Epigenetic Regulation of αA-crystallin in High Myopia-Induced Dark Nuclear Cataract

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Abstract

**Purpose:** To assess the etiology of early-onset dark nucleus in high-myopic patients and its relationship with the epigenetic regulation of αA-crystallin (CRYAA).

**Methods:** We reviewed clinical data from patients who underwent cataract surgery at our center in 2012. Lens epithelial samples were collected during capsulorhexis, whereas young lens epithelium was donated. Cataract type and severity were graded according to the Lens Opacity Classification System III (LOCS III). DNA methylation was analyzed by pyrosequencing the CpG islands of the CRYAA promoter in the following groups: Age-Related Cataract (ARC) Nuclear Color (NC) 2–3; High-Myopic Cataract (HMC) NC2–3; ARC NC5–6; HMC NC5–6; and in young lenses graded NC1. We analyzed CRYAA expression by real-time polymerase chain reaction (PCR), reverse transcription PCR, and immunohistochemistry.

**Results:** The odds ratio of dark nucleus in high-myopic patients was 5.16 (95% confidence interval: 3.98–6.69; p<0.001). CpG islands in lens epithelial CRYAA promoter in the HMC NC5–6 Group exhibited the highest methylation of all the groups, but no statistically significant differences were evident between the HMC NC2–3 and ARC NC2–3 Groups. Likewise, CRYAA mRNA and protein levels in the HMC NC5–6 Group were significantly lower than the ARC NC5–6 Group and high-myopic controls.

**Conclusions:** High myopia is a risk factor for dark nucleus. Downregulation of CRYAA via the hypermethylation of CpG islands in its promoter could underlie the earlier onset of dark nucleus in high-myopic patients.

Introduction

High myopia, defined as myopia exceeding –6.00 diopters or axis length ≥ 26 mm, is a disorder that affects almost the entire human eye, from the anterior pole to the posterior pole. High myopia can be associated with high refractive errors, cataract, open-angled glaucoma, and retinopathy, all of which are recognized as significant problems of subspecialties related to ophthalmology [1], and more difficulties are encountered during surgery in such cases.

As the incidence of high myopia is much higher in Asia than elsewhere [2], it is likely that high-myopic cataract (HMC) is more common in Asia [3]. Several population-based studies [3–6] have examined the association between high myopia and cataract. For example, data from the Blue Mountains Eye Study [4] suggested that high myopia is associated with both nuclear and posterior subcapsular cataract. In another series of studies [2,7], it was reported that nuclear cataract is strongly associated with high-axial myopia. Praveen et al. [7] also reported that cataract density is higher in patients with high myopia than in other groups of patients. However, high myopia is not associated with posterior subcapsular or cortical cataract.

We have noticed that some patients with high myopia develop dark nuclear cataract at an earlier age than high-myopic patients without a dark nucleus or patients with age-related cataract (ARC). Patients with high-myopic dark nuclear cataract show rapid cataract progression at a relatively young age, they usually have a brown or dark nucleus, and may have a longer axis. As the largest cataract surgery center in Eastern China, we perform cataract surgery on more than 2,000 patients with high myopia each year, and those with high-myopic dark nuclear cataract are referred to our center much earlier than other patients because of rapid deterioration in their vision. However, cataract surgery is very hazardous in such patients because of the high risk of posterior capsular tear, vitreous prolapse [8], persistent postoperative corneal edema, and retinal detachment. Therefore, it is particularly important to determine the etiology of this disorder.

Alpha-crystallin, composed of two subunits, αA- and αB-crystallin, is the most abundant structural protein in the human lens [9,10]. The biologic role of α-crystallin is to bind to lens proteins that denature over time, thereby maintaining long-term lens transparency. This is particularly important in the lens, as there is no protein turnover in the inner part [11–13] and, once
hypothesized that hypermethylation of CpG islands in the high myopia show earlier-onset and higher-density cataract. We silencing [16].

islands in a promoter causes heterochromatin formation and gene electrostatic nature of chromatin. Hypermethylation of CpG methylation and histone modification. In DNA methylation, DNA–transcription factor affinity through two mechanisms: DNA methylation and histone modification. In DNA methylation, addition of a methyl group to cytosine changes the electrostatic nature of chromatin. Hypermethylation of CpG islands in a promoter causes heterochromatin formation and gene silencing [16].

Very few studies have addressed the reasons why patients with high myopia show earlier-onset and higher-density cataract. We hypothesized that hypermethylation of CpG islands in the CRELA promoter underlies the etiology of HMC as well. To test our hypothesis, we examined the differences in characteristics among high-myopic eyes with dark nuclear cataract, high-myopic cataract.

Methods

The Ethics Committee of the Eye and ENT Hospital, Fudan University, Shanghai, China, approved our collection and use of anterior capsule membranes from patients undergoing cataract surgeries. All samples were collected after obtaining written informed consent from patients, and the consent procedure was also approved by the ethics committee. All procedures adhered to the Declaration of Helsinki for research involving human subjects.

Clinical data review

We identified patients with/without dark nuclear cataract (NC5–6/NC2–4) among those with ARC and HMC using our follow-up tables for 2012. We retrieved clinical data for these patients and calculated the prevalence of cataract of LOCS III Grade NC2–3 in different age groups among age-related cataract and high-myopic cataract. Statistically significant differences were evident in the 40–55, 56–65, and 66–75 years age groups (Pearson’s chi-squared test; “*” p<0.05).

doi:10.1371/journal.pone.0081900.g001

Prevalence of Grade Nuclear Color 5–6 cataracts in different age groups among age-related cataract and high-myopic cataract. Statistical significant differences were evident in the 40–55, 56–65, and 66–75 years age groups (Pearson’s chi-squared test; “*” p<0.05).

doi:10.1371/journal.pone.0081900.t001

| Category | NC5–6 | NC2–4 |
|----------|-------|-------|
| ARC      | 85    | 4833  |
| HMC      | 198   | 2181  |

Table 1. Number of patients with a dark nucleus (Nuclear Color 5–6) in each group.

*age-related cataract.

**high-myopic cataract.

**nuclear color.

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Ophthalmologists independently and simultaneously graded each cataract.

Consecutive patients with nuclear cataract and a length (≥26 mm) or normal (21–23.99 mm) axis admitted to our clinic between September 2012 and January 2013 were considered for this study. Patients with glaucoma, uveitis, previous trauma, or other ocular diseases, and those with diabetes or malnutrition, were excluded from this study. Patients were then subdivided into four groups based on LOCS III grade: HMC NC2–3 (n = 7); ARC NC2–3 (n = 7); and ARC NC5–6 (n = 8). The average LOCS III grades for these groups were 2.57, 5.45, 2.57, and 5.50, respectively.

Lens anterior capsule membrane samples were obtained by continuous curvilinear capsulorhexis during cataract surgery; only samples without blood or iris tissue contamination or damage to other intraocular structures were used. We also collected lens anterior capsule membrane samples from nine individuals aged 30–40 years classified as NC1 without cataract. These samples were obtained from eyes of normal axial length donated to the Eye Bank of the Eye and ENT Hospital, and were used as normal controls. The interval between death and the collection of cadaveric eyes was < 4 hours. Anterior capsule membranes were carefully collected from the center of lenses. We then compared groups defined by axial length and LOCS III scores.

Pyrosequencing

Pyrosequencing was performed as previously described [14,18]. The polymerase chain reaction (PCR) was performed with 0.2 M of each primer for biotin (forward: TGGGGA-TATGTAGTTATTTTGTAGGAG; reverse: CTAACCCC-CAACCCCATACGAT) following bisulfite conversion. PCR products were bound to Streptavidin Sepharose® High Performance (Amersham Biosciences, Uppsala, Sweden). Sepharose® beads containing the immobilized PCR products were purified, washed, and denatured with 0.2 M NaOH, then reanimated with a Pyrosequencing Vacuum Prep Tool (Qiagen, Venlo, Limburg, The Netherlands). Next, 0.5 µM of the pyrosequencing primer (CAACCCCATACGATC) was annealed to the purified single-stranded PCR products, and the resulting products (10 µl) were sequenced using the Pyrosequencing PSQ 96 HS System.
For each locus, methylation status was analyzed individually as a T/C single nucleotide polymorphism using Pyro-Q CpG software (Qiagen).

Reverse transcription polymerase chain reaction and relative quantitative real-time polymerase chain reaction

As previously described [14,19], CRYAA mRNA levels were assayed as follows. For reverse transcription polymerase chain reaction (RT-PCR), total RNA was extracted with TRIzol®

![Figure 2](https://example.com/figure2.png)

**Figure 2.** CpG island methylation in the αA-crystallin promoter was similar in age-related cataracts and high-myopic cataracts classified as Lens Opacity Classification System III Grade Nuclear Color 2–3. (A) Representative Pyrosequencing results of the Control, Age-Related Cataract (ARC) Nuclear Color (NC) 2–3 and High-Myopic Cataract (HMC) NC2–3 Groups. (B) Anterior segment photos of patients in each group. (C) The six selected CpG islands in the ARC NC2–3 (44.8 ± 8.3%) and HMC NC2–3 (43.7 ± 3.8%) Groups displayed higher methylation compared with the Control Group (34.6 ± 6.1%). (D) Percentage of methylation at each individual CpG site in each group. (*Statistically significant differences were found between the Control and ARC NC2–3 Groups; **Statistically significant differences were found between the Control and HMC NC2–3 Groups; ***All p < 0.05. However, no statistically significant differences were found between the two cataract groups.)

doi:10.1371/journal.pone.0081900.g002
Reagent (Life Technologies Corporation, Carlsbad, CA, USA). Quantification and purity of total RNA was measured by A260/A280 absorption. A total of 1 µg RNA was reverse-transcribed using the First Strand cDNA Synthesis Kit. The primers used in RT-PCR were as follows: CRYAA, forward: 5′-CCTGCTGCGCTCGTTGTGCT-3; CRYAA, reverse: 5′-ACTCCGTGTGGATCGG-3. Representative immunohistochemistry images of the Control, Age-Related Cataract Nuclear Color (NC) 2–3, and High-Myopic Cataract NC2–3 Groups. (F) Mean staining density in each group (*all p < 0.05). doi:10.1371/journal.pone.0081900.g003

Figure 3. Expression of αA-crystallin was similar in age-related cataracts and high-myopic cataracts classified as Lens Opacity Classification System III Grade Nuclear Color 2–3. Alpha-crystallin expression in the lens epithelium of each group was detected by (A) real-time polymerase chain reaction (PCR) also showing levels of expression in the lens epithelium of each group was detected by (A) real-time polymerase chain reaction (PCR; also showing levels of αA-crystallin relative to β-actin) and (B) reverse transcription PCR (C, D, E) Representative immunohistochemistry images of the Control, Age-Related Cataract Nuclear Color (NC) 2–3, and High-Myopic Cataract NC2–3 Groups. (F) Mean staining density in each group (*all p < 0.05).

Statistical analysis

Data are expressed as means ± standard deviation. Logistic regression was used to analyze the association between high myopia and dark nuclear cataract. Continuous parameters were compared by one-way analysis of variance followed by the least significant difference test. P-values < 0.05 were considered statistically significant. All analyses were performed using SPSS software (version 11.0; SPSS Inc., Chicago, IL, USA).

Results

High myopia is a risk factor for dark nuclear cataract

Table 1 shows the number of HMC and ARC patients who presented with dark nuclear cataract (NC5–6) and underwent cataract surgery at our center in 2012. High myopia was associated with a significantly greater risk of dark nuclear cataract (odds ratio [OR]: 5.16; 95% confidence interval [CI]: 3.98–6.69; p < 0.001). The mean patient ages in the ARC NC2–4, HMC NC2–4, ARC NC5–6, and HMC NC5–6 Groups were 69.9 ± 16.4, 68.7 ± 13.8, 71.4 ± 11.6, and 59.6 ± 8.3 years, respectively. Of all four groups, the mean age at cataract onset was significantly lower in the HMC NC5–6 Group (all p < 0.05). Figure 1 shows the prevalence of NC5–6 cataract in different age groups of ARC and HMC patients. Statistically significant differences were evident for the 40–55, 56–65, and 66–75 years age groups (all p < 0.05). The male: female ratio was 1124:1057 in the HMC NC2–4 Group and 108:90 in the HMC NC5–6 Group. Gender was not associated with the prevalence of NC5–6 cataract in HMC patients (p > 0.05).

General characteristics of the samples

The general characteristics of patients in each group are summarized in Table 2. The mean age of patients in the HMC NC5–6 Group was lower than that of patients in the ARC NC2–3, HMC NC2–3, and ARC NC5–6 Groups. There were also more men in the HMC NC5–6 Group than in the other four groups (all p > 0.05).

CpG island methylation in the αA-crystallin promoter was similar in age-related cataract and high-myopic cataract classified as Lens Opacity Classification System III Grade Nuclear Color 2–3

Pyrosequencing, a real-time DNA sequencing method, was used to determine single-base variations caused by CpG methylation of the CRYAA promoter in each group. Figure 2 shows the
methylation status of the \( \alpha \)-crystallin promoter in the Control, ARC NC2–3, and HMC NC2–3 Groups. Methylation of the six selected CpG islands was significantly greater in the ARC NC2–3 (44.8\%\( \pm \)3.3%) and HMC NC2–3 (43.7\%\( \pm \)3.8%) Groups than in the Control Group (34.6\%\( \pm \)6.1%; both \( p < 0.05 \)). However, methylation of the \( \alpha \)-crystallin promoter was not significantly different between the ARC NC2–3 and HMC NC2–3 Groups.

Expression of \( \alpha \)-crystallin was similar in age-related cataract and high-myopic cataract classified as Lens Opacity Classification System III Grade Nuclear Color 2–3

To determine the expression of \( \alpha \)-crystallin, we conducted real-time PCR, RT-PCR, and IHC using lens anterior capsule tissue samples from each group. Real-time PCR and RT-PCR (Figure 3A and B) showed that \( \alpha \)-crystallin mRNA expression in the ARC NC2–3 and HMC NC2–3 Groups (64.8\%\( \pm \)5.9% and 52.8\%\( \pm \)6.1%, respectively; both \( p < 0.05 \)). However, methylation of the \( \alpha \)-crystallin promoter was not significantly different between the ARC NC2–3 and HMC NC2–3 Groups.

CpG island methylation in the \( \alpha \)-crystallin promoter was significantly greater in high-myopic cataract than in age-related cataract classified as Lens Opacity Classification System III Grade Nuclear Color 5–6

Figure 4 shows that, among patients classified as LOCS III Grade NC5–6, CpG island methylation in the \( \alpha \)-crystallin promoter was significantly greater in the HMC Group (57.6\%\( \pm \)2.6%) than in the ARC Group (48.1\%\( \pm \)3.4%); \( p = 0.01 \), even though mean patient age in the HMC group was lower (Tables 1: \( p < 0.05 \) and Table 2: \( p > 0.05 \)). Methylation was also significantly greater in the HMC NC5–6 Group than in the axial length-matched cataract Control Group (i.e., HMC NC2–3; \( p < 0.001 \)). In fact, it was higher in the HMC NC5–6 Group than in all other groups (all \( p < 0.05 \)). Therefore, in patients with high myopia, dark nuclear cataract develops at an earlier age and the lens epithelium exhibits greater methylation of the \( \alpha \)-crystallin promoter compared with axial length-matched or age-matched cataract controls.

AlphaA-crystallin expression in the lens epithelium was significantly lower in the HMC NC5–6 Group than in the ARC NC5–6 Group

We next performed real-time PCR, RT-PCR, and IHC using lens epithelial cell samples from the ARC NC5–6 and HMC NC5–6 Groups. Real-time PCR and RT-PCR (Figure 5A and B)
expression was lowest in the HMC NC5–6 Group (all p < 0.05). Likewise, of all five experimental groups, epithelium was significantly lower in the HMC NC5–6 Group showed that (79.8
epithelium was significantly lower in the High-Myopic Cataract Nuclear Color (NC) 5–6 Group (both p < 0.05).

Discussion

High myopia is a common eye disease in Asian populations. It is closely associated with many pathologic changes of the eye, including marked fundus changes, open-angled glaucoma [23,24], and the early onset of cataract [25,26]. However, in the last 50 years, few studies have tried to elucidate the etiology of HMC, especially of dark nuclear cataract. In this study, we found that high myopia is a risk factor for dark nuclear cataract, and that Crysralin expression was also significantly downregulated in patients with high-myopic dark nuclear cataract than in patients with ARC of the same LOCS III grade. Overall, our findings suggest that downregulation of CRYAA associated with DNA hypermethylation is involved in the development of high-myopic dark nuclear cataract.

Increasing clinical evidence suggests that high myopia is a risk factor for dark nuclear cataract. For example, epidemiologic studies conducted in India [2,7] revealed that mean cataract density was higher in patients with high myopia. However, they did not report whether high myopia was associated with dark nuclear cataract. At our clinic, we frequently encounter patients with high myopia, who often present with brown or dark nuclear cataract at a relatively young age. Thus, we believed that the incidence of dark nuclear cataract to be higher in patients with high myopia than in patients with ARC. Therefore, we reviewed the clinical data of patients who underwent cataract surgery at our clinic in 2012, and found that high myopia was associated with a significantly greater risk of dark nuclear cataract (OR: 5.16; 95% CI: 3.98–6.69; p < 0.001).

Increased oxygen tension caused by earlier vitreous liquefactions in patients with high myopia relative to patients with ARC could be an important cause of the earlier onset of dark nuclear cataract in HMC patients, according to studies of cataract formation after vitrectomy [27,28]. Of note, increased oxygen tension around the lens exposes lens proteins to marked oxidative stress in high-myopic patients [28,29]. Additionally, studies comparing the oxidative consumption of glutathione in the lenses of HMC and ARC patients [30,31] showed that, compared with healthy controls, the lenses of patients with cataract had lower glutathione levels, with the lowest levels found in the lenses of high-myopic patients. Also, elevated malondialdehyde levels were found in the cataractous lenses and vitreous of myopic patients compared with nonmyopic patients with cataract [30]. Therefore, the lens needs to maintain a healthy defense system to protect against oxidative stress, particularly in patients with high myopia.

Alpha-crystallin, the best-characterized structural protein of the human lens, reportedly acts as a chaperone under conditions of oxidative stress, thereby maintaining the transparency of the lens [10,32]. However, in this study, we observed a significant decrease in αA-crystallin expression in patients with high-myopic dark nuclear cataract compared with patients with ARC of similar severity. This reduction in αA-crystallin expression may increase the susceptibility of lens proteins to oxidative stress associated with aging.

Epigenetic regulation of gene expression was recently shown to be associated with many eye diseases, including age-related macular degeneration [33], glaucoma [34], and retinoblastoma [35]. Additionally, we reported [14] that hypermethylation of CpG islands in the CRYAA promoter downregulates αA-crystallin expression in ARC. The role of hypermethylation in the downregulation of CRYAA expression was confirmed by treating lens epithelial cells with a demethylating agent, which restored CRYAA expression [14]. Therefore, we wondered whether the epigenetic regulation of CRYAA expression also functioned in the pathogenesis of HMC. Our analysis showed that hypermethylation of CpG islands in the CRYAA promoter was much greater in patients with high-myopic dark nuclear cataract than in patients with ARC of a similar LOCS III grade. Consistent with this, we found that much less αA-crystallin was synthesized in the lenses of patients with high-myopic dark nuclear cataract.

It is established that fewer reducing agents, such as glutathione [36] and ascorbic acid, are present in the nuclear region of the lens than in the cortical region, leading to a localized reduction in antioxidative capacity. Consequently, a greater proportion of proteins are denatured as a result of oxidative stress in this region. In the lenses of the young, a sufficient number of chaperones (the majority αA-crystallin) are expressed to bind these denatured...
proteins, preserving the transparency of the lens. However, in high-myopic patients, much less αA-crystallin is synthesized to pass into the nucleus and restore these denatured proteins. This may accelerate the oxidative modification of proteins in the nucleus, resulting in the development of high-myopic dark nuclear cataract. Figure 6 summarizes the pathogenesis of high-myopia-induced dark nuclear cataract.

While collecting lens anterior capsule membranes over a period of 4 months, it became apparent that patients with high-myopic dark nuclear cataract were younger than other groups of patients. These patients visited our clinic at an earlier age because of a rapid deterioration in their visual acuity. The significantly higher prevalence of Grade NC5–6 cataract in the younger age groups of HMC patients further indicated that dark nuclear cataract occurs at an earlier age in patients with HMC than in patients with ARC. In contrast, patients with HMC without a dark nucleus were admitted for cataract surgery at up to 80 years of age, often with cataract classified as LOCS III Grade NC2. Our clinical experience is also supported by our current data, which show that methylation of CpG islands in the CRYAA promoter is significantly lower in high-myopic patients with soft nuclear cataract (NC2–3) compared with patients with dark nuclear cataract, but similar to that in ARC classified as NC2–3. Therefore, we believe that two types of HMC may exist. The first is characterized by the early onset and rapid progression of nuclear cataract, and typically presents with a dark nucleus. The second is characterized by gradual progression, and may retain a soft nucleus classified as NC2–3, even in older patients. Further studies are needed to elucidate the causes of this phenomenon. On the other hand, although the proportion of men was higher in the samples we collected, gender was not associated with the prevalence of dark nucleus in this study. Larger, population-based studies are still warranted in future.

In conclusion, our data indicate that high myopia is a risk factor for dark nuclear cataract, and that hypermethylation of CpG islands in the CRYAA promoter leading to downregulation of αA-crystallin, a major component of antioxidative mechanisms within the lens, functions in the pathogenesis of high-myopic dark nuclear cataract. These results provide a new direction for research into HMC.

Acknowledgments

The authors thank Fan Fan and Xin Liu for collecting the lens anterior capsular membranes.

Author Contributions

Conceived and designed the experiments: XJZ Y. Lu. Performed the experiments: XJZ PZ KKZ. Analyzed the data: XJZ PZ. Contributed reagents/materials/analysis tools: JY Y. Luo. Wrote the paper: XJZ.

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