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Journal Title: PLoS ONE
Volume: Volume 12, Number 12
Publisher: Public Library of Science | 2017-12-04, Pages e0188725-e0188725
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1371/journal.pone.0188725
Permanent URL: https://pid.emory.edu/ark:/25593/s7njf

Final published version: http://dx.doi.org/10.1371/journal.pone.0188725

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Accessed July 26, 2019 10:47 AM EDT
RESEARCH ARTICLE

Genome-wide admixture and association study of subclinical atherosclerosis in the Women’s Interagency HIV Study (WIHS)

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Abstract

Cardiovascular disease (CVD) is a major comorbidity among HIV-infected individuals. Common carotid artery intima-media thickness (cCIMT) is a valid and reliable subclinical measure of atherosclerosis and is known to predict CVD. We performed genome-wide association (GWA) and admixture analysis among 682 HIV-positive and 288 HIV-negative Black, non-Hispanic women from the Women’s Interagency HIV study (WIHS) cohort using a combined and stratified analysis approach. We found some suggestive associations but none of the SNPs reached genome-wide statistical significance in our GWAS analysis. The top GWAS SNPs were rs2280828 in the region intergenic to mediator complex subunit 30 (MED30) gene among HIV-positive women, and rs2907092 in the catenin delta 2 (CTNND2) gene among HIV-positive women, and rs7529733 in the region intergenic to family with sequence similarity 5, member C and regulator of G-protein signaling 18 (FAM5C | RGS18) genes among HIV-negative women. The most significant local European ancestry associations were in the region intergenic to the zinc finger and SCAN
GWAS and admixture mapping of cIMT among HIV in WIHS

approached to obtain informed consent for sharing of their data. Thus, we are not currently permitted to release their genomic or phenotypic data. This has been consistent with other genomic studies in the WIHS cohort. Data are available from the WIHS WDMAC Institutional Data Access/Ethics Committee for researchers who meet the criteria for access to confidential data. Data requests can be made by contacting WIHS WDMAC at JHSPH, wdmac@jhu.edu.

Funding: Data in this manuscript were collected by the Women’s Interagency HIV Study (WIHS). The contents of this publication are solely the responsibility of the authors and do not represent the views of the National Institutes of Health (NIH). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. WIHS (Principal Investigators): UAB-MS WIHS (Michael Saag, Mirjam-Colette Kemph, and Deborah Konkle-Parker), U01-AI-103401; Atlanta WIHS (Ighovwerha Ofotokun and Gina Wingood), U01-AI-103408; Bronx WIHS (Kathryn Anastos), U01-AI-035004; Brooklyn WIHS (Howard Minkoff and Deborah Gustafson), U01-AI-031834; Chicago WIHS (Mardge Cohen and Audrey French), U01-AI-034993; Metropolitan Washington WIHS (Seble Kassaye), U01-AI-034994; Miami WIHS (Margaret Fischl and Lisa Metsch), U01-AI-103397; UNC WIHS (Adara Adimora), U01-AI-033930; Connie Wofsy Women’s HIV Study, Northern California (Ruth Greenblatt, Bradley Assuizetar, and Phyllis Tien), U01-AI-034989; WIHS Data Management and Analysis Center (Stephen Gange and Elizabeth Tien), U01-AI-034993; Southern California WIHS (Joel Milam), U01-HD-032632 (WIHS I - WIHS IV). The WIHS is funded primarily by the National Institute of Allergy and Infectious Diseases (NIAID), with additional co-funding from the Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD), the National Cancer Institute (NCI), the National Institute on Drug Abuse (NIDA), and the National Institute on Mental Health (NIMH). Targeted supplemental funding for specific projects is also provided by the National Institute of Dental and Craniofacial Research (NIDCR), the National Institute on Alcohol Abuse and Alcoholism (NIAAA), the National Institute on Deafness and other Communication Disorders (NIDCD), and the NIH Office of Research on Women’s Health. WIHS data collection is also supported by the UCSD CTSA and UL1-TR000454 (Atlanta CTSA). The cardiovascular subsurveys in WIHS are also supported by 1R01HL126543, 1R21HL120394, 1R01HL083760, 1R01HL095140. The analysis in this study was supported by the pilot project funds domain containing 5D gene and NADH: ubiquinone oxidoreductase complex assembly factor 1 (ZSCAN5D | NDUF1) pseudogene on chromosome 19 among all women, in the region intergenic to vomeronasal 1 receptor 6 pseudogene and zinc finger protein 845 (VNR6P | ZNF845) gene on chromosome 19 among HIV-positive women, and in the region intergenic to the SEC23-interacting protein and phosphatidic acid phosphatase type 2 domain containing 1A (SEC23IP | PPAPDC1A) genes located on chromosome 10 among HIV-negative women. A number of previously identified SNP associations with cIMT were also observed and included rs2572204 in the ryanodine receptor 3 (RYR3) and an admixture region in the secretion-regulating guanine nucleotide exchange factor (SERGEO) gene. We report several SNPs and gene regions in the GWAS and admixture analysis, some of which are common across HIV-positive and HIV-negative women as demonstrated using meta-analysis, and also across the two analytic approaches (i.e., GWA and admixture). These findings suggest that local European ancestry plays an important role in genetic associations of cIMT among black women from WIHS along with other environmental factors that are related to CVD and may also be triggered by HIV. These findings warrant confirmation in independent samples.

Introduction

Early diagnosis and treatment of human immunodeficiency virus (HIV) has increased the life expectancy of infected individuals [1], and shifted the focus to the management of other comorbidities such as cardiovascular disease (CVD). CVD risk is higher among HIV-positive individuals compared to HIV-negative individuals [2–4]. HIV infection as well as certain antiretroviral therapies (ART) were shown to independently increase the risk of CVD [2]. Moreover, disparities in CVD risk by race and sex prevail, specifically, among HIV-positive individuals [5, 6]. Higher rates of acute myocardial infarction were observed among Blacks as compared to Whites, and among women as compared to men [3–5]. Similarly, ischemic stroke rates were reported to be higher among HIV-positive women as compared to HIV-positive men [7].

Efforts to prevent CVD among HIV-positive individuals have led to the evaluation of subclinical atherosclerotic measures such as carotid artery intima-media thickness (cIMT). CIMT can predict future CVD events among HIV-negative individuals [8]. CIMT is a non-invasive measure obtained using B-mode ultrasound, with the common carotid artery measurements considered the most reliable and reproducible of all CIMT segments [9, 10]. Common CIMT (cCIMT) measurements were reported to be thicker among HIV-positive as compared to matched HIV-negative individuals, and also independently associated with HIV infection and treatment [11, 12]. Racial and sex differences have shown greater cCIMT thickness among Blacks as compared to Whites, and among women as compared to men beyond traditional CVD risk factors [13–15].

The role of genetics as a risk factor for subclinical atherosclerosis, specifically in genome-wide association studies (GWAS) of cCIMT, has been explored among the HIV-negative [16] and non-Black populations [16–18]. A recent study on subclinical atherosclerosis that conducted a genome-wide association analysis, reported two SNPs—rs7412 in the apolipoprotein E (APOE) gene and rs143873045 in the KN Motif and Ankyrin Repeat Domains 2 (KANK2) gene that were found significantly associated with cCIMT among HIV-negative Blacks [19]. Among HIV-positive individuals, the only GWAS in relation to cCIMT has been conducted
by our group, among White men from the Fat Redistribution and Metabolic Change in HIV Infection (FRAM) Study [20]. We observed two single nucleotide polymorphisms (SNPs) in tight linkage disequilibrium (LD), rs2229116 and rs7177922, in the ryanodine receptor 3 (RYR3) gene on chromosome 15 to be associated with cCIMT. The association of rs2229116 was replicated in another group of HIV-positive White men from the Multicenter AIDS Cohort Study (MACS) [21]. Given the consistent association with cCIMT in HIV-positive White men, we further evaluated the association of SNPs within the RYR3 gene among a racially diverse group of HIV-positive women from the Women’s Interagency HIV Study (WIHS) [22]. The RYR3 gene association with cCIMT was observed among different SNPs than those reported earlier, rs2572204 being most significant among Black HIV-positive women. Haplotype blocks observed among the Black HIV-positive women compared to the other two racial/ethnic groups were smaller and indicated that SNP associations are possibly different based on LD, warranting further evaluation specifically in the context of cCIMT related admixture.

Admixture mapping enables the identification of genetic variations differentially distributed in ancestral populations associated with a disease—the premise being that associated variants are observed on chromosomal segments from ancestral populations with higher prevalence of the disease/trait [23]. Thus, this approach takes advantage of the extended LD present in the recently admixed population to discover these variations. Admixture mapping has been considered a feasible and more powerful tool in contrast to GWAS among individuals of mixed continental ancestry [24], and advances in statistical techniques and their applications in a genome-wide set of markers has greatly increased the ability to detect disease loci. We recently performed genome-wide admixture mapping in relation to cCIMT among HIV-negative Black individuals from two large prospective cohorts and found a region in the secretion regulating guanine nucleotide exchange factor (SERGEF) gene that reached genome-wide significance [25]. Likewise, we have shown various genomic regions associated with clinical events in African Americans through admixture mapping [26].

Currently, methods based on principal components are commonly used to control for population stratification in genetic association studies [27]. Recent studies have focused on admixture mapping and related techniques to incorporate ancestry information in association tests among admixed populations [28–32]. However, there is a lack of consensus in approaches and whether accounting for local and global ancestries alone or together would sufficiently control for both type I and type II error rates [31, 32]. In this study, we performed genome-wide admixture and association analysis to determine single SNP and local European ancestry associations with cCIMT among Black women from the WIHS cohort.

Results

The HIV and CVD related characteristics of the 970 Black women included in the current study are presented in Table 1. In this study, 70% of the women were HIV-positive, and had similar cCIMT values as well as percent global European ancestry as compared to HIV-negative women. CVD risk factors including hypertension, diabetes, and current smoking were similar across the two groups. HIV-positive women were older, had lower CD4 cell count and HDL cholesterol levels, and were more likely to take medications for hypertension compared to HIV-negative women.

Genome-wide association results

The genome-wide association results for cCIMT are presented for all women and after further stratification by HIV status, as shown in the Manhattan plots (Fig 1A–1C). None of the SNPs
reached the genome-wide significance level (S1 Table), but we present the top 5 SNP-cCIMT associations for all three groups (Table 2) and the regional plots for the top SNPs in each group (S1A–S1C Fig). The lambda values for each of the GWAS models were 1.005 for the combined model, 1.009 for the HIV-positive model, and 1.013 for the HIV-negative model (Fig 2A–2C). We also present the admixture analyses results for gene regions that include these significant GWAS SNPs to determine if local European ancestry at these regions is associated with cCIMT in Table 2. The top hit among all women was rs2280828 located in the intergenic region of mediator complex subunit 30 and exostosin glycosyltransferase 1 (MED30 | EXT1) genes on chromosome 8 ($\beta = -0.0216$, $p = 8.01 \times 10^{-7}$). The same SNP was also significantly associated with cCIMT among HIV-positive women ($\beta = -0.0271$, $p = 2.89 \times 10^{-7}$). The most significant SNP associated with cCIMT among HIV-positive women was rs2907092 in the catenin delta 2 (CTNND2) gene on chromosome 5 ($\beta = 0.0250$, $p = 2.37 \times 10^{-7}$). Local European ancestry at the corresponding gene region of this SNP was also associated with cCIMT ($\beta = 0.0089$, $p = 0.04$). Local European ancestry was also associated with cCIMT at the gene region that includes the GWAS identified rs4761669 intergenic to the transmembrane and coiled-coil domain family 3 gene and the NADH: ubiquinone oxidoreductase subunit A12 (TMCC3 | NDUFA12) genes on chromosome 12 ($\beta = -0.0084$, $p = 0.04$). Among HIV-negative women, rs7529733 intergenic to family with sequence similarity 5, member C and regulator of G-protein signaling 18 genes (FAM5C | RGS18) on chromosome 1 was the top SNP associated with cCIMT ($\beta = 0.0431$, $p = 2.92 \times 10^{-7}$).

Local ancestry association results

The local European ancestry (LEA) association with cCIMT is presented in Table 3 for the top 5 gene regions for all women and then separately for HIV-positive and HIV-negative women, and the corresponding Manhattan plots are presented in Fig 1D–1F. The regional plots for the top regions are presented in S1D–S1F Fig (all regions in S2 Table). The lambda values for each of the admixture models were 1.001 for the combined model, 1.103 for the HIV-positive model, and 1.000 for the HIV-negative model (Fig 2D–2F). We also present the top GWAS hit located within the top ancestry gene regions if it was significant at $\alpha$ of 0.05 (Table 3). LEA at 17 gene regions reached genome-wide significance in relation to cCIMT among all women. These regions were largely located on chromosome 19 over a 3.07 Mb region but also included...
one gene region on chromosome 6 spanning 106.38 kb. There were no gene regions that reached genome-wide significance among HIV-positive women. Among HIV-negative women, a total of 48 LEA gene regions achieved genome-wide significance and were largely located on chromosome 10 over a span of 4.23 Mb. A single gene region each on chromosomes 2, 4, and 9 also achieved genome-wide significance among HIV-negative women. The most significant local European ancestry association among all women was observed in the region intergenic to the zinc finger and SCAN domain containing 5D gene and NADH: ubiquinone oxidoreductase complex assembly factor 1 pseudogene (ZSCAN5D | NDUF1) on chromosome 19 (β = 0.0134, p = 8.87x10^-5). The region on chromosome 6 in the leucine rich repeat containing 1 gene (LRRC1) achieved genome-wide significance among all women but was also observed among the top 5 regions in HIV-positive women (β = 0.0138, p = 8.42x10^-4). The region intergenic to the vomeronasal 1 receptor 6 pseudogene and zinc finger protein 845 gene (VN1R6P | ZNF845) on chromosome 19 was most significant among HIV-positive women (β = 0.0141, p = 6.86x10^-4). Among HIV-negative women, the top association was observed in the region intergenic to the SEC23 interacting protein gene and phosphatidic acid phosphatase type 2 domain containing 1A gene located on chromosome 10 (SEC23IP | PPAPDC1A) (β = -0.0290, p = 1.71x10^-5).

**Meta-analysis and previously significant SNPs**

Previously significant SNPs associated with cCIMT that were found significant in the current GWAS and LEA association analysis are reported in the **Tables 4 and 5**, respectively. A total of 25 SNPs in the GWAS and 31 SNPs in the admixture analysis were significantly associated with cCIMT. The most significant SNPs across the combined and stratified GWAS analyses
Table 2. Genome-wide association (GWA) results for common carotid artery intima media thickness and the corresponding admixture (LEA) regions for the top 5 hits among participants from the Women's Interagency HIV Study (WIHS) cohort.

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Position</th>
<th>SNP</th>
<th>MAF-CEU/YRI</th>
<th>Gene</th>
<th>Location</th>
<th>Beta</th>
<th>P</th>
<th>Gene region</th>
<th>First SNP</th>
<th>Last SNP</th>
<th>Beta</th>
<th>P</th>
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</tr>
<tr>
<td>8</td>
<td>118762436</td>
<td>rs2280828</td>
<td>C: 0.009/0.08</td>
<td>MED30/EXT1</td>
<td>Intergenic</td>
<td>-0.0216</td>
<td>8.0E-07</td>
<td>118.554–118.808</td>
<td>rs1688962</td>
<td>rs4876390</td>
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<td>11</td>
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<td>159.84–159.846</td>
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<td>3.3E-06</td>
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<td>2.9E-07</td>
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<td>1.7E-06</td>
<td>111.499–111.75</td>
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<td>2.2E-06</td>
<td>169.075–169.607</td>
<td>rs1885634</td>
<td>rs7764005</td>
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SNP: single nucleotide polymorphisms, LEA: local European ancestry

https://doi.org/10.1371/journal.pone.0188725.t002
were—rs445925 in the apolipoprotein C-1 (APOC1) gene among all women ($\beta = -0.0081$, $p = 0.0018$), rs2572204 in the ryanodine receptor 3 (RYR3) among HIV-positive women ($\beta = 0.0153$, $p = 0.0027$), and rs854571 in paraoxonase 1 (PON1) gene in the HIV-negative women ($\beta = 0.0107$, $p = 0.02$). The most significant SNPs in the LEA association included rs1436023 in LOC105375856 gene in the combined analysis ($\beta = -0.0099$, $p = 0.005$), the region between 17.809 to 18.035 Mb in the SERGEF gene significantly associated with cCIMT among the HIV-positive women ($\beta = -0.0093$, $p = 0.02$) whereas the SNP rs11120748 in the CD55 molecule (Cromer blood groups) (CD55) gene was significant among HIV-positive women ($\beta = -0.0156$, $p = 0.02$).

Meta-analysis results for both GWAS and LEA associations reflect the results observed in the combined analyses (S1 and S2 Tables).

**Discussion**

The current study presents the genome-wide admixture and association findings for cCIMT among Black HIV-positive and HIV-negative women from the WIHS cohort. We ran separate analyses for the two methods but also noted whether any of the GWAS SNPs within the top admixture regions, and any of the admixture regions including the top GWAS SNPs were significant for cCIMT. We observed SNPs in the GWA that showed suggestive associations ($p < 10^{-6}$) with cCIMT (included in Table 2 in bold). The comparison to the admixture results yielded two significant regions observed among HIV-positive women (included in Table 2 in bold). Conversely, several regions with LEA associations revealed GWAS SNPs that were nominally significant. Meta-analysis results suggest that some SNPs and gene regions are common
Table 3. Genome-wide admixture (LEA) and the corresponding genome wide association (GWA) results for the top 5 hits among participants from the Women’s Inter-agency HIV Study (WIHS) cohort.

<table>
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<th>Chromosome</th>
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<td>rs73934760</td>
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<td>Intron</td>
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<td>0.005*</td>
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<td><strong>HIV-negative</strong></td>
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SNP: single nucleotide polymorphism.
* Significant (P<0.05) after adjusting for number of SNPs in each region.

https://doi.org/10.1371/journal.pone.0188725.t003
across HIV-positive and HIV-negative women and may therefore play a role in atherosclerosis beyond HIV associated risk factors. Lastly, we observed associations with SNPs that have been previously reported to have significant associations with cCIMT.

The top GWAS hit rs2280828 observed in the combined analysis was in the region intergenic to the MED30 and EXT1 genes on chromosome 8, approximately 50 kb from the EXT1 gene. The EXT1 gene encodes for a transmembrane glycoprotein on the endoplasmic reticulum (ER) of the cell which is involved in the synthesis and polymerization of heparan sulfate [33, 34]. Heterozygous loss of function mutation in the EXT1 and EXT2 genes have been postulated to improve endothelial function by increasing the bioavailability of nitric oxide (NO) [35]. Calcium ion concentrations in the cell regulate the activity of nitric oxide synthase via calmodulin, indirectly influencing the release of nitric oxide when shear stress is exerted on the endothelium [36]. The RYR3 receptor encoded by the RYR3 gene is a calcium-ion channel...
present on the ER and is involved in calcium homeostasis [37]. Thus, EXT1 may play a role in atherosclerosis by regulating endothelial function in conjunction with RYR3. On the other hand, the MED30 gene is part of a large group of genes that encode the mediator complex proteins, constituting a multi-subunit complex, known to regulate the RNA polymerase II enzyme [38]. Silencing of this gene was shown to inhibit the transcription of HIV, in part via the HIV-Tat protein [38]. The HIV-Tat protein causes endothelial dysfunction and brings about an inflammatory response as well as triggers loss of calcium through the RYRs on the ER [39]. These factors are part of the biological mechanisms that could contribute to atherosclerosis in HIV. Therefore, MED30 gene-related functions may play a role in mitigating the downstream

Table 5. Admixture (LEA) results for common carotid-artery intima media thickness association with previously significant single nucleotide polymorphisms (SNPs).

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Position</th>
<th>SNP</th>
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<th>Beta</th>
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https://doi.org/10.1371/journal.pone.0188725.t005
effects of HIV in atherosclerosis and other cardiovascular complications. Spliced variants of the gene have been observed in circulating endothelial progenitor cells and are hypothesized to have potential diagnostic or therapeutic roles in cardiovascular disease in the future [40].

Rs2907092 was the top GWAS hit among HIV-positive women and is located in the intronic region of the CTNND2 gene on chromosome 5. This gene encodes for the δ-catenin protein, a neuronal protein that is also expressed in the vascular endothelium [41]. The protein binds cadherins to the cytoskeleton and is involved in pathways and mechanisms that modulate vascular remodeling. They are responsible for endothelial cell motility, angiogenesis and wound repair regulated by inflammatory responses and are specifically observed during pathologic states. This gene showed a strong association with cerebral malaria in an environmental correlation analysis (ECA), whereby the involved mechanisms such as luminal blockade by infected cells, increased shear stress, presence of reactive oxygen species (ROS) and the ensuing inflammatory response follow a pattern that is similar to atherosclerosis [42]. ECA identifies genes that are enriched as a result of local environmental effects within a population that has likely been isolated for a long period of time. As a result, the CTNND2 gene may likely have an admixture component, which was precisely noted in our admixture results.

The most significant GWAS SNP among HIV-negative women was rs7529733 in the regions intergenic to the FAM5C and RGS18 genes on chromosome 1. The FAM5C gene has been associated with myocardial infarction [43] and functional studies have shown that it increases the production of ROS, nuclear factor-kappaB (NF-κB) activity, and expression of the intercellular and vascular cell adhesion molecules, ICAM-1 and VCAM-1 [44]. These factors induce vascular inflammation and facilitate monocyte adhesion to the endothelium. HIV proteins such as HIV-Tat trigger this mechanism indirectly by activating the monocyte chemotactant protein-1 (MCP-1) resulting in migration of the monocytes and their transformation into macrophages, steps integral to the atherosclerotic process [39]. The regulator of G-protein signaling 18 GTPase-activating protein is encoded by the RGS18 gene. This protein is involved in the activation and inhibition of platelets through the G-α-q subunit of the heterotrimeric G-protein [45]. The RGS18 protein is in turn regulated by thrombin resulting in the phosphorylation of one of its residues, and subsequent release of calcium ions to induce platelet aggregation. A similar process is involved during platelet inhibition whereby prostacyclin and NO regulate RGS18 resulting in reduced intracellular calcium levels. Thus, the RGS18 effect on platelets is mediated through calcium homeostasis and therefore genetic interactions between RGS18 and RYR3 gene are possible, and may play a substantial role in atherosclerosis as platelets are involved in its inflammatory response.

The top admixture mapping results for the combined, and stratified analyses among HIV-positive women revealed genes that are pseudogenes, and which have not yet been evaluated for associations with human complex diseases or traits. The second top hit involves the galanin-like peptide encoded by the GALP gene on chromosome 19. It is a neuropeptide of the galanin family and has been shown to play a role in homeostatic processes of the central nervous system (CNS) [46]. These include but are not limited to hormonal regulation of food intake with the potential to develop obesity and other weight gain related complications like diabetes. Additionally, a splice variant of the peptide called alarin was observed to have vasoactive properties and induced vasoconstriction and decrease in water retention in the epidermis. Diabetes and obesity are risk factors for atherosclerosis and CVD, and vascular perturbations in the microvasculature are known to induce downstream effects that play a role in the atherosclerotic process. The TRPM6 gene located on chromosome 9 was associated with cCIMT among HIV-positive women. This gene encodes for the transient receptor potential melastatin 6 channel which acts as both a channel and an enzyme involved in the regulation of magnesium (Mg^{2+}) in the body [47]. TRPM6 and TRPM7 were both found in the vascular smooth
muscle cells and were shown to regulate hypertension via Mg$^{2+}$ homeostasis. Reduced Mg$^{2+}$ increases blood pressure by increasing the reactivity and contractility of the blood vessels and is also responsible for endothelial dysfunction, inflammation and vascular remodeling. All the above-mentioned mechanisms are involved in the ongoing process of atherosclerosis and subsequent CVD and therefore TRPM6 may play a part in the development of CVD. Among HIV-negative women, the top LEA region was on chromosome 10 and located in the intergenic region of the SEC23IP and PPAPDC1A genes. The rs1571099 SNP in PPAPDC1A was significantly associated with interventricular septal wall thickness among African Americans in a recent GWAS using the Candidate Gene Association Resource (CARe) Study [48]. This gene is involved in pathways related to innate immunity, phagocytosis and triacylglycerol synthesis [49]. Therefore, this gene may also play a role in atherosclerosis, albeit an indirect one.

Replication revealed associations with some of the cCIMT-SNPs found significant in earlier studies. Previously, we had reported the association of two SNPs in RYR3 –rs2229116 and rs7177922 that had achieved genome-wide significance among HIV-positive men of European ancestry [20], with successful replication of rs2229116 in another group of HIV-positive White men [21]. However, we did not find any association with these SNPs in the current analysis. Instead, we found another SNP –rs2572204 in RYR3 that was associated with cCIMT among HIV-positive women of WIHS. We reported the same association among Black HIV-positive women in our previous candidate-gene study on RYR3 in WIHS. The different SNP associations within this gene may either be because of the differences in race or sex between the two studies. WIHS women are also noted to have different heart disease risk because of differences in their immunosuppression or duration of ART as compared to HIV-positive men [50, 51]. Nonetheless, RYR3 appears to play a role in HIV-related atherosclerosis risk based on some of the possible gene (or protein) interactions found significant in this study. The admixture association of the SERGEF gene in the HIV-positive women provides further support for the role of this gene in atherosclerosis. However, the direction of association was opposite to that observed in the HIV-negative population in our previous study. The gene encodes a guanine nucleotide exchange factor that may play a role in atherosclerosis through immune or inflammatory mechanisms via guanosine-triphosphatases and serotonin [25]. We speculate that lower cCIMT associated with SERGEF in the HIV-positive women may be a result of immuno-suppression because of ART or HIV itself or in conjunction with the gene as it appears to mediate atherosclerosis through immune-related mechanisms.

Population stratification is an issue that needs to be addressed during GWAS to avoid spurious associations, more so in admixed populations. Literature on genetic association studies among admixed populations has persistently examined different approaches to account for population structure. An initial and basic approach was genomic control [52] which was improved upon by Price et al. [27] who suggested controlling for principal components. This is currently the mostly widely used approach in GWAS and other genetic association studies among admixed populations. However, recent studies argue that adjusting for global ancestry is not sufficient and does not account for ancestry at the marker level for each individual SNP which is commonly referred to as local ancestry [28]. To this effect, joint testing of admixture and association was proposed and demonstrated an increase in statistical power and accuracy of the results compared to existing methods [30, 31]. In the current study, we compared the performance of two different methods based on level of ancestry adjustment; the methods are described in the supplementary methods section (S1 File) and the results are presented in S3 Table. The comparison shows that the power is highest for the QSUM test that adjusts for both local and global ancestries using the MIXSCORE program. Pasaniuc et al. reported that the QATT test performed better for quantitative traits but also noted that the QSUM test
might perform better given a set of conditions where high signals of admixture associations were observed [30]. The BMIX test as proposed by Shriner et al. performs even better than the QSUM test with a power of ~72% to detect phenotype-genotype associations in the overall analysis [31]. Both results support the hypothesis that joint testing with adjustment for both local and global ancestries is more powerful than the current association tests in use. Therefore, we performed a secondary analysis using BMIX and present our results in the supplemental S4 Table. Of note, the top SNPs in Table 2 showed similar significance after adjustment for local ancestry using BMIX (if available). Since BMIX uses the local ancestry information available through admixture mapping, as such the two analyses are not independent of each other. In our study, we did not conduct the primary GWAS analyses using the BMIX method as we could only estimate local ancestry in 473,732 SNPs, and so adjustment with local ancestry would have dramatically reduced the number of informative SNPs available for GWAS. Nevertheless, this approach of adjustment for both local and global ancestry has not been extensively explored and needs further validation to ensure broader application in future genetic studies among admixed populations and also tease shared traits across ancestries [53].

The advantage of using the WIHS cohort is that a higher proportion of women in the cohort were African American, especially among the HIV-positive subpopulation. It also presented us with an opportunity to evaluate the common CIMT association exclusively among women. However, we recognize that the homogeneity of the population also reduces the generalizability of our findings. As the WIHS participants are relatively young and the incidence of clinical CVD events is low, we could not assess the association of CVD events at the genome-wide level. Therefore, we assessed cCIMT which is a reliable and reproducible subclinical measure of atherosclerosis that has been shown to predict future CVD events. Most of the HIV-positive women were also on combination ART including multiple antiretroviral drugs which prevented us from evaluating the independent effect of specific ART agents on cCIMT. Furthermore, our admixture analysis only included a subset of the SNPs that were part of the GWAS (N = 473,732), and so we were limited in our ability to assess local ancestry associations at the excluded SNPs. This also limited our ability to adjust for both local and global ancestry during GWAS using the joint admixture and association testing method. We acknowledge that the sample size was small for a genome-wide analysis, however, WIHS is currently the largest prospective cohort of HIV-positive women in the U.S. with the cCIMT measurements since cCIMT is not readily available, specifically among HIV-positive women. The heterogeneity index $I^2$ (S1 and S2 Tables) also shows that there are some consistent results (both in direction of association and significance) but also inconsistent ones between HIV-positive and HIV-negative groups. These results may suggest differences in underlying biological mechanism, i.e. some of the genes or pathways maybe specific in the context of HIV (virus itself or treatment) and others similar in both with or without infection, or they could be artifacts as a result of the small sample size. While we cannot conclude the reason of heterogeneity in the two groups, the suggestive results should be examined in other populations.

In summary, we report several novel associations in relation to common CIMT among a group of Black HIV-positive and HIV-negative women observed through genome-wide association and admixture analysis. The results suggest that GWAS results do not completely capture associations in regions of local ancestry and vice-a-versa among admixed populations and both approaches may still be necessary to evaluate genotype-phenotype associations in specific settings. These findings need to be confirmed in larger studies with local ancestry available for a denser set of markers to allow equal comparisons across GWAS and admixture results. Future studies may also benefit from combining data from other HIV cohorts and performing a meta-analysis to detect novel or stronger associations.
Methods

Ethics statement

The parent WIHS study and this sub-study conformed to the procedures for informed written consent approved by institutional review boards (IRB) at all sponsoring organizations and to human-experimentation guidelines set forth by the United States Department of Health and Human Services, and finally reviewed and approved by the University of Alabama Institute Review Board.

Study population

WIHS is a large prospective cohort of HIV-positive and risk-matched HIV-negative women enrolled across six sites in the United States [54, 55]. The women in the study were enrolled during the initiation phase in 1994/95 (2054 HIV-positive and 569 HIV-negative), the expansion phase in 2001/02 (737 HIV-positive and 406 HIV-negative), and most recently during 2011/2012 to replace the women who had died during follow-up (270 HIV-positive and 93 HIV-negative). The study was designed to comprehensively evaluate the impact of HIV infection in U.S. women, and provide comparisons with the HIV-negative group having similar characteristics and a high risk of acquiring the infection themselves. The women in the expansion phase were included to examine the risks and benefits associated with ART. A vascular disease sub-study was initiated in April 2004 to examine the effect of HIV and ART on subclinical atherosclerosis over time, focusing specifically on inflammation, coagulation and lipid pathways [50]. The measurements for cCIMT were obtained among 1331 HIV-positive and 534 HIV-negative women in WIHS through the vascular sub-study. In the current study, 970 Black, non-Hispanic women (682 HIV-positive and 288 HIV-negative) with complete genotype, phenotype and covariate information were included, after confirming the race/ethnicity for these women using the principal components method.

Data collection. The collection and processing of data in WIHS was standardized with centralized training of all staff before initiation of enrollment. Interviews with structured questionnaires were conducted to obtain participant-reported information on demographic, lifestyle, socio-economic and clinical factors. In addition, anthropometric measures such as weight and height; physiologic measures such as blood pressure (BP); laboratory measures such as markers of HIV, and lipid profile were obtained using standardized protocols; and gynecologic examinations were conducted. Follow-up every six months included collection of interview related information and medication use [54].

CIMT images were obtained via high-resolution ultrasound by sonographers trained centrally at the University of Southern California Atherosclerosis Research Unit Core Imaging and Reading Center using a standard protocol across all study sites [56]. CIMT was measured by the automated computerized edge-tracking method (patents 2005, 2006, 2011) [57]. The coefficient of variation was 1.8% (intraclass correlation = 0.98; n = 113) for repeated CIMT measurements with the initial images guiding the repeat scans. The complete ultrasound included: 1) standardized measurements of the far-right wall of the distal common carotid artery for CIMT, stiffness and lesions, and 2) scanning the proximal internal carotid, external carotid and right carotid bulb at the bifurcation for lesions [50]. In the current study, we utilized the common carotid artery measurements from the distal far right wall. The mean right wall cCIMT was log base 10 transformed for normalization.

Genotyping. DNA isolation, quantification and the microplate reading were performed using the Pure-gene DNA isolation kit (Gentra Systems, Minneapolis, MN), the PICO Green dsDNA quantification kit (Molecular Probes, Eugene, OR) and the Perkin Elmer HTS7000
BioAssay Reader, respectively [58]. All samples were stored at -20 °C. The genome-wide scan was performed using the Illumina HumanOmni2.5-quad beadchip (NCBI build 37, hg19, Illumina, San Diego).

**Statistical analysis**

Both the genome-wide and ancestry association analyses were initially conducted for all individuals followed by stratified analyses among HIV-positive and HIV-negative Black women. Common CIMT was associated with all CVD risk factors except HDL cholesterol, and was also not associated with ART or CD4 cell count in bivariate analysis. However, HIV related risk factors such as ART and CD4 cell counts have been previously associated with cCIMT and were therefore included in the final analysis.[11, 50] Analyses among all and the HIV-positive individuals were adjusted for age, CD4 cell count, ART use, hypertension (systolic BP ≥ 140 mmHg, diastolic BP ≥ 90 mmHg, antihypertensive medication use or self-report), diabetes mellitus (anti-diabetic medication use, fasting glucose ≥ 126 mg/dl, HgbA1C ≥ 6.5%, or self-report), current smoking, low-density lipoprotein (LDL), and high-density lipoprotein (HDL) cholesterol. Covariate adjustment for HIV-negative individuals was the same except for ART use.

**Quality control of genomic data and GWA analysis.** Multivariable linear regression analysis using an additive model was performed in PLINK (v1.9) with default settings [59]. In the current study, genotype information for a total of 1,590,141 SNPs was available. Quality control resulted in the removal of 2 SNPs because of missingness at >10%, 6,433 because of Hardy-Weinberg equilibrium < 0.001, and 459,405 because of minor allele frequency (MAF) < 5%, leaving a total of 1,124,301 SNPs to be included in the genome-wide association analysis (S5 Table). Genetic ancestry components were evaluated with principal component (PC) analysis using 168 ancestry informative markers (AIMs) as previously described [60]. Briefly, the number of PCs which distinguished the major racial/ethnic groups in the sample was sought by visual inspection of scatter plots of orthogonal PCs (i.e., PC 1 versus PC2, PC2 versus PC3). This procedure was repeated until no discernible clustering of individuals by their self-reported race/ethnicity was possible (data not shown) [60, 61]. Based on the scree plot (S2 Fig), GWAS analysis was adjusted for 2 principal components (PCs) in addition to the other covariates, described above. We also tested for genomic inflation of the GWAS results by obtaining the lambda values as well as QQ plots for both the combined as well as stratified models. Correction for multiple testing was performed using a method proposed by Shriner et al [31]. The method estimates the effective number of tests by fitting an autoregressive model to the additively coded genotypes and choosing the order of the model based on the lowest Akaike information criteria (AIC). The total number of independent tests estimated using this method were 769,204, resulting in a threshold significance level of $6.50 \times 10^{-8}$.

**Local ancestry estimation.** Local ancestry was estimated using the Local Ancestry in adMixtured Populations using Linkage Disequilibrium (LAMP-LD) program [62]. The program incorporates a dense set of markers to determine ancestry at each SNP for each individual based on haplotype sets from ancestral populations. Phased haplotype data for the reference populations was obtained from HapMap phase II and III data from Utah residents with North-West European ancestry (CEU) and Yorubans from Ibadan, Nigeria (YRI) for African ancestry (HapMap phase 3, release 2). The reference haplotype data was used to determine parameters for the Hidden Markov Model (HMM) with 15 state spaces. The local ancestry was estimated using the HMM parameters within a 300 SNP-long window-based framework. Each chromosome was analyzed separately and the local ancestry coded as the number of European ancestry alleles at each SNP (i.e., 0, 1, or 2 where “0” means both chromosomal regions had
African ancestry, “1” means the chromosomal region is admixed, one with European ancestry and the other with African ancestry, and “2” means both chromosomal regions had European ancestry). Before ancestry estimation, the physical coordinates for the remaining 1,124,301 SNPs were converted from hg19 to hg18 to match the coordinates with the HapMap phased data using the UCSC table browser [63]. As a result, 119 SNPs were lost during conversion. Furthermore, 648,814 SNPs were not available in HapMap, 1,259 had ambiguous A/T or G/C SNPs with MAF > 35% and could not be resolved for strand annotation, and 63 were duplicates; this resulted in 473,732 autosomal SNPs remaining for inclusion in the admixture estimation and association analyses (S5 Table).

A SNP most significantly associated with the clinical events along with the block of neighboring SNPs with similar number of European ancestry (“0”, “1” or “2”) were denoted as a “LEA region”. Additionally, blocks of same ancestry LEA regions were further defined based on whether the SNPs were in the genic or intergenic regions. A genic region included the 5’ and 3’ untranslated regions, the exons and the introns; the intergenic regions were those that fell between two genic regions as described above. Gene nomenclatures were obtained from the National Center for Biotechnology Information’s (NCBI) Reference Sequence database through the web ANNOVAR program [64].

Local ancestry association with cCIMT. Association testing in relation to cCIMT was performed using the PLINK (v1.9) software with default settings [59]. We conducted an additive linear regression analysis to evaluate the association of local ancestry coded as homozygous European ancestry (11), heterozygous European/African ancestry (12) and homozygous African ancestry (22), controlling for global European ancestry in the model. Since ancestry estimates are highly correlated, it is necessary to calculate the total number of effective independent tests to arrive at a threshold genome-wide significance level. We calculated the total number of independent tests using the method proposed by Shriner et al for admixture analysis as well [31, 65]. The total number of independent tests in our analyses was 146 resulting in a threshold significance level of 3.42 x 10^-4.

Meta-analysis. Meta-analysis was performed on the results from the GWAS and admixture analyses separately. The coefficients obtained from the stratified analyses of HIV-positive and HIV-negative women were combined to obtain a pooled estimate. The analysis was conducted using PLINK (v1.9) for quantitative traits with both fixed- and random-effects analysis, including the detection of heterogeneity of the results (between HIV-positive and HIV-negative) using the heterogeneity index I^2 (S1 and S2 Tables).

Evaluation of previously significant cCIMT SNPs
We identified 377 SNPs based on previous cCIMT-genetic association studies (S6 Table). Of these, 145 SNPs were obtained from a combination of candidate gene, genome-wide (GWAS) or meta-analysis studies [16, 19, 20, 22, 66–87], 232 SNPs from a GWAS conducted by Melton et. al. [17]. We assessed the GWAS results and LEA regions that contained these SNPs and report the associations for those found significant at P<0.05 in our cCIMT analysis.

Supporting information

S1 File. Supplemental methods. Comparing test performance after ancestry adjustment. (DOCX)

S1 Table. Genome-wide association (GWA) results for cCIMT in the combined, HIV-positive, and HIV-negative models, including the meta-analysis results. (XLSX)
S2 Table. Genome-wide admixture (LEA) results for cCIMT in the combined, HIV-positive, HIV-negative models, including the meta-analysis results.
(XLSX)

S3 Table. Comparing test performance after adjusting for global, local or both ancestries in the association tests. QATT: quantitative Armitage’s trend test adjusting for global ancestry, QSNP1: quantitative single nucleotide polymorphism association adjusting for local ancestry, QADM: quantitative admixture test, QSUM: sum of QSNP1 and QADM, BMIX: joint admixture and association test using a Bayesian approach, Score: minus log 10 p-values.
(DOCX)

S4 Table. Joint association and admixture (LEA) results for cCIMT in the combined, HIV-positive, and HIV-negative models using the BMIX method.
(XLSX)

S5 Table. Quality control of the genome wide data from the Women’s Interagency HIV study (WIHS). SNP: single nucleotide polymorphism, MAF: minor allele frequency.
(DOCX)

S6 Table. List of SNPs previously significantly associated with cCIMT in candidate gene, genome-wide association or meta-analysis studies. *Publications before the year 2004 are represented in two reviews by Manolio TA 2004 and Pollex RL 2006. Two or more publications are referenced for some SNPs. Several SNP or gene associations were common across studies but are only presented once in the table. Publications in bold were the only studies that included a HIV-positive population.
(XLSX)

S1 Fig. Regional plots for the topmost hit from the genome-wide association and admixture analyses. a) All women (GWAS), b) HIV-positive women (GWAS), c) HIV-negative women (GWAS), d) All women (admixture), e) HIV-positive women (admixture), and f) HIV-negative women (admixture). Negative log10 p-values and recombination rates are plotted against SNPs and their chromosomal positions.
(PDF)

S2 Fig. Scree plot showing the eigenvalues plotted against the principal components.
(PDF)

Acknowledgments
Data in this manuscript were collected by the Women’s Interagency HIV Study (WIHS). The contents of this publication are solely the responsibility of the authors and do not represent the official views of the National Institutes of Health (NIH).

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

We would also like to thank Dr. Daniel Shriner for his input with the analysis.

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References


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