Birth Order Differences in Education Are Environmental in Origin

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Abstract

Siblings share many environments and much of their genetics. Yet, siblings turn out different. Intelligence and education are influenced by birth order, where earlier-born outperform later-born. We investigate whether birth order differences in education are caused by genetic differences. Using data that spans two generations, combining registry, survey, and genotype information, this study is based on the Norwegian Mother, Father and Child Cohort Study (MoBa). We show that there are no genetic differences by birth order as captured by polygenic scores (PGSs) for educational attainment. Furthermore, we show that earlier-born have lower birth weight than later-born, indicating worse in utero environments. Educational outcomes are higher for earlier-born children when we control for PGSs and in utero variables. Finally, we consider environmental influences, such as maternal age, parental educational attainment, and sibling genetic nurture. We show that birth order differences are not genetic in origin, but their environmental cause remains elusive.

Introduction

Sibling differences receive less attention than sibling similarities, although they make up an important part of the picture of social inequality. Even for socio-economic outcomes, inequality within families (i.e., among adult siblings) can be as large as inequality between families 1. While evidence indicates a large role for environmental influences in producing differences among siblings in the same family, documenting specific, systematic influences has been more elusive.

Birth order has long been offered as an example of a systematic source of environmental differentiation within families. Empirically, firstborn siblings have slightly higher intelligence 2–4, educational achievement 5,6, and income 7,8 than their siblings born later. These differences are routinely interpreted as reflecting causal mechanisms that are social environmental in origin. For example, some work suggests that parental resources are more diluted for laterborn siblings as a result of competing demands for parental attention 9,10, while other work considers whether the presence of older siblings adversely affects the cognitive environment in which younger siblings are raised 11,12.
Even though birth order differences in achievements are routinely presumed to reflect environmental causes, there are several reasons to worry they may not. First, many studies do not compare siblings in the same family, for which genetic differences across families lurk as an uncontrolled confounder. Even when they do, birth-order studies often define birth order by the rearing family or by siblings who share a mother. Genetic differences by birth order may result from paternal genetic differences by siblings who do not share a father. Even when studies intend to include only full siblings, this is usually based only on maternal self-report. Sometimes mothers are unsure themselves who the father is or misreport paternity for other reasons. For half-siblings with different fathers, paternal genetic differences could be associated with birth order, as mothers who have children with multiple partners typically have children with less educated fathers for their laterborn children.

Second, genetic differences may also be induced by fertility decision-making. Genetic differences among siblings are sometimes described as a "lottery" and parents may be more likely to have an additional child if their already-born children evince desirable traits consistent with a favorable draw from this "lottery". Later-born children may exhibit a "regression to the mean" phenomenon of less propitious genetic endowments for these same traits.

Third, mutations rise with parental age, in particular with paternal age at conception. There is evidence of advanced paternal age increases the probability of offspring autism, as well as the rate of several other health-related traits. Within families, of course, birth order and parent age are always monotonically related.

In addition, maternal age also may influence in-utero environments, such as the rate of antibody attacks. Beyond this, maternal nutrition, stress, medical professional visits, and other health-related behaviors may vary according to birth order, causing differences in birth weight and other outcomes which influence intelligence and educational attainments later in life.

If birth order differences are not environmental in origin, this would not only have implications for our understanding of family dynamics, but it would undermine various causal inference strategies that effectively assume genetic differences among full siblings are independent. We examine whether birth order effects on education are influenced by genetic differences as
measured by polygenic scores for educational attainment. We do so with data that combines registry, survey, and genotype information from families in Norway (see Methods). We also consider birth weight and birth length, as indicators of combined genetic and in-utero influences, and we also examine maternal age. Furthermore, we look at non-transmitted alleles of parents, whose influence on development is sometimes referred to as "genetic nurture".30–32.

In the current study we address three research questions: 1) Is there a difference in genetics associated with educational attainment between birth orders? 2) Are there differences in birth weight and birth length between birth orders? 3) Does the putatively socially-based effect of birth order remain after accounting for genetic differences and in-utero-variables? After having established that birth order differences are environmental in origin, we follow up by investigating further what may cause these differences, considering whether birth order differences vary by polygenic scores, family background, and non-transmitted parental- and sibling alleles.

Results

We begin by analysing the full population of Norway using administrative data to see whether there are birth order differences among children and adults. To better parallel subsequent analyses with genomic data, we exclude participants whose parents were not born in Norway (see Methods). For the population of children we study (born 1994-2009), we examined performance on national tests by computing the child’s mean score of tests in three subjects measured at three times (grades 5, 8, and 9; standardized within subject, age, and cohort; N=301,795). For the adult population (b. 1945-1988), our outcome is completed years of education at age 30 (standardized within cohort, N=2,067,878). We use family-level fixed effects when estimating the bivariate association between birth order and these outcomes, meaning we are comparing siblings within the same family. We control for sex and maternal age, and run the models separately by sibship size.
Fig. 1 | Birth order and educational differences in the population. a,b Results from family-fixed effects linear regression models run separately by sibship size, with controls for sex and maternal age and cluster-robust standard errors. Firstborns serve as the reference category. All point estimates presented with 95 % CI. In a) children part of the population (N=301,795) where the outcome is the mean of national test score standardized within test, year of test, and birth cohort. b) Parental part of the population (N=2,067,878), where the outcome is educational attainment at age 30, standardized within birth cohort.

Consistent with other studies, we find that firstborn siblings have better educational outcomes than their later-born siblings. Figure 1 shows the magnitude of these associations. The top panel shows lower test scores for each successive birth order for all family sizes from 2-5 siblings. Most of these differences are present in the first test scores we observe (fifth grade), but the gaps do grow modestly from the last scores in our data (ninth grade) (see Supplementary Figure A1). The bottom panel shows similar patterns for educational attainment in the adult population.
**Polygenic score.** Figure 2 shows the relationship between birth order and the polygenic score for educational attainment among families with 2 and 3 children, which are the vast majority of sibships in Norway. Sample sizes for this and subsequent analyses using genetic data are much smaller than those used in the previous figure as these data are only available for a portion of the population (see Methods). We provide separate results for the child sample (panels a and b; N = 24,507; 2,705) and their parents (panels c and d, N = 43,316; 3,897). Within each sample, we provide estimates for models comparing siblings between families, and for models comparing siblings within families. In addition to sex and cohort, between-family models adjust for the ten principal components of the GWAS data to address potential confounding by ancestral differences (i.e., population stratification).
Fig. 2 | Polygenic Score and birth order. a, b, c, d, Association between educational attainment polygenic score and birth order. Results from linear regression models run separately by sibship size, with controls for sex. Firstborns serve as the reference category. All point estimates presented with 95% CI. In a, c, between-family estimate, adjusted for 10 principal components. In b, d, family-level fixed effects models with cluster-robust standard errors. In a, b, children part of the sample (N = 24,507 (a); 2,705 (b)); c, d, parental part of the sample (N = 43,316 (c); 3,897 (d)).

a)
In all panels in Figure 2, confidence intervals overlap zero for birth orders two and three, meaning that we find no differences in the polygenic score for educational attainment among different birth orders. Consequently, the genetic differences captured by that polygenic score cannot explain the observed relationship between birth order and educational outcomes.

**Birth weight and birth length.** In Figure 3, we turn to birth order differences in birth weight and birth length, available in the child part of the sample only (N=2,705). We adjust for child sex, gestational age, and the polygenic score for education.
Fig. 3 | Birth length, birth weight and birth order. a,b Results from family-level fixed effects linear regressions run separately by sibship size, with dummies for birth order. Children part of the sample (N=2,705). Cluster robust standard errors, 95 % CI. Firstborns serve as the reference category. In a, birth length, with different control variables, black point: sex; yellow triangle: sex, gestational age, educational attainment polygenic score; blue rectangle: sex, gestational age, educational attainment polygenic score, mothers age at birth. In b, birth weight, with controls, black point: sex; yellow triangle: sex, gestational age, educational attainment polygenic score and birth length; blue rectangle: sex, gestational age, educational attainment polygenic score, birth length, mothers age at birth.
On average, laterborn children are both longer and heavier than firstborn children. Differences in length account for roughly half of the difference in weight. Further analyses indicate that the differences in mean length or weight are not due to differences only in especially low- or high-weight births (see Supplementary Figure A2). We find nearly identical results in the between-family models (see Supplementary Figure A3). As an indication of advantaged birth weight environment, then, it appears that later born children are actually advantaged relative to their first born siblings.

**Maternal age and birth spacing.** We considered also the possibility that birth order differences were due to maternal age, as maternal age has been variously posited to influence cognitive development via both biological and social mechanisms. To confound an observed firstborn advantage, mothers age would need to be inversely associated with test scores. When we simultaneously model within- and between-family effects, we find maternal age to be positively associated between families and even more so within families (see Supplementary Table A1). Within families, maternal age differences correspond to birth spacing differences, so our finding is that birth order differences decline as spacing increases. While this pattern is consistent with explanations of birth order differences rooted in differential parental investment and overall intellectual climate in the family, it is not consistent with ideas of their being an in utero or other biological advantages to being born to a younger mother.

**Educational attainment/achievement with controls.** In Figure 4, we show birth order differences within families after adjusting for all the aforementioned measures simultaneously: sex, gestational age, polygenic score for education, birth weight, birth length, and mothers age at birth. We show results for birth order differences in educational achievement (children, panel a, N=2,933) and educational attainment at age 30 (parents, panel b, N = 3,365). In all models, point estimates are negative for laterborn children compared to firstborn. Although controlling for mothers’ age at birth increase the uncertainty of estimates and confidence intervals overlap zero (blue rectangle in Fig. 4), all point estimates remain negative, and some even increase.
Fig. 4 | Educational achievement/attainment and birth order. a,b Results from family-level fixed effects linear regressions run separately by sibship size, with dummies for birth order where firstborns are the reference. Cluster robust standard errors, 95% CI. In a, children part of the sample (N = 2,933) with mean of national test score as the outcome. Controls for sex (black circle); sex, gestational age, educational attainment polygenic score (yellow triangle); sex, gestational age, birth weight, birth length, educational attainment polygenic score (blue rectangle). In b, parental part of the sample (N = 3,365) with educational attainment at age 30 as the outcome. Controls for sex (black circle); sex, educational attainment polygenic score (yellow triangle), and sex, educational attainment polygenic score, mothers age (blue rectangle).
Taken together, these analyses rule out various scenarios by which birth order differences in educational outcomes are either confounded or the result of non-environmental causes. The availability of multigenerational genotyped data with extensive phenotyping does not contradict an understanding of birth order differences as environmental in origin.

**Interrogating Environmental Origins.** Given this, we next turned to see what additional clues the genomic data may offer for how these environmental influences operate. First, we examined whether birth order differences varied by polygenic score. Muslimova et al.\(^3^4\) posit that birth order differences may be strongest among children with the highest genetic potential for achievement, as these children may be able to capitalize best on increased parental investment. In the child sample, we observed no difference (see Supplementary Table A3 and Supplementary Figure A4). In the adult sample, we find a borderline statistically significant pattern opposite the expected direction: a smaller firstborn advantage among participants with higher polygenic scores. If that is true, higher genetic potential could mitigate the processes that produce birth order differences. Either way, there is no indication in our data that firstborn advantage only or more strongly exists among those with higher polygenic scores.

Second, we also considered whether, net of polygenic score, birth order differences are moderated by family background. Contradictory hypotheses exist in the literature. One is that advantaged environments would have greater disparities in effective parental investment and so larger birth order differences, or that advantaged environments would have more compensatory investment in lower-performing children, which would imply smaller birth order differences see\(^3^5\). Our results here were consistent with the former of these scenarios (Supplementary Table A2): differences were larger in families in which parents had higher educational attainment, including within-families and net of polygenic score. Still, the moderation according to family background is very small compared to the overall birth order difference (Supplementary Figure A5)

Third, we examined the influence of other family member’s polygenic scores on child achievement, net of the child’s own score. Recent findings of "genetic nurture" have documented relationships between non-transmitted parental alleles and child attainment and achievement\(^3^1,3^6\). Given that mothers are typically more involved in childrearing than fathers--not infrequently to a substantial degree--the finding of some past research that maternal non-transmitted alleles matter more for attainments than paternal non-transmitted alleles is perhaps
unsurprising\textsuperscript{30}. We did find this pattern in our data (see Supplementary Table A4). However, when we include the polygenic score for the siblings, this accounts for more than half of the magnitude of the parental polygenic scores, and results in maternal non-transmitted alleles being non-significant. The sibling score itself is borderline statistically significant. Nevertheless, the result points to the importance of considering that genetic nurture may reflect sibling influence to a greater extent than has been previously appreciated.

**Discussion**

Our starting point for this investigation was the common presumption that birth order differences in educational outcomes reflect environmental rather than genetic mechanisms. As we noted, there are multiple potential reasons for genetic variation by birth order. Historically the problem has been exacerbated by many studies using between-family samples and methods\textsuperscript{37}, but the possibility of genetic differences remains even in studies that compare siblings in the same family. The only way to assess decisively whether genetic differences may confound birth order studies is with data that contains genetic and comprehensive phenotypic information on many siblings. Using data that spans two generations of Norwegian siblings, we document no genetic differences by birth order as captured by polygenic scores for educational attainment. Similar findings have recently been reported in a preprint using data from the United Kingdom\textsuperscript{34}.

We examined other indicators of non-environmental origins of birth order differences: birth weight, birth length, and maternal age. Our analyses demonstrate that none of these provide any leverage for explaining the robust advantage that earlier-born children have in test scores and ultimate educational attainment in our data. Indeed, laterborn children are advantaged in birth weight and birth length, consistent with some earlier research\textsuperscript{26,38,39}, and so if anything they might have better in utero environments, and yet have worse educational outcomes. These results strengthen the conclusion that birth order differences in educational outcomes originate in post-birth environments\textsuperscript{4,13}. A happy side consequence of this finding is that various methodologies that use siblings in causal inferences, i.e. studies using PGS within-family to alleviate population stratification\textsuperscript{40,41}, and social science studies using siblings to capture the omnibus effect of (social background), are not confounded in ways they would be if birth order differences were due to genetic differences.

Given that birth order differences are not genetic, they constitute one of the clearest-cut examples of environmentally-induced social inequality within families. Understanding how they
come about offers a window that may be more generally instructive for mechanisms that lead to inequalities among people with similar familial and sociodemographic backgrounds. To this end, we find that the birth order differences in test scores are already mostly present at our earliest time of observation (5th grade, age 9/10) and grow only modestly from then to 9th grade. This clarifies that the environmental differences resulting in birth order differences largely manifest in the first decade of life. The difference in educational attainment among adults is similar in magnitude, suggesting that these differences resulting from childhood environments have lifelong effects.

The two main environmental theories on birth order effects are linked to parental resources and sibling interactions. According to the resource dilution theory, economic and parental resources deplete as more household members arrive. In societies where resources are abundant like Norway, cultural resources are thought to be more important than economic (Conley 2005). According to the theory, as earlier-born receive more of the parental cultural resources such as personal attention and help with homework than laterborns do, they excel in educational performance. While this pattern is consistent with the finding that birth order differences decline with increased spacing between births, it is not consistent with our results that maternal genetic endowments do not matter any more than paternal endowments, given that mothers usually contribute more to child-rearing than fathers. Other studies have found stronger effects of maternal genetics, so more research will be needed to reach any decisive conclusion.

As for sibling interactions, these have been most prominently raised in variations of "confluence theory", which proposes that siblings generally have negative effects on one another's cognitive development. Because firstborns have less exposure to being reared with siblings, and may also benefit from the opportunity to teach younger siblings, the negative influence of siblings is proposed to be least for firstborns. Our finding that the apparent influence of parental alleles shrinks when sibling polygenic scores are included, and that sibling polygenic scores have a borderline significant effect in their own right, indicates that some of what has hitherto been called "genetic nurture" could in fact be "sibling genetic nurture".

Sibling effects have received less attention as compared to parental resources when it comes to explaining birth order effects. That said, Gibbs et al propose a conditional resource dilution (CRD) model, opening up for institutional- and family-level variation influencing birth order
effects. In the CRD, brothers and sisters, and especially older siblings, may provide, rather than compete for resources (p. 741).

Our study does have some limitations. First, the sample size is limited, especially for the within-family analyses based on MoBa as we have relatively few families with two or more full siblings. Statistical power curtails our present capacity to draw more decisive conclusions for some of our analyses, although expansions of genotyping in MoBa may substantially improve power for future research. Second, polygenic scores are based on common alleles, and we are not able to tell if there are differences in rarer variants. If rarer variants in the form of de novo mutations were importantly associated with birth order, however, the most obvious explanation would have involved parental age, which is contradicted by our findings regarding maternal age. Third, polygenic scores for educational attainment still only account for a limited portion of the overall heritability of educational attainment, and they will likely improve in the future. Last, we have no direct measures of parental and sibling behaviors, which obviously thwarts the ability to interrogate either as putative causes. Amidst the excitement for what incorporating genomic information may bring to the social sciences, we should not lose sight that the available phenotyping of population datasets often still leaves much to be desired.

In addition, even though it is easy to talk about birth order differences like they pose a unary puzzle, in fact the importance of different environmental mechanisms may shift in different societal and historical contexts. For the cohorts in our child sample, the vast majority of Norwegian children attend child care from age 1, with the overall attendance being 90 % for children aged 1-5 in 2010 \(^46\). In such a context, mothers may be spending less time with their children than perhaps at any point in history, which might mean our data could be unpropitious for observing a maternal role in birth order effects. Scandinavian data has often been used in research on family dynamics research because of these countries’ exceptional population registers. Societies with exceptional data may also be exceptional in other respects. Moreover, Scandinavian population register data sources do not contain large-scale data on actual behaviors, resources, and practices within families, and there may simply be no way of getting to the bottom of birth order effects without such richer data. While our results may help put to rest lurking ideas that birth order differences are not environmental, the question of what about environments produce these differences remains far from resolved.
Methods

Population data and variables from administrative registries

The data in this paper are from several sources. We begin with data from administrative registries covering the full population of Norway. The registries are of very high quality, and do not suffer from attrition, and have few registration errors. From the Central Population Registry we identify all family linkages and demographic variables, like birth cohort, sibship size, birth order, and mothers age at birth. We identify sibship size and birth order according to birth year within each mother. We also identify any sibships with multiple births, and exclude them as the assignment of birth order is less clear cut, and family dynamics may be different in multiple birth sibships. To better mirror our genomic sample we remove individuals born outside of Norway. Current genomic methods do not allow for us to include persons of non-European ancestry, and we restrict the data based on the population registry based on this related criteria, i.e. we only include Norwegian-born to Norwegian parents. For the adult part of the population, we restrict birth cohorts 1945-1988.

We link the Central Population Registry to the National Educational database (NuDB) 47, also covering the full population. For the children part of our analysis, we use standardized tests as the outcome. From NuDB we use data from national tests conducted in 5th, 8th, and 9th grade in reading, mathematics, and English. We standardize each test to a z-score within test, year of test assessment, and birth cohort. Thereafter, we calculate a mean of all available test scores for each child, which serves as our outcome variable for the children in the sample. For the adult part of our analysis, the outcome is educational attainment at age 30 in years following the International Standard Classification of Education (ISCED 2011) 47,48. The variable is continuous as we transform it to being measured in years, using how many years it takes to complete the level of education attained following normal progression, according to ISCED. We standardize this variable too to compare effect sizes between outcomes. After restricting our sample and removing people with missing information etc., our population-based data has an N of 301,795 for the child birth cohorts, and 2,067,878 for the adult birth cohorts.

The Norwegian Mother, Father and Child Cohort Study (MoBa)

The prepared population data are linked to The Norwegian Mother, Father and Child Cohort Study (MoBa) 49,50, which we use for the analysis with genomic and in-utero variables. MoBa is
population-based pregnancy cohort study conducted by the Norwegian Institute of Public Health. Participants were recruited from all over Norway from 1999-2008, with the sample unit being pregnancy. The women consented to participation in 41% of the pregnancies. The cohort includes 114,500 children, 95,200 mothers and 75,200 fathers. The current study is based on version 12 of the quality-assured data files. The establishment of MoBa and initial data collection was based on a license from the Norwegian Data Protection Agency and approval from The Regional Committees for Medical and Health Research Ethics. The MoBa cohort is now based on regulations related to the Norwegian Health Registry Act. The current study was approved by The Regional Committees for Medical and Health Research Ethics. Some 98 110 individuals in around 32 000 trios are currently genotyped. This interim release is known as “MoBa Genetics”, and is comprised of several separately genotyped and imputed batches due to partial funding for genotyping on a per-project basis. The current release is a merger of all subprojects after quality control and imputation. Details are available here: https://github.com/folkehelseinstituttet/mobagen.

MoBa-participants are sampled on pregnancy independent of previous pregnancies 51, meaning that children in MoBa could have siblings younger or older not included in the sample. However, as we identify family linkages and birth order from the population registry, we observe full sibship sizes and birth orders independently of the information in MoBa. The sampling of births within birth order is therefore random. As we construct sibship size variables from registries with observations from the birth of the parents and up until 2018, we are confident that we observe completed fertility histories for the vast majority of the sample, as most have reached ages where fertility rates are very low. For the parental part of the MoBa-sample, we use the central population registry to link them to their parents (i.e. the grandparents of children in MoBa). By doing this, we can establish sibship size, birth order, and other demographic information from the Central Population Registry, and use that to investigate birth order differences taking into account genomic information also for the parental part of the MoBa-sample. Our observational window from the NuDB register is up until 2018. In a few cases the parents in MoBa have yet to have reached age 30 at our latest observation year 2018. Here, we take the latest observed age available, the lowest age being 27.

Variables from MoBa

MoBa contains genomic information for parents and children which allows us to create Polygenic Scores (PGS) for each individual based on a Genome-Wide-Association study (GWAS) for
educational attainment of 1,1 million people. For the MoBa-sample we conducted quality control using PLINK (v 1.90). Before we perform quality control in PLINK, we remove families with any individuals born in countries outside of Europe, The United States, Canada, New Zealand, or Australia. We also remove families with any multipartnered fertility parents in MoBa. Thresholds for genotyping call rate were set to 98 %, minor allele frequency (MAF) 5 %, and deviations from Hardy-Weinberg equilibrium was $<10^{-4}$ with mid p-value adjustment. We remove individuals with poor genotype quality with a threshold missing rate of 5 %, as well as those with heterozygosity rates which deviate ±3 standard deviations from the sample mean. We remove ancestry outliers in plink with an identity-by-state binominal test (PPC) set to 0.05, MAF 0.01 and compare the 1-5 nearest neighbors in the data. We remove families with a z-score for any neighbor below 4 standard deviations.

We use Pseudocons to create the non-transmitted part of parental genomes. We use PRSice to make polygenic scores for parents, children, and non-transmitted alleles for the parents using the same parameters. We use all available SNPs, genome-wide significant or not, and clump with a 250 kb window. Clump-r2 was set to 0.1, excluding SNPs with higher linkage disequilibrium. The polygenic score is standardized.

In addition to genomic information, MoBa has several survey waves. Birth length and birth weight were self-reported from the mother when the child was 6 months old.

After doing quality control on the genomic data, and removing people according to what we have just described, 79 057 individuals are left in the sample, 29 815 children, 24 032 fathers and 25 210 mothers. However, while we use this sample for between-family analysis in some figures, we rely mostly on within-family analysis, i.e. family fixed-effects with clustering on mothers and robust standard errors. Here, our sample size is smaller, as at least two children born to the same mother are needed to estimate the models, and we need both siblings to be genotyped (N=2,933).

Data availability: The consent given by the participants does not open for storage of data on an individual level in repositories or journals. Researchers who want access to data sets for replication should submit an application to datatilgang@fhi.no. Access to data sets requires approval from The Regional Committee for Medical Research Ethics in Norway and a formal contract with MoBa.
Acknowledgments:
This research is part of the EQOP (ERC Consolidator grant 818425 Socioeconomic gaps in language development and school achievement: Mechanisms of inequality and opportunity). The Norwegian Mother, Father and Child Cohort Study is supported by the Norwegian Ministry of Health and Care Services and the Ministry of Education and Research. We are grateful to all the participating families in Norway who take part in this ongoing cohort study. We thank the Norwegian Institute of Public Health (NIPH) for generating high-quality genomic data. This research is part of the HARVEST collaboration, supported by the Research Council of Norway (#229624). We also thank the NORMENT Centre for providing genotype data, funded by the Research Council of Norway (#223273), South East Norway Health Authority and KG Jebsen Stiftelsen. We thank Per Minor Magnus, Pål Njølstad and Ole Andreassen who headed the aforementioned projects. We further thank the Center for Diabetes Research, the University of Bergen for providing genotype data and performing quality control and imputation of the data funded by the ERC AdG project SELECTionPREDISPOSED, Stiftelsen Kristian Gerhard Jebsen, Trond Mohn Foundation, the Research Council of Norway, the Novo Nordisk Foundation, the University of Bergen, and the Western Norway health Authorities (Helse Vest). Isungset thank Robbee Wedow for making genomics less hard.
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