



HAL
open science

Association of longitudinal alcohol consumption trajectories with coronary heart disease: a meta-analysis of six cohort studies using individual participant data

Dara O’neill, Annie R. Britton, Mary K Hannah, Marcel Goldberg, Diana Kuh, Kay Tee Khaw, Steven Bell

► To cite this version:

Dara O’neill, Annie R. Britton, Mary K Hannah, Marcel Goldberg, Diana Kuh, et al.. Association of longitudinal alcohol consumption trajectories with coronary heart disease: a meta-analysis of six cohort studies using individual participant data. *BMC Medicine*, 2018, 16 (1), pp.124. 10.1186/s12916-018-1123-6 . inserm-02057158

HAL Id: inserm-02057158

<https://inserm.hal.science/inserm-02057158>

Submitted on 5 Mar 2019

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

RESEARCH ARTICLE

Open Access



Association of longitudinal alcohol consumption trajectories with coronary heart disease: a meta-analysis of six cohort studies using individual participant data

Dara O'Neill^{1*} , Annie Britton², Mary K. Hannah³, Marcel Goldberg⁴, Diana Kuh^{2,5}, Kay Tee Khaw⁶ and Steven Bell⁷

Abstract

Background: Studies have shown that alcohol intake trajectories differ in their associations with biomarkers of cardiovascular functioning, but it remains unclear if they also differ in their relationship to actual coronary heart disease (CHD) incidence. Using multiple longitudinal cohort studies, we evaluated the association between long-term alcohol consumption trajectories and CHD.

Methods: Data were drawn from six cohorts (five British and one French). The combined analytic sample comprised 35,132 individuals (62.1% male; individual cohorts ranging from 869 to 14,247 participants) of whom 4.9% experienced an incident (fatal or non-fatal) CHD event. Alcohol intake across three assessment periods of each cohort was used to determine participants' intake trajectories over approximately 10 years. Time to onset for (i) incident CHD and (ii) fatal CHD was established using surveys and linked medical record data. A meta-analysis of individual participant data was employed to estimate the intake trajectories' association with CHD onset, adjusting for demographic and clinical characteristics.

Results: Compared to consistently moderate drinkers (males: 1–168 g ethanol/week; females: 1–112 g ethanol/week), inconsistently moderate drinkers had a significantly greater risk of incident CHD [hazard ratio (HR) = 1.18, 95% confidence interval (CI) = 1.02–1.37]. An elevated risk of incident CHD was also found for former drinkers (HR = 1.31, 95% CI = 1.13–1.52) and consistent non-drinkers (HR = 1.47, 95% CI = 1.21–1.78), although, after sex stratification, the latter effect was only evident for females. When examining fatal CHD outcomes alone, only former drinkers had a significantly elevated risk, though hazard ratios for consistent non-drinkers were near identical. No evidence of elevated CHD risk was found for consistently heavy drinkers, and a weak association with fatal CHD for inconsistently heavy drinkers was attenuated following adjustment for confounding factors.

Conclusions: Using prospectively recorded alcohol data, this study has shown how instability in drinking behaviours over time is associated with risk of CHD. As well as individuals who abstain from drinking (long term or more recently), those who are inconsistently moderate in their alcohol intake have a higher risk of experiencing CHD. This finding suggests that policies and interventions specifically encouraging consistency in adherence to lower-risk drinking guidelines could have public health benefits in reducing the population burden of CHD. The absence of an effect amongst heavy drinkers should be interpreted with caution given the known wider health risks associated with such intake.

Trial registration: ClinicalTrials.gov, [NCT03133689](https://clinicaltrials.gov/ct2/show/study/NCT03133689).

Keywords: Alcohol, Coronary heart disease, IPD meta-analysis, Longitudinal design

* Correspondence: d.oneill@ucl.ac.uk

¹CLOSER, Department of Social Science, Institute of Education, University College London, London, UK

Full list of author information is available at the end of the article



Background

The relationship between alcohol consumption and coronary heart disease (CHD) is of scientific and public health interest, yet it remains a subject of debate. Studies have found evidence both for and against the possibility of an association [1, 2]. The concept of a potentially cardio-protective effect of moderate drinking compared to non-drinking or heavier consumption, termed the U/J-shaped curve, has been particularly controversial [3–5]. Some clinical evidence suggests that alcohol may affect different pathways thought to influence CHD risk, including hypertension [6, 7], body mass index (BMI) [8] and lipid levels [9, 10]. However, this purported effect continues to be disputed [11], which poses challenges in the formation of health-care policy and can hinder wider public understanding of the health impact of lifestyle choices.

Much of the discussion around the evidence base for the alcohol–CHD association has focussed on design limitations in observational studies, such as the failure to distinguish between non-drinkers and former drinkers [12, 13]. The decision to stop drinking could be influenced by the onset of ill health, and such sick quitters could potentially bias estimates of disease risk in lifelong abstainers if not analysed independently [14]. Studies have most commonly used single baseline measures of alcohol intake and that drinking behaviours can change over time has, therefore, not typically been reflected in the alcohol epidemiology literature [15, 16].

Recent efforts have been made to establish long-term trajectories of alcohol intake, enabling differentiation between patterns of drinking that fluctuate over time. Different trajectories have been found to have distinct patterns of association with intermediate markers of cardiovascular health, including carotid intima media thickness [17], pulse wave velocity [18] and inflammatory markers [19], but this work has yet to link these drinking typologies to CHD events directly. More commonly, studies with longitudinal assessments of drinking have used average intake, typically between only two measurement occasions, in evaluations of CHD risk [20], but such aggregation can mask consumption variation over time. The importance of capturing variability is evident from previous work that has shown how isolated episodes of heavy drinking can offset the potentially protective effects of moderate drinking [2]. Failure to account for stability in alcohol intake levels may bias risk estimates [21]. In the current study, we have used an alcohol intake trajectory approach, previously employed in the study of intermediate CHD markers [17–19], to address this research gap. We have drawn data from multiple cohort studies to investigate whether longitudinal trajectories of alcohol consumption differ in their association with total CHD incidence (fatal or non-fatal). Furthermore, since research has suggested that the cardio-protective effect of moderate drinking may be less evident

with fatal CHD outcomes [22], particularly in comparison to heavier intake [23], a secondary aim of this work was to examine how the longitudinal trajectories are specifically associated with mortality due to CHD.

Methods

Sample, design and cohort selection

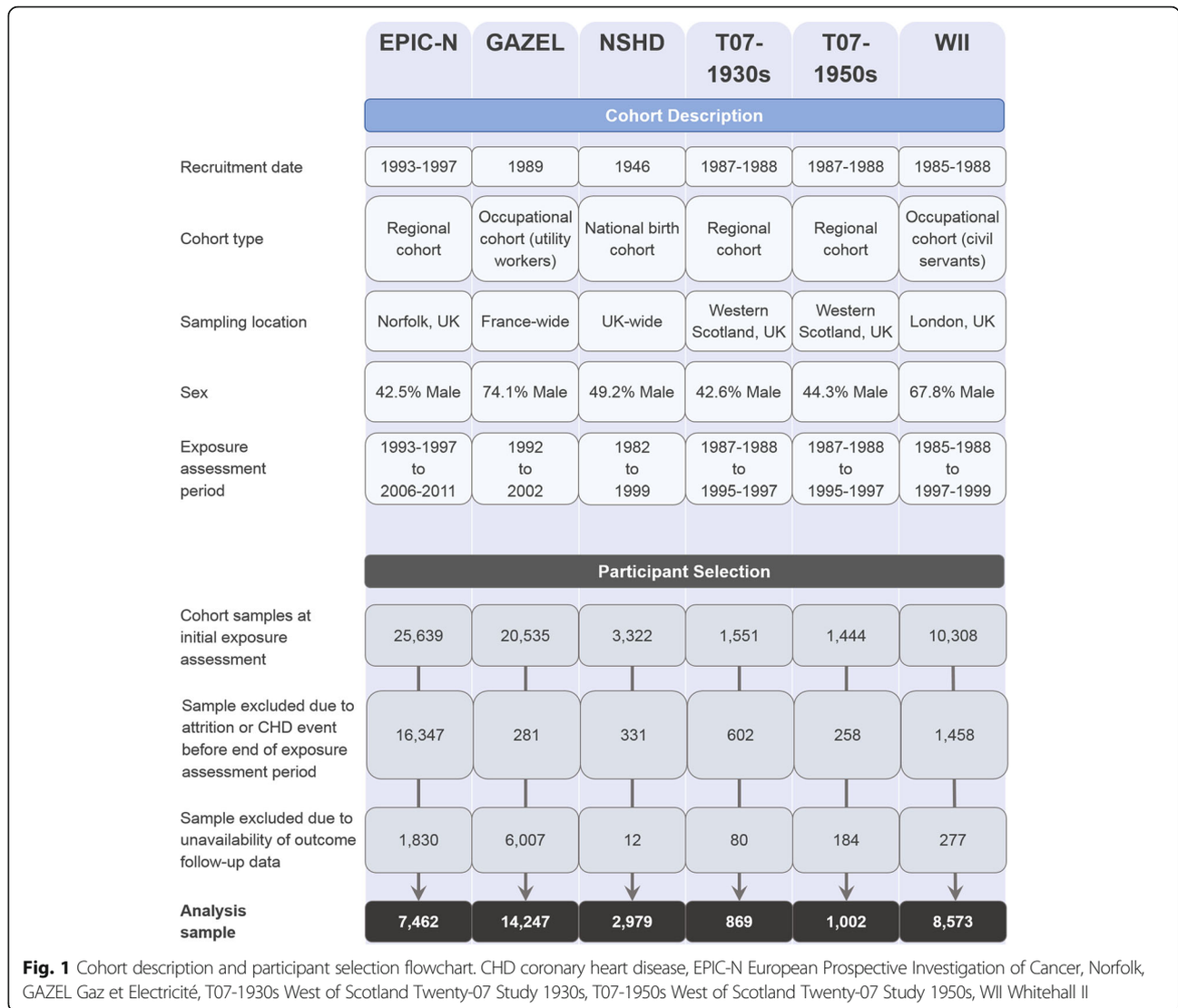
Data were obtained from five British cohort studies: the European Prospective Investigation of Cancer, Norfolk Cohort (EPIC-N) [24]; the Medical Research Council's National Survey of Health and Development 1946 (NSHD) [25]; West of Scotland Twenty-07: 1930s (T07-1930s) [26]; West of Scotland Twenty-07: 1950s (T07-1950s) [26] and Whitehall II (WII) [27]. Further data were obtained from an additional French cohort: Gaz et Electricité (GAZEL) [28]. Descriptions of each cohort are provided in Fig. 1 and complete cohort profiles are available via the above citations. The cohorts were chosen for their coverage of relevant variables and design similarity. They each included prospective alcohol intake data across three assessments covering an approximate 10-year interval, as well as pertinent covariate and verified CHD outcome data. Prior to commencement of the analysis, additional harmonisation was performed for all cohort datasets to maximise consistency in variable names and definitions. The study design was pre-registered on [ClinicalTrials.gov](https://www.clinicaltrials.gov) (identifier NCT03133689), and a STROBE statement is provided in Additional file 1 (Section S1).

The combined dataset initially comprised 62,799 participant records in total (cohort-specific counts are provided in Fig. 1). The exposure variable was measured across three assessment points covering a decade on average, with the last assessment point treated as the study baseline from which time-to-event outcomes were measured. Of the initial sample, 19,277 participants were excluded due to attrition or having experienced a CHD event prior to the study baseline. A further 8390 participants were not included due to incomplete data linkage. Following these exclusion criteria (further details of which are provided in Fig. 1), a sample of 35,132 (62.1% male) participants remained and these comprised the analytic sample.

Measures

Outcomes

The primary endpoint was CHD incidence, as ascertained from linked health records and survey data. Non-fatal CHD data were available for NSHD, but the time to event from the end of the exposure period was not ascertainable, so this study was omitted from the analysis of the incident CHD end point. Mortality due to CHD was examined as a secondary outcome in supplementary analyses, and all cohorts contributed data to this analysis. CHD events were previously coded using the codebook of the International Statistical Classification of Diseases and Related Health



Problems (ICD) [29]: ICD-9: 410–414 Ischaemic heart disease and ICD-10: I20-I25 Ischaemic heart diseases. For the Twenty-07 cohorts, non-fatal CHD events were identified using the Royal College of General Practitioners’ codebook [30] (codes 1940, 1945 and 1950). Survival time was calculated for all participants as time (in years) between the end of the alcohol assessment period and date of CHD event, death from non-CHD causes, study dropout or last date of data linkage (study specific), whichever occurred first. Additional details of study enrolment and follow-up procedures are available in Additional file 1 (Section S10).

Exposure

Trajectories of self-reported alcohol intake were derived using weekly alcohol intake measurements collected across three consecutive waves of each cohort study. The resultant trajectories comprised a decade of follow-up on average. Standard drink definitions were used to define alcohol

(ethanol) content in reported drinks (half-pints of beer or cider, small glasses of wine, and single servings of spirits): 8 g ethanol in the British cohort data and 10 g in the French GAZEL data [31, 32]. Reported consumption at each measurement occasion was categorised according to UK drinking guidelines at the time of data collection, which recommended a maximum intake level for lower-risk drinking of 168 g of ethanol a week for males and 112 g of ethanol a week for females [33]. Although recently published UK guidelines have proposed identical thresholds for males and females [34], the analytic report upon which they are based identified risk functions for both CHD morbidity and mortality that notably differed between males and females [35], particularly at higher levels of consumption [23]. The focus in the present study was on stability of adherence to lower-risk drinking guidelines over time, and we consequently categorised participants according to longitudinal profiles as defined in Table 1. Consistently moderate

Table 1 Drinker type definitions with observed counts and percentages (within sex and overall)

Drinker type	Weekly alcohol intake	N (%)		
		Male	Female	Total
Consistent non-drinker	0 g at each wave of data collection	807 (4.6)	1335 (12.4)	2142 (7.5)
Former drinker	0 g at last wave but intake >0 g at any earlier wave	1831 (10.4)	2249 (21.0)	4080 (14.4)
Consistently moderate	Male: 1–168 g at each wave	7249 (41.0)	4161 (38.8)	11,410 (40.2)
	Female: 1–112 g at each wave			
Inconsistently moderate	Male: 1–168 g for most but not all waves	3599 (20.4)	2037 (19.0)	5636 (19.8)
	Female: 1–112 g for most but not all waves			
Consistently heavy	Male: >168 g at each wave	2216 (12.5)	349 (3.3)	2565 (9.0)
	Female: >112 g at each wave			
Inconsistently heavy	Male: >168 g for most but not all waves	1979 (11.2)	598 (5.6)	2577 (9.1)
	Female: >112 g for most but not all waves			

drinkers were used as the reference category [14]. Drinkers with inconsistent levels of alcohol intake were defined according to their modal intake, i.e. their most frequent level of intake. For example, where a participant had an equal number of heavy and moderate drinking periods, they were categorised as inconsistently heavy drinkers. This ensured that participants who occasionally drank heavily were not grouped with participants who consistently adhered to lower-risk drinking guidelines.

Covariates

Known demographic and lifestyle risk factors for CHD were selected for inclusion in the modelling, including sex and age. Socioeconomic position was defined using the participant's occupational status, categorised as low (non-skilled or semi-skilled), intermediate (mid-level) or high (professional or executive) [36]. Smoking status was assessed, with participants categorised as current, ex- or non-smokers. To account for variability in the alcohol intake assessment interval, the time difference between initial and final assessments was calculated for each participant. Additional clinical data were obtained on BMI (measured as kg/m²) and self-reported high blood pressure or use of antihypertensive medication (yes/no). All of the covariates were assessed at the commencement of the follow-up period for CHD (the occasion of the third and final alcohol assessment), which we have defined as the current study's baseline.

Statistical analysis

Prior to undertaking inferential analyses, multiple imputation by chained equations was completed using the R 'mice' package (v2.30) to address missing covariate and exposure data. Altogether, 100 imputations were performed for both the incident and fatal CHD analyses, ensuring congruence between the imputation and substantive models. Outcome data with the Nelson–Aalen hazard [37] were used but not imputed.

The modelling was performed as individual participant data (IPD) meta-analyses, accounting for the clustering of participants within each cohort. Both one- and two-step approaches are available and can give comparable results under particular conditions [38]. However, the one-step approach, in which all data are analysed simultaneously with clustering incorporated as a random effect term, is thought to be less prone to bias in pooled effect estimates and standard errors [39] and to be the preferred approach where covariate adjustments are required or where inter-study heterogeneity may be present [40, 41]. Consequently, one-step IPD meta-analysis was performed using hierarchical (mixed effects) Cox regression modelling incorporating a random effect term for cohort membership with maximum likelihood estimations. Models were developed iteratively: an initial model accounting for age, sex and intake assessment interval (partially adjusted for confounding), followed by an extended model that additionally included smoking status and socioeconomic status covariates (maximally adjusted for confounding). Supplementary modelling extended the adjustment further, including potential mediators, to examine clinical pathways (maximally adjusted for confounding and mediation). Schoenfeld residuals were plotted to ascertain that the proportional hazards assumption had not been violated (available in Additional file 1: Section S2).

Given most existing work in this area has employed single one-off measures of alcohol intake, for comparative purposes, an initial IPD meta-analysis was undertaken in this study using participants' final intake measurement prior to the outcome follow-up period (i.e. at this study's baseline). This single measure categorisation allowed a distinction to be made between different intake levels (none, moderate or heavy), but not stability of intake over time or discontinuation of drinking. This analysis was followed by the modelling of the primary exposure, the longitudinal drinking trajectory categorisation. Additional stratified analyses were also completed to explore specific

characteristics of the alcohol–CHD relationship. Research has suggested that the association of alcohol with cardiovascular risk may differ between older and younger populations [42], so age-stratified modelling of the longitudinal drinker typology was also performed (aged ≤ 55 vs > 55 years at this study's baseline). Further stratified analyses were undertaken to explore sex-specific effects. Finally, sensitivity analyses were conducted to determine the impact of modelling assumptions on this study's main results.

The statistical analyses were performed in R (v3.4.1; R Foundation for Statistical Computing, Vienna, Austria). All statistical significance testing was two-tailed, using an inference threshold of $p < 0.05$.

Results

Sample characteristics

Descriptive statistics, for the overall sample and stratified by drinker type, are presented in Table 2. Additional descriptive statistics, stratified by cohort, are provided in Additional file 1 (Section S3). Statistics on data missingness are also reported in Table 2, and further detail is provided in Additional file 1 (Section S4).

Across the drinker types, mean age ranged from 57.1 years (standard deviation, SD = 6.4) for the inconsistently moderate drinkers to 61.9 (SD = 9.1) for the former drinkers. Heavy drinkers were most likely to be male (consistently heavy 86.4%; inconsistently heavy 76.8%), whereas abstainers were more likely to be female (consistent non-drinker 62.3%; former drinker 55.1%). Heavy drinkers had the highest proportion reporting past or current smoking (consistently heavy 50.7%; inconsistently heavy 50.4%). Consistently moderate drinkers were most likely to be of high socioeconomic position (47.8%), followed by both consistently and inconsistently heavy drinkers (45.9% and 46.5%). Conversely, consistent non-drinkers had the highest proportion in a low socioeconomic position (23.9%). BMI showed little variation between drinker types (all had means of 26 kg/m²). Known hypertension was least common amongst consistently moderate drinkers (26.2%) and most common amongst inconsistently heavy drinkers (31.6%) and former drinkers (31.7%). The mean assessment interval covered by the drinking trajectories was similar across all drinker types (range 10.7–11.8 years).

Crude outcome statistics are also provided in Table 2. In the pooled sample, 4.9% of participants experienced an incident CHD (fatal or non-fatal) event during the follow-up. This was lowest for consistently heavy drinkers (3.8%) and highest for former-drinkers (6.1%). The mean follow-up time was 12.6 years (SD = 4.3). In total, 397,264.4 person-years at risk were captured, with mean person-years varying from 11.0 years (former drinkers) to 13.9 years (consistently heavy drinkers). The

overall CHD incidence rate was 4.3 CHD cases per 1000 person-years.

The proportion of individuals dying due to CHD during the follow-up was 0.9%. This varied between drinker types, from 0.6% for the consistently heavy group to 1.3% amongst consistent non-drinkers. The mean follow-up time was 13.7 years (SD = 4.1). In combination, 480,124.7 person-years were captured for this outcome, with the mean person-years again lowest for former drinkers (11.9 years) but highest for inconsistently moderate drinkers (14.8 years). The overall rate of fatal CHD was 0.7 cases per 1000 person-years.

Single intake measure categorisation

In a series of hierarchical Cox regression models with alcohol intake defined according to a single intake measurement just prior to the outcome follow-up period, no discernible difference in incident CHD risk was observed between heavy and moderate drinkers. However, those who reported no intake at this most recent measurement point had an increased risk of CHD compared to those who drank but did so within the recommended limits [model maximally adjusted for confounding: hazard ratio (HR) = 1.26, 95% confidence interval (CI) = 1.11–1.43]. The estimates are illustrated in Fig. 2 and reported in full in Additional file 1 (Section S5a).

Longitudinal intake trajectories

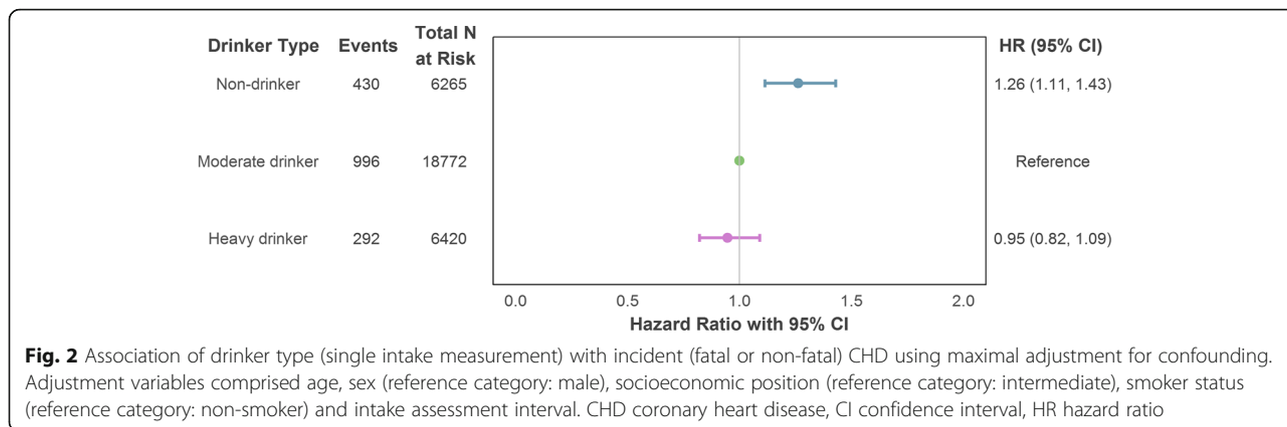
When modelling overall CHD risk using the longitudinal intake typology with adjustment for age, sex and intake assessment interval, both consistent non-drinkers (HR = 1.51, 95% CI = 1.25–1.82) and former drinkers (HR = 1.35, 95% CI = 1.16–1.57) showed greater risk of incident CHD compared to participants who reported persistently moderate intake. A smaller but still significant effect was also found for inconsistently moderate drinkers (HR = 1.21, 95% CI = 1.04–1.40). The effects remained statistically significant after additional adjustment for smoking status and socioeconomic position (detailed in Fig. 3). No differences in risk for heavy drinking, consistent or otherwise, were found.

When potential mediators, BMI and hypertension, were included in the modelling, the drinker type effects were attenuated, with the effect for inconsistently moderate drinkers becoming non-significant (HR = 1.16, 95% CI = 1.00–1.34). Full details of the modelling steps are provided in Additional file 1 (Section S5a), including the associations of each covariate with CHD onset risk. Older age, male sex, history (current or past) of smoking, higher BMI and

Table 2 Descriptive results: overall sample

Variable	Level	Drinker type					Overall		
		Consistent non-drinker	Former drinker	Consistently moderate drinker	Inconsistently moderate drinker	Consistently heavy drinker		Inconsistently heavy drinker	Unknown
Record count, N		2142	4080	11,410	5636	2565	2577	6722	35,132
Age, Mean (SD)		58.9 (8.1)	61.9 (9.1)	59.7 (8.1)	57.1 (6.4)	58.0 (5.7)	57.9 (6.6)	58.6 (7.3)	59.1 (7.7)
Sex, N (%)	Male	807 (37.7)	1831 (44.9)	7249 (63.5)	3599 (63.9)	2216 (86.4)	1979 (76.8)	4140 (61.6)	21,821 (62.1)
	Female	1335 (62.3)	2249 (55.1)	4161 (36.5)	2037 (36.1)	349 (13.6)	598 (23.2)	2582 (38.4)	13,311 (37.9)
Smoker, N (%)	No	1351 (63.1)	2383 (58.4)	6933 (60.8)	3124 (55.4)	1218 (47.5)	1234 (47.9)	2280 (33.9)	18,523 (52.7)
	Current smoker	321 (15.0)	494 (12.1)	1008 (8.8)	764 (13.6)	540 (21.1)	473 (18.4)	1022 (15.2)	4622 (13.2)
	Ex-smoker	407 (19.0)	1129 (27.7)	3322 (29.1)	1630 (28.9)	759 (29.6)	825 (32.0)	1335 (19.9)	9407 (26.8)
	Unknown	63 (2.9)	74 (1.8)	147 (1.3)	118 (2.1)	48 (1.9)	45 (1.7)	2085 (31.0)	2580 (7.3)
Socioeconomic position, N (%)	High	510 (23.8)	1353 (33.2)	5454 (47.8)	2279 (40.4)	1178 (45.9)	1198 (46.5)	2400 (35.7)	14,372 (40.9)
	Intermediate	1104 (51.5)	1953 (47.9)	4924 (43.2)	2708 (48.0)	1184 (46.2)	1158 (44.9)	3005 (44.7)	16,036 (45.6)
	Low	512 (23.9)	733 (18.0)	980 (8.6)	612 (10.9)	190 (7.4)	207 (8.0)	1263 (18.8)	4497 (12.8)
	Unknown	16 (0.7)	41 (1.0)	52 (0.5)	37 (0.7)	13 (0.5)	14 (0.5)	54 (0.8)	227 (0.6)
BMI, mean (SD)		26.2 (4.9)	26.3 (4.4)	25.6 (3.5)	26.0 (3.9)	26.1 (3.3)	26.2 (3.7)	26.5 (4.1)	26 (3.9)
High blood pressure, N (%)	No	1498 (69.9)	2784 (68.2)	8409 (73.7)	4010 (71.1)	1790 (69.8)	1759 (68.3)	3376 (50.2)	23,626 (67.2)
	Yes	641 (29.9)	1292 (31.7)	2990 (26.2)	1620 (28.7)	774 (30.2)	815 (31.6)	1551 (23.1)	9683 (27.6)
	Unknown	3 (0.1)	4 (0.1)	11 (0.1)	6 (0.1)	1 (~0.0)	3 (0.1)	1795 (26.7)	1823 (5.2)
Intake interval, mean (SD)		11.4 (2.6)	11.8 (2.4)	11.4 (2.2)	11.2 (2.4)	10.7 (1.7)	11.1 (2.3)	12.6 (2.8)	11.5 (2.4)
CHD (all) during follow-up, N (%)	No	1803 (84.2)	3549 (87.0)	10,260 (89.9)	4775 (84.7)	2375 (92.6)	2229 (86.5)	4748 (70.6)	29,739 (84.6)
	Yes	129 (6.0)	250 (6.1)	560 (4.9)	264 (4.7)	98 (3.8)	107 (4.2)	310 (4.6)	1718 (4.9)
	Unknown	210 (9.8)	281 (6.9)	590 (5.2)	597 (10.6)	92 (3.6)	241 (9.4)	1664 (24.8)	3675 (10.5)
CHD (all) person years, mean (SD)		12.7 (4.5)	11.0 (4.7)	12.4 (4.4)	13.8 (3.7)	13.9 (3.4)	13.5 (3.8)	12.1 (4.5)	12.6 (4.3)
CHD (fatal) during follow-up, N (%)	No	2113 (98.6)	4031 (98.8)	11,328 (99.3)	5593 (99.2)	2549 (99.4)	2550 (99.0)	6630 (98.6)	34,794 (99.0)
	Yes	27 (1.3)	49 (1.2)	78 (0.7)	40 (0.7)	15 (0.6)	25 (1.0)	91 (1.4)	325 (0.9)
	Unknown	2 (0.1)	0 (0.0)	4 (~0.0)	3 (0.1)	1 (~0.0)	2 (0.1)	1 (~0.0)	13 (~0.0)
CHD (fatal) person years, mean (SD)		13.9 (4.2)	11.9 (4.6)	13.2 (4.3)	14.8 (3.2)	14.6 (3.1)	14.3 (3.5)	13.9 (4.0)	13.7 (4.1)

BMI body mass index, kg/m² kilogram per metre squared, CHD coronary heart disease, SD standard deviation, N count



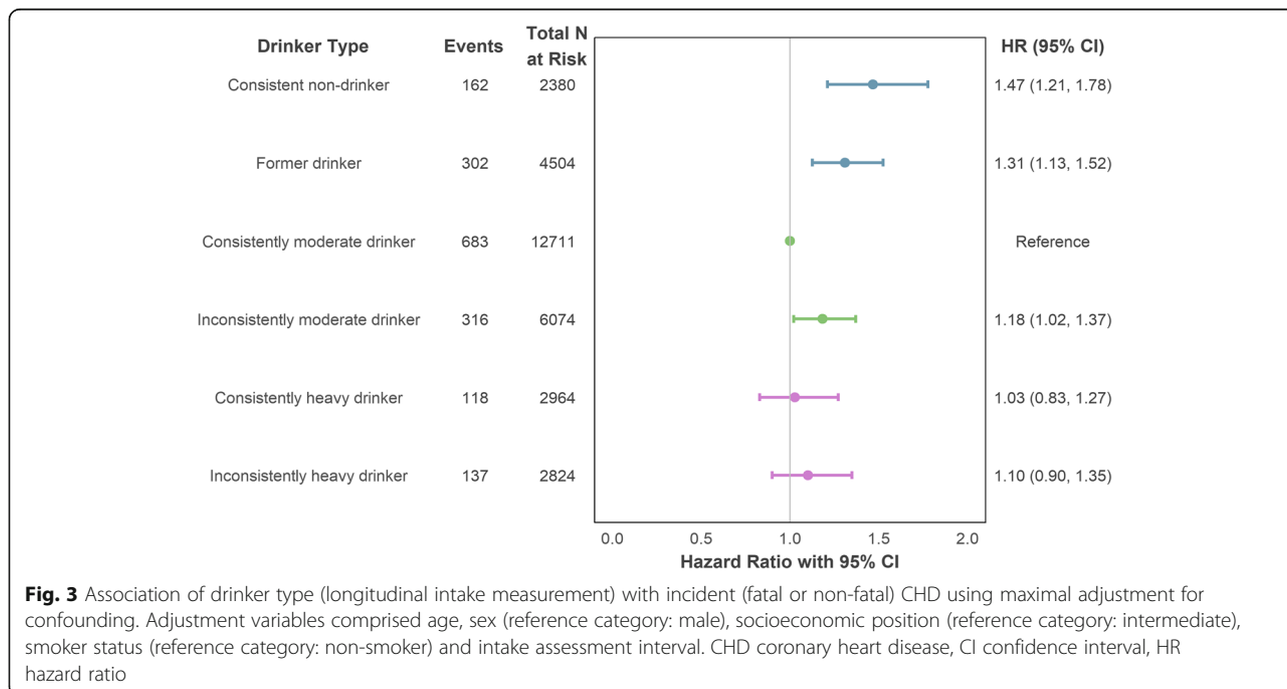
high blood pressure were all significantly associated with increased risk of CHD.

Stratified analyses

In age-stratified analyses of the longitudinal trajectory exposure, participants aged up to 55 years and those aged above showed comparable associations with the incident CHD outcome (visualised in Fig. 4). Consistent non-drinkers (aged ≤55: HR = 1.97, 95% CI = 1.29–3.02; aged > 55: HR = 1.38, 95% CI = 1.11–1.71) and former drinkers (aged ≤55: HR = 1.60, 95% CI = 1.09–2.37; aged > 55: HR = 1.27, 95% CI = 1.08–1.51) both demonstrated significantly greater CHD risk compared to consistently moderate drinkers. However, inconsistently moderate drinkers in the older age group also had an increased risk of incident CHD (HR = 1.25, 95% CI = 1.06–1.48), a

finding not replicated in the younger subsample. Further details are provided in Additional file 1 (Section S6a).

Further stratified analyses were performed to assess if the alcohol–CHD association differed by sex, again using the longitudinal intake categories (illustrated in Fig. 5). Amongst male participants, former drinkers were at significantly greater risk of incident CHD compared to consistently moderate drinkers after maximal adjustment for confounding factors (HR = 1.29, 95% CI = 1.06–1.56). After equivalent adjustment in the female stratum, both former drinkers (HR = 1.38, 95% CI = 1.07–1.78) and consistent non-drinkers (HR = 1.91, 95% CI = 1.43–2.55) showed increased risk compared to their consistently moderate counterparts. A full table of results is provided in Additional file 1 (Section S7a).



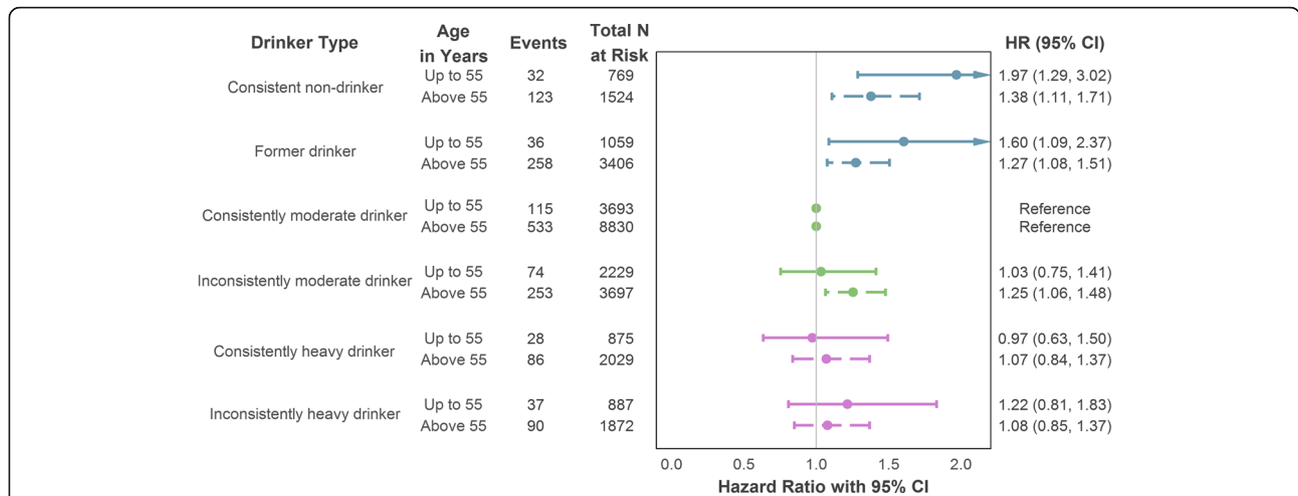


Fig. 4 Age-stratified association of drinker type (longitudinal intake measurement) with incident (fatal or non-fatal) CHD using maximal adjustment for confounding. Adjustment variables comprised age, sex (reference category: male), socioeconomic position (reference category: intermediate), smoker status (reference category: non-smoker) and intake assessment interval. CHD coronary heart disease, CI confidence interval, HR hazard ratio

CHD mortality

When analyses were replicated using fatal CHD as the outcome, most results were comparable to those obtained when using all incident CHD events. For the longitudinal intake trajectories, and in contrast to the incident CHD analysis, inconsistently moderate drinkers did not have a greater CHD mortality risk compared to the consistently moderate reference group (HR = 1.04, 95% CI = 0.72–1.52). Only former drinkers had a significantly elevated risk of fatal CHD (HR = 1.54, 95% CI = 1.07–2.22) after maximal adjustment for confounding factors, but the HR for consistent non-drinkers was near identical (HR = 1.52, 95% CI = 0.97–2.38), implying that again both drinker types were at

elevated risk of fatal CHD. Inconsistently heavy drinkers did show some evidence of having an increased risk of experiencing a fatal CHD event in the lesser-adjusted model (HR = 1.53, 95% CI = 0.99–2.37), but it did not achieve statistical significance and was attenuated after additional adjustment for smoking status and socioeconomic status (HR = 1.36, 95% CI = 0.87–2.11). Full model details are provided in Additional file 1 (Section S5b).

Age-stratified analyses revealed similar patterns of association as with the pooled (non-stratified) analysis. In sex-stratified analyses, however, some differences were observed, with only female consistent non-drinkers having an elevated risk of fatal CHD after adjustment for covariates

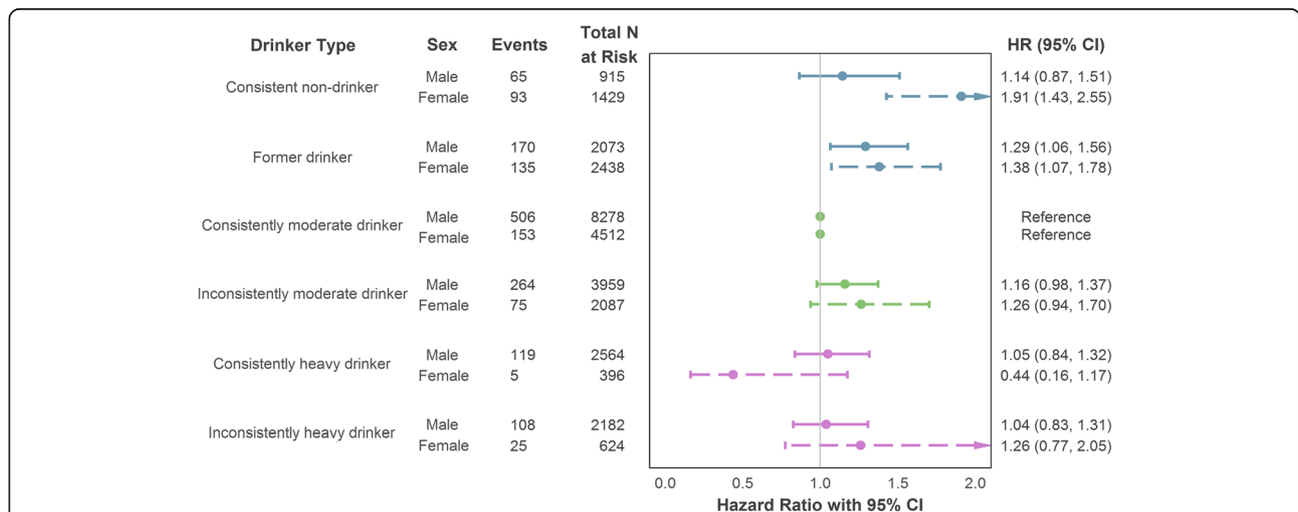


Fig. 5 Sex-stratified association of drinker type (longitudinal intake measurement) with incident (fatal or non-fatal) CHD using maximal adjustment for confounding. Adjustment variables comprised age, socioeconomic position (reference category: intermediate), smoker status (reference category: non-smoker) and intake assessment interval. CHD coronary heart disease, CI confidence interval, HR hazard ratio

(HR = 2.62, 95% CI = 1.25–5.49). Additional detail on age-stratified and sex-stratified analyses are included in Additional file 1 (Sections S6b and S7b, respectively).

Sensitivity analyses

As the GAZEL cohort was the only non-UK data source included in this study, the longitudinal modelling was replicated with this cohort's data omitted to verify that its inclusion did not introduce bias. Results obtained using only UK data sources were essentially unchanged from the findings attained when all cohorts were included (details are provided in Additional file 1: Section S8). To identify the impact of the imputation model implemented in the primary analyses, the modelling was also performed using complete case data only. The point estimates and significance of effects were essentially unchanged from the imputed data modelling (see Additional file 1: Section S9).

Discussion

In this study, we utilised prospectively collected longitudinal data on alcohol consumption from six cohorts to examine the association of 10-year drinking trajectories and risk of developing and/or dying from CHD. Through iterative modelling that accounted for heterogeneity across the datasets and potential confounders of the alcohol–CHD association, our work has shown that incident CHD risk is significantly higher amongst both non-drinkers and former drinkers compared to drinkers who always adhered to lower-risk intake guidelines. We have also demonstrated that the stability of such adherence is pertinent. Participants who mostly drank moderately, but not persistently so, had greater risk of incident CHD compared to their consistently moderate drinking counterparts. In terms of CHD mortality, former drinkers and consistent non-drinkers were again found to be at higher risk, although the effect for the persistent abstainers was somewhat attenuated after adjustment for smoking status and socioeconomic status. We found no evidence that heavy drinking was associated with risk of CHD, and reasons for this are discussed below. Overall, the findings from this study support the notion of a cardioprotective effect of moderate alcohol intake relative to non-drinking. However, crucially, stability in the level of alcohol consumption over time appears to be an important modifier of this association.

The use of repeat measurements of alcohol consumption in lieu of a one-time assessment has enabled us to measure the stability of consumption over time and to address the call for research on the role of intake trajectories in CHD onset [43]. Through this approach, we have demonstrated how intermittent adherence to lower-risk drinking guidelines, i.e. an inconsistently moderate intake, is associated with an increased risk of incident CHD. This provides some support for the proposal that variability in

alcohol intake levels can offset the potential protective effects of moderate drinking [2, 20]. An association was found between inconsistently heavy drinkers and fatal CHD, although the wide confidence bounds and weakening of the association following maximal adjustment for confounding factors limits interpretation of this effect. It may be that unstable drinking patterns reflect wider lifestyle changes across the life course, and possibly even the impact of periods of ill health or life stress. The effects were further attenuated when adjustment was made for clinical characteristics, namely BMI and hypertension, suggesting that these may both act as potential pathways through which unstable drinking trajectories are associated with CHD. The impact of BMI could also reflect the role of other lifestyle choices, such as diet and exercise.

Access to prospectively recorded alcohol intake data across multiple assessment times has also allowed the current study to distinguish recent abstainers from longer-term non-drinkers in a manner that helps reduce the potential for recall bias. Such bias can occur where drinking behaviour is retrospectively measured at a single time point [44], a technique commonly used in alcohol epidemiology research. In line with the sick-quitter hypothesis [14], former drinkers were found in the present study to have an elevated risk of both incident and fatal CHD. These effects were attenuated following adjustment for the clinical covariates, suggesting that poor health may explain former drinkers' increased likelihood of developing CHD and perhaps may even have motivated the decision to abstain itself. Consistent non-drinkers, however, did also have a significant risk of incident CHD after adjustment for potential confounders, and although the error bounds were wider, their CHD mortality estimate was equivalent to that of former drinkers, implying that short- and long-term abstinence are both associated with an increased risk of CHD.

Despite our finding of parity in CHD risk amongst non-drinkers and former drinkers in the pooled sample analyses, previous research has suggested that there may be age-dependent differences in this association. However, this observation was based on studies in which abstinence was determined retrospectively from a single baseline assessment [42], in contrast to the repeated measures design used in the current study. When we stratified our sample by age, the associations between both abstainer groups and incident CHD risk was comparable for younger (≤ 55 years) and older (> 55 years) participants. As similar results were also observed for risk of fatal CHD, our findings challenge the argument that there are age-dependent differences between long-term and more recent abstainers, yet the wide confidence bounds around the fatal CHD risk estimates for those aged 55 or below arguably restricts such inferences. A divergence between the age groups was found for inconsistently moderate drinkers. Such drinkers in the

older subsample had a significantly elevated risk of incident CHD, an effect that was not evident in the younger group. Older participants may have been more likely to experience lifestyle changes that influenced their drinking habits. Retirement, for example, is known to co-occur with increases in alcohol intake [45, 46], particularly amongst existing drinkers [47].

It has also been suggested that the J-shaped association between alcohol consumption and CHD may be more pronounced in women than men [23], a theory that our study supports in part. Whilst both male and female former drinkers had significantly increased risk of incident CHD, only female consistent non-drinkers showed such an elevated risk. Female non-drinkers (both long term and more recent abstainers) were similarly at risk of fatal CHD, even after maximal adjustment for confounding factors. Research has also suggested that alcohol intake may increase oestrogen levels in women, which in turn act as a protective factor against CHD [48]. Male former drinkers also showed significantly greater risk of CHD mortality than consistently moderate drinkers after accounting for age and other characteristics, but this difference was attenuated once the estimates were adjusted for lifestyle behaviours such as smoking. This suggests that these additional covariates may play a greater role than drinking in the occurrence of fatal CHD events for males. Previous literature has proposed that smoking can offset any alcohol-related differences in CHD mortality risk amongst men [49].

In the present study, no association with CHD risk was found for consistently heavy drinkers. Stable patterns of heavy drinking may reflect continued good health across the assessment interval [50], the converse of the sick quitter type. Statistically significant associations between high levels of alcohol intake and CHD onset risk have been observed in some previous research [21, 51], but not persistently so [52–54]. Although our study identified heavy drinkers across all cohorts, only a limited number were in the female sample, potentially limiting statistical power in their analysis, and by extension, in the non-stratified analysis. This issue of small counts for female heavy drinkers has similarly constrained earlier work in this area [1]. Particularly heavy drinkers may be under-represented in the datasets utilised in this study, which could have biased downwards the estimate of association between heavy intake and cardiovascular risk. If further data are available, it may be possible to explore alternative intake thresholds and validate the present study's findings. Similarly, additional data may enable the disaggregation of CHD phenotypes, which could provide more nuanced insights into how heavy drinking is associated with different variants of the disease [55]. Consequently, the interpretation of the absence of an effect amongst heavy drinkers in the current study should be done cautiously, particularly in

light of the known wider health impact of heavy alcohol intake levels [56].

There are additional limitations to our study that warrant consideration. For example, selection bias may have occurred [57], in which participants dropped out of the cohort studies before the outcome assessment period. It is possible that some heavy drinkers could have experienced adverse health outcomes at a younger age and discontinued their research participation. Particularly heavy drinkers are already known to be under-sampled in population-level surveys [32, 58], so caution is required in drawing inferences about such elevated intake levels. Similarly, information on alcohol intake prior to the exposure assessment period was not consistently available, so the long-term abstainers modelled in this current study may include some participants who ceased drinking prior to recruitment. Given that the current work included only cohort studies for which we had access to individual-level data, the concept of availability bias [59] is also pertinent. Access to additional datasets may help further validate our findings. Such increased sample sizes would also permit more detailed examination than was possible in the current study into the intake variance that occurs amongst drinkers who are inconsistent in their adherence to lower-risk drinking guidelines. Relatedly, the identification of drinking trajectories in the present study was based on drinking volume only and so we were not equipped to look at the role of episodic heavy drinking [60]. Further clarification of the alcohol–CHD association may be achieved where sufficient data are available on other characteristics of consumption, such as drinking frequency. All cohorts included in the current study used self-report for determining alcohol intake; although this is vulnerable to estimation errors, research has shown that drinking data collected through this method remains valid and reliable [44, 61]. A further design consideration in interpreting the current study's results is the harmonisation of data across the different cohort datasets. Establishing equivalent variable definitions in the harmonisation of data constrains the level of detail and raises the possibility of residual confounding. For example, it was not possible to establish a more nuanced smoking variable due to data availability and so there is a possibility of residual confounding by smoking intensity. Relatedly, although an equal number of intake measurements was used across cohorts to establish intake trajectories, the observed time intervals varied (see Section S2 of Additional file 1). While adjustment was made through inclusion of assessment interval length in the regression modelling, it remains possible that limitations in the cohort data harmonisation may have introduced bias. Although country-specific drink conversions were used to calculate alcohol intake [31], there remains potential differences between GAZEL and the other cohorts, such as the possible influence of dietary differences for which residual

confounding could also have occurred [62]. The French paradox, for example, implies that there is an inverse relationship between saturated fat intake and CHD onset risk that is specific to France [63], a relationship in which alcohol debatably plays a role [64]. However, sensitivity analyses showed that the exclusion of GAZEL data did not modify the current study findings. Moreover, the use throughout this study of mixed-effects modelling has helped account for data clustering and thereby helped improve the validity of the results obtained.

Conclusions

In summary, the present study has utilised longitudinal alcohol intake data pooled from multiple cohort sources to establish trajectories of drinking behaviour and assess their association with risk of incident and fatal CHD. The study has demonstrated that recent and more long-term abstainers are at elevated risk of developing CHD, although the effect for persistent abstainers may be confined to females only. The trajectory approach used in this work has also enabled us to show that stability of alcohol intake levels amongst those who do not abstain is pertinent to risk of CHD onset. Drinkers who mostly, but inconsistently, adhered to moderate drinking levels, particularly if aged over 55 years, were found to have elevated risk of incident CHD. There was also some indication that variability in drinking levels amongst heavier drinkers were associated with increased likelihood of CHD mortality, although that effect was attenuated by adjustment for other demographic and lifestyle characteristics. No evidence of elevated risk amongst consistently heavy drinkers was found but this was potentially attributable to under-representation of such drinkers in the sampled data. These findings, nonetheless, illustrate that longitudinal alcohol trajectories have added utility in identifying at-risk drinker types beyond what is possible with single assessments of alcohol consumption. Our findings provide additional insight into the potential cardioprotective effect of moderate alcohol intake, and indicate that consistency of intake levels is a relevant consideration in cardiovascular risk assessments, and in related health education efforts.

Additional file

Additional file 1: Supplementary Materials. (PDF 3002 kb)

Abbreviations

BMI: Body mass index; CHD: Coronary heart disease; CI: Confidence interval; EPIC-N: European Prospective Investigation of Cancer, Norfolk; GAZEL: Gaz et Electricité; HR: Hazard ratio; ICD: International Statistical Classification of Diseases and Related Health Problems; IPD: Individual participant data; NSHD: Medical Research Council's National Survey of Health and Development 1946; SD: Standard deviation; T07-1930s: West of Scotland Twenty-07 Study 1930s; T07-1950s: West of Scotland Twenty-07 Study 1950s; WII: Whitehall II

Acknowledgments

We thank all participants, researchers and support staff involved in the EPIC-N, NSHD, West of Scotland Twenty-07, WII and GAZEL studies.

Funding

This work has been undertaken as part of the Alcohol Life-course Project (<http://www.ucl.ac.uk/alcohol-lifecourse>), which is funded by the UK Medical Research Council/Alcohol Research UK (MR/M006638/1) and the European Research Council (ERC-StG-2012-309337_AlcoholLifeCourse).

EPIC-N is supported by programme grants from the UK Medical Research Council (G0401527 and G1000143) and Cancer Research United Kingdom (C864/A8257) with additional support from the Stroke Association, British Heart Foundation, Research Into Ageing and the Academy of Medical Science.

GAZEL was partly supported by Agence Nationale De La Recherche (ANR-08-BLAN-0028-01), Agence Française de Sécurité Sanitaire de l'Environnement et du Travail (AFSSET-EST08-35), Electricité de France-Gaz de France and the TGIR Cohortes Santé 2008 Program.

NSHD is supported by the UK Medical Research Council (MR_UC_12019/01). Twenty-07 is funded by the UK Medical Research Council (MC_A540_53462 and MC_UU_12017/13). WII is supported by the UK Medical Research Council (MR/K013351/1 and G0902037), the British Heart Foundation (RG/13/2/30098) and the US National Institutes of Health (R01HL36310 and R01AG013196). The funding bodies had no role in the design of this study, the collection, analysis, and interpretation of the data, or in writing the manuscript.

Availability of data and materials

The datasets that support the findings of this study are available to bona fide researchers from the individual cohort studies upon application.

Authors' contributions

DON, AB and SB devised the study, designed the study protocol, analysed the data, interpreted the output and wrote the manuscript. MKH, MG, DK and KTK provided data access and critical revisions of the manuscript. All authors read and approved the final version.

Ethics approval and consent to participate

All data used and analysed in this study were secondary data that had previously been anonymised by the individual cohort studies. Ethical approval for the EPIC-N study was received from the Norwich District Health Authority Ethics Committee. The GAZEL study received approval from the Commission Nationale Informatique et Liberté. The NSHD was granted ethical approval from the Greater Manchester Local Research Ethics Committee and the Scotland A Research Ethics Committee. Approval for the Twenty-07 Study was received from the ethical sub-committee of the West of Scotland Medical Committee and the University of Glasgow's Ethics Committee for Non-Clinical Research Involving Human Subjects. The WII study received approval from the University College London Medical School Committee on the Ethics of Human Research. All participants in these studies gave written informed consent.

Consent for publication

Not applicable

Competing interests

The authors declare that they have no competing interests.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details

¹CLOSER, Department of Social Science, Institute of Education, University College London, London, UK. ²Research Department of Epidemiology and Public Health, University College London, London, UK. ³MRC/CSO Social and Public Health Sciences Unit, Institute of Health and Wellbeing, University of Glasgow, Glasgow, UK. ⁴Inserm UMS 011, Villejuif, France and Paris Descartes University, Villejuif, France. ⁵UK MRC Unit for Lifelong Health & Ageing at UCL, London, UK. ⁶Cambridge Institute of Public Health, University of Cambridge, Cambridge, UK. ⁷Department of Public Health and Primary Care, University of Cambridge, Cambridge, UK.

Received: 16 March 2018 Accepted: 10 July 2018

Published online: 22 August 2018

References

- Roerecke M, Rehm J. Chronic heavy drinking and ischaemic heart disease: a systematic review and meta-analysis. *Open Heart*. 2014;1:e000135.
- Ruidavets J-B, Ducimetière P, Evans A, Montaye M, Haas B, Bingham A, et al. Patterns of alcohol consumption and ischaemic heart disease in culturally divergent countries: the prospective epidemiological study of myocardial infarction (PRIME). *BMJ*. 2010;341:c6077.
- Stockwell T, Chikritzhs T. Commentary: another serious challenge to the hypothesis that moderate drinking is good for health? *Int J Epidemiol*. 2013;42:1792–4.
- Thompson PL. J-curve revisited: cardiovascular benefits of moderate alcohol use cannot be dismissed. *Med J Aust*. 2013;198:419–22.
- de Gaetano G, Costanzo S. Alcohol and health: Praise of the J curves. *J Am Coll Cardiol*. 2017;70:923 LP–925.
- Roerecke M, Kaczorowski J, Tobe SW, Gmel G, Hasan OSM, Rehm J. The effect of a reduction in alcohol consumption on blood pressure: a systematic review and meta-analysis. *Lancet Public Health*. 2017;2:e108–20.
- Chen L, Davey Smith G, Harbord RM, Lewis SJ, Smith GD, Harbord RM, et al. Alcohol intake and blood pressure: a systematic review implementing a Mendelian randomization approach. *PLoS Med*. 2008;5:0461–71.
- Cho Y, Shin S-Y, Won S, Relton CL, Davey Smith G, Shin M-J. Alcohol intake and cardiovascular risk factors: a Mendelian randomisation study. *Sci Rep*. 2015;5:18422.
- Holmes MV, Asselbergs FW, Palmer TM, Drenos F, Lanktree MB, Nelson CP, et al. Mendelian randomization of blood lipids for coronary heart disease. *Eur Heart J*. 2015;36:539–50.
- Tabara Y, Ueshima H, Takashima N, Hisamatsu T, Fujiyoshi A, Zaid M, et al. Mendelian randomization analysis in three Japanese populations supports a causal role of alcohol consumption in lowering low-density lipoprotein cholesterol levels and particle numbers. *Atherosclerosis*. 2016;254:242–8.
- Caetano R. Does a little drinking make your heart grow stronger? *J Stud Alcohol Drugs*. 2017;78:341–3.
- Fillmore KM, Kerr WC, Stockwell T, Chikritzhs T, Bostrom A. Moderate alcohol use and reduced mortality risk: systematic error in prospective studies. *Addict Res Theory*. 2006;14:101–32.
- Rimm EB, Moats C. Alcohol and coronary heart disease: drinking patterns and mediators of effect. *Ann Epidemiol*. 2007;17:53–7.
- Rehm J, Irving H, Ye Y, Kerr WC, Bond J, Greenfield TK. Are lifetime abstainers the best control group in alcohol epidemiology? On the stability and validity of reported lifetime abstinence. *Am J Epidemiol*. 2008;168:866–71.
- Roerecke M. On bias in alcohol epidemiology and the search for the perfect study. *Addiction*. 2017;112:217–8.
- Knott CS, Bell S, Britton A. The stability of baseline-defined categories of alcohol consumption over the adult life course: a 28-year prospective cohort study. *Addiction*. 2018;113:34–43.
- Britton A, Hardy R, Kuh D, Deanfield J, Charakida M, Bell S. Twenty-year trajectories of alcohol consumption during midlife and atherosclerotic thickening in early old age: findings from two British population cohort studies. *BMC Med*. 2016;14:111.
- O'Neill D, Britton A, Brunner EJ, Bell S. Twenty-five-year alcohol consumption trajectories and their association with arterial aging: a prospective cohort study. *J Am Heart Assoc*. 2017;6:e005288.
- Bell S, Mehta G, Moore K, Britton A. Ten-year alcohol consumption typologies and trajectories of C-reactive protein, interleukin-6 and interleukin-1 receptor antagonist over the following 12 years: a prospective cohort study. *J Intern Med*. 2017;281:75–85.
- Embersson JR, Shaper AG, Wannamethee SG, Morris RW, Whincup PH. Alcohol intake in middle age and risk of cardiovascular disease and mortality: accounting for intake variation over time. *Am J Epidemiol*. 2005;161:856–63.
- Britton A, Marmot MG, Shipley MJ. How does variability in alcohol consumption over time affect the relationship with mortality and coronary heart disease? *Addiction*. 2010;105:639–45.
- Rehm J, Sempos CT, Trevisan M. Average volume of alcohol consumption, patterns of drinking and risk of coronary heart disease - a review. *J Cardiovasc Risk*. 2003;10:15–20.
- Roerecke M, Rehm J. The cardioprotective association of average alcohol consumption and ischaemic heart disease: a systematic review and meta-analysis. *Addiction*. 2012;107:1246–60.
- Day N, Oakes S, Luben R, Khaw K-T, Bingham S, Welch A, et al. EPIC-Norfolk: study design and characteristics of the cohort. *Br J Cancer*. 1999;80(Suppl 1):95–103.
- Kuh D, Pierce M, Adams J, Deanfield J, Ekelund U, Friberg P, et al. Cohort profile: updating the cohort profile for the MRC National Survey of health and development: a new clinic-based data collection for ageing research. *Int J Epidemiol*. 2011;40:e1–9.
- Benzeval M, Der G, Ellaway A, Hunt K, Sweeting H, West P, et al. Cohort profile: west of Scotland Twenty-07 study: health in the community. *Int J Epidemiol*. 2009;38:1215–23.
- Marmot M, Brunner E. Cohort profile: the Whitehall II study. *Int J Epidemiol*. 2005;34:251–6.
- Goldberg M, Leclerc A, Zins M. Cohort profile update: The GAZEL cohort study. *Int J Epidemiol*. 2015;44:77g.
- World Health Organization. The ICD-10 classification of mental and behavioural disorders: clinical descriptions and diagnostic guidelines. Geneva: World Health Organization; 1992.
- The Royal College of General Practitioners. The classification and analysis of general practice data. Occasional paper no. 26. 2nd ed. London: The Royal College of General Practitioners; 1986.
- Kalinowski A, Humphreys K. Governmental standard drink definitions and low-risk alcohol consumption guidelines in 37 countries. *Addiction*. 2016;111:1293–8.
- Britton A, O'Neill D, Bell S. Underestimating the alcohol content of a glass of wine: the implications for estimates of mortality risk. *Alcohol Alcohol*. 2016;51:609–14.
- Royal Colleges of Physicians Psychiatrists and General Practitioners. Alcohol and the heart in perspective: Sensible limits reaffirmed. *J R Coll Physicians Lond*. 1995;29:266–71.
- Department of Health. UK chief medical officers' low risk drinking guidelines. London: Department of Health; 2016.
- Holmes J, Angus C, Buykx P, Ally AK, Stone T, Meier P, et al. Mortality and morbidity risks from alcohol consumption in the UK: analyses using the Sheffield alcohol policy model (v.2.7) to inform the UK chief medical officers' review of the UK lower risk drinking guidelines. Sheffield: ScHARR, University of Sheffield; 2016.
- Office for National Statistics. Standard occupational classification 2010. Basingstoke: Palgrave Macmillan; 2010.
- White IR, Royston P. Imputing missing covariate values for the cox model. *Stat Med*. 2009;28:1982–98.
- Thompson S, Kaptoge S, White I, Wood A, Perry P, Danesh J, et al. Statistical methods for the time-to-event analysis of individual participant data from multiple epidemiological studies. *Int J Epidemiol*. 2010;39:1345–59.
- Debray TPA, Moons KGM, van Valkenhoef G, Fthimiou O, Hummel N, Groenwold RHH, et al. Get real in individual participant data (IPD) meta-analysis: a review of the methodology. *Res Synth Methods*. 2015;6:293–309.
- Thomas D, Radji S, Benedetti A. Systematic review of methods for individual patient data meta-analysis with binary outcomes. *BMC Med Res Methodol*. 2014;14:79.
- Tierney JF, Vale C, Riley R, Smith CT, Stewart L, Clarke M, et al. Individual participant data (IPD) meta-analyses of randomised controlled trials: guidance on their use. *PLoS Med*. 2015;12:e1001855.
- Zhao J, Stockwell T, Roemer A, Naimi T, Chikritzhs T. Alcohol consumption and mortality from coronary heart disease: an updated meta-analysis of cohort studies. *J Stud Alcohol Drugs*. 2017;78:375–86.
- Maggs JL, Staff J. No benefit of light to moderate drinking for mortality from coronary heart disease when better comparison groups and controls included: a commentary on Zhao et al. (2017). *J Stud Alcohol Drugs*. 2017;78:387–8.
- Bell S, Britton A. Reliability of a retrospective decade-based life-course alcohol consumption questionnaire administered in later life. *Addiction*. 2015;110:1563–73.
- Zins M, Guéguen A, Kivimaki M, Singh-Manoux A, Leclerc A, Vahtera J, et al. Effect of retirement on alcohol consumption: longitudinal evidence from the French GAZEL cohort study. *PLoS One*. 2011;6:e26531.
- Halonen JI, Stenholm S, Pulakka A, Kawachi I, Aalto V, Pentti J, et al. Trajectories of risky drinking around the time of statutory retirement: a longitudinal latent class analysis. *Addiction*. 2017;112:1163–70.
- Wang X, Steier JB, Gallo WT. The effect of retirement on alcohol consumption: results from the US health and retirement study. *Eur J Pub Health*. 2014;24:485–9.

48. Grønbaek M. The positive and negative health effects of alcohol- and the public health implications. *J Intern Med.* 2009;265:407–20.
49. Xu W-H, Zhang X-L, Gao Y-T, Xiang Y-B, Gao L-F, Zheng W, et al. Joint effect of cigarette smoking and alcohol consumption on mortality. *Prev Med.* 2007;45:313–9.
50. Ng Fat L, Cable N, Shelton N. Worsening of health and a cessation or reduction in alcohol consumption to special occasion drinking across three decades of the life course. *Alcohol Clin Exp Res.* 2015;39:166–74.
51. Bobak M, Malyutina S, Horvat P, Pajak A, Tamosiunas A, Kubinova R, et al. Alcohol, drinking pattern and all-cause, cardiovascular and alcohol-related mortality in Eastern Europe. *Eur J Epidemiol.* 2016;31:21–30.
52. Dai J, Mukamal KJ, Krasnow RE, Swan GE, Reed T. Higher usual alcohol consumption was associated with a lower 41-y mortality risk from coronary artery disease in men independent of genetic and common environmental factors: the prospective NHLBI twin study. *Am J Clin Nutr.* 2015;102:31–9.
53. Huang C, Zhan J, Liu Y-J, Li D-J, Wang S-Q, He Q-Q. Association between alcohol consumption and risk of cardiovascular disease and all-cause mortality in patients with hypertension: a meta-analysis of prospective cohort studies. *Mayo Clin Proc.* 2014;89:1201–10.
54. Bell S, Daskalopoulou M, Rapsomaniki E, George J, Britton A, Bobak M, et al. Association between clinically recorded alcohol consumption and initial presentation of 12 cardiovascular diseases: population based cohort study using linked health records. *BMJ.* 2017;356:j909.
55. Bell S, Hemingway H. Alcohol and cardiovascular disease: Author's response. *BMJ.* 2017;356:j1340.
56. Molina PE, Gardner JD, Souza-smith FM, Annie M, Molina PE, Gardner JD, et al. Alcohol abuse: critical pathophysiological processes and contribution to disease burden. *Physiology.* 2014;29:203–15.
57. Naimi TS, Stockwell T, Zhao J, Xuan Z, Dangardt F, Saitz R, et al. Selection biases in observational studies affect associations between 'moderate' alcohol consumption and mortality. *Addiction.* 2017;112:207–14.
58. Britton A, Ben-Shlomo Y, Benzeval M, Kuh D, Bell S. Life course trajectories of alcohol consumption in the United Kingdom using longitudinal data from nine cohort studies. *BMC Med.* 2015;13:47.
59. Riley RD. Commentary: like it and lump it? Meta-analysis using individual participant data. *Int J Epidemiol.* 2010;39:1359–61.
60. Roerecke M, Greenfield TK, Kerr WC, Bondy S, Cohen J, Rehm J. Heavy drinking occasions in relation to ischaemic heart disease mortality - an 11–22 year follow-up of the 1984 and 1995 US national alcohol surveys. *Int J Epidemiol.* 2011;40:1401–10.
61. Batty GD, Shipley M, Tabák A, Singh-Manoux A, Brunner E, Britton A, et al. Generalizability of occupational cohort study findings. *Epidemiology.* 2014;25:932–3.
62. Zhang X-Y, Shu L, Si C-J, Yu X-L, Liao D, Gao W, et al. Dietary patterns, alcohol consumption and risk of coronary heart disease in adults: a meta-analysis. *Nutrients.* 2015;7:6582–605.
63. Chiva-Blanch G, Arranz S, Lamuela-Raventos RM, Estruch R. Effects of wine, alcohol and polyphenols on cardiovascular disease risk factors: evidences from human studies. *Alcohol Alcohol.* 2013;48:270–7.
64. Biagi M, Bertelli AAE. Wine, alcohol and pills: what future for the French paradox? *Life Sci.* 2015;131:19–22.

Ready to submit your research? Choose BMC and benefit from:

- fast, convenient online submission
- thorough peer review by experienced researchers in your field
- rapid publication on acceptance
- support for research data, including large and complex data types
- gold Open Access which fosters wider collaboration and increased citations
- maximum visibility for your research: over 100M website views per year

At BMC, research is always in progress.

Learn more biomedcentral.com/submissions

