Genetic Variation Associated with Differential Educational Attainment in Adults Has Anticipated Associations with School Performance in Children

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Abstract

Genome-wide association study results have yielded evidence for the association of common genetic variants with crude measures of completed educational attainment in adults. Whilst informative, these results do not inform as to the mechanism of these effects or their presence at earlier ages and where educational performance is more routinely and more precisely assessed. Single nucleotide polymorphisms exhibiting genome-wide significant associations with adult educational attainment were combined to derive an unweighted allele score in 5,979 and 6,145 young participants from the Avon Longitudinal Study of Parents and Children with key stage 3 national curriculum test results (SATS results) available at age 13 to 14 years in English and mathematics respectively. Standardised (z-scored) results for English and mathematics showed an expected relationship with sex, with girls exhibiting an advantage over boys in English (0.433 SD (95%CI 0.395, 0.470), p = 1.79 × 10⁻⁴⁰) and more similar results (though in the opposite direction) in mathematics (0.042 SD (95%CI 0.007, 0.050), p = 0.01) increases in standardised SATS score for English and mathematics respectively. Educational attainment is a complex multifactorial behavioural trait which has not had heritable contributions to it fully characterised. We were able to apply the results from a large study of adult educational attainment to a study of child exam performance marking events in the process of learning rather than realised adult end product. Our results support evidence for common, small genetic contributions to educational attainment, but also emphasise the likely life-course nature of this genetic effect. Results here also, by an alternative route, suggest that existing methods for child examination are able to recognise early life variation likely to be related to ultimate educational attainment.

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Introduction

Although evidence from twin and family studies suggests that psycho-social traits such as educational attainment are moderately heritable [1–6], identifying genetic variants reliably associated with such traits has proven particularly challenging. This is likely due to the use of samples that are too small and has been responsible for numerous false positives [2,7,8].

In efforts to address this, Rietveld et al. conducted a genome-wide association study (GWAS) of educational attainment in 126,559 individuals, which is much larger than previous social-science genetic studies [9]. This study identified three single nucleotide polymorphisms (SNPs) (rs9320913, rs11584700 and rs48512966) which were associated with educational attainment in adults. These variants are located in independent loci and whilst the mechanism of their association is unknown, they have been advanced as possible targets for further studies of either biological function or where intermediate phenotypes for educational attainment are available.

Effect sizes at these three confirmed signals are small (~0.02% of variance explained by the strongest of these – equating to ~1 month in total schooling experience per allele) and sit in a wider...
context where the genome-wide linear polygenic score from all SNPs accounts for only around 2% of variation in educational attainment. This architecture compares to that seen in other polygenic complex traits such as BMI and height which have single largest common genetic effects explaining up to 0.4% of observed variance and which have total common variant contributions that are far stronger [10–12]. Despite this, these results are the first to report a replicable genetic association with educational attainment and merit further dissection.

A natural extension to Rietveld et al. [9] is to follow-up the initial analysis in a collection whose phenotypic measurement is more intermediate or marks more formative features to ultimate educational attainment and in a study where known population characteristics contribute to the validation of the original signal of association. Whilst often not available in large samples, an ideal situation in this case would be to reassess the original genetic association in an ethnically homogeneous, large, population-based collection with a standardised method for the assessment of educational performance and potentially one from an earlier part of the lifetime. Not only would this provide a form of validation for the original association signal, but improved measurement would theoretically heighten analytical power. Further to this, association would also provide an alternative source of evidence suggesting that early life assessments are able to capture meaningful information about the likely formative events going on to be important for ultimate educational attainment.

This investigation set out to generate an allele score from genome-wide significant SNPs derived from the original association study for educational attainment and assess its association with refined measures of educational performance. To do this, this investigation derived this allele score (also known as a genetic risk score) and assessed whether this score-based summary of genetic variation is associated with educational attainment measured by key stage 3 national curriculum test results at age 13 to 14 years. The data used comes from an ethnically homogeneous collection from the South West of the UK, the Avon Longitudinal Study of Parents and Children (ALSPAC).

Results

The mean English and mathematics non-standardised scores recorded within the ALSPAC collection were level 5.63 (SD = 1.09) for English and level 6.18 (SD = 1.32) for mathematics respectively. These data were available for 10,323 English scores and 10,683 mathematics scores at age 13 to 14 years. The total number of children with data available for the analyses (i.e. with genotypic data, covariables and outcomes) was 5,979 for English, and 6,145 for mathematics.

English and mathematics z-scores showed an expected relationship with sex, with girls exhibiting an advantage over boys in English on average (0.433 SD (95% CI 0.395, 0.470), p<10^{-10}) and attaining more similar exam results to boys in mathematics (boys higher than girls, 0.042 SD (95% CI 0.004, 0.080), p = 0.0303) (Figure 1).

The SNPs rs9320913, rs11584700 and rs4851266 were observed within the ALSPAC sample set at minor allele frequencies of 0.483, 0.216 and 0.393 respectively. These all passed quality control for imputation and were imputed with frequencies of 0.483, 0.216 and 0.393 respectively. These all observed within the ALSPAC sample set at minor allele (boys higher than girls, 0.042 SD (95% CI 0.004, 0.080), p = 0.0303) (Figure 1).

The SNPs rs9320913, rs11584700 and rs4851266 were observed within the ALSPAC sample set at minor allele frequencies of 0.483, 0.216 and 0.393 respectively. These all passed quality control for imputation and were imputed with RSQR scores of >0.99 [13]. Mean unweighted allele score was 2.19 and attended an approximately normal distribution (Figure 2).

English z-score was associated with the unweighted allele score with each additional educational attainment increasing allele being associated with a 0.041 SD increase in SATS z-score (95% CI 0.020, 0.063), p = 0.0002. Whilst there was weaker evidence of association for mathematics, each additional educational attainment increasing allele was associated with a 0.028 SD increase in SATS z-score (95% CI 0.007, 0.050), p = 0.0103 (Figure 2).

A proportion (58.3%) of participants in this study had mothers involved in the original discovery study for adult educational attainment [9]. Neither taking into account maternal allele score in analyses nor looking only at offspring whose mothers were not part of the original discovery study had a substantive effect on the results shown above (Table 1). The relationships were also maintained when controlling for the first four principal components (PC) of genetic population structure in ALSPAC. We found no evidence for differences in the observed effects by sex. Results from sex-specific analyses can be found in Table S2.

SNP-specific effects were underpowered given the relatively small amount of variance explained by singular SNPs. However these analyses showed evidence for an association between rs9320913 and both the English and mathematics z-scores (Table S3).

Discussion

Our results suggest that there is an association between a child's educational performance at age 13 to 14 years and their unweighted allele score created from the three SNPs (rs9320913, rs11584700 and rs4851266) found to be associated with adult educational attainment by Rietveld et al. [9]. This is of particular interest as we are not only able to reposition the phenotype initially used in the discovery to one of intermediate nature (or process during the lifetime), but we have been able to demonstrate that in a homogeneous population using commonly available, standardised, exam results. Relationships between genotypic variation and SATS z-scores were consistent across a series of testing scenarios taking into account the presence of mothers in the original data set.

This work sits in a wider context of research attempting to understand the aetiology of extremely complex traits which have traditionally been recognised predominantly as social phenomena. Educational attainment is critically important as it is strongly associated with social and health related outcomes and there is a recognised gradient in relationships between education and health status [9,14–17]. To this complicated and controversial field, estimates of the genetic contribution to educational attainment have suggested that up to 40% of the variance in existing measures may be explained by genetic factors [3,14,18]. Some twin studies [4–6] have suggested that genetic variation accounts for up to 60% of the variation in educational attainment, though these high heritability estimates may be inflated by genetic interactions [19].

Previous work addressing this directly attempted to collect a harmonized measure of educational attainment in large numbers to try to unpick the contribution of common genetic variation [9]. This work coded study-specific measurements pertinent to education using the International Standard Classification of Education (1997) scale [20] and derived variables marking an individual's years of schooling or whether college education had been completed. Whilst successful in identifying genetic correlates, this work is of course limited by the distal nature of the measurements employed. Work here has been able to give evidence that there is likely to be a quantifiable genetic contribution to not only attained educational experience, but performance within childhood educational experience.

A 1 SD increase in the English and mathematics z-scores is approximately equivalent to increases of 1.09 levels and 1.32 levels in the non-standardised English and mathematics scores respec-
This corresponds to a child's non-standardised English score increasing by ~0.045 levels and their non-standardised mathematics score increasing by ~0.038 levels for each additional educational attainment increasing allele. To put this into context, if a child's English level increases by 2 national curriculum levels between the ages of 11 and 14, then an increase of 0.045 levels is approximately equivalent to the increase that would be expected over 3.5 weeks within this period. Put another way, these differences approximate to a tenth of that seen across the sexes for performance in English at this age.

Whilst it is not possible to make direct comparisons between the effect sizes reported here and those elsewhere owing to differing phenotypic measures, it is possible to make some comparison of variance explained. For total years of schooling, the strongest reliable genetic association concerning a single SNP was that of rs9320913 which, in a previous study, explained 0.02% of phenotypic variance [9]. The strongest effect on English score in this study was for the same variant and in this case it explained 0.18% of observed variance (0.10% in mathematics score, Table S4). In this study allele score explained 0.27% of the variance in English score and 0.10% in mathematics score.

Overall, the influence of the allele score appeared to be consistent across the assessment of English and mathematics. This does have some impact on the interpretation of these effects in that this may be suggesting a basal or developmental origin rather than an acquired performance change; however it is difficult to speculate. Finer examination of these effects does suggest that to some extent the evidence for a child's English z-score increasing with their allele score is stronger than that for mathematics. It is difficult to suggest why this might be the case, however a potential scenario may follow that if the allele score assessed here is equal in its effects across disciplines, then observed differences may be the result of measurement inconsistencies across the subjects or the result of the subjects being affected by different environmental influences. That being said, the presence of association between the same genetic variation and both measurements of ultimate adult educational attainment and childhood educational performance does provide another source of evidence (in addition to phenotypic...
correlation between these educational attainment measures) to the
question of whether existing examinations are able to capture
variation in early life relevant to ultimate educational attainment.
In a field where few previous GWAS have yielded evidence for
common genetic contributions [21,22] our results increase the
evidence that with appropriate sample design or measurement,
relationships may be found. Findings here go some way to validate
the association between the three SNPs found by Rietveld et al. [9]
and educational attainment suggesting that these observations are
less likely to be a function of chance, artefact or residual
population stratification. Furthermore, we have reinforced the
assertion that it is feasible to use a more distal phenotype in order
to gain a larger sample size, but have also provided the reciprocal
argument in that the use of more refined measures of educational
attainment can show these effects in smaller samples and that they
are persistent in earlier years. It would of course be interesting to
investigate whether the allele score made available from this
original work can be applied to other alternative measures of
educational attainment or cognitive ability in different population-
based sample sets.

Materials and Methods

Study population and ethics statement
ALSPAC is a transgenerational longitudinal cohort study
investigating factors that influence health and development. The
study recruited pregnant women living in the former county of
Avon, UK with estimated delivery dates between April 1991 and
December 1992 [23,24]. Children from 14,541 pregnancies were
enrolled initially, rising to 15,247 pregnancies by the age of 18
years. Data has been collected for a wide range of phenotypes as
well as genetic and biological samples. Information on data
available can be found in the Data Dictionary (http://www.
bristol.ac.uk/alspac/researchers/data-access/data-dictionary/).
Ethical approval for the study was obtained from the ALSPAC
Ethics and Law Committee and the Local Research Ethics
Committees.

Education data
Educational attainment was assessed by a child’s English and
mathematics results in their key stage 3 national curriculum tests.
Key stage 3 national curriculum tests (informally known as SATS)
were statutory for children age 13 to 14 (Year 9) in maintained
schools in England until 2008 [25,26]. The children’s test results
were obtained by data linkage of ALSPAC with the National Pupil
Database (NPD).

The national curriculum measures achievement by levels, where
the average level of attainment is level 2 at age 6 to 7, level 4 at age
10 to 11, and level 5 or 6 at age 13 to 14 [27]. The English test
could only be taken at one tier whereas the mathematics test could
be taken at one of four different tiers, chosen based on a child’s
expected level of achievement. A child could be awarded a level N,
3, 4, 5, 6 or 7 for English and a level N, 2, 3, 4, 5, 6, 7 or 8 for
mathematics. For mathematics the possible levels that they could
be awarded depended on the tier that they took. A level N is given
when a child does not reach the lowest level for that tier.

Due to different tiers existing for mathematics it is necessary to
adjust the marks according to tier before using them in analyses.
We adjusted the marks using the method described in Levacic et
al. [28]. The marks are adjusted so that, for example, a score of
level 4.5 corresponds to being awarded a level 4 and having a
mark that is halfway between the boundaries for a level 4 and a
level 5. In creating these scores we extended the definition of the

| Table 1. Relationships between SATS z-scores and unweighted allele score within the ALSPAC study. |
|---------------------------------------------------------------|
| **Regression model** | **Beta** | **95%CI** | **P-value** | **Number of observations** |
|----------------------|----------|-----------|-------------|---------------------------|
| **English z-score**  |          |           |             |                           |
| OLS of English z-score on child’s allele score (all children) | 0.041 | 0.020, 0.063 | 0.0002 | 5,979 |
| OLS of English z-score on child’s allele score, adjusting for maternal allele score (all children) | 0.041 | 0.011, 0.071 | 0.0076 | 4,008 |
| OLS of English z-score on child’s allele score (children without maternal genome-wide data) | 0.037 | 0.0003, 0.074 | 0.0483 | 1,971 |
| OLS of English z-score on child’s allele score (all children), controlling for first four PCs of genetic population structure in ALSPAC | 0.041 | 0.020, 0.063 | 0.0002 | 5,979 |
| **Mathematics z-score** |          |           |             |                           |
| OLS of mathematics z-score on child’s allele score (all children) | 0.028 | 0.007, 0.050 | 0.0103 | 6,145 |
| OLS of mathematics z-score on child’s allele score, adjusting for maternal allele score (all children) | 0.040 | 0.010, 0.071 | 0.0094 | 4,106 |
| OLS of mathematics z-score on child’s allele score (children without maternal genome-wide data) | 0.015 | −0.022, 0.052 | 0.4286 | 2,039 |
| OLS of mathematics z-score on child’s allele score (all children), controlling for first four PCs of genetic population structure in ALSPAC | 0.028 | 0.007, 0.050 | 0.0102 | 6,145 |

All models include sex and age as covariables.
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national curriculum levels to be a continuous variable taking values from 0 to 8 for English and 0 to 9 for mathematics, aligned with the discrete level values. By alignment we mean that, for example, a continuous level with a value ≥5 and <6 corresponded to the child originally being awarded a level 5. The mathematics scores were all correctly aligned with the level awarded. We dropped the English scores for 18 children whose scores were not aligned with the level that they were awarded. This may be due to errors in the data or school appeals against the level awarded resulting in a change of level. Due to a lack of normality in residuals, scores were inverse rank transformed and standardised and their z-scores were used in the analysis. We report results as standard deviation (SD) changes in SATS score throughout.

**Genetic data**

9,912 ALSPAC children were genotyped using the Illumina HumanHap550 quad genome-wide SNP genotyping platform by the Wellcome Trust Sanger Institute (Cambridge, UK) and the Laboratory Corporation of America (Burlington, NC, USA). Individuals with incorrect sex assignments; extreme heterozygosity (<0.320 and >0.345 for Sanger data and <0.310 and >0.330 for LabCorp data); disproportionate levels of individual missingness (>3%); evidence of cryptic relatedness (>10% identity-by-descent) or non-European ancestry were excluded. The resulting data set consisted of 8,365 individuals. Of 609,203 SNPs, those with a minor allele frequency of 0.5% or less were excluded. The resulting data set (release 22) was used in the original educational attainment GWAS conducted by Rietveld et al. [9]. These three genetic variants have been confirmed as single variants, genome-wide independent signals [9]. Our allele score takes the values 0, 1, …, 6. Each unit increase in the allele score corresponds to an effect size from Rietveld et al. [9]. We repeated these three regressions with the mathematics z-score as the outcome variable, rather than the English z-score.

Whilst ALSPAC is an ethnically homogeneous population and genome-wide data are quality controlled for population structure, we also performed a further version of the first analysis adjusting for the first four PCs derived from genome-wide common variant data to assess the contribution of population structure estimated according to Price et al. [29].

We lastly assessed potential differences by sex by using a likelihood ratio test to compare a model with no interaction term to a model with an interaction term fitted between sex and allele score. We performed this test for the English and mathematics z-scores separately.

**Supporting Information**

**File S1** Supplementary information. (DOCX)

**Table S1** Genome-wide meta-analysis results for educational attainment in a sample excluding mothers from the Avon Longitudinal Study of Parents and Children. (DOCX)

**Table S2** Sex specific estimates of the relationship between English and mathematics SATS z-scores and allele score in the ALSPAC study. (DOCX)

**Table S3** Estimates of the relationship between English and mathematics SATS z-scores and individual SNPs rs9320913, rs11304700 and rs4851266 in the ALSPAC study. (DOCX)

**Table S4** Percentage of variation in the English and mathematics scores explained by the child's allele score and the individual SNPs. The regression models do not include any covariables. (DOCX)
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