ORIGINAL RESEARCH ARTICLE



Mild-to-Moderate Kidney Dysfunction and Cardiovascular Disease: Observational and Mendelian Randomization Analyses

Liam Gaziano, PhD*; Luanluan Sun, PhD*; Matthew Arnold, PhD*; Steven Bello, PhD; Kelly Cho, MD, MPH; Stephen K. Kaptoge, PhD; Rebecca J. Song, MPH; Stephen Burgess, PhD; Daniel C. Posner, PhD; Katja Mosconi, PhD; Cassianne Robinson-Cohen, PhD; Amy M. Mason PhD; Thomas R. Bolton, PhD; Ran Tao, PhD; Elias Allara, MD, PhD; Petra Schubert, MPH; Lingyan Chen, PhD; James R. Staley, PhD; Natalie Staplin, PhD; PhD; Servet Altay, MD; Pilar Amiano, PharmD, MSc; Volker Arndto, PhD; Johan Ärnlövo, PhD; Elizabeth L.M. Barro, PhD; Cecilia Björkelund, MD; Jolanda M.A. Boer, PhD; Hermann Brenner, MD; Edoardo Casiglia, MD; Paolo Chiodini[®], PhD; Jackie A. Cooper, MSc; Josef Coresh[®], MD, PhD; Mary Cushman[®], MD; Rachel Dankner, MD; Karina W. Davidson, PhD; Renate T. de Jongh, MD, PhD; Chiara Donfrancesco, PhD; Gunnar Engström, MD; Heinz Freisling, PhD; Agustín Gómez de la Cámara[®], PhD; Vilmundur Gudnason[®], MD; Graeme J. Hankey[®], MD; Per-Olof Hansson[®], MD, PhD; Alicia K. Heath[®], PhD; Ewout J. Hoorn[®], MD, PhD; Hironori Imano[®], MD; Simerjot K. Jassal, MD; Rudolf Kaaks, PhD; Verena Katzke, PhD; Jussi Kauhanen, MD; Stefan Kiechl, MD; Wolfgang Koenig[®], MD; Richard A. Kronmal[®], PhD; Cecilie Kyrø[®], PhD; Deborah A. Lawlor, PhD; Börje Ljungberg[®], MD; Conor MacDonald[®], PhD; Giovanna Masala[®], MD; Christa Meisinger, MD; Olle Melander, MD; Conchi Moreno Iribas, PhD; Toshiharu Ninomiya, PhD; Dorothea Nitsch, MD, PhD; Børge G. Nordestgaard, MD. PhD; Charlotte Onland-Moret, PhD; Luigi Palmieri, PhD; Dafina Petrova, PhD; Jose Ramón Quirós Garcia, PhD; Annika Rosengren, MD; Carlotta Sacerdote, MD; Masaru Sakurai, MD; Carmen Santiuste, MD; Matthias B. Schulze, PhD; Sabina Sieri, PhD; Johan Sundström, MD; Valérie Tikhonoff, MD, PhD; Anne Tjønneland, MD; Tammy Tong, PhD; Rosario Tumino, MD; Ioanna Tzoulaki, PhD; Yvonne T. van der Schouw[®], PhD; W.M. Monique Verschuren, PhD; Henry Völzke, MD; Robert B. Wallace, MD; S. Goya Wannamethee, PhD; Elisabete Weiderpass, PhD; Peter Willeit, MD, PhD; Mark Woodward, PhD; PhD; Kazumasa Yamagishi, MD; Raul Zamora-Ros, PhD; Elvis A. Akwo, MD PhD; Saiju Pyarajan, PhD; David R. Gagnon[®], MD, MPH, PhD; Philip S. Tsao[®], PhD; Sumitra Muralidhar, PhD; Todd L. Edwards[®], PhD; Scott M. Damrauer, MD; Jacob Joseph, MD, PhD; Lisa Pennells, PhD; Peter W.F. Wilson, MD; Seamus Harrison, MD, PhD; Thomas A. Gaziano, MD, PhD; Michael Inouye, PhD; Colin Baigent, MD, PhD; Juan P. Casas, MD, PhD; Claudia Langenberg, PhD; Nick Wareham, MD; Elio Riboli, MD; J.Michael Gaziano, MD, MPH+; John Danesh, MD, PhD+; Adriana M. Hung, MD, MPH+; Adam S. Butterworth, PhD+; Angela M. Wood, PhD†; Emanuele Di Angelantonio , MD, PhD†; on behalf of the Emerging Risk Factors Collaboration/EPIC-CVD/Million Veteran Program‡

Correspondence to: Emanuele Di Angelantonio, MD, PhD, Department of Public Health and Primary Care, Heart and Lung Research Institute, University of Cambridge, Papworth Road, Cambridge CB2 0BB, UK; or Health Data Science Centre, Human Technopole, V.le Rita Levi-Montalcini 1, 20157 Milano, Italy. Email ed303@medschl.cam.ac.uk or emanuele.diangelantonio@fht.org

^{*}L. Gaziano, L. Sun, and M. Arnold contributed equally.

t J.M. Gaziano, J. Danesh, A.M. Hung, A.S. Butterworth, A.M. Wood, and E. Di Angelantonio contributed equally.

[‡]A list of Emerging Risk Factors Collaboration Investigators/EPIC-CVD/Million Veteran Program is provided in the Supplemental Material.

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BACKGROUND: End-stage renal disease is associated with a high risk of cardiovascular events. It is unknown, however, whether mild-to-moderate kidney dysfunction is causally related to coronary heart disease (CHD) and stroke.

METHODS: Observational analyses were conducted using individual-level data from 4 population data sources (Emerging Risk Factors Collaboration, EPIC-CVD [European Prospective Investigation into Cancer and Nutrition—Cardiovascular Disease Study], Million Veteran Program, and UK Biobank), comprising 648 135 participants with no history of cardiovascular disease or diabetes at baseline, yielding 42 858 and 15 693 incident CHD and stroke events, respectively, during 6.8 million person-years of follow-up. Using a genetic risk score of 218 variants for estimated glomerular filtration rate (eGFR), we conducted Mendelian randomization analyses involving 413718 participants (25 917 CHD and 8622 strokes) in EPIC-CVD, Million Veteran Program, and UK Biobank.

RESULTS: There were U-shaped observational associations of creatinine-based eGFR with CHD and stroke, with higher risk in participants with eGFR values <60 or >105 mL·min⁻¹·1.73 m⁻², compared with those with eGFR between 60 and 105 mL·min⁻¹·1.73 m⁻². Mendelian randomization analyses for CHD showed an association among participants with eGFR <60 mL·min⁻¹·1.73 m⁻², with a 14% (95% CI, 3%–27%) higher CHD risk per 5 mL·min⁻¹·1.73 m⁻² lower genetically predicted eGFR, but not for those with eGFR >105 mL·min⁻¹·1.73 m⁻². Results were not materially different after adjustment for factors associated with the eGFR genetic risk score, such as lipoprotein(a), triglycerides, hemoglobin A1c, and blood pressure. Mendelian randomization results for stroke were nonsignificant but broadly similar to those for CHD.

CONCLUSIONS: In people without manifest cardiovascular disease or diabetes, mild-to-moderate kidney dysfunction is causally related to risk of CHD, highlighting the potential value of preventive approaches that preserve and modulate kidney function.

Key Words: cardiovascular diseases ■ coronary disease ■ kidney diseases ■ stroke

Clinical Perspective

What Is New?

- In people without manifest cardiovascular disease or diabetes, there is a nonlinear causal relationship between kidney function and coronary heart disease.
- Even mildly reduced kidney function is causally associated with higher risk of coronary heart disease with a possible risk threshold for eGFR value of ≈75 mL·min⁻¹·1.73 m⁻².
- The effect of reduced kidney function on coronary heart disease is independent of traditional cardiovascular risk factors.

What Are the Clinical Implications?

- Preventive approaches that can preserve and modulate kidney function can help prevent cardiovascular diseases.
- Given the nonlinear causal relationship, it may be a
 preferable strategy to identify individuals in the population with mild-to-moderate kidney dysfunction
 and target them for renoprotective interventions
 alongside routine strategies to reduce cardiovascular risk.

hronic kidney disease (CKD), a major public health burden, affects >10% of the adult population globally. 1.2 Kidney failure is associated with a high risk of cardiovascular disease (CVD) and all-cause mortality. 3-5 Strong associations have also been reported

Nonstandard Abbreviations and Acronyms

CHD Coronary heart diseaseCKD Chronic kidney diseaseCVD Cardiovascular diseases

eGFR Estimated glomerular filtration rate **EPIC-CVD** European Prospective Investigation into

Cancer and Nutrition - Cardiovascular

Disease Study

GRS Genetic risk score

MVP Million Veteran Program

UKB UK Biobank

between non-dialysis-dependent CKD and these outcomes in both people without manifest CVD and patients with ischemic CVD, heart failure, high blood pressure, or diabetes. These observations have led to guideline recommendations that patients with CKD should be regarded as being at very high risk of CVD. 8,9

It is not known, however, whether mild-to-moderate kidney dysfunction is causally relevant to CVD or whether the increase in CVD risk associated with kidney dysfunction is related to changes in known risk factors, such as blood pressure and dyslipidemia, which seem to be a direct result of kidney dysfunction. 10-12 An approach to help evaluate the causal relevance of kidney dysfunction to CVD is Mendelian randomization. Mendelian randomization uses genetic variants specifically related to a particular exposure to compare

genetically defined population subgroups with different average levels of the exposure. The independent segregation of alleles at conception means that these genetically defined subgroups should not differ systematically with respect to confounding variables, creating a natural experiment analogous to a randomized trial. Therefore, compared with conventional observational analyses, Mendelian randomization analyses provide more reliable insights into causal relationships between risk factors and disease outcomes.^{13,14}

Previous Mendelian randomization analyses that have assumed a linear dose-response relationship between kidney function and CVD have reported null associations. However, observational analyses have reported U-shaped associations of CVD risk with creatinine-based estimated glomerular filtration rate (eGFR), a measure of kidney function. Therefore, drawing on multiple large-scale population bioresources, we evaluated the causal relevance of eGFR to coronary heart disease (CHD) and stroke, using Mendelian randomization methods tailored to nonlinear relationships, 16-20 which require concomitant information on eGFR, genetic determinants of eGFR, and first-ever CVD outcomes in the same individuals.

METHODS

The data, code, and study material that support the findings of this study are available from the corresponding author on reasonable request.

Study Design and Study Overview

This study involved interrelated components (Figure 1). First, we characterized observational associations between eGFR and incident CHD or stroke, using data from the Emerging Risk Factors Collaboration,²¹ EPIC-CVD (European Prospective Investigation into Cancer and Nutrition-Cardiovascular Disease Study),22 Million Veteran Program (MVP),23 UK Biobank (UKB),24 collectively involving 648 135 participants, who had serum creatinine measurements but no known CVD or diabetes at baseline. Second, we constructed a genetic risk score (GRS) for eGFR by computing a weighted sum of eGFR-associated index variants reported in a discovery genome-wide association study from the CKDGen consortium comprising 567 460 participants with European ancestry, 25 none of whom were from MVP, EPIC-CVD, or UKB. Third, we used this GRS to conduct Mendelian randomization analyses in a total of 413718 participants (ie, EPIC-CVD, MVP, UKB), with concomitant individuallevel information on genetics, serum creatinine, and disease outcomes. Fourth, to assess the potential for interference by horizontal pleiotropy²⁶ and explore potential mechanisms that could mediate associations between eGFR and CVD outcomes, we studied our GRS for eGFR in relation to several established and emerging risk factors for CVD.

Data Sources

Information on each of the data sources used in the analysis is provided in the Expanded Methods in the Supplemental Material. In brief, Emerging Risk Factors Collaboration, a

global consortium of population cohort studies with harmonized individual-participant data for multiple CVD risk factors, has included 47 studies with available information on serum creatinine and diabetes status at recruitment.21 EPIC-CVD, a case-cohort study embedded in the pan-European EPIC prospective study of >500000 participants, has recorded data on serum creatinine and imputed genome-wide array data from 21 of its 23 recruitment centers.²² MVP, a prospective cohort study recruited from 63 Veterans Health Administration medical facilities throughout the United States, has recorded serum creatinine, and imputed genome-wide array data are available for a large subset of its participants.23 UKB, a prospective study of 22 recruitment centers across the United Kingdom, has cohort-wide information on serum creatinine and imputed genome-wide array data.24 Relevant ethical approval and participant consent were already obtained in all studies that contributed data to this work.

Estimation of Kidney Function

Kidney function was estimated using creatinine-based eGFR, calculated using the Chronic Kidney Disease Epidemiology Collaboration equation.²⁷ Creatinine concentration was multiplied by 0.95 for studies in which measurements were not standardized to isotope-dilution mass spectrometry.^{25,28} In a subset of participants with available data, kidney function was also defined using the Chronic Kidney Disease Epidemiology Collaboration cystatin C-based equation²⁹ and albuminuria measured as spot urine albumin-to-creatinine ratio (Expanded Methods).

Observational Analyses

Primary outcomes were incident CHD and stroke. Details of end-point definitions for each study are provided in Table S1. Participants in the contributing studies were eligible for inclusion in the present analysis if they met all of the following criteria: (1) aged 30 to 80 years at recruitment; (2) had recorded information on age, sex, circulating creatinine, and diabetes status; (3) had a creatinine-based eGFR of <300 mL·min $^{-1}$ ·1.73 m $^{-2}$; (4) did not have a known history of CVD or diabetes at baseline; (5) had complete information on the risk factors of smoking status, systolic blood pressure, total cholesterol, high-density lipoprotein cholesterol, and body mass index; and (6) had at least 1 year of follow-up data after recruitment.

Hazard ratios for associations of creatinine-based eGFR with incident CHD and stroke were calculated using Cox regression, stratified by sex and study center, and when appropriate, adjusted for traditional vascular risk factors (defined here as age, systolic blood pressure, smoking status, total cholesterol, high-density lipoprotein cholesterol, and body mass index) on a complete-case basis. To account for the EPIC-CVD case-cohort design, Cox models were adapted using Prentice weights.³⁰ To avoid overfitting models, studies contributing <20 incident events to the analysis of a particular outcome were excluded from the analysis. Fractional polynomials were used to characterize nonlinear relationships of creatinine-based eGFR with risk of CHD and stroke, adjusted for age and CVD risk factors.31 Study-specific estimates for each outcome were pooled across studies using multivariable random-effects meta-analysis, using a reference point of 90 mL·min⁻¹·1.73 m⁻². When information on urinary biomarkers in UKB was available,

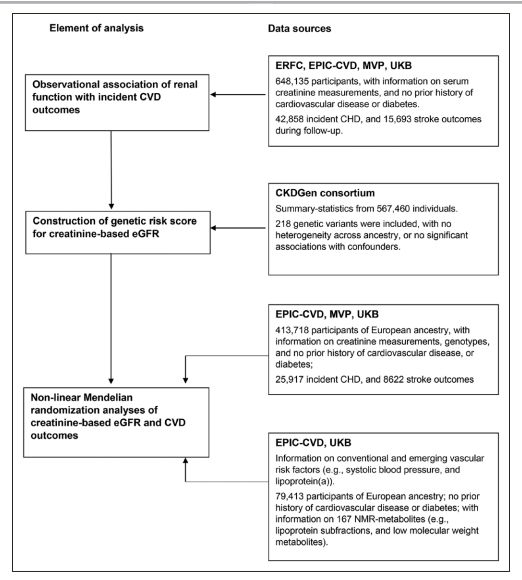


Figure 1. Study design and overview.

CHD indicates coronary heart disease; CKDGen, CKD Genetics consortium; CVD, cardiovascular disease; eGFR, estimated glomerular filtration rate; EPIC-CVD, European Prospective Investigation into Cancer and Nutrition—Cardiovascular Disease; ERFC, Emerging Risk Factors Collaboration; MVP, Million Veteran Program; NMR, nuclear magnetic resonance; and UKB, UK Biobank.

participants were grouped into tenths on the basis of levels of urinary albumin-to-creatinine ratio to assess the shapes of associations between urinary biomarkers and CVD risk, using participants without albuminuria as the reference group.³²

GRS for Kidney Function

Using individual-participant data from EPIC-CVD, MVP, and UKB, we calculated a GRS³³ weighted by the conditional effect estimated of the genetic variants associated ($P\!\!<\!5\!\times\!10^{-8}$) with creatinine-based eGFR in CKDGen,²⁵ a global genetics consortium that has published genome-wide association study summary statistics for creatinine-based eGFR. Of the 262 variants associated with creatinine-based eGFR, 37 were excluded because of ancestry heterogeneity as reported in CKDGen,²⁵ 4 were excluded because of associations ($P\!\!<\!5\!\times\!10^{-8}$) with vascular risk factors as reported in previous genome-wide association studies (ie, smoking

status, alcohol consumption, and education attainment),³⁴ and 3 were excluded because of missingness in at least 1 of the contributing studies, leaving 218 variants for the primary GRS for creatinine-based eGFR.

In sensitivity analysis, we constructed 2 restricted GRSs using 126 and 121 genetic variants that were likely to be relevant for kidney function on the basis of their associations with cystatin C-based eGFR³⁵ and blood urine nitrogen,²⁵ respectively. Sensitivity analysis was also conducted using a GRS that included all 262 transancestry eGFR-associated index variants. Furthermore, to evaluate traits that could mediate or confound (through horizontal pleiotropy) the associations between genetically predicted eGFR and outcomes, we tested associations of GRSs for eGFR with a range of cardiovascular risk factors in UKB and EPIC-CVD and with 167 metabolites measured using targeted high-throughput nuclear magnetic resonance metabolomics (Nightingale Health Ltd) in UKB.

Mendelian Randomization Analyses

To account for the nonlinear relationship between eGFR and risk of CVD outcomes in observational analyses, we performed a stratified Mendelian randomization analysis using methods described previously. 16-20 For each participant, we calculated the residual eGFR by subtracting the genetic contribution determined by the GRS from observed eGFR. Participants were grouped on the basis of their residual eGFR into 5-unit categories between 45 and $<105 \text{ mL}\cdot\text{min}^{-1}\cdot 1.73 \text{ m}^{-2}$, plus $<45 \text{ and } \ge 105 \text{ mL}\cdot\text{min}^{-1}\cdot 1.73$ m⁻². By stratifying on residual eGFR, we compared individuals in the population who would have an eGFR in the same category if they had the same genotype and reduced the potential influence of collider bias. We then calculated Mendelian randomization estimates for each eGFR category using the ratio method with the GRS as an instrumental variable, adjusting for age, agesquared, sex, study center, and the first 10 principal components. Stratum-specific estimates were combined across studies using fixed-effect meta-analysis and plotted as a piecewise-linear function of eGFR, with pointwise confidence intervals calculated by resampling the stratum-specific estimates. Sensitivity analyses used non-parametric doubly-ranked stratification method. Detailed methods describing statistical analysis are in the Expanded Methods. Analyses used STATA 15.1 and R 3.6.1.

RESULTS

Among the 648 135 participants without history of CVD or diabetes at baseline, the mean age was 57 years, 57% were men, and 4.4% had creatinine-based eGFR <60 mL·min $^{-1}\cdot1.73$ m $^{-2}$ (Table 1, Tables S2 and S3). During 6.8 million person-years of follow-up, there were 42 858 incident CHD outcomes and 15 693 strokes. Up to 413 718 participants of European ancestry from EPIC-CVD, MVP, and UKB contributed to the main genetic analyses (Figure 1). Distributions of serum creatinine concentration and creatinine-based eGFR were broadly similar across studies (Figures S1 and S2).

Observational Associations of eGFR With Cardiovascular Outcomes

For both CHD and stroke, there were U-shaped associations of creatinine-based eGFR. Compared with participants with creatinine-based eGFR values between 60 and 105 mL·min⁻¹·1.73 m⁻², risks of both CHD and stroke were higher in people with eGFR <60 or >105 mL·min⁻¹·1.73 m⁻² (Figure 2, Figure S3). The shapes of these associations did not change substantially after adjustment for several traditional risk factors (Figure 2). Associations were similar in men and women, in clinically relevant subgroups (ie, smokers, people with obesity, or hypertension; Figure S4), in the different studies contributing to this analysis (Figure S5), and when participants with a history of diabetes or missing information on cardiovascular risk factors were included (Figures S6 and S7). Similar associations were also observed for ischemic stroke (Figure S3).

For the 338044 participants in UKB with available data on serum cystatin C and urinary albumin-to-creatinine ratio, there were broadly similar associations of CHD or stroke with cystatin C-based eGFR as creatinine-based eGFR equations, but only when eGFR values were lower than ≈90 mL·min⁻¹·1.73 m⁻². However, there was no evidence of higher risk of CHD in participants with cystatin C-based eGFR values > 105 mL·min⁻¹·1.73 m⁻² (Figure S8), in contrast with creatinine-based eGFR values >105 mL·min⁻¹·1.73 m⁻². Levels of urinary microalbumin and urinary albumin-to-creatinine ratio showed approximately linear associations with risk of CHD and stroke, which were somewhat attenuated after adjustment for traditional risk factors (Figure S9). Compared with participants with a creatinine-based eGFR of 75 to <90 mL·min⁻¹·1.73 m⁻² and without albuminuria, participants with albuminuria had higher risk of CHD and stroke (Figure S10).

Mendelian Randomization of Genetically Predicted eGFR With Cardiovascular Outcomes

The GRS for eGFR (Table S4) explained 2.0% of variation in creatinine-based eGFR in EPIC-CVD, 2.2% in MVP, and 3.2% in UKB. A 1 SD increase in the GRS for eGFR was associated with 0.18 SD higher creatinine-based eGFR (Table S5, Figure S11). The GRS for eGFR was not associated with body mass index, diabetes, smoking status, or low-density lipoprotein cholesterol concentrations but showed modest associations with lipoprotein(a), triglycerides, blood pressure, and hemoglobin A1c measurement (Figure S11). Modest associations were also observed between the GRS for eGFR and triglyceriderelated lipoprotein subclasses in a subset of participants with available data (Figure S12).

In nonlinear Mendelian randomization analysis, we observed a curvilinear relationship between genetically predicted eGFR and CHD (Figure 3). Among participants with eGFR <60 mL·min⁻¹·1.73 m⁻², each 5 mL·min⁻¹·1.73 m⁻² lower genetically predicted eGFR was associated with 14% (95% CI, 3%-27%) higher risk of CHD (Table 2). There was no clear evidence of association among participants with eGFR >75 mL·min⁻¹·1.73 m⁻² (Figure 3). Similar, but not statistically significant, associations were observed for stroke (Table 2, Figure 3). Overall, stratumspecific localized average causal estimates and nonlinear Mendelian randomization estimates were compatible across the studies contributing to this analysis (Table S6, Figure S13). Findings were supported in analyses using the non-parametric doubly-ranked stratification (Table S7, Figure S14). Similar associations were observed in analyses that further adjusted for systolic blood pressure, lipoprotein(a), hemoglobin A1c, and triglycerides (Figure S15), included participants with a history of diabetes at baseline (Figure S16), or used ischemic stroke as the stroke outcome (Figure S17). Results were also similar

Table 1. Study-Level and Participant-Level Characteristics of the Contributing Data Sources

ERFC	EPIC-CVD	ИКВ	MVP
47 cohorts from 19 countries	21 centers from 8 European countries	England, Scotland, and Wales	United States
1964-2008	1990-2002	2006-2010	2011-Present
129601	20 985	350193	147356
58.3 (8.9)	56.3 (9.0)	56.3 (8.1)	57.9 (11.9)
68 278 (52.7)	9670 (46.1)	155 284 (44.3)	128610 (87.3)
26.3 (4.3)	26.1 (4.0)	27.1 (4.6)	29.0 (5.5)
135 (20)	138 (21)	137 (19)	130 (16)
38381 (29.6)	6233 (29.7)	36 422 (10.4)	14394 (9.77)
1.4 (0.4)	1.4 (0.4)	1.5 (0.4)	1.3 (0.4)
5.8 (1.1)	6.2 (1.2)	5.8 (1.1)	4.8 (1.0)
0.94 (0.22)	0.82 (0.23)	0.81 (0.18)	1.0 (0.4)
11 121 (8.6)	3113 (14.8)	44303 (12.7)	17 988 (12.2)
32 971 (25.4)	9400 (44.8)	165 603 (47.3)	41 461 (28.1)
44 654 (34.5)	5524 (26.3)	100351 (28.7)	46 200 (31.4)
30 751 (23.7)	2306 (11.0)	33 895 (9.7)	29 552 (20.1)
10105 (7.8)	642 (3.1)	6041 (1.7)	12 155 (8.2)
84.5 (16.6)	92.1 (14.8)	91.2 (13.1)	84.9 (18.1)
10390 (8.0)	7638 (36.4)	13863 (4.0)	10967 (7.4)
4838 (3.7)	3572 (17.0)	4544 (1.3)	2739 (1.8)
	47 cohorts from 19 countries 1964–2008 129601 58.3 (8.9) 68 278 (52.7) 26.3 (4.3) 135 (20) 38 381 (29.6) 1.4 (0.4) 5.8 (1.1) 0.94 (0.22) 11 121 (8.6) 32 971 (25.4) 44 654 (34.5) 30 751 (23.7) 10 105 (7.8) 84.5 (16.6) 10 390 (8.0)	47 cohorts from 19 countries 1964–2008 1990–2002 129601 20 985 58.3 (8.9) 56.3 (9.0) 68 278 (52.7) 9670 (46.1) 26.3 (4.3) 135 (20) 138 (21) 38 381 (29.6) 6233 (29.7) 1.4 (0.4) 1.4 (0.4) 5.8 (1.1) 0.94 (0.22) 0.82 (0.23) 11 121 (8.6) 3113 (14.8) 32971 (25.4) 44654 (34.5) 5524 (26.3) 30 751 (23.7) 10 105 (7.8) 84.5 (16.6) 92.1 (14.8) 10 390 (8.0) 7638 (36.4)	47 cohorts from 19 countries 21 centers from 8 European countries 2006–2010 1964–2008 1990–2002 2006–2010 129 601 20 985 350 193 58.3 (8.9) 56.3 (9.0) 56.3 (8.1) 68 278 (52.7) 9670 (46.1) 155 284 (44.3) 26.3 (4.3) 26.1 (4.0) 27.1 (4.6) 135 (20) 138 (21) 137 (19) 38 381 (29.6) 6233 (29.7) 36 422 (10.4) 1.4 (0.4) 1.4 (0.4) 1.5 (0.4) 5.8 (1.1) 6.2 (1.2) 5.8 (1.1) 0.94 (0.22) 0.82 (0.23) 0.81 (0.18) 11 121 (8.6) 3113 (14.8) 44 303 (12.7) 32 971 (25.4) 9400 (44.8) 165 603 (47.3) 44 654 (34.5) 5524 (26.3) 100 351 (28.7) 30 751 (23.7) 2306 (11.0) 33 895 (9.7) 10 105 (7.8) 642 (3.1) 6041 (1.7) 84.5 (16.6) 92.1 (14.8) 91.2 (13.1) 10 390 (8.0) 7638 (36.4) 13 863 (4.0)

Data are n, n (%), or mean (SD). Participants with a history of diabetes or cardiovascular diseases at recruitment, or incomplete information on creatinine, body mass index, systolic blood pressure, smoking status, high-density lipoprotein cholesterol, or total cholesterol were excluded. ERFC indicates Emerging Risk Factors Collaboration; EPIC-CVD, European Prospective Investigation into Cancer and Nutrition–Cardiovascular Disease Study; MVP, Million Veteran Program; and UKB, LIK Richank

using GRS for cystatin C-based eGFR, blood urine nitrogen, or variants associated with creatinine-based eGFR regardless of ancestry heterogeneity (Figure S18).

DISCUSSION

In analyses combining genetic, biomarker, and clinical data in $\approx\!\!640\,000$ participants, our study has suggested that, in people without manifest CVD or diabetes, even mildly reduced kidney function is causally associated with a higher risk of CVD outcomes. Our results provide novel causal insights and highlight the wider potential value of preventive approaches that can preserve and modulate kidney function.

First, our study estimated a dose-response curve for genetically predicted eGFR and CHD, identifying an eGFR value of ≈75 mL·min⁻¹·1.73 m⁻² as a possible risk threshold. Therefore, the causal relationship of kidney function with CHD is nonlinear in shape, in contrast with those for blood pressure and low-density lipoprotein cholesterol, which each have approximately log-linear relationships with CHD risk across their range of values. In contrast with population-wide strategies to improve blood pressure and low-density lipoprotein cholesterol levels, this finding implies that it may be a preferable strategy to identify those in the population with mild-to-moderate kidney dysfunction and target them

for renoprotective interventions alongside routine strategies to reduce cardiovascular risk. For example, the use of renoprotective interventions, such as renin angiotensin aldosterone system inhibitors³⁶ and inhibitors of sodiumglucose cotransporter 2, might provide a potential means to do so.³⁷ Our findings encourage additional evaluation of such agents in patients with CKD without manifest CVD or diabetes.^{38,39}

Second, we found that our GRS for eGFR was modestly associated with several established and emerging CVD risk factors, including plasma concentration of proatherogenic lipids (eg, lipoprotein(a), triglycerides, and triglyceride-related lipoprotein subclasses), hemoglobin A1c values, and blood pressure, consistent with previous studies. 11,40 However, adjustment for such factors did not materially alter the associations between eGFR and atherosclerotic CVD, indicating that they are unlikely to mediate or confound the associations between genetically predicted kidney dysfunction and CHD or stroke and limiting the likelihood that results are subject to influences of horizontal pleiotropy. These results suggest that the effect of reduced kidney function on CVD is independent of traditional cardiovascular risk factors and underscores the potential importance of direct preservation of renal function to prevent CVD, in addition to control of known risk factors.

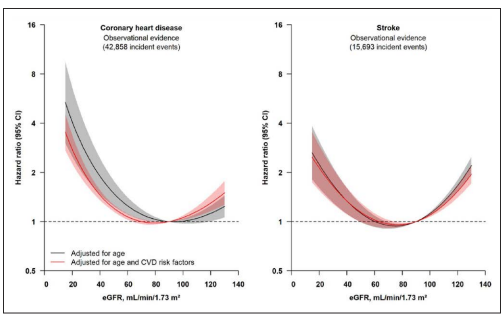


Figure 2. Observational associations of eGFR levels with risk of coronary heart disease and stroke (n=648 135).

Participants with missing information on age and CVD risk factors (systolic blood pressure, total and high-density lipoprotein cholesterol, body mass index, and smoking status) were excluded from the analyses. Hazard ratios were estimated using Cox regression, adjusting for age and CVD risk factors (systolic blood pressure, total and high-density lipoprotein cholesterol, body mass index, and smoking status), and stratified by sex and study center. The reference point is 90 mL·min⁻¹·1.73 m⁻². Shaded regions indicate 95% CIs. CVD indicates cardiovascular disease; and eGFR, estimated glomerular filtration rate.

Third, our data help to resolve controversies about the relevance to CHD of higher-than-average eGFR. In contrast with the observation that higher-than-average creatinine-based eGFR values are associated with higher CHD risk at >105 mL·min⁻¹·1.73 m⁻², we found that genetically predicted higher eGFR values were not associated with CHD risk in this same group. This discordance implies different pathophysiological meanings of creatinine-based

eGFR values >105 mL·min⁻¹·1.73 m⁻² (which may represent a transient state of hyperfiltration before progression to poorer kidney function and CKD) and genetically predicted higher eGFR values (which represent a lifelong tendency toward exposure to better kidney function). This explanation is supported by our findings showing that the association between higher creatinine-based eGFR values and higher CHD risk was principally in participants

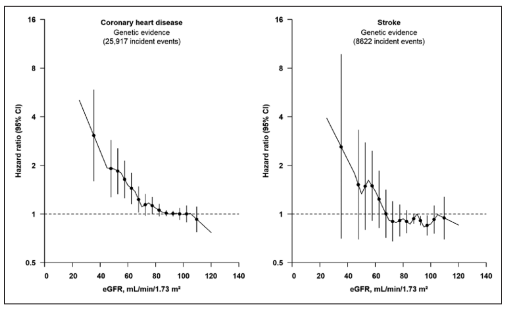


Figure 3. Associations of genetically predicted eGFR with risk of coronary heart disease and stroke (n=413718).

The reference point is 90 mL·min⁻¹·1.73 m⁻². Gradients at each point of the curve represent the localized average causal effect on coronary heart disease or stroke per 5 mL·min⁻¹·1.73 m⁻² change in genetically predicted eGFR. The vertical lines represent 95% CIs. Analyses were adjusted for age, age-squared, sex, study center, and the first 10 principal components of ancestry. eGFR indicates estimated glomerular filtration rate.

Table 2. Mendelian Randomization Estimates per 5 mL·min⁻¹·1.73 m⁻² Lower Genetically Predicted eGFR With Risk of Coronary Heart Disease and Stroke

eGFR, mL·min ⁻¹ ·1.73 m ⁻²	Mean eGFR, mL·min ⁻¹ ·1.73 m ⁻²	No. of participants	Coronary heart disease		Stroke	
			No. of events	HR (95% CI)	No. of events	HR (95% CI)
<60	51.2	14818	1749	1.14 (1.03–1.27)	491	1.19 (0.97–1.47)
60 to <75	68.9	53 256	4314	1.08 (1.01-1.15)	1318	1.14 (1.01-1.28)
75 to <90	83.3	123664	8229	1.05 (1.00-1.10)	2565	0.97 (0.89-1.06)
90 to <105	96.9	176180	9973	1.01 (0.96-1.05)	3582	1.06 (0.98–1.14)
≥105*	109.7	45800	1652	0.91 (0.82-1.02)	666	0.95 (0.79-1.14)

Hazard ratios are shown per 5 mL·min⁻¹·1.73 m⁻² lower genetically predicted eGFR and are adjusted for age, age-squared, sex, study center, and the first 10 principal components. eGFR indicates estimated glomerular filtration rate; and HR, hazard ratio.

*HRs in the group with eGFR >105 were shown per 5 mL·min⁻¹·1.73 m⁻² higher genetically predicted eGFR. Mean eGFR within each stratum was weighted by the number of participants from each contributing study, and HR estimates within each stratum were meta-analyzed using inverse variance weighting and fixed effects.

who had albuminuria (and, therefore, preexisting kidney damage) at entry into the study.

Fourth, our results are broadly consistent with a causal relationship between eGFR and stroke. The lack of statistically significant findings in our Mendelian randomization analysis for stroke outcomes principally reflects the lower power of our study to evaluate a GRS with stroke compared with CHD. It may also be attributable to pathogenetic heterogeneity in stroke diagnoses (eg, cardioembolic, small vessel disease, and hemorrhagic subtypes may be less driven by atherosclerotic pathology than other ischemic stroke subtypes).^{41,42}

Our study had major strengths, including a large sample size, access to individual-participant data, use of multiple genetic causal inference methods tailored to the evaluation of nonlinear disease associations, and an updated GRS that explains more variation in eGFR than previous analyses.14 However, there are also potential limitations. First, Mendelian randomization assumptions state that the only causal pathway from the genetic variants to the outcome is through eGFR. Although we assessed the potential for interference by horizontal pleiotropy, there is the possibility of residual confounding by unrecognized effects of genotypes on other risk factors and by adaptation during early life to compensate for genetically lower eGFR. Second, to reduce the scope for confounding by ancestry (population stratification), our analyses were limited to participants of European ancestries. This limitation means that our findings might not be applicable to other populations, and additional studies on this topic are needed, especially in non-European ancestry populations. Third, although serum creatinine is used routinely for estimating eGFR, true measurement of GFR requires the use of inulin, iohexol, or iothalamate. Assay of serum creatinine is liable to interference from other serum components (eg, bilirubin and glucose)43,44 and autoimmune activation⁴⁵ and is sensitive to changes in individuals' muscle mass (eg, sarcopenia). Assessment of cystatin C, an analyte that enables an alternative calculation of eGFR without the potential limitations of creatinine, was available only in a subset of the participants we studied. However, our genetic analyses restricted to genetic variants additionally associated with other biomarkers of kidney function showed results consistent with those for creatinine-based eGFR. Last, we used the 2009 Chronic Kidney Disease Epidemiology Collaboration equation to calculate eGFR. However, our analysis was limited to populations with European ancestry, in which the 2009 and 2021 Chronic Kidney Disease Epidemiology Collaboration equations provide similar estimates of eGFR.⁴⁶

CONCLUSIONS

In people without manifest CVD or diabetes, mild-tomoderate kidney dysfunction was causally related to cardiovascular outcomes, highlighting the potential cardiovascular benefit of preventive approaches that improve kidney function.

ARTICLE INFORMATION

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Affiliations

Massachusetts Veterans Epidemiology Research and Information Center (MAV-ERIC), VA Boston Healthcare System, Boston, MA (L.G., K.C., R.J.S., D.C.P., P.S., J.J., J.P.C., J.M.G.). BHF Cardiovascular Epidemiology Unit, Department of Public Health and Primary Care (L.G., L.S., S. Bell, S.K.K., S. Burgess, K.M., A.M.M., T.R.B., E.A., L.C., J.R.S., P.W., L. Pennells, S.H., M.I., J.D., A.S.B., A.M.W., E.D.A.), BHF Centre of Research Excellence, School of Clinical Medicine, Addenbrooke's Hospital (A.M.M., S. Burgess, J.D., A.M.W., A.S.B., E.D.A.), Heart and Lung Research Institute, University of Cambridge, Cambridge UK (L.G., S. Bell, S.K.K., S. Burgess, K.M., A.M.M., E.A., L. Pennells, M.I., J.D., A.S.B., A.M.W., E.D.A.). Medical Research Council Biostatistics Unit (A.M.M., S. Burgess), Stroke Research Group, Department of Clinical Neurosciences (S. Bell), NIHR Blood and Transplant Research Unit in Donor Health and Behaviour (S. Bell, T.R.B., E.A., J.D., A.S.B., A.M.W., E.D.A.), MRC Epidemiology Unit, School of Clinical Medicine (C.L., N.W.), University of Cambridge, UK. Division of Aging (K.C., S.P., J.P.C. J.M.G.), Division of Cardiovascular Medicine (J.J., T.A.G.), Brigham and Women's Hospital, Harvard Medical School, Boston, MA. Department of Epidemiology, Boston University School of Public Health, MA (R.J.S.). Division of Nephrology, Department of Medicine (C.R.-C., E.A.A.), Department of Biostatistics (R. Tao), Vanderbilt University Medical Center, Nashville, TN. Medical Research Council Population Health Research Unit, Clinical Trial Service Unit and Epidemiological Studies Unit (N.S., C.B.), Cancer Epidemiology Unit (T.T.), Nuffield Department of Population Health, University of Oxford, UK. Department of Cardiology, Trakya University School of Medicine, Edirne, Turkey (S.A.). Ministry of Health of the Basque Government, Sub Directorate for Public Health and Addictions of Gipuzkoa, San Sebastián, Spain (P.A.). Biodonostia Health Research Institute, Epidemiology of Chronic and Communicable Diseases Group, San Sebastián, Spain (P.A.). Centro de Investigación Biomédica

en Red de Epidemiología y Salud Pública (CIBERESP), Madrid, Spain (P.A., A.G.d.I.C., D.P., C. Santiuste). Division of Clinical Epidemiology and Aging Research (V.A.), Department of Cancer Epidemiology (S.K.J., R.K., V.K.), German Cancer Research Center (DKFZ), Heidelberg, Germany. Division of Family Medicine and Primary Care, Department of Neurobiology, Care Sciences and Society (NVS), Karolinska Institutet, Stockholm, Sweden (J.A., H.B.), School of Health and Social Studies, Dalarna University, Falun, Sweden (J.A.). Wellbeing & Preventable Chronic Diseases (WPCD) Division, Menzies School of Health Research, Charles Darwin University, Darwin, NT, Australia (E.L.M.B.). Baker Heart and Diabetes Institute, Melbourne, VIC, Australia (E.L.M.B., M.I.). Institute of Medicine, School of Public Health and Community Medicine (C.B.), Institute of Medicine, Department of Molecular and Clinical Medicine (P.-O.H., A.R.), Sahlgrenska Academy, University of Gothenburg, Sweden. National Institute for Public Health and the Environment (RIVM), Bilthoven, the Netherlands (J.M.A.B., W.M.M.V.). Network Aging Research (NAR), Heidelberg University, Germany (H.B.). Studium Patavinum (E.C.), Department of Medicine (V.T.), University of Padua, Italy. Dipartimento di Salute Mentale e Fisica e Medicina Preventiva, Università degli Studi della Campania 'Luigi Vanvitelli', Caserta, Italy (P.C.). William Harvey Research Institute, NIHR Barts Biomedical Research Centre, Queen Mary University of London, UK (J.A.C.). Johns Hopkins Bloomberg School of Public Health, Baltimore, MD (J.C.). Larner College of Medicine, The University of Vermont, Burlington (M.C.). The Gertner Institute for Epidemiology and Health Policy Research, Sheba Medical Center, Tel Hashomer, Israel (R.D.). School of Public Health, Department of Epidemiology and Preventive Medicine, Tel Aviv University, Ramat Aviv, Tel Aviv, Israel (R.D.). Feinstein Institutes for Medical Research, Northwell Health, Manhasset, New York, NY (R.D., K.W.D.). Amsterdam University Medical Centers, VUMC, the Netherlands (R.T.d.J.). Department of Cardiovascular, Endocrine-metabolic Diseases and Aging, Istituto Superiore di Sanità, Rome, Italy (C.D., L. Palmer), Department of Clinical Sciences, Malmö, Lund University, Sweden (G.E., O.M.). International Agency for Research on Cancer (IARC), World Health Organization, Lyon, France (H.F., E.W.). 12 Octubre Hospital Research Institute, Madrid, Spain (A.G.d,I,C.). Faculty of Medicine, University of Iceland, Reykjavik, Iceland and Icelandic Heart Association, Kopavogur, Iceland (V.G.). Medical School Faculty of Health & Medical Sciences, The University of Western Australia, Perth, WA, Australia (G.J.H.). Region Västra Götaland, Sahlgrenska University Hospital, Department of Medicine Geriatrics and Emergency Medicine/Östra, Gothenburg, Sweden (P.-O.H., A.R.). School of Public Health (A.K.H., I.T., E.R.), The George Institute for Global Health (M.W.), Imperial College London, UK. Department of Internal Medicine, Division of Nephrology and Transplantation, Erasmus MC, University Medical Center Rotterdam, the Netherlands (E.J.H.). Public Health, Department of Social Medicine, Osaka University Graduate School of Medicine, Suita, Japan (H.I.) University of Eastern Finland (UEF), Kuopio, Finland (J.K.). Department of Neurology & Neurosurgery, Medical University of Innsbruck, Innsbruck, Austria (S.K.). Clinical Epidemiology Team, Institute of Health Economics, Medical University of Innsbruck, Innsbruck, Austria (S.K., P.W.). Institute of Epidemiology and Medical Biometry, University of Ulm, Germany (W.K.). Deutsches Herzzentrum München, Technische Universität München, Germany (W.K.). German Centre for Cardiovascular Research (DZHK), Partner Site Munich Heart Alliance (W.K.). School of Public Health, University of Washington, Seattle (R.A.K.). Danish Cancer Society Research Center, Copenhagen, Denmark (C.K., A.T.). Medical Research Council Integrative Epidemiology Unit, University of Bristol, UK (D.A.L.). Population Health Science, Bristol Medical School, UK (D.A.L.). Department of Surgical and Perioperative sciences, Urology and Andrology, Umeå University, Sweden (B.L.). University Paris-Saclay, UVSQ, Inserm, Villejuif, France (C. MacDonald). Institute for Cancer Research, Prevention and Clinical Network (ISPRO), Florence, Italy (G.M.). Helmholtz Zentrum München, Munich, Germany (C. Meisinger). Navarra Public Health Institute, IdiSNA, Pamplona, Spain (C.M.I.). Red de Investigación en Servicios de Salud en Enfermedades Crónicas (REDISSEC), Pamplona, Spain (C.M.I.). Graduate School of Medical Sciences, Kyushu University, Fukuoka, Japan (T.N.). London School of Hygiene & Tropical Medicine, UK (D.N.). Herlev and Gentofte Hospital (B.G.N.), Frederiksberg Hospital B.G.N.), Copenhagen University Hospital, Copenhagen, Denmark. Department of Clinical Medicine, Faculty of Health and Medical Sciences (B.G.N.), Department of Public Health (A.T.), University of Copenhagen, Denmark, Julius Center for Health Sciences and Primary Care, University Medical Center Utrecht, Utrecht University, the Netherlands (C.O.-M., Y.T.v.d.S., W.M.M.V.). Escuela Andaluza de Salud Pública (EASP), Granada, Spain (D.P.). Instituto de Investigación Biosanitaria ibs.GRANADA, Granada, Spain (D.P.). Consejería de Sanidad del Principado de Asturias Oviedo, Asturias, Spain (J.R.Q.G.). Unit of Cancer Epidemiology, Città della Salute e della Scienza University-Hospital, Turin, Italy (C. Sacerdote). Department of Social and Environmental Medicine, Kanazawa Medical University, Uchinada, Japan (M.S.). Department of Epidemiology, Murcia Regional Health Council, IMIB-Arrixaca, Spain (C. Santiuste). German Institute of Human Nutrition Potsdam-Rehbruecke, Nuthetal, Germany (M.B.S.). German Center for Diabetes Research (DZD), Neuherberg, Germany (M.B.S.). Institute of Nutritional Science, University of Potsdam, Germany (M.B.S.). Fondazione IRCCS Istituto Nazionale dei Tumori di Milano, Milano, Italy (S.S.). Department of Medical Sciences, Uppsala University, Sweden (J.S.). Hyblean Association for Epidemiological Reserach AIRE - ONLUS, Ragusa, Italy (R.T.). Universitätsmedizin Greifswald, Institut für Community Medicine, Abteilung SHIP/ Klinisch-Epidemiologische Forschung, Germany (H.V.). College of Public Health, University of Iowa (R.B.W.). University College London, UK (S.G.W.). The George Institute for Global Health, Camperdown, NSW, Australia (M.W.). Department of Public Health Medicine, Faculty of Medicine, and Health Services Research and Development Center, University of Tsukuba, Japan (K.Y.). Unit of Nutrition and Cancer, Epidemiology Research Program, Catalan Institute of Oncology, Bellvitge Biomedical Research Institute (IDIBELL), L'Hospitalet de Llobregat (Barcelona), Spain (R.Z.-R.). Center for Data and Computational Sciences, VA Boston Healthcare System, Boston, MA (S.P.). Department of Biostatistics, Boston University School of Public Health, MA (D.R.G.). VA Pal Alto Epidemiology Research and Information Center for Genomics, VA Palo Alto Health Care System, CA (P.S.T.). Medicine (Cardiovascular Medicine), Stanford University of School of Medicine, CA (P.S.T.). Office of Research and Development, Veterans Health Administration, Washington, DC (S.M.). Department of Veterans Affairs, Tennessee Valley Health Care System, Vanderbilt University, Nashville (T.L.E.). Medicine/Epidemiology, Vanderbilt Genetics Institute, Vanderbilt University Medical Center, Nashville, TN (T.L.E.). Department of Surgery, Corporal Michael Crescenz VA Medical Center and Perelman School of Medicine, University of Pennsylvania, Philadelphia (S.M.D.). Internal Medicine, VA Atlanta Healthcare System, Decatur, GA (P.W.F.W.). Emory University School of Medicine (Cardiology), Emory University, Atlanta, GA (P.W.F.W.). Center for Health Decision Science, Harvard T.H. Chan School of Public Health, Boston, MA (T.A.G.). Health Data Research UK Cambridge, Wellcome Genome Campus and University of Cambridge, UK (M.I., J.D., A.S.B., A.M.W., E.D.A.), The Alan Turing Institute, London, UK (M.I.). Computational Medicine, Berlin Institute of Health at Charité - Universitätsmedizin Berlin, Germany (C.L.). Department of Human Genetics, Wellcome Sanger Institute, Hinxton, UK (J.D.). Division of Nephrology & Hypertension, Department of Medicine, Tennessee Valley Health Care System and Vanderbilt University Medical Center, Nashville (A.M.H.). Cambridge Centre for AI in Medicine, UK (A.M.W.). Health Data Science Centre, Human Technopole, Milan, Italy (E.D.A.).

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Supplemental Material

Expanded Methods Tables S1-S7 Figures S1-S18

REFERENCES

- Global, regional, and national burden of chronic kidney disease, 1990– 2017: a systematic analysis for the Global Burden of Disease Study 2017. Lancet. 2020;395:709–733. doi: 10.1016/S0140-6736(20)30045-3
- Levey AS, Coresh J. Chronic kidney disease. Lancet. 2012;379:165–180. doi: 10.1016/S0140-6736(11)60178-5
- Di Angelantonio E, Chowdhury R, Sarwar N, Aspelund T, Danesh J, Gudnason V. Chronic kidney disease and risk of major cardiovascular dis-

- ease and non-vascular mortality: prospective population based cohort study. BMJ. 2010;341:c4986. doi: 10.1136/bmj.c4986
- Fox CS, Matsushita K, Woodward M, Bilo HJG, Chalmers J, Heerspink HJL, Lee BJ, Perkins RM, Rossing P, Sairenchi T, et al. Associations of kidney disease measures with mortality and end-stage renal disease in individuals with and without diabetes: a meta-analysis. *Lancet*. 2012;380:1662–1673. doi: 10.1016/S0140-6736(12)61350-6
- Baigent C, Burbury K, Wheeler D. Premature cardiovascular disease in chronic renal failure. *Lancet*. 2000;356:147–152. doi: 10.1016/S0140-6736(00)02456-9
- Webster AC, Nagler EV, Morton RL, Masson P. Chronic kidney disease. Lancet. 2017;389:1238–1252. doi: 10.1016/S0140-6736(16)32064-5
- Matsushita K, van der Velde M, Astor BC, Woodward M, Levey AS, de Jong PE, Coresh J, Gansevoort RT. Association of estimated glomerular filtration rate and albuminuria with all-cause and cardiovascular mortality in general population cohorts: a collaborative meta-analysis. *Lancet*. 2010;375:2073– 2081. doi: 10.1016/S0140-6736(10)60674-5
- Visseren FLJ, Mach F, Smulders YM, Carballo D, Koskinas KC, Bäck M, Benetos A, Biffi A, Boavida J-M, Capodanno D, et al. 2021 ESC Guidelines on cardiovascular disease prevention in clinical practice. Eur Heart J. 2021;42:3227–3337. doi: 10.1093/eurheartj/ehab484
- Grundy SM, Stone NJ, Bailey AL, Beam C, Birtcher KK, Blumenthal RS, Braun LT, de Ferranti S, Faiella-Tommasino J, Forman DE, et al. 2018 AHA/ ACC/AACVPR/AAPA/ABC/ACPM/ADA/AGS/APhA/ASPC/NLA/ PCNA Guideline on the management of blood cholesterol: a report of the American College of Cardiology/American Heart Association Task Force on Clinical Practice Guidelines. Circulation. 2019;139:e1082-e1143. doi: 10.1161/CIR.000000000000000625
- Said S, Hernandez GT. The link between chronic kidney disease and cardiovascular disease. J Nephropathol. 2014;3:99–104. doi: 10.12860/jnp. 2014.19
- Yu Z, Coresh J, Qi G, Grams M, Boerwinkle E, Snieder H, Teumer A, Pattaro C, Köttgen A, Chatterjee N, et al. A bidirectional Mendelian randomization study supports causal effects of kidney function on blood pressure. *Kidney Int.* 2020;98:708–716. doi: 10.1016/j.kint.2020.04.044
- Bulbul MC, Dagel T, Afsar B, Ulusu NN, Kuwabara M, Covic A, Kanbay M. Disorders of lipid metabolism in chronic kidney disease. *Blood Purif*. 2018;46:144–152. doi: 10.1159/000488816
- Davey Smith G, Ebrahim S. 'Mendelian randomization': can genetic epidemiology contribute to understanding environmental determinants of disease?.
 Int J Epidemiol. 2003;32:1–22. doi: 10.1093/ije/dyq070
- Charoen P, Nitsch D, Engmann J, Shah T, White J, Zabaneh D, Jefferis B, Wannamethee G, Whincup P, Mulick Cassidy A, et al. Mendelian randomisation study of the influence of eGFR on coronary heart disease. *Sci Rep.* 2016;6:28514. doi: 10.1038/srep28514
- Morris AP, Le TH, Wu H, Akbarov A, van der Most PJ, Hemani G, Smith GD, Mahajan A, Gaulton KJ, Nadkarni GN, et al. Trans-ethnic kidney function association study reveals putative causal genes and effects on kidneyspecific disease aetiologies. *Nat Commun.* 2019;10:29. doi: 10.1038/ s41467-018-07867-7
- Staley JR, Burgess S. Semiparametric methods for estimation of a nonlinear exposure-outcome relationship using instrumental variables with application to Mendelian randomization. *Genet Epidemiol.* 2017;41:341–352. doi: 10.1002/gepi.22041
- Sun YQ, Burgess S, Staley JR, Wood AM, Bell S, Kaptoge SK, Guo Q, Bolton TR, Mason AM, Butterworth AS, et al. Body mass index and all cause mortality in HUNT and UK Biobank studies: linear and non-linear mendelian randomisation analyses. *BMJ*. 2019;364:I1042. doi: 10.1136/bmj.I1042
- Biddinger KJ, Emdin CA, Haas ME, Wang M, Hindy G, Ellinor PT, Kathiresan S, Khera AV, Aragam KG. Association of habitual alcohol intake with risk of cardiovascular disease. *JAMA Netw Open.* 2022;5:e223849. doi: 10.1001/jamanetworkopen.2022.3849
- Arvanitis M, Qi G, Bhatt DL, Post WS, Chatterjee N, Battle A, McEvoy JW. Linear and nonlinear mendelian randomization analyses of the association between diastolic blood pressure and cardiovascular events: the J-curve revisited. *Circulation*. 2021;143:895–906. doi: 10.1161/CIRCULATIONAHA.120.049819
- Emerging Risk Factors Collaboration/EPIC-CVD/Vitamin D Studies Collaboration. Estimating dose-response relationships for vitamin D with coronary heart disease, stroke, and all-cause mortality: observational and Mendelian randomisation analyses. *Lancet Diabetes Endocrinol*. 2021;9:837–846. doi: 10.1016/S2213-8587(21)00263-1
- 21. The Emerging Risk Factors Collaboration. Danesh J, Erqou S, Walker M, Thompson SG, Tipping R, Ford C, Pressel S, Walldius G, Jungner I, Folsom AR, et al, The Emerging Risk Factors Collaboration: analysis of individual

- data on lipid, inflammatory and other markers in over 1.1 million participants in 104 prospective studies of cardiovascular diseases. *Eur J Epidemiol.* 2007;22:839–869. doi: 10.1007/s10654-007-9165-7
- Danesh J, Saracci R, Berglund G, Feskens E, Overvad K, Panico S, Thompson S, Fournier A, Clavel-Chapelon F, Canonico M, et al. EPIC-Heart: the cardiovascular component of a prospective study of nutritional, lifestyle and biological factors in 520,000 middle-aged participants from 10 European countries. Eur J Epidemiol. 2007;22:129-141. doi: 10.1007/s10654-006-9096-8
- Gaziano JM, Concato J, Brophy M, Fiore L, Pyarajan S, Breeling J, Whitbourne S, Deen J, Shannon C, Humphries D, et al. Million veteran program: a mega-biobank to study genetic influences on health and disease. J Clin Epidemiol. 2016;70:214–223. doi: 10.1016/j.jclinepi.2015.09.016
- Sudlow C, Gallacher J, Allen N, Beral V, Burton P, Danesh J, Downey P, Elliott P, Green J, Landray M, et al. UK biobank: an open access resource for identifying the causes of a wide range of complex diseases of middle and old age. PLoS Med. 2015;12:e1001779. doi: 10.1371/journal.pmed.1001779
- Wuttke M, Li Y, Li M, Sieber KB, Feitosa MF, Gorski M, Tin A, Wang L, Chu AY, Hoppmann A, et al. A catalog of genetic loci associated with kidney function from analyses of a million individuals. *Nat Genet*. 2019;51:957– 972. doi: 10.1038/s41588-019-0407-x
- Burgess S, Davey Smith G, Davies NM, Dudbridge F, Gill D, Glymour MM, Hartwig FP, Holmes MV, Minelli C, Relton CL, et al. Guidelines for performing Mendelian randomization investigations. Wellcome Open Res. 2019;4:186. doi: 10.12688/wellcomeopenres.15555.2
- Levey AS, Stevens LA, Schmid CH, Zhang YL, Castro AF, Feldman HI, Kusek JW, Eggers P, Van Lente F, Greene T, et al. A new equation to estimate glomerular filtration rate. *Ann Intern Med.* 2009;150:604–612. doi: 10.7326/0003-4819-150-9-200905050-00006
- Levey AS, Coresh J, Greene T, Marsh J, Stevens LA, Kusek JW, Van Lente F. Expressing the Modification of Diet in Renal Disease Study equation for estimating glomerular filtration rate with standardized serum creatinine values. Clin Chem. 2007;53:766–772. doi: 10.1373/clinchem.2006.077180
- Shlipak MG, Matsushita K, Arnlov J, Inker LA, Katz R, Polkinghorne KR, Rothenbacher D, Sarnak MJ, Astor BC, Coresh J, et al. Cystatin C versus creatinine in determining risk based on kidney function. N Engl J Med. 2013;369:932–943. doi: 10.1056/NEJMoa1214234
- Prentice RL. A case-cohort design for epidemiologic cohort studies and disease prevention trials. *Biometrika*. 1986;73:1–11.doi:10.1093/biomet/73.1.1
- White IR, Kaptoge S, Royston P, Sauerbrei W. Meta-analysis of non-linear exposure-outcome relationships using individual participant data: a comparison of two methods. Stat Med. 2019;38:326–338. doi: 10.1002/sim.7974
- 32. KDIGO 2012 Clinical Practice Guideline for the Evaluation and Management of Chronic Kidney Disease. Kidney Int Suppl. 2013;3:1–150.
- Burgess S, Thompson SG. Use of allele scores as instrumental variables for Mendelian randomization. *Int J Epidemiol.* 2013;42:1134–1144. doi: 10.1093/ije/dyt093
- Kamat MA, Blackshaw JA, Young R, Surendran P, Burgess S, Danesh J, Butterworth AS, Staley JR. PhenoScanner V2: an expanded tool for searching human genotype-phenotype associations. *Bioinformatics*. 2019;35: 4851–4853. doi: 10.1093/bioinformatics/btz469
- Stanzick KJ, Li Y, Schlosser P, Gorski M, Wuttke M, Thomas LF, Rasheed H, Rowan BX, Graham SE, Vanderweff BR, et al. Discovery and prioritization of variants and genes for kidney function in >1.2 million individuals. *Nat Commun*. 2021;12:4350. doi: 10.1038/s41467-021-24491-0
- Lewis EJ, Hunsicker LG, Bain RP, Rohde RD. The effect of angiotensinconverting-enzyme inhibition on diabetic nephropathy. The Collaborative Study Group. N Engl J Med. 1993;329:1456–1462. doi: 10.1056/ NEJM199311113292004
- Bhatt DL, Szarek M, Steg PG, Cannon CP, Leiter LA, McGuire DK, Lewis JB, Riddle MC, Voors AA, Metra M, et al. Sotagliflozin in patients with diabetes and recent worsening heart failure. N Engl J Med. 2021;384:117–128. doi: 10.1056/NEJMoa2030183
- Heerspink HJL, Stefánsson BV, Correa-Rotter R, Chertow GM, Greene T, Hou F-F, Mann JFE, McMurray JJV, Lindberg M, Rossing P, et al. Dapagliflozin in patients with chronic kidney disease. N Engl J Med. 2020;383:1436–1446. doi: 10.1056/NEJMoa2024816
- Zannad F, Ferreira JP, Pocock SJ, Anker SD, Butler J, Filippatos G, Brueckmann M, Ofstad AP, Pfarr E, Jamal W, et al. SGLT2 inhibitors in patients with heart failure with reduced ejection fraction: a meta-analysis of the EMPEROR-Reduced and DAPA-HF trials. *Lancet*. 2020;396:819– 829. doi: 10.1016/S0140-6736(20)31824-9
- Aguilar-Ramirez D, Alegre-Díaz J, Herrington WG, Staplin N, Ramirez-Reyes R, Gnatiuc L, Hill M, Romer F, Torres J, Trichia E, et al. Association of kidney function with NMR-quantified lipids, lipoproteins, and metabolic measures

- in Mexican adults. *J Clin Endocrinol Metab.* 2021;106:2828–2839. doi: 10.1210/clinem/dgab497
- Dichgans M, Pulit SL, Rosand J. Stroke genetics: discovery, biology, and clinical applications. *Lancet Neurol.* 2019;18:587–599. doi: 10.1016/S1474-4422(19)30043-2
- Sun L, Clarke R, Bennett D, Guo Y, Walters RG, Hill M, Parish S, Millwood IY, Bian Z, Chen Y, et al. Causal associations of blood lipids with risk of ischemic stroke and intracerebral hemorrhage in Chinese adults. *Nat Med.* 2019;25:569–574. doi: 10.1038/s41591-019-0366-x
- Bargnoux A-S, Kuster N, Cavalier E, Piéroni L, Souweine J-S, Delanaye P, Cristol J-P. Serum creatinine: advantages and pitfalls. J Lab Precis Med. 2018;3:71. doi: 10.21037/jlpm.2018.08.01
- Delanaye P, Cavalier E, Cristol JP, Delanghe JR. Calibration and precision of serum creatinine and plasma cystatin C measurement: impact on the estimation of glomerular filtration rate. *J Nephrol.* 2014;27:467–475. doi: 10.1007/s40620-014-0087-7
- Altay S, Onat A, Ozpamuk-Karadeniz F, Karadeniz Y, Kemaloglu-Oz T, Can G. Renal "hyperfiltrators" are at elevated risk of death and chronic diseases. BMC Nephrol. 2014;15:160. doi: 10.1186/1471-2369-15-160
- Inker LA, Eneanya ND, Coresh J, Tighiouart H, Wang D, Sang Y, Crews DC, Doria A, Estrella MM, Froissart M, et al. New creatinine- and cystatin c-based equations to estimate GFR without race. N Engl J Med. 2021;385:1737– 1749. doi: 10.1056/NEJMoa2102953
- 47. Hellwege JN, Velez Edwards DR, Giri A, Qiu C, Park J, Torstenson ES, Keaton JM, Wilson OD, Robinson-Cohen C, Chung CP, et al. Mapping eGFR loci to the renal transcriptome and phenome in the VA Million Veteran Program. *Nat Commun.* 2019;10:3842. doi: 10.1038/s41467-019-11704-w
- Hunter-Zinck H, Shi Y, Li M, Gorman BR, Ji S-G, Sun N, Webster T, Liem A, Hsieh P, Devineni P, et al. Genotyping array design and data quality control in the Million Veteran Program. Am J Hum Genet. 2020;106:535–548. doi: 10.1016/j.ajhg.2020.03.004
- Das S, Forer L, Schönherr S, Sidore C, Locke AE, Kwong A, Vrieze SI, Chew EY, Levy S, McGue M, et al. Next-generation genotype imputation service and methods. *Nat Genet*. 2016;48:1284–1287. doi: 10.1038/ng.3656
- Auton A, Brooks LD, Durbin RM, Garrison EP, Kang HM, Korbel JO, Marchini JL, McCarthy S, McVean GA, Abecasis GR. A global reference for human genetic variation. *Nature*. 2015;526:68–74. doi: 10.1038/nature15393
- Loh PR, Danecek P, Palamara PF, Fuchsberger C, Reshef YA, Finucane HK, Schoenherr S, Forer L, McCarthy S, Abecasis GR, et al. Reference-based phasing using the Haplotype Reference Consortium panel. *Nat Genet*. 2016;48:1443–1448. doi: 10.1038/ng.3679
- Fang H, Hui Q, Lynch J, Honerlaw J, Assimes TL, Huang J, Vujkovic M, Damrauer SM, Pyarajan S, Gaziano JM, et al. Harmonizing genetic ancestry and self-identified race/ethnicity in genome-wide association studies. *Am J Hum Genet*. 2019;105:763–772. doi: 10.1016/j.ajhg.2019.08.012
- Bycroft C, Freeman C, Petkova D, Band G, Elliott LT, Sharp K, Motyer A, Vukcevic D, Delaneau O, O'Connell J, et al. The UK Biobank resource with deep phenotyping and genomic data. *Nature*. 2018;562:203–209. doi: 10.1038/s41586-018-0579-z
- 54. Allen NE, Arnold M, Parish S, Hill M, Sheard S, Callen H, Fry D, Moffat S, Gordon M, Welsh S, et al. Approaches to minimising the epidemiological impact of sources of systematic and random variation that may affect biochemistry assay data in UK Biobank. Wellcome Open Res. 2020;5:222. doi: 10.12688/wellcomeopenres.16171.2
- McCarthy S, Das S, Kretzschmar W, Delaneau O, Wood AR, Teumer A, Kang HM, Fuchsberger C, Danecek P, Sharp K, et al. A reference panel of 64,976 haplotypes for genotype imputation. *Nat Genet*. 2016;48:1279–1283. doi: 10.1038/ng.3643
- Howie BN, Donnelly P, Marchini J. A flexible and accurate genotype imputation method for the next generation of genome-wide association studies. PLoS Genet. 2009;5:e1000529. doi: 10.1371/journal.pgen.1000529
- Inker LA, Schmid CH, Tighiouart H, Eckfeldt JH, Feldman HI, Greene T, Kusek JW, Manzi J, Van Lente F, Zhang YL, et al. Estimating glomerular filtration rate from serum creatinine and cystatin C. N Engl J Med. 2012;367:20–29. doi: 10.1056/NEJMoa1114248
- Burgess S, Ference BA, Staley JR, Freitag DF, Mason AM, Nielsen SF, Willeit P, Young R, Surendran P, Karthikeyan S, et al. Association of Ipa variants with risk of coronary disease and the implications for lipoprotein(a)-lowering therapies: a mendelian randomization analysis. *JAMA Cardiol*. 2018;3:619–627. doi: 10.1001/jamacardio.2018.1470
- Tian H, Mason AM, Liu C, Burgess S. Relaxing parametric assumptions for non-linear Mendelian randomization using a doubly-ranked stratification method [published online ahead of print June 28, 2022]. bioRxiv. doi: 10.1101/2022.06.28.497930