1	Proxy gene-by-environment Mendelian randomization study confirms a causal effect of maternal
2	smoking on offspring birthweight, but little evidence of long-term influences on offspring health
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19	Key Words: gene × environment, Mendelian randomization, proxy, maternal smoking, pregnancy

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Abstract Objective To validate a novel proxy gene-by-environment (G×E) Mendelian randomization (MR) approach by replicating the previously established effect of maternal smoking heaviness in pregnancy on offspring birthweight, and then use GxE MR to investigate the effect of smoking heaviness in pregnancy on offspring health outcomes in later life and grandchild's birthweight. Design A proxy G×E MR using participants' genotype (i.e. rs16969968 in CHRNA5) as a proxy for their mother's genotype. Setting UK Biobank. Participants 289,684 white British men and women aged 40-69 in UK Biobank. Main outcome measures Participants' birthweight and later life outcomes (height, body mass index, lung function, asthma, blood pressure, age at menarche, years of education, fluid intelligence score, depression/anxiety, happiness), and birthweight of female participants' first child. Results In our proof of principle analysis, each additional smoking-increasing allele was associated with a 0.018 (95% confidence interval (CI): -0.026, -0.009) kg lower birthweight in the "maternal smoking during pregnancy" stratum, but no meaningful effect (-0.002kg; 95% CI: -0.008, 0.003) in the "maternal non-smoking during pregnancy" stratum (interaction P-value=0.004). We found little evidence of an effect of maternal smoking heaviness on participants' later life outcomes. We found the differences in associations of rs16969968 with grandchild's birthweight between grandmothers who did versus did not smoke were heterogeneous (interaction P-value=0.042) among female participants who did (-0.020kg per allele; 95% CI: -0.044, 0.003) versus did not (0.007kg per allele; 95% CI: -0.005, 0.020) smoke in pregnancy. **Conclusions** Our study demonstrated how offspring genotype can be used to proxy for mothers' genotype in G×E MR. We confirmed the previously established causal effect of maternal smoking on offspring birthweight but found little evidence of an effect on long-term health outcomes in the

- offspring. For grandchild's birthweight, the effect of grandmother's smoking heaviness in pregnancy
- 45 may be modulated by maternal smoking status in pregnancy.
- 46 (word count: 300)

47 WHAT IS ALREADY KNOWN TO THIS TOPIC Heavier maternal smoking in pregnancy causes lower offspring birthweight 48 49 Maternal smoking in pregnancy is also associated with offspring outcomes in later life and 50 grandchild's birthweight, but it is not known whether these associations are causal 51 Understanding the transgenerational causal effects of maternal smoking heaviness in pregnancy is 52 important to inform public health policies 53 WHAT THIS STUDY ADDS 54 The proxy gene-by-environment Mendelian randomization approach can be used to explore 55 maternal effects on offspring phenotypes when maternal genetic information is unavailable The approach confirmed the causal effect of smoking on offspring birthweight. 56 57 Maternal smoking status in pregnancy modulates the effect of grandmother's smoking heaviness in pregnancy on grandchild's birthweight, highlighting the importance of smoking cessation before 58 59 pregnancy in each generation

Introduction

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The developmental origins of health and disease hypothesis proposes that early life experiences, including those in utero, can have long-term health effects, and maternal pregnancy exposures are important to long-term health of offspring (1). Heavier maternal smoking in pregnancy is known to be causally associated with lower offspring birthweight (2-6), but its other effects in offspring are less clear. Multivariable regression in observational data showed that heavier maternal smoking during pregnancy was associated with offspring being shorter (7) and more overweight/obese (8, 9), and having higher blood pressure (10), but had mixed associations with age at menarche (11), respiratory (12), cognitive (13), and mental health (14). Heavier maternal smoking in pregnancy has also been associated with higher grandchild's birthweight in certain subpopulations (15-17). It is unclear whether these associations reflect a causal effect of maternal smoking in pregnancy, as they may be due to residual confounding. Some studies have assessed this using paternal smoking as a 'negative control' since an effect via uterine environment would be observed in mothers but not fathers, such that similar-magnitude associations would indicate confounding via shared familial, social, environmental and genetic factors (2, 5, 18). Negative control studies suggest little evidence of a causal effect on offspring body mass index (BMI) (2, 5, 8), blood pressure (19, 20) and depression (21). Mendelian randomization (MR) provides an alternative way to explore this question by using single nucleotide polymorphisms (SNPs) as instrumental variables (IVs) for an exposure of interest. MR is less prone to confounding as germline genetic variants are randomly allocated at meiosis and not influenced by subsequent socioeconomic and health behaviours (22, 23). MR has been applied in a gene-by-environment (G×E) framework (24, 25), which requires variation in the strength of the gene-exposure association across strata of another factor. If there is a causal effect of the IV on the outcome via the exposure of interest, then we would expect the association of the IV with the outcome to vary in proportion to the gene-exposure association. The rs1051730/rs16969968

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(CHRNA5) SNPs, previously robustly associated with smoking heaviness amongst smokers (26), have been widely used as IVs for smoking heaviness in GxE MR studies (3, 27-29). A causal effect of the smoking heaviness IV on an outcome should be seen amongst ever but not amongst never smokers if the effect is via smoking heaviness rather than other pathways (24, 25). G×E MR has also been used to assess cross-generational causal effects. A smoking heaviness IV has been associated with lower offspring birthweight amongst mothers who smoked in pregnancy but not amongst mothers who did not smoke in pregnancy, suggesting the genetic instrument affects birthweight through maternal smoking (3). It is usually difficult to investigate transgenerational associations due to a lack of data across the generations of interest. Thus, previous work has sought to test transgenerational associations using available traits as proxies for unmeasured traits of interest. A Norwegian cohort aimed to examine whether women's smoking in adulthood was related to their mothers' smoking habits (that were not recorded) and hence used maternal smoking-related mortality as a proxy (30). Recently, a casecontrol by proxy approach has been proposed (31). Participants' genotypes were used to proxy unavailable parental genotypes, and their associations were tested against parental diagnosis of Alzheimer's disease in UK Biobank (31), since Alzheimer's disease was much more prevalent in the parents than the participants (aged between 40 and 69 at baseline in 2006-2010 (32)). Our study aimed to demonstrate how an analogous approach can be used within a G×E MR framework to test maternal-offspring effects when maternal genotype is not available, using offspring genotype as a proxy for the maternal genotype. First, we performed a proof of principle analysis to demonstrate this approach, testing the previously established finding that maternal smoking in pregnancy leads to lower offspring birthweight. Second, we tested for causal effects of maternal smoking on offspring later life outcomes. Finally, we tested for a causal effect of grandmother's smoking on grandchild's birthweight.

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Methods Study population Our study was conducted using UK Biobank, a population-based cohort of more than 500,000 men and women in the UK. This study collected a large and diverse range of data from physical measures, questionnaires and hospital episode statistics (32). Of 463,013 participants of European descent with genetic data passing initial quality control (i.e. genetic sex same as reported sex, XX or XY in sex chromosome and no outliers in heterozygosity and missing rates) (33), 289,684 participants (54% women) of white British descent were eligible for inclusion in our analyses (Supplementary Figure 1). We refer to the UK Biobank participants as generation one (G1), and their parents and offspring as G0 and G2, respectively. Genetic IV for maternal smoking The rs16969968 SNP located in CHRNA5 has been robustly associated with smoking heaviness (26). Ideally, we would use the maternal rs16969968 as an IV for the heaviness of maternal smoking, but in UK Biobank parental genetic data are not available. Hence, we used rs16969968 of the UK Biobank participants as a proxy for that of their mothers', coded as the number of smoking heaviness increasing alleles. Smoking phenotypes We used participants' answers to the question "Did your mother smoke regularly around the time when you were born?" as a proxy for G0 smoking during pregnancy. Participants were also asked to report their smoking status (current/former/never). We derived a binary ever versus never measure of smoking status by combining current and former smokers. For female participants with at least one live birth, we derived a measure denoting whether they smoked during the pregnancy of their first child (see Supplementary Methods). Outcomes in participants (G1)

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We used baseline data measured at the UK Biobank initial assessment center. Anthropometric traits included participants' birthweight (kg, self-reported), standing height (cm) and BMI (kg/m², constructed from standing height and weight). To assess lung function, forced vital capacity (L) and forced expiratory volume in 1-second (L) were measured by spirometry. Participants reported whether they had had asthma via the question "Has a doctor ever told you that you have had any of the following conditions?" (with an option of asthma) (34). Systolic and diastolic blood pressure (mmHg) were measured twice using a digital monitor or a manual sphygmomanometer if the digital monitor could not be employed, and we took the average of the two readings. Female participants reported their age at menarche. We derived years of education based on qualifications achieved by participants, as described previously (35). We included follow-up data of a subset of participants to define intelligence and depression/anxiety. Fluid intelligence score was generated as an unweighted sum of the number of correct answers given to 13 questions, and we used the earliest score if we had data at multiple time points (36). We defined depression/anxiety cases as participants that either answered "Yes" to "Have you ever seen a general practitioner (GP) for nerves, anxiety, tension or depression?" or "Have you ever seen a psychiatrist for nerves, anxiety, tension or depression?", or had hospital episode coded using ICD-10 (37). Happiness was assessed via a question – "In general how happy are you?", with six categories ranging from "extremely happy" to "extremely unhappy". Outcomes in participants' offspring (G2) The female participants with at least one live birth were asked to report their first child's birthweight. Male participants were not asked to report the birthweight of their offspring. Statistical analyses Proof of principle analysis: testing the causal effect of maternal (G0) smoking heaviness in pregnancy on participants' (G1) birthweight In this proof of principle analysis, we seek to replicate the finding, previously established using GXE

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MR and many other methods (6), that heavier maternal smoking causes lower offspring birthweight. We use our proxy GxE approach, where participants' (G1) genotype is used as a proxy for their mothers' (G0) genotype. To assess whether rs16969968 affects participants' birthweight via G0 smoking in pregnancy, we stratified our G1 sample by G0 smoking status during pregnancy, and then tested the associations of rs16969968 with birthweight in each stratum using multivariable linear regression. Since birth precedes smoking initiation, participants' genotype cannot affect birthweight through their own smoking heaviness, which means we do not need to consider smoking status of participants (Figure 1A). We included participants' sex as a covariate to reduce variation in their birthweight and the first ten principal components to control for population stratification. We assumed an additive genetic effect and identified the strength of interaction between strata using Cochran's Q test for heterogeneity. Testing for causal effects of GO smoking in pregnancy on G1 later life outcomes We use the proxy GxE MR approach to test for causal effects of maternal (G0) smoking heaviness on offspring (G1) height, BMI, lung function, asthma, blood pressure, age at menarche, education, intelligence, depression/anxiety and happiness. In contrast to our proof of principle example where participants smoking in adulthood cannot influence their birthweight, participants' rs16969968 could affect these outcomes via both maternal (G0) and participants' (G1) smoking heaviness (Figure 1B). To assess whether rs16969968 may affect these outcomes via maternal versus participants' smoking, we stratified on both maternal and participants' smoking status. In each stratum, we examined associations of rs16969968 with height, BMI, lung function, blood pressure, age at menarche, education and intelligence using linear regression, asthma and depression/anxiety using logistic regression, and happiness using ordinal logistic regression. We included participants' age at baseline, sex and the first ten genetic principal components as covariates. Height and age at menarche manifest around the time of puberty such that participants' own smoking can only affect these if they started smoking before these outcomes are determined. We

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conducted sensitivity analyses for these outcomes stratifying G1 participants according to whether they were ever smokers before achieving their adulthood height (assuming age at 17 for men and 15 for women (38)) or their age at menarche. Testing for causal effects of GO smoking in pregnancy on grandchild's (G2) birthweight To test for a causal effect of participants mothers' smoking on birthweight of participants' offspring, we stratified G1 women based on their own and their mothers' smoking status during pregnancy, as rs16969968 could affect G2 birthweight through both G0 and G1 smoking heaviness (Figure 1C). Within each stratum, we assessed associations of rs16969968 with G2 birthweight using linear regression, adjusting for the first ten genetic principal components. We estimated the strength of interaction between G0 smokers and G0 non-smokers within each G1 stratum. We also calculated a difference (39) in those associations between G0 smokers and G0 non-smokers within each G1 stratum, and estimated the strength of interaction between two differences to investigate whether G1 smoking status modulates the effect of rs16969968 on G2 birthweight. Our G × E MR may be vulnerable to collider bias (29, 40, 41), as we stratified on smoking status (Supplementary Figure 2). Therefore, we tested associations of rs16969968 with G0 and G1 smoking status and potential confounders available in UK Biobank (see Supplementary Methods). We also tested observational associations of maternal (G0) smoking status with offspring (G1) smoking status and all outcomes for comparison with our MR results. Analyses were performed using R version 3.5.1 (R Foundation for Statistical Computing, Vienna, Austria). Patient and public involvement The current research was not informed by patient and public involvement because it used secondary data. However, future research following on from our findings should be guided by patient and public opinions.

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Results

Characteristics of participants across sex are shown in Supplementary Table 1. Each additional smoking-increasing allele of participants' rs16969968 was associated with a 1.02 (95% confidence interval [CI]: 1.01, 1.03; P-value = 5×10⁻³) higher odds of their mothers' smoking in pregnancy, a 0.98 (95% CI: 0.97, 0.99; P-value = 7×10⁻⁴) lower odds of being an ever (versus never) smoker themselves, and a 1.06 (95% CI: 1.04, 1.09; P-value = 3×10^{-7}) higher odds that female participants were a smoker (versus non-smoker) in their own pregnancy. We found little evidence of an association between rs16969968 and potential confounders, with small associations for participants' age and years of education in some strata (Supplementary Table 2). Our proof of principle analysis found that, amongst participants whose mothers smoked in pregnancy, each additional smoking-increasing allele was associated with a 0.018kg lower birthweight (95% CI: -0.026, -0.009) after adjustment for covariates (Figure 2). Amongst participants whose mothers did not smoke in pregnancy, we found little evidence for an association of rs16969968 with birthweight (-0.002kg [95% CI: -0.008, 0.003]), and we observed heterogeneity between these associations (interaction P-value = 0.004). Figure 3 showed estimates of rs16969968 on the 12 outcomes in the UK Biobank participants. Overall, within each stratum, the estimates were broadly consistent between those whose mothers smoked and those whose mothers did not, except for height among participants who never smoked (all interaction P-values were in Supplementary Table 3). Each additional smoking-increasing allele was associated with a 0.115cm lower height (95% CI: -0.200, -0.030) among never smokers whose mothers smoked in pregnancy, but a 0.002cm lower height (95% CI: -0.057, 0.053) among never smokers whose mothers did not smoke in pregnancy (interaction P-value = 0.029). However, this difference was not observed amongst ever smokers (Figure 3A). We obtained largely consistent results in sensitivity analyses (Supplementary Figure 3). Figure 4 showed estimates of rs16969968 on grandchild's birthweight. Among female participants

who did not smoke in pregnancy, each additional smoking-increasing allele was associated with a 0.007kg higher grandchild's birthweight difference (95% CI: -0.005, 0.020) between grandmothers who did versus did not smoke in pregnancy. However, this difference was -0.020kg per allele (95% CI: -0.044, 0.003) among female participants who smoked in pregnancy. These two differences were heterogeneous (-0.028kg per allele [95% CI: -0.055, -0.001]; interaction P-value=0.042).

The directions of observational estimates were consistent with our MR estimates for both participants' and their child's birthweight. Our observational analyses also found associations of maternal smoking in pregnancy with offspring later life outcomes, where smoking in pregnancy was associated with lower height, higher BMI, poorer lung function, higher risk of asthma, earlier age at menarche, higher blood pressure, and poorer cognitive and mental health (Supplementary Table 4).

Discussion

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Principle findings and comparison with the literature In this study, we have demonstrated how G×E MR can be used to test transgenerational causal effects of maternal smoking heaviness in pregnancy using participants' genotype as a proxy for their mothers' genotype. Our proof of principle analysis identified an effect of heavier maternal smoking on lower offspring birthweight, consistent with previous studies (2-6). Our MR study also confirmed previously established causal effects of participants' smoking on their own health, where heavier smoking reduced BMI (27) and lead to impaired lung function (42), but found little evidence of an effect on asthma risk (43) or blood pressure (28). Our tests of effects of maternal smoking heaviness on offspring later life health outcomes were not conclusive, given a lack of precision for many of our MR estimates. We found little evidence of an effect on BMI, lung function, asthma, blood pressure, cognition, depression/anxiety or happiness. These findings were consistent with negative control studies for BMI (2, 8), blood pressure (19, 20) and depression/anxiety (21), although our estimation of interactions is not directly quantitatively comparable to their estimation of effects of ever/never smoking or smoking heaviness categories in observational studies. Our MR results found little evidence to support findings from our own and previous observational studies indicating maternal smoking led to poorer lung function (44), higher risk of asthma (45, 46), and lower happiness in offspring (47). This may be due to residual confounding in observational associations, or because of low statistical power in MR. Previous studies did not use the same cognition measurement approaches as used UK Biobank, making our results for this outcome less comparable. We observed lower offspring adulthood height according to maternal smoking in never smokers but not in ever smokers, which could be a chance finding given we tested multiple outcomes. We found little evidence of an effect of maternal smoking in pregnancy on offspring age at menarche. However, we did find an effect of rs16969968 on age at menarche across strata of

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smoking status (of both the participant and participants' mother) suggesting that rs16969968 may have horizontal pleiotropic effects on age at menarche (e.g. via smoking outside of pregnancy). Future MR studies could examine this (25). Our observational results were consistent with previous observational studies (15-17) by showing a positive association of grandmother's (G0) smoking in pregnancy with grandchild's (G2) birthweight after adjusting for mother's (G1) smoking in pregnancy. Although our G×E MR was vulnerable to insufficient statistical power, we did find evidence that female G1 smoking in pregnancy modulates the effect of G0 smoking heaviness in pregnancy on G2 birthweight, consistent with previous observational findings (15-17). These results highlight the importance of both grandmother's and maternal smoking in pregnancy for fetal growth, which could have implications for public health interventions aiming to reduce the prevalence of low birthweight. Strength of weakness of this study We now discuss some limitations of this work. First, our proxy GXE MR used offspring genotype as a proxy for maternal genotype and offspring rs16969968 contains 50% information from fathers. This may cause regression dilution bias in each stratum, where the measurement error in the SNP biases associations towards the null (48). However, we checked the extent that this might affect our results, by comparing the associations of participant's rs16969968 with their own birthweight versus their child's birthweight for smokers during pregnancy, and found little difference (-0.005kg (95% CI: -0.020, 0.009)) between them. Second, we stratified on smoking status which rs16969968 was weakly associated with. Stratification on colliders (between rs16969968 and outcomes) may bias our MR estimates (see Supplementary Figure 2) (40, 41). Additionally, we used a highly selected sample related to smoking (49) and had missing data in outcomes. These may also make our MR estimates vulnerable to selection bias (50). However, previous evidence (29, 51) and our genetic associations with measured confounders indicated that these selection effects may not be large enough to have a considerable impact on our MR estimates. Third, rs16969968 predicts life-course smoking heaviness

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and not just in pregnancy. Women who smoked in pregnancy may also smoke outside of pregnancy. Therefore, the effect of maternal smoking might be via other pathways such as poor oocyte quality for offspring birthweight, or postnatal maternal smoking (e.g. passive smoke exposure) for adulthood outcomes among offspring (52). Fourth, both participants' and their mothers' (G0) smoking status may be misclassified. Participants were asked to report whether their mother smoked around the time of their birth and we used this as our measure of G0 smoking in pregnancy. This means that G0 smokers might have smoked during all their pregnancy, part of their pregnancy or started smoking shortly after giving birth. Effects of smoking heaviness in pregnancy may vary according to the duration and pregnancy period during which a woman smoked. For instance, previous work found that smoking in the first trimester was not associated with lower birthweight in offspring suggesting that later stages may be more important for fetal growth (3, 15). Similarly, participants reported their smoking status at baseline, but this may not reflect their smoking status at an important time point for a given outcome. For, instance, participants' height and age at menarche can only be affected by their own smoking behaviour if they started smoking before achieving adult height or the onset of puberty. We performed sensitivity analyses for height and age at menarche using estimates of participants smoking status before these outcomes. For height, this assumed that men and women achieved their adult height at 17 and 15 years old (38), respectively, as this information was not available in UK Biobank. Fifth, we tested several hypotheses which increases the probability that our identified associations may be due to chance. Finally, our study may lack statistical power due to small sample sizes in strata and the low power of tests for interactions (53). We were unable to account for grandchild's sex in our models assessing the impact of grandmother's smoking in pregnancy since that is unavailable in UK Biobank, which may also reduce our statistical power. MR studies with larger sample sizes and hence greater statistical power are needed to further investigate transgenerational effects of smoking heaviness, together with studies in which both maternal and offspring genotype are known.

Conclusion

G×E MR demonstrates how offspring genotype can be used to proxy for maternal genotype to investigate causal effects of maternal smoking heaviness in pregnancy when maternal genotype is unavailable. We demonstrated our proxy GxE approach by replicating the previously identified effect of heavier smoking on lower offspring birthweight. We found little evidence of a causal effect of maternal smoking heaviness on offspring's later life outcomes. Finally, we found evidence that the effect of grandmother's smoking in pregnancy on grandchild's birthweight may be modulated by mother's smoking status in pregnancy. Further studies with larger sample sizes are needed to improve statistical power.

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Acknowledgments This research has been conducted using the UK Biobank Resource under Application Number 16729 (dataset ID 11148 and 21753). **Footnotes** Contributors: QY contributed to the design of the study, performed all analyses, wrote the first version of the manuscript, critically reviewed and revised the manuscript and approved the final version of the manuscript as submitted. LACM contributed to the design of the study, critically reviewed and revised the manuscript and approved the final version of the manuscript as submitted. GDS conceptualized the study, contributed to the design of the study, critically reviewed and revised the manuscript and approved the final version of the manuscript as submitted. GDS is the guarantor. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted. Funding: This work was supported by the University of Bristol and UK Medical Research Council [grant number MC UU 12013/1]. LACM is funded by a University of Bristol Vice-Chancellor's Fellowship. QY is funded by a China Scholarship Council PhD Scholarship. The funders had no role in the design, analyses, interpretation of results, writing of the paper, or decision of publication. Competing interests: All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi disclosure.pdf and declare: no support from any organization for the submitted work other than detailed above; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work. Ethical approval: The UK Biobank received ethical approval from the research ethics committee (REC reference for UK Biobank 11/NW/0382) and participants provided written informed consent.

Data sharing: The data reported in this paper are available by applying directly to the UK Biobank. All code used to produce the results can be accessed at https://github.com/MRCIEU/MR-maternal-smoking. Git tag v0.1 corresponds to the version presented here.

Transparency: The lead author (the manuscript's guarantor) affirms that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as originally planned (and, if relevant, registered) have been explained.

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doi: 10.1016/j.ntt.2012.09.004

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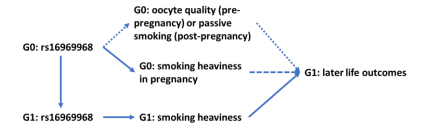
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Figure 1. Directed acyclic graphs (DAGs) of this study

(A)



(B)



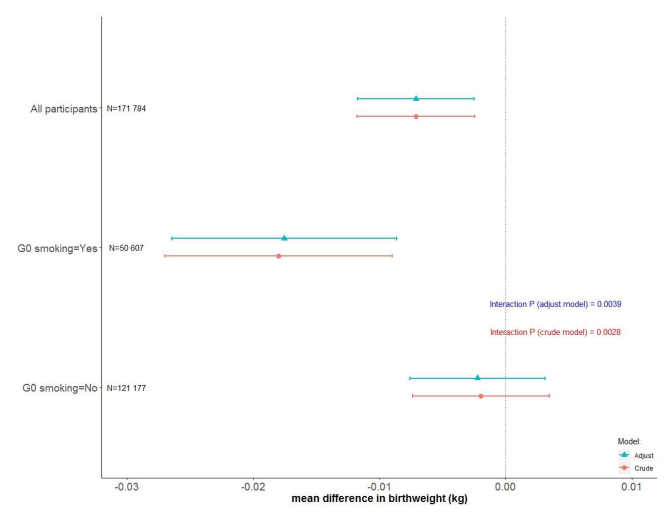
(C)



Generation (G)0: UK Biobank participants' mother; G1: UK Biobank participants themselves; G2: First offspring of UK Biobank participants.

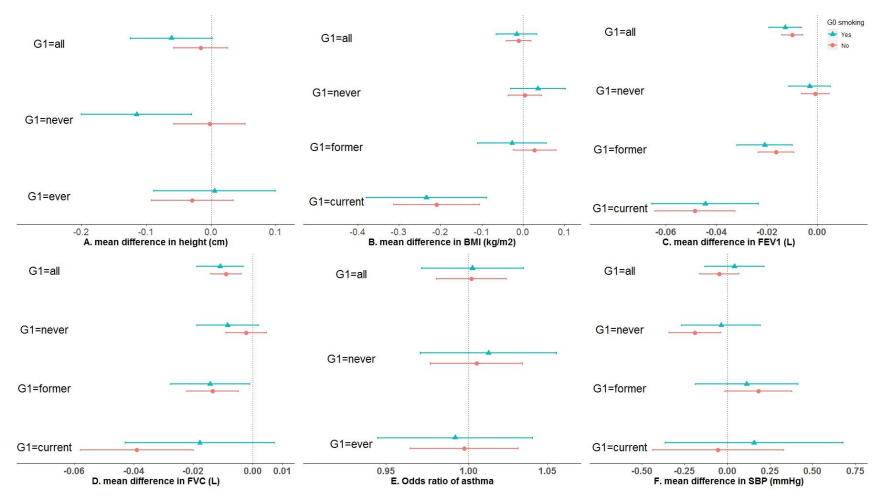
- A) Assessing the effect of G0 smoking heaviness on G1 birthweight: We used G1 rs16969968 as a proxy for G0 rs16969968 and stratified on G0 smoking status in pregnancy. There is no backdoor path (54) via G1 smoking heaviness since G1 cannot smoke before they were born. Maternal smoking outside of pregnancy might influence the outcome (52), e.g. via oocyte quality, causing an alternate path between rs16969968 and G1 birthweight (shown as).
- B) Assessing the effect of G0 smoking on G1 later life outcomes: Besides the paths described in (A), there is a backdoor path from G1 rs16969968 via G1 life-course smoking heaviness to the outcomes. To estimate the effect of G0 smoking heaviness in pregnancy (shown as ---->), we need to block this backdoor path by further stratifying on G1 smoking status.
- C) Assessing the effect of G0 smoking on G2 birthweight: Besides the paths described in (A), there is a backdoor path from G1 rs16969968 via G1 smoking heaviness in pregnancy to the outcomes. To estimate the effect of G0 smoking heaviness in pregnancy (shown as ---->), we need to block this backdoor path by further stratifying on G1 smoking status in pregnancy. G1 pre-pregnancy smoking might influence G2 birthweight (shown as ---->).
- See further DAGs in the Supplementary Figure 2 illustrating potential sources of bias due to conditioning on a collider.

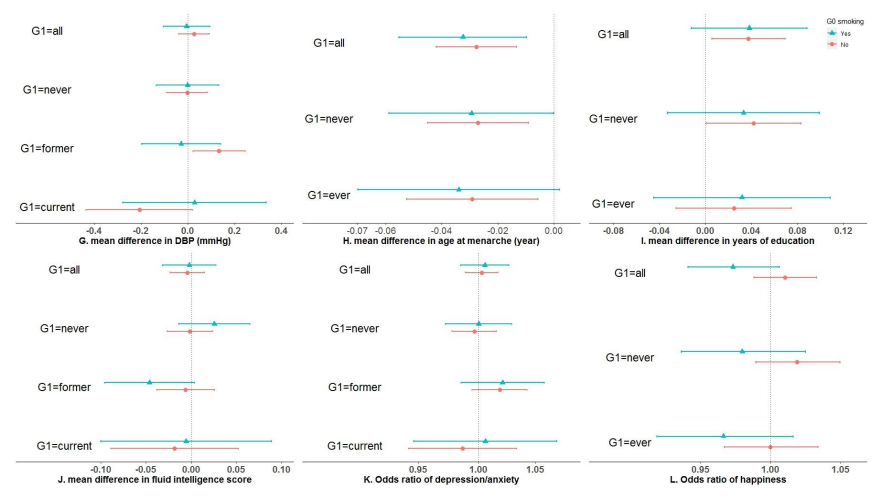
Figure 2. The associations of rs16969968 of UK Biobank participants with their own birthweight by their mothers' smoking status during pregnancy



Generation (G)0: UK Biobank participants' mother; G1: UK Biobank participants themselves. Estimates are the mean difference of G1 birthweight per each smoking-heaviness increasing allele of rs16969968. Associations are adjusted for sex of participants and the first ten principal components. The number of participants was listed for each analysis.

Figure 3. The associations of rs16969968 with 12 outcomes in UK Biobank participants by their mothers' smoking status during pregnancy and their own smoking status



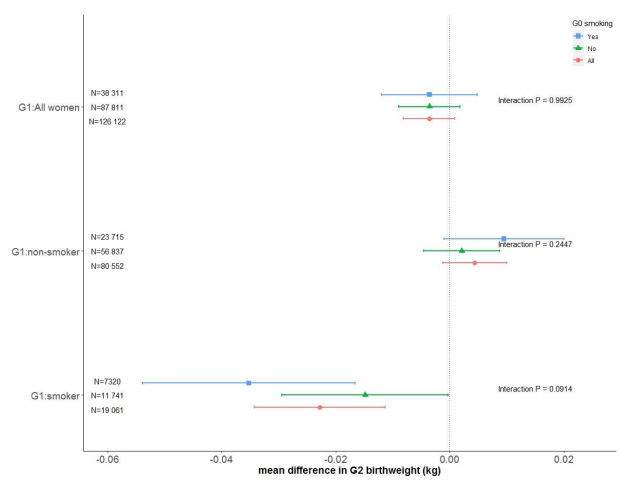


Generation (G)0: UK Biobank participants' mother; G1: UK Biobank participants themselves. Estimates are the mean difference (or change in odds) of G1 outcome per each smoking-heaviness increasing allele of rs16969968. We adjusted for age and sex of participants for outcomes except for menarche, and the first ten principal components for all 12 outcomes. We combined G1 current and former smokers into ever smokers for height, menarche, education, asthma and happiness to enlarge sample sizes given smoking cessation may not have a rapid impact on them.

Abbreviations: BMI, body mass index, DBP, diastolic blood pressure; FEV₁, forced expiratory volume in 1-second; FVC, forced vital capacity; SBP, systolic

blood pressure.

Figure 4. The associations of rs16969968 of UK Biobank women participants with their first child's birthweight by their mothers' and their own smoking status during pregnancy, after adjusting for the first ten genetic principal components



 Generation (G)0: UK Biobank participants' mother; G1: UK Biobank participants themselves; G2: First offspring of UK Biobank participants. Estimates are the mean difference of G2 birthweight per each smoking-heaviness increasing allele of rs16969968. Interactions are tested between G0 smokers (blue line) and non-smokers (green line) with their P-values presented. All women in G1 included G1 smokers, G1 non-smokers and G1 women whose smoking status in pregnancy was missing.