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**DOI**

[10.1038/s41586-024-07148-y](https://doi.org/10.1038/s41586-024-07148-y)

**Publication date**

2024

**Document Version**

Final published version

**Published in**

Nature: international weekly journal of science

**Citation (APA)**

Karjalainen, M. K., Karthikeyan, S., Oliver-Williams, C., Sliz, E., Allara, E., Fung, W. T., Surendran, P., van den Akker, E. B., & More Authors (2024). Genome-wide characterization of circulating metabolic biomarkers. *Nature: international weekly journal of science*, 628(8006), 130-138.  
<https://doi.org/10.1038/s41586-024-07148-y>

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# Genome-wide characterization of circulating metabolic biomarkers


<https://doi.org/10.1038/s41586-024-07148-y>

Received: 7 November 2022

Accepted: 1 February 2024

Published online: 06 March 2024

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Genome-wide association analyses using high-throughput metabolomics platforms have led to novel insights into the biology of human metabolism<sup>1–7</sup>. This detailed knowledge of the genetic determinants of systemic metabolism has been pivotal for uncovering how genetic pathways influence biological mechanisms and complex diseases<sup>8–11</sup>. Here we present a genome-wide association study for 233 circulating metabolic traits quantified by nuclear magnetic resonance spectroscopy in up to 136,016 participants from 33 cohorts. We identify more than 400 independent loci and assign probable causal genes at two-thirds of these using manual curation of plausible biological candidates. We highlight the importance of sample and participant characteristics that can have significant effects on genetic associations. We use detailed metabolic profiling of lipoprotein- and lipid-associated variants to better characterize how known lipid loci and novel loci affect lipoprotein metabolism at a granular level. We demonstrate the translational utility of comprehensively phenotyped molecular data, characterizing the metabolic associations of intrahepatic cholestasis of pregnancy. Finally, we observe substantial genetic pleiotropy for multiple metabolic pathways and illustrate the importance of careful instrument selection in Mendelian randomization analysis, revealing a putative causal relationship between acetone and hypertension. Our publicly available results provide a foundational resource for the community to examine the role of metabolism across diverse diseases.

Large genome-wide association studies (GWASs) coupled with metabolic profiling platforms have successfully identified many loci associated with circulating metabolic traits<sup>1–7,12–16</sup>. For example, studies combining genomics with detailed metabolic profiling from a high-throughput metabolomics platform based on nuclear magnetic resonance spectroscopy<sup>17</sup> have enabled the identification of dozens of loci for traits associated with circulating lipid, lipoprotein and fatty acid and small molecules such as amino acids<sup>2,4,5,9,18,19</sup>. These studies have provided novel insights into the biology of human metabolism and have guided large-scale epidemiological studies, such as Mendelian randomization analyses to infer causal relationships<sup>17</sup>. Here, using the same NMR metabolomics platform from Nightingale Health with an updated quantification version, we considerably extend our previous GWAS<sup>4</sup> of 123 circulating metabolic traits in up to around 25,000 participants to study 233 traits in more than 135,000 participants.

## Genetic discovery

GWAS was performed under the additive model separately in each of 33 cohorts (Supplementary Table 1). Subsequent meta-analysis involved 233 metabolic traits (Supplementary Table 2), including 213 lipid and lipoprotein parameters or fatty acids and 20 non-lipid traits (amino acids, ketone bodies and glycolysis/gluconeogenesis, fluid balance and inflammation-related metabolites). After variant filtering and quality control, up to 13,389,637 imputed autosomal single-nucleotide polymorphisms (SNPs) were included in the meta-analysis for up to 136,016 participants.

In the meta-analysis, we detected genome-wide significant associations for all 233 metabolic traits (Supplementary Data Figs. 1–3 and Supplementary Tables 4 and 5) with extensive pleiotropy and polygenicity. We detected 276 broad regions (defined as a  $\pm 500$ -kb region around the set of genome-wide significant SNPs) associated with at least one metabolic trait (Fig. 1a and Supplementary Table 4). Eighty-six of these regions were associated with just a single metabolic trait, whereas most regions harboured associations with multiple traits (Fig. 1b,c) up to a maximum of 214 associated traits at the well-characterized lipid-associated *APOE* region. The lipid, lipoprotein and fatty acid traits were mostly demonstrably polygenic, with 60 traits having associations at more than 50 loci, 137 traits (64.3%) having associations at 20–50 loci, and 16 traits (7.5%) having associations at fewer than 20 loci (Supplementary Tables 5 and 6). Most non-lipid traits had substantially fewer associated loci (13 with fewer than 20 associated traits; 65% of all 20 non-lipid traits), including 3 glucose metabolism-related traits (lactate, pyruvate and glycerol) having fewer than 5 associated loci, whereas glycoprotein A and some amino acids had associations at 20–33 loci and creatine had associations at 49 loci (Supplementary Tables 5 and 6). The non-lipid traits accounted for most of the regions with a single associated trait ( $n = 67$ ; 78%), and the majority ( $n = 163$ ; 57%) of the regions with non-lipid trait associations had fewer than 5 associated metabolic traits in total. By contrast, the lipid, lipoprotein and fatty acid trait-associated regions ( $n = 186$ ) were generally more pleiotropic with 75% ( $n = 140$ ) of the regions being associated with 5 or more traits. The pleiotropy difference is owing to the fact that lipoprotein metabolism is a continuum, with genes often affecting several

particle categories, whereas the non-lipid traits are often affected by more distinct processes and enzymatic modifications, thus leading to less pleiotropy<sup>3,9,20</sup>. Within the 276 regions, we found 8,795 lead SNP–lead trait associations corresponding to 1,447 unique lead SNPs (Supplementary Table 5). After resolving independent signals on the basis of pairwise linkage disequilibrium (LD), we concluded that the 276 broad regions involved at least 443 independent loci. We estimated the genome-wide common variant heritability for 223 traits also available in UK Biobank (Supplementary Table 6). Median trait heritability was 0.29, of which only around a quarter was explained by the lead SNPs, supporting the high polygenicity of many of the traits.

## Ancestry-stratified analyses

To investigate the generalizability of our associations across ethnic groups and search for additional ancestry-specific association signals, we conducted ancestry-stratified analyses of South Asian (five cohorts, 11,340 participants), East Asian (one cohort, 4,435 participants), Finnish (six cohorts, 27,577 participants) and non-Finnish European (21 cohorts, 92,664 participants) cohorts (Supplementary Table 7). To investigate the generalizability further, we additionally performed a post hoc comparison of the associations to a small population with African ancestry ( $n = 1,405$ ). Associations were strongly positively correlated across ancestry groups (Extended Data Fig. 1), suggesting that associations are broadly transferable across ancestries. Stronger correlations were seen between Finnish and non-Finnish Europeans ( $r = 0.96$ ) compared with East Asians and South Asians ( $r \approx 0.7$ ) and Africans ( $r \approx 0.4$ ). For some loci, effect estimates were notably stronger in one ancestry than another. The number of genome-wide significant associations was strongly related to sample size, ranging from 7,002 associations in the non-Finnish Europeans to 331 in the East Asian and 97 in the African participants. We did not detect any novel genomic regions in any of the ancestry groups beyond the 276 discovered in the ancestry-combined meta-analysis, suggesting that substantially larger sample sizes of participants with non-European ancestry will be required in future studies.

## Associations in UK Biobank

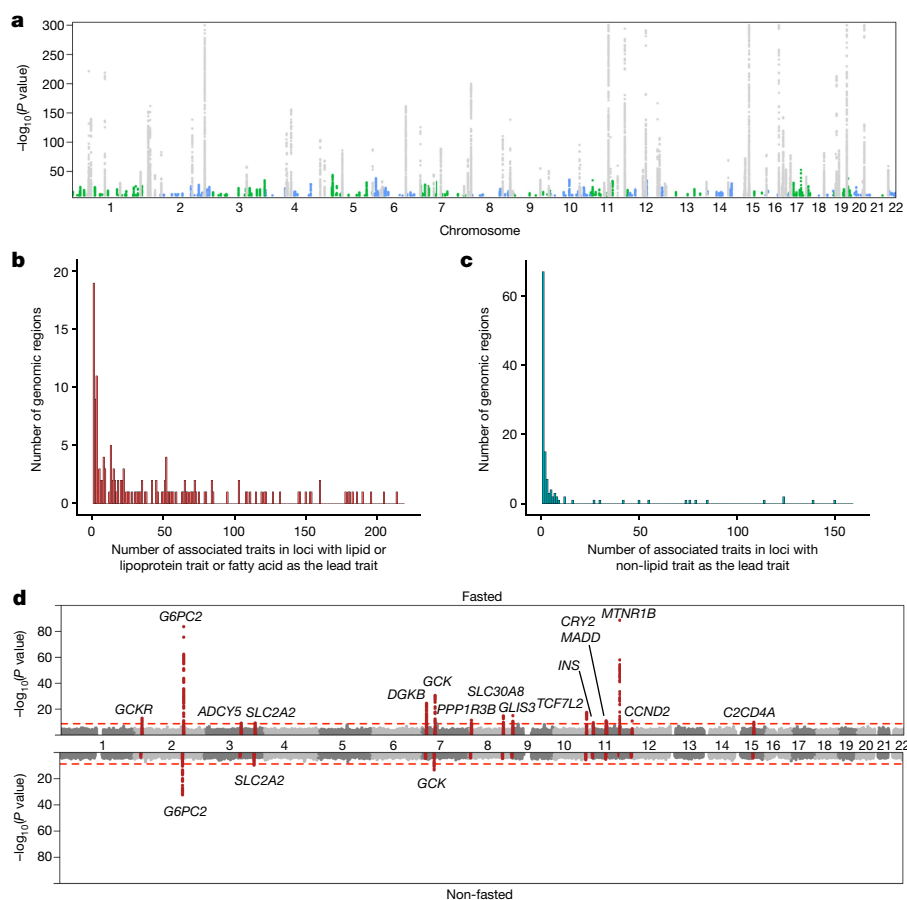
The availability of NMR data from the UK Biobank resource<sup>21</sup> (March 2021 release) enabled us to check for associations of the lead variants in an independent population and to assess the effects of participant characteristics and sample-related factors on our associations. Of the 8,502 lead SNP–metabolic trait pairs that could be tested in up to 115,078 UK Biobank participants with European ancestry, 5,442 (64.0%) associated at  $P < 5 \times 10^{-8}$ , and a further 772 (9.1%; 328 unique SNPs) associated at  $P < 1 \times 10^{-5}$  (Supplementary Table 8). When we performed further stratified analyses in cohorts with different sample types (serum,  $n = 90,223$ ; plasma,  $n = 45,793$ ), and in fasted ( $n = 68,559$ ) and non-fasted ( $n = 58,112$ ) cohorts, we detected that, in addition to subtle differences in population ancestry between the studies, sample type and fasting status were probably the major drivers of non-replication. The UK Biobank NMR measurements were performed on EDTA plasma samples, whereas the current meta-analysis involved predominantly serum samples. For example, several of the non-replicating associations with phenylalanine were in coagulation-related loci (for example, *KLKB1*, *F12*, *KNG1* and *FGB*) but these signals were absent in UK Biobank (Extended Data Fig. 2 and Supplementary Table 8)—therefore we speculate that the removal of clotting factors in the preparation of serum could reveal associations with phenylalanine via coagulation. Two loci (*NHLRC1*, lead SNP rs73726535; *TXNRD1*, lead SNP rs191631370) also had associations for phenylalanine in UK Biobank that were absent in the current meta-analysis. Similarly, we found associations with glucose that did not replicate in the UK Biobank, including a well-known association at the melatonin receptor 1B gene<sup>22</sup> (*MTNR1B*), a key regulator

in glucose metabolism (rs10830963; meta-analysis  $P$  value =  $1.5 \times 10^{-60}$ ; UK Biobank  $P$  value = 0.60). The UK Biobank predominantly includes non-fasted samples, but the current meta-analysis mainly consists of cohorts (26 cohorts) with fasted samples (Supplementary Table 1), and our fasting-stratified meta-analysis suggested that some of the glucose associations were driven by cohorts with predominantly fasted samples (Fig. 1d and Extended Data Fig. 3) and are thus absent in UK Biobank. In addition to *MTNR1B* rs10830963 ( $P$  values  $2.9 \times 10^{-89}$  and 0.57 in meta-analysis of fasted and non-fasted cohorts, respectively), the association of which was also previously shown to be absent in non-fasting samples<sup>23</sup>, *GLIS* family zinc finger 3 (*GLIS3*; a known diabetes risk gene<sup>24</sup> with a role in pancreatic beta cell biology) rs10974438 represents another example of an association that was not robustly replicated in UK Biobank (meta-analysis  $P$  value =  $4.0 \times 10^{-14}$ ; UK Biobank  $P$  value = 0.001) and was characterized by the absence of signals in the non-fasted cohorts ( $P$  values  $1.1 \times 10^{-15}$  and 0.14 in meta-analysis of fasted and non-fasted cohorts; Extended Data Fig. 3).

Many of the metabolic trait associations differed by sample type and fasting status, although comparisons with the overall associations are complicated by the reduced power of the stratified analyses. For example, associations of several lipoprotein subclass measures were substantially affected by fasting status at loci with central roles in lipid biology, such as *LPA* and *ANGPTL3* (Supplementary Table 9). Similarly, multiple loci had greater than twofold higher or lower effect estimates in cohorts using serum compared to those using plasma (Supplementary Table 9). These differences were detected both for lipid and non-lipid traits, with some associations being notably augmented by removal of plasma samples. However, only 7 additional loci (beyond the 276 initially associated genomic regions) were detected in analyses stratified by sample type and 10 were detected in fasting-stratified analyses (Supplementary Tables 10 and 11)—for example, *C2CD4A* rs10083587 for glucose and *KAT5* rs12787843 for creatinine, both of which showed associations only in fasting cohorts ( $P$  values  $1.3 \times 10^{-10}$  and  $7.6 \times 10^{-10}$ , respectively). We note that the effects of the sample type and fasting status require careful consideration when interpreting the results of GWAS of metabolic traits and conducting downstream analyses, such as Mendelian randomization studies using trait-associated variants as instruments.

## Novel loci and candidate genes

We conducted extensive manual curation to prioritize 231 likely causal genes with clear biological relevance to the associated traits at 297 (67.0%) of the 443 loci (Methods). As some regions were extremely complex and pleiotropic owing to overlapping genetic associations of up to 11 independent lead variants with heterogeneous associations across the metabolic traits, we characterized these loci in detail to pinpoint potential multiple probable causal genes within each locus (Supplementary Table 5). For example, in a 7.6-Mb region on chromosome 16 with 139 associated metabolic traits, we identified 6 distinct biologically relevant potential causal genes: lecithin-cholesterol acyltransferase (*LCAT*; associated with multiple lipoprotein subclass measures), solute carrier family 7 member 6 (*SLC7A6*; associated with acetate and creatinine), pyruvate dehydrogenase phosphatase regulatory subunit (*PDPR*; associated with pyruvate and amino acids), alanyl-tRNA synthetase 1 (*AARS*; associated with pyruvate and amino acids), tyrosine aminotransferase (*TAT*; associated with tyrosine) and haptoglobin (*HP*; associated with a range of lipoprotein subclass measures, fatty acids, cholesterol, apolipoprotein B (apoB) and glycoprotein acetylation). This locus exemplifies the complexity of the metabolic trait-associated loci. For additional loci without an obvious biological candidate, we assigned a further 39 probable causal genes on the basis of SNP function or the presence of probable functional (missense, stop gained or splice region) variants in strong LD ( $r^2 \geq 0.8$ ) with the lead variant (Supplementary Table 5).



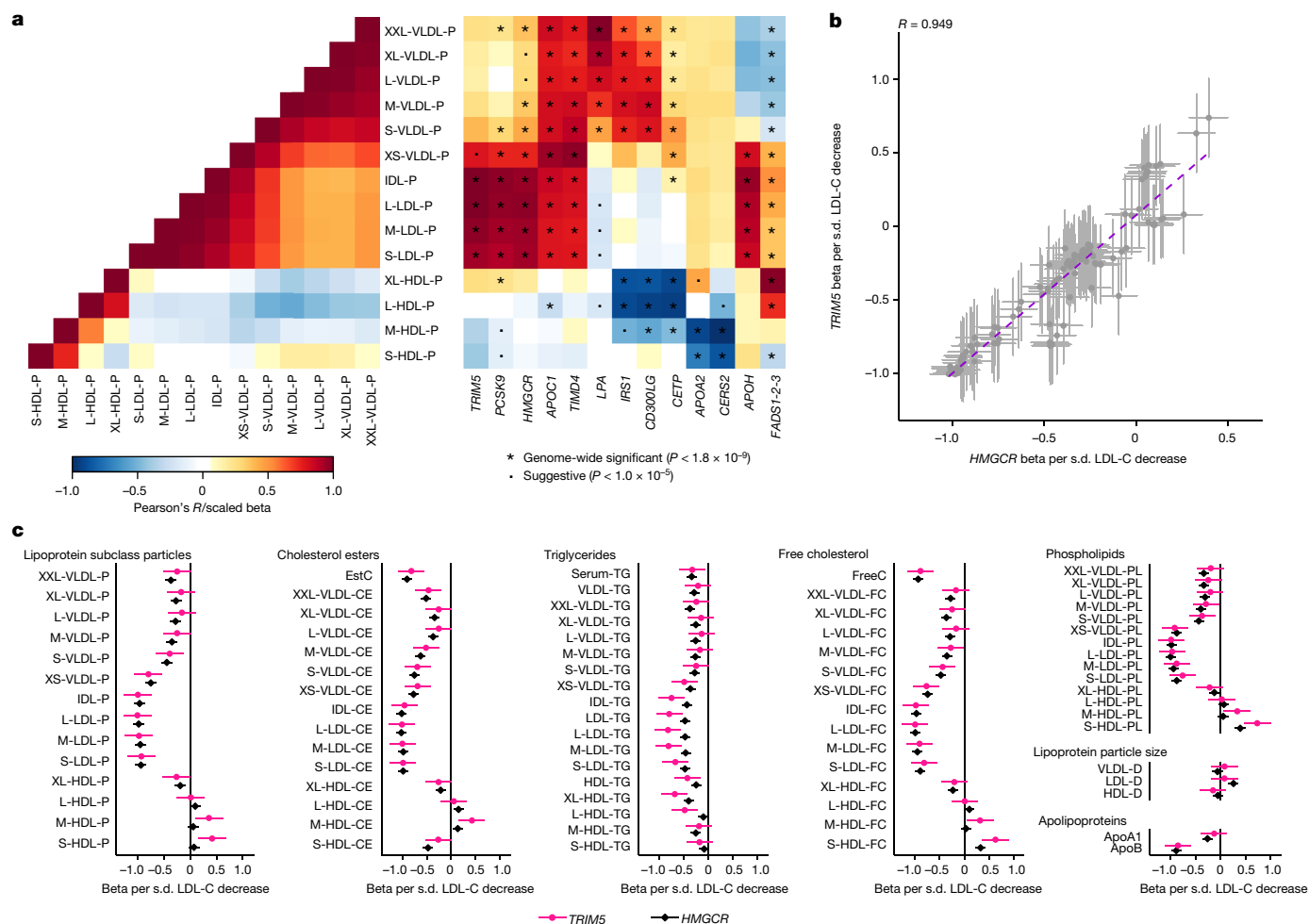
**Fig. 1 | Results of the GWAS meta-analysis of 233 metabolic traits. a**, Manhattan plot summarizing the metabolic trait associations from inverse variance-weighted GWAS meta-analysis. Loci that do not overlap with those identified in the previous large-scale NMR metabolomics GWAS<sup>4,5</sup> are shown in blue and green. Only genome-wide significant SNPs (two-sided  $P < 1.8 \times 10^{-9}$ ) are shown and  $-\log_{10}(P\text{value})$  were capped at 300. **b,c**, Numbers of associated metabolic traits at the 276 associated genomic regions are shown separately for genomic

regions in which the lead trait was a lipid, lipoprotein or fatty acid trait (**b**; 155 loci; median 24 traits per locus) and for those in which the lead trait was a non-lipid trait (**c**; 121 loci; median one trait per locus). **d**, Results of the GWAS for glucose for the fasted (top; total  $n = 68,559$ ) and non-fasted (bottom; total  $n = 58,112$ ) cohorts. The red line indicates the threshold for genome-wide significance. The 500-kb regions around lead SNPs in the fasted cohorts are highlighted.

We performed an extensive comparison of the discovered associations to previously reported genetic associations of metabolic traits and traditional clinical lipids (high density lipoprotein C (HDL-C), low density lipoprotein C (LDL-C), triglycerides and total cholesterol; Supplementary Table 5). In comparison to previous large-scale NMR metabolomics GWAS<sup>4,5</sup>, we identified 212 additional associated genomic regions (Supplementary Table 4). These included 138 novel genomic regions for the lipoprotein, lipid and fatty acid traits, and 113 novel regions associated with the non-lipid traits. New associations for several lipoprotein subclass measures were detected in loci previously associated with clinical lipids, such as the locus containing low density lipoprotein receptor adapter protein 1 gene (*LDLRAP1*), which is involved in cholesterol metabolism. This locus was previously known to be associated with LDL-C, triglycerides and total cholesterol<sup>25–27</sup>, and we found associations at this locus with several lipoprotein subclass measures, lipids and fatty acids (Supplementary Table 5). Locus containing the sterol *O*-acyltransferase 2 gene (*SOAT2*; which functions in cholesterol metabolism) represents another example of a novel biologically plausible locus associated with the lipoprotein and lipid traits. Our analyses also identified genetic associations with detailed lipoprotein subclass measures in loci that have not previously been reported to be associated with traditional clinical lipids. Compared with the largest trans-ancestry study of clinical lipids to date<sup>27</sup>, we detected associations at twelve additional

loci (Supplementary Table 5) for the lipid and lipoprotein traits (corresponding to 6.5% of all lipoprotein and lipid trait-associated regions); gene encoding type 2 lactosamine  $\alpha$ -2,3-sialyltransferase (*ST3GAL6*; which functions in glycolipid metabolism) represents an example of a biologically plausible gene associated with multiple lipoprotein subclass measures and lipids.

Novel loci were also detected for small molecules such as phenylalanine and glutamine. For phenylalanine, we detected associations at 13 loci. Novel phenylalanine-associated loci include both a well-known metabolic trait-associated locus (*FADS1–FADS2*) and two novel, biologically plausible loci (*GSTA2* and *SLC2A4RG*). For example, *SLC2A4RG* encodes *SLC2A4* regulator, a transcription factor involved in the activation of solute carrier family 2 member 4 (*SLC2A4*, also known as *GLUT4*), a key regulator of glucose transport. For glutamine, we detected associations at 26 loci. Of note, seven of the loci were associated only with glutamine (*GLS*, *PLCL1*, *SFXN1*, *KCNK16*, *MED23*, *SLC25A29* and *PCK1*). Thus, these associations are likely to represent biology local to glutamine, most of the loci having biologically plausible candidate genes with roles in glutamine metabolism (*GLS*), amino acid transport (*SFXN1* and *SLC25A29*) or glucose and gluconeogenesis-related pathways (*PCK1* and *KCNK16*). *KCNK16*, a known type 2 diabetes susceptibility gene that encodes the potassium channel subfamily K member 16, a pancreatic potassium channel, represents an example of a novel glutamine-associated locus with a role in glucose biology<sup>28,29</sup>.



**Fig. 2 | Effects of SNPs across lipoprotein and lipid traits. a**, Heat maps of the correlation structure of lipoprotein subclass particle concentrations (left) and the association landscapes of exemplar SNPs (right). In the heat maps, pairwise correlations of lipoprotein subclass particle concentrations (calculated in FINRISK 1997; left) and effect estimates for the SNP-metabolic trait associations (right) are represented as a colour range. The SNP effect sizes were scaled relative to the absolute maximum effect size in each locus. Each column represents a single SNP, and each row corresponds to a single metabolic measure. Two-sided tests were used. **b, c**, Scatter plot (**b**) and forest plots (**c**)

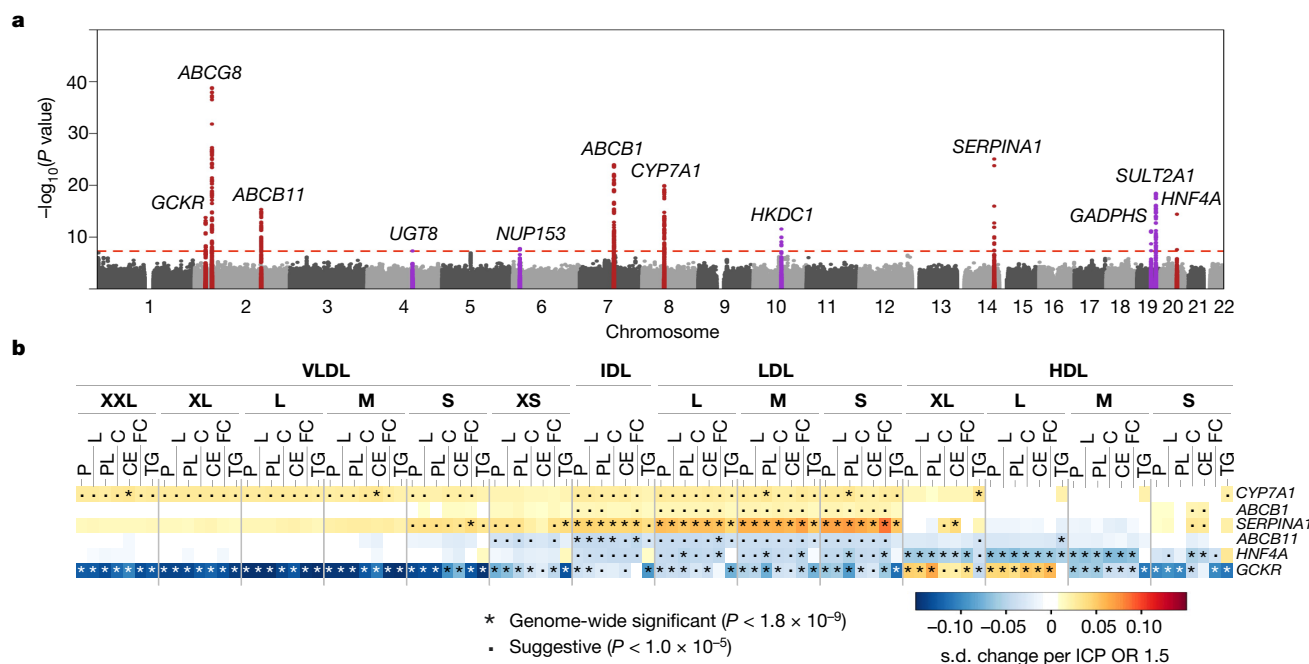
of the effect estimates (betas and 95% confidence intervals) for  $TRIM5$  and  $HMGCR$  lead SNPs (rs11601507 and rs12916, respectively;  $n = 136,016$  individuals) across the lipoprotein and lipid traits. **b**, A best fit regression line (purple dashed line) and an estimate of Pearson's correlation coefficient  $R$  (for betas of 116 SNP-trait pairs) are shown. The effect estimates (s.d. units) were scaled relative to a one-s.d. decrease in LDL cholesterol. VLDL, LDL, IDL and HDL are classified into different particle sizes (in order of decreasing size: XXL, XL, L, M, S and XS). Detailed descriptions of the metabolic traits and abbreviations are shown in Supplementary Table 2. ApoA1, apolipoprotein A1.

## Effects of apoB variants

To provide insights into the distinct ways in which lipid loci can affect the continuum of lipoprotein metabolism, we characterized clusters of genes with similar metabolic association profiles. The effect estimates were scaled by dividing all effect estimates of a given SNP using the strongest association effect estimate across all metabolic associations in each locus. In this way, the scaled effect estimates for all SNPs were between  $-1$  and  $1$ , and the statistical strength of an association affects the clustering less, and more emphasis is given to the association landscape in guiding the clustering. We concentrated on 134 loci with nominal evidence ( $P < 0.05$ ) of an association with apoB, as recent studies have highlighted the predominant role of apoB in coronary artery disease aetiology<sup>30–32</sup>. The clustering of the loci produced at least seven major clusters of loci (Extended Data Figs. 4 and 5). The top cluster in Extended Data Fig. 4 is very similar to the previously observed epidemiological association profile with type 2 diabetes risk<sup>33</sup> and adiposity<sup>34</sup>. The second cluster of loci (Extended Data Figs. 4 and 5) primarily shows increasing triglyceride-rich very low density lipoprotein (VLDL) particles and decreasing

large HDL particles. The genes in this cluster, such as *LPL*, *MLXIPL* and *ANGPTL4*, relate to triglyceride metabolism, and glucose metabolism, exemplified by *GCK*, *GCKR* and *INSR*. The other clusters associate primarily with LDL particles and generally less with other lipoproteins. The lowest cluster includes biologically relevant genes that are known to affect LDL cholesterol in circulation, including *APOB*, *LDLR*, *PCSK9*, *SORT1* and *HMGCR*. Despite the strong correlation structure within the lipid and apolipoprotein traits, we identified several loci with association patterns that do not follow the between-trait correlation structure (Fig. 2a and Extended Data Figs. 4 and 5). For example, some loci (*APOC1* and *TIMD4*) are strongly associated with all the apoB-containing particles (VLDL, intermediate density lipoprotein (IDL) and LDL), whereas other loci are predominantly associated with IDL and LDL particles (*PCSK9*, *HMGCR* and *TRIM5*), with VLDL and the largest HDL particles (*IRS1* and *CD300LG*), or with medium and small HDL particles (*APOA2* and *CERS2*). Several SNPs also exhibit discordant associations within highly correlated metabolic traits (for example, *LPA* and *APOH* SNPs within apoB-containing particles and *FADS* cluster SNP within both apoB-containing and HDL particles; Fig. 2a).





**Fig. 3 | Metabolic trait-associated variants are associated with ICP.**

**a,b**, Manhattan plot of the GWAS of intrahepatic cholestasis of pregnancy (ICP) (**a**) and heat map of loci associated with metabolic traits and ICP (**b**). Twelve loci were associated with ICP in the FinnGen study (1,460 cases, 172,286 controls). **a**, The 500-kb regions flanking the lead SNPs are highlighted, and the nearest gene is indicated for each signal. The ICP GWAS was performed with scalable and accurate implementation of generalized mixed model (SAIGE). Loci that overlap with the loci identified in the NMR meta-analysis are indicated in red. **b**, Loci that are likely to have shared causal variants with the metabolic traits

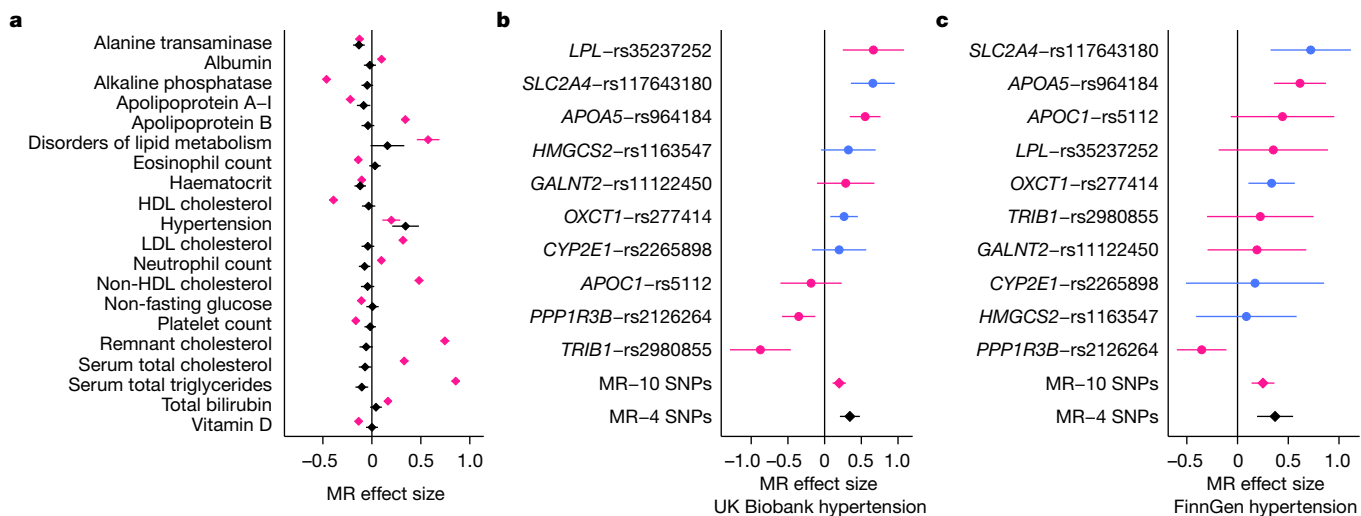
Metabolic profiles of 84 novel loci that were not identified in the previous NMR GWASs<sup>2,4,5</sup> were characterized here using the clustering approach (Extended Data Figs. 4 and 5). As the approach we have taken uses scaled effect estimates, our results are not directly comparable to previous studies which have used unscaled effect estimates<sup>9</sup> or numbers of associations per lipoprotein type<sup>5</sup> in clustering. Even though many loci, such as the master regulator genes *PCSK9* and *LDLR*, clustered mostly similarly as reported previously<sup>5,9</sup>, the genetically calibrated approach applied here can specifically add to the understanding of the detailed metabolic effects of less well-known lipid-associated loci as their metabolic association patterns have not been previously characterized. The tripartite motif-containing protein 5 gene (*TRIM5*) is an example of a poorly characterized locus associated with 42 lipoprotein and lipid traits (Supplementary Table 5). *TRIM5* is best known for its role in antiviral host defence<sup>35</sup>, but variants near *TRIM5* have also been associated with several traits related to liver biology, such as levels of liver enzymes<sup>36</sup>, and have recently been reported to associate with risk of coronary artery disease<sup>37</sup>. Notably, the metabolic effects on the lipoprotein and lipid traits of the lead *TRIM5* variant (rs11601507, p.Val112Ile) appear similar to those of the *HMGR* variant rs12916 (Fig. 2b,c), the metabolic effects of which are concordant with those of statin therapy<sup>38–40</sup>. The mechanism by which *TRIM5* affects lipid and lipoprotein levels and predisposes to coronary artery disease is unclear and it has been speculated to be related to innate immunity<sup>41</sup>. A recent study using a mouse model of nonalcoholic fatty liver disease suggested that *TRIM5* may mediate degradation of DEAD-box protein 5, which could affect mTORC1 signalling and the LDL receptor pathway, consequently affecting lipid accumulation and inflammation<sup>42</sup>. Irrespective of the pathophysiological mechanism, our findings raise the possibility that inhibition of *TRIM5* could provide an alternative therapeutic pathway for reducing the risk of cardiovascular disease via lowering

the concentrations of circulating atherosclerotic apoB-containing lipoprotein particles similar to PCSK9-inhibition therapies that are useful for statin intolerant individuals or for statin users requiring further risk reduction. Although we specifically chose the *TRIM5* association for further investigation, our clustering analysis suggests there are several other novel loci worthy of further in-depth investigation.

## Metabolic trait variants and diseases

To investigate the roles of the metabolic trait-associated variants in disease, we scanned all the disease and trait associations of the 1,447 lead SNPs in the (1) FinnGen study (data freeze 7, up to 309,154 participants, 3,095 phenotypes), a dataset linking genomic information from Finnish participants to digital health care data<sup>43</sup>, and (2) curated collections of published GWASs, including PhenoScanner<sup>44,45</sup> and GWAS catalog<sup>46</sup> (Supplementary Table 5). In addition, we scanned the SNPs for association with gene expression and protein levels.

Most ( $n = 1,279$ ) of the 1,447 lead SNPs had previously reported associations ( $P < 5 \times 10^{-8}$ ) with traits or diseases, including directly relevant outcomes such as use of statin medication and hypercholesterolaemia (Supplementary Table 5). Most of the SNPs ( $n = 1,270$ ) were also associated with messenger RNA (mRNA) or protein levels (Supplementary Table 5), indicating that at least some of the associations are likely mediated by direct or indirect effects of SNPs on mRNA or protein levels. Seven metabolic trait-associated loci (*GSKR*, *ABCG8*, *ABCB11*, *ABCB1*, *CYP7A1*, *SERPINA1* and *HNF4A*) were associated ( $P < 5 \times 10^{-8}$ ) with risk of intrahepatic cholestasis of pregnancy (ICP) in FinnGen (Fig. 3a and Supplementary Table 12), of which all except *ABCG8* showed robust evidence of colocalization or shared regional associations with the metabolic trait associations (Supplementary Table 13). ICP is a cholestatic disorder with onset in the second or third trimester of pregnancy,



**Fig. 4 | Mendelian randomization suggests a causal association between acetone and hypertension.** **a**, Effect estimates (betas per s.d. increase in acetone) from Mendelian randomization (MR) analysis performed under the inverse variance-weighted model are shown for the UK Biobank outcomes that were significant ( $P < 4.88 \times 10^{-6}$ ) with the full (pleiotropic,  $n = 10$  instrument SNPs, pink) or strict (non-pleiotropic,  $n = 4$  instrument SNPs, black) set of instruments. Betas and  $P$  values are shown in Supplementary Table 15. **b,c**, Effect estimates (betas per s.d. increase in acetone) in Mendelian

randomization analysis with hypertension as the outcome in the UK Biobank (**b**; 104,824 cases with hypertension, 367,542 controls) and FinnGen (**c**; 70,651 cases with hypertension, 223,663 controls) datasets. Single-SNP Mendelian randomization effect estimates and 95% confidence intervals are shown, with the SNPs in the strict instrument coloured blue and the other SNPs coloured pink. Mendelian randomization effect estimates are shown with pink and black diamonds for the full instrument (all ten SNPs) and strict instrument (four non-pleiotropic SNPs), respectively.

that is characterized by pruritus and elevated concentrations of serum aminotransferases and bile acids. ICP increases the risk of meconium staining of amniotic fluid, preterm delivery, fetal bradycardia, fetal distress and fetal loss<sup>47</sup>. The genetic background of ICP is poorly characterized with few published GWAS<sup>7,48</sup> and the metabolic effect of the ICP loci has not been characterized. Compared with results of a recent ICP GWAS that included data from meta-analysis of an earlier FinnGen release (data freeze 4) and two other cohorts<sup>48</sup>, associations at nine loci (*GCKR*, *ABCG8*, *ABCB11*, *ABCB1*–*ABCB4*, *CYP7A1*, *SERPINA1*, *GAPDH*–*TMEM147*, *SULT2A1* and *HNF4A*) were replicated here and three novel loci (*UGT8*, *NUP153* and *HKDC1*) were additionally identified. Rare coding variants at two of the loci, within the *ABCB11* and *ABCB4* genes, have additionally been previously reported in ICP<sup>49,50</sup>. A pathway analysis of the ICP-associated loci showed that biological processes related to bile acid, glucose and lipid metabolism were enriched for ICP (Supplementary Table 14), consistent with the metabolic trait associations. For some loci (*CYP7A1*, *ABCB1* and *SERPINA1*), the most profound associations were detected for IDL and LDL particles, whereas two loci (*HNF4A* and *GCKR*) were more pleiotropic, with effects across both apoB-containing and HDL particles (Fig. 3b). At three of the loci (*CYP7A1*, *ABCB1* and *SERPINA1*) the ICP-predisposing alleles were associated with higher concentrations of IDL and LDL subclass measures, whereas the direction of the association was reversed for others (*GCKR*, *ABCB11* and *HNF4A*). This information may be useful when considering these genes as therapeutic targets, as targets that adversely influence atherosclerotic lipids in pregnant women may be undesirable, despite the relatively short treatment period. By characterizing the associations of ICP-associated loci with metabolic traits in detail, we exemplify the value of combining the metabolic association information with disease associations to clarify the metabolic underpinnings of poorly understood conditions.

## Mendelian randomization

Finally, we exploited the absence of UK Biobank from our GWAS meta-analysis to perform a two-sample Mendelian randomization analysis to investigate associations of genetically predicted levels of the 20 non-lipid traits with 460 Phecodes and 52 quantitative traits from

the UK Biobank. Initial Mendelian randomization analyses using all lead variants for each trait as genetic instruments identified 503 significant associations ( $P < 4.88 \times 10^{-6}$ ) under the inverse variance-weighted model, including positive associations between glucose and diabetes, creatinine and renal failure, and amino acids with diabetes (Supplementary Tables 15 and 16), all of which represent well-known causal relationships. Less well-characterized relationships included a positive association between genetically predicted lactate levels and benign neoplasm of uterus. This potentially causal association is concordant with a recent GWAS that linked genetic tendency to gain muscle mass with uterine fibroids<sup>51</sup>. We also found an inverse association between genetically predicted circulating glycine levels and blood pressure, which is supported by a strong observational association with hypertension<sup>52</sup> and by genetic data<sup>53</sup>. This finding suggests a potential mediator for the previously reported inverse association of glycine levels with myocardial infarction<sup>54</sup>. These examples highlight the value of linking data on genetics, metabolic traits and disease outcomes at scale to identify novel causal relationships between metabolic traits and disease.

Restricting the analyses to less pleiotropic variants (associated with fewer than 5 metabolic traits), the association estimates were on average considerably weaker with less between-variant heterogeneity (median absolute beta, 0.058 versus 0.152; Q-statistic, 34.2 versus 385.6, Extended Data Fig. 6), suggesting that pleiotropy was driving many of the initial Mendelian randomization associations. Results using two alternative thresholds for variant pleiotropy (fewer than three metabolic trait associations and fewer than seven metabolic traits associations) were very similar (Supplementary Table 17), suggesting that the findings are not sensitive to the choice of threshold. This clearly emphasizes that pleiotropy should be carefully considered when selecting instrument SNPs for Mendelian randomization to avoid false interpretations about potential causal relationships.

As an example, the Mendelian randomization results for acetone were substantially affected by the inclusion of more pleiotropic SNPs in the instrument (Fig. 4). Acetone is a ketone body that is produced primarily in the liver during fasting and which has been associated with several cardiometabolic conditions including heart failure<sup>55</sup> and diabetes<sup>56</sup> in biochemical and epidemiological studies. In the GWAS, we identified

associations for acetone at ten loci (only one associated locus—*APOA5*—was identified in the previous NMR GWAS meta-analysis<sup>4</sup>), and Mendelian randomization yielded 20 robust associations (Fig. 4a). These included associations with triglycerides, HDL cholesterol and remnant cholesterol, probably reflecting the inclusion of well-known lipid loci (*LPL*, *APOA5*, *TRIB1*, *APOC1*, *GALNT2* and *PPPIR3B*) in the instrument. The less pleiotropic instrument for acetone included only four loci: 3-hydroxy-3-methylglutaryl-CoA synthase 2 (*HMGS2*), 3-oxoacid CoA-transferase 1 (*OXCT1*), cytochrome P450 family 2 subfamily E member 1 (*CYP2E1*) and *SLC2A4*, all of which have direct roles in ketone body or glycaemic-related pathways. Using these 4 variants only, the positive association with hypertension (OR per s.d. higher genetically predicted acetone level = 1.41,  $P = 6.9 \times 10^{-7}$ ) was robust (Fig. 4a,b) and was also replicated in FinnGen (OR 1.45,  $P = 4.5 \times 10^{-5}$ ) (Fig. 4c). Consistent with these results, acetone has recently been suggested as a biomarker for hypertension<sup>57</sup>. It should be noted that previous studies using the NMR metabolomics platform had incorrectly labelled acetone as acetoacetate, which was detected and corrected in 2020 and later versions of the platform. The discovery regarding this potential causal relationship between acetone and hypertension is noteworthy, since the data on the role of ketogenic diets in hypertension are suggestive but inconclusive<sup>58,59</sup> and ketone bodies have also emerged as potential therapeutic agents for coronary disease<sup>60</sup>. This finding concurs with preclinical and human studies that link interventions that alter levels of ketone bodies, such as ketogenic diets and ketone salt supplementation, with changes in blood pressure<sup>61,62</sup>, leading to suggestions that ketone bodies could be a promising potential therapeutic strategy for hypertension and other cardiovascular diseases<sup>60,63</sup>. The mechanisms by which ketone bodies influence risk of hypertension are currently unclear, with both indirect (for example, obesity and diabetes) and direct (for example, sympathetic nervous system activity, vasodilation and cardiac endothelial cell proliferation) pathways being suggested<sup>64–66</sup>. A recent study in the UK Biobank demonstrated that some loci and pathways associated with the non-lipid NMR traits are highly pleiotropic, with the less pleiotropic variants often reflecting biology more proximal to the traits<sup>67</sup>. This is also in line with our findings as demonstrated by the identification of several pleiotropic triglyceride-related genes that are associated with acetone levels, as well as four less pleiotropic acetone-associated loci with direct links to pathways related to ketone biology. These results accentuate that genetic pleiotropy can be common for metabolic measures, even for some non-lipid traits, and that careful selection of variants for Mendelian randomization is crucial to avoid bias due to pervasive pleiotropy.

## Limitations

The predominance of participants of European ancestries (27 out of 33 cohorts) meant that we had limited power to detect associations in other ancestry groups. However, our ancestry-stratified comparisons suggested that the associations discovered were broadly transferable across ancestries. Future larger studies of diverse ancestries, including African ancestries, will be required to better understand genetic regulation of metabolism on a global scale. Our NMR-based study was also limited in the number of metabolic traits analysed compared with studies using mass spectrometry, a complementary method that can simultaneously measure thousands of metabolites. Although mass spectrometry is more sensitive, NMR is analytically more robust, high-throughput and low cost, thus our study includes more than sixfold more participants than the largest GWAS of mass spectrometry-based circulating metabolites<sup>68</sup> enabling much deeper characterization of the genetic regulation. Furthermore, mass spectrometry cannot provide the detailed analysis of lipoprotein subclasses that is available from NMR platforms. Another limitation is that although we identified differences in genetic associations according to fasting status and sample type, the mechanisms explaining these

differences remain suggestive and require further investigations. These differences suggest that caution should be used when interpreting heritability estimates across different studies, such as UK Biobank. Furthermore, we have described the detailed metabolic associations of genetic loci associated with ICP, and it should be noted that many of the ICP-associated loci are known to be associated with liver function enzymes or bilirubin, increased levels of which are included in diagnostic criteria for ICP. However, the presence of pruritus (itching) is required for ICP to be diagnosed, and the ICP cases defined through hospital discharge registries included in the GWAS should therefore represent true symptomatic cases.

## Conclusion

Through this large-scale, genome-wide meta-analysis including more than 136,000 participants, we identified more than 8,000 genetic associations of circulating metabolic biomarkers involving over 400 loci. The fivefold increase in sample size and doubling of the number of metabolic traits compared to our previous GWAS meta-analysis of NMR metabolic traits led to a marked increase in the number of significant associations (62 associated loci previously<sup>4</sup>), leading to a substantial improvement in understanding of genetic regulation of systemic metabolism. Key features of our meta-analysis are the inclusion of participants from 33 cohorts, which enables the discovery of many new robust associations with evidence from independent datasets. Through internal comparisons across these datasets and external comparison with UK Biobank, we have highlighted the important role that sample and participant characteristics, such as sample type and fasting status, can have in revealing or masking genetic associations, with significant consequences for biological interpretation and downstream analyses. Our extensive manual curation to identify highly probable causal genes at nearly 300 associated loci provides a useful resource to further biological understanding of the associations and allows high-confidence identification of causal genes for disease associations that colocalize. For the remaining loci, our results provide a starting point for identification of genes that have so far not been known to be involved in metabolic regulation. Our comparison of the fine-grained metabolic associations across the lipoprotein measures enables the identification of clusters of genes with similar metabolic profiles, suggesting *TRIM5* as a potential therapeutic target for lowering pro-atherogenic lipid levels, and therefore cardiovascular diseases, as the metabolic profile of *TRIM5* aligns well with genes that affect LDL cholesterol intake to hepatocytes through the LDL receptor. By making the summary statistics publicly available, we provide a valuable resource for Mendelian randomization studies and have illustrated the potential pitfalls of using pleiotropic variants as genetic instrumental variables. Finally, we have illustrated the potential to use these findings to shed light on inadequately characterized diseases by examining the metabolic effects of genetic variants associated with ICP, a disease with a largely unknown genetic background.

## Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-024-07148-y>.

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## Methods

### NMR metabolomics

In this work, we expand our previous GWAS of 123 human metabolic traits in ~25,000 individuals<sup>4</sup> to include additional cohorts and a more comprehensive panel of metabolic traits. Up to 233 serum or plasma metabolic traits were quantified in 33 cohorts (total sample size up to 136,016) using an updated quantification version of the same NMR metabolomics platform<sup>17</sup> as in the previous study. The NMR metabolomics platform provides data of lipoprotein subclasses and their lipid concentrations and compositions, apoA1 and apoB, cholesterol and triglyceride measures, albumin, various fatty acids and low-molecular-weight metabolites—for example, amino acids, glycolysis-related measures and ketone bodies. In this work, the metabolic traits were quantified in the following cohorts (described in detail in Supplementary Notes and Supplementary Table 1): Avon Longitudinal Study of Parents and Children (ALSPAC), China Kadoorie Biobank (CKB), Estonian Genome Center of University of Tartu Cohort (EGCUT), The Erasmus Rucphen Family study (ERF), European Genetic Database (EUGENDA), FINRISK 1997 (FR97), FINRISK 2007 (FR07, that is, DILGOM), The INTERVAL Bioresource (INTERVAL), CROATIA-Korcula Study (KORCULA), LifeLines-DEEP (LLD), Leiden Longevity Study (LLS), eight subcohorts from the London Life Sciences Prospective Population Study (LOLIPOP), The Metabolic Syndrome in Men study (METSIM), The Netherlands Epidemiology of Obesity Study (NEO), The Netherlands Study of Depression and Anxiety (NESDA), Northern Finland Birth Cohort 1966 (NFBC1966), NFBC1986, The Netherlands Twin Register (NTR), Oxford Biobank (OBB), Orkney Complex Disease Study (ORCADES), PROspective Study of Pravastatin in the Elderly at Risk (PROSPER), three subcohorts from the Rotterdam Study (RS), TwinsUK (TUK), and The Cardiovascular Risk in Young Finns Study (YFS). Most of the cohorts consisted of individuals of European ancestry (six Finnish and 21 non-Finnish), and six cohorts had individuals of Asian ancestry (one Han Chinese and five South Asian). All participants gave informed consent and all studies were approved by the ethical committees of the participating centres.

Detailed description of the NMR method is given in the Supplementary Notes.

### Genome-wide association study

A GWAS was performed for 233 metabolic traits (Supplementary Table 2) in each of 33 cohorts (Supplementary Table 1), leading to inclusion of up to 136,016 individuals with both NMR metabolic trait measurements and genome-wide SNP data available. Pregnant individuals or those using lipid-lowering medication were excluded from the study. SNPs were imputed using the Haplotype Reference Consortium release 1.1 or the 1000 Genomes Project phase 3 release, and GWAS was performed under the additive model separately in each cohort (details in Supplementary Table 3). Before analyses, the metabolic trait distributions were adjusted for age, sex, principal components and relevant study-specific covariates (see Supplementary Table 3), and inverse rank normal transformation of trait residuals was performed. The cohorts were combined in fixed-effect meta-analysis with METAL<sup>69</sup>, and the SNPs were filtered to those present in at least seven cohorts. The NMR metabolic traits are highly correlated and using the Bonferroni correction to account for multiple testing would result in an overconservative threshold for genome-wide significance. We therefore used the number of principal components (28) explaining >95% variation in the metabolic traits defined in the largest cohort, INTERVAL, to correct for multiple testing, and our genome-wide significance threshold was set to  $P < 1.8 \times 10^{-9}$  (standard genome-wide significance level,  $P < 5 \times 10^{-8}$ , divided by 28). After the primary GWAS, fasting- and sample type-stratified analyses were performed for the 233 metabolic traits. In these analyses 26 of the cohorts were classified as fasted ( $n = 68,559$ ), six cohorts were classified as non-fasted

( $n = 58,112$ ), seventeen cohorts were classified as having serum samples ( $n = 90,223$ ) and sixteen cohorts had plasma samples ( $n = 45,793$ ; see Supplementary Table 1). To define associated loci across the metabolic traits, we defined a 500-kb window flanking each SNP meeting the significance threshold, pooled together these windows from all metabolic traits for each chromosome, and iteratively merged the windows. As this approach can lead to inclusion of multiple independent signals within these loci, we further defined potential independent signals that reside within the defined loci based on pairwise LD ( $r^2$  cut-off of 0.3, defined in INTERVAL and FINRISK97) of all the lead SNPs within each locus. Regional association plots were created in LocusZoom, v. 1.4. We assigned the associated lead SNPs to the most likely causal genes based on two criteria: (1) we prioritized genes with clear biological relevance to the associated metabolic traits; and (2) if no biologically plausible causal gene was detected and the lead SNP was a functional variant (missense, splice region or stop gained) or in high LD ( $r^2 > 0.8$  in INTERVAL) with such a variant, the gene with the functional variant was assigned as the most likely candidate gene. If criteria 1 and 2 were not fulfilled, the nearest gene was indicated as the candidate gene.

### Ancestry-specific analyses

We conducted ancestry-stratified analyses within our primary discovery meta-analysis for South Asian (five cohorts, 11,340 participants), East Asian (one cohort, 4,435 participants), all European (27 cohorts, 120,241 participants), Finnish (six cohorts, 27,577 participants) and non-Finnish European (21 cohorts, 92,664 participants) participants. For these ancestry-specific analyses, we used the standard threshold for genome-wide significance ( $P < 5 \times 10^{-8}$ ). To also compare to participants with African ancestry, we conducted an African-specific subgroup analysis using the UK Biobank dataset (March 2021 release). Using self-reported ethnicity information (Field 21000: Ethnicity background) from the baseline questionnaire, 1,405 participants with African ancestry were identified as having Caribbean (code 4001), African (code 4002), or any other Black background (code 4003). Variant QC was performed by excluding SNPs with minor allele frequency <1%, INFO score <0.4, and variants in complex LD regions. LD thinning was performed with  $r^2 < 0.1$ , a window size of 1,000 and a step size of 80. Related individuals were identified and excluded using relatedness data provided by the UK Biobank (Field 22021: Genetic kinship to other participants). Outliers of the first 6 genetic principal components computed on the unrelated samples were removed from the analysis. NMR metabolic traits were adjusted for age, sex, fasting status and 10 genetic principal components, and trait residuals were inverse rank normal-transformed. Associations between SNPs and metabolic traits were tested using PLINK 2.0.

### Replication in publicly available data

UK Biobank SNP–metabolic trait summary statistics were downloaded ([https://gwas.mrcieu.ac.uk/datasets/?gwas\\_id\\_icontains=met-d](https://gwas.mrcieu.ac.uk/datasets/?gwas_id_icontains=met-d)) from the IEU Open GWAS Project<sup>70</sup>. These summary statistics were derived from the publicly available March 2021 release of the UK Biobank data in which the metabolic traits were measured with a similar NMR technology (newer version of the Nightingale Health platform) as in our study. The data were used to compare the association of our lead SNP–metabolic trait pairs within the 276 associated regions. Two thresholds were used to define an association in the UK Biobank data: the standard genome-wide significance level ( $P < 5 \times 10^{-8}$ ) and the suggestive level of significance ( $p < 1 \times 10^{-5}$ ).

### Heritability and variance explained

We used GCTA-GREML<sup>71</sup> v. 1.94 to estimate common variant heritability for each trait using an independent dataset, specifically the UK Biobank phase 1 NMR release. This research was conducted using the UKBB Resource under application number 7439. We randomly selected

10,000 unrelated UK Biobank participants of European ancestry with available NMR data and filtered imputed variants to minor allele frequency >0.005, missingness <0.1 and Hardy–Weinberg equilibrium  $P$  value <10<sup>−6</sup>. We removed technical variation from the traits using methods described previously<sup>72</sup>, and adjusted the traits for age, sex, lipid-lowering medication usage and the first 10 genetic principal components of ancestry. Traits were rank inverse normal-transformed prior to GREML analysis. Variance explained by the lead SNPs for each trait was estimated as described before<sup>73</sup>.

### Comparing to previous associations

We performed an extensive comparison of our metabolic trait associations to previous GWASs of metabolic traits. Our comparisons were divided into three groups: (1) comparison to results of previously published large GWAS of circulating NMR traits<sup>4,5</sup>; (2) comparison with loci associated with clinical lipids (including those from the UK Biobank September 2019 version 3 release)<sup>21,25–27,74</sup>; and (3) comparison with an extensive list of associations from previous metabolite and metabolomic studies<sup>11,13,53,75–87</sup>. The comparisons were performed by indicating: (1) co-located known variants; (2) any known associations within a 500-kb flank of a lead SNP; or (3) known associations in LD ( $r^2 > 0.3$ , defined in INTERVAL) with a lead SNP. Since we used the UK Biobank for replication, we did not compare the associations to those from studies that used UK Biobank NMR metabolomics as a single cohort without validation cohorts<sup>67,88</sup>.

In addition to comparing to previous metabolic trait associations, we screened previous disease and trait associations ( $P$  value cut-off  $5 \times 10^{-8}$ ) of the lead SNPs using PhenoScanner, v2<sup>44,45</sup>, and NHGRI-EBI GWAS Catalog<sup>46</sup> (associations downloaded on 30 March 2023 using the gwasrapidd R package, v. 0.99.14<sup>89</sup>). In addition, we screened the FinnGen<sup>43</sup> data freeze 7 summary statistics of 3,095 disease endpoints for overlapping associations ( $P$  value cut-off  $5 \times 10^{-8}$ ). Associations with gene expression and protein levels were screened using PhenoScanner, v2<sup>44,45</sup>.

### Metabolic effects of lipoprotein loci

To compare the metabolic effects of lipoprotein, lipid and apolipoprotein-associated variants, the effect estimates were visualized as colour-coded heat maps. To allow comparison of SNP effects, the estimates were scaled relative to the highest absolute value of the estimate for each SNP. In this analysis, we included lead SNPs at the 276 initially defined regions that were associated with any of the lipoprotein lipids or apolipoproteins at genome-wide significance and nominally associated ( $P < 0.05$ ) with apoB. We used these criteria to restrict the analysis to SNPs associated with apoB, because apoB is known to be a causal part of lipoprotein metabolism for cardiovascular disease<sup>30–32</sup>. To exclude signals with similar effects across the metabolic traits due to the same causal gene, we included only a single SNP from the initially defined genomic regions that had multiple independent signals if the patterns of metabolic traits associations were similar ( $R > 0.5$ ). In the heat maps each line represents a single SNP, each column corresponds to a single metabolic measure, and the scaled effect estimates for the SNP-metabolite associations are visualized with a colour range. Directions of effects are shown in relation to the allele associated with increased apoB. To group SNPs with similar effects together, dendrograms were constructed based on hierarchical clustering of the scaled SNP effects. Heat maps were constructed using the heatmap.2 function of the gplots v. 3.0.3 R package. Pearson correlations were assessed in R, v. 4.0.0.

### Intrahepatic cholestasis of pregnancy

We assessed overlap of our metabolic trait associations with ICP using summary statistics from the FinnGen study<sup>43</sup> data freeze 7 (O15\_ICP; 1,460 cases, 172,286 controls). ICP cases were defined through hospital discharge registry, ICD10 code O26.6 and ICD9 codes 6467A and 6467X.

Using the nearest genes at each associated locus, we performed gene ontology (GO) enrichment analysis to search for enriched biological process and molecular function GO terms<sup>90,91</sup>. We assessed colocalizations of association signals using the hypothesis prioritization for multi-trait colocalization (HyPrColoc) R library, v. 1.0, in which an efficient deterministic Bayesian algorithm is used to detect colocalization across vast numbers of traits simultaneously<sup>92</sup>. We searched for colocalization at single causal variants and shared regional associations. To visualize SNP effects across lipid and lipoprotein traits, heat maps were constructed using the heatmap.2 function of the gplots v. 3.0.3 R package. The following SNPs were included in the heat maps: *GCKR*-rs1260326, *ABCB11*-rs10184673, *ABCB1*-rs17209837, *CYP7A1*-rs9297994, *SERPINA1*-rs28929474 and *HNF4A*-rs1800961. Effects of the metabolic trait-associated SNPs were scaled relative to an odds ratio of 1.5 for ICP.

### Mendelian randomization

Two-sample Mendelian randomization was performed using 20 NMR non-lipid metabolic traits (including amino acids (alanine, glutamine, glycine, histidine, isoleucine, leucine, valine, phenylalanine and tyrosine), ketone bodies (acetate, acetone and 3-hydroxybutyrate), and glycolysis/gluconeogenesis (glucose, lactate, pyruvate, glycerol and citrate), fluid balance (albumin and creatinine) or inflammation-related (glycoprotein acetylation) metabolic traits) as exposures and 460 Phecodes and 52 quantitative traits from the UK Biobank<sup>21</sup> as outcomes. We defined two sets of instruments for the analyses that are referred to as full and strict instruments. As initial instruments we used the 334 lead variants (a single instrument SNP per each defined associated locus) associated with these traits ('full instruments'). To avoid potential bias due to pleiotropy, we also selected a subset of 193 variants ('strict instruments') that had fewer than 5 associations across all 233 metabolic traits. Our threshold of 5 associations was based on empirical assessment of the distribution of per-variant trait associations. To investigate the sensitivity of the Mendelian randomization analyses to the choice of threshold, we also tested using fewer than 3 associations and fewer than 7 associations. We defined disease outcomes in UK Biobank using a curated list of major Phecodes available in the PheWAS R package<sup>93,94</sup>. To restrict our analysis to major disease outcomes, we discarded any sub-categories (that is, Phecodes with 4 or more characters) and removed outcomes with fewer than 100 events across up to 367,542 unrelated UK Biobank participants with European ancestry. The resulting 460 diseases were grouped into 15 broad domains: circulatory system, dermatologic, digestive, endocrine/metabolic, genitourinary, haematopoietic, infectious diseases, mental disorders, musculoskeletal, neoplasms, neurological, pregnancy complications, respiratory, sense organs and symptoms. We also analysed 52 quantitative traits available in UK Biobank, including blood pressure, lung function measures, blood cell traits and clinical chemistry biomarkers. In our replication analysis (acetone as the exposure and hypertension as the outcome), we used essential hypertension from the FinnGen study<sup>43</sup> data freeze 7 as the outcome (hypertension essential, I9\_HYPTESS; 70,651 cases, 223,663 controls). Cases were defined through hospital discharge registry, ICD10 code I10, ICD9 codes 4019X and 4039A, ICD8 codes 40199, 40299, 40399, 40499, 40209, 40100, 40291, 40191 and 40290.

We performed univariable Mendelian randomization using the inverse variance-weighted method for each instrument<sup>95</sup>. We also performed sensitivity analyses using Mendelian randomization–Egger regression to account for unmeasured pleiotropy<sup>96</sup> and weighted median regression to assess robustness to invalid genetic instruments<sup>97</sup>. Our primary analyses were based on fixed-effect models, but as sensitivity analyses we used random-effect models to account for between-variant heterogeneity, which we quantified using the I-squared statistic. The Mendelian randomization analyses were performed using the MendelianRandomization package v. 0.5.1<sup>98</sup> or the TwoSampleMR

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package v. 0.5.3<sup>99</sup>. Single-SNP Mendelian randomization estimates were based on the Wald ratio. We considered the fixed-effects inverse variance-weighted method as the main Mendelian randomization model but report the results of all models in Supplementary Table 15. To account for multiple testing, associations with  $P < 4.88 \times 10^{-6}$  were considered significant (Bonferroni correction to account for testing of 20 metabolic traits with 512 outcomes).

## FinnGen study

In the present study, we used GWAS summary statistics of 3,095 disease endpoints from FinnGen data freeze 7. Full description of the FinnGen study<sup>43</sup> and data analysis steps is provided in the Supplementary Notes. FinnGen contributors are listed in Supplementary Table 18.

## Statistics and reproducibility

The meta-analyses were conducted independently by two investigators in two different centres (University of Oulu, Finland and University of Cambridge, UK), and the summary statistics were compared to verify consistency of results.

## Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

## Data availability

Full GWAS summary statistics are publicly available through the NHGRI-EBI GWAS catalogue (GCST90301941–GCST90302173) and <https://www.phpc.cam.ac.uk/ceu/lipids-and-metabolites/>. Individual-level raw metabolic data from the INTERVAL study can be requested as instructed in <https://www.phpc.cam.ac.uk/ceu/lipids-and-metabolites/>. For access to individual-level genotype and phenotype data for the other studies included in this meta-analysis, please see Supplementary Table 1 for details of websites or references of the individual studies. The NMR metabolomics platform, including the proprietary analysis software, is protected by the intellectual property rights of Nightingale Health, therefore the NMR spectra are not in the possession of the authors and cannot be made publicly available. Source data are provided with this paper.

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**Acknowledgements** Please see the Supplementary Notes for acknowledgements and funding.

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**Competing interests** During the course of the project P.S. became a full-time employee of GlaxoSmithKline. V.S. has received an honorarium from Sanofi for consulting. V.S. also has an ongoing research collaboration with Bayer (outside the present study). A.M. is an employee of Genentech and a holder of Roche stock. N.v.Z. is currently employed by AstraZeneca PLC and is a shareholder in AstraZeneca. R.L.-G. is a part-time contractor of Metabolon Inc. During the course of the project J.Z. became a full-time employee of Novartis. A.I.d.H. is currently an employee of AbbVie. C.M. is funded by the Chronic Disease Research Foundation (CDRF). T.D.S. is co-founder and shareholder of ZOE Ltd. M.I.M. is an employee of Genentech and a holder of Roche stock. J.D. serves on scientific advisory boards for AstraZeneca, Novartis and UK Biobank, and has received multiple grants from academic, charitable and industry sources outside of the submitted work. A.S.B. reports institutional grants outside of this work from AstraZeneca, Bayer, Biogen, BioMarin, Bioverativ, Novartis, Regeneron and Sanofi. The other authors declare no competing interests.

## Additional information

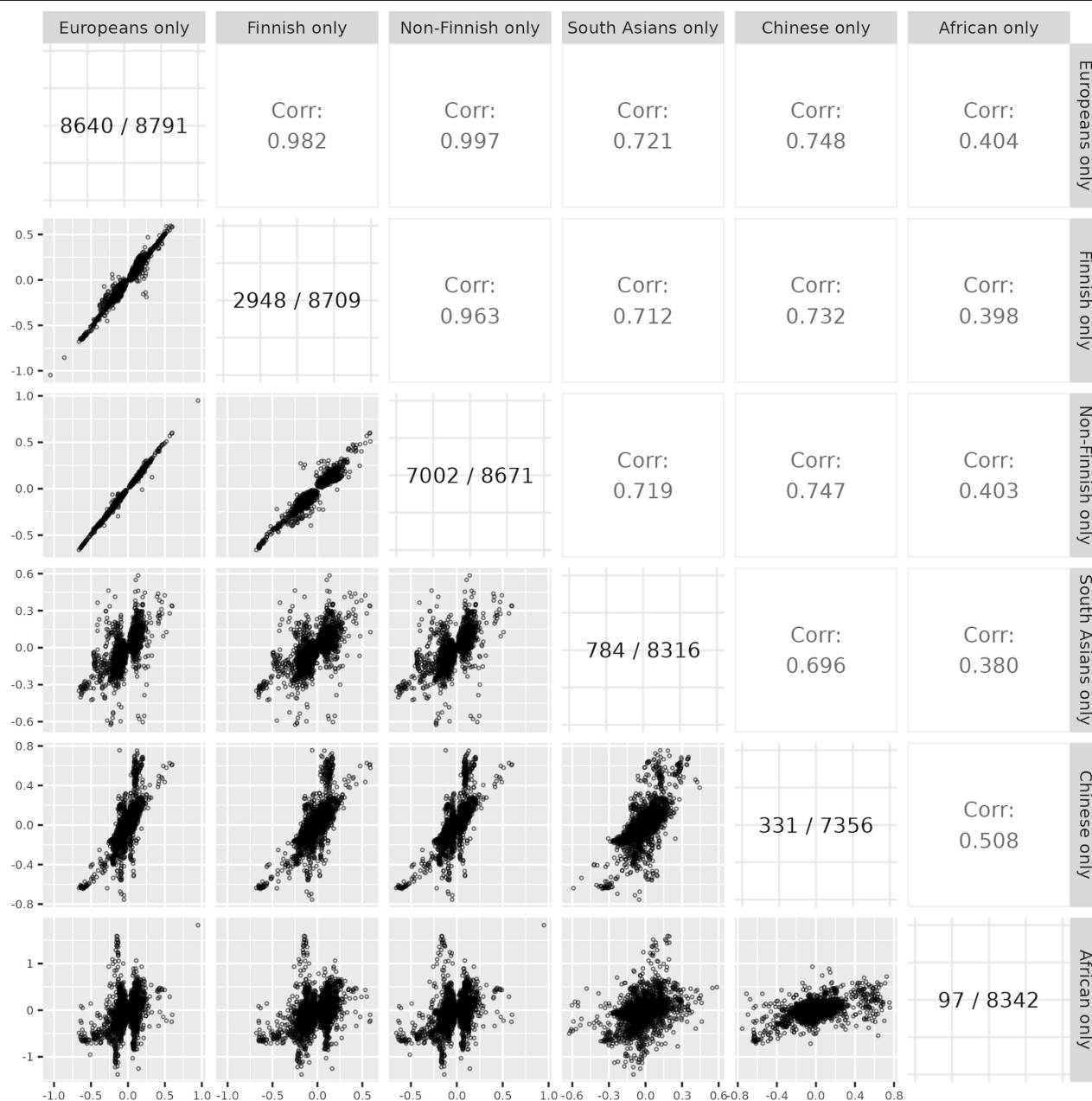
**Supplementary information** The online version contains supplementary material available at <https://doi.org/10.1038/s41586-024-07148-y>.

**Correspondence and requests for materials** should be addressed to Minna K. Karjalainen.

**Peer review information** Nature thanks Themistocles Assimes, Christopher Whelan and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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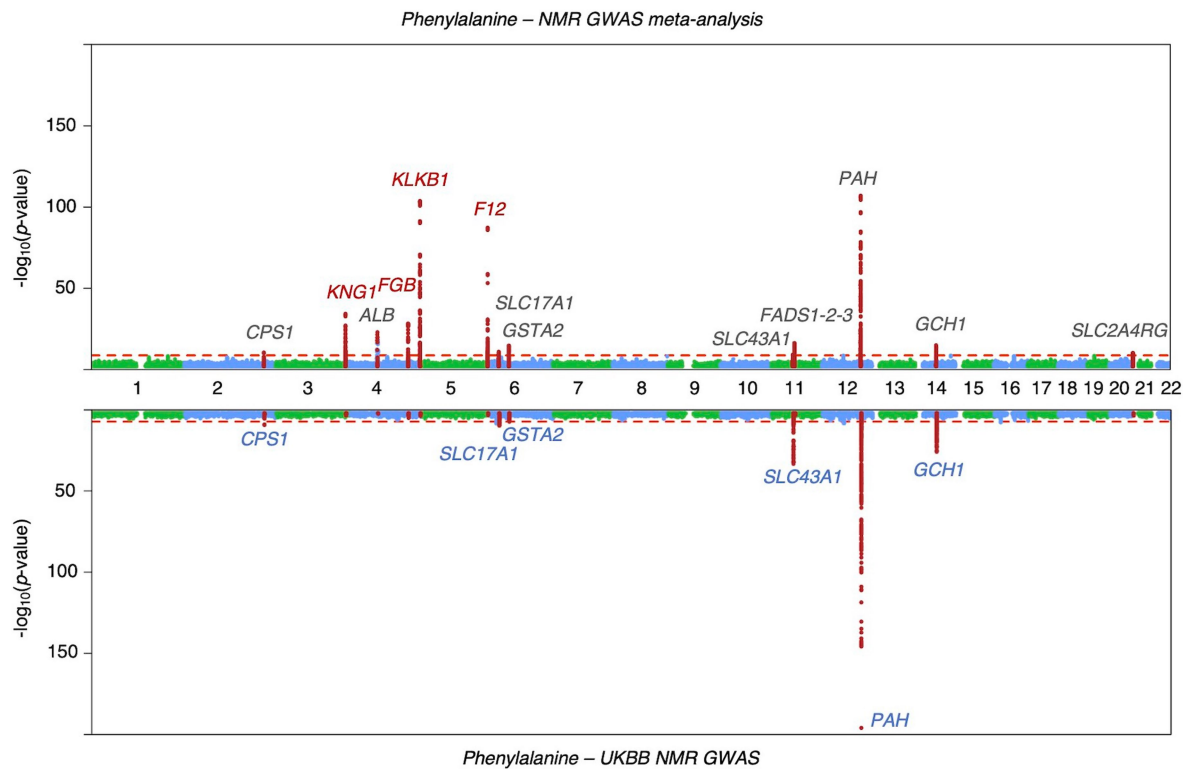




# Extended Data Fig. 1 | Comparisons of effect sizes across ancestries.

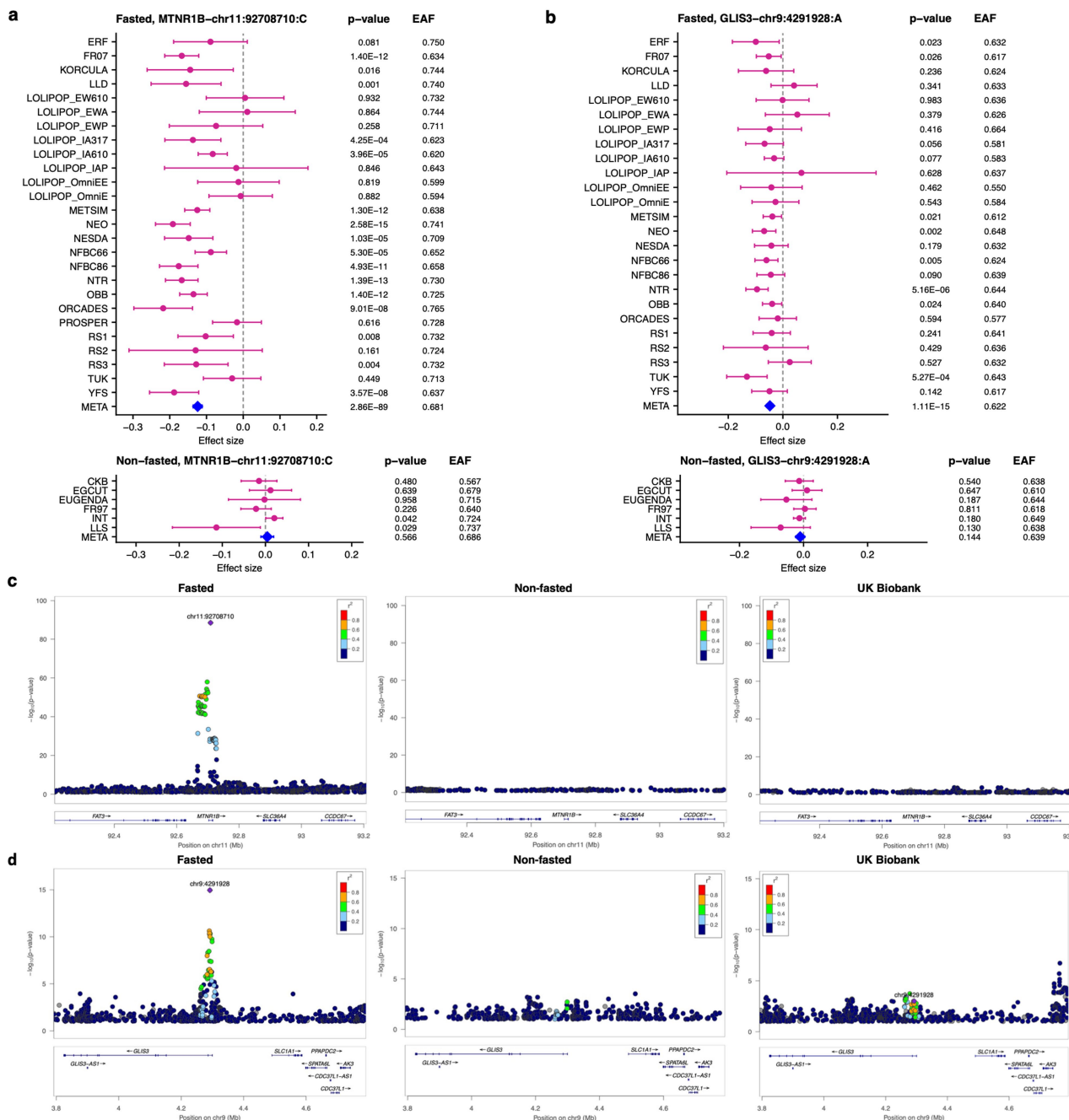
Scatter plots (lower left) show SNP effect sizes of the lead SNP – metabolic trait pairs within the 276 associated genomic regions across different ancestries (Europeans,  $n = 120,251$ ; Finnish,  $n = 27,577$ ; non-Finnish Europeans,  $n = 92,664$ ; South Asians,  $n = 11,340$ ; Han Chinese,  $n = 4,435$ ; Africans,  $n = 1,405$ ). Pearson

correlation (Corr) values ( $r$ ) from each comparison are shown (upper right). The diagonal squares show the number of the 8,795 associations that could be tested in each ancestry group (denominator) and the number that reached the traditional level of genome-wide significance ( $p < 5 \times 10^{-8}$ ) (numerator).



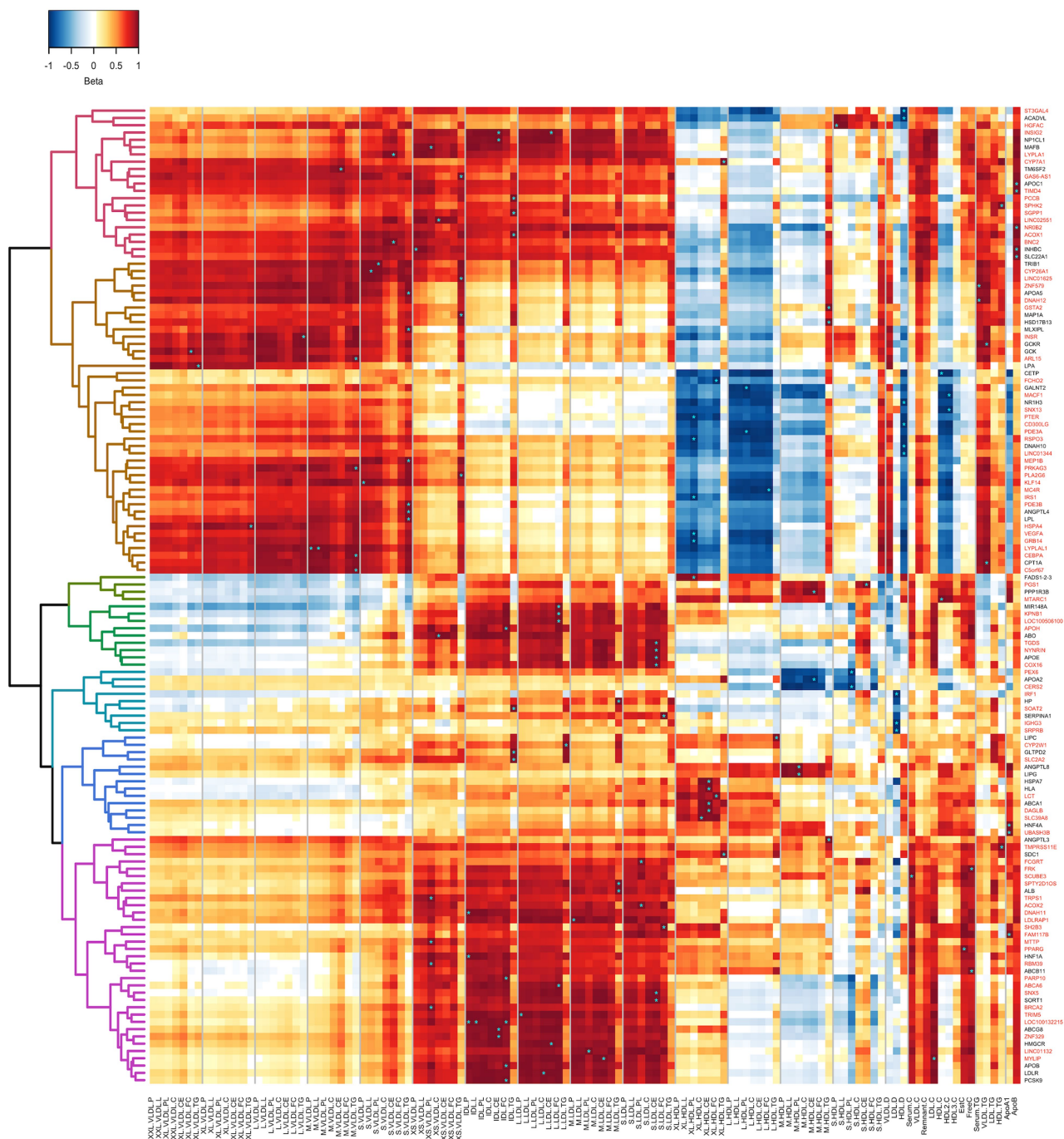
**Extended Data Fig. 2 | Mirrored Manhattan plot showing the results of genome-wide association study of phenylalanine in the NMR GWAS meta-analysis and UK Biobank.** The top panel of the mirrored Manhattan plot shows the NMR inverse variance weighted GWAS meta-analysis results ( $n = 136,016$ ) and the bottom panel the UKBB results ( $n = 115,025$ ). The red lines indicate the

thresholds for genome-wide significance (top panel  $p < 1.8 \times 10^{-9}$ ; bottom panel  $p < 5 \times 10^{-8}$ ). 500-kb regions around lead SNPs in the NMR GWAS are highlighted. Loci indicated in red have roles in coagulation-related pathways. Loci indicated in blue were genome-wide significant in both NMR GWAS meta-analysis and UK Biobank.



**Extended Data Fig. 3 | Examples of glucose associations for fasted and non-fasted cohorts.** The forest plots in panels a and b show examples of two lead SNPs in which glucose associations were significant in the fasted cohorts (top;  $n = 68,559$ ) and non-significant in the non-fasted cohorts (bottom;  $n = 58,112$ ). The associations were analyzed by inverse variance weighted GWAS meta-analysis. These associations were absent in the UK Biobank. Effect sizes (betas and 95% confidence intervals), effect allele frequencies (EAF) and

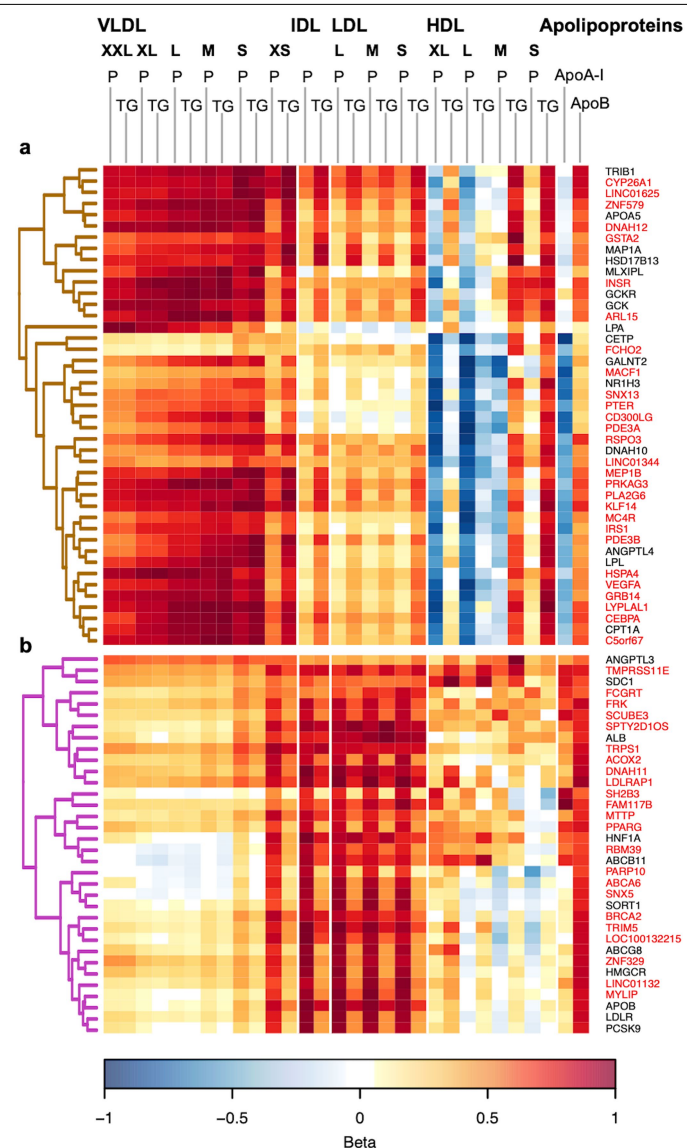
p-values are indicated for each cohort. Cohort acronyms can be found in Supplementary Table 1. Panels c and d show regional association plots of the *MTNR1B* (c) and *GLIS3* (d) loci in the fasted (left) and non-fasted (center) cohorts and in UK Biobank (right). SNPs with  $p < 0.1$  are shown. 500-kb flanking regions around each lead SNP are shown. The linkage disequilibrium values ( $r^2$ ) are based on the 1000Genomes European population.



#### Extended Data Fig. 4 | Heat map of lipoprotein and lipid associations.

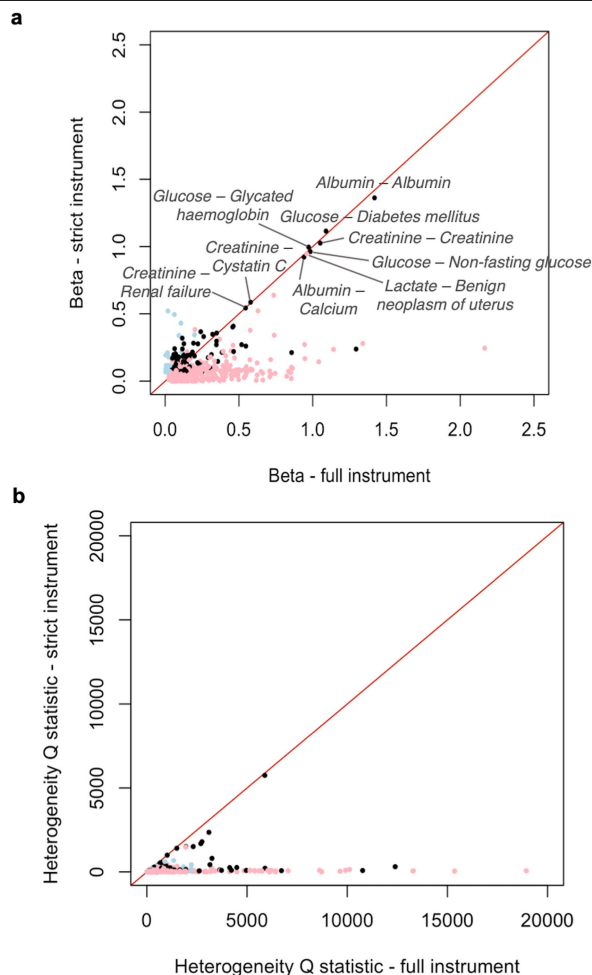
Lipoprotein and lipid associated loci with similar association patterns across the lipoprotein measures were grouped together in a dendrogram based on hierarchical clustering of the SNP effects. The apolipoprotein B associated loci ( $n = 134$ ,  $p < 0.05$ ) were included since apolipoprotein B represents a causal part of the lipoprotein metabolism for cardiovascular disease. The heat map illustrates the resemblances of the association landscapes; each row represents a single SNP, each column corresponds to a single metabolic measure, and the

scaled effect estimates from inverse variance weighted GWAS meta-analysis for the SNP-metabolite associations are visualized with a colour range. Effect sizes were scaled relative to the absolute maximum effect size (beta) in each locus. Loci that were not identified in the previous large-scale NMR metabolomics GWAS are indicated by red. Traits with absolute maximum effects in each locus are indicated by asterisks. For clarity, two of the clusters (brown and purple) are highlighted in Extended Data Fig. 5.



**Extended Data Fig. 5 | A zoomed heat map of lipoprotein and lipid associations.** The full heat map including all the loci and a full set of lipoprotein traits is shown in Extended Data Fig. 4. For clarity, two of the clusters are highlighted here. For details, see legend for Extended Data Fig. 4. Panels a and b corresponding to the brown and purple branches of the dendrogram shown in the full-sized heat map, respectively. Effect sizes were scaled relative to the absolute maximum effect size (beta) in each locus. In the heat map, each row represents a single SNP, each column corresponds to a single metabolic measure, and the effect estimates for the SNP-metabolite associations are visualized with a colour range. Loci highlighted in red were not identified in the previous NMR metabolomics GWAS.





**Extended Data Fig. 6 | Influence of pleiotropy on Mendelian randomization estimates.** The effect estimates (absolute betas) (panel a) and heterogeneity Q statistics (panel b) from the Mendelian randomization (MR) analyses using the full (pleiotropic) and strict (non-pleiotropic) MR instruments are shown. MR was performed using a fixed-effects inverse-variance weighted method. Associations that were significant ( $p < 4.88 \times 10^{-6}$ ) using either the full or strict instrument or both were included; some of the significant exposure-outcome associations are indicated. Estimates indicated in light pink and light blue were not significant with the strict and full instruments, respectively. For clarity, very large beta ( $>5$ ) and Q values ( $>60,000$ ) were excluded from the plots.

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Software and code

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Data collection	The nuclear magnetic resonance spectroscopy (NMR) data were acquired using commercial software by Nightingale Health Ltd., quantification version 2016.
Data analysis	BOLT-LMM v. 2.3.2; GCTA v. 1.91.1 and 1.94; GEMMA 0.97; Plink v. 2.0; RegScan v. 0.5; Rvtest v. 2.0.6; SNPTEST v. 2.5.1, 2.5.2 and 2.5.4; METAL v. 2011; LocusZoom v. 1.4; R v. 3.4.3 and 4.0.0; R packages gplots 3.0.3, HyPrColoc 1.0, MendelianRandomization 0.5.199 and TwoSampleMR 0.5.3.

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## Human research participants

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Reporting on sex and gender	We utilized data from studies that comprised both men and women. We did not perform any sex-specific analyses. Biological sex was used as a covariate in the analyses as indicated in the manuscript.
Population characteristics	In the principal analyses, we utilized data from 33 cohorts/subcohorts. In addition, we used data from two biobanks/biobank-related projects. Details of the participants are summarized in the Supplementary Tables and Supplementary Notes. We used age, sex, population stratification and other relevant study-specific factors as covariates as indicated in the manuscript and Supplementary Tables.
Recruitment	The studies are predominantly population-based cohorts recruited from a variety of settings, including primary care registries, household surveys and blood donors.
Ethics oversight	The studies were approved by appropriate local ethics committees; the committees are indicated in the Supplementary Notes. The following committees approved the studies: Avon Longitudinal Study of Parents and Children: ALSPAC Law and Ethics committee; China Kadoorie Biobank: Oxford Tropical Research Ethics Committee, the Ethical Review Committees of the Chinese Centre for Disease Control and Prevention, Chinese Academy of Medical Sciences, and the Institutional Review Board (IRB) at Peking University; Estonian Genome Center, Institute of Genomics, University of Tartu: Research Ethics Committee of the University of Tartu; Erasmus Rucphen Family study: Medical ethics committee of the Erasmus Medical Center, Rotterdam, the Netherlands; European Genetic Database: Institutional review board in Radboud UMC (Commissie Mensgebonden Onderzoek Radboudum), ethics committees in Cologne and Nijmegen; FINRISK: The Coordinating Ethics Committee of the Helsinki and Uusimaa Hospital District; The INTERVAL Bioresource: the National Research Ethics Service Committee East of England - Cambridge East; CROATIA-Korcula: Ethics committees of the Medical School of the University of Zagreb, the Medical School of the University of Split and the National Health Service, Lothian, Scotland; LifeLines-DEEP: The University Medical Center Groningen review board; Leiden Longevity Study: Ethical committee of the Leiden University Medical Center; London Life Sciences Prospective Population Study: National Research Ethics Service; The Metabolic Syndrome in Men study: Ethics Committee of the University of Eastern Finland and Kuopio University Hospital in Kuopio, Finland; The Netherlands Study of Depression and Anxiety: Ethical review boards of the participating research centers in Amsterdam, Leiden, and Groningen; Northern Finland Birth Cohort: Northern Ostrobothnia Hospital District Ethical Committee; The Netherlands Twin Register: Central Ethics Committee on Research Involving Human Subjects of the VU University Medical Center, Amsterdam; Oxford Biobank: South Central - Oxford C Research Ethics Committee; Orkney Complex Disease Study: Research Ethics Committees in Orkney, Aberdeen (North of Scotland REC), and South East Scotland REC, NHS Lothian; PROspective Study of Pravastatin in the Elderly at Risk: The institutional ethics review boards of centers of Cork University (Ireland), Glasgow University, Scotland, and Leiden University Medical Center, the Netherlands; Rotterdam Study: Medical Ethics Committee of the Erasmus MC and the Dutch Ministry of Health, Welfare and Sport; TwinsUK: St. Thomas' Hospital Research Ethics Committee; The Cardiovascular Risk in Young Finns Study: Ethics Committee of the wellbeing services county of Southwest Finland.

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Sample size	All participants with available genotype and metabolite data were included. Total sample size was up to 136,016.
Data exclusions	The quality control steps including participant/data exclusions are described in the manuscript. Pregnant individuals, statin users and data outliers were excluded in relevant cohorts. Genetic variants that did not meet quality control thresholds were excluded.
Replication	This is a meta-analysis, and as such includes internal replication. Replication of the findings was further investigated in the UK Biobank for those genetic variants and metabolic traits that were available in the UK Biobank data (8,502 of 8,795 lead SNP - metabolic trait pairs).

Randomization

This is a genome-wide association study and thus the randomization was due to genetic variants.

Blinding

This was not a clinical trial so there was no requirement for blinding. Due to the sheer size of the genetic data, the investigators were blinded to the genotype group allocation.

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