





Association of Chromosome 9p21 With Subsequent Coronary Heart Disease Events A GENIUS-CHD Study of Individual Participant Data

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ORIGINAL ARTICLE

Association of Chromosome 9p21 With Subsequent Coronary Heart Disease Events

A GENIUS-CHD Study of Individual Participant Data

BACKGROUND: Genetic variation at chromosome 9p21 is a recognized risk factor for coronary heart disease (CHD). However, its effect on disease progression and subsequent events is unclear, raising questions about its value for stratification of residual risk.

METHODS: A variant at chromosome 9p21 (rs1333049) was tested for association with subsequent events during follow-up in 103357 Europeans with established CHD at baseline from the GENIUS-CHD (Genetics of Subsequent Coronary Heart Disease) Consortium (73.1% male, mean age 62.9 years). The primary outcome, subsequent CHD death or myocardial infarction (CHD death/myocardial infarction), occurred in 13 040 of the 93 115 participants with available outcome data. Effect estimates were compared with case/control risk obtained from the CARDIoGRAMplusC4D consortium (Coronary Artery Disease Genome-wide Replication and Meta-analysis [CARDIoGRAM] plus The Coronary Artery Disease [C4D] Genetics) including 47 222 CHD cases and 122 264 controls free of CHD.

RESULTS: Meta-analyses revealed no significant association between chromosome 9p21 and the primary outcome of CHD death/myocardial infarction among those with established CHD at baseline (GENIUS-CHD odds ratio, 1.02; 95% CI, 0.99–1.05). This contrasted with a strong association in CARDIoGRAMPlusC4D odds ratio 1.20; 95% CI, 1.18–1.22; *P* for interaction <0.001 compared with the GENIUS-CHD estimate. Similarly, no clear associations were identified for additional subsequent outcomes, including all-cause death, although we found a modest positive association between chromosome 9p21 and subsequent revascularization (odds ratio, 1.07; 95% CI, 1.04–1.09).

CONCLUSIONS: In contrast to studies comparing individuals with CHD to disease-free controls, we found no clear association between genetic variation at chromosome 9p21 and risk of subsequent acute CHD events when all individuals had CHD at baseline. However, the association with subsequent revascularization may support the postulated mechanism of chromosome 9p21 for promoting atheroma development.

Riyaz S. Patel, MD* Amand F. Schmidt, PhD* Vinicius Tragante, PhD* et al

6

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The full author list is available on page 169.

Key Words: chromosome **=** genetic variation **=** myocardial infarction **=** risk factor **=** secondary prevention

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https://www.ahajournals.org/journal/ circgen Sing a case-control approach, a large number of common genetic variants have now been associated with coronary heart disease (CHD) through genome-wide association studies, in an effort largely led by the CARDIOGRAMPlusC4D consortium (Coronary Artery Disease Genome-wide Replication and Meta-analysis [CARDIOGRAM] plus The Coronary Artery Disease [C4D] Genetics).¹ Among these variants, the chromosome 9p21 locus was the first to be discovered and the variant with the largest individual effect and is the most widely replicated genetic risk factor for CHD.²⁻⁴ Multiple studies including case-control and prospective cohort studies in general populations have reliably confirmed its effect on risk of CHD among otherwise healthy individuals.⁵

However, it is uncertain whether variants at the 9p21 locus also affect risk of recurrent or subsequent events, including mortality in those with established CHD. Elucidation of this hypothesis would help to better understand its mechanism and estimate its incremental value for stratification of residual risk. Prior studies have shown conflicting results, although most have been underpowered. A literature-based meta-analysis indicated a null association of chromosome 9p21 variants with subsequent CHD events but was based on summary, not individual level data, with varying outcome definitions.^{6,7}

The new collaborative GENIUS-CHD (Genetics of Subsequent Coronary Heart Disease) consortium, described in this issue of the journal, was established to investigate genetic determinants of disease progression following an index CHD event.⁸

In this article, we use the GENIUS-CHD resource to: (1) examine the association of variants at the 9p21 locus on risk of subsequent CHD events in individuals with established CHD; (2) compare these to the association between chromosome 9p21 and any CHD observed in the CARDIo-GRAMPlusC4D consortium; and (3) explore the potential impact on these estimates of biases that might affect genetic association studies of disease outcome and prognosis.

METHODS

In accordance with Transparency and Openness Promotion Guidelines, the data, analytic methods, and study materials will be made available to other researchers for purposes of reproducing the results or replicating the procedure. Participating studies received local institutional review board approval and included patients who had provided informed consent at the time of enrollment. The central analysis sites also received waivers from their local institutional review board for collating and analysing summary level data from these individual studies. Details about the GENIUS-CHD consortium and study inclusion criteria have been published separately in this issue of the journal,⁸ whereas for this study full details about data sources, genetic variant selection, outcomes and statistical analyses are available in the Data Supplement.

RESULTS

In total, 49 studies from the GENIUS-CHD consortium contributed to the federated analysis resulting in a sample size of 103357 individuals of European descent with established CHD and available genotype data at the 9p21 locus. Of these, 93115 individuals had available data for the primary composite outcome of subsequent CHD death/myocardial infarction (MI), of whom 13040 experienced these events. Contributing study details are provided in Table. Participant characteristics are representative for populations with established CHD with a weighted mean age of 62.9 years; 73.1% male. As expected, risk factor prevalence was high in this population, including diabetes mellitus (24.4%), hypertension (59.1%), and current smoking (25.7%). Statin use at enrollment varied by study, ranging from 5.2% to 97.3%, with a median of 61.5% (Table).

The rs1333049 single nucleotide polymorphism was genotyped in 42 studies, with the remaining 7 studies using highly correlated proxies (R^2 >0.90); rs10757278 (4 studies) or rs4977574 (3 studies) when the primary single nucleotide polymorphism was unavailable. Genotyping details are provided in Table I in the Data Supplement. For rs1333049, the average risk allele frequency across the participating studies was 0.518 ranging from 0.453 to 0.587 (Figure I in the Data Supplement).

From CARDIOGRAMplusC4D, after excluding 6 cohorts which had contributed data to both consortia, data were available for association with chromosome 9p21 from 41 studies, including 47222 cases with CHD and 122264 controls free of any CHD.

Power to detect different effect sizes, including the effect size identified in CARDIoGRAMplusC4D, using a 2-sided alpha of 0.05, are provided in Table II in the Data Supplement.

Chromosome 9p21 Association With Subsequent CHD Events

Study-specific results for the association between chromosome 9p21 and risk of the primary outcome of CHD death or MI among individuals with established CHD at baseline, adjusted for age and sex are presented in Figure II in the Data Supplement.

The per-allele odds ratio (OR) for the primary outcome during follow-up was 1.02 (95% CI, 0.99–1.05). The effect estimate again for the primary outcome, based on a time to event analysis and using a Cox regression model, was also similar with a hazard ratio of 1.02 (95% CI, 0.99–1.04; Figure III in the Data Supplement).

In contrast, a meta-analysis of CARDIOGRAMIplus C4D data (excluding studies also contributing data to

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Table. Overview of	Overview of Studies Contributing to Chromosome 9p21 Analysis and Participant Characteristics	Chromosom	e 9p21 Ai	nalysis a	nd Part.	icipant Chara	cteristics									
Alias	Cohort	Total N genotyped	Study Design	CHD Type	Male, %	Age, y, SD	BMI (SD)	Diabetes mellitus, %	Smoking, %	Systolic BP, SD	Total Cholesterol, SD	Statin use, %	Creatinine, SD	Prior Revasc, %	Prior MI, %	PubMED ID
4C	Clinical Cohorts in Coronary disease Collaboration (4C)	1538	Cohort	CAD	62.1	62.2 (11.95)	30.2 (5.67)	23.4	:	133.9 (23.7)	4.69 (1.10)	26.4	99.3 (83.2)	22.6	15.5	:
AGNES	Arrhythmia Genetics in the Netherlands	1316	Cohort	ACS	79.3	57.7 (10.81)	26.5 (3.87)	7.6	59.3	÷	5.28 (1.04)	9.8	:	:	:	20622880
ANGES	Angiography and Genes Study	588	Cohort	Mixed	65.5	64.1 (9.55)	28.1 (4.36)	30.8	14.7	÷	4.84 (0.84)	69.4	83.0 (32.0)	42.4	24.7	21640993
ATVB	Italian Atherosclerosis, Thrombosis and Vascular Biology Group	1465	Cohort	ACS	90.4	40.0 (4.40)	26.8 (4.07)	8.4	78.7	132.3 (20.6)	5.76 (1.39)	56.2	:	÷	÷	21757122
CABGenomics	Coronary Artery Bypass Genomics	1542	Cohort	Mixed	80.1	64.7 (10.08)	29.7 (5.71)	10.1	11.2	:	4.21 (0.95)	75.2	:	:	42.8	25649697
CDCS	Coronary Disease Cohort Study	1800	Cohort	ACS	71.5	67.5 (11.96)	27.3 (4.66)	15.4	5.8	129.2 (21.6)	5.00 (1.09)	46.5	100.5 (40.0)	26.9	30.3	20400779
COROGENE	Corogene Study	1489	Cohort	ACS	70.9	64.7 (11.87)	27.6 (4.84)	18.2	34.4		4.63 (0.99)	5.2	84.0 (44.3)	:	:	21642350
CTMM	Circulating Cells	605	Cohort	Mixed	68.9	63.0 (9.83)	27.6 (4.45)	20.7	20.7	135.4 (19.1)	4.43 (1.05)		86.4 (34.9)	:	30.1	23975238
CURE	Cure-Genetics Study	4242	RCT	ACS	59.3	64.7 (10.99)	27.9 (4.44)	19.9	22.6	135.7 (21.9)	:	:	93.0 (33.9)	13.9	31.8	11102254
EGCUT	Estonian Biobank	2408	Cohort	CAD	51.0	67.1 (10.88)	28.9 (5.16)	18.7	19.2	135.6 (18.0)	5.64 (1.17)	27.3	:	15.7	36.0	24518929
EMORY	Emory Cardiovascular Biobank	2411	Cohort	Mixed	70.1	64.5 (11.06)	:	30.7	9.8	:	4.49 (1.02)	76.0	99.0 (45.1)	61.7	27.9	20729229
ERICO	Estratégia de Registro de Insuficiência Coronariana	438	Cohort	ACS	55.5	63.8 (13.36)	27.0 (5.06)	39.1	31.0	99.2 (38.4)	:	23.8	:	11.3	25.9	23644870
FINCAVAS	Finnish Cardiovascular Study	1671	Cohort	Mixed	69.4	60.9 (11.03)	27.8 (4.35)	18.4	24.3	140.2 (22.1)	4.74 (0.90)	57.3	90.8 (66.8)	32.6	39.0	16515696
FRISCII	FRISCII Study	3106	RCT	ACS	69.4	66.2 (9.80)	26.8 (3.87)	12.7	27.1	143.3 (22.4)	5.80 (1.12)	12.3	90.6 (18.8)	12.1	27.2	10475181
GENDEMIP	Genetic Determination of Myocardial Infarction in Prague	1267	Cohort	ACS	75.8	56.4 (8.63)	28.6 (4.68)	18.8	60.8	137.0 (20.8)	5.51 (1.17)	16.6	:	29.7	41.6	23249639
GENEBANK	Cleveland Clinic Genebank Study	2345	Cohort	Mixed	74.3	61.5 (11.06)	29.4 (5.44)	11.8	16.8	132.7 (21.1)	4.46 (0.93)	71.8	:	65.3	56.1	21475195
GENESIS-PRAXY	Gender and Sex Determinants of Cardiovascular Disease: From Bench to Beyond- Premature Acute Coronary Syndrome	784	Cohort	ACS	69.2	48.3 (5.62)	:	13.8	44.2	139.5 (26.5)	4.85 (1.18)	93.1	75.9 (19.7)	11 .3	11.4	22607849

(Continued)

Prior Prior Revasc, MI, PubMED % ID	13.7 63.2 22717531	0.2 1.3	31.4 48.9	80.1 20231156	0.4 32.0 20231156	41.3 16287954	6.6 20691829	47.6 21372283	50.1 39.7 28444280	2.8 21.6 24829374	0.1 22216169	48.7 57.4 11258203	1.3 10.6 26086777	41.5 40.8 24262617	28.2 21.9	15.1 20.6 19332184	17.2 12771003
Creatinine, Rev SD	94.8 (27.2) 1:	108.0 (64.6) C	101.6 (34.5) 3	102.8 (61.9)	105.0 (40.0) 20.	100.1 (16.7) 4	99.6 (66.6)		89.9 (37.5) 50	82.4 (26.5) 42.	87.3 (34.6)	88.7 (38.5) 44	91.6 (36.3)	101.0 (68.1) 4	89.8 (21.1) 28	85.6 (26.3) 1!	87.8 (27.8)
Statin use, %	72.1	50.8	66.8	79.3	16.8	76.9	38.7	52.8	88.3	77.4	38.9	58.9	80.7	82.5	92.4	79.7	46.0
Total Cholesterol, SD	4.70 (0.92)	4.62 (1.02)	4.37 (0.83)	5.33 (1.19)	5.19 (1.29)	5.03 (0.98)	4.96 (1.12)	:	5.02 (1.06)	4.46 (0.84)	5.24 (1.18)	4.94 (0.99)	5.04 (1.05)	4.57 (1.02)	5.53 (1.03)	5.42 (1.23)	5.98 (1.19)
Systolic BP, 5	129.5 (20.3)	126.7 (NA)	136.2 (19.7)	138.8 (25.3)	137.8 (27.1)	136.8 (19.8)	141.8 (24.4)	148.6 (18.1)	148.1 (23.8)	119.9 (15.5)	138.3 (21.8)	142.3 (24.1)	138.9 (27.4)	129.4 (14.2)	131.9 (19.0)	135.6 (21.8)	117.0 (15.6)
Smoking, %	64.4	:	14.9	49.9	69.5	20.3	10.2	12.8	27.5	32.4	29.0	23.9	48.1	26.8	19.5	35.2	28.2
Diabetes mellitus, %	13.3	77.9	77.8	81.3	15.0	11.4	20.3	23.9	36.9	18.5	34.4	44.3	22.2	43.0	6.9	22.8	12.0
BMI (SD)	:	29.7 (5.64)	30.3 (5.43)	27.0 (4.35)	28.0 (5.15)	27.4 (3.80)	29.5 (6.08)	:	26.3 (4.46)	27.0 (3.26)	29.0 (4.68)	27.5 (3.89)	24.8 (3.79)	28.0 (4.18)	28.6 (5.00)	28.2 (4.51)	26.5 (3.82)
Age, y, SD	65.2 (8.45)	71.1 (10.62)	69.1 (9.20)	65.7 (12.01)	64.4 (12.04)	61.2 (9.32)	61.2 (11.05)	68.7 (9.38)	68.3 (10.25)	58.6 (8.13)	64.0 (11.15)	63.8 (9.85)	62.4 (11.86)	67.6 (10.49)	65.3 (11.07)	62.6 (10.95)	62.7 (10.29)
Male, %	86.7	62.0	66.5	75.4	69.1	81.8	66.7	56.6	71.6	84.6	75.5	76.5	75.0	74.4	73.0	69.5	78.3
CHD Type	Mixed	CAD	CAD	ACS	ACS	ACS	Mixed	CAD	Mixed	Mixed	Mixed	Mixed	ACS	Mixed	Mixed	ACS	ACS
Study Design		Cohort	Cohort	Cohort	Cohort	RCT	Cohort	RCT	Cohort	Cohort	Cohort	Cohort	Cohort	Cohort	Cohort	RCT	Cohort
Total N genotyped		1003	2000	669	1086	6223	6763	2145	704	1147	4330	2175	603	1380	393	9814	783
Cohort	Genetic Mapping for Assessment of Cardiovascular Risk	Genetics of Diabetes Audit and Research in Tayside Scotland (I)	Genetics of Diabetes Audit and Research in Tayside Scotland (P)	Global Registry of Acute Coronary Events-Belgium	Global Registry of Acute Coronary Events - UK	Incremental Decrease in End Points Through Aggressive lipid Lowering	Intermountain Heart Collaborative Study	International Verapamil SR Trandolopril Study Genetic Substudy	Krakow-GENIUS-CHD	Karola Study	Leipzig (LIFE) Heart Study	The Ludwigshafen Risk and Cardiovascular Health Study	North East Poland Myocardial Infarction Study	Neapolis Campania Italia	Ottawa Heart Genomics Study	The Study of Platelet Inhibition and Patient Outcomes	Post Myocardial
Alias	GENOCOR	GoDARTS incident	GoDARTS prevalent	GRACE_B	GRACE_UK	IDEAL	INTERMOUNTAIN	INVEST	JUMC	KAROLA	LIFE-Heart	LURIC	NE_POLAND	NEAPOLIS	OHGS	PLATO	PMI

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Networkersby 987 RCI ACS 28.8 (0.4.06) 189 29.7 (0.6.1) 20.7 (0.7 (0.7) 20.7 (0.7 (0.7)		Cohort	Total N genotyped	Study Design		Male, %	Age, y, SD	BMI (SD)	Diabetes mellitus, %	Smoking, %	Systolic BP, SD		Statin use, %		Prior Revasc, %	Prior MI, %	PubMED ID
Programmeric function 439 RC1 Cup 651 5.4 (3.31) 2.6 (3.81) 103 103 (3.23.2) 256 Reunamericen in the Action optimericania 1022 Cohont MC2 759 19.11.10 27.3 (4.43) 19.8 30.4 466 1006 (23.6) 23.2 Reunamericania Tex Action optimericania 102 Cohont ACS 79.3 56.4 (3.0) 19.8 30.4 466 1006 (23.6) 23.2 Reunamericania Tex Action optimericania 243 19.8 30.4 47.3 (0.9) 75.7 92.3 (2.1) 47.3 97.3 24.4 97.4 109.4 (23.2) 25.2 Reunamericanic 2435 Cohont XC 29.3 (7.2) 26.4 (3.0) 19.4 137.4 (19.8) 473.0 (9.7) 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 27.2 <	POPular	The Popular study	266	RCT	ACS	74.3	63.8 (10.40)	:	18.9	27.5	145.0 (22.1)	4.25 (0.64)	80.7	92.7 (26.8)	33.1	43.7	26542508
Recurrance and hith mutuantion 1052 Cohort ACS 73 81.4 13.8 30.4 46.6 100.6 (28.6) 28.2 Intermutantion Profenses 1150 Cohort ACS 70.7 59.3 (7.1) 26.8 (4.02) 18.2 50.0 131.8 (20.6) 6.28 (1.16)	PROSPER	Prospective Study of Pravastatin in the Elderly at Risk	439	RCT	CAD	6.9	75.4 (3.31)	26.4 (3.87)	10.3	16.2	150.0 (21.6)	5.55 (0.83)	:	109.4 (23.2)	26.0	85.9	10569329
Stocholm Heart I150 Cohort ACS 70.7 59.3 (7.21) 26.8 (4.02) 18.2 500 131.8 (2.06) 6.28 (1.16) <th< td=""><td>RISCA</td><td>Recurrance and Inflammation in the Acute Coronary Syndromes Study</td><td>1052</td><td>Cohort</td><td>ACS</td><td>75.9</td><td>61.9 (11.40)</td><td>27.2 (4.43)</td><td>19.8</td><td>30.4</td><td>:</td><td>:</td><td>46.6</td><td>100.6 (28.6)</td><td>28.2</td><td>27.9</td><td>18549920</td></th<>	RISCA	Recurrance and Inflammation in the Acute Coronary Syndromes Study	1052	Cohort	ACS	75.9	61.9 (11.40)	27.2 (4.43)	19.8	30.4	:	:	46.6	100.6 (28.6)	28.2	27.9	18549920
Second Manifectations 2445 cond Manifectations 2446 1374 (19.0) 1374 (19.0) 233 (22.1) 333 (21.1) <	SHEEP	Stockholm Heart Epidemiology Program	1150	Cohort	ACS	70.7	59.3 (7.21)	26.8 (4.02)	18.2	50.0	131.8 (20.6)	6.28 (1.16)	:	:	:	:	17667644
Subilization of Attraction of Darapladib Therapy trail 9287 RCT Mixed 82.0 64.7 (9.10) 29.9 (4.97) 38.4 21.4 131.7 (16.1) 97.3 746 Darapladib Therapy trail Darapladib Therapy trail 2229 Cohort ACS 75.3 63.6 (10.62) 29.6 (5.9) 30.5 21.3 97.3 21.5 Taraniational Research trail 1974 Cohort ACS 73.3 63.6 (10.62) 29.6 (5.9) 30.5 21.3 21.4 39.7 117.8 (18.3) 21.5 Taraniational Research interction Patients 1974 Cohort ACS 72.4 59.8 (12.04) 29.5 (50.6) 28.6 37.4 117.8 (18.3) 21.5 77.1 27.0 Interction Patients 1974 Cohort ACS 72.4 59.8 (12.04) 29.5 (50.6) 28.7 70.3 117.8 (18.3) 21.5 70.3 117.5 70.5 77.0 77.0 77.0 77.0	SMART	Second Manifestations of Arterial Disease	2485	Cohort	Mixed	82.2	60.2 (9.26)	3 (3	16.6	24.4	137.4 (19.8)	4.73 (0.96)	75.7	92.3 (22.7)	:	43.6	10468526
Texpen 2729 Cohort ACS 75.3 63.6 (10.62) 29.6 (5.59) 30.5 21.3 57.1 21.5 Terating to New Targets 5104 RCT CAD 81.3 61.3 (8.73) 28.6 (4.59) 148 13.4 130.9 (16.8) 451 (0.61) 7.03 104.8 (17.3) Tarnalizional Research 1974 Cohort ACS 72.4 59.8 (12.04) 29.5 (5.96) 28.6 37.4 117.8 (18.3) 89.0 112.5 (76.7) 27.0 Investigating nocietying Disparities 1974 Cohort ACS 72.4 59.8 (12.04) 29.5 (5.96) 28.6 37.4 117.8 (18.3) 89.0 112.5 (76.7) 27.0 Investigating nocietying Disparities 1073 Cohort Mxed 75.6 57.4 (3.4) 21.5 23.1 117.8 (18.3) 87.0 112.5 (76.7) 27.0 InterAct Coronary 1073 Cohort Mxed 75.6 57.4 (3.4) 21.5<	STABILITY	Stabilization of Atherosclerotic Plaque by Initiation of Darapladib Therapy trial	9287	RCT	Mixed	82.0	64.7 (9.10)	29.9 (4.97)	38.4	21.4	131.7 (16.1)	:	97.3	:	74.6	58.6	24678955
Translational Research FIC CAD 81.3 61.3 (8.73) 28.6 (4.59) 14.8 13.4 130.9 (16.8) 4.51 (0.61) 70.3 104.8 (17.3) Translational Research 1974 Cohort ACS 72.4 59.8 (12.04) 295 (5.96) 28.6 37.4 117.8 (18.3) 89.0 112.5 (76.7) 27.0 Investigating Underlying Disparities 107.3 Cohort ACS 72.4 59.8 (12.04) 295 (5.96) 28.6 37.4 117.8 (18.3) 89.0 112.5 (76.7) 27.0 Investigating Underlying Disparities 107.3 Cohort ACS 72.4 59.8 (12.04) 29.5 (5.9) 28.6 37.4 117.8 (18.3) 89.0 112.5 (76.7) 27.0 In Acute Wooradial Indecktoonany 1073 Cohort Mixed 75.4 54.4 (10.26) 27.2 (4.34) 21.5 23.1 87.0 (1.10) 27.1 94.7 (24.8) Underbin Study 150.6 <		Texgen	2729	Cohort	ACS	75.3	63.6 (10.62)	29.6 (5.59)	30.5	21.3	:	:	57.1	:	21.5	16.7	21414601
Tanslational Research 1974 Cohort ACS 72.4 59.8 (12.04) 29.5 (5.96) 28.6 37.4 117.8 (18.3) 89.0 112.5 (76.7) 27.0 Investigating Inderlying Disparities Inderlying Disparities 89.0 112.5 (76.7) 27.0 27.0 Inderlying Disparities Inderlying Disparities 89.0 112.5 (76.7) 27.0 27.0 Inderlying Disparities Vaceth Connary 1073 Cohont Nixed 75.6 65.4 (10.26) 21.2 (4.34) 21.5 23.1 4.76 (1.18) 64.0 91.9 (42.9) Utrecht Connary 1073 Cohont Nixed 75.4 64.1 (9.6) 21.2 (4.34) 21.1 4.76 (1.18) 64.0 91.9 (42.9) Utrecht Connary 1073 Cohont Nixed 75.4 64.1 (9.6) 21.2 23.1 17.76 (1.18) 7.76 (1.8) 7.74 (1.8) Planmacogenetics 1500 Cohont Nixed 75.4 6		Treating to New Targets	5104	RCT	CAD	81.3	61.3 (8.73)	28.6 (4.59)	14.8	13.4	130.9 (16.8)	4.51 (0.61)	70.3	104.8 (17.3)	:	57.3	15755765
RBIO Utrecht Coronary 1073 Cohort Mixed 75.6 65.4 (10.26) 27.2 (4.34) 21.5 23.1 4.76 (1.18) 64.0 91.9 (42.9) Utrecht Cardiovascular 1500 Cohort Mixed 75.4 64.1 (9.96) 153.4 (21.4) 5.50 (1.10) 27.1 94.7 (24.8) Pharmacogenetics 1500 Cohort Mixed 75.4 64.1 (9.96) 153.4 (21.4) 5.50 (1.10) 27.1 94.7 (24.8) Verona Heart Study 907 Cohort AIX 80.9 61.3 (9.78) 26.9 (3.57) 18.5 69.2 5.37 (1.10) 47.1 96.7 (32.2) 17.4 Vorarlberg Institute for 1318 Cohort CAD 80.9 61.3 (9.78) 26.9 (3.77) 18.5 69.2 5.37 (1.10) 47.1 96.7 (32.2) 17.4 Vorarlberg Institute for 1318 Cohort CAD 73.1 64.5 (10.44) 27.4 (4.14)	HdWi	Translational Research Investigating Underlying Disparities in Acute Myocardial Infarction Patient's Health Status	1974	Cohort	ACS	72.4	59.8 (12.04)	29.5 (5.96)	28.6	37.4	117.8 (18.3)	:	0.08	112.5 (76.7)	27.0	18.3	21772003
Utrecht Cardiovascular 1500 Cohort Mixed 75.4 64.1 (9.96) 153.4 (21.4) 5.50 (1.10) 27.1 94.7 (24.8) Pharmacogenetics Study 907 Cohort AD 80.9 61.3 (9.78) 18.5 69.2 5.37 (1.10) 27.1 94.7 (24.8) Venona Heart Study 907 Cohort CAD 80.9 61.3 (9.78) 26.9 (3.57) 18.5 69.2 5.37 (1.10) 47.1 96.7 (32.2) 17.4 Voranberg Institute for 1318 Cohort CAD 73.1 64.5 (10.44) 27.4 (4.14) 30.6 18.9 137.2 (19.2) 49.7 88.4 (32.7) 21.1 Vascular Investigation and Treatment Study AD 73.1 64.5 (10.44) 27.4 (4.14) 30.6 18.9 137.2 (19.2) 545 (1.14) 96.7 (32.2) 21.1	JRBIO	Utrecht Coronary Biobank	1073	Cohort		75.6	65.4 (10.26)	27.2 (4.34)	21.5	23.1	:	4.76 (1.18)	64.0	91.9 (42.9)	:	28.9	:
Verona Heart Study 907 Cohort CAD 80.9 61.3 (9.78) 26.9 (3.57) 18.5 69.2 5.37 (1.10) 47.1 96.7 (32.2) 17.4 * Vorarlberg Institute for vascular Investigation 1318 Cohort CAD 73.1 64.5 (10.44) 27.4 (4.14) 30.6 18.9 137.2 (19.2) 5.45 (1.14) 49.7 88.4 (32.7) 21.1 * Vascular Investigation and Treatment Study 2 2 21.4 30.6 18.9 137.2 (19.2) 5.45 (1.14) 49.7 88.4 (32.7) 21.1 * Mascular Investigation and Treatment Study 2		Utrecht Cardiovascular Pharmacogenetics Study	1500	Cohort		75.4	64.1 (9.96)	:	:	÷	153.4 (21.4)	5.50 (1.10)	27.1	94.7 (24.8)	:	÷	25652526
Voralberg Institute for Vascular Investigation 1318 Cohort CAD 73.1 64.5 (10.44) 27.4 (4.14) 30.6 18.9 137.2 (19.2) 5.45 (1.14) 49.7 88.4 (32.7) 21.1 Vascular Investigation and Treatment Study and Treatment Study and Treatment Study as 2.71 21.1 30.6 18.9 137.2 (19.2) 5.45 (1.14) 49.7 88.4 (32.7) 21.1		Verona Heart Study	907	Cohort	CAD	80.9	61.3 (9.78)	26.9 (3.57)	18.5	69.2	:	5.37 (1.10)	47.1	96.7 (32.2)	17.4	59.6	10984565
Wareaw ACS Ganatic 660 Cohort ACS 74.5 63.6/11.72) 28.1.(1.72) 21.0 1.2.0 1.2.7.8/22.6/1.02) 03.5.(11.1) 03.5		Vorarlberg Institute for Vascular Investigation and Treatment Study	1318	Cohort	CAD	73.1	64.5 (10.44)	27.4 (4.14)	30.6	18.9	137.2 (19.2)	5.45 (1.14)	49.7	88.4 (32.7)	21.1	31.0	24265174
	WARSAW ACS	Warsaw ACS Genetic Registry	669	Cohort	ACS	74.5	63.6 (11.72)	28.1 (4.72)	21.9	42.0	127.8 (22.6)	4.99 (1.07)	:	93.5 (41.4)	:	18.6	:
WTCCC WTCCC CAD Study 1924 Cohort Mixed 79.3 60.0 (8.13) 27.6 (4.20) 11.7 12.8 143.6 (22.0) 5.28 (0.98) 71.6 67.1 7	CC	WTCCC CAD Study	1924	Cohort			60.0 (8.13)	27.6 (4.20)	11.7	12.8	143.6 (22.0)	5.28 (0.98)	71.6	:	67.1	72.0	17634449

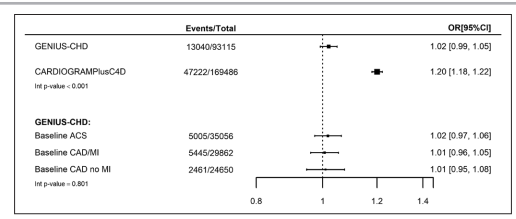


Figure 1. Association between chromosome 9p21 and subsequent coronary heart disease (CHD) events in all participants with baseline CHD (GENIUS-CHD [Genetics of Subsequent Coronary Heart Disease]) compared with association in CHD cases and CHD-free controls (CARDIoGRAM-PlusC4D).

For the CARDIOGRAMPlusC4D consortium (Coronary Artery Disease Genome wide Replication and Meta-analysis [CARDIOGRAM] plus The Coronary Artery Disease [C4D] Genetics) meta-analysis estimate, 6 studies (LURIC, LIFE-Heart [The Leipzig Heart Study], GoDARTS [Genetics of Diabetes Audit and Research in Tayside Scotland], OHGS [Ottawa Heart Genomics Study], PROSPER [Prospective Study of Pravastatin in the Elderly at Risk], WTCCC [Welcome Trust Case Control Consortium]) were excluded as they were also included in GENIUS-CHD. Estimates for GENIUS-CHD are also presented by subtype of CHD at baseline, including acute coronary syndrome (ACS), stable coronary artery disease (CAD) without prior myocardial infarction (MI; CAD/no MI), and stable CAD with prior MI (CAD/MI). All estimates were adjusted for age and sex.

GENIUS-CHD), revealed a per-allele OR for a CHD event similar to that reported previously (OR, 1.20; 95% CI, 1.18–1.22). There was evidence of statistical heterogeneity between the estimates (interaction P<0.001), Figure 1.

Subgroup Analyses

We found minimal evidence for heterogeneity in effect estimates when stratifying by CHD subtype at baseline (interaction *P* value 0.801), with no clear evidence for an effect of chromosome 9p21 genetic variation on subsequent CHD death or MI in individuals enrolled with acute coronary syndrome (OR, 1.02; 95% CI, 0.97–1.06), those with coronary artery disease with a prior MI (OR, 1.01; 95% CI, 0.96–1.05), and those with coronary artery disease without prior MI (OR, 1.01; 95% CI, 0.95–1.08, Figure 1). We further examined the effect of chromosome 9p21 on the primary outcome in prespecified subgroup analyses. We noted a borderline nominally significant interaction with sex, suggesting a greater risk among women with the chromosome 9p21 risk allele, for subsequent CHD death/MI (interaction *P* value = 0.04), whereas nonsignificant trends were noted for greater risk in those without hypertension (*P* value=0.08) or without renal impairment (*P* value=0.17). There were minimal differences in effect estimates by other patient level characteristics including age and diabetes mellitus or according to statin or antiplatelet use or left ventricular impairment at baseline (Figure IV in the Data Supplement).

Similarly, when stratified by study level features, we observed minimal evidence for heterogeneity in effect estimates by study size, geographic region, study design, or length of follow-up (Figure V in the

Subsequent Event	Events/Total		OR[95%CI]
MI	9355/86946	⊷	1.02 [0.99, 1.05]
Revascularization	16818/78732	⊦∎⊣	1.07 [1.04, 1.09]
Heart failure	5955/67665	⊢ ∎	0.97 [0.93, 1.01]
Ischaemic stroke	1311/33591		1.01 [0.93, 1.10]
Any stroke	3133/77317	r - 1	0.99 [0.94, 1.04]
Any CVD	26973/87657	- ∎•	1.04 [1.02, 1.07]
CHD death	6168/81064	r- # -1	1.00 [0.96, 1.04]
CVD death	6610/82012	⊢ ∎	0.97 [0.94, 1.01]
All cause death	14550/99363	H	1.00 [0.97, 1.02]
	0.8	 1 1.2	1.4

Figure 2. Association between chromosome 9p21 and secondary outcomes in participants with baseline CHD, within GENIUS-CHD (Genetics of Subsequent Coronary Heart Disease).

All meta-analysis estimates were adjusted for age and sex. CHD indicates coronary heart disease; CVD, cardiovascular disease; MI, myocardial infarction; and OR, odds ratio. Data Supplement). However, when ordered by date of first enrollment, there was no evidence for variation in effect by time of enrollment (Figure II in the Data Supplement).

Secondary Outcomes

We additionally examined the association between chromosome 9p21 and other subsequent events available for this analysis within the GENIUS-CHD Consortium, listed in Table III in the Data Supplement, with summary estimates provided in Figure 2. Of note, the per-allele effect of risk variants at chromosome 9p21 on subsequent revascularization during follow-up was 1.07 (95% CI, 1.04–1.09). The effect on the composite outcome of any cardiovascular disease, which includes revascularization, was also significant at 1.04 (95% CI, 1.02–1.07). However, there was no clear evidence of association for the remaining secondary outcomes, with only a marginal trend to protection for both subsequent heart failure (OR, 0.97; 95%, CI 0.93–1.01) and cardiovascular disease death (OR, 0.97; 95% CI, 0.94–1.01), as shown in Figure 2.

Selection Bias

To explore the potential for index event bias, we looked for differences in associations between chromosome 9p21 and known cardiovascular risk factors in the United Kingdom Biobank, among the subset of participants with established CHD, compared with the full UKB cohort (Table IV in the Data Supplement). Although there were differences between the groups in the prevalence or values of the tested risk factors, we did not find clear evidence to indicate a distortion in associations between chromosome 9p21 and age, blood pressure, diabetes mellitus, or smoking. There was, however, a small difference for body mass index, with a greater statistical association between the chromosome 9p21 risk allele and lower body mass index identified in those with established CHD than in the general population (nominal interaction P value 0.02. Table IV in the Data Supplement).

We also observed that the chromosome 9p21 risk allele frequency in those surviving with CHD, both in UKB (0.529) and in GENIUS-CHD (0.518, Figure I in the Data Supplement), was higher than the general population in the UKB (0.481) and European reference populations from the 1000 Genomes (Phase 3),⁹ (0.472). This difference in frequency confirms the association of chromosome 9p21 with CHD and also indicated absence of a crude survival bias with loss of large numbers of risk allele carriers to fatal events before entry into CHD cohorts. We did, however, observe a trend to an age association in those with established CHD, as well as the general population in the UKB, with lower chromosome 9p21 risk allele frequencies with advancing age, relative to younger carriers (Figure VI in the Data Supplement).

DISCUSSION

In this study, we examined the effect of genetic variation at the chromosome 9p21 locus on risk of subsequent events in 103357 individuals with established CHD using the newly formed GENIUS-CHD consortium.8 We found that (1) in contrast to the known strong association with CHD observed in CARDIo-GRAMPlusC4D, there was a markedly attenuated and nonsignificant association with subsequent CHD events in GENIUS-CHD; (2) effect estimates in GENIUS-CHD were broadly consistent in stratified analyses based on features related to study design, patient characteristics, and type of index CHD event; and (3) exploratory analyses suggested that selection biases were unlikely to explain the discrepancy. However, we did find evidence of an association between these variants and a secondary outcome of future revascularization events. Our findings, taken together with those from others, support the view that chromosome 9p21 promotes CHD through progressive stable atheroma rather than through development of an unstable phenotype.

The chromosome 9p21 locus is the most widely replicated genetic risk locus for CHD identified to date, with an estimated 15% to 35% increased risk in carriers of the variant allele in prospective population and casecontrol studies.⁵ However, studies examining the effect on subsequent CHD events in people with known CHD at baseline have reported conflicting results.^{10–14} Our group previously examined this in a literature-based meta-analysis, based on 15 studies with median sample size of 1750 individuals, accruing 25163 cases of established CHD, and reported no clear evidence of an effect of variants at chromosome 9p21 on the risk of subsequent events.⁶ An analysis by the CHARGE consortium (The Cohorts for Heart and Aging Research in Genomic Epidemiology) of 2953 MI survivors also reported no association with subsequent mortality.⁷ However, the limited size of most prior studies and the limitations of literature meta-analyses indicate that many possible explanations, including errors in risk allele coding and selection biases, could not be adequately explored, precluding meaningful interpretations for any mechanistic or clinical implications.

The emergence of the GENIUS-CHD Consortium has now permitted a robust evaluation of the role of chromosome 9p21 in subsequent CHD event risk, revealing a clear lack of association with a common composite coronary end point. This is in marked contrast to findings from studies comparing cases to CHD-free controls, as confirmed through meta-analysis of CAR-DIoGRAMPlusC4D data. Furthermore, we were able to add to previous findings by showing that the type of CHD at baseline, whether acute coronary syndrome or stable CHD with or without prior MI, does not alter this association. We also interrogated several widely proposed explanations that could account for our findings through prespecified subgroup analyses and confirmed that most of these, specifically older age, medication use at baseline (statin or antiplatelet), study size or follow-up duration, did not appreciably alter the association findings. Our finding of a possible interaction with sex, warrants further investigation but should be considered hypothesis-generating given the borderline evidence of an interaction.

Selection bias (ie, index event bias or collider-stratification bias) could potentially explain reversed or attenuated associations in disease progression studies like this, operating by inducing relationships between (otherwise independent) risk factors through the selection of individuals with disease.^{15,16} Specifically, individuals surviving a first event consequent on exposure to a particularly strong risk factor may have lower levels of exposure to other individually weaker, independent risk factors, which can then attenuate the association of the risk factor of interest with subsequent events. However, the distribution of common risk factors by chromosome 9p21 genotype did not differ when compared between the general population and the subset with CHD in the UKB, using interaction tests. The only exception was for body mass index, a potentially differential association with chromosome 9p21 in those with CHD compared with the general population was noted. However, the effect size was small in both populations and on its own is unlikely to indicate presence of substantial index event bias.

Selection bias may also theoretically occur by focusing on subjects surviving a first event, where chromosome 9p21 risk allele carriers at risk of fatal CHD events are lost before enrollment into CHD cohorts, thereby diluting the future impact of the variant on subsequent CHD events. In this scenario, we would expect a lower risk allele frequency in those surviving CHD and entering CHD cohorts, but we found no evidence for this. Among those with CHD in the UKB, and among the whole UKB cohort, we did find a progressive loss of risk allele carriers with increasing age, consistent with prior findings of a greater association with CHD, among younger individuals in case-control studies.⁵ Given patients with CHD are generally older, it is possible that a subtle survival bias may still be influencing our findings, although all analyses were adjusted for age. However, based on simulation modeling, sample size, and projected single nucleotide polymorphism effect size, we and others have previously estimated that selection biases are only minimally operating in this context and would be unlikely to account for our observed findings.^{17,18} Although our

findings potentially argue against important selection biases in the analysis for the primary outcome, they are relatively insensitive assessments and may not fully elucidate such biases.

Possible biological explanations could also exist for our findings. Pathological studies indicate differences between chronic stable atherosclerotic plagues that cause ischemia through progressive vessel occlusion and vulnerable plaques with thin caps, prone to sudden plaque rupture, unheralded MI, and coronary deaths.¹⁹ In a seminal study dissecting the phenotype of CHD, a lack of effect for chromosome 9p21 and MI was noted, when both cases and controls had underlying atherosclerosis.²⁰ Our group and others have in parallel shown that chromosome 9p21 robustly associates with atherosclerotic phenotypes,²¹ whereas functional studies have also implicated this region with molecular activity that drives atheroma.²² Furthermore, in this study, we show that the only outcome positively associated with chromosome 9p21 is incident revascularization, perhaps reflecting more severe atherosclerosis burden. Collectively, these data support the concept that chromosome 9p21 promotes progressive atheroma formation and does not confer risk via plaque rupture.

In this context, it is worth noting that chromosome 9p21 associates more robustly with CHD in case-control studies than in prospective cohort studies.⁷ The difference, as proposed by others, could hypothetically be accounted for by incidence-prevalence bias, with chromosome 9p21 carriers more likely to survive a CHD event and thus be over represented among CHD cases (the opposite to survival bias described above).⁷ This becomes more likely as stated above if chromosome 9p21 drives a more progressive and stable atheroma phenotype. If this holds true, then among survivors with established CHD, one might expect that chromosome 9p21 carriers could hold a small favorable advantage over those who experience CHD in its absence, due instead to other more dangerous or vulnerable characteristics, and despite undergoing more subsequent revascularization, these chromosome 9p21 carriers do not experience more dangerous or fatal events.

These findings have important implications. Clinically, they indicate that a degree of caution should be applied when considering or evaluating patients for chromosome 9p21 to predict disease progression or residual risk. They also highlight the need to appreciate important biases that may inflate or attenuate association findings in the setting of subsequent events for individuals with established disease. Mechanistically, these findings support existing and emerging efforts seeking to elucidate the mechanism of the most robust genetic discovery for CHD in recent decades.

There are important limitations to consider. First, among individuals in GENIUS with established CHD,

the timing of the first CHD event or age of onset was often unknown, so we could not account for this variable in our analyses. However, the lack of association in the acute coronary syndrome studies, which had documented timing of the first event, suggests this did not impact the findings. Second, we had limited information on whether subsequent revascularization events were late staged procedures, which would count as part of the index CHD event or unplanned and symptom driven and thereby a true subsequent event, which may have diluted the effect estimate. Third, although we did not observe a specific interaction for statin or aspirin use, we cannot rule out an effect of combined or additional medication usage attenuating the association signal, given the high prevalence of secondary prevention drug use in this setting compared with general population cohorts. Fourth, our analyses were restricted to participants of European descent as most of the included studies only recruited these individuals, and so we were markedly underpowered to explore associations in other ethnic groups. Unfortunately, this remains a wider problem of genetic research and global efforts are ongoing to address this imbalance. Finally, variability of follow-up duration across studies is an analytical challenge and could have impacted our findings, through misclassification. However, a sensitivity analysis stratifying on the follow-up duration of individual studies (<5 or $5 \ge$ years) revealed minimal evidence (P=0.62) of heterogeneity in effect estimates (Figure V in the Data Supplement), suggesting that this is unlikely to have influenced our findings significantly as effect estimates were concordant across studies with different lengths of follow-up. Our major strengths, however, include the size of the study and the large number and types of subsequent events and an effort to examine for selection biases. We also sought to mitigate potential miscoding of the risk allele, given rs1333049 is a palindromic single nucleotide polymorphism, and also the risk allele C changes from being a minor allele in population cohorts to the major allele in CHD cohorts. Finally, this analysis benefitted from the collective expertise and input of over 170 investigators and analysts, many of whom have previously reported on chromosome 9p21.

In conclusion, using the newly formed GENIUS-CHD consortium, we demonstrate that variation at chromosome 9p21 shows no clear association with risk of subsequent CHD events when all individuals have established CHD at baseline. This is in marked contrast to prior case-control studies examining odds of CHD presence compared with disease-free controls. We could not account for the attenuation of effect in terms of selection biases or subgroup effects. However, we did find a greater risk for incident revascularization in those with established CHD, and although residual bias may be at play, our findings collectively support the view that chromosome 9p21 promotes CHD through progressive stable atheroma rather than through development of an unstable phenotype.

ARTICLE INFORMATION

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REFERENCES

 Nikpay M, et al. A comprehensive 1,000 genomes-based genomewide association meta-analysis of coronary artery disease. *Nat Genet*. 2015;47:1121–1130. doi: 10.1038/ng.3396

- Helgadottir A, et al. A common variant on chromosome 9p21 affects the risk of myocardial infarction. *Science*. 2007;316:1491–1493. doi: 10.1126/science.1142842
- McPherson R, et al. A common allele on chromosome 9 associated with coronary heart disease. *Science*. 2007;316:1488–1491. doi: 10.1126/science.1142447
- Samani NJ, et al; WTCCC and the Cardiogenics Consortium. Genomewide association analysis of coronary artery disease. N Engl J Med. 2007;357:443–453. doi: 10.1056/NEJMoa072366
- Palomaki GE, et al. Association between 9p21 genomic markers and heart disease: a meta-analysis. JAMA. 2010;303:648–656. doi: 10.1001/jama.2010.118
- Patel RS, et al. Genetic variants at chromosome 9p21 and risk of first versus subsequent coronary heart disease events: a systematic review and meta-analysis. J Am Coll Cardiol. 2014;63:2234–2245. doi: 10.1016/j.jacc.2014.01.065
- Dehghan A, et al. Genome-wide association study for incident myocardial infarction and coronary heart disease in prospective cohort studies: the CHARGE consortium. *PLoS One*. 2016;11:e0144997. doi: 10.1371/journal.pone.0144997
- Patel RS, et al. Subsequent event risk in individuals with established coronary heart disease: design and rationale of the GENIUS-CHD consortium. *Circ Genom Precis Med.* 2019;12:e002470.
- Genomes Project C, et al. A global reference for human genetic variation. Nature. 2015;526:68–74.
- Ellis KL, et al. A common variant at chromosome 9P21.3 is associated with age of onset of coronary disease but not subsequent mortality. *Circ Cardiovasc Genet.* 2010;3:286–293. doi: 10.1161/CIRCGENETICS.109.917443
- Gong Y, et al. Chromosome 9p21 haplotypes and prognosis in white and black patients with coronary artery disease. *Circ Cardiovasc Genet*. 2011;4:169–178. doi: 10.1161/CIRCGENETICS.110.959296
- Horne BD, et al. Association of variation in the chromosome 9p21 locus with myocardial infarction versus chronic coronary artery disease. *Circ Cardiovasc Genet*. 2008;1:85–92. doi: 10.1161/CIRCGENETICS.108.793158
- Virani SS, et al. Chromosome 9p21 single nucleotide polymorphisms are not associated with recurrent myocardial infarction in patients with established coronary artery disease. *Circ J.* 2012;76:950–956.
- Wauters E, et al. Influence of 23 coronary artery disease variants on recurrent myocardial infarction or cardiac death: the GRACE Genetics Study. *Eur Heart J.* 2013;34:993–1001. doi: 10.1093/eurheartj/ehs389
- Dahabreh IJ, et al. Index event bias as an explanation for the paradoxes of recurrence risk research. JAMA. 2011;305:822–823. doi: 10.1001/jama.2011.163
- Cole SR, et al. Illustrating bias due to conditioning on a collider. Int J Epidemiol. 2010;39:417–420. doi: 10.1093/ije/dyp334
- Anderson CD, et al. The effect of survival bias on case-control genetic association studies of highly lethal diseases. *Circ Cardiovasc Genet*. 2011;4:188–196. doi: 10.1161/CIRCGENETICS.110.957928
- Hu YJ, et al. Impact of selection bias on estimation of subsequent event risk. Circ Cardiovasc Genet. 2017;10.
- 19. Falk E, et al. Update on acute coronary syndromes: the pathologists' view. *Eur Heart J.* 2013;34:719–728. doi: 10.1093/eurheartj/ehs411
- Reilly MP, et al.; Myocardial Infarction Genetics Consortium; Wellcome Trust Case Control Consortium. Identification of ADAMTS7 as a novel locus for coronary atherosclerosis and association of ABO with myocardial infarction in the presence of coronary atherosclerosis: two genome-wide association studies. *Lancet*. 2011;377:383–392. doi: 10.1016/S0140-6736(10)61996-4
- Chan K, et al. Association between the chromosome 9p21 locus and angiographic coronary artery disease burden: a collaborative meta-analysis. J Am Coll Cardiol. 2013;61:957–970. doi: 10.1016/j.jacc.2012.10.051
- Visel A, et al. Targeted deletion of the 9p21 non-coding coronary artery disease risk interval in mice. *Nature*. 2010;464:409–412. doi: 10.1038/nature08801