New Polygenic Risk Score to Predict High Myopia in Singapore Chinese Children

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Purpose: The purpose of this study was to develop an Asian polygenic risk score (PRS) to predict high myopia (HM) in Chinese children in the Singapore Cohort of Risk factors for Myopia (SCORM) cohort.

Methods: We included children followed from 6 to 11 years old until teenage years (12–18 years old). Cycloplegic autorefraction, ultrasound biometry, Illumina Human-Hap 550, or 550 Duo Beadarrays, demographics, and environmental factors data were obtained. The PRS was generated from the Consortium for Refractive Error and Myopia genomewide association study ($n = 542,934$) and the Strabismus, Amblyopia, and Refractive Error in Singapore children Study ($n = 500$). The Growing Up in Singapore Towards healthy Outcomes Cohort study ($n = 339$) was the replication cohort. The outcome was teenage HM ($\leq -5.00$ D) with predictive performance assessed using the area under the curve (AUC).

Results: Mean baseline age $\pm$ SD was $7.85 \pm 0.84$ ($n = 1004$) and 571 attended the teenage visit; 23.3% had HM. In multivariate analysis, the PRS was associated with a myopic spherical equivalent with an incremental $R^2$ of 0.041 (95% confidence interval [CI] $= 0.010$, $0.073$; $P < 0.001$). AUC for HM (0.77 [95% CI $= 0.71$–0.83]) performed better ($P = 0.02$) with the PRS compared with a model without (0.72 [95% CI $= 0.65$, 0.78]). Children at the top 25% PRS risk had a 2.34-fold-greater risk of HM (95% CI $= 1.53$, 3.55; $P < 0.001$).

Conclusions: The new Asian PRS improved the predictive performance to detect children at risk of HM.

Translational Relevance: Clinicians may use the PRS with other predictive factors to identify high risk children and guide interventions to reduce the risk of HM later in life.
Introduction

The prevalence of myopia is increasing globally and is particularly high in urbanized East Asian countries where up to 80 to 97% of young adults have myopia.\textsuperscript{1–3} High myopia (HM) is associated with potentially blinding ocular complications, including myopic macular degeneration and retinal detachment.\textsuperscript{4}

Myopia is a complex trait, arising from environmental factors, which include educational attainment and intensity, that are likely mediated by increased near work, and lack of outdoor time.\textsuperscript{5–9} Both genetic variation and gene-environment interactions are also important risk factors for myopia.\textsuperscript{10–13}

Several large-scale genomewide association studies (GWAS) in Europeans and Asians have identified hundreds of loci associated with refractive error and myopia.\textsuperscript{14–16} These loci have enabled calculation of polygenic risk scores (PRS) that provide overall risk of individual genetic susceptibility to myopia. The PRS aggregates the effect of several genetic influences using single nucleotide polymorphisms (SNPs) allowing the estimation of specific individual risk at birth.\textsuperscript{17} The PRS in adult populations of European ancestry was able to explain 7.9% and 12.1% of the interindividual variation in spherical equivalent (SE) refractive error and self-reported age of myopia onset (area under the receiver-operating characteristic curve [AUC] of 0.75 to predict moderate myopia [MM],\textsuperscript{15,16} but it remains unclear if these findings are generalizable to populations of other ancestry and demographics. The Avon Longitudinal Study of Parents and Children (ALSPAC; \(n = 1516\)) found that individuals in the top 10% of the PRS distribution (HM \(≥−5\) D) had a 6.1-fold (95% CI = 3.4–10.9) higher risk than the remaining individuals.\textsuperscript{17} In a retrospective analysis in the ALSPAC (\(n = 2048\)), the combination of parental myopia and a genetic risk score gave the best performance to predict SE in children aged 7 and 15 years (\(P < 0.001\)).\textsuperscript{18}

Although these studies demonstrate the utility of PRS to distinguish myopia risk in large-scale studies of mainly European ancestry, few studies have examined the generalizability of a PRS in East Asian children with the trans-ancestry portability of PRS remaining poor. In this study, we leverage data from Chinese Singaporean children (\(n = 1004\)) and summary statistics from the largest GWAS of myopia in Europeans\textsuperscript{16} to date, to generate a new PRS in East Asians.

We aim to develop a novel Asian PRS and use this PRS to predict HM in Singapore Chinese children in the Singapore Cohort of Risk Factors for Myopia (SCORM) cohort.

Methods

Singapore Cohort of Risk Factors for Myopia

SCORM is a prospective cohort whereby children from grades 1 to 3 were recruited from 3 Singapore schools in 1999 and 2001 (\(n = 1979\)), and has been described previously.\textsuperscript{19–21} Briefly, children were excluded if they had serious medical or eye disorders, such as congenital cataract. Institutional review board (IRB)/ethics committee approval was obtained. All human research was conducted according to the Declaration of Helsinki. Written informed consent was obtained after the nature of the study was explained.

Seven annual follow-up visits were conducted in the schools and children aged 12 to 18 years were seen at the last teenage follow-up (2007). Cycloplegic refraction was measured at every visit. After the instillation of 1 drop of 0.5% proparacaine, cycloplegia was achieved with 3 drops of 1% cyclopentolate instilled at 5-minute intervals. Subsequently, cycloplegic autorefraction was performed with a table-mounted autorefractor (model RK5; Canon, Japan) at least 30 minutes after the last eye drop. SE was calculated as sphere +1/2 cylinder. Individuals were further grouped into HM (\(≤−5.00\) D) and MM (\(−3.00\) D to \(>−5.00\) D). Axial length (AL) measurements were obtained after instillation of 1 drop of 0.5% proparacaine; contact ultrasound biometry was performed (Echoscan model US-800, probe frequency 10 mHz: Nidek Co., Ltd., Tokyo Japan).

Questionnaires were administered to ascertain information, such as number of books read per week and number of parents with myopia.\textsuperscript{20} Time outdoors (hours per week) in teenagers was recorded separately for school weekdays and school weekends using the Sydney Myopia Study questionnaire, and was defined as the sum of outdoor leisure and outdoor sporting activities.\textsuperscript{22}

Genotyping was performed using the Illumina HumanHap 550 or 550 Duo Beadarrays\textsuperscript{23} data array for Chinese children (\(n = 1004\)) in 2006, with 571 children (57.4%) continuing through the last teenage follow-up. Quality control of the genotype data was achieved by excluding SNPs with call rates <95%, excluding minor allele frequencies (MAF) <0.05 and using the Hardy-Weinberg equilibrium (HWE) test \(P < 10^{−6}\). The East Asia (EAS) reference population in the 1000 Genomes reference panel was used.
Multidimensional scaling analysis of SCORM. The genotype data from the SCORM cohort \( n = 1004 \) was combined with data from the 1000 Genomes (phase 1, version 3) comprised of 2504 individuals from 26 populations. Multidimensional scaling (MDS) analysis was performed on the combined set of 3508 individuals and 568,974 HapMap3 SNPs that were filtered on minor allele frequency < 0.05, Hardy-Weinberg equilibrium test \( P < 10^{-6} \) and genotype call rate < 0.01. Shown are the first two components from the MDS analysis.

Strabismus, Amblyopia, and Refractive Error Study

Strabismus, Amblyopia, and Refractive Error Study (STARS) is a population-based survey of Chinese families with children aged 6 to 72 months residing in the southwestern and western region of Singapore. Details of the study design and methodology have been previously described. Data on 550 children were included in this study.

Growing Up in Singapore Towards Healthy Outcomes Cohort Study

Growing Up in Singapore Towards Healthy Outcomes (GUSTO) cohort study consists of offspring of ethnic Chinese (60%), Malay and Indian pregnant
women, aged ≥18 years who attended the first trimester antenatal clinic at the National University Hospital (NUH) and the KK Women’s and Children’s Hospital (KKH) in 2009 to 2010. In this study, we only included data for Chinese children from the 9-year visit. Myopia was measured with cycloplegic refraction. Risk factors for myopia, including near work, time outdoors, mother’s education, and parental myopia, were ascertained using questionnaires. The study was approved by the SingHealth Centralized Institutional Review Board and National Health Group’s Domain Specific Review Board. Written informed consent was obtained from the parents after the nature of the study was explained.

Genotyping was performed using Infinium OmniExpressExome array. Quality control included excluding SNPs with call rates < 95%, MAF < 0.05, and HWE test $P < 10^{-6}$. GUSTO Chinese participants were compared with the EAS reference population in the 1000 Genome reference panel, where variants with different allele codings than 1000 Genome as well as SNPs with frequencies that differ more than 0.20 were removed. Following quality control, a total of 529,083 SNPs were available for analysis, and were pre-phased using SHAPEIT version 2.837 with family trio information. The SNP data was then imputed to the 1000 Genome reference panel using the Sanger Imputation Service for imputation with “PBWT, no pre-phasing” pipeline. Quality control of the imputed data retained non-monomorphic (MAF > 0), biallelic SNPs with HWE test $P > 10^{-6}$, MAF > 0.05, and INFO score > 0.50.

### Polygenic Risk Score

We used the input GWAS summary statistics from Hysi et al., a large GWAS study on refractive error conducted in individuals of European ancestry. The SNPs were selected using the MAF (equal to or higher than 0.01) in Chinese children and retained if they were directly associated, or in high linkage disequilibrium (LD) with SNPs significantly associated ($P < 5\times10^{-8}$) with refractive error in European adults. SNPs with associations that were less statistically significant in the European analysis were not considered. Presence of multiple redundant signals in our subjects of Chinese descent was subsequently verified and eventually rectified with LD pruning as implemented in the PLINK software. In all cases, an independent (i.e. not used for the analyses whose results are described here) population of ethnic Chinese ancestry, the STARS ($n = 500$), was used to calculate MAF and LD between pairs of SNP markers. SNPs were selected if they had a probability of conditional association $P < 10^{-8}$.

Because the original meta-analysis association was $z$-score based, linear logistic regression coefficients (in standardized units) were calculated using effective population sizes using the formula:

$$
\beta = \frac{\text{Zscore}}{(N_{\text{eff}} \times p \times (1 - p))}
$$

where Zscore is the reported meta-analysis Z-score, $N_{\text{eff}}$ is the effective population number, and $p$ is the MAF at the locus.

A PRS was calculated using the 655 SNPs for each individual in SCORM as the sum of risk alleles weighted by the effect sizes described above using the PRSice version 2.3.1.e (https://github.com/choishingwan/PRSice) software. To determine if the inclusion of additional SNPs in the PRS calculation improved the prediction accuracy in SCORM, we applied three recently developed methods (SBayesR, SBayesS, and SBayesRS), which have been shown the perform better than LDpred and clumping and thresholding (P + T) approaches. These three approaches are genomewide Bayesian methods that take as input GWAS summary statistics from Hysi et al. and an LD reference panel. Each method effectively shrinks SNPs effect sizes while maximizing the variance explained by binning SNPs into a mixture of normally distributed priors while accounting for LD. Shrunken sparse LD matrices generated by Lloyd-Jones et al. (downloaded from: https://cnsgenomics.com/software/gctb/#Download) were used as the LD reference panel, which were built using 1.09 million HapMap3 SNPs from a subset of 50,000 unrelated Europeans from the UKBiobank. Each method was run with the default parameters: -pi 0.95, 0.02, 0.02, and 0.01; gamma 0, 0.01, and 0.1, 1; chain-length 50,000; burn-in 20,000; out-freq 10, and using the –exclude-mhc flag. The PRS was calculated for each individual in SCORM using a total of 683,970 HapMap3 SNPs and multiplying the best guess genotypes by the effect sizes reweighted by SBayesR, SBayesS, and SBayesRS using the PLINK – score function. The PRSs were subsequently standardized to have a mean of zero and variance of one to aid in the interpretation of results.

### Statistical Analysis

Last teenage follow-up visit (children aged 12 to 18 years) SE and AL measurements (dependent variables) were tested for association with the standardized PRS by multivariable linear regression. First, we tested a model without the PRS (basic model), including age, sex, mother’s education, school, and 10 genotyping principal components. The 10 principal components are the principal components derived from the
Polygenic Risk Score to Predict High Myopia

Polygenic Risk Score

The PRS and cycloplegic auto-refraction data were available for a total of 1004 Chinese children in SCORM (mean age at baseline ± standard deviation [SD] = 7.85 ± 0.84). Of these, 571 attended the last follow-up visit with 22.8% having MM and 23.3% having HM. The mean SE at the last visit was −2.99 ± 2.57 D and mean AL at last visit was 24.75 ± 1.24 mm. The distribution of the PRS generated from the 655 SNPs in SCORM is illustrated in Figure 2.

A model without the PRS (basic model), including age, sex, mother’s education, school, and 10 genotyping principal components explained 4.0% and 10.8% of SE and AL variance, respectively (Table 1). Adding the PRS to the basic model showed an incremental R² of 0.041 (95% CI = 0.010, 0.073) for SE (Fig. 3); that is, inclusion of the PRS in the basic model showed statistically significant improvement (ANOVA P < 0.001) in the prediction (i.e. increase in adjusted R² of 4.1%) of SE. The PRS had an incremental R² of 0.022 for AL (95% CI = −0.001, 0.046; ANOVA P < 0.001; see Fig. 3). The incremental R² values for parental myopia and number of books read per week were lower (R² ≤ 1%). The inclusion of parental myopia and time outdoors to the basic model showed statistically significant improvement over the basic model, although, in both cases, the incremental R² was less than that observed with the inclusion of the PRS in the model (see Table 1). We found a small increase in the prediction accuracy of SE (i.e. incremental R² of 4.1% vs. 6.7% with SbayesRS).
Table 1. Multivariable Linear Regression Models of Polygenic Risk Score, Parental Myopia, Time Outdoors, and Association With Teenage Spherical Equivalent and Axial Length, to Determine the Degree of Improvement in Prediction Accuracy in SCORM (n = 1004)

<table>
<thead>
<tr>
<th></th>
<th>Spherical Equivalent (D)</th>
<th>Axial Length (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adjusted R², Full Model</td>
<td>Incremental R², Full Model</td>
</tr>
<tr>
<td>Basic</td>
<td>0.040</td>
<td>–</td>
</tr>
<tr>
<td>Basic + PRS</td>
<td>0.082</td>
<td>0.041</td>
</tr>
<tr>
<td>Basic + Parental myopia</td>
<td>0.051</td>
<td>0.011</td>
</tr>
<tr>
<td>Basic + Time outdoors</td>
<td>0.076</td>
<td>0.035</td>
</tr>
<tr>
<td>Basic + books read per week</td>
<td>0.040</td>
<td>0</td>
</tr>
</tbody>
</table>

aBasic model for spherical equivalent included age, sex, mother’s education, school and 10 genotyping principal components.

bBasic model for axial length included age, sex, height, mother’s education, school, and 10 genotyping principal components.

and AL (i.e. incremental R² of 2.2% vs. 3.7% with SbayesRS and SbayesS) when 683,970 HapMap3 SNPs were used in the calculation of the PRS versus our main approach that uses 655 SNPs from a GWAS of refractive error in Europeans.

The prediction accuracy of the full multivariable model was better, explaining between 11.9% and 15.7% of the variance for SE and AL, respectively (Table 2g), as compared to lower prediction for parental myopia (R² = 8.3% for SE; R² = 13.5% for AL). Time outdoors provided similar accuracy to the PRS in relation to SE (R² = 11.9%) and AL (R² = 15.7%). Number of books read per week, an indicator of near work, was not significant associated with SE or AL (P > 0.05; data not shown).

The PRS was associated with both an altered risk to time to HM (HR = 1.49; R² = 12.8%) and time to MM (HR = 1.39; R² = 12.5%) with good R² (full multivariable model; Table 3). An increase in the amount of time outdoors corresponded to a decreased risk in time to HM (HR = 0.58) and time to MM (HR = 0.79). Parental myopia had similar risk and only slightly lower R². Books read per week, an indicator of near work, was not significant associated with time to MM (P = 0.17), but was nominally significantly associated with time to HM (P = 0.03; data not shown).
Prediction accuracy (incremental $R^2$) of polygenic risk scores, time outdoors, and parental myopia in SCORM ($n = 1004$). Error bars represent 95% confidence intervals. Incremental $R^2$ values represent the increase in $R^2$ obtained by adding the PRS as predictor to a model with covariates. Multivariate models were adjusted for baseline age, sex, mother’s education, school, time outdoors, number of books read per week, and 10 genotyping principal components. AL at last visit was additionally adjusted for baseline height.

The AUC approach was used to assess the ability for the PRS to distinguish between HM ($n = 133$) and MM ($n = 129$, versus $n = 109$ no myopia controls), where an AUC of 1 and 0.5 represents a PRS with perfect and no discriminatory power, respectively. A model without the PRS (model 1: age, time outdoors, and parental myopia) showed an AUC of 0.72 (95% CI = 0.65, 0.78) for HM and 0.60 (95% CI = 0.53, 0.67) for MM. The AUC showed statistically significant performance improvement for HM but not MM when the PRS was added to the model (model 2: age, time outdoors, parental myopia, and PRS) with AUC for HM of 0.77 (95% CI = 0.71–0.83, DeLong’s test $P = 0.02$; Fig. 4) and 0.62 (95% CI = 0.55–0.69; DeLong’s test $P = 0.36$).
Table 2. Multivariable Linear Regression Models of Polygenic Risk Score and Teenage Spherical Equivalent and Axial Length in SCORM (n = 1004)

<table>
<thead>
<tr>
<th></th>
<th>Spherical Equivalent (D)</th>
<th>Axial Length (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Univariate</td>
<td>Multivariable</td>
</tr>
<tr>
<td><strong>Polygenic risk score</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>β</td>
<td>−0.21</td>
<td>−0.22</td>
</tr>
<tr>
<td>Standard error</td>
<td>0.04</td>
<td>0.04</td>
</tr>
<tr>
<td>95% CI</td>
<td>−0.30, −0.13</td>
<td>−0.30, −0.14</td>
</tr>
<tr>
<td>P</td>
<td>$6.21 \times 10^{-7}$</td>
<td>$1.49 \times 10^{-7}$</td>
</tr>
<tr>
<td>$R^2$</td>
<td>0.041</td>
<td>0.119</td>
</tr>
<tr>
<td><strong>Parental myopia</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>β</td>
<td>−0.32</td>
<td>−0.23</td>
</tr>
<tr>
<td>Standard error</td>
<td>0.09</td>
<td>0.09</td>
</tr>
<tr>
<td>95% CI</td>
<td>−0.49, −0.15</td>
<td>−0.41, −0.05</td>
</tr>
<tr>
<td>P</td>
<td>$2.10 \times 10^{-4}$</td>
<td>0.01</td>
</tr>
<tr>
<td>$R^2$</td>
<td>0.022</td>
<td>0.083</td>
</tr>
<tr>
<td><strong>Time outdoors</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>β</td>
<td>0.16</td>
<td>0.21</td>
</tr>
<tr>
<td>Standard error</td>
<td>0.04</td>
<td>0.04</td>
</tr>
<tr>
<td>95% CI</td>
<td>0.08, 0.24</td>
<td>0.13, 0.29</td>
</tr>
<tr>
<td>P</td>
<td>$1.08 \times 10^{-4}$</td>
<td>$8.77 \times 10^{-7}$</td>
</tr>
<tr>
<td>$R^2$</td>
<td>0.024</td>
<td>0.119</td>
</tr>
</tbody>
</table>

Caption: Association effect sizes ($\beta$); 95% confidence interval (95% CI); $P$ value ($P$); Adjusted $R^2$.

*a* Multivariate models were adjusted for baseline age, sex, mother’s education, school, time outdoors, number of books read per week and 10 genotyping principal components. Axial length at last visit was additionally adjusted for baseline height.

*b* Multivariate models were adjusted for baseline age, sex, mother’s education, school, polygenic risk score, number of books read per week and 10 genotyping principal components. Axial length at last visit was additionally adjusted for baseline height.

for MM. A model with only the PRS (model 3: AUC of 0.64 [95% CI = 0.57–0.71] for HM and 0.57 [95% CI = 0.49–0.64] for MM) was more predictive than a model with only parental myopia (model 4: AUC of 0.61 [95% CI = 0.55–0.67] for HM and 0.56 [95% CI = 0.50–0.63] for MM), but the best model included both the PRS and parental myopia together with age and time outdoors (model 2). We observed a small improvement in the AUC for MM (e.g. 0.65 [95% CI = 0.58–0.72] with SbayesS) when the PRS was generated using 683,970 HapMap3 SNPs versus our main approach, but no significant improvement in the AUC was found for HM (Supplementary Fig. S1).

We also analyzed the distribution of the PRS in our four myopia groups: HM (SE $\leq −5.00$ D), MM ($−5.00$ D $< SE \leq −3.00$ D), myopia ($−3.00$ D $< SE \leq −0.5$ D), and no myopia (SE $> −0.50$ D). The PRS varied significantly across the four myopia groups ($P < 0.001$), where the average PRS increased with the severity of myopia. In particular, we found the PRS to be significantly higher in individuals with HM than children with no myopia ($P < 0.001$). Further, we found that individuals in the upper percentile of the PRS distribution had increased odds of HM and MM. Individuals with PRS in the top 25% had 2.34 (95% CI = 1.53, 3.55; $P < 0.001$) and 1.76 (95% CI = 1.20, 2.59; $P < 0.001$) times higher odds of HM and MM, respectively, as compared to individuals in the remaining 75% of the PRS distribution (Fig. 5). Similarly, individuals with time outdoors in the top 25% had 0.60 (95% CI = 0.37, 0.95; $P = 0.032$) times lower odds of HM, as compared to the remaining individuals. Books read per week could not distinguish between HM and MM (data not shown). The results were similar when comparing the top and bottom 50% of the PRS and time outdoors distribution. Individuals with PRS in the top 50% had 1.50 (95% CI = 1.07, 2.09; $P = 0.02$) and 1.77 (95% CI = 1.19, 2.64; $P < 0.001$) times higher odds of MM and HM, respectively, as compared to individuals in the bottom 50%. Further, individuals with time outdoors in the top 50% had 0.68 (95% CI = 0.49, 0.95; $P = 0.02$) and 0.49 (95% CI = 0.33, 0.73; $P < 0.001$) times lower odds of MM and HM, respectively, as compared to individuals in the bottom 50%. Again, the number
### Table 3. Cox Proportional Hazard Regression Models of Time to Moderate Myopia and Time to High Myopia in SCORM (n = 1004)

<table>
<thead>
<tr>
<th></th>
<th>Univariate</th>
<th>Multivariable</th>
<th>n</th>
<th>Univariate</th>
<th>Multivariable</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time to High Myopia (−5.00 D)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Polygenic risk score</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.42</td>
<td>1.49</td>
<td>672</td>
<td>1.33</td>
<td>1.39</td>
<td>672</td>
</tr>
<tr>
<td>HR</td>
<td>1.42</td>
<td>1.49</td>
<td>672</td>
<td>1.33</td>
<td>1.39</td>
<td>672</td>
</tr>
<tr>
<td>Standard error</td>
<td>0.08</td>
<td>0.08</td>
<td></td>
<td>0.06</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>95% CI</td>
<td>1.21, 1.67</td>
<td>1.26, 1.75</td>
<td></td>
<td>1.18, 1.49</td>
<td>1.24, 1.56</td>
<td></td>
</tr>
<tr>
<td>$P$</td>
<td>$1.97 \times 10^{-5}$</td>
<td>$1.84 \times 10^{-6}$</td>
<td></td>
<td>$1.19 \times 10^{-6}$</td>
<td>$1.25 \times 10^{-8}$</td>
<td></td>
</tr>
<tr>
<td>$R^2$</td>
<td>0.026</td>
<td>0.128</td>
<td></td>
<td>0.034</td>
<td>0.125</td>
<td></td>
</tr>
<tr>
<td><strong>Parental myopia</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.80</td>
<td>1.61</td>
<td>672</td>
<td>1.52</td>
<td>1.34</td>
<td>672</td>
</tr>
<tr>
<td>HR</td>
<td>1.80</td>
<td>1.61</td>
<td>672</td>
<td>1.52</td>
<td>1.34</td>
<td>672</td>
</tr>
<tr>
<td>Standard error</td>
<td>0.19</td>
<td>0.21</td>
<td></td>
<td>0.12</td>
<td>0.13</td>
<td></td>
</tr>
<tr>
<td>95% CI</td>
<td>1.25, 2.61</td>
<td>1.07, 2.41</td>
<td></td>
<td>1.19, 1.93</td>
<td>1.03, 1.74</td>
<td></td>
</tr>
<tr>
<td>$P$</td>
<td>$1.71 \times 10^{-3}$</td>
<td>0.02</td>
<td></td>
<td>$7.60 \times 10^{-4}$</td>
<td>0.03</td>
<td></td>
</tr>
<tr>
<td>$R^2$</td>
<td>0.016</td>
<td>0.106</td>
<td></td>
<td>0.018</td>
<td>0.089</td>
<td></td>
</tr>
</tbody>
</table>

| **Time outdoors<b>** | 0.71 | 0.58 | 672 | 0.89 | 0.79 | 672 |

<table>
<thead>
<tr>
<th></th>
<th>Univariate</th>
<th>Multivariable</th>
<th>n</th>
<th>Univariate</th>
<th>Multivariable</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HR</strong></td>
<td>0.71</td>
<td>0.58</td>
<td>672</td>
<td>0.89</td>
<td>0.79</td>
<td>672</td>
</tr>
<tr>
<td><strong>Standard error</strong></td>
<td>0.09</td>
<td>0.10</td>
<td></td>
<td>0.06</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td><strong>95% CI</strong></td>
<td>0.59, 0.85</td>
<td>0.47, 0.70</td>
<td></td>
<td>0.79, 1.00</td>
<td>0.70, 0.90</td>
<td></td>
</tr>
<tr>
<td><strong>$P$</strong></td>
<td>$1.41 \times 10^{-4}$</td>
<td>$8.30 \times 10^{-8}$</td>
<td></td>
<td>0.04</td>
<td>$2.80 \times 10^{-4}$</td>
<td></td>
</tr>
<tr>
<td><strong>$R^2$</strong></td>
<td>0.023</td>
<td>0.128</td>
<td></td>
<td>0.006</td>
<td>0.125</td>
<td></td>
</tr>
</tbody>
</table>

**Caption:** Hazard ratios (HR); 95% Confidence interval (95% CI); $P$ value ($P$); Adjusted $R^2$.

<sup>a</sup>Multivariate models were adjusted for baseline age, sex, mother’s education, school, time outdoors, number of books read per week and 10 genotyping principal components.

<sup>b</sup>Multivariate models were adjusted for baseline age, sex, mother’s education, school, polygenic risk score, number of books read per week and 10 genotyping principal components.

of books read could not distinguish between HM and MM.

**Growing Up in Singapore Towards Healthy Outcomes Cohort Study**

The association between PRS and SE and AL were tested for replication in an independent dataset from the GUSTO cohort. Genetic and cycloplegic auto-refraction measurement information was available in a total of 339 Chinese children from GUSTO (9-year-olds). Among these children, 10.9% and 2.4% had MM and HM, respectively.

The prediction accuracy of the full multivariable model in the GUSTO cohort was similar to SCORM (Fig. 6), indicating good replicability across cohorts of similar Chinese ancestry. For example, in the full multivariable model (which included sex, time outdoors, and near-work), the association between the PRS and SE was of similar effect size in SCORM ($\beta = -0.22; R^2 = 11.9\%$) and in GUSTO ($\beta = -0.24; R^2 = 4.6\%; P < 0.001$), albeit with a much lower variation explained in GUSTO for SE at 9-year-olds. Time outdoors was associated with teenage SE in SCORM ($\beta = 0.21; R^2 = 11.9\%$) but was not significantly associated with SE at 9-year-olds in GUSTO ($P = 0.18$). The association between the PRS and AL in the full multivariable model (which included sex, height, time outdoors, and near-work) was of similar effect size in SCORM ($\beta = 0.17; R^2 = 15.7\%$) and in GUSTO ($\beta = 0.19; R^2 = 16.7\%; P < 0.001$). Adding the PRS to the basic model (which included sex as a covariate for SE, and sex and height as covariates for AL) in GUSTO showed an incremental $R^2$ of 4.9% for SE (ANOVA $P < 0.001$) and 3.3% for AL (ANOVA $P < 0.001$); that is, inclusion of the PRS to the basic model showed statistically significant improvement in GUSTO.

**Discussion**

In our study, we found that our newly developed PRS based on the latest GWAS results was associated with HM. The model (model 2), including age,
Figure 4. Receiver Operating Characteristic (ROC) curve for detecting high myopia (≤ −5.00 D) and moderate myopia (≤ −3.00 D) versus no myopia controls using the polygenic risk score in SCORM. ROC curve for high myopia (≤ −5.00 D, n = 133) and moderate myopia (≤ −3.00 D, n = 129) versus no myopia controls (n = 109) with PRS, age, time outdoors, and parental myopia as predictors. The area under the curve (AUC) and 95% confidence interval corresponds to the PRS, age, time outdoors, and parental myopia model. The black line represents an AUC of 0.5.

Figure 5. Odds ratios for teenage high myopia (≤ −5.00 D) and moderate myopia (≤ −3.00 D) for children classified as high risk using the polygenic risk score and time outdoors in SCORM (n = 1004). Odds ratios were calculated with a 2 × 2 table of myopia level (high myopia, ≤ −5.00 D and moderate myopia, ≤ −3.00 D) versus high-risk group where error bars represent 95% confidence intervals.

Polygenic risk score | Time outdoors
---|---
Odd Ratio | 3.0
SE(≤−5.00D) | SE(≤−3.00D)

Figure 6. Association effects size (β) from multivariate linear regression of spherical equivalent (SE) and axial length (AL) on polygenic risk score (PRS) in SCORM (n = 1004) and GUSTO (n = 339).

Our full multivariate model showed that the PRS explained 11.9% of the variance for SE, whereas parental myopia only explained 8.3%. We also found that the PRS explained about 13% of the phenotypic variance of time to HM. The PRS alone was able to incrementally explain 2.2% to 4.1% of the phenotypic variance for teenage AL and SE, respectively. These results were lower than previous studies in subjects of European ancestry that reported a genetic risk score explaining 7.9% \(^{15}\) (Rotterdam Study I–III; n = 10,792) and 12.1% \(^{16}\) of the interindividual variation in SE and self-reported age of myopia onset in a group of adults (AUC of 0.75 to predict MM). Thus, we hypothesize that the performance of the PRS may be better, as

time outdoors, parental myopia, and the PRS, had the best predictive ability of HM. Clinicians may use the PRS with other predictive factors to identify high risk children and guide interventions to reduce the risk of HM later in life.
the SNPs selected on the basis of effects in adults are likely to perform better in predicting SE in adults. In a previous study from the ALSPAC mother’s cohort \((n = 1516 \text{ aged 24 to 51 years})\), the best PRS result was obtained for SE combined with age of onset of spectacle wear.\(^{17}\) The author’s compared a genomewide approach using LDpred to a clumping and thresholding \((P + T)\) approach applied to 1.1 million HapMap3 SNPs using a variety of \(P\) value thresholds to maximize the refractive error variance explained by the PRS. They found that a PRS generated from the top GWAS signals \((R^2 = 6.3\% \text{ with } 7372 \text{ top SNPs with } P < 0.01 \text{ identified by the } P + T \text{ approach})\) was less accurate than one generated from the 1.1 million SNPs \((R^2 = 11.2\% \text{ by the } \text{LDpred approach})\). To our knowledge, this is one of the highest prediction \(R^2\) for refractive error to date.

Similar to our results, the ALSPAC birth cohort study \((n = 2048; \text{ age } 7–15 \text{ years})\) reported that the highest \(R^2\) value achieved using a genetic risk score and parental myopia was less than \(\sim 7\%\) for SE.\(^{18}\) The ALSPAC birth cohort study results were the first to demonstrate an independent effect of parental myopia and genetic risk on myopia in children. In our study, we found that the PRS, time outdoors, and parental myopia each have an independent effect on SE. This indicates that parental myopia may be capturing the risk of myopia from common environmental factors, and, in particular, is replicating the result from the ALSPAC birth cohort study\(^{18}\) in Singaporean Chinese children from SCORM. Thus, combining genetic testing with parental myopia may provide more accurate prediction of later HM and guide treatment decisions. For example, younger children with both parents having myopia may be targeted for more aggressive treatment to avoid progression to HM during their teenage years.

Our results showed acceptable accuracy of prediction of the PRS combined with parental myopia to predict HM, supporting the role of genetic factors in the progression of myopia. The transferability of the PRS across ancestry divergent populations has been shown to be influenced by differences in allele frequencies of casual variants, the magnitude and direction of effect sizes, and variation in the patterns of linkage disequilibrium between the training and target populations.\(^{34}\) Population-specific causal variants and gene-environment interactions may also contribute to differences in prediction performance between populations. Further, empirical and simulation studies have shown that the prediction accuracy of a genetic predictor decreases with greater genetic distance between the training and target samples.\(^{35}\) Therefore, there is some expectation that the prediction \(R^2\) generated in the present study will be lower than those observed in studies where the training and target samples are both of European ancestry; that is, this study does not expect to reach 11% to 12% prediction accuracy as demonstrated in other European studies. However, future large-scale GWAS studies in East Asians will likely close this gap.

We also found that the PRS alone (model 3) performed better in the prediction of HM than the conventional measurement of parental myopia (model 4). However, combining both the PRS and parental myopia with the other predictors (model 2) showed better predictive performance than using parental myopia alone to determine the risk of developing HM. Previously, genetic factors have mostly been measured by the number of myopic parents.\(^{10–12}\) Nevertheless, in our study, including both the PRS and number of myopic parents improved the AUC. The PRS explained higher phenotypic variance of the SE at adolescence, showing that parental myopia may be a less effective proxy for genetic factors then the PRS. For example, myopic parents may raise their children in a myopia-induced environment, with parental myopia being an imperfect proxy for genetic factors, reflecting gene-environmental interactions, such as with time outdoors, whereas the PRS allows a more accurate measurement of pure genetic risk.

In the current study, the best predictive performance was found when the PRS was combined with age, time outdoors, and parental myopia \((\text{AUC} = 0.77)\). Although there are differences in the HM rates in Asians compared with Europeans, the predictive performance of the AUC was considered acceptable, although not excellent. It is also important to note that the PRS alone had an AUC of 0.64 for HM and 0.57 for MM, showing that accuracy of prediction for the clinical setting must improve further. Thus, further studies using large-scale GWAS studies of myopia in East Asians are necessary to ascertain the predictive performance of the PRS before translation to clinical practice.

As our PRS may predict children who develop HM later, future gene testing may be implemented using the PRS with early identification of myopic children at risk of developing HM before the irreversible elongation of the eye sets in. These children may benefit from personalized counselling as well as treatment regimens to slow the progression of myopia to HM. Genetic predictions may be combined with information on child’s age, parental myopia, and other lifestyle predictors, such as time outdoors, to prevent the progression of myopia to HM and later visually disabling complications. Clinically significant results should be communicated to families to aid in their decision making and
we also conducted time to event analyses that allowed the analyses of children who may have been lost to follow up, considering the time they remained in the study (Cox models).

The approach used to generate the PRS in our study included 655 SNPs in the PRS calculation. This approach may be limited due to the exclusion of potentially informative SNPs. Our results were consistent with those observed by Mojarrad et al. where a PRS generated from the top GWAS signals ($R^2 = 6.3\%$ with 7372 top SNPs with $P < 0.01$ identified by the $P + T$ approach) was less accurate than one generated from the 1.1 million SNPs ($R^2 = 11.2\%$ by the LDpred approach). Nevertheless, the improvement observed in the incremental $R^2$ with the genomewide Bayesian methods in our study was relatively small. This was confirmed in the AUC/ROC analysis, which did not show a notable improvement for classification of individuals with MM or HM.

Another limitation was that SNPs for the PRS in the current study were obtained from a GWAS conducted in Europeans. The genetic structure between European and Asian subjects has different haplotypic structure arising from LD and different frequency of genetic risk factors between these populations. Empirical and theoretical studies have shown that there is an expected decrease in performance (i.e. lower incremental $R^2$) of the PRS when transferred across ancestries. The difference in genetic structure of the European and Asian population may have contributed to lower predictive ability of the PRS in our Asian cohort (SCORM and GUSTO). Further, there have also been an increasing number of mixed marriages, so the PRS based only on the highly significant SNPs may not perform as well in other ethnic groups or admixed ethnicity groups. There is therefore a strong need for future large-scale GWAS studies of myopia in East Asians and other non-European ancestries.

**Conclusion**

We found that adding the PRS to other clinical information, such as child’s age, time outdoors, and parental myopia, improves the prediction of HM risk ($AUC = 0.77$) in teenagers. The PRS alone performed well in the prediction of HM, with children in the highest PRS risk percentile having increased odds of developing HM. Our findings suggest the potential clinical value of utilizing information on this new Asian PRS together with parental myopia to improve the predictive performance to detect children at risk of HM. Clinicians may use the PRS with other predictive factors to identify high risk children and guide...
interventions to reduce the risk of HM later in life. Further predictive studies with genetic loci from GWAS studies of myopia in East Asians, larger sample sizes, and detailed analyses of ocular and lifestyle factors may be important to increase the predictive performance to a level acceptable for use in clinical application.

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* CL and IK considered joint first authors.

References


