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Loss of Cardioprotective Effects at the ADAMTS7 Locus as a Result of Gene-Smoking Interactions

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1 Loss of cardio-protective effects at the *ADAMTS7* locus due to gene-smoking interactions

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21 Running Title: Gene*Smoking interaction, ADAMTS7 locus & CHD risk

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138 ABSTRACT

Background Common diseases such as coronary heart disease (CHD) are complex in etiology. The interaction of genetic susceptibility with lifestyle factors may play a prominent role. However, geneenvironment interactions for CHD have been difficult to identify. Here, we investigate interaction of smoking behavior, a potent lifestyle factor, with genotypes that have been shown to associate with CHD risk.

144 **Methods** We analyzed data on 60,919 CHD cases and 80,243 controls from 29 studies for gene-145 smoking interactions for genetic variants at 45 loci previously reported to associate with CHD risk. 146 We also studied 5 loci associated with smoking behavior. Study specific gene-smoking interaction 147 effects were calculated and pooled using fixed-effects meta-analyses. Interaction analyses were 148 declared to be significant at a *P-value* < 1.0×10^{-3} (Bonferroni correction for 50 tests).

Results We identified novel gene-smoking interaction for a variant upstream of the *ADAMTS7* gene. Every T allele of rs7178051 was associated with lower CHD risk by 12% in never-smokers (P-value: 1.3x10⁻¹⁶) compared to 5% in ever-smokers (P-value: 2.5x10⁻⁴) translating to a 60% loss of CHD protection conferred by this allelic variation in people who smoked tobacco (*Interaction P-value*: 8.7x10⁻⁵). The protective T allele at rs7178051 was also associated with reduced *ADAMTS7* expression in human aortic endothelial cells and lymphoblastoid cell lines. Exposure of human coronary artery smooth muscle cells to cigarette smoke extract led to induction of *ADAMTS7*.

156 **Conclusion** Allelic variation at rs7178051 that associates with reduced *ADAMTS7* expression 157 confers stronger CHD protection in "never-smokers" compared to "ever-smokers". Increased 158 vascular *ADAMTS7* expression may contribute to the loss of CHD protection in smokers.

159 Key words: Gene-smoking interaction, gene-environment interaction, coronary heart disease,160 ADAMTS7, smoking.

- 161 Word count: 269
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164 Clinical Perspective

165 1) What is new?

- Using data on 60,919 CHD cases and 80,243 controls, this study conducted geneenvironment interaction analyses to investigate effect modification by smoking behavior at
 established CHD and smoking related loci.
- Cardio-protective effects associated with allelic variation at the *ADAMTS7* locus were attenuated by 60% in people who smoked tobacco compared to those who did not smoke.
- Allelic variation at *ADAMTS7* associated with reduced CHD risk was associated with reduced
 ADAMTS7 expression in human aortic endothelial cells and lymphoblastoid cell lines.
- Exposure of human coronary artery smooth muscle cells to cigarette smoke extract led to induction of *ADAMTS7*.

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177 2) What are the clinical implications?

- These human genomic data provide new insights into potential mechanisms of CHD in
 cigarette smokers.
- Findings from this study also point towards the directional impact of the *ADAMTS7* locus on
 CHD.
- ADAMTS7 and its substrates have a specific role in cigarette smoking related CHD.
- Inhibition of ADAMTS7 is a novel potential therapeutic strategy for CHD that may have
 particular benefits in individuals who smoke cigarettes.
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194 INTRODUCTION

Coronary heart disease (CHD) is a complex disorder resulting from the interplay of lifestyle 195 and genetic factors.^{1, 2} Yet, gene-environment interactions for CHD have been difficult to identify. 196 Cigarette smoking is one of the strongest lifestyle risk factors for CHD but the underlying molecular 197 198 mechanisms of CHD in humans who smoke remain uncertain.³⁻⁵ Cigarette smoking accounts for one-fifth of all CHD events globally and is responsible for ~1.6 million deaths attributable to CHD 199 annually.6 Genome-wide association studies (GWAS) have improved our understanding on the 200 genetic predisposition to both CHD and smoking behavior.⁷⁻¹⁰ Joint or interactive effects of genetic 201 variation and smoking behavior in the etiology of CHD, however, remain poorly understood. GWAS 202 can provide new opportunities to investigate gene-smoking interactions. 203

We hypothesized that genetic predisposition to CHD is modified by cigarette smoking at loci discovered by GWAS to be associated with either CHD or smoking behavior. We conducted a focused experiment at 50 main-effect loci (45 for CHD and 5 for smoking behavior) using genetic data and information on smoking behavior in 60,919 CHD cases and 80,243 controls from 29 different studies. We report novel findings on gene-smoking interactions in CHD.

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221 METHODS

222 Summary of study Design

All studies participating in the CARDIoGRAMplusC4D consortium⁷⁻⁹ that had information 223 available on smoking status, CHD risk and genotypes at the 50 CHD and smoking behavior-224 225 associated loci were invited to participate. The current study had five inter-related components (Supplementary Figure-1). First, as part of the quality control, we investigated the association of 226 smoking status with CHD risk within each study. Second, we performed an updated analysis of all 227 the SNPs (± 50 KB) at the 45 established CHD loci to identify the variant with the strongest CHD 228 association in our study population at each established CHD locus. Effect estimates from each study 229 in association with CHD risk were obtained and pooled to identify the strongest CHD associated 230 variant ("lead variant"). Third, we investigated gene-smoking interactions at these 45 CHD loci and at 231 5 loci related to smoking behavior. Fourth, for loci demonstrating differential CHD associations by 232 smoking status, we mapped the interaction region, examined linkage disequilibrium (LD) across the 233 region and performed conditional analyses to identify independent genetic signals. Finally, for loci 234 exhibiting gene-smoking interaction in CHD, we assessed eQTL data for association of variants with 235 expression of local genes in available datasets and examined expression of these genes in multiple 236 cell types that play prominent roles in smoking-CHD pathobiology. 237

238 Harmonization of phenotypes and genotypes

Summary level estimates for each study were shared via a secure FTP site. We used 239 "ever-smoking" as a primary exposure and data were harmonized by uniformly characterizing 240 participants in each study into two categories, "ever-smokers" and "never-smokers". "Ever-smokers" 241 were defined as those who had smoked more than 100 cigarettes in a lifetime. For case-control 242 studies, information on "ever smoking" status collected at the time of enrollment was used for the 243 current analyses; whereas for prospective cohort studies, information on smoking status obtained at 244 the baseline visit was used for the current investigation. CHD was defined based on evidence from 245 angiography or history of verified myocardial infarction (MI), percutaneous coronary interventions 246 (PCI) or coronary artery bypass grafting (CABG) as published in CARDIoGRAMplusC4D projects.⁷⁻⁹ 247 Genotype data generated through GWAS (directly genotyped or imputed) or cardio-metabochip 248 (directly genotyped only) arrays were obtained from each study and all genetic data were aligned 249 using the build-37 reference panel. Imputed SNPs were removed if they had any of the following: (i) 250 a minor allele frequency of <1%; (ii) info score of <0.90; or (iii) confidence score <0.90. For each 251 252 study, GWAS data were imputed using the Phase II CEU HapMap reference population.¹¹ Standard quality control criteria were applied by each participating study, as described previously.⁷ All participating studies in the CARDIoGRAMplusC4D consortium were approved by their locally relevant institutional review boards and all participants gave written informed consent before their enrollment in each study.⁷⁻⁹

257 STATISTICAL ANALYSIS

258 Gene-smoking interaction analyses

Initial quality control and association of established CHD loci with CHD risk: As part of an initial 259 quality control, effect estimates from each study were obtained for "ever-smoking" status and CHD 260 risk using a case-control logistic regression model adjusted for age and sex. Each participating study 261 also assessed and, if needed, controlled for population stratification by including principal 262 components as covariates in the regression model as described earlier.⁷⁻⁹ To identify variant(s) with 263 the most significant association with CHD risk at established CHD loci in our study population, 264 logistic regression analyses were conducted by each participating study for all the SNPs flanking 265 266 (±50 kb) the lead variant previously reported at each CHD locus. Effect estimates and standard errors were obtained and meta-analyzed using a fixed-effects inverse variance approach. All lead 267 variants identified through these analyses were further investigated for gene-smoking interactions in 268 CHD. One lead variant per locus was selected for primary gene-smoking interaction analyses. 269

270 <u>Investigation of the APOE locus</u>: Although APOE has been recently established as a GWAS locus,⁷ 271 previous studies prior to GWAS have suggested that CHD risk is higher among carriers of the ϵ 4 272 allele at the APOE locus in smokers than in non-smokers.¹²⁻¹⁴ Because the ϵ 2, ϵ 3 and ϵ 4 alleles at 273 the APOE locus are not captured by the GWAS platform, we specifically conducted genotyping for 274 rs429358 and rs7412 variants to capture the three epsilon (ϵ) alleles in 13,822 participants (including 275 7,286 first-onset myocardial infarction cases) in the PROMIS study.¹⁵

276 <u>Gene-smoking interaction analyses at CHD and smoking loci</u>: To assess gene-smoking interactions, 277 analyses were conducted within each study, adjusted for age, sex and other study specific 278 covariates (e.g., principal components), and variants were analyzed in association with CHD 279 separately in "ever-smokers" and "never-smokers". Results from the two groups were then used to 280 test for interaction within each study. For the 50 variants, an interaction test statistic was calculated 281 within each study using the following equation as adapted from *Teslovich TM* et.al.¹⁶

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$$\frac{(\beta n - \beta e)}{\sqrt{SEn^2 + SEe^2}}$$

where β_n and β_e are the beta coefficients for the SNP in never-smokers and ever-smokers respectively, *SEn* and *SEe* are the standard errors for the log-ORs estimated for never-smokers and ever-smokers, respectively. Study specific interaction beta(s) and se(s) were calculated within each study and were pooled across studies using a fixed-effects meta-analysis. Interaction analyses were declared to be significant at a P-value of <1.0x10⁻³ (Bonferroni correction for 50 tests).

Conditional analyses on chr.15q25.1: At chr.15q25.1, we observed two variants exhibiting gene-288 smoking interactions for CHD. The proximity of these two signals raised the possibility that the 289 observed interactions may represent a single interaction locus across the entire region. To 290 investigate this possibility we undertook conditional analyses using an approximate conditional and 291 joint analyses approach, also known as GCTA (Genome-wide Complex Trait Analysis), as described 292 previously.¹⁷⁻²² Briefly, this method leverages summary-level statistics from a meta-analysis and uses 293 LD corrections between SNPs estimated from a reference sample. Such an approach has been 294 shown to yield similar results to that obtained from conditional analyses conducted on individual 295 participant data and has been successfully implemented in several other studies that have fine-296 mapped loci for other complex traits.¹⁷⁻²² Using this approach, we first conducted separate 297 conditional analyses at the chr.15q25.1 locus to identify main-effect variant(s) independently 298 299 associated with CHD and smoking behavior, respectively. We used the meta-analyzed data for CHD main effects in the CARDIoGRAMplus4D consortium to identify SNPs independently associated with 300 CHD risk and we used the genetic meta-analysis data from the Tobacco and Genetics Consortium 301 302 (TGC) in 140,000 participants to identify variants independently associated with smoking behavior. We then estimated the effects of these independent variants on CHD risk stratified by smoking 303 status and mutually adjusted the effects of these variants for each other. 304

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306 Analysis of eQTLs and regulatory features at the chr15q25.1 gene-smoking interaction locus

307 <u>eQTL analyses:</u> We mined publicly available databases to identify genotype-related expression 308 differences (eQTLs) in *ADAMTS7* and the *CHRNB4-A3-A5* gene cluster in order to understand the 309 directionality of the association of expression of these genes with CHD and smoking behavior. 310 Specifically, we investigated data available from the GTEx consortium,²³ the HapMap consortium 311 (restricted to European populations), and the Multiple Tissue Human Expression Resource 312 (MuTHER).²⁴ We also analyzed expression data in 147 donor HAoEC lines.²⁵ We used a nominal P-313 value of 0.002 to account for multiple testing involved in the eQTL analyses. 314 <u>Regulatory features of the chr. 15q25.1 region:</u> Data from ENCODE²⁶ were explored as described in 315 eMethods. ChIP-seq experiments were performed on confluent HCASMC (Cell Applications 350-05a 316 & Lonza CC-2583; cultured in SmGM-2 BulletKit media; Lonza) as described.²⁷ TCF21 (Abcam 317 ab49475), Jun (Santa Cruz Biotechnology sc-1694), JunD (Santa Cruz Biotechnology sc-74), and 318 CEBP (Santa Cruz Biotechnology sc-150) transcription factor binding was interrogated and H3K27ac 319 data were acquired using the same ChIP protocol with an anti-H3K27ac antibody (Abcam; ab4729). 320 Reads were aligned to the human genome (GRCh37p13) using STAR.²⁸

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322 Analyses of ADAMTS7 and CHRNB4-A3-A5 gene expression in vascular cells and tissues

<u>ADAMTS7 and CHRNB4-A3-A5 gene expression in vascular cells:</u> ADAMTS7 and CHRNB4-A3-A5 mRNA levels were measured in cultured human coronary artery smooth muscle cells (HCASMC; Lonza CC-2583, Lonza Walkersville, MD), human coronary artery endothelial cells (HCAEC, Lonza CC-2585), human aortic smooth muscle cells (HAoSMC, Lonza CC-2571), human aortic endothelial cells (HAoEC, Lonza CC-2535), human aortic adventitial fibroblasts (HAoAF, Lonza CC-7014), and human acute monocytic leukemia cell line (THP-1, ATCC TIB-202). Further details are provided in eMethods.

ADAMTS7 and CHRNB4-A3-A5 gene expression in response to cigarette smoke extract: HCASMC 330 331 were grown to confluence and cigarette smoke extract experiments performed at passage-7. Cigarette smoke extract was custom-prepared by Arista Laboratories (Richmond, VA). Briefly, the 332 condensate was generated by smoking Marlboro Red King Size Hard Pack cigarettes on an 333 analytical smoke machine under International Organization for Standardization smoking conditions. 334 The smoke condensate was collected on 92 mm filter pads and extracted from each pad in DMSO 335 by shaking to obtain a solution of ~20 mg/mL final concentration of the total particulate matter. 336 Serum starved (24 hrs) HCASMC were treated with 0.5% or 1.0% cigarette smoke extract (v/v) for 4, 337 338 12, and 24 hrs in serum reduced conditions (0.5% FBS in DMEM). Details on RNA preparation and 339 g-PCR are provided in **Supplementary Methods**.

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344 **RESULTS**

345 Description of the participating studies

Of the 37 studies participating in the CARDIoGRAMplusC4D consortium, information on "ever-smoking" was available in 30 studies, yielding a total sample size of 60,919 CHD cases and 80,243 controls. All studies recruited participants of European ancestry, except PROMIS (South Asian),¹⁵ LOLIPOP (South Asian)²⁹ and FGENTCARD (Lebanese).³⁰ Number of CHD cases and controls and percentages that were "ever-smokers" are provided in **Supplementary Table 1**. As expected, in all the participating studies, association of "ever-smoking" status with CHD risk was directionally consistent with an increased risk of CHD (**Supplementary Figure 2**).

353 New variants associated with CHD at established loci

Supplementary Figure 3 and Supplementary Table 2 present effect estimates for the 354 355 association with CHD for (i) the most significant variant that we identified at known CHD loci in the current CARDIoGRAMplusC4D consortium analysis as well as for (ii) the top SNP previously 356 reported at each of these established CHD loci. Of the 45 established CHD loci, we identified 32 for 357 which we discovered a more statistically significant SNP in association with CHD risk in our dataset 358 than the prior reported top variant. All of these 32 SNPs were in moderate to high LD ($r^2 > 0.6$) with 359 the previously published variants.⁷⁻⁹ In our primary gene-smoking interaction analyses, at each of the 360 CHD loci, we, therefore, used the SNP with the most significant CHD association (Supplementary 361 Figure 3 and Supplementary Table 2). Because the smoking behavior phenotype (captured as 362 cigarettes per day [CPD]) was not available in all CARDIoGRAMplusC4D studies, we used the top 363 variant previously reported for CPD¹⁰ at each locus (**Supplementary Figure 4**). 364

365 Analyses of the APOE locus.

The effect of rs6857, the lead CHD variant at the APOE locus, was similar in "ever-366 367 smokers" compared to "never-smokers" (Supplementary Table 3). Specifically, the CHD OR for the T allele at rs6857 was found to be 1.10 (P-value 7.93x10⁻⁴) in "never-smokers" (12,159 CHD cases 368 369 and 22,932 controls) which was quantitatively similar to the CHD OR of 1.09 (P-value: 8.68x10⁻⁵) observed in "ever-smokers" (23,753 CHD cases and 24,019 controls) (interaction P-value: 0.76) 370 (Supplementary Figure 5a). Investigation in the PROMIS study of the APOE epsilon genotypes 371 yielded consistent findings; the OR for CHD among £4 carriers in "never-smokers" was 1.13 372 compared to the CHD OR of 1.07 observed in "ever-smokers" (interaction P-value: 0.82) 373 (Supplementary Figure 5a). 374

375 Novel gene-smoking interaction effects on CHD at chromosome 15q25.1

Of the 50 loci, we identified effect-modification by "ever-smoking" status on CHD risk for the lead variants at two distinct loci, rs7178051, in proximity of *ADAMTS7* (an established CHD locus), and rs1051730, in proximity of *CHRNB4-A3-A5* (an established smoking behavior locus) (**Supplementary Table 3**). Although associated with different traits and located in distinct LD blocks, these two variants reside ~224 KBs apart on chr.15q25.1 and are in weak linkage disequilibrium (LD) ($r^2 = 0.22$), raising the question of whether these two chr.15q25.1 gene-smoking interactions on CHD are independent of each other.

383 At the ADAMTS7 CHD locus, the T allele at the rs7178051 variant was found to be more strongly and inversely associated with CHD risk in never-smokers (OR: 0.88; P-value: 7.02x10⁻¹⁶) 384 compared to a much weaker effect in ever-smokers (OR: 0.95; P-value: 8.64x10⁻⁴) (P-value of 385 interaction: 8.57x10⁻⁵) (Table 1). Thus, the protective impact of the rs7178051 T allele observed in 386 never-smokers was halved in people who smoked (Figure-1). This difference is not related to power 387 differences within strata because for this variant, there were less data available in the never-smoking 388 group (21,232 CHD cases and 38,713 controls) compared to the ever-smoking group (39,585 CHD 389 cases and 40,749 controls). There was no substantial evidence of heterogeneity for the interaction 390 beta across the participating studies (Heterogeneity chi-squared = 36.23 (d.f. = 25); P-value for the 391 χ^2 test of heterogeneity = 0.06; I^2 = 31.0%; tau-squared (τ^2 = 0). We further conducted sensitivity 392 analyses using a random effect model; the results remained unchanged and the interaction beta 393 394 remained significant (Supplementary Figure 5b). Although the frequency of rs7178051 was 39% in Europeans compared to 28% in South Asians, further analyses stratified by ancestry (i.e., European 395 versus non-Europeans) showed similar results (Supplementary Figure 5c). Other variants 396 discovered through prior CHD GWAS at the ADAMTS7 locus (e.g., rs7173743, rs4380028, 397 rs3825807) were in moderate to high LD ($r^2 > 0.50$) with rs7178051 and were also found to display a 398 similar pattern of gene-smoking interaction effects (Table 1). 399

At the CHRNB4-A3-A5 smoking locus, the A allele at the rs1051730 variant had an inverse 400 401 trend (not significant after adjustment) of association with CHD in never-smokers (OR: 0.96; P-value: 1.56x10⁻²) and a positive trend (not significant after adjustment) in ever-smokers (OR: 1.03; P-value: 402 1.53x10⁻²) (P-value of interaction: 2.37x10⁻⁴) (Table 1 and Supplementary Table 3). For this 403 variant, data on 20,559 CHD cases and 38,198 controls were available in the never-smoking group 404 whereas 38,923 CHD cases and 40,371 controls were available in the ever-smoking group. Similar 405 gene-smoking interaction patterns were observed for other variants (e.g., rs2036527, rs8034191) 406 that have been previously reported for CPD behavior at the CHRNB4-A3-A5 gene cluster (Table 1). 407

408 Further interrogation of the chr15g21.1 region encompassing rs7178051 and rs1051730 across three distinct LD blocks (Figure 1) revealed multiple additional variants for which we 409 observed gene-smoking interactions in CHD (Table 1 and Figure 1). Indeed, several SNPs (e.g., 410 rs7178051, rs10083696, rs7176187, rs6495335, rs4887077) had genome-wide significant 411 associations with CHD in "never-smokers" but had weaker and less significant associations with 412 CHD in "ever-smokers" (Figure 1). Alleles clustered specifically around ADAMTS7 rather than at the 413 CHRNB4-A3-A5 genes appear to be protective of CHD in "never-smokers" but have attenuated 414 protective effects in "ever-smokers" (Figure 2). 415

416 Conditional analyses

To investigate the possibility that the two chr.15q25.1 gene-smoking interactions might represent a single interaction locus across the entire region we undertook an approximate conditional and joint analyses¹⁷⁻²² using summary data derived from CARDIoGRAMplus4D for CHD and from the TGC for smoking behavior. In-addition to rs7178051, we identified one other variant, rs11072794 in low LD with rs7178051 (r²=0.20) that was associated independently with CHD (Figure 3a; red triangles) (Figure 3b & Supplementary Figure 6b; red triangles). We also confirmed two variants (rs1051730 and rs684513) located in two different LD blocks that were independently associated with smoking behavior in the TGC data¹⁰ (Figure 3d & Supplementary Figure 6b; grey circles).

426 In analyses of the CHD variants, both rs7178051 and rs11072794 remained strongly associated with CHD after adjusting for the top CPD variants (rs1051730 and rs684513) (Figure 3d, 427 428 red triangles) whereas their weak association with CPD was lost after adjusting for the top CPD variants (Figure 3d; grey circles); e.g., the P-value for rs7178051 association with CPD was 1x10⁻⁵ 429 in unadjusted analyses but attenuated to 0.55 after adjusting for rs1051730 and rs684513. In 430 analyses of the CPD variants, both rs1051730 and rs684513 remained strongly associated with CPD 431 after adjusting for the top CHD variants (rs7178051 and rs11072794) (Figure 3b, grey circles) 432 whereas their weak association with CHD was lost after adjusting for the top CHD variants (Figure 433 **3b**, red triangles). As expected, conditional analyses that included all four of these variants resulted 434 in a null association of the region with both CHD and CPD (Supplementary Figure 6b). To 435 underscore the validity of the conditional approach using summary data, we used individual 436 participant data from an expanded PROMIS sample involving 9,025 MI cases and 8,506 controls. 437 438 We found that the OR conferred by allelic variation at rs7178051 remained associated with MI risk 439 independent of the two CPD variants (rs1051730 and rs684513) and rs11072794 (the second CHD

440 SNP) (**Supplementary Figure 6c**). Conversely, the apparent effect of allelic variation at rs1051730 441 (the top CPD variant) on CHD risk was lost when we adjusted for the other three variants, 442 rs7178051, rs11072794 and rs684513 (**Supplementary Figure 6c**).

443 Next, using summary level data we examined the association of each of these four variants with CHD risk separately in "ever-smokers" and "never-smokers" while mutually adjusting for the 444 other three variants (Figure 4 & Supplementary Figure 7). In these analyses, only allelic variation 445 at rs7178051 was found to have independent genome-wide significant effects on CHD in never-446 smokers. rs7178051 was also the only one of these four variants with significant differences in the 447 effect estimate for gene-CHD associations between the two smoking groups (P-value for the χ^2 test 448 of heterogeneity: 5.4x10⁻⁵) after adjusting for the effects of other variants (rs11072794, rs1051730 449 and rs684513). These conditional analyses suggest that (a) variants located near the ADAMTS7 450 gene but not CHRNB4-A3-A5 genes have independent effects on CHD, (b) a single independent 451 gene-smoking interaction signal for CHD exists on chr.15g.25.1 which is localized at the ADAMTS7 452 CHD locus (marked by rs7178051) and (c) an apparent interaction signal observed at the nearby 453 CHRNB4-A3-A5 CPD locus (marked by rs1051730) is not independent of the ADAMTS7 454 (rs7178051) interaction signal. 455

To assess the robustness of conditional analyses methodology that uses summary level data 456 (i.e., GCTA)¹⁷⁻²², we conducted sensitivity analyses in the PROMIS dataset (9,025 MI cases and 457 8,506 controls). We assessed the association of rs7178051 (top CHD SNP) and rs1051730 (top 458 CPD SNP) after mutually adjusting for each other by conducting (i) standard logistic regression using 459 individual participant data and (ii) summary level data in PROMIS using the GCTA method 460 (Supplementary Table 4). The top CHD SNP was found associated with CHD risk in PROMIS 461 independent of the top CPD variant using both the methods, in-contrast the effect on CHD of the top 462 CPD SNP attenuated sharply when adjusted for the top CHD SNP - the effect estimates obtained 463 464 using the two methods were very similar (Supplementary Table 4).

Finally, to further demonstrate that the gene-smoking interaction effect in CHD at rs7178051 is independent of the *CHRNB4-A3-A5* CPD locus, we conducted sensitivity analyses in the PROMIS study by restricting our gene-environment interaction analysis to subjects who do not carry the minor alleles of rs1051730 and rs684513 (the two SNPs associated with CPD) at the *CHRNB4-A3-A5* locus. The T allele at the rs7178051 variant was associated with CHD only in never-smokers (OR: 0.88; P-value: 0.01) compared to a weaker and non-significant association in ever-smokers (OR: 0.94; P-value: 0.21) (**Supplementary Table 5**). The effect estimates obtained in each of the 472 categories defined by smoking status in PROMIS were similar to the effect estimates obtained in our
473 overall meta-analyses that utilized data in all participants (Supplementary Table 5).

474 Analysis of eQTLs and regulatory features at the chr15q25.1 gene-smoking interaction locus.

We mined publicly available eQTL data from the HapMap consortium.¹¹ GTEx consortium²³ 475 476 and the MuTHER consortium²⁴ as well as data from 147 HAoEC lines²⁵ to examine the association between mRNA expression of ADAMTS7 and CHRN genes with CHD, CPD and gene-smoking 477 interaction SNPs at the chr15q25.1 locus. SNP-mRNA associations with p-values <0.002 (correction 478 for 20 tests) are presented (Figure 5). The top two CHD variants (rs7178051, rs11072794) are 479 associated with reduced ADAMTS7 expression (e.g., rs11072794 p=6.01x10⁻²¹ in MuTHER LCL, 480 n=850; and rs7178051 p=0.0029 in HAoEC, n=147) but have no association with expression of 481 CHRN genes in any cell or tissue examined. In contrast, the top two CPD variants (rs1051730 and 482 rs684513) were associated with CHRN gene expression (e.g., rs1051730 p=6.9x10⁻⁷ for CHRNA5 in 483 GTEx skeletal muscle and nerve tissue) but have no association with ADAMTS7 in these cells or 484 tissues. These findings complement conditional analyses suggesting that gene-CHD and gene-485 smoking interaction effects on CHD are likely mediated by ADAMTS7 whereas the smoking behavior 486 effect appears to be mediated through the CHRNA3-5 gene cluster. 487

In analysis of data from the ENCODE project²⁶ and for human aortic tissue in NIH 488 Roadmap Epigenomics project, ADAMTS7 was associated with RNAseq reads and an active 489 transcription mark, H3K36me3, whereas CHRN genes had low/absent RNAseq reads and were 490 positive for repressive marks, H3K27me3 and H3K9me3 (Supplementary Figure 8). In HCASMC 491 492 ChIPseq data, rs7178051 the top CHD and gene-smoking CHD interacting SNP, is located in a region with active regulatory marks H3K4me1 and H3K4me3 as well as transcription factor binding 493 site for TCF21, an important HCASMC transcription factor also associated with CAD. This ChIPseq 494 pattern was observed also in human aortic tissue (Figure 6). These regulatory data suggest active 495 transcription of ADAMTS7, but not CHRN genes, in vascular cells and aortic tissue and reveal that 496 rs7178051, the lead gene-smoking CHD interaction SNP, overlaps active transcription marks and 497 transcription factor binding regions in HCASMC. 498

499 <u>ADAMTS7 and CHRNB4-A3-A5 expression in vascular cells and their response to cigarette smoke</u> 500 <u>extract</u>

501 In order to explore which genes at the chr15q25.1 locus are expressed in CHD-relevant 502 vascular cells, we performed q-PCR of *ADAMTS7* and the *CHRNB4-A3-A5* genes in primary human vascular cells and in the THP1 human monocyte cell line **(Supplementary Figure 9 & Figure 5)**. Whilst *ADAMTS7* mRNA was expressed abundantly in all vascular cell types, mRNA was below detection or expressed at a very low level for each of the genes in the *CHRNB4-A3-A5* cluster in any of these cell types **(Supplementary Figure 9)**. Next, we explored the effect of cigarette smoke extract on gene expression in HCASMC, a cell type of particular relevance to vascular responses to cigarette smoke products^{31, 32} as well as to *ADAMTS7* vascular functions in atherosclerosis and CHD.³³ In primary HCASMC, cigarette smoke extract exposure increased *ADAMTS7* mRNA levels by over 2-fold **(Figure 5)** but did not affect expression of the *CHRN* genes (not shown). Thus, in contrast to *CHRN* genes, *ADAMTS7* is both expressed and modulated by cigarette smoke extract in 512 CHD-relevant vascular cells providing biological support for *ADAMTS7*, but not CHRN genes, in the 513 gene-smoking interaction at chr15q25.1.

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529 DISCUSSION

530 We conducted a gene-environment interaction study at fifty loci associated with either CHD or smoking behavior and found evidence of effect-modification of genotype-related CHD risk by 531 smoking-behavior at the chr.15q21.1 CHD locus. Specifically, we observed highly significant 532 attenuation of the cardio-protective effects associated with alleles at this locus in people who 533 smoked cigarettes. Conditional analyses identified an LD block located at the ADAMTS7 gene that 534 accounted for both the main effect on CHD as well as the gene-smoking interactions in CHD. Data 535 from expression and cell studies support our genetic analysis, suggesting that the underlying 536 537 mechanism relates to genotype differences in the effect of smoking on expression of ADAMTS7 in vascular tissue. 538

539 Our findings have novel mechanistic and clinical implications. These human genomic data provide new insights into the mechanism of CHD in cigarette smokers. Identification of gene-540 smoking interaction at the chr15q21.1 locus suggests a specific role in smoking-related CHD for 541 ADAMTS7 and its substrates, vascular matrix and vascular smooth muscle cell biology more 542 broadly. Such insights can help to prioritize translational strategies for smoking-related CHD and 543 present opportunities to study lifestyle interventions and pharmacological strategies to lower CHD in 544 individuals who smoke cigarettes. Thus, inhibition of ADAMTS7 represents a novel potential 545 therapeutic strategy for CHD that may have particular benefits in individuals who smoke cigarettes. 546 All smokers should receive counseling for smoking cessation yet such broad public health strategies 547 have failed to reach or impact smoking behavior in a large portion of nicotine-addicted individuals. 548 Our data provides a human genomic context for consideration of targeting specific genetically at-risk 549 individuals via intensified preventive strategies and development of novel pharmacological 550 treatments. 551

Our study also represents a realistic strategy for performing gene-environment interaction 552 studies using contemporary genetic data. We illustrate that identifying joint effects of genetic and 553 lifestyle factors in CHD requires very large sample sizes, yet such analyses are biologically 554 informative when studies are adequately powered. In this context, an important observation in our 555 large sample is the lack of effect modification by smoking behavior on CHD at the APOE locus, a 556 previously reported smoking interaction locus.¹²⁻¹⁴ This finding is consistent with a recent meta-557 analysis that found no evidence of effect modification by smoking for APOE genotypes and CHD 558 risk.³⁴ These studies raise concerns that most prior gene-environment interactions studies in CHD 559 560 have been prone to the same biases (i.e., limited statistical power and false positive associations) as

561 candidate gene studies investigating main effects in the pre-GWAS era. The present study differs 562 from previous studies by being much larger and, importantly, it includes genomic and functional 563 follow-up data supporting the plausibility of the observed gene-environment interaction.

564 ADAMTS7 (or the A disintegrin and metalloproteinase with thrombospondin motifs-7) is a member of the ADAMTS family of secreted zinc metalloproteases.^{35, 36} We previously discovered 565 and replicated genetic variation at the ADAMTS7 locus in association with coronary atherosclerosis 566 and MI.⁷⁻⁹ Both in vivo and in vitro studies suggest that ADAMTS7 modulates VSMC phenotype 567 switching and migration and that this may be mediated via cartilage oligomeric matrix protein 568 (COMP) or thrombospondin-1 (TSP-1),^{32,33} i.e. putative ADAMTS7 substrates expressed in vascular 569 tissue. Genetic variation at ADAMTS7, however, has no relationship with traditional risk factors or 570 mechanistic biomarkers; hence the directional impact of ADAMTS7 expression on CHD risk and the 571 underlying biological mechanisms have been unclear.³² 572

Our gene-smoking interaction analyses provide novel insights into the directional impact of 573 the ADAMTS7 locus on CHD, the underlying mechanisms of CHD in smokers, and how such 574 findings ultimately might translate towards achieving health benefits in society. Our human eQTL 575 interrogations reveal that common alleles that relate to lower CHD risk at the ADAMTS7 locus are 576 also associated with reduced ADAMTS7 expression, implying an atherogenic role of the gene. This 577 is supported by our recent in vivo experimental studies; Adamts7 deficiency protected against diet-578 induced atherosclerosis in both the Ldlr^{-/-} and ApoE^{-/-} mouse models, reduced neointima formation 579 following arterial injury, and decreased VSMC migration *in vitro*.³³ In our smoking-stratified analyses, 580 we observed CHD protective effect which was attenuated in smokers. Thus, smoking exposure may 581 overcome the genetic effect of protective alleles that act by reducing ADAMTS7 expression. Such a 582 possibility is supported by our HCASMC data that reveals increased ADAMTS7 expression in 583 HCASMC exposed to cigarette smoke extract. These human genome-smoking studies are the first to 584 585 implicate a specific locus as causal in mediating increased risk of CHD in smokers. Additional 586 translational studies are needed to establish the precise mechanisms of atheroprotection for alleles 587 at the ADAMTS7 locus, how cigarette smoking impacts these genetic effects, and whether deletion or inhibition of ADAMTS7 in vivo attenuates the specific acceleration of atherosclerosis conferred by 588 cigarette smoking. 589

590 Strengths and limitations of our study merit consideration. This is a large study that 591 conducted gene-smoking interaction analyses in CHD by using GWAS data. We observed 592 substantial heterogeneity across study samples in our initial quality control analyses of "ever-

smoking" status with CHD risk. This is similar, however, to the heterogeneity reported in a recent 593 meta-analysis that pooled risk ratios from all the past prospective studies with information on 594 association of "ever-smoking" with incident CHD events.⁵ We recognize that other smoking related 595 phenotypes are important e.g., "current smoking" may have a more pronounced role than "ever-596 smoking" in plague rupture and thrombosis in patients with MI. We were however unable to 597 distinguish between "former" versus "current" smokers within "Ever Smokers" in our current 598 analyses; furthermore we were not able to analyze graded exposure to cigarette smoking such as 599 "pack-years". Given the use of multiple studies and meta-analyses of data, we used only one 600 analytical approach to investigate gene-smoking interactions. This approach, however, was feasible 601 and powerful in this large-scale consortium setting. While we used a fixed effects approach in our 602 meta-analyses, a random effects meta-analysis yielded qualitatively similar results (data not shown). 603 The lack or replication is partially offset by a large sample size, consistency across study cohorts 604 605 and racial groups and supplemental genomic and experimental evidence supporting biological plausibility. This approach is also consistent with recent recommendations³⁷ which favor use of a 606 607 powerful discovery experiment using all data rather than reducing power by splitting available dataset for discovery and validation. While our in vitro studies support a role for ADAMTS7 in the 608 gene-smoking interaction, it will be important to confirm that Adamts7 deficiency protect against 609 cigarette-smoke acceleration of atherosclerosis in rodent models. 610

Our interaction analyses, conditional analyses, eQTL interrogations and cell studies 611 612 suggest that ADAMTS7, but not the CHRNB4-A3-A5 gene cluster, is likely causal at 15q21.1 for gene-smoking interaction effects in CHD. Yet, analyses are not definitive. Although top interacting 613 SNPs and CHD SNPs (e.g., rs7178051) were associated with ADAMTS7, but not CHRNB4-A3-A5, 614 expression in LCLs, large-scale eQTL data and allele specific expression data (e.g., via RNA 615 sequencing) are not available for vascular tissues limiting causal inference. In our small HCAEC 616 datasets, we did however find that alleles at rs7178051 associate with ADAMTS7 expression but not 617 with any CHRNB4-A3-A5 genes suggesting, at least in one vascular cell type, that the gene-smoking 618 619 interaction is mediated via ADAMTS7.

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625 Conclusions

We provide novel evidence for allelic variation exhibiting gene-smoking interaction in CHD at the chr.15g21.1 locus. The protective effect conferred by variation at this locus in never-smokers is markedly attenuated in people who are ever-smokers. Stepwise conditional analyses, gene expression data in vascular cells, eQTL interrogation, and cigarette smoke extract exposure in HCASMC suggest that ADAMTS7 accounts for both the gene-smoking interaction in CHD and the CHD main effect on chr.15g21.1. Our findings reveal interactions of genetic variants and key lifestyle determinants in the etiology of CHD, provide new insights into the potential mechanisms of CHD in cigarette smokers, and facilitate precision medicine advances in cigarette-smoking related CHD. Our work motivates future large-scale studies investigating joint effects of genes and environment in CHD using existing complex-disease consortia datasets and genome-wide discovery approaches. This will provide opportunities to detect additional and novel loci displaying gene-environment interactions revealing genetic contexts for targeting intensive lifestyle interventions and novel therapeutics.

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Variant	Association	allele	LD with rs7178051*	LD with rs1051730^	Never Smokers						Ever Smokers				
					N cases	N controls	N Total	Beta (SE)	P-value	N case	N s controls	N Total	Beta (SE)	P-value	P-value interaction
*rs7178051 ⁴	CHD (NPR)	T/C	-	0.22	21232	38713	59945	-0.13 (0.01)	1.30E-16	3958	5 40749	80334	-0.05 (.01)	2.49E-04	8.57E-05
†rs1051730 ¹⁶	SB (known)	A/G	0.22	-	20559	38198	58757	-0.04 (0.02)	0.02	3892	3 40371	79294	0.03 (0.01)	0.02	2.37E-04
Other variants	on chr.15q25.1 v	with signi	ficant gene-sm	oking interaction	ns on CHD				11					11	I
rs71737431	CHD (Known)	C/T	0.61	0.18	21050	37955	59005	-0.11 (0.01)	2.73E-13	3904	4 39559	78603	-0.04 (0.01)	8.60E-04	9.29E-05
rs10083696 ²	CHD (Novel)	A/G	1.0	0.22	19721	36206	55927	-0.11 (0.02)	1.60E-12	3880	7 40018	78825	-0.05 (0.01)	2.72E-04	5.15E-05
rs7176187 ³	CHD (Novel)	T/C	1.0	0.24	21232	38713	59945	-0.12 (0.01)	7.02E-16	3958	5 40749	80334	-0.04 (0.01)	8.64E-04	6.93E-05
rs6495335⁵	CHD (Novel)	G/T	1.0	0.22	20144	37217	57361	-0.13 (0.02)	2.39E-15	3644	8 38203	74651	-0.04 (0.01)	1.69E-03	9.51E-04
rs43800286	CHD (Known)	T/C	1	0.22	21232	38713	59945	-0.12 (0.01)	2.20E-15	3958	5 40749	80334	-0.04 (.01)	1.03E-03	5.44E-04
rs38258077	CHD (Known)	G/A	0.52	0.43	17137	28633	45771	-0.09 (0.02)	2.82E-08	3007	1 29014	59086	-0.03 (0.01)	0.04	2.6E-03
rs3813565 ⁸	CHD (NPR)	T/G	0.43	0.56	19466	35830	55296	-0.08 (0.02)	5.08E-07	3664	2 37759	74401	-0.01 (0.01)	0.42	3.05E-04
rs11638490 ⁹	CHD (NPR)	T/C	0.44	0.51	20465	37897	58362	-0.08 (0.01)	6.90E-08	3853	3 39690	78223	-0.01 (0.01)	0.28	2.25E-04
rs1107279111	CHD (NPR)	A/C	0.44	0.51	19289	35944	55233	-0.08 (0.02)	2.83E-07	3524	5 36635	71880	005 (0.01)	0.68	1.06E-04
rs92269212	CHD (NPR)	A/C	0.44	0.50	20559	38198	58757	-0.08 (0.01)	2.81E-07	3892	3 40371	79294	-0.01 (0.01)	0.29	2.75E-04
rs1163837213	CHD (NPR)	T/C	0.44	0.50	21232	38713	59945	-0.08 (0.01)	6.92E-08	3958	5 40749	80334	-0.01 (0.01)	0.23	3.16E-04
rs4887077 ¹⁴	CHD (NPR)	T/C	0.44	0.50	21232	38713	59945	-0.08 (0.01)	4.71E-08	3958	5 40749	80334	-0.02 (0.01)	0.20	3.92E-05
rs12899135 ¹⁵	CHD (NPR)	G/A	0.39	0.56	20377	37440	57817	-0.07 (0.02)	3.97E-06	3838	2 39181	77563	-0.01 (0.01)	0.58	4.54E-04
rs68451318	SB (Known)	C/G	0.01	0.10	12517	21054	33572	-0.01 (0.02)	0.67	2464	1 24487	49129	0.03 (0.02)	0.18	0.08
rs2036527 ¹⁹	SB (Known)	A/G	0.17	0.90	20559	38198	58757	-0.04 (0.02)	0.02	3892	3 40371	79294	0.03 (0.01)	0.02	2.14E-04
rs10519203 ²⁰	CHD (NPR)	G/A	0.19	0.93	21232	38713	59945	-0.04 (0.01)	5.93E-03	3958	5 40749	80334	0.03 (0.01)	0.03	1.27E-04
rs8034191 ²¹	SB (Known)	C/T	0.19	1.0	19251	32131	51382	-0.05 (0.02)	2.62E-03	3492	5 34047	68972	0.02 (0.01)	0.06	3.91E-05

1098 Table-1. Novel genotype-smoking interaction findings in coronary heart disease at the chromosome 15q25.1 locus

1100 1101 CHD = coronary heart disease; SB = smoking behavior; NPR: Not a previously reported variant with disease risk *lead variant in association with CHD in our dataset; † lead variant in association with SB

1-21 each number refers to the physical location of the variant in figure-

1111 Figure Legends

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Figure 1. (a) Regional association analyses at the chromosome 15q25.1 locus in association with CHD risk stratified by smoking status. Association P-values for genetic variants with CHD risk in "never-smokers" (green squares) and "ever-smokers" (red triangles). (b) Longitudinal bars represent gene-smoking CHD interaction P-values at the chromosome 15q25.1 locus; bars in blue are Pvalues for variants listed in Table-1 and each variant has been assigned a unique identification number based on its physical location; (c) LD-blocks at the 15q25.1 locus visualized through HAPLOVIEW using LD estimates in the HapMAP-2 CEU reference population.

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Figure 2. Several variants at chromosome 15q21.1 have stronger effects on CHD risk in "neversmokers" compared to "ever-smokers". Variants with the strongest interaction P-value are displayed.

Figure 3. Step-wise conditional analysis of genetic variation at the chromosome 15q21.1 locus with CHD (red triangles) and smoking behavior (cigarettes per day, CPD; grey circles). At the chromosome 15q21.1 locus, analyses adjusted for rs7178051 and rs11072794 completely attenuated the gene-CHD associations whereas gene-smoking remained unchanged. Analyses adjusted for rs1051730 and rs684513 completely attenuated the gene-smoking associations whereas gene-CHD effect remained unchanged.

Figure 4. Analyses mutually adjusted for rs7178051, rs11072794, rs1051730 and rs684513 at 15q21.1 on CHD and smoking behavior; gene-CHD interaction analyses were only found significant for rs7178051. Analyses on the left panel show associations of rs7178051, rs11072794, rs1051730 and rs684513 with CHD risk mutually adjusted for each other. Analyses on the right panel show associations of rs7178051, rs11072794, rs1051730 and rs684513 with smoking behavior mutually adjusted for each other.

Figure 5. (a) *ADAMTS7* and *CHRNB4-A3-A5* mRNA levels were measured in HCASMC. Cells were cultured to confluence, total RNA was extracted and cDNA generated. q-PCR was performed for *ACTB, GAPDH, TBP, ADAMTS7, CHRNB4, CHRNA3, CHRNA5* (95°C 15s, 60°C 1min). Delta Cts were calculated as follows: $(Ct_{ACTB} + Ct_{GAPDH} + Ct_{TBP})/3 - Ct_{TARGET GENE})$. Fold changes are derived from delta delta Cts based on formula FC = 2^{-dCt}. (b) Confluent HCASMC were exposed to cigarette smoke extract. Serum starved (x24 hrs.) confluent HCASMC were treated with 0.5% or 1.0% cigarette smoke extract (v/v) for 4, 12, and 24 hrs. in serum reduced conditions (0.5% FBS in DMEM). Total RNA was extracted, cDNA generated preparation and q-PCR performed for

ADAMTS7 by Taqman and normalized to *GAPDH*. The Average Ct for ADAMTS7 at baseline was 28.25. Results were presented as means ± SEM, and data were analyzed using Student's t-Test. (c) expression and eQTL Data from the GTEx consortium, the HapMap consortium (restricted to European populations), the Multiple Tissue Human Expression Resource (MuTHER) and in 147 donor HAoEC lines. Association of the independent lead variants identified in our conditional analyses with expression of *ADAMTS7* and genes in the *CHRNB4-A3-A5* cluster. A P-value threshold of 0.002 was set to account for multiple testing involved in the eQTL analyses.

Figure 6. Genome browser view of regulatory features at rs7178051 on Chr15q21.1. ChIP-seq experiments were performed on confluent HCASMC for TCF21, Jun, JunD, CEBP and H3K4me1, H3K27me3, H3K27ac. DNAasel hypersensitivity data for human AoSMC were acquired from the ENCODE project. Human aortic tissue H3K4me1, H3K9me3, H3K27me3, and H3K36me3 ChIP-seq data were acquired from the NIH Roadmap Epigenomics Project. HCASMC = human coronary artery smooth muscle cells; AoSMC = human aortic smooth muscle cells.

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