Leveraging population admixture to explain missing heritability of complex traits

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Abstract
Despite recent progress on estimating the heritability explained by genotyped SNPs ($h_g^2$), a large gap between $h_g^2$ and estimates of total narrow-sense heritability ($h^2$) remains. Explanations for this gap include rare variants, or upward bias in family-based estimates of $h^2$ due to shared environment or epistasis. We estimate $h^2$ from unrelated individuals in admixed populations by first estimating the heritability explained by local ancestry ($h^\gamma_2$). We show that $h^\gamma_2 = 2F_{STC} \theta (1 - \theta) h^2$, where $F_{STC}$ measures frequency differences between populations at causal loci and $\theta$ is the genome-wide ancestry proportion. Our approach is not susceptible to biases caused by epistasis or shared environment. We examined 21,497 African Americans from three cohorts, analyzing 13 phenotypes. For height and BMI, we obtained $h^2$ estimates of 0.55 ± 0.09 and 0.23 ± 0.06, respectively, which are larger than estimates of $h_g^2$ in these and other data, but smaller than family-based estimates of $h^2$.

**Introduction**

Understanding the genetic architecture of complex human phenotypes is a fundamental question to the field of genetics, with broad implications for identifying genes related to disease and predicting individual risk profiles. A central element of this problem is estimating narrow-sense heritability ($h^2$), the fraction of phenotypic variation in a population determined by genetic variation under an additive model. While the last decade of genome-wide association studies (GWAS) produced thousands of novel loci associated with hundreds of phenotypes, the sum of their effects explain only a small fraction of the estimated heritability for most phenotypes. The gap between $h^2_{g_{GWAS}}$ and $h^2$ is called the “missing heritability” and several explanations for this difference have been posited, including upward bias in estimates of $h^2$. The objective of this work is to develop a method for estimating $h^2$ that (1) does not require closely related individuals, (2) can be applied to both quantitative and case-control phenotypes, and (3) is able to localize narrow-sense heritability to individual chromosomes or other genomic segments.

Current approaches to heritability estimation proceed by phenotyping many closely related individuals with a known genetic relationship, such as monozygotic (MZ) and dizygotic (DZ) twins. Yang et al. avoided the use of related individuals by applying linear mixed models to estimate the heritability explained by genotyped SNPs ($h_g^2$). $h_g^2$ corresponds to the fraction of phenotypic variation that could be captured by $h_{g_{GWAS}}^2$ under an additive model if GWAS sample sizes were infinitely large. While current estimates of $h_g^2$ are often much larger than $h_{g_{GWAS}}^2$, they are typically only slightly more than half of $h^2$. One reason $h_g^2$ is less than $h^2$ is because $h_g^2$ does not include the contribution of variants poorly tagged by the genotyping platform, such as rare variants. Another reason for the difference in heritability estimates is that existing methods for estimating $h^2$ can be biased, since they rely on related individuals. As a result, epistatic interactions between SNPs, gene environment interactions, and the shared environmental factors of related individuals can all lead to inflated estimates of $h^2$. We recently showed that by jointly using related and unrelated individuals it is possible to obtain less biased estimates of $h^2$. However, the joint fit will
still lead to inflated estimates of $h^2$ in the presence of shared environment\textsuperscript{11}, and cannot be applied to case-control phenotypes.

In this work we propose a new approach for estimating $h^2$, which takes as input the phenotypes and genotypes of admixed individuals such as African Americans. We show via analytical derivation as well as extensive simulation over both simulated and real genotype data that heritability explained by local ancestry ($h_γ^2$) is related to the total narrow sense heritability $h^2$ via the equation $h_γ^2 = 2F_{STC}(1-\theta)h^2$, where $F_{STC}$ is a specific measure of weighted allele frequency differences between ancestral populations at causal loci (see Online Methods) and $\theta$ is the fraction of European ancestry\textsuperscript{14,15}. Since our approach does not use closely related individuals it is free from bias due to epistasis, gene environment interactions, and shared environment effects. Unlike previous work in which $h^2$ estimates could not be obtained for case-control phenotypes\textsuperscript{11}, our current approach can obtain estimates of $h^2$ for both quantitative and case-control phenotypes, achieving goals (1) and (2). Furthermore, unlike previous methods that provide genome-wide estimates, we are able to estimate $h^2$ for a particular genomic region, such as a chromosome, achieving goal (3). Our approach can be applied to all existing and future GWAS of admixed populations, without requiring additional expensive and time-consuming collections of large numbers of MZ and DZ twins.

We applied this approach to 21,497 African Americans from the NHLBI CARe, WHI-SHARE, and AAPC projects, analyzing 12 quantitative phenotypes and 1 case-control phenotype. For height and BMI, we obtained $h^2$ estimates of $0.55 \pm 0.09$ and $0.23 \pm 0.06$, respectively, which are larger than estimates of $h_g^2$ in these and other data sets but smaller than twin-based estimates of $h^2$, consistent with inflation in twin-based estimates because of shared environment or epistasis. We also estimated the heritability of height for each chromosome and found a significant correlation between chromosome length and heritability (p-value < 0.003).

**Results**

**Overview of method**

We consider three approaches to estimating heritability for a phenotype with a narrow-sense heritability of 80%. First, the classic approach to estimating heritability is to divide the phenotypic covariance of related individuals by the fraction of the genome they share IBD\textsuperscript{13}. In this instance, the phenotypic covariance of pairs of related individuals will be 0.80 times the fraction of genome shared IBD (Figure 1a). The second approach, developed by Yang et al.\textsuperscript{10}, is to estimate the genetic relationship of unrelated individuals over genotyped SNPs and applied a linear mixed model with the genetic relationship matrix to estimate phenotype. To illustrate this approach we simulated 2 million independent pairs of individuals, regressing their normalized genetic similarity over the product of their normalized phenotypes giving a regression coefficient of $0.79\pm0.014$ (Figure 1b). This Haseman-Elston regression\textsuperscript{16} shows how genetic similarity of unrelated individuals can be used to estimate heritability of genotyped SNPs ($h_g^2$). In general, the heritability explained by genotyped SNPs is less than the total narrow-sense heritability ($h^2$) since phenotypic variation determined by poorly tagged SNPs such as rare variants will not be captured\textsuperscript{10}.
The approach used in this work is similar to that of Yang et al.\textsuperscript{10}, but instead of using genotypes to estimate genetic similarity we use the number of copies of local ancestry in an admixed population. A crucial element of our approach is that the phenotypic variation described by variation in local ancestry ($h^2_\gamma$) is a function of all causal variation, not just that tagged by SNPs on the genotyping platform. This is because local ancestry tags both common and rare variation. To illustrate this approach we simulated 4 million unrelated admixed individuals from ancestral populations with genetic distance $F_{STC} = 0.08$ and an equal proportion of ancestry from each ancestral population $\theta = 0.5$ (see Online Methods). Applying Haseman-Elston regression to regress the product of normalized phenotypes against genetic similarity of local ancestry, we observe a regression coefficient $0.033 \pm 0.007 \approx 2 F_{STC} \theta (1-\theta) h^2 = 0.032$, corresponding to $h^2 = 0.83$ (s.e. = 0.18) (Figure 1c). The Haseman-Elston regression used in generating these figures is for illustrative purposes (as in Figure 3 of \textsuperscript{10}). In practice, we use a mixed model approach due to its lower standard errors\textsuperscript{10}.

We first construct a local ancestry based kinship matrix $K_\gamma$, which is constructed similarly to the genotype-based kinship matrix $K$ in previous methods\textsuperscript{10}, but with local ancestry substituted for genotypes at each SNP. We use a variance components approach to estimate the phenotypic variance explained by variation in local ancestry ($\sigma^2_\gamma$) and the residual phenotypic variance ($\sigma^2_e$)\textsuperscript{10,17}. We included genome-wide ancestry proportion $\theta$ and the top five principal components as fixed effects when fitting the mixed model (see Online Methods). The heritability explained by local ancestry is given by $h^2_\gamma = \frac{\sigma^2_\gamma}{\sigma^2_\gamma + \sigma^2_e}$. Finally, to estimate $h^2$, we use the formula $h^2 = 2 F_{STC} \theta (1-\theta) h^2$, where $F_{STC}$ is a specific measure of weighted allele frequency differences between ancestral populations at causal loci (see Online Methods). For dichotomous phenotypes we applied the same approach, but converted the observed scale estimates to a liability scale estimate of heritability using \textsuperscript{18}, and the published disease prevalence in African Americans. In our previous work\textsuperscript{11}, this conversion was not possible because non-randomly ascertainment individuals in multiple relatedness classes (e.g. siblings, first cousins, avuncular) were studied, and there is currently no method for accounting for ascertainment in such complex pedigrees. A complete description of the approach, along with an analytical derivation, is given in Online Methods.

**Simulations with Simulated Genotypes**

We first verified the analytical derivations and examined the properties of the approach under a simple simulation framework. We simulated the genotypes and local ancestry of 4,000 unrelated diploid individuals at 1,000 SNPs from a two-way admixed population with causal variant genetic distance $F_{STC}$, and either normally or uniformly distributed ancestry proportion $\theta$. Each local ancestry segment contained exactly one SNP and all segments were generated independently. Phenotypes were simulated under an additive model with heritability $h^2$ in which a proportion $r$ of the 1,000 SNPs was causal (see Online Methods). We applied our method to estimate heritability over a range of values of $F_{STC}$, $\theta$, $r$, and $h^2$. For each parameter setting we estimated heritability from 2,000 independent simulated data sets. The results shown in Table 1 show that our heritability estimates are accurate across a
range of parameter settings, confirming our analytical derivation. Results for additional parameter settings are shown in Supplementary Table 1.

The results also demonstrate the relationship between $h^2_\gamma$ and the parameters $F_{STC}$, $\theta$, and $h^2$. For a fixed value of $r$, phenotypes with a larger $h^2$ will have larger genetic effects resulting in larger $h^2_\gamma$. When ancestral populations are genetically distant (larger $F_{STC}$), variants are more likely to have a different frequency in the ancestral populations resulting in a concomitant increase in $h^2_\gamma$. Increasing the variance of $\theta$ results in a larger standard error around the heritability estimates.

**Simulations with Real Genotypes**

We made several simplifying assumptions in the above simulations that do not hold in real data sets. These include a single SNP per ancestry block, no genotyping error, no local ancestry inference error, no LD, a normal or uniform distribution of ancestry proportion, continuous phenotypes, and that the effect size distribution of common and rare variants used in computing $F_{STC}$ was identical. To address these complexities, we took the approach of using real genotypes and simulating phenotypes. We simulated continuous and case-control phenotypes over 5,129 individuals (excluding close relatives) from the CARe cohort (see Online Methods). Although phenotypes were generated from SNPs sampled across all genotyped SNPs, we only used local ancestry information from every 5th SNP.

We tried a range of parameters for $h^2$. Instead of simulating phenotypes under an infinitesimal model, we sampled a proportion of causal variants $r$. We could not alter ancestry proportion $\theta$, since this is fixed in the real data set. However, we altered the effect size distribution of SNPs according to their value of $F_{STC}$.

The data did not contain a sufficient number of genotyped variants that were rare in both populations to simulate rare versus common effects. Instead we examined SNPs common in both populations (common) vs. SNPs rare in at least one population (uncommon). Only common variants were used in constructing the kinship matrix, and so uncommon variants will only contribute to $h^2_g$ via LD. The common SNPs had an $F_{STC}$ of 0.15, while the uncommon SNPs had an $F_{STC}$ of 0.25. We simulated phenotypes with a different proportion of phenotypic variance from uncommon variants ($\alpha$). When $\alpha$ is different from 0, the kinship matrix variant and causal variant frequencies are different. The results in Table 2 show that simulations involving a large proportion of causal variants not included in the kinship matrix (high $\alpha$) had a lower value of $h^2_g$ than $h^2$ because the common variants did not completely capture the phenotypic variance driven by the uncommon variants. The parameter $\alpha$ also determines the study wide $F_{STC}$ according to $F_{STC} = (0.15(1-\alpha) + 0.25\alpha)$ (see Online Methods). The results shown in Table 2 use the correct value of $\alpha$, and hence the estimates of $h^2$ are unbiased. However, if we incorrectly assume that $\alpha=0$ when it does not, then $h^2$ will be biased by factor of $0.15(1-\alpha) + 0.25\alpha)/0.15$. We describe this (and other potential sources of bias) in detail in the Discussion.

Setting individuals with the lowest $P\%$ of phenotypes as cases and all other as controls generated dichotomous phenotypes with prevalence $P$. The small number of individuals

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prevented simulation of case-control ascertainment, which may produce downward bias for low prevalence diseases in very large studies (see Supplementary Table 9 in [19]). Those biases are expected to be small in the prostate cancer data analyzed here because of the high prevalence of prostate cancer and moderate sample size. For large sample sizes, replacing mixed model based estimates with Haseman-Elston regression estimates will alleviate the issue of ascertainment bias.

The results in Table 2 also demonstrate that complexities such as genotyping error, LD, or errors in local ancestry inference in African Americans do not introduce bias into the heritability estimates when phenotypes are generated under a non-infinitesimal mixture model. This may not be the case for other admixed populations such as Latinos[21] (see Discussion).

**Application to WHI, CARE, and AAPC cohorts**

We applied our method to 21,497 African-American individuals from the WHI, CARE, and AAPC cohorts over a total of 12 quantitative phenotypes and 1 case-control phenotype (see Online Methods). Local ancestry was inferred using the HAPMIX, SABER+, and RFMix methods, which are extremely accurate in African Americans ($r^2=0.98$ or greater)[22,23,24].

For each phenotype we estimated $h_g^2$, $h_e^2$ and by extension $h^2$. For $h_g^2$ and $h_e^2$ we used the GCTA software package applied to the genotypes and local ancestry at each SNP respectively[17]. For those phenotypes measured in both cohorts we compute the inverse variance-weighted mean and standard error. For each phenotype we also list previously published estimates of heritability from family studies using twins and African-American estimates where available ($h_{pub}^2$). The results are shown in Table 3, and published African-American estimates are marked for reference. Estimates from European populations may not be directly comparable if the genetic or environmental bases for the phenotype differ substantially.

Several phenotypes, including height, BMI, HDL, TG, PC, and WBC (conditioned on ancestry at the Duffy antigen locus FY; see below), had $h^2$ estimates lower than family-based estimates. This could be due to the phenotype-specific effects of epistasis, gene environment interaction, and/or shared environmental factors that can inflate family based estimates[12,13]. In our recent work using an extended genealogy inclusive of more distantly related individuals we also found height and BMI estimates lower than previous heritability estimates, providing further evidence of inflation[11]. The lower estimates could also reflect a difference in the heritability between African Americans and the previous study populations. There were no statistically significant differences in $h^2$ estimates between the cohorts.

Yang et al. proposed an adjustment to account for the incomplete coverage of genotyping platforms[10]. We applied this approach in the CARE data (see Supplementary Table 3), and observed an increase in $h_g^2$ of less than 1% in all phenotypes. We include genome-wide ancestry proportion as a fixed effect in our mixed model. If there exists an environmental factor that affects phenotype and is correlated with ancestry, our heritability estimates will discount this environmental effect leading to higher estimates of heritability. Specifically, it will remove the variance of the environmental factor that can be explained by ancestry from...
the environmental component ($\sigma^2_e$) in the denominator of the heritability estimate
\[ h^2 = \frac{\sigma^2_g}{\sigma^2_g + \sigma^2_e} \] (see Online Methods).

Differences between our heritability estimates and those of previous studies can also be due to differences between the value of $F_{ST C}$ we used in this study and the true value of $F_{ST C}$ for the phenotype in question. Based on recent evidence that rare variants unlikely to contribute to a large proportion of phenotypic variation\textsuperscript{25,26}, we computed an $F_{ST C}$ of 0.182 over the common variants (MAF > 5%) in African-Americans. However, this estimate drops to 0.165 for low-frequency variants (MAF < 5%) and 0.054 for rare variants (MAF < 1%). Estimates of heritability assuming a rare variant only phenotype model would be more than three times as large as from a common variant only phenotype model. Therefore, if rare variants contribute substantially to phenotypic variation or if balancing or negative selection constrained the genetic distance at causal variants, then our estimates of heritability will be biased downward (see Discussion).

Positive selection acting at causal variants could induce such a bias in $F_{ST C}$, and we included WBC as a positive control for this type of bias. A SNP in the DARC gene (FY, Duffy null allele) is highly differentiated between CEU and YRI, likely due to its protective effect against vivax malaria\textsuperscript{27}. It is also a SNP of large effect size for WBC\textsuperscript{28}. Therefore, the average $F_{ST C}$ at causal variants for WBC, is much higher than the value 0.182 estimated from common variants (see Online Methods). The $h^2$ estimate of WBC is 3.42 due to the effect of this positive selective pressure. Ancestry at the FY locus accounts for \sim 20% of the phenotypic variation in WBC\textsuperscript{28}. By including ancestry at FY as a fixed effect (WBC|FY) we obtain an $h^2$ estimate of 0.19, which is lower than the published estimate of 0.48.

We perform a sensitivity analysis to assess whether this type of bias is likely to be problematic. Since strong positive selection is unusual\textsuperscript{29}, we consider a single locus under positive selection. We estimate bias as a function of $F_{ST C}$ at the locus and the variance explained by the locus. The results in Supplementary Table 4 show that only for extreme values of both locus $F_{ST C}$ and heritability will there be significant bias in heritability due to positive selection. As an example we consider the 8q24 locus in prostate cancer, which contains causal SNPs that are highly differentiated SNPs between African and European ancestors, producing an admixture-mapping peak\textsuperscript{30}. However, because this locus explains less than 2% of the heritability of prostate cancer, even exceedingly strong population differentiation at this locus will not substantially bias our overall results.

**Partitioning heritability across the genome**

Our method is also capable of estimating the total narrow sense heritability attributable to a particular genomic region. This is accomplished by constructing the kinship matrix using just those ancestry segments in the region of interest and applying the variance component model to the phenotype of interest using the region-specific kinship matrix (see Online Methods). We partitioned the heritability for each of the phenotypes from the CARe data set across each of the chromosomes\textsuperscript{31}. We applied weighted linear regression to determine the relationship between heritability and chromosome length (see Online Methods). The results
for height are presented in Figure 2 and the full results are provided in Supplementary Table 2. We find a strong correlation between chromosome length and the heritability of height (Pearson correlation = 0.513, weighted p-value = 0.0028). logHDL, BMI, and SBP, also produced significant results (weighted p-value < 0.03, 0.02, 0.02 respectively). Other phenotypes had standard errors too large to produce meaningful results. To address this, we averaged the heritability from each chromosome across all phenotypes (using WBC|FY instead of WBC) and we observed a significant correlation between chromosome length and mean chromosomal heritability (Pearson correlation=0.686, weighted p-value <0.0002).

Discussion

We developed a method for estimating narrow sense heritability from unrelated individuals by leveraging the two ancestral genomes in recently admixed populations, such as African Americans. We used a population genetic approach to derive the relationship between heritability and variation in local ancestry in admixed populations. Theory and simulations confirm that under an infinitesimal phenotypic model our approach produces unbiased estimates of heritability. Since the individuals are distantly related, our approach will not produce heritability estimates inflated by epistasis, gene environment interactions, or shared environmental effects.

Our method is also able to partition total narrow-sense heritability ($h^2$) along genomic segments such as chromosomes, as we have shown by application to the phenotypes in the CAREe data set. This is distinct from recent work that instead partitioned the heritability explained by genotyped SNPs ($h_g^2$) across chromosomes\textsuperscript{31-33}. While a previous method has also partitioned $h^2$ along chromosomes\textsuperscript{34,35}, it relies on the use of siblings, leading to very large standard errors, and is limited by the coarseness of shared IBD segments (which extend for tens of megabases). Our approach is limited by the coarseness of local ancestry segments (which extend for megabases) and thus cannot be applied at the level of individual genes.

We applied our method to an African-American population in this study. Application to more complex admixed populations such as Latinos will have to account for the reduced accuracy in local ancestry inference\textsuperscript{21} to avoid downward bias. Restricting to two ancestry categories (e.g. Native American vs. non-Native American ancestry)\textsuperscript{36} is one approach to handle multi-way admixture, but it may be possible to extend our derivation to multi-way admixture. There is evidence that African Americans have a small proportion of admixture from Native American populations (0.5%)\textsuperscript{24}, but this very small proportion is unlikely to significantly change our results. Substantial errors in the assumed population genetic structure would perturb the values of $F_{STC}$ and $\theta$, and resulting $h^2$ estimates would be biased in proportion to these errors. Application to sex chromosomes can be adapted from the approach taken in\textsuperscript{31}, but must be analyzed separately due to the differences in admixture proportion of European ancestry on autosomes and sex chromosomes.

In our previous work we found that heritability estimates from related individuals followed a pattern consistent with biases due to shared environment\textsuperscript{11}. In this work we found that a linear additive model, implicitly including both rare and common variants, typically
explained less phenotypic variation than that predicted in family studies. These new estimates of narrow-sense heritability are less susceptible to bias and provide additional evidence that family based estimates are inflated. Unlike [11], we were able to obtain estimates for both quantitative and case-control traits. We also found that chip based additive models explained less phenotypic variation than our estimates. In the meta-analyzed phenotypes common to CARe and WHI the average of these estimates were 24.7% and 31.1% respectively. Rare variants and poorly tagged common variants are the most likely explanation for the difference between these two estimates. We discuss other possible explanations below.

Our method does produce biased estimates when model assumptions are violated. Specifically, if the genetic distance we estimated over common variants (0.182; see Online Methods) differs from the distribution over causal variants, our method can be either inflated or deflated. If selection were acting on the causal variants their $F_{STC}$ could be higher or lower depending on the direction of selection. In the case of positive selection in one of the ancestral groups but not the other, the true value of $F_{STC}$ will be larger than our genome-wide estimate and so our $h^2$ estimate will be inflated. For example, estimates for white blood cell count were larger than $h^2_{pub}$ due to strong selective pressure at the Duffy locus[27,37]. However, strong positive selection is believed to be rare in recent human evolution[29]. If a large proportion of phenotypic variance is due to rare variants then incorrect estimates of $F_{STC}$ may induce bias. However, previous reports suggest that rare variation explains a small proportion of total heritability[25,26].

The application of our approach to two large cohorts of African Americans revealed a difference between previously published family-based estimates of the heritability of height and BMI and our estimates. This suggests that there is a significant contribution of non-additive genetic effects or shared environmental effects that differ between MZ and DZ twins. The future application of our method to large-scale studies of African Americans will both provide a mechanism of estimating the total narrow sense heritability of phenotypes as well as determining the genetic architecture of complex phenotypes.

### Online Methods

Given a set of $M$ admixed individuals with two ancestral populations ($P_0$ and $P_1$), let the local ancestry for individual $i$ at SNP $s$, $y_{is} \in \{0,1,2\}$, be the number alleles inherited from a $P_1$ ancestor. We use a mixed model approach to estimate $h^2_g$ the contribution of variation in local ancestry to phenotypic variation for the phenotype $Y=y_1, y_2, \ldots y_M$. We first construct a local ancestry based kinship matrix $K_\gamma$, which is constructed similarly to the genotyped-based kinship matrix $K$, but with local ancestry substituted for genotypes at each SNP. We then find the parameters $\sigma^2_\gamma$ and $\sigma^2_\epsilon$ which maximize the likelihood of the mixed model $Y \sim N(0, K_{\gamma}\sigma^2_\gamma + I\sigma^2_\epsilon)$. The heritability explained by local ancestry is given by $h^2_{\gamma}$. Finally, we use the formula $h^2_g = h^2_{\gamma} / 2F_{STC}(1 - \theta)$ to estimate $h^2$. 

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**Definition of \( h^2 \)**

Heritability is the ratio of genetic variance to the sum of genetic and environmental variance:

\[
h^2 = \frac{\sigma_g^2}{\sigma_g^2 + \sigma_e^2}.
\]

In this case we are defining these elements with respect to an admixed population. For a given phenotype, both \( \sigma_g^2 \) and \( \sigma_e^2 \) can vary between the ancestral European and African populations. For example, \( \sigma_g^2 \) will vary with ancestry if the minor allele frequency at causal variants is systematically larger in one of the two populations. It is also possible for ancestry to be associated with environmental factors. In this case, by conditioning on genome-wide ancestry, our method will remove the environmental variance that can be explained by ancestry and estimate the heritability of the component of phenotype that cannot be predicted by genome-wide ancestry, thereby increasing the heritability estimate.

**Estimation of \( h^2 \)**

We use a variance components approach to determine the phenotypic variance described by local ancestry \( h^2 \) using \( \theta \) as a fixed effect to prevent confounding from environmental factors associated with ancestry. This method is equivalent to recent methods used to determine the phenotypic variance described by genotyped SNPs (\( h^2_g \)), replacing genotypes with inferred local ancestry\(^{10} \).

**Derivation of relationship between \( h^2 \) and \( h^2 \)**

Let \( i \) denote (diploid) individuals and \( s \) index SNPs. Individual \( i \) is assigned global ancestry proportion \( \theta_i \) from some distribution \( F(\cdot) \) with mean \( E[\theta_i] = \theta \) and variance \( \sigma^2_{\theta} \). Given \( \theta_i \), an individual is assigned maternal and paternal local ancestries \( \gamma_{i,s,M} \) and \( \gamma_{i,s,P} \) at each SNP (0 or 1 copies of European ancestry), from Bernoulli distribution \( \text{Ber}(\theta_i) \). Given local ancestries \( \gamma_{i,s,M}, \gamma_{i,s,P} \) and allele frequencies \( p_{s,0}, p_{s,1} \) at SNP \( s \) in populations 0 and 1, individuals are assigned maternal genotypes \( g_{i,s,M} = \gamma_{i,s,M} Z_{i,s,1} + (1 - \gamma_{i,s,M}) Z_{i,s,0} \) where \( Z_{i,s,0} \sim \text{Ber}(p_{s,0}) \) and \( Z_{i,s,1} \sim \text{Ber}(p_{s,1}) \), and similarly for paternal genotypes. The diploid genotype \( g_{i,s} = g_{i,s,P} + g_{i,s,M} \) (0, 1 or 2), and the diploid local ancestry \( \gamma_{i,s} = \gamma_{i,s,P} + \gamma_{i,s,M} \) (0, 1 or 2).

We define \( E[g_{i,s}] = \mu_{g,s} \) and \( \text{Var}[g_{i,s}] = \sigma^2_{g,s} \) and the normalized genotype \( \overline{g}_{i,s} = \frac{g_{i,s} - \mu_{g,s}}{\sigma_{g,s}} \), where

\[
\mu_{g,s} = 2(\mu p_{s,1} + (1 - \mu) p_{s,0}) \quad (1)
\]

\[
\sigma^2_{g,s} = 2[\mu(1 - \mu)(p_{s,1} - p_{s,0})^2 + (\mu p_{s,1} (1 - p_{s,1}) + (1 - \mu)p_{s,0}(1 - p_{s,0})] \quad (2)
\]

Similarly, we define \( E[\gamma_{i,s}] = \mu_{\gamma} \) and \( \text{Var}[\gamma_{i,s}] = \sigma^2_{\gamma} \) and the normalized local ancestry at each locus \( \overline{\gamma}_{i,s} = \frac{\gamma_{i,s} - \mu_{\gamma}}{\sigma_{\gamma}} \), where

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Although though equation (4) may not be strictly true (e.g. in a population where all individuals have 1 European parent and 1 African parent), it is approximately true for African Americans\textsuperscript{22}. Furthermore, $\sigma^2_\gamma$ can be estimated empirically, and we do so in this work. We model the phenotype of individual $i$ as

$$y_i = \sum_s \beta_s \bar{y}_{i,s} + \varepsilon_i \quad (5)$$

where $\varepsilon_i \sim N(0, \sigma^2_\varepsilon)$, $\text{Var}[y_i] = 1$, $E[y_i] = 0$, the effect size of SNP $s$ is $\beta_s$, and $h^2 = \sum_s \beta_s^2$. By substitution and algebra we get

$$\bar{y}_{i,s} = \frac{\theta_i - \mu_{s,0}}{\sigma_{s,\gamma}} = \frac{1}{\sigma_{s,\gamma}} \left( \sigma_{s,\gamma} \gamma_{i,s} - 2 \mu p_s - 2(1 - \mu)p_s0 \right) + \frac{1}{\sigma_{s,\gamma}} \theta (Z_{i,s,1} - Z_{i,s,0}) + \frac{1}{\sigma_{s,\gamma}} \theta (Z_{i,s,1} - p_s) + \frac{1}{\sigma_{s,\gamma}} \theta (Z_{i,s,0} - p_s)$$

$$= \frac{\sigma_{s,\gamma}}{\sigma_{s,\gamma}} \gamma_{i,s} (Z_{i,s,1} - Z_{i,s,0}) + \frac{2}{\sigma_{s,\gamma}} \theta (Z_{i,s,1} - p_s0) + (1 - \theta)(Z_{i,s,0} - p_s)$$

Plugging into equation (5), we get

$$y_i = \sum_s \beta_s \bar{y}_{i,s} + \varepsilon_i$$

$$= \sum_s \beta_s \sigma_{s,\gamma} \gamma_{i,s} (Z_{i,s,1} - Z_{i,s,0}) + \sum_s \beta_s \frac{2}{\sigma_{s,\gamma}} \theta (Z_{i,s,1} - p_s0) + (1 - \theta)(Z_{i,s,0} - p_s0) + \varepsilon_i$$

$$= \sum_s \beta_s \sigma_{s,\gamma} \gamma_{i,s} (Z_{i,s,1} - Z_{i,s,0}) + \delta_i \quad (7)$$

Note that $\delta_i$ does not depend on local ancestry, which allows us to compute the heritability due to local ancestry $h^2_{\gamma}$ as:

$$h^2_{\gamma} = \text{Var}[E[y_i | \gamma_{i,1}, \ldots, \gamma_{i,N}]]$$

$$= \text{Var}[\sum_s \beta_s \sigma_{s,\gamma} \gamma_{i,s} (p_s1 - p_s0)] \approx \sum_s [\beta_s \sigma_{s,\gamma} (p_s1 - p_s0)]^2$$

$$= 2\theta (1 - \theta) \sum_s [\beta_s \sigma_{s,\gamma} (p_s1 - p_s0)]^2$$

We define $F_{\text{STC}}$ as a measure genetic distance between ancestral populations weighted by the square of effect size $\beta$:

$$F_{\text{STC}} = \sum_s \frac{\beta_s^2 (p_s1 - p_s0)^2}{\sigma_{s,\gamma}^2}$$

This results in a final relationship

$$\mu_{\gamma} = 2\theta \quad (3)$$

$$\sigma^2_{\gamma} = 2\theta (1 - \theta) \quad (4)$$
In practice we do not know the effect size of every SNP and must make simplifying assumptions about their distribution in order to estimate $F_{STC}$. First consider a simple phenotypic model in which genotypic effect size $\beta_s$ is independent of $p_{s,o}$ and $p_{s,1}$. Then

$$h^2 = \sum_s \beta_s^2 \approx NE\{\beta_s^2\},$$  \hspace{1cm} (11)

where $N$ is the number of SNPs. Then equation (8) becomes

$$h^2 = 2\theta(1 - \theta)NE\{\beta_s^2\}E\left[\frac{(p_{s,1} - p_{s,0})^2}{\sigma_{g,s}^2}\right]$$
$$\quad = h^2 2\theta(1 - \theta)E\left[\frac{(p_{s,1} - p_{s,0})^2}{\sigma_{g,s}^2}\right]$$  \hspace{1cm} (12)

The $F_{STC}$ in equation (12) is a genome-wide measure of genetic difference between the ancestral populations. This is related to the classic parameter $F_{ST}$ when all variants are causal (i.e. the infinitesimal model).

$$F_{ST} = E\left[\frac{(p_{s,1} - p_{s,0})^2}{\sigma_{g,s}^2}\right]$$  \hspace{1cm} (13)

Now consider a more complex model in which the effect size of SNPs can fall into one of $L$ classes such that the effect size distribution is a function of the class $L$. These classes could be, for example, rare and common variants (used in this work). We defined the genetic distance between ancestral populations within each class as $F_{STL}$ and the phenotypic variance explained by SNPs in this class as $h^2_L$. Again substituting into equation (8) we have,

$$h^2 = 2\theta(1 - \theta)F_{STC}h^2 \sum_{s \in L} \left[\frac{\beta_s^2(p_{s,1} - p_{s,0})}{\sigma_{g,s}^2}\right]$$
$$\quad = 2\theta(1 - \theta)\frac{h^2}{h^2} \sum_{L} h^2_L F_{STL}$$  \hspace{1cm} (14)

Therefore $F_{STC} = \sum \frac{h^2}{h^2} F_{STL}$, a weighted measure of genetic distance in each class.

To obtain an estimate of $h^2$ we must estimate $\theta$, $F_{STC}$, and $h^2_L$. The parameter $\theta$ is estimated from local ancestry inference. The parameter $F_{STC}$ is estimated from assumptions about the variance explained by SNPs in each genotypic class combined with external reference panels.\textsuperscript{45,46}
**Definition and Estimation of F_{STC}**

As shown in the equations above we are defining $F_{ST}$ to be the weighted average (across all SNPs $s$) of ratios $(p_{s,1} - p_{s,0})^2 / \hat{\sigma}_{g,s}^2$. While this is similar to standard versions of $F_{ST}$, a ratio of averages is recommended instead when the goal is to draw population genetic inferences. If the distribution of SNPs effect sizes is not a function $F_{ST}$ then this would be the appropriate definition for our heritability estimation approach. However, recent work has shown that rare variants are unlikely to contribute to a large proportion of phenotypic variation. As has been reported previously, the average of ratios estimate will shrink when including many rare variants in the estimate. This is reflected in the 1000 Genomes based estimate of $F_{ST}=0.07$, which used an average of ratios. Therefore, $F_{ST}$ will produce a biased estimate of heritability because for the variance explained by rare variants is different from the variance explained by common variants. To account for this we defined a parameter $F_{STC}$, which is a weighted measure of genetic distance between ancestral populations.

In practice we defined $F_{STC}$ as the average $F_{ST}$ within each class $L$ of SNPs ($F_{STL}$), weighted by the proportion of phenotypic variance explained by that class:

$$F_{STC} = \sum_L \frac{h^2}{h^2} F_{STL} \quad (15)$$

Consider a situation in which $L$ contains two classes, rare and common SNPs, with $F_{ST}$ 0.054 and 0.182 respectively. If rare variants explained 10% of the heritability and common variants explain 90% of the heritability, then $F_{STC}=0.1692$. We estimated $F_{STC}$ over the HapMap3 data set by using CEU and YRI as proxies for the ancestral populations of African-Americans, using an admixture proportion of 18.3% European ancestry, and assuming distribution of causal variant frequencies. We estimated a value of 0.182 assuming causal variant MAF > 5% (which we used in this work), 0.165 assuming MAF < 5%, and 0.054 assuming MAF < 1%.

**Simulations with Simulated Genotypes**

In order to examine the properties of our approach, we first applied our method to data generated under a simple simulation framework for generating genotypes, local ancestries, and phenotypes of individuals from an admixed population. Allele frequencies $p_{A1}, p_{A2}, \ldots, p_{AN}$ of $N$ SNPs from an ancestral population were drawn uniformly from $[0.1-0.9]$. Allele frequencies of SNPs from $P_0$ were drawn from a beta distribution with parameters $p(1 - F_{STC}) / F_{STC}$ and $(1-p)(1 - F_{STC}) / F_{STC}$ for each SNP $s$, and similarly for $P_1$. The parameter $F_{STC}$ determines the genetic distance between the two populations. The global proportion of $P_0$ ancestry $\theta_1, \theta_2, \ldots, \theta_M$ for each of $M$ individuals was drawn either uniformly from $[0.4,0.6]$, from the normal distribution $\mathcal{N}(0.5,0.1)$, or fixed at 0.5. Local ancestry for individual $i$ at SNP $s$ ($\gamma_{is}$), was generated by two draws from binomial distribution with parameter $\theta_s$. The genotypes from individual $i$ at SNP $s (g_{is})$ were then generated by drawing from the binomial distributions with allele frequencies specified by the local ancestry for each SNP.
that individual at that SNP. That is, if the individual had two copies of ancestry from \( P_0 \) at SNP \( s \) then two draws from a binomial with parameter \( p_{0s} \) were used. To create a phenotype we first selected \( Nr \) causal variants where \( r \) is the proportion of causal variants. Effect sizes were drawn from the normal distribution \( N(0, h^2/(Nr)) \) and the genetic element of the phenotype was generated by taking the inner product of the causal variants, normalized to have mean 0 and variance 1, and the effect sizes for the variants. Normally distributed random noise was added such that the total heritability in the population was \( h^2 \).

**Simulations with Real Genotypes**

We split the genotypes from 5,129 distantly related CARe individuals into two groups. The *common* group contained those SNPs with MAF > 5% in both CEU and YRI. The *uncommon* group contained all other SNPs (i.e. MAF < 5% in either or both of CEU and YRI). The genotype kinship matrix \( K \) was constructed over the common SNPs and the local ancestry kinship matrix \( K_\gamma \) was constructed using the local ancestry called at every 5th common SNP.

We simulated a phenotype by first selecting a proportion \( r \) of causal variants at random from the *common* and *uncommon* SNPs, leaving \( N_c \) *common* causal and \( N_n \) *uncommon* causal SNPs. We then selected a fraction of phenotypic variance \( \alpha \) explained by the *uncommon* SNPs. At \( \alpha=0.0 \) *uncommon* variants had no effect and the genetic basis of the phenotype was entirely determined by *common* variants. We then chose effect sizes for each *common* and *uncommon* SNP by drawing from normal distributions \( N(0, (1-\alpha)h^2/(N_c)) \) and \( N(0, (\alpha)h^2/(N_n)) \) respectively. The genetic element of the phenotype was generated by taking the inner product of the causal variants, normalized to have mean 0 and variance 1 in the admixed population, and the effect sizes for the variants. Normally distributed random noise was added such that the total heritability in the population was \( h^2 \). The \( F_{STC} \) of the *common* and *uncommon* SNPs was 0.15 and 0.25 respectively. The study \( F_{STC} \) used to estimate heritability was the weighted mean 0.15(1-\( \alpha \)) + 0.25\( \alpha \) as described in the derivation above. Setting individuals with the lowest \( P\% \) of phenotypes as cases and all other as controls generated dichotomous phenotypes with prevalence \( P \).

**Data set approvals**

The CARe project has been approved by the Committee on the Use of Humans as Experimental Subjects (COUHES) of the Massachusetts Institute of Technology, and by the Institutional Review Boards of each of the nine parent cohorts.

The WHI project has been approved by the Human Subjects Committees at the WHI Clinical Coordinating Center (FHCRC) and at the 40 WHI Field Centers.

The AAPC project has been approved by the Institutional Review Board of the University of Southern California. The 11 studies contributing to the AAPC each received approval for the use of specimens from their patients.
CARe data set

Affymetrix 6.0 genotyping and QC filtering of African-American samples from the CARe cardiovascular consortium was performed as described previously\textsuperscript{50}. After QC filtering for each of ARIC, CARDIA, CFS, JHS and MESA cohorts and subsequent merging, 8,367 samples and 770,390 SNPs remained. To limit relatedness among samples we restricted all analyses to a subset of 5,129 samples in which all pairs have genome-wide relatedness of 0.05 or less and had between 5% and 45% European ancestry. We performed local ancestry inference using the HAPMIX software with the CEU and YRI HapMap populations as reference ancestral populations. We examined seven phenotypes from the CARe cohort, height, body mass index (BMI), log transformed high density lipoprotein cholesterol (logHDL), low density lipoprotein cholesterol (LDL), white blood cell count (WBC), diastolic blood pressure (DBP), and systolic blood pressure (SBP). For each phenotype we included age, sex, study center, proportion of European ancestry, and the top 5 principal components as fixed effects. A detailed description of the phenotypes can be found here\textsuperscript{51}.

WHI Data Set

Affymetrix 6.0 genotyping and QC filtering of African-American samples from the Women’s Health Initiative (WHI) SNP Health Association Resource (SHARe) was performed as described previously\textsuperscript{52}. The dataset includes extensive phenotypic and genotypic data on 12,008 African American and Hispanic women aged 50-79 enrolled in one or more components of the WHI program. We included only African American samples and to limit relatedness among samples we restricted all analyses to a subset of 8,153 samples in which all pairs have genome-wide relatedness of 0.05 or less. We performed local ancestry inference using the SABER\textsuperscript{+} software with the CEU and YRI HapMap populations as reference ancestral populations. We examined 10 phenotypes from the WHI cohort (BMI), log transformed high density lipoprotein cholesterol (logHDL), low density lipoprotein cholesterol (LDL), white blood cell count (WBC), log transformed triglycerides (logTG), glucose, log transformed insulin (logInsulin), QT interval duration (QT-INTERVAL), C-reactive protein (CRP), diastolic blood pressure (DBP), and systolic blood pressure (SBP). For each phenotype we included age and proportion of European ancestry as fixed effects. A detailed description of the phenotypes can be found here\textsuperscript{52}.

African American Prostate Cancer Data Set (AAPC)

IlluminaHuman1M-Duov3_B genotyping and QC filtering of African-American samples from the African American Prostate Cancer Study (AAPC) from a total of 11 participating studies was performed as described previously\textsuperscript{53,54,55}. The cleaned dataset includes 9,641 African American subjects and 1,001,899 autosomal SNPs. To limit relatedness among samples we restricted all analyses to a subset of 8,215 samples in which all pairs have genome-wide relatedness of 0.05 or less. We performed local ancestry inference using the RFMix\textsuperscript{24} with the CEU and YRI HapMap populations as reference ancestral populations. We examined prostate cancer (PC) outcome for each subject. There were 4207 cases and 4008 controls after QC. Due to the admixture signal at the 8q24 locus\textsuperscript{54}, we also estimated heritability removing 8q24 from the SNPs used to estimate the kinship (PC|8q24). For each
phenotype we included age and the top 10 principal components as fixed effects. For conversion to the liability scale we used a prevalence of 5%.

**Partitioning Heritability across the genome**

To estimate the heritability for a particular genomic segment we compute the genetic relatedness matrix as defined in Yang et al\(^{10}\), replacing genotypic for local ancestry calls, and restricting to just those SNPs contained in the region of interest. Given a partitioning of segments along the genome (in our case 22 segments), it is possible to fit them individually or jointly. We attempted both approaches, but found that the joint fit produced a numerical instability in the optimization algorithm preventing convergence. Thus all results reported for the single chromosome analyses are provided by individual and not joint estimates.

We performed both weighted and standard linear regression to assess the relationship between the heritability explained by a chromosome and the length of the chromosome. The weighted version accounts for the differences in number of SNPs contained in longer and shorter chromosomes and the weighting factor we used was the length of the chromosome in centimorgans.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.
Relationships between genetic distance and phenotype for a trait with heritability = 80%. (a) The phenotypic covariance of pairs of individuals at different expected fractions of genome shared IBD is 0.8*%IBD. (b) Regression of genetic distance estimated from genetic variation against the product phenotypes normalized to have mean 0.0 and variance 1.0 has coefficient 0.79 (se = 0.014). (c) Regression of genetic distance estimated from local ancestry variation against normalized phenotypes has coefficient 0.033 (s.e. = 0.007) $\approx 2F_{STC}(1-\theta)h^2 = 0.032$, corresponding to $h^2 = 0.83$ (s.e. = 0.18).
Figure 2.
Estimated heritability of height for each chromosome in the CARe data set. The numbers adjacent to each point are the chromosomes. We plot the regression line of $h^2$ per chromosome regressed on chromosome length. We find a strong correlation between chromosome length and height heritability (Pearson correlation = 0.513, weighted p-value = 0.0028).
Table 1

Results of local ancestry based heritability estimation from simulated genotypes and simulated phenotypes over a range of population and disease architectures. Mean heritability estimates and standard errors are reported from 2,000 simulations for each choice of parameters.

| $h^2$ | $F_{ST}$ | $r$ | $\hat{h}^2$ |
|-------|---------|-----|---------|
| 0.8   | 0.30    | 1.0 | 0.802(0.003) |
| 0.8   | 0.30    | 0.1 | 0.802(0.005) |
| 0.8   | 0.15    | 1.0 | 0.800(0.005) |
| 0.8   | 0.15    | 0.1 | 0.804(0.006) |
Table 2

Results of heritability simulations over 5,129 African American individuals from the CARe cohort. Average estimates and standard errors of heritability explained from genotyped SNPs ($\hat{h}^2$), and our local ancestry based estimate of heritability explained from all SNPs ($\tilde{h}^2$) are reported from 2,500 simulations for representative choices of 4 parameters: true heritability ($h^2$), proportion of causal variants ($r$), prevalence ($P$) (NA for continuous phenotypes), and proportion of heritability from uncommon variants ($\alpha$).

| $h^2$ | $r$ | $P$ | $\alpha$ | $\hat{h}^2$ | $\tilde{h}^2$ |
|------|------|-----|---------|-------------|-------------|
| 0.8  | 0.01 | NA  | 0.0     | 0.797(0.001) | 0.800(0.004) |
| 0.8  | 0.01 | NA  | 0.0     | 0.797(0.001) | 0.793(0.005) |
| 0.5  | 0.01 | NA  | 0.0     | 0.499(0.001) | 0.498(0.003) |
| 0.5  | 0.01 | NA  | 0.0     | 0.499(0.001) | 0.501(0.004) |
| 0.8  | 0.01 | NA  | 0.25    | 0.689(0.003) | 0.689(0.003) |
| 0.8  | 0.01 | NA  | 0.25    | 0.691(0.002) | 0.691(0.002) |
| 0.8  | 0.01 | NA  | 0.25    | 0.691(0.002) | 0.691(0.002) |
| 0.8  | 0.01 | NA  | 0.50    | 0.625(0.002) | 0.625(0.002) |
| 0.8  | 0.01 | NA  | 0.50    | 0.637(0.003) | 0.637(0.003) |
| 0.8  | 0.01 | NA  | 1.0     | 0.498(0.003) | 0.498(0.003) |
| 0.8  | 0.01 | NA  | 1.0     | 0.498(0.003) | 0.498(0.003) |
Heritability estimates of phenotypes from 21,497 African Americans from the WHI, CARe, and AAPC cohorts. Meta shows the inverse variance weighted meta-analysis for those phenotypes contained in both WHI and CARe data sets. $h^2_{g,pub}$ is the previously published estimates of $h^2_g$ and $h^2_{pub}$ is the previously published estimates of $h^2$ from family studies. Published heritability studies of African Americans are denoted with a *.

### The heritability explained by genotyped SNPs ($\hat{h}^2_g$) in WHI and CARe.

| Phenotype | WHI $\hat{h}^2_g$ | s.e. | CARe $\hat{h}^2_g$ | s.e. | Meta $\hat{h}^2_g$ | s.e. | $\hat{h}^2_{g,pub}$ |
|-----------|------------------|------|-------------------|------|------------------|------|------------------|
| height    | 0.461            | 0.058| 0.378             | 0.029| 0.395            | 0.026| 0.450            |
| BMI       | 0.198            | 0.055| 0.078             | 0.065| 0.148            | 0.042| 0.148*           |
| Log(HDL)  | 0.316            | 0.057| 0.224             | 0.066| 0.277            | 0.043| 0.128*           |
| LDL       | 0.294            | 0.056| 0.156             | 0.067| 0.238            | 0.043| 0.101*           |
| WBC       | 0.725            | 0.051| 0.848             | 0.091| 0.755            | 0.044| 0.211*           |
| WBC|FY     | 0.188            | 0.054| 0.167             | 0.097| 0.183            | 0.047| NA               |
| Log(TG)   | 0.226            | 0.056| NA                | NA   | NA               | NA   | NA               |
| Glucose   | 0.16             | 0.063| NA                | NA   | NA               | NA   | 0.108*           |
| log(Insulin) | 0.086    | 0.051| NA                | NA   | NA               | NA   | 0.098*           |
| QT-interval| 0.251           | 0.098| NA                | NA   | NA               | NA   | NA               |
| Log(CRP)  | 0.295            | 0.056| NA                | NA   | NA               | NA   | NA               |
| DBP       | 0.148            | 0.053| 0.170             | 0.066| 0.157            | 0.041| NA               |
| SBP       | 0.162            | 0.054| 0.189             | 0.066| 0.173            | 0.042| 0.243*           |

### The total narrow sense heritability ($\hat{h}^2$) as derived from $\hat{h}^2_g$ in WHI and CARe.

| Phenotype | WHI $\hat{h}^2$ | s.e. | CARe $\hat{h}^2$ | s.e. | Meta $\hat{h}^2$ | s.e. | $\hat{h}^2_{pub}$ |
|-----------|------------------|------|------------------|------|------------------|------|------------------|
| height    | 0.611            | 0.135| 0.503            | 0.120| 0.550            | 0.090| 0.770*           |
| BMI       | 0.252            | 0.097| 0.208            | 0.085| 0.227            | 0.064| 0.470*           |
| Log(HDL)  | 0.418            | 0.117| 0.470            | 0.146| 0.438            | 0.091| 0.520*           |
| LDL       | 0.395            | 0.116| 0.333            | 0.140| 0.370            | 0.089| 0.530*           |
| WBC       | 3.267            | 0.322| 3.703            | 0.447| 3.415            | 0.261| 0.480*           |
The total narrow sense heritability (\(h^2\)) as derived from \(h^2_{\gamma}\) in WHI and CARe.

| Phenotype  | WHI \(h^2\) s.e. | CARe \(h^2\) s.e. | Meta \(h^2\) s.e. | \(h^2_{\gamma,\text{psh}}\) |
|------------|------------------|--------------------|-------------------|-----------------|
| WBC|FY   | 0.172 | 0.084 | 0.247 | 0.166 | 0.187 | 0.075 | NA |
| Log(TG)   | 0.225 | 0.094 | NA    | NA    | NA    | NA    | 0.40^{0.06} |
| Glucose   | 0.104 | 0.087 | NA    | NA    | NA    | NA    | 0.29^{0.11} |
| log(Insulin) | 0.105 | 0.077 | NA    | NA    | NA    | NA    | 0.28^{0.11} |
| QT-Interval | 0.336 | 0.164 | NA    | NA    | NA    | NA    | 0.41^{0.12} |
| Log(CRP)  | 0.542 | 0.139 | NA    | NA    | NA    | NA    | 0.56^{0.13} |
| DBP       | 0.179 | 0.088 | 0.238 | 0.119 | 0.200 | 0.071 | 0.13^{0.06} |
| SBP       | 0.187 | 0.092 | 0.233 | 0.117 | 0.205 | 0.072 | 0.17^{0.06} |

The complete AAPP results.

| Phenotype  | \(\text{AAPC} h^2\) s.e. | \(h^2_{\gamma,\text{psh}}\) s.e. | \(h^2_{\gamma,\text{psh}}\) s.e. |
|------------|--------------------------|-------------------------------|-------------------------------|
| PC         | 0.182 | 0.040 | NA | 0.328 | 0.093 | 0.58^{0.04} |
| PC|8q24 | 0.174 | 0.040 | NA | 0.315 | 0.092 | NA |
Table 4

Number of individuals for each phenotype in the CARe and WHI data sets. The AAPC data set contained 4207 PC cases and 4008 controls.

| Phenotype   | WHI  | CARe |
|-------------|------|------|
| height      | 8109 | 5024 |
| BMI         | 8153 | 5026 |
| Log(HDL)    | 8014 | 4928 |
| LDL         | 7979 | 4794 |
| WBC         | 8035 | 3367 |
| WBC|FY      | 8035 | 3367 |
| Log(TG)     | 8015 | NA   |
| Glucose     | 6826 | NA   |
| log(Insulin)| 7749 | NA   |
| QT-Interval | 4143 | NA   |
| Log(CRP)    | 8014 | NA   |
| DBP         | 8153 | 5030 |
| SBP         | 8153 | 5029 |