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ABSTRACT

Objectives: To investigate whether associations of smoking with depression and anxiety are likely to be causal, using a Mendelian randomisation approach.

Design: Mendelian randomisation meta-analyses using a genetic variant (rs16969968/rs1051730) as a proxy for smoking heaviness, and observational meta-analyses of the associations of smoking status and smoking heaviness with depression, anxiety and psychological distress.

Participants: Current, former and never smokers of European ancestry aged ≥16 years from 25 studies in the Consortium for Causal Analysis Research in Tobacco and Alcohol (CARTA).

Primary outcome measures: Binary definitions of depression, anxiety and psychological distress assessed by clinical interview, symptom scales or self-reported recall of clinician diagnosis.

Results: The analytic sample included up to 58,176 never smokers, 37,428 former smokers and 32,028 current smokers (total N=127,632). In observational analyses, current smokers had 1.85 times greater odds of depression (95% CI 1.65 to 2.07), 1.71 times greater odds of anxiety (95% CI 1.54 to 1.90) and 1.69 times greater odds of psychological distress (95% CI 1.56 to 1.83) than never smokers. Former smokers also had greater odds of depression, anxiety and psychological distress than never smokers. There was evidence for...
positive associations of smoking heaviness with depression, anxiety and psychological distress (ORs per cigarette per day: 1.03 (95% CI 1.02 to 1.04), 1.03 (95% CI 1.02 to 1.04) and 1.02 (95% CI 1.02 to 1.03) respectively). In Mendelian randomisation analyses, there was no strong evidence that the minor allele of rs16969968/rs1051730 was associated with depression (OR=1.00, 95% CI 0.95 to 1.05), anxiety (OR=1.02, 95% CI 0.97 to 1.07) or psychological distress (OR=1.02, 95% CI 0.98 to 1.06) in current smokers. Results were similar for former smokers.

Conclusions: Findings from Mendelian randomisation analyses do not support a causal role of smoking heaviness in the development of depression and anxiety.

INTRODUCTION

Smoking is highly comorbid with both depression and anxiety across many different populations. Furthermore, there is evidence to suggest that tobacco control interventions may not be as effective in populations with mental health conditions; for example, recent trends in the USA suggest that, since 2004, smoking rates have declined less rapidly in individuals with anxiety than in the general population. Given the profound public health burden of both tobacco-related disease, and depression and anxiety, understanding this relationship is of great importance.

Unfortunately, it is difficult to infer causal links between smoking and depression and anxiety from observational data, due to confounding. There may be factors associated with both smoking and depression and anxiety, such as other substance use (eg, alcohol), socioeconomic adversity and education which cannot be fully accounted for in observational studies. In addition, even if a causal association does exist, the direction of the relationship between smoking and depression and anxiety is unclear. Prospective studies have provided evidence that depressive symptoms are associated with increased likelihood of smoking initiation, while smoking cessation appears to be associated with a short-term increase in depressive symptoms during their quit attempt among a subgroup of smokers, and these individuals have poor smoking cessation outcomes. This evidence is consistent with the popular belief that cigarette smoking can reduce anxiety and improve mood, particularly among those experiencing anxiety or low mood (the self-medication hypothesis). However, there is also a growing body of evidence suggesting that smoking may contribute to the development of these conditions and that smoking cessation is associated with improvements in mental health, including depression and anxiety, compared to continued smoking.

Plausible biological mechanisms through which constituents of tobacco smoke may cause depression and anxiety have been described. In animal studies, for example, there is evidence that nicotine administration produces dysregulation in the hypothalamic-pituitary-adrenal system, which leads to hypersecretion of cortisol and changes in the activity of associated monoamine neurotransmitters. These systems function to regulate the biological and psychological reactions to stressors. Similarly, human data have demonstrated elevated cortisol levels in smokers compared to non-smokers. Constituents of tobacco smoke inhibit the activity of monoamine oxidase, enzymes that are involved in the breakdown of monoamines (including dopamine, serotonin and norepinephrine); this effect appears to normalise following smoking cessation. Animal studies also indicate that both drugs of abuse (including nicotine) and environmental stressors appear to trigger changes in midbrain dopaminergic function. Consequently, prolonged smoking may act to sensitise stress response systems, weakening adaptive coping responses and making smokers more susceptible to emotional distress in response to environmental stressors.

Mendelian randomisation methods allow us to investigate causal relationships in humans by using genetic variants as proxies for exposures of interest. The principle of Mendelian randomisation relies on the basic (but approximate) laws of Mendelian genetics (segregation and independent assortment). When these principles hold, genetic variants, at a population level, will not be associated with the confounding factors that generally distort conventional observational studies. In addition, as an outcome measure cannot alter the genotype that an individual is born with, these analyses should not be biased by reverse causality. A genetic variant, single nucleotide polymorphism (SNP) number rs16969968, in the CHRNA5-CHRNA3-CHRNB4 (CHRNA5-A3-B4) nicotinic receptor subunit gene cluster on chromosome 15 has demonstrated robust association with smoking heaviness within smokers.

The rs16969968 variant is functional and leads to an amino acid change (D398N) in the nicotinic receptor α5 subunit protein. The minor (risk) allele of this variant is associated with an average increase in smoking amount of one cigarette per day in smokers, and even more strongly associated with increases in cotinine (a metabolite of nicotine) levels. However, given the known role of the variant in altering receptor function, it is likely that the greater variance explained for cotinine levels is due to this measure better capturing total tobacco exposure, and not because the variant directly affects nicotine metabolism. There is also good evidence that the rs16969968 variant, unlike smoking heaviness, does not associate with confounding factors that may distort associations with health outcomes, for example, socioeconomic status and education level.

The rs16969968 variant (or its proxy rs1051730) has been used as an instrument for smoking heaviness in Mendelian randomisation studies to demonstrate that smoking causally lowers body mass index and that maternal smoking during pregnancy lowers offspring birth weight (see figure 1 for an illustration of the Mendelian randomisation approach). Using the rs1051730 variant, two recent studies have applied Mendelian randomisation to examine the causal
relationship of smoking with depression and anxiety. In one large Norwegian population, the rs1051790 variant was not associated with depression or anxiety in smokers; in a British cohort, the rs1051790 variant was associated with decreased depression during pregnancy in women who smoked prior to pregnancy. These findings are not consistent with a causal role of smoking in increasing depression or anxiety. To test the robustness of these findings, we performed a Mendelian randomisation meta-analysis combining data from 25 studies (n=127,632) in the consortium for Causal Analysis in Tobacco and Alcohol (CARTA).

METHODS

Study populations

We used data on individuals aged ≥16 years and of self-reported European ancestry from 25 studies from the CARTA consortium: the 1958 Birth Cohort (1958BC), the Avon Longitudinal Study of Parents and Children (ALSPAC, including both mothers and children), the British Regional Heart Study (BRHS), the British Women’s Heart and Health Study (BWHHS), the Caerphilly Prospective Study (CaPS), the Christchurch Health and Development Study (CHDS), Cohorte Lauzanico (CoLaus), the English Longitudinal Study of Ageing (ELSA), the National FINRISK Study (FINRISK), Generation Scotland: the Scottish Family Health Study (GS:SFHS), Genomics of Overweight Young Adults (GOYA) females, the Helsinki Birth Cohort Study (HBCS), Health2006, Health2008, the second wave of the Nord-Trøndelag health study (HUNT 2), Inter99, the Northern Finland Birth Cohorts (NFBC1966 and NFBC1986), the National Health and Nutrition Examination Survey (NHANES), the MRC National Survey of Health and Development (NSHD), the Netherlands Twin Registry (NTR), Patch 2, the Rotterdam Study, the Saguenay Youth Study-Parents (SYS-P) and the Whitehall II study. Further details of these studies are provided in online supplementary material.

Genotype

Within each study, individuals were genotyped for one of two SNPs in the CHRNA5-A3-B4 nicotinic receptor subunit gene cluster, rs16969968 or rs1051790. These SNPs are in perfect linkage disequilibrium with each other in Europeans (R²≈1.00 in HapMap 3, http://hapmap.ncbi.nlm.nih.gov/) and therefore represent the same genetic signal. Where studies had data available for both SNPs, we used the SNP that was genotyped in the largest number of individuals. Details of genotyping methods within each study are provided in online supplementary material.

Measures of depression, anxiety and psychological distress

Depression and anxiety were assessed by clinical interview, symptom scales or self-reported recall of clinician diagnosis (see table 1). As some of the scales do not distinguish between symptoms of depression and anxiety, we used the term ‘psychological distress’ to refer a composite phenotype.

To compare measures across studies, we created two case definitions for each of depression, anxiety and psychological distress (see table 2). According to case definition 1, individuals were classified as depressed or anxious if they self-reported clinician diagnosis of depression or anxiety, met clinical criteria for depression (excluding bereavement where known) or anxiety, or were above previously published cut points for depression or anxiety on symptom scales. Individuals were classified as having psychological distress if they met case definition 1 for depression or anxiety, or if they were above a cut point on a general scale for psychiatric symptoms. As not all scale measures used for assessing mental health have published cut-offs for defining cases, we created a second definition. According to case definition 2, individuals were classified as depressed or anxious if they were above the 90th centile for the specific depression or anxiety scales, and psychologically distressed if they were above the 90th centile on either the depression or anxiety scales or above the 90th centile on the general scales of psychiatric symptoms. Where both case definitions 1 and 2 were available within a study, case definition 1 was used. Full details of the measures and cut points used are provided in online supplementary table S1.

For the majority of studies (k=17), diagnoses were based on current depression and anxiety (at the time of measurement). Where current diagnoses were not available, diagnoses of depression or anxiety in the previous 12 months or lifetime diagnoses were used. For lifetime diagnoses, if information on age at first diagnosis was collected, individuals reporting diagnoses prior to 16 years of age were excluded.

Figure 1 Diagram of Mendelian randomisation analysis of smoking and depression/anxiety. The genetic variant rs16969968/rs1051790 is associated with smoking heaviness but should not be associated with the confounders of the association between smoking heaviness and depression/anxiety. In addition, there is no pathway from depression and anxiety to the genetic variant (reverse causality).
Symptom scales were also used as continuous measures of depression, anxiety and psychological distress (see online supplementary table S2). To compare across studies, these were converted to z-scores within each study. Most measures of depression, anxiety and psychological distress were strongly right skewed. However, standard transformations (e.g., log, square root) did not greatly improve distributions in several of the samples. Therefore, z-scores were constructed using the untransformed data in all samples (z-score = (individual score – sample mean) / sample SD).

### Smoking status

Smoking status was self-reported (either by questionnaire or interview) at the same time as mental health assessment for all studies, with the exception of 1958BC, CoLaus and HBCS which used smoking status and depression/anxiety data collected up to 3 years apart (see online supplementary material). Individuals were classified as never, former, current or ever (i.e., current and former combined) cigarette smokers. Where information on smoking frequency was available, current smokers were restricted to individuals smoking at least one cigarette per day. Where information on pipe and cigar smoking was available, individuals reporting being current or former smokers of pipes or cigars but not cigarettes were excluded from all analyses.

For studies with adolescent populations (ALSPAC children and NFBC1986), analyses were restricted to current daily smokers who reported smoking at least one cigarette per day.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Measures of depression, anxiety and psychological distress in the CARTA studies</th>
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</thead>
<tbody>
<tr>
<td>Study</td>
<td>Psychological Distress</td>
</tr>
<tr>
<td>1958BC</td>
<td>CIS-R</td>
</tr>
<tr>
<td>ALSPAC children</td>
<td>CIS-R</td>
</tr>
<tr>
<td>ALSPAC mothers</td>
<td>EPDS or CCEI</td>
</tr>
<tr>
<td>BRHS</td>
<td>EPDS</td>
</tr>
<tr>
<td>BWHS</td>
<td>GHQ-30</td>
</tr>
<tr>
<td>CaPS</td>
<td>CIDI (previous 12 months)</td>
</tr>
<tr>
<td>CHDS</td>
<td>CIDI (previous 12 months)</td>
</tr>
<tr>
<td>CoLaus</td>
<td>DIGS</td>
</tr>
<tr>
<td>ELSA</td>
<td>CES-D or clinician diagnosis of anxiety</td>
</tr>
<tr>
<td>FINRISK</td>
<td>Clinician diagnosis (previous 12 months)</td>
</tr>
<tr>
<td>Generation Scotland</td>
<td>GHQ-28</td>
</tr>
<tr>
<td>GOYA females</td>
<td>Clinician diagnosis (since giving birth)</td>
</tr>
<tr>
<td>HBCS</td>
<td>CES-D or STAI</td>
</tr>
<tr>
<td>Health2006</td>
<td>SCL-90-R</td>
</tr>
<tr>
<td>Health2008</td>
<td>SCL-90-R</td>
</tr>
<tr>
<td>HUNT</td>
<td>HADS</td>
</tr>
<tr>
<td>Inter99</td>
<td>SCL-90-R</td>
</tr>
<tr>
<td>NFBC1996</td>
<td>SCL-25</td>
</tr>
<tr>
<td>NFBC1986</td>
<td>YSR</td>
</tr>
<tr>
<td>NHANES</td>
<td>DIS (lifetime diagnosis)</td>
</tr>
<tr>
<td>NSHD</td>
<td>GHQ-28</td>
</tr>
<tr>
<td>NTR</td>
<td>ASR</td>
</tr>
<tr>
<td>Patch 2</td>
<td>ASR</td>
</tr>
<tr>
<td>Rotterdam</td>
<td>CES-D or M-CIDI</td>
</tr>
<tr>
<td>SYS-P</td>
<td>CES-D or DSM instrument</td>
</tr>
<tr>
<td>Whitehall II</td>
<td>GHQ-30</td>
</tr>
</tbody>
</table>

All scales measure current depression and anxiety unless otherwise stated. Clinician diagnosis was assessed by self-reported recall in all studies.

ALSPAC, Avon Longitudinal Study of Parents and Children; ASR, Adult Self Report; BC, Birth Cohort; BRHS, British Regional Heart Study; BWHS, British Women’s Heart and Health Study; CARTA, Causal Analysis Research in Tobacco and Alcohol; CCEI, Crown-Crisp Experiential Index; CES-D, Centre for Epidemiologic Studies Depression; CHDS, Christchurch Health and Development Study; CIDI, Composite International Diagnostic Interview; CIS-R, Computerised interview schedule-revised; CoLaus, Cohorte Lausannoise; DIGS, Diagnostic Interview for Genetic Studies; DIS, Diagnostic Interview Schedule; ELSA, English Longitudinal Study of Ageing; EPDS, Edinburgh Postnatal Depression Scale; GHQ, General Health Questionnaire; GOYA, Genomics of Overweight Young Adults; HADS, Hospital Anxiety and Depression Scale; HBCS, Helsinki Birth Cohort Study M-CIDI, Munich version of CIDI; NIHANES, National Health and Nutrition Examination Survey; NSHD, National Survey of Health and Development; NTR, the Netherlands Twin Registry; SCID, Structured Clinical Interview for DSM-III-R diagnosis; SCIDI/NP, Structured Clinical Interview for DSM-IV Axis disorders non-patient edition; SCL, symptoms checklist; STAI, State Trait Anxiety Inventory; YSR, Youth Self Report.
cigarette per day (current smokers) and individuals who had never tried smoking (never smokers).

Smoking heaviness in current smokers, measured as cigarettes smoked per day, was collected in some studies as a continuous variable and in some studies as a categorical variable. Further details of the smoking measures collected within each study are provided in the online supplementary material.

**Statistical analysis**

Analyses were conducted within each contributing study using Stata or R software, following the same analysis plan. Analyses were restricted to individuals with full data on depression and anxiety outcomes, smoking status and rs16969968/rs1051730 genotype.

Sex-adjusted and age-adjusted associations of smoking status (never, former, current, ever) and smoking heaviness with binary measures of depression, anxiety and psychological distress were assessed using logistic regression. For the smoking status analysis, never smokers were used as the reference group. The smoking heaviness analysis was restricted to current daily smokers, and ORs represent differences in odds of the outcome measure per additional cigarette consumed per day. These analyses were restricted to studies with continuous measures of cigarettes per day.

Within each study, genotype frequencies were tested for deviation from Hardy-Weinberg equilibrium (HWE) using a \( \chi^2 \) exact test. Mendelian randomisation analyses of the association between rs16969968/rs1051730 and binary measures of depression, anxiety and psychological distress were assessed using logistic regression and adjusted for age and sex. Analyses were performed stratified by smoking status (never, former, current and ever), because the variant only influences smoking heaviness in individuals who smoke. The analysis in never smokers is a test of a key assumption of Mendelian randomisation: that the gene only operates on the outcome through its effects on smoking heaviness (i.e., no pleiotropy). If rs16969968/rs1051730 only operates on an outcome measure through smoking heaviness, no association should be observed in never smokers. An additive genetic model was assumed, so ORs represent the difference in odds of the outcome per additional copy of the minor (risk) allele. As a secondary analysis, Mendelian randomisation analyses were performed of the association of rs16969968/rs1051730 with z-scores of symptoms scales for depression, anxiety and psychological distress using linear regression stratified by smoking status. These analyses were adjusted for age and sex and additionally for use of depression or anxiety medication where available. For studies with a survey (NHANES) or family-based design (SYS-P), appropriate methods were used to adjust SEs (see online supplementary material for further information).

**Table 2** Case definitions for depression, anxiety and psychological distress

<table>
<thead>
<tr>
<th></th>
<th>Case definition 1</th>
<th>Case definition 2</th>
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<tbody>
<tr>
<td>Depression</td>
<td>Self-report of clinical diagnosis OR</td>
<td>Scoring &gt;90th centile on specific depression scale</td>
</tr>
<tr>
<td></td>
<td>Meeting clinical criteria for depression OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Scoring above published cut point on specific depression scale OR</td>
<td></td>
</tr>
<tr>
<td>Anxiety</td>
<td>Self-report of clinical diagnosis OR</td>
<td>Scoring &gt;90th centile on specific anxiety scale OR</td>
</tr>
<tr>
<td></td>
<td>Meeting clinical criteria for anxiety disorders OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Scoring above published cut point on specific anxiety scale OR</td>
<td></td>
</tr>
<tr>
<td>Psychological distress</td>
<td>Depression or anxiety as defined above OR</td>
<td>Depression or anxiety as defined above OR</td>
</tr>
<tr>
<td></td>
<td>Scoring above published cut point on general scale for psychiatric symptoms OR</td>
<td>&gt;90th centile on general scale for psychiatric symptoms OR</td>
</tr>
</tbody>
</table>

Results from individual studies were meta-analysed in Stata (V.11) using the ‘metan’ command. Where there was evidence of heterogeneity between studies (I^2>50%), both fixed and random effects analyses were performed. Within the Mendelian randomisation analyses, the Cochran Q statistic was used to test for interactions between genotype and smoking status on the outcome measures.

Analyses were also performed stratified by sex because there is some evidence from observational studies that the association between smoking and mental health outcomes may differ by sex.\(^{42,45}\) Sex differences in the association between genotype and outcomes measures were tested for using meta-regression after taking into account potential differences by smoking status.
RESULTS

Descriptive statistics
In total, data on up to 127,632 individuals were available for analysis, including 58,176 never smokers, 37,428 former smokers and 32,028 current smokers. Overall, 45% of the combined study population was male. The median age within the contributing studies ranged between 16 and 68 years. The mean prevalence of depression, anxiety and psychological distress (using case definition 1) was 12.5% (range 6.1–37.5%), 10.2% (range 2.6–19.9%) and 17.4% (range 8.9–27%), respectively. Descriptive statistics for each of the study populations are provided in the online supplementary table S3.

The minor allele frequency for rs16969968/rs1051730 ranged between 0.31 and 0.39 (see online supplementary table S4). There was no strong evidence for deviation from HWE in any of the studies (p values all ≥0.06).

Observational analysis
Levels of depression, anxiety and psychological distress differed by smoking status (see figure 2). In age-adjusted and sex-adjusted analyses, current smokers had 1.85 times (95% CI 1.65 to 2.07, p<0.001) greater odds of depression, 1.71 times (95% CI 1.54 to 1.90, p<0.001) greater odds of anxiety and 1.69 times (95% CI 1.56 to 1.83, p<0.001) greater odds of psychological distress than never smokers. Former smokers had 1.22 times (95% CI 1.14 to 1.30, p<0.001) greater odds of depression, 1.23 times (95% CI 1.12 to 1.36, p<0.001) greater odds of anxiety and 1.17 times (95% CI 1.11 to 1.25, p<0.001) greater odds of psychological distress than never smokers.

Among smokers, smoking heaviness was positively associated with levels of depression, anxiety and psychological distress (figure 3). In age-adjusted and sex-adjusted analyses, a one cigarette per day increase in smoking heaviness was associated with a 1.03-fold (95% CI 1.02 to 1.04, p<0.001) increase in the odds of having depression, a 1.03-fold (95% CI 1.02 to 1.04, p<0.001) increase in the odds of having anxiety and a 1.02-fold (95% CI 1.02 to 1.03, p<0.001) increase in the odds of having psychological distress.

As there was evidence of between-study heterogeneity for analyses of both smoking status and smoking heaviness, random effects meta-analyses are presented. However, results from fixed effects meta-analyses were similar (data not shown). Individual study estimates for observational analyses are provided in online supplementary figures S1 and S2.

Mendelian randomisation analysis
There was no clear evidence that rs16969968/rs1051730 was associated with binary measures of depression in never (OR per minor allele 1.02, 95% CI 0.97 to 1.06, p=0.47), former (OR 1.00, 95% CI 0.95 to 1.05, p=0.99), current (OR 1.00, 95% CI 0.95 to 1.05, p=0.99) or ever (OR 1.01, 95% CI 0.98 to 1.05, p=0.58) smokers (see figure 3). Similarly, there was no clear evidence that rs16969968/rs1051730 was associated with binary measures of anxiety in former (OR 1.02, 95% CI 0.97 to 1.08, p=0.44), current (OR 1.02, 95% CI 0.97 to 1.07, p=0.42) or ever (OR 1.03, 95% CI 0.99 to 1.07, p=0.16) smokers. However, in never smokers there was some evidence of a positive association between the minor allele of rs16969968/rs1051730 and anxiety (OR 1.05, 95% CI 1.01 to 1.10, p=0.03). For psychological distress there was a similar pattern, with no strong evidence for an association between rs16969968/rs1051730 in smokers, but some evidence of a positive association in never smokers.
For all outcomes, there was no clear statistical evidence for a difference in the effect of rs16969968/rs1051730 between never, former and current smokers (p values for heterogeneity between never, former and current smokers from Cochran Q test all >0.57). Individual study estimates for observational analyses are provided in online supplementary figure S3.

Results were similar for continuous measures of symptoms of depression, anxiety and psychological distress (see online supplementary figures S4 and S5). There was no clear evidence for associations of rs16969968/rs1051730 with continuous outcomes in any of the smoking categories.

Finally, there was no clear evidence for sex differences in either observational or Mendelian randomisation analyses for associations between smoking or smoking-related genotype and depression or anxiety (data available on request).

**DISCUSSION**

In the largest Mendelian randomisation study on the association of smoking with depression and anxiety conducted to date, we found no evidence to suggest that smoking causes either depression or anxiety. Despite higher levels of depression, anxiety and psychological distress in current and former smokers compared to never smokers, and a positive association between the number of cigarettes smoked per day and depression and anxiety, there was no clear evidence for associations between the CHRNA5-A3-B4 variant and these outcomes in smokers. If heavier smoking were to cause depression or anxiety, we would expect to see an increased risk of depression or anxiety per copy of the minor allele of rs16969968/rs1051730, which increases smoking heaviness, in current smokers and potentially also in former smokers, but no difference in risk for never smokers. In our meta-analyses, ORs for the effect of rs16969968/rs1051730 in current, former and ever smokers were all close to the null, with the CIs for these estimates all overlapping the null. In addition, we found no evidence to suggest that the variant was differentially associated with depression or anxiety according to smoking status.

Our results are consistent with those of the two previous Mendelian randomisation studies, which did not find evidence that smoking increases depression or anxiety or ante-natal depression. Both of these studies were included in this meta-analysis and the HUNT study made up more than half of the study sample in some analyses. However, exclusion of either of these samples did not make a substantial difference to effect estimates (see online supplementary figures S6 and S7). These findings suggest that previous findings linking smoking to higher levels of depression and anxiety may be due to residual confounding, a shared vulnerability to both mental disorders and...
The rs16969968/rs1051730 variant associates with smoking heaviness within smokers but is not an instrument for smoking status (ever smoking vs never smoking). Therefore we cannot rule out the possibility that being a smoker, rather than smoking heaviness, could influence likelihood of depression or anxiety. However, we do see an observational association between smoking heaviness and depression and anxiety (figure 3), and a dose-dependent relationship between an exposure and an outcome strengthens support for a causal association. We might therefore expect to observe an association between rs16969968/rs1051730 and depression or anxiety if smoking were to cause these conditions. Furthermore, while rs16969968/rs1051730 is the strongest genetic contributor to smoking behaviour identified to date, this variant only explains a fraction of the estimated 50% of total variation in smoking behaviour within a population at any one time that is due to genetic factors. Further signals for smoking heaviness have been identified in the same gene cluster, in other nicotinic receptor units and in other genes, such as those related to nicotine metabolism like CYP2A6. Combining these variants in genetic risk scores for smoking behaviour in Mendelian randomisation studies will be an important future direction for validation of these results.

We would not expect to see an effect of rs16969968/rs1051730 on depression or anxiety in never smokers, because the variant is not associated with smoking heaviness within these individuals. Thus, this group can be used to test potential bias due to pleiotropy (that the gene affects more than one exposure) in Mendelian randomisation analyses. We did observe some evidence for an association between the variant and anxiety in never smokers, a finding previously reported by the HUNT study. Removal of HUNT from this analysis did not affect the point estimate, suggesting that this association is not driven solely by the data from this study (see online supplementary figure S6). However, using case definition 2 (where available) for anxiety in preference to case definition 1 slightly attenuated this association in never smokers (see online supplementary figure S8).

While this may be a chance finding, it is possible that rs16969968/rs1051730 or a variant in linkage disequilibrium with this variant, may affect anxiety directly, not through tobacco consumption. There is some suggestion from animal studies that nicotinic acetylcholine receptors may play a role in anxiety (eg, mice lacking the α4 subunit show increases in anxiety-related behaviour). However, there is currently little evidence for this association in humans, and rs16969968/rs1051730 has not been identified in genome-wide association studies of depression or anxiety to date.

It is important to note that stratifying by the measured exposure variable in Mendelian randomisation studies can lead to collider bias. In this specific analysis, if both the genetic variant and anxiety cause individuals to smoke, then stratifying on smoking could, in theory,
induce an association between the variant and anxiety.13 We do not think that collider bias is likely to be a major issue in these analyses because rs16969968/rs1051730 does not appear to be associated with smoking initiation in this sample (see online supplementary figure S9) or in previous studies.29 30 There is, however, as reported previously in a few specific populations,31 62 63 some evidence that the minor allele of rs16969968/rs1051730 is associated with smoking cessation; ORs of being a current compared to a former smoker were 1.08-fold higher per copy of the minor allele (95% CI 1.06 to 1.11) in this sample.

Strengths and limitations
The key strength of this study is the large sample size, using data on over 125 000 individuals from 25 different populations. Despite this, we did not have the power to rule out the possibility of a causal effect. A substantial increase in sample size would be required to be confident that what we observe is a true null association in smokers. We hope that our estimates may be combined with those of further studies addressing the same question in future meta-analyses, to provide more definitive answers.

One of the main limitations is the use of broad definitions of depression and anxiety rather than clinical definitions, which were not available in all studies. It is possible that use of more precise phenotypic measures of depression and anxiety based solely on clinical criteria could yield stronger results because non-differential misclassification of a binary outcome is likely to attenuate associations towards the null.2 64 However, we showed the expected observational associations between smoking and depression, anxiety and psychological distress. In addition, we used two case definitions and performed a sensitivity analysis using case definition 2 in preference (where both were available) which produced similar results (see online supplementary figure S8). Sensitivity analyses performed excluding lifetime definitions of depression or anxiety also produced similar results (see online supplementary figure S10). Finally, restricting our analyses to those studies with questionnaire based on clinical criteria or self-report of doctor diagnosis produced consistent results, although these analyses were underpowered (see online supplementary figure S11).

Although we analysed depression and anxiety separately, these conditions are highly comorbid in the general population65 66 and symptom scale questionnaires are not adequate to distinguish between them.67 68 In addition, the definition of anxiety we used encompassed general anxiety disorder, panic and phobias. It is possible that these conditions have different aetiologies.9 Therefore, we cannot make inferences about specific anxiety disorders from these results. Furthermore, the sample encompasses a wide age range, so it is unlikely that this analysis would be able to capture any age-specific effects of smoking on depression and anxiety.

CONCLUSION
In conclusion, our Mendelian randomisation analyses do not support a causal role of smoking heaviness among smokers in the development of depression and anxiety. While we cannot directly address the question of whether smoking initiation (ie, starting smoking) plays a causal role in relation to these outcomes, we expect that if it did we would also see a dose-dependent relationship between smoking heaviness and depression and anxiety. We see such an association in our observational analyses, but no strong evidence for this in our Mendelian randomisation analyses. Future research should focus on the possible role of depression and anxiety in increasing susceptibility to smoking. As larger genome-wide association studies of depression and anxiety emerge, it is likely that genetic variants will be identified that can be utilised in Mendelian randomisation analyses for this purpose.

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Contributors MRM and GDS conceived the study. AET and MEF drafted the analysis protocol and co-ordinated the data analysis. AET conducted the final meta-analyses. AET and MEF contributed to writing the final version of the manuscript. MEF and GDS performed the meta-analysis. All authors commented on a draft of the manuscript and approved the final version.

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Competing interests MRM: Medical Research Council, LTL: British Heart Foundation, LJH: New Zealand Health Research Council, JK: Jim and Mary Carney Charitable Trust, New Zealand Health Research Council, CCM: European Research Council, DIB, JVM: NWO, European Research Council, National Institutes of Health, HT: Dutch Medical Research Council, TP: Canadian Institutes of Health Research, MK: Medical Research Council, NIH: Lung, Lung and Blood Institute and Economic and Social Research Council); financial relationships with any organisations that might have an interest in the submitted work in the previous 3 years (LJH: Australian National Health and Medical Research Council, PMV: Swiss National Science Foundation, MP: Eli Lilly and Lundbeck advisory board membership, Swiss National Science Foundation, TK: Pfizer consultancy on tobacco dependence, Juho Vainio Foundation, The Finnish Medical Society, TSA: Gene Diet Interactions in Obesity project, HT: KOC University, Pfizer), YBS: grants from NIHR, MRC, POLUK, BHF, AET, MRM and MEF are members of the UK Centre for Tobacco and Alcohol Studies, a UKCRC Public Health Research Centre of Excellence. Funding from British Heart Foundation, Cancer Research UK, Economic and Social Research Council, Medical Research Council, and the National Institute for Health Research, under the auspices of the UK Clinical Research Collaboration is gratefully acknowledged. LTL is supported by BHF programme grant RG/08/013/25942. LP is funded by an MRC Population Health Science Fellowship (MR/J012165/1). TSA was supported by the Gene Diet Interactions in Obesity (GENDINOB, www.gendinob.dk) postdoctoral fellowship grant. LLNH was supported by the Health Insurance Foundation (grant No. 2010 B 131). The work of SSM was funded by the Netherlands Genomics Initiative (NGL)/Netherlands Organization for Scientific Research (NWO) (grant-numbers 050-060-810 NCHA); the work of HT was supported by the EU FP7 NWO-VID1 (grant 017-106-370). LTL is supported financially by the Economic and Social Research Council International Centre for Life Course Studies in Society and Health (RES-596-28-0001).

Patient consent Obtained.

Ethics approval 1958BC: Ethics approval for the study was obtained from the South-East Multi-Centre Research Ethics Committee (Ref: 01/144) and the Joint UCL/UCLH Committees on the Ethics of Human Research (Committee A) Ref: 08/H0714/40. ALSpac: Ethics approval for the study was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committee. BRHS: The BRHS has local (from each of the districts in which the study was based) and multicentre ethical committee approvals. Ethical approval was granted for the BWIHSs from the London Multi-Centre Research Ethics Committee and 23 Local Research Ethics Committees. BWHHs: Ethical approval was granted for the BWIHSs from the London Multi-Centre Research Ethics Committee and 23 Local Research Ethics Committees. CaPS: Ethics approval was obtained from the South Glamorgan Area Health Authority, the Gwent REC, and the South Wales Research Ethics Committee D. CHDS: All phases of the study have received ethical approval from the regional and Country Ethics Committees. The ethics of this study were approved by the Institutional Ethics Committee of the University of Lausanne. ELSA: ELSA has been approved by the National Research Service. FINRISK: The 2002 and 2007 FINRISK surveys have been approved by the Coordinating Ethics Committee of the Helsinki University Hospital District. Generation Scotland: Ethical approval for the study was given by the NHS Tayside committee on research ethics (reference 05/s1401/89). GOYA females: The study was approved by the regional scientific ethics committee and by the Danish Data Protection Board. HBCS: The research plan of the HBCS was approved by the Institutional Review Board of the National Public Health Institute. Health2006/Health2008/Inter99: The studies have been approved by the Ethical Committee of Copenhagen. HUNT: Use of data in the present study was approved by the Regional Committee for Medical Research Ethics (Reference nr. 2013/1127/REK midt). NFBC: The University of Oulu Ethics Committee and the Ethical Committee of Northern Ostrobothnia Hospital District have approved the study. NIHANES: Data collection for NIHANES was approved by the NCHS Research Ethics Review Board. Analysis of de-identified data from the survey is exempt from the federal regulations for the protection of human research participants. Analysis of restricted data of de-identified data from the survey is exempt from the federal regulations for the protection of human research participants. Analysis of restricted data.
residence of the patients. Rotterdam: The Medical Ethics Committee of Erasmus Medical Center Rotterdam approved the study. SYS-P: The Research Ethics Committee of the Chicoutimi Hospital in Quebec, Canada approved the study protocol. Whitehall: Ethical approval for the Whitehall II study was obtained from the University College London Medical School committee on the ethics of human research.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement 1958BC: This study makes use of data generated by the Wellcome Trust Case-Control Consortium. A full list of investigators who contributed to generation of the data is available from the Wellcome Trust Case-Control Consortium website. The 1958 birth cohort data can be accessed via the UK Data Service (http://ukdataservice.ac.uk/). ALSPAC: Data used for this submission will be made available on request to the ALSPAC executive committee (alspac-exec@bristol.ac.uk). The ALSPAC data management plan (available here: http://www.bristol.ac.uk/alspac/researchers/data-access/) describes in detail the policy regarding data sharing, which is through a system of managed open access. BRHS: We welcome proposals for collaborative projects and data sharing (http://www.ucl.ac.uk/pcph/research-groups-themes/brhs-pub). For general data sharing enquiries, please contact Lucy Lennon (l.lennon@ucl.ac.uk). BHWS: All BHWS data collected is held by the research team based at London School of Hygiene and Tropical Medicine, for ongoing analysis. If you would like to collaborate with the BHWS team, contact the study coordinator, AA (antonette.amuzuz@lshtm.ac.uk). Data and biological samples provided to the collaborators can only be used for the purposes originally stated and must not be used in any other way without re-application to the BHWS team. No data should be passed to any third party unless they were stipulated in the original application. CaPS: Data used for the Caerphilly Prospective Study (CaPS) was made available by the CaPS access committee (Chair: Professor Kay Tew Khaw). More information about its managed access procedure is available on the study website (http://www.brisc.ac.uk/social-community-medicine/people/project/1392). CHDS: Data contributed for this submission are available on request from the CHDS (john.horwood@otago.ac.nz). Colaus/PsyCoLaus: Data from the Colaus/PsyCoLaus study can be requested according to the procedure described on the CoLaus website (http://www.colaus.ch/en/cls_home/cls_pro_home/cls-research-3.htm). ELSA: ELSA data are made available through the EDSOS website (http://www.elsa-project.ac.uk/availableData). FINRISK: Data used for this submission will be made available on request to the FINRISK Management Group, according to the given ethical guidelines and Finnish legislation. Generation Scotland: Data is available on request (access@generationscotland.org). GOYA females: An anonymized copy of the data used for this submission will be made available on request to the GOYA analysts after permission have been given by the DNBC executive committee (www.dnbc.dk). HBCS: Data used for this submission will be made available on request to the HBCS executive committee (johan.eriksson@helsinki.fi). Health2006/Health2008/Inter99: Data used for this submission can be made available on request to the Research Centre for Prevention and Health (http://www.region.dk/rcfsm/Menu/). Please contact LLNH (lis.lotte.nystrup.husemoen@regionh.dk) or AL (allan.linneberg@regionh.dk). HUNT: Data used from the HUNT Study for this submission will be made available on request to the HUNT Data Access Committee (hunt@medisin.ntnu.no). The HUNT data access information (http://www.ntnu.edu/hunt/data) describes in detail the policy regarding data availability. NFBC: Data used for this submission can be made available on request to Tuula Ylitalo (tuula.ylitalo@oulu.fi), Minna Mannikko (minna.mannikko@oulu.fi) or M-RJ (m.jarvelin@imperial.ac.uk). NHANES: NHANES data can be accessed here: (http://www.cdc.gov/nchs/nhanes.htm). The genotype used in this analysis is a restricted variable. Applications for access to these data must be made through the Research Data Center: (http://www.cdc.gov/rdc/). NSHD: The NSHD data are made available to researchers who submit data requests (tomcricha.switinfo@ucl.ac.uk). More information is available in the full policy documents (http://www.nshd.mrc.ac.uk/data.aspx). Managed access is in place for this study to ensure that use of the data are within the bounds of consent given previously by participants, and to safeguard any potential threat to anonymity since the participants are all born in the same week. NTR: Data used for this submission will be made available on request to the NTR committee (ntr@psy.vu.nl). SYS-P: Data used for this submission will be made available on request to the SYS principal investigators (http://www.saguenay-youth-study.org) (scientific community/collaborative interests). Whitehall: Data from the Whitehall II study are made publicly available as described in the Whitehall II data sharing policy (http://www.ucl.ac.uk/whitehallII/datasharing).

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