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Alcohol consumption and risk of dementia: 23 year follow-up of Whitehall II cohort study

Séverine Sabia,1,2 Aurore Fayosse,1 Julien Dumurgier,3 Aline Dugravot,1 Tasnime Akbaraly,2,4,5 Annie Britton,2 Mika Kivimäki,2 Archana Singh-Manoux1,2

ABSTRACT

OBJECTIVE
To examine the association between alcohol consumption and risk of dementia.

DESIGN
Prospective cohort study.

SETTING
Civil service departments in London (Whitehall II study).

PARTICIPANTS
9087 participants aged 35-55 years at study inception (1985/88).

MAIN OUTCOME MEASURES
Incident dementia, identified through linkage to hospital, mental health services, and mortality registers until 2017. Measures of alcohol consumption were the mean from three assessments between 1985/88 and 1991/93 (midlife), categorised as abstinence, 1-14 units/week, and >14 units/week; 17 year trajectories of alcohol consumption based on five assessments of alcohol consumption between 1985/88 and 2002/04; CAGE questionnaire for alcohol dependence assessed in 1991/93; and hospital admission for alcohol related chronic diseases between 1985/88 and 1991/93 (midlife), categorised as alcohol related hospital admission (4.28, 2.72 to 6.73) were also included in the analyses, resulting in potential bias due to selection in results.9 10 This is particularly likely to promote cognitive health at older ages.

WHAT IS ALREADY KNOWN ON THIS TOPIC
Moderate alcohol consumption is thought to be associated with a lower risk of dementia; the association of alcohol with cognitive outcomes appears to be J-shaped or U-shaped, with harmful effects of both abstinence and excessive alcohol consumption.

The evidence is far from robust, however, as excessive alcohol consumption is not included in current guidelines to prevent or delay dementia onset.

Inconsistency in findings stems from the fact that most studies assess alcohol consumption in late life, which may not reflect lifetime consumption, and selection bias is likely to affect these findings as studies used face-to-face assessment of cognitive status.

WHAT THIS STUDY ADDS
The results show a greater risk of dementia in those who abstain from alcohol or consume >14 units/week, with risk increasing in a linear fashion at higher levels of consumption.

Data on hospital admission for chronic disease caused by high alcohol consumption showed a fourfold higher risk of dementia in these people.

The study also found support for a mediating role of cardiometabolic disease; some of the excess risk of dementia in abstainers was explained by greater risk of cardiometabolic disease in this group.

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Introduction

Excessive alcohol consumption is a leading risk factor for several chronic diseases and mortality.1 1 With continuously increasing life expectancy and the expected tripling of dementia prevalence by 2050,1 understanding the impact of alcohol consumption on aging outcomes is important.4 Moderate alcohol consumption has been suggested to lower the risk of dementia, and the association of alcohol consumption with cognitive outcomes is thought to be J-shaped or U-shaped.5 7 However, several issues remain unresolved that might explain why alcohol consumption is not listed in the most recent guideline on modifiable risk factors for the prevention of dementia.8 Firstly, as most studies rely on face-to-face assessment for dementia diagnosis, people who drop out of the study or die during follow-up are not included in the analyses, resulting in potential bias due to selection in results.2 10 This is particularly likely in relation to excessive alcohol consumption, which is known to be associated with greater mortality and drop-out rates.9 Secondly, most studies on aging...
RESEARCH

assess alcohol consumption in late life,5-7 which may not reflect lifetime consumption, and this may be critically important for dementia as it involves neuropathological changes over many years, perhaps decades. The tendency to reduce alcohol consumption at older ages may bias results11 and prevent accurate analyses of the quantity of alcohol consumed.5 7

Thirdly, most studies use a single assessment of alcohol consumption, which is prone to measurement error. Furthermore, indicators of heavy drinking other than the reported frequency and units of alcohol consumed may add to understanding dementia but are rarely used.12 In addition, the mechanisms underlying the association between alcohol and cognitive aging remain unclear.5-7 Non-moderate alcohol consumption is associated with a higher risk of cardiometabolic disease,1 13-15 which is itself associated with a higher risk of dementia,4 16 suggesting a potential role of these diseases in the association between alcohol consumption and dementia.

To tackle some of these limitations, we used repeat data spanning nearly three decades to investigate the association between alcohol consumption and risk of dementia, assessed through linkage to electronic health records for all participants irrespective of their continued participation in follow-up. We examined associations of dementia with alcohol consumption in midlife, alcohol dependence, hospital admission for alcohol related disease, and trajectories of alcohol consumption over 17 years. In addition, we examined whether cardiometabolic disease modifies the association between alcohol consumption and dementia.

**Methods**

**Study population**

The Whitehall II study is an ongoing cohort study of men and women originally employed by the British civil service in London based offices.17 A total of 10 308 adults (6895 men and 3413 women, aged 35-55) were recruited during 1985-88. Since baseline, follow-up clinical examinations have taken place about every four or five years, with each wave taking two years to complete; the last one was for 2015/16. Written informed consent from participants and research ethical approvals were renewed at each contact.

**Measures**

**Alcohol consumption**


**Midlife alcohol consumption**—The mean age of participants assessed for alcohol consumption in midlife was 50.3 years (fig 1 and appendix table S1). To reduce measurement error we used the mean of consumption measured in 1985/88, 1989/90, and 1991/93 waves (n=7908)*.

**Dementia cases during follow-up:** 1991/93 wave to 31 March 2017 (n=397)
Mean follow-up=23.2 years (SD=4.4 years)

**Dementia cases during follow-up:** 1991/93 wave to 31 March 2017 (n=328)
Mean follow-up=23.4 years (SD=4.1 years)

**Dementia cases during follow-up:** 2002/04 wave to 31 March 2017 (n=396)
Mean follow-up=12.9 years (SD=2.7 years)

**Cardiometabolic disease during follow-up:** 1991/93 wave to dementia diagnosis, or date of death or 31 March 2017 (n=2985)
Dementia cases during follow-up: 1991/93 wave to 31 March 2017: (n=371) (of which 208 occurred after a first cardiometabolic disease and 163 among healthy participants)

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Fig 1 | Flow chart of study. Of 10 308 baseline participants, the following were excluded: *1221 (77 died before 1991/93 and 1144 had <2 alcohol measures during 1985/88, 1989/90, and 1991/93 waves); 12339 (77 died before 1991/93 and 2262 did not have data on CAGE or covariates at 1991/93 wave); €169 (77 died before 1991/93 and 92 did not have data on covariates during follow-up (missing covariates at a specific wave were replaced by data from closest wave)); 1381 (14 had dementia and 491 died before 2002/04, and 876 had <2 alcohol measures over 1985/88, 1989/90, 1991/93, 1997/99, and 2002/04 waves); \$1559 (77 died before 1991/93, 1144 had <2 alcohol measures over 1985/88, 1989/90, and 1991/93 waves, and 338 had prevalent cardiometabolic disease)
We classified participants who reported no alcohol consumption over the five previous years in 1985/88 and no alcohol consumption in the previous year in 1989/90 and 1991/93 as 10 year abstainers (n=269, 3.0%). Participants who reported alcohol cessation in the previous five years in 1985/88 and those consuming alcohol in 1985/88 or 1989/90 but not in 1991/93 were categorised as former drinkers (n=172, 1.9%). Participants who reported consuming alcoholic beverages in the previous year but not in the last week at all three waves were classified as occasional drinkers (n=862, 9.5%). As these three categories had similar hazards of dementia (see appendix table S2), we combined them into a single category of “abstinence.” Among drinkers, we categorised average alcohol consumption (1985/88 to 1991/93) as 1-14 units/week and >14 units/week to reflect alcohol guidelines in the United Kingdom.18

Alcohol consumption trajectory from midlife to early old age—We constructed trajectories of alcohol consumption between 1985/88 and 2002/04 (mean age 44.8 and 61.2, respectively; fig 1 and appendix table S1) using our defined categories of alcohol consumption data from 1985/88, 1989/90, 1991/93, 1997/99, and 2002/04 based on 8927 participants who were alive and free of dementia in 2002/04 and who had at least two assessments of alcohol consumption. Trajectories were determined using group based trajectory modelling, fitted using the traj command in Stata.19 We chose the trajectories based on model fit statistics (bayesian information criterion values and average posterior probabilities) and judgment about whether they adequately addressed the research question (appendix table S3).20 The identified trajectories (appendix figure S1) were long term abstinence, decreased alcohol consumption, long term consumption of 1-14 units/week, increased consumption, and long term consumption of >14 units/week.

Alcohol dependence
Alcohol dependence was measured by the CAGE questionnaire in 1991/93. This brief four item scale (felt need to Cut down on drinking, Annoyed by people criticising your drinking, Guilty about drinking, Need a drink first thing (Eye-opener)).21 It is a validated screening instrument for alcohol dependence that was originally developed for general practice settings and has been found to identify problems with alcohol.22

Hospital admission for alcohol related chronic disease
Using the national hospital episode statistics database we identified hospital admissions attributable to alcohol related chronic disease according to the ICD codes (international classification of diseases, ninth and 10th revisions) defined by the Centers for Disease Control and Prevention23 (appendix table S4).

Dementia
We used comprehensive tracing of electronic health records for dementia ascertainment based on three databases: the national hospital episode statistics, the Mental Health Services Data Set, and the mortality register. Record linkage was until 31 March 2017, using ICD-10 codes F00-F03, F05.1, G30, and G31. The UK National Health Service (England, Scotland, and Wales) provides most of the healthcare, including outpatient and inpatient care; private medical insurance, held by around 12% of the UK population (1997 figures).24 is mainly used for elective surgery rather than for chronic conditions such as dementia. The Mental Health Services Data Set is a national database that contains information on people in contact with mental health services in hospitals, outpatient clinics, and the community. Mortality data were drawn from the British national mortality register (National Health Services Central Registry). The tracing exercise was carried out using the unique NHS identification number given to each UK resident. The date of dementia was set at the first record of dementia diagnosis in any of the three databases used for ascertainment.

To assess the validity of the method of dementia ascertainment, we used a mixed model with a backward time scale and determined trajectories of global cognitive score based on performance in three cognitive domains (memory, reasoning, fluency), assessed five times between the 1997/99 and 2015/16 waves.25 These results show an accelerated decline in global cognition in the 10 years before the dementia diagnosis (appendix figure S2), as has been shown in studies that use a “gold standard” method for dementia ascertainment.26

Covariates
Sociodemographic variables included age, sex, ethnicity (white, non-white), marital status (married/cohabiting, other), and socioeconomic status using occupational position (three categories: high, intermediate, and low, representing income and status at work) and education (five categories: less than primary school (age <11), lower secondary school (age <16), higher secondary school (age <18), and university degree and higher).

Health behaviours were assessed over the follow-up period by questionnaire and included smoking (current, former, never), hours of moderate to vigorous physical activity (using questions on moderately energetic (eg, dancing, cycling, leisurely swimming) and vigorous (eg, running, hard swimming, playing squash) physical activity at baseline and then using a modified version of the Minnesota leisure time physical activity questionnaire),25 and dietary behaviour (frequency of fruit and vegetables consumed in a week).

Health related covariates included systolic blood pressure, total cholesterol level, prevalent diabetes...
mellitus (determined by fasting blood glucose ≥7.0
mmol/L, reported doctor diagnosed diabetes, use of
antidiabetic drugs, or hospital records), body mass
index (categorised as <20, 20-24.9, 25-29.9, and ≥30
kg/m²) assessed by a trained nurse at the clinical
examination through blood tests and standard clinical
protocols, cardiovascular diseases (including coronary
heart disease, stroke, atrial fibrillation, and heart
failure) identified using linkage to national hospital
records, self reported use of drugs for cardiovascular
disease, and the 30 item general health questionnaire
on anxiety and depression symptoms.27

Statistical analysis
Three sets of analyses were undertaken (fig 1 and
appendix table S1). Cox regression was used in all
analyses, with age as the timescale to model the
associations with hazard of incident dementia. Participants were censored at date of record of
dementia, death, or 31 March 2017, whichever came
first. Models were first adjusted for sociodemographic
factors, then additionally for health behaviours, and
finally for health status.

Analysis of midlife alcohol consumption
We examined the association of alcohol consumption
in midlife (mean of 1985/88, 1989/90, and 1991/93),
alcohol dependence in 1991/93, and hospital
admission for alcohol related chronic diseases from
Covariates were assessed in 1991/93 (or the closest
wave if missing), apart for the analysis on hospital
admission for alcohol related chronic diseases, for
which all covariates were time varying. We first
examined the shape of the association of midlife
alcohol consumption and dementia using restricted
cubic spline regressions with Harrell knots,28 and Stata
command xblc39 with 14 units/week consumption
as the reference. In subsequent Cox regression we
assessed the association of alcohol consumption
categories (abstinence, 1-14 units/week, and >14
units/week) with risk of dementia. In those reporting
consumption of >14 units/week, we used spline
regressions to examine whether there was a linear
trend in the association of alcohol consumption with
dementia and then estimated the risk of dementia for
every increment of 7 units/week.

Trajectories of alcohol consumption from midlife to
early old age
The aim of these analyses was to examine the
association of long term alcohol consumption with risk
of dementia using trajectories of alcohol consumption
between 1985/88 and 2002/04 waves; the follow-up
started in 2002/04 and covariates were drawn from the
2002/04 wave (or the closest wave if missing).

Role of cardiometabolic disease in association
between alcohol consumption and dementia
In subsequent analyses we examined the mediating
role of cardiometabolic disease (stroke, coronary
heart disease, atrial fibrillation, heart failure, and
diabetes) over the follow-up period in the association
between alcohol consumption and risk of dementia. These analyses were carried out using multistate
models (package mstate, R software) in participants
free of dementia and cardiometabolic disease in
1991/93. The models allow simultaneous estimation
of the risk associated with alcohol consumption
in three transitions: from healthy state to incident
cardiometabolic disease, from cardiometabolic disease
to incident dementia, and from healthy state to incident
dementia in those free of cardiometabolic disease.
Age was used as the timescale, and the analyses were
adjusted for sociodemographic factors.

Sensitivity analysis
Five sets of sensitivity analyses were undertaken.
Firstly, we examined the association between alcohol
consumption in midlife and risk of mortality in the
total sample using Cox regression and then in those
without a diagnosis of dementia during the follow-
up period using multistate models. We used the
modified Fine and Gray competing risks method to
address the possibility that those who died could have
developed dementia had they lived longer.26 Secondly,
we repeated analysis on the shape of the association
between alcohol consumption and risk of dementia
excluding abstainers in order to check for potential
distortion of the association due to abstainers. Thirdly,
to examine whether alcohol consumption at age
50, 60, and 70 was similarly associated with risk of
dementia, we extracted data on alcohol consumption
at these ages for each participant across the data
waves, allowing a five year margin either way for each
age category, and we assessed the association with
incident dementia using Cox models; start of follow-
up and covariates were from age 50, 60, and 70,
respectively. Fourthly, we examined the associations
of type of alcohol consumed using mean midlife
restricted cubic splines adjusted for sociodemographic
variables and mutually adjusted for types of alcohol
consumed. Finally, to examine whether poor
sensitivity of our method of dementia ascertainment
biased our results, we simulated scenarios with
differential misclassification (ie, the hypothesis that
proportion of misclassified dementia cases depends
on midlife alcohol consumption categories) using a
SAS macro provided by Fox et al.11 We allowed the
sensitivity to range between 60% and 90% and the
specificity between 97% and 100% using a trapezoidal
probability density function. We first analysed the
impact of potential differential misclassification on
the hazard ratio estimate associated with abstinence
compared with alcohol consumption of 1-14 units/
week. To account for differential misclassification, we
used separate sensitivity (specificity) distributions in the
two alcohol consumption groups (abstinence and
1-14 units/week). We simulated two scenarios: first
where the correlation of the two sensitivity (specificity)
distributions in the alcohol consumption groups was

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0.8 and then 0.6 (a correlation of 1 would correspond to non-differential misclassification). We repeated these analyses to assess the impact of differential misclassification on the comparison between alcohol consumption of >14 units/week compared with 1-14 units/week. Cox regression analyses were undertaken using STATA 15, multistate models using R, and sensitivity analyses on dementia misclassification using SAS 9.4.

**Patient involvement** Participants of the Whitehall II study were not involved in setting the research question or the outcome measures, nor were they involved in developing plans for recruitment, design, or implementation of the study. No participants were asked advice on interpretation or writing up of results. However, all results are disseminated to study participants through newsletters and our website, which has a participant portal, www.ucl.ac.uk/whitehallII/participants/.

**Results** Among the 10 231 participants alive in 1991/93, 9087 had at least two measurements of alcohol consumption between 1985/88 and 1991/93 and complete data on covariates (fig 1). Among these participants, a total of 397 cases of dementia were recorded over a mean follow-up of 23.2 (SD 4.4, range 0.08-25.6) years. Mean age at dementia diagnosis was 75.6 (SD 5.8; interquartile range 72.2-80.0; range 53.4-85.9) years, with the first case recorded in 1995 and 72% (287 cases out of 397) of the cases recorded in the last five years of follow-up. Greater age (hazard ratio 1.21 per 1 year older age at study baseline, 95% confidence interval 1.18 to 1.23), female sex (1.57, 1.29 to 1.92), education less than secondary school diploma (1.68, 1.38 to 2.05), and low occupational position (2.39, 1.94 to 2.95) were associated with a greater hazard of dementia. Table 1 presents the characteristics of the study population (n=9087). Abstainers were more likely to be women, non-white, and from the lower socioeconomic group. They also had a worse cardiometabolic profile (table 1). Participants in the 1-14 units/week alcohol consumption group were more likely to drink wine and those in the >14 units/week group to drink beer (table 1).

**Midlife alcohol consumption** Age at dementia diagnosis was 76.1, 75.7, and 74.4 years (P=0.13) in the abstinence, 1-14 units/week, and >14 units/week groups, respectively. As no evidence was found of an interaction between alcohol consumption and age (P=0.76), or sex (P=0.92), or occupational position (P=0.95) in associations with dementia, we combined these subgroups in the analyses.

Figure 2 shows the association between alcohol consumption in midlife and risk of dementia in analysis adjusted for sociodemographic factors; age at which alcohol consumption was assessed did not modify this association (fig 2 and appendix table S5). Abstinence was associated with a higher risk of dementia when the

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**Table 1 | Characteristics of study population in 1991/93.** Values are numbers (percentages) unless stated otherwise.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>No dementia status at end of follow-up</th>
<th>Average alcohol consumption during 1985/88, 1989/90, and 1991/93 waves</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of participants</td>
<td>8690</td>
<td>397</td>
</tr>
<tr>
<td>Mean (SD) (years)</td>
<td>50.0 (6.0)</td>
<td>55.8 (4.7)</td>
</tr>
<tr>
<td>Women</td>
<td>2708 (31.2)</td>
<td>166 (41.8)</td>
</tr>
<tr>
<td>Non-white</td>
<td>789 (9.1)</td>
<td>53 (13.4)</td>
</tr>
<tr>
<td>Mean age at dementia diagnosis</td>
<td>75.6 (SD 5.8; interquartile range 72.2-80.0; range 53.4-85.9)</td>
<td>72.2 (SD 5.8; interquartile range 72.2-80.0; range 53.4-85.9)</td>
</tr>
<tr>
<td>Mean (SD) systolic blood pressure (mm Hg)</td>
<td>121.0 (14.0)</td>
<td>124.3 (15.2)</td>
</tr>
<tr>
<td>Mean (SD) total cholesterol (mmol/L)</td>
<td>6.4 (1.2)</td>
<td>6.7 (1.3)</td>
</tr>
<tr>
<td>Mean (SD) moderate to vigorous physical activity (hrs)</td>
<td>4.7 (8.8)</td>
<td>3.6 (8.8)</td>
</tr>
<tr>
<td>Mean (SD) age at which hospital admission for alcohol related disease(s)</td>
<td>1307 (15.0)</td>
<td>74 (16.7)</td>
</tr>
<tr>
<td>Mean (SD) moderate to vigorous physical activity (hrs)</td>
<td>3.9 (4.3)</td>
<td>3.5 (3.8)</td>
</tr>
<tr>
<td>Weight gain (kg)</td>
<td>5.1 (5.4)</td>
<td>5.1 (5.4)</td>
</tr>
<tr>
<td>Body mass index ≥30</td>
<td>813 (9.4)</td>
<td>60 (15.1)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>202 (2.3)</td>
<td>20 (5.0)</td>
</tr>
<tr>
<td>Mean (SD) total cholesterol (mmol/L)</td>
<td>6.4 (1.2)</td>
<td>6.7 (1.3)</td>
</tr>
<tr>
<td>Mean (SD) systolic blood pressure (mm Hg)</td>
<td>121.0 (14.0)</td>
<td>124.3 (15.2)</td>
</tr>
<tr>
<td>Cardiovascular disease</td>
<td>187 (2.2)</td>
<td>17 (4.3)</td>
</tr>
<tr>
<td>Cardiovascular disease drugs</td>
<td>795 (8.1)</td>
<td>75 (18.9)</td>
</tr>
<tr>
<td>General health questionnaire score</td>
<td>3.1 (5.2)</td>
<td>3.2 (5.3)</td>
</tr>
</tbody>
</table>

*All data are drawn from 1991/93, baseline of study population.

†Based on those with available data (n=9769, number of cases=328), cases defined as CAGE score ≥2.

‡Numbers differ with those in table 2 as values are for those with available data on five year alcohol consumption.
reference was alcohol consumption of 14 units/week; in these analyses alcohol consumption >14 units/week was associated with an increased risk of dementia in a linear fashion (among those drinking >14 units/week, P for non-linearity=0.97 using spline regressions). In a model adjusted for sociodemographic factors alcohol abstinence was associated with a greater risk of dementia (hazard ratio 1.47, 1.15 to 1.89) compared with alcohol consumption of 1-14 units/week (table 2). Among those drinking >14 units/week, a 7 unit increase in alcohol consumption was associated with a 17% (95% confidence interval 4% to 32%) increase in risk of dementia. Additional adjustment for health behaviours and health related variables did not attenuate the observed associations. Use of time varying health related factors (data not shown) did not change the results: the hazard ratio for abstainers was 1.44 (1.11 to 1.85) and for each 7 unit increase in consumption among those drinking >14 units/week was 1.18 (1.05 to 1.34).

Compared to those with a CAGE score of 0, a higher risk of dementia was observed in those with a CAGE score >2 (hazard ratio 2.19, 1.29 to 3.71). This association was slightly attenuated in the fully adjusted model (table 2). In analysis adjusted for sociodemographic factors, one hospital admission or more for alcohol related chronic disease over follow-up was associated with a 4.3 times higher risk of dementia (95% confidence interval 2.7 to 6.7). This hazard ratio reduced to 3.0 (1.9 to 4.7) in fully adjusted analysis (table 2).

Trajectories of alcohol consumption between midlife and early old age
We identified five trajectories of alcohol consumption (appendix figure S1): long term abstinence (9% of participants were in this group), decreased consumption (6%), long term consumption 1-14 units/week group (59%), increased consumption (11%), and long term consumption >14 units/week (14%). Compared with participants in the long term consumption 1-14 units/week group, those with long term abstinence (1.74, 1.31 to 2.30), decreased consumption (1.55, 1.08 to 2.22), and long term consumption >14 units/week (1.40, 1.02 to 1.93) had a higher risk of dementia. These associations remained after adjustment for behavioural and health related factors (table 3).

Role of cardiometabolic disease in association between midlife alcohol consumption and dementia
The mediating role of cardiometabolic disease was examined using multistate models (figs 3 and 4). Among those without cardiometabolic disease, compared with alcohol consumption of 1-14 units/week, the hazard ratio for dementia associated with abstinence was 1.33 (0.88 to 2.02) and with consumption >14 units/week was 1.28 (0.85 to 1.92) (fig 3); corresponding hazard ratios in the entire population, when cardiometabolic disease was not taken into consideration, were 1.44 (1.44 to 1.86) and 1.33 (0.88 to 2.02) (fig 3); corresponding hazard ratios in the entire population, when cardiometabolic disease was not taken into consideration, were 1.47 (1.15 to 1.89) and 1.08 (0.82 to 1.43; table 2). In participants consuming >14 units/week (fig 4), cardiometabolic disease did not seem to play a role, as the hazard ratio associated with each increase of 7 units/week was 1.16 (0.96 to 1.41), similar to that observed when cardiometabolic diseases were not taken into account (1.17, 1.04 to 1.32; table 2).

Sensitivity analysis
Analysis using Cox regression showed that compared with alcohol consumption of 1-14 units/week, the
Table 2 | Association between alcohol consumption and risk of dementia

<table>
<thead>
<tr>
<th>Variables</th>
<th>No of cases/total No</th>
<th>Hazard ratio (95% CI)</th>
<th>Adjusted for sociodemographic variables†</th>
<th>Additionally adjusted for behavioural factors‡</th>
<th>Fully adjusted§</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average alcohol consumption in midlife (1985/88, 1989/90, and 1991/93); cases=397/9087; mean follow-up 23.2 (SD 4.4) years</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abstinence</td>
<td>98/1303</td>
<td>1.47 (1.15 to 1.89)*</td>
<td>1.48 (1.15 to 1.91)*</td>
<td>1.45 (1.12 to 1.86)*</td>
<td></td>
</tr>
<tr>
<td>1-14 units/week</td>
<td>229/5552</td>
<td>1 (ref)</td>
<td>1 (ref)</td>
<td>1 (ref)</td>
<td></td>
</tr>
<tr>
<td>&gt;14 units/week</td>
<td>70/2232</td>
<td>1.08 (0.82 to 1.43)</td>
<td>1.05 (0.80 to 1.39)</td>
<td>1.02 (0.77 to 1.35)</td>
<td></td>
</tr>
<tr>
<td>Among those drinking &gt;14 units/week</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Per 7 units/week increase</td>
<td>70/2232</td>
<td>1.17 (1.04 to 1.32)*</td>
<td>1.19 (1.05 to 1.34)*</td>
<td>1.18 (1.04 to 1.30)*</td>
<td></td>
</tr>
<tr>
<td>CAGE score in 1991/93**: cases=328/7969; mean follow-up 23.4 (SD 4.1) years</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>253/5727</td>
<td>1 (ref)</td>
<td>1 (ref)</td>
<td>1 (ref)</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>42/1429</td>
<td>0.93 (0.67 to 1.29)</td>
<td>0.93 (0.66 to 1.30)</td>
<td>0.90 (0.64 to 1.26)</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>18/574</td>
<td>1.03 (0.64 to 1.66)</td>
<td>1.01 (0.62 to 1.63)</td>
<td>0.96 (0.59 to 1.56)</td>
<td></td>
</tr>
<tr>
<td>3/4</td>
<td>15/239</td>
<td>2.19 (1.29 to 3.71)*</td>
<td>2.13 (1.25 to 3.61)*</td>
<td>1.98 (1.15 to 3.38)*</td>
<td></td>
</tr>
<tr>
<td>Hospital admission for alcohol related disease from 1991††; cases=460/10 139; mean follow-up 23.0 (SD 5.0) years</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No (0/1)</td>
<td>295/1156</td>
<td>1 (ref)</td>
<td>1 (ref)</td>
<td>1 (ref)</td>
<td></td>
</tr>
<tr>
<td>Yes (x2)</td>
<td>33/813</td>
<td>1.37 (0.95 to 1.97)</td>
<td>1.34 (0.93 to 1.93)</td>
<td>1.27 (0.88 to 1.84)</td>
<td></td>
</tr>
<tr>
<td>At least one during follow-up</td>
<td>20/193</td>
<td>4.28 (2.72 to 6.73)*</td>
<td>3.70 (2.34 to 5.86)*</td>
<td>2.95 (1.85 to 4.71)*</td>
<td></td>
</tr>
</tbody>
</table>

*P<0.05.
†Adjusted for age (time scale), sex, ethnicity, education, occupational position, and marital status.
‡Additionally adjusted for physical activity, smoking status, and fruit and vegetable consumption.
§Fully adjusted for systolic blood pressure, total cholesterol, diabetes, body mass index, general health questionnaire score, cardiovascular disease, and cardiovascular disease drugs.
¶Participants with at least two measures between 1985/88 and 1991/93 (78.8% had three measures and 21.2% had two measures).
**CAGE was used for first time in 1991/93.
††All data entered as time varying covariates.

Discussion

In this longitudinal study, multiple approaches to examine the association between alcohol consumption and dementia present converging evidence on three key findings on abstinence, excessive alcohol consumption, and the role of cardiometabolic disease. First, the risk of dementia was higher in those abstaining from alcohol in midlife. Alcohol consumption trajectories from midlife to early old age supported these findings—both long term abstainers and those reporting decreased alcohol consumption had an increased risk of dementia. Second, alcohol consumption >14 units/week increased the risk of dementia in a linear fashion; an excess risk that was evident when alcohol consumption was assessed at ages 50, 60, and 70 years. Data using hospital admission for chronic disease caused by high alcohol consumption showed a four times higher risk of dementia, supporting findings on the neurotoxic effects of alcohol consumption >14 units/week. Thirdly, multistate models showed that part of the excess risk of dementia in abstainers was attributable to the greater risk of cardiometabolic disease in this group. Taken together, these results suggest that abstention and excessive alcohol consumption are associated with an increased risk of dementia, although the underlying mechanisms are likely to be different in the two groups. Overall, no evidence was found that alcohol consumption between 1 unit/week and 14 units/week increases the risk of dementia.

Strengths and limitations of this study

The present study has several strengths. Repeat assessment of alcohol consumption allowed us to assess mean midlife alcohol consumption in order to minimise biases due to measurement error, examine associations with dementia of trajectories of alcohol consumption between midlife and early old age, and examine whether age modifies associations between alcohol consumption and dementia. These features, along with a mean follow-up period of 23 years, allowed a comprehensive assessment of the association of alcohol consumption with dementia. Besides measurement error, studies that recruit participants at older ages are not able to assess the excess risk in those who change their alcohol consumption with age. We were also able to examine the shape of the...
association between alcohol consumption >14 units/week and dementia, which was similar to that reported in a recent meta-analysis.7 Dose-response assessment by meta-analysis can be problematic for heavy alcohol consumption as the estimate is constrained to the mean or median consumption in the high alcohol consumption category. Finally, we used multistate models to examine the role of cardiometabolic disease and we undertook further analyses to take the competing risk of mortality into account where results were similar to those obtained using Cox regression, increasing confidence in our main findings.

The study findings need to be interpreted keeping in mind the observational nature of the data. A key limitation, as in other observational studies, is the measurement of alcohol consumption using self-reports. It is possible that systematic reporting biases affected findings, although comparison of alcohol consumption reported by the participants of the Whitehall II study suggests patterns similar to those in several other UK cohort studies. The use of multiple approaches, including the measures of hospital admission for alcohol related chronic disease with converging findings, suggest that our results on excessive alcohol consumption are robust.

The ascertainment of dementia based on linkage to electronic health records has advantages and disadvantages. A recent study32 reported that passive assessment of dementia through UK hospital records has high specificity but modest sensitivity (78%) owing to milder cases of dementia being missing, as also found in the Mayo Clinic Study of Aging and the Adult Changes study.33 In addition to hospital records, we used other sources of dementia diagnosis, such as the UK mental health database, which is likely to improve the sensitivity of dementia diagnosis. Accordingly, our analyses, which simulated differential misclassification scenarios, show the results to be robust. Our findings are also in accordance with previous findings from the Whitehall II study suggesting that both alcohol abstinence and high alcohol consumption are associated with accelerated cognitive decline.34 The advantage of ascertainment through linkage to health records is that it allows analysis of everyone recruited to the study rather than only those who continue to participate in the study over a long follow-up and are available for an in-person ascertainment of dementia. A further difficulty of face-to-face assessment is that people can develop dementia and die between two assessments, which prevents them from being categorised as having dementia. Such bias is particularly likely with risk factors that also affect mortality, as is the case with excessive alcohol consumption.

Comparison with other studies
A recent meta-analysis of observational studies concluded that light to moderate alcohol consumption is associated with a reduced risk of dementia, whereas both abstinence and heavy drinking are associated with a higher risk of dementia.7 In seven of the 10 studies included in the analysis, the mean follow-up period was less than 10 years, with alcohol consumption being assessed later in life and thus potentially modified by health related problems.11 In two of the three other studies, the CAIDE35 and the Finnish Twin36 cohorts, although a U-shaped association was suggested, results were not robust owing to the small number of cases in the extreme alcohol consumption categories. In a study based on the Swedish Twin Registry with 43 years of follow-up, where dementia was assessed through electronic health records, a
quadratic association was found whereby both no alcohol consumption and high alcohol consumption were associated with an increased risk of dementia, although the excess risk in abstainers did not reach statistical significance and the excess risk of high consumption began at 12 g/day (corresponding to around 10.5 units/week). The study also found, as in our investigation, a reduced risk of dementia for moderate wine consumption and a linear increased risk of dementia in those consuming spirits. Furthermore, a recent study based on a nationwide dataset of patients admitted to hospital in France between 2008 and 2013 reported that those with a hospital admission record for alcohol use disorders had a 3.3-times higher risk of dementia in multivariate analysis, providing further support for our findings.

We, as with others, observed an increased risk of dementia in alcohol abstainers, a finding subject to much debate. As studies usually assess alcohol consumption only once, excess risk might be driven by the inclusion of former drinkers in the same group as abstainers. Our analyses using repeat data on alcohol consumption across midlife suggest that former drinking might not explain the excess dementia risk in abstainers, although we cannot exclude the possibility that those who report alcohol abstinenbe in midlife were heavy drinkers in young adulthood or misreported their alcohol consumption. We accounted for several sociodemographic and health-related characteristics in the analysis, but residual confounding cannot be excluded as an explanation for the higher risk of dementia among abstainers.

Indeed, this group is particular in that it is composed mainly of women from the lower socioeconomic group with higher prevalence of cardiometabolic risk factors and disease at baseline, a pattern that has also been observed in other studies. A recent study of nearly 600,000 people, for example, found a J-shaped association between alcohol consumption and cardiovascular disease, with a weekly alcohol consumption of 100 g (12.5 units) being associated with the lowest risk of cardiovascular disease and a higher disease risk observed in those consuming smaller amounts of alcohol. Moderate alcohol consumption has been hypothesised to benefit cardiovascular health through favourable impacts on lipid profile, inflammation level, endothelial function, and insulin sensitivity. In agreement with this, a meta-analysis of interventional studies (alcohol use versus a period of no alcohol use) reported that moderate alcohol consumption had favourable effects on levels of high density lipoprotein cholesterol, apolipoprotein A1, adiponectin, and fibrinogen, potentially underlying the apparent neuroprotective effects of moderate alcohol consumption. Our multistate models lent partial support for a mediating role of cardiometabolic disease in the association between alcohol abstention and increased risk of dementia; the hazard ratio of dementia associated with alcohol abstinence, compared with moderate consumption, was reduced to 1.33 (95% confidence interval 0.88 to 2.02) in those without cardiometabolic disease, compared with 1.47 (1.15 to 1.89) in the entire population. Nevertheless, studies using other approaches such as mendelian randomisation and brain imaging data from a subsample of the Whitehall II study suggest linear adverse effects of alcohol consumption. The present findings on alcohol abstinence should therefore not motivate people who do not drink to start drinking given the known detrimental effects of alcohol consumption across midlife were heavy drinkers in young adulthood or misreported their alcohol consumption. We accounted for several sociodemographic and health-related characteristics in the analysis, but residual confounding cannot be excluded as an explanation for the higher risk of dementia among abstainers.

### Table 4 | Association between midlife alcohol consumption and risk of dementia in competing risk analysis

<table>
<thead>
<tr>
<th>Variables</th>
<th>Dementia cases/total</th>
<th>Hazard ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abstinence</td>
<td>98/1303</td>
<td>1.47 (1.15 to 1.89)*</td>
</tr>
<tr>
<td>1-14 units/week</td>
<td>229/5552</td>
<td>1 (ref)</td>
</tr>
<tr>
<td>&gt;14 units/week</td>
<td>70/2232</td>
<td>1.08 (0.82 to 1.43)</td>
</tr>
<tr>
<td>Among those drinking &gt;14 units/week</td>
<td></td>
<td>1.17 (1.04 to 1.32)*</td>
</tr>
<tr>
<td>Per 7 unit increase</td>
<td>70/2232</td>
<td>1.16 (1.07 to 1.25)*</td>
</tr>
</tbody>
</table>

*P<0.05.
†Modified Fine and Gray competing risk analysis. Cases=397/9087, mean follow-up=23.2 (SD 4.4) years. Adjusted for age (time scale), sex, ethnicity, education, occupational position, and marital status.
in a linear fashion. In addition, long term exposure to alcohol consumption above this limit increased the risk of dementia by 50% compared with long term moderate consumption (1-14 units/week). These results support the recent downward revision of UK guidelines that moved the recommended alcohol consumption limit to 14 units/week in men compared with 21 units/week before, bringing them in line with women. Analyses on alcohol dependence scale and hospital admission for alcohol related chronic disease strengthen the evidence that excessive alcohol consumption is a risk factor for dementia. The negative impact of heavy alcohol intake on the risk of dementia has been suggested to involve nutritional deficiency, the direct neurotoxic effects of ethanol, and the indirect negative impacts through increased risk of diabetes, hypertension, and stroke.

Conclusion

Given the number of people living with dementia is expected to triple by 2050 and the absence of a cure, prevention is key. We show that both long term alcohol abstinence and excessive alcohol consumption may increase the risk of dementia. The UK guidelines suggest an alcohol threshold of 14 units/week but many countries use a much higher threshold to define excessive consumption. The present study encourages the use of a lower threshold of alcohol consumption in such guidelines, applicable over the adult life course, in order to promote cognitive health.

We thank the participating civil service departments and their welfare, personnel, and establishment officers; the British Occupational Health and Safety Agency; the British Council of Civil Service Unions; the participating civil servants in the Whitehall II study; and the Whitehall II study team, which comprises research scientists, statisticians, study coordinators, nurses, data managers, administrative assistants, and data entry staff.

Contributors: ASM and SS developed the hypothesis and study design. SS, AF, and AD performed the statistical analysis. SS wrote the first and successive drafts of the manuscript. All authors conceived and designed the study, analysed and interpreted the data, and drafted or critically revised the manuscript for important intellectual content. SS, AF, and AD obtained funding for the Whitehall II study. SS, AD, and ASM had full access to the data and take responsibility for the integrity of the data and the accuracy of the data analysis. SS is the guarantor. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

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Competing interests: All authors have completed the ICMJE uniform disclosure form at http://www.icmje.org/coi_disclosure.pdf (available on request from the corresponding author) and declare no other support from any organisation for the submitted work or the grants reported in the funding section, 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. The sponsors had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; and preparation, review, or approval of this manuscript.

Ethical approval: This study was approved by University College London Hospital Committee on the Ethics of Human Research (reference No 85/0938). Written informed consent from participants and research ethics approvals were renewed at each contact.

Data sharing: Data, protocols, and other metadata of the Whitehall II study are available to the scientific community. Please refer to the Whitehall II study data sharing policy at www.ucl.ac.uk/whitehallll/data-sharing.

Transparency: The lead author (SS) 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 planned have been explained.

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18. UK Chief Medical Officers. UK Chief Medical Officers’ Low Risk Drinking Guidelines 2016.
Supplementary information: figures S1-S5 and tables S1-S7