Genetic variants linked to education predict longevity

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Educational attainment is associated with many health outcomes, including longevity. It is also known to be substantially heritable. Here, we used data from three large genetic epidemiology cohort studies (Generation Scotland, n = ~17,000; UK Biobank, n = ~115,000; and the Estonian Biobank, n = ~6,000) to test whether education-linked genetic variants can predict lifespan length. We did so by using cohort members’ polygenic profile score for education to predict their parents’ longevity. Across the three cohorts, meta-analysis showed that a 1 SD higher polygenic education score was associated with ~2.7% lower mortality risk for both mothers (total n_deaths = 79,702) and ~2.4% lower risk for fathers (total n_deaths = 97,630). On average, the parents of offspring in the upper third of the polygenic score distribution lived 0.55 y longer compared with those of offspring in the lower third. Overall, these results indicate that the genetic contributions to educational attainment are useful in the prediction of human longevity.

Significance

Individuals with more education tend to live longer. Genetic variants have been discovered that predict educational attainment. We tested whether a “polygenic score” based on these genetic variants could make predictions about people’s lifespan. We used data from three cohort studies (including >130,000 participants) to examine the link between offspring polygenic score for education and parental longevity. Across the studies, we found that participants with more education-linked genetic variants had longer-living parents; compared with those with the lowest genetic education scores, those with the highest scores had parents who lived on average 6 months longer. This finding suggests the hypothesis that part of the ultimate explanation for the extended longevity of better-educated people is an underlying, quantifiable, genetic propensity.


Supporting Information

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1605334113/-/DCSupplemental.

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general cognitive ability (13, 14, 26), itself a well-replicated phenotypic (27) and genetic (28) correlate of longevity.

The Current Study

In this study, we tested whether the genetic variants associated with educational attainment are associated with longevity. We thus assessed the extent to which the genetic contributions to educational outcomes, which are preexisting and non-social, are related to a key health outcome. To do so, we used the established technique of testing for associations between genotyped subjects and their phenotyped relatives (in this case, the lifespan of parents) (29).

Offspring genetic variants, such as the Alzheimer’s-linked APOE e4 allele, have also been linked to parental longevity in candidate gene studies (30) and more recently in a GWAS (31; see ref. 32 for a similar analysis of epigenetic markers). Moreover, higher genetic risk for conditions, such as cardiovascular disease, diabetes, and Alzheimer’s disease, has been related to earlier parental mortality (33). Because the expected allelic effect of one allele in parents is 0.5 alleles in offspring (31), precise predictions can be made of the effect of alleles and polygenic scores on traits in the offspring themselves.

Here, we used summary data from an independent GWAS of educational attainment (15) to create polygenic profile scores (34). These scores quantify the extent to which each participant carried the genetic variants known to be associated with higher educational attainment (in the GWAS, education was measured as the number of years of education). We then linked these polygenic profile scores to data on the participants’ parents’ age at death. Our hypothesis was that offspring with polygenic profiles for higher educational attainment would have longer-living parents. We did not make a specific prediction about whether any effect would be stronger in fathers or mothers. We performed the analysis in three large, independent cohorts to test the replicability of the result, and meta-analytically combined the three estimates. The cohorts were Generation Scotland (35, 36) (n = 17,542), UK Biobank (37) (n = 116,425), and the Estonian Biobank (38) (n = 7,950).

As a sensitivity analysis, we tested whether our results still held when taking into account parental fertility: that is, when including as a numerical covariate the number of siblings that each participant reported. This was because of a possible biasing effect whereby parents with higher numbers of offspring, and thus linearly proportionate greater likelihood of the parental phenotype being included in the study, might have different genetic propensities for educational attainment. Finally, we compared the predictive value of the educational polygenic profile score for parental mortality with the predictive values for a number of other polygenic profile scores indexing phenotypes that are known to relate to mortality risk.

Results

A summary of the parental data, including number of deaths, for each of the three cohorts is presented in Table 1, and the cohorts are described more fully in Materials and Methods.

Polygenic Profile Score Analysis. The polygenic scores for educational attainment were built using the previous GWAS results (15) and applied to the participants in Generation Scotland, UK Biobank, and the Estonian Biobank. Fig. 1 provides descriptive data for each sample, showing the age at mortality of each parent depending on each decile of the education polygenic risk score; in general, higher polygenic scores were associated with older age at death. However, this illustration only includes parents who had died. To take into account all of the data, we calculated the associations between offspring polygenic scores and parental longevity using Cox proportional hazard models (Table 2). For mothers, the HRs were not significantly different from zero in the smaller samples, but were highly significant in UK Biobank. For fathers, the results were significant in all three samples. In all cases, the point estimate was in the hypothesized direction: higher polygenic profile score was associated with lower parental mortality risk.

We combined the scores across the three cohorts, separately for mothers and for fathers, using a fixed-effects meta-analysis. The meta-analytic results showed that a 1 SD higher polygenic profile score for education was associated with an −2.5% lower mortality risk in mothers [HR = 0.976, 95% CI = (0.968, 0.983), P = 8.21 × 10−11] and fathers [HR = 0.973, 95% CI = (0.967, 0.979), P < 1.73 × 10−15]. A forest plot is shown in Fig. 2. Splitting the education polygenic profile score into tertiles, we calculated that, in the UK Biobank sample, mothers of children in the highest tertile lived on average 40.8 y beyond age 40, compared with 40.2 y in the lowest tertile (a difference of 0.6 y). The corresponding values for fathers were 35.1 y beyond age 40 in the highest tertile and 34.6 y in the lowest (a difference of 0.5 y).

We next ran the sensitivity analysis including number of siblings as a covariate in the model predicting parental mortality from offspring genotype. Again, this analysis was run in UK Biobank

<table>
<thead>
<tr>
<th>Table 1. Descriptive statistics for parents across the three samples</th>
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<tbody>
<tr>
<td><strong>Parent</strong></td>
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<tr>
<td></td>
</tr>
<tr>
<td><strong>Mother</strong></td>
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<tr>
<td></td>
</tr>
<tr>
<td><strong>Father</strong></td>
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</table>
alone. The inclusion of this covariate made little difference to the UK Biobank results reported above: the association between educational polygenic profile score and mother’s longevity was HR per 1 SD higher polygenic score = 0.978, 95% CI = (0.970, 0.985), \( P = 3.29 \times 10^{-10} \Delta \text{HR from original model} = 0.002 \); the association with father’s longevity was HR = 0.977, 95% CI = (0.970, 0.984), \( P = 9.86 \times 10^{-12} \Delta \text{HR} = 0.002 \).

Other Polygenic Scores. To provide context for the effect size of the education polygenic profile score’s association with parental mortality, we tested associations of parental longevity with a series of other offspring polygenic profile scores. We ran these analyses only in UK Biobank, using the same methods as those for the educational polygenic profile score analyses. The polygenic profile scores were for five known phenotypic predictors of mortality risk: height, body mass index (BMI), cardiovascular disease, major depressive disorder (MDD), and smoking (39–43).

The hazard ratios for predicting mortality for either parent from offspring polygenic profiles for height and for MDD were both near to 1 and nonsignificant (\( P > 0.28 \)). For mothers, there was no significant relation with offspring smoking genetic risk (\( P = 0.07 \)). However, scores for BMI, cardiovascular disease and, for fathers, smoking, made significant predictions of mortality risk (\( P < 5.85 \times 10^{-06} \)). The effect sizes for each of these genetic predictors were similar to that for education (approximately a 2% difference in mortality risk per 1 SD difference in the score) (Table S1). Thus, the score composed of genetic variants weighted toward their relation to educational attainment made similar-sized predictions of longevity risk to genetic scores weighted toward alleles linked to other well-established risk factors for mortality. Note that a number of other polygenic associations with mortality were addressed in the UK Biobank sample in a previous study, using somewhat different methods (33).

**Discussion**

This study found that offspring polygenic profiles for education were robustly associated with parental longevity: those with more genetic variants related to better educational qualifications had longer-living parents. We tested the study’s principal hypothesis across three large cohorts, totaling over 130,000 participants. The associations were of broadly similar effect size in all three cohorts. Meta-analytically, there was a substantial and strongly significant overall prediction, which was similar for males and for females: individuals with 1 SD higher polygenic profile score for a college degree had parents who were at \( \sim 2.5\% \) lower risk of mortality. Put another way, parents with offspring in the upper third of the polygenic score distribution lived an average of 0.55 y longer than those in the lower third. The results—which were comparable to the effect sizes from other known predictors of mortality, such as cardiovascular disease and smoking, and which were bolstered by the finding of a moderate-sized genetic correlation between the two variables—suggest the hypothesis that the ultimate reason education predicts mortality is, in part, because of an underlying, quantifiable, genetic propensity.

Why do genetic variants related to educational attainment predict parental mortality? There are a number of possible mechanisms—both genetically and environmentally mediated—that might explain the result. First, these genetic variants might improve cognitive or personality phenotypes, such as intelligence, motivation, and conscientiousness, thus improving educational attainment; the higher quality of life and environment afforded by a better education might then improve health and reduce mortality risk. The effects of the genetic score might manifest directly on the parents’ behavior (to the extent that they are shared between parents and offspring), or have indirect effects via greater offspring resources and ability to care for aging parents (44). Our analysis could not test between these direct and indirect possibilities.

**Table 2. Results from Cox proportional hazard models predicting parental mortality risk from offspring education polygenic profile score**

<table>
<thead>
<tr>
<th>Parent</th>
<th>Cohort</th>
<th>n Offspring</th>
<th>n Parental deaths</th>
<th>HR (95% CI)</th>
<th>SE</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mother</td>
<td>Generation Scotland</td>
<td>16,670</td>
<td>6,330</td>
<td>0.954 (0.907, 1.001)</td>
<td>0.024</td>
<td>0.056</td>
</tr>
<tr>
<td></td>
<td>UK Biobank</td>
<td>115,323</td>
<td>69,990</td>
<td>0.976 (0.968, 0.984)</td>
<td>0.004</td>
<td>1.52 \times 10^{-10}</td>
</tr>
<tr>
<td></td>
<td>Estonian Biobank</td>
<td>5,929</td>
<td>2,682</td>
<td>0.979 (0.940, 1.018)</td>
<td>0.020</td>
<td>0.280</td>
</tr>
<tr>
<td></td>
<td>Meta-analysis</td>
<td>137,922</td>
<td>79,702</td>
<td>0.976 (0.968, 0.983)</td>
<td>0.004</td>
<td>8.21 \times 10^{-10}</td>
</tr>
<tr>
<td>Father</td>
<td>Generation Scotland</td>
<td>16,390</td>
<td>8,467</td>
<td>0.932 (0.891, 0.973)</td>
<td>0.021</td>
<td>0.0007</td>
</tr>
<tr>
<td></td>
<td>UK Biobank</td>
<td>111,334</td>
<td>85,419</td>
<td>0.975 (0.969, 0.981)</td>
<td>0.003</td>
<td>2.05 \times 10^{-13}</td>
</tr>
<tr>
<td></td>
<td>Estonian Biobank</td>
<td>6,097</td>
<td>3,744</td>
<td>0.942 (0.909, 0.975)</td>
<td>0.017</td>
<td>0.0003</td>
</tr>
<tr>
<td></td>
<td>Meta-analysis</td>
<td>133,821</td>
<td>97,630</td>
<td>0.973 (0.967, 0.979)</td>
<td>0.004</td>
<td>1.73 \times 10^{-18}</td>
</tr>
</tbody>
</table>

Hazard ratios are expressed per SD of polygenic profile score. Meta-analytic rows are in bold. All models adjusted for offspring sex, genotyping array, and SNP principal components as described in Materials and Methods.
Second, the genetic variants related to education might also affect other variables that themselves lower mortality risk. This could occur in the absence of a causal pathway involving education itself. For example, individuals with long-term illnesses are at greater risk of educational failure (23), and also tend to perform more poorly on tests of cognitive ability (45); such health complaints, which might partly be genetic in origin, could also increase mortality risk. Third, unlike the above mechanisms that all posit a mediated mechanism (that is, genetic variants related to educational attainment have effects on some factor that leads to additional longevity), biological pleiotropy (46) might play a role: in this view, the genetic variants reflect a general “system integrity” (47), whereby genotypes related to better physical health (and thus lower mortality risk) are also related to better neural health (and thus better educational performance).

Testing the relative contributions of each of the three possible mechanisms described above—which are not mutually exclusive—will require a finer-grained analysis of the complex pathways that link the education-associated genetic variants to longevity, via mediating traits, conditions, and behaviors. In this study, we compared the size of the association of longevity with the educational polygenic score with that for polygenic scores for height, BMI, cardiovascular disease, major depressive disorder, and smoking. Future analyses could get closer to the mechanism by assessing the degree of overlap and generality among genetic predictors of longevity. Because GWAS studies provide a deeper knowledge of the specific, causal genetic variants that are linked to education and to longevity (15–17, 31), we will be able to address their biological and social mechanisms in greater detail, improving our understanding of precisely why scores for, for example, height make no prediction of longevity but those for education do. In any case, regardless of the underlying mechanisms, the polygenic profile score for education showed predictive value. As GWAS sample sizes for education and related variables increase, and more genetic variants are uncovered (48), we would expect steadily to obtain improved genetic predictions of longevity.

The longevity prediction made by the polygenic profile for educational attainment was substantially smaller than that for the phenotype of educational attainment. This is to be expected for two reasons. First, polygenic profiles only explain small amounts of variance in their respective phenotypes because of the power of the original GWAS studies to detect SNPs with significant associations (as noted, we expect this to improve with larger, future GWAS studies), and do not include nonadditive genetic variants that may also be important in explaining heritability. Second, educational attainment is far from completely heritable (12), being influenced by social and environmental factors that may also be predictive of parental mortality. However, the finding that the longevity-predicting power of a genetic profile for education (a variable often thought of as “social”) compares favorably in effect size to polygenic profiles for cardiovascular disease and BMI (variables that are medical in nature) supports the importance of educational attainment as a general indicator of health and social status.

Our method, using parental longevity as an outcome variable predicted by offspring genotype, allowed considerably higher power compared with studies of genotype and mortality in the same individuals; such data are more difficult to collect because they require follow-up of genotyped individuals until their own death. This method, combined with the large sample sizes, our replication and meta-analysis, and the inclusion of parents who were still alive as censored data points in the proportional hazard models, substantially improved our results’ robustness. Our effect size estimates for the main analysis were similar in samples from the United Kingdom and from Estonia, indicating that the education-related genetic variants make predictions across different cultures (although further replication in other groups will be necessary). Finally, our results appeared robust to parental fertility: they were only slightly altered after adjustment for number of siblings.

Although the effects found here were broadly consistent across cultures, the samples were not fully representative of the populations from which they were drawn. All samples were restricted to individuals of White European ancestry. Whereas this reduces bias due to population stratification within each sample, it does make the results less generalizable, and samples of participants with different genetic backgrounds may show different results. In addition, self-selection effects (49) mean that those with more education, higher intelligence, higher socioeconomic status, higher conscientiousness, and closer proximity to testing centers were probably more likely to participate. The concomitant restriction of range potentially led to downward bias in our effect sizes. A more subtle consequence of self-selection is that many of the above characteristics might make these self-selected individuals more likely to benefit, in terms of health or other life outcomes such as longevity, from higher educational attainment (that is, their genetic propensities for education may interact with other traits). However, in such a conceptualization, the educational variants are still the ultimate explanation for some of the variance in longevity.

No measures of parental educational attainment were available in our samples, precluding an analysis testing whether there were any incremental associations of the polygenic score beyond phenotypic education, or whether any effects of the genetic score were entirely mediated by educational attainment. The advantages of our parental-proxy method are noted above, but we may have underestimated the effects: we would expect exactly double the effect size for polygenic prediction of an individual’s own longevity from their genetic profile (31). Finally, although we adjusted for each study participant’s number of siblings to control for fertility differences, by definition all of the individuals whose ages at mortality were analyzed in the present study (i.e., the parents) had children. It remains possible that the associations studied here would be different for individuals with no children, who may have also had systematically different polygenic profiles for education.
Materials and Methods

Educational Attainment "Discovery" GWAS. The educational attainment polygenic scores were built using summary data from the largest GWAS meta-analysis of educational attainment to date (15). To reduce the possibility of sample overlap or cryptic relatedness affecting the polygenic scores, the GWAS data were reanalyzed after excluding all United Kingdom-based cohorts for predictions into the independent United Kingdom cohorts. Similarly, the Estonian Biobank data were excluded from a second reanalysis of the GWAS data. Data from 23andMe, used in the original meta-analysis, were not available for the calculation of polygenic scores.

Independent Sample 1: Generation Scotland: The Scottish Family Health Study. Participants. Generation Scotland: the Scottish Family Health Study (35, 36) is a cohort study of participants recruited in the Glasgow, Tayside, Ayrshire, Arran, and northeast areas of Scotland. Initially, 7,953 probands aged 35–65 y were recruited either through their general medical practitioner (95% of probands) or via direct publicity and word-of-mouth. Their family members were also invited to take part, resulting in a final sample of 24,084 participants with an age range of 18–100 y.

All components of Generation Scotland received ethical approval from the National Health Service Tayside Committee on Medical Research Ethics (Research Ethics Committee Reference no. 05/S1401/89). Generation Scotland has been granted Research Tissue Bank status by the Tayside Committee on Medical Research Ethics (Research Ethics Committee Reference no. 10/S1402/ 20), providing generic ethical approval for a wide range of uses within medical research.

Genotyping. Generation Scotland participants were genotyped with either the HumanOmniExpressExomeV1-2_A or HumanOmniExpressExome-Exome-V1. A Quality control was carried out in PLINK v1.9b2c (50, 51). SNPs were removed if they had a missingness rate >2% or a Hardy–Weinberg Equilibrium test at \( P < 10^{-6} \), leaving a total of 561,125 autosomal SNPs for analysis. Duplicate samples were removed. Individuals were removed based on gender mismatch and missingness (>2% of genotypes missing). The subsequent dataset was combined with the 1,092 individuals of the 1000 Genomes population (52) before principal components were calculated using GCTA (53). Outliers, defined by being more than 6 SDs away from the mean of the first two principal components, were removed (54). This left a sample of 20,032 participants. Individuals who appeared in both the UK Biobank and Generation Scotland studies \((n = 174)\) were excluded from the latter study. After merging with the available covariate data, 17,542 participants had age at death or censoring information in at least one parent.

Independent Sample 2: UK Biobank. Participants. Data stem from the baseline wave of the UK Biobank Study (37) (www.ukbiobank.ac.uk). Analyses were performed under data application 8304 and 10279. The UK Biobank sample was substantially larger than our other two studies: it contains around 500,000 community dwelling men and women in the United Kingdom, who were recruited between 2006 and 2010. Here, we used data from 116,425 participants (aged 40–73 y) who had genetic data and at least one parent’s longevity data available for analyses. Ethical approval for UK Biobank was granted by the Research Ethics Committee (11/NW/0382).

Genotyping. Details on the UK Biobank genotyping procedure and quality-control steps that were included for the current analyses have been reported previously (31). Briefly, of the 152,729 participants with genetic data available as of August 2015, 116,425 were retained after exclusions based on SNP missingness, relatedness, gender mismatch, non-British ancestry, and previously reported quality control failure for the UK BiLEVE study.

Independent Sample 3: Estonian Biobank. Participants. The Estonian Biobank (38) is the population-based biobank of the Estonian Genome Centre of the University of Tartu (EGCUT). For this study, 51,380 volunteer participants (aged 18–103 y) were recruited between 2002 and 2011. The cohort included ~5% of the adult population from all counties of Estonia. At recruitment, the participants completed an extensive questionnaire on health, lifestyle, and genealogy and provided a blood sample. Approval for the Estonian Biobank was given by the Research Ethics Committee of the University of Tartu. All participants signed a broad informed consent form at recruitment.

Genotyping. In total, DNA samples from >16,000 participants have been genotyped with various genome-wide arrays. In 2011, the subset of individuals selected to be genotyped with the Illumina OmniExpress chip, intentionally included 1,200 individuals who had died by that time, as well as 500 women and 250 men who were 80 y old or older at that time. The rest of this genotyped sample (in total 7,950 subjects after removing close relatives) consists of random population controls.

Parental Longevity Phenotype. In all three independent cohorts, parental longevity was assessed for both mothers and fathers. To account for premature deaths as a result of external causes, such as accidents, and in particular, the Second World War (55), we excluded individuals who died prematurely (<40 y). Parents who were alive at the baseline wave of the respective studies were treated as censored observations. Age at censoring was calculated as the cohort’s baseline year of assessment minus the parent’s year of birth. Parents whose age at censor was <40 y were excluded. In Generation Scotland, a small number of outliers \((n = 26)\) with an age-at-death/censor >100 y were removed.

Statistical Analyses. LD score regression (genetic correlation). To assess the genetic correlation between the two primary phenotypes of interest, we used LD score regression (56), which allows genetic correlations to be calculated using GWAS summary data alone (without raw genotype or phenotype data). It does not matter for LD score regression whether there is sample overlap between the studies. We calculated the genetic correlation \(\rho_{g}\) using the summary data for educational attainment from the GWAS that also served as the basis for our polygenic profile score (see below) and using the summary data for parental longevity (specifically, the Martingale residuals from Cox proportional hazards models of parental lifespan) from a recent GWAS in the UK Biobank sample (31). LD score regression was used with all its default settings.

Polygenic scoring. The results from the educational attainment GWAS analyses were carried forward into polygenic score models in the three independent cohorts using the PLRscie software (57). We built polygenic profile scores based on the genotyped SNP data in Generation Scotland \((n_{\text{SNP}} = 561,125)\), UK Biobank \((n_{\text{SNP}} = 672,491)\), and EGCUT \((n_{\text{SNP}} = 628,325)\). The optimal threshold determined from the previous analysis (15), specifically \(P < 1.00\) (that is, inclusion of all SNPs) was used for all analyses in all samples.

We tested whether the polygenic profile score was related to the participant’s (that is, the offspring’s) own educational duration. To do this, we included the score in a linear regression analysis predicting the offspring’s educational attainment along with covariates of age, sex, and 15 SNP-based principal components to account for population stratification. The polygenic profile score was significantly related to years of education in Generation Scotland \((n = 17,814\); standardized \( \beta = 0.132\), \( SE = 0.008\), \( P = 2.20 \times 10^{-13}\)\), UK Biobank \((n = 119,167\); standardized \( \beta = 0.106\), \( SE = 0.003\), \( P = 2.20 \times 10^{-16}\)\), and EGCUT \((n = 7,959\); standardized \( \beta = 0.118\), \( SE = 0.011\), \( P = 1.77 \times 10^{-26}\)). Note that, for Generation Scotland, a kinship matrix was fitted to account for the structure of relatedness (as in the main analysis; see below) and for UK Biobank, years of education were calculated in accordance with the protocol used in the previous GWAS meta-analysis (15). Overall, the polygenic profile score had a small, positive relation to offspring’s own educational duration.

The respective effect sizes for the SNP risk alleles were multiplied by the respective \( \beta \) of the polygenic profile score for each participant in the three cohorts. These scores are proxy measures (~50% accurate) of the parental genetic risk scores.

Polygenic prediction of mortality. The educational attainment polygenic scores were modeled against the age at death for the parents of the independent cohorts. Cox proportional hazards models were run, using a pedigree-derived kinship matrix to account for relatedness in Generation Scotland. Covariates included sex, and the first 15 SNP-based principal components to account for population stratification. Analyses were conducted in R using the “survival” (58), “kinship2” (59), and “coxme” (60) packages.

Meta-analysis. Fixed-effects meta-analysis was performed across the three cohorts, separately for mothers and fathers, using the metafor package for R (61).
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