Testing for non-linear causal effects using a binary genotype in a Mendelian randomisation study: application to alcohol and cardiovascular traits

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Summary

Background
Mendelian randomisation studies have so far restricted attention to linear associations relating the genetic instrument to the exposure, and the exposure to the outcome. In some cases, however, observational data suggest a non-linear association between exposure and outcome. For example, alcohol consumption is consistently reported as having a U-shaped association with cardiovascular events. In principle Mendelian randomisation could address concerns that the apparent protective effect of light-to-moderate drinking might reflect ‘sick-quitters’ and confounding.

Methods
The Alcohol-ADH1B Consortium was established to study the causal effects of alcohol consumption on cardiovascular events and biomarkers, using the single nucleotide polymorphism rs1229984 in ADH1B as a genetic instrument. To assess non-linear causal effects in this study we propose a novel method based on estimating local average treatment effects for discrete levels of the exposure range, then testing for a linear trend in those effects. Our method requires an assumption that the instrument has the same effect on exposure in all individuals. We conduct simulations examining the robustness of the method to violations of this assumption, and apply the method to the Alcohol-ADH1B Consortium data.

Results
Our method gave a conservative test for non-linearity under realistic violations of the key assumption. We found evidence for a non-linear causal effect of alcohol intake on several cardiovascular traits.
Conclusions

We believe our method is useful for inferring departure from linearity when only a binary instrument is available. We estimated non-linear causal effects of alcohol intake which could not have been estimated through standard instrumental variable approaches.

Key words: Mendelian randomisation; Instrumental variables; Causal inference; Local average treatment effects; Alcohol consumption; Cardiovascular disease.
Key messages

- Mendelian randomisation studies have so far restricted attention to linear associations relating the genetic instrument to the exposure, and the exposure to the outcome, but this may not always be appropriate. For example, alcohol consumption is consistently reported as having a U-shaped association with cardiovascular events in observational studies.

- We propose a novel Mendelian randomisation method based on estimating local average treatment effects for discrete levels of the exposure range, then testing for a linear trend in those effects.

- Our method gave a conservative test for non-linearity under realistic violations of the key assumption in simulations, and we believe our method is useful for inferring departure from linearity when only a binary instrument is available.

- We found evidence for a non-linear causal effect of alcohol intake on several cardiovascular traits in the Alcohol-ADH1B Consortium, using the single nucleotide polymorphism rs1229984 in ADH1B as a genetic instrument.
Introduction

Recent years have seen an increasing number of Mendelian randomisation (MR) analyses that examine causal relationships between heritable exposures, such as levels of circulating biomarkers, and outcomes such as multifactorial diseases, for example coronary heart disease and type 2 diabetes.\textsuperscript{1,2,3} In principle MR reduces problems of confounding and abolishes reverse causation by using a genetic proxy for the exposure in an instrumental variable (IV) analysis.\textsuperscript{4}

To date, applications of MR have been limited to linear (or log-linear) models for the associations between gene and exposure, and between exposure and outcome. In part this is because linear models have a natural interpretation which may be useful even if the true relationship is non-linear.\textsuperscript{5} Furthermore, many of the associations between genetic variants and complex traits discovered to date have appeared to be linear.\textsuperscript{6} However in learning about causal relationships it is clearly of value to identify and characterise non-linear effects when they are present, bearing in mind that the existence and extent of such relationships may depend on the measurement scale. In particular, non-linear associations may translate into opposing effects (protective as well as harmful) according to the level of the exposure. Such opposing effects have been observed in many observational studies examining the relationship between alcohol consumption and cardiovascular events.\textsuperscript{7} Specifically, light-to-moderate levels of alcohol consumption have been associated with decreased risk of cardiovascular events relative to non-drinkers, with increased risk only occurring at higher levels of consumption. This apparent protective effect of light-to-moderate alcohol consumption could be explained by several different mechanisms, and corresponding ‘J’ or ‘U’ shaped associations have been observed with cardiovascular risk factors including low-density lipoprotein particles,\textsuperscript{8} abdominal adiposity,\textsuperscript{9} C-reactive protein (CRP),\textsuperscript{10,11} and
triglycerides (TG). Similar observational associations were seen in our earlier analyses of ADH1B Consortium data (Holmes et al, Supplementary Appendix, Figure S3).

As these observational findings suggest that light-to-moderate consumption may be cardio-protective, it is of great interest to consumers, suppliers and policy makers to establish whether this pattern is causal. Confounding is plausible, since socio-economic groups that drink moderately may have other lifestyle factors that directly lead to lower rates of disease, and the relationship between confounders and alcohol may themselves be non-linear. Evidence for reverse causation is also well established, with those developing ill-health or commencing medication more likely to reduce or quit alcohol consumption (the ‘sick-quitters’ phenomenon).

Alcohol consumption is influenced by genetic variants that affect alcohol metabolism. Heritability of alcoholism has been estimated at 40-60%, and variants in ALDH2, ADH1B and ADH1C that encode for liver enzymes have been associated with decreased intake, via increased metabolism of alcohol to acetaldehyde or decreased acetaldehyde clearance, both leading to unpleasant side effects. In particular, ADH1B has been shown to be robustly associated with alcohol consumption and has been used in MR analyses to explore the causal effect of alcohol consumption on coronary heart disease risk factors.

We recently established a large consortium (the ‘Alcohol-ADH1B Consortium’) of genetic association studies of European descent that used a single nucleotide polymorphism (SNP) in ADH1B, rs1229984, as the instrument to assess the impact of alcohol consumption on cardiovascular events and risk factors. This consortium showed that carrying the rs1229984 A-allele was associated with non-drinking, lower alcohol consumption, and lower incidence
of binge drinking, which expands the previous associations of this variant with alcohol traits. Using a genetic association analysis, the consortium also showed that ADH1B carriers had a more favourable cardiovascular profile and a reduced risk of CHD. However, because of the existing literature on non-linear effects of alcohol consumption on cardiovascular events and the lack of appropriate methods to account for non-linear associations within IV analyses, we did not initially conduct an MR analysis in the Alcohol-ADH1B Consortium.

Approaches have been proposed for non-linear IV analysis in the econometric literature, but they cannot be used in this context because we use a single SNP as the IV. In the present paper, we develop new methods to conduct non-linear IV analysis using a single binary instrument, and also evaluate the impact of the key assumption of our method. We then apply our method to the data from the Alcohol-ADH1B Consortium to assess whether the causal effect of alcohol on cardiovascular traits is indeed non-linear and whether this implies a non-zero optimal level of consumption for cardiovascular health, which has clear implications for public health.

Material and methods

Data

The Alcohol-ADH1B Consortium is a collaboration of studies in which the associations between an allele of the ADH1B gene and twenty-two cardiovascular biomarkers, risk of coronary heart disease, stroke and type 2 diabetes have been examined. Here our analyses are restricted to the 22 studies (18 cohorts, 2 nested case-control studies, 1 randomized trial and 1 case-control study) with individual participant data originating from Europe (n = 16) and North America (n = 6). Analysis was restricted to individuals of European descent.
The principal alcohol trait was weekly volume of alcohol in British units (1 British unit is equivalent to 0.57 US units or 10 ml [7.9g ethanol]), which we derived using questionnaire data from each study. For studies in which this variable was not already present, we either calculated weekly volume of alcohol by summing over the individual components of beverage-specific drink questions (available in 20 of the 22 studies), or by converting alcohol recorded in grams/week into British units. The units/week were log transformed, after incrementing by one to allow for individuals reporting zero weekly alcohol consumption, resulting in a normally distributed phenotype that had homoscedastic residual error after regressing on the \( ADH1B \) genotype.

Here we considered a subset of outcomes for which a non-linear causal association was either postulated from subject-matter knowledge, or suggested by the observational data available from the Alcohol-\( ADH1B \) Consortium (all \( P < 0.001 \) for the quadratic term in a quadratic model): systolic blood pressure (SBP), non-high density lipoprotein cholesterol (non-HDL-C), TG, high density lipoprotein cholesterol (HDL-C), body mass index (BMI), waist circumference (WC), CRP and interleukin 6 (IL-6). Outcomes were log transformed towards normality when appropriate (TG, CRP and IL-6).

The \( rs1229984 \) polymorphism in \( ADH1B \) was directly genotyped in all studies and coded as 0/1 according to the carriage of at least one minor allele. This coding was adopted owing both to the low prevalence of the \( rs1229984 \) A-allele (average carriage of \( rs1229984 \) A-alleles in the analysis sample: 7.7%) and the stronger association observed with alcohol dependence and other alcohol-related traits under a dominant model compared to a recessive model.\(^{24} \)
Full details of participating studies, phenotype definition and genotyping are reported elsewhere\textsuperscript{13} and are summarised in Table S1 in the Supplementary Data.

**Linear instrumental variable analysis**

We used standard two-stage least squares (2SLS) to estimate a linear causal effect of log(weekly units of alcohol + 1) (hereafter, log-alcohol) on continuous cardiovascular outcomes. That is, we fitted the first stage linear regression

\[
x_i = \beta_{\text{XG}} g_i + \beta_{\text{XZ}} z_i + \epsilon_{X_i}
\]

where \(x_i\) is log-alcohol for subject \(i\), \(g_i\) is a binary code for the \texttt{rs1229984} genotype, \(z_i\) is a vector of covariates and \(\epsilon_{X_i}\) are residual errors assumed to be independent and identically distributed with mean zero. Regression coefficients \(\beta_{\text{XG}}\) and \(\beta_{\text{XZ}}\) were estimated as fixed effects. We used the fitted model to predict \(\hat{x}_i\) then estimated the alcohol-outcome association \(\beta_{XY}\) from the regression

\[
y_i = \beta_{\text{YX}} \hat{x}_i + \beta_{\text{YZ}} z_i + \epsilon_{Y_i}
\]

where \(y_i\) is the continuous cardiovascular outcome for subject \(i\) and \(\epsilon_{Y_i}\) are residual errors assumed to be independent and identically distributed with mean zero. A 95\% confidence interval for \(\hat{\beta}_{XY}\) was derived by nesting the 2SLS within a bootstrap resampling procedure.
using 10 000 bootstrap samples. As covariates we included in both regressions a fixed effect for each study, and fixed effects for age and sex.

**Non-linear causal effects**

To test for non-linearity of the causal $X$-$Y$ association we consider local average treatment effects (LATEs) in subgroups of $X$. First we coarsen $X$ into a discrete and rescaled variable 

$$X^* = \left[ \frac{X}{\beta_{X^*}} \right]$$

with finite support, assumed without loss of generality to be \{0,\ldots,J\} for fixed $J$. $G$ is an instrument for $X^*$ if it is independent of the remainder $X - X^*$ (see Figure 1); this is not generally true but it can be tested in applications. Under linear models we can obtain an estimate of the causal effect of $X^*$ on $Y$, but this effect can also be represented as a weighted sum of LATEs, which are causal effects among the individuals whose exposures $X^*$ are changed from one level to the next by the genetic instrument.

More precisely, let $Y_i(j)$ denote the potential outcome for subject $i$ obtained by setting, possibly contrary to fact, the exposure $X_i^* = j$. Moreover let $X_i^*(0)$ and $X_i^*(1)$ be the possibly counterfactual values of the exposure obtained by setting the binary instrument to 0 and 1 respectively. Then the LATE at exposure level $j$ is defined as

$$\tau_j = E[Y_i(j) - Y_i(j-1) \mid X_i^*(1) \geq j > X_i^*(0)]$$

that is, the average treatment effect among those whose exposure would be at least $j$ if their instrument were set to 1, and whose exposure would be less than $j$ if their instrument were set to 0. Identification of LATEs requires the further assumption of monotonicity, that is either
$X^*_i(1) - X^*_i(0) \geq 0$ or $X^*_i(1) - X^*_i(0) \leq 0$ for all subjects $i$, implying that the instrument either does not decrease the exposure in all subjects, or does not increase it in all subjects.

[Figure 1 here]

If we could estimate the LATEs $\tau_j$ then testing them for equality would provide a direct test of linearity of the causal effect. Here we propose an assumption that allows this to be performed. Assume that the causal effect of the instrument on the discretised exposure is exactly 1 in each subject:

$$X^*_i(1) - X^*_i(0) = 1 \forall i$$

This is a stronger version of the monotonicity assumption. In fact, this assumption will hold if the first stage linear model is a true structural model for $X$, with no unmeasured confounders of the $G$-$X$ association, or modifiers of the effect of $G$ on $X$. Under this assumption (and noting that $X$ has been rescaled so that a one unit change in $X^*$ corresponds to the expected exposure change with genotype), every subject contributes to a LATE, since for every $i$ there is a $j$ such that $X^*_i(1) \geq j > X^*_i(0)$, in fact $X^*_i(1) = j = X^*_i(0) + 1$. That is, the instrument moves each subject from one level of $X^*$ to the next: in the randomised trials terminology, all subjects are compliers.

It is now possible to assign each subject to the estimation of a LATE, based on the observed data. Since $X^*_i(1) = j = X^*_i(0) + 1$ if and only if $X^*_i = j$ and $G_i = 1$ or $X^*_i = j - 1$ and $G_i = 0$, we can write the LATE as
\[
\tau_j = E[Y_i(j) - Y_i(j - 1) \mid X_i^* (1) \geq j > X_i^* (0)]
\]
\[
= E[Y_i(j) \mid X_i^* = j, G_i = 1, G_i = 0] - E[Y_i(j - 1) \mid X_i^* = j, G_i = 1, G_i = 0]
\]
\[
= E[Y_i(j) \mid X_i^* = j, G_i = 1] - E[Y_i(j - 1) \mid X_i^* = j, G_i = 0]
\]

which may be estimated using ordinary linear regression (possibly with adjustment for relevant covariates) restricted to the subjects having \( X_i^* = j \) and \( G_i = 1 \) or \( X_i^* = j - 1 \) and \( G_i = 0 \).

Having estimated a LATE (with its standard error) for each level of \( X^* \), the estimates may be tested for equality using standard methods of meta-analysis. In particular, we use meta-regression to test for a linear trend in the LATEs. A linear model relating LATEs to the exposure levels \( j \)

\[
E(\tau_j) = \gamma_1 + \gamma_2 j
\]

would apply if the underlying causal model were quadratic

\[
E(Y) = \gamma_0 + \gamma_1 j + \frac{1}{2} \gamma_2 j^2
\]

The coefficient \( \gamma_2 \) is zero if the LATEs are equal, which is the case when the causal effect of \( X \) on \( Y \) is linear. Then the mean LATE, calculated by fixed-effects meta-analysis of the estimated LATEs, is an alternative measure of the linear causal effect of \( X \). Rejection of \( \gamma_2 = 0 \) implies a non-linear causal effect; a quadratic form is not directly implied but such a model could be hypothesised, up to its intercept term, from the fitted meta-regression. The
estimation of a linear model relating LATEs to the exposure levels is a simple but powerful way to investigate departures from linearity, as any such departures are captured by a single parameter. However, alternative models could be fitted to characterise the dose-response relationship more flexibly. For example, a piecewise constant model relating the LATEs to the exposure levels would correspond to a linear spline model relating the exposure to the outcome. This could be detected by a test of Cochran’s Q on the estimated LATEs.

This procedure requires rescaling of $X$ by the effect size $\beta_{XG}$ of the instrument. However the true value of $\beta_{XG}$ is unknown and it must be estimated. To account for sampling uncertainty in $\hat{\beta}_{XG}$ we nest the entire LATE and meta-regression procedure within a bootstrap resampling procedure, using 10 000 bootstrap samples, to obtain proper confidence intervals on the meta-regression estimates $\hat{\gamma}_1, \hat{\gamma}_2$. Our procedure for testing departure from linearity of the causal effect of $X$ on $Y$ is summarised in Box 1.

[Box 1 here]

Beyond a test for departure from linearity, we are interested in identifying the way the causal effect changes with increasing alcohol consumption and, in particular, the nadir of the curve which could be conceived as an ‘optimal’ level of consumption regarding cardiovascular traits. As we cannot estimate the intercept term in the fitted quadratic model, we cannot predict the absolute value of the outcome for a given level of alcohol consumption, so we focus on the difference in outcome relative to zero alcohol consumption. For those outcomes with evidence of non-linearity we predict this at four values of alcohol consumption (3.04, 12.15, 31.90 and 84.52 units/week), which are the medians of observed values in the categories representing low (>0-7 units/week), moderate (7-21 units/week), heavy (21-70
units/week) and very heavy (70+ units/week) alcohol consumption in the analysis of Holmes et al. By differentiation of the hypothesised quadratic function we estimate three additional features of the curve: i) the ‘optimal’ level of alcohol consumption; ii) the difference in outcome at the optimal alcohol consumption relative to zero alcohol consumption; iii) the level of alcohol consumption required to have an outcome level equivalent to that at zero alcohol consumption. Confidence intervals for all the estimates are obtained by nesting the estimation within the bootstrap resampling procedure outlined above. In the bootstrap samples we left truncated the nadir of alcohol consumption at zero.

All analyses were conducted using R version 2.13.

Simulations

We conducted simulations to assess the proposed approach in terms of bias and coverage under various data generating models. Full details and results are given in the Supplementary Data. In brief, we simulated data in which there was no causal X-Y association, in which the association was linear, and in which there was a quadratic causal association, allowing throughout for quadratic effects of confounders. We assessed robustness to the assumption of individual level homogeneity of the genetic effect using additional simulations of $\beta_{XG}$ heterogeneity and $G-U$ interaction at both the individual and subgroup level.

We observed that the LATE estimates were essentially unbiased with generally good coverage properties under null, linear and quadratic models, and that the test for a non-linear effect was slightly conservative. Together the results suggest that this method is a useful extension to standard approaches in the non-linear setting. Reasonable levels of individual-
level heterogeneity in $\beta_{xg}$ or between-subgroup heterogeneity in $\beta_{xg}$ were not found to lead to significant bias in the estimates. High levels of interaction between $G$ and $U$ led to bias in the estimates, but such interactions may be unlikely in practice.

**Results**

We investigated the potential non-linear effects of log-alcohol on each of the outcomes in the Alcohol-ADH1B Consortium using the proposed procedure. Some issues relating to the inclusion of multiple studies in the Consortium are discussed in the Supplementary Data.

Age- and sex-adjusted study-specific estimates of the association between rs1229984 and log-alcohol are presented in Figure S17 of the Supplementary Data. These study-specific estimates have (inverse-variance-weighted) mean -0.235 and SD 0.121, indicating some degree of between-study variability. However, in our simulations (see Supplementary Data) a similar degree of heterogeneity between known subgroups (scenario ‘f’ with $\gamma = 0.1$) was not found to result in bias to either the LATE intercept or slope, with slightly conservative confidence intervals for each.

To examine whether $G = rs1229984$ is a valid instrument for discretised $X^*$, assuming that it is valid for the continuous measure $X = \log$-alcohol, we examined the correlation between $G$ and the remainder $X - X^*$; these should be independent for $G$ to be a valid instrument for $X^*$. We observed a weak but significant correlation (Pearson’s $r = -0.013$, 95% CI: -0.020, -0.006). We hypothesised that this residual correlation was due to the large number of individuals reporting drinking zero weekly units of alcohol ($\log$-alcohol = 0), because these
individuals have a residual $X - X^* = 0$ and are also more likely to have $G = 0$. When individuals with log-alcohol = 0 were excluded from the analysis the correlation between $G$ and the remainder $X - X^*$ was close to zero (Pearson’s $r = 0.001$, 95% CI: -0.007, 0.009). We therefore re-analysed the data after excluding individuals with log-alcohol = 0, but obtained very similar results to those from the full sample. Because it is necessary to retain individuals reporting zero drinking to meet the objectives of the analysis, we only report results using the full sample.

The results of the LATE-based analysis for each of the outcomes are presented in Table 1 along with the standard linear IV analysis. We illustrate our approach in more detail using SBP as an example, following the steps in Box 1. We estimated $\hat{\beta}_{XG} = -0.244$ assuming a common genetic effect across all studies. Discretising log-alcohol into units of -0.244 gave an integer exposure $X^*$ with range [-26, 0]. We then estimated the LATE at each value of $X^*$. For example, for $j = -11$ (corresponding to a log-alcohol of $-11 \times -0.244 = 2.684$, or $\exp(2.684) - 1 = 13.6$ units/week) we selected the subjects with $X^* = -11$ and $rs1229984 = 1$, or $X^* = -12$ and $rs1229984 = 0$. Linear regression of SBP on $X^*$, on these subjects only, and adjusting for study, age and sex, gave $\tau_{-11} = -1.55$. That is, in subjects whose $X^*$ was changed from -12 to -11 by the SNP, their SBP was decreased by 1.55 mmHg.

[Table 1 here]

Rescaling by $\hat{\beta}_{XG} = -0.244$, subjects whose log-alcohol was changed from $-12 \times -0.244 = 2.928$ to $-11 \times -0.244 = 2.684$ (i.e. whose weekly units of alcohol consumption was changed from $\exp(2.928) - 1 = 17.7$ to $\exp(2.684) - 1 = 13.6$) by the SNP
have their SBP decreased by 1.55 mmHg. Alternatively, a one unit increase in log-alcohol at this level of alcohol consumption (e.g. from 2.684 to 3.684, or from \(\exp(2.684) - 1 = 13.6\) to \(\exp(3.684) - 1 = 38.8\) units/week – a considerable increase) was associated with an increase in SBP of \(-1.55/0.244 = 6.35\) mmHg.

The full graph of estimated LATEs for SBP is shown in Figure 2. Negative LATEs represent decreasing SBP with log-alcohol whilst positive LATEs represent increasing SBP, so a LATE trend crossing zero from negative to positive indicates a nadir. Fixed effects meta-analysis of these effects gave a mean LATE of 4.9 (95% CI: 2.6, 7.5), which is effectively a complier average treatment effect and similar to the linear IV estimate of 5.2 (95% CI: 3.2, 7.3). Meta-regression of the estimated LATEs on \(X^+\) gave a slope of 3.3 (95% CI: 1.0, 5.5). This provided strong evidence (Z-test \(p = 0.004\)) that the LATEs were not constant across values of log-alcohol; that is, there was a non-linear association between log-alcohol and SBP.

[Figure 2 here]

Full results for the remaining outcomes are provided in Table 1. As indicated by the LATE slope there was evidence of a non-linear causal effect for SBP, non-HDL-C, BMI, WC and CRP (all \(P \leq 0.01\)). For other outcomes there was no evidence of a non-linear causal effect (HDL-C, IL-6 and triglycerides, all \(P > 0.4\), though note that power is lower for IL-6 due to the relatively smaller sample size). In these cases we recommend that the linear IV results are employed as fewer assumptions are required in their estimation. It should also be noted that the linear IV estimates and the mean LATEs were similar for each of the outcomes, albeit with the latter having wider CIs.
Table 2 shows the predicted difference in each outcome relative to zero alcohol consumption for 3.04, 12.15, 31.90 and 84.52 units/week of alcohol consumption under the fitted quadratic functions. All outcomes, with the exception of SBP, were predicted to be lower at 3.04 units/week (‘low’ alcohol consumption) than at zero alcohol consumption, though each confidence interval included the possibility of no true difference. By 31.90 units/week (‘heavy’ alcohol consumption) all outcomes were predicted to be higher than at zero alcohol consumption, though each confidence interval, with the exception of SBP, again included the possibility of no true difference. By 84.52 units/week (‘very heavy’ alcohol consumption) all the confidence intervals excluded the possibility of no true difference.

[Table 2 here]

Table 2 also shows the additional estimated features of the hypothesised quadratic functions. For all outcomes the optimal level of alcohol consumption was estimated to be greater than zero, ranging from 1.0 units/week (SBP) to 3.5 units/week (CRP). However, only for non-HDL-C did the confidence interval exclude the possibility that zero consumption may be optimal. Correspondingly, the estimated difference in outcome at the optimal alcohol consumption level relative to zero consumption was negative for each outcome, though only for non-HDL-C did the confidence interval exclude the possibility of no true difference. The level of alcohol consumption required to have an outcome level equivalent to that at zero consumption was estimated as ranging from 2.8 units/week (SBP) to 19.4 units/week (CRP), though for all outcomes the confidence intervals were very wide. These results are illustrated for non-HDL-C, for which the strongest evidence of non-linearity was observed, in Figure 3. However, the precise values of our quantitative results should be interpreted with some
caution as the quadratic causal model that we fit may not be sufficiently flexible to fully characterise the dose-response relationship.

[Figure 3 here]

**Discussion**

We have proposed a method based on estimating LATEs that allows a basic estimation of local causal effects of a continuous exposure when using a binary instrument. Our method requires an assumption of homogeneous individual treatment effects of the instrument on the exposure, but our simulations found the estimates obtained under our approach to be largely unbiased and with good coverage properties under a variety of heterogeneous effects of instrument on exposure.

The local effects we estimate are within discretised units of the exposure, with the size of those units depending on the gene-exposure association. This is not a scale with a generally useful interpretation, and different genetic instruments could lead to different discrete units with different definitions of local causal effects. We therefore emphasise the ability to test for a non-linear causal effect and draw qualitative conclusions about the shape of that effect, and we suggest that a strictly quantitative interpretation of the estimated parameters should be viewed with some caution. Further work is required in investigating alternative models relating the LATEs to the exposure levels in order to provide greater flexibility for characterising the dose-response relationship.
Using this approach we detected evidence for a non-linear causal effect of log-alcohol on several cardiovascular traits in a large collaborative study, which would not have been possible using standard IV approaches. For each outcome that exhibited evidence of a non-linear causal effect, our results suggested that the level of alcohol consumption associated with the lowest value of the cardiovascular traits to lie between 1.0 and 3.5 units/week. However, only for non-HDL-C do we have strong evidence that the optimal level of consumption truly differs from zero.

As the cardiovascular traits considered in this analysis were observed concurrently with the level of alcohol consumption in many of the studies within the ADH1B Consortium, a conventional analysis would be at risk of bias due to reverse causality (for example, someone with high SBP reducing their alcohol intake so that they are observed to have a low level of consumption). A Mendelian randomisation analysis removes the possibility of such reverse causality, which is a significant strength of the present study.

For our estimated effects to be interpreted causally we need the standard assumptions underlying MR analysis to hold. Of particular concern in the present application is the exclusion restriction that $G$ has no effect on $Y$ other than through $X$. We have only considered one aspect of alcohol consumption (weekly units), but if the polymorphism in $ADH1B$ reduces alcohol consumption generally then other aspects, such as frequency of binge drinking, may also be associated with the instrument. If such other aspects have a causal effect on the outcome independently of weekly units then the exclusion restriction would not hold. The strong correlation between weekly units and other aspects of alcohol consumption makes a significant violation of this assumption unlikely. However, further research is required in this area.
Although we limited our analyses to individuals of European descent and adjusted for study in all our analyses, there may be residual population stratification of the variant which could lead to backdoor pathways from the instrument to the outcome. The restriction to individuals of European descent may also reduce the generalisability of our findings beyond such populations.

An inherent aspect of our approach is the need for a large sample with a sufficiently strong association between the gene and the exposure. If the gene-exposure association is very weak then the exposure will be discretised into many bins, none of which will contain sufficient subjects for the LATEs to be estimated. Many MR studies are now conducted on large samples in order to improve power to detect causal effects, but our approach requires large samples across a sufficient range of the exposure in order to detect non-linearities. This problem is compounded when studying binary outcomes, as each bin should contain a sufficient number of events. Therefore we have restricted our attention to continuous outcomes in this paper, but we recognise that here the key interest is in the nature of the causal relationship with cardiovascular disease events, which cannot be readily deduced from the associations with different risk factors. Further work in this area is required.

We believe our method is useful for inferring departure from linearity when only a binary instrument is available. Although there is clearly greater scope for bias than in standard IV analysis, we did not infer non-linear effects for several of the cardiovascular outcomes we considered, suggesting some degree of specificity using our method. More robust inference of non-linear causal effects may be possible from polychotomous or continuous instruments, such as gene scores constructed from multiple SNPs.²⁸,²⁹ Such instruments will allow the identification of non-linear models with many parameters, though IV estimation of parametric non-linear models has been found to be dependent on the choice of parametric model.²³ A further
key issue is whether the exposures predicted by those instruments cover a sufficient range to capture the non-linear features of the causal effects. If this is not the case, then it may be necessary to pursue approaches based on local effects similar to the one for binary instruments that we have discussed here.

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Table 1. Comparison of linear and non-linear instrumental variable estimates for selected cardiovascular traits in the Alcohol-ADH1B Consortium.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>n</th>
<th>Linear IV approach</th>
<th>Non-linear IV approach</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Estimate</td>
<td>95% CI</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>95% CI</td>
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<tr>
<td></td>
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<tr>
<td>SBP (mmHg)</td>
<td>78172</td>
<td>5.2</td>
<td>3.2, 7.3</td>
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<tr>
<td>Non-HDL-C (mmol/L)</td>
<td>60140</td>
<td>0.13</td>
<td>-0.02, 0.28</td>
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<tr>
<td>HDL-C (mmol/L)</td>
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<td>-0.02</td>
<td>-0.07, 0.03</td>
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<td>BMI (kg/m²)</td>
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<td>0.2, 1.2</td>
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<td>WC (cm)</td>
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<td>1.3, 4.4</td>
</tr>
<tr>
<td>CRP&lt;sup&gt;C&lt;/sup&gt; (mg/l)</td>
<td>63367</td>
<td>0.17</td>
<td>0.03, 0.31</td>
</tr>
<tr>
<td>IL-6&lt;sup&gt;C&lt;/sup&gt; (pg/ml)</td>
<td>23535</td>
<td>0.30</td>
<td>0.16, 0.45</td>
</tr>
<tr>
<td>TG&lt;sup&gt;C&lt;/sup&gt; (mmol/L)</td>
<td>63667</td>
<td>0.01</td>
<td>-0.06, 0.07</td>
</tr>
</tbody>
</table>

SBP, systolic blood pressure; Non-HDL-C, non-high density lipoprotein cholesterol; HDL-C, high density lipoprotein cholesterol; BMI, body mass index; WC, waist circumference; CRP, C-reactive protein; IL-6, interleukin 6; TG, triglycerides.

<sup>A</sup>Derived using 10,000 bootstrap samples. <sup>B</sup>Approximate Z-test using the bootstrap standard error. <sup>C</sup>Log transformed prior to analysis.
Table 2. Predicted difference in cardiovascular traits relative to zero alcohol consumption at several levels of alcohol consumption and predicted curve features in the Alcohol-ADH1B Consortium. Only calculated for traits with evidence of non-linearity.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Difference in outcome (95% CI(^a))</th>
<th>Level of alcohol consumption</th>
<th>Difference in outcome at optimal alcohol consumption (95% CI(^a))</th>
<th>Level of alcohol consumption with outcome equal to that at zero (95% CI(^a))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3.04 units/week(^c)</td>
<td>12.15 units/week(^c)</td>
<td>31.90 units/week(^c)</td>
<td>84.52 units/week(^c)</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>0.1 (-5.5, 6.1)</td>
<td>5.2 (-2.6, 12.4)</td>
<td>22.8 (12.2, 34.6)</td>
<td>1.0 (0.0, 3.6)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>13.9</td>
<td>22.1</td>
<td>34.6</td>
</tr>
<tr>
<td>Non-HDL-C (mmol/L)</td>
<td>-0.39 (-0.79, 0.15)</td>
<td>-0.15 (-0.72, 0.40)</td>
<td>1.30 (0.45, 3.2)</td>
<td>3.2 (0.7, 6.0)</td>
</tr>
<tr>
<td></td>
<td>0.06</td>
<td>0.47</td>
<td>1.10</td>
<td>2.16</td>
</tr>
<tr>
<td>BMI (kg/m(^2))</td>
<td>-0.6 (-2.2, 0.8)</td>
<td>0.2 (-2.0, 1.6)</td>
<td>2.3 (0.0, 6.3)</td>
<td>-0.6 (-2.3, 0.0)</td>
</tr>
<tr>
<td></td>
<td>1.6 (-0.8, 3.8)</td>
<td>1.2 (1.2, 3.6)</td>
<td>10.1 (0.0, 48.4)</td>
<td></td>
</tr>
<tr>
<td>WC (cm)</td>
<td>-0.6 (-4.7, 3.5)</td>
<td>1.9 (-3.9, 7.8)</td>
<td>5.7 (-0.6, 11.5)</td>
<td>1.5 (0.0, 5.4)</td>
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<tr>
<td></td>
<td></td>
<td>12.5</td>
<td>19.2</td>
<td></td>
</tr>
<tr>
<td>CRP(^b) (mg/l)</td>
<td>-0.29 (-0.68, 0.15)</td>
<td>-0.15 (-0.68, 0.22)</td>
<td>0.83 (0.15, 3.5)</td>
<td>-0.30 (-0.75, 0.00)</td>
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<td>0.15</td>
<td>0.50</td>
<td>19.4 (0.0, 66.0)</td>
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</tbody>
</table>

SBP, systolic blood pressure; Non-HDL-C, non-high density lipoprotein cholesterol; BMI, body mass index; WC, waist circumference; CRP, C-reactive protein.
A Derived using 10,000 bootstrap samples. B Log transformed prior to analysis. C Weekly units of alcohol values are medians of observed values in categories representing low (1-7 units/week), moderate (7-21 units/week), heavy (21-70 units/week) and very heavy (70+ units/week) alcohol consumption in the analysis of Holmes et al. 13
Box 1. Summary of proposed method for testing for a non-linear causal effect.

1. For the observed data and for each of \( K \) bootstrap samples:
   1.1. Regress \( X \) on \( G \) for all subjects, giving estimated regression coefficient \( \hat{\beta}_{XG} \)
   1.2. Discretise \( X \) into units of \( \hat{\beta}_{XG} \), that is derive the discrete variable \( X^* = \frac{X}{\hat{\beta}_{XG}} \)
   1.3. For each discrete value of \( j \):
      1.3.1. Regress \( Y \) on \( X^* \) using only the subjects for which \( X^*_i = j \) and \( G_i = 1 \), or \( X^*_i = j - 1 \) and \( G_i = 0 \). Among these subjects there is no variation in \( X^* \) that is not explained by \( G \).
      1.3.2. This yields \( \hat{\tau}_j \), the estimated local average treatment effect (LATE) for level \( j \) of \( X^* \)
      1.3.3. Rescale \( \hat{\tau}_j \) by \( \hat{\beta}_{XG} \) to the original scale of \( X \)
   1.4. Obtain the mean LATE by fixed-effects meta-analysis of \( \hat{\tau}_j \)
   1.5. Meta-regress \( \hat{\tau}_j \) on \( j \) to obtain the intercept and slope of the LATEs, corresponding to a quadratic causal model
2. Obtain empirical confidence intervals on the mean LATE and the LATE intercept and slope from the bootstrap samples
Figure legends

Figure 1. Directed acyclic graphs encoding a) the standard Mendelian randomisation assumptions: (i) G is associated with X, ii) G is not associated with confounders U of the X-Y association, and iii) G affects Y only via its association with X; b) how these are assumptions are affected by the discretisation of X in the proposed non-linear Mendelian randomisation approach.

Figure 2. Local average treatment effects (LATEs) of log(weekly units of alcohol + 1) on systolic blood pressure. Circular markers are LATEs; bars are 95% pointwise confidence intervals; dashed line is estimated mean LATE; solid line is estimated linear LATE trend; dotted line is linear IV estimate using the ratio method (virtually indistinguishable from the estimated mean LATE).

Figure 3. Predicted difference in non-high density lipoprotein cholesterol (non-HDL-C) relative to zero alcohol consumption across the range of values of observed alcohol consumption, with estimated optimal level of alcohol consumption (3.2 (95% confidence interval (CI): 0.7, 6.0) units/week), estimated difference in non-HDL-C relative to zero alcohol consumption at optimal level (-0.39 (95% CI: -0.85, -0.03) mmol/L), and estimated level of alcohol consumption with the same level of non-HDL-C as at zero (16.9 (95% CI: 2.1, 48.2) units/week) indicated.
References


Testing for non-linear causal effects using a binary genotype in a Mendelian randomisation study: application to alcohol and cardiovascular traits

Richard J Silverwood, Michael V Holmes, Caroline E Dale, Debbie A Lawlor, John C Whittaker, George Davey Smith, David A Leon, Tom Palmer, Brendan J Keating, Luisa Zuccolo, Juan P Casas, Frank Dudbridge, on behalf of the Alcohol-ADH1B Consortium

Supplementary material

Simulations – Methods

We conducted simulations to assess the proposed approach in terms of bias and coverage at a variety of sample sizes and under different X-Y associations. We performed additional simulations to explore the effects of $\beta_{xg}$ heterogeneity and $G-U$ interaction at both the individual and subgroup level.

The simulation parameters were chosen to resemble those observed in the Alcohol-ADH1B Consortium, and a plausible degree of confounding was applied. In each simulation the data were generated according to

$$Y \sim N(\beta_1X + \beta_2X^2 - 2U + U^2, S^2)$$

with the following specifications remaining constant:
Thus, we allow for a quadratic causal effect of $X$ on $Y$, but also for quadratic effects of confounders. The following simulations were conducted:

a) Sample size. The effect of sample size was examined using the null model with $(\beta_1, \beta_2) = (0, 0)$. Sample size was varied within \{5000, 10000, 50000, 100000\}. $X$ was simulated using $X \sim N(-0.2G + 0.5U, 1)$.

b) $X$-$Y$ associations. A variety of $X$-$Y$ associations were explored using a sample size of 10000. Linear associations were generated using $(\beta_1, \beta_2) = \{(-5, 0), (-2, 0), (-1, 0), (1, 0), (2, 0), (5, 0)\}$; quadratic associations were generated using $(\beta_1, \beta_2) = \{(2, -5), (1, -2), (2, -1), (-2, 1), (-1, 2), (-2, 5)\}$. $X$ was again simulated using $X \sim N(-0.2G + 0.5U, 1)$.

c) Individual-level $\beta_{XG}$ heterogeneity. The key assumption of our approach is that $\beta_{XG}$ is constant for all individuals. The degree of variability introduced into these simulations regarding individual-level $\beta_{XG}$ heterogeneity was informed by the observation that within the ADH1B Consortium the study-specific estimates of the association between rs1229984 and log-alcohol had mean -0.235 and SD 0.121, along with the assumption that individual-level heterogeneity is likely to be somewhat greater than this. We defined $X$ using $X \sim N((-0.2 + \alpha)G + 0.5U, 1)$ where $\alpha \sim N(0, \gamma^2)$ was simulated independently for each subject within a sample size of 10,000 and $\gamma$ varied within \{0, 0.1, 0.2, 0.3, 0.5, 1\}. Thus, for $\gamma = 0.1$ $\beta_{XG}$ is expected to lie between -0.4 and 0 in 95% of subjects, and approximately 61% of subjects are
expected to be compliers in the sense that they are moved by the instrument to the adjacent bin of size $\beta_{XG} = -0.2$. Simulations were conducted using the quadratic data generating model with $(\beta_1, \beta_2) = (-1, 2)$.

d) Individual-level $G-U$ interaction. We defined $X$ using

$$X \sim N(-0.2G + 0.5U + \gamma GU, 1)$$

where $\delta$ varied within \{0, 0.1, 0.2, 0.3, 0.5, 1\} in a sample of size 10,000. This means that the effect of $G$ on $X$ is allowed to vary by levels of the confounder $U$. Simulations were conducted using the quadratic data generating model with $(\beta_1, \beta_2) = (-1, 2)$.

e) $\beta_{XG}$ heterogeneity between unknown subgroups. The degree of variability introduced into these and subsequent simulations regarding $\beta_{XG}$ heterogeneity between known or unknown subgroups were informed by the observation that within the ADH1B Consortium the study-specific estimates of the association between rs1229984 and log-alcohol had mean -0.235 and SD 0.121. We defined $X$ using

$$X \sim N((-0.2 + \alpha)G + 0.5U, 1)$$

where $\alpha \sim N(0, \gamma^2)$ took the same value within subgroups of 1000 subjects within an overall sample size of 10,000 and $\gamma$ varied within \{0, 0.01, 0.02, 0.03, 0.05, 0.1\}. Thus, for $\gamma = 0.1$ $\beta_{XG}$ is expected to lie between -0.4 and 0.0 in 95% of subgroups. Simulations were conducted using the quadratic data generating model with $(\beta_1, \beta_2) = (-1, 2)$. Data were analysed as previously described, with all subgroups pooled together.

f) $\beta_{XG}$ heterogeneity between known subgroups. $X$ was again defined using

$$X \sim N((-0.2 + \alpha)G + 0.5U, 1)$$

where $\alpha \sim N(0, \gamma^2)$ now took the same value within subgroups of 10,000 subjects within an overall sample size of 100,000 and $\gamma$ varied
between \{0, 0.01, 0.02, 0.03, 0.05, 0.1\}. Simulations were again conducted using the quadratic data generating model with \((\beta_1, \beta_2) = (-1, 2)\). Data were analysed with all subgroups pooled together and adjustment for subgroup using indicator variables.

g) **G-U interaction which varies between unknown subgroups.** In addition to individual-level G-U interaction, the degree of interaction may also vary between subgroups in a population. We investigated this issue in the situation where the subgroups between which the G-U interaction varied are unknown. We defined \(X\) using

\[ X \sim N(-0.2G + 0.5U + \delta GU, 1) \]

where \(\delta \sim N(0, \gamma^2)\) took the same value within subgroups of 1000 subjects within an overall sample size of 10,000 and \(\gamma\) varied within \{0, 0.01, 0.02, 0.03, 0.05, 0.1\}. Simulations were conducted using the quadratic data generating model with \((\beta_1, \beta_2) = (-1, 2)\). Analysis proceeded as previously described, with data from all subgroups pooled together.

When regressing \(Y\) on \(X^*\) we only estimated a LATE if there were at least 5 subjects with \(X^*_i = j\) and \(G_i = 1\) and 5 subjects with \(X^*_i = j - 1\) and \(G_i = 0\), as regressions using smaller numbers of subjects were found to lead to anomalous estimated LATEs with spuriously high precision. A thousand simulations of each specification were conducted, with percentile bootstrap confidence intervals derived using 1000 bootstrap samples within each simulation. Standard 2SLS and quadratic OLS estimates were also calculated for comparison.

The analysis was conducted using R version 2.13.1
Simulations - Results

a) Sample size. Results for varying sample size are presented in Figure S1. The linear IV estimate, mean LATE, LATE intercept and LATE slope were unbiased even with a sample size of 5000. The variability in the mean LATE and LATE slope was similar to that of the linear IV estimate. The quadratic OLS estimates were significantly biased at all sample sizes. The corresponding coverages are shown in Figure S2. The linear IV estimate, mean LATE and LATE intercept coverages varied between 94% and 96%, even at smaller sample sizes. The coverage of the LATE slope was somewhat conservative (around 97%) at smaller sample sizes.

b) X-Y associations. In the linear data generating models the linear IV estimate, mean LATE, LATE intercept and LATE slope were all unbiased (Figure S3). The mean LATE and LATE slope again had similar variability to the linear IV estimate. The quadratic OLS estimates were biased in all scenarios. The coverages of the linear IV estimate, mean LATE, and LATE intercept were generally between 94% and 96%, but the coverage of the LATE slope was approximately 96% across the range of linear coefficients (Figure S4). This suggests that our procedure gives a slightly conservative test of no non-linear effect. In the quadratic data generating models the LATE intercepts and LATE slopes were unbiased for all combinations of coefficients but the quadratic OLS estimates were again heavily biased (Figure S5). The coverages of the LATE intercept and LATE slope were generally between 94% and 96%, though the LATE slope coverage was as high as 97.5% for \((\beta_1, \beta_2) = (-2, 5)\) (Figure S6).

c) Individual-level \(\beta_{XG}\) heterogeneity. For small values of \(\gamma\) the LATE intercept and LATE slope displayed little bias (Figure S7). However, for values of \(\gamma\) of 0.2 and greater noticeable bias began to appear, particularly in the LATE slope. By \(\gamma = 1\) the
bias was so great that the 95% range of estimates of the LATE slope did not include
the target value. Both the LATE intercept and LATE slope had acceptable levels of
coverage for values of $\gamma$ up to 0.3 (Figure S8). For values of $\gamma$ larger than 0.5 the
coverage of the LATE slope was drastically reduced. However, it should be noted that
the larger values of $\gamma$ in these simulations do represent very extreme cases (e.g. for
$\gamma = 1$ $\beta_{XG}$ is expected to lie between -2.2 and 1.8 in 95% of subjects, so in almost
50% of subjects the gene would have an effect on alcohol consumption in the opposite
direction to that assumed). Thus, given the reliance of our method on the assumption
of $\beta_{XG}$ homogeneity, the observed bias and reduced coverage is not unexpected.

d) *Individual-level G-U interaction.* Bias was observed for all values of $\delta$ greater than
zero for both the LATE intercept and the LATE slope (Figure S9). The bias in the
LATE intercept increased as $\delta$ increased, but the bias in the LATE slope decreased
slightly at $\delta = 1$. The coverage of the parameters correspondingly generally deviated
further from 95% as $\delta$ increased (Figure S10), reaching 74% for the LATE intercept
and 59% for the LATE slope at $\delta = 0.5$. Coverage for the LATE intercept reduced
further to 14% at $\delta = 1$, but improved slightly for the LATE slope.

e) $\beta_{XG}$ heterogeneity between unknown subgroups. At no values of $\gamma$ was there any
appreciable bias in either the LATE intercept or the LATE slope (Figure S11). For all
values of $\gamma$ the LATE intercept had coverage close to 95%, but there was again some
evidence that the coverage of the LATE slope was slightly conservative (Figure S12).

f) $\beta_{XG}$ heterogeneity between known subgroups. At no values of $\gamma$ was there any
evidence of bias in either the LATE intercept or the LATE slope (Figure S13).
Coverage was approximately 96\% for the LATE intercept and 97-99\% for the LATE slope (Figure S14).

g) \textit{G-U interaction which varies between unknown subgroups.} There was no bias in either parameter at $\gamma = 0.1$, but bias in both parameters, most noticeably in the LATE slope, increased as $\gamma$ increased further (Figure S15). Coverage was approximately appropriate for the LATE intercept as far as $\gamma = 0.5$, though it was $<90\%$ for the LATE slope from $\gamma = 0.3$ onwards (Figure S16). By $\gamma = 1$ the coverages of the LATE intercept and slope were 84\% and 23\% respectively.

These results suggest that the LATE estimates are essentially unbiased with generally good coverage properties under null, linear and quadratic models. Reasonable levels of individual-level heterogeneity in $\beta_{xG}$ were not found to lead to significant bias in the estimates. Low levels of between-subgroup heterogeneity in $\beta_{xG}$ were also not found to lead to significant bias, whether or not the heterogeneity was adjusted for in the analysis. High levels of interaction between $G$ and $U$ led to bias in the estimates, but such interactions may be unlikely in practice. For example, in simulation d) a $\delta$ of 0.2 biased the LATE intercept upwards by 30\% and the LATE slope upwards by 16\%. This level of interaction means that the effect of $G$ on $X$ can be expressed as $-0.2 + 0.2U$. As $U \sim N(0,1)$, 95\% of the values of $U$ will lie between -2 and 2 (approximately). Thus the effect of $G$ on $X$ will lie between -0.6 and 0.2 approximately 95\% of the time, depending on the value of $U$. Such a wide range of genetic effects, including a reversal of sign, may well be deemed implausible.

Overall, our simulations indicate that the LATE method is a useful extension to standard approaches in the non-linear setting.
Multiple studies

When data from multiple studies are available we must decide whether to estimate LATEs within each study, combining them to draw an overall conclusion, or to estimate LATEs on the combined data. In general, weak instrument bias is reduced by performing the MR analysis on the combined data.2 But because our approach relies on the genetic effect size $\beta_{XG}$ being constant for all individuals, we need to consider whether this assumption is tenable across studies as well as within each study.

If $\beta_{XG}$ varies across studies then clearly we should perform the procedure separately in each study. This however raises two problems. Firstly, the restriction to subjects having particular genotype-exposure combinations in each bin of $X^+$ leads to small sample sizes for estimating some LATEs, leading to large standard errors on some $\hat{\tau}_j$ and considerable uncertainty on the final inference of non-linearity. This problem occurs particularly when one genotype is rare, or when $\beta_{XG}$ is small, leading to narrow bin definitions, both of which apply to the Alcohol-ADH1B Consortium. Secondly, different $\beta_{XG}$ across studies leads to different bin sizes across studies and different local causal effects being estimated. It is not clear how such effects should be combined into an overall inference on non-linearity.

For these reasons we assumed that $\beta_{XG}$ is constant within and between studies and performed simulations to assess robustness to that assumption. As in the standard IV analysis, covariates for study, age and sex were included in both the estimation of $\beta_{XG}$ and the LATEs in order to reduce the potential for confounding of $\beta_{XG}$ and the LATEs.
Table S1. Design and genotyping characteristics of the studies included in the analysis.

<table>
<thead>
<tr>
<th>Study</th>
<th>Study design</th>
<th>Sampling Frame</th>
<th>Number with DNA in this analysis</th>
<th>Number contributing to one or more of our analyses</th>
<th>Year of blood sampling used for DNA extraction</th>
<th>Genotyping method</th>
<th>Country</th>
<th>HWE P value (exact significance probability)</th>
<th>Call rate (%)</th>
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<td>Cohort</td>
<td>Community</td>
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<td>0.012</td>
<td>97</td>
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<td>NPHS II</td>
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<td>2000</td>
<td>TaqMan</td>
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<td>UK</td>
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<td>99.3</td>
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<td>Community</td>
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<td>1993-1998</td>
<td>IBC 50k CardioChip</td>
<td>USA</td>
<td>3.15E-25</td>
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*aNumber of study members non-missing for weekly volume of alcohol, rs1229984 polymorphism in ADH1B, age, sex, and one or more of the outcomes (systolic blood pressure, non-high density lipoprotein cholesterol, high density lipoprotein cholesterol, body mass index, waist circumference, C-reactive protein, interleukin 6 and triglycerides).
Figure S1. Observational quadratic model linear parameter estimates (brown) and quadratic parameter estimate (purple), two-stage least squares estimates (red), mean local average treatment effects (LATEs) (blue), LATE intercepts (green) and LATE slopes (orange) for different sample sizes in the null data generating model. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S2. 95% confidence interval coverage for the two-stage least squares estimates (red), mean LATEs (blue), LATE intercepts (green) and LATE slopes (orange) for different sample sizes in the null data generating model. Horizontal solid line represents the target value (95%).
Figure S3. Observational quadratic model linear parameter estimates (brown) and quadratic parameter estimate (purple), two-stage least squares estimates (red), mean LATEs (blue), LATE intercepts (green) and LATE slopes (orange) for different linear coefficients in the linear data generating model. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S4. 95% confidence interval coverage for the two-stage least squares estimates (red), mean LATEs (blue), LATE intercepts (green) and LATE slopes (orange) for different linear coefficients in the linear data generating model. Horizontal solid line represents the target value (95%).
Figure S5. Observational quadratic model linear parameter estimates (brown) and quadratic parameter estimate (purple), two-stage least squares estimates (red), mean LATEs (blue), LATE intercepts (green) and LATE slopes (orange) for different combinations of linear and quadratic coefficients in the quadratic data generating model. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S6. 95% confidence interval coverage for the LATE intercepts (green) and LATE slopes (orange) for different combinations of linear and quadratic coefficients in the quadratic data generating model. Horizontal solid line represents the target value (95%).
Figure S7. LATE intercepts (green) and LATE slopes (orange) for different degrees of individual-level heterogeneity of $\beta_{xg}$ in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S8. 95% confidence interval coverage for LATE intercepts (green) and LATE slopes (orange) for different degrees of individual-level heterogeneity of $\beta_{xy}$ in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Horizontal solid line represents the target value (95%).
Figure S9. LATE intercepts (green) and LATE slopes (orange) for different degrees of individual-level G-U interaction in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S10. 95% confidence interval coverage for LATE intercepts (green) and LATE slopes (orange) for different degrees of individual-level G-U interaction in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Horizontal solid line represents the target value (95%).
Figure S11. LATE intercepts (green) and LATE slopes (orange) for different degrees of heterogeneity of \( \beta_{xg} \) in unknown subgroups in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Points represent means and bars represent 95\% of the data. Horizontal solid lines represent target values.
Figure S12. 95% confidence interval coverage for LATE intercepts (green) and LATE slopes (orange) for different degrees of heterogeneity of $\beta_{sg}$ in unknown subgroups in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Horizontal solid line represents the target value (95%).
Figure S13. LATE intercepts (green) and LATE slopes (orange) for different degrees of heterogeneity of $\beta_{xg}$ in known subgroups in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S14. 95% confidence interval coverage for LATE intercepts (green) and LATE slopes (orange) for different degrees of heterogeneity of $\beta_{xg}$ in known subgroups in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Horizontal solid line represents the target value (95%).
Figure S15. LATE intercepts (green) and LATE slopes (orange) for different degrees of G-U interaction which varies between unknown subgroups in the quadratic data generating model with linear coefficient $\beta = -1$ and quadratic coefficient $\gamma = 2$. Points represent means and bars represent 95% of the data. Horizontal solid lines represent target values.
Figure S16. 95% confidence interval coverage for LATE intercepts (green) and LATE slopes (orange) for different degrees of G-U interaction which varies between unknown subgroups in the quadratic data generating model with linear coefficient = -1 and quadratic coefficient = 2. Horizontal solid line represents the target value (95%).
Figure S17. Study-specific estimates of the association between rs1229984 and log-alcohol.
References
