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Liver function and risk of type 2 diabetes: bidirectional Mendelian randomization study

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Figure 2
Liver dysfunction and type 2 diabetes (T2D) are consistently associated. However, it is currently unknown whether liver dysfunction contributes to, results from or is merely correlated with T2D due to confounding. We used Mendelian randomization (MR) to investigate the presence and direction of any causal relation between liver function and T2D risk including up to 64,094 T2D cases and 607,012 controls. Several biomarkers were used as proxies of liver function [i.e. alanine aminotransferase (ALT), aspartate aminotransferase (AST), alkaline phosphatase (ALP), and gamma-glutamyl transferase (GGT)]. Genetic variants strongly associated with each liver function marker were used to investigate the effect of liver function on T2D risk. In addition, genetic variants strongly associated with T2D risk and with fasting insulin were used to investigate the effect of predisposition to T2D and insulin resistance, respectively, on liver function. Genetically predicted higher circulating ALT and AST were related to increased risk of T2D. There was a modest negative association of genetically predicted ALP with T2D risk and no evidence of association between GGT and T2D risk. Genetically predisposition to higher fasting insulin, but not to T2D, was related to increased circulating ALT. Since circulating ALT and AST are markers of NAFLD, these findings provide some support for insulin resistance resulting in NAFLD, which in turn increases T2D risk.
INTRODUCTION

Observational studies have repeatedly reported that liver dysfunction and type 2 diabetes (T2D) are associated (1-3). Since the liver plays a core role in the regulation of glucose homeostasis, it is hypothesized that liver dysfunction might increase T2D risk by exacerbating hepatic insulin resistance, leading to overstimulation of hepatic gluconeogenesis (4). Alternatively, it is suggested that insulin resistance and T2D might disturb liver function, possibly via an effect of chronic inflammation and immunological changes (5, 6), as well as by directly upregulating hepatic lipogenesis (4). It is currently unknown whether liver dysfunction contributes to, or results from T2D development, or whether there is a genuine bidirectional relationship, where insulin resistance facilitates fat accumulation in the liver, which in turn leads to hepatic insulin resistance and increased fasting glucose (7-9). As most evidence to date is from observational studies, it is possible that the association between liver dysfunction and T2D reflects underlying common causes, such as obesity or lifestyle characteristics.

Plasma concentration of liver enzymes (i.e. alanine aminotransferase (ALT), aspartate aminotransferase (AST), alkaline phosphatase (ALP), and gamma-glutamyl transferase (GGT)) are routinely measured clinical markers that represent different dimensions of liver dysfunction. ALT, located in the cytosol, and AST, located in the mitochondria, are released from damaged hepatic cells into the blood after hepatocellular injury or death. ALT and AST are potentially useful surrogates for alcohol-induced liver disease and non-alcoholic fatty liver disease (NAFLD), defined as hepatic steatosis in the absence of excessive alcohol consumption. ALP is present in the ducts of the liver and GGT is located on liver cell membranes. The combined elevation of ALP and GGT can indicate obstructive or cholestatic liver disease, where bile is not properly transported from the liver because of obstruction of the bile duct. GGT is also an indicator of alcohol use (10).
Classical observational studies show that plasma concentrations of these enzymes, even within the normal range, are positively associated with type 2 diabetes (1-3). Mendelian randomization (MR), where genetic variants that are strongly associated with a risk factor of interest are used to test its causal effect on an outcome, can help to distinguish causal effects from associations due to confounding or reverse causality (11, 12). Previous MR studies do not support a link between circulating GGT or ALP on T2D risk or glycemic status in Europeans (13, 14) or of ALT on T2D risk or glycemic status in Chinese (15). In contrast, MR studies reported some evidence of a positive relation of circulating GGT on T2D risk in South Koreans (16) and on insulinaemia in Europeans (17) and of circulating ALT on T2D risk in Europeans (14). To the best of our knowledge, no MR study has investigated the effect of predisposition to T2D or to insulin resistance on liver function markers.

We have used the largest available datasets to interrogate the potential effect of liver dysfunction, proxied by multiple biomarkers (ALT, AST, ALP, and GGT), on T2D risk (64,094 T2D cases and 607,012 controls), as well as on related outcomes (fasting glucose, insulin and lipids). In addition, we have investigated whether predisposition to T2D and insulin resistance affect circulating liver function markers (ALT, AST, ALP, and GGT).
RESEARCH DESIGN AND METHODS

Study design

We explored the relationship of four liver function markers (plasma concentration of ALT, AST, ALP, and GGT) with T2D (primary outcome) and with six related metabolic traits (secondary outcomes) reflecting hyperglycemia, assessed by fasting glucose, insulin resistance, assessed by fasting insulin, and dyslipidemia, assessed by LDLc, HDLc, total cholesterol, triglycerides, using two approaches – multivariable regression and MR. We also used MR to investigate whether predisposition to T2D and to insulin resistance are likely to have an impact on circulating ALT, AST, ALP, and GGT. The hypotheses, study design and data sources used are detailed in Figure 1.

Data sources

Participant-level data

The UCL-LSHTM-Edinburgh-Bristol (UCLEB) consortium consists of 12 prospective observational studies comprising over 30,000 participants (18). For the present study, data from up to seven UCLEB studies were included in multivariable and MR analyses: the British Regional Heart Study (BRHS) (19), British Women’s Heart and Health Study (BWHHS) (20), Caerphilly Prospective Study (CaPS) (21), Edinburgh Artery Study (EAS) (22), English Longitudinal Study of Ageing (ELSA) (23), Whitehall II study (WHII) (24), MRC National Survey of Health and Development (NSHD) (25). Full details of the studies included in the UCLEB consortium have been described previously (18). For the multivariable analyses of liver marker - T2D associations, we used data from up to 6,593 individuals (728
T2D cases) from up to five UCLEB studies (BRHS, BWHHS, EAS, CaPS, and NSHD). For the MR analyses of the effect of liver function on T2D risk, we used up to 11,790 individuals (1,202 T2D cases) from up to seven UCLEB studies (BRHS, BWHHS, CaPS, EAS, ELSA, NSHD, and WHII) where information on genotypes and a liver function marker/s, and/or information on genotypes and outcome measure/s were available.

The **Fenland study** is a UK population-based study based in the East Cambridgeshire and Fenland areas and has been described in detail elsewhere (26). Data from Fenland study participants were included in both the multivariable (up to 9,968 individuals) and the MR analyses of the liver marker-lipid outcome associations (up to 9,982 individuals) except for AST.

Full details of the exposure, outcome and confounder variables available in each UCLEB and Fenland study are given in **Supplementary table 1** and participant characteristics, in **Supplementary tables 2a and 2b**.

**Summary-level data**

For the multivariable analysis, we pooled our individual participant-level results (from UCLEB and Fenland studies) with those from two published meta-analysis of the association of liver function markers and T2D risk: Kunutsor *et al.* (2) for ALT and AST (up to 60,359 participants including 3,890 incident T2D cases) and Fraser *et al.* (27) for GGT (up to 63,285 including 2,805 incident T2D cases).

For the MR analysis, we also used publicly available summary level data from genome-wide association studies (GWAS) for the association of single nucleotide polymorphisms (SNPs) with exposures and outcomes from the relevant GWAS studies/consortia. Summary statistics for the association between SNP and liver function markers were extracted from Chambers *et al.* (28), including 61,089 individuals with information of ALT, ALP, AST, and
GGT plasma concentration. Summary data for the association between SNPs and T2D was extracted from a consortium (29) including 62,892 cases and 596,424 controls, mostly of European descent. In cases where a liver function SNP could not be found in this latest T2D GWAS, data was extracted from a previous T2D GWAS from the DIAGRAM consortium including 34,840 cases and 114,981 controls (30). Summary statistics for the association of SNPs with fasting glucose and with fasting insulin were obtained from MAGIC (the Meta-Analyses of Glucose and Insulin-related traits Consortium) consortium (31, 32), which included up to 133,010 and 108,557 nondiabetic participants of European ancestry, respectively. Summary data for the association of SNPs with LDLc, HDLc, total cholesterol, and triglycerides were extracted from the GLGC (Global Lipids Genetics Consortium) consortium (33), including 188,577 individuals mostly of European ancestry. We excluded Fenland participants who had been included in the meta-analyses of the GLGC consortium to avoid having duplicated information from these participants.

**Definition of diabetes**

In UCLEB studies, T2D definition varied by study and included self-report, medical history review, use of glucose lowering medication, or having a fasting glucose value of $\geq 7$ mmol/L (18). For the MR analysis, we used both prevalent and incident cases of type 2 diabetes to maximise power, as the prevalent diabetes cases cannot influence genetic variation, which is fixed at conception.

In the GWAS, criteria for defining T2D differed across studies and included previous diagnosis, fasting glucose $\geq 7$ mmol/L, treatment with glucose-lowering medication, or self-reported T2D status (29, 30, 34).

**Genotyping and quality control**
All studies in UCLEB consortium were genotyped using the Illumina CardioMetabochip (Illumina, San Diego, CA, USA). Details on the genotyping and imputation quality control criteria used have been previously published (18, 35). Genotyping and imputation of missing genotypes in the published GWAS studies used here are described in the original publications (28, 30-33, 36, 37).

**Statistical analyses**

All continuous variables in the UCLEB and Fenland studies that were not normally distributed were natural log transformed. All continuously measured traits were standardised within each study to allow comparison between studies. In any published GWAS that did not report effects in standard deviation units, effects were standardised based on the GWAS reported standard deviation or, where this was not available, the median standard deviation across UCLEB studies.

For analyses involving fasting glucose and fasting insulin we removed individuals with type 2 diabetes (defined as a clinical diagnosis, fasting glucose values ≥ 7.0 mmol/L or individuals who were taking glucose lowering drugs). We also excluded individuals on lipid lowering medication from the multivariable and MR.

Analyses were performed using Stata/SE version 14.0 (StataCorp, Brownsville, TX) and R version 3.4.4 (R Foundation for Statistical Computing, Vienna, Austria).

**Multivariable analysis**

We estimated the association between each liver function marker and T2D in each UCLEB study using logistic regression. For continuous outcomes we used linear regression models. We adjusted the logistic/linear regression models for age, sex (if relevant), and for as
many potential confounders available in each study from the following: BMI, waist circumference, alcohol consumption, smoking, and social class. We then used DerSimonian and Laird random-effects model meta-analysis to combine these estimates with results from the published meta-analyses of Kunutsor et al (2) and Fraser et al (27) for the associations between ALT, AST and GGT with T2D, excluding any UCLEB studies that had contributed to the published meta-analyses. In the previously published meta-analyses, all (27) or some (2) of the participating studies expressed T2D results as hazard ratios. Given the overall proportion of T2D cases was lower than 10%, hazard ratios and odds ratios were assumed to approximate to the same measure of T2D relative risk in our meta-analysis.

**Mendelian randomization analysis**

For the MR analysis, we used multiple genetic variants robustly associated with circulating ALT, AST, ALP, or GGT as genetic instruments for each liver function marker to investigate their effect on T2D and other outcomes. We also applied MR analysis to assess the effect of predisposition to T2D and fasting insulin (a marker of insulin resistance) on circulating ALT, AST, ALP, or GGT by using genetic variants robustly associated with T2D risk and fasting insulin. We used a “two-sample” analysis strategy in which the genetic variant-exposure and genetic variant-outcome associations are estimated from different data sources with comparable populations (38).

**Selection of genetic instruments for liver function markers**

Genetic instruments for each liver function marker were defined as independent SNPs (R² < 0.3) associated with each liver function marker at genome-wide levels of significance (P < 5 x10⁻⁸) from GWAS in the NIHR GWAS catalogue (available at www.ebi.ac.uk/gwas/home, last accessed on July 2017). At the time the study was conducted, there was limited GWAS data available for AST. Therefore, we conducted a genome-wide
association analysis in up to 6,647 individuals from five UCLEB studies (BRHS, BWHHS, CaPS, EAS, and ET2DS) to identify novel variants for AST, as this biomarker had only been assessed in one previous GWAS with fewer than 1,000 individuals (37). We selected GWAS studies, which had been primarily conducted in populations of European ancestry. In total, we selected 4, 3 (including the two novel SNPs identified from our own GWAS), 15, and 26 independent SNPs associated with ALT, AST, ALP, and GGT, respectively (Supplementary table 3). UCLEB data were excluded from the MR analysis of AST (as an exposure) to avoid any bias from Winner’s curse (39).

Selection of genetic instrument for type 2 diabetes and fasting insulin

We used multiple independent SNPs strongly associated (P < 5 x10^-8) with T2D risk (n = 139 SNPs) (29) and with fasting insulin (n = 14 SNPs) (32). T2D SNPs were identified as reported by the original GWAS publication (29). Fasting insulin SNPs were selected from data published by Scott et al (32) using the R package ‘TwoSampleMR’ excluding any correlated SNPs (R^2 > 0.001) (40).

Main analysis

In the main Mendelian randomization analyses, we used the conventional inverse variance weighted (IVW) estimator. The IVW method consists of a weighted regression of the SNP-outcome regression coefficients on the SNP-exposure regression coefficients constraining the intercept to be zero. IVW weights are the inverse of the variance of the SNP-outcome regression coefficients. For a dichotomous outcome such as a T2D status, the regression coefficient of the SNP-outcome association is a log odds ratio from a logistic regression model. The resulting regression coefficient from the IVW regression represents an increase/decrease in the outcome per unit increase in the exposure.
Sensitivity analyses

We performed several sensitivity analyses to test whether the MR IVW estimates are likely to be biased by unbalanced horizontal pleiotropic effects (i.e. due to genetic variants that affect the outcome independently of the exposure of interest). We used MR-Egger regression method (41) and the weighted median estimator (42), both of which are more robust to pleiotropic genetic variants, to test the extent to which any unbalanced pleiotropy may have biased the IVW result. The MR-Egger method is similar to the IVW except that the model allows the intercept to vary. The intercept of the MR-Egger regression will reflect the average pleiotropic effect across genetic variants and the slope coefficient will provide an estimate of the causal effect provided that the InSIDE (Instrument Strength Independent of Direct Effect) assumption holds, which requires that there is no correlation between SNP-exposure association and any direct (pleiotropic) effects of SNP on outcome (41). In contrast, the weighted median estimator gives consistent estimates even if up to 50% of weight in the analysis comes from invalid genetic instruments (42). For the MR analysis of the effect of insulin resistance, proxied by fasting insulin, on liver markers, we have performed an additional sensitivity analysis, in which we used multivariable MR (43) to adjust results by BMI (32). In all analyses with GWAS data, we have excluded SNPs with C/G or A/T genotypes and minor allele frequency > 0.42 due to ambiguity problems they could introduce when harmonising SNP-exposure and SNP-outcome datasets (44).
RESULTS

Genetic instruments for liver function markers, T2D risk and fasting insulin

Results for the association of genetic instruments with the respective liver function marker are given in Table 1. Most of the genetic instruments previously identified in GWAS consortia replicated (had consistent direction and magnitude) in UCLEB and Fenland, seven were not available in UCLEB but were replicated in Fenland, seven were null or in the opposite direction in UCLEB but consistent in Fenland, and one was null in Fenland (but consistent in UCLEB). In all cases, confidence intervals in UCLEB and Fenland included the point estimate from the published GWAS.

For the known variant for AST (rs17109512), only P-values were provided by the original GWAS and, therefore, it was not possible to compare effect estimates with that from the previous GWAS. The two novel variants for AST, identified in our GWAS conducted in five UCLEB studies, were replicated in independent data sources (Table 1 and Supplementary table 4). When meta-analysing UCLEB and Fenland studies, 4, 13 and 22 variants were replicated from 4, 15 and 26 variants associated with ALT, ALP and GGT, and the remaining variants were directionally consistent with the previous reports (data only shown for UCLEB and Fenland separately).

Results for the association of the 139 T2D genetic instruments with T2D risk are given in Supplementary table 5. Results for the association of the 14 fasting insulin genetic instruments with fasting insulin (in SD log pmol/L) are given in Supplementary table 6.

Association of genetic variants related to liver function markers with potential confounders of the exposure-outcome association
Some individual SNPs used as instruments in the MR analysis were associated with potential confounders of the exposure-outcome association (e.g. age, BMI, waist circumference, waist-hip ratio, and other liver function markers). However, when combined into a single instrument using fixed-effect meta-analysis, there was no strong evidence that genetic instruments were associated with potential confounders, except for the ALT instrument that was also associated with AST and GGT, and the AST instrument that was associated with ALT (Supplementary tables 7a and 7b).

**Multivariable analysis between liver function markers and T2D and related continuous outcomes**

Pooled results from multivariable analyses across the relevant UCLEB and Fenland studies and published meta-analyses are given in Figure 2A. Most liver function markers were positively associated with T2D risk. The OR for T2D was 1.27 [95% CI: 1.15, 1.40] for ALT, 1.06 [95% CI: 0.97, 1.17] for AST, 1.25 [95% CI: 1.10, 1.43] for ALP and 1.61 [95% CI: 1.15, 2.26] for GGT (per standard unit increase in the liver function marker). Heterogeneity across studies was low for ALT and ALP (I² = 0%) but high for AST (I² = 81%) and GGT (I² = 93%) (Figure 2A).

In meta-analyses of the continuous outcomes, ALT was positively associated with insulin and triglycerides. AST was positively associated with insulin and HDLc. ALP was negatively related with HDLc. GGT was positively associated with all continuous outcomes (Supplementary table 8).

**Effect of liver function markers on T2D and related continuous outcomes using MR**

Pooled results from the MR across UCLEB, Fenland and GWAS studies are given in Figure 2A and Supplementary table 8. In the main MR analysis (IVW), the OR for T2D was
1.45 [95% CI: 1.10, 1.92] for ALT, 1.25 [95% CI: 1.14, 1.38] for AST, 0.91 [95% CI: 0.86, 0.97] for ALP and 0.92 [95% CI: 0.80, 1.06] for GGT (per each standard unit increase in the liver function marker) (Figure 2A). The other MR methods (MR-Egger and weighted median) used as sensitivity analyses were consistent with IVW estimates for ALT, AST and ALP. The inverse point estimate for the IVW association between GGT and T2D changed in direction when using the MR-Egger method. There was no clear evidence of unbalanced horizontal pleiotropy in any liver function marker – T2D associations based on the intercept for MR-Egger method (all P-values ≥0.38). Heterogeneity between UCLEB and GWAS estimates was low for all liver markers. For AST, only GWAS data were available for analysis (Figure 2A).

In meta-analyses of the continuous outcomes, there was some evidence across different MR methods that genetically predicted AST and ALP were negatively related to HDLc, LDLc and total cholesterol (Supplementary table 8). There was some evidence of unbalanced horizontal pleiotropy of ALT instruments in relation to HDLc, LDLc, total cholesterol and triglycerides (Supplementary table 9).

**Effect of T2D and insulin resistance on liver function markers using MR**

Overall, findings from MR analysis did not consistently support a reverse causal effect of T2D on any of the liver function markers assessed (ALT, AST, ALP, or GGT). In the main MR analysis (IVW), higher predisposition to T2D (each increase in 1 log odds) was related to an increase of 0.06 SD units of ALT [95% CI: 0.02, 0.09], but not of AST [0.01; 95% CI: -0.04, 0.07], ALP [-0.04; 95% CI: -0.11, 0.02] or GGT [0.03; 95% CI: -0.01, 0.06]. Results for ALT were substantially attenuated when using MR methods that are more robust to pleiotropic variants: 0.01 [95% CI: -0.06, 0.08] for MR-Egger and 0.03 [95% CI: -0.02, 0.07] for the weighted median estimator (Figure 2B).
On the other hand, there was evidence from the main MR findings that higher insulin resistance, proxied by fasting insulin, increases circulating ALT. Results were consistent with the weighted median and BMI-adjusted estimates. MR-Egger point estimates substantially differed, but 95% CI were very wide. Results for the other liver markers were less consistent across different MR methods and 95% CI were wide (Table 2).
DISCUSSION

More than a century ago a link between liver disease and diabetes was described (45). Since then, multiple observational studies have repeatedly reported that liver dysfunction and T2D are associated (1-3), as broadly replicated by our multivariable analysis using clinical biomarkers (i.e. circulating ALT, AST, ALP, and GGT) as proxies of liver dysfunction. Nevertheless, it is still unclear whether this association reflects causation, and if so, whether liver dysfunction represents a cause or a consequence of T2D.

Our study expands on previous MR analyses investigating the effect of liver dysfunction on T2D risk by including a more comprehensive set of liver function markers routinely used in clinical practise in the largest available datasets and by applying bi-directional MR to investigate whether predisposition to T2D and to insulin resistance might instead lead to liver dysfunction.

Our findings from the MR analyses show evidence that genetic predisposition to higher circulating ALT and AST is related to higher risk of T2D. No strong evidence of a causal effect of genetically predicted GGT on T2D, and evidence of a modest negative effect of genetically predicted ALP on T2D were found. Genetic predisposition to T2D did not appear to influence blood concentration of any of the studied liver function markers (ALT, AST, ALP, and GGT), whereas genetic predisposition to insulin resistance, proxied by fasting insulin, seems to increase ALT (effects on other liver markers are uncertain).

Our results are broadly consistent with two previous MR studies, of largely European origin participants (using a study sample that partially overlaps with ours), which reported strong evidence for a positive effect of ALT on T2D (14), but not for ALP (14) or GGT (13, 14). Results for ALP (14) were directionally consistent with our findings, but we were better powered to test for the association between ALP and T2D risk given the substantially larger
number of T2D cases and controls included in our analyses. In non-European populations, MR studies suggest that ALT does not relate to T2D risk in Chinese adults (15), but that higher GGT increases T2D risk in Koreans (16). However, the latter result might be explained by statistical overfitting since instruments were selected from a GWAS that included the population used in the MR analyses (~ 20% of the GWAS discovery sample).

The different liver biomarkers reflect different aspects of liver dysfunction. High circulating ALT and AST are widely used proxies of NAFLD, while high circulating ALP and GGT (in combination) are more related to obstructive or cholestatic liver disease. The positive association between genetically predicted liver function markers and T2D in MR analyses was robust for ALT and AST, but not apparent for ALP or GGT, which suggests that NAFLD might be the primary type of liver dysfunction driving these associations.

In agreement with that, a genetic variant (rs738409) in perfect linkage disequilibrium (R2 = 1.0 for 1000 Genomes European population - GRCh37) with one of our instruments for AST (rs738408) has been previously reported to be associated with computed tomography (CT) measured hepatic steatosis (46, 47). NAFLD is the most common cause of chronic liver disease in Western countries owing to the rapid increase in obesity prevalence (48, 49). NAFLD affects 70% of patients with type 2 diabetes (T2D) in contrast to around 20% of the general population (48, 49).

To our knowledge previous MR studies have not examined the causal effect of predisposition to T2D or insulin resistance on liver dysfunction. We found supportive evidence that insulin resistance increases circulating ALT. Our combined findings that insulin resistance (but not T2D) may cause elevated ALT (marker of NAFLD), and that ALT and AST are in turn related to increased T2D risk (but not insulin resistance) are consistent with the twin-cycle hypothesis (8), which postulates that there is a vicious cycle between hepatic insulin resistance and β cell dysfunction. According to the twin-cycle hypothesis, elevated insulin (due to insulin resistance) stimulates de novo lipogenesis in the liver, which promotes hepatic insulin
resistance leading to overstimulation of hepatic gluconeogenesis and increased fasting glucose. The resulting increased output of triglycerides and glucose by the liver to the circulation would impair beta cell function, eventually leading to type 2 diabetes (9).

On the other hand, it is worth emphasising that the relation between ALT/AST and T2D risk might be explained by factors other than NAFLD since circulating ALT/AST are not specific markers of NAFLD and can also increase in response to liver injury from other causes, such as drug toxicity, infection and alcohol consumption (50, 51). In addition, there is some evidence that hepatic triglyceride accumulation by itself may not necessarily cause metabolic changes increasing the risk of cardiometabolic complications (52) and that there might be multiple T2D subtypes that differ in terms of disease presentation and responsiveness to interventions (53), which we were unable to tease out due to the predominant use of summary level data. Finally, given the key roles of ALT and AST in the intermediary metabolism of glucose and amino acids, we cannot fully discard that they might be directly implicated in T2D development.

Although MR can substantially improve causal inference in Epidemiological studies (54), it is important to note that the reliability of MR findings depend on the three core assumptions of instrumental variable (IV) analysis, which require that genetic instruments are strongly associated with the exposure (IV1), are not related to exposure-outcome confounders (IV2) and only influence the outcome through the exposure (IV3).

To avoid violations of IV1, we have selected genetic variants strongly associated with liver function markers (P < 5*10^{-8}), that broadly replicated in independent datasets. It should also be noted that our genetic instruments were selected to be strongly associated with each liver function marker in the largest available GWAS (P < 5*10^{-8}) and that some variants relevant to specific forms of liver dysfunction might not have met our inclusion criteria, as is the case of variants nearby TM6SF2, previously reported to be associated with NAFLD in a GWAS (55) and with alcohol-related cirrhosis in an exome-wide association study (56). To
minimise the risk of population stratification, which could violate IV2, we mostly restricted our analyses to individuals of European ancestry. IV3 could be violated in the presence of horizontal pleiotropy. We have attempted to address that by examining the relation of the genetic variants with several social, behavioural and metabolic phenotypes and by using methods that are more robust to violations of this assumption in sensitivity analyses (i.e. MR-Egger and weighted median estimator). Overall, there was no strong suggestion that horizontal pleiotropy could have biased our results. Importantly, our liver function instruments were not associated with general adiposity measures such as BMI, and therefore it is unlikely that these associations are driven by general adiposity. However, it is important to note that we cannot fully discard that our genetic instruments may be associated with exposure-outcome confounders that we have not tested for and that the sensitivity analyses used (i.e. MR-Egger and weighted median) are of limited use for exposures instrumented by few genetic variants (as in the analyses of ALT and AST as exposures).

Two-sample MR makes the additional assumption that data from two independent (but comparable) populations are used. In our study, there was some overlap between participants used to estimate SNP-exposure and SNP-outcome associations. However, this is unlikely to bias the study results since the overlap is very low in proportion to the overall sample size (< 7% of T2D cases in the main analysis) (57).

Finally, effect estimates for the relation of each liver function marker on T2D risk should be interpreted with caution given these biomarkers are unlikely to be the causal factors for T2D risk, but rather proxies of liver function.

In conclusion, MR findings indicate that increased circulating ALT and AST is related with higher T2D risk, while increased circulating ALP is associated with lower T2D risk. In addition, higher fasting insulin (but not predisposition to T2D) is related to higher circulating ALT. Since circulating ALT and AST are markers of NAFLD, these findings provide some support for insulin resistance resulting in NAFLD, which in turn increases T2D risk.
Funding sources

This work was supported by funds from the UK Medical Research Council (Grant number: G0801456) and a British Heart Foundation grant (AA/18/7/34219). MCB, TRG and DAL work in a unit that receives funding from the UK Medical Research Council (Grant number: MC_UU_00011/1-6 and MC_UU_00011/1-4). MCB is supported by a Skills Development Fellowship from the UK Medical Research Council (Grant number MR/P014054/1). XZ is supported by a scholarship from the China Scholarship Council (Grant number: 201406220101). ADH is an NIHR Senior Investigator. AW and DK are funded by the UK Medical Research Council [MC UU 12019/1]. DAL is a UK National Institute of Health Research Senior Investigator (Grant number: NF-SI-0611-10196).

The UCLEB (UCL-LSHTM-Edinburgh-Bristol) consortium is supported by BHF Programme Grant RG/10/12/28456 and the UCL Hospitals NIHR Biomedical Research Centre.

BRHS is supported by British Heart Foundation grants (RG/08/013/25942, RG/13/16/30528). The British Heart Foundation had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; preparation, review, or approval of the manuscript; and decision to submit the manuscript for publication. The authors acknowledge the British Regional Heart Study team for data collection.

BWHHS is supported by the British Heart Foundation (PG/13/66/30442). Data on mortality and cancer events were routinely provided from NHS Digital to the BWHHS under data sharing agreement MR104a- Regional Heart Study (Female Cohort). British Women’s Heart and Health Study data are available to bona fide researchers for research purposes. Please refer to the BWHHS data sharing policy at www.ucl.ac.uk/british-womens-heart-health-study.
**CaPS** was funded by the Medical Research Council and undertaken by the former MRC Epidemiology Unit (South Wales). The CaPS DNA bank was established with funding from a MRC project grant. The CaPS data archive is maintained by the University of Bristol. MRC Integrative Epidemiology Unit, Bristol is supported by MRC grants (MR_UU_12013/1, MR_UU_12013/5 and MR_UU_12013/8).

The **EAS** is supported by the British Heart Foundation and by the Chief Scientist Office of Scotland.

**ELSA** is supported by the National Institute on Aging (NIA/NIH) USA (grant number 5 R01 AG017644-16) and a consortium of the United Kingdom government departments coordinated by the Economic and Social Research Council (ESRC).

The **WHII study** is supported by grants from the Medical Research Council (K013351), British Heart Foundation (RG/07/008/23674), Stroke Association, the US National Heart Lung and Blood Institute (5RO1 HL036310), the US National Institute on Aging (5RO1AG13196) the US Agency for Healthcare Research and Quality (HS06516); and the John D. and Catherine T. MacArthur Foundation Research Networks on Successful Midlife Development and Socio-economic Status and Health.

**NSHD** is funded by the UK Medical Research Council [MC_UU_12019/1].

The **Fenland Study** is funded by the Wellcome Trust and the Medical Research Council (Grant number: MC_U106179471 and MC_UU_12015/1).
The funders had no role in the design, analyses, interpretation of results or writing of the paper. The views expressed in this paper are those of the authors and not necessarily any of the funders.

Acknowledgments

The UCLEB (UCL-LSHTM-Edinburgh-Bristol) consortium consists of 12 studies: Northwick Park Heart Study II (NPHS II), British Regional Heart Study (BRHS), Whitehall II Study (WHII), English Longitudinal Study of Ageing (ELSA), Medical Research Council National Survey of Health and Development (NSHD), 1958 Birth cohort (1958BC), Caerphilly prospective study (CaPS), British Women's Heart and Health Study (BWHHS), Edinburgh Artery Study (EAS), Edinburgh Heart Disease Prevention Study (EHDPS), Edinburgh Type 2 Diabetes Study (ET2DS) and Asymptomatic Atherosclerosis Aspirin Trial (AAAT). We thank all participants and staff who have contributed to those studies that contributed to the present study (BRHS, BWHHS, CaPS, EAS, ELSA, ET2DS, NSHD, WHII, Fenland study, and NFBC1966). We thank Naveed Sattar for the insightful comments on the interpretation of findings.

Data and Resource Availability

Data on type 2 diabetes have been contributed by the T2D GWAS published by Xue et al (2018) and have been downloaded from http://cnsgenomics.com/data.html. Data on glycaemic traits have been contributed by MAGIC investigators and have been downloaded from www.magicinvestigators.org. Data on lipid traits have been contributed by Global Lipids Genetics Consortium and have been downloaded from http://csg.sph.umich.edu/abecasis/public/lipids2013/. Data on anthropometric traits have been
contributed by Genetic Investigation of ANthropometric Traits (GIANT) consortium and have been downloaded from http://www.broadinstitute.org/collaboration/giant/index.php/GIANT_consortium_data_files. UCLEB data is available upon request.

Author contributions

DAL designed the study and initial analysis plan, with further input from NMGDS, MCB, and TG. DAL, JE, AW, DK, TS, XZ, CL, JL, JCC, WZ, and TRG collected and/or managed data from one or more of the contributing studies, NMGDS and MCB undertook statistical analyses with supervision from DAL and TG. NMGDS, DAL and MCB wrote the first draft of the paper with all other co-authors contributing to revisions.

Conflict of Interest statement

DAL has received support from Medtronic and Roche Diagnostics for biomarker research unrelated to this study. TRG receives support from GlaxoSmithKline, Biogen and Sanofi for research unrelated to this study. All other authors report no competing interests.

Guarantor Statement

NMGDS and MCB are the guarantors of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

References


49. Sattar N, Forrest E, Preis D. Non-alcoholic fatty liver disease. BMJ. 2014;349:g4596.


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<th>SNP</th>
<th>Liver function marker</th>
<th>Locus*</th>
<th>Effect size in SD units in UCLEB studies† (95% CI)</th>
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* Mapped gene for each SNP is given as reported in the original GWAS publication.
† All SNP-liver function marker associations in the UCLEB studies and the Fenland study were adjusted for age and sex (if relevant). Individual study estimates in UCLEB were combined using fixed effect meta-analyses.
‡ Effect sizes extracted from Chambers et al., 2011 (28). GWAS effect sizes for ALT, ALP and GGT were converted to SD units using the median SD from the UCLEB and Fenland studies.
ALP: alkaline phosphatase; ALT: alanine aminotransferase; AST: aspartate aminotransferase; GGT: gamma-glutamyl transferase; GWAS: genome-wide association study; N/A: not applicable; SNP: single nucleotide polymorphism; SD: standard deviation; UCLEB consortium: UCL-LSHTM-Edinburgh-Bristol consortium
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Results correspond to mean difference (in S.D. units) of log10 liver function marker (95% CI) per one S.D increase in fasting insulin (in pmol/L). ALT: alanine aminotransferase; AST: aspartate aminotransferase; ALP: alkaline phosphatase; GGT: gamma-glutamyl transferase; IVW: inverse variance weighted method; S.D.: standard deviation.
Figure legends

Figure 1. Study design and data sources used to investigate the effect of liver dysfunction (proxied by biomarkers: ALT, AST, ALP, and GGT) on type 2 diabetes or secondary outcomes (fasting glucose, fasting insulin, LDL-C, HDL-C, total cholesterol, and triglycerides) (A) and the effect of predisposition to type 2 diabetes or insulin resistance on circulating liver function biomarkers (B)

As shown in figure 1A, the multivariable association of liver function markers with T2D risk (or related outcomes) was estimated by meta-analysing results from each data source using logistic regression models (or linear regression models in the case of secondary outcomes) with participant-level data from relevant studies within UCLEB consortium (BRHS, BWHHS, MRC-NHS) and summary-level data from the published meta-analyses of Kunutsor et al (2013) and Fraser et al (2009). We also estimated the association of liver function markers with T2D risk (or secondary outcomes) using a Mendelian randomization approach. In Mendelian randomization analysis, we used different data sources to estimate the SNP-liver function marker association (UCLEB consortium — BRHS, BWHHS and MRC-NHS —, Fenland study, and GWAS of liver function markers — Chambers et al (2011)) —, and SNP-T2D risk association (UCLEB — BRHS, BWHS, CaPS, EAS, ELSA, MRC-NHS, and WHII —, and GWAS consortium) or SNP-secondary outcomes. As shown in figure 1B, the summary-level data for the association of SNP-T2D risk and SNP-fasting insulin for the reverse MR was extracted from GWAS consortia, and the association of SNP-liver function marker was extracted from Chambers et al (2011). ALP: alkaline phosphatase; ALT: alanine aminotransferase; AST: aspartate aminotransferase; BMI: body mass index; BRHS: British Regional Heart Study; BWHHS: British Women’s Heart and Health Study; CaPS: Caerphilly Prospective Study; DIAGRAM consortium: Diabetes Genetics Replication And Meta-analysis consortium; EAS: Edinburgh Artery Study; ELSA: English Longitudinal Study of Ageing; GGT: gamma-glutamyl transferase; GWAS: genome-wide association study; HDL-C: high density lipoprotein-cholesterol; LDL-C: low density lipoprotein-cholesterol; MRC-NHS: National Survey of Health and Development; SNPs: single nucleotide polymorphisms; T2D: type 2 diabetes; UCLEB consortium: UCL-LSHTM-Edinburgh-Bristol consortium; WHII: Whitehall II study.

Figure 2. Multivariable and Mendelian randomization analysis of the effect of liver function on T2D (A) and Mendelian randomization analysis of the effect of T2D on liver function markers (B).

Results from Figure 1A correspond to odds ratio of T2D per unit increase in standardized liver function markers (and 95% confidence interval). Results from Figure 1B correspond to change in standardized liver function markers per unit increase in log odds of T2D (and 95% confidence interval). I-squared indicates between-study heterogeneity and is only presented when estimates for more than one study were available. ALT: alanine aminotransferase; AST: aspartate aminotransferase; ALP: alkaline phosphatase; GGT: gamma-glutamyl transferase; SNPs: single nucleotide polymorphisms; T2D: type 2 diabetes.