Genetic and observational evidence: No independent role for cholesterol efflux over static high-density lipoprotein concentration measures in coronary heart disease risk assessment

Sanna Kuusisto1,2,3,4, Minna K. Karjalainen1,2,4,5, Therese Tillin6, Antti J. Kangas7, Michael V. Holmes8, Mika Kähönen9, Terho Lehtimäki10, Jorma Viikari11,12, Markus Perola13,14,15, Nishi Chaturvedi6, Veikko Salomaa15, Olli T. Raitakari16,17,18, Marjo-Riitta Järvelin2,4,19,20,21, Johannes Kettunen1,2,4,15 & Mika Ala-Korpela1,2,3,4

From the 1Computational Medicine, Faculty of Medicine, University of Oulu, Oulu, Finland; 2Center for Life Course Health Research, Faculty of Medicine, University of Oulu, Oulu, Finland; 3NMR Metabolomics Laboratory, School of Pharmacy, University of Eastern Finland, Kuopio, Finland; 4Biocenter Oulu, University of Oulu, Oulu, Finland; 5Northern Finland Birth Cohorts, Arctic Biobank, Infrastructure for Population Studies, Faculty of Medicine, University of Oulu, Oulu, Finland; 6MRC Unit for Lifelong Health and Ageing at UCL, Institute of Cardiovascular Science, University College London, London, UK; 7Nightingale Health Plc., Helsinki, Finland; 8Medical Research Council Population Health Research Unit, University of Oxford, Oxford, UK; 9Department of Clinical Physiology, Tampere University Hospital and Finnish Cardiovascular Research Center Tampere, Faculty of Medicine and Health Technology, Tampere University, Tampere, Finland; 10Department of Clinical Chemistry, Fimlab Laboratories and Finnish Cardiovascular Research Center Tampere, Faculty of Medicine and Health Technology, Tampere University, Tampere, Finland; 11Department of Medicine, University of Turku, Turku, Finland; 12Division of Medicine, Turku University Hospital, Turku, Finland; 13Research Programs Unit, Diabetes and Obesity, University of Helsinki, Helsinki, Finland; 14Estonian Genome Center, University of Tartu, Tartu, Estonia; 15Department of Public Health and Welfare, Finnish Institute for Health and Welfare, Helsinki, Finland; 16Centre for Population Health Research, University of Turku and Turku University Hospital, Turku, Finland; 17Research Centre of Applied and Preventive Cardiovascular Medicine, University of Turku, Turku, Finland; 18Department of Clinical Physiology and Nuclear Medicine, Turku University Hospital, Turku, Finland; 19Unit of Primary Health Care, Oulu University Hospital (OYS), Oulu, Finland; 20Department of Epidemiology and Biostatistics, MRC-PHE Centre for Environment and Health, School of Public Health, Imperial College London, London, UK; and 21Department of Life Sciences, College of Health and Life Sciences, Brunel University London, London, UK

Abstract. Kuusisto S, Karjalainen MK, Tillin T, Kangas AJ, Holmes MV, Kähönen M, et al. Genetic and observational evidence: No independent role for cholesterol efflux over static high-density lipoprotein concentration measures in coronary heart disease risk assessment. J Intern Med. 2022;292:146–153.

Background. Observational findings for high-density lipoprotein (HDL)-mediated cholesterol efflux capacity (HDL-CEC) and coronary heart disease (CHD) appear inconsistent, and knowledge of the genetic architecture of HDL-CEC is limited.

Objectives. A large-scale observational study on the associations of HDL-CEC and other HDL-related measures with CHD and the largest genome-wide association study (GWAS) of HDL-CEC.

Participants/methods. Six independent cohorts were included with follow-up data for 14,438 participants to investigate the associations of HDL-related measures with incident CHD (1,570 events). The GWAS of HDL-CEC was carried out in 20,372 participants.

Results. HDL-CEC did not associate with CHD when adjusted for traditional risk factors and HDL cholesterol (HDL-C). In contradiction, almost all HDL-related concentration measures associated consistently with CHD after corresponding adjustments. There were no genetic loci associated with HDL-CEC independent of HDL-C and triglycerides.
Conclusion. HDL-CEC is not unequivocally associated with CHD in contrast to HDL-C, apolipoprotein A-I, and most of the HDL subclass particle concentrations.

Keywords: cholesterol efflux, coronary heart disease, genome-wide association study, HDL, observational cohort study, triglycerides

Introduction
A functional attribute of high-density lipoprotein (HDL) particles, cholesterol efflux capacity (CEC), associates inversely with incident cardiovascular events in observational studies, independent of HDL cholesterol (HDL-C) [1, 2]. However, findings appear inconsistent [3, 4], and it is currently unknown whether HDL-CEC plays a causal role in cardiovascular disease. No direct randomized controlled trials exist, and Mendelian randomization analyses with reliable genetic instruments are yet to be conducted. Nevertheless, apolipoprotein A-I (apoA-I) infusion therapies have failed to show any clinical benefit [5–7], and a recent Mendelian randomization analysis did not support a cardioprotective role for apoA-I [8]. These studies provide indirect evidence against the causality of HDL-CEC since the physiological concept of HDL-mediated cholesterol efflux is based on the rationale that apoA-I is the key molecular component to promote cholesterol efflux from arterial wall macrophages [9].

To date, one genome-wide association study (GWAS) is available with results in 5,293 individuals for four different experimental measures of HDL-CEC, depicting different efflux pathways, indicating the involvement of five well-known lipid loci [10]. For the most commonly used HDL-CEC measure in cardiovascular studies, J774 stimulated HDL-CEC [1, 3, 4, 11], only two loci were detected, and these associations are not independent of HDL-C and triglycerides [10].

To clarify the role of J774 stimulated HDL-CEC in cardiovascular disease, we (1) combined three prospective cohorts \( n = 14,438 \) to study the association of HDL-CEC and various other HDL-related measures with coronary heart disease (CHD; incident events \( n = 1,570 \)) and (2) performed a GWAS of HDL-CEC in five independent cohorts of 20,372 participants.

Methods
Study populations
The cohorts are characterized in Table 1 (see Study populations in the Supplement for details). The studies were approved by the ethics committees of the study sites and written informed consent was obtained from all participants.

HDL-CEC, lipoprotein, and lipid analyses
HDL-CEC, four HDL subclass particle concentrations, standard lipoprotein lipids, and apoA-I were analyzed by nuclear magnetic resonance (NMR) spectroscopy [11, 12]. This methodology has been widely used in epidemiological and genetic studies over the last 10 years [12]. The HDL-CEC values correspond to the most commonly used assay to quantify HDL-CEC, that is, the use of cAMP-treated J774 macrophages with radiolabeled cholesterol (J774 stimulated HDL-CEC) [11]. The HDL-CEC values as well as all the static HDL-related measures are directly estimated from the NMR spectral data points with a distinct regression model. Each metabolic measure therefore has its own specific and optimized quantification routine. The estimation of HDL-CEC does not rely directly on any other HDL (or lipoprotein) measure, and there is no direct simple formula between HDL-CEC and the other HDL measures [11, 12]. The correlations of the (NMR-based) HDL-CEC with other HDL and lipoprotein measures are weak—modest at best—and agree with those using the in vitro HDL-CEC measurements via cAMP-treated J774 cell assays [11].

Statistical analyses in the follow-up studies
Prospective information on CHD events were available for DILGOM2007, FINRISK1997, and SABRE with follow-up times of 8, 15, and 20 years, respectively. These cohorts had complete data for 14,438 participants with 1,570 incident

© 2022 The Authors. Journal of Internal Medicine published by John Wiley & Sons Ltd on behalf of Association for Publication of The Journal of Internal Medicine.
Table 1. Characteristics of the cohorts

|                  | FINRISK1997 (n = 7,603) | DILGOM2007 (n = 4,884) | SABRE (n = 3,268) | YFS2007 (n = 2,160) | NFBC1986 (n = 5,604) | NFBC1966 (n = 5,692) |
|------------------|-------------------------|------------------------|-------------------|-------------------|---------------------|---------------------|
| Age, years       | 48 (37–59)              | 54 (42–64)             | 52 (46–58)        | 39 (33–42)        | 16 (16–16)          | 31 (31–31)          |
| Female, n (%)    | 3825 (50)               | 2602 (53)              | 465 (14.2)        | 1186 (55)         | 2801 (51)           | 2950 (52)           |
| BMI, kg/m²       | 26.1 (23.5–29.1)        | 26.5 (24.0–30.0)       | 25.7 (23.7–28.2)  | 25.3 (22.7–28.4)  | 20.5 (19.0–22.6)    | 24.0 (21.9–26.6)    |
| Systolic BP, mmHg| 134 (121–149)           | 135 (122–149)          | 122 (112–134)     | 119 (111–130)     | 115 (107–124)       | 124 (115–133)       |
| Diastolic BP, mmHg| 82 (74–90)              | 79 (72–87)             | 78 (71–85)        | 75 (68–83)        | 68 (63–73)          | 77 (70–84)          |
| Total cholesterol, mmol/L | 5.25 (4.55–6.03) | 4.30 (3.74–4.87) | 3.93 (3.37–4.53) | 5.16 (4.56–5.92) | 4.29 (3.81–4.88) | 5.21 (4.53–6.07) |
| Triglycerides, mmol/L | 1.16 (0.84–1.63) | 1.02 (0.78–1.37) | 0.92 (0.73–1.19) | 1.11 (0.81–1.61) | 0.86 (0.67–1.12) | 1.02 (0.75–1.44) |
| LDL cholesterol, mmol/L | 1.89 (1.49–2.30) | 1.57 (1.27–1.91) | 1.40 (1.13–1.70) | 1.92 (1.58–2.33) | 1.45 (1.19–1.75) | 1.89 (1.51–2.34) |
| HDL cholesterol, mmol/L | 1.54 (1.30–1.81) | 1.44 (1.23–1.69) | 1.01 (0.87–1.17) | 1.60 (1.34–1.90) | 1.43 (1.24–1.63) | 1.59 (1.32–1.90) |
| HDL-CEC, %        | 21.83 (20.66–23.44)    | 20.37 (19.32–21.73)    | 21.19 (20.12–22.28) | 21.85 (20.84–22.88) | 20.43 (19.36–21.37) | 21.19 (20.16–22.25) |

Notes: Data are median (25th and 75th percentiles) or percentage, when appropriate. The number of individuals in each characteristic may vary slightly depending on data availability. Individuals with data available for the required variables were included in each analysis. HDL-CEC, HDL-mediated cholesterol efflux, expressed as an estimated percentage of cholesterol effluxed from total cholesterol present in the cells, as described in Kuusisto et al. [11]. Abbreviations: BMI, body mass index; HDL, high-density lipoprotein; HDL-CEC, HDL-mediated cholesterol efflux capacity; LDL, low-density lipoprotein.
events (prevalent CHD events, outliers [11], and missing data were removed; see Study populations in the Supplement) and were used to study the associations of HDL-CEC and other HDL-related biomarkers with CHD. Data were analyzed by Cox proportional hazard regression models in each cohort and combined via random-effect meta-analysis due to obvious heterogeneity in HDL-CEC associations (Fig. 1).

**Genome-wide association study**

A GWAS of HDL-CEC was performed in the Finnish cohorts (Table 1) under the additive model, followed by fixed-effect meta-analysis (in 20,372 participants) (see Genetic analyses in the Supplement). For comparative purposes, we performed a GWAS of HDL-C using the same cohorts and participants. In addition to the primary analysis, HDL-CEC was also analyzed adjusting for HDL-C and serum triglycerides to directly compare with the findings by Low-Kam et al. [10]. We also analyzed the associations of the lead single nucleotide polymorphisms (SNPs) of those loci found by Low-Kam et al. [10] (see Replication of previously described associations in the Supplement).

**Discussion**

This is the first study including multiple cohorts with consistent measurements of J774 stimulated HDL-CEC with the same method. Our large-scale genetic and observational results—in comparison to multiple HDL-related concentration measures—suggest that J774 stimulated HDL-CEC does not unequivocally have an independent role in cardiometabolic risk assessment. These findings fit well with those from other recent studies indicating that HDL-CEC appears to be a more heterogeneous measure of incident [3, 4] and prevalent [13] cardiovascular outcomes than other HDL-related biomarkers. Our results also denote more variability between the individual cohorts for the associations of HDL-CEC and CHD than for those of the other studied HDL-related measures. While there is a rather contrasting difference in the heterogeneity of the associations for HDL-CEC and the static HDL measures, the heterogeneity (and its variability) in the observational studies as such is not unexpected. HDL-CEC estimates functionality that in vivo is most likely a sum of many naturally involved lipoprotein components, that is, it is not inherently HDL specific like the static HDL measures. Thus, the cohort-specific confounding (e.g., by lifestyle, socioeconomic factors, or
Fig. 1 Associations of high-density lipoprotein (HDL)-mediated cholesterol efflux capacity (CEC) and various other HDL-related measures with coronary heart disease (CHD). Data were meta-analyzed across cohorts by random-effects meta-analysis. Hazard ratios are per 1-standard deviation (SD) higher HDL measure. The HDL subclasses were defined by particle size as follows: very large (XL-HDL, average particle diameter 14.3 nm), large (L-HDL, 12.1 nm), medium (M-HDL, 10.9 nm), and small HDL (S-HDL, 8.7 nm). Total HDL refers to the sum of all the four HDL subclass particle concentrations. Open and closed black diamonds indicate $p \geq 0.01$ and $p < 0.01$ to denote evidence in favor of an association based on the Bonferroni correction of five independent tests ($p = 0.05/5 = 0.01$) due to the highly correlated nature of HDL-related measures. 

Traditional risk factors (TRF) included: age, sex, log body mass index, smoking, geographical region (ethnicity in SABRE), diabetes, mean arterial pressure, cardiovascular treatment, and serum concentrations of low-density lipoprotein cholesterol and log(triglycerides). In the third panel, HDL-CEC was adjusted for TRF + HDL-C and other HDL-related measures for TRF + HDL-CEC.
Results of genome-wide association studies of high-density lipoprotein (HDL)-mediated cholesterol efflux capacity (CEC) and HDL cholesterol (HDL-C). Genome-wide association studies of HDL-CEC and HDL-C were performed in five Finnish cohorts (total n = 20,372). Each dot represents a single single nucleotide polymorphism (SNP); 500-kb regions flanking the lead SNPs in the associated loci are highlighted. The level of genome-wide significance (p < 5 × 10⁻⁸) is indicated by a red dashed line. Altogether two and 13 genome-wide significant loci were detected for HDL-CEC and HDL-C, respectively.

Baseline health status affecting the associations of the static HDL measures and HDL-CEC cannot generally be assumed to be the same. In addition, the potential effects of reverse causation (i.e., when the early stages of the disease process influence the exposure) may vary for different measures [14].

The GWAS for HDL-CEC performed here is the largest to date with almost four times more participants than in the previous study (20,372 vs. 5,293) [10]. In the primary GWAS, two genetic loci were identified—namely, LIPC and CETP—but those were abolished when HDL-CEC was adjusted for circulating HDL-C and triglyceride concentrations. Previous evidence on the genetic determinants of HDL-CEC is limited to the study by Low-Kam et al. [10]. For J774 stimulated HDL-CEC, they detected two associated loci (CETP and APOE/C1/C2/C4), but these were not independent of HDL-C and triglycerides. The CETP locus was replicated in our GWAS analyses, but the association was abolished when adjusted for HDL-C and triglycerides.

We detected 13 associated loci in the GWAS of HDL-C—performed in the same individuals as the GWAS of HDL-CEC—suggesting an adequate sample size to detect strong associations. The LIPC and CETP loci associated with HDL-CEC in our primary analysis are highly pleiotropic (Table S3), and these associations were abolished via adjustments for circulating HDL-C and triglycerides. The genetic pleiotropy of HDL-CEC is not unexpected, since HDL-CEC is not a single protein biomarker with a clear genetic coding region, but a measure of HDL function, affected inherently by the complexity of overall lipoprotein metabolism.

This large-scale multicohort study for HDL-CEC was made possible due to the recent development of a new cost-effective NMR-based method to estimate HDL-CEC directly from serum samples [11]. The estimated HDL-CEC values have been shown to correspond to those from in vitro experiments, but it cannot be ruled out that the methodology could partly contribute to the heterogeneity between diverse study populations. The results should thus be interpreted with care. It should also be noted that this method is a proxy for radiolabeled cholesterol efflux assay performed in cAMP-treated J774 macrophages, and it may not be taken to represent other efflux models. The current GWAS for HDL-CEC with over 20,000 participants—approximately four times more than in the previous GWAS [10]—did not find genetic associations independent of HDL-C and triglycerides. However, it cannot be ruled out that an even larger GWAS might distinguish HDL-CEC specific genetic loci or that they might exist for other HDL-CEC pathways not represented by J774 stimulated HDL-CEC [11].
would hold in humans [16]. On the contrary, apoA-1 infusion therapies have not led to reductions in cardiovascular outcomes, despite increasing HDL-CEC [7]. Furthermore, recent Mendelian randomization analyses have shown that circulating apoA-I concentrations are unlikely to be causally related to CHD [8, 17], providing indirect evidence that therapeutic modification of HDL-CEC may not be beneficial.

The current large-scale genetic and observational analyses temper enthusiasm for an independent role of HDL-CEC either in disease prediction or as a causal entity in cardiovascular disease. However, further genome-wide data to potentially allow for robust genetic instruments for HDL-CEC and Mendelian randomization analyses against various outcomes would be beneficial alongside large-scale randomized controlled trials. The results and interpretations here regarding HDL function refer only to cholesterol efflux as defined in cAMP-treated J774 macrophages, and no extrapolation to other efflux models or other types of HDL functionality should be done.

Author contributions
Sanna Kuusisto, Minna K. Karjalainen, and Mika Ala-Korpela conceived the idea and drafted the manuscript. Therese Tillin, Antti J. Kangas, Mika Kähönen, Terho Lehtimäki, Jorma Viikari, Markus Perola, Nishi Chaturvedi, Veikko Salomaa, Olli T. Raitakari, Marjo-Riitta Järvelin, and Mika Ala-Korpela contributed epidemiological, genetic, or clinical data. Sanna Kuusisto, Minna K. Karjalainen, Therese Tillin, and Antti J. Kangas performed statistical analyses. Michael V. Holmes and Johannes Kettunen contributed to the genetic analyses and formulation of the study concept. All authors provided feedback on the writing of the manuscript. Mika Ala-Korpela acts as the guarantor for the work.

Conflict of interests
Antti J. Kangas is employed by Nightingale Health Plc. (Helsinki, Finland), a company providing metabolic profiling services. Veikko Salomaa has consulted for Sanofi and received a modest honorarium from this company. He also has ongoing research collaboration with Bayer Ltd. (all unrelated to the present study). All other authors report no conflict of interests.

Funding
This study was funded by the EU, Academy of Finland, Social Insurance Institution of Finland, Juho Vainio Foundation, Paavo Nurmi Foundation, Finnish Foundation for Cardiovascular Research, Finnish Cultural Foundation, Sigrid Juselius Foundation; Emil Aaltonen Foundation, Yrjö Jahnsson Foundation, Novo Nordisk Foundation, and others. The funders had no role in study design or reporting.

References
1 Saleheen D, Scott R, Javad S, Zhao W, Rodrigues A, Picataggi A, et al. Association of HDL cholesterol efflux capacity with incident coronary heart disease events: a prospective case-control study. Lancet Diabetes Endocrinol. 2015;3(7):507–13. https://doi.org/10.1016/S2213-8587(15)00126-6
2 Rohatgi A, Khera A, Berry JD, Givens EG, Ayers CR, Wedin KE, et al. HDL cholesterol efflux capacity and incident cardiovascular events. N Engl J Med. 2014;371(25):2383–93. https://doi.org/10.1056/NEJMoa1409065
3 Khera AV, Demler OV, Adelman SJ, Collins HL, Glynn RJ, Ridker PM, et al. Cholesterol efflux capacity, high-density lipoprotein particle number, and incident cardiovascular events: an analysis from the JUPITER trial (justification for the use of statins in prevention: an intervention trial evaluating rosuvastatin). Circulation. 2017;135(25):2494–504. https://doi.org/10.1161/CIRCULATIONAHA.116.025678
4 Cahill LE, Sacks FM, Rimm EB, Jansen MK. Cholesterol efflux capacity, HDL cholesterol, and risk of coronary heart disease: a nested case-control study in men. J Lipid Res. 2019;60(8):1457–64. https://doi.org/10.1194/jlr.P093823
5 Nicholls SJ, Puri R, Ballantyne CM, Jukema JW, Kastelein JJP, Koenig W, et al. Effect of infusion of high-density lipoprotein mimetic containing recombinant apolipoprotein A-I milano on coronary disease in patients with an acute coronary syndrome in the MILANO-PILOT trial: a randomized clinical trial. JAMA Cardiol. 2018;3(9):806–14. https://doi.org/10.1001/jamacardio.2018.2112
6 Nicholls SJ, Andrews J, Kastelein JJP, Merkely B, Nissen SE, Ray KK, et al. Effect of serial infusions of CER-001, a pre-β high-density lipoprotein mimetic, on coronary atherosclerosis in patients following acute coronary syndromes in the CER-001 atherosclerosis regression acute coronary syndrome trial: a randomized clinical trial. JAMA Cardiol. 2018;3(9):815–22. https://doi.org/10.1001/jamacardio.2018.2121
7 Rader DJ. Apolipoprotein A-I infusion therapies for coronary disease: two outs in the ninth inning and swinging for the fences. JAMA Cardiol. 2018;3(9):799–801. https://doi.org/10.1001/jamacardio.2018.2168
8 Karjalainen MK, Holmes MV, Wang Q, Anufrieva O, Kähönen M, Lehtimäki T, et al. Apolipoprotein A-I concentrations and risk of coronary artery disease: a Mendelian randomization study. Atherosclerosis. 2020;299:66–73. https://doi.org/10.1016/j.atherosclerosis.2020.02.002
9 Gille A, D’Andrea D, Tortorici MA, Hartel G, Wright SD. CSL112 [apolipoprotein A-I [human]] enhances cholesterol efflux similarly in healthy individuals and stable
atherosclerotic disease patients. *Arterioscler Thromb Vasc Biol.* 2018;38(4):953–63. https://doi.org/10.1161/ATVBAHA.118.310538

10 Low-Kam C, Rhainds D, Lo KS, Barhdadi A, Boulé M, Alem S, et al. Variants at the APOE /C1/C2/C4 locus modulate cholesterol efflux capacity independently of high-density lipoprotein cholesterol. *J Am Heart Assoc.* 2018; 7(16):e009545. https://doi.org/10.1161/JAHA.118.009545

11 Kuusisto S, Holmes MV, Ohukainen P, Kangas AJ, Karsikas M, Tiainen M, et al. Direct estimation of HDL-mediated cholesterol efflux capacity from serum. *Clin Chem.* 2019;65(8):1042–50. https://doi.org/10.1373/clinchem.2018.299222

12 Würtz P, Kangas AJ, Soininen P, Lawlor DA, Davey Smith G, Ala-Korpela M. Quantitative serum nuclear magnetic resonance metabolomics in large-scale epidemiology: a primer on -Omic technologies. *Am J Epidemiol.* 2017;186(9):1084–96. https://doi.org/10.1093/aje/kwx016

13 Josefs T, Wouters K, Tietge UJF, Annema W, Dullaart RPF, Vaisar T, et al. High-density lipoprotein cholesterol efflux capacity is not associated with atherosclerosis and prevalence of cardiovascular outcome: the CODAM study. *J Clin Lipidol.* 2020;14:122–32.e4. https://doi.org/10.1016/S1933-2874(19)30319-8

14 Davey Smith G, Lawlor DA, Harbord R, Timpson N, Day I, Ebrahim S. Clustered environments and randomized genes: a fundamental distinction between conventional and genetic epidemiology. *PLoS Med.* 2007;4(12):e352. https://doi.org/10.1371/journal.pmed.0040352

15 Rader DJ, Tall AR. The not-so-simple HDL story: is it time to revise the HDL cholesterol hypothesis? *Nat Med.* 2012;18(9):1344–6. https://doi.org/10.1038/nm.2937

16 Kontush A. HDL and reverse remnant-cholesterol transport (RRT): relevance to cardiovascular disease. *Trends Mol Med.* 2020;26(12):1086–100. https://doi.org/10.1016/S1471-4914(20)30184-2

17 Richardson TG, Sanderson E, Palmer TM, Ala-Korpela M, Ference BA, Davey Smith G, et al. Evaluating the relationship between circulating lipoprotein lipids and apolipoproteins with risk of coronary heart disease: a multivariable Mendelian randomisation analysis. *PLoS Med.* 2020;17(3):e1003062. https://doi.org/10.1371/journal.pmed.1003062

**Correspondence:** Mika Ala-Korpela, Computational Medicine, Center for Life Course Health Research, Faculty of Medicine, P.O. Box 5000, FI-90014, University of Oulu, Oulu, Finland. Email: mika.ala-korpela@oulu.fi

**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Figure S1:** Manhattan plot showing the results of genome-wide association study of HDL-CEC with adjustment for HDL-C and triglycerides in the five Finnish population cohorts.

**Figure S2:** Regional association plots of the HDL-CEC associated loci, *LIPC* and *CETP*.

**Figure S3:** Regional association plots of the *LIPC* and *CETP* loci in the GWAS of HDL-C.

**Figure S4:** Forest plot showing the causal estimates for coronary artery disease.

**Table S1:** SNPs associated with HDL-CEC in the Finnish populations.

**Table S2:** Associations of previously reported CEC-associated SNPs with CEC in the Finnish populations.

**Table S3:** Phenotype associations of the HDL-CEC associated SNPs.