# Phenome-Wide Association Study (PheWAS) for Detection of Pleiotropy within the Population Architecture using Genomics and Epidemiology (PAGE) Network

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#### **Abstract**

Using a phenome-wide association study (PheWAS) approach, we comprehensively tested genetic variants for association with phenotypes available for 70,061 study participants in the Population Architecture using Genomics and Epidemiology (PAGE) network. Our aim was to better characterize the genetic architecture of complex traits and identify novel pleiotropic relationships. This PheWAS drew on five population-based studies representing four major racial/ethnic groups (European Americans (EA), African Americans (AA), Hispanics/Mexican-Americans, and Asian/Pacific Islanders) in PAGE, each site with measurements for multiple traits, associated laboratory measures, and intermediate biomarkers. A total of 83 single nucleotide polymorphisms (SNPs) identified by genome-wide association studies (GWAS) were genotyped across two or more PAGE study sites. Comprehensive tests of association, stratified by race/ethnicity, were performed, encompassing 4,706 phenotypes mapped to 105 phenotype-classes, and association results were compared across study sites. A total of 111 PheWAS results had significant associations for two or more PAGE study sites with consistent direction of effect with a significance threshold of p<0.01 for the same racial/ethnic group, SNP, and phenotype-class. Among results identified for SNPs previously associated with phenotypes such as lipid traits, type 2 diabetes, and body mass index, 52 replicated previously published genotype-phenotype associations, 26 represented phenotypes closely related to previously known genotype-phenotype associations, and 33 represented potentially novel genotype-phenotype associations with pleiotropic effects. The majority of the potentially novel results were for single PheWAS phenotype-classes, for example, for CDKN2A/B rs1333049 (previously associated with type 2 diabetes in EA) a PheWAS association was identified for hemoglobin levels in AA. Of note, however, GALNT2 rs2144300 (previously associated with high-density lipoprotein cholesterol levels in EA) had multiple potentially novel PheWAS associations, with hypertension related phenotypes in AA and with serum calcium levels and coronary artery disease phenotypes in EA. PheWAS identifies associations for hypothesis generation and exploration of the genetic architecture of complex traits.

Citation: Pendergrass SA, Brown-Gentry K, Dudek S, Frase A, Torstenson ES, et al. (2013) Phenome-Wide Association Study (PheWAS) for Detection of Pleiotropy within the Population Architecture using Genomics and Epidemiology (PAGE) Network. PLoS Genet 9(1): e1003087. doi:10.1371/journal.pgen.1003087

Editor: Greg Gibson, Georgia Institute of Technology, United States of America

Received June 5, 2012; Accepted September 12, 2012; Published January 31, 2013

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Funding: The Population Architecture Using Genomics and Epidemiology (PAGE) program is funded by the National Human Genome Research Institute (NHGRI), supported by U01HG004803 (CALiCo), U01HG004798 (EAGLE), U01HG004802 (MEC), U01HG004790 (WHI), and U01HG004801 (Coordinating Center), and their respective NHGRI ARRA supplements. The contents of this paper are solely the responsibility of the authors and do not necessarily represent the official views of the NIH. The complete list of PAGE members can be found at http://www.pagestudy.org. The data and materials included in this report result from a collaboration between the following studies: The "Epidemiologic Architecture for Genes Linked to Environment (EAGLE)" is funded through the NHGRI PAGE program (U01HG004798-01 and its NHGRI ARRA supplement). Genotyping services for select NHANES III SNPs presented here were also provided by the Johns Hopkins University under federal contract number (N01-HV-48195) from NHLBI and from the University of Washington's Center for Ecogenetics and Environmental Health

(CEEH) pilot study funded by the National Institute of Environmental Health Sciences grant 5 P30 ES007033-12. The study participants derive from the National Health and Nutrition Examination Surveys (NHANES), and these studies are supported by the Centers for Disease Control and Prevention. The findings and conclusions in this report are those of the authors and do not necessarily represent the views of the Centers for Disease Control and Prevention. The Multiethnic Cohort study (MEC) characterization of epidemiological architecture is funded through the NHGRI PAGE program (U01HG004802 and its NHGRI ARRA supplement). The MEC study is funded through the National Cancer Institute (R37CA54281, R01 CA63, P01CA33619, U01CA136792, and U01CA98758). Funding support for the "Epidemiology of putative genetic variants: The Women's Health Initiative" study is provided through the NHGRI PAGE program (U01HG004790 and its NHGRI ARRA supplement). The WHI program is funded by the National Heart, Lung, and Blood Institute; NIH; and U.S. Department of Health and Human Services through contracts N01WH22110, 24152, 32100-2, 32105-6, 32108-9, 32111-13, 32115, 32118-32119, 32122, 42107-26, 42129-32, and 44221. The authors thank the WHI investigators and staff for their dedication, and the study participants for making the program possible. A full listing of WHI investigators can be found at: http://www.whiscience.org/publications/ WHI\_investigators\_shortlist.pdf. Funding support for the Genetic Epidemiology of Causal Variants Across the Life Course (CALiCo) program was provided through the NHGRI PAGE program (U01HG004803 and its NHGRI ARRA supplement). The following studies contributed to this manuscript and are funded by the following agencies: The Atherosclerosis Risk in Communities (ARIC) Study is carried out as a collaborative study supported by National Heart, Lung, and Blood Institute contracts N01-HC-55015, N01-HC-55016, N01-HC-55018, N01-HC-55019, N01-HC-55020, N01-HC-55021, and N01-HC-55022. The Coronary Artery Risk Development in Young Adults (CARDIA) study is supported by the following National Institutes of Health, National Heart, Lung, and Blood Institute contracts: N01-HC-95095, N01-HC-48047, N01-HC-48049, N01-HC-48049, N01-HC-48050, N01-HC-45134, N01-HC-05187, and N01-HC-45205. The Cardiovascular Health Study (CHS) is supported by contracts HHSN268201200036C, N01-HC-85239, N01-HC-85079 through N01-HC-85086, N01-HC-35129, N01 HC-15103, N01 HC-55222, N01-HC-75150, N01-HC-45133, and grant HL080295 from the National Heart, Lung, and Blood Institute (NHLBI), with additional contribution from the National Institute of Neurological Disorders and Stroke (NINDS). Additional support was provided through AG-023629, AG-15928, AG-20098, and AG-027058 from the National Institute on Aging (NIA). The Strong Heart Study (SHS) is supported by NHLBI grants U01 HL65520, U01 HL41642, U01 HL41652, U01 HL41654, and U01 HL65521. The opinions expressed in this paper are those of the author(s) and do not necessarily reflect the views of the Indian Health Service. Assistance with phenotype harmonization, SNP selection and annotation, data cleaning, data management, integration and dissemination, and general study coordination was provided by the PAGE Coordinating Center (U01HG004801-01 and its NHGRI ARRA supplement). The National Institutes of Mental Health also contributes to the support for the Coordinating Center. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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#### Introduction

Phenomic approaches are complementary to the more prevalent paradigm of genome-wide association studies (GWAS), which have provided some information about the contribution of genetic variation to a wide range of diseases and phenotypes [1]. While a typical GWAS evaluates the association between the variation of hundreds of thousands, to over a million, genotyped single nucleotide polymorphisms (SNPs) and one or a few phenotypes, a common limitation of GWAS is the focus on a pre-defined and limited phenotypic domain. An alternate approach is that of PheWAS, which utilizes all available phenotypic information and all genetic variants in the estimation of associations between genotype and phenotype [1]. By investigating the association between SNPs and a diverse range of phenotypes, a broader picture of the relationship between genetic variation and networks of phenotypes is possible.

A challenge for PheWAS is the availability of large studies with genotypic data that are also linked to a wide array of high quality phenotypic measurements and traits for study. Biorepositories linked to electronic medical records (EMR) have been an initial resource for PheWAS, but these EMR-based studies are often limited to phenotypes and traits commonly collected for clinical use and may represent sets of limited racial/ethnic diversity [2,3]. While there is no U.S. national, population-based cohort [4], several diverse, population-based studies exist with tens of thousands of samples linked to detailed survey, laboratory, and medical data. These large population-based studies have limitations [5], but collectively [6] they offer an opportunity to perform a PheWAS of unprecedented size and diversity.

To capitalize on the potential for collaborative discovery among some of the large population-based studies of the U.S., the National Human Genome Research Institute (NHGRI) funded the Population Architecture using Genomics and Epidemiology (PAGE) network. PAGE includes eight extensively characterized, large population-based epidemiologic studies where data were collected across multiple racial/ethnic groups, supported by a coordinating center [7], providing an exceptional opportunity to pursue PheWAS with a large number of SNPs, and thousands of phenotypic measurements including a wide range of common diseases, risk factors, intermediate biomarkers and quantitative traits in diverse populations. Herein, we illustrate the feasibility and utility of the PheWAS approach for large

population-based studies and demonstrate that PheWAS provides information on, and exposes the complexity of, the relationship between genetic variation and interrelated and independent phenotypes. We have found PheWAS results that replicate previously identified genotype-phenotype associations with the exact phenotype in previous associations or closely related phenotypes, as well as a series of novel genotype-phenotype associations. This data exploration method exposes a more complete picture of the relationship between genetic variation and phenotypic outcome. PheWAS provides the unbiased, high throughput design achieved by GWAS in the genome and phenotype domains simultaneously. This approach changes the paradigm of phenotypic characterization and allows for exploratory research in both genomics and phenomics.

#### Results

Data from five PAGE study sites were available for this PheWAS: Epidemiologic Architecture for Genes Linked to Environment (EAGLE) using data from the National Health and Nutrition Examination Surveys (NHANES); the Multiethnic Cohort Study (MEC); the Women's Health Initiative (WHI); and two studies of the Causal Variants Across the Life Course (CALiCo) group: the Cardiovascular Health Study (CHS) and Atherosclerosis Risk in Communities (ARIC). Text S1 provides full information on study design, phenotype measurement, and genotyping for each study. These studies collectively include four major racial/ethnic groups: European Americans (EA), African Americans (AA), Hispanics/Mexican Americans (H), and Asian/ Pacific Islanders (API). All PAGE study sites included both males and females, except for WHI (which includes only women). Table 1 provides an overview of the sample sizes by PAGE study site as well as the number of SNPs and phenotypes available for this PheWAS. Sample size and the number of phenotypes varied across studies, and the sample size for various phenotypes within each study varied dependent on the number of individuals for which a given phenotype was measured. The number of phenotypes available for this PheWAS ranged within studies from 63 (MEC) to 3,363 (WHI). Study sites also had differing numbers of genotyped SNPs, and Table S1 contains the list of all SNPs available for two or more sites in this study, arranged by previously associated phenotypes. The PAGE network has focused on characterization of well-replicated variants across multiple race/

#### **Author Summary**

In phenome-wide association studies (PheWAS) all potential genetic variants in a dataset are systematically tested for association with all available phenotypes and traits that have been measured in study participants. By investigating the relationship between genetic variation and a diversity of phenotypes, there is the potential for uncovering novel relationships between single nucleotide polymorphisms (SNPs), phenotypes, and networks of interrelated phenotypes. PheWAS also can expose pleiotropy, provide novel mechanistic insights, and foster hypothesis generation. This approach is complementary to genome-wide association studies (GWAS) that test the association between hundreds of thousands, to over a million, single nucleotide polymorphisms and a single phenotype or limited phenotypic domain. The Population Architecture using Genomics and Epidemiology (PAGE) network has measures for a wide array of phenotypes and traits, including prevalent and incident status for clinical conditions and risk factors, as well as clinical parameters and intermediate biomarkers. We performed tests of association between a series of genome-wide association study (GWAS)-identified SNPs and a comprehensive range of phenotypes from the PAGE network in a high-throughput manner. We replicated a number of previously reported associations, validating the PheWAS approach. We also identified novel genotypephenotype associations possibly representing pleiotropic effects.

ethnicities, so each study independently genotyped a set of SNPs with previously reported associations with phenotypes such as body mass index, C-reactive protein, and lipid levels.

Tests of association assuming an additive genetic model were performed independently by each PAGE study site for each SNP and each phenotype, stratified by race/ethnicity. The last column of Table 1 presents the total number of comprehensive associations with and without a p-value cutoff of 0.01, showing the proportion of significant results for this many tests of association. The total number of tests of association ranged from >20,000

(MEC) to >1 million (WHI) reflecting the variability in both the number of phenotypes available for study as well as the number of SNPs genotyped by each PAGE study site. As expected, the total number of significant tests of association (p<0.01) represented a fraction of the total number of tests performed.

Results from these tests of association were then compared across study sites to identify overlapping significant associations, as these results most likely represent robust findings. To facilitate determining overlapping significant associations, similar phenotypes that existed across more than one study were binned into 105 distinct phenotype-classes. For some phenotypes, the specific phenotype existed across more than one PAGE study, such as for the phenotype "Hemoglobin", where hemoglobin measurements were available for ARIC, CHS, EAGLE, and WHI. Other groups of phenotypes binned within phenotype-classes were within similar phenotypic domains but were not represented in exact same form across studies. Table S2 contains a list of the study level phenotypes, the study from which the phenotype is available, and the phenotype-class for each phenotype that overlapped with another study.

The same or similar phenotypes may or may not have been collected by each PAGE study. Thus, the number of studies that were available for comparison of results across studies varied from one phenotype-class to another phenotype-class. Table 2 presents the number of results where at least two of five independent studies had SNP-phenotype associations with p<0.01 for single phenotype-class and single race/ethnicity group, compared to the total number of SNP-phenotype association tests performed. For example, >8,500 tests of association for the same SNP and same phenotype were available from two PAGE study sites whereas only 906 and 58 tests of association were available from four and five PAGE study sites, respectively. There were 3 results where two or more of the groups had a SNP-phenotype association p<0.01 for a single phenotype class across 5 groups represented.

For this PAGE-wide PheWAS, tests of association were considered significant across PAGE study sites where two or more phenotypes in the same phenotype-class in the same racial/ethnic group passed a significance threshold of p<0.01 with a consistent direction of genetic effect. Based on these criteria, a total of 111

Table 1. Study Descriptions.

				Maximu	ım Sampl	e Size <sup>3</sup>		Mini	mum S	ample	Size <sup>3</sup>			
Study <sup>1</sup>	PMID <sup>2</sup>	Age Range	Sex	EA	AA	н	API	EA	AA	н	API	# SNPs <sup>4</sup>	# Phenotypes <sup>5</sup>	Total # of Tests <sup>6</sup> (# p<0.01)
ARIC	[36]	45-64	M/F	11,068	4,007	NA	NA	17	7	NA	NA	69	612	138,207 (2,378)
CHS	[37]	65–100 at baseline	M/F	4,487	820	NA	NA	151	116	NA	NA	46	341	34,829 (550)
EAGLE	[38]	12-95	M/F	2,628	2,107	2,071	NA	7	16	15	NA	236	327	359,508 (6,496)
MEC	[39]	45-75	M/F	3,893	4,749	6,863	6,810	33	27	40	13	74	63	23,310 (212)
WHI	[40]	50–79 at baseline	F	13,334	4,274	2,023	927	14	5	7	5	94	3,363	1,123,366 (14,068)

Abbreviations: European American (EA), African American (AA), Hispanic/Mexican American (H), Asian/Pacific Islander (API),

<sup>1</sup>Data from PAGE studies available for this PheWAS include: Atherosclerosis Risk in Communities (ARIC), Cardiovascular Health Study (CHS), Epidemiologic Architecture for Genes Linked to Environment (EAGLE), Multiethnic Cohort (MEC), and Women's Health Initiative (WHI). PAGE study sites and study design descriptions are in Text S1.
<sup>2</sup>Pubmed ID of study description manuscript for each study.

<sup>3</sup>Maximum sample size and Minimum sample size are dependent both on who was genotyped and who had a specific phenotype measured. Not all phenotypic measurements were available for all participants within each study.

<sup>4</sup>This is the total number of SNPs available for each study. Table S1 has the list of these SNPs for each study, genotyped across two or more studies.

<sup>5</sup>This includes the number of phenotypes transformed and untransformed, as well as categorical phenotypes divided into binary phenotypes, full description in Materials and Methods.

<sup>6</sup>Total number of tests of association calculated for each study, in parenthesis is the total number of associations with p<0.01. doi:10.1371/journal.pgen.1003087.t001

**Table 2.** The number of SNP-Phenotype tests of association for phenotype-classes varies by PAGE study site genotype and phenotype overlap.

Number of PAGE study sites <sup>1</sup>	Number of Total Tests of Association <sup>2</sup>	Number of Significant Tests of Association <sup>3</sup>
2	8680	45
3	3295	42
4	906	21
5	58	3*
Total	12939	111

<sup>1</sup>Number of PAGE study sites where both the SNP and phenotype were available for a given phenotype class.

PheWAS associations were identified (Table S3). Overall, among the 111 significant PheWAS associations identified, 52 PheWAS results replicated previously published genotype-phenotype associations (Table S4), 26 represented phenotype-classes closely related to previously known genotype-phenotype associations (Table 3), and 33 represented novel genotype-phenotype associations (Table 4).

#### Known Associations—Validating the PheWAS Approach

Almost half of the PAGE PheWAS results (52/111; 48%) replicated previously known genotype-phenotype associations. These replicated results serve as positive controls and demonstrate that the high-throughput PheWAS approach is feasible and valid. As an example, low-density lipoprotein cholesterol (LDL-C) has previously been associated with rs4420638 near APOE/APOC1/ C1P1/C2/C4 in European Americans [8,9]. In the PAGE PheWAS, a significant association between the same SNP and LDL-C phenotypes of the "LDL-C" phenotype-class in European Americans as reported in the literature [8,9] was observed in two PAGE study sites, with the same direction of effect  $(\beta)$  as well as a third PAGE site with near significant results: ARIC  $(p=1.27\times10^{-15},\ \beta=-5.75),\ CHS\ (p=7.89\times10^{-12},\ \beta=-7.06),$ and WHI (p = 0.06,  $\beta$  = -4.15). Figure 1 shows the significant PheWAS LDL-C results, as well as other associations considered significant for rs4420638 across PAGE study sites for other phenotype-classes in a similar racial/ethnic group passed a significance threshold of p<0.01 with a consistent direction of genetic effect.

#### **Related Associations**

Approximately one-fourth of the PAGE PheWAS results (26/111; 23%) represented SNP-phenotype associations in phenotype-classes closely related to previously known genotype-phenotype associations. For example, rs10757278 near *CDKN2A/CDKN2B* has been robustly associated with myocardial infarction (MI) [10,11]. In this PheWAS, rs10757278 was associated with the "Cardiac" phenotype-class, but also with the related phenotype-classes of "Artery Treatment" and "Angina". Specifically, rs10757278 was associated with phenotypes in the Artery Treatment phenotype-class, such as "percutaneous transluminal coronary angioplasty" (WHI, p =  $2.86 \times 10^{-6}$ ,  $\beta = -0.17$ , EA), and "coronary bypass surgery" (CHS, p =  $9.60 \times 10^{-3}$ ,  $\beta = -0.26$ , EA). The SNP rs10757278 was also associated with phenotypes in the Angina phenotype-class, such as presence or absence of angina (WHI, p =  $6.59 \times 10^{-3}$ ,  $\beta = -0.14$ , EA) and the phenotype "Ever

see a doctor because of chest pain?" (ARIC,  $p=4.44\times10^{-3}$ ,  $\beta=-0.31$ , EA). Replication of association of this SNP with previously known phenotypes were also found with the phenotype-class "Cardiac", with phenotypes such as "MI (Y/N)" (WHI,  $p=1.39\times10^{-4}$ ,  $\beta=-0.11$ , EA), and "MI status at baseline (Y/N)" (CHS,  $p=6.35\times10^{-3}$ ,  $\beta=-0.18$ , EA). Significant PheWAS associations at p<0.01 for rs10757278 are plotted by phenotype in Figure 2, as well as additional results at p<0.05.

Another example of PheWAS associations for phenotype-classes closely related to known genotype-phenotype associations existed for rs599839 near the CELSR2/PSRC1/SORT1 gene cluster. The SNP rs599839 has been associated with serum LDL cholesterol levels [8,12-14], and coronary artery disease [13,15]. In our PheWAS, associations were found for the "LDL-C" phenotypeclass, as well the coronary artery disease related "Angina" and lipid related "HDL-C" phenotype-classes, including specific phenotypes such as "angina, presence or absence of" (WHI,  $p = 2.10 \times 10^{-4}$ ,  $\beta = 0.25$ , EA), and "HDL-C"  $p = 1.23 \times 10^{-3}$ ,  $\beta = -0.04$ , AA). As expected, a significant association was also identified for the LDL-C level related phenotype "LDL-C (mg/dl)" (ARIC,  $p = 5.25 \times 10^{-22}$ ,  $\beta = 6.42$ EA). Significant PheWAS associations at p<0.01 for rs599839 are plotted by phenotype in Figure 3, as well as additional results at p < 0.05.

#### **Potentially Novel Associations**

PheWAS results were considered novel, if the significant phenotype-class associations varied substantially from the previously reported GWAS and candidate gene studies. Approximately one-third of the PAGE PheWAS results (33/111; 30%) represented novel genotype-phenotype-class associations. Further research will be required to determine the further validity of these exploratory results.

The most statistically significant of the novel phenotype-class associations identified by this PheWAS include multiple associations involving phenotype-classes for hematologic traits in African Americans (Figure 4). SNPs rs599839 (CELSR2/PSRC1), rs10923931 (NOTCH2), rs2228145 (IL6R), rs2144300 (GALNT2), rs10757278 (CDKN2A,CDKN2B), and rs7901695 (TCF7L2) were each associated with white blood cell count phenotypes among AA (significant p-values ranging  $7.96\times10^{-3}$  to  $9.99\times10^{-15}$ ). IL6R rs2228145 was also associated with neutrophils and lymphocyte numbers in AA with p-values ranging from  $2.44\times10^{-4}$  to  $4.66\times10^{-10}$ . These SNPs were previously associated with LDL-C, total cholesterol levels, and coronary artery disease (rs599839)

<sup>&</sup>lt;sup>2</sup>Total number of tests of association, by number of PAGE study sites, where both the SNP and phenotype were available for a given phenotype class.

<sup>3</sup>Number of tests of association that was significant (p<0.01) for two or more PAGE study sites for a single phenotype class and SNP, taking into account matching

<sup>&</sup>lt;sup>3</sup>Number of tests of association that was significant (p<0.01) for two or more PAGE study sites for a single phenotype class and SNP, taking into account matching direction of effect when phenotypically relevant.

<sup>\*</sup>Three results where two or more of the groups had a SNP-phenotype association p<0.01 for a single phenotype class across 5 groups represented. The most replicated novel results across studies were for SNPs rs599839, rs10923931, and rs2228145 with hematologic traits. doi:10.1371/journal.pgen.1003087.t002

 Table 3. PheWAS Tests of Association: Related Associations.

Nearest Gene	SNP ID	CA¹, (CAF²)	Phenotypes <sup>3</sup>	Associated Phenotype Class <sup>4</sup>	Ethnicity <sup>5</sup>	P-Values <sup>6</sup>	Beta(SE) <sup>7</sup>	Sample Size <sup>8</sup>	Substudies <sup>9</sup>	Substudy Count <sup>10</sup>	Previously Associated Phenotype <sup>11</sup>	References <sup>12</sup>
CELSR2, PSRC1	RS599839	A (0.8)	Ever hospitalized for chest pain? (Y/N)	ANGINA	EA	2.57e-3	0.55(0.18)	268	ARIC	3	Coronary Artery Disease, LDL cholesterol, Total cholesterol	18262040, 18193043, 18179892, 17634449, 18193044
			Angina status at baseline (Y/N)			8.79e-3	0.21(0.08)	4482	CHS			
			Angina (Y/N)			2.10e-4	0.25(0.07)	13308	MHI			
			Ever had pain/ discomfort in your chest (Y/N)			1.06e-4	0.21(0.05)	4477	СНЅ			
CDKN2A, CDKN2B	, RS1333049	C (0.5)	Chest pain or discomfort? (Y/N)	ANGINA	EA	4.69e-3	0.08(0.03)	9338	ARIC	3	Coronary Artery Disease, Type 2 17554300, 17634449 Diabetes, Hypertension	17554300, 17634449
			See a doctor because of chest pain? (Y/N)			8.80e-3	0.32(0.12)	572	ARIC			
			Ever had pain/discomfort in your chest (Y/N)			9.72e-3	0.11(0.04)	4472	CHS			
			Angina (Y/N)			7.75e-3	0.14(0.05)	13331	MHI			
LPL	RS6586891	A (0.6)	LN1 Total triglycerides in mmol/l	Serum Triglycerides	EA	7.76e-7	-0.02(4.75e-3) 9326	) 9326	ARIC	ю	HDL Cholesterol	18193043
			Triglyceride (mg/dl)			3.45e-3	-5.18(1.77)	4469	CHS			
			LN1 core triglyceride (mg/dl)			6.68e-3	-0.06(0.02)	922	MHI			
			Total Triglycerides (mmol/l)			3.50e-4	-0.06(0.02)	9326	ARIC			
			LN1 Triglycerides (mg/dl)			2.45e-3	-0.03(9.65e-3) 4469	4469	CHS			
CDKN2A, CDKN2B	, RS10757278	A (0.5)	Ever had pain/ discomfort in your chest (Y/N)	ANGINA	EA	5.96e-3	-0.12(0.04)	4474	СНЅ	3	Myocardial infarction	17478679
			Angina (Y/N)			6.59e-3	-0.14(0.05)	13313	MHI			
			Chest pain or discomfort? (Y/N)			9.73e-3	-0.07(0.03)	11047	ARIC			
			See a doc because of chest pain? (Y/N)			4.44e-3	-0.31(0.11),	722	ARIC			
			Rose angina (Y/N)			8.17e-3	0.16(0.06)	11040	ARIC			
LDLR	RS6511720	G (0.9)	Total cholesterol (mmol/I)	Serum Cholesterol	EA	6.84e-16	0.18(0.02)	10891	ARIC	2	LDL Cholesterol	18193043
			Total cholesterol (mg/dl)			2.64e-5	12.05(2.85)	924	MHI			
			LN1 Total cholesterol (mmol/l)	<b>&gt;</b>		4.15e-17	0.03(3.30e-3)	10891	ARIC			
			High cholesterol requiring pills ever (Y/N)			10.00e-12	0.42(0.06)	12431	MHI			

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Nearest Gene	SNP ID	CA <sup>1</sup> , (CAF <sup>2</sup> )	Phenotypes <sup>3</sup>	Associated Phenotype Class <sup>4</sup>	Ethnicity <sup>5</sup>	P-Values <sup>6</sup>	Beta(SE) <sup>7</sup>	Sample Size <sup>8</sup>	Substudies <sup>9</sup>	Substudy Count <sup>10</sup>	Previously Associated Phenotype <sup>11</sup>	References <sup>12</sup>
			LN1 Total cholesterol (mg/dl)			1.87e-5	0.05(0.01)	924	MHI			
PCSK9	RS11591147	(66.0) D	Total cholesterol (mmol/l)	Serum Cholesterol	EA	1.13e-16	0.46(0.06)	10833	ARIC	2	LDL Cholesterol, lipids	18193044
			Total cholesterol (mg/dl)			8.47e-3	18.62(7.06)	924	MHI			
			LN1 total cholesterol (mg/dl)			4.67e-3	0.09(0.03)	924	MHI			
			High cholesterol requiring pills ever (Y/N)			4.72e-6	0.92(0.20)	12419	MHI			
			LN1 total cholesterol (mmol/l)			1.59e-18	0.07(8.39e-3)	10833	ARIC			
APOE, APOC1, APOC4, APOC2	RS4420638	A (0.8)	High cholesterol requiring pills ever (Y/N)	Serum Cholesterol	EA	1.18e-6	-0.22(0.04)	12430	WHI	2	LDL Cholesterol, lipids, Alzheimer's disease, Coronary Artery Disease, C-reactive protein, Sporadic late onset Alzheimer's disease, ApoB, Triglycerides, Total cholesterol	; 17474819, 17463246, 18193043, 18802019, 17998437, 19197348, 19567438
			Total cholesterol (mmol/l)			2.31e-12	-0.14(0.02)	9323	ARIC			
			LN1 Total cholesterol (mmol/l)			1.96e-13	-0.02(3.00e-3) 9323	9323	ARIC			
			High cholesterol requiring pills ever (Y/N)			1.18e-6	-0.22(0.04)	12430	MHI			
			Total cholesterol (mmol/l)			2.31e-12	-0.14(0.02)	9323	ARIC			
			LN1 Total cholesterol (mmol/l)			1.96e-13	-0.02 (3.00e-3)	9323	ARIC			
TMEM18	RS6548238	C (0.8)	Hip circumference cm	Hip Circumference	EA	5.99e-3	0.62(0.22)	4456	CHS	2	Body Mass Index	19079261
			Hip circumference cm			7.12e-3	0.59(0.22)	13272	WHI			
			LN1 hip circumference cm			6.85e-3	5.84e-3(2.16e-3)4456	3)4456	CHS			
			LN1 hip circumference cm			3.93e-3	5.60e-3(1.94e-3)13272	3)13272	WHI			
TMEM18	RS6548238	C (0.9)	High blood pressure ever diagnosed?	Hypertension	AA	7.69e-3	0.19(0.07)	3946	ARIC	2	Body Mass Index	19079261
			Pills for hypertension now (Y/N)			7.17e-3	0.18(0.07)	4122	MHI			
			Hypertension category: never hypertensive (Y/N)			1.03e-3	-0.22(0.07)	4076	MHI			
			Hypertension ever (Y/N)			5.85e-4	0.23(0.07)	4206	MHI			
			Pills for hypertension ever (Y/N)			6.30e-3	0.19(0.07)	4186	MHI			

Table 3. Cont.

Nearest Gene	SNP ID	CA¹, (CAF²)	Phenotypes <sup>3</sup>	Associated Phenotype Class <sup>4</sup>	Ethnicity <sup>5</sup>	P-Values <sup>6</sup>	Beta(SE) <sup>7</sup>	Sample Size <sup>8</sup>	Substudies <sup>9</sup>	Substudy Count <sup>10</sup>	Previously Associated Phenotype <sup>11</sup>	References <sup>12</sup>
			Hypertension category: treated hypertensive (Y/N)			6.54e-3	0.19(0.07)	4076	MHI			
FTO	RS3751812	T (0.4)	Waist circumference (cm)	WaistSize	EA	3.88e-3	1.33(0.46)	2360	EAGLEIII	2	Body Mass Index	17658951
			LN1 waist circumference cm			3.68e-8	0.01(1.99e-3)	13282	MHI			
			LN1 waist circumference (cm)			2.73e-3	0.01(4.99e-3)	2360	EAGLEIII			
			Waist circumference cm			2.32e-8	1.03(0.18)	13282	MHI			
CELSR2, PSRC1	RS599839	A (0.3)	Re-calibrated HDL(3) cholesterol in mg/dl	HDL	AA	3.07e-3	-1.00(0.34)	3023	ARIC	2	Coronary Artery Disease, LDL cholesterol, Total cholesterol	18262040, 18193043, 18179892, 17634449, 18193044
			LN1 high-density lipoprotein-3 (mg/dl)			2.55e-3	-0.03(9.54e-3) 974	974	WHI			
			LN1 HDL Cholesterol in mg/dl	=		4.60e-3	-0.02(8.74e-3) 3024	3024	ARIC			
			High-density lipoprotein-2 (mg/dl)			2.94e-3	-1.09(0.36)	974	WHI			
			LN1 re-calibrated HDL(3) cholesterol in mg/dl			2.06e-3	-0.03(9.10e-3) 3023	3023	ARIC			
			HDL cholesterol (mg/dl)			1.51e-3	-2.20(0.69)	982	MHI			
			LN1 high-density lipoprotein-2 (mg/dl)			2.43e-3	-0.06(0.02)	974	WHI			
			LN1 HDL cholesterol (mg/dl)			1.23e-3	-0.04(0.01)	982	MHI			
			High-density lipoprotein-3 (mg/dl)			2.70e-3	-1.20(0.40)	974	MHI			
CDKN2A, CDKN2	RS1333049	C (0.5)	Percutaneous transluminal coronary angioplasty (Y/N)	Artery Treatment EA	EA	2.93e-6	0.17(0.04)	13331	WHI	2	Coronary Artery Disease, Type 2 17554300, 17634449 Diabetes, Hypertension	17554300, 17634449
			Coronary bypass surgery (Y/N)			6.61e-3	0.28(0.10)	4446	CHS			
CDKN2A, CDKN2B	RS1333049	C (0.5)	Myocardial infarction status Cardiac at baseline (Y/N)	Cardiac	EA	8.24e-3	0.17(0.06)	4477	CHS	2	Coronary Artery Disease, Type 2 17554300, 17634449 Diabetes, Hypertension	17554300, 17634449
			Myocardial infarction (Incident or Prevalent))			2.03e-3	0.22(0.07)	4477	CHS			
			Myocardial infarction (Y/N)			8.78e-5	0.11(0.03)	13331	MHI			
APOA1, APOC3, APOA4, APOA5	RS964184	G (0.3)	LN1 serum cholesterol (mg/dl)	Serum Cholesterol	MA	2.18e-3	0.02(7.61e-3),	2034	EAGLEIII	2	HDL Cholesterol, lipids, Triglycerides, Triglyceride levels	18193043

Table 3. Cont.

Nearest Gene	SNP ID	CA¹, (CAF²)	Phenotypes <sup>3</sup>	Associated Phenotype Class <sup>4</sup>	Ethnicity <sup>5</sup>	ese	Beta(SE) <sup>7</sup>	Sample Size <sup>8</sup>	dies	Substudy Count <sup>10</sup>	Previously Associated Phenotype <sup>11</sup>	References <sup>12</sup>
			LN1 serum cholesterol (mmol/l)			2.91e-3	0.02(6.36e-3)	2040	EAGLEIII			
			Serum cholesterol (mmol/l)			2.50e-3	0.12(0.04)	2034	EAGLEIII			
			High cholesterol requiring pills ever (Y/N)			3.39e-3	0.29(0.10)	1859	IHM			
			Serum cholesterol (mmol/l)			3.44e-3	0.12(0.04)	2040	EAGLEIII			
			LN1 serum cholesterol (mg/dl)			2.98e-3	0.02(7.64e-3)	2040	EAGLEIII			
			Serum cholesterol (mg/dl)			3.42e-3	4.50(1.53)	2040	EAGLEIII			
			LN1 serum cholesterol: si (mmol/l)			2.10e-3	0.02(6.35e-3)	2034	EAGLEIII			
			Serum cholesterol (mg/dl)			2.50e-3	4.64(1.53)	2034	EAGLEIII			
APOA1, APOC3, APOA4, APOA5	RS964184	G (0.1)	LN1 total cholesterol (mg/dl) Serum Choles	) Serum Cholesterol	EA	1.53e-3	0.03(0.01)	924	WHI	2	HDL Cholesterol, lipids, Triglycerides, Triglyceride levels	18193043
			Total cholesterol (mg/dl)			3.02e-3	7.46(2.51)	924	MHI			
			Serum cholesterol (mmol/l)			8.29e-3	0.13(0.05)	2592	EAGLEIII			
			High cholesterol requiring pills ever (Y/N)			2.06e-14	0.35(0.05)	12433	WHI			
			Serum cholesterol (mg/dl)			8.28e-3	4.89(1.85)	2592	EAGLEIII			
TIMD4, HAVCR1	RS1501908	C (0.6)	LN1 serum cholesterol (mmol/l)	Serum Cholesterol	EA	9.63e-3	0.01(5.48e-3)	2582	EAGLEIII	2	LDL Cholesterol	19060906
			LN1 total cholesterol in mmol/l			4.04e-4	8.64e-3(2.44e-3)9323	)9323	ARIC			
			Serum cholesterol (mmol/l)			9.35e-3	0.09(0.03)	2584	EAGLEIII			
			Serum cholesterol (mg/dl)			9.40e-3	3.45(1.33)	2584	EAGLEIII			
			Total cholesterol in mmol/l			7.27e-4	0.05(0.02)	9323	ARIC			
			LN1 serum cholesterol (mmol/l)			8.93e-3	0.01(5.43e-3)	2584	EAGLEIII			
			LN1 serum cholesterol (mg/dl)			9.83e-3	0.02(6.52e-3)	2582	EAGLEIII			
			LN1 serum cholesterol (mg/dl)			9.17e-3	0.02(6.49e-3)	2584	EAGLEIII			
	RS6511720	G (0.9)	Total cholesterol in mmol/l	Serum Cholesterol	AA	1.50e-10	0.25(0.04)	3786	ARIC	2	LDL Cholesterol	18193043

Table 3. Cont.

Nearest		٦		Associated Phenotype				Sample		Substudy	Previously Associated	
Gene	SNP ID	(CAF <sup>2</sup> )	Phenotypes <sup>3</sup>		Ethnicity <sup>5</sup>	P-Values <sup>6</sup>	Beta(SE) <sup>7</sup>	Size	Substudies <sup>9</sup>	Count 10	Phenotype <sup>11</sup>	References <sup>12</sup>
			LN1 total cholesterol (mg/dl)			8.37e-4	0.04(0.01)	1009	WHI			
			High cholesterol requiring pills ever (Y/N)			2.21e-3	0.31(0.10)	4041	MH			
			Total cholesterol (mg/dl)			1.36e-3	8.31(2.59)	1009	WHI			
			LN1 total cholesterol in mmol/l			6.45e-11	0.04(5.85e-3)	3786	ARIC			
ABCG8	RS6544713	C (0.7)	High cholesterol requiring pills ever (Y/N)	Serum Cholesterol	EA	7.94e-5	-0.14(0.04)	12438	MHI	2	LDL Cholesterol	19060906
			Total cholesterol in mmol/l			2.23e-9	-0.09(0.02)	10826	ARIC			
			LN1 total cholesterol in mmol/l			2.13e-9	-0.01(2.34e-3) 10826	10826	ARIC			
APOB	RS754523	T (0.7)	LN1 total cholesterol in mmol/l	Serum Cholesterol	EA	3.01e-10	-0.02(2.49e-3) 9314	9314	ARIC	2	LDL Cholesterol	19750184
			High cholesterol requiring pills ever (Y/N)			4.76e-4	-0.13(0.04)	12427	MH			
			Total cholesterol in mmol/l			5.52e-10	-0.10(0.02)	9314	ARIC			
APOE, APOC1, APOC4, APOC2	RS4420638	A (0.8)	LN1 dietary cholesterol (mg) (dietary consumption)	Cholesterol MG	EA	1.48e-3	0.03(9.34e-3)	9194	ARIC	2	LDL Cholesterol, lipids, 17474819, 17463246, Alzheimer's disease, Coronary 18193043, 18802019, Artery Disease, C-reactive protein,17998437, 19197348, Sporadic late onset Alzheimer's 19567438 disease, ApoB, Triglycerides, Total cholesterol	17474819, 17463246, 18193043, 18802019, 1,17998437, 19197348, 19567438
			Dietary cholesterol (mg)			9.48e-3	5.84(2.25)	13291	WHI			
			LN1 dietary cholesterol (mg)			4.35e-3	0.03(9.66e-3)	13291	MHI			
			Dietary cholesterol (mg)			3.28e-3	6.94(2.36)	9194	ARIC			
CDKN2A, CDKN2B	RS10757278	A (0.5)	Coronary bypass surgery (Y/N)	Artery Treatment EA	EA	9.60e-3	-0.26(0.10)	4448	CHS	2	Myocardial infarction	17478679
			Percutaneous transluminal coronary angioplasty (Y/N)			2.86e-6	-0.17(0.04)	13313	HW			
FTO	RS8050136	A (0.4)	LN1 hip circumference cm	Hip Circumference	EA	3.11e-7	7.67e-3(1.50e-3)13272	3)13272	WHI	2	Obesity, Type 2 diabetes	17554300, 19079260, 17463249, 17463248, 18372903, 18159244, 19056611, 17658951

Table 3. Cont.

Nearest Gene	SNP ID	CA¹, (CAF²)	Phenotypes <sup>3</sup>	Associated Phenotype Class <sup>4</sup>	Ethnicity <sup>5</sup>	P-Values <sup>6</sup>	Beta(SE) <sup>7</sup>	Sample Size <sup>8</sup>	Substudies <sup>9</sup>	Substudy Count <sup>10</sup>	Previously Associated Phenotype <sup>11</sup>	References <sup>12</sup>
			LN1 hip girth to nearest cm			1.26e-8	7.35e-3(1.29e-3)9334	3)9334	ARIC			
			Hip girth to nearest cm			7.26e-9	0.81(0.14)	9334	ARIC			
			Hip circumference cm			6.06e-7	0.84(0.17)	13272	WHI			
FTO	RS8050136	A (0.4)	LN1 waist hip ratio	Waist Hip Ratio	EA	7.15e-3	1.46e-3(5.43e-4)13269	t)13269	MHI	2	Obesity, Type 2 diabetes	17554300, 19079260, 17463249, 17463248, 18372903, 18159244, 19056611, 17658951
			Waist-to-hip ratio			1.71e-7	6.11e-3(1.17e-3)9333	1)9333	ARIC			
			LN1 waist-to-hip ratio			1.75e-7	3.21e-3(6.10e-4)9333	1)9333	ARIC			
H0	RS8050136	A (0.4)	LN1 waist circumference cm	Waist Size	EA	2.88e-8	0.01(1.99e-3)	13284	WHI	2	Obesity, Type 2 diabetes	17554300, 19079260, 17463249, 17463248, 18159244, 19056611, 17658951
			LN1 waist girth to nearest cm	E C		3.63e-12	0.01(2.01e-3)	9334,9334 ARIC	ARIC			
			Waist girth to nearest cm			1.29e-12	1.40(0.20)	13284	ARIC			
			Waist circumference cm			1.91e-8	1.03(0.18)		WHI			
IGF2BP2	RS4402960	G (0.7)	LN1 circulating glucose value in mg/dl	Plasma Serum Glucose	EA	7.38e-3	-8.10e- 3(3.02e-3)	9339	ARIC	2	Type 2 Diabetes	17463249, 17463248, 17463246, 19401414
			Baseline glucose (mg/dl)			4.80e-3	-2.07(0.73)	4468	CHS			
			LN1 baseline glucose (mg/dl)	(lþ)		2.28e-3	-0.01(4.84e-3) 4468	4468	CHS			
TCF7L2	RS7903146	C (0.7)	LN1 circulating glucose (mg/dl)	Plasma Serum Glucose	EA	8.39e-3	-0.02(8.95e-3) 918	918	WHI	2	Type 2 Diabetes	19734900, 17668382, 17463246, 17463248, 18372903, 17293876, 17460697, 19056611, 19401414
			LN1 circulating glucose value in mg/dl	ne		2.09e-7	-0.02(2.99e-3) 10739	10739	ARIC			
			Circulating glucose value in mg/dl	C		1.73e-5	-2.04(0.48)	10739	ARIC			

phenotype-classes closely related to phenotypes among known associations. Significance was defined as a test of association with p<0.01 observed in two or more PAGE studies for the same SNP, phenotype-class, and race/ethnicity and consistent direction of effect when relevant. For each, the nearest gene(s), the SNP is number, coded allele (CA) and frequency (CAF), associated phenotype-s, phenotype-class, race/ethnicity, p-values, genetic effect/beta values (standard error; SE), sample sizes, substudies, number of substudies with results passing our p-value cutoff, the previously associated phenotype for that SNP, and references for the previously associated phenotypes are given. Related associations that met the criteria for PheWAS significance are given here, sorted by the number of PAGE site replications for a given phenotype-class. Related associations were defined as SNPs significantly associated in this PheWAS with

<sup>9</sup>studies with the significant result, in order of the associated phenotypes. <sup>11</sup>Total number of studies with at least one result passing p-value threshold for specific phenotype-class and SNP.

Coded Allele.

<sup>&</sup>lt;sup>2</sup>Coded allele frequency.

Associated phenotypes for individual results.

<sup>&</sup>lt;sup>4</sup>Phenotype-class.

Race/ethnicity for association, abbreviations: African American (AA), European American (EA), Mexican American/Hispanic (H).

 $<sup>^{6}</sup>$ P-Values of results that passed p = 0.01 threshold in order of the associated phenotypes.

Beta and standard error in order of the associated phenotypes. <sup>8</sup>Sample size in order of the associated phenotypes.

<sup>&</sup>lt;sup>11</sup>Previously reported associated phenotypes for SNP.

<sup>&</sup>lt;sup>12</sup>Pubmed ÍD's for previously associated phenotypes. doi:10.1371/journal.pgen.1003087.t003

 Table 4. PheWAS Tests of Association: Novel Associations.

County   C													
PSRC    RS59983   A (0.3)   White blood cell (keel/ml)   White blood cell (keel/ml)   Count   Count	Nearest Gene	SNP ID	CA¹ (CAF²)	Phenotypes <sup>8</sup>	Associated Phenotype Class <sup>3</sup>	Ethnicity <sup>5</sup>	P- Values <sup>6</sup>	Beta (SE) <sup>7</sup>	Sample Size <sup>9</sup>	Substudies <sup>10</sup>	Substudy Count <sup>11</sup>	Previously Associated Phenotype <sup>4</sup>	References <sup>12</sup>
My white blood count (x1000)cubic   5.62e4   0.05(0.01)   777   CHS	CELSR2,PSRC1		A (0.3)	White blood cell (kcell/ml)	White Blood Count	AA	2.14e-3	0.88(0.29)	4147	WHI	es es	Coronary Artery Disease, LDL cholesterol, Total cholesterol	18262040, 18193043, 18179892, 17634449, 18193044
LN white blood count (x,1,000/cubic   56264   0.05(0.01)   777   CH5				White blood count			3.69e-7	0.29(0.06)	3049	ARIC			
LNI white blood cell (kcell/ml)   1,28e-4   0,03(6,51e-3) 3,049   ARIC				LN1 white blood count (×1,000/cubic mm)	O.		5.62e-4	0.05(0.01)	777	CHS			
LN1 white blood count   Count   White Blood   AA   9.59e-4   0.03(8.51e-3)   3049   ARC				LN1 white blood cell (kcell/ml)			9.99e-15	0.05(6.03e-3)	4147	WHI			
12   RS10923931 G (0.77)   LN1 white blood cell count   Count   Count   Count   Count   Count   Count   RS2228145   G (0.77)   LN1 white blood cell count   Count   Count   Count   RS2228145   G (0.78)   LN1 white blood count   Count   Count   Count   Count   RS2228145   G (0.78)   G				LN1 white blood count			1.28e-8	0.04(7.65e-3)	3049	ARIC			
White blood count   White blood cell count   Cou	NOTCH2	RS10923931	G (0.7)	LN1 white blood cell count	White Blood Count	AA	9.59e-4	0.03(8.51e-3)	2083	EAGLEIII		Type 2 Diabetes, Type I Diabetes	18372903
White blood cell count   White blood cell count   Lise+   0.18(0.05)   3051   ARIC				LN1 white blood cell count			9.59e-4	0.03(8.51e-3)	2083	EAGLEIII			
White blood cell count   White blood cell count   LIN white blood cell count   Count   LIN white blood cell count   LIN white blood cell count   Cou				White blood count			8.74e-4	0.18(0.05)	3051	ARIC			
LN1 white blood cell (kcell/ml)				White blood cell count			2.75e-4	0.25(0.07)	2083	EAGLEIII			
White blood cell count   LIX white blood cell count				LN1 white blood cell (kcell/ml)			1.36e-10	0.04(6.14e-3)	4215	WHI			
R52228145 A (0.9)   White blood count				White blood cell count			2.75e-4	0.25(0.07)	2083	EAGLEIII			
R52228145 A (0.9)   White blood count Co				LN1 white blood count			1.25e-3	0.02(7.14e-3)	3051	ARIC			
White blood count (×1,000/cubic mm)   White blood count (×1,000/cubic mm)   S.61e-4   -0.06(0.02)   783   CHS	IL6R	RS2228145	A (0.9)	White blood count	White Blood Count	AA	4.17e-8	-0.36(0.06)	3806	ARIC	3	C-reactive Protein	20186139
LN1 white blood count (×1,000/cubic mm)   White blood cell count   X 1,000/cubic mm   2.83e4   -0.35(0.10)   2.038   EAGLEIII     LN1 white blood cell count   X 1,000/cubic mt   2.83e4   -0.04(0.01)   2.038   EAGLEIII     LN1 white blood cell count   X 1,57e4   -0.04(0.01)   2.038   EAGLEIII     LN1 white blood cell count   X 1,57e4   -0.04(0.01)   2.038   EAGLEIII     LN1 white blood cell count   X 1,57e4   -0.04(0.01)   2.038   EAGLEIII     Smoked at least 100 cigarettes ever (Y/N)   Ever Smoked   EA   7.57e-3   0.08(0.03)   13222				White blood cell count			2.83e-4	-0.35(0.10)	2038	EAGLEIII			
White blood cell count         2.83e-4         -0.35(0.10)         2038         EAGLEIII           LN1 white blood count         8.42e-10         -0.05(8.70e-3)         3806         ARIC           LN1 white blood cell count         3.57e-4         -0.04(0.01)         2038         EAGLEIII           RS2815752         T (0.08)         Ever smoked cigarettes ever (V/N)         Ever Smoked         EA         7.57e-3         0.08(0.03)         9339         ARIC,WHI         2         Body Mass Index           N)         N)         N)         N         7.48e-4         0.09(0.03)         13222         RIC,WHI         2         Body Mass Index				LN1 white blood count (×1,000/cubic mm)	(T		5.61e-4	-0.06(0.02)	783	CHS			
LN1 white blood count   LN1 white blood cell count   S.57e-4   -0.04(0.01)   2038   EAGLEIII   S.57e-4   S.57e-3   S.57e-3   S.57e-4   S.57e-3   S.57e-3   S.57e-4   S.57e-3   S.57e-3				White blood cell count			2.83e-4	-0.35(0.10)	2038	EAGLEIII			
LN1 white blood cell count   3.57e-4   -0.04(0.01)   2038   EAGLEIII				LN1 white blood count			8.42e-10	-0.05(8.70e-3)		ARIC			
LN1 white blood cell count   3.57e-4   -0.04(0.01)   2038   EAGLEIII				LN1 white blood cell count			3.57e-4	-0.04(0.01)	2038	EAGLEIII			
RS2815752 T (0.08) Ever smoked cigarettes (Y/N) Ever Smoked EA 7.57e-3 0.08(0.03) 9339 ARIC,WHI 2 Body Mass Index Smoked at least 100 cigarettes ever (Y/ 7.48e-4 0.09(0.03) 13222 N)				LN1 white blood cell count			3.57e-4	-0.04(0.01)	2038	EAGLEIII			
ever (Y/ 7.48e-4 0.09(0.03)	NEGR1		T (0.08)	Ever smoked cigarettes (Y/N)	Ever Smoked	EA	7.57e-3	0.08(0.03)	9339	ARIC,WHI	2	Body Mass Index	19079261
				Smoked at least 100 cigarettes ever (Y N)	<i>h</i>			0.09(0.03)	13222				

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References <sup>12</sup>	15820616, 1654245, 16644865, 17003362, 17416766, 17416766, 17916900, 17916900, 17916900, 17916900, 17916900, 17916900, 18239642, 18239642, 18252935, 19006106, 19106168, 19106168, 19106168, 19106168, 19106168, 19249263, 19339011, 19339016, 19339016, 19452524, 19542000, 19572000, 19572000, 19573000, 19573000, 1957300,		
Refe			
Previously Associated Phenotype <sup>4</sup>	C-reactive Protein		
Substudy Count <sup>11</sup>	7		
Substudies <sup>10</sup>	ARIC	EAGLEIII	ARIC
Sample Size <sup>9</sup>	3071	1964	3071
Beta (SE) <sup>7</sup>	0.12(0.04)	0.24(0.09)	0.26(0.09)
P- Values <sup>6</sup>	5.43e-3	9.44e-3	2.74e-3
Ethnicity <sup>5</sup>	<b>\{</b>		
Associated Phenotype Class <sup>3</sup>	Carotene	2	
Phenotypes <sup>8</sup>	Consumption)	LN1 carotenes (dietary consumption)	LN1 alpha carotene mcg (dietary consumption)
CA <sup>1</sup> (CAF <sup>2</sup> )	C (0.06)		
OI ANS	RS1800795		
Nearest Gene	ادو ادو		

Table 4. Cont.

Nearest Gene	SNP ID	CA¹ (CAF²)	Phenotypes <sup>8</sup>	Associated Phenotype Class <sup>3</sup>	Ethnicity <sup>5</sup>	P- Values <sup>6</sup>	Beta (SE) <sup>7</sup>	Sample Size <sup>9</sup>	Substudies <sup>10</sup>	Substudy Count <sup>11</sup>	Previously Associated Phenotype <sup>4</sup>	References <sup>12</sup>
IL6R	RS2228145	A (0.9)	LN1 lymphocytes	Lymphocytes	AA	5.12e-10	0.06(0.01)	3769	ARIC	2	C-reactive Protein	20186139
			Lymphocytes (percent of 100 cells)			5.39e-5	2.96(0.73)	949	EAGLEIII			
			Lymphocytes			6.63e-10	2.35(0.38)	3769	ARIC			
			LN1 lymphocytes (percent of 100 cells)			3.88e-5	0.08(0.02)	949	EAGLEIII			
IL6R	RS2228145	A (0.9)	LN1 neutrophils	Neutrophils	AA	1.54e-8	-0.06(9.81e-3)	3769	ARIC	2	C-reactive Protein	20186139
			LN1 Segmented neutrophils			2.44e-4	-0.06(0.02)	949	EAGLEIII			
			Neutrophils			4.66e-10	-2.61(0.42)	3769	ARIC			
			Segmented neutrophils			7.74e-5	-3.12(0.79)	949	EAGLEIII			
CELSR2,PSRC1	RS599839	A (0.8)	Currently of on a special diet? (Y/N)	Dieting	EA	9.48e-3	0.12(0.05)	9328	ARIC	2	Coronary Artery Disease, LDL cholesterol, Total cholesterol	18262040, 18193043, 18179892, 17634449, 18193044
			Low-fat or low cholesterol diet (Y/N)			5.87e-3	0.08(0.03)	13068	WHI			
GALNT2	RS2144300 C (0.8)	C (0.8)	LN1 FEV(3) over FEV(6) (Lung function)	FEV3	AA	4.90e-4	-0.11(0.03)	3090	ARIC	2	Coronary Heart Disease, HDL cholesterol, Triglycerides	18193043
			FEV(3) (liters) (Lung function)			1.13e-3	-0.10(0.03)	3090	ARIC			
			LN1 FEV3 at 3.0 seconds Largest value (Lung Function)			8.82e-3	-0.03(0.01)	1953	ARIC			
			FEV(3) over FEV(6) (Lung Function)			7.93e-4	-2.26(0.67)	3090	EAGLEIII			
			LN1 FEV(3) (liters) (Lung Function)			8.43e-4	-0.03(7.92e-3)	3090	ARIC			
GALNT2	RS2144300	C (0.8)	Pills for hypertension ever (Y/N)	Hypertension	AA	8.27e-3	0.15(0.06)	4180	WHI	2	Coronary Heart Disease, HDL cholesterol, Triglycerides	18193043
			High blood pressure ever diagnosed? (Y/N)			1.61e-3	0.24(0.08)	3142	ARIC			
GALNT2	RS2144300 C (0.4)	C (0.4)	LN1 serum total calcium: (mmol/l)	Serum Calcium	EA	8.10e-3	– 2.36e-3(8.92e-4) 2587	t) 2587	EAGLEIII	2	Coronary Heart Disease, HDL cholesterol, Triglycerides	18193043
			LN1 serum calcium (mg-dl)			1.47e-3	-1.84e-3(5.80e-4) 9098	8606 (t	ARIC			
			Serum total calcium			5.95e-3	0.03(0.01)	2585	EAGLEIII			

Table 4. Cont.

				Associated							Previously	
Nearest Gene	SNP ID	CA¹ (CAF²)	Phenotypes <sup>8</sup>	Phenotype Class <sup>3</sup>	Ethnicity <sup>5</sup>	P- Values <sup>6</sup>	Beta (SE) <sup>7</sup>	Sample Size <sup>9</sup>	Substudies 10	Substudy Count <sup>11</sup>	Associated Phenotype <sup>4</sup>	References <sup>12</sup>
			Serum total calcium (mmol/l)			7.47e-3	-7.97e-3(2.98e-3) 2587	3) 2587	EAGLEIII			
			LN1 serum total calcium (mmol/l)			6.28e-3	-2.39e-3(8.75e-4) 2585	t) 2585	EAGLEIII			
			LN1 serum total calcium			6.40e-3	-3.09e-3(1.13e-3) 2585	3) 2585	EAGLEIII			
			Serum calcium (mg-dl)			1.71e-3	-0.02(6.24e-3)	8606	ARIC			
			Serum total calcium (mmol/l)			5.95e-3	-7.98e-3(2.90e-3) 2585	3) 2585	EAGLEIII			
GALNT2	RS2144300	C (0.8)	LN1 white blood cell (kcell/ml)	White Blood Count	AA	3.32e-6	-0.04(7.53e-3)	4210	WHI	2	Coronary Heart Disease, HDL cholesterol, Triglycerides	18193043
			LN1 white blood count (×1,000/cubic mm)			7.96e-3	-0.05(0.02)	785	CHS			
GALNT2	RS2144300	C (0.4)	Coronary artery bypass graft (cabg) (Y/N)	Artery Treatment EA	t EA	2.46e-3	0.24(0.08)	13152	WHI	2	Coronary Heart Disease, HDL cholesterol, Triglycerides	18193043
			Aortic Aneurysm repair (Y/N)			5.49e-3	0.57(0.20)	4250	CHS			
LIPG	RS2156552	A (0.2)	Doctor ever told you had goiter? (Y/N)	Thyroid Goiter	EA	8.60e-3	-1.91(0.25)	2273	EAGLEIII	2	HDL Cholesterol	18193043
			Age told had goiter/thyroid disease			7.57e-3	-2.97(1.11)	763	ARIC			
CETP	RS3764261 T (0.3)	Т (0.3)	Age at first period category: 11 years	Menstruation	EA	4.30e-3	-0.11(0.04)	13276	WHI	2	HDL Cholesterol, LDL cholesterol, Waist circumference	18193043, 19359809
			Age when menstruation began			2.04e-3	0.11(0.03)	5705	WHI			
			LN1 age when menstruation began			5.43e-4	9.19e-3(2.66e-3)	5705	ARIC			
			Age at first period category: 10 years			6.81e-3	-0.17(0.06)	13276	ARIC			
FADS1, FADS2, FADS3	RS174547	C (0.3)	Platelet count	Platelet Count	EA	7.37e-3	3.26(1.22)	9174	ARIC	2	HDL Cholesterol, Triglycerides, Total cholesterol, LDL cholesterol	20037589, 19060906
			LN1 platelet count (kcell/ml)			2.28e-3	0.01(3.30e-3)	13140	WHI			
			LN1 platelet count			1.68e-3	0.01(3.90e-3)	9174	ARIC			
LDLR	RS6511720	(6.0) D	Max % arterial stenosis (1–24%) (Arterial measurement)	Artery	EA	4.19e-3	-0.22(0.08)	4460	CHS	2	LDL Cholesterol	18193043
			Arterial plaque in any site (Y/N)			9.19e-3	0.13(0.05)	8269	ARIC			
			Max % arterial stenosis (25–49%) (Arterial measurement)			4.91e-3	0.21(0.07)	4460	CHS			
PCSK9	RS11591147 G (0.996)	G (0.996)	Coronary artery bypass graft (Y/N)	Artery Treatment AA	t AA	2.11e-4	-2.23(0.60)	4274	WHI	2	LDL Cholesterol, lipids	18193044

Table 4. Cont.

		5	,	Associated Phenotype	•	4	Beta	Sample	;	Substudy		;
Nearest Gene	SNP ID	(CAF²)	Phenotypes <sup>8</sup>	Class³	Ethnicity	Values	(SE)′	Size	Substudies 10	Count	Phenotype <sup>4</sup>	References ' 2
			Heart or arterial surgery? (Y/N)			1.53e-3	-1.97(0.62)	3909	ARIC			
APOC4, APOC2 APOC4, APOC2	RS4420638	A (0.8)	Circulating glucose value in mg/dl	Plasma Serum Glucose	EA	8.86e-4	1.89(0.57)	9336	ARIC	2	LDL Cholesterol, 17474819, lipids, Alzheimer's 17463246, disease, Coronary 18193043, Artery Disease, C- 18802019, reactive protein, 17998437, Sporadic late onset 19197348, Alzheimer's disease, 19567438 ApoB, Triglycerides,	17474819, 17463246, 18193043, 18802019, 17998437, 19197348,
			Baseline glucose (mg/dl)			4.65e-3	2.67(0.94)	4470	CHS			
			LN1 circulating baseline glucose (mg/dl)			5.67e-3	0.02(6.23e-3)	4470	CHS			
			LN1 circulating glucose value in mg/dl			6.81e-4	0.01(3.62e-3)	9336	ARIC			
CELSR2, PSRC1,SORT1	RS646776	G (imputed)	Are you currently on a special diet (Y/N)	Dieting	EA	4.20e-3	-0.14(0.05)	9331	ARIC	2	LDL Cholesterol, Myocardial infarction (early onset)	18193044, 18262040
			Low-fat or low cholesterol diet (Y/N)			8.78e-3	-0.08(0.03)	13051	MHI			
APOB	RS562338	Т (0.6)	Are you following a special diet (Y/N)	Dieting	AA	4.82e-3	-0.34(0.12)	817	CHS	2	LDL Cholesterol, Total cholesterol, Type 2 Diabetes	18193043
			Currently of diet (Y/N)			5.07e-3	-0.19(0.07)	3149	ARIC			
CDKN2A,CDKN2B RS10757278	RS10757278	A (0.8)	LN1 white blood count	White Blood Count	AA	5.05e-4	-0.03(7.32e-3)	3877	ARIC	2	Myocardial infarction	17478679
			White blood count			3.24e-3	-0.16(0.05)	3877	ARIC			
			LN1 white blood cell (kcell/ml)			3.80e-3	-0.02(6.96e-3)	4214	WHI			
CDKN2A,CDKN2B RS2383207	RS2383207	A (imputed)	A (imputed) LN1 dietary vitamin b12 (mcg)	VitaminB12	EA	4.34e-3	0.02(5.69e-3)	13254	WHI	2	Myocardial infarction	17478679
			Dietary vitamin b12 (mcg)			3.35e-3	0.14(0.05)	13254	WHI			
			LN1 vitamin b12 (micrograms)			3.95e-3	0.02(6.98e-3)	9197	ARIC			
ANGPTL3	RS1748195	G (0.6)	LN1 hemoglobin (g/dl)	Hemoglobin	AA	7.66e-3	-9.61e-3(3.60e-3) 2085	2085	EAGLEIII	2	Triglycerides, lipids 18193043	18193043
			LN1 hemoglobin (gm/dl)			9.57e-3	-4.64e-3(1.79e-3) 4195	4195	WHI			
			Hemoglobin (g/dl)			6.28e-3	-0.14(0.05)	2085	EAGLEIII			
ADAMTS9	RS4607103	C (0.8)	LN1 min lumen diameter (arterial measurement)	Artery	EA	7.00e-3	0.07(0.03)	120	ARIC	2	Type 2 Diabetes	18372903
			LN1 average near and far wall max common carotid artery (mm) (Arterial measurement)			6.22e-3	6.32e-3(2.31e-3)	4460	CHS			

Table 4. Cont.

Nearest Gene	SNP ID	CA¹ (CAF²)	Phenotypes <sup>8</sup>	Associated Phenotype Class <sup>3</sup>	Ethnicity <sup>5</sup>	P- Values <sup>6</sup>	Beta (SE) <sup>7</sup>	Sample Size <sup>9</sup>	Substudies <sup>10</sup>	Substudy Count <sup>11</sup>	Previously Associated Phenotype <sup>4</sup>	References <sup>12</sup>
			Ave near and far wall max common carotid artery (mm) (Arterial measurement)			7.91e-3	0.01(5.02e-3)	4460	CHS			
ADAMTS9	RS4607103	C (0.8)	Smoked at least 100 cigarettes ever? (Y/N)	Ever Smoked	EA	1.38e-3	-0.09(0.03)	13214	WHI	2	Type 2 Diabetes	18372903
			Ever smoked? (Y/N)			4.11e-3	-0.14(0.05)	4477	CHS			
IGF28P2	RS4402960	G (0.7)	Currently of diet? (Y/N)	Dieting	EA	3.14e-4	-0.15(0.04)	9331	ARIC	2	Type 2 Diabetes	17463249, 17463248, 17463246, 19401414
			Diabetic or ADA diet? (Y/N)			9.63e-4	-0.23(0.07)	12958	WHI			
TCF71.2	RS7901695	C (0.5)	LN1 white blood count, white blood count	White Blood Count	AA	2.75e-3	-0.02(6.77e-3)	3053	ARIC	2	Type 2 Diabetes	17554300, 17463249, 17463246, 17668382
			LN1 white blood cell (kcell/ml)			7.03e-3	-0.14(0.05)	3053	ARIC			
						7.73e-3	-0.02(5.76e-3)	4182	MHI			
TSPAN8/LGR5	RS7961581	C (0.3)	Pulse rate (beats/min) (age 5+ years)	HeartRate	EA	8.38e-3	-1.33(0.50)	2570	EAGLEIII	2	Type 2 Diabetes	18372903
			LN1 heart rate per minute			8.17e-3	-6.20e-3 (2.34e-3)	9314	ARIC			
NOTCH2	RS10923931 G (0.7)	G (0.7)	Hypertension ever (Y/N)	Hypertension	AA	3.07e-3	-0.14(0.05)	4206	WHI	2	Type 2 Diabetes, Type I Diabetes	18372903
			History of high blood pressure from baseline questionnaire (Y/N)			2.00e-3	-0.14(0.04)	4736	MEC			
			History of high blood pressure (Y/N)			3.50e-3	-0.14(0.05)	4362	MEC,			
			Hypertension interested category: never Hypertensive (Y/N)			2.33e-3	-0.15(0.05)	4076	MH			
			Hypertension interested category: treated Hypertensive (Y/N)			2.92e-3	-0.14(0.05)	4076	IH <sub>W</sub>			
			Pills for hypertension ever (Y/N)			8.18e-4	-0.16(0.05)	4186	WHI			
			Pills for hypertension now (Y/N)			2.82e-3	-0.14(0.05)	4122	WHI			

Table 4. Cont.

Nearest Gene	SNP ID	CA¹ (CAF²)	Phenotypes <sup>8</sup>	Associated Phenotype Class <sup>3</sup>	P- Ethnicity <sup>5</sup> Values <sup>6</sup>	P- Values <sup>6</sup>	Beta (SE) <sup>7</sup>	Sample Size <sup>9</sup>	Substudies <sup>10</sup>	Substudy Count <sup>11</sup>	Previously Associated Phenotype <sup>4</sup>	References <sup>12</sup>
FTO	RS8050136	A (0.4)	Age at first period interested category: 14 years	Menstruation	EA	5.26e-3	-0.10(0.04)	13275	WHI	2	Obesity, Type 2 diabetes	17554300, 19079260, 17463249, 17463248, 18159244, 19056611, 17658951
			Age when menstruation began			1.18e-4	-0.14(0.04)	4917	ARIC			
			LN1 age when Menstruation began			6.09e-5	-0.01(2.74e-3)	4917	ARIC			
CELSR2, PSRC1, RS646776 SORT1	RS646776	G (imputed)	G (imputed) Hospitalized for chest pain? (Y/N)	ANGINA	EA	3.05e-3	-0.55(0.18)	568	ARIC	2	LDL Cholesterol	19060911, 19198609, 18193044, 18262040, 19060910
			Angina (Y/N)			3.17e-4	-0.25(0.07)	13289	WHI			
CDKN2A, CDKN2B	RS1333049 C (0.2)	C (0.2)	Hemoglobin (g/dl)	Hemoglobin	<b>AA</b>	5.40e-3	0.15(0.05)	2092	EAGLEIII	2	Coronary Artery Disease, Type 2 Diabetes, Hypertension	17554300, 17634449
			Hemoglobin (g/dl)			8.21e-3	0.26(0.10)	786	CHS			
			LN1 Hemoglobin (g/dl)			4.07e-3	0.01(3.95e-3)	2092	EAGLEIII			

Novel associations that met the criteria for PheWAS significance are given here, sorted by the most to least number of PAGE study sites available. Related associations were defined as SNPs significantly associated in this PheWAS with phenotype-classes closely related to phenotypes among known associations. Significance was defined as a test of association with p<0.01 observed in two or more PAGE studies for the same SNP, phenotype class, and race/ ethnicity, p-values, genetic effect/ ethnicity, and consistent direction of effect when relevant. For each, the nearest gene(s), the SNP rs number, coded allele (CA) and frequency (CAF), associated phenotypes, phenotype-class, race/ethnicity, p-values, genetic effect/ beta values (standard error; SE), sample sizes, substudies, number of substudies with results passing our p-value cutoff, the previously associated phenotype for that SNP, and references for the previously associated phenotypes are

Coded Allele.

<sup>&</sup>lt;sup>2</sup>Coded allele frequency.

<sup>&</sup>lt;sup>3</sup>Associated phenotypes. <sup>4</sup>Phenotype-class.

<sup>\*</sup>Race/ethnicity for association, abbreviations: African American (AA), European American (EA), Mexican American/Hispanic (H). \*P-Values of results that passed p = 0.01 threshold in order of the associated phenotypes.

Beta and standard error in order of the associated phenotypes.

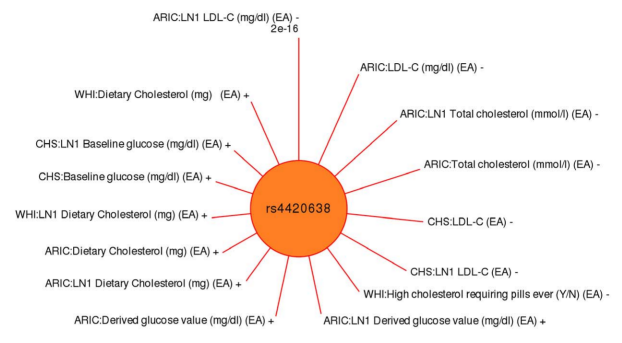
<sup>&</sup>lt;sup>3</sup>Sample size in order of the associated phenotypes

<sup>&</sup>lt;sup>9</sup>Studies with the significant result, in order of the associated phenotypes.

<sup>&</sup>lt;sup>10</sup>Total number of studies with at least one result passing p-value threshold for specific phenotype class and SNP.

<sup>&</sup>lt;sup>12</sup>Pubmed ID's for previously associated phenotypes. doi:10.1371/journal.pgen.1003087.t004

# APOE/APOC1/C1P1/C2/C4, rs4420638, Coded Allele A



**Figure 1. PheWAS associations for rs4420638 near** *APOC1.* SNP rs4420638 has previously been associated with LDL cholesterol levels, triglycerides, Alzheimer's disease, coronary artery disease, and sporadic late onset Alzheimer's. The length of the lines correspond to −log10(p-value), and the lines are plotted clockwise starting at top for the association with the smallest p-value. Lines are labeled with the study-specific phenotype, the PAGE study, racial/ethnic group, and direction of effect (+ or −). Red lines represent associations at p<0.01. "LN1" indicates the phenotype had 1 added to the variable, and then the variable was natural log transformed. The PheWAS phenotypes significantly associated with this SNP varied, with known associations for LDL cholesterol levels, as well as the related phenotypes "Total cholesterol (mmol/l)" and "Dietary cholesterol (mg)", and novel phenotypes such as "Baseline glucose (mg/dl)". doi:10.1371/journal.pgen.1003087.g001

[8,12-14]; type 2 diabetes (rs10923931) [16]; C-reactive protein (rs2228145) [17]; coronary heart disease, HDL-C and triglycerides (rs2144300) [13]; MI (rs10757278) [11]; and type 2 diabetes (rs7901695) in EA [18-20]. It is likely that the majority of the significant findings for three of the SNPs on chromosome 1 [rs599839 (CELSR2/PSRC1), rs10923931 (NOTCH2), rs2228145 (IL6R)] are not truly novel given that these variants are likely in linkage disequilibrium with the white blood cell count-associated Duffy null allele (DARC rs2814778) [21,22] in African Americans. Of note is *GALNT2* rs2144300 (p =  $3.32 \times 10^{-6}$  in WHI and 7.96×10<sup>-3</sup> in CHS), located outside the 90 Mb region known to be associated with white blood cell counts in African Americans [21] and possibly representing a novel genotype-phenotype association for this trait. Also for chromosome 1, novel associations were identified in African Americans at p<0.01 for the phenotypeclass "Hemoglobin" and ANGPTL3 rs1748195, previously associated with triglycerides in European-descent populations [13,19].

Of the remaining hematologic trait associations identified that were not on chromosome 1, rs10757278 near *CDKN2A/B* on chromosome 9 and *TCF7L2* rs7901695 on chromosome 10 were both associated with white blood cell count, neither of which were previously reported in GWAS for this trait [21,22]. For *CDKN2A/B* rs1333049, a SNP previously associated with type 2 diabetes, coronary artery disease, and hypertension in European-descent populations [15,23] p<0.01 associations were identified for the phenotype-class of Hemoglobin. Finally, a novel association in European Americans was noted between *FADS1* rs174547, a SNP previously associated with LDL-C [13,19], and the phenotype-class of "Platelet Count" at p<0.01.

Aside from hematologic traits, the most significant novel association identified in this PheWAS was identified for phenotypes in the phenotype-class "Forced Expiratory Volume in 3 Seconds (FEV3)" and GALNT2 rs2144300 in African Americans (p-values ranging from  $8.82 \times 10^{-3}$  to  $4.90 \times 10^{-4}$ ). GALNT2 rs2144300, previously associated with HDL-C in European Americans and African Americans [13,24], has not previously been associated with lung function or asthma quantitative traits. Interestingly, GALNT2 rs2144300 was also associated with phenotypes in the "Hypertension" phenotype-class among African Americans in this PheWAS Specifically the phenotypes were "High blood pressure ever diagnosed?" (ARIC,  $p = 1.61 \times 10^{-3}$ ,  $\beta = 0.24$ ) and "Pills for hypertension ever?" (WHI,  $8.27 \times 10^{-3}$ ,  $\beta = 0.15$ ). Indeed, *GALNT2* rs2144300 displayed the most suggestion of pleiotropy among all the SNPs tested in this study. In addition to the associations identified in African Americans, rs2144300 was associated with phenotypes in the phenotype-classes "Serum Calcium" (p-values ranging from  $1.47 \times 10^{-4}$  to  $8.10 \times 10^{-3}$ ) and "Artery Treatment", specifically the phenotypes "Coronary artery bypass graft (CABG)" (WHI,  $p = 2.46 \times 10^{-3}$ ,  $\beta = 0.24$ ) and "Aortic aneurysm repair" (CHS,  $5.49 \times 10^{-3}$ ,  $\beta = 0.57$ ) in European Americans. Significant PheWAS associations at p<0.01 for rs2144300 are plotted by phenotype in Figure 5, as well as additional results at p < 0.05.

The remaining significant novel PheWAS results have identified potentially pleiotropic effects for SNPs previously associated with lipid traits, type 2 diabetes, inflammation, myocardial infarction, and body mass index. The lipid trait-associated SNPs were associated with the "Menstruation" phenotype-class (specifically age at menarche) in European Americans (CETP rs3764261), the

## CDKN2A/CDKN2B, rs10757278, Coded Allele A

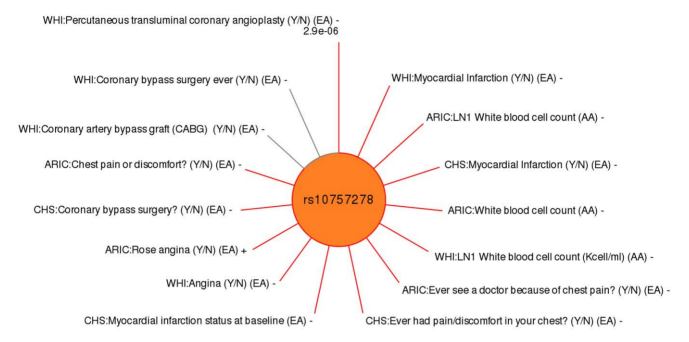


Figure 2. PheWAS associations for rs10757278 near *CDKN2A/CDKN2B*. SNP rs10757278 was previously associated with myocardial infarction (MI). Associations are plotted clockwise starting at top for the association with the smallest p-value and the length of the line corresponds to  $-\log 10(p-value)$ . Lines are labeled with the study-specific phenotype, the PAGE study, racial/ethnic group, and direction of effect (+ or -). Red lines represent associations at p<0.01, and results with p<0.05 are also plotted in grey to show trends for additional phenotypes. "LN1" indicates the phenotype had 1 added to the variable, and then the variable was natural log transformed. The PheWAS phenotypes significantly associated with this SNP varied, from MI (known), to coronary artery disease and MI related phenotypes such as presence or absence of "percutaneous transluminal coronary angioplasty", "angina", and "coronary bypass surgery". doi:10.1371/journal.pgen.1003087.g002

"Dieting" phenotype-class (APOB rs562338 in African Americans and CELSR2/PSRC1/SORT1 rs599839 and rs646776 in European Americans), "Thyroid Goiter" in European Americans (LIPG rs2156552), "Artery Measurements" in European Americans (LDLR rs6511720) and "Artery Treatment" in African Americans (PCSK9 rs11591147), "Plasma Serum Glucose Levels" (APOE/APOC1/ APOC4/APOC2/APOC3 rs4420638) in European Americans, and the "Angina" phenotype-class in European Americans (CELSR2/ PSRC1/SORT1 rs646776). For the type 2 diabetes-associated SNPs, the PheWAS-identified associations were observed for the phenotype-classes of "Dieting" (IGFBP2 rs4402960) in European Americans, "Artery" and "Ever Smoked" (ADAMTS9 rs4607103) in European Americans, "Hypertension" (NOTCH2 rs10923931) in African Americans, "Heart Rate" (LGR5 rs7961581) in European Americans, and "Menstruation" (specifically age at menarche) in European Americans (FTO rs8050136). Like type 2 diabetesassociated ADAMTS9 rs4607103, BMI-associated NEGR1 rs2815752 was associated with the phenotype-class of "Ever Smoked" in European Americans. The final two PheWAS-identified significant associations involved nutrient based phenotype-classes: MI-associated CDKN2A/B rs2383207 was associated with the phenotype-classes of "Vitamin B12" in European Americans, and inflammation-associated IL6 rs1800795 was associated with the phenotype-class of "Carotene" in African Americans.

#### Discussion

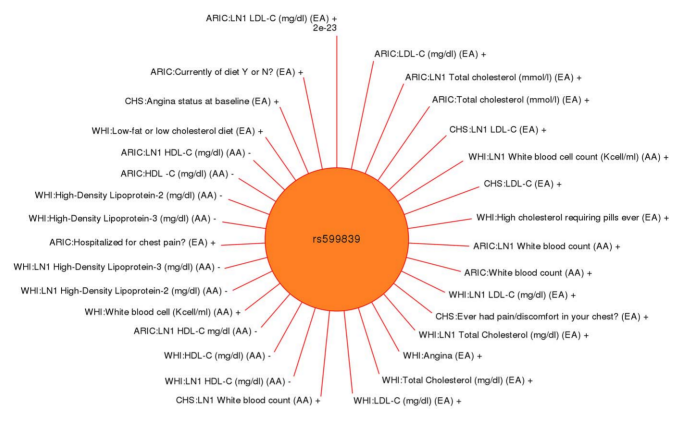
The PheWAS results herein present the result of tests of association between a large number of SNPs and an extensive

range of phenotypes and traits available within five studies of the PAGE network. For this first PAGE PheWAS analysis we have emphasized associations that replicated across two or more independent PAGE studies for the same phenotype class and same race/ethnicity. Most of the robust findings reported here represent previously known genotype-phenotype relationships, but a tantalizing few also represent potentially novel pleiotropic relationships.

The 33 novel results presented here are intriguing, but it is important to emphasize that these first-pass analyses are considered hypothesis-generating, exploratory, and require additional scrutiny before the findings are further considered for follow-up, unlike the directed a priori hypothesis-testing analyses within PAGE that involve SNPs hypothesized to be associated with specific phenotypes. Further analysis of PheWAS results will be on an individual result basis and will include careful phenotype harmonization for traits and outcomes that cross two or more PAGE studies, as well as considerable investigation of the possible effect of covariates such as age, sex, and environmental exposure(s) on the association between genetic variation and phenotypic outcome.

One of the many challenges for the interpretation of PheWAS results is dissecting the genetic effect observed among correlated phenotypes. In some cases, the relationship is likely attributable to a common biological process with known genetic contribution (e.g., body mass index and waist circumference). In other cases, the networks that exist between intermediary and/or outcome related phenotypes add complexity to interpreting association results. For instance, genetic variation may impact the variation of a single

## CELSR2/PSRC1, rs599839, Coded Allele A



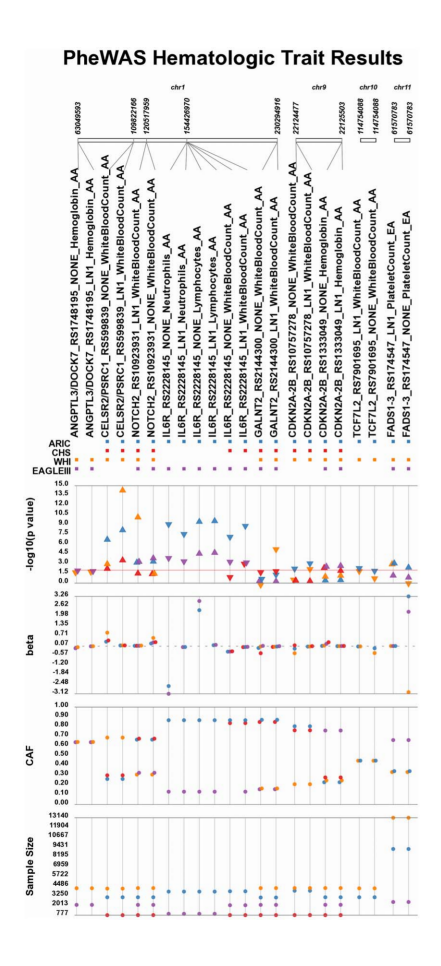
**Figure 3. PheWAS associations for rs599839 near** *CELSR2/PSRC1*. This SNP has previously published associations with serum LDL cholesterol levels, total cholesterol, and coronary artery disease. Genotype-phenotype associations are plotted clockwise starting at top for the association with the smallest p-value. The length of the line corresponds to −log10(p-value), the longer the line the more significant the result. The study race/ ethnicity/and phenotype for each tests of association are listed. Red lines represent associations at p<0.01, and results with p<0.05 are also plotted in grey to show trends for additional phenotypes. "LN1" indicates the phenotype had 1 added to the variable, and then the variable was natural log transformed. The PheWAS phenotypes significantly associated with this SNP varied, from LDL cholesterol levels (previously published), to lipid level-related phenotypes such as "High cholesterol requiring pills ever". In the case of coronary artery disease, phenotypes with significant results that were related to coronary artery disease included "Ever had pain/discomfort in your chest", and "Hospitalized for chest pain". doi:10.1371/journal.pgen.1003087.g003

phenotype, but variation in that phenotype could then result in changes in other downstream phenotypes indirectly. Examples of added complexity include obesity leading to impaired immune function [28], and metabolic syndrome where there is a spectrum of risk factors that are all associated with increased risk of cardiovascular disease and type 2 diabetes [25]. As a result, significant associations between a genetic variant and many phenotypes could represent a network or cascade of events. This is a potential interpretation of results found for SNP rs10923931 (NOTCH2) in AA, where type 2 diabetes was the previously reported association for this SNP and the novel result was found for hypertension, and type 2 diabetes and hypertension are often a co-occurrence. Further analysis of individual PheWAS results is necessary to conclusively establish the impact of the relationship between phenotypes on significant SNP-phenotype associations.

With the large number of phenotype-genotype associations calculated, there will be an increase in type 1 error due to multiple testing. A Bonferroni correction could be used within each individual study to choose a cutoff for significance that controls for multiple hypothesis testing. However, this would not take into account the correlations that exist between the phenotypes in these studies that impact the assumption of independence between tests as well as the correlations between the genotypes.

For our first PAGE PheWAS analysis, we chose to seek replication of results across studies and required the same direction of effect as one approach to reduce the false discovery rate. Significant results can still be found by chance across more than one study. Multiple challenges arise when attempting to get a metric of the type 1 error rate across multiple studies. First, as with individual studies, correlations between phenotypes and previous associations for the SNPs are still present. Also, there are varying type 1 error rates depending on the number of studies available for seeking replication. Quantification of how many results were found with a p-value cutoff, and without a p-value cutoff, depending on the number of studies where replication could be sought (2, 3, 4, or 5) provides some information about the number of significant results we found, in Table 2. Table 1 has the total number of results with and without p-value cutoff for individual studies. It is important to note that in cases where replication could be sought in more than two studies, there were cases where the result replicated in 3 or more studies, further increasing our confidence in the result.

A potential limitation of this study is the granularity of phenotypes within our phenotype classes. The phenotypes within some phenotype classes are the same or extremely similar, such as white blood cell count measurements across studies. However, the phenotype class "Artery Treatment" is broad in terms of the types



**Figure 4. PheWAS results for blood cell counts and hemoglobin levels.** Eleven novel genotype-phenotype-class associations were identified for white blood cell counts and hemoglobin levels collectively. The top track indicates the chromosomal location of each SNP, below that track is a SNP/Phenotype identification track containing the SNP ID, as well as the phenotype, phenotype transformation if present (LN1 = ln(1+variable)), and the race-ethnicity for the test population (AA or EA). The next track is a "presence/absence" track, box presence indicates if the SNP was present for ARIC (blue), CHS (red), WHI (orange), or EAGLE (purple). The next tracks are as follows: -log10(p-value), where the each p-value is plotted, the direction of the triangle indicates the direction of effect (triangle pointed up is positive, triangle pointed down is negative), base of the triangle corresponds to the location of the p-value, solid red line is positioned at p-value = 0.01; The next track is magnitude of effect (beta) dotted grey line is positioned at the null; Next are coded allele frequencies (CAF) for each study; Final track is sample size for each test of association. doi:10.1371/journal.pgen.1003087.q004

of phenotypes included, such as presence/absence aortic aneurysm repair and presence/absence of angioplasty of the coronary arteries. For some classes, the replicated results encompass more variation in the phenotypes captured, compared to other results. As a result, significant associations between a genetic variant and all phenotypes in a network may be present. PheWAS is an exploratory and hypothesis generating exercise, thus the choice was made to have a broader match for some groups of phenotypes in order to allow for those phenotypes to be part of the exploration of the data. In addition, misclassification of phenotypes when matching is possible, and thus can limit identification of significant associations across studies. Other potential limitations include sample size/power, study heterogeneity, and the SNPs selected for study. As shown in Table 1, there is much variability across independent PAGE studies. While each PAGE study is sizeable, individual tests of association may be underpowered depending on the availability of the genetic variant, phenotype class, and race/ ethnicity. Tests of association that failed to reach statistical significance may represent underpowered genotype-phenotype relationships and will require larger epidemiologic or clinic-based samples to identify. In regards to the potential impact of heterogeneity, we have some cases where replication existed in only two or three studies out of those where replication could be sought. In some instances this may be due to power, but this also may reflect the heterogeneity between studies, such as how various phenotypes are measured in individual studies and variation in mean age across the different studies. Finally, SNPs were originally selected for this study to replicate known genotype-phenotype associations and to generalize them to diverse populations. A comprehensive set of genome-wide "agnostic" SNPs may uncover additional pleiotropic or novel genotype-phentoype relationships not tested here.

Despite the the limitations present for this PheWAS, there are multiple strengths within our study. We have had the opportunity to perform a PheWAS of substantial size with an unprecedented diversity of high quality phenotypic measurements and traits, across multiple races/ethnicities. In addition, because of this PheWAS was conducted across multiple independent studies, we were able to identify the most robust genotype-phenotype relationships across studies

### GALNT2, rs2144300, Coded Allele C

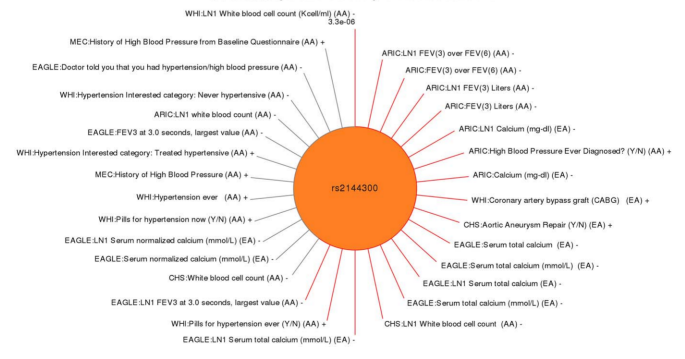
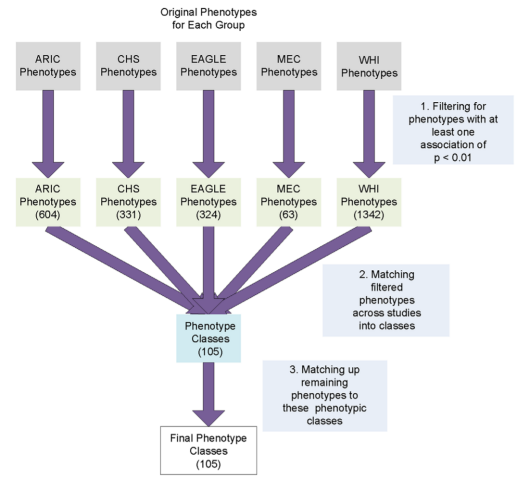


Figure 5. PheWAS associations for rs2144300 within *GALNT2*. The previously published associations for this SNP were with triglyceride and HDL cholesterol levels. Genotype-phenotype associations are plotted clockwise starting at top for the association with the smallest p-value. The length of the line corresponds to  $-\log 10$ (p-value), the longer the line the more significant the result. The study race/ethnicity/and phenotype for each tests of association are listed. Red lines represent associations at p<0.01, and results with p<0.05 are also plotted in grey to show trends for additional phenotypes. The novel PheWAS phenotypes significantly associated with this SNP varied, including white blood cell counts, forced vital capacity at three seconds (FEV3), and serum calcium levels. doi:10.1371/journal.pgen.1003087.g005



**Figure 6. Workflow for phenotype matching, to develop the 105 phenotype classes.** A MySQL database was used to filter the data from five studies for any results with p<0.01 to generate lists of the unique phenotypes for each individual PAGE study. The number of phenotypes that passed this significance threshold for each of the four groups was 604 (ARIC), 331 (CHS), 63 (MEC), 324 (EAGLE), 1,342 (WHI). Note that during the binning process, a smaller number of phenotypes are listed in Figure 6 than the total number of phenotypes referred to in the manuscript for the actual associations, in the phenotype matching process we only took into account distinct phenotypes regardless of whether or not they were transformed or untransformed or if they were categorical phenotypes binned into case/control phenotypes. Next, resulting phenotypes were then manually matched up between ARIC, CHS, MEC, EAGLE and WHI using and knowledge about the phenotypes and the known focus of specific PAGE study questions (such as arterial measurements including degree of arterial stenosis). In the last step, phenotypes from all studies, regardless of significance from genotype-phenotype tests of association, were matched to the already-defined phenotype classes using the criteria described above.

doi:10.1371/journal.pgen.1003087.g006

#### Conclusion

This initial PheWAS within PAGE has presented challenges in terms of generating high-throughput tests of association across large epidemiologic studies as well as the synthesis of the resulting data and its eventual interpretation. Even with these limitations, this PheWAS demonstrates the utility of investigating the relationship between genetic variation and an extensive range of phenotypes by validating known genotype-phenotype associations as well as identifying novel genotype-phenotype associations, revealing complex phenotypic relationships and perhaps actual pleiotropy. The utility of this hypothesis-generating approach will continue to improve over time as more samples, variants, and phenotypes/traits across diverse populations are available for study in PAGE and other genomic resources. Larger, richer datasets coupled with methods development promise to more fully reveal the complex nature of genetic variation and its relationship with human diseases and traits.

#### **Methods**

#### **Study Populations**

All studies were approved by Institutional Review Boards at their respective sites (details are given in Text S1). The Population Architecture using Genomics and Epidemiology (PAGE) study includes the following epidemiologic collections: Atherosclerosis Risk in Communities (ARIC), Coronary Artery Risk in Young Adults (CARIDA), Cardiovascular Health Study (CHS), the Multiethnic Cohort (MEC), the National Health and Nutrition Examination Surveys (NHANES), Strong Heart Study (SHS), and Women's Health Initiative (WHI). For this PheWAS, data were available from ARIC, CHS, MEC, NHANES III, NHANES 1999–2002, and WHI (Table 1). The PAGE study design is described in Matise et al [26] and the PAGE PheWAS study design is described in Pendergrass et al [1].

**Table 5.** Example phenotype-classes and binned subphenotypes within phenotype-classes.

Phenotype Class	Study	Sub-phenotype binned within the phenotype-class
Asthma	ARIC	Asthma ever diagnosed?
Asthma	ARIC	Chest wheeze, whistle alot?
Asthma	ARIC	Chest wheeze, whistle, otherwise?
Asthma	ARIC	Age at 1st wheezing attack
Asthma	ARIC	Age asthma started
Asthma	ARIC	Age asthma stopped
Asthma	ARIC	Short of breath wheezing attack?
Asthma	CHS	Inhaled steroids for asthma
Asthma	CHS	Asthma confirmed by doctor
Asthma	CHS	Current asthma diagnosis by doctor
Asthma	EAGLE	Doctor ever told you had: asthma
Asthma	MEC	Asthma: History of Asthma, Hayfever, Skin Allergy, Food Allergy or Any Other Allergy from Baseline Questionnaire
CRP	CHS	C-reactive protein, adjusted original values (mg/l)
CRP	CHS	C-reactive protein, original values (mg/l)
CRP	EAGLE	Serum C-reactive protein (mg/dL)
Hemoglobin	ARIC	Hemoglobin
Hemoglobin	CHS	hemoglobin (g/dl)
Hemoglobin	EAGLE	Hemoglobin (g/dL)
Hemoglobin	WHI	Hemoglobin (gm/dl)
Hemoglobin	EAGLE	Mean cell hemoglobin concentration
Hemoglobin	EAGLE	Mean cell hemoglobin: SI (pg)

Presented below are examples of phenotypes binned into three phenotype classes, "Asthma", "BMI", and "CRP". Table S2 contains the complete list of matched phenotypes across studies within phenotype-classes, for all phenotype-classes used within this study. doi:10.1371/journal.pqen.1003087.t005

#### SNP Selection and Genotyping

All SNPs considered for genotyping in PAGE were candidate gene or GWAS-identified variants for phenotypes and traits available in the epidemiologic collections accessed by PAGE study sites. Cohorts and surveys were genotyped using either commercially available genotyping arrays (Affymetrix 6.0, Illumina 370CNV BeadChip), and/or custom mid- and low-throughput assays (TaqMan, Sequenom, Illumina GoldenGate or BeadXpress). Quality control was implemented at each PAGE study site independently. Study specific genotyping details are described in Text S1.

In this PheWAS, data were available for SNPs previously associated with HDL-C, LDL-C, and triglycerides [27], body mass index, obesity [28], type 2 diabetes, glucose, insulin [29], and measures of inflammation (C-reactive protein), among other diseases/traits. A total of 83 SNPs overlapped across at least PAGE study sites: ten were specifically selected for body mass index traits replication, three for C-reactive protein, six for coronary/cardiac traits, three for gout/kidney, 41 for lipids, and 20 for type 2 diabetes. Table S1 lists these SNPs, along with references reporting phenotypic associations from the NHGRI GWAS catalog [30] and the open access database of GWAS results of Johnson et al. 2009 [31]. The NHGRI GWAS catalog was most recently accessed in October, 2011. If no references were available from either of those two sources, a PubMed search was performed to retrieve relevant citations.

#### Statistical Methods

All tests of association were performed independently by each PAGE study site using the following analysis protocol: Linear or logistic regressions were performed for continuous or categorical dependent variables, respectively, assuming an additive genetic model (0, 1, or 2 copies of the coded allele). For variables with multiple categories, binning was used to create new variables of the form "A versus not A" for each category, and logistic regression was used to model the new binary variable. Linear regressions were repeated following a y to log (y+1) transformation of the response variable with +1 added to all continuous measurements before transformation to prevent variables recorded as zero from being omitted from analysis. All analyses were stratified by race-ethnicity.

Test of association were calculated for the number of SNPs and phenotypes listed in Table 1. The software used to calculate the associations for each study was as follows: ARIC (StatSoftware), CHS (R [32]), MEC (SAS), MEC (SAS v9.2), WHI (R), EAGLE (SAS v9.2 using the Analytic Data Research by Email (ANDRE) portal of the CDC Research Data Center in Hyattsville, MD).

All association results from the tests of association were reported in standardized templates designed by the PAGE coordinating center to facilitate data sharing. All results were then imported into a relational database (MySQL). The database was also used to match previously reported GWAS data with the SNPs analyzed in this study.

#### Plotting Significant Results

The software PheWAS-View was developed for data visualization of the PheWAS results as well as for plotting "Sun Plots" [33]. Synthesis-View [34,35] was also used to present results within this manuscript. Both software packages are freely available software for academic users: http://ritchielab.psu.edu/ritchielab/software,

and can be used with a web interface at: http://visualization.ritchielab.psu.edu/.

#### Matching Phenotypes

A total of 105 phenotype-classes were developed to manually match related phenotypes across studies. To bin related phenotypes into classes the following steps were used as visualized in Figure 6: First, using a MySQL database, the data from EAGLE, MEC, CHS, ARIC, and WHI were independently filtered for any tests of association results at p<0.01, and then lists of the unique phenotypes for each individual PAGE study were generated. The number of phenotypes that passed this significance threshold for each of the four groups was 604 (ARIC), 331 (CHS), 63 (MEC), 324 (EAGLE), 1,342 (WHI). Resulting phenotypes were then manually matched up between ARIC, CHS, MEC, EAGLE and WHI using knowledge about the phenotypes and the known focus of specific PAGE study survey questions (such as bone fracture questions used primarily for collecting information about osteoporosis). For some phenotypes, the specific phenotype existed clearly across more than one PAGE study, such as for the phenotype "Hemoglobin", where hemoglobin measurements were present for ARIC, CHS, EAGLE, and WHI. Other groups of phenotypes that fell within similar phenotypic domains but were not represented in the same form across studies were also collected into phenotype classes. One example is the phenotypes grouped together for the phenotype class of "Allergy". EAGLE collected specific quantitative data from allergy skin testing and had survey questions about the presence of allergies in participants. ARIC and MEC did not have skin allergy testing, but did have survey questions about the presence of allergies. Thus these allergy phenotypes were grouped together. Finally, phenotypes from all studies, regardless of significance from genotype-phenotype tests of association, were matched to the already-defined phenotype classes using the criteria described above. A phenotype that matched a phenotype class but was not associated with a SNP at the significance threshold of p<0.01 for a single study would still be included in the phenotype-class list. Using these criteria, a second curator reviewed the resultant phenotypes and phenotype classes for consistency and accuracy. To provide examples of the phenotype-classes, and which subphenotypes were matched with phenotype-classes, we show three phenotype-class examples in Table 5, and Table S2 contains the matched phenotypes across studies within the phenotype-classes for all phenotype-classes used within this study.

It is important to note resources that can be used for further investigation of the phenotypes listed in Table S2, as well as in the results presented in this paper. The following study websites contain additional information about all collected study information, including how those phenotypes were collected:

- ARIC http://www.cscc.unc.edu/aric/
- CHS http://www.chs-nhlbi.org/CHSData.htm, https://biolincc.nhlbi.nih.gov/static/studies/chs/Other\_Documents.htm
- WHI https://cleo.whi.org/data/Pages/home.aspx
- EAGLE http://www.cdc.gov/nchs/nhanes/ nhanes\_questionnaires.htm/
- MEC http://www.crch.org/multiethniccohort/ mec\_questionnaires.htm

#### References

1. Pendergrass SA, Brown-Gentry K, Dudek SM, Torstenson ES, Ambite JL, et al. (2011) The use of phenome-wide association studies (PheWAS) for exploration of

#### Criteria for Significance of Association

After creating phenotype-classes, significant PheWAS tests of association for single genotype-phenotype associations across PAGE studies were identified using a database query. Our criteria for considering a PheWAS test of association significant included a threshold of p<0.01 observed in two or more PAGE studies for the same SNP, phenotype class, and race/ethnicity and consistent direction of effect.

A total of 111 PheWAS tests of association met our criteria for significance (Table S3). Significant results were then binned based on class of association: known, related, and novel. In this PheWAS, Known Associations are positive controls and represent previously reported genotype-phenotype associations. Related Associations are SNPs significantly associated in this PheWAS with phenotypes judged to be closely related to phenotypes among Known Associations found here and the literature. Novel Associations are significant PheWAS results where 1) the association does not match a known association and 2) the phenotype for the PheWAS association is not within a similar phenotypic domain as the phenotype of known association.

#### **Ethics Statement**

All participating studies were approved by their respective IRBs, and all study participants signed informed consent forms.

#### **Supporting Information**

**Table S1** The list of all SNPs available for two or more sites in this study, arranged by previously associated phenotypes. (XLSX)

**Table S2** A list of the study level phenotypes, the study from which the phenotype is available, and the phenotype-class for each phenotype that overlapped with another study. (XLSX)

**Table S3** The expanded results for the 111 PheWAS associations identified in this study. (XLSX)

**Table S4** The 52 PheWAS results that replicated previously published genotype-phenotype associations. (XLSX)

**Text S1** Information on study design, phenotype measurement, and genotyping for each study. (DOCX)

#### **Acknowledgments**

The PAGE consortium thanks the staff and participants of all PAGE studies for their important contributions.

#### **Author Contributions**

Conceived and designed the experiments: SAP CK SB DCC MDR JLA CLA ED MDF CAH LAH C-NH RDJ LLM TCM KRM LM AR. Performed the experiments: SAP KB-G SD AF EST RG SB YL SLP. Analyzed the data: SAP KB-G SD AF EST RG SB YL SLP RW. Contributed reagents/materials/analysis tools: SAP SD AF EST RG SB GH CK LRW YL PB SLP. Wrote the paper: SAP JLA CLA SB PB ED MDF CAH GH LAH C-NH RDJ CK LLM YL SLP TCM KRM LM AR RW LRW DCC MDR. Developed the software for the plots in this publication: SAP SD.

novel genotype-phenotype relationships and pleiotropy discovery. Genetic epidemiology  $35\colon 410{-}422.$ 

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