Examining the Lancet Commission risk factors for dementia using Mendelian randomisation

Roopal Desai,1 Amber John,1 Rob Saunders,2 Natalie L Marchant,3 Joshua E J Buckman,2 Georgina Charlesworth,1 Verena Zuber,4,5 Joshua Stott 1

ABSTRACT

Background Dementia incidence is increasing across the globe and currently there are no disease-modifying pharmaceutical treatments. The Lancet Commission on dementia identified 12 modifiable risk factors which explain 40% of dementia incidence. However, whether these associations are causal in nature is unclear.

Objective To examine the modifiable risk factors for dementia as identified in the Lancet Commission review using Mendelian randomisation (MR) to establish if, based on genetic evidence, these associations with different dementia subtypes are causal in nature.

Methods Publicly available genome-wide association study data were used for 10 risk factors and Alzheimer’s disease (AD), frontotemporal dementia and dementia with Lewy bodies. Two-sample MR using the inverse variance weighted method was conducted to test for causal relationships. Weighted median MR and MR-Egger were used to test for pleiotropic effects.

Results Genetic proxied risk for higher levels of smoking (OR: 0.80 (95% CI: 0.69; 0.92), p=0.002), obesity (OR: 0.87 (95% CI: 0.82; 0.92), p<0.001) and blood pressure (OR: 0.90 (95% CI: 0.82; 0.99), p=0.035) appeared to be protective against the risk of AD. Post hoc analyses indicated these associations had pleiotropic effects with the risk of coronary artery disease. Genetic proxied risk of educational attainment was found to be inconsistently associated with the risk of AD.

Conclusions and implications Post hoc analysis indicated that the apparent protective effects of smoking, obesity and blood pressure were a result of survivor bias. The findings from this study did not support those presented by the Lancet Commission. Evidence from causal inference studies should be considered alongside evidence from epidemiological studies and incorporated into reviews of the literature.

INTRODUCTION

Dementia incidence and prevalence rates are steadily increasing across the globe as lifespans increase and the global population ages.1 The quest for pharmaceutical interventions that prevent or modify dementia has thus far proved unsuccessful. Therefore, the spotlight of attention has fallen on lifestyle risk factors, the modification of which may have a role in delaying or preventing dementia. To this end, the Lancet Commission report on Dementia prevention, intervention and care2 has provided the most comprehensively reviewed evidence for potentially modifiable risk factors for dementia. The report highlighted the importance of nine risk factors which were categorised as falling under early-life, mid-life and late-life.3 Less education was identified as an early-life risk factor (<45 years); hearing loss, hypertension and obesity as mid-life risk factors (age 45–65 years) and smoking, depression, physical inactivity, low social contact and diabetes as late-life risk factors (age >65 years). This was further extended to 12 by including the mid-life risk factor of traumatic brain injury and excessive alcohol consumption and the later-life risk factors of air pollution.4 A population attributable fraction (PAF) for each risk factor was calculated, and the authors report that together these 12 risk factors have a combined PAF of approximately 40%. This can be interpreted as: if a causal relationship between the risk factors and dementia...
exists, then elimination of these risk factors would result in a 40% reduction in the number of dementia cases. The report used the most robust information available, pooling data from epidemiological studies and randomised controlled studies (RCTs). However, it was limited due to assumptions underlying the PAF calculation and the use of all-cause dementia as the outcome.

A key assumption in the PAF calculation and interpretation is that the relationship between the risk factor and outcome is causal. However, although a mixture of study designs examining risk factors was included in the Lancet Commission review, the bulk of the evidence comes from epidemiological studies. A limitation to reporting PAF statistics based on epidemiological studies is that causal relationships cannot be established in these types of observational studies. One reason for this is that observed associations can arise as an artefact of unmeasured confounders. Another reason is that reverse causation cannot be ruled out and this may be particularly problematic when the disease in question has a long preclinical period as is the case with dementia. Therefore, one of the key messages of the Lancet commission report (modifying the 12 lifestyle risk factors may prevent or delay up to 40% of dementias in the population) is based on the assumption that the results of the studies included in the report have provided evidence to support causality.

The second limitation is the use of all-cause dementia as an outcome. Although there are benefits to using all-cause dementia as a generic outcome in terms of power and maximising the number of included studies, a limitation is that each subtype of dementia is a different disorder with differing underlying pathologies. As such it seems likely that each subtype of dementia may be differentially affected by a particular risk factor. For example, evidence from meta-analyses suggests that obesity may pose a greater risk for the development of Alzheimer’s disease (AD) than for vascular dementia. These potentially differential effects could not be accounted for in the Lancet Commission report.

One way to address the first of these limitations is by using Mendelian randomisation (MR). MR is a statistical technique that combines genetic data with epidemiological data to overcome the limitations of confounders and reverse causation and is considered analogous to the RCT. Within the population, some individuals are more at risk of experiencing certain exposures. For example, some people may be at greater risk of obesity by virtue of their genetic make-up. This risk is ‘allocated’ by genetic meiosis and determined at conception and therefore this natural assortment of risk alleles is considered equivalent to the random allocation process of the RCT. Although population stratification and assortative mating may have an impact on human DNA, reverse causality can be largely ruled out because it is unlikely that the exposure individuals experience throughout the course of their lives will have an impact on their genes. Thus, MR studies can be used to estimate the causal effect of the exposure on the outcome given three assumptions are met. These assumptions underpinning MR analysis are the following: the genetic instrument is associated with the exposure phenotype; the genetic instrument is not associated with any confounders; and the genetic instrument exerts its impact on the outcome via the exposure. Given these assumptions are met, MR studies can be useful tools in determining and providing evidence of causal relationships.

To date, evidence coming from MR studies investigating the relationship between risk factors and dementia has been mixed. One potential reason for the mixed findings is the discordant application of instrument selection across studies and subsequent analysis pipelines. Therefore, we aim to conduct a comprehensive MR analysis of as many as possible of the modifiable risk factors for dementia as identified by the Lancet Commission using a standardised rigorous pipeline to allow for the comparability of results. In addition, to address the limitation of using all-cause dementia as a generic outcome we aim to conduct the analysis for three different subtypes of dementia: AD dementia, dementia with Lewy bodies (DLB) and frontotemporal dementia (FTD), for which genome-wide association study (GWAS) data exist. We aim to use the most recent GWAS releases based on the largest sample sizes thus creating the most up-to-date and comprehensive MR analysis on risk factors for dementia.

METHOD
Study design
GWAS data were available for 10 out of the 12 Lancet Commission risk factors for dementia, and data were not available for traumatic brain injury or pollution. Two-sample Mendelian randomisation analysis was used to investigate potential causal relationships between 10 of the Lancet Commission risk factors for all-cause dementia and 3 different subtypes of dementia: AD (using 4 different GWAS datasets), DLB and FTD. We followed the reporting guidelines provided by the Strengthening the Reporting of Observational Studies in Epidemiology—Mendelian Randomisation.

Data sources
The current study used publicly available summary statistics and therefore no additional ethics or approvals were required. We searched the GWAS catalogue (https://www.ebi.ac.uk/gwas/) to find the latest GWAS studies for dementia outcomes and the Lancet Commission risk factors for dementia. Publicly available GWAS summary statistics were obtained for each of the 10 risk factors and the 3 dementia-related outcomes (table 1).

MENDELIAN RANDOMISATION ANALYSIS

Genetic instruments
For each of the 10 risk factors, single nucleotide polymorphisms (SNPs) were selected based on the genome-wide significance threshold of $p < 5 \times 10^{-8}$. The exposure dataset was merged with the outcome dataset by aligning the effect alleles to the forward strand. The overall $F$ statistic for each exposure was calculated and used as a measure of instrument strength. Only instruments with an $F$ statistic value $> 10$ were used in the analyses to minimise any effects of weak instrument bias. Once the SNPs which met the threshold for genome-wide significance were identified, these SNPs were clumped to remove SNPs in linkage disequilibrium using a threshold of $r^2 < 0.01$ and pruned by selecting the SNP which had the lowest $p$ value. The association between each individual SNP and the dementia outcome was calculated using the Wald ratio, and these individual estimates were combined to provide an overall estimate using the inverse variance weighting (IVW) method. To test for potential pleiotropic effects, a sensitivity analysis using MR-Egger and weighted median MR was conducted. Cochran’s $Q$ statistic was used to estimate heterogeneity and as an additional test of horizontal pleiotropy.

As 10 tests were conducted per dementia outcome, a Bonferroni-corrected $p$ value was used and set at $p < 0.005$ to account for multiple comparisons. All estimates were calculated using standardised betas (log OR) and then transformed to the OR. For risk factors measured on a continuous scale, ORs estimated the increased risk per 1 SD increment in the risk factor.

Post hoc analyses were carried out to test a further hypothesis that the effects of obesity, blood pressure and smoking on AD were a result of survivor bias. Multivariable MR was
Table 1  GWAS datasets used in the study

<table>
<thead>
<tr>
<th>Phenotypes</th>
<th>Study/dataset</th>
<th>Cohort/consortium</th>
<th>Maximum number of participants</th>
<th>Underlying population</th>
<th>Phenotype assessment</th>
<th>Summary data</th>
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<tbody>
<tr>
<td><strong>Early-life exposure</strong></td>
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<tr>
<td>Education</td>
<td>Okbay et al, 201624,25</td>
<td>64 separate cohorts</td>
<td>293723</td>
<td>European</td>
<td>Educational attainment was measured by the number of years of schooling completed and assessed in cohorts of participants aged 30 years or above.</td>
<td>Mean=14.3 years (SD=3.6)</td>
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<tr>
<td><strong>Mid-life exposures</strong></td>
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<tr>
<td>Alcohol consumption</td>
<td>Evangelou et al, 201916,27</td>
<td>UK Biobank, Alcohol Genome-Wide Consortium and Cohorts for Heart and Ageing Research in Genomic Epidemiology Plus consortia</td>
<td>480842</td>
<td>European</td>
<td>Self-reported weekly and monthly alcohol consumption.</td>
<td>Mean=19.8 g of alcohol per day (SD=21.2)</td>
</tr>
<tr>
<td>Blood pressure (systolic)</td>
<td>UK Biobank</td>
<td>UK Biobank</td>
<td>475142</td>
<td>European</td>
<td>Average of automated readings of systolic blood pressure.</td>
<td>Mean=138.0 mm Hg (SD=19.3)</td>
</tr>
<tr>
<td>Age-related hearing loss</td>
<td>Kalra et al, 202019,30</td>
<td>UK Biobank</td>
<td>330759</td>
<td>European</td>
<td>Self-reported four hearing loss traits: 1. Do you have difficulty with your hearing? 2. Do you find it difficult to follow a conversation if there is background noise (such as TV, radio, children playing)? 3. Do you use a hearing aid most of the time? 4. Do you get or have you had noises (such as ringing or buzzing in your head or in one or both ears that lasts for more than 5 min at a time?</td>
<td>Trait 1: n=23 333 (4.7%) Trait 2: n=123 089 (37.8%) Trait 3: n=10 322 (5.3%) Trait 4: n=7214 (7.1%)</td>
</tr>
<tr>
<td>Obesity</td>
<td>Pulit et al, 201931,32</td>
<td>UK Biobank</td>
<td>694649</td>
<td>European</td>
<td>Body fat distribution in the form of measurements of waist:hip ratio adjusted for body mass index.</td>
<td>SD=0.0199</td>
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<tr>
<td><strong>Late-life exposures</strong></td>
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<tr>
<td>Depression</td>
<td>Howard et al, 201833,34</td>
<td>UK Biobank</td>
<td>322580</td>
<td>European</td>
<td>Depression categorised into three groups: 1. Broad depression: measured by self-reported help-seeking behaviour for ‘nerves, anxiety, tension or depression’. 2. Probable major depression: measured by self-reported depressive symptoms with ‘impairment’. 3. Major depression: assessed by hospital admission records.</td>
<td>Cases, n=2 676 (16.8%)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>Scott et al, 201735,36</td>
<td>Diabetes Genetics Replication and Meta-analysis Consortium</td>
<td>159208</td>
<td>European</td>
<td>Type 2 diabetes assessed based on the prescription medication or laboratory evidence of hyperglycaemia.</td>
<td>Cases, n=26 766 (16.8%)</td>
</tr>
<tr>
<td>Loneliness</td>
<td>Day et al, 201837,38</td>
<td>UK Biobank</td>
<td>452302</td>
<td>European</td>
<td>Self-reported traits: (1) Do you often feel lonely? (2) Composite of: including yourself, how many people are living together in your household? And: How often do you visit friends or family or have them visit you? (3) How often are you able to confide in someone close to you?</td>
<td>Trait 1: n=80 134 (18.0%) Trait 2: n=2 426 (0.8%) Trait 3: n=64 506 (21.3%)</td>
</tr>
<tr>
<td>Physical activity</td>
<td>Klimentidis et al, 201839,40</td>
<td>UK Biobank</td>
<td>377234</td>
<td>European</td>
<td>Self-reported activity levels and accelerometer readings.</td>
<td>Vigorous physical activity ≥3 vs 0 days/week, n=98 060 (37.6%)</td>
</tr>
<tr>
<td>Smoking</td>
<td>Wootton et al, 201931,42</td>
<td>UK Biobank</td>
<td>462690</td>
<td>European</td>
<td>Self-reported measures of lifetime smoking including smoking duration, heaviness and cessation.</td>
<td>Mean=0.359, lifetime smoking score SD=0.694</td>
</tr>
<tr>
<td><strong>Outcomes</strong></td>
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<tr>
<td>Late-onset AD (clinical diagnosis and family history)</td>
<td>Bellenguez et al, 202215,43</td>
<td>European Alzheimer &amp; Dementia Biobank</td>
<td>788989</td>
<td>European</td>
<td>Clinical diagnosis of AD plus self-report of at least one first-degree biological relative with a diagnosis of AD.</td>
<td>Cases, n=1 112 326 (14.1%)</td>
</tr>
<tr>
<td>Late-onset AD</td>
<td>Kunkle et al, 201912,44</td>
<td>International Genomics of Alzheimer’s Project</td>
<td>94437</td>
<td>European</td>
<td>Clinical diagnosis or evidenced by autopsy reports.</td>
<td>Cases, n=35 274 (37.4%)</td>
</tr>
<tr>
<td>Late-onset AD</td>
<td>Lambert et al, 201311,45</td>
<td>International Genomics of Alzheimer’s Project</td>
<td>74046</td>
<td>European</td>
<td>Clinical diagnosis or evidenced by autopsy reports.</td>
<td>Cases, n=17 008 (31.4%)</td>
</tr>
</tbody>
</table>

Continued
**Sensitivity analyses**
A sensitivity analysis for each of the 10 risk factors was conducted using 3 additional GWAS datasets for AD. The first of these was a GWAS dataset published by Kunkle et al.\(^1\) and the second was a dataset published by Lambert et al.\(^1\) The third dataset was the GWAS published by Schwartzentruber et al.,\(^1\) which combines existing GWAS data on AD with a proxy measure of familial genetic risk.

The main analysis was conducted using data from the largest GWAS on AD and comprised a sample size of 788,989 participants.\(^1\) This dataset offered the greatest power to detect causal effects; however, the inclusion of a proxy diagnosis of AD may be vulnerable to disease misclassification and thereby add additional heterogeneity to the results. The Kunkle et al.\(^1\) and Lambert et al.\(^1\) datasets were used in the sensitivity analysis; although comprising much smaller samples (n=94,437 and n=74,046, respectively), they both used more reliable methods of disease classification. Schwartzentruber et al.,\(^1\) had a sample size of 408,942 and relied on mostly proxy reports of family history of AD diagnosis was used as an additional dataset to establish the overall replicability of the results.

All analyses were conducted using RStudio (2020) V.1.3.1093 and the iegwasr and the MendelianRandomization (V.0.5.0)\(^1\) R packages. Code was available from: https://github.com/koop-hub/Examining-the-Lancet-Commission-Risk-Factors-for-Dementia-Using-Mendelian-Randomization.

**RESULTS**
Ten established risk factors for dementia as identified in the Lancet Commission report\(^7\) were investigated for their causal effect on three dementia outcomes using two-sample IVW MR. The number of SNPs for each exposure and the test of instrument strength (F statistic) are reported in figures 1–3.

**Alzheimer’s disease**
Genetically determined higher educational attainment was found to increase the risk of dementia (OR: 1.93 (95% CI: 1.03; 3.54), p=0.03). However, this effect did not meet the criteria for statistical significance after a Bonferroni correction was applied. The mid-life risk factor of genetically determined obesity was found to have a protective effect against the genetic risk of AD with the risk of AD reducing by 13% (OR: 0.87 (95% CI: 0.82; 0.92), p<0.001) with each standard increment in BMI-adjusted waist:hip ratio. The mid-life risk factor of genetically determined lifetime smoking was found to have a protective effect on the risk of AD with a 20% reduction in the risk of AD associated with 1 SD increase in lifetime smoking index (OR: 0.80 (95% CI: 0.69; 0.92), p=0.002). Genetically determined higher SBP was also found to be protective against the genetic risk of AD; however, the estimate did not meet the threshold for Bonferroni-corrected significance (OR: 0.90 (95% CI: 0.82; 0.99), p=0.035).

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**Table 1 Continued**

<table>
<thead>
<tr>
<th>Phenotypes</th>
<th>Study/dataset</th>
<th>Cohort/consortium</th>
<th>Underlying population</th>
<th>Phenotype assessment</th>
<th>Summary data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum number of participants</td>
<td>EU Biobank</td>
<td>408,942</td>
<td>European</td>
<td>Number of first-degree relatives with a diagnosis of AD.</td>
<td>Cases, n=53,042 (13.0%)</td>
</tr>
<tr>
<td>Dementia-Using-Mendelian-Randomization.</td>
<td>UK Biobank</td>
<td>942</td>
<td>European</td>
<td>Number of first-degree relatives with a diagnosis of AD.</td>
<td>Cases, n=2,391 (39.2%)</td>
</tr>
<tr>
<td></td>
<td>Chia et al., 2021</td>
<td>6618</td>
<td>European</td>
<td>Consensus clinical diagnosis based on established criteria.</td>
<td>Cases, n=2591 (39.2%)</td>
</tr>
<tr>
<td></td>
<td>Ferrari et al., 2014</td>
<td>47,46</td>
<td>European</td>
<td>Cohort recruited from 44 international research groups</td>
<td>Cases, n=17,283 (11.1%)</td>
</tr>
<tr>
<td></td>
<td>Nelson et al., 2017</td>
<td>155,437</td>
<td>European</td>
<td>Self-reported angina or other evidence of chronic coronary heart disease including myocardial infarction and/or revascularisation.</td>
<td>Cases, n=74,046 (39.2%)</td>
</tr>
<tr>
<td></td>
<td>Cohen et al., 2018</td>
<td>12,46</td>
<td>European</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Chia et al., 2021</td>
<td>15,46</td>
<td>European</td>
<td></td>
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<tr>
<td></td>
<td>Ferreira et al., 2014</td>
<td>12,46</td>
<td>European</td>
<td></td>
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<tr>
<td></td>
<td>Nelson et al., 2017</td>
<td>155,437</td>
<td>European</td>
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None of the other risk factors investigated in the current study were found to have a significant effect on the genetic risk of AD (Figure 1). Pleiotropy was investigated by conducting MR-Egger analyses. The pattern of results was similar to those attained from the IVW method (online supplemental file 1).

Sensitivity analyses
For the early-life risk factor of education, contrary to the results from the main analysis the two datasets which measured AD using clinical diagnosis found greater education to be protective against the risk of AD. For the mid-life risk factor of obesity, in keeping with the results from the main analysis, the results from all three additional datasets replicated the protective direction of effect. For the mid-life risk factor of SBP, in keeping with the main analysis all three additional datasets indicated a protective direction of effect. In the case of the late-life risk factor of smoking, the results were mixed.

MVMR analyses
The effects of obesity, smoking and blood pressure were found to be in a direction contrary to the findings reported in the Lancet Commission. That is, the genetic risk of higher obesity, smoking and blood pressure were protective factors against the genetic risk of AD. Given these results, post hoc analyses were conducted using a multivariable MR approach to ascertain if these findings were driven by the pleiotropic effect of CAD. The multivariable MR analyses indicated that the apparent protective effect of all three of these risk factors was driven by the pleiotropic effect of CAD. When CAD was added to the model, the effects of obesity, blood pressure and smoking were attenuated. In the case of blood pressure (OR: 0.95 (95% CI: 0.84; 1.06), p=0.36) and smoking (OR: 0.90 (95% CI: 0.77; 1.06), p=0.20), these effects became non-significant. In the case of obesity (OR: 0.90 (95% CI: 0.84; 0.96), p=0.002) although the effect remained significant, it was greatly attenuated. This suggests that these apparent protective effects on AD are due to the pleiotropic effects of CAD.

Table: Risk Factor SNPs F Estimate [95% CI]

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>SNPs</th>
<th>F</th>
<th>Estimate [95% CI]</th>
</tr>
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<tbody>
<tr>
<td><strong>Early Life</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Education</td>
<td>67</td>
<td>45.5</td>
<td>1.99 [0.24, 16.23]</td>
</tr>
<tr>
<td><strong>Mid Life</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alcohol Consumption</td>
<td>69</td>
<td>58</td>
<td>0.60 [0.18, 2.03]</td>
</tr>
<tr>
<td>Blood Pressure</td>
<td>213</td>
<td>45.6</td>
<td>1.12 [0.82, 1.52]</td>
</tr>
<tr>
<td>Hearing Loss</td>
<td>29</td>
<td>43.4</td>
<td>0.75 [0.50, 1.18]</td>
</tr>
<tr>
<td>Obesity</td>
<td>538</td>
<td>55.5</td>
<td>0.75 [0.57, 0.99]</td>
</tr>
<tr>
<td><strong>Late Life</strong></td>
<td></td>
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</tr>
<tr>
<td>Depression</td>
<td>71</td>
<td>42.2</td>
<td>0.99 [0.64, 1.55]</td>
</tr>
<tr>
<td>Diabetes</td>
<td>38</td>
<td>62.5</td>
<td>1.02 [0.86, 1.22]</td>
</tr>
<tr>
<td>Loneliness</td>
<td>14</td>
<td>35</td>
<td>0.22 [0.03, 1.47]</td>
</tr>
<tr>
<td>Physical Activity</td>
<td>7</td>
<td>38</td>
<td>0.07 [0.00, 10.45]</td>
</tr>
<tr>
<td>Smoking</td>
<td>119</td>
<td>42.6</td>
<td>0.83 [0.49, 1.42]</td>
</tr>
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</table>

Figure 1 Forest plot showing inverse variance weighted causal effect estimates of modifiable risk factors on Alzheimer’s disease. The number of single nucleotide polymorphisms in each instrumental variable is reported with the corresponding F statistic.

Figure 2 Forest plot showing inverse variance weighted causal effect estimates of modifiable risk factors on dementia with Lewy bodies. The number of single nucleotide polymorphisms in each instrumental variable is reported with the corresponding F statistic.
Other dementias

Dementia with Lewy bodies

The MR analysis using the 10 exposure risk factors was repeated using DLB as the outcome. Neither educational attainment nor any of the other nine risk factors had a significant effect in relation to DLB (figure 2). Pleiotropy was investigated by conducting MR-Egger analyses. The pattern of results was similar to those attained from the IVW method (online supplemental file 1).

Frontotemporal dementia

The MR analysis using the 10 exposure risk factors was repeated using FTD as the outcome (figure 3). Pleiotropy was investigated by conducting MR-Egger analyses. The pattern of results was similar to those attained from the IVW method (online supplemental file 1).

DISCUSSION

This study aimed to examine 10 of the 12 risk factors for dementia as identified by the Lancet Commission review using MR and a rigorous pipeline analysis to establish if these associations are causal in nature. We used the most recent GWAS data for AD, which is based on the largest sample size. We found apparent protective effects of increased obesity, blood pressure and smoking. Additional post hoc analyses indicated that these apparent protective effects may be due to pleiotropic effects of CAD. There was inconsistency in the direction of effect for educational attainment dependent on the outcome of AD GWAS. None of the other risk factors were found to be significantly associated with the risk of AD. We did not find evidence that any of the 10 risk factors were significant causal risk factors for DLB or FTD.

Contrary to the findings of the Lancet Commission, there was evidence that increased obesity, blood pressure and levels of smoking were protective factors in the risk of developing AD. The direction of this effect was replicated using the three additional AD datasets. The results for blood pressure showed a similar pattern with estimates from all four AD datasets indicating a protective direction. In the case of smoking analyses from two out of the four datasets indicated a protective effect. In a post hoc analysis, we tested the hypothesis that these relationships were a result of survivor bias and driven through the pleiotropic pathway of CAD. Survivor bias has been highlighted as a threat to causal inference in MR studies. This is particularly of relevance where the outcome affects an older population as is the case for most dementias. The underlying threat coming from survivor bias in these types of studies is that those individuals who live long enough to develop an age-related disease are the ones who have not died earlier from other morbidities. Thus, what appears to be a link with dementia may be an association with longevity. To test this, we conducted a post hoc multivariable MR and found that the effects of smoking and blood pressure attenuated to a non-significant effect when conditioning on CAD. In the case of obesity although the effect remained significant, it was greatly reduced. This attenuation in effect supports the hypothesis that these results were due to survivor bias. Although we demonstrated survivor bias through the pleiotropic pathway of CAD, it should be noted that other potentially life-limiting diseases such as cancer may also have pleiotropic pathways between the risk factors in question and AD. In addition, alternative explanations for counterintuitive findings should not be discounted. For example, in the case of smoking there is evidence that smoking can be both a risk or a protective factor depending on the dementia type. Another factor that should be considered is whether the effects of demographic and indirect effects are adequately controlled for. Growing evidence suggests that GWAS studies from purportedly unrelated individuals do not completely exclude genetic variation due to demographic elements (eg, population stratification and assortative mating). This uncontrolled genetic variation can lead to a potential bias in the estimate of the direct effect between genetic variants and the phenotype in question. Within-sibship GWAS analyses more accurately account for demographic effects and should be used in future studies to further explore the associations between risk factors and dementia.

Educational attainment is well established from observational studies as having a protective role in dementia risk. However, our results were inconsistent with opposite effect directions found across different datasets. These inconsistent findings replicate a previous study which found that when proxy information on AD diagnosis is included in the GWAS, as in the Bellenguez et al (2022) data, then greater educational attainment is associated with an increased risk of AD. One explanation for this finding is that the heterogeneity with the phenotype is greatly increased when a proxy diagnosis is used. Proxy diagnoses rely on
individual participants knowing the difference between demen-
tias which may not be reliable. This reliance on lay diagnosis is
potentially vulnerable to misclassification thereby increasing the
heterogeneity of the SNPs that are associated with the outcome.
None of the other risk factors examined were significantly asso-
ciated with any of the dementia outcomes.

We were not able to completely rule out that the null findings
from this study are a consequence of limited power. In addition
analyses (online supplemental file 2, online supplemental
file 1) we found evidence that genetic risk of higher BMI, SBP
and smoking has strong positive causal associations with CAD.
These positive associations suggest that if there had been similar
effects of these risk factors on AD, we would have been able to
detect them. Notwithstanding the limitations of MR studies and
that of interpreting null findings, our study did not find evidence
to support the risk factors as proposed by the Lancet Commiss-
ion are causal in nature.

Limitations and further research
A strength of the current study was running the analysis for
different subtypes of dementia. However, one limitation was
that only three were analysed. Thus, we were not able to provide
a comprehensive analysis for all common subtypes of dementia.
One obvious omission is that of vascular dementia. At the time
of completing these analyses, no GWAS studies were available
for the genetic risk of vascular dementia. In time there may be
GWAS studies for vascular dementia and future similar studies
should aim to include this as an outcome. On a related note, with
some of the rarer forms of dementia such as posterior cortical
atrophy (PCA) although GWAS data do exist for this phenotype,
the sample sizes are relatively small. For example, one study only
included a sample of 302 PCA cases20 and this was considered
by the study authors not sufficiently powered to include in the
current analysis. Another limitation of the present study was the
inability to stratify the data by APOE-e4 carrier status and sex.
APOE-e4 is a risk factor for AD,21 and carrier status may interact
with risk factors in a way that produces different outcomes in
carriers versus non-carriers.22 For example, APOE-e4 status is
associated with slower cognitive decline in women with coronary
artery risk factors,22 and in APOE-e4 carriers with AD treated
for hypertension.23 A further limitation of this study is that the
analysis was conducted using datasets of European population
samples. Therefore, it is not known if these results are general-
isable to other ethnic populations. Currently, the large GWAS
datasets required for MR analysis for different ethnicities are
not available. Again, with time and more data, these GWAS may
increase in sample sizes and diversity of population sample and
allow researchers to conduct similar research on rarer forms of
dementia and different ethnic groups.

Conclusions
The results of our study found limited genetic evidence for causal
links between 10 of the risk factors identified by the Lancet
Commission and dementia. There are limitations to results
from MR studies, such as that of survivor bias, and therefore
the inferences that can be drawn from such studies. However,
the results from this study do not align with the findings of
the Lancet Commission. Given the limitations to both design
types, evidence from causal inference studies should be consid-
ered alongside evidence from traditional epidemiological studies
and incorporated in reviews of the literature and when applying
statistics at a population level.

Contributors

RD, VZ and JS came up with the initial conceptualisation, study
design and analysis plan. RD, VZ and RS conducted the data analysis with RD taking
the lead and VZ and RS validating the analysis. RD was responsible for writing
the manuscript with intellectual input and critical evaluation from all coauthors. JS and
VZ provided supervision. RD and JS are the guarantors of the study.

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None declared.

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Data availability statement

Data are available in a public, open access repository.

Supplemental material

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