Lens

Dysregulation of Autophagy Occurs During Congenital Cataract Development in β A3 Δ G91 Mice

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Citation: Boateng AK, Joseph R, Srivastava OP. Dysregulation of autophagy occurs during congenital cataract development in β A3 Δ G91 mice. *Invest Ophthalmol Vis Sci.* 2024;65(4):4. https://doi.org/10.1167/iovs.65.4.4 **PURPOSE.** To examine lens phenotypic characteristics in β A3 Δ G91 mice and determine if β A3 Δ G91 affects autophagy in the lens.

METHODS. We generated a $\beta A3\Delta G91$ mouse model using CRISPR/Cas9 methodology. Comparative phenotypic and biochemical characterizations of lenses from postnatal day 0 (P0), P15, and 1-month-old $\beta A3\Delta G91$ and wild-type (WT) mice were performed. The methodologies used included non-invasive slit-lamp examination, reverse transcription– quantitative polymerase chain reaction (RT-qPCR), western blot, and immunohistochemical (IHC) analyses to determine the levels of autophagy-related genes and proteins. Transmission electron microscopy (TEM) analysis of lenses was performed to assess organelle degradation and the presence of autophagic vesicles. TUNEL staining was used to determine apoptosis in the lens.

RESULTS. Relative to WT lenses, 1-month-old $\beta A3\Delta G91$ mice developed congenital nuclear cataract and microphthalmia and showed an early loss of endoplasmic reticulum (ER) in the cortex and attenuation of nuclei degradation. This observation was confirmed by TEM analysis, as was the presence of autophagic vesicles in $\beta A3\Delta G91$ lenses. Comparative IHC and RT-qPCR analyses showed relatively higher levels of autophagy markers (ubiquitinated proteins and p62, LC3, and LAMP2 proteins) in $\beta A3\Delta G91$ lenses compared to WT lenses. Additionally, $\beta A3\Delta G91$ lenses showed relatively greater numbers of apoptotic cells and higher levels of cleaved caspase-3 and caspase-9.

CONCLUSIONS. The deletion of G91 in β A3 Δ G91 mice leads to higher levels of expression of autophagy-related proteins and their transcripts relative to WT lenses. Taken together, G91 deletion in β A3/A1-crystallin is associated with autophagy disruption, attenuation of nuclei degradation, and cellular apoptosis in the lens, which might be congenital cataract causative factors.

Keywords: β A3/A1-crystallin, congenital cataract, autophagy, lens

T he crystalline lens is composed of a single layer of lens epithelial cells (LECs) that differentiate at the equator into lens fiber cells throughout life to maintain lens transparency.¹ During the differentiation of epithelial cells to fiber cells, transitioning lens epithelial cells undergo programmed degradation of organelles to produce an organelle-free zone (OFZ).^{2,3} Autophagy is a major cellular event involved in the clearance of organelles as well as misfolded or aggregated proteins in the lens,⁴⁻⁶ and impairment of this process could result in cataract formation.^{7,8}

Congenital cataracts cause approximately 10% of childhood blindness worldwide, occurring at a rate of three to six per 10,000 live births worldwide⁹ and three to four visually significant cataracts per 10,000 live births in the United States.¹⁰ Heterogeneous congenital cataracts, inherited as autosomal recessive, autosomal dominant, or X-linked traits, represent ~8% to 25% of non-syndromic cataracts.⁹ Approximately one-third of pediatric cataract mutations are hereditary,¹¹ and among these mutations, ~50% are found in crystallins, ~25% in connexins, and the remainder in other structural genes and transcription factors.¹² Several mutations in β A3/A1-crystallin have been identified that cause autosomaldominant congenital cataracts,¹² most of which are clustered at two sites; one occurs at the exon 3 splice site and the other is an exon 4 in-frame deletion.^{13–23} Among the CRY β A1 mutations, one common β A3 Δ G91 mutation has been shown to cause congenital nuclear cataract in several ethnically disparate families.^{13,16–22,24}

In vertebrates, crystallins are classified as α -, β -, or γ -crystallins. Even though these crystallins have been studied extensively, β -crystallins remain elusive in their structural significance and functions. This is partly because of the existence of their several subunits and tendency to form oligomers.²⁵ β -crystallins are further subdivided into three basic (β B-) and four acidic (β A-) proteins.^{25,26} Of all β -crystallins, β A3/A1-crystallin remains a unique lens structural protein because it produces two protein isoforms (β A1- and β A3-crystallins) from a single *CRY* β A1 mRNA via an alternative translation initiation site by leaky ribosomal scanning.²⁷ The two proteins share sequence homology except that β A1-crystallin is 17 amino acids shorter at the N-terminal region than β A3-crystallin.²⁷ Structurally, β A3/

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A1-crystallin is encoded by six exons, out of which four (exons 3–6) code for four Greek key motifs that fold into two similar domains, connected by a short peptide.²⁷ The distinctive Greek key pattern of β - and γ -crystallins is required to maintain lens transparency and the refractive power of the lens.²⁸

In rodents, β A3/A1-crystallin mRNA is first detected around embryonic day 12.5 in the posterior lens vesicle, suggesting its role in early lens development.²⁹ Despite a robust expression in the lens, β A3/A1-crystallin is also expressed in the retinal pigment epithelium (RPE), retinal astrocytes, and the brain.²⁹⁻³² Although the function of β A3/A1-crystallin as a lysosomal resident protein and an autophagy regulator is well studied in the retina,^{30–32} its similar function in the lens, other than as a structural protein, remains unclear. Interestingly, studies have shown that an absence of β A3/A1-crystallin in murine lenses leads to autophagy disruption and cataract development.^{31–34}

Among the β A3/A1-crystallin mutations that cause human congenital cataracts, there is the commonly occurring in-frame deletion of a highly conserved Gly-91 within the second Greek key motif ($\beta A3 \Delta G91$). This autosomal dominant congenital nuclear cataract, discovered in several ethnically disparate families worldwide, causes a dysfunctional β A3/A1-crystallin due to the loss of its structural integrity. It is evidenced by the recombinant G91-deleted β A3/A1-crystallin, which has shown defective refolding, loss of solubility, and oligomerization with other crystallins.^{19,35} In order to explore the molecular mechanism of β A3 Δ G91induced congenital cataract, we have generated a $\beta A3 \Delta G91$ knock-in mouse model using CRISPR/Cas9 methodology. Here, we report comparative phenotypic characteristics of lenses of β A3 Δ G91 mice and wild-type (WT) mice and show that the mutation leads to the disruption of autophagy. Our mutant mice showed nuclei and mitochondria being retained in the OFZ and exhibited elevated levels of autophagy markers and cellular apoptosis relative to WT mice. This finding strongly suggests that autophagy is associated with cataract development of $\beta A3 \Delta G91$ mice lenses relative to WT mice lenses.

MATERIALS AND METHODS

Ethics Statement

All animal experiments were conducted in accordance with protocols approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Alabama (UAB) at Birmingham (protocol no. 08393), and all procedures adhered to the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Mice were kept under a 12-hour/12-hour light/dark cycle in a controlled, pathogenfree facility of the University of Alabama at Birmingham.

Generation of β A3 Δ G91 Mice

The $\beta A3 \Delta G91$ mouse model was generated at the UAB Transgenic and Genetically Engineered Models (TGEMs) Core Facility using CRISPR/Cas9 methodology. Briefly, a CRISPR guide RNA was designed using CRISPOR.tefor.net to target exon 4 on the mouse Cryba1 locus adjacent to Gly-91. A symmetrical 200-bp single-stranded DNA (ssDNA) repair template was designed to introduce the desired mutation, as well as an additional silent mutation creating a second Taq1-restriction site for genotyping. The CRISPR guide RNA and Cas9 protein were complexed in vitro before injection at a concentration of 50-ng/µL guide RNA, 50-ng/µL Cas9 protein, and 50-ng/µL ssDNA repair template into 0.5day mouse embryos. Embryos surviving microinjection were transferred into pseudo-pregnant C57BL/6 recipient mice to develop to term. Desired timed-mating was performed between female C57BL/6 mice and male founder β A3 Δ G91 mice to generate autosomal-dominant β A3 Δ G91 mice. All mice were bred in a C57BL/6 strain background.

PCR and DNA Analysis

Genomic DNA was extracted from mouse tail tissues using a DNA isolation kit from QIAGEN (Germantown, MD, USA). For initial verification of the knock-in, PCR products were sequenced in the UAB Heflin Center for Genomic Sciences. PCR primers used for genotyping were $\beta A3\Delta G91$ (forward: 5'-AGGTCTTCTCCCTTAGGCCA-3') and $\beta A3\Delta G91$ (reverse: 5'-ACTAGAAGAGGGCACGGGAT-3'). Following PCR amplification, the restriction enzyme digestion was performed using Taq1-v2 endonuclease (New England Biolabs, Ipswich, MA, USA), and analyzed by using a 6% polyacrylamide gel.

Lens Imaging

The lenses from postnatal day 0 (P0), P15, and P30 β A3 Δ G91 and WT mice were enucleated and imaged using a microscope with a charge-coupled device (CCD) camera attachment. One-month-old β A3 Δ G91 and WT mice were subjected to minimal anesthesia containing a ketamine-xylazine cocktail (0.05 mL/20g mouse weight, by intraperitoneal injection). Their eyes were dilated with 0.5% tropicamide eye drops and were examined within 5 minutes with a hand-held slit-lamp biomicroscope equipped with a camera.

PCR Analyses

RNA was extracted from mice lenses (n = 6 from WT and $\beta A3\Delta G91$ mice) using the TRIzol method, and 1 µg of total RNA from each lens was reverse transcribed using an Invitrogen cDNA Synthesis Kit (Thermo Fisher Scientific, Waltham, MA, USA). The housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (*GAPDH*) served as an endogenous control for the quantitation of autophagy-related genes. Primers used in the reverse transcription–quantitative polymerase chain reaction (RT-qPCR) analyses are listed in Table 1.

Protein Extraction and Mass Spectrometry Analyses

Lens proteins were extracted from WT and $\beta A3\Delta G91$ mice and separated into water-soluble (WS), water-

TABLE 1. Primer Sequences for QPCR Analyses

Primers	Primer Sequences (5'-3')		
LC3B	Forward: 5'-ATCCCAGTAGGTTCTCCTGC-3'		
	Reverse: 5'-CGCTCTATAATCACCCGCC-3'		
p62	Forward:5'-GGACCCATCTACAGAGGCTG-3'		
	Reverse: 5'-ACTGGATCGTGTCCAGAGCA-3'		
LAMP2	Forward: 5'-GGACAGTATTCTACAGCCCAGGAG-3'		
	Reverse: 5'-CCAGCATAGGTCTTTCTTCTGCC-3'		
CRYβA1	Forward: 5'-TGGAGTGTGGCGCAATCATA-3'		
	Reverse: 5'-GAAGTCTGGGCGTGAGATCC-3'		

TABLE 2. Mass Spectrometric Analyses: Normalized Spectral Counts^{*} for CRY β A1/A3 in WT Versus β A3 Δ G91 Lenses

_	WT	β A3 ∆ G91	Difference (βA3∆G91 – WT)
Water-soluble	17.29	7.45	-9.84
Water-insoluble– urea-soluble	9.23	6.19	-3.05
Water-insoluble– urea-insoluble	2.72	6.70	3.98

* As described in the Materials and Methods section.

insoluble-urea-soluble (WI-US), and water-insolubleurea-insoluble (WI-UI) fractions, following previously described methods.⁵² Protein concentrations in these fractions were determined using the Pierce BCA Protein Assay Kit (Thermo Fisher Scientific). Lens proteins were electrophoretically separated by SDS-PAGE using 12% polyacrylamide gels. Following mass spectrometry (MS) analyses,⁵² with a confidence score set to at least 95%, spectral counts were computed by summing the number of MS/MS identifications for β A3/A1-crystallin. Spectrum count normalization in Scaffold was used (Proteome Software, Portland, OR, USA). Briefly, the normalized spectral count for β A3/A1-crystallin was calculated as $\frac{average spectral count between WT and \betaA3\Delta G91}{total spectral count per group}$ spectrum Normalized spectral counts for β A3/A1-crystallin are listed in Table 2.

Immunoblotting

Isolated lenses (n = 6, from WT and $\beta A3 \Delta G91$ mice) were homogenized in radioimmunoprecipitation assay (RIPA) buffer (Sigma-Aldrich, St. Louis, MO, USA) containing a protease inhibitor cocktail tablet (Roche, Indianapolis, IN, USA). Protein concentration was measured using a BCA Protein Assay Kit. Between 50 and 80 µg of lens protein was electrophoretically separated using 12% polyacrylamide gels, and the protein bands were transferred to a polyvinylidene fluoride (PVDF) membrane (Bio-Rad, Hercules, CA, USA). The blots were blocked with 5% non-fat dry milk in TBS for 1 hour and incubated with primary antibodies (LC3, anti- β A3-crystallin) at 4°C overnight. After three washes with TBST (Tris-buffered saline + 0.1% Tween 20), blots were incubated with a secondary IRDye 800CW antibody, and protein bands were visualized using LI-COR Odyssey system (LI-COR, Lincoln, NE, USA). GAPDH was used as the loading control (Cell Signaling Technology, Danvers, MA, USA). Full images of immunoblots are provided in the Supplementary Figures.

Histology and Immunofluorescence

Eyes from P0, P15, and 1-month-old $\beta A3\Delta G91$ and WT mice were enucleated and immediately embedded in Tissue-Tek O.C.T compound (Sakura Finetek USA, Torrance, CA, USA). They were kept on dry ice for 1 hour and then transferred to -80° C. Eyes were then cut into 16-µm sections using a Leica cryostat (Leica Biosystems, Deer Park, IL, USA). Lens sections were fixed in 4% paraformaldehyde (PFA) for 10 minutes. For all immunofluorescence studies, the sections were blocked in 5% goat serum and 1% BSA in PBS for 1 hour and incubated overnight with a desired primary antibody. The primary antibodies used in this study included anti- β A3-crystallin (ab151722; Abcam, Waltham, MA, USA); anti-p62/SQSTM (55274-I-AP;

y antibody. The primary antibodies col, and fluore uded anti- β A3-crystallin (ab151722; a Zeiss Axiople USA); anti-p62/SQSTM (55274-I-AP; a CCD camera.

Proteintech, Rosemont, IL, USA); anti-ubiquitin (58395S; Cell Signaling Technology); anti-LC3 (A17424; Cell Signaling Technology); anti-LAMP2 (66301-1-Ig; Proteintech); anticleaved caspase-3 (9664; Cell Signaling Technology); and anti-cleaved caspase-9 (9507; Cell Signaling Technology). The sections were washed with PBS and incubated with fluorophore-labeled secondary antibodies (Thermo Fisher Scientific) for 1 hour in the dark at room temperature. The lens tissues were washed in phosphate-buffered saline with 0.1% Tween 20 (PBST) and counterstained with Hoechst 33342 or 4',6-diamidino-2-phenylindole (DAPI)-containing mounting medium. Photographs were taken with a Nikon AX-R laser confocal microscope (Nikon, Tokyo, Japan) or, in some cases, with a Zeiss Axioplan 2 fluorescence microscope equipped with a CCD camera (Carl Zeiss Microscopy, Jena, Germany) at the UAB Vision Science Research Center core facility. Signal intensities were quantified using Image J (National Institutes of Health, Bethesda, MD, USA).

Cell Culture Studies

Lens epithelial cells from WT and $\beta A3\Delta G91$ mice were cultured as previously described.³³ Briefly, lenses from 4to 5-week-old mice were extracted and incubated in 0.25% trypsin for 3 hours. Cells were then recovered by centrifuging at 1500 rpm and cultured in M199 medium containing 2% fetal bovine serum (HyClone, Logan, UT, USA). For all immunofluorescence studies, cells were plated on eightwell chambered slides. The immunostaining procedure was performed as described above. To induce autophagy in vitro, lens epithelial cells from WT and $\beta A3\Delta G91$ mice were serum starved for 18 hours. In the same way, autophagy inhibition was carried out by treating cells with bafilomycin A1 (100 nM) for 18 hours.

Transmission Electron Microscopy

Lenses from 1-month-old WT and $\beta A3 \Delta G91$ mice were fixed in a modified Karnovsky's fixative (2% PFA with 2% glutaraldehyde in 0.1-M phosphate buffer; Microscopy Science, Hatfield, PA, USA) for 1 hour at room temperature and then kept at 4°C overnight. The fixed lenses were rinsed several times with 0.1-M sodium cacodylate buffer (pH 7.4) and post-fixed with 1% OsO₄ in cacodylate buffer for 1 hour in the dark. After several rinses in cacodylate buffer, lenses were dehydrated through a series of graded 50% to 100% ethyl alcohol. Next, the tissues were placed in a 1:1 mixture of propylene oxide and embedding resin overnight. The following day, the lenses were transferred into a fresh change of 100% embedding media and left overnight to polymerize in a 60°C embedding oven. Ultrathin sections (70- to 80-nm thickness) were cut using a Leica Ultracut UCT 7 ultramicrotome and stained with uranyl acetate and lead citrate. The lens sections were imaged on an FEI 120 kv Spirit TEM (FEI Company, Hillsboro, OR, USA), and images were collected using a BioSprint 29 CCD camera (AMT Imaging, Woburn, MA, USA).

Apoptosis Assay

Apoptotic cells in lens tissues were detected using a Roche Fluorescein In Situ Cell Death Detection Kit. TUNEL staining was performed according to the manufacturer's protocol, and fluorescent images of the sections were taken with a Zeiss Axioplan 2 fluorescence microscope equipped with a CCD camera.

Statistical Analysis

Data were recorded as mean \pm SD, and analyzed using R 4.0.5 for Windows (R Foundation for Statistical Computing, Vienna, Austria). Significant differences were evaluated by analysis of independent Student's *t*-test (two-tailed). Statistical significance was defined as *P* < 0.05. All experiments were performed and repeated in triplicate.

RESULTS

Generation and Characterization of β A3 Δ G91 Mice

CRISPR/Cas 9 technology was used to generate the $\beta A3\Delta G91$ knock-in mouse model. The knock-in mutation was verified via DNA sequencing, which confirmed the

3-bp (GAG) deletion that resulted in the deletion of Gly-91 in F0 founder mice, as shown by dashes highlighted in pink in Figure 1A. In these mice, an additional silent mutation was introduced to induce a second T^{CGA} site (Fig. 1A, black arrowhead), in addition to the first (black arrow), which is originally present in the genome. The T^CGA sequence is recognized and cleaved by the Taq1 endonuclease, producing three nucleotide fragments in $\beta A3 \Delta G91$ mice instead of two fragments in the WT mice during genotyping (Fig. 1B). After successfully generating the mouse model, we used the hand-held slit lamp to examine 1-monthold mice for cataract development. β A3 Δ G91 mice exhibited a dense nuclear cataract and microphthalmia (Fig. 1C, right panel) compared to lenses of WT mice (Fig. 1C, left panel). Upon gross inspection, the anterior chamber and ocular adnexa appeared normal. Whole lens tissues and LECs were extracted from 1-month-old WT and $\beta A3 \Delta G91$

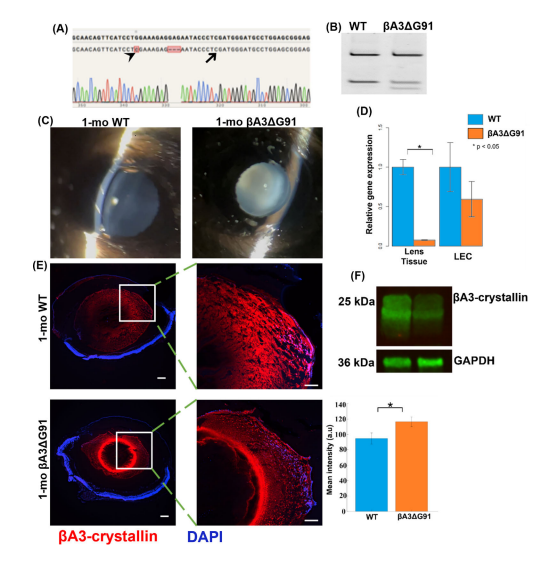


FIGURE 1. Phenotypic characterization of $\beta A_3 \Delta G_91$ mice. (**A**) DNA from the tail was amplified with primers spanning the target region on exon 4 of the *CRY* $\beta A_1/A_3$ gene. Sequence analysis verifies the GAG deletion that leads to the glycine-91 deletion and the additional silent mutation, creating a second Taq1 restriction site for genotyping (*black arrowbead*). (**B**) Genotyping results show two and three bands for WT and $\beta A_3 \Delta G_{91}$ mice, respectively. (**C**) Silt-lamp photographs of 1-month-old mice revealed that the mutant mice had a dense nuclear cataract (*rigbt*) whereas the WT mice had a clear lens. (**D**) Significantly lower levels of *CRY* βA_1 mRNA seen in the whole lens tissue is demonstrated by RT-qPCR analysis (*left grapb*). (**E**) IHC revealed a relatively smaller lens with puncta in the outer cortex, and a bright ring-like pattern formed in the inner cortex of $\beta A_3 \Delta G_{91}$ lenses (shown in $4 \times$ and $20 \times$, respectively). *Scale bar*: 200 µm. **P* < 0.05 versus age-matched WT. (**F**) Western blots of RIPA-soluble lens homogenates probed with the anti- $\beta A_3 \Delta G_{91}$ lenses relative to WT lenses. GAPDH was used as a loading control.

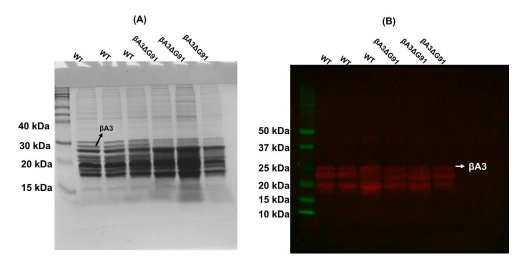


FIGURE 2. Coomassie blue staining and western blot analysis of RIPA buffer–solubilized proteins from lenses of WT and $\beta A3\Delta G91$ mice. (**A**) Three individual lenses from both types of mice were solubilized in identical volumes of RIPA buffer, and identical volumes of the preparations were analyzed by SDS-PAGE using 12% polyacrylamide gels. (**B**) During western blot analysis, the anti- $\beta A3$ antibody (see Fig. 1F) was utilized.

mice to determine $CRY\beta A1$ gene expression by RT-qPCR (Fig. 1D). In the whole lens tissue, $CRY\beta A1$ mRNA was significantly downregulated in β A3 Δ G91 lenses compared to the WT lenses. Additionally, our immunohistochemical (IHC) analyses using anti- β A3-crystallin antibody showed punctate staining in the outer cortex and a bright ring-like pattern in the inner cortex of the mutant lens sections (Fig. 1E, lower panel), which was not present in the WT lenses (Fig. 1E, upper panel). Interestingly, immunoblots of the RIPA-soluble lens protein fractions corroborated our qPCR studies by showing decreased levels of β A3/A1-crystallin expression in $\beta A3 \Delta G91$ lenses relative to the WT lenses (Fig. 1F). The β A3-crystallin immunoreactive ring-structure during IHC (Fig. 1E, lower panel) suggested the possibility that $\beta A3 \Delta G91$ protein becomes water insoluble in the cortex, and it causes the protein to accumulate in the form of a ring (perhaps at the outer edge of nuclear cataract), which was absent in the WT lenses. This was supported by the insolubility of recombinant $\beta A3 \Delta G91$ protein as described in a previous report.¹⁹ Further, we speculate that the intense ring in the nuclear region of $\beta A3 \Delta G91$ lenses compared to the WT lenses (Fig. 1E) could represent insolubilization of the crystallin and could be a cataractogenic factor. As described above, the western blot analysis with anti- β A3-crystallin antibody of the soluble fraction showed that the immunoreactive β A3-crystallin band was found to be at relatively lower levels compared to the lenses from WT mice (Fig. 1F). We suspected that the decreased expression of β A3-crystallin in our western blot was due to either degradation or insolubilization or both of the crystallin in β A3 Δ G91 lenses relative to WT lenses. To further examine this, we comparatively analyzed the WS and WI protein fractions of the two types of lenses by western blot analysis and mass spectrometry. As shown in Fig. 2A, the Coomassie-blue-stained gel protein profiles of the RIPA buffer-soluble proteins showed the loss of a 25-kDa β A3 crystallin band in β A3 Δ G91 lenses relative to WT lenses (identified by an arrow). On western blot analysis with the same anti- β A3 crystallin antibody as used in Figure 1F, the level of the immunoreactive band of 25 kDa was relatively reduced in β A3 Δ G91 lenses relative to WT lenses (identified by an arrow in Fig. 2B). Additionally, the comparative mass spectrometric analyses of WS, WI-US, and WI-UI proteins of 1-month-old β A3 Δ G91 and WT mice showed that, relative to WT lenses, β A3 Δ G91 lenses exhibited relatively greater levels of insolubility of β A3-crystallin in WI-UI protein fractions (Table 2). Together, the data suggest that β A3-crystallin undergoes both degradation and insolubilization in β A3 Δ G91 lenses relative to WT lenses.

Delayed Nuclei Degradation in Lenses of $\beta A3\Delta G91$ Mice Relative to WT Mice

Following epithelial to fiber cell differentiation, the fiber cells lose their organelles (e.g., nuclei, mitochondria, endoplasmic reticulum [ER]), to form an organelle-free zone (OFZ) in the nuclear region.^{2,3} Lenses of 1-month-old β A3 Δ G91 mice exhibited attenuation of nuclei in the OFZ region relative to WT lenses during hematoxylin and eosin (H&E) staining and the presence of extensive vacuoles in the outer and inner cortex (Figs. 3A, 3B). The DAPI staining further confirmed the attenuation of nuclei in the OFZ region (Figs. 3C, 3D). IHC analysis of lens sections used anti-HSPA5 antibody (heat shock protein family A member 5, an Endoplasmic Reticulum chaperone that localizes in its lumen, and is involved in the folding and assembly of proteins). At 1 month of age, immunostaining of the ER with HSPA5 antibody was diminished in β A3 Δ G91 lenses relative to WT lenses, suggesting a loss of ER in the mutant lenses (Figs. 3E, 3F). Together, the results suggest that the degradation of nuclei was impaired in the OFZ zone. In contrast, the ER levels were reduced in the same region, possibly due to the usage of the ER membranes in autophagosome formation during autophagy.³⁶⁻³⁸

The above result of the attenuation of nuclei degradation in $\beta A3\Delta G91$ lenses relative to WT lenses was further corroborated by TEM analyses of the different regions of the lens (Fig. 4). In the lens epithelium, the mitochon-

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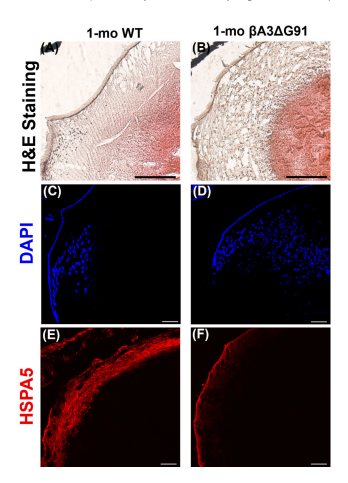


FIGURE 3. Impaired nuclei degradation in $\beta A3\Delta G91$ relative to WT lenses. (**A**, **B**) H&E staining of WT and $\beta A3\Delta G91$ lenses revealed the presence of several nuclei scattered throughout the lens. *Scale bar*: 200 µm. (**C**-**F**) Immunofluorescence with DAPI and HSPA5. (**C**, **D**) $\beta A3\Delta G91$ contained more nuclei scattered throughout the lens than the WT lens. (**E**, **F**) There was a decrease in HSPA5 intensity, even though the area covered by the signal was similar in both WT and $\beta A3\Delta G91$ lenses.

dria (M) and ER were equally present in both WT lenses (Fig. 4A) and $\beta A3 \Delta G91$ lenses (Fig. 4B), but the epithelium of $\beta A3 \Delta G91$ lenses showed the presence of extensive vacuoles (Fig. 4B). These organelles were also found in the outer cortex of both WT and β A3 Δ G91 lenses (Figs. 4C, 4D). However, fewer ER were detected in the β A3 Δ G91 lenses (Fig. 4D) than in the WT lenses. Interestingly, mitochondria present in the WT lenses were mostly elongated, whereas the mutants contained a combination of both round and elongated mitochondria. The round mitochondria might represent their degenerative nature. In the inner cortex (Figs. 4E-4H), relatively more condensed nuclei (N) forming inclusion bodies, disrupted nuclear envelopes, and chromatin seepage through breaks in the nuclear envelope into the cytosol were observed in β A3 Δ G91 lenses (Figs. 4F, 4H).^{39,40} The inner cortex of WT lenses exhibited signs of mitochondrial degradation, including discontinuous double membrane and loss of cristae (Figs. 4E, 4G), whereas $\beta A3 \Delta G91$ had intact mitochondria (Figs. 4F, 4H). In the nuclear region, the $\beta A3\Delta G91$ lenses (Fig. 4J) exhibited disorganization of fiber cells and membranes in the regions proximal to the opacity.

Impaired Autophagy in β A3 Δ G91 Lenses Relative to WT Lenses

Broadly, the four major sequential steps of autophagy are as follows: (1) After induction by a stress signal (e.g., starvation, oxidative stress, toxins), ubiquitination of proteins occurs. (2) p62, a cargo receptor for autophagy, binds to ubiquitinated proteins. (3) Autophagosome formation is initiated, and, concomitantly, a cytosolic form of LC3 (LC3-I, a 17-kDa microtubule-associated soluble protein 1A/1B light chain) is conjugated to phosphatidylethanolamine to form a LC3-phosphatidylethanolamine conjugate (LC3-II), which is recruited to autophagosomal membranes. (4) Autophagosome fusion with lysosomal constituents (e.g., lysosomal membrane proteins and enzymes) to form the autolysosome; during autophagy, autophagosomes engulf and degrade cytoplasmic components, including cytosolic proteins and organelles.⁴¹ To examine whether autophagy might be linked to the attenuation of nuclei degradation in β A3 Δ G91 lenses relative to WT lenses, we determined the expression of autophagy markers at both the transcriptional and translational levels during the specific autophagic steps described above.

Differential Expression of Autophagy Markers With Cataract Progression in β A3 Δ G91 Mice

We examined mice lenses of $\beta A3 \Delta G91$ and WT mice at P0, P15, and P30 (1 month) for progressive cataract development and expression of an early autophagy marker. As expected, in contrast to clear lenses of WT mice, $\beta A3 \Delta G91$ mice exhibited a bilateral nuclear cataract at P0 (Fig. 5B) that progressively increased in size at P15 (Fig. 5D) and P30 (Fig. 5F) and showed microphakia. The lenses of the two types of mice at identical ages (i.e., at P0, P15, and P30) were also examined by IHC for the expression of p62 as an early autophagy marker. The expression of p62 at P0 was relatively elevated and scattered as puncta throughout both outer and inner cortical and nuclear regions in β A3 Δ G91 lenses (Fig. 5H) relative to a lower expression in the WT lenses (Fig. 5G). In the P15 lenses, a greater level of immunoreactive proteins with anti-p62 antibody was observed which were mostly confined in the inner cortical and nuclear regions of the β A3 Δ G91 lenses relative to WT lenses (Figs. 5I, 5J). The elevated p62 immunoreactivity throughout the lens at P0 in β A3 Δ G91 lenses suggests that the autophagy process occurs in both epithelial and fiber cells during early differentiation but is disrupted relative to WT lenses. The disruption of autophagy is further supported by Figures 3 and 4, which show the attenuation of nuclei and mitochondrial degradation in β A3 Δ G91 lenses relative to WT lenses. The elevated expression pattern of p62 in β A3 Δ G91 was not surprising, because at P0 epithelial-to-fiber cell differentiation occurs throughout the lens; therefore, elevated p62 was observed in both cortical and nuclear regions. However, at the P15 and P30 stage, an active autophagic process to degrade organelles occurs in the outer and inner cortical regions to generate an OFZ, so its disruption is represented by elevated p62 in the cortical regions. The above results show that disruption of the autophagy process is an ongoing early event during epithelial-to-fiber cell differentiation along with early opacity development. Apparently, autophagosomes are also formed and fuse with the opaque region

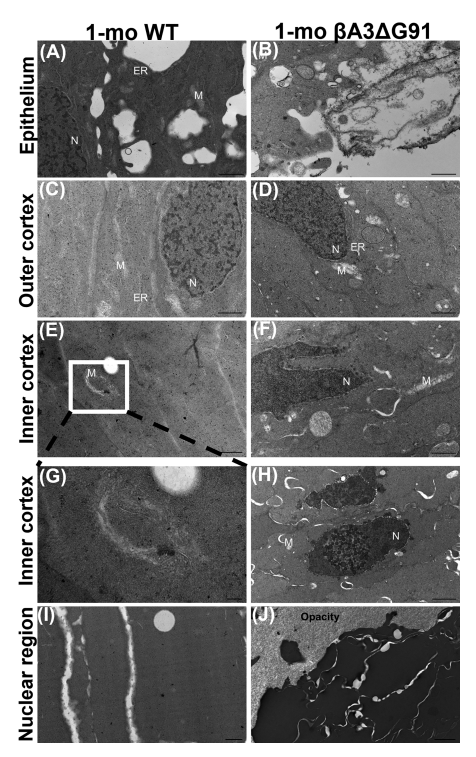


FIGURE 4. TEM analysis showing attenuation of nuclei and mitochondrial degradation in lenses of $\beta A3\Delta G91$ mice relative to WT mice. (**A**, **B**) Epithelium, nuclei, ER, and mitochondria were present in both types of lenses. *Scale bar*: 1 µm. (**C**, **D**) In the outer cortex, less ER was spotted in the $\beta A3\Delta G91$ than in the WT lens. *Scale bar*: 1 µm. (**E**–**H**) In the inner cortex, the $\beta A3\Delta G91$ nuclear envelope was broken at some ends and chromatin exited into the cytosol. Some nuclear materials were packaged into inclusion bodies, forming a beaded appearance. The WT lens began to show mitochondria degradation. No ER was seen in either type of lens. *Scale bar*: 1 µm; inset, 400 nm. (**I**, **J**) In the nuclear region, adjacent fiber cells surrounding the region of opacity in $\beta A3\Delta G91$ were highly disorganized. The WT lens was free of organelles or any opacity. *Scale bar*: 1 µm.

as supported by our TEM results in lenses of 1-month-old $\beta A3\Delta G91$ mice (Figs. 8H, 8I). This led to our hypothesis that the opacity development might involve non-degradation of autophagosomes containing intact organelles, which

might be due to lack of vacuolar (V)-ATPase activity to acidify lysosomes by the mutated β A3/A1-crystallin. Such lysosomal acidification is a normal process of the native β A3-crystallin.

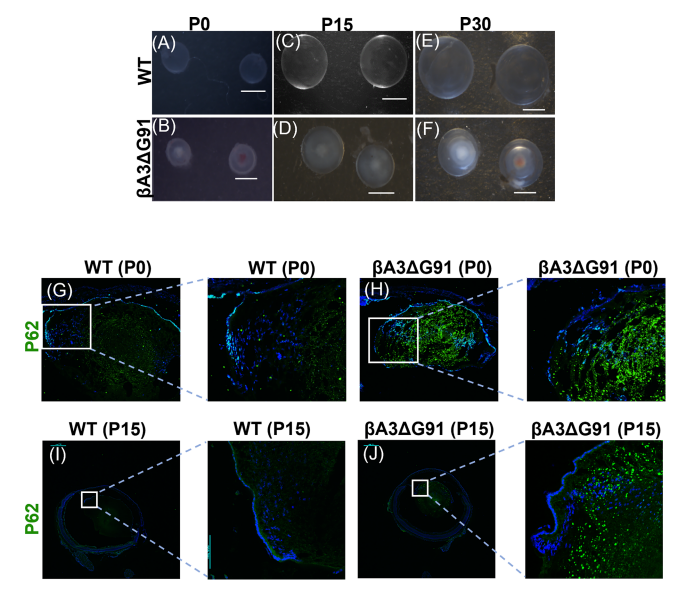


FIGURE 5. Differential expression of autophagy marker, p62, with cataract progression at P0, P15, and P30 (1 month) in β A3 Δ G91 and WT mice. (**A**–**F**) Images of WT and β A3 Δ G91 lenses at P0, P15, and P30. The lenses showed cataract development only in β A3 Δ G91 mice at P0 which progressed in size at P15 and P30. (**G**, **H**) IHC analysis using anti-p62 antibody of lenses from WT and β A3 Δ G91 mice at P0. (**I**, **J**) IHC analysis using anti-p62 antibody of lenses relative to the WT lenses.

Ubiquitination of Proteins and p62 Upregulation in β A3 Δ G91 Lenses

One hallmark of the beginning of the autophagic process in a cell is the ubiquitination of proteins. Ubiquitin binds to misfolded proteins or superfluous organelles and tags them for degradation. In cases where autophagy is impaired, ubiquitin and p62 accumulate within cells.⁴² On the other hand, p62 is an autophagy receptor that recognizes and binds to polyubiquitinated proteins to activate autophagy through its interaction with LC3.⁴³ To confirm whether aberrant organelle degradation in β A3 Δ G91 lenses relative to WT lenses occurred due to impaired autophagy, these lenses were first examined for the presence of ubiquitinated proteins. IHC analyses showed that ubiquitin immunostaining was about three times greater in the β A3 Δ G91 lenses relative to the WT lenses (Figs. 6A, 6B). Similarly, p62 puncta, which were mostly localized to the outer cortex, were also at relatively higher levels in β A3 Δ G91 lenses (Figs. 6E, 6F) than the WT lenses (Figs. 6C, 6D). Additionally, p62 was localized at relatively higher levels in both outer and inner cortex and also within the immunoreactive β A3 ring region (Figs. 1E, 6E, 6F). The expression of p62 was also examined in cultured LECs from both types of mice. Under normal serum conditions, p62 immunofluorescence was relatively about three times higher in β A3 Δ G91 LECs (Fig. 6I) than in the WT LECs (Fig. 6G), suggesting that p62 is upregulated in the former LECs.⁴⁴ Upon treatment with bafilomycin A1 (100-nM), p62 immunostaining intensities were significantly higher than under the normal serum conditions in both types of cells, but their levels were comparable (Figs. 6H, 6J). This showed that the effect of bafilomycin A1 treatment was similar between WT and mutant LECs. This is because bafilomycin A1 has been shown to inhibit the later stages

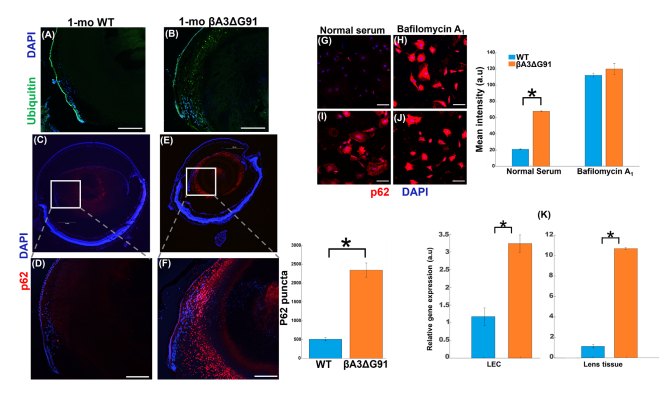


FIGURE 6. Ubiquitination of proteins and p62 upregulation in $\beta A3\Delta G91$ relative to WT lenses. (**A**, **B**) Immunohistochemical analyses of ubiquitin in mouse lens sections. Ubiquitin expression was higher and seen as a band of speckles in the $\beta A3\Delta G91$ relative to the WT lenses (10×). (**C–F**) Immunofluorescence analyses of p62 puncta demonstrated a significant increase in $\beta A3\Delta G91$ relative to WT lenses (4×, 20×). *Scale bar*: 500 µm. (**G–J**) Immunofluorescence analyses of p62 in LECs showed increased p62 intensity in $\beta A3\Delta G91$ LECs relative to WT under normal serum conditions. When cells were treated with bafilomycin A1, p62 was similarly upregulated in both WT and $\beta A3\Delta G91$ LECs. *Scale bar*: 100 µm. **P* < 0.05 versus age-matched WT. (**K**) p62 mRNA analyses showed 10-fold and threefold increases in p62 transcripts in $\beta A3\Delta G91$ whole lenses and LECs, respectively.

of autophagy (inhibiting lysosomal acidification by inhibiting V-ATPase).⁴⁵ Additionally, when RT-qPCR quantified the levels of *p62* transcripts in both whole lens tissues and LECs, threefold and ninefold upregulation in the β A3 Δ G91 relative to WT were observed in the LECs and lens tissues, respectively (Fig. 6K).

Elevated Levels of LC3 and LAMP2 Suggest Disruption of Autophagy

Labeling of LC3 serves as a way to track the binding of p62 and subsequent recruitment of autophagosomes and is therefore considered a good indicator of autophagosome formation. In cells, quantifying LC3-positive puncta is a gold-standard assay to assess autophagosome count.^{46,47} Similarly, lysosome-associated membrane glycoprotein 2 (LAMP2) is a membrane glycoprotein that functions in the protection, maintenance, and adhesion of the lysosome.48,49 To further expand our assessment that the accumulation of ubiquitin and p62 (Fig. 7) was due to impaired autophagosome formation or degradation by lysosomes, we examined the expression of LC3 and LAMP2 as markers for autophagosome and lysosome formation, respectively. LC3 exhibited a weak immunoreaction in the inner cortex but relatively strong staining in the nuclear region, forming a ring-like pattern in the β A3 Δ G91 lens (Figs. 7D, 7F) relative to the WT lens (Figs. 7A, 7C). A similar pattern was seen with LAMP2 immunoreaction in β A3 Δ G91 lenses relative to WT lenses, and LAMP2 was again colocalized in the ring-like structure (Figs. 7E, 7F). When LECs of $\beta A3 \Delta G91$ and WT lenses were cultured under normal serum conditions, the β A3 Δ G91 LECs exhibited significantly higher immunofluorescence of LC3 and LAMP2 (Figs. 7J, 7K) relative to the LECs from normal lenses (Figs. 7G, 7H). Also, the evidence of relatively more yellow puncta in $\beta A3 \Delta G91$ (Fig. 7L) suggests the fusion of LC3 (autophagosomes) and LAMP2 (lysosomes) and a lack of autophagosome degradation. When the cells were grown under serum-starvation conditions to activate autophagy, LC3 and LAMP2 immunofluorescence intensities decreased in both groups of cells compared to normal serum conditions but were still relatively higher in the mutant LECs than the WT LECs (Figs. 7M, 7N, 7P, 7Q). The levels of LAMP2 and LC3 in LECs under normal serum and serumstarvation conditions were quantified as shown in bar graphs on the right side of the immune-stained LECs in Figure 7. The expression levels of LC3 and LAMP2 in lens tissues were also determined by RT-qPCR, which showed greater levels of LAMP2 in β A3 Δ G91 lenses relative to WT lenses (Fig. 7V).

Another criterion used to assess autophagosome formation and degradation is the measurement of LC3-I to LC3-II conversion via western blot analysis. LC3-II is generated by the conjugation of cytosolic LC3-I to phosphatidylethanolamine (PE) to form LC3-II on the surface of nascent autophagosomes.⁵⁰ Results of the western blot analysis using 1-month-old lens homogenates showed that the ratio of LC3-II to LC3-I was higher in the mutant lenses

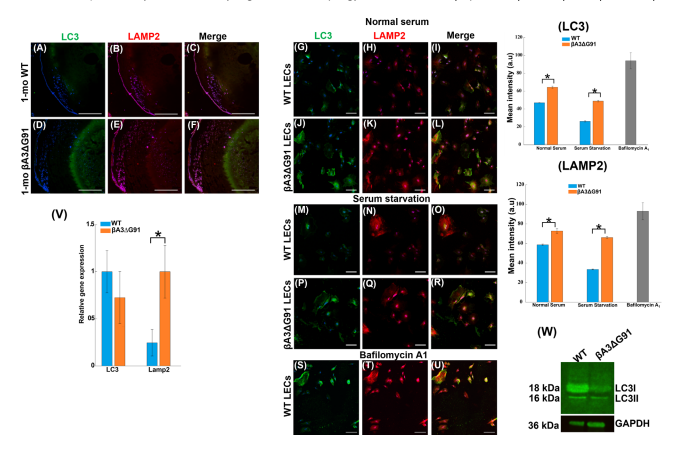


FIGURE 7. Elevated levels of LC3 and LAMP2 in β A3 Δ G91 relative to WT lenses suggest disruption of autophagy. (**A**–**F**) Immunohistochemical analyses of LC3 and LAMP2 in mouse lens sections. LC3 and LAMP2 staining was higher in the β A3 Δ G91 lenses than in the WT lenses (20×). (**G**–**L**) Under normal serum conditions, β A3 Δ G91 LECs had higher LC3 and LAMP2 staining intensities relative to WT LECs. Yellow fluorescence in β A3 Δ G91 indicated the presence of more undigested autolysosomes. (**M**–**R**) When cells were starved of serum, LC3 and LAMP2 were generally higher in the β A3 Δ G91 LECs relative to the WT LECs. (**S**–**U**) When WT LECs were treated with bafilomycin A1, LC3 and LAMP2 immuno-intensities increased. *Scale bar*: 100 µm. (**V**) RT-qPCR analyses showed increased LAMP2 mRNA in the β A3 Δ G91 lenses. **P* < 0.05 versus age-matched WT. (**W**) Western blot analyses with anti-LC3 antibody showed a higher ratio of LC3-II to LC3-I in β A3 Δ G91 relative to WT lenses.

compared to the WT lenses, suggesting normal formation but impeded breakdown of autophagosomes (Fig. 7W).

Autophagic Vesicles Accumulate in β A3 Δ G91 Lenses

The above results showed that autophagy markers were relatively elevated, suggesting impaired autophagy in $\beta A3 \Delta G91$ lenses relative to WT lenses, but further structural assessment of autophagy was carried out by examining the presence of double-membrane autophagic vesicles by TEM analysis. In this study, we comparatively examined by TEM the cortex and nuclear regions of β A3 Δ G91 and WT lenses for the presence of vacuoles, autophagosomes, and autolysosomes. Relative to the WT lens epithelium (Fig. 8A), the β A3 Δ G91 lens epithelium exhibited several vesicles with double and single membranes (Figs. 8B, 8C, red arrows), representing autophagosomes and autolysosomes, respectively, and also large numbers of vacuoles. These vesicles seem to contain either cytoplasmic materials or cellular debris, which were absent in the WT epithelial cells. Double-membrane autophagic vesicles are characteristic of autophagosomes and single-walled vesicles result from the fusion of autophagosomes to lysosomes to form autolysosomes. In the cortex of $\beta A3 \Delta G91$ lenses, the accumulation

of autophagic vesicles was observed (Fig. 8E, red arrows), but just a few virtually empty vesicles were seen in the cortex of WT lenses (Fig. 8D). The large vesicles in the β A3 Δ G91 lens contained heterochromatin, membrane whorls, undegraded organelles, and double membrane-bound structures (Fig. 8F). A lysosome was observed lying at the boundary of the vesicle (Fig. 8F, white arrow). Interestingly, the $\beta A3 \Delta G91$ nuclei were condensed and formed inclusion/apoptotic bodies (Fig. 8E, circular dashes),³⁹ along with disrupted nuclear envelopes and chromatin seepage (Fig. 8E, white arrow). The nuclear region of $\beta A3\Delta G91$ lenses contained several double-membrane vesicles (autophagosomes) and inclusion bodies that possibly fused with the nuclear opaque (cataract) region (Figs. 8H, 8I) but were absent in the same region of WT lenses (Fig. 8G). In summary, the findings of the comparative analyses of the elevated autophagic markers and double-membraned vesicles suggest that autophagy was disrupted in the β A3 Δ G91 lenses relative to WT lenses.

Dysregulation of Autophagy Triggers Apoptosis in β A3 Δ G91 Lenses

Our qPCR analyses showed that, relative to WT lenses, the $\beta A3\Delta G91$ lenses exhibited downregulation of B-cell lymphoma 2 (Bcl-2; results not shown) and the presence

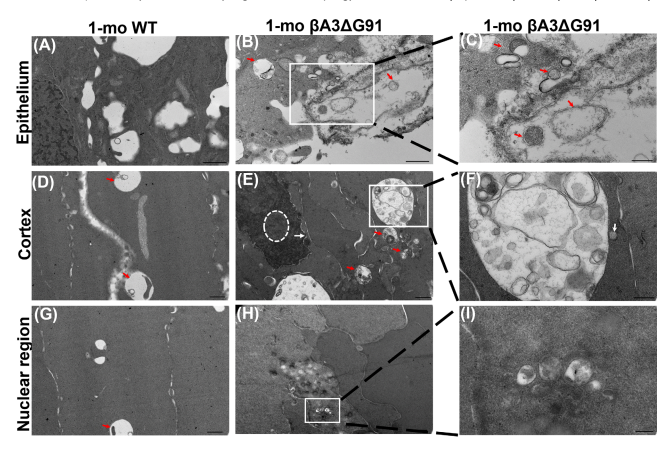


FIGURE 8. TEM analyses showed accumulation of autophagic vesicles in $\beta A3 \Delta G91$ versus WT lenses. (**A**–**C**) Several autophagosomes and autophagosomes were observed in the epithelium of $\beta A3 \Delta G91$ lenses compared to WT lenses. *Scale bar*: 1 µm. (**D**–**F**) Several autophagic vesicles containing cytoplasmic materials, membrane whorls, and heterochromatin were seen in the cortex of $\beta A3 \Delta G91$ but not WT lenses. *Scale bar*: 1 µm; inset, 400 nm. (**G**–**I**) Nuclear remnants and autophagic vesicles fused with the opaque (cataract) region in $\beta A3 \Delta G91$ lenses. *Scale bar*: 1 µm; inset, 200 nm. *Red arrows* indicate autophagosomes or autolysosomes. The *white arrow* points to a lysosome attached to an autophagosome.

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of inclusion/apoptotic bodies, which suggested the possibility of apoptosis in the mutant lenses. Bcl-2 has been shown to be an anti-apoptotic protein that plays a role in regulating cell survival.⁵¹ Additionally, impaired nuclei degradation in the lens can also cause unwanted organelles to build up and become toxic to the lens cells, which could also trigger apoptosis. During apoptosis, the release of activated caspases occurs. Therefore, we carried out a TUNEL assay and determined the levels of cleaved caspase-3 and caspase-9 in the two types of lenses. TUNEL staining of 1-monthold lenses showed more than three times higher apoptotic cells in the outer cortex and the nuclear region in $\beta A3 \Delta G91$ lenses compared to the WT lenses (Figs. 9A, 9B). This finding was supported by elevated levels of cleaved caspase-3 and caspase-9 in the LECs (Figs. 9D, 9E, 9G, 9H) and lens tissues (Figs. 9J, 9K, 9M, 9N) of β A3 Δ G91 lenses relative to WT lenses.

DISCUSSION

The focus of this study was to determine whether the delayed nuclei degradation in lenses of $\beta A3\Delta G91$ mice relative to WT mice is associated with autophagy disruption, and if the disruption is evidenced by the relatively elevated levels of autophagy markers and the presence of double-membrane autophagosomes in $\beta A3\Delta G91$ lenses. The ratio-

nale for the study was as follows. Our previous studies of both the complete β A3/A1-crystallin knockout mouse (β A3KO) model³³ and lens-specific β A3/A1 knockout mouse model $(\beta A3cKO)^{52}$ exhibited accumulation of undegraded organelles in the cortex and autophagy disruption. Also, β A3/A1-crystallin has been reported to be a retinal lysosomal resident protein in astrocytes, RPE cells, and ganglion cells,³⁰⁻³² and it modulates a specific autophagic-signaling pathway via V-ATPase.³¹ Together, these findings suggest that β A3/A1-crystallin and V-ATPase proton pump interaction plays an essential role in regulating and maintaining lysosomal acidic pH. Based on the retinal reports, we have hypothesized that β A3/A1-crystallin acts similarly in the lens, and either its absence in β A3KO and β A3cKO lenses or its dysfunction due to G91-deletion in lenses would disrupt the autophagy process. As stated above, the dysfunctional nature of $\beta A3 \Delta G91$ is supported by in vitro studies of the recombinant β A3 Δ G91, showing its altered structure, precipitation, and inability to oligomerize with other crystallins.^{19,35} More importantly, G91Del in β A3/A1-crystallin resulted in congenital cataract development among families in various countries.13-19

The reason for the dysfunctional β A3 Δ G91 in vivo is that G91 is located at a conserved key site that stabilizes the structures of both β - and γ -crystallins (members of a super family). Both crystallins contain two similar consecu-

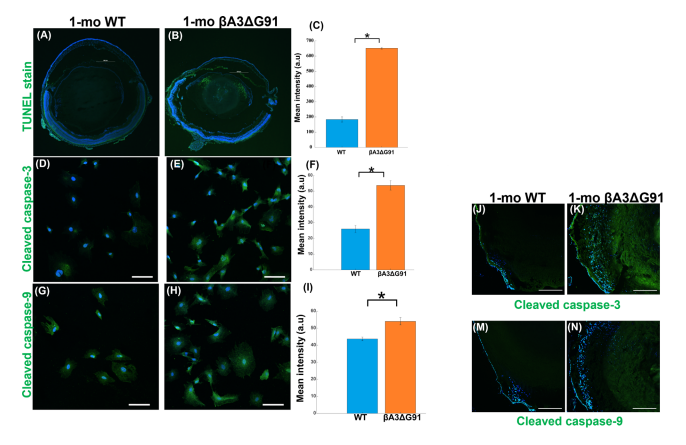


FIGURE 9. Dysregulation of autophagy triggers apoptosis in β A3 Δ G91 lenses. (**A–C**) TUNEL stain. Several TUNEL-positive cells appear as puncta, co-localizing with nuclei in β A3 Δ G91 relative to WT lenses (4×). *Scale bar*: 500 µm. (**D–F**) Expression of cleaved caspase-3 was higher in β A3 Δ G91 relative to WT LECs. (**G–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**G–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression of cleaved caspase-9 was higher in β A3 Δ G91 relative to WT LECs. (**F–I**) Expression in the β A3 Δ G91 lenses compared to the WT lenses. (**M**, **N**) Higher immunofluorescence intensity for cleaved caspase-9 in the β A3 Δ G91 relative to the WT lenses.

tive N-terminal domains with Greek key motifs 1 and 2. Similarly, the C-terminal domain contains Greek key motifs 3 and $4.^{19,53,54}$ Each Greek key motif is comprised of four consecutive β strands (a–d) that intercalate to form two β sheets.¹⁹ The G91 deletion in β A3-crystallin results in the loss of G91 at the edge of strand c in the second motif of the N-terminal domain, affecting the conserved structural feature known as the tyrosine corner. This tyrosine corner is known to stabilize the connection between b and c β strands and is present in most β -sandwich domains, where they appear to stabilize the fold topology in β - and γ -crystallins.^{53,54} Therefore, G91 deletion leads to major structural destabilization rendering a dysfunctional β A3-crystallin, as evidenced by the results of recombinant β A3-crystallin.^{19,35}

 β A3 Δ G91 mice developed congenital nuclear cataract (Fig. 5B), which potentially results from (1) attenuation of nuclei and mitochondrial degradation in the lens inner cortex due to autophagy disruption, (2) cellular apoptosis, or (3) β A3 Δ G91 insolubilization. Relative to WT lenses, the undegraded nuclei were seen in the OFZ of β A3 Δ G91 lenses during nuclear staining by H&E and DAPI staining (Fig. 3) and during TEM analysis of lens sections (Fig. 4). Interestingly, the relative expression of HSPA5 (an ER marker) in β A3 Δ G91 lenses was rather decreased, which could be indicative of an impaired protein quality control system. Evidence shows that, when autophagy is disrupted, there is a higher burden of misfolded proteins, which can over-

whelm the protein folding and quality control machinery.⁵⁵ Because HSPA5 is a chaperone that facilitates the clearance of misfolded proteins, its decreased levels may indicate an inability to cope with the increased load of misfolded proteins, which may be linked to autophagy disruption.^{55,56} The higher ubiquitination of proteins in β A3 Δ G91 lenses relative to WT lenses provides further evidence for the increased levels of damaged/unfolded proteins in the mutant lenses.

Because past reports and our recent reports of β A3KO³³ and β A3cKO⁵² mouse models have implicated the attenuation of lens organelle degradation in autophagy, this prompted us to investigate in this study the effects of dysfunctional β A3 Δ G91 on autophagy in the lenses of β A3 Δ G91 versus WT mice. The following relative results suggest autophagy disruption in the β A3 Δ G91 lenses: (1) attenuation of nuclei and mitochondrial degradation (Figs. 3, 4), (2) elevated autophagy markers (Figs. 6–8), (3) accumulation of autophagosomes and autolysosomes with undigested organelles (Fig. 8), (4) activated cellular apoptosis (Fig. 9), and (5) β A3-crystallin insolubilization (Table 2).

 β A3 Δ G91 lenses showed relatively higher levels of ubiquitinated proteins (Fig. 6). The ubiquitination of target proteins (a posttranslational modification process) acts either as a signal for the substrates to be degraded by proteasome or lysosome or as an indicator for modulating their non-proteolytic processes.⁴⁴ This is evidenced by reported crosstalk between autophagy and the ubiquitin-proteasome system.⁵⁷ Relative to WT lenses, p62 was upregulated at both transcriptional and translational levels in both $\beta A3 \Delta G91$ lenses and LECs (Figs. 6, 7). The increased levels of p62 are indicative of defective autophagic flux in various diseases, and p62 is also an autophagy substrate that is used as a reporter of autophagic activity.43 Similarly, LC3-II levels were increased in β A3 Δ G91 lenses relative to WT lenses (Fig. 7). LC3-II, a standard marker for autophagosomes, is generated by the conjugation of cytosolic LC3-I to PE on the surface of nascent autophagosomes.^{46,47} As LC3 is specifically associated with autophagosomes and autolysosomes, quantifying LC3-positive puncta is considered a gold-standard assay for assessing the numbers of autophagosomes in cells.⁴⁶ LAMP2 was also relatively upregulated in β A3 Δ G91 lenses and LECs (Fig. 7). LAMP1 and LAMP2 comprise 50% of all lysosomal membrane proteins, and are thought to be responsible in part for maintaining lysosomal integrity, pH, and catabolism. LAMP2 gene mutation causes lysosomal storage disease in humans.^{46,49} In embryonic fibroblasts, disruption of both LAMP1 and LAMP2 is associated with Danon disease, including accumulation of autophagic vacuoles in heart and skeletal muscles.58 Similar vacuoles were also observed in the epithelial cells of $\beta A3 \Delta G91$ lenses relative to those of WT cells (Fig. 8). The autophagosomes, large double-membrane vesicles enclosing cytoplasmic components that are targeted for degradation during autophagy, were also observed during TEM analysis of β A3 Δ G91 lenses (Fig. 8). The double-membrane autophagosomes are the hallmark of the catabolic pathway during autophagy.³⁸ Interestingly, the TEM images in Figure 8 show the presence of autophagosomes in the epithelium, cortex, and nuclear region. One intriguing finding from TEM images was the fusion of autophagosomes with the nuclear opaque region, representing its potential role in cataract development. Currently, the identity of the autophagosomes that fuse with the opaque region is unknown, and whether the fusion plays a role in the nuclear opacity development in the $\beta A3 \Delta G91$ lenses is also unknown. This led to our hypothesis that the opacity development in β A3 Δ G91 lenses is associated with non-degradation of autophagosomes which might be due to lack of V-ATPase activity to acidify lysosomes. Such lysosomal acidification is a normal process by the native β A3-crystallin in the retina.³¹

The lenses of $\beta A3\Delta G91$ mice also exhibited activated apoptosis in the nuclear region relative to WT lenses. IHC analyses showed that cleaved caspase-3 and caspase-9 were relatively upregulated in the $\beta A3\Delta G91$ LECs (Fig. 9). Apoptosis has been suggested as a decisive process for cell death during autophagy.⁵⁹ Because caspase-9 on activation cleaves caspase-3 and caspase-7, which are effector caspases that carry out apoptosis via DNA fragmentation, cell shrinkage, and membrane blebbing, it is difficult to determine the levels of the two caspases by western blot analysis. Therefore, the use of IHC is preferable to western blot analysis for determining the levels of caspase-3 and caspase-9.

In summary, the results from our study provide evidence that our novel mouse model shares a similar phenotype with $\beta A3\Delta G91$ human patients. Similar to total and conditional *CRY* $\beta A1$ knockout mice, $\beta A3\Delta G91$ lenses showed impaired autophagy. Based on the association of $\beta A3\Delta G91$ with autophagy dysregulation, further investigations are warranted to explore the potential role of *CRY* $\beta A1$ as an autophagy regulator via the V-ATPase modulator in the lens.

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