GENERAL ARTICLE

Blood copper and risk of cardiometabolic diseases: a Mendelian randomization study

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Abstract

Observational evidence links higher blood levels of copper with higher risk of cardiovascular diseases. However, whether those associations reflect causal links or can be attributed to confounding is still not fully clear. We investigated causal effects of copper on the risk of cardiometabolic endpoints (stroke, coronary artery disease [CAD] and type 2 diabetes) and cardiometabolic risk factors in two-sample Mendelian randomization (MR) studies. The selection of genetic instruments for blood copper levels relied on meta-analysis of genome-wide association studies in three independent studies (European Prospective Investigation into Cancer and Nutrition-Potsdam study, Prospective investigation of the Vasculature in Uppsala Seniors study, Queensland Institute of Medical Research studies). For the selected instruments, outcome associations were drawn from large public genetic consortia on the respective disease endpoints (MEGASTROKE, Cardiogram, DIAGRAM) and cardiometabolic risk factors. MR results indicate an inverse association for genetically higher copper levels with risk of CAD (odds ratio [95% confidence interval] = 0.92 [0.86–0.99], P = 0.022) and systolic blood pressure (beta [standard error (SE)] = −0.238 [0.121]; P = 0.049). Multivariable MR incorporating copper and systolic blood pressure into one model suggested systolic blood pressure as mediating factor between copper and CAD risk. In contrast to previous observational evidence
establishing higher blood copper levels as risk-increasing factor for cardiometabolic diseases, this study suggests that higher levels of genetically predicted copper might play a protective role for the development of CAD and systolic blood pressure.

**Graphical Abstract**

![Graphical abstract](image)

**Introduction**

Blood levels of copper have been associated with higher risk of chronic diseases such as cardiovascular disease (CVD) and stroke in several observational studies (1,2). Although serum concentrations of copper are strictly regulated by compensatory mechanisms within certain ranges of nutritional intake, under inflammatory conditions, aging or age-related disorders inducing oxidative stress, serum concentrations of copper are increased (3,4). However, it is still not fully clear whether these changes might causally impact disease risk.

To investigate causal associations between risk factors and disease outcomes, genetic variants can be used as instrumental variables in Mendelian randomization (MR). The principle is based on Mendel’s second law of inheritance, stating that the assignment of alleles is random during meiosis. Therefore, genetic variants are generally not associated with possible confounders (e.g. environmental exposures or lifestyle factors) in exposure-disease associations (5). A previous MR study found an inverse association between higher blood copper levels and risk of ischemic heart disease (IHD) (odds ratio [OR]: 0.94; 95% confidence interval [CI]: 0.90–0.98, \( P = 0.022 \); Tables 2, Supplementary Material, Table S3). However, there were no MR studies on other cardiometabolic disease outcomes. Therefore, we first aimed to use human genetics to study the associations of blood copper on the risk of type 2 diabetes, coronary artery disease (CAD) and stroke. Second, we investigated related risk factors that may mediate associations between copper and disease endpoints.

**Results**

**Selection of genetic instruments by meta-analysis of GWAS in EPIC-Potsdam, PIVUS and QIMR**

We conducted genome-wide association study (GWAS) on serum copper levels within European Prospective Investigation into Cancer and Nutrition (EPIC)-Potsdam data (Supplementary Material, Fig. S1) and meta-analyzed these data with published GWAS datasets on blood copper levels (7,8) (Supplementary Material, Fig. S2). We identified two genome-wide significant hits for blood copper levels. One was located in an intergenic region near selenium binding protein 1 (SELENBP1; lead single nucleotide polymorphism [SNP]: rs17564336). The other was located in the intronic region of CP (rs34951015) (Table 1, Supplementary Material, Table S1). Both instruments had F-statistics higher than 10. A complete list of genome-wide significant SNPs can be found in Supplementary Material, Table S2.

**Causal estimates for copper on risk of chronic diseases and cardiometabolic risk factors**

We could not find indication for a causal effect of copper levels on risk of type 2 diabetes or stroke (Tables 2, Supplementary Material, Table S3 and S4, Supplementary Material, Fig. S3). However, we found a significant inverse association for copper and CAD (beta \( \beta = -0.079 \) [0.035], OR [95% CI] =0.92 [0.86–0.99], \( P = 0.022 \); Tables 2, Supplementary Material, Table S4, Supplementary Material, Fig. S3).

In MR analyses of copper and cardiometabolic risk factors, we found indication for an inverse association with systolic blood pressure (beta \( \beta = -0.238 \) [0.121], \( P = 0.049 \); Tables 2, Supplementary Material, Table S3 and S4, Supplementary Material, Fig. S3).

Multivariable MR indicated a direct causal effect of systolic blood pressure on CAD (beta \( \beta = 0.034 \) [0.003], OR [95% CI] =1.03 [1.03–1.04], \( P = 1.81 \times 10^{-10} \)), whereas for copper no significant association was observed after adjusting for systolic blood pressure (beta \( \beta = -0.019 \) [0.027], OR [95% CI] =0.98 [0.93–1.04], \( P = 0.4935 \). (Fig. 1, Supplementary Material, Table S5) There was indication for weak instruments for copper with conditional F-statistic of 1.18. When we corrected for weak instruments in the multivariable MR, we received comparable estimates for systolic blood pressure (beta [95% CI] =0.034 [0.027–0.041], OR [95% CI] =1.04 [1.03–1.04] and copper (beta [95% CI] =-0.028 [−0.091–0.025], OR [95% CI] =0.97 [0.91–1.03]).

**Discussion**

Our two-sample MR approach indicated an inverse association of genetically higher copper on risk of CAD. Similarly, we obtained an inverse association for genetically higher copper on systolic blood pressure. Multivariable MR suggested systolic blood pressure as mediating factor between copper and CAD risk.
Copper and cardiometabolic risk factors and diseases

Prospective studies on copper levels and cardiometabolic outcomes are scarce. In EPIC-Potsdam, we found a positive association between serum copper and risk of CVD and type 2 diabetes in an observational setting, the latter, however, not significant. In multivariable MR, we found a direct association between copper and risk of myocardial infarction and higher risk of arterial fibrillation (15) in MR approach. However, Cu levels of copper may be a consequence of increased transcription of the Cu/Cp gene region, suggesting that besides copper itself, also its transport protein ceruloplasmin (Cp) with high affinity. Therefore, increased blood levels of copper may be a consequence of increased transcription of Cp in liver tissues (3). One of our genetic instruments (rs34951015) is located in the Cp encoding gene region, suggesting that besides copper itself, also its transport protein might be involved in biological pathways linking copper to cardiometabolic risk. The SNP represents a novel hit for systolic blood pressure on CAD risk, attenuating the association between copper and CAD risk.

Biological mechanisms linking copper to CAD and systolic blood pressure

In the circulation, 80–95% of total blood copper is bound to ceruloplasmin (Cp) with high affinity. Therefore, increased blood levels of copper may be a consequence of increased transcription of Cp in liver tissues (3). One of our genetic instruments (rs34951015) is located in the Cp encoding CP gene region, suggesting that besides copper itself, also its transport protein might be involved in biological pathways linking copper to cardiometabolic risk. The SNP represents a novel hit for blood copper levels and is in LD to a potentially functional CP variant in the promoter region upstream of the CP gene, previously associated with Cp levels (rs11708215: $r^2 = 0.35, D' = 0.99$ (11)) (15). Cp was associated with increased risk for myocardial infarction (16), CVD mortality (10,17) and stroke (17) in observational setting and higher risk of arterial fibrillation (15) in MR approach. Based on GTEx data, CP is mainly expressed in liver, artery-aorta and artery-tibial tissues (18). Our CP instrument rs34951015 represents an eQTL for CP in artery-tibia ($P = 1.3 \times 10^{-10}$) and artery-aorta ($P = 0.007$) tissues (18).

### Table 1. Independent GWAS hits for copper at genome-wide significance ($< 5 \times 10^{-8}$) from Meta-analysis of EPIC-Potsdam, PIVUS and QIMR

| SNP          | Chr/bp   | Consequence/ Gene | CADDa | Effect allele | Other allele | N     | EAF  | P-value       | Direction  | F-statistic (56) |
|--------------|----------|-------------------|-------|---------------|--------------|-------|------|---------------|-------------|-----------------|
| rs17564336   | 1:151349174 | Intergenic/ near SELENBP1 | 1.751 | T             | G            | 6937  | 0.298| $1.79 \times 10^{-17}$ | --         | 73.54           |
| rs34951015   | 3:148926315 | Intronic/CP       | 0.419 | A             | G            | 6937  | 0.099| $9.57 \times 10^{-10}$ | +++        | 37.81           |

CADD, combined annotation dependent depletion; EAF, effect allele frequency.
aExtracted from Ensembl database (11).
bFrom three different studies: EPIC-Potsdam, PIVUS and QIMR.

### Table 2. Causal estimates for copper and cardiometabolic disease endpoints and related risk factors from MR analysis

| Outcome                  | IVWa       | Cochran’s Q statistic (P-value) |
|--------------------------|------------|---------------------------------|
| Type 2 diabetes          | 0.028 (0.026), 0.275 | 2.814 (0.093)                  |
| Coronary artery disease  | −0.079 (0.035), 0.022 | 2.215 (0.137)                  |
| Stroke                   | 0.034 (0.038), 0.366 | 2.233 (0.135)                  |
| BMI                      | 0.009 (0.008), 0.238 | 11.794 (0.001)                 |
| LDL                      | 0.003 (0.009), 0.729 | 2.221 (0.136)                  |
| HDL                      | −0.001 (0.009), 0.928 | 2.537 (0.111)                  |
| Systolic blood pressure  | −0.238 (0.121), 0.049 | 0.538 (0.463)                  |
| Diastolic blood pressure | −0.075 (0.069), 0.280 | 0.044 (0.833)                  |
| HbA1c                    | −0.007 (0.021), 0.728 | −                  |
| Blood glucose            | 0.004 (0.010), 0.661 | 0.006 (0.936)                  |

aValues are beta (SE); P-values are from fixed effects models.
bWald ratios as only one SNP (rs17564336) was available in the outcome dataset.
Besides selenium, SELENBP1 may also bind copper. Extracellular SELENBP1 was identified in blood after myocardial infarction or during cardiac surgery. Our instruments show furthermore associations with red blood cell traits (rs17564336) and histidine (rs34951015), that may convey vertical pleiotropic pathways on the way to CAD development. Indeed, we found inverse associations between copper and systolic blood pressure which might represent a mediating pathway between copper levels and the reduction of CAD risk. Copper is an essential trace element acting as co-factor for several copper-dependent enzymes like lysyl oxidase and copper-zinc superoxide dismutase that convey important biological functions like cross-linking of collagen and elastin in blood vessels and protection against free radicals, respectively. Hence, copper-deficiency has been reviewed in the context of development of CVD, dysregulation of microcirculation and impaired lipid metabolism. Therefore, our finding of an inverse association of higher blood copper with systolic blood pressure and CAD risk fits with the detrimental effects of copper deficiency.

**Strengths and limitations**

We used genetic instruments that showed reliable associations in at least two of the three GWAS datasets and showed sufficient strength (>10) with F-statistics of 73.5 for rs17564336 and 37.8 for rs34951015 (relevance assumption). However, as we had only two instruments for the analysis, sensitivity analyses such as MR-Egger that can be used to assess directional horizontal pleiotropy could not be applied. Hence, we cannot exclude potential horizontal pleiotropy by other pathways that were not considered in the MR analyses (exclusion restriction assumption). Database look-ups did not reveal genome-wide significant associations between the two instruments and sociodemographic or lifestyle factors that might confound the copper outcome association (independence assumption).

By pooling three different GWAS on copper levels involving up to 6937 individuals to select genetic instruments, we generated the largest genetic data source for blood copper values to date. However, different sample materials were used for copper measurements (serum, whole blood, erythrocytes) and varying transformations of copper values and model adjustments were conducted in the individual GWAS. We tried to overcome limitations due to different analytical approaches by applying the Stouffer Method for meta-analysis, which is robust to different effect estimate scales as it simply requires P-value, sample size and estimated direction of effect for each study. However, due to the use of resulting pooled z-scores, neither the beta estimates for associations with copper nor the effect sizes from the MR analysis have interpretable units. Still, the results are of value in terms of effect direction.

Moreover, as the liver is the main homeostatic organ for copper and liver content is not accurately reflected by copper levels in plasma or urine, blood levels of copper might only weakly correlate with available copper in the organism. Similarly, we cannot use biopsies to determine liver copper content in epidemiological research. In addition, we quantified the total copper content in the different blood fractions and we cannot distinguish between the amount that is bound to Cp and the loosely bound or free copper content.

Our study might be underpowered to detect causal effects for other cardiometabolic risk factors besides systolic blood pressure, where we had a power of 71% (33). Furthermore, we tested multiple phenotypes and the associations for CAD and systolic blood pressure were only nominally significant (<0.05). Using a strict cut-off for multiple testing of 0.017 (0.05/3) or 0.007 (0.05/7) would result in no significant association for disease endpoints or related risk factors, respectively. Although those findings may represent false positives, the result of copper and CAD is in accordance with the previous MR study (6), reinforcing each other. Furthermore, we used the advantage of multivariable MR to study copper and systolic blood pressure in one model. However, we recognized an imbalance of suitable genetic instruments between both exposures of one for copper versus 415 for systolic blood pressure, resulting in a low conditional F-statistic for copper. Although we corrected for weak instruments and received comparable estimates, this analysis should be interpreted cautiously. Furthermore, the outcome summary statistics for blood pressure traits are adjusted for BMI, which might have biased the results due to unmeasured confounding (34). However, these summary statistics represent the largest datasets for blood pressure.
pressure traits (35) and are not available without BMI adjustment, hence the relationship between copper, blood pressure and CAD needs further investigation. Finally, our findings are restricted to mainly Europeans, and therefore, cannot be generalized to other ethnicities. For CAD, also Asian individuals contributed to the meta-analysis (36), which might be considered as limitation, as individuals included in two-sample MR analyses should come from the same ethnic background. Nevertheless, this should not increase the likelihood of finding an association when there is none (37). Hence, our results on copper and CAD might still be informative in terms of effect direction of causality (38).

**Conclusion**

In contrast to previous observational evidence establishing higher blood copper levels as risk-increasing factor for cardiometabolic diseases, this study suggests that genetically higher blood copper might play a protective role for CAD development and systolic blood pressure. Whether systolic blood pressure mediates the association between copper and CAD needs further investigation.

**Materials and Methods**

**Study populations**

**EPIC-Potsdam study.** The EPIC-Potsdam study consists of 27 548 participants recruited between 1994 and 1998 from the general population in Potsdam and surroundings, Germany. The baseline examination involved a personal interview, interviewer-conducted anthropometric measurements and a blood sample collection (39). We used a random sample within the EPIC-Potsdam study, described in detail previously (40). Briefly, a sub-cohort of 2500 individuals was randomly selected from 26 444 participants who provided blood samples at baseline. Of these, participants with prevalent diabetes, myocardial infarction or stroke were excluded. Further exclusion criteria were missing genetic data and missing data on copper measurements, leaving 2045 individuals for analyses in the sub-cohort (Supplementary Material, Fig. S4).

All participants provided written informed consent. The EPIC-Potsdam study was approved by the ethics committee of the State of Brandenburg, Germany. All procedures were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

**PIVUS study.** The PIVUS study aims to investigate the predictive power of different measurements of vascular characteristics for future cardiovascular events (41). Briefly, individuals at the age of 70 years living in the community of Uppsala, Sweden were randomly selected from the community register and invited to participate in the study. Of 2025 invited, 1016 participants were recruited. For the present analysis, GWAS data of 946 individuals were used. The PIVUS study was approved by the Ethical Committee of Uppsala University and each participant gave written informed consent.

**QIMR studies.** Evans et al. investigated the association between genetic variants and copper levels from erythrocytes in adult twins from different studies conducted at the QIMR, Australia (7). For the present analysis, we utilized updated GWAS data from 3946 individuals. Participants in the QIMR studies gave written informed consent, and the studies were approved by the Human Research Ethics Committee of the QIMR.

**Summary-level data for cardiometabolic outcomes.** All data resources for cardiometabolic outcomes and cardiometabolic risk factors include individuals with European ancestry and are summarized in Supplementary Material, Table S6. In brief, genetic associations with diabetes were obtained from the D1Abetes Genetics Replication And Meta-analysis (DIAGRAM) consortium including 74 124 T2D cases and 824 006 controls without BMI adjustment (48.0% women; mean age 59.4 years) (42). Genetic associations with CAD came from the meta-analysis of UK Biobank SOFT CAD GWAS with Coronary ARtery Disease Genome wide Replication and Meta-analysis (CARDioGRAM) plus The Coronary Artery Disease (CARD) Genetics (CARDioGRAMplusC4D) 1000 Genomes-based GWAS and the Myocardial Infarction Genetics and CARDioGRAM Exome (36). The SOFT CAD phenotype incorporates self-reported angina or other evidence of chronic coronary heart disease and 10 801 cases and 137 371 controls (36).

Associations with stroke were obtained from MEGASTROKE (cases = 40 585, controls = 406 111) (43). For the risk factors BMI, high-density lipoprotein (HDL), LDL and blood glucose, we used UK Biobank data (up to 361 194 individuals; 461 460 for BMI) (44). For blood pressure phenotypes, we used GWAS data from the International Consortium of Blood Pressure (mean age ∼ 54.5 years, ~51% women), UK Biobank and UK BiLEVE (mean age: 56.7 years, 54% women) with up to 757 601 individuals (35). For HbA1c, we used data provided by the MAGIC consortium including 46 368 non-diabetic individuals (52% women) (45).

**DNA-extraction, genotyping and quality control within EPIC-Potsdam, PIVUS and QIMR.** In EPIC-Potsdam, DNA was extracted from buffy coats using the chemagic DNA Buffy Coat Kit on a Chemagic Magnetic Separation Module I (PerkinElmer chemagen Technologies, Baesweiler, Germany) according to the manufacturer’s manual. Samples were genotyped by three different genotyping arrays: Human660W-Quad_v1_A (n = 363), HumanCoreExome-12v1-0_B (n = 675, two datasets) and Illumina InfiniumOmniExpressExome-8v1-3_A DNA Analysis BeadChip (n = 1377). Genotyping and quality control of the Human660W-Quad_v1_A and HumanCoreExome-12v1-0_B chips was described elsewhere (46). Genotyping using the Illumina InfiniumOmniExpressExome-8v1-3_A DNA Analysis BeadChip was performed in the Life and Brain Center in Bonn, Germany. A detailed description of genotyping, quality control and imputation was previously published (47). Briefly, pre-/phasing and imputation were conducted using Eagle2 (48) and the Michigan Imputation Service (49) with The Haplotype Reference Consortium (release 1.1) as reference panel (50). Imputation was conducted in four separate datasets (one for each chip or two for the HumanCoreExome-12v1-0_B chip) using minima3 (49). Imputed files were merged, keeping the minimal R² score from the four files. SNPs were filtered by R² keeping those with values bigger than 0.6. Overall, we had genotype data for n = 2253 samples from the sub-cohort available (Supplementary Material, Fig. S1).

Genotyping of PIVUS samples was previously described (8). Briefly, samples were genotyped with illumina Infinium Omni Express bead microarray and imputed according to the Haplo-type Reference Consortium reference panel.
QMR genotypes were determined using Illumina 317, 370 or 610 K chips. Quality control steps and imputation of untyped HapMap 2 SNPs was previously described (51). For the present analysis, we used updated genetic data imputed to 1000 Genomes reference panel.

**Blood collection and serum copper measurements in EPIC-Potsdam, PIVUS and QIMR.** In EPIC-Potsdam, the method published by Kopp et al. (52) was employed (4) for trace element profiling. Detailed information can be found in the online supplement. In brief, 50 μl of serum sample was diluted with 440 μl of a diluent solution as described in (52). The analysis was conducted via inductively coupled plasma tandem mass spectrometry (ICP-MS/MS) (Agilent ICP-QQQMS 8800, Agilent Technologies, Waldbronn, Germany).

In PIVUS, copper levels were determined in whole blood. The analysis was performed using inductively coupled plasma sector field mass spectrometry, after microwave-assisted digestion with nitric acid (53).

In QIMR, copper levels were measured in erythrocytes (7). Briefly, the erythrocytes were thawed at room temperature and diluted 1:20 in ammonia/ethylene-diaminetetraacetic acid (EDTA) solution containing rhodium as an internal standard. Copper concentrations were measured by ICP-MS on a PerkinElmer Elan 5000 mass spectrometer (PerkinElmer, Inc., Wellesley, MA, USA) or a Varian UltraMass (Varian Inc., Palo Alto, CA, USA) (7).

**Statistical analysis**

In EPIC-Potsdam, Statistical Analysis System (SAS) Enterprise Guide 7.1 with SAS version 9.4 (SAS Institute Inc., Cary, NC, USA) was used for data management and preparation. QTool v1.4 was used for genetic data filtering. SNP test v2.5.2 (54) was used for GWAS.

For meta-analysis of GWAS results, we used METAL (55). For MR analyses we used R (version 3.6.3 (2020-02-29)) and the TwoSampleMR (v0.5.5) (38) and MVMR (v0.2) (56) R packages.

**Genome-wide association study in EPIC-Potsdam, PIVUS and QIMR.** In EPIC-Potsdam, SNPs were filtered by SNP missing rate (removed ≥0.05), minimum allele frequency (removed out of interval [0.05–0.5]) and Hardy–Weinberg equilibrium (removed -log10(P-value) ≥3). We performed exploratory single variant association analysis using n = 5339559 markers as exposures. We considered a P-value as genome-wide significant at 5 × 10^{-8}. Copper levels were natural log-transformed to normalize the right-skewed distributions and standardized. We assumed an additive genetic model, adjusted for age at recruitment and sex. Variants were mapped to Ensembl annotation version 104 (GRCh37) (11,57).

In the PIVUS study, we performed GWAS by applying additive genetic models using inverse rank normalized copper levels and adjusted for sex, cholesterol, triglycerides and two principal components from multi-dimensional scaling of the genetic relationship matrix to account for population structure (8).

In QIMR data, copper levels were log-transformed and adjusted for analysis batch, hemoglobin concentration in the thawed sample and analytical quality control data to generated standardized residuals for subsequent GWAS (7). GWAS on standardized residuals were adjusted for sex and age, using an additive model accounting for within family relatedness (7).

**Meta-analysis of GWAS results and instrument selection.** We performed meta-analysis of the three different GWAS datasets by applying the Stouffer Method (30) in METAL (55) because different approaches for copper measurement and analysis were used across studies, and effect sizes were therefore not on the same scale. Results were filtered by keeping only SNPs that were available in at least two of the three studies. Based on those, we selected genome-wide significant (P < 5 × 10^{-8}) instrumental variables and performed clumping according to linkage disequilibrium (LD). Therefore, SNPs within a window of 10,000 kb and being in LD as defined by r > 0.001 were removed. The SNPs with the lowest P-values were retained.

**Two-sample MR analyses using on copper levels and cardiometabolic outcomes.** We conducted a univariable two-sample MR study using copper levels as exposures on cardiometabolic diseases and risk factors. Effect estimates of the association between genetic instruments and copper were obtained from z-scores (31) from meta-analysis and effect estimates of the SNP-outcome association were used from public summary GWAS data (Supplementary Material, Table S6) (35,36,42–45). Data were harmonized for the direction of effects between exposure and outcome associations. We used an inverse variance weighted (IVW) meta-analysis of each SNP specific Wald ratio (SNP-outcome estimate divided by SNP-exposure estimate) using fixed effects and random effects models (the latter as sensitivity analysis). Heterogeneity was assessed by the Cochran’s Q statistic. We obtained the F-statistic as indicator of instrument strength (58).

To assess whether systolic blood pressure might mediate the association between copper and CAD, we conducted multivariable MR (59). Genetic instruments for each exposure were selected based on strict genome-wide significance threshold of 5 × 10^{-8} and default clumping threshold of r > 0.001 for a combined list of instruments. Then we obtained all SNP effects on the respective other exposure. The extraction of the outcome GWAS results and the data harmonization was conducted as described before. We calculated conditional F-statistics to evaluate the presence of weak instruments and adjusted for those by minimizing the Q-statistic allowing for heterogeneity using ‘qhet_mvmr’ function from the MVMR package (56). To assess the correlation between systolic blood pressure and copper we used EPIC-Potsdam data. 95% CIs for the effect estimates were determined using non-parametric bootstrap with 1000 iterations.

**Data Availability**

The EPIC-Potsdam datasets analyzed during the current study are not publicly available due to data protection regulations. In accordance with German Federal and State data protection regulations, epidemiological data analyses of EPIC-Potsdam may be initiated upon an informal inquiry addressed to the secretariate of the Human Study Center (Office.HSZ@dife.de). Each request will then have to pass a formal process of application and review by the respective PI and a scientific board.

Summary-statistics of this meta-analysis of GWAS on blood copper values are available through the GWAS catalog.

Summary-level data for genetic associations with type 2 diabetes, CAD and stroke have been contributed by the DIAGRAM consortium (http://diagram-consortium.org/downloads.html; accessed 13.08.2019), the Cardiogram consortium (http://www.cardiogramplusc4d.org/; accessed 12.08.2019) and MEGASTROKE (https://www.megastroke.org/; accessed 05.02.2020). The MEGASTROKE project received funding from sources specified at http://www.megastroke.org/acknowledgments.html. GWAS-Summary data from cardiometabolic risk factors were accessed...
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Supplementary Material

Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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Author contributions

S.J. and M.B.S had access to all data for this study and take responsibility for the manuscript contents. All authors approved the final version to be published.

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