recombinase resulted in removal of the neo cassette, leaving the loxP site adjacent to exon 1 of the Cryaa gene, and producing a functional but less active mutant allele [42,55]. Two lines of R49C<sub>neo</sub> mice, R49C<sub>neo</sub>KI 3 and R49C<sub>neo</sub>KI 4, were generated from a single ES cell clone, and the results obtained from these two lines were very similar. The knock-in mice were viable and bred normally. Heterozygous WT/R49C<sub>neo</sub> mice were intercrossed to produce R49C<sub>neo</sub>/R49C<sub>neo</sub> homozygous offspring. A control mouse line was also generated from an ES cell clone that had the neomycin cassette inserted,
but lacked the R49C αA-crystallin mutation. These mice did not have the altered lens phenotype, and served as a useful control. We speculated based on previous work by others [58-60] that the neor gene might exhibit diminished activity of mutant R49C gene that would provide novel insights into the biology of R49C αA-crystallin in vivo. To determine the effect of the R49C neo gene on α-crystallin protein expression, lenses were analyzed by gel permeation chromatography. Figure 2 shows the chromatography profile of water-soluble lens proteins of WT/WTneo and WT/R49Cneo mice measured by absorbance measurements at 280 nm. The analysis showed a 30% decrease in expression of total α-crystallin protein in WT/R49Cneo heterozygous lens as compared with wild type lenses. The expression of β-crystallins also decreased whereas γ-crystallin expression was not appreciably affected. As compared with WT/R49C lenses [42], α-crystallin decreased more in the WT/R49Cneo lenses.

Lens opacities were confirmed in R49Cneo mutant mice by slit lamp analysis (Figure 3). By 3 months, R49Cneo mice showed evidence of opacities in posterior and nuclear regions. Over time, the cataract progressed to a nuclear cataract and then to an all-over opacity that included the cortical fibers (Figure 3A–D). Heterozygous mice at each age showed variable lens opacities ranging from clear (stage 0) at <2 months, stage 0 to stage 2 at 2–3 months, and clear to complete opacity (stage 4) at >4 months of age. Wild type mice did not show any of these abnormalities. At least four mice were examined for each genotype at a given age (Table 1A). The neomycin cassette was deleted by breeding homozygous R49Cneo mice with Cre-Elα transgenic mice. These neomycin-deleted R49C mutant mice had a more severe lens phenotype, with homozygous R49C/R49C knock-in neo-deleted mice exhibiting an opacity that covered three-quarters of the young lens (Figure 3E and 3F). Overall, cataract progression was more rapid in R49C/R49C mice than in R49Cneo/R49Cneo mice.

Examination of αA-crystallin soluble protein revealed that some of these changes were due to protein insolubility (Figure 4). Mice with nuclear opacity ranging from stage 1 to stage 3 (by slit lamp analysis) were separated into water-soluble and insoluble fractions. Water-soluble frac-

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**Figure 1**

Plasmid construct used to generate R49Cneo gene knock-in mice. The 5' and 3' arms of the αA-crystallin gene (Cryaa) were cloned into a vector containing the floxed neomycin (neo) cDNA. Mutagenesis was performed to mutate amino acid arginine 49 of αA-crystallin to cysteine (R49C). The asterisk above exon 1 indicates the mutation. The numbered blue rectangles indicate exons. The filled triangles are loxP sites and X denotes the XhoI site. Mouse embryonic stem (ES) cells SCC-10 were electroporated with the mutant plasmid, and clones testing positive for neo were identified and used to generate R49Cneo αA-crystallin knock-in mice (WT/R49Cneo). One clone containing wild type (WT) αA-crystallin cDNA and neo was also analyzed and used to generate mice with the neo allele but no mutation (WT/WTneo).

**Figure 2**

FPLC analysis of crystallins in WT/WTneo and WT/R49Cneo mouse lenses. UV absorption profile of watersoluble mouse lens proteins separated from WT/WTneo (red) and WT/R49Cneo (blue) by gel permeation chromatography. Proteins were pooled from six lenses of each genotype at ~2-month-old. mAU represents milli absorbance units at 280 nm.
**Figure 3**

Lens phenotypes in R49C\textsuperscript{neo} mice. (A-D) Eyes were dilated and examined by slit lamp. (A) Wild type mice (5 months old) had clear lenses (stage 0). (B) Heterozygous WT/R49C\textsuperscript{neo} \( \alpha \text{A-crystallin} \) knock-in mice (3 months old) had opacity in the posterior and nuclear regions of the lens (stage 2). (C) Homozygous R49C\textsuperscript{neo}/R49C\textsuperscript{neo} \( \alpha \text{A-crystallin} \) knock-in mice (3 months old) had a nearly complete cataract (stage 3–4). (D) R49C\textsuperscript{neo}/R49C\textsuperscript{neo} \( \alpha \text{A-crystallin} \) knock-in mouse lenses showed a complete cataract at 5 months (stage 4). (E, F) Cataract in lens of a newborn R49C/R49C homozygous mouse with deletion of the neo\textsuperscript{r} gene by Cre-recombinase. Slit lamp image (E) shows a severe nuclear opacity at birth (stage 3). (F) Higher magnification of the lens shown in (E) shows the nuclear opacity covering ~70% of the lens.

**Table 1: Phenotypic changes in wild type (WT), WT/R49C\textsuperscript{neo} heterozygous, and R49C\textsuperscript{neo}/R49C\textsuperscript{neo} homozygous mice.**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Phenotype</th>
<th>Age Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nuclear and posterior changes by slit lamp biomicroscopy</td>
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<td>2–3 months</td>
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<td>WT</td>
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<td>0/4</td>
<td>2/9</td>
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<tr>
<td>WT/R49C\textsuperscript{neo}</td>
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<td>14/22</td>
<td>7/7</td>
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<tr>
<td>R49C\textsuperscript{neo}/R49C\textsuperscript{neo}</td>
<td>4/5</td>
<td>5/7</td>
<td>13/14</td>
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<table>
<thead>
<tr>
<th>Genotype</th>
<th>Phenotype</th>
<th>Age Group</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
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<td>2–3 months</td>
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<tr>
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<td>1/4</td>
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<td>WT/R49C\textsuperscript{neo}</td>
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<tr>
<td>R49C\textsuperscript{neo}/R49C\textsuperscript{neo}</td>
<td>4/5</td>
<td>3/4</td>
<td>6/8</td>
</tr>
</tbody>
</table>

* For each age group, the total number of mice analyzed is shown in the denominator, and the number of mice demonstrating the phenotype is shown in the numerator. The final columns show the total number of mice of each genotype that demonstrate a phenotype, with the proportion given in parentheses.
tions were analyzed by immunoblotting. An excess of protein extract was loaded so that protein would be detectable even in the homozygous lenses. Protein insolubility was proportional to nuclear cataract formation by slit lamp analysis.

Examples of the posterior lens changes observed in R49C<sub>neo</sub> αA-crystallin knock-in lens sections are shown in Figures 5 and 6. Posterior cataract was evident in 3-month-old R49C<sub>neo</sub>/R49C<sub>neo</sub> homozygous mice. At 9 months, WT/R49C<sub>neo</sub> heterozygous mouse lenses showed severe posterior rupture, curling up of the posterior capsule, and migration of cells to the posterior lens (Figure 6).

At 3 weeks, histological examination of lenses revealed the presence of both small and large swollen cells or vacuoles in the lens cortical fibers of WT/R49C<sub>neo</sub> mice, which were even more evident in R49C<sub>neo</sub> homozygous mouse lenses (Figure 7). Early fiber formation appeared to occur normally. Swollen fiber cells begin to appear at 3 weeks postnatal in WT/R49C<sub>neo</sub> heterozygous mice. Swollen fiber cells were confined to a distinct band of cortical fibers ~100 μm from the lens surface in both young and old mice. Fiber cells on either side of the swollen fibers appear to be unaffected. These swollen cells appeared to
be formed by membrane rupture and fusion of cytoplasmic contents from multiple cells. Extensive undulations and interdigitations of these membranes as well as large separations between fiber cells occurred in homozygous mouse lenses. These gaps are evident in fiber cells that have detached from the capsule (Figure 6C). The proportion of swollen cells and posterior changes in WT/R49C<sup>neo</sup> and R49C<sup>neo</sup>/R49C<sup>neo</sup> mice examined by histology is shown in Table 1B.

The number of swollen fiber cells increased in lenses of older homozygous mice, and extended towards the center of the lens (Figure 8). At 10 months old, a large area of aberrant fiber cells was observed. Fiber cells near the lens surface appeared normal. Newly synthesized cortical fibers appeared to form normally, but vacuoles were apparent in a band of fibers in the deep cortex 100 μm from lens surface, and increased dramatically towards the lens center.

Immunofluorescence analysis with an antibody to membrane intrinsic protein (MIP) revealed that the swollen cells are enclosed by fiber cell membranes (Figure 9). Swollen cells were not found in wild type littermates or in WT<sup>neo</sup> mice. Opacification was related to defects in membrane structure of cortical lens fiber cells. Lenses were further examined by cryoimmuno electron microscopy with an antibody to αA-crystallin. Wild type lenses showed smooth and linear plasma membranes between the fiber cells, and αA-crystallin was restricted to the cytoplasm (Figure 10A). In contrast, fiber cell membranes of heterozygous WT/R49C<sup>neo</sup> lenses were non-linear with extensive undulations, and significant gaps between the cells. A greater proportion of the αA-crystallin immunoreactivity was associated with fiber cell membranes in the heterozygous lenses (Figure 10B) than in wild type lenses.

**Discussion**

In human patients with a missense mutation in exon 1 of the gene encoding αA-crystallin (CRYAA), autosomal dominant nuclear cataract was shown to segregate in a four generation Caucasian family [1]. The C to T transition in the first base of codon 49 of CRYAA results in the non-conservative substitution of arginine 49 to cysteine (R49C) in αA-crystallin protein [1]. Patients with cataracts are heterozygous for this mutation, which suggests that both wild type and R49C mutant αA-crystallin subunits are expressed in these lenses in addition to wild type αB-crystallin. Unfortunately, photographic documentation of the cataracts is not available, preventing a detailed pheno-typic analysis. It was shown previously that the αA-crystallin R49C mutation causes lens opacities in a knock-in mouse model, due to protein insolubilization and cell death [42,55]. Here a R49C<sup>neo</sup> construct was used that

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**Figure 7**

**Morphological alterations in cortical fiber cell membranes of R49C<sup>neo</sup> lenses.** Lens sections were stained with haematoxylin/eosin. (A) Normal appearance of fiber cells in the lens equatorial region in 3-week-old wild type lenses. (B) Swollen fiber cells and separations between fiber cells in 3-week-old WT/R49C<sup>neo</sup> heterozygous lenses. (C) Gaps between fiber cells in 3-week-old R49C<sup>neo</sup>/R49C<sup>neo</sup> homozygous lenses.
causes a milder lens phenotype to show that R49C αA-crystallin disrupts normal fiber cell organization and structure.

Homozygous mice with deletion of the floxed neor gene (R49C/R49C) have a more drastic phenotype, with smaller eyes and smaller lenses than wild type and heterozygous (WT/R49C) mutant mice [55]. This implies that the neor gene is suppressing expression of the R49C-αA mutant protein, and is supported by FPLC analysis of soluble lens proteins. Indeed, previous studies indicate that selectable markers inserted in non-coding regions can affect gene expression at both the DNA and RNA level, resulting in reduced protein expression [58-60].

The lenses of WT/R49Cneo and R49Cneo/R49Cneo mice displayed remarkable posterior lens defects and accumulation of swollen cells in the lens cortex, and developed nuclear cataracts. Both posterior lens cell defects and number of swollen fiber cells increased with age and with R49Cneo dosage, with more defects in R49Cneo/R49Cneo mice than in WT/R49Cneo mice. The appearance of swollen fiber cells in the lens cortex may be an early marker of structural perturbation in the R49Cneo lenses. The swollen fiber cells in deep cortical fibers of R49Cneo lenses were evident as early as 3 weeks postnatal and increased with age, such that more than half of the lens was covered with swollen fiber cells at 10 months.

Histological and immunofluorescence analysis of the R49Cneo knock-in lenses suggests that the lens opacities observed in R49Cneo mice are associated with membrane defects in the deep cortical fiber cells approximately 100 μm from the lens surface that result in gaps between adjacent fiber cells. Visible swelling or vacuolar areas are often limited to one or more segments of a fiber cell; hence it is likely that there are many more aberrant fiber cells than those observed in a single mid-sagittal section. Vacuolated cells indicate destruction of the fiber cells. Further studies are necessary to determine whether these vacuolar or swollen cells are formed by membrane rupture and fusion of cytoplasmic contents from multiple cells. This type of fiber cell swelling has been reported in diabetic and other models of cataract [61,62]. Membrane defects have also been shown to be associated with lens opacities induced by dexamethasone and mechanical stress in cultured lenses, both of which are associated with the loss of cadherin junctions [17,63]. The interaction between αA-crystallin and fiber cell membranes is well-established [16,64,65], and has been shown to increase with lens...
aging and in cataracts [66]. In the current study, membrane alterations in WT/R49C neo mouse lenses were confirmed by ultrastructural analysis. Cryoimmuno electron microscopic analysis of heterozygous R49C neo lenses with an antibody specific to \(\alpha\)-crystallin showed loss of membrane linearity in the cortical fiber cells, undulations of the membranes, and significant gaps between cells. Furthermore, more \(\alpha\)-crystallin was membrane-associated in the heterozygous knock-in lens fiber cells than in the wild type controls. Our studies also showed that the solubility of \(\alpha\)-crystallin is decreased in the R49C neo lenses, which may have a profound effect on its binding to lens fiber cell membranes [67].

Evidence in the literature suggests that loss of \(\alpha\)-crystallin function through gene knockout or mutation alters protein homeostasis by increasing the abundance of aggregation-prone proteins. This may result in altered protein-protein interactions, and protein insolubility [55,68]. Correct protein conformation is essential for normal cell function, and proteotoxic stress due to protein aggregation and loss of protein homeostasis is known to be associated with aging and disease [69].

In the present work, no significant defects were found in WT/WT neo and WT neo/WT neo lenses. In targeted gene deletion studies in mice, \(\alpha\)A-/- heterozygous mouse lenses do not show lens opacification, whereas \(\alpha\)A-/ homozygous lenses develop cataracts at an early age [24]. Most of the defects in \(\alpha\)A-cristallin knockout (\(\alpha\)A-/) mouse lenses are due to protein insolubility in the lens nucleus and cell death in the lens epithelium [24,31]; no defects in fiber cell elongation or fiber cell morphology were reported. Our previous study showed that lens defects in R49C/R49C knock-in mice are the result of a drastic alteration of lens fiber cell morphology, although early fiber cell elongation is normal [55]. In the present study, we did not observe as strong an effect on lens fiber morphology in young R49C neo mice as in R49C/R49C mice, but did observe varied posterior lens defects beginning at an early postnatal age, as well as posterior migration of nuclei at older ages and posterior rupture. Such posterior migration of epithelial cells has been reported in human posterior subcapsular cataracts [70,71]. A disorganization of the postequatorial fiber zone was also observed, which has been reported in human posterior subcapsular cataracts. Thus, the R49C neo mouse model system appears to be an effective model for human posterior subcapsular cataracts.

Conclusion

In summary, the results of the present work, combined with previously published data, provide evidence that \(\alpha\)-crystallin plays a role in the structure of fiber cell membranes, and is involved in the formation of hereditary cataracts. The present study provides insight into the \textit{in vivo} effects of the R49C mutation of \(\alpha\)-crystallin on the lens, and extends the findings of previous studies [1,42]. In mice, expression of the R49C \(\alpha\)-crystallin mutant protein in the presence of the neo gene causes posterior cataracts with posterior migration of epithelial cell nuclei, and demonstrates that the mutant protein affects the integrity of lens fiber cells in the deep cortex. These changes are milder than those observed when the neo gene is deleted. By studying homozygous R49C neo/R49C neo mice that begin to lose transparency around 2 months of age, compared with the typical onset at birth in R49C/R49C mice, the author has obtained additional information about the progression of cataracts. The R49C neo mice described in the present study are one of only a few models available.
to study hereditary posterior cataract, a common human lens pathology.

Competing interests
The author declares no competing interests.

Authors' contributions
The author performed slit lamp biomicroscopy, microscopic analysis of lens sections, and drafted the manuscript and subsequent revisions.

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