Increased Association of Deamidated αA-N101D with Lens Membrane of Transgenic αAN101D vs. Wild Type αA Mice: Potential Effects on Intracellular Ionic Imbalance and Membrane Disorganization

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Research article

Keywords: Lens, Crystallins, Deamidation, Post-translational modifications, Transgenic Mice, Cataract

DOI: https://doi.org/10.21203/rs.2.17769/v3

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Abstract

We have generated two mouse models, in one by inserting the human lens αAN101D transgene in CRYαA_{N101D} mice, and in the other by inserting human wild-type αA-transgene in CRYαA_{WT} mice. The CRYαA_{N101D} mice developed cortical cataract at about 7-months of age relative to CRYαA_{WT} mice. The objective of the study was to determine the following relative changes in the lenses of CRYαA_{N101D} vs. CRYαA_{WT} mice: age-related changes with specific emphasis on protein insolubilization, relative membrane-association of αA_{N101D} vs. WTαA proteins, and changes in intracellular ionic imbalance and membrane organization.

Methods: Lenses of varying ages from CRYαA_{WT} and CRYαA_{N101D} mice were compared for an age-related protein insolubilization. The relative lens membrane-association of the αAN101D- and WTαA proteins in the two types of mice was determined by immunohistochemical-, immunogold-labeling-, and western blot analyses. The relative levels of membrane-binding of recombinant αA_{N101D}- and WTαA proteins was determined by an *in vitro* assay, and the levels of intracellular Ca^{2+} uptake and Na, K-ATPase mRNA were determined in the cultured epithelial cells from lenses of the two types of mice.

Results: Compared to the lenses of CRYαA_{WT}, the lenses of CRYαA_{N101D} mice exhibited: (A) An increase in age-related protein insolubilization beginning at about 4-months of age. (B) A greater lens membrane-association of αAN101D- relative to WTαA protein during immunogold-labeling- and western blot analyses, including relatively a greater membrane swelling in the CRYαA_{N101D} lenses. (C) During *in vitro* assay, the greater levels of binding αAN101D- relative to WTαA protein to membranes was observed. (D) The 75% lower level of Na, K-ATPase mRNA but 1.5X greater Ca^{2+} uptake was observed in cultured lens epithelial cells of CRYαA_{N101D} than those of CRYαA_{WT} mice.

Conclusions: The results show that an increased lens membrane association of αA_{N101D}-relative WTαA protein in CRYαA_{N101D} mice than CRYαA_{WT} mice occurs, which causes intracellular ionic imbalance, and in turn, membrane swelling that potentially leads to cortical opacity.

Background

Although the cornea is the primary refractive tissue performing 70-80% of refraction of the eye, the major function of the lens is in accommodation and to partly help in the refraction. The lens accommodative function gradually diminishes with age and is almost completely lost at age of > 50 years. The lens transparency plays an important role in focusing light on to the retina, but this role is gradually lost as it develops age-related opacity. Several unique factors maintain lens transparency for up to > 60 year of our lifetime. These include: cellular homeostasis among only two types of cells (epithelial and fiber cells) [1], an orderly terminal differentiation of epithelial to fiber cells with precise organelles loss [2], the unique interactions among crystallins [3], with almost no protein turnover [4], the specialized lens metabolism [5], specific interactions among α-crystallin and membrane [6], the precise maintenance of intracellular and
extracellular ionic concentrations [7], the low levels of cellular water and oxygen in the lens inner cortex and nuclear regions [8], and a unique membrane lipid composition [9]. Alterations among some of these lens unique factors play direct or indirect roles in pathogenesis of cataracts (e.g., pediatric- and age-related cataracts). However, additional cataract-causative factors are also identified, which include mutations in crystallins [10], oxidative insults of crystallins, the loss of redox balance of glutathione [11], extensive truncations of α-, β-, and γ-crystallins [12-20], a variety of post-translational modifications with deamidation as being the most abundant [21-25], and the loss of membrane integrity [7, 26, 27]. These factors individually or in combination also cause lens opacity through altered lens cellular structures and contents, ionic imbalance, increased water and oxygen levels, loss of natural interactions among crystallins, and crystallins’ unfolding, degradation and cross-linking.

Our focus in this study is the potential roles of deamidation of Asn$_{101}$ of αA crystallin to Asp that introduces negative charges and shown to alter their hydrophobicity, tertiary structures, crystallin-crystallin interactions, and leads to aggregation and cross-linking [21-27]. In this study, the deamidation of Asn$_{101}$ to Asp in in a mouse model was studied to determine phenotypic and molecular changes within the lens due to deamidation of a single nucleotide change in CRYAA crystallin gene. This site was chosen because our past study showed that only deamidation of Asn localized at specific sites in crystallins (e.g., deamidation of N101 but not of N123 residues in αA-crystallin [24], and of N146 but not of N78 of αB-crystallin) exhibited the above-described deamidation-induced effects [25]. To show the potential effects of deamidation in vivo, we have generated mouse models by inserting the human lens αA-N101D transgene in CRYαA$_{N101D}$ mice, and human lens wild-type αA-transgene in CRYαA$_{WT}$ mice (to act as a control). The CRYαA$_{N101D}$ mice developed cortical cataract at about 7-months of age relative to CRYαA$_{WT}$ mice [28, 29]. This model showed for the first time that in vivo expression of the deamidated αAN101D caused cortical lens opacity, which was due to the disruption of fiber cell structural integrity and protein insolubilization as aggregation [28]. The comparative RNA sequencing and Ingenuity Pathway Analyses (IPA) of lenses from 2- and 4-months old CRYαA$_{N101D^-}$ and CRYαA$_{WT}$ mice showed that the genes belonging to cellular assembly and organization, cell cycle and apoptosis networks were altered in αA$_{N101D}$ lenses [29]. This was accompanied with several cellular defects in αA$_{N101D}$ lenses that included defective terminal differentiation (increased proliferation and decreased differentiation) of epithelial cells to fiber cells, and reduced fiber cells denucleation and expressions of Rho A and Na, K-ATPase (the major lens membrane-bound molecular transporter) [29]. The findings also suggested the potential role of lens intracellular ionic imbalance as the major reason for the development of cataract [29]. The above findings suggested that the altered intracellular ionic imbalance could be due to potential loss of membrane integrity that caused cortical opacity at about 7-months of age in the CRYαA$_{N101D}$ mouse model. Therefore, the focus of the present study was to determine whether an increased membrane-association of αA$_{N101D}$ potentially compromises membrane integrity, causes an ionic imbalance, and leads to cataract development.

**Methods**
Ethics Statement

All animal experiments were performed per protocols approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Alabama at Birmingham (Protocol no. 130208393). Mice were housed in a pathogen-free environment at the facility of the University of Alabama at Birmingham.

Materials

Unless stated otherwise, the molecular biology-grade chemicals were purchased from Millipore-Sigma (St. Louis, MO, USA) or Fisher (Atlanta, GA, USA) companies. The Rabbit polyclonal anti-human aquaporin-0 (AQP0) antibody were purchased from Alpha Diagnostics (San Antonio, TX, USA). Additional commercial sources of various chemicals and antibodies used in the study are described throughout the text.

Generation of Transgenic Mice

The mouse model that expresses a human αA-crystallin gene in which Asn-101 was replaced with Asp is referred to as α\(\text{AN}_{101}\)D-transgenic mouse model. This model has been considered to be “deamidated” in this study, and the mice expressing αAN101D-transgene is referred here as CRYα\(\text{AN}_{101}\)D mice. Both mouse models (human lens α\(\text{AN}_{101}\)D transgenic- and human wild-type αA-transgenic mouse models were generated in Dr. Om Srivastava’s laboratory [28]. αAN101D protein expression constituted about 14% and 14.2% of the total αA in the lens WS-and WI-proteins of the α\(\text{AN}_{101}\)D transgenic mice, respectively [28]. The mouse lenses were extracted after the mice were euthanized using the CO\(_2\) procedure as per approved method by the Institutional Animal Care and Use Committee (IACUC) of the University of Alabama at Birmingham (Protocol no.130208393). Adult (2–3 months) wild type mice (C57BL6) were obtained from the university breeding colony. Animals were kept under a 12/12 h light–dark cycle and had ad libitum access to food and water. We have used three mice from each group of CRYα\(\text{A}_{\text{WT}}\) mice control and α\(\text{AN}_{101}\)D mice in all the experiments described below.

Isolation of Water Soluble (WS)- and Water Insoluble (WI)-Proteins from Mouse Lenses

The WS- and WI-protein fractions from lenses of desired ages of CRYα\(\text{A}_{\text{WT}}\) and CRYα\(\text{AN}_{101}\)D mice were prepared as previously described by us [28]. All procedures were performed at 5°C unless specified otherwise. The lenses were removed under a dissecting microscope and placed in 5°C-cold buffer A (5 mM Tris-HCl, 1 mM EDTA, 1 mM DTT, pH 7.8, and protease inhibitor cocktail [Roche Life Science, Indianapolis, IN]), and centrifuged at 14,000 x g for 15 min at 5°C to separate the WS- and WI- protein fractions. The supernatant (WS-protein fraction) was collected, and next the pellet (WI-protein fraction) was resuspended in buffer A, centrifuged as above. The recovery of WS- and WI-protein fractions was repeated twice after centrifugation, and the WS supernatants after each centrifugation steps were pooled. The final WI-protein pellet was solubilized in 5 mM Tris-HCl, pH 7.5, containing 8 M urea, 5 mM EDTA, and 5 mM EGTA. The 8 M urea concentration was diluted to 4 M urea with buffer A prior to centrifugation as above. The protein concentrations in these fractions were determined by using a kit (Pierce Biotechnology-Thermo Fisher) using bovine serum albumin as a standard.
Membrane Isolation from Mouse Lenses

The membranes from lenses of 1- and 6-month-old CRYαA<sub>WT</sub> and CRYαA<sub>N101D</sub> mice were prepared as described previously [30, 31]. Lenses of identical ages from both types of mice were homogenized in buffer B (0.05 M Tris-HCl, pH 8.0 containing 5 mM EDTA, 1 mM DTT, 150 mM NaCl, and protease inhibitor cocktail [Roche, Indianapolis, IN]), and the preparations were centrifuged at 100,000 x g for 30 min using Beckman TL 100 centrifuge with a TLA 100.3 rotor. The supernatant was collected, and the pellets were washed twice with the above buffer B and centrifuged as above. This was followed by three additional washes with buffer B containing 8 M urea and centrifugation as above after each wash. Next, the pellet was washed twice with water and centrifuged as above. The pellet was then washed with 0.1 M cold (5°C) NaOH [30, 31]. A final wash of pellet was with water and centrifugation as above to recover the purified lens membrane preparations as pellets.

Purification of Recombinant WTαA- and aA<sub>N101D</sub>-crystallins, their Conjugation with Alexa Fluor 350 and Membrane Binding

The WTαA- and aA-N101D mutant proteins were expressed in E.coli and purified by a Ni-affinity column chromatographic method as previously described by us [28]. Each protein was labeled with Alexa-350 using a protein labeling kit as suggested by the manufacturer (Molecular Probes, Carlsbad, CA). The binding of Alexa 350-conjugated recombinant WT αA- and aA-N101D mutant proteins to mouse lens membrane (isolated from C57BL non-transgenic mice)) was determined as previously described [32, 33]. During the binding assay, the purified lens membrane (containing 2.5 mg protein; isolated from 1 to 3-month old non-transgenic C57 mice) was incubated with increasing but identical concentrations of either Alexa-labelled WT αA- or aA<sub>N101D</sub> proteins at 37°C for 6 h. Next, the incubated preparations were centrifuged at 14,000 X g and the supernatant and pellet (membrane fraction) recovered. After washing the membrane fraction with water and centrifugation as above, the relative levels of fluorescence of membranes incubated with WT αA- and aA<sub>N101D</sub> mutant proteins were determined using Perkin Elmer Multiplate Reader (Model Victor1420-04).

Determination of Intracellular Ca<sup>2+</sup> in Epithelial Cells in Culture from Lenses of CRYαA<sub>WT</sub> - and CRYαA<sub>N101D</sub> Mice

To culture epithelial cells, six 5-months old lenses from CRYαA<sub>WT</sub> - and CRYαA<sub>N101D</sub> mice were excised and incubated with 0.25% trypsin at 37°C for 2.5 h in an incubator with 5% CO<sub>2</sub>-humified air. Next, the lens cells in trypsin solution were centrifuged at 1200 rpm for 3 min, and trypsin (in the supernatant) was discarded. The lens epithelial cells (recovered as pellet) were suspended in medium 199 (Thermo Fisher Scientific, Grand Island, NY) containing 10% fetal calf serum (Hyclone, Logan, Utah) and 1% antibiotic-antimycotic solution (Thermo Fisher Scientific, Grand Island, NY) in 12-well plates (Corning, Franklin Lakes, NJ). After 24 h, the unattached cells were discarded by washing with the above medium. The old medium was replaced with fresh medium after every 48 h, and the cells were allowed to grow for 7 to 10 days until confluent. Next, the confluent cells were trypsinized and seeded in 12-well plates for
intercellular Ca\(^{2+}\) determination and were allowed to grow for 24 h. The cells were washed with medium 199 without phenol red, incubated in calcium orange dye (Thermo Fisher Scientific, Grand Island, NY) at a final concentration of 4 μM for 30 min at room temperature as instructed in the manufacturer's protocol. After 30 min, the cells were washed with the above medium, and Ca\(^{2+}\) indicator was examined under a microscope (Leica DMI 4000B) using a Texas Red filter.

**Western Blot and Immunohistochemical Analyses**

The WS- and WI-proteins and membrane fractions isolated from lenses were analyzed for their immunoreactivity with anti-aquaporin-0 antibody (to visualize the membrane intrinsic protein), and Mouse anti-His monoclonal antibody ([Novagen, Madison, WI], to visualize WT\(\alpha\)A and \(\alpha\)\(_{N101D}\)) during Western blot analysis. The SDS-PAGE analysis was carried out as described by Laemmli [34].

The confocal immunohistochemical analysis of lens axial sections of WT\(\alpha\)A and \(\alpha\)\(_{N101D}\) was carried out as previously described by us [28]. The analysis was performed at the High-Resolution Imaging core facility of the University of Alabama at Birmingham.

**Localization by Immunohistochemical-Transmission Electron Microscopic Method**

The analysis was performed at the High-Resolution Imaging core facility of the University of Alabama at Birmingham. His-tagged \(\alpha\)\(_{WT}\)- and \(\alpha\)\(_{N101D}\)-crystallins were localized in lens cells by an Aurion immunogold method and the reagents used were Aurion Conventional Immunogold reagents (Electron Microscopy Science [PA]). Lenses of desired ages were fixed in phosphate-buffered saline, pH 7.4 containing 4% paraformaldehyde and 0.05% glutaraldehyde (Electron Microscopy Sciences, Hatfield, PA) for 2 h at room temperature, and then overnight at 4°C. The fixed lenses were washed with water (Millipore, Billerica, MA). Samples were dehydrated by ascending ethanol gradient series followed by infiltration overnight at 4°C with absolute ethanol: London Resin (LR) white (1:1). Next, the samples were incubated overnight with pure LR white resin on a rotating platform. The lenses were removed and transferred to gelatin capsules containing fresh LR white and allowed to polymerize for 24 h at 45-50°C. Ultra-thin (silver gold to light gold) LR white lens sections were collected on nickel mesh grids. The color of sections was silver-gold to light gold, and based on their color, the thickness was estimated to between 70-80 nm. For immunogold-labeling, the protocol as described in Electron Microscopy Sciences (Hatfield, PA) was precisely followed. To inactivate aldehyde group's present after aldehyde fixation, the samples on grids were incubated on 0.05 M glycine in PBS buffer for 10-20 minutes. Next, the grids were transferred onto drops of the matching Aurion blocking solution for 15 min, and then were washed for 15 min in incubation solution (PBS containing 0.1% bovine serum albumin and 15 mM NaN3, pH 7.3). This was followed by a 2X wash in incubation buffer, each time for 5 min. The grids were incubated with two primary antibodies (Mouse anti-His monoclonal antibody and Rabbit anti-aquaporin-0 polyclonal antibody for 1 h. In controls, the primary antibodies were omitted. The grids were then washed 6X (5 minutes each time) with the incubation solution and transferred to following secondary antibody conjugates {(goat anti-Rabbit EM grade conjugate 25 nm diameter) and (goat anti- mouse EM grade
conjugate 10nm diameter) and were incubated for 30 minutes to 2 h. The grids were washed on drops of incubation solution for 6X (5 min each time). The grids were washed twice with PBS for 5 min, post-fixed in 2% glutaraldehyde in PBS for 5 min, and finally washed with distilled water and contrasted according to standard procedures. Lens sections were imaged using a FEI 120kv Spirit TEM (FEI-Thermo Fisher), and images were collected using an AMT (AMT-Woburn, MA) digital camera.

**RNA Extraction and Reverse Transcription-Quantitative Polymerase Chain Reaction (RT-PCR [qPCR])**

RNA was extracted with Trizol reagent (Invitrogen) from cultured lens epithelial cells from CRYαA_{N101D} and CRYαA_{WT} mice, and all the samples were analyzed in triplicates. Real-time PCR quantifications were performed using the BIO-RAD iCycler iQ system (Bio-Rad, Hercules, CA), using a 96-well reaction plate for a total volume of 25 μL. RNA was extracted as described above. Primers were designed using Primer3 for the following genes:

- **Atp1a2 Forward-** 5'CGGGAGCCATAAGGGTTTGT 3', and **Atp1a2 Reverse-** 5'GCACTGACTTGGCTGTTGTG 3'

The ACTB gene was used for normalization. The reaction mixture included 12.5 μL of Real-Time SYBR Green PCR master mix, 2.5 μL of reverse transcription product, 1 μL of forward and reverse primer and 8 μL of DNase/RNase free water. The reaction mixtures were initially heated to 95°C for 10 min to activate the polymerase, followed by 40 cycles, which consisted of a denaturation step at 95°C for 15 sec, annealing at 55°C for 60 sec and an elongation step at 72°C. The qRT-PCR data were analyzed by the comparative ΔCt method.

**Results**

**Age-Related Protein Insolubilization in Lenses of CRYαA_{N101D} and CRYαA_{WT} Mice**

To determine at what age there is change in the protein profiles in lenses of CRYαA_{N101D} and CRYαA_{WT} mice occurred, a comparative analysis of WS-proteins and WI-proteins from the lenses of the two types of mice of different ages was carried out (Fig. 1). The WS- and WI-proteins from lenses of different ages (1-, 3-, 4-, 5- and 7-months) were analyzed by SDS-PAGE. The WS-protein profiles from the lenses of the CRYαA_{N101D} and CRYαA_{WT} mice were almost identical until 3-months of age, except lens preparations from ages of 4-, 5- and 7-months of CRYαA_{N101D} mice exhibited relatively greater levels of aggregated protein of M_r >30 kDa and higher relative to same-aged lenses from CRYαA_{WT} mice (Lanes 4 and 5 in Figure 1B). Additionally, on quantification, the relatively increasing levels of WS-proteins showed age-related water insolubilization beginning at 4-months of age in the lenses of αA_{N101D} mice (Table 1). Between 4- to 7-months of age, relatively about 5 to 10% higher proteins became insoluble in lenses of CRYαA_{N101D}. To determine changes in individual crystallins due to their insolubilization, the WS-protein fraction from 7-month-old lenses was fractionated by a size-exclusion HPLC using a G-4000PWXL column (Tosoh Biosciences, fractionation range of protein with M_r's between 1X10^4 to 1X10^7 Da). The comparative protein elution profiles at 280 nm of 7-month old lenses of αA_{N101D}-mice showed an
increased protein in the void volume peak (representing WS-HMW proteins), and reduced β- and γ-crystallin peaks relative to lenses of CRYαA<sub>WT</sub> mice (the differences shown in green in Figure 2A). The void volume peak in WS-protein fraction was also higher in the 7-month old lenses relative to 1-month old lenses of α<sub>N101D</sub>-mice (Results not shown), suggesting a relatively increased HMW protein aggregate formation with aging. On western blot analysis of the individual column fractions nos. 6 to 9 (constituting the void volume-HMW-protein peak) with an anti-His antibody, the levels of His-immunoreactive protein were higher in 7-month old CRYαA<sub>N101D</sub> lenses compared to the identical aged CRYαA<sub>WT</sub> lenses (Figure 2B). Additionally, because the immunoreactive peak in the WT lenses was in the fractions no. 8 and 9, whereas it was in the fractions no. 7 and 8 in the α<sub>N101D</sub> lenses that suggested that the HMW proteins of α<sub>N101D</sub> lenses showed a high molecular weights relative to the HMW proteins from WT lenses. On quantification of Western blot images with Image J (Figure 2C); the intensity of the immunoreactive HMW proteins of α<sub>N101D</sub> was about 20% greater relative to WT lenses. Together, the results suggested a greater aggregation with higher M<sub>r</sub> of the HMW-proteins in CRYαA<sub>N101D</sub> lenses compared to the identical aged CRYαA<sub>WT</sub> lenses.

Identification Proteins Present in Water Insoluble-Urea Soluble (WI-US) - and Water Insoluble Urea Insoluble (WI-UI) Protein Fractions of Lenses of CRYαA<sub>WT</sub> and CRYαA<sub>N101D</sub> Mice

To identify the insolubilized proteins in WTαA vs. α<sub>N101D</sub> lenses, the WI-proteins from 5-month-old mice were further fractionated into WI-US- and WI-UI-protein fractions and were examined by SDS-PAGE (Figure 3), followed by their protein compositional analysis by mass spectrometry. SDS-PAGE analysis showed that both WI-US- and WI-UI-protein fractions from CRYαA<sub>N101D</sub> lenses contained greater levels of protein species including aggregated proteins (M<sub>r</sub> > 30 kDa) [Identified as ‘a’, and ‘c’ in Figure 3] relative to the same fractions from lenses of CRYαA<sub>WT</sub> mice (Identified as ‘b’ and ‘d’ in Figure 3). The mass spectrometric analysis was carried out at the following two levels: (i) In the first level analysis, determination of the total protein compositions in the WI-US- and WI-UI protein fractions of the two types of lenses (Supplemental Tables A [Comparative protein compositions of WI-US-fractions of CRYαA<sub>N101D</sub> and WTαA lenses], and Supplemental Table B [Comparative protein compositions of WI-UI-fractions of α<sub>N101D</sub> and WTαA lenses]). (ii) In the second level analysis, the protein compositions of protein aggregates (M<sub>r</sub> > 30 kDa) in WI-US-fraction of αAN101D lenses (Identified as ‘a’ in Figure 3), and WI-US-protein fraction of WTαA lenses (Identified as ‘b’ in Figure 3) [Supplemental Table C]. Similarly, the compositions of protein aggregates (M<sub>r</sub> > 30 kDa) in WI-UI-fraction of α<sub>N101D</sub> lenses (Identified as ‘c’ in Figure 3), and WI-US-fraction of WTαA lenses (Identified as ‘d’ in Figure 3) were determined [Supplemental Table D]. The rationale of the two levels of analysis was to determine the relative proteins compositions due to the greater insolubilization of proteins in CRYαA<sub>N101D</sub> lenses relative to CRYAAWT lenses (Figure 1, Table1). Our expectation was that the level 1 comparative examination would identify the total proteins that underwent insolubilization, and existed in the US- and UI-protein fractions, whereas the level 2 analysis would selectively identify those proteins that formed aggregates (M<sub>r</sub> > 30 kDa) in the
US- and UI-fractions. The rationale was that the information would implicate potential roles of specific crystallins in the aggregation and therefore, in the cataractogenic mechanism.

(i) Comparative Protein Compositions in WI-US Fractions of Lenses from CRYαA<sub>N101D</sub> and CRYαA<sub>WT</sub> Mice

The proteins detected in the WI-US-protein fractions of CRYαA<sub>N101D</sub> lenses but were absent in the WT lenses are described in Supplemental Table A. Together, the results show that the WI-US fraction of CRYαA<sub>N101D</sub> lenses was enriched in several histones, which could be due to the lack of denucleation relative to WT lenses. Absence of Retinal dehydrogenase in transgenic lens fraction.

(ii) Comparative Protein Compositions of WI-UI-Fractions of Lenses from CRYαA<sub>N101D</sub> and CRYαA<sub>WT</sub> Mice

The proteins present in the WI-UI-protein fractions of CRYαA<sub>N101D</sub> lenses but were absent in WT lenses are described in Supplemental Table B. In summary, the results again show that the majority of histones that existed in CRYαA<sub>N101D</sub> lenses were absent in the WT lenses, which could be due to the lack of denucleation in the lenses of former mice. Also, specifically αB- and βB2-crystallin became insoluble as their levels were higher even in the WI-UI-fraction of lenses of CRYαA<sub>N101D</sub> relative to WT lenses.

(iii) Compositions of Aggregated Proteins (M<sub>r</sub> >30 kDa) in WI-US- and WI-UI-Fractions of Lenses from CRYαA<sub>N101D</sub> and CRYαA<sub>WT</sub> Mice

As noted above, the purpose of the second level of mass spectrometric analysis was to elucidate the comparative compositions of aggregated proteins (M<sub>r</sub> >30 kDa) in WI-US- and WI-UI-protein fractions of CRYαA<sub>N101D</sub> and CRYαA<sub>WT</sub> lenses [Supplemental Tables C and D]. On comparison, the major proteins present as aggregates (M<sub>r</sub> > 30 kDa) in WI-US fraction of CRYαA<sub>N101D</sub> but absent in CRYαA<sub>WT</sub> were (Supplemental Table C): βB3- and γC-crystallins, collagen alpha-1(IV) chain and -alpha-2(IV) chain and nestin. In contrast, the exclusively present major proteins in WI-US fraction of CRYαA<sub>WT</sub> were: γC-, γD-, γE-, γF-crystallins. The above list describes the selective proteins that were water insoluble-urea soluble and became the part of the complexes with M<sub>r</sub> > 30 kDa in CRYαA<sub>N101D</sub> lenses. The greater abundance of αA-, and βB1-crystallins in the aggregated form suggested their potential involvement in the aggregation process along with βB3- and γC-crystallins.

On comparison of major proteins that existed in WI-UI protein fraction as > 30 kDa aggregates in CRYαA<sub>N101D</sub> not in the CRYαA<sub>WT</sub> included [Supplemental Table D]: γB-, γD- and γE-crystallins, and nestin. In the WI-UI fraction, the greater abundance of proteins in CRYαA<sub>N101D</sub> compared to CRYαA<sub>WT</sub> were: αA-crystallin and lens fiber intrinsic proteins. Together, the results showed that the proteins that remained urea insoluble and were possibly associated with the membrane of CRYαA<sub>N101D</sub> lenses included: γB-, γD- and γE-crystallins, and nestin (Nestin is an intermediate filament protein).
Increased Association of αAN101D with Lens Membrane in the Outer Cortical Fiber Cells relative WTαA in CRYYAAWT lenses

Our previous report [28] showed an increased levels and abnormal deposition of αA_N101D within the outer cortical region in CRYαA_N101D lenses compared CRYαA_WT lenses. This suggested a relatively greater membrane binding of αA_N101D, which was further investigated in experiments as described below.

(i) Immunohistochemical Analyses of Lenses from CRYαA_N101D and CRYαA_WT Mice

The purpose of the experiments was to determine relative levels of αAN101D and WTαA in the outer cortical regions of CRYαA_N101D vs. CRYαA_WT lenses. This was examined by immunohistochemical analysis of 5-months old lenses of the two types of mice using anti-His monoclonal (for detection of WTαA and αA_N101D [green fluorescence]) - and polyclonal anti-aquaporin 0 (for membrane detection [red fluorescence])-antibodies (Figure 4). The axial sections (at 10X magnification) showed an irregular and greater deposition of His-tagged αA (Green) in the lens outer cortex of CRYαA_N101D mice (Shown by an arrow in Figure 4A) relative to CRYαA_WT mice (Shown by an arrow in Figure 4B). Similarly, the equatorial sections (at 40X magnification) also exhibited a greater immunoreactive green fluorescence in the outer cortex of the CRYαA_N101D lens relative to the CRYαA_WT lens (shown by arrows in Figure 4C and D). Together, the results suggested the abnormally greater levels of association of αA_N101D in the outer cortical regions, and potentially with the fiber cell membranes in the CRYαA_N101D lenses relative to those of CRYαA_WT lenses.

(ii) Relative Membrane-Association of WTαA- and αA_N101D in Lenses of CRYαA_N101D and CRYαA_WT Mice

The rationale for the next experiment was that if the greater membrane-association of αA_N101D occurs in vivo in CRYαA_N101D lenses compared to CRYαA_WT lenses, the difference in their levels could also be determined in the purified membrane fractions by western blot analysis. The expectation was that following the step-wise membrane purification by using 8M urea (to dissociate non-covalently-bound membrane proteins), and by the final wash with 0.1N NaOH (to remove non-membranous extrinsic proteins) [30, 31], the purified membrane would show relative levels of membrane-association of αA_N101D vs. WTαA in the two types of mice. To normalize the levels of the relative association during the membrane preparations, two lenses of 1-month-old and two lenses from 6-month old from CRYαA_N101D and CRYαA_WT mice were identically processed, using identical volumes of buffers at each steps during membrane purification (See Methods). Next, Western blot analysis using anti-His- and anti-aquaporin 0-antibodies were used to determine the relative levels of membrane-association of WTαA and αAN101D at different purification steps (Results not shown). To simplify the western blot results of fractions recovered during the sequential steps of membrane purification, only the results of immunoblots with anti-His antibody but not with anti-aquaporin-0 are shown in Figure 5. However, the western blot profiles with anti-aquaporin-0 were almost identical to anti-His antibody results. Relative membrane-association of WTαA- and αAN101D proteins in lenses of CRYαA_N101D and CRYαA_WT mice. A, B, E and F show SDS-
PAGE analysis followed by Coomassie blue-stained gels exhibiting relative levels of protein bands in preparations at different membrane purification steps in lenses at two different age groups (1 and 6 months). The C and D (1-month old lenses) and G and H (6-months old lenses) corresponded to samples of A, B, E and F (Coomassie blue-stained gels), and show the Western blot results with anti-His antibody (green fluorescence). The levels of immunoreactive green fluorescence with His-tagged αA in lenses of 1-month old lenses (Left panel: WTαA protein [C] and αAN101D protein [D]) and 6-month old lenses (Right panel: WTαA protein[G] and αAN101D protein [G]) are shown. Additionally, in both left and right upper panels, the lanes 1, 2 and 3 show the WS-protein fractions recovered after first, second and third consecutive washes in buffer A to solubilize WS-proteins, respectively. Lanes 4 and 5 represent the urea soluble-protein fractions recovered during two consecutive washes of WI-protein pellet (containing membranes) with buffer B containing 8M urea, respectively. Lane 6 represents the 0.1N NaOH-solubilized proteins from membranes and the lane 7 from both 1- and 6-month old lenses (left and right panels) show the purified lens membrane preparations. Similarly, lanes 7 and 8 from 6-month old lenses (right panel) represent purified membrane preparation. Lane 9 of 6-month old lenses represents the crude lens WS-homogenate. The results show that the green fluorescence representing WTαA in CRYαA<sub>WT</sub> mice was entirely disappeared on urea solubilization in 1- and 6-month old lenses (lanes 1 to 5 in both left and right panels), whereas it was still present in these lenses until 0.1N NaOH wash (lane 6 in left and right panels). In contrast, the green fluorescence still existed in lane 6 of membranes from 1- and 6-month-old CRYαA<sub>N101D</sub> lenses. Together, the results suggest that αA<sub>N101D</sub> was tightly bound and at the higher levels to lens membrane of CRYαA<sub>N101D</sub> lenses relative to CRYαA<sub>WT</sub> lenses. On Image J-quantification of the Western blots (I and J), the lanes 4 and 5 (urea soluble fractions) of 1-month old lenses showed higher levels (2.5X) of immunoreactivity with anti-His antibody in the CRYαA<sub>N101D</sub> lenses (shown in red) compared to those from CRYαA<sub>WT</sub> lenses (blue). Similarly, in J, among the lanes 4 and 5 containing same fractions from 6-month old lenses (as described in 1-month old lenses), the lane 5 showed a greater immunoreactive level of CRYαA<sub>N101D</sub> lenses (red) compared to CRYAAWT lenses (blue). Additionally, the lane 6 (representing membrane remaining after two urea washes, right panel) of 6-month CRYαA<sub>N101D</sub> lenses exhibited about 2X greater immunoreactivity than CRYαA<sub>WT</sub> lenses (Quantification results not shown). Together, the results show that relative to CRYαA<sub>WT</sub>, higher levels of CRYαA<sub>N101D</sub> were tightly associated with the lens membranes of 1- and 6-month old CRYαA<sub>N101D</sub> mice.

(iii) Relative Membrane-Binding of Alexa 350-Labeled Recombinant WTαA- and αA<sub>N101D</sub> Crystallins

To examine whether αA<sub>N101D</sub> show a greater binding affinity to the lens membrane relative to WTαA-crystallin, the binding of the two recombinant proteins to purified lens membrane was determined. The recombinant WTαA- and αAN101D proteins were labeled with Alexa 350 using a protein labeling kit by the procedure described by the manufacturer (Molecular Probes, Thermo fisher Scientific). The two labeled-proteins were purified by a size-exclusion HPLC column and were analyzed by SDS-PAGE. Figure 6A shows the Coomassie blue-stained WT αA (lane 1), αA<sub>N101D</sub> protein (lane 2), and the purified lens membrane from non-transgenic C57 mice (lane 3). The Figure 6B shows the images of the two Alexa 350-labeled proteins under a UV
trans-illuminator (Lane 1: Images of Alexa 350-labeled WTαA, and lane 2: Alexa 350-labeled αAN101D). During the binding assay, the purified lens membrane (containing 2.5 mg protein isolated from 1 to 3-month old non-transgenic C57BL mice) was incubated with increasing but identical concentrations of either Alexa-labelled WT αA- or αAN101D proteins at 37°C for 6 h (See details in Methods). A relatively higher levels (> 1.5X) of binding of αAN101D protein relative to WTαA protein with membrane preparation was observed (Figure 6C). The values reported are the average of triplicate assays.

(iv) Immunogold-Labeling for Relative Localization of αA-WT and αAN101D in Lens Membranes of CRYαAN101D and CRYαAWT Mice

To ascertain the relative levels association αAN101D vs. WTαA to the lens membrane in vivo, the immunogold-labeling experiment was carried out (See details in Methods). (A) and (B) in Figure 7 show lens membranes from CRYαAN101D and CRYαAWT mice at 500 nm magnification, and (C) and (D) from these lenses at 100 nm magnification, respectively. The bigger gold particles (25 nm, red arrows) the smaller gold particles (10 nm, yellow arrows) represented the aquaporin-0 and the His-tagged αAN101D and WTαA, respectively. As shown in the representative images in (A) to (D), the 25 nm gold particles (representing aquaporin-0, identified by red arrows) were bound to membranes. On counting the membrane-associated 10 nm particles (representing His-tagged αAN101D and WTαA), almost the same numbers of the particle were found to be associated with membranes of both CRYαAN101D and CRYαAWT lenses, suggesting that the His-tagged αAN101D and WTαA were bound to the membranes of the two types of lenses. Our previous study [28] showed that αAN101D protein constituted about 14% and 14.2% of the total αA- crystallin in the WS- and WI-proteins, respectively in the lenses of CRYαAN101D mice. Therefore, an argument can be made that although an almost equal number of 10 nm and 25 nm particles were associated with membranes of the two type of lenses, a higher number of gold particle representing αAN101D relative to WTαA were associated with the membrane.

Another interesting observation was that the membranes of CRYαAN101D lenses were about 2X more swollen relative to those of CRYαAWT lenses [Figure 7, compare (A) to (B) and (C) to (D)]. The width of the membrane was quantified using Image J as shown in Figure 7E. The swelling could represent water intake within the lens cells due to the potential ionic imbalance in the CRYαAN101D lenses compared to CRYαAWT lenses. Such a possibility of ionic imbalance was further determined as described below.

Na, K-ATPase and Ca²⁺ Levels in Cultured Epithelial Cells from Lenses of CRYαAN101D and CRYαAWT Mice

Sodium-potassium-adenosine triphosphatase (Na, K-ATPase) has been recognized for its role in regulating electrolyte concentrations in the lens, and the electrolyte balance is vital to lens transparency [35, 36]. In addition, calcium has been reported to control both sodium and potassium permeability through lens membranes [37]. In our previous study [29], we showed that the expression of Na,K-ATPase at the protein level was drastically reduced in CRYαAN101D lenses relative to CRYαAWT lenses. Next, the
levels of Na, K-ATPase mRNA, and Ca\(^{2+}\) levels were determined in cultured epithelial cells from lenses of CRY\(\alpha\)A\(_{N101D}\) and CRY\(\alpha\)A\(_{WT}\) mice. Both (A) and (B) in Figure 8 show intracellular Ca\(^{2+}\) levels in the presence of calcium orange in cultured epithelial cells from CRY\(\alpha\)A\(_{N101D}\) and CRY\(\alpha\)A\(_{WT}\), respectively. Only a few CRY\(\alpha\)A\(_{N101D}\) epithelial cells showed the Ca\(^{2+}\) uptake, which was possibly due to our previous finding that the lens cells contained only about 14% of \(\alpha\)AN101D mutant protein [28]. In this experiment, 100 cells from the cultures of two types of lenses were counted. On quantification by Image J, the number of cells exhibiting calcium orange uptake were 1.5X greater in cells of CRY\(\alpha\)A\(_{N101D}\) lenses relative to cells from CRY\(\alpha\)A\(_{WT}\) lenses (Figure 8B). On the determination of levels of mRNA of Na, K-ATPase in these cells, its level was 75% lower in the CRY\(\alpha\)A\(_{N101D}\) lens cells than CRY\(\alpha\)A\(_{WT}\) lens cells (Figure 8C).

**Discussion**

Several past studies have shown *in vitro* effects of deamidation of crystallins on their structural properties including those in \(\alpha\)A-, and \(\alpha\)B-crystallins [21-25]. It has been reported that deamidation of Asn and Gln was the major modification identified in several human cataractous and aged lenses and these totaled 66% of the modification in the water-soluble and water-insoluble protein fractions that was analyzed by 2D LC/MS [38]. The mass spectrometric analysis found that there is negligible (less than 1%) deamidation at \(\alpha\)A\(_N101\) site in both aged and cataractous human lenses [39]. These studies suggested that because of low levels of deamidation of \(\alpha\)A- at N101 to D in normal and cataractous lenses, the \(\alpha\)AN101D might not play a significant role in cataract development. However, additional studies suggest otherwise. For example, our *in vitro* studies showed significant altered structural and functional properties of \(\alpha\)A-crystallin on deamidation of N101 residue but not of N123 residue [24]. We also showed that the WS-protein fraction from 50-70, year-old human donors contained \(\alpha\)A- fragments with deamidation of N101 to D [40]. This finding is significant because recent studies have also shown an increasing role of crystallin fragments in cataract development [41, 42]. In the present study, the cortical cataract development in mice on the introduction of \(\alpha\)A-N101D transgene further show significance of deamidation of this site and altered changes in the lens. However, the exact *in vivo* molecular mechanism of \(\alpha\)AN101D-induced crystallin’s aggregation is yet to be fully understood.

Previously we showed that the three recombinant deamidated \(\alpha\)A-mutants (N101D, N123D, and N101D/N123D) exhibited reduced levels of chaperone activity, alterations in secondary and tertiary structures, and larger aggregates relative to WT-\(\alpha\)A-crystallin [24, 25]. Among the above three mutants, the maximally affected and altered properties were observed in the recombinant \(\alpha\)AN101D mutant [25]. Additionally, our recent results show that *in vitro*, the deamidated \(\alpha\)A-, and \(\alpha\)B-crystallins facilitated greater interaction with \(\beta\)A3-crystallin, leading to the formation of larger aggregates, which might contribute to the lens cataractogenic mechanism [43]. As a further extension of our previous studies (28, 29), in some studies the 7-month old lenses were chosen because of the development of cortical cataract at about 7-month of age in the \(\alpha\)A\(_{N101D}\) mice relative to \(\alpha\)A\(_{wt}\) mice. In other experiments, lenses from 5-month old of both types of mice were used to determine the progression of phenotypic changes in lenses to determine their significance in cataractogenic mechanism.
The present study show that the introduction of αAN101D trans-gene in a mouse model resulted in the following major *in vivo* effects in lenses of CRYαAN101D- relative to CRYαAWT mice: (A) An age-related difference in protein profiles with an increasing association αAN101D with Wl-protein fraction suggesting its insolubilization after 4-months of age. (B) The WS-HMW protein fraction showed a higher level of proteins with a greater $M_r$. (C) Mass spectrometric analysis showed preferential insolubilization of αA-, αB-, γD- and γE-crystallins, and nestin, which remained insoluble even in 8M urea. (D) The tight association of αAN101D with membranes relative to WTαA, which could not be fully dissociated with 8M urea treatment. (E) In vitro, αAN101 showed greater affinity and binding to lens membranes relative WTαA. (F) The greater number of immunogold-labeled αAN101 relative WTαA binding to membrane along with relatively greater swelling of lens membranes, suggesting the potential water uptake due to intracellular ionic imbalance, and (G) The ionic imbalance was suggested by the greater Ca$^{2+}$ uptake and 75% reduction in mRNA levels of Na, K-ATPase in the epithelial cells cultured from CRYαAN101D lenses relative to those from CRYαAWT lenses. Our mass spectrometry analysis showed that retinal dehydrogenase was absent in the N101D mice. It has been shown earlier that *Aldh1a1* (–/–) knock-out mice developed lens opacification later in life (44). Retinal dehydrogenase 1 may protect the lens against cataract formation by detoxifying aldehyde products on lipid peroxidation in both cornea and lens. It has been shown that antimalarial drug chloroquine which binds and inhibit retinal dehydrogenase 1 (45) induce cataract in rats (46). Together, these findings suggest altered membrane integrity (possibly due to greater levels of αAN101D binding to membrane than WTαA) resulting in intracellular ionic imbalance in CRYαAN101D lenses, which could play a major role in the cortical cataract development.

Among the lens crystallins, only α-crystallin show an association with the membrane in both normal and cataractous lenses [6, 47-51]. Lens membranes contain both a high-affinity saturable and low-affinity non-saturable α-crystallin-binding sites [47, 51-53]. Alpha-crystallin binding to native membranes was enhanced on stripping of extrinsic proteins from the lens membrane surface to expose lipid moieties [32, 33], which contradicted a previous report that the crystallin mostly interacts with lens membrane MP26 protein [54]. Even after stripping extrinsic membrane proteins by alkali-urea treatment, the full-length αA-, and αB-crystallins remained associated with membranes of both bovine and human lenses [6]. Additionally, αB-crystallin showed three-fold higher binding to lens membrane relative to αA-crystallin, and their binding was affected by the residual membrane-associated proteins, suggesting that their binding behaviors were affected by an intrinsic lens peptides [6]. A large-scale association of proteins with cell membranes in the lens nucleus (mostly in the barrier region) occurs after middle age in human lenses [49], and such association was enhanced by mild thermal stress [50]. The *in vitro* studies further supported this because the binding capacity of α-crystallin from older lenses to lipids increased with age and decreased in diabetic donors who were treated with insulin [51]. This implied that under diabetic conditions, abnormal binding of α-crystallin to lens membrane occurred. Such information in the literature about membrane binding of native vs. post-translationally modified crystallins including the deamidated αAN101D species is presently lacking. Therefore, the results of the present study showing relatively increased binding of αAN101D relative to WTαA are highly significant.
The RNA sequence and IPA data of our previous study [29] further support the findings of the present study. This study [29] showed that the genes belonging to gene expression, cellular assembly, and organization, and cell cycle and apoptosis networks were altered, and specifically, the tight junction-signaling and Rho A signaling were among the top three canonical pathways that were affected in the CRYαAN101D lenses relative to CRYαAWT lenses. The present study showed an increased association of αAN101D to membrane, and this could lead to potential ionic imbalance affecting tight junction assembly and RhoA GTPase expression. This in turn causes increased proliferation and decreased of differentiation and denucleation of epithelial cells, and an accumulation of nuclei and nuclear debris in the lens anterior inner cortex and fiber cell degeneration. Some of these phenotypic changes could be cause or effects, but together could be responsible for the age-related cortical cataract development in CRYαAN101D lenses.

To maintain ionic balance within lens cells, a permeability barrier close to the surface of the lens is responsible for the continuous sodium extrusion via Na, K-ATPase-mediated active transport [35-37]. Without an active sodium extrusion, lens sodium and calcium contents are shown to increase resulting in lens swelling that leads to loss of lens transparency [35]. Similarly, an excessive intracellular Ca\(^{2+}\) levels can be detrimental to lens cells, and its increased levels play an important role in development of cortical cataract [37]. Therefore, homeostasis of Na\(^{+}\), K\(^{+}\), Ca\(^{2+}\) and other ions within the lens has been recognized as of fundamental importance in lens pathophysiology. These have been altered as shown in our present and our previous studies [29]. It is also possible that the increased Ca\(^{2+}\) levels could in turn lead to calpain activation and proteolysis of crystallins, which will be investigated in future.

Similar to our study, other studies have shown that an increased membrane binding of α- crystallin in the pathogenesis of different forms of cataracts. The high molecular weight complexes (HMWCs), comprised of α-crystallin and other crystallins, accumulate with aging and show a greater membrane binding capacity than native α-crystallin [51]. Other mutants of αA-crystallin, like the αAN101D mutant, also exhibit a greater membrane binding than corresponding wild-type species [55]. For example, in the αAR116C-associated congenital cataracts, an increased membrane binding capacity along with changes in complex polydispersity, and the reduction of subunit exchange were considered potential factors in the cataract pathogenesis [55]. Similarly, αA-crystallin R49C neo mutation influenced the architecture of lens fiber cell membranes and caused posterior and nuclear cataracts in mice [56].

Interactions between proteins and the cell membrane are an integral aspect of many biological processes, which are influenced by compositions of both membrane lipids and protein structure [57]. Reports have shown the age-related lipid compositional changes in the lens membrane, which might affect α-crystallin binding, i.e., in the nucleus of the human lenses, the levels of glycerophospholipids declined steadily by age 40 as opposed to the levels of ceramides and dihydroceramides increased approximately 100 fold during middle-age [58, 59]. Further, it has been shown that because of the elevation of sphingolipid levels with species, age, and cataract, lipid hydrocarbon chain order, or stiffness increased. Therefore, the increased membrane stiffness caused an increase in light-scattering, reduced calcium pump activity,
altered protein-lipid interactions, and perhaps slow fiber cell elongation [60]. Presently, whether similar changes occur in α\textsubscript{AN101D} lenses are not known.

Alpha A- and αB-crystallins differently associate with the cellular membrane, i.e. αA-crystallin may interact exclusively with membrane phospholipids, and thereby unaffected by the presence of extrinsic proteins on the membrane, whereas these proteins may act as conduits for αB-crystallin to bind to the membrane [59]. Presently, the specific binding mechanism of αAN101D to the membrane and age-related changes in lipid composition in lenses CRYα\textsubscript{AN101D} vs. CRYα\textsubscript{WT} are unknown, and these are presently the focus of our investigations.

Conclusions

The results presented in this study suggest that an increased association of αAN101D relative WTαA with the lens membrane causes a possible loss of membrane integrity, leading to an ionic imbalance, and in turn, to membrane swelling, cellular disorganization and finally cortical opacity. Our future study will determine the specific binding site in the αAN101D relative WTαA, and changes in the membrane compositions that might facilitate the increased binding of the deamidated crystallin with the membrane.

Declarations

Ethics approval and consent to participate

No human subjects were involved in the study. All animal experiments were performed per protocols approved by the Institutional Animal Care and Use Committee (IACUC) of the University of Alabama at Birmingham (Protocol no. 130208393). Mice were housed in a pathogen-free environment at the facility of the University of Alabama at Birmingham.

Consent for publication

Not applicable

Availability of data and materials

All data generated or analyzed during this study are included in this unpublished article [and its supplementary information files].

Competing interests

No competing interest

Funding

This study was funded by NIH grants, EY- 06400 (OS) and P30EY003039.
Authors’ contributions

KS and RJ conducted experiments, analyzed and interpreted the data and wrote the manuscript. LW conducted the mass spectrometric analysis and analyzed and interpreted the results. OS with the help of RJ analyzed and interpreted the data and have written and edited the manuscript.

Source of Animals

The CRYαA\textsubscript{N101D} mouse model was generated by inserting the human lens αA-N101D transgene and the CRYαA\textsubscript{WT} mouse was generated by inserting human wild-type αA-transgene. Both mouse models were generated in Dr. Om Srivastava’s laboratory. The details of the methodology are described in reference no. 28 (Gupta R, Asomugha CO, Srivastava OP. The common modification in αA-crystallin in the lens, N\textsubscript{101D} is associated with increased opacity in a mouse model. J Biol Chem. 2011; 286:11579-592)

Acknowledgements

Authors thank Ms. Rebecca Vance for the help with handling the CRYαA\textsubscript{N101D} and CRYαA\textsubscript{WT} mice. Thanks also goes to the High Resolution Imaging Facility, Targeted Metabolomics & Proteomics Laboratory and the Ocular Phenotyping and Molecular Analysis Core Facility of Vision Science Research Center at the University of Alabama at Birmingham.

ARRIVE Guidelines

Minimum Standard of Reporting Checklist

Experimental design and statistics

The method section contains the information as described in the guideline.

Resources

The description of all the resources used are included in the Method section.

Availability of data and materials

All the data are available in the corresponding author’s notebooks and his computer.

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Supplemental File Legends

Supplemental Tables A to D:

Table A: Water Insoluble-Urea Soluble (WI-US)-Protein Fraction of Alpha A-WT lenses.

Table B: Water Insoluble-Urea Insoluble (WI-UI)-Protein Fraction of AlphaA N101D lens

Table C: Water insoluble-urea soluble alphaAN101D (Mr >30 kDa)

Table D: Water insoluble-urea insoluble alpha A N101D (Mr >30 kDa)

Supplementary Methods

Miscellaneous Methods

Mass Spectrometric Analysis

The mass spectrometric analysis was carried out at the Targeted Metabolomics and Proteomics Laboratory of the University of Alabama at Birmingham.

(A) In-Gel Digestion, NanocHiPLC-Tandem Mass Spectrometry, and Protein Pilot 4.5 Search Queries
Following SDS-PAGE analysis, the desired gel bands were excised, and an overnight wash removed the excess stain with 50% of 100 mM ammonium bicarbonate/50% acetonitrile. Next, the disulfide bonds were reduced by 25 mM dithiothreitol at 50°C for 30 min, and the alkylation of the free thiols groups was carried out with 55 mM iodoacetamide for 30 min in the dark. The excess alkylation agent was removed and the gel pieces were washed twice with a 100 mM ammonium bicarbonate for 30 min, and was evaporated to dryness in a SpeedVac (Savant, ThermoFisher Scientific, Atlanta, GA) before the addition of 12.5 ng/µl of trypsin (Promega Gold Mass Spectrometry Grade, Madison, WI), and incubated overnight at 37°C. Peptides were extracted twice for 15 min from the gel pieces using a 1:1 mixture of 1% formic acid and acetonitrile, then the extracts were and evaporated to dryness, and the samples were resuspended in 30 µl of a 0.1% formic acid before mass spectrometric analysis.

An aliquot (5 µL) of each digest was loaded onto a Nano cHiPLC 200 µm x 0.5 mm ChromXP C18-CL 3 µm 120Å reverse-phase trap cartridge (Eksigent, Dublin, CA) at 2 µL/min using an Eksigent autosampler (Eksigent, Dublin, CA). After washing the cartridge for 4 min with 0.1% formic acid in ddH2O, the bound peptides were flushed onto a Nano cHiPLC column [200 µm x 15 cm. ChromXP C18-CL 3 µm 120Å] with a 45 min linear (5-50%) acetonitrile gradient in 0.1% formic acid at 1000 µl/min using an Eksigent Nano1D+LC (Eksigent, Dublin, CA). The column was washed with 90% acetonitrile-0.1% formic acid for 10 min and then re-equilibrated with 5% acetonitrile-0.1% formic acid for 10 min. The SCIEX 5600 Triple-TOF mass spectrometer (AB-Sciex, Toronto, Canada) was used to analyze the protein digest. The IonSpray voltage was 2300 V, and the declustering potential was 80 V. Ion spray, and curtain gases were set at 10 psi and 25 psi, respectively, and the interface heater temperature was 120°C. Eluted peptides were subjected to a time-of-flight survey scan from 400-1250 m/z to determine the top twenty most intense ions for MS/MS analysis. Product ion time-of-flight scans at 50 msec were carried out to obtain the tandem mass spectra of the selected parent ions over the range from m/z 100-1500. Spectra are centroided and de-isotoped by Analyst software, version TF (Applied Biosystems). A β-galactosidase-trypsin digest was used to confirm the mass accuracy of the mass spectrometer. The tandem mass spectrometry data were processed to provide protein identifications using an in-house Protein Pilot 4.5 search engine (SCIEX) using the Mus musculus (Mouse) UniProt protein database and using a trypsin digestion parameter. All proteins that had at least one peptide with a confidence score of 95% or higher were considered as potential candidates.

(B) In-Solution Digestion Protocol for Mass Spectrometric Analysis

All of the reagents used were freshly prepared before their use. A 100-µL aliquot of the protein sample (1 mg) in 100 mM Tris buffer (pH 7.8) containing the 6M urea was mixed with the reducing reagent (DTT) and incubated for 1 h at room temperature. Next, 20 µL of the alkylating reagent (iodoacetamide) was added and incubated for 1 h at room temperature, and then 20 µL of the reducing agent was added to consume any unreacted iodoacetamide and allowed to stand at room temperature for 1 h. The urea concentration was reduced to ~0.6 M by diluting the reaction mixture with 775 µL of water, 100-µL trypsin solution (20 µg of stock trypsin) was added to bring protease-to-substrate ration 1-to-50, and the digestion was carried out overnight at 37°C. The trypsin digestion was stopped by adjusting the pH to
<6.0 by adding concentrated acetic acid. The digest was analyzed directly or concentrated by evaporation. As needed, the samples were desalted using a C18 ZipTip™ (Millipore Corporation, Bedford, MA) using manufacturer's instructions. The mass spectrometric analysis was carried out as described above.