The Inheritance of Peripapillary Atrophy

Paul R. Healey,1 Paul Mitchell,2 Clare E. Gilbert,2 Anne J. Lee,1 Dongliang Ge,3 Harold Snieder,3,4 Timothy D. Spector,1 and Christopher J. Hammond4,5

PURPOSE. To estimate the relative importance of genes and environment in peripapillary atrophy type beta (β-PPA) in a classic twin study.

METHODS. Female twin pairs (n = 506) aged 49 to 79 years were recruited from the St. Thomas’ UK Adult Twin Registry. Peripapillary atrophy was identified from masked grading of stereoscopic optic disc photographs. Structural equation modeling was performed using Mx with polychoric correlations of β-PPA and refractive error (divided into deciles).

RESULTS. β-PPA prevalence was 25.1% and did not vary with zygosity. Case-wise concordance for right eyes was 0.76 (95% CI, 0.57–0.88) for monozygotic (MZ) and 0.37 (95% CI, 0.15–0.56) for dizygotic (DZ) pairs. Multivariate modeling suggested additive genetic effects and individual environment, with no shared environment or dominant genetic effect. β-PPA heritability was 0.70 (95% CI, 0.54–0.85), and spherical equivalent 0.88 (95% CI, 0.85–0.91); age had no significant effect on variance. The genetic correlation between β-PPA and spherical equivalent was −0.21. However, only 3% of the genetic variance of β-PPA was explained by genetic factors in common with refractive error, with 67% explained by specific genetic factors for β-PPA. Of the 30% of variance explained by unique environmental factors, only 3% was explained by these factors in common with environmental factors involved in refractive error.

CONCLUSIONS. The presence of β-PPA, a frequent ocular finding known to be associated with open-angle glaucoma, appears to be under strong genetic control, with only a small amount of this genetic effect shared with genes involved in myopia.


Choroidal atrophy surrounding the optic disc (known as either perifoveal or parapapillary atrophy) is a relatively common clinical finding. Jonas et al.1 have developed a classification of two zones: an area of visible sclera and large choroidal vessels, adjacent to the optic disc border, termed zone beta, and a more peripheral area of irregular hypo- and hyperpigmentation, termed zone alpha.1 This classification into β- and α-peripapillary atrophy (β-PPA and α-PPA) has now been accepted into common use, with reports of an association with glaucoma made principally with β-PPA. β-PPA atrophy has been reported to be associated with both the prevalence2–7 and progression8–11 of open-angle glaucoma (OAG). The presence of β-PPA12 and its enlargement over time12,13 have been reported to precede and predict the onset of glaucoma in ocular hypertensive eyes.

In examining the heritability of OAG, it is important to examine the inheritance of OAG risk factors. A familial aggregation of risk factors for OAG suggested that optical disc signs may be stronger determinants of glaucoma in relatives than intraocular pressure (IOP).14–16

The frequency of β-PPA has been reported to increase with increasing levels of myopia.15 However, in these eyes, the etiology of the peripapillary morphology may not be uniform. Congenital anomalies of peripapillary tissue alignment and changes induced by scleral stretching may resemble age-related atrophic β-PPA.15,16 Myopia is also associated with OAG,17 suggesting that true atrophic β-PPA may occur as well.

OAG and myopia are both known to have genetic and environmental components, although the environmental component of neither is well understood. The relative genetic and environmental components of β-PPA and its relationship to both myopia and environmental factors are unknown. Such information may give insight into the relative contribution of inheritance in β-PPA and the relationship between β-PPA, myopia, and OAG. This study was conducted to analyze the relative importance of genes and environment on β-PPA in a large sample of adult twins.

METHODS

Subjects

The study population consisted of 506 female twin pairs aged between 50 and 79 years, recruited from the Twins UK Adult Twin Registry, held at St. Thomas’ Hospital in London. Registrants were ascertained from the general population of the United Kingdom through national media campaigns before the planning of any eye studies. Details of the registry have been described previously.20 Study subjects were examined between January 1998 and July 1999. Twelve pairs of twins were unable to attend for reasons of ill health or refusal to participate. Zygosity was determined in 506 twin pairs by standardized questionnaire21 and confirmed by DNA short tandem repeat fingerprinting in approximately 40% of twin pairs in whom zygosity was uncertain. This procedure was used if there was any doubt about true zygosity or when the answers to the standardized questionnaire were not definitely monozygotic (MZ) or dizygotic (DZ).

Measurements

Examination included visual acuity testing using the ETDRS logMar (Early Treatment Diabetic Retinopathy Study logarithm of the minimum angle of resolution) chart and undilated refraction with an automatic refractor (Humphrey 670; Carl Zeiss Meditec, Dublin, CA). Spherical equivalent (SE) refraction was calculated in each eye. SE data were reproducible, and their distribution has been reported.22 After pupil dilatation, nonsimultaneous stereoscopic photographs centered...
on the optic nerve were taken of both eyes of all the twins. A camera (Kowa-Europe, Dusseldorf, Germany) was used with a 30° field setting (Ektachrome 64 film; Eastman-Kodak, Rochester, NY). The same individual took all photographs, using the same camera, and the film was processed by the same laboratory. Photographs were examined using stereoscopic viewing spectacles on an x-ray viewing light box. The best stereo pair was digitized with a slide scanner (Coolscan III; Nikon, Tokyo, Japan). Images of 1280 by 960 pixels were created at a magnification of 135%.

Grading of Peripapillary Atrophy

The images were graded for the presence of peripapillary atrophy by one of two graders masked to the pairing and zygosity of subjects. The classification described by Jonas et al. was used to identify peripapillary atrophy. No deviations from this system were allowed. If present, \(\beta\)-PPA bordered the peripapillary scleral ring. It was characterized by visible large choroidal vessels and sclera, due to a denuded Bruch’s membrane and loss of retinal pigment epithelium. It was distinguished from \(\alpha\)-PPA, characterized by irregular retinal hypo- or hyperpigmentation, which bordered zone beta or the scleral ring when the beta zone was absent. Examples of these two types of peripapillary atrophy are shown in Figure 1. \(\beta\)-PPA was graded as present, uncertain, absent, ungradable, or image missing and entered into an electronic database. Where graders disagreed or when an image was graded as uncertain, adjudication was made by one of the authors (PRH).

Analytical Approach

Data analysis was based on quantitative genetic modeling\(^22,23\) using the Mx program.\(^24\) In short, the technique is based on the comparison of the covariances (or correlations) within MZ and DZ twin pairs. Familial aggregation of a trait or disease suggests that genetic factors may be involved in etiology, but it does not exclude the possibility of shared environmental factors. Twin modeling assumes that MZ and DZ twin pairs share the same common (family) environment—the “equal environment” assumption—which has been tested and largely found to be true.\(^25\) As MZ pairs share the same segregating genes but DZ pairs share only half, any greater similarity between MZ pairs allows an estimation of this additional gene-sharing, using the equal-environment assumption. To quantify the genetic and environmental contributions to a dichotomous variable, such as the presence or absence of \(\beta\)-PPA, an underlying, continuous liability to \(\beta\)-PPA is assumed, which is affected by multiple genetic and environmental factors.\(^20\) Structural equation modeling uses variance-covariance matrix algebra to separate the observed phenotypic variance into additive (G) or dominant (D) genetic components and common (C) or unique (E) environmental components (E also includes measurement error). MZ twins share the same genetic component, DZ twins 0.5, and MZ twins the D genetic component, whereas DZ twins share only 0.25 of D, since there is a one-in-four chance that two siblings will share a dominant allele transmitted from one parent. Thus, the known twin-twin relationships were used in maximum likelihood methods to estimate the best-fitting model that fits the variance-covariance data obtained in the study. Dividing each of these components by the total variance yields the different standardized components of variance—for example, the heritability, which can be defined as the ratio of additive genetic variance to total phenotypic variance.

As \(\beta\)-PPA is a bivariate variable (yes/no) and refractive error is measured as a continuous variable (mean SE), to include both variables in a single model, polychoric correlations were estimated by using a saturated model, with refractive error divided into deciles to recode it.
from a continuous measurement to a categorical one, allowing it to be modeled with the dichotomous \( \beta \)-PPA data. The univariate models were extended to include the bivariate cases of SE and \( \beta \)-PPA, to allow assessment of the extent to which any correlation between these two variables could be explained by common genes.

**Model Fitting Procedure**

Model fitting is used to try to explain the data in the most economic (parsimonious) manner. A backward-elimination procedure from the full model was used to develop a final model for the variance components G, C, and D in which the pattern of variances and covariances was explained by as few parameters as possible. Submodels were compared with the full model by hierarchic \( \chi^2 \) tests. The difference in \( \chi^2 \) values between submodel and full model is itself approximately distributed as a \( \chi^2 \), with degrees of freedom (df) equal to the difference in df of submodel and full model. Statistical software data handling and preliminary analyses were performed with commercial software (Stata Corp., College Station, TX). Research adhered to the tenets of the Declaration of Helsinki, and local ethics committee approval and informed consent were obtained.

**RESULTS**

Optic disc photographs were gradable in 1813 eyes of 944 subjects (93.5%) comprising 209 MZ twin pairs and 263 DZ twin pairs.

\( \beta \)-PPA was present in at least one eye of 254 subjects (25.1%). The ocular prevalence was 22.9% in the right eyes and 21.7% in the left eyes. \( \beta \)-PPA was bilateral in 150 subjects or 59% of those with this sign. Figure 1 shows examples of \( \beta \)-PPA subtypes in MZ and DZ eyes. \( \beta \)-PPA prevalence did not vary significantly with zygosity (Tables 1 and 2).

**TABLE 1. \( \beta \)-PPA Concordance in Twin Pairs**

<table>
<thead>
<tr>
<th></th>
<th>( \beta )-PPA</th>
<th>No PPA</th>
<th>Total</th>
</tr>
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<tbody>
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<tr>
<td>( \beta )-PPA</td>
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<td>24</td>
<td>156</td>
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<tr>
<td>No PPA</td>
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<td>30</td>
<td>44</td>
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<tr>
<td>Total</td>
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<td>54</td>
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<td>51</td>
<td>188</td>
</tr>
<tr>
<td>No PPA</td>
<td>34</td>
<td>28</td>
<td>62</td>
</tr>
<tr>
<td>Total</td>
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<td>79</td>
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**Figure 2.** The prevalence of \( \beta \)-PPA by degree of myopia in diopters (D) using SE refraction, for right and left eyes.
DISCUSSION

The prevalence of β-PPA in this twin study is similar to that reported in singletons. In a very large study of normal subjects in Japan, β-PPA was found in 26%. Smaller clinic-based estimates of β-PPA frequency generally lie between 12% and 20%. This study confirms the relationship between β-PPA and myopia that has been reported in population-based studies, which may partially explain the differences in prevalence estimates between studies. The similar prevalence of β-PPA in MZ and DZ twins suggests that there is no genetic association between β-PPA and being a member of a set of twins. These findings are in agreement with a systematic study of disease- and lifestyle-related characteristics between twins and singletons that suggests that generalization from twins to singleton populations is valid.

Our data suggest that β-PPA has a strong heritable component. A heritability of 75% compares with 90% for myopia alone in this cohort, 58% for cortical cataract, and 45% for age-related maculopathy. Lower heritability of degenerative conditions such as ARM and cataract implies more population variation attributable to environmental factors. If one supposes β-PPA to be (as its name suggests) a feature caused by degeneration of the peripapillary tissues, then it is surprising that the heritability is closer to refractive error, which is generally occurs at a young age. It does not appear that common genes for myopia and β-PPA are the cause, as we estimated only 3% of the genetic variance to have a common genetic basis. The explanation may relate to the possibility that β-PPA has several etiologies.

Two recent studies of children have suggested that in myopic eyes, peripapillary morphology may be partly congenital or acquired very early in life. One study reported a significant (34%) prevalence of myopic peripapillary crescents in young Japanese children. In contrast, the prevalence of chorioretinal atrophy was only 16%. In the second study, Singaporean children were examined. It reported an association between increasing frequency of myopic temporal crescents and increasing myopic refraction and axial length but not an association with age. The hypothesis of a multifactorial etiology for myopic β-PPA is further strengthened by an angiographic study of myopic peripapillary morphology in adults. This study suggested that there may be two components to clinical β-PPA in these eyes: one related to stretching of the sclera and the other to vascular changes.

Histologic studies of peripapillary atrophy in adults also suggest several distinct entities: simple misalignment of the choroid and RPE, malpositioning of the embryonic fold between the RPE and outer retina, and age-related atrophy of the peripapillary RPE and rods, usually surrounded by an area of RPE degeneration. In one study, atrophy was found in 15 of 17 otherwise normal eyes, suggesting that it may be a normal aging change. It varied from complete absence of the RPE to a loss of RPE pigmentation.

It is not possible to distinguish clinically between these three histologic entities of peripapillary atrophy. A clinical distinction can be made between areas of severe or complete loss of RPE pigment and areas with irregular pigmentation. A peripapillary area through which sclera and large choroidal...
vessels can be seen is termed zone beta peripapillary atrophy. Areas of irregular hypo- or hyperpigmentation adjacent to the optic disc border or zone beta (when present) are termed zone alpha peripapillary atrophy.7

Clinically prevalent β-PPA is not nearly as common as histologic atrophy, even taking into account misclassification of congenital peripapillary misalignment.7 This disparity may be partly because the β-PPA is obscured by the overlying nerve fiber layer.35,39 Because there is no clear relationship between β-PPA defined clinically and true atrophy as defined histologically, one must presume a large potential for classification bias when using the clinical finding to classify the histologic state. The β-PPA, as ascertained in all clinical studies reported to date, may consist of both true atrophy and congenital malformations of the retinal layers. Were it possible to distinguish the two, the congenital variant of β-PPA might be highly heritable, whereas the atrophic variant may be more influenced by environmental factors. There may also be a difference between the relationships of each type of β-PPA with glaucoma. This hypothesis could be examined further by comparison of β-PPA heritability in younger adult and pediatric twin studies, as well as comparison with one of the several large studies of childhood refractive error, which are currently under way.40,41

CONCLUSIONS

This study provides the first information about the heritability of β-peripapillary atrophy and its relationship to refractive error. We estimate that the genetic effects are important with a heritability adjusted for refraction of 70%. We confirmed an association with myopia but suggest that this not due to common genes. Rather, it is predominately due to the co-inheritance of genes specific for β-peripapillary atrophy. Although the relationship between the clinical sign and histologic phenotype needs further clarification, it raises the possibility of a genetic basis for optic disc morphologic characteristics associated with glaucoma.

Acknowledgments

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References


