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1 Association analyses based on false discovery rate implicate 243

2 susceptibility loci for coronary artery disease

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Genome-wide association studies (GWAS) in coronary artery disease (CAD) have identified 113 114 66 loci at 'genome-wide significance' ($p < 5 \times 10^{-8}$) but a much larger number of putative loci at a false discovery rate (FDR) of 5%¹⁻⁴. Here, we leverage an interim release of UK Biobank 115 (UKBB) data to evaluate the validity of the FDR approach. We tested a CAD phenotype 116 117 inclusive of angina (SOFT; N_{cases}=10,801) as well as a stricter definition without it (HARD; 118 N_{cases}=6,482) and selected the former for conducting a meta-analysis with the two most recent CAD GWASs²⁻³. This approach identified 13 new loci at genome-wide significance, 12 119 120 of which were in our previous 5% FDR list², and provided strong support that the remaining 121 FDR loci represent genuine signals. The set of 304 independent variants at 5% FDR in this 122 study explain 21.2% of CAD heritability and identified 243 loci that implicate pathways in 123 blood vessel morphogenesis as well as lipid metabolism, nitric oxide signaling and 124 inflammation.

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Previous GWAS studies of CAD risk¹⁻⁴ have interrogated a large number of cases and controls 127 128 but remain less well-powered than GWAS of quantitative traits⁵. UKBB was established to 129 improve understanding of the causes of common diseases including CAD, a leading health 130 problem around the world⁶. In addition to self-reported disease outcomes and extensive 131 health and life-style questionnaire data, the 502,713 participants are being tracked through 132 their NHS records and national registries (including cause of death and Hospital Episode Statistics). In July 2015, UKBB released genotypes imputed to the 1000 Genomes panel for 133 152,249 participants profiled with a SNP array harboring 820,967 variants comprising 134 135 common variants optimized for imputation, validated rare coding variants and sets of 136 phenotype-associated variants or their proxies (e.g. GWAS catalogue).

We set up The UKBiobank-CardioMetabolic-Consortium CHD working group to assess the use
of self-reported and hospital record data on CAD in UKBB and define the relevant case and
control subgroups to undertake genetic analyses of CAD risk.

140 The July 2015 release of UKBB comprises 10,801 genotyped individuals with an inclusive CAD 141 phenotype ('SOFT') that incorporates self-reported angina or other evidence of chronic 142 coronary heart disease, of which 6,482 have a more stringently defined CAD phenotype 143 ('HARD') of myocardial infarction and/or revascularisation (Fig. 1a). After QC we analysed the 144 SOFT and HARD cases separately against 137,914 controls for 9,149,595 variants present either in the CARDIoGRAMplusC4D 1000-Genomes GWAS² or the MIGen/CARDIoGRAM 145 Exome-chip study³⁻⁴. The SOFT definition was selected for the primary analysis based on 146 147 power calculations (**Supplementary Table 1**). We found 4 (SOFT and HARD), 1 (SOFT only) and 148 2 (HARD only) variants reaching genome-wide significance, all located in known CAD loci 149 (Supplementary Figure 1).

150 We then meta-analysed the UKBB data for each CAD definition with each of the two published 151 data sets (Supplementary Figure 2) applying a double genomic control correction. For both 152 the SOFT and HARD definitions, we validated all 66 known CAD loci (72 independent variants 153 with $p < 1.2xx10^{-3}$) with 43 and 37 respectively reaching genome-wide significance in this 154 study (Supplementary Table 2). Outside the known CAD loci (1 Mb window centred on the 155 published lead SNP) we found 9 new signals (in both SOFT and HARD) reaching genome-wide 156 significance (Table 1 and Fig. 2). The anticipated increase in power with the SOFT definition 157 (Supplementary Table 1) was attenuated by an inflation of the lambda statistic 158 (Supplementary Table 3), potentially due to a combination of larger sample size (i.e. 159 polygenicity) and a less homogeneous phenotype in the SOFT definition. Overall, there was 160 strong concordance between corresponding signals for SOFT and HARD (Fig. 1b, Supplementary Table 4); subsequent analyses were undertaken using the SOFT meta-analysisresults.

To look for additional signals beyond the 9 that reached genome-wide significance (**Fig. 2**) we performed an FDR analysis and selected 23 suggestive signals at 1% FDR (p < 1.55x10⁻⁶; **Supplementary Table 4**) outside known CAD loci which we validated in an independent sample of up to 4,412 cases and 3,910 controls from the German MI-Family-Studies V and VI and a Greek case-control study (**Supplementary Table 5**). In total, we identified 13 new genome-wide significant CAD loci in the combined discovery and replication sample (**Table 1**, **Supplementary Table 6**).

In our recent large-scale GWAS², we reported 162, mainly common, variants at an FDR 170 171 discovery cutoff of 5% showing conditional independent associations with the P_{ioint} test in 172 GCTA⁷. Twelve of the 13 new sentinel SNPs were present or had a proxy (r²>0.8) among these 173 162 variants². Fig. 3 shows a strong linear relationship between association signals for these 162 variants in the earlier² and current analysis, with overall greater significance levels in the 174 175 current meta-analysis. As expected, we observed an excess of small p-values for this set of 176 variants in the UK Biobank alone (Supplementary Figure 3a). Monte Carlo simulations show 177 that the expected number of replicated variants in the UK Biobank data is 56 (95%CI 42 – 69) 178 (Supplementary Figure 3b) and we found 58 variants after allowing for multiple testing (q-179 values < 0.05). This further confirms the validity of extended lists of associated variants based on FDR criteria. We therefore defined a new FDR list of association signals by performing an 180 approximate joint association analysis with the GCTA software⁷ as described elsewhere² using 181 the 11,427 SNPs with 5%FDR. We identified 304 independent variants at P_{ioint} < 10⁻⁴, clustering 182 183 in 243 putative CAD loci (Supplementary Table 7). The new 5%FDR set overlaps by 122 SNPs 184 with the old set (75.3%; including proxies at an $r^2 > 0.8$). We then assessed heritability using 185 the independent set of 304 SNPs and obtained a heritability estimate of 21.2%. The 186 contribution to this heritability estimate of the 13 new loci (Table 1) was 1.03% whereas the 187 new and known genome-wide significant CAD loci together explained 8.53% of CAD heritability. To further assess the validity and utility of the 5%FDR set, we tested the ability to 188 189 predict CAD using genetic risk scores (GRS) based on either the 5%FDR SNPs (GRS1) or only 190 CAD variants reaching genome-wide significance (GRS2; **Online Methods**) in an independent sample, EPIC-CVD⁸, comprising 7910 CHD cases and 12958 controls. In a model with age and 191 192 sex, GRS1 increased the C-index by 0.25% compared to GRS2 (Supplementary Table 8). GRS1 193 improved the point estimates of the HR compared to GRS2 mainly in the second (from 0.9116 to 0.8314) and fourth quintile (from 1.0437 to 1.176), Supplementary Figure 4. 194

195 We then explored the biology of the 13 new genome-wide significant CAD risk loci; 196 Supplementary Figure 5 shows regional association plots. Supplementary Figure 6 provides 197 in silico functional annotation (Online Methods) for each lead variant and its proxies (1000 198 Genomes). We found compelling evidence to implicate candidate genes ITGB5, TGB1, PDE5A, 199 ARHGEF26, FN1, CDH13, and HNF1 (detailed in Supplementary Note). The risk allele of 200 rs150512726 (proxy for rs142695226; Table 1), causes a 3 amino acid deletion within the 201 cytoplasmic tail of integrin subunit beta 5 (ITGB5), part of a heterodimer which regulates the 202 activation of latent TGFB1 (Transforming growth factor beta 1)⁹⁻¹⁰. The intronic variant (rs8108632; Table 1) we identified in TGFB1, further implicates the TGFB1 pathway in CAD 203 risk. TGFB1 is known to have important roles in endothelium and vascular smooth muscle¹¹ 204 205 but has not been widely studied in atherosclerosis, though a recent study implicates TGF? signalling downstream of CDKN2B in the CDKN2BAS cardiovascular risk locus¹². eQTL analyses 206 suggested candidate CAD risk genes (TDRKH, FN1, ARHGEF26, PDE5A, ARNTL, and CDH13) in 207 208 six new loci (Supplementary Table 9). For example, the lead variant rs7678555 (Table 1) was 209 found to be a strong eQTL (p=8.1x10⁻¹³) for PDE5A only in aorta from CAD patients 210 (STARNET¹³; Supplementary Table 9) although its regulatory potential was modest using 211 functional prediction tools (Online methods). PDE5A encodes a cGMP-specific 212 phosphodiesterase which is important for smooth muscle relaxation in the cardiovascular 213 system where it regulates nitric-oxide-generated cGMP¹⁴. Furthermore, mining eQTL data in 214 tissues from CAD patients (STARNET) showed several other instances of eSNPs (TDRKH, FN1, *CDH13*; **Supplementary Table 9**) having no effect in tissues from non-CAD patients (GTEx¹⁵), 215 216 highlighting the need to expand efforts to map regulatory elements in disease tissues.

217 Other candidate genes fit with emerging data on atherosclerosis mechanisms. For example, a knockout mouse for ARHGEF26 on a hyperlipidemic background resulted in reduced 218 atherosclerosis and plaques with reduced macrophage content¹⁶. Similarly, *FN1* expression is 219 220 increased in plaques and mouse models have demonstrated a causal role for fibronectin-1 in 221 the development and progression of atherosclerosis¹⁷⁻¹⁸. Finally, we undertook a phenome 222 scan to assess pleiotropy (**Supplementary Table 10**). Several of the new lead SNPs (or a proxy) 223 had robust associations ($p < 5x10^{-8}$) with traditional CAD risk factors such as LDL-cholesterol 224 (HNF1A and FN1), blood pressure (PRDM8/FGF5) and BMI (SNRPD2).

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We next evaluated the broader functional relationships among genes associated with variants (N=11,427) at 5%FDR. The 5%FDR set was annotated for eQTLs which, when present, were mainly found in atherosclerotic aortic wall (25%) or internal mammary artery (22%) of CAD patients (STARNET¹³; **Supplementary Table 9**). In GTEx¹⁵, eQTLs were mainly found in subcutaneous fat (**Supplementary Table 9**; **Supplementary Figure 7**).

Prior pathway analyses of GWAS CAD loci have highlighted genes involved in lipid metabolism,
 cellular movement, and processes such as tissue morphology and immune cell trafficking¹.

233 Analysis of 357 genes, selected as either eQTLs and/or the nearest gene to a 5%FDR 234 independent variant in this study (N=304), with the Ingenuity Knowledge base confirmed the 235 above findings¹ highlighting cardiovascular system development and function ($p = 1.31 \times 10^{-1}$ ¹⁶), organismal development ($p = 1.31x10^{-16}$) and survival ($p = 1.52x10^{-16}$) as the most 236 237 significant processes. In addition to canonical pathways related to lipid metabolism, 238 extracellular matrix, inflammation and nitric oxide production, the 357 gene set showed enrichment for angiogenesis and signalling by the pro-angiogenic growth factor VEGF 239 (Supplementary Figure 8). We also applied DEPICT¹⁹ with the full distribution of 5%FDR 240 241 signals (**Online Methods**) to search for enriched gene sets. Blood vessel development, which includes angiogenesis, was in the top 10 ($p < 6.67 \times 10^{-12}$) DEPICT Grouped-GeneSets 242 (GO:0001568; Fig. 4, Supplementary Figure 9, Supplementary Table 11). 243

Ingenuity built 5 networks out of the 357 genes with the largest three integrating 12 of the
new candidate CAD risk genes with 67 candidate genes in known CAD loci (Supplementary
Table 12). In total, the 5 networks comprise 66.4% of the 357 genes.

247 This is the largest CAD genetic study to assess simultaneously common and rare (MAF < 248 1%)/low-frequency (MAF 1-5%) variants. In total, 101 low-frequency and 3 rare variants 249 reached genome-wide significance among all 5%FDR markers (N=11,427). This apparent 250 paucity in rare variants which has also been reported for type 2 diabetes²⁰, is likely due to lack of power compared to studies of quantitative traits e.g. a study of adult height in ~700,000 251 individuals has reported 32 rare variants⁵. As expected, lower-frequency variants tend to have 252 253 stronger effects compared to common variants (Supplementary Figure 10) with the exception 254 of rs2891168 in CDK2NB-AS1 (MAF 48.7%; OR 1.19; Supplementary Table 13). The intergenic 255 variant rs186696265 which had the largest OR (1.62) in our study is known to affect LDL cholesterol levels²¹. 256

Our findings highlight the importance of the FDR approach to define an extended list of associated variants. As we have previously proposed¹⁻², suggestive association signals in wellpowered GWAS such as this one can substantially improve our knowledge of disease architecture at only a modest penalty implied by the 5%FDR. We have demonstrated the potential value of the new 5%FDR list in improving prediction of CAD risk and implicating new networks underlying CAD pathophysiology. This extended list of candidate genes provides a powerful resource for functional studies.

- 264
- 265 URLs
- 266 www.ukbiobank.ac.uk/
- 267 <u>GWAS catalogue: https://www.ebi.ac.uk/gwas/</u>
- 268 <u>GTEx portal: http://www.gtexportal.org/home/</u>
- 269 <u>PhenoScanner: http://www.phenoscanner.medschl.cam.ac.uk/</u>
- 270 Ingenuity Knowledge Base: http://www.ingenuity.com/science/knowledge-
- 271 <u>base?utm_source=Blog&utm_medium=link&utm_campaign=Doug%20Bassett%20ASHG%20</u>
- 272 <u>2014</u>

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318 AUTHOR CONTRIBUTIONS

- 319 Writing group (wrote and edited manuscript): C.P.N., A.G., A.S.B., S.K., T.R.W., E.M., I.N., 320 J.C.H., O.G., H.S., M.F., J.D., N.J.S., H.W., P.D. All authors contributed and discussed the results, 321 and commented on the manuscript. Data generation & cohorts: A.S.B., O.G., T.J., L.Z., S.E.H., 322 E.A., T.L.A., E.P.B., J.C.C., R.C., P.C., R.M.C., R.E., E.E., P.W.F., C.G., D.G., A.H., J.M.M.H., E.I., 323 A.K., T.K., T.K., T.L., X.L., Y.L., W.M., R.McP., A.M., M.Pujades-R., A.F.S., M.J.S., P.A.Z., R.J.F.L., 324 E.Z., J.E., G.D., H.S., J.D., N.J.S., H.W., P.D. Phenotype data (UK Biobank, replication): C.P.N., 325 A.S.B., I.N., J.C.H., O.G., B.D.K., J.S.K., R.J.F.L., R.S.P., M.R., M.T., I.T., E.Z., J.E., G.D., H.S., J.D., 326 N.J.S., H.W., P.D. Statistical analysis: C.P.N., A.G., A.S.B., S.K., T.J., M.F. Functional annotation: 327 C.P.N., S.K., T.W., A.S.B., R.E., A.R., E.E.S., J.L.M.B. Biological and clinical enrichment and 328 pathway analyses: E.M., P.D.
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342

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394 Figure legends

395 Figure 1. (a) Diagram depicting the CAD phenotype definition in UK Biobank. HARD CAD 396 defined as fatal or non-fatal myocardial infarction (MI), PTCA (percutaneous transluminal 397 coronary angioplasty), or coronary artery bypass grafting (CABG). SOFT CAD includes HARD 398 CAD as well as chronic ischaemic heart disease (IHD) and angina. UK Biobank self-reported 399 data: 'Vascular/heart problems diagnosed by doctor' or 'Non-cancer illnesses that self-400 reported as angina or heart attack'. Self-reported surgery defined as either PTCA, CABG or 401 triple heart bypass. HESIN hospital episodes data and death registry data using diagnosis and 402 operation - primary and secondary cause: MI defined as hospital admission or cause of death due to ICD9 410-412, ICD10 I21-I24, I25.2; PTCA is defined as hospital admission for PTCA 403 404 (OPCS-4 K49, K50.1, K75); CABG is defined as hospital admission for CABG (OPCS-4 K40 – 405 K46); Angina or chronic IHD defined as hospital admission or death due to ICD9 413, 414.0, 406 414.8, 414.9, ICD10 I20, I25.1, I25.5-I25.9. (b) Radar plot highlighting the proportions (%) of 407 signals between the HARD and SOFT CAD phenotype definitions based on the 5%FDR results (Supplementary Table 4); MAF = minor allele frequency, $p < 5x10^{-8}$ marks variants reaching 408 genome-wide significance, OR = odds ratio (OR > 1.05 corresponds to 85% power to detect a 409 410 signal (alpha < 0.05) in the SOFT analysis). The results for all six subgroups of variants assessed 411 did not differ statistically between the two phenotype definitions (p>0.1)

Figure 2. Transposed Manhattan plot showing the SOFT meta-analysis results under an additive model. The *P*-values are truncated at $-\log_{10}(P) = 20$. The red dotted lines are at GWAS (*P*=5x10⁻⁸) and 5% FDR significance (*P*=6.28x10⁻⁵). The known CAD risk loci are shown in black (**Supplementary Table 2**); *KSR2* and *ZNF507-LOC400684* had reached genome-wide significance under a recessive model². The exome chip markers are shown with an *. The 13 417 novel CAD loci which reached genome-wide significance in our study (including replication
418 data; Table 1), are written in brown font.

Figure 3. Single marker p-value comparison of the 5% FDR variants in the published 419 CARDIoGRAMplusC4D 1000Genomes CAD GWAS meta-analysis² and current FDR study. Of 420 421 the 162 variants which had p $<5x10^{-5}$ in the CAD 1000Genomes GWAS, 116 had a match or good proxy ($r^2 > 0.8$) in the new FDR list (red circles). SNPs in green (n=7) were present in the 422 423 earlier FDR list and reached genome-wide significance in the current analysis. 424 Figure 4. Heat map showing the DEPICT gene set enrichment results with zoom-in on a subset of the results. 556 gene sets are included which had evidence of enrichment at 1% FDR. The 425 426 x– axis shows the gene name, which is predicted to be included in the reconstituted gene set 427 indicated in the y – axis. The color red indicates higher Z-score, where Z-score is a value 428 representing each gene's inclusion in DEPICT's reconstituted gene sets. Clustering was made 429 based on complete linkage method. Highlighted pathways in the cluster, include 430 angiogenesis, blood vessel development and morphogenesis.

431

432 Online Methods

433 Phenotype Definitions & Power calculation

UKBB recruited 502,713 individuals aged 40-69 years from England, Scotland and Wales
between 2006 and 2010 (94% of self-reported European ancestry). HARD CAD was defined
as fatal or non-fatal myocardial infarction (MI), percutaneous transluminal coronary
angioplasty (PTCA), or coronary artery bypass grafting (CABG). SOFT CAD includes all HARD
CAD as well as chronic ischemic heart disease (IHD) and angina. Controls were defined as
patients which were not a SOFT case after exclusions (listed below). All conditions were
defined by either self-reported, hospital episode or death registry data.

Exclusions were made for aneurysm and atherosclerotic cardiovascular disease using
hospital admissions, or cause of death, codes ICD9 414.1, ICD 10 I25.0, I25.3, I25.4, and not
having MI, PTCA, CABG, Angina or chronic IHD as defined above.

444 Susceptibility effect sizes in MI cases and an inclusive CAD definition were very similar in 445 the earlier GWAS². We hypothesized that the detailed clinical information in UKBB might 446 enhance the search for novel loci by further broadening the CAD phenotype to increase 447 sample size.

449 GWAS and meta-analyses

478

- 450 All participants gave written consent for participation in genetic studies, and the protocol of each study was approved by the corresponding local research ethics committee or 451 452 institutional review board. Participating cohorts in the 1000 Genomes and Exome GWAS studies are described elsewhere^{2,3}. UK Biobank (UKBB samples) were excluded due to 453 454 withdrawn consent, sex mismatches (n=182), Biobank/Believe QC exclusions (n=406) and 455 sample relatedness (n=3,481) determined as Kinship>0.088. GWAS analysis in UKBB was restricted to variants with results available in the published GWAS² or Exome³⁻⁴ dataset. 456 457 Further exclusions included poorly imputed (info<0.4) or monomorphic variants, duplicate variants across data sets, variants that deviated strongly from Hardy-Weinberg Equilibrium 458 in European ancestry controls ($p<1x10^{-9}$), variants with an effect allele frequency in 459 460 European ancestry samples that differed strongly (i) from 1000G European panel, (ii) from 461 GWAS/Exome data, (iii) between arrays (UKBB vs UK-BiLEVE), and (iv) across genotyping 462 batches. Variants that did not produce a valid result or estimated extreme log odds ratios (|beta|>4) were also excluded after analysis. Cluster plots lead variants and of proxies 463 464 were visually inspected.
- 465 We ran the GWAS under an additive frequentist mode of inheritance for each variant using 466 the dosages from the imputed data, adjusting for array (UK Biobank vs UK BiLEVE) and the 467 first five principal components using SNPTEST. Age and sex were not adjusted for to 468 maximize the power to detect associations with diseases that have a prevalence <10%²². 469 Population stratification was assessed and standard errors were adjusted using the 470 genomic inflation statistic (λ).
- 471 Association summary statistics (after λ correction) from the UKBB were combined with the 472 1000 Genomes (1000G) imputed GWAS results² and the Exome results³ via two separate 473 fixed-effect inverse-variance weighted meta-analysis implemented in GWAMA²³. We 474 applied post meta-analysis λ correction in each instance. We identified 36,460 variants 475 present in both the 1000G imputed GWAS and the Exome results. We retained the variants 476 from the 1000G imputed GWAS if the median info score was 1, otherwise we retained the 477 results from the Exome data.

479 Comparison of SOFT vs HARD peak variant lists at 5% q-value

480 The false discovery rate (FDR) following the meta-analysis with UKBB was assessed using a step-up procedure in the qqvalue Stata program²⁴ as it is well controlled under positive 481 regression-dependency conditions. We used the Simes method to generate q-values for 482 483 the 8.9M variants. The p-value cut-off for a q-value of 5% for HARD was 7.24x10⁻⁵ and SOFT was 6.28x10⁻⁵. Peak SNPs were identified in a 1cM window. There is an exact overlap of 484 485 155 variants between the 2 peak variant lists, however, using the 1cM window the overlap 486 increases to 206 variants. Both the lists were annotated and classified into 6 categories 487 (exome chip, indels, Odds Ratio (OR)>1.05, p<5e-8, MAF<5% and exonic). The proportions 488 were calculated in each of the 6 categories and plotted as a radar plot (Fig. 1b). Monte 489 Carlo simulations were used to assess the post-hoc power of the UKBB interim data to 490 replicate the 155 variants. The 1000G GWAS effect sizes ("betas") are expected to be subject to winner's curse inflation so were shrunken (towards the null) by application of 491 the FIQT procedure²⁵. Effect sizes for firmly established CAD loci were systematically lower 492 493 for SOFT compared to the HARD phenotype (Supplementary Table 1) noting that HARD 494 closely corresponds to the CAD phenotype in reference 2. Betas were therefore further 495 shrunken by a factor log $(1.059)/\log(1.072) = 0.82$ (Supplementary Table 1). 10,000 496 replicates were then randomly drawn from the vector of shrunken betas and the 497 corresponding UKBB standard errors, to allow for variation in genotype call rates, 498 imputation quality and allele frequency and to calculate Wald association statistics. 499 Multiple testing of 155 variants was allowed for by controlling the FDR to 5% with a stepup procedure encoded in the *multproc*²⁶ Stata[™] program. The average expected number 500 501 of replicated variants was 56 (95%CI 42 – 69). Testing the 5% FDR variants (Supplementary 502 Table 7) in UKBB with a model adjusted for age and sex gave concordant results to the 503 unadjusted model (data not shown).

505 GCTA & Heritability analysis

506 We used the GCTA software⁷ to perform joint association analysis in (SOFT) meta-analysis 507 results. This approach fits an approximate multiple regression model using summary-level 508 meta-analysis statistics and LD corrections estimated from a reference panel (here the 509 UKBB sample). We adopted a chromosome-wide stepwise selection procedure to select 510 variants and estimate their joint effects at i) a genome-wide significance level (pJoint \leq 511 5x10⁻⁸) in the totality of meta-analysed variants (n[~] 9M; Supplementary Figure 10, 512 **Supplementary Table 11**) and ii) a Bonferroni-corrected pJoint<1x10⁻⁴ corresponding to the number of independent LD bins ($r^2 < 0.1$) in the 5% FDR variant list (n=11,427; 513 514 Supplementary Table 6).

- 515 Heritability calculations were based on a multifactorial liability-threshold model, 516 implemented in the INDI-V²⁷ calculator (<u>http://cnsgenomics.com/shiny/INDI-V/</u>), under 517 the assumption of a baseline population risk (K) of 0.0719²⁸ and a twins heritability (H_L^2) of 518 0.4. Multiple regression estimates from the GCTA joint association analysis were used to 519 estimate heritability for the 304 independent CAD risk variants within the 5% FDR list.
- 520

504

521 Genetic risk score analysis

GRS analysis was undertaken in the EPIC-CVD study⁸ which comprises 7910 CAD cases and 522 12958 controls (Supplementary Note). We considered either all known and new lead CAD 523 524 risk variants reaching genome-wide significance (GRS2; Supplementary Table 2 and Table 525 1) or the 304 variants in the 5% FDR set (GRS1; **Supplementary Table 7**). We used variants 526 with an INFO score filter of 0.4 in EPIC-CVD and replaced missing ones with proxies (r2 > 527 0.8 in 1000 Genomes European participants). GRS1 comprised 280 variants and GRS2 71. 528 The raw GRS was obtained by summing the dosages of these variants for all individuals. 529 We then fitted a Prentice weighted cox regression model for each GRS, adjusting for age 530 and sex, to obtain survival forecasts and calculate the C indices. Statistical analyses were 531 performed using R 3.3.3 and STATA 13.1. Variant extraction was done using qctool 1.4.

532

533 Functional annotation

eQTLs: For associations between the 304 independent variants (5% FDR) and gene expression traits we searched for expression quantitative trait loci (eQTLs) in the Stockholm-Tartu Atherosclerosis Reverse Network Engineering Task (STARNET) RNA-seq dataset¹³ and the Genotype-Tissue Expression¹⁵ (GTEx) portal. eQTLs were included if the best eSNP (i.e. the variant with the most significant association with gene expression in cis) was in high LD (r²>0.8) with the CAD lead SNP. 540 Regulatory elements: We functionally annotated each of the 13 lead variants and their proxies $(r^2>0.8)$ using HaploregV4²⁹. Overlap with regulatory elements including 541 542 chromosome state segmentation, DNase hypersensitivity, and transcription factor binding (TFB) as determined by the ENCODE³⁰ and Roadmap Epigenome projects³¹, and predicted 543 effects on TFB based on regulatory motifs from TRANSFAC³² and JASPAR³³ were identified 544 using HaploregV4¹⁹ and the UCSC genome browser. Variants were then scored using three 545 different bioinformatics tools that help prioritise causal disease variants. Combined 546 Annotation Dependent Depletion (CADD)³⁴ incorporates a range pathogenicity prediction 547 tools to provide a genome-wide score (C-score) for each test variant from its pre-calculated 548 549 database of ~8.6 billion genetic variants. High scores indicate variants that are not 550 stabilized by selection and are more likely to be disease-causing and low scores indicate 551 evolutionary stable non-damaging variants. The top 10% of likely functional variants will have a C-score >10 and top 1% of variants will have a C-score >20. Genome-wide 552 annotation of variants (GWAVA)³⁵ predicts the functional impact of noncoding variants 553 554 based on genomic and epigenomic annotations and provides scores between 0 and 1 with 555 higher scores indicating variants that are more likely to be functional. RegulomeDB³⁶ 556 annotates and scores variants in seven categories based datasets such as ENCODE. Scores 557 of 1-2 variants likely to affect TFB, 3 less likely to affect binding, 4-6 relate to variants with 558 minimal binding evidence and 7 is for variants with no regulatory annotation.

559 Phenome-scan: look ups in other common traits were performed using the PhenoScanner560 database as described in ref 37.

562 Pathway analysis

561

563 **DEPICT:** DEPICT¹⁹ is a computational tool which performs gene set enrichment analyses to 564 prioritize genes in associated GWAS loci with probabilistically predefined gene sets based on Gene Ontology terms, canonical pathways, protein-protein interaction subnetworks 565 566 and rodent phenotypes; reconstituted gene sets are detailed in refs 19 and 38. Input to 567 our analysis were the 11,427 CAD variants (FDR 5%) of which 11,311 were annotated in 568 DEPICT. We constructed loci as previously described (beta version 1.1, release 194, 569 www.broadinstitute.org/mpg/depict). Analysis was performed with default parameters 570 (50 repetitions to compute FDRs, 500 permutations to adjust for biases, such as gene 571 length). The 11,311 variants were collapsed to 288 loci which were used in the gene set 572 enrichment analyses. Correlated gene sets were grouped together based on gene 573 membership to expedite data interpretation.

574 Ingenuity: Genes were selected using 304 independent SNPs (5% FDR) based on eQTLs 575 (Supplementary Table 9) and physical proximity (included overlapping genes on opposite 576 strands or at equal distance from the SNP). Spliced ESTs and putative transcripts were not 577 included. Network analysis was performed using the Ingenuity Pathway Analysis software 578 (www.ingenuity.com). We considered molecules and or relationships available in The IPA 579 Knowledge Base (IKB) for human OR mouse OR rat and set the confidence filter to Experimentally Observed OR High (Predicted). Networks were generated with a maximum 580 581 size of 70 genes and up to 10 networks were allowed. Networks are ranked according to 582 their degree of relevance to the 'eligible' molecules in the query data set. The network 583 score is based on the hypergeometric distribution and is calculated with the right-tailed 584 Fisher's Exact Test. The significance p-value associated with enrichment of functional 585 processes is calculated using the right-tailed Fisher Exact Test by considering the number

of query molecules that participate in that function and the total number of molecules that 586 587 are known to be associated with that function in the IKB. 588 589 590 Data Availability: Meta-analysis summary statistics for the variants considered in this 591 study for association with CAD (SOFT definition) are available at http://www.cardiogramplusc4d.org/data-downloads/. 592 593 594 595 596

597 Method References

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- 637

	Markername	CHR	POS (hg19)	EA	EAF	Functional	UKBB+CoG/Exome			Meta analysis	
Locus Name						Evidence	OR (95% CI)	P-value	FDR Q-value	OR (95% CI)	P-value
TDRKH	rs11810571	1	151762308	G	0.849	eQTL/coding	1.060 (1.039 , 1.082)	2.21x10 ⁻⁸	8.05x10 ⁻⁵	1.057 (1.036 , 1.079)	4.24x10 ⁻⁸
FN1	rs1250229*	2	216304384	Т	0.256	eQTL/coding	1.072 (1.052 , 1.092)	1.85x10 ⁻¹³	2.05x10 ⁻⁹	1.071 (1.051 , 1.091)	2.77x10 ⁻¹³
RHOA	rs7623687	3	49448566	А	0.855	none	1.074 (1.049 , 1.100)	3.72x10 ⁻⁹	1.62x10⁻⁵	1.076 (1.052 , 1.101)	3.44x10 ⁻¹⁰
UMPS/ITGB5	rs142695226	3	124475201	G	0.138	eQTL/coding	1.069 (1.045 , 1.094)	1.00x10 ⁻⁸	3.98x10⁻⁵	1.071 (1.048 , 1.095)	1.53x10 ⁻⁹
ARHGEF26	rs12493885*	3	153839866	С	0.886	eQTL	1.074 (1.047 , 1.101)	3.29x10 ⁻⁸	1.15x10 ⁻⁴	1.073 (1.047 , 1.101)	3.16x10 ⁻⁸
PRDM8/FGF5	rs10857147	4	81181072	Т	0.275	none	1.056 (1.036 , 1.075)	8.96x10 ⁻⁹	3.60x10 ⁻⁵	1.054 (1.036 , 1.073)	5.66x10 ⁻⁹
PDE5A/MAD2L1	rs7678555	4	120909501	С	0.30	eQTL	1.049 (1.031 , 1.069)	1.43x10 ⁻⁷	4.25x10 ⁻⁴	1.052 (1.034 , 1.070)	1.32x10 ⁻⁸
HDGFL1	rs6909752	6	22612629	Α	0.351	none	1.051 (1.034 , 1.069)	5.59x10 ⁻⁹	2.35x10 ⁻⁵	1.051 (1.034 , 1.068)	2.19x10 ⁻⁹
ARNTL	rs3993105	11	13303071	Т	0.704	none	1.048 (1.030 , 1.067)	1.06x10 ⁻⁷	3.33x10 ⁻⁴	1.048 (1.031 , 1.066)	4.77x10 ⁻⁸
HNF1A	rs2244608	12	121416988	G	0.355	coding	1.053 (1.035 , 1.070)	2.32x10 ⁻⁹	1.06x10 ⁻⁵	1.053 (1.035 , 1.070)	7.74x10 ⁻¹⁰
CDH13	rs7500448	16	83045790	А	0.752	eQTL	1.061 (1.040 , 1.082)	5.14x10 ⁻⁹	2.18x10 ⁻⁵	1.063 (1.043 , 1.083)	4.76x10 ⁻¹⁰
TGFB1	rs8108632	19	41854534	Т	0.488	none	1.049 (1.031 , 1.067)	5.88x10 ⁻⁸	1.95x10 ⁻⁴	1.048 (1.031 , 1.066)	4.04x10 ⁻⁸
SNRPD2	rs1964272	19	46190268	G	0.510	none	1.045 (1.028 , 1.063)	2.29x10 ⁻⁷	6.15x10 ⁻⁴	1.047 (1.030 , 1.064)	2.46x10 ⁻⁸

638	Table 1 - Novel variants reaching gend	me-wide significance (P<5x10 ⁻⁸) in the combined (discovery and re	plication) SOFT meta-analysis
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639 *Exome marker

640 EA: effect allele; EAF: Effect allele frequency; CoG = CARDIoGRAMplusC4D 1000G GWAS; Exome = Exome array analysis; UKBB = UK Biobank;

641 Discovery sample comprised 71,602 cases and 260,875 controls (for exome markers 53,135 and 215,611 respectively); Replication sample

642 comprised up to 4412 cases and 3910 controls. Functional evidence for the locus is given where the lead variant or a variant in high LD (r²>0.8)

643 is a coding change, has evidence as an expression quantitative trait locus (eQTL), or both. Further details of functional evidence are provided in

644 Supplementary Table 7 and Supplementary Figure 6.

645





Figure 2

	-log10(P-V	alue)						
20 I	10	5E-08	FDR 5%	0	SNP	EAF	OR P-Value	GENES (NEARBY)
					, rs11591147*	0.98	1.25 2.80E-10	PCSK9
	-	THE OWNER AND ADDRESS			- rs56170783	0.92	1.11 2.10E-12	PPAP2B
		1	10 - CT		- rs7528419	0.78	1.11 3.80E-27	SORT1
					- rs6689306	0.85	1.06 4.24E-08 1.05 1.50E-09	IL SR
					- rs67180937	0.68	1.07 8.50E-14	MIA3
					- rs16986953*	0.07	1.11 4.80E-10	AK097927
		-			rs585967	0.84	1.07 2.80E-08	APOB
		· -+			rs4299376*	0.32	1.06 5.70E-10	ABCG5/ABCG8
-			1.1		rs/568458	0.45	1.06 2.40E-13	7EB2 / AC074093 1
					rs114123510	0.03	1 13 2 90E-19	WDR12
					rs1250229*	0.26	1.07 2.77E-13	FNI
			The second second		rs13003675	0.36	1.04 1.70E-06	KCNJ13/GIGYF2
					- rs7623687	0.86	1.08 3.44E-10	RHOA
			1.00		120016240	0.14	1.07 1.53E-09	UMPS/IIGB5
				111115	1512493885*	0.89	1.07 3.16E-08	ARHGEE26
		1			, rs72627509	0.20	1.06 8.10E-08	REST/NOA1
			-300		rs10857147	0.28	1.05 5.66E-09	PRDM8
					rs7678555	0.30	1.05 1.32E-08	MAD2L1
		1.1.1		4	, rs6841581	0.15	1.07 4.60E-10	EUNRA GUOX1A3
	-				rs77335401	0.12	1.05 7.60E-05	SLC22A4 / SLC22A5
		1	- And all		, rs742115	0.48	1.04 2.90E-05	ADTRP / C6orf105
					, rs9349379*	0.41	1.11 1.00E-35	PHACTR1
				· / ·	rs6909752	0.35	1.05 2.19E-09	HDGFL1
			-		re4472337	0.80	1.08 2.80E-08	ANKSIA
					- rs56015508	0.79	1.06 1.10E-07	KCNK5
		A NEWS	State of Lot of		, rs12202017	0.70	1.07 6.00E-14	TCF21
				φ.	rs10455872	0.06	1.31 1.70E-49	SLC22A3/LPAL2/LPA/PLG
	·		:=-		re112370447	0.18	1.08 3.40E-13 1.05 9.60E=07	7022
			-	111112/	· rs11556924	0.66	1.07 6.30E-13	ZC3HC1
			- here		, rs3918226	0.07	1.13 1.60E-12	NOS3
					rs2083636	0.74	1.05 6.40E-08	LPL
		1 -			, rs2954029"	0.54	1.06 5.20E-13 1.191.30E-101	TRIB1
		-	1		rs111245230*	0.04	1.12 8.30E-07	SVEP1
			T	· · ·	' · rs507666*	0.19	1.08 1.30E-12	ABO
		1	14.75		,, rs1887318	0.43	1.06 4.10E-12	KIAA1462
				;	/, rs1870634	0.65	1.06 5.50E-13	CXCL12
			-	o //	// rs11191416	0.35	1.08 5.60E-09	CYP17A1/CNNM2/NT5C2
				111111	rs10840293	0.55	1.05 6.90E-09	SWAP70
					, rs201267813	0.07	1.05 1.20E-03	11p15_MRVI1 / CTR9
		1	- Long	₽,;'	rs3993105	0.70	1.05 4.77E-08	ARNTL
					re964184	0.31	1.06 2.00E-11 1.05 4 70E-06	ZNE259 / APO45 / APO41
					, rs2229357*	0.76	1.05 3.40E-06	LRP1
			- 70	= ,4	rs2681472	0.19	1.07 7.60E-11	ATP2B1
			-	::::::::::::::::::::::::::::::::::::::	rs10774625*	0.49	1.07 9.20E-14	SH2B3
				1	, rs11830157	0.38	1.03 1.70E-03	HNE14
					- rs11057830*	0.15	1.07 4.20E-09	SCARB1
		7. aufr		••••••••••••••••••••••••••••••••••••••	- rs1924981	0.33	1.05 1.90E-07	FLT1
		1-		· · · · · · · · · · · · · · · · · · ·	, rs11617955	0.89	1.09 4.10E-10	COL4A1/COL4A2
			1	÷.,	, rs10139550	0.42	1.05 1.80E-09	HHIPL1 SMAD3
	***				, . rs7164479	0.58	1.07 6.40E-18	ADAMTS7
			- 5	2 /	, rs2083460	0.88	1.07 1.40E-07	MFGE8 / ABHD2
			a martin a paint		· rs2071382	0.46	1.06 7.10E-13	FURIN / FES
					· ·rs24/616*	0.68	1.04 1.00E-06	CDH12
				11111°3*,	rs113348108	0.31	1.05 5.80E-08	SMG6
			1.1	· ·	rs9897596	0.52	1.04 3.10E-06	RAI1 / PEMT / RASD1
		1.00			· rs4643373	0.72	1.05 1.20E-06	UBE2Z
			A CONTRACTOR		- rs8068952*	0.23	1.07 1.40E-09	BCAS3 PMAIP1 / MC4R
				TILLES.	rs116843064*	0.23	1.17 2.90E-07	ANGPTL4
			1	₽./	, rs6511720*	0.88	1.14 7.90E-22	LDLR
			-		- rs10417115	0.06	1.07 2.30E-05	ZNF507 / LOC400684
		1	Laborer		- rs8108632	0.49	1.05 4.04E-08	TGFB1
		1 1		20	rs1964272	0.51	1.05 2.46E-08	SNRPD2
			and different	· · · · · · · · ·	rs28451064	0.12	1.14 2.60E-23	gene_desert / KCNE2
					rs180803	0.98	1.18 7.10E-10	POM121L9P / ADORA2A
				_				

Figure 3





