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Author manuscript *Nature*. Author manuscript; available in PMC 2012 May 01.

Published in final edited form as: *Nature*. ; 478(7367): 103–109. doi:10.1038/nature10405.

Genetic Variants in Novel Pathways Influence Blood Pressure and Cardiovascular Disease Risk

The International Consortium for Blood Pressure Genome-Wide Association Studies

Abstract

Blood pressure (BP) is a heritable trait¹ influenced by multiple biological pathways and is responsive to environmental stimuli. Over one billion people worldwide have hypertension (BP

140 mm Hg systolic [SBP] or 90 mm Hg diastolic [DBP])². Even small increments in BP are associated with increased risk of cardiovascular events³. This genome-wide association study of SBP and DBP, which used a multi-stage design in 200,000 individuals of European descent, identified 16 novel loci: six of these loci contain genes previously known or suspected to regulate BP (*GUCY1A3-GUCY1B3; NPR3-C5orf23; ADM; FURIN-FES; GOSR2; GNAS-EDN3*); the other 10 provide new clues to BP physiology. A genetic risk score based on 29 genome-wide significant variants was associated with hypertension, left ventricular wall thickness, stroke, and coronary artery disease, but not kidney disease or kidney function. We also observed associations with BP in East Asian, South Asian, and African ancestry individuals. Our findings provide new insights into the genetics and biology of BP, and suggest novel potential therapeutic pathways for cardiovascular disease prevention.

Genetic approaches have advanced the understanding of biological pathways underlying inter-individual variation in BP. For example, studies of rare Mendelian BP disorders have identified multiple defects in renal sodium handling pathways⁴. More recently two genome-wide association studies (GWAS), each of >25,000 individuals of European-ancestry, identified 13 loci associated with SBP, DBP, and hypertension^{5,6}. We now report results of a new meta-analysis of GWAS data that includes staged follow-up genotyping to identify additional BP loci.

Primary analyses evaluated associations between 2.5 million genotyped or imputed single nucleotide polymorphisms (SNPs) and SBP and DBP in 69,395 individuals of European ancestry from 29 studies (Supplementary Materials Sections 1–3, Supplementary Tables 1–2). Following GWAS meta-analysis, we conducted a three-stage validation experiment that made efficient use of available genotyping resources, to follow up top signals in up to 133,661 additional individuals of European descent (Supplementary Fig. 1 and Supplementary Materials Section 4). Twenty-nine independent SNPs at 28 loci were

Author contributions

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Note added in proof: Since this manuscript was submitted, Kato et al published a BP GWAS in East Asians that identified a SNP highly correlated to the SNP we report at the *NPR3-c5orf23* locus²⁸.

Full author contributions and roles are listed in the Supplementary Materials Section 19.

significantly associated with SBP, DBP, or both in the meta-analysis combining discovery and follow up data (Fig. 1, Table 1, Supplementary Figs 2–3, Supplementary Tables 3–5). All 29 SNPs attained association $P < 5 \times 10^{-9}$, an order of magnitude beyond the standard genome-wide significance level for a single stage experiment (Table 1).

Sixteen of these 29 associations were novel (Table 1). Two associations were near the *FURIN* and *GOSR2* genes; prior targeted analyses of variants in these genes suggested they may be BP loci^{7,8}. At the *CACNB2* locus we validated association for a previously reported⁶ SNP rs4373814 and detected a novel independent association for rs1813353 (pairwise r² =0.015 in HapMap CEU). Of our 13 previously reported associations^{5,6}, only the association at *PLCD3* was not supported by the current results (Supplementary Table 4). Some of the associations are in or near genes involved in pathways known to influence BP (*NPR3*, *GUCY1A3-GUCY1B3*, *ADM*, *GNAS-EDN3*, *NPPA-NPPB*, and *CYP17A1*; Supplementary Fig. 4). Twenty-two of the 28 loci did not contain genes that were *a priori* strong biological candidates.

As expected from prior BP GWAS results, the effects of the novel variants on SBP and DBP were small (Fig. 1 and Table 1). For all variants, the observed directions of effects were concordant for SBP, DBP, and hypertension (Fig. 1, Table 1, Supplementary Fig. 3). Among the genes at the genome-wide significant loci, only *CYP17A1*, previously implicated in Mendelian congenital adrenal hyperplasia and hypertension, is known to harbour rare variants that have large effects on BP⁹.

We performed several analyses to identify potential causal alleles and mechanisms. First, we looked up the 29 genome-wide significant index SNPs and their close proxies ($r^2>0.8$) among *cis*-acting expression SNP (eSNP) results from multiple tissues (Supplementary Materials Section 5). For 13/29 index SNPs, we found association between nearby eSNP variants and expression level of at least one gene transcript ($10^{-4} > p > 10^{-51}$, Supplementary Table 6). In 5 cases, the index BP SNP and the best eSNP from a genome-wide survey were identical, highlighting potential mediators of the SNP-BP associations.

Second, because changes in protein sequence are strong *a priori* candidates to be functional, we sought non-synonymous coding SNPs that were in high LD ($r^2 > 0.8$) with the 29 index SNPs. We identified such SNPsat 8 loci (Table 1, Supplementary Materials Section 6, Supplementary Table 7). In addition we performed analyses testing for differences in genetic effect according to body mass index (BMI) or sex, and analyses of copy number variants, pathway enrichment, and metabolomic data, but we did not find any statistically significant results (Supplementary Materials Sections 7–9, Supplementary Tables 8–10).

We evaluated whether the BP variants we identified in Europeans were associated with BP in individuals of East Asian (N=29,719), South Asian (N=23,977), and African (N=19,775) ancestries (Table 1, Supplementary Tables 11–13). We found significant associations in individuals of East Asian ancestry for SNPs at 9 loci and in individuals of South Asian ancestry for SNPs at 6 loci; some have been reported previously (Supplementary Tables 12 and 15). The lack of significant association for individual SNPs may reflect small sample sizes, differences in allele frequencies or LD patterns, imprecise imputation for some

ancestries using existing reference samples, or a genuinely different underlying genetic architecture. Because of limited power to detect effects of individual variants in the smaller non-European samples, we created genetic risk scores for SBP and DBP incorporating all 29 BP variants weighted according to effect sizes observed in the European samples. In each non-European ancestry group, risk scores were strongly associated with SBP ($P=1.1\times10^{-40}$ in East Asian, $P=2.9\times10^{-13}$ in South Asian, $P=9.8\times10^{-4}$ in African ancestry individuals) and DBP ($P=2.9\times10^{-48}$, $P=9.5\times10^{-15}$, and $P=5.3\times10^{-5}$, respectively; Supplementary Table 13).

We also created a genetic risk score to assess association of the variants in aggregate with hypertension and with clinical measures of hypertensive complications including left ventricular mass, left ventricular wall thickness, incident heart failure, incident and prevalent stroke, prevalent coronary artery disease (CAD), kidney disease, and measures of kidney function, using results from other GWAS consortia (Table 2, Supplementary Materials Sections 10-11, Supplementary Table 14). The risk score was weighted using the average of SBP and DBP effects for the 29 SNPs. In an independent sample of 23,294 women¹⁰, an increase of 1 standard deviation in the genetic risk score was associated with a 21% increase in the odds of hypertension (95% CI 19%-28%; Table 2, Supplementary Table 14). Among individuals in the top decile of the risk score, the prevalence of hypertension was 29% compared with 16% in the bottom decile (odds ratio 2.09, 95% CI 1.86-2.36). Similar results were observed in an independent hypertension case-control sample (Table 2). In our study, individuals in the top compared to bottom quintiles of genetic risk score differed by 4.6 mm Hg SBP and 3.0 mm Hg DBP, differences that approach population-averaged BP treatment effects for a single antihypertensive agent¹¹. Epidemiologic data have shown that differences in SBP and DBP of this magnitude, across the population range of BP, are associated with an increase in cardiovascular disease risk³. Consistent with this and in line with findings from randomized trials of BP-lowering medication in hypertensive patients^{12,13}, the genetic risk score was positively associated with left ventricular wall thickness ($P=6.0 \times 10^{-6}$), occurrence of stroke ($P=3.3 \times 10^{-5}$) and CAD ($P=8.1 \times 10^{-29}$). The same genetic risk score was not, however, significantly associated with chronic kidney disease or measures of kidney function, even though these renal outcomes were available in a similar sample size as for the other outcomes (Table 2). The absence of association with kidney phenotypes could be explained by a weaker causal relation of BP with kidney phenotypes than with CAD and stroke. This finding is consistent with the mismatch between observational data that show a positive association of BP with kidney disease, and clinical trial data that show inconsistent evidence of benefit of BP lowering on kidney disease prevention in patients with hypertension¹⁴. Thus, several lines of evidence converge to suggest that BP elevation may in part be a consequence rather than a cause of sub-clinical kidney disease.

Our discovery meta-analysis (Supplementary Fig. 2) suggests an excess of modestly significant $(10^{-5} < P < 10^{-2})$ associations likely arising from common BP variants of small effect. By dividing our principal GWAS dataset into non-overlapping discovery (N \approx 56,000) and validation (N \approx 14,000) subsets, we found robust evidence for the existence of such undetected common variants (Supplementary Fig. 5, Supplementary Materials Section 12).

Nature. Author manuscript; available in PMC 2012 May 01.

We estimate¹⁵ that there are 116 (95% CI 57–174) independent BP variants with effect sizes similar to those reported here, which collectively explain \approx 2.2% of the phenotypic variance for SBP and DBP, compared with 0.9% explained by the 29 associations discovered thus far (Supplementary Fig. 6, Supplementary Materials Section 13).

Most of the 28 BP loci harbour multiple genes (Supplementary Table 15, Supplementary Fig. 4), and although substantial research is required to identify the specific genes and variants responsible for these associations, several loci contain highly plausible biological candidates. The *NPPA* and *NPPB* genes at the *MTHFR-NPPB* locus encode precursors for atrial- and B-type natriuretic peptides (ANP, BNP), and previous work has identified SNPs, modestly correlated with our index SNP at this locus, that are associated with plasma ANP, BNP, and BP¹⁶. We found the index SNP at this locus was associated with opposite effects on BP and on ANP/BNP levels, consistent with a model in which the variants act through increased ANP/BNP production to lower BP¹⁶ (Supplementary Materials Section 14).

Two other loci identified in the current study harbour genes involved in natriuretic peptide and related nitric oxide signalling pathways,^{17,18} both of which act to regulate cyclic guanosine monophosphate (cGMP). The first locus contains *NPR3*, which encodes the natriuretic peptide clearance receptor (NPR-C). *NPR3* knockout mice exhibit reduced clearance of circulating natriuretic peptides and lower BP¹⁹. The second locus includes *GUCY1A3* and *GUCY1B3*, encoding the alpha and beta subunits of soluble guanylatecyclase (sGC); knockout of either gene in murine models results in hypertension²⁰.

Another locus contains *ADM*, encoding adrenomedullin, which has natriuretic, vasodilatory, and BP-lowering properties²¹. At the *GNAS-EDN3* locus, *ZNF831* is closest to the index SNP, but *GNAS* and *EDN3* are two nearby compelling biological candidates (Supplementary Fig. 4, Supplementary Table 15).

We identified two loci with plausible connections to BP via genes implicated in renal physiology or kidney disease. At the first locus, *SLC4A7* is an electro-neutral sodium bicarbonate co-transporter expressed in the nephron and in vascular smooth muscle²². At the second locus, *PLCE1* (phospholipase-C-epsilon-1 isoform) is important for normal podocyte development in the glomerulus; sequence variation in *PLCE1* has been implicated in familial nephrotic syndromes and end-stage kidney disease²³.

Missense variants in two genes involved in metal ion transport were associated with BP in our study. The first encodes a His/Asp change at amino acid 63 (*H63D*) in *HFE* and is a low penetrance allele for hereditary hemochromatosis²⁴. The second is an Ala/Thr polymorphism located in exon 7 of *SLC39A8*, which encodes a zinc transporter that also transports cadmium and manganese²⁵. The same allele of *SLC39A8* associated with BP in our study has recently been associated with high-density lipoprotein (HDL) cholesterol levels²⁶ and BMI²⁷ (Supplementary Table 15).

In conclusion, we have shown that 29 independent genetic variants influence BP in people of European ancestry. The variants reside in 28 loci, 16 of which were novel, and we confirmed association of several of them in individuals of non-European ancestry. A risk score derived from the 29 variants was significantly associated with BP-related organ

Nature. Author manuscript; available in PMC 2012 May 01.

damage and clinical cardiovascular disease, but not kidney disease. These loci improve our understanding of the genetic architecture of BP, provide new biological insights into BP control and may identify novel targets for the treatment of hypertension and the prevention of cardiovascular disease.

Methods summary

Supplementary Materials provide complete methods and include the following sections: study recruitment and phenotyping, adjustment for antihypertensive medications, genotyping, data quality control, genotype imputation, within-cohort association analyses, meta-analyses of discovery and validation stages, stratified analyses by sex and BMI, identification of eSNPs and nsSNPs, metabolomic and lipidomic analyses, CNV analyses, pathway analyses, analyses for non-European ancestries, association of a risk score with hypertension and cardiovascular disease, estimation of numbers of undiscovered variants, measurement of natriuretic peptides, and brief literature reviews and GWAS database lookups of all validated BP loci.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

A number of the participating studies and authors are members of the CHARGE and Global BPgen consortia. Many funding mechanisms by NIH/NHLBI, European, and private funding agencies contributed to this work and a full list is provided in Section 21 of the Supplementary Materials.

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Authors

Georg B. Ehret^{1,2,3*}, Patricia B. Munroe^{4*#}, Kenneth M. Rice^{5*}, Murielle Bochud^{2*}, Andrew D. Johnson^{6,7*}, Daniel I. Chasman^{8,9*}, Albert V. Smith^{10,11*}, Martin D. Tobin¹², Germaine C. Verwoert^{13,14,15}, Shih-Jen Hwang^{6,16,7}, Vasyl Pihur¹, Peter Vollenweider¹⁷, Paul F. O'Reilly¹⁸, Najaf Amin¹³, Jennifer L Bragg-Gresham¹⁹, Alexander Teumer²⁰, Nicole L. Glazer²¹, Lenore Launer²², Jing Hua Zhao²³, Yurii Aulchenko¹³, Simon Heath²⁴, Siim Sõber²⁵, Afshin Parsa²⁶, Jian'an Luan²³, Pankaj Arora²⁷, Abbas Dehghan¹³, ¹⁴, ¹⁵, Feng Zhang²⁸, Gavin Lucas²⁹, Andrew A. Hicks³⁰, Anne U. Jackson³¹, John F Peden³², Toshiko Tanaka³³, Sarah H. Wild³⁴, Igor Rudan^{35,36}, Wilmar Igl³⁷, Yuri Milaneschi³³, Alex N. Parker³⁸, Cristiano Fava³⁹, ⁴⁰, John C. Chambers¹⁸, ⁴¹, Ervin R.

p.b.munroe@qmul.ac.uk; cnewtoncheh@partners.org

- ⁶Framingham Heart Study, Framingham, MA, USA
- ⁷National Heart Lung, and Blood Institute, Bethesda, MD, USA
- ⁸Division of Preventive Medicine, Brigham and Women's Hospital, 900 Commonwealth Avenue East, Boston MA 02215, USA
- ⁹Harvard Medical School, Boston, MA, USA
- ¹⁰Icelandic Heart Association, Kopavogur, Iceland
- ¹¹University of Iceland, Reykajvik, Iceland
- ¹²Department of Health Sciences, University of Leicester, University Rd, Leicester LE1 7RH, UK
- ¹³Department of Epidemiology, Erasmus Medical Center, PO Box 2040, 3000 CA, Rotterdam, The Netherlands
- ¹⁴Department of Internal Medicine, Erasmus Medical Center, Rotterdam, The Netherlands
- ¹⁵Netherlands Consortium for Healthy Aging (NCHA), Netherland Genome Initiative (NGI), The Netherlands
- ¹⁶Center for Population Studies, National Heart Lung, and Blood Institute, Bethesda, MD, USA
- ¹⁷Department of Internal Medicine, Centre Hospitalier Universitaire Vaudois, 1011 Lausanne, Switzerland

¹⁸Department of Epidemiology and Biostatistics, School of Public Health, Imperial College London, Norfolk Place, London W2 1PG,

UK ¹⁹Center forStatistical Genetics, Department of Biostatistics, University of Michigan School of Public Health, Ann Arbor, MI 48103, USA ²⁰Interfaculty Institute for Genetics and Functional Genomics, Ernst-Moritz-Arndt-University Greifswald, 17487 Greifswald,

- ²³MRC Epidemiology Unit, Institute of Metabolic Science, Cambridge CB2 0QQ, UK
- ²⁴Centre National de Génotypage, Commissariat à L'Energie Atomique, Institut de Génomique, Evry, France
- ²⁵Institute of Molecular and Cell Biology, University of Tartu, Riia 23, Tartu 51010, Estonia
- ²⁶University of Maryland School of Medicine, Baltimore, MD, USA, 21201, USA

²⁷Center for Human Genetic Research, Cardiovascular Research Center, Massachusetts General Hospital, Boston, Massachusetts, 02114, USA 28Department of Twin Research & Genetic Epidemiology, King's College London, UK

²⁹Cardiovascular Epidemiology and Genetics, Institut Municipal d'Investigacio Medica, Barcelona Biomedical Research Park, 88 Doctor Aiguader, 08003 Barcelona, Spain ³⁰Institute of Genetic Medicine, European Academy Bozen/Bolzano (EURAC), Viale Druso 1, 39100 Bolzano, Italy -Affiliated

Institute of the University of Lübeck, Germany ³¹Department of Biostatistics, Center for Statistical Genetics, University of Michigan, Ann Arbor, Michigan, 48109, USA

³²Department of Cardiovascular Medicine, The Wellcome Trust Centre for Human Genetics, University of Oxford, OXford, OX3 7BN, UK

- ³³Clinical Research Branch, National Institute on Aging, Baltimore MD 21250, USA
- ³⁴Centre for Population Health Sciences, University of Edinburgh, EH89AG, UK

³⁵Centre for Population Health Sciences and Institute of Genetics and Molecular Medicine, College of Medicine and Vet Medicine, University of Edinburgh, EH8 9AG, UK

³⁶Croatian Centre for Global Health, University of Split, Croatia

³⁷Department of Genetics and Pathology, Rudbeck Laboratory, Uppsala University, SE-751 85 Uppsala, Sweden ³⁸Amgen, 1 Kendall Square, Building 100, Cambridge, MA 02139, USA

- ³⁹Department of Clinical Sciences, Lund University, Malmö, Sweden
- ⁴⁰Department of Medicine, University of Verona, Italy
- ⁴¹Ealing Hospital, London, UB1 3HJ, UK

¹Center for Complex Disease Genomics, McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University School of Medicine, Baltimore, MD 21205, USA

²Institute of Social and Preventive Medicine (IUMSP), Centre Hospitalier Universitaire Vaudois and University of Lausanne, Bugnon 17, 1005 Lausanne, Switzerland

³Cardiology, Department of Specialties of Internal Medicine, Geneva University Hospital, Rue Gabrielle-Perret-Gentil 4, 1211 Geneva 14, Switzerland

contributed equally

⁴Clinical Pharmacology and The Genome Centre, William Harvey Research Institute, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London EC1M 6BQ, UK

[#]to whom correspondence should be addressed: aravinda@jhmi.edu; m.j.caulfield@qmul.ac.uk; levyd@nhlbi.nih.gov;

Department of Biostatistics, University of Washington, Seattle, WA, USA

Germany ²¹Cardiovascular Health Research Unit, Departments of Medicine, Epidemiology and Health Services, University of Washington,

Seattle, WA, USA ²²Laboratory of Epidemiology, Demography, Biometry, National Institute on Aging, National Institutes of Health, Bethesda,

Fox⁴², Meena Kumari⁴³, Min Jin Go⁴⁴, Pim van der Harst⁴⁵, Wen Hong Linda Kao⁴⁶, Marketa Sjögren³⁹, D. G. Vinay⁴⁷, Myriam Alexander⁴⁸, Yasuharu Tabara⁴⁹, Sue Shaw-Hawkins⁴, Peter H. Whincup⁵⁰, Yongmei Liu⁵¹, Gang Shi⁵², Johanna Kuusisto⁵³, Bamidele Tayo⁵⁴, Mark Seielstad⁵⁵, ⁵⁶, Xueling Sim⁵⁷, Khanh-Dung Hoang Nguyen¹, Terho Lehtimäki⁵⁸, Giuseppe Matullo⁵⁹,⁶⁰, Ying Wu⁶¹, Tom R. Gaunt⁶², N. Charlotte Onland-Moret⁶³, ⁶⁴, Matthew N. Cooper⁶⁵, Carl G.P. Platou⁶⁶, Elin Org²⁵, Rebecca Hardy⁶⁷, Santosh Dahgam⁶⁸, Jutta Palmen⁶⁹, Veronique Vitart⁷⁰, Peter S. Braund⁷¹,⁷², Tatiana Kuznetsova⁷³, Cuno S.P.M. Uiterwaal⁶³, Adebowale Adeyemo⁷⁴, Walter Palmas⁷⁵, Harry Campbell³⁵, Barbara Ludwig⁷⁶, Maciej Tomaszewski⁷¹,⁷², Ioanna Tzoulaki^{77,78}, Nicholette D. Palmer⁷⁹, CARDIoGRAM consortium⁸⁰, CKDGen

⁴³Genetic Epidemiology Group, Epidemiology and Public Health, UCL, London, WC1E 6BT, UK

- ⁵¹Epidemiology & Prevention, Division of Public Health Sciences, Wake Forest University School of Medicine, Winston-Salem, NC 27157, USA ⁵² ⁵²Division of Biostatistics and Department of Genetics, School of Medicine, Washington University in St. Louis, Saint Louis,
- Missouri 63110, USA
- 53 Department of Medicine, University of Eastern Finland and Kuopio University Hospital, 70210 Kuopio, Finland
- ⁵⁴Department of Preventive Medicine and Epidemiology, Loyola University Medical School, Maywood, IL, USA
- ⁵⁵Department of Laboratory Medicine & Institute of Human Genetics, University of California San Francisco, 513 Parnassus Ave. San Francisco CA 94143, USA ⁵⁶Genome Institute of Singapore, Agency for Science, Technology and Research, Singapore, 138672, Singapore
- ⁵⁷Centre for Molecular Epidemiology, Yong Loo Lin School of Medicine, National University of Singapore, Singapore, 117597, Singapore ⁵⁸Department of Clinical Chemistry, University of Tampere and Tampere University Hospital, Tampere, 33521, Finland
- ⁵⁹Department of Genetics, Biology and Biochemistry, University of Torino, Via Santena 19, 10126, Torino, Italy
- ⁶⁰Human Genetics Foundation (HUGEF), Via Nizza 52, 10126, Torino, Italy
- ⁶¹Department of Genetics, University of North Carolina, Chapel Hill, NC, 27599, USA
- ⁶²MRC Centre for Causal Analyses in Translational Epidemiology, School of Social & Community Medicine, University of Bristol, Bristol BS8 2BN, UK
- ⁶³Julius Center for Health Sciences and Primary Care, University Medical Center Utrecht, Heidelberglaan 100, 3508 GA Utrecht, The Netherlands

⁶⁵Centre for Genetic Epidemiology and Biostatistics, University of Western Australia, Crawley, WA, Australia

⁶⁶HUNT Research Centre, Department of Public Health and General Practice, Norwegian University of Science and Technology, 7600 Levanger, Norway

- Sahlgrenska Academy, University of Gothenburg, 40530 Gothenburg, Sweden ⁶⁹Centre for Cardiovascular Genetics, University College London, London WC1E 6JF, UK
- ⁷⁰MRC Human Genetics Unit and Institute of Genetics and Molecular Medicine, Edinburgh, EH2, UK
- ⁷¹Department of Cardiovascular Sciences, University of Leicester, Glenfield Hospital, Leicester, LE3 9QP, UK
- ⁷²Leicester NIHR Biomedical Research Unit in Cardiovascular Disease, Glenfield Hospital, Leicester, LE3 9QP, UK

⁷³Studies Coordinating Centre, Division of Hypertension and Cardiac Rehabilitation, Department of Cardiovascular Diseases,

University of Leuven, Campus Sint Rafaël, Kapucijnenvoer 35, Block D, Box 7001, 3000 Leuven, Belgium ⁷⁴Center for Research on Genomics and Global Health, National Human Genome Research Institute, Bethesda, MD 20892, USA ⁷⁵Columbia University, NY, USA

⁷⁶Department of Medicine III, Medical Faculty Carl Gustav Carus at the Technical University of Dresden, 01307 Dresden, Germany ⁷⁷Epidemiology and Biostatistics, School of Public Health, Imperial College, London, W2 1PG, UK

⁴²Department of Medicine, University of Mississippi Medical Center, USA

⁴⁴Center for Genome Science, National Institute of Health, Seoul, Korea

⁴⁵Department of Cardiology, University Medical Center Groningen, University of Groningen, The Netherlands

⁴⁶Departments of Epidemiology and Medicine, Johns Hopkins University, Baltimore MD, USA

⁴⁷Centre for Cellular and Molecular Biology (CCMB), Council of Scientific and Industrial Research (CSIR), Uppal Road, Hyderabad 500 007, India ⁴⁸Department of Public Health and Primary Care, University of Cambridge, CB1 8RN, UK

⁴⁹Department of Basic Medical Research and Education, and Department of Geriatric Medicine, Ehime University Graduate School of Medicine, Toon, 791-0295, Japan ⁵⁰Division of Community Health Sciences, St George's University of London, London, SW17 0RE, UK

⁶⁴Complex Genetics Section, Department of Medical Genetics -DBG, University Medical Center Utrecht, 3508 GA Utrecht, The Netherlands

⁶⁷MRC Unit for Lifelong Health & Ageing, London, WC1B 5JU, UK

⁶⁸Occupational and Environmental Medicine, Department of Public Health and Community Medicine, Institute of Medicine,

⁷⁸Clinical and Molecular Epidemiology Unit, Department of Hygiene and Epidemiology, University of Ioannina School of Medicine, Ioannina, Greece

⁹Wake Forest University Health Sciences, Winston-Salem, NC 27157, USA

⁸⁰A list of consortium members is supplied in the Supplementary Materials

Page 9

Consortium⁸⁰, KidneyGen Consortium⁸⁰, EchoGen consortium⁸⁰, CHARGE-HF consortium⁸⁰, Thor Aspelund¹⁰,¹¹, Melissa Garcia²², Yen-Pei C. Chang²⁶, Jeffrey R. O'Connell²⁶, Nanette I, Steinle²⁶, Diederick E, Grobbee⁶³, Dan E, Arking¹, Sharon L, Kardia⁸¹, Alanna C. Morrison⁸², Dena Hernandez⁸³, Samer Najjar⁸⁴, ⁸⁵, Wendy L. McArdle⁸⁶, David Hadley⁵⁰,⁸⁷, Morris J. Brown⁸⁸, John M. Connell⁸⁹, Aroon D. Hingorani⁹⁰, Ian N.M. Day⁶², Debbie A. Lawlor⁶², John P. Beilby^{91,92}, Robert W. Lawrence⁶⁵, Robert Clarke⁹³, Rory Collins⁹³, Jemma C Hopewell⁹³, Halit Ongen³², Albert W. Dreisbach⁴², Yali Li⁹⁴, J H. Young⁹⁵, Joshua C. Bis²¹, Mika Kähönen⁹⁶, Jorma Viikari⁹⁷, Linda S. Adair⁹⁸, Nanette R. Lee⁹⁹, Ming-Huei Chen¹⁰⁰, Matthias Olden¹⁰¹,¹⁰², Cristian Pattaro³⁰, Judith A. Hoffman Bolton¹⁰³, Anna Köttgen¹⁰⁴,¹⁰³, Sven Bergmann¹⁰⁵, ¹⁰⁶, Vincent Mooser¹⁰⁷, Nish Chaturvedi¹⁰⁸, Timothy M. Frayling¹⁰⁹, Muhammad Islam¹¹⁰, Tazeen H. Jafar¹¹⁰, Jeanette Erdmann¹¹¹, Smita R. Kulkarni¹¹², Stefan R. Bornstein⁷⁶, Jürgen Grässler⁷⁶, Leif Groop¹¹³,¹¹⁴, Benjamin F. Voight¹¹⁵, Johannes Kettunen¹¹⁶,¹²⁶, Philip Howard¹¹⁷, Andrew Taylor⁴³, Simonetta Guarrera⁶⁰, Fulvio Ricceri⁵⁹, ⁶⁰, Valur Emilsson¹¹⁸, Andrew Plump¹¹⁸, Inês Barroso¹¹⁹, ¹²⁰, Kay-Tee

- ⁸⁷Pediatric Epidemiology Center, University of South Florida, Tampa, FL, USA
- ⁸⁸Clinical Pharmacology Unit, University of Cambridge, Addenbrookes Hospital, Hills Road, Cambridge CB2 2QQ, UK
- ⁸⁹University of Dundee, Ninewells Hospital & Medical School, Dundee, DD1 9SY, UK
- ⁹⁰Genetic Epidemiology Group, Department of Epidemiology and Public Health, UCL, London WC1E 6BT, UK
- ⁹¹Pathology and Laboratory Medicine, University of Western Australia, Crawley, WA, Australia
- ⁹²Molecular Genetics, PathWest Laboratory Medicine, Nedlands, WA, Australia
- ⁹³Clinical Trial Service Unit and Epidemiological Studies Unit, University of Oxford, Oxford, OX3 7LF, UK
- ⁹⁴Department of Epidemiology and Biostatistics, Case Western Reserve University, 2103 Cornell Road, Cleveland, OH 44106, USA 95 Department of Medicine, Johns Hopkins University, Baltimore, USA
 - ⁹⁶Department of Clinical Physiology, University of Tampere and Tampere University Hospital, Tampere, 33521, Finland

- ⁹⁹Office of Population Studies Foundation, University of San Carlos, Talamban, Cebu City 6000, Philippines
- ¹⁰⁰Department of Neurology and Framingham Heart Study, Boston University School of Medicine, Boston, MA, 02118, USA
- ¹⁰¹Department of Internal Medicine II, University Medical Center Regensburg, 93053 Regensburg, Germany
- 102 Department of Epidemiology and Preventive Medicine, University Medical Center Regensburg, 93053 Regensburg, Germany ¹⁰³Department of Epidemiology, Johns Hopkins University, Baltimore MD, USA
- ¹⁰⁴Renal Division, University Hospital Freiburg, Germany
- ¹⁰⁵Département de Génétique Médicale, Université de Lausanne, 1015 Lausanne, Switzerland
- 106 Swiss Institute of Bioinformatics, 1015 Lausanne, Switzerland
- ¹⁰⁷Division of Genetics, GlaxoSmithKline, Philadelphia, Pennsylvania 19101, USA
- 108 International Centre for Circulatory Health, National Heart & Lung Institute, Imperial College, London, UK

- ¹¹⁰Department of Community Health Sciences & Department of Medicine, Aga Khan University, Karachi, Pakistan
- ¹¹¹Medizinische Klinik II, Universität zu Lübeck, Lübeck, Germany
- ¹¹²Diabetes Unit, KEM Hospital and Research Centre, Rasta Peth, Pune-411011, Maharashtra, India
- ¹¹³Department of Clinical Sciences, Diabetes and Endocrinology Research Unit, University Hospital, Malmö, Sweden ¹¹⁴Lund University, Malmö 20502, Sweden

- ¹¹⁶Department of Chronic Disease Prevention, National Institute for Health and Welfare, FIN-00251 Helsinki, Finland
- 126FIMM, Institute for Molecular Medicine, Finland, Biomedicum, P.O. Box 104, 00251 Helsinki, Finland

- ¹¹⁹Wellcome Trust Sanger Institute, Hinxton, CB10 1SA, UK

¹²⁰University of Cambridge Metabolic Research Labs, Institute of Metabolic Science Addenbrooke's Hospital, CB2 OQQ, Cambridge, UK

⁸¹Department of Epidemiology, School of Public Health, University of Michigan, Ann Arbor, MI 48109, USA

⁸²Division of Epidemiology, Human Genetics and Environmental Sciences, School of Public Health, University of Texas at Houston Health Science Center, 12 Herman Pressler, Suite 453E, Houston, TX 77030, USA

⁸³Laboratory of Neurogenetics, National Institute on Aging, Bethesda, MD 20892, USA

⁸⁴Laboratory of Cardiovascular Science, Intramural Research Program, National Institute on Aging, NIH, Baltimore, Maryland, USA ⁸⁵Washington Hospital Center, Division of Cardiology, Washington DC, USA

⁸⁶ALSPAC Laboratory, University of Bristol, Bristol, BS8 2BN, UK

⁹⁷Department of Medicine, University of Turku and Turku University Hospital, Turku, 20521, Finland

⁹⁸Department of Nutrition, University of North Carolina, Chapel Hill, NC, 27599, USA

¹⁰⁹Genetics of Complex Traits, Peninsula Medical School, University of Exeter, UK

¹¹⁵Program in Medical and Population Genetics, Broad Institute of Harvard and MIT, Cambridge, Massachusetts, 02139, USA

¹¹⁷ William Harvey Research Institute, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London EC1M 6BQ, UK ¹¹⁸Merck Research Laboratory, 126 East Lincoln Avenue, Rahway, NJ 07065, USA

Khaw⁴⁸, Alan B. Weder¹²¹, Steven C. Hunt¹²², Yan V. Sun⁸¹, Richard N. Bergman¹²³, Francis S. Collins¹²⁴, Lori L. Bonnycastle¹²⁴, Laura J. Scott³¹, Heather M. Stringham³¹, Leena Peltonen¹¹⁹, ¹²⁵, ¹²⁶, ¹²⁷, Markus Perola¹²⁵, Erkki Vartiainen¹²⁵, Stefan-Martin Brand¹²⁸,¹²⁹, Jan A. Staessen⁷³, Thomas J. Wang⁶,¹³⁰, Paul R. Burton¹²,⁷², Maria Soler Artigas¹², Yanbin Dong¹³¹, Harold Snieder¹³², ¹³¹, Xiaoling Wang¹³¹, Haidong Zhu¹³¹, Kurt K. Lohman¹³³, Megan E. Rudock⁵¹, Susan R Heckbert¹³⁴,¹³⁵, Nicholas L Smith¹³⁴,¹³⁶,¹³⁵, Kerri L Wiggins¹³⁷, Ayo Doumatey⁷⁴, Daniel Shriner⁷⁴, Gudrun Veldre²⁵,¹³⁸, Margus Viigimaa¹³⁹,¹⁴⁰, Sanjay Kinra¹⁴¹, Dorairajan Prabhakaran¹⁴², Vikal Tripathy¹⁴², Carl D. Langefeld⁷⁹, Annika Rosengren¹⁴³, Dag S. Thelle¹⁴⁴, Anna Maria Corsi¹⁴⁵, Andrew Singleton⁸³, Terrence Forrester¹⁴⁶, Gina Hilton¹, Colin A. McKenzie¹⁴⁶, Tunde Salako¹⁴⁷, Naoharu Iwai¹⁴⁸, Yoshikuni Kita¹⁴⁹, Toshio Ogihara¹⁵⁰, Takayoshi Ohkubo¹⁴⁹,¹⁵¹, Tomonori Okamura¹⁴⁸, Hirotsugu Ueshima¹⁵², Satoshi Umemura¹⁵³, Susana Eyheramendy¹⁵⁴, Thomas Meitinger¹⁵⁵, ¹⁵⁶, H.-Erich Wichmann¹⁵⁷, ¹⁵⁸, ¹⁵⁹, Yoon

127Broad Institute, Cambridge, Massachusetts 02142, USA

¹²¹Division of Cardiovascular Medicine, Department of Internal Medicine, University of Michigan Medical School, Ann Arbor, MI, USA 122Cardiovascular Genetics, University of Utah School of Medicine, Salt Lake City, UT, USA

¹²³Department of Physiology and Biophysics, Keck School of Medicine, University of Southern California, Los Angeles, California 90033, USA ¹²⁴National Human Genome Research Institute, National Institutes of Health, Bethesda, Maryland 20892, USA

¹²⁵National Institute for Health and Welfare, 00271 Helsinki, Finland

¹²⁸Leibniz-Institute for Arteriosclerosis Research, Department of Molecular Genetics of Cardiovascular Disease, University of Münster, Münster, Germany ¹²⁹Medical Faculty of the Westfalian Wilhelms University Muenster, Department of Molecular Genetics of Cardiovascular Disease,

University of Muenster, Muenster, Germany ¹³⁰Division of Cardiology, Massachusetts General Hospital, Boston, MA, USA

¹³¹Georgia Prevention Institute, Department of Pediatrics, Medical Collegeof Georgia, Augusta, GA, USA

¹³²Unit of Genetic Epidemiology and Bioinformatics, Department of Epidemiology, University Medical Center Groningen, University of Groningen, Groningen, The Netherlands ¹³³Department of Biostatical Sciences, Division of Public Health Sciences, Wake Forest University School of Medicine, Winston-

Salem, NC 27157, USA

Department of Epidemiology, University of Washington, Seattle, WA, 98195, USA

¹³⁵ Group Health Research Institute, Group Health Cooperative, Seattle, WA, USA ¹³⁶Seattle Epidemiologic Research and Information Center, Veterans Health Administration Office of Research & Development,

Seattle, WA 98108, USA ¹³⁷Department of Medicine, University of Washington, 98195, USA

¹³⁸Department of Cardiology, University of Tartu, L. Puusepa 8, 51014 Tartu, Estonia

¹³⁹Tallinn University of Technology, Institute of Biomedical Engineering, Ehitajate tee 5, 19086 Tallinn, Estonia

¹⁴⁰Centre of Cardiology, North Estonia Medical Centre, Sütiste tee 19, 13419 Tallinn, Estonia

¹⁴¹Division of Non-communicable disease Epidemiology, The London School of Hygiene and Tropical Medicine London, Keppel Street, London WC1E 7HT, UK

²South Asia Network for Chronic Disease, Public Health Foundation of India, C-1/52, SDA, New Delhi 100016, India

¹⁴³Department of Emergency and Cardiovascular Medicine, Institute of Medicine, Sahlgrenska Academy, University of Gothenburg, 41685 Gothenburg, Sweden

¹⁴⁴Department of Biostatistics, Institute of Basic Medical Sciences, University of Oslo, 0317 Oslo, Norway

¹⁴⁵ Tuscany Regional Health Agency, Florence, Italy

¹⁴⁶Tropical Medicine Research Institute, University of the West Indies, Mona, Kingston, Jamaica

¹⁴⁷University of Ibadan, Ibadan, Nigeria

¹⁴⁸Department of Genomic Medicine, and Department of Preventive Cardiology, National Cerebral and Cardiovascular Research Center, Suita, 565-8565, Japan 149Department of Health Science, Shiga University of Medical Science, Otsu, 520-2192, Japan

¹⁵⁰Department of Geriatric Medicine, Osaka University Graduate School of Medicine, Suita, 565-0871, Japan

¹⁵¹Tohoku University Graduate School of Pharmaceutical Sciences and Medicine, Sendai, 980-8578, Japan

¹⁵²Lifestyle-related Disease Prevention Center, Shiga University of Medical Science, Otsu, 520-2192, Japan

¹⁵³Department of Medical Science and Cardiorenal Medicine, Yokohama City University School of Medicine, Yokohama, 236-0004, Japan 154Department of Statistics, Pontificia Universidad Catolicade Chile, Vicuña Mackena 4860, Santiago, Chile

¹⁵⁵ Institute of Human Genetics, Helmholtz Zentrum Munich, German Research Centre for Environmental Health, 85764 Neuherberg, Germany ¹⁵⁶Institute of Human Genetics, Klinikum rechts der Isar, Technical University of Munich, 81675 Munich, Germany

Shin Cho⁴⁴, Hyung-Lae Kim⁴⁴, Jong-Young Lee⁴⁴, James Scott¹⁶⁰, Joban S. Sehmi¹⁶⁰,⁴¹, Weihua Zhang¹⁸, Bo Hedblad³⁹, Peter Nilsson³⁹, George Davey Smith⁶², Andrew Wong⁶⁷, Narisu Narisu¹²⁴, Alena Stan áková⁵³, Leslie J. Raffel¹⁶¹, Jie Yao¹⁶¹, Sekar Kathiresan¹⁶²,²⁷, Chris O'Donnell¹⁶³,²⁷,⁹, Stephen M. Schwartz¹³⁴, M. Arfan Ikram¹³,¹⁵, W. T. Longstreth Jr.¹⁶⁴, Thomas H. Mosley¹⁶⁵, Sudha Seshadri¹⁶⁶, Nick R.G. Shrine¹², Louise V. Wain¹², Mario A. Morken¹²⁴, Amy J. Swift¹²⁴, Jaana Laitinen¹⁶⁷, Inga Prokopenko⁵¹,¹⁶⁸, Paavo Zitting¹⁶⁹, Jackie A. Cooper⁶⁹, Steve E. Humphries⁶⁹, John Danesh⁴⁸, Asif Rasheed¹⁷⁰, Anuj Goel³², Anders Hamsten¹⁷¹, Hugh Watkins³², Stephan J.L. Bakker¹⁷², Wiek H. van Gilst⁴⁵, Charles S. Janipalli⁴⁷, K. Radha Mani⁴⁷, Chittaranjan S. Yajnik¹¹², Albert Hofman¹³, Francesco U.S. Mattace-Raso¹³, ¹⁴, Ben A. Oostra¹⁷³, Ayse Demirkan¹³, Aaron Isaacs¹³, Fernando Rivadeneira¹³, ¹⁴, Edward G Lakatta¹⁷⁴, Marco Orru¹⁷⁵,¹⁷⁶, Angelo Scuteri¹⁷⁴, Mika Ala-Korpela¹⁷⁷,¹⁷⁸,¹⁷⁹, Antti J Kangas¹⁷⁷, Leo-Pekka Lyytikäinen⁵⁸, Pasi Soininen¹⁷⁷, ¹⁷⁸, Taru Tukiainen¹⁸⁰, ¹⁸¹, ¹⁷⁷, Peter Würtz¹⁷⁷, ¹⁸, ¹⁸⁰, Rick Twee-Hee Ong⁵⁶, ⁵⁷, ¹⁸², Marcus Dörr¹⁸³, Heyo K. Kroemer¹⁸⁴, Uwe Völker²⁰, Henry Völzke¹⁸⁵, Pilar Galan¹⁸⁶, Serge Hercberg¹⁸⁶, Mark Lathrop²⁴, Diana Zelenika²⁴, Panos Deloukas¹¹⁹, Massimo Mangino²⁸, Tim D. Spector²⁸, Guangju Zhai²⁸, James F. Meschia¹⁸⁷, Michael A. Nalls⁸³, Pankaj Sharma¹⁸⁸, Janos Terzic¹⁸⁹, M.

¹⁷³Department of Medical Genetics, Erasmus Medical Center, Rotterdam, The Netherlands

¹⁵⁷Institute of Epidemiology, Helmholtz Zentrum Munich, German Research Centre for Environmental Health, 85764 Neuherberg, Germany ¹⁵⁸Chair of Epidemiology, Institute of Medical Informatics, Biometry and Epidemiology, Ludwig-Maximilians-Universität, 81377

Munich, Germany 159Klinikum Grosshadern, 81377 Munich, Germany

¹⁶⁰National Heart and Lung Institute, Imperial College London, London, UK, W12 0HS, UK

¹⁶¹Medical Genetics Institute, Cedars-Sinai Medical Center, Los Angeles, CA, USA

¹⁶²Medical Population Genetics, Broad Institute of Harvard and MIT, 5 Cambridge Center, Cambridge MA 02142, USA

¹⁶³National Heart, Lung and Blood Institute and its Framingham Heart Study, 73 Mount Wayte Ave., Suite #2, Framingham, MA 01702, USA

¹⁶⁴Department of Neurology and Medicine, University of Washington, Seattle, USA

¹⁶⁵Department of Medicine (Geriatrics), University of Mississippi Medical Center, Jackson, MS, USA

¹⁶⁶ Department of Neurology, Boston University School of Medicine, USA

¹⁶⁷ Finnish Institute of Occupational Health, Finnish Institute of Occupational Health, Aapistie 1, 90220 Oulu, Finland

¹⁶⁸Wellcome Trust Centre for Human Genetics, University of Oxford, UK ¹⁶⁹Lapland Central Hospital, Department of Physiatrics, Box 8041, 96101 Rovaniemi, Finland

¹⁷⁰Center for Non-Communicable Diseases Karachi, Pakistan

¹⁷¹ Atherosclerosis Research Unit, Department of Medicine, Karolinska Institute, Stockholm, Sweden

¹⁷²Department of Internal Medicine, University Medical Center Groningen, University of Groningen, The Netherlands

¹⁷⁴Gerontology Research Center, National Institute on Aging, Baltimore, MD 21224, USA

¹⁷⁵Istituto di Neurogenetica e Neurofarmacologia, Consiglio Nazionale delle Ricerche, Cittadella Universitaria di Monserrato, Monserrato, Cagliari, Italy 176Unita' Operativa Semplice Cardiologia, Divisione di Medicina, Presidio Ospedaliero Santa Barbara, Iglesias, Italy

¹⁷⁷Computational Medicine Research Group, Institute of Clinical Medicine, University of Oulu and Biocenter Oulu, 90014 University of Oulu, Oulu, Finland ¹⁷⁸NMR Metabonomics Laboratory, Department of Biosciences, University of Eastern Finland, 70211 Kuopio, Finland

¹⁷⁹Department of Internal Medicine and Biocenter Oulu, Clinical Research Center, 90014 University of Oulu, Oulu, Finland

¹⁸⁰Institute for Molecular Medicine Finland FIMM, 00014 University of Helsinki, Helsinki, Finland

¹⁸¹Department of Biomedical Engineering and Computational Science, School of Science and Technology, Aalto University, 00076 Aalto, Espoo, Finland ¹⁸²NUS Graduate School for Integrative Sciences & Engineering (NGS) Centre for Life Sciences (CeLS), Singapore, 117456,

Singapore

Department of Internal Medicine B, Ernst-Moritz-Arndt-University Greifswald, 17487 Greifswald, Germany

¹⁸⁴Institute of Pharmacology, Ernst-Moritz-Arndt-University Greifswald, 17487 Greifswald, Germany

¹⁸⁵ Institute for Community Medicine, Ernst-Moritz-Arndt-University Greifswald, 17487 Greifswald, Germany

¹⁸⁶U557 Institut National de la Santé et de la Recherche Médicale, U1125 Institut National de la Recherche Agronomique, Université Paris 13, Bobigny, France ¹⁸⁷Department of Neurology, Mayo Clinic, Jacksonville, FL, USA

¹⁸⁸ Imperial College Cerebrovascular Unit (ICCRU), Imperial College, London, W6 8RF, UK

¹⁸⁹Faculty of Medicine, University of Split, Croatia

J. Kranthi Kumar⁴⁷, Matthew Denniff⁷¹, Ewa Zukowska-Szczechowska¹⁹⁰, Lynne E. Wagenknecht⁷⁹, F. Gerald R. Fowkes¹⁹¹, Fadi J. Charchar¹⁹², Peter E.H. Schwarz¹⁹³, Caroline Hayward⁷⁰, Xiuqing Guo¹⁶¹, Charles Rotimi⁷⁴, Michiel L, Bots⁶³, Eva Brand¹⁹⁴, Nilesh J. Samani⁷¹,⁷², Ozren Polasek¹⁹⁵, Philippa J. Talmud⁶⁹, Fredrik Nyberg⁶⁸,¹⁹⁶, Diana Kuh⁶⁷, Maris Laan²⁵, Kristian Hveem⁶⁶, Lyle J. Palmer¹⁹⁷, ¹⁹⁸, Yvonne T. van der Schouw⁶³, Juan P. Casas¹⁹⁹, Karen L. Mohlke⁶¹, Paolo Vineis²⁰⁰,⁶⁰, Olli Raitakari²⁰¹, Santhi K. Ganesh²⁰², Tien Y. Wong²⁰³,²⁰⁴, E Shyong Tai²⁰⁵,⁵⁷,²⁰⁶, Richard S. Cooper⁵⁴, Markku Laakso⁵³, Dabeeru C. Rao²⁰⁷, Tamara B. Harris²², Richard W. Morris²⁰⁸, Anna F. Dominiczak²⁰⁹, Mika Kivimaki²¹⁰, Michael G. Marmot²¹⁰, Tetsuro Miki⁴⁹, Danish Saleheen^{170,48}, Giriraj R. Chandak⁴⁷, Josef Coresh²¹¹, Gerjan Navis²¹², Veikko Salomaa¹²⁵, Bok-Ghee Han⁴⁴, Xiaofeng Zhu⁹⁴, Jaspal S. Kooner¹⁶⁰,⁴¹, Olle Melander³⁹, Paul M Ridker⁸,²¹³,⁹, Stefania Bandinelli²¹⁴, Ulf B. Gyllensten³⁷, Alan F. Wright⁷⁰, James F. Wilson³⁴, Luigi Ferrucci³³, Martin Farrall³², Jaakko Tuomilehto²¹⁵, ²¹⁶, ²¹⁷, ²¹⁸, Peter P. Pramstaller^{30, 219}, Roberto Elosua^{29, 220}, Nicole Soranzo^{119, 28}, Eric J.G. Sijbrands^{13, 14}, David Altshuler²²¹,¹¹⁵, Ruth J.F. Loos²³, Alan R. Shuldiner²⁶,²²², Christian Gieger¹⁵⁷, Pierre Meneton²²³, Andre G. Uitterlinden¹³,¹⁴,¹⁵, Nicholas J. Wareham²³, Vilmundur Gudnason^{10,11}, Jerome I. Rotter¹⁶¹, Rainer Rettig²²⁴, Manuela Uda¹⁷⁵, David P.

²⁰⁵Department of Medicine, Yong Loo Lin School of Medicine, National University of Singapore, Singapore, 119074, Singapore ²⁰⁶Duke-National University of Singapore Graduate Medical School, Singapore, 169857, Singapore

¹⁹⁰Department of Internal Medicine, Diabetology, and Nephrology, Medical University of Silesia, 41-800, Zabrze, Poland ¹⁹¹Public Health Sciences section, Division of Community Health Sciences, University of Edinburgh, Medical School, Teviot Place, Edinburgh, EH8 9AG, UK

¹⁹²School of Science and Engineering, University of Ballarat, 3353 Ballarat, Australia

¹⁹³Prevention and Care of Diabetes, Department of Medicine III, Medical Faculty Carl Gustav Carus at the Technical University of Dresden, 01307 Dresden, Germany 194 University Hospital Münster, Internal Medicine D, Münster, Germany

¹⁹⁵Department of Medical Statistics, Epidemiology and Medical Informatics, Andrija Stampar School of Public Health, University of Zagreb, Croatia 196 AstraZeneca R&D, 431 83 Mölndal, Sweden

¹⁹⁷Genetic Epidemiology & Biostatistics Platform, Ontario Institute for Cancer Research, Toronto

¹⁹⁸Samuel Lunenfeld Institute for Medical Research, University of Toronto, Canada

¹⁹⁹Faculty of Epidemiology and Population Health, London School of Hygiene and Tropical Medicine, UK

²⁰⁰Department of Epidemiology and Public Health, Imperial College, Norfolk Place London W2 1PG, UK

²⁰¹Research Centre of Applied and Preventive Cardiovascular Medicine, University of Turku and the Department of Clinical Physiology, Turku University Hospital, Turku, 20521, Finland ²⁰²Department of Internal Medicine, Division of Cardiovascular Medicine, University of Michigan Medical Center, Ann Arbor,

Michigan, USA ²⁰³Singapore Eye Research Institute, Singapore, 168751, Singapore

²⁰⁴Department of Ophthalmology, National University of Singapore, Singapore, 119074, Singapore

²⁰⁷Division of Biostatistics, Washington University School of Medicine, Saint Louis, MO, 63110, USA

²⁰⁸Department of Primary Care & Population Health, UCL, London, UK, NW3 2PF, UK

²⁰⁹BHF Glasgow Cardiovascular Research Centre, University of Glasgow, 126 University Place, Glasgow, G12 8TA, UK

²¹⁰Epidemiology Public Health, UCL, London, UK, WC1E 6BT, UK

²¹¹Departments of Epidemiology, Biostatistics, and Medicine, Johns Hopkins University, Baltimore MD, USA

²¹²Division of Nephrology, Department of Internal Medicine, University Medical Center Groningen, University of Groningen, The Netherlands ²¹³Division of Cardiology, Brigham and Women's Hospital, 900 Commonwealth Avenue East, Boston MA 02215, USA

²¹⁴Geriatric Rehabilitation Unit, Azienda Sanitaria Firenze (ASF), Florence, Italy

²¹⁵National Institute for Health and Welfare, Diabetes Prevention Unit, 00271 Helsinki, Finland

²¹⁶Hjelt Institute, Department of Public Health, University of Helsinki, 00014 Helsinki, Finland

²¹⁷South Ostrobothnia Central Hospital, 60220 Seinäjoki, Finland

²¹⁸Red RECAVA Grupo RD06/0014/0015, Hospital UniversitarioLa Paz, 28046 Madrid, Spain

²¹⁹Department of Neurology, General Central Hospital, 39100 Bolzano, Italy

²²⁰CIBER Epidemiología y Salud Pública, 08003 Barcelona

²²¹Department of Medicine and Department of Genetics, Harvard Medical School, Boston, Massachusetts 02115, USA

²²²Geriatric Research and Education Clinical Center, Veterans Administration Medical Center, Baltimore, MD, USA

²²³U872 Institut National de la Santé et de la Recherche Médicale, Centre de Recherche des Cordeliers, Paris, France

²²⁴Institute of Physiology, Ernst-Moritz-Arndt-University Greifswald, 17487 Greifswald, Germany

Strachan⁵⁰, Jacqueline C.M. Witteman¹³,¹⁵, Anna-Liisa Hartikainen²²⁵, Jacques S. Beckmann¹⁰⁵,²²⁶, Eric Boerwinkle²²⁷, Ramachandran S. Vasan⁶,²²⁸, Michael Boehnke³¹, Martin G. Larson^{6,229}, Marjo-Riitta Järvelin^{18,230,231,232,233}, Bruce M. Psaty^{21,135*}, Gonçalo R Abecasis^{19*}, Aravinda Chakravarti^{1*#}, Paul Elliott¹⁸,^{233*}, Cornelia M. van Duijn¹³,^{234*}, Christopher Newton-Cheh²⁷,^{115*#}, Daniel Levy⁶,¹⁶,^{7*#}, Mark J. Caulfield^{4*#}, Toby Johnson^{4*}

²²⁵Institute of Clinical Medicine/Obstetrics and Gynecology, University of Oulu, Finland

 ²²⁶Service of Medical Genetics, Centre Hospitalier Universitaire Vaudois, 1011 Lausanne, Switzerland
²²⁷Human Genetics Center, 1200 Hermann Pressler, Suite E447 Houston, TX 77030, USA

 ²²⁸Division of Epidemiology and Prevention, Boston University School of Medicine, Boston, MA, USA
²²⁹Department of Mathematics, Boston University, Boston, MA, USA
²³⁰Institute of Health Sciences, University of Oulu, BOX 5000, 90014 University of Oulu, Finland
²³¹Enter and Enter and Ent ²³¹Biocenter Oulu, University of Oulu, BOX 5000, 90014 University of Oulu, Finland

²³²National Institute for Health and Welfare, Box 310, 90101 Oulu, Finland

²³³MRC-HPA Centre for Environment and Health, School of Public Health, Imperial College London, Norfolk Place, London W2 1PG, UK ²³⁴Centre of Medical Systems Biology (CMSB 1-2), NGI Erasmus Medical Center, Rotterdam, The Netherlands

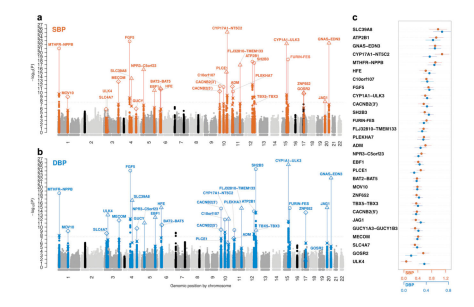


Fig. 1.

Genome-wide $-\log_{10} P$ -value plots and effects for significant loci. Genome-wide $-\log_{10} P$ -value plots are shown for systolic (SBP: panel a) and diastolic (DBP: panel b). SNPs within loci reaching genome-wide significance are labeled in red for SBP and blue for DBP (±2.5Mb of lowest *P*-value) and lowest *P*-values in the initial genome-wide analysis as well as the results of analysis including validation data are labeled separately. The lowest *P*-values in the initial GWAS are denoted as an X. The range of different sample sizes in the final meta-analysis including the validation data are indicated as: circle (96–140k), triangle (>140–180k), and diamond (>180–220k). SNPs near unconfirmed loci are in black. The horizontal dotted line is *P*=2.5 × 10⁻⁸. Panel c shows the effect size estimates and 95% confidence bars per BP-increasing allele of the 29 significant variants for SBP (red) and DBP (blue). Effect sizes are expressed in mmHg/allele. GUCY = *GUCY1A3-GUCY1B3*.

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Table 1

Summary association results for 29 BP SNPs

shown. New genome-wide significant findings (17 SNPs) are presented in the top half of the table, data on 12 previously published signals are presented Summary association statistics, based on combined discovery and follow-up data, for 29 independent SNPs in individuals of European ancestry are in the lower half.

Locus	Index SNP	Chr	Position	CA/NCA	CAF	nsSNP	eSNP		SBP			DBP			NTH
								Beta	P-value	Effect in EA/SA/A	Beta	P-value	Effect in EA/SA/A	Beta	P-value
0110W	rs2932538	-	113,018,066	G/A	0.75	$\mathbf{Y}(\mathbf{p})$	Y(p)	0.388	$1.2^{*}10^{-9}$	-/+/+	0.24	9.9^*10^{-10}	-/*+/+	0.049	$2.9^{*}10^{-7}$
atita. F	rs13082711	с	27,512,913	T/C	0.78	Y(p)	Y(p)	-0.315	1.5^*10^{-6}	+/-/-	-0.238	3.8^*10^{-9}	+/-/-	-0.035	3.6^*10^{-4}
MECOM Trouger	rs419076	б	170,583,580	T/C	0.47		ı.	0.409	1.8^*10^{-13}	+/+/+	0.241	$2.1^* 10^{-12}$	-/+/+	0.031	3.1^*10^{-4}
nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate nau Rate Rate Rate Rate Rate Rate Rate Rate	rs13107325	4	103,407,732	T/C	0.05	Y	Y(+)	-0.981	3.3^*10^{-14}	+/+/2	-0.684	2.3^*10^{-17}	+/+/¿	-0.105	$4.9^{*}10^{-7}$
GUCYIA3-GUCYIB3	rs13139571	4	156,864,963	C/A	0.76			0.321	$1.2^{*}10^{-6}$	+/-/+	0.26	2.2^*10^{-10}	+/-/+	0.042	2.5^*10^{-5}
RPR3-CSorf23	rs1173771	S.	32,850,785	G/A	0.6		ı	0.504	1.8^*10^{-16}	+/+/* +	0.261	9.1^*10^{-12}	-/+/* +	0.062	3.2^*10^{-10}
n BMC	rs11953630	S.	157,777,980	T/C	0.37		ı.	-0.412	3.0^*10^{-11}	+/+/+	-0.281	3.8^*10^{-13}	+/+/+	-0.052	1.7^*10^{-7}
Э. Д 20 Д 2 М	rs1799945	9	26,199,158	G/C	0.14	Y	ı	0.627	7.7^*10^{-12}	-/+/+	0.457	1.5^*10^{-15}	-/+/+	0.095	$1.8^* 10^{-10}$
ta BAT2-BAT5	rs805303	9	31,724,345	G/A	0.61	Y(p)	$\mathbf{Y}^{(+)}$	0.376	1.5^*10^{-11}	¿/-/-	0.228	3.0^*10^{-11}	+/-/-	0.054	1.1^*10^{-10}
CACNB2(5')	rs4373814	10	18,459,978	G/C	0.55		ı.	-0.373	4.8^*10^{-11}	-/+/+	-0.218	4.4^*10^{-10}	-/+/-	-0.046	8.5^*10^{-8}
PLCE1	rs932764	10	95,885,930	G/A	0.44	,	ı	0.484	7.1^*10^{-16}	-/+/+	0.185	$8.1^{*}10^{-7}$	-/+/+	0.055	9.4^*10^{-9}
ADM	rs7129220	Ξ	10,307,114	G/A	0.89		,	-0.619	3.0^*10^{-12}	+/-/%	-0.299	6.4^*10^{-8}	+/-/¿	-0.044	1.1^*10^{-3}
FLJ32810-TMEM133	rs633185	Ξ	100,098,748	G/C	0.28		ı	-0.565	1.2^*10^{-17}	+/+/*+	-0.328	$2.0^{*}10^{-15}$	-/+/* +	-0.07	5.4^*10^{-11}
FURIN-FES	rs2521501	15	89,238,392	T/A	0.31		Y(-)	0.65	5.2^*10^{-19}	+/+/*+	0.359	$1.9^{*}10^{-15}$	+/+/* +	0.059	$7.0^{*}10^{-7}$
GOSR2	rs17608766	17	42,368,270	T/C	0.86		Y(+)	-0.556	1.1^*10^{-10}	+/-/+	-0.129	0.017	+//+	-0.025	0.08

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Locus	Index SNP	Chr	Position	CA/NCA	CAF	nsSNP	eSNP		SF	SBP		DBP	4	H	NTH
								Beta	P-value	Effect in EA/SA/A	Beta	P-value	Effect in EA/SA/A	Beta	P-value
JAGI	rs1327235	20	10,917,030	G/A	0.46			0.34	$1.9^{*}10^{-8}$	+/+/*+	0.302	1.4^*10^{-15}	+/*+/*+	0.034	$4.6^{*}10^{-4}$
GNAS-EDN3	rs6015450	20	57,184,512	G/A	0.12	Y(p)	ı	0.896	3.9^*10^{-23}	+/+/	0.557	5.6^*10^{-23}	+/*+/?	0.11	$4.2^{*}10^{-14}$
MTHFR-NPPB	rs17367504	-	11,785,365	G/A	0.15		Y(-/r)	-0.903	8.7^*10^{-22}	+/+/+	-0.547	3.5^*10^{-19}	+/+/+	-0.103	2.3^*10^{-10}
ULK4	rs3774372	3	41,852,418	T/C	0.83	Y	Y(r/p)	-0.067	0.39	+/-/-	-0.367	9.0^*10^{-14}	+/+/+	-0.017	0.18
FGF5	rs1458038	4	81,383,747	T/C	0.29			0.706	1.5^*10^{-23}	+/+/*+	0.457	8.5^*10^{-25}	+/*+/*+	0.072	$1.9^{*}10^{-7}$
CACNB2(3')	rs1813353	10	18,747,454	T/C	0.68		ı	0.569	2.6^*10^{-12}	+/+/+	0.415	2.3^*10^{-15}	+/+/+	0.078	6.2^*10^{-10}
C10orf107	rs4590817	10	63,137,559	G/C	0.84		Y(r)	0.646	4.0^*10^{-12}	-/+/-	0.419	1.3^*10^{-12}	-/-/-	0.096	9.8^*10^{-9}
CYP17A1-NT5C2	rs11191548	10	104,836,168	T/C	0.91		Y(-)	1.095	6.9^*10^{-26}	+/*+/*+	0.464	9.4^*10^{-13}	+/*+/*+	0.097	$1.4^{*}10^{-5}$
PLEKHA7	rs381815	Ξ	16,858,844	T/C	0.26	ı	ı	0.575	5.3^*10^{-11}	+/+/*+	0.348	5.3^*10^{-10}	+/-/*+	0.062	3.4^*10^{-6}
ATP2BI	rs17249754	12	88,584,717	G/A	0.84		ı	0.928	1.8^*10^{-18}	-/*+/*+	0.522	1.2^*10^{-14}	-/*+/*+	0.126	1.1^*10^{-14}
SH2B3	rs3184504	12	110,368,991	T/C	0.47	Y	Y(+)	0.598	3.8^*10^{-18}	+/-/-	0.448	3.6^*10^{-25}	+//	0.056	2.6^*10^{-6}
TBX5-TBX3	rs10850411	12	113,872,179	T/C	0.7		ı	0.354	$5.4^{*}10^{-8}$	-/+/-	0.253	5.4^*10^{-10}	-/-/-	0.045	$5.2^{*}10^{-6}$
CYPIAI-ULK3	rs1378942	15	72,864,420	C/A	0.35		$\mathbf{Y}^{(+)}$	0.613	5.7^*10^{-23}	+/+/*+	0.416	2.7^*10^{-26}	-/+/*+	0.073	$1.0^{*}10^{-8}$
ZNF652	rs12940887	17	44,757,806	T/C	0.38		Y(-)	0.362	$1.8^* 10^{-10}$	+/-/+	0.27	2.3^*10^{-14}	+//+	0.046	1.2^*10^{-7}
Y indicates the BP index SNP is a nsSNP, $Y(p)$ indicates a proxy SNP is a nsSNP. $Y(+)$: indicates BP index SNP is the strongest known eSNP for a transcript; $Y(-)$: indicates BP index SNP is an eSNP but	SNP is a nsSNP	, Y(p) i	ndicates a proxy	SNP is a ns	SNP. Y(⊦	+): indicaté	ssBP index	SNP is th	le strongest k	nown eSNP for a transc	sript; Y(-)	: indicates BP ind	index SNP is an eSNF	NP but	

Nature. Author manuscript; available in PMC 2012 May 01.

not strongest known eSNP for any transcript. Y(r): indicates BP index SNP is strongest known eSNP in a regional SNP-RTPCR experiment. Y(p): indicates a proxy SNP (r² > 0.8) to BP SNP is an eSNP but not the strongest known eSNP. Observed effect directions in East Asian (EA), South Asian (SA), and African (A) ancestry individuals are coded + or - if concordant or discordant with directions in European ancestry results;

* denotes significance controlling the FDR at 5% over 58 tests per ancestry (Supplementary Tables 5 and 12). Effect size estimates (beta) correspond to mmHg per coded allele for SBP and DBP and ln(odds) per coded allele for HTN.

CA = coded allele; NCA = non-coded allele; CAF = coded allele frequency; ? denotes missing data. Genomic positions use NCBI Build 36 coordinates.

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Table 2

Genetic risk score and cardiovascular outcome association results

Association of genetic risk score (using all 29 SNPs at 28 loci, parameterised using the average of SBP and DBP effects [=(SBP effect + DBP effect)/2] from the discovery analysis), tested in results from other GWAS consortia.

		Effect	SE			Contrast	Contrast top vs. bottom	m	
Phenotype	Source	(per SD of g	(per SD of genetic risk score)) <i>P</i> -value	# SNPs	quintiles	deciles		N case/control or total
Blood pressure phenotypes									
SBP [mmHg]	WGHS	1.645	0.098 (a)	$6.5*10^{-63}$	29	4.61	5.77 (<i>(a)</i>	23,294
DBP [mmHg]	WGHS	1.057	0.067 (a)	8.4*10 ⁻⁵⁷	29	2.96	3.71 (<i>(a)</i>	23,294
Prevalent hypertension	WGHS	0.211	0.018 (b)	3.1*10 ⁻³³	29	1.80	2.09 ((q)	5,018/18,276
Prevalent hypertension	BRIGHT	0.287	0.031 (b)	$7.7*10^{-21}$	29	2.23	2.74 ((q)	2,406/1,990
Dichotomous endpoints									
Incident heart failure	CHARGE-HF	0.035	0.021 (c)	0.10	29	1.10	1.13 ((c)	2,526/18,400
Incident stroke	NEURO-CHARGE	0.103	0.028 (c)	0.0002	28	1.34	1.44 (<i>(c)</i>	1,544/18,058
Prevalent stroke	UK-US Stroke Collaborative Group(SCG)	0.075	0.037 (b)	0.05	29	1.23	1.30 ((q)	1,473/1,482
Stroke (combined, incident and prevalent)	CHARGE & SCG	NA	NA NA	3.3*10 ⁻⁵	NA	NA	NA I	NA	3,017/19,540
Prevalent CAD	CARDIoGRAM	0.092	0.010 (b)	$1.6*10^{-19}$	28	1.29	1.38 ((q)	22,233/64,726
Prevalent CAD	C4D ProCARDIS	0.132	0.022 (b)	$2.2*10^{-9}$	29	1.45	1.59 ((q)	5,720/4,381
Prevalent CAD	C4D HPS	0.083	0.027 (b)	0.002	29	1.26	1.34 ((q)	2,704/2,804
Prevalent CAD (combined)	CARDIoGRAM & C4D	0.100	(<i>q</i>) 600.0	$8.1*10^{-29}$	29	1.32	1.42 ((q)	30,657/71,911
Prevalent chronic kidney disease	CKDGen	0.014	0.015 (b)	0.35	29	1.04	1.05 ((q)	5,807/61,286
Prevalent microalbuminuria	CKDGen	0.008	(<i>b</i>) 0.019 (<i>b</i>)	0.68	29	1.02	1.03 ((q)	3,698/27,882
Continuous measures oftarget organ damage	mage								

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		Effect	SE				Contrast top vs. bottom	top vs. bot	tom	
Phenotype Blood pressure phenotypes	Source	(per SD of genetic risk score)	enetic risk s	core)	<i>P</i> -value	# SNPs	P-value #SNPs quintiles deciles	deciles		N case/control or total
Left ventricular mass [g]	EchoGen	0.822	0.317 (a)	<i>(a)</i>	0.01	29	2.30	2.89 <i>(a)</i>	(a)	12,612
Left ventricular wall thickness[cm]	EchoGen	600.0	0.002 (a) $6.0*10^{-6}$	<i>(a)</i>	$6.0*10^{-6}$	29	0.03	0.03 (a)	<i>(a)</i>	12,612
Serum creatinine	KidneyGen	-0.001	0.001 (d)	(<i>p</i>)	0.24	29	1.00	1.00 1.00 (<i>d</i>)	(p)	23,812
eGFR (4 parameter MDRD equation)	CKDGen	-0.0001	-0.0001 0.0009 (d)	<i>(p)</i>	0.93	29	1.00	1.00 (d)	(p)	67,093
Urinary albumin/creatinine ratio	CKDGen	0.005	0.007 (d)	(<i>p</i>)	0.43	29	1.01	1.02 (d)	(p)	31,580

 $^{(b)}$ Units are ln(odds) per SD of genetic risk score, or odds ratio between top/bottom quintiles or deciles.

(c) Units are ln(hazard) per SD of genetic risk score, or hazard ratio between top/bottom quintiles or deciles.

 $^{(d)}$ Units are ln(phenotype) per SD of genetic risk score, or phenotypic ratio between top/bottom quintiles or deciles.