



Clinical and Translational Research

Investigating the causal associations between five anthropometric indicators and nonalcoholic fatty liver disease: Mendelian randomization study

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Specialty type: Medicine, research and experimental

Provenance and peer review: Unsolicited article; Externally peer reviewed.

Peer-review model: Single blind

Peer-review report's scientific quality classification

Grade A (Excellent): 0
Grade B (Very good): B
Grade C (Good): 0
Grade D (Fair): D
Grade E (Poor): 0

P-Reviewer: Ulasoglu C, Turkey

Received: October 25, 2023

Peer-review started: October 25, 2023

First decision: December 31, 2023

Revised: January 14, 2024

Accepted: February 6, 2024

Article in press: February 6, 2024

Published online: March 6, 2024



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Abstract

BACKGROUND

Although the etiology of nonalcoholic fatty liver disease (NAFLD) has not been thoroughly understood, the emerging roles of anthropometric indicators in assessing and predicting the risk of NAFLD have been highlighted by accumulating evidence.

AIM

To evaluate the causal relationships between five anthropometric indicators and NAFLD employing Mendelian randomization (MR) design.

METHODS

The Anthropometric Consortium provided genetic exposure data for five anthropometric indicators, including hip circumference (HC), waist circumference (WC), waist-to-hip ratio (WHR), body mass index (BMI), and body fat percentage (BF). Genetic outcome data for NAFLD were obtained from the United Kingdom Biobank and FinnGen Consortium. Genome-wide significant single nucleotide polymorphisms were chosen as instrumental variables. Univariable MR (UVMR) and multivariable MR (MVMR) designs with analytical approaches, including inverse variance weighted (IVW), MR-Egger, weighted median (WM), and weighted mode methods, were used to assess the causal relationships between anthropometric indicators and NAFLD.

RESULTS

Causal relationships were revealed by UVMR, indicating that a higher risk of NAFLD was associated with a per-unit increase in WC [IVW: odds ratio (OR) = 2.67, 95%CI: 1.42-5.02, $P = 2.25 \times 10^{-3}$], and BF was causally associated with an increased risk of NAFLD (WM: OR = 2.23, 95%CI: 1.07-4.66, $P = 0.033$). The presence of causal effects of WC on the decreased risk of NAFLD was supported by MVMR after adjusting for BMI and smoking. However, no causal association between BF and NAFLD was observed. In addition, other causal relationships of HC, WHR (BMI adjusted), and BMI with the risk of NAFLD were not retained after FDR correction.

CONCLUSION

This study establishes a causal relationship, indicating that an increase in WC is associated with a higher risk of NAFLD. This demonstrates that a suitable decrease in WC is advantageous for preventing NAFLD.

Key Words: Anthropometric indicator; Waist circumference; Nonalcoholic fatty liver disease; Mendelian randomization; Genetic variant

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Core Tip: Previous studies have demonstrated the potential significance of anthropometric indicators in the development of nonalcoholic fatty liver disease (NAFLD). Nevertheless, inconsistencies exist in the results of these studies, and the causal association remains unclear. Abdominal obesity, measured by waist circumference (WC), is a risk factor for NAFLD, as demonstrated by previous studies. Nevertheless, many of these studies were cross-sectional or considered only a single measurement, neglecting a comprehensive evaluation of changes in WC over time and the effect of long-term development and lifestyle changes. Consequently, establishing a causal relationship between anthropometric indicators and NAFLD requires further robust evidence.

Citation: Xiao XP, Dai YJ, Zhang Y, Yang M, Xie J, Chen G, Yang ZJ. Investigating the causal associations between five anthropometric indicators and nonalcoholic fatty liver disease: Mendelian randomization study. *World J Clin Cases* 2024; 12(7): 1215-1226

URL: <https://www.wjgnet.com/2307-8960/full/v12/i7/1215.htm>

DOI: <https://dx.doi.org/10.12998/wjcc.v12.i7.1215>

INTRODUCTION

Excessive fat deposition in liver cells characterizes nonalcoholic fatty liver disease (NAFLD). Currently, due to lifestyle changes and insufficient daily physical activity, NAFLD has emerged as the most severe chronic disease in society[1]. Reports indicate that NAFLD affects a quarter of the global population, with a prevalence rate of 24% and a tendency toward younger age groups[2,3]. Although the etiology of NAFLD has not been thoroughly understood, the emerging roles of anthropometric indicators in assessing and predicting the risk of NAFLD have been highlighted by accumulating evidence[4]. Numerous previous observational studies have reported relationships between the risk of NAFLD and noninvasive quantitative measurements of the body, such as anthropometric indicators, which comprise height, weight, hip circumference (HC), waist circumference (WC), waist-to-hip ratio (WHR), body mass index (BMI), and body fat percentage (BF)[4,5]. Contradictory findings have been obtained in some other studies, indicating no relationship between anthropometric indicators and the risk of NAFLD[6,7]. The causal relationships between anthropometric indicators and NAFLD risk remain undetermined, considering these inconsistent findings and the absence of randomized controlled studies.

Mendelian randomization (MR) is a novel epidemiological tool that employs genetic data to investigate the causal relationship between exposure and outcome[8]. Generally, genetic variants are independent of disease state and are randomly assigned to offspring through the maternal generation. Consequently, the limitations of conventional observational design can be overcome, and biases such as potential confounders and reverse causality can be minimized[9,10]. Previous studies have demonstrated the use of MR to investigate causal relationships between NAFLD and numerous diseases, including cardiovascular disease and psoriasis[11,12].

In this study, the causality of five anthropometric indicators, including HC, WC, WHR (BMI adjusted), BMI, and BF, with the risk of NAFLD, was investigated using genetic summary statistics with Univariable MR (UVMR) and multivariable MR (MVMR) frameworks to establish a foundation for the prevention and management of this high-burden disease.

MATERIALS AND METHODS

Study design

In this MR analysis, the causal effects of various modifiable exposures on outcomes were estimated using single nucleotide polymorphisms (SNPs) as instrumental variables (IVs). However, three basic assumptions must be satisfied [13]. First, IVs must exhibit a high correlation with the exposure factor (correlation assumption) to minimize weak IV bias [14]. Second, the outcome should only be influenced by the identified IVs through exposure and not through other factors, expressed as "no horizontal pleiotropy" (exclusionary assumption) [15]. Third, to identify genuine causal associations, IVs should be independent of confounders in exposure-outcome associations (independence assumption) [16].

A two-stage design was used in this study, with the first phase employing bidirectional UVMR analyses to analyze the causal relationship between the five anthropometric measures and NAFLD. In the second phase, MVMR analyses were conducted to establish causality after adjusting for potential confounders, such as BMI and smoking. These findings were validated using databases from various sources. Figure 1 shows the study design.

Exposure data sources and IVs selection

Five anthropometric measures, including HC, WC, WHR (BMI adjusted), BMI, and BF, were chosen as exposure points. Shungin *et al* [17] reported a genome-wide association study (GWAS) that included up to 224459 individuals, from which summary statistics for HC, WC, and WHR (BMI adjusted) were extracted [17]. The summary data for BMI were obtained from another GWAS, which involved 700000 participants [18]. Genetic summary data for BF were obtained from GWAS data published by Neale Lab in 2017, which included 331117 participants (<http://www.nealelab.is/uk-biobank>). All exposure datasets were conducted in European ancestry.

The threshold was set at $P < 5 \times 10^{-08}$ to choose SNPs related to anthropometric metrics as IVs. Furthermore, SNPs with strong linkage disequilibrium ($R^2 < 0.001$, window size = 10000 kb) were deleted to avoid possible bias. Significantly heterogeneous SNPs were excluded, and the remaining anthropometric-associated SNPs were selected as valid IVs using the heterogeneity test. In some of the exposure-outcome analyses, some SNPs were excluded due to the lack of available proxies, as they were not present in the outcome (Supplementary Table 1). Finally, a total of 807 anthropometric-associated SNPs were available, including 18 HC-associated SNPs, 46 WC-associated SNPs, 32 WHR (BMI adjusted)-associated SNPs, 485 BMI-associated SNPs, and 226 BF-associated SNPs (Supplementary Table 1).

Outcome data sources

Concerning the outcome datasets, genetic associations for NAFLD agent-related SNPs were extracted from the United Kingdom biobank, which included all European subjects, comprising 8434 cases and 770180 controls [19]. The GWAS findings were harmonized with those of GWAS exposed to the same effect allele. Effects allele frequencies (minor allele frequency ≤ 0.5) were employed to harmonize the palindromic SNPs. For IVs that could not be found in the results, mutins were made to include suitable proxy SNPs ($r^2 \geq 0.8$) by searching for them on the website (<http://www.mulinlab.org/vportal/index.html>).

Ethical approval

This study was conducted using public open data from the MRC-IEU database and the FinnGen research project; thus, no ethical review was necessary.

Statistical analyses

UVMR analysis: Inverse variance weighted (IVW), weighted median (WM), weighted model, and MR Egger methods were used to evaluate the causal effects of HC, WC, WHR (BMI adjusted), and BF (per SD) on the risk of NAFLD. The IVW model, which was defined as the primary approach, can provide causal effect estimates with optimal precision when all IVs are valid and minimize the impact of heterogeneity. To assess weak IV bias, the variance (R^2) and approximated F -statistic for per-exposure explained by IVs were calculated. This evaluation, which is based on the inherent flaws in IV selection, is generally considered to be sufficiently instrumental for F -statistics greater than 10 [20,21]. The formula used is $F = R^2 \times (N - 2) / (1 - R^2)$, where R^2 represents the variance of exposure explained by each IV [22]. In addition, FDR correction was implemented to control false positives that tend to occur due to multiple testing.

MVMR analyses: Smoking and BMI were the primary confounders associated with exposure (anthropometric indicators) and outcome (NAFLD). Therefore, to determine the independent effects of anthropometric indicators on the risk of NAFLD development, MVMR analyses were employed. In MVMR analyses, MR-PRESSO models were employed to correct for horizontal pleiotropy in causal effects, estimate heterogeneity, and exclude potential outliers. A significance level of $P < 0.05$ in the MVMR analysis was considered indicative of statistical significance.

Sensitivity analysis: To ensure the stability of the study's results, several sensitivity analyses were conducted. First, the nonexistence of horizontal pleiotropy, a fundamental premise for satisfying causal inference, was analyzed using the MR-PRESSO approach. Findings with a P value greater than 0.05 indicated the nonexistence of horizontal pleiotropy [23]. Second, Cochran's Q statistic was employed to eliminate heterogeneity of IVs, with P values over 0.05 indicating the absence of heterogeneity in the IVs [24]. Furthermore, leave-one-out (LOO) analyses were performed to exclude single SNPs exerting a substantial influence on the findings [25]. An online tool "mRnd" was used to calculate the statistical efficacy (power). R programming software (version 4.0.3) with the "MR," "TwoSampleMR," "MVMR," and "MRPRESSO" packages was used to perform all statistical analyses and data visualization.

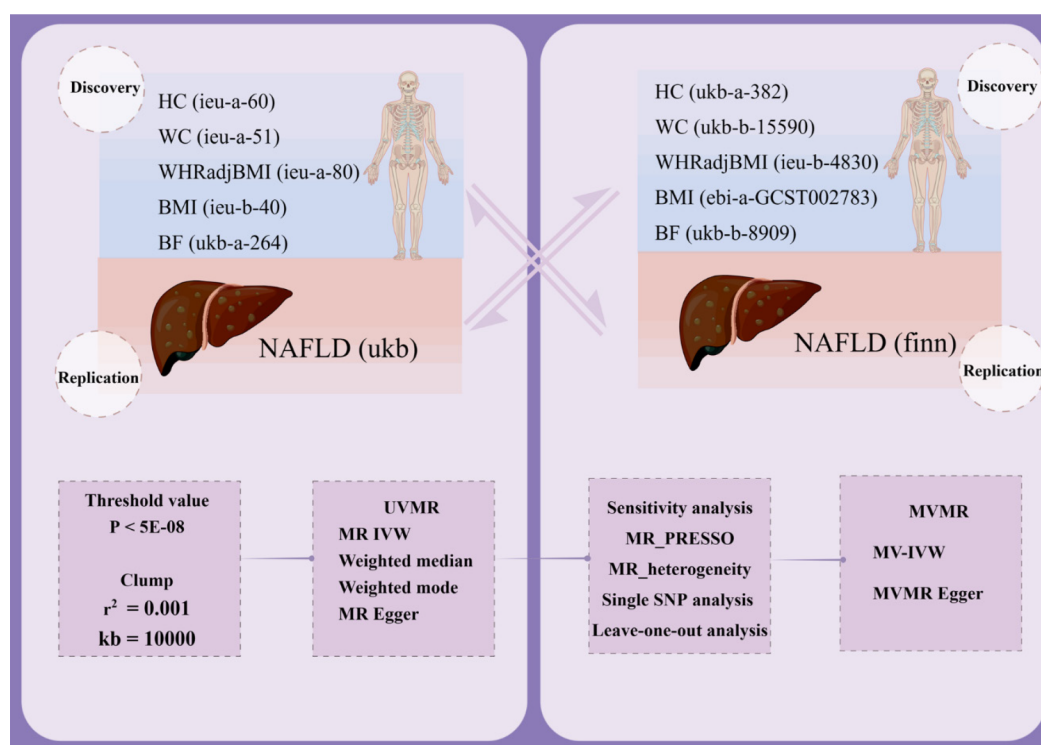


Figure 1 The study design of causal associations between anthropometric indicators and risk of non-alcoholic fatty liver disease. NAFLD: Non-alcoholic fatty liver disease; MR: Mendelian randomization; IVW: The inverse variance weighting method; MR-PRESSO: The Mendelian randomization pleiotropy residual sum and outlier; MVMR: Multivariable Mendelian randomization; SNPs: Single nucleotide polymorphisms; HC: Hip circumference; WC: Waist circumference; WHR: Waist-to-hip ratio; BMI: Body mass index; BF: Body fat percentage; adj BMI: Adjusting body mass index. Drawing by Figdraw (<https://www.figdraw.com/>).

Replication in another European-based population

Anthropometric indicators replication: Published anthropometric indicators GWAS data were extracted from other consortiums ($P < 5 \times 10^{-08}$, $R^2 < 0.001$, window size = 10000 kb). The GWAS data for HC were obtained from Neale Lab's 2017 summary database, which included 336639 individuals in this study (<http://www.nealelab.is/uk-biobank>). Summary data for WC and BF were derived from the MRC-IEU database and exported from the GWAS pipeline using the United Kingdom Biobank's pheasant-derived variables. Furthermore, the Within Family GWAS consortium was the source of data for WHR (BMI adjusted). In this genetic epidemiological study, sample analyses of related individuals (such as siblings or parent-child triples) were performed (<https://www.withinfamilyconsortium.com>). The BMI data were obtained from the GWAS study conducted by Chittani *et al*[26], which included 236781 individuals[26]. All data were extracted from published studies, and consequently, no additional ethical review was necessary. Figure 1 shows the details of the exposure and outcome data.

NAFLD replication: Outcome data were obtained from FinnGen, a large cohort study comprising genomic and health data from 500000 Finland BioBank individuals (<https://www.finnngen.fi>). Similar allele frequency distributions to other European populations were observed in Finland; however, significant strengths, including uncommon variants in intricate phenotypes and unique group genetic history, were noted[27].

RESULTS

Baseline characteristics

A total of 807 anthropometric indicator-associated SNPs, including 18 HC-associated SNPs, 46 WC-associated SNPs, 32 WHR (BMI adjusted)-associated SNPs, 485 BMI-associated SNPs, and 226 BF-associated SNPs, were identified from three independent GWAS analyses (Supplementary Table 1). Among the 807 anthropometric indicator-associated SNPs, all F-statistics were greater than 10, indicating a low likelihood of weak IVs among the included SNPs. In addition, the selected IVs explained approximately 0.84% (HC), 1.13% (WC), 1.56% [WHR (BMI adjusted)], 4.85% (BMI), and 3.60% (BF) of the phenotype variances. Low degrees of sample overlap between exposure and outcome were indicated by the sample overlap results, with overlap rates within datasets being less than 15%.

Stage 1 causal associations between anthropometric indicators and NAFLD risk

All findings were based on the IVW random-effects model because evidence of horizontal pleiotropy was not found ($P > 0.05$), but heterogeneity was present ($P < 0.05$). The findings of the IVW approach revealed causal relationships between

five anthropometric indicators and NAFLD: HC ($OR_{IVW} = 1.70$, 95%CI: 1.27-2.27, $P = 3.29 \times 10^{-04}$; $OR_{WM} = 1.76$, 95%CI: 1.25-2.48, $P = 1.20 \times 10^{-03}$), WC ($OR_{IVW} = 1.80$, 95%CI: 1.42-2.27, $P = 1.24 \times 10^{-06}$; $OR_{WM} = 1.88$, 95%CI: 1.40-2.51, $P = 2.42 \times 10^{-05}$), WHR (BMI adjusted) ($OR_{IVW} = 1.42$, 95%CI: 1.20-1.69, $P = 6.47 \times 10^{-05}$; $OR_{WM} = 1.35$, 95%CI: 1.06-1.71, $P = 1.40 \times 10^{-03}$), BMI ($OR_{IVW} = 1.60$, 95%CI: 1.43-1.79, $P = 1.49 \times 10^{-16}$; $OR_{WM} = 1.64$, 95%CI: 1.38-1.94, $P = 1.16 \times 10^{-08}$), and BF ($OR_{IVW} = 1.67$, 95%CI: 1.37-2.04, $P = 2.50 \times 10^{-07}$; $OR_{WM} = 1.91$, 95%CI: 1.50-2.43, $P = 1.36 \times 10^{-07}$; **Figure 2A** and **Supplementary Table 2**). Furthermore, sensitivity analyses using the LOO approach indicated that no single SNPs drove these findings after the stepwise elimination of individual SNPs (all error lines were on either side of zero; **Supplementary Figures 1-5**). Considering the positive false due to multiple testing, significant causal relationships between HC, WC, WHR (BMI adjusted), BMI, BF, and NAFLD were still observed in the results after FDR correction ($P < 0.01$; **Supplementary Figure 6**).

Causal associations between NAFLD and anthropometric indicators

No significant genetic predictive correlation was observed when NAFLD was tested as an exposure factor for inverse correlation, as indicated by estimates derived from the IVW method ($P > 0.05$). This further supports the notion that reversed causal relationships with NAFLD do not confound anthropometric indicators (**Supplementary Table 2**). In addition, the result of no significant genetic prediction of associations between NAFLD and anthropometric indicators was supported by sensitivity analysis.

Replication of results within different GWAS of exposures and outcomes

With the above IVs selection standard, 1052 SNPs associated with anthropometric indicators, including 390 HC-associated SNPs, 209 WC-associated SNPs, 15 WHR-associated SNPs, 79 BMI-associated SNPs, and 359 BF-associated SNPs, were selected for causal estimation in the replicated exposures (**Supplementary Table 1**). For HC, WC, WHR (BMI adjusted), BMI, and BF, the F -statistics were all over 10, indicating that the chosen IVs were sufficiently robust and not susceptible to the influence of weak IVs. Furthermore, the statistical power of these replication exposures was similarly calculated, and the selected IVs were shown to explain approximately 5.50% (HC), 3.38% (WC), 0.99% [WHR (BMI adjusted)], 2.17% (BMI), and 4.60% (BF) of the phenotype variances.

Subsequently, the two sources of exposure (discovery and replication) were analyzed in a bidirectional MR analysis with outcomes from various GWAS (United Kingdom biobank and FinnGen). The causal relationship between an increased risk of NAFLD and higher WC, WHR (BMI adjusted), BMI, and BF was supported by our cross-validation comparison of four sets of MR findings; sensitivity analysis findings remained consistent. Although the existence of heterogeneity between SNPs was indicated by the evidence, the overall level of pleiotropy based on the MR-Egger intercept was not significant. In our inverse MR analysis of anthropometric indicators (discovery database) with NAFLD (FinnGen research project), the presence of NAFLD resulted in a decrease in BMI, indicating that the previously observed causal relationship between BMI and NAFLD was influenced by reverse causality bias. In addition, FDR correction was equally performed on the validated results, resulting in the loss of some weak causal relationships. The findings of the four MR analyses supported the fact that an increase in WC ($OR_{IVW} = 1.80$, 95%CI: 1.42-2.27, $P = 1.24 \times 10^{-06}$; $OR_{WM} = 1.88$, 95%CI: 1.40-2.51, $P = 2.42 \times 10^{-05}$) and BF ($OR_{IVW} = 1.67$, 95%CI: 1.43-1.79, $P = 2.50 \times 10^{-07}$; $OR_{WM} = 1.91$, 95%CI: 1.50-2.43, $P = 1.36 \times 10^{-07}$) would result in a higher risk of NAFLD after the exclusion of reverse causality (**Figure 2B-D** and **Supplementary Tables 3-5**). **Supplementary Figure 6** shows the IVW results of the FDR correction.

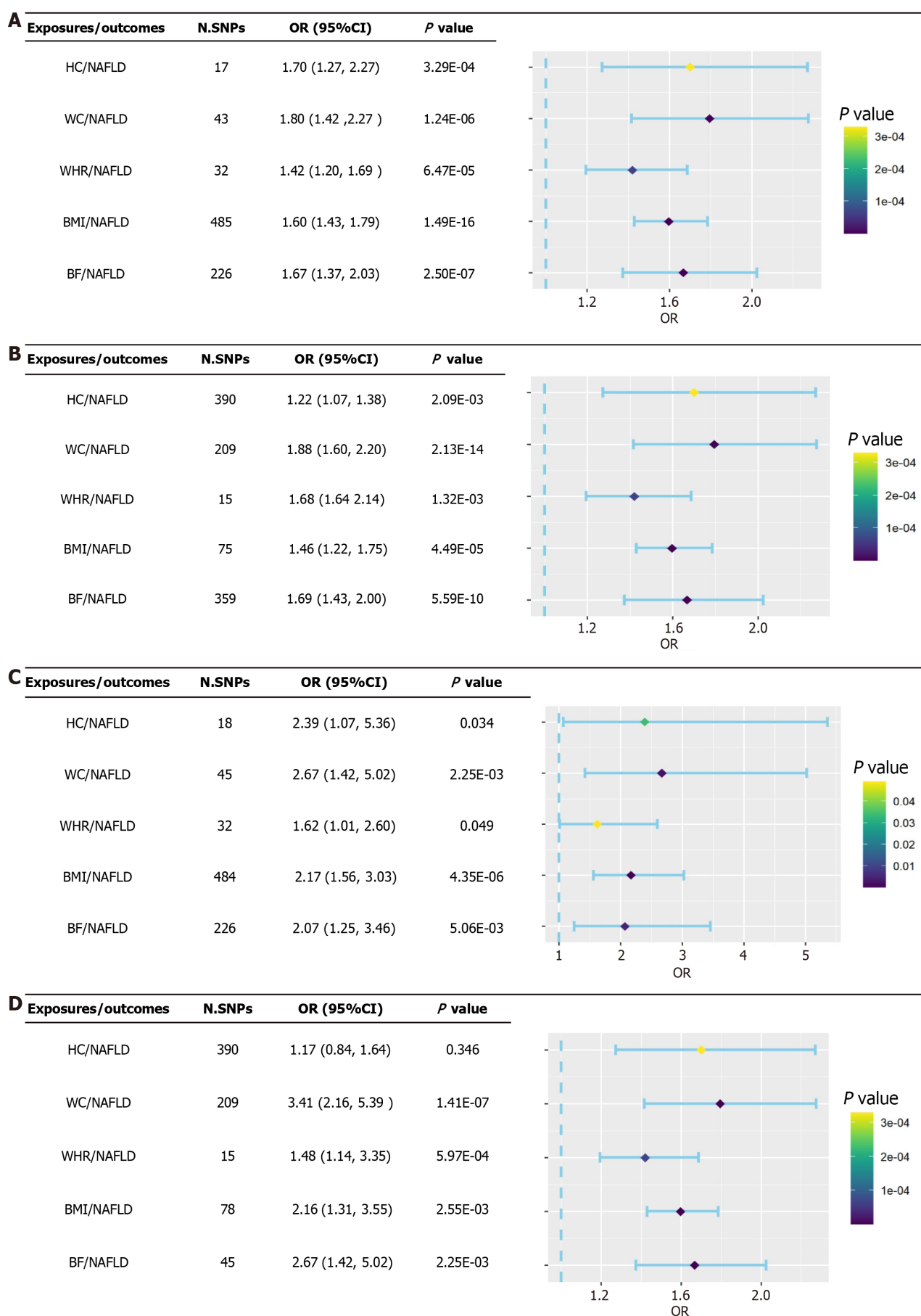
MVMR analysis of potential effects between WC and NAFLD, BF and NAFLD

After adjusting for BMI and smoking, the findings of the MVMR demonstrated that the relationship between the genetically determined increase in WC and an increase in the risk of NAFLD remained significant ($OR = 1.58$, 95%CI: 1.06-2.35, $P = 0.025$), indicating that WC may be a factor for increased risk of NAFLD. Nevertheless, a causal relationship between BF and the risk of NAFLD was not supported by the MVMR findings ($OR_{IVW} = 1.02$, 95%CI: 0.60-1.73, $P = 0.941$; **Figure 2E**). Furthermore, causal relationships between HC, WHR (BMI adjusted), BMI, and the risk of NAFLD were not supported by the cross-validation findings.

DISCUSSION

The primary causes of the increased global burden of NAFLD are changes in lifestyle and dietary habits, and identifying risk factors is particularly crucial for disease prevention and control[28]. Previous studies have demonstrated that anthropometric indicators may primarily be risk factors for developing NAFLD[29]. Nevertheless, conflicting findings exist from various studies, and the causal relationships still need to be clarified[4,30]. Risky relationships between WC and BF for NAFLD were observed from repeated-validation TSMR. After excluding the confounding effects of BMI and smoking, the direct causal effect of BF on NAFLD could not be pursued in subsequent MVMR studies.

As a critical indicator of abdominal obesity (AO) patterns, a strong relationship exists between WC and the early risk of all-cause mortality. An increase of 10 cm in WC for an individual results in an increase of 11% points in the risk of all-cause mortality[31]. Furthermore, positive relationships with the risk of morbidity and mortality of major chronic diseases were exhibited by WC[32]. The importance of WC as an indicator of population health and long-term adverse outcomes was emphasized by these results. In addition, a relationship between WC and NAFLD was observed. A study in an adolescent population indicated that AO was a predictor of pulmonary fibrosis in children and adolescents with NAFLD, with WC as the primary measure[33]. Furthermore, a Korean study discovered a direct relationship between a larger WC and an increased risk of developing NAFLD[33]. This result was supported by a study of the Iranian population[34]. Although numerous studies have demonstrated that AO is a risk factor for NAFLD, with WC being the



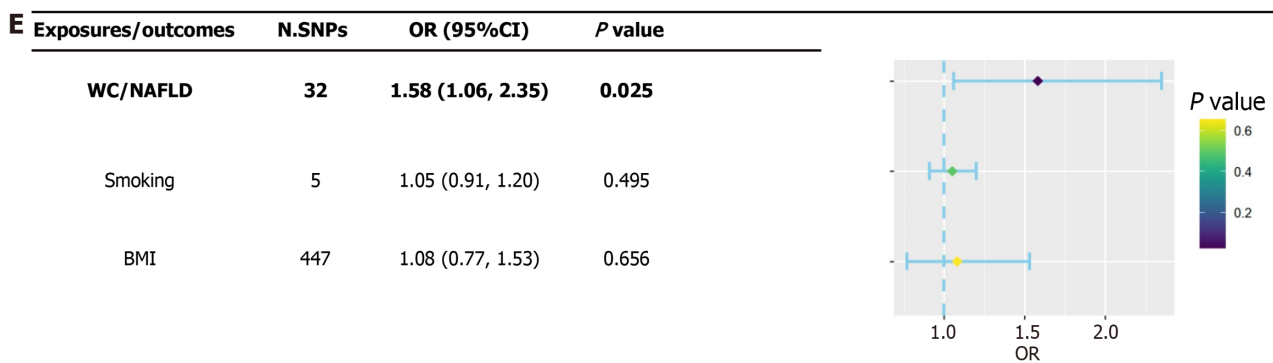


Figure 2 Forest plot of causal associations of anthropometric indicators on the risk of nonalcoholic fatty liver disease. A: Forest plot of causal associations of anthropometric indicators (discovery dataset) on the risk of nonalcoholic fatty liver disease (NAFLD) (United Kingdom biobank); B: Forest plot of causal associations of anthropometric indicators (discovery dataset) on the risk of NAFLD (FinnGen); C: Forest plot of causal associations of anthropometric indicators (replication dataset) on the risk of NAFLD (United Kingdom biobank); D: Forest plot of causal associations of anthropometric indicators (replication dataset) on the risk of NAFLD (FinnGen); E: Causal associations of waist circumference on the risk of NAFLD after adjusting for body mass index and smoking. HC: Hip circumference; WC: Waist circumference; WHR: Waist-to-hip ratio; BMI: Body mass index; BF: Body fat percentage; NAFLD: Nonalcoholic fatty liver disease; SNPs: Single nucleotide polymorphisms; OR: Odds ratio.

primary measure, most of them have been cross-sectional studies or single-measurement findings and the long-term dynamic changes in an individual's WC in response to changes in growth, development, and lifestyle, among other things, have not been sufficiently evaluated. From a genetic viewpoint, our results indicated a causal relationship between greater WC and higher NAFLD. The biological relationship can be explained by several possible mechanisms. On the one hand, AO reflects the excessive accumulation of visceral adiposity (VAT), which increases WC; on the other hand, an increase in WC results in a substantial collection of VAT, enabling the continuous secretion of pro-inflammatory cytokines, such as tumor necrosis factor- α . This activates inflammatory signaling pathways, thereby mediating metabolic homeostasis and insulin sensitivity of the organism[35-37].

BF measures the amount of fat in a person's body and has been identified as a crucial risk factor for cardiometabolic [38]. BF is strongly related to the risk of NAFLD, as indicated by the Rotterdam study, and this relationship is more visible in the female population. Furthermore, in a longitudinal relationship investigating the effect of BF changes on NAFLD incidence and remission, Kim *et al*[39] demonstrated that increased BF was longitudinally associated with increased risk of NAFLD and negatively associated with NAFLD regression[39]. From the two replication datasets, our combined TSMR findings indicated that increased BF substantially increased the risk of NAFLD. After excluding the effects of BMI and smoking, MVMR rejected a direct causal impact of BF on NAFLD.

Furthermore, the relationships of HC, WHR (BMI adjusted), and BMI with NAFLD have been analyzed in numerous studies. A cross-sectional study of the prevalence of NAFLD in pathologically overweight women in South India revealed that BMI, BF, and BWP are three indicators that can be used to a large degree as indicators of the development of NAFLD [40]. Improved cardiovascular metabolism is associated with HC. The development of NAFLD may be triggered by excessive HC, as it has been closely associated with muscle mass, and improving muscle mass will reduce insulin resistance and decrease the likelihood of NAFLD[41,42]. Indicators commonly employed to evaluate nutritional conditions and obesity, such as BMI and WHR (BMI adjusted), primarily reflect the whole-body fat problem rather than VAT. WC is more reflective of the volume of VAT than BMI[43]. Consequently, some studies support visceral obesity with WC and BF as primary measures of a crucial risk factor for NAFLD[44]. Based on European GWAS pooled data, a previous study reported partially different findings compared with our findings, indicating that WHR may be a potential risk factor for NAFLD[45]. Several factors could explain this difference. First, the WHR (BMI adjusted) data used in the GWAS were corrected for BMI, indicating that changes in WHR independent of BMI might not be fully captured by the selection of WHR (BMI adjusted) SNPs. Second, the causal inferences included fewer SNPs, resulting in limited phenotypic differences and reduced statistical power to detect a true causal association between WHR and NAFLD risk. Furthermore, differences in study populations and sample sizes could have contributed to the inconsistent findings. To address these challenges, cross-repeated validation was performed using two independent GWAS datasets to eliminate bias resulting from data selection and enhance the credibility of our findings.

Several limitations require attention. First, the exposure and outcome datasets were obtained from European ancestry; therefore, these findings may not be generalizable to other populations with different genetic backgrounds. Further studies are required to validate these results in other ethnic populations. Second, although the *F*-statistic can be used to evaluate the first hypothesis, verifying the second and third hypotheses is generally challenging and may lead to potential bias. Third, the causal relationship between anthropometric indicators and NAFLD in different gender/age groups cannot be investigated due to the lack of personal demographic information on anthropometric indicators.

Although previous studies have extensively investigated the relationship between AO and NAFLD, there are limited studies on the association between other anthropometric measures[45,46]. This study expands and validates these results by incorporating a wider range of anthropometric measures, offering additional supporting evidence for a causal association in early-stage NAFLD. Compared with previous studies, this study has numerous notable advantages. Although the association between WC, BMI, and BMI-adjusted WHR and NAFLD has been partially examined in

previous studies, evidence for a causal association between other anthropometric indicators (such as HC and BF) and NAFLD remains limited. To the best of our knowledge, the causal association between the five major anthropometric indicators and NAFLD was comprehensively assessed for the first time in this study using UVMR and MVMR approaches. This study provides valuable causal evidence and directionality for the early prediction and diagnosis of NAFLD. The results of this study were replicated using GWAS data from a European population to enhance the reliability of the findings and ensure robust conclusions. Data from two distinct sources were examined in this replication, thereby increasing the potential for identifying new opportunities. In our preliminary analysis, several unidirectional causal associations, such as HC, WC, WHR (BMI adjusted), BMI, and BF, were identified. However, nonreproducible causal relationships between four indicators (HC, WHR, BMI, BF) and NAFLD were excluded from our conclusion to enhance the reproducibility of our approach and reinforce the strength of our conclusions.

CONCLUSION

This study demonstrates that genetically determined increased WC maintains a positive and causal association with NAFLD, even in the presence of confounders, including BMI and smoking. This underscores the potential of WC as a reliable indicator for the early identification and diagnosis of NAFLD.

ARTICLE HIGHLIGHTS

Research background

This study offers the first comprehensive assessment of causal associations between five anthropometric measures and nonalcoholic fatty liver disease (NAFLD) by using both Univariate Mendelian randomization (MR) and Multivariable MR methods. Considering the possibility of potential chance in the results, we additionally selected another exposure and outcome Genome-wide association study data from European population for replication, including a cross-analysis of two different sources of data.

Research motivation

Although the etiology of NAFLD has not been thoroughly understood, the emerging roles of anthropometric indicators in assessing and predicting the risk of NAFLD have been highlighted by accumulating evidence. Numerous previous observational studies have reported relationships between the risk of NAFLD and noninvasive quantitative measurements of the body, such as anthropometric indicators, which comprise height, weight, hip circumference, waist circumference (WC), waist-to-hip ratio, body mass index (BMI), and body fat percentage. Contradictory findings have been obtained in some other studies, indicating no relationship between anthropometric indicators and the risk of NAFLD. The causal relationships between anthropometric indicators and NAFLD risk remain undetermined, considering these inconsistent findings and the absence of randomized controlled studies.

Research objectives

This study demonstrates that genetically determined increased WC maintains a positive and causal association with NAFLD, even in the presence of confounders, including BMI and smoking. This underscores the potential of WC as a reliable indicator for the early identification and diagnosis of NAFLD.

Research methods

MR is a novel epidemiological tool that employs genetic data to investigate the causal relationship between exposure and outcome. Generally, genetic variants are independent of disease state and are randomly assigned to offspring through the maternal generation. Consequently, the limitations of conventional observational design can be overcome, and biases such as potential confounders and reverse causality can be minimized. Previous studies have demonstrated the use of MR to investigate causal relationships between NAFLD and numerous diseases, including cardiovascular disease and psoriasis.

Research results

Genetically determined increased WC maintains a positive and causal association with NAFLD, even in the presence of confounders, including BMI and smoking. Several limitations require attention. First, the exposure and outcome datasets were obtained from European ancestry; therefore, these findings may not be generalizable to other populations with different genetic backgrounds. Further studies are required to validate these results in other ethnic populations. Second, although the F-statistic can be used to evaluate the first hypothesis, verifying the second and third hypotheses is generally challenging and may lead to potential bias. Third, the causal relationship between anthropometric indicators and NAFLD in different sex/age groups cannot be investigated due to the lack of personal demographic information on anthropometric indicators.

Research conclusions

This study demonstrates that genetically determined increased WC maintains a positive and causal association with

NAFLD, even in the presence of confounders, including BMI and smoking. This underscores the potential of WC as a reliable indicator for the early identification and diagnosis of NAFLD.

Research perspectives

Future studies should consider using WC as an auxiliary measurement for identifying NAFLD.

ACKNOWLEDGEMENTS

The authors express their gratitude to the Figdraw website and the CHiPlot website for their provision of scientific drawing tools.

FOOTNOTES

Co-first authors: Xian-Pei Xiao and Yong-Jun Dai.

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Author contributions: Xiao XP and Dai YJ conceived and designed the study; Zhang Y and Yang M conducted preliminary analysis of the original data; Dai YJ and Xie J completed the drawing of the charts; Xiao XP, Dai YJ, Zhang Y and Yang M performed data analysis and result validation; Xiao XP wrote the paper. All authors read and approved the final manuscript. Xiao XP proposed, designed this study, conducted data analysis, and wrote the first draft of the paper. Dai YJ participated in proposing, designing this study, performing data analysis, and drawing graphs. Both authors have made vital and integral contributions to the completion of the project and are therefore eligible to be co-first authors of the paper. As co-corresponding authors, Yang ZJ and Chen G played an important and indispensable role in experimental design, data interpretation, and manuscript preparation. Funding for this project was applied for and obtained by Chen G. Yang ZJ conceived, designed and supervised the entire project process. The collaboration of Yang ZJ and Chen G was critical to the publication of this and other manuscripts still in preparation.

Supported by Science and Technology Research Project of Sichuan Administration of Traditional Chinese Medicine, No. 2023MS419.

Institutional review board statement: No need for ethical approval as used of anonymous open data.

Conflict-of-interest statement: All the authors report no relevant conflicts of interest for this article.

Data sharing statement: The data and material that support the findings of this study are available from public datasets that could be found in IEU OPEN GWAS and FinnGen research project.

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Country/Territory of origin: China

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S-Editor: Li L

L-Editor: A

P-Editor: Zhao S

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