Nuclear Cataract Shows Significant Familial Aggregation in an Older Population after Adjustment for Possible Shared Environmental Factors

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PURPOSE. To quantify the association between siblings in age-related nuclear cataract, after adjusting for known environmental and personal risk factors.

METHODS. All participants (proband) in the Salisbury Eye Evaluation (SEE) project and their locally resident siblings underwent digital slit lamp photography and were administered a questionnaire to assess risk factors for cataract including: age, gender, lifetime sun exposure, smoking and diabetes history, and use of alcohol and medications such as estrogens and steroids. In addition, blood pressure, body mass index, and serum antioxidants were measured in all participants. Lens photographs were graded by trained observers masked to the subjects’ identity, using the Wilmer Cataract Grading System. The odds ratio for siblings for affectedness with nuclear cataract and the sibling correlation of nuclear cataract grade, after adjusting for covariables, were estimated with generalized estimating equations.

RESULTS. Among 307 probands (mean age, 77.6 ± 4.5 years) and 434 full siblings (mean age, 72.4 ± 7.4 years), the average sibship size was 2.7 per family. After adjustment for covariables, the probability of development of nuclear cataract was significantly increased (odds ratio [OR] = 2.07, 95% confidence interval [CI]: 1.30–3.00) among individuals with a sibling with nuclear cataract (nuclear grade ≥ 3.0). The final fitted model indicated a magnitude of heritability for nuclear cataract of 35.6% (95% CI: 21.0%–50.3%) after adjustment for the covariables.

CONCLUSIONS. Findings in this study are consistent with a genetic effect for age-related nuclear cataract, a common and clinically significant form of lens opacity. (Invest Ophthalmol Vis Sci. 2004;45:2182–2186) DOI:10.1167/iovs.03-1163

A practical difficulty in understanding the genetics of nuclear cataract, as with all age-related diseases, is that unaffected status at the time of examination may mean that the individual is truly unaffected or that he or she has simply not manifested the phenotype yet. Misattribution can thus be a serious problem in all but the oldest populations. An additional difficulty in performing studies to ascertain the degree of heritability of age-related cataract in the United States is that general access to cataract surgical services leads to systematic censorship of the most affected persons in the population, with attendant loss of power and possible introduction of bias.

In the current investigation, we studied the heritability of nuclear cataract in a cohort of older sibships recruited through the Salisbury Eye Examination (SEE) on Maryland’s Eastern Shore. We attempted to overcome some of the mentioned difficulties in studying the genetics of age-related cataract. To distinguish between familial aggregation of cataract due to environmental and genetic causes, information was collected on major personal and environmental cataract risk factors, including cigarette, alcohol, exogenous estrogen and other medication use, lifetime ultraviolet-B light exposure, serum antioxidant levels, history of diabetes and other relevant medical conditions, and body mass index (BMI). To minimize misattribution due to as yet unaffected individuals who will eventually manifest the phenotype, we chose to work in an older population, with the minimum age for probands of 72 years. Finally, to reduce the impact of censored data from bilaterally pseudophakic individuals, we relied on previous study photographs and records from operating ophthalmologists to assign cataract grades for individuals without a native lens in either eye at the time of enrollment in the study.

MATERIALS AND METHODS

The Salisbury Eye Examination (SEE) initially recruited 2520 persons aged 65 to 84 years on a population basis from Medicare roles in the Salisbury area. This cohort has been observed with regular measure-
ment of various cataract risk factors through a total of four rounds of visits to date, with lens photographs having been obtained at rounds 1, 2 and 4. At rounds 3 and 4, all surviving subjects (n = 1504) were administered a family history questionnaire, and were eligible for the SEE Cataract Genetics (SEECAT) study if they had one or more full or half siblings living within 100 miles of Salisbury or Baltimore (Fig. 1). Consent was obtained from eligible SEE participants (proband) to contact their siblings. These siblings were then sent letters describing the study and containing stamped, self-addressed postcards that they could return to study headquarters if they did not want to be contacted further. Siblings from whom no card was received after 2 weeks were contacted by telephone and administered the family history questionnaire. Interested eligible siblings were invited to study headquarters.

After giving informed consent, all probands and siblings gave full medical and ophthalmologic histories and also answered questionnaires regarding their age; gender; lifetime ultraviolet-B exposure (methodology documented elsewhere); medical history including diabetes; and use of tobacco, alcohol, estrogen supplements, and other prescription and over-the-counter drugs. In addition, height was measured in stocking feet with a height board, and weight was obtained in kilograms on a digital scale. Seated blood pressure was measured in the right arm with a mercury sphygmomanometer. Three 7-mL EDTA tubes of blood were drawn by sterile venipuncture, two for later DNA analysis, and one that was centrifuged (average of two readings) in the right arm with a mercury sphygmomanometer. Three 7-mL EDTA tubes of blood were drawn by sterile venipuncture, two for later DNA analysis, and one that was centrifuged (average of two readings) in the right arm with a mercury sphygmomanometer. Three 7-mL EDTA tubes of blood were drawn by sterile venipuncture, two for later DNA analysis, and one that was centrifuged (average of two readings) in the right arm with a mercury sphygmomanometer. Three 7-mL EDTA tubes of blood were drawn by sterile venipuncture, two for later DNA analysis, and one that was centrifuged (average of two readings) in the right arm with a mercury sphygmomanometer.

**FIGURE 1.** Participants in the Salisbury Eye Evaluation Study of Cataract Genetics (SEECAT).

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![Diagram of Family Aggregation of Nuclear Cataract](image)

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Statistical Methods

Nuclear cataract was treated in both quantitative (Wilmer decimal grade 0–4.0) and binary fashion (with an individual defined as “affected” if the photograph grade was ≥3 or the assigned grade was “present” in either eye). The degree of family association for nuclear cataract was assessed in the binary analysis by the odds ratio (OR)
The parameters \( h \) that the heritability is twice the residual sibling correlation:

\[
equation \text{residual correlation between siblings} \\
\text{corr}(y_{ij}, y_{ik}) = 2 \rho \\
\text{where } \rho \text{ is the residual correlation.}
\]

In the analysis of the quantitative measure, we assumed a linear model, \( E(y_{ij} | x_{ij}, z_{ij}) = \beta x_{ij} + \gamma z_{ij} + \epsilon_{ij} \), with constant residual correlation between siblings \( \text{corr}(y_{ij}, y_{ik}) = \rho \). Note that the heritability is twice the residual sibling correlation: \( h^2 = 2 \rho \).

The parameters \( \beta \) and \( \gamma \) were estimated by generalized estimating equations (GEE) using the package \textsc{ggepca} version 0.2-4 (available at http://cran.r-project.org/src/contrib/Descriptions/ggepca.html) with the R statistical system version 1.7.1.18

In the analysis of the binary measure, we followed the approach of Liang and Beaty19 and assumed a logistic model, \( \logit (\text{Pr}(z_{ij} = 1 | x_{ij})) = \theta x_{ij} + \gamma z_{ij} \), with constant log odds ratio in \( \text{OR}(z_{ij} = 1 | x_{ij} = 0, z_{ij} = 1) = \gamma \). The parameters \( \theta \) and \( \gamma \) were again estimated by GEE, as described by Liang et al.20

Calculations were performed in R version 1.7.1, using the package \textsc{geesb}, available from the authors.

With both analyses, standard errors were obtained by the robust sandwich estimator.16 Covariates were chosen by stepwise selection. A covariate was retained in the model if the corresponding probability was less than 0.1.

**RESULTS**

A total of 307 probands and their 434 full siblings (total \( n = 741 \)) participated in the study, forming a total of 274 sibships of one to eight individuals and an average size of 2.7. (Fig. 1) The number of sibships is smaller than the number of probands, because some of the individuals recruited from the SEE study as probands were siblings of other probands. Probands were significantly older than their siblings, and were more likely to have nuclear cataract, bilateral pseudophakia and several other risk factors for cataract, though none of these differences except the greater nuclear cataract prevalence remained significant after age adjustment (Table 1). Approximately 30% of subjects were black. Cataract grades could be assigned for 64% and 72% of bilaterally pseudophakic/aphakic probands and siblings, respectively (Table 1). Among persons with nuclear cataract, 49% had pure nuclear, 23% nuclear mixed with posterior subcapsular cataract (PSC), 17% nuclear mixed with cortical, and 11% nuclear mixed with PSC and cortical opacity.

When probands and siblings were considered together in a regression model with nuclear cataract grade as a continuous outcome, age was found to be positively associated with the presence of nuclear cataract, as was female gender, white race, and a history of smoking. BMI, alcohol use, presence of diabetes, use of steroids and exogenous estrogens, and the various serum antioxidants were not significantly associated with nuclear cataract (Table 2).

The odds of being affected by nuclear cataract was elevated by twofold among those with an affected sibling (OR 2.08, 95% CI 1.39–3.10), a significant association that persisted after adjusting for personal and environmental risk factors for cataract measured in our study (OR 2.07, 95% CI 1.30–3.50; Table 3). The heritability of nuclear cataract in this population was 35.6% (95% CI: 21.0%–50.3%), suggesting that some 36% of the variance in nuclear cataract grade can be attributed to genetic causes (Table 3).

**DISCUSSION**

Our results provide evidence that genetics plays a significant role in nuclear cataract, the most prevalent type of cataract in European-derived populations,21 and the cataract subtype most commonly requiring surgery.22 The finding that nuclear cataract is genetically as well as environmentally determined in consistent with previous population-based investigations21 and twin studies.10 The fact that several independent lines of investigation in different populations have supported the heritability of nuclear cataract in this population was 35.6% (95% CI: 21.0%–50.3%), suggesting that some 36% of the variance in nuclear cataract grade can be attributed to genetic causes (Table 3).

**TABLE 1.** Characteristics of Participants in the Salisbury Eye Evaluation Study (Probands) and Their Siblings, with Regard to Demographics, Cataract Risk Factors, Nuclear Cataract Status, and Pseudophakia

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Probands</th>
<th>Siblings</th>
<th>Unadjusted Difference</th>
<th>Age-Adjusted Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age (y)</td>
<td>77.6 ± 4.5</td>
<td>72.4 ± 7.4</td>
<td>&lt;0.001</td>
<td>—</td>
</tr>
<tr>
<td>Gender (female)</td>
<td>180/307 (58.6)</td>
<td>254/454 (58.5)</td>
<td>1.0</td>
<td>—</td>
</tr>
<tr>
<td>Race (black)</td>
<td>86/307 (28.0)</td>
<td>119/434 (27.4)</td>
<td>0.016</td>
<td>0.34</td>
</tr>
<tr>
<td>Mean body mass index</td>
<td>28.4 ± 5.5</td>
<td>29.4 ± 5.6</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Smoking status (%) (never/former/current)</td>
<td>45/42/12</td>
<td>43/47/10</td>
<td>0.43</td>
<td>0.86</td>
</tr>
<tr>
<td>Alcohol status (%) (never/former/current)</td>
<td>30/48/21</td>
<td>29/45/25</td>
<td>0.51</td>
<td>0.49</td>
</tr>
<tr>
<td>Systolic or diastolic hypertension (%)</td>
<td>198/302 (66)</td>
<td>239/434 (55)</td>
<td>0.005</td>
<td>0.065</td>
</tr>
<tr>
<td>Diabetes (%)</td>
<td>77/302 (25)</td>
<td>83/434 (19)</td>
<td>0.046</td>
<td>0.097</td>
</tr>
<tr>
<td>Current or recent steroid use (%)</td>
<td>22/276 (6.8)</td>
<td>41/432 (9)</td>
<td>0.59</td>
<td>0.24</td>
</tr>
<tr>
<td>Bilateral pseudophakia (%)</td>
<td>78/307 (25.4)</td>
<td>69/434 (15.9)</td>
<td>0.002</td>
<td>0.85</td>
</tr>
<tr>
<td>Able to assign cataract grade (among pseudophakics)</td>
<td>50/78 (64.1)</td>
<td>50/69 (72.5)</td>
<td>0.29</td>
<td>0.61</td>
</tr>
<tr>
<td>Nuclear cataract in either eye (%)*</td>
<td>141/274 (51)</td>
<td>121/406 (30)</td>
<td>&lt;0.001</td>
<td>0.036</td>
</tr>
</tbody>
</table>

*Includes persons with nuclear cataract grade ≥3.0 on SEE round 3 photographs and also persons determined to be “affected” by nuclear cataract on the basis of clinical records and/or old photographs.

**TABLE 2.** Association of Various Covariates with Nuclear Cataract Grade among a Population of Older Persons and Their Siblings in Salisbury, Maryland

<table>
<thead>
<tr>
<th>Covariate</th>
<th>Beta Coefficient (95% Confidence Interval)</th>
<th>Standard Error</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (per year)</td>
<td>0.045 (0.036–0.054)</td>
<td>0.0046</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Female gender</td>
<td>0.212 (0.092–0.332)</td>
<td>0.061</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>White race</td>
<td>0.289 (0.150–0.428)</td>
<td>0.071</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Smoking (per pack-year)</td>
<td>0.0025 (0.0005–0.0045)</td>
<td>0.0010</td>
<td>0.014</td>
</tr>
</tbody>
</table>

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Table 3. Odds Ratio of Development of Nuclear Cataract among Persons with a Sibling Having Nuclear Cataract Compared with Those without Affected Siblings and Adjusted and Unadjusted Heritability of Nuclear Cataract

<table>
<thead>
<tr>
<th>Nuclear Cataract</th>
<th>Unadjusted</th>
<th>Adjusted*</th>
</tr>
</thead>
<tbody>
<tr>
<td>OR</td>
<td>2.08 (1.39–3.10)</td>
<td>2.07 (1.30–3.30)</td>
</tr>
<tr>
<td>Heritability</td>
<td>54.9% (31.5–78.3)</td>
<td>35.6% (21.0–50.3)</td>
</tr>
</tbody>
</table>

* Adjusted for age, gender, race, body mass index, use of tobacco and alcohol, blood pressure, diabetes, use of steroids, total serum carotenoids, and lifetime ultraviolet-B light exposure. Only age, gender, race, and smoking status were significantly associated in the model and were used for the final adjustment.

Heritability of nuclear cataract suggests that this finding is comparatively robust. The 36% heritability estimate for nuclear cataract in our study, an index of the proportion of cataract variation controlled by genetic causes, is comparable to that which has been reported in other studies. Hammond et al. reported a heritability figure of 48% in their twin study of nuclear cataract, and Heibl et al. estimated that a single major gene may account for 35% of nuclear cataract variation in the Beaver Dam Eye Study population.

In the regression model used in the present study, advancing age, female gender, white race, and smoking history were associated with nuclear cataract grade. All these factors have been identified previously in epidemiologic studies. Certain other covariates that have been found to be positively or negatively associated with nuclear cataract in the past, including alcohol intake, BMI, serum antioxidant status, and steroid use, were not found to predict nuclear cataract outcome significantly in our models. The list of covariates found to be predictive in our model included factors that have been most robust across numerous studies, whereas the associations with nutrition, alcohol intake, and BMI have been either less consistent or of uncertain directionality. Although steroid use has consistently been associated with development of cataract in many studies, regular users of steroids were relatively rare in this population.

The shortcomings of this study in providing reliable information about the heritability of nuclear cataract must be acknowledged. In the first place, our population was selected at baseline to be 65 years and older, and had reached a mean age in the mid-70s by the time of the present study. This is more than a decade older than the population studied by Hammond et al., for example, with a mean age of 62 years. There are both advantages and disadvantages to working with a population in the eighth and ninth decades of life in performing genetic studies on ocular diseases of aging. Although the chances of misclassification of affection status due to a younger individual having failed yet to manifest the phenotype of interest is reduced; however, the impact of environmental as opposed to genetic influences may be magnified in an older population. In addition, the effect of bilateral pseudophakia, where the opportunity directly to examine the lens phenotype has been lost, is increased among older individuals. Finally, the effects of survivorship, whereby persons living longer may be healthier than those dying younger, and of bias introduced by the inability of sicker and more physically impaired individuals to participate, are exaggerated in an older group.

In an attempt to reduce the impact of one of these problems inherent to cataract studies in an older population, the high prevalence of bilateral pseudophakia, we assigned grades for approximately one of eight probands in our study, based on photographs from previous rounds of SEE and preoperative clinical notes by ophthalmic surgeons in the area. Although this effort was successful in obtaining information for most of the bilaterally pseudophakic subjects, several assumptions were made: that a grade ≥2+ (on a scale of 4) or the notation of "dense" by a clinician was equivalent to a grade of ≥3.0 in a standardized cataract grading scale as implemented by trained graders and that clinical notations across different nonstandardized ophthalmologists and over time were equivalent. Obviously, these assumptions are not entirely valid and introduced an unavoidable degree of noise into our primary outcome measures. We decided that this inevitable lack of precision was nonetheless acceptable, given the alternative of reduced power, and possible introduction of bias, from the systemic censoring of a large number of the most severely affected persons in the population that would have been necessary without some imputation strategy. Our use of the current approach is to some extent validated by the fact that our results were similar in magnitude and direction whether the estimated grades were included or not, although confidence intervals were of course somewhat tighter with the use of the additional data.

The rate of participation among eligible subjects in this study was modest, and in the case of siblings of SEE participants, fell below 50%. It is possible that the low participation resulted in bias, though estimates of heritability, the principle outcome for the study, would be biased only if persons whose cataract status was concordant with that of their siblings were more or less likely to participate than those with discordant phenotypes. Because we have no information regarding the lens status of nonparticipating siblings, we are unable to exclude this possibility, but it is not immediately obvious that factors would lead to such differential participation between concordant and discordant sibships. The relatively low rates of participation also resulted in a smaller total sample size and thus wider confidence intervals around our heritability estimate.

Finally, the conclusions reached by this study regarding genetic influences on nuclear cataract must to some extent depend on the assumptions inherent in the statistical models used. In the first place, by defining the residual correlation between siblings as a measure of the association due to genetic influences, we have implicitly assumed that there are no additional, unmeasured personal or environmental risk factors, the sharing of which between siblings may actually have accounted for residual correlation not due to measured factors. This assumption is in some ways difficult either to defend or attack. By definition, putative unknown risk factors cannot be enumerated or their existence discounted. It can only be said that in the present study we assessed nearly all factors shown in the past two decades of cataract epidemiology to impact on the development and progression of lens opacity. It is also possible that genetic influences on nuclear cataract is mediated through an individual’s response to the effects of smoking and other such risk factors—that is, through gene–environment interactions. In a model such as the one used here, adjusting on such factors may actually lead to an underestimate of the heritability of nuclear cataract. The statistical model used in this report has no way to account for the impact of possible gene–environment interactions.

Studies of the genetics of age-related cataract are in their infancy. Though a small number of genes have been reported to be associated with cataract in limited populations, particularly in Japan (galactokinase, glutathione-S-transferase), the reported impact been modest, and these findings have generally not been replicated in other populations. Though known, Mendelian-inherited forms of congenital cataract provide several potential candidate genes for age-related cataract; however, given the very different phenotype and age of onset between these two types of cataract, the applicability of these candidates is far from clear. The immediate future direction of cataract genetic studies will probably involve genome-wide scans, based either on individual studies or consortiums.
between studies in an effort to increase power. Large, population-based studies of cataract in which investigators have not concentrated on the collection of families, but have obtained blood samples, are likely to be in the position to contribute best through case-control studies of positional candidates in areas highlighted by genome-wide scans. It is to be hoped that such efforts will take the field from where it is now, where most information about cataract has come from traditional, population-based studies not organized to explore genetic issues, toward a better understanding of the genes contributing to cataract development. The minimal data currently available from genetic studies of other ocular diseases of aging, such as age-related macular degeneration, suggest that such genes may be numerous, and their individual impact modest.

In view of such a complex model, what likely practical benefits can we expect from cataract genetic studies? It seems unlikely that this ubiquitous, late-onset disease with an existing surgical cure will provide a practical forum for resource-intensive strategies such as gene therapy or population or clinic-based screening of potentially affected persons. A more likely scenario is that studies of the cataract genetic studies will eventually yield knowledge of the protein pathways involved in lens opacity, so that discovery of anticataract agents may proceed in a rational fashion, rather than through the current process of hit or miss. To be practical, such agents probably should be delivered on a mass basis as supplements, which sets a high standard indeed for safety and low cost. However, the power of supplementation—a high standard indeed for safety and low cost. However, the process of hit or miss. To be practical, such agents probably should be delivered on a mass basis as supplements, which sets a high standard indeed for safety and low cost. However, the last century of medical progress provides at least two examples of the power of supplementation—fluoridated water and iodized salt—interventions that should serve as models for future work in the area of cataract prevention.

References