Heritability of Strabismus: Genetic Influence Is Specific to Eso-Deviation and Independent of Refractive Error

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Strabismus represents a complex oculomotor disorder characterized by the deviation of one or both eyes and poor vision. A more sophisticated understanding of the genetic liability of strabismus is required to guide searches for associated molecular variants. In this classical twin study of 1,462 twin pairs, we examined the relative influence of genes and environment in comitant strabismus, and the degree to which these influences can be explained by factors in common with refractive error. Participants were examined for the presence of latent (‘phoria’) and manifest (‘tropia’) strabismus using cover-uncover and alternate cover tests. Two phenotypes were distinguished: eso-deviation (esophoria and esotropia) and exo-deviation (exophoria and exotropia). Structural equation modeling was subsequently employed to partition the observed phenotypic variation in the twin data into specific variance components. The prevalence of eso-deviation and exo-deviation was 8.6% and 20.7%, respectively. For eso-deviation, the polychoric correlation was significantly greater in monozygotic (MZ) (r = 0.65) compared to dizygotic (DZ) twin pairs (r = 0.33), suggesting a genetic role (p = .003). There was no significant difference in polychoric correlation between MZ (r = 0.55) and DZ twin pairs (r = 0.53) for exo-deviation (p = .86), implying that genetic factors do not play a significant role in the etiology of exo-deviation. The heritability of an eso-deviation was 0.64 (95% CI 0.50–0.75). The additive genetic correlation for eso-deviation and refractive error was 0.13 and the bivariate heritability (i.e., shared variance) was less than 1%, suggesting negligible shared genetic effect. This study documents a substantial heritability of 64% for eso-deviation, yet no corresponding heritability for exo-deviation, suggesting that the genetic contribution to strabismus may be specific to eso-deviation. Future studies are now needed to identify the genes associated with eso-deviation and unravel their mechanisms of action.

Keywords: twins, heritability, ophthalmology, strabismus

Comitant strabismus (heterotropia or squint) is a complex disorder characterized by constant or intermittent ocular misalignment that often results in reduced or absent binocular vision and subsequent amblyopia of the deviated eye.
The condition affects 2–6% of the general population and is the most common cause of poor vision in childhood (Graham, 1974; Robaei et al., 2006; Williams et al., 2008). Comitant strabismus is not associated with systemic abnormality and is marked by non-restrictive, non-paralytic ocular misalignment of the same magnitude in all directions of gaze. There are two primary forms of horizontal comitant strabismus (hereafter referred to as ‘strabismus’): esotropia (convergent strabismus in which one or both eyes turn inwards) and exotropia (divergent strabismus in which one or both eyes turn outwards). In contrast, the more common form of ocular motor status, heterophoria, represents a ‘latent’ type of strabismus in which binocular vision functions normally except under certain conditions such as tiredness or illness, or that which may give rise to asthenopic (eye-strain) symptoms. Like manifest strabismus, heterophoria can be classified by the direction of horizontal deviation (esophoria — inwards, exophoria — outwards). Binocular vision status, the type and magnitude of deviation and the differentiation of heterophoria and heterotropia may be assessed with the cover test. It is important to distinguish that although disease-causing genes and loci have been identified for incomitant strabismus, often referred to as congenital cranial dysinnervation disorders (Engle, 2007; Gutowski et al., 2003; Traboulsi, 2007), currently we have limited understanding of the more prevalent comitant type, the focus of the current study.

Notwithstanding the lack of definitive data in the literature on the etiological foundation for strabismus, it is often assumed to be a multifactorial anomaly with both genetic and environmental risk factors. The tendency for the disorder to run in families, which could result from either genetic or environmental factors, has been recognized for centuries, with Hippocrates reporting in 400 BC that ‘children of parents having distorted eyes squint for the most part’ (Jones, 1886). More recent studies have confirmed a higher prevalence of strabismus among families of an affected individual than among the general population (Abrahamsson et al., 1999; Cross, 1975; Podgor et al., 1996; Waardenburg, 1954; Ziakas et al., 2002). Twin studies, which allow genetic and environmental influence to be distinguished, have shown higher concordances for strabismus within identical (monozygotic [MZ]) twin pairs compared to their non-identical (dizygotic [DZ]) counterparts, thus supporting a genetic contribution to the disorder (Wilmer & Backus, 2009). Recently, Wilmer and Backus published a critical review and meta-analysis of twin strabismus literature. Based on 3,096 4- to 7-year-old twin pairs in studies that met acceptable methodological standards, the meta-analysis estimated a high 92% genetic contribution to strabismus (Wilmer & Backus, 2009). However, due to insufficient subtyping in prior studies, the authors were unable to examine convergent (eso) and divergent (exo) subtypes of strabismus separately nor did prior studies contain enough information to evaluate whether strabismus and refractive error share a common genetic basis.

Environmental risk factors have also been observed to predict one’s susceptibility to strabismus. Neonatal risk factors such as low birth weight and prematurity may be associated with the disorder developing in childhood (Bremner et al., 1998; Chew et al., 1994; Keith & Kitchen, 1983; Robaei et al., 2006; Torp-Pedersen et al., 2010; Williams et al., 2008). This is supported by the fact that twins, commonly born premature and with low birth weight, have a higher prevalence of strabismus than the general population (Wilmer & Backus, 2009). Moreover, DZ twins, who share a common prenatal environment, demonstrate higher strabismus concordance than full siblings (even after accounting for different prevalences between twins and singletons) (Wilmer & Backus, 2009). Several studies have found that maternal smoking during pregnancy predicts disrupted visual development (Chew et al., 1994; Christianson, 1980; Hakim & Tielisch, 1992). Our previous work from the Twins Eye Study in Tasmania (TEST) similarly found that maternal smoking predicts delayed age at first crawling or walking, potentially due to adverse impacts on vision (Ponsonby et al., 2007). Additionally, in the Danish National Birth Cohort (DNBC) of 96,842 children, large head circumference and congenital abnormalities — possibly affected by environmental factors — were associated with increased risk of strabismus (Torp-Pedersen et al., 2010).

We evaluated the heritability of strabismus in a large twin population. To our knowledge, this is the first large study to utilize clinical data (rather than parental reporting) or to distinguish between eso- and exo-deviations. Furthermore, this work is novel in examining the genetic association between refractive error and strabismus and the magnitude of potential pleiotropic effects.

**Methods**

**Participants**

Twin pairs were identified from three cohorts: the TEST (Mackey et al., 2009), the Australian Twins Eye Study (ATES) (Wright & Martin, 2004), and the TwinsUK Adult Twin Registry based at St Thomas’ Hospital in London (Spector & Williams, 2006). The vast majority of subjects were Caucasian. The TEST and ATES cohorts consisted of younger twin pairs of both sexes in contrast to the predominantly older, female demographic of the UK Twins group (the Registry was originally established to study osteoporosis). Recruitment was conducted without providing subjects with knowledge of specific hypotheses or eye studies being conducted, reducing ascertainment bias. The relevant ethics committees of the Royal Victorian Eye and Ear Hospital, University of Tasmania, and St Thomas’ Hospital approved the study. The study adhered to the tenets of the Declaration of Helsinki.
Examination Protocol

Subjects underwent a comprehensive clinical examination that included: assessment of binocular alignment, anterior segment examination, corneal pachymetry, intraocular pressure measurement, autorefractio, and a mydriatic optic disc assessment. A detailed ocular history was obtained including history of amblyopia, family history of strabismus, previous ocular surgery, and age at first wearing glasses. A cover–uncover and alternating cover test was performed on all subjects at standard near (33 cm) and far (6 m) test distances (Mackey et al., 2009). The presence of any ocular misalignment (manifest or latent) was recorded or, alternatively, in the absence of deviation, orthophoria was noted. Subjects with strabismus on examination associated with previous history of trauma or ocular surgery to correct the misalignment were excluded from the study. Stereoscopic depth perception was measured using the TNO test. Spherical equivalent and total astigmatism was recorded, in dioptres (D), for each eye. For all twin pairs, zygosity was confirmed by DNA analysis of short tandem-repeat and subsequent Genome-Wide Association Studies.

Statistical Analysis

Familial aggregation of a specific trait or disease occurs when relatives who share genes and/or environmental factors are more phenotypically similar than unrelated individuals. The twin study provides an ideal means for partitioning such familial aggregation into the effect of genes and shared environment on a phenotype of interest. The simplest method is to determine whether identical (MZ) twins raised together are more similar to each other for a given trait than non-identical (DZ) twins raised together. If a genetic basis for the trait exists, one would expect MZ twins to exhibit greater trait concordance or correlation as they share 100% of their genetic material, compared to DZ twins who on average share only 50%.

Data management and statistical tests were performed in the R statistical environment (R Development Core Team, 2010). Two phenotypes were examined: eso-deviation (esophoria and esotropia) and exo-deviation (exophoria and exotropia). A continuous ‘liability’ model was employed utilizing multiple thresholds to account for the categorical nature of the data. In this way an underlying normally distributed risk to strabismus in the population is assumed, which is affected by multiple genetic and environmental factors (Falconer & Mackay, 1996). Thus, each phenotype was classified in an ordinal manner: normal score = 0, phoria score = 1, and tropia score = 2. Structural equation modeling (SEM) was subsequently employed to fit the data to a saturated model in which polygenic correlations and thresholds were freely estimated. Genetic and environmental parameters that best fit the observed twin covariances were estimated directly from the raw categorical data using OpenMx software (Boker et al., 2011). Potential sex differences in the genetic and environmental influences on strabismus were investigated by equating twin pair correlations in a stepwise manner for five zygosity groups: MZ males, MZ females, DZ males, DZ females, and DZ opposite-sex twins. ‘Quantitative’ sex differences were initially examined by equating correlations between MZ males and MZ females, and DZ males and DZ females (Neale & Cardon, 1992). ‘Qualitative’ sex differences were examined by equating correlations between same- and opposite-sex DZ pairs.

To determine heritability, a series of hierarchical models were fitted to the raw data and the significance of each assessed. Each of the nested sub-models was then compared to the full model by χ² tests. Twice the difference between log-likelihood values of the nested and full models is asymptotically distributed as χ² with degrees of freedom (df) equal to the difference in parameters being estimated. The Akaike information criterion (AIC) was used to determine the best-fitting mode by evaluating model parsimony (i.e., the best goodness-of-fit combined with the fewest latent variables). The model with the lowest AIC suggests the best fit. Results were adjusted for the covariate effects of age, sex, and refractive error (modeled as fixed effects). The observed phenotypic variation in the twin data may be partitioned into specific variance components: additive (A) genetic, dominant (D) genetic, and common (C) and unique (E) environmental variance components (the latter also including measurement error). Heritability is defined as the proportion of total phenotypic variation due to genetic effects ((A + D)/(A + D + C + E)) (Sanfilippo et al., 2010).

For comparison between strabismus and refractive error, polygenic correlations were estimated 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 categorical strabismus data. The univariate models were extended to include the bivariate case of spherical equivalent and strabismus, to allow assessment of the extent to which any correlation between these two variables could be explained by common genes.

Results

Data were available for 1,462 twin pairs (2,924 individuals) in total: 956 pairs (389 MZ, 567 DZ) from the Australian cohort (mean age 20.8 (±12.7), range 5–90 years) and 506 pairs (226 MZ, 280 DZ) from the UK group (mean age 62.2 (±5.7), range 49–79 years). In the saturated model, liability thresholds could be equated across birth order and zygosity without any loss in fit, indicating that the prevalence of strabismus is not contingent on birth order or zygosity. No quantitative or qualitative sex differences were observed as polygenic correlations could also be equated across zygosity groups stratified by sex without significant loss of model fit. Therefore, subsequent analyses were...
performed with males and females grouped within zygosity. Table 1 shows the prevalence of strabismus for the two phenotypes examined. For eso-deviation, the polychoric correlation showed a non-significant additive genetic correlation, $r_g = 0.13 \pm 0.45$. Additionally, the additive genetic correlation could be constrained to zero without significant loss of model fit, thus indicating that eso-deviation and refraction predominantly do not share common genes. In the bivariate model, the heritability specific to eso-deviation was 59% and that common to both eso-deviation and refraction was estimated to be 64%. Our study represents the first attempt to quantify genetic and environmental variance in strabismus using a clinical data set based on twin subjects. Familial clustering of comitant strabismus has long been observed and an underlying heritable component for the disorder suspected. However, until recently no attempt has been made to quantify the genetic and environmental components of this relatively common childhood visual dysfunction. The reason for discerning these relative contributions is to improve both our understanding of its etiology and our ability to identify those at risk at an earlier stage. Given the distinctly larger genetic contribution to eso-deviation (as defined in this study), we feel this is an important group to focus limited resources on in future gene-finding efforts.

While there is a relatively large body of twin strabismus literature extending over the past 30 years (Wilmer & Backus, 2009), few studies have attempted to evaluate these data in accordance with modern twin analytical research techniques (Neale & Cardon, 1992). Based on family history and familial concordance, several studies have found that the magnitude of familial contribution to strabismus varies based on the type of deviation, with eso-deviation being most significant (Abrahamsson et al., 1999; Matsuo et al., 2002; Ziakas et al., 2002). Ziakas et al. found that a family

**Table 1**

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Number</th>
<th>Prevalence % (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eso-deviation</td>
<td>252</td>
<td>8.6 (7.7–9.7)</td>
</tr>
<tr>
<td>Exo-deviation</td>
<td>605</td>
<td>20.7 (19.3–22.2)</td>
</tr>
</tbody>
</table>

**Table 2**

Contingency Tables for MZ and DZ Twin Pairs

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Twin 1</th>
<th>Twin 2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normal</td>
<td>Phoria</td>
</tr>
<tr>
<td>Eso-deviation</td>
<td>529</td>
<td>13</td>
</tr>
<tr>
<td>Phoria</td>
<td>12</td>
<td>9</td>
</tr>
<tr>
<td>Tropia</td>
<td>16</td>
<td>3</td>
</tr>
<tr>
<td>Exo-deviation</td>
<td>403</td>
<td>55</td>
</tr>
<tr>
<td>Phoria</td>
<td>38</td>
<td>60</td>
</tr>
<tr>
<td>Tropia</td>
<td>21</td>
<td>4</td>
</tr>
</tbody>
</table>

**Table 3**

Model-Fitting Results from Heritability Analyses

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Model</th>
<th>$\Delta \chi^2$</th>
<th>$\Delta df$</th>
<th>AIC</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eso-deviation</td>
<td>ACE</td>
<td>-3,843.35</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>AE</td>
<td>-3,845.35</td>
<td>1</td>
<td>1.00</td>
<td>&lt;.001</td>
</tr>
<tr>
<td></td>
<td>CE</td>
<td>-3,837.13</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Exo-deviation</td>
<td>ACE</td>
<td>-2,384.73</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>AE</td>
<td>-2,365.1</td>
<td>1</td>
<td>&lt;.001</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>CE</td>
<td>-2,376.37</td>
<td>1</td>
<td>1.00</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>E</td>
<td>-2,234.55</td>
<td>2</td>
<td>&lt;.001</td>
<td>-</td>
</tr>
</tbody>
</table>

Note: The model in bold represents the best-fitting model for that phenotype. Abbreviations: A, additive genetic; C, common environment; E, unique environment; $\Delta \chi^2$, chi-square goodness-of-fit statistic; $\Delta df$, change in degrees of freedom between sub-model & full model; p, probability that change in $\chi^2$ is zero; AIC, Akaike information criterion.

Discussion

Our study has shown that genetic factors are important in the etiology of eso-deviation, while supporting a greater role for environmental factors in the development of exo-deviation. After adjustment for the effects of age, sex, and refractive error, the heritability of comitant eso-deviation was estimated to be 64%. Our study represents the first attempt to quantify genetic and environmental variance in strabismus using a clinical data set based on twin subjects. Familial clustering of comitant strabismus has long been observed and an underlying heritable component for the disorder suspected. However, until recently no attempt has been made to quantify the genetic and environmental components of this relatively common childhood visual dysfunction. The reason for discerning these relative contributions is to improve both our understanding of its etiology and our ability to identify those at risk at an earlier stage. Given the distinctly larger genetic contribution to eso-deviation (as defined in this study), we feel this is an important group to focus limited resources on in future gene-finding efforts.

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history of strabismus was most frequently reported in cases of hypermetropic accommodative esotropia (26.1% first-degree relatives affected) compared to 4% of first-degree relatives of index cases with exotropia (Ziakas et al., 2002). Greater familial aggregation of esotropia is supported by Podgor et al.’s large cohort study of over 40,000 families, which determined that the odds of esotropia more than doubled for a child if his or her sibling had esotropia (OR 2.6, 95% CI 1.6–4.2), in contrast to a non-significant association observed for exotropia (Podgor et al., 1996). In a recent review and meta-analysis employing SEM techniques to quantify variance components, Wilmer and Backus retrospectively analyzed Podgor’s data in conjunction with two other studies (Orlebeke & Koole, 1999; Richter, 1967) selected to satisfy inclusion criteria (Wilmer & Backus, 2009). Genetic modeling revealed a strong effect of genes on the etiology of strabismus, with heritability estimated at 92% for the combined data sets. The primary reason these studies were included in the meta-analysis was that the method of ascertainment was specified. Similarly, we used unselected twin pair volunteers from a population-based sample, thus attempting to minimize potential ascertainment bias that might produce inflated concordance estimates if twin pairs were selected on the basis of one member being affected. Wilmer and Backus also analyzed phoria data from their own unselected sample and found that genetic factors played no role across a mostly sub-clinical range of latent deviations, with a best-fitting CE model (C = 0.61, E = 0.39).

Phenotypes encompassing heterophorias were included to increase statistical power in our study. The clinical relevance of such a definition may be argued, although our reasoning reflects difficulties encountered in the characterization of endophenotypes for categorical disorders in other fields, psychiatry being a classic example. Bipolar disorder has been shown to be highly heritable mental illness (McGuffin et al., 2003), with linkage studies supporting a complex polygenic basis for the condition (Craddock & Jones, 1999). Additionally, phenotypic heterogeneity associated with diaphanous diagnostic boundaries of the bipolar disorder spectrum has been recognized as a major obstacle in the genetic dissection of the illness (Hasler et al., 2006; MacQueen et al., 2005). Consequently, it has been suggested that genetic and clinical work might consider whether all phenotypic variants of the bipolar spectrum are an expression of a continuous underlying genetic liability (Edvardsen et al., 2008). In a similar way, if strabismus and orthophoria were considered to occupy opposing ends of a clinical spectrum of anomalous ocular deviation, perhaps the same genes influence latent in addition to manifest deviation, and it is thus useful for analysis to group the conditions.

It is likely that we were underpowered to properly assess heritability for strabismus independently in this study. Even though data were available for nearly 1,500 twin pairs, this may be fewer than necessary to reject the CE model when the true model is AE. Neale et al. (1994) constructed power curves for threshold genetic models in the classic twin study. With a prevalence for esotropia of ~5% in our study and ~1,500 twin pairs, we would have 80% power to detect an additive genetic variance component of ~0.75 (Neale et al., 1994). An alternative way to consider this is that if esotropia were similarly heritable to eso-deviation (i.e., A = 0.64), we would require over 2,000 twin pairs (~2,300) to resolve an additive genetic component and reject the CE model (80% power, prevalence ~5%). Other limitations of the present work include its retrospective nature and the potential for recall bias. Additionally, and due to statistical power considerations, stratification of esotropia into subgroups (e.g., infantile, fully/partially accommodative, etc.) while ideal, would lead to such high margins of error as to be uninterpretable. Further large prospective twin studies, with accurate ascertainment of the different subtypes of strabismus, will be important for future genetic linkage or association studies to target those subtypes with the highest genetic risk. Our study nevertheless provides clear guidance on the importance of distinguishing esodeviations from exo-deviations in genetic studies. Moreover, our study also rejects the possibility that the genetic contribution to strabismus is due primarily to heritable refractive anomalies.

The mode of strabismus inheritance and the identification of associated genetic variants has thus far proven difficult. Genome-wide linkage analysis of comitant strabismus has yielded relatively few chromosomal susceptibility loci. An initial study found loci at chromosomes 7p22.1 (Parikh et al., 2003) and a more recent study (Shaaban et al., 2009), extending previous work by Fujiwara et al. (2003), identified loci at 4q28.3 and 7q3.1. This second study, conducted in 55 Japanese families with at least two cases of comitant strabismus, did not distinguish these cases based on the direction of their deviation. Our findings suggest that future linkage studies could improve statistical power for detecting genetic loci by focusing on esotropia. Additionally, it may be of value to include individuals with esophoria in such work.

The prevalence of manifest strabismus in this study (9.1%) is considerably greater than that reported (2.5–4.0%) in non-twin Caucasian populations (Kvarnstrom et al., 2001; Robaei et al., 2006), but is consistent with that reported in twins (Wilmer & Backus, 2009). A likely explanation for this difference concerns the representativeness of twin data for the general population (Sanfilippo et al., 2011). While the majority of ocular biometric traits have been shown to be similar among twins and singletons, this may not be the case for strabismus. Certainly, it is well accepted that on average, twins are more likely to be born premature and have lower birth weights than singleton pregnancies (Kiely, 1990; Min et al., 2000). Low birth weight has been found to be associated with a three- to four-fold increased risk of strabismus (Bremer et al., 1998; Robaei et al., 2006), potentially accounting for higher prevalence in our study.
In summary, our classic twin study has shown that genetic factors are significant in the etiology of comitant eso-deviation, but not of exo-deviation, and that the heritability of eso-deviation is largely independent of refractive error. These findings provide a more fine-tuned understanding of genetic factors in strabismus. In the clinic, they support an emphasis on genetic risk factors for eso-deviations and environmental risk factors for exo-deviations. In the lab, they suggest the importance of directed efforts to identify strabismus susceptibility loci specific to eso-deviations. Identifying such loci and clarifying their mechanisms of action could improve our understanding of eso-deviations and point the way toward novel treatments and preventative measures.

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Disclosure of Interests

No conflicting relationship exists for any author.

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