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# Estimating heritability of complex traits in admixed populations with summary statistics — Source link 🗹

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# Estimating heritability and its enrichment in tissue-specific gene sets in admixed populations

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## Abstract

The increasing size and diversity of genome-wide association studies provide an exciting opportunity to study how the genetics of complex traits vary among diverse populations. Here, we introduce covariate-adjusted LD score regression (cov-LDSC), a method to accurately estimate genetic heritability  $(h_a^2)$  and its enrichment in both homogenous and admixed populations with summary statistics and in-sample LD estimates. In-sample LD can be estimated from a subset of the GWAS samples, allowing our method to be applied efficiently to very large cohorts. In simulations, we show that unadjusted LDSC underestimates  $h_q^2$  by 10% - 60% in admixed populations; in contrast, cov-LDSC is robust to all simulation parameters. We apply cov-LDSC to genotyping data from approximately 170,000 Latino, 47,000 African American and 135,000 European individuals. We estimate  $h_g^2$  and detect heritability enrichment in three quantitative and five dichotomous phenotypes respectively, making this, to our knowledge, the most comprehensive heritability-based analysis of admixed individuals. Our results show that most traits have high concordance of  $h_g^2$  and consistent tissue-specific heritability enrichment among different populations. However, for age at menarche, we observe population-specific heritability estimates of  $h_g^2$ . We observe consistent patterns of tissue-specific heritability enrichment across populations; for example, in the limbic system for BMI, the per-standardized-annotation effect size  $\tau^*$  is  $0.16 \pm 0.04$ ,  $0.28 \pm 0.11$  and  $0.18 \pm 0.03$  in Latino, African American and European populations respectively. Our results demonstrate that our approach is a powerful way to analyze genetic data for complex traits from underrepresented populations.

## Author summary

Admixed populations such as African Americans and Hispanic Americans bear a disproportionately high burden of disease but remain underrepresented in current genetic studies. It is important to extend current methodological advancements for understanding the genetic basis of complex traits in homogeneous populations to individuals with admixed genetic backgrounds. Here, we develop a computationally efficient method to answer two specific questions. First, does genetic variation contribute to the same amount of phenotypic variation (heritability) across diverse populations? Second, are the genetic mechanisms shared among different populations? To answer these questions, we use our novel method to conduct the first comprehensive heritability-based analysis of a large number of admixed individuals. We show that there is a high degree of concordance in total heritability and tissue-specific enrichment between different ancestral groups. However, traits such as age at menarche show a noticeable differences among populations. Our work provides a powerful way to analyze genetic data in admixed populations and may contribute to the applicability of genomic medicine to admixed population groups.

## Introduction

It is important for human geneticists to study how genetic variants that influence phenotypic variability act across different populations worldwide [1,2]. With increasingly large and diverse genetic studies, it is now becoming feasible to assess how the genetic mechanisms of complex traits act across populations. However, to date, most genome-wide association studies (GWAS) have been focused on relatively homogenous continental populations, and in particular those of European descent [3]. Non-European populations, particularly those with mixed ancestral backgrounds such as African Americans and Latinos, have been underrepresented in genetic studies. Many statistical methods to analyze genetic data assume homogeneous populations. In order to ensure that the benefits of GWAS are shared beyond individuals of homogeneous continental ancestry, statistical methods for admixed populations are needed [4].

Among methods to analyze polygenic complex traits in homogeneous populations,

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summary statistics-based methods such as linkage disequilibrium score regression (LDSC) [5, 6] and its extensions [7–9] have become particularly popular due to their computational efficiency, relative ease of application, and their applicability without raw genotyping data [10]. These methods can be used to estimate SNP-heritability, the proportion of phenotypic variance explained by genotyped variants [5, 11–13], distinguish polygenicity from confounding [5], establish relationships between complex phenotypes [7], and model genome-wide polygenic signals to identify key cell types and regulatory mechanisms of human diseases [6, 9, 14].

Summary statistics-based methods for polygenic analysis frequently rely on linkage 22 disequilibrium (LD) calculations. For LD score regression, the LD information needed is 23 the LD score for each SNP, defined to be the sum of its pairwise correlations  $(r^2)$  with 24 all other SNPs. For homogeneous populations there is usually a reference panel of 25 individuals with matching ancestry that can be used to approximate the in-sample LD. For studies with heterogeneous or admixed ancestry, however, even when reference 27 panels are available, they may not be representative of the precise populations used in the genetic study. For example Latino populations in different regions worldwide may share the same ancestral continental populations, but with dramatic differences in admixture proportions and timing of the admixture event [15]. A generic reference 31 panel cannot easily capture these differences and hence cannot produce accurate LD scores that can be widely used for all Latino populations. Moreover, the structure of LD 33 in heterogenous and admixed populations is complex and includes longer-range 34 correlations that are absent or negligible in homogeneous populations. Thus, while LD scores computed from a matching reference panel reflect the appropriate matching LD for summary statistics computed in a homogeneous population, it has not been clear what the appropriate matching LD is for summary statistics computed in a heterogenous or admixed population, and so LDSC has only been recommended to be applied in homogeneous populations. 40

Here, we evaluate the heritability estimates using LDSC in admixed population and observe systematic underestimation. We then introduce covariate-adjusted LD score regression (cov-LDSC) to estimate heritability and partitioned heritability in admixed populations. We apply our approach to 8, 124 Latinos from a type 2 diabetes study (the Slim Initiative in Genomic Medicine for the Americas, SIGMA) [16] as well as 161, 894 Latino, 46,844 African American, and 134,999 European research participants from a personal genetics company (23andMe). We analyze three quantitative phenotypes (body mass index (BMI), height, and age at menarche), and five dichotomous phenotypes (type 2 diabetes (available in the SIGMA cohort only), left handedness, morning person, motion sickness, and nearsightedness).

One powerful component of LDSC is that it can be used to test whether a particular 51 genome annotation -- for example, sets of genes that are specifically expressed within a 52 candidate tissue or cell type -- capture more heritability than expected by chance [9,11]. 53 We demonstrate that cov-LDSC can be applied in the same way to identify trait-relevant 54 tissue and cell types in admixed and homogenous populations with well-calibrated type 55 I error. We examine height, BMI and morning person since these traits had sufficient statistical power [6] for cell-type enrichment analyses in the 23 and Me cohort. We 57 observe a high level of consistency among enriched tissue types, highlighting that the underlying biological processes are shared among studied populations. This heritability 59 enrichment analysis of hundreds of genome annotations in cohorts of over 100,000 individuals would have been challenging with existing genotype-based methods [17–19]. 61

## Results

### **Overview of methods**

In this work, we extended the LDSC-based methods to heterogeneous and admixed populations by introducing covariate-adjusted LDSC (cov-LDSC). We first showed 65 through derivations that the appropriate matching LD for summary statistics computed in a heterogeneous or admixed population is in-sample LD computed on genotypes that 67 have been adjusted for the same covariates (e.g. principal components) included in the summary statistics (S1 Appendix). In cov-LDSC, we compute these covariate-adjusted 69 LD scores and then use LDSC to estimate heritability and its enrichment (Methods). 70 We showed that, unlike LDSC, cov-LDSC produces accurate estimates of heritability 71 with summary statistics from admixed populations (Methods, Fig 1). Furthermore, 72 heritability can be partitioned to identify key gene sets that have disproportionately 73 high heritability. While access to the genotype data of the GWAS samples is required to 74 compute the covariate-adjusted LD scores, LD can be estimated on a random subset of 75

the individuals, preserving the computational efficiency of LDSC and allowing for its application to very large studies. Individual cohorts can also release the in-sample covariate-adjusted LD scores as well as the summary statistics to avoid privacy concerns associated with genotype-level information to facilitate future studies.

#### **Robustness of LD score estimation**

To demonstrate the effect of admixture on the stability of LD score estimates, we first 81 calculated LD scores with genomic window sizes ranging from 0-50 cM in both 82 European (EUR, N = 503) and admixed American (AMR, N = 347) populations from 83 the 1000 Genomes Project [20]. As window size increases, we expect the mean LD score 84 to plateau because LD should be negligible for large enough distance. If the mean LD score does not plateau, but continues to rise with increasing window size, then one of two possibilities may apply: (1) the window is too small to capture all of the LD; (2)87 the LD scores are capturing long-range pairwise SNP correlations arising from 88 admixture. If this increase is non-linear then there is non-negligible distance-dependent 89 LD, violating LDSC assumptions. Examining unadjusted LD scores, we observed that 90 in the EUR population [5], the mean LD score estimates plateaued at windows beyond 91 1-cM in size, as previously reported. However, in the AMR population the mean LD 92 score estimates continued to increase concavely with increasing window size. In contrast, 93 when we applied cov-LDSC with 10 PCs to calculate covariate adjusted LD scores, we observed that LD score estimates plateaued for both EUR and AMR at a 1-cM and 20-cM window size respectively (< 1% increase per cM, S1 Table). This suggested that cov-LDSC was able to correct the long-range LD due to admixture and yielded stable 97 estimates of LD scores (Method, S1 Fig), and also that cov-LDSC was applicable in homogeneous populations (S1 Table). The larger window size for the AMR population 99 was needed due to residual LD caused by recent admixture. We next tested the 100 sensitivity of the LD score estimates with regard to the number of PCs included in the 101 cov-LDSC. We observed that in the AMR panel, LD score estimates were unaffected by 102 adding PCs and by increasing window sizes above 20-cM (S2 Fig). 103

#### Simulations with simulated genotypes

To assess whether cov-LDSC produces less biased estimates of  $h_a^2$ , we simulated 105 genotypes of two admixed populations (African American and Latino, **Methods**). We 106 simulated genotypes of 10,000 unrelated diploid admixed individuals for approximately 107 400,000 common SNPs on chromosome 2 in a coalescent framework using 108 msprime [21] (Methods). First, we tested LDSC and cov-LDSC with different 109 admixture proportions between two ancestral populations, and a quantitative phenotype 110 with a  $h_a^2$  of 0.4 using an additive model (**Methods**). We observed that as the 111 proportion of admixture increased,  $\widehat{h}_g^2$  for LDSC increasingly underestimated true  $h_g^2$  by 112 as much as 18.6%. In marked contrast, cov-LDSC produced consistently less biased 113 estimates regardless of admixture proportion for both Latinos (S3 Fig(a)) and African 114 Americans (S4 Fig). Since both simulated admixed populations would lead to the same 115 conclusions, we performed the subsequent simulations in the Latino individuals only. 116

Second, we varied the percentage of causal variants from 0.01% to 50% in a polygenic quantitative trait with  $h_g^2 = 0.4$  in a population with a fixed admixture proportion of 50%. LDSC again consistently underestimated  $h_g^2$  by 12% – 18.6%. In contrast, cov-LDSC yielded less biased estimates regardless of the percentage of causal variants (S3 Fig(b)).

Third, we assessed the robustness of LDSC and cov-LDSC for different assumed total  $h_g^2$  (0.05, 0.1, 0.2, 0.3, 0.4 and 0.5). At each  $h_g^2$  value, LDSC underestimated by 11.5% – 19.6%. For cov-LDSC, we observed that the standard error increased with  $h_g^2$ , 124 but point estimates remained less biased (S3 Fig(c)). 125

Fourth, we included an environmental stratification component aligned with the first <sup>126</sup> PC of the genotype data (**Methods**), and concluded that cov-LDSC was also robust to <sup>127</sup> confounding (S3 Fig(d)). <sup>128</sup>

Finally, to assess the performance of cov-LDSC in polygenic binary phenotypes, we simulated genotype data for a binary trait with a prevalence of 0.1 assuming a liability threshold model (**Methods**). We showed that cov-LDSC provided less biased estimates in case-control studies with the same four simulation scenarios (S5 Fig). In contrast, LDSC underestimated heritability for binary phenotypes in the same way as it did for quantitative phenotypes.

#### Simulation results with real genotypes

We next examined the performance of both unadjusted LDSC and cov-LDSC on real 136 genotypes of individuals from admixed populations. We used genotype data from the 137 SIGMA cohort, which includes 8,214 Mexican and other Latino individuals. Using 138 ADMIXTURE [22] and populations from the 1000 Genomes Project [20] as reference 139 panels, we observed that each individual in the SIGMA cohort had different admixture 140 proportions (S6 Fig). As in the AMR panel, we observed that using a 20-cM window, 141 LD score estimates plateaued in the SIGMA cohort (S7 Fig, S2 Table), and were 142 unaffected by different numbers of PCs (S8 Fig). When we assumed a non-infinitesimal, 143 additive model with 1% of all SNPs to be causal and  $h_a^2 = 0.4$ , we observed that 144 cov-LDSC  $h_g^2$  estimates produced less biased estimates using a 20-cM window with 10 145 PCs (S9 Fig). We subsequently used a 20-cM window and 10 PCs in all simulations. 146

We observed that cov-LDSC yielded less biased  $h_a^2$  estimates in simulated traits 147 where we varied the number of causal variants and total heritability compared to the 148 original LDSC (Fig 2(a)-(b)). In contrast, LDSC underestimated heritability by as 149 much as 62.5%. To examine the performance of cov-LDSC in the presence of 150 environmental confounding factors, we simulated an environmental stratification 151 component aligned with the first PC of the genotype data, representing European v.s. 152 Native American ancestry. In this simulation scenario, cov-LDSC still provided less 153 biased  $h_a^2$  estimates (Fig 2(c)). Intercepts of all the simulation scenarios were less than 154 the genomic control inflation factor (GC), suggesting that polygenicity accounts for a 155 majority of the increase in the mean  $\chi^2$  statistic compared to potential confounding 156 biases (S10 Fig(a)-(c), S3 Table). 157

Thus far, we have used cov-LDSC by calculating LD scores on the same set of 158 samples that were used for association studies (in-sample LD scores). In practical 159 applications, computing LD scores on the whole data set can be computationally 160 expensive and difficult to obtain, so we investigated computing LD scores on a subset of 161 samples. To investigate the minimum number of samples required to obtain accurate 162 in-sample LD scores, we computed LD scores on subsamples of 100, 500, 1,000 and 163 5,000 individuals from a GWAS of 10,000 simulated genotypes (S11 Fig). We repeated 164 these analyses in simulated phenotypes in the SIGMA cohort. We subsampled the 165

SIGMA cohort, and obtained less biased estimates when using as few as 1,000 samples (Fig 2(d)). We therefore recommend computing in-sample LD scores on a randomly chosen subset of at least 1,000 individuals from a GWAS in our approach.

#### Assessing power and bias in tissue type specific analysis

Following Finucane et al [9], we extended cov-LDSC so that we can assess enrichment in 170 and around sets of genes that are specifically expressed in tissue and cell-types 171 (cov-LDSC-SEG). To test whether cov-LDSC can produce robust results with properly 172 controlled type I error, We calculated the in-sample LD scores using LDSC and 173 cov-LDSC, respectively, using a 20-cM window and 10 PCs in cov-LDSC for all 53 174 baseline and limbic system annotations. We used PLINK2 [23] for association test and 175 performed tissue type specific enrichment analysis using both LDSC and cov-LDSC for 176 limbic system conditioning on all 53 baseline annotations. We reported the number of 177 significant tests out of 1,000 simulations in each scenario. We observed no inflation in 178 false-positive rate (FPR) at 0.05 for both LDSC and cov-LDSC under null (i.e., no 179 enrichment). The greatest gains in power were observed in cases where there were 180 modest enrichment ( $\langle 2 \times \rangle$ ). We showed that cov-LDSC-SEG was better powered to 181 detect tissue type specific signals compared to LDSC-SEG (S12 Fig). 182

### Application to SIGMA and 23andMe cohorts

We next used cov-LDSC to estimate  $h_q^2$  of height, BMI and T2D phenotypes, measured 184 within the SIGMA cohort (Methods, Table 1). We estimated  $h_q^2$  of height, BMI and 185 T2D to be  $0.38 \pm 0.08$ ,  $0.25 \pm 0.06$  and  $0.26 \pm 0.07$ , respectively. These results were 186 similar to reported values from UK Biobank [24] and other studies [17,25] for European 187 populations. Although estimands differed in different studies (Methods), we noted 188 that without cov-LDSC, we would have obtained severely deflated estimates (**Table 1**). 189 To confirm that our reported heritability estimates were robust under different model 190 assumptions, we applied an alternative approach based on REML in the linear mixed 191 model framework implemented in GCTA [17]. To avoid biases introduced from 192 calculating genetic relatedness matrices (GRMs) in admixed individuals, we obtained a 193 GRM based on an admixture-aware relatedness estimation method REAP [26] 194

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(Methods). GCTA-based results were similar to reported  $h_g^2$  estimates from cov-LDSC, indicating our method was able to provide reliable  $h_g^2$  estimates in admixed populations (Table 1). We noted, however, that the GCTA-based results would be computationally expensive to obtain on the much larger datasets, for example the 23andMe cohort described below.

We next applied both LDSC and cov-LDSC to 161,894 Latino, 46,844 African 200 American and 134,999 European research participants from 23 and Me. We analyzed 201 three quantitative and four dichotomous phenotypes (Methods, S4 Table). In this 202 setting, we noted that if different individuals were included in different traits of 203 interests, one would need to re-compute the GRM for each trait when using 204 genotype-based methods such as GCTA [17] or BOLT-REML [19]. Whereas for 205 cov-LDSC we do not require complete sample overlap between LD reference panel and 206 summary statistics generation. Thus one would only need to compute 207 covariate-adjusted baseline LD score once for each cohort. This makes cov-LDSC a 208 more computationally attractive strategy for estimating heritability and its enrichment 209 in large cohorts. We used a 20-cM window and 10 PCs in LD score calculations for both 210 populations (S13 Fig, S5 Table). LDSC and cov-LDSC produced similar heritability 211 estimates in the European population, whereas in the admixed populations, LDSC 212 consistently provided low estimates of  $h_g^2$  (S6 Table). For each phenotype, we estimated 213  $h_q^2$  using the same population-specific in-sample LD scores. Intercepts of all the traits 214 were substantially less than the genomic control inflation factor ( $\lambda_{ac}$ ), suggesting that 215 polygenicity accounts for a majority of the increase in the mean  $\chi^2$  statistics (S7 Table). 216 For most phenotypes, the reported  $h_q^2$  was similar among the three population groups 217 with a notable exception for age at menarche (Fig 3, S8 Table). This suggested possible 218 differences (two-sample t-test  $p = 7.1 \times 10^{-3}$  between Latinos and Europeans) in the 219 genetic architecture of these traits between different ancestral groups. It has been long 220 established that there is population variation in the timing of menarche [27-29]. Early 221 menarche might influence the genetic basis of other medically relevant traits since early 222 age at menarche is associated with a variety of chronic diseases such as childhood 223 obesity, coronary heart disease and breast cancer [30, 31]. These results highlighted the 224 importance of including diverse populations in genetic studies in order to enhance our 225 understanding of complex traits that show differences in their genetic heritability. 226

#### Tissue type specific analysis

We applied stratified cov-LDSC to sets of specifically expressed genes [9] (SEG) to 228 identify trait-relevant tissue and cell types in traits included in the 23 and Me cohort 229 across European, Latino, and African American populations. We only tested height, 230 BMI and morning person, which were the three traits that had heritability z-scores 231 larger than seven in at least two populations [6] (S9 Table). We also performed 232 inverse-variance weighting meta-analysis across the three populations (S10 Table). 233 Across different populations, BMI showed consistent enrichment in central nervous 234 system gene sets. In the European population, most of the enrichments recapitulated 235 the results from the previous analysis using UK Biobank [9]. We found similar but fewer 236 enrichments in Latinos and African Americans, most likely due to smaller sample sizes. 237 The most significantly enriched tissue types for BMI in all three populations were limbic 238 system ( $\tau^*_{\text{EUR}} = 0.18, \tau^*_{\text{LAT}} = 0.16, \tau^*_{\text{AA}} = 0.28, \tau^*_{\text{meta}} = 0.18$ ), entorhinal cortex 239  $(\tau^*_{\text{EUR}} = 0.18, \tau^*_{\text{LAT}} = 0.15, \tau^*_{\text{AA}} = 0.24, \tau^*_{\text{meta}} = 0.17)$ , and cerebral cortex 240  $(\tau^*_{\text{EUR}} = 0.16, \tau^*_{\text{LAT}} = 0.14, \tau^*_{\text{AA}} = 0.15, \tau^*_{\text{meta}} = 0.15);$  none of the three effects 241 were significantly different across populations. When we compared the enrichment for 242 all of the tissues between population pairs, we observed that they have significant 243 non-zero concordance correlation coefficient ( $\rho_{\text{EUR-LAT}} = 0.78 (0.72 - 0.83);$ 244  $\rho_{\text{EUR-LAT}} = 0.32 (0.21 - 0.42))$  (Fig 4(a)-(e), S11 Table). The sizes of these three 245 brain structures have been shown to be correlated with BMI using magnetic resonance 246 imaging data [32]. The midbrain and the limbic system are highly involved in the food 247 rewarding signals through dopamine releasing pathway [33]. Furthermore, the 248 hypothalamus in the limbic system releases hormones that regulate appetite, energy 249 homeostasis and metabolisms, like leptin, insulin, and ghrelin [33, 34]. For height, 250 similar to previously reported associations [9], we also identified enrichments in the gene 251 sets derived from musculoskeletal and connective tissues. In the meta-analysis, the three 252 most significant enrichments were cartilage ( $\tau^*_{\text{EUR}} = 0.21$ ,  $\tau^*_{\text{LAT}} = 0.19$ ,  $\tau^*_{\text{AA}} = 0.24$ , 253  $\tau^*_{\text{meta}} = 0.20$ ), chondrocytes ( $\tau^*_{\text{EUR}} = 0.21$ ,  $\tau^*_{\text{LAT}} = 0.15$ ,  $\tau^*_{\text{AA}} = 0.11$ , 254  $\tau^*_{\text{meta}} = 0.17$ ), and uterus ( $\tau^*_{\text{EUR}} = 0.17$ ,  $\tau^*_{\text{LAT}} = 0.15$ ,  $\tau^*_{\text{AA}} = 0.16$ ,  $\tau^*_{\text{meta}} = 0.16$ ). 255 A heterogeneity test revealed no difference across three populations ( $I^2 < 70\%$  and 256 p-value > 0.05). The concordance correlation coefficients were 257

 $\rho_{\text{EUR-LAT}} = 0.91 (0.89 - 0.93)$  between European and Latio; 258  $\rho_{\text{EUR-AA}} = 0.60 (0.50 - 0.68)$  between European and African American (Fig 4(f)-(j), 259 S11 Table). The importance of these tissues and their roles in height have been 260 addressed in the previous pathway analysis, expression quantitative trait loci (eQTLs) 261 and epigenetic profiling [35, 36]. Previous studies have shown that the longitudinal 262 growth of bones is partly controlled by the number and proliferation rate of 263 chondrocytes on the growth plate which is a disc of cartilages [37]. For the morning 264 person phenotype, we found enrichments in many brain tissues in Europeans, 265 concordant with a previous study [38]. Entorhinal cortex ( $\tau^*_{\text{EUR}} = 0.16$ ,  $\tau^*_{\text{LAT}} = 0.22$ , 266  $\tau^*_{\text{meta}} = 0.18$ ), cerebral cortex ( $\tau^*_{\text{EUR}} = 0.15$ ,  $\tau^*_{\text{LAT}} = 0.22$ ,  $\tau^*_{\text{meta}} = 0.18$ ), and 267 brain ( $\tau^*_{\text{EUR}} = 0.17$ ,  $\tau^*_{\text{LAT}} = 0.19$ ,  $\tau^*_{\text{meta}} = 0.18$ ) were enriched in both Latinos and 268 Europeans. Evidence showed that circadian rhythm was controlled by the 269 suprachiasmatic nucleus, the master clock in our brain, and also the circadian oscillator 270 that resides in neurons of the cerebral cortex [39–41]. We also found unique enrichments 271 of esophagus muscularis and the esophagus gastroesophageal junction in the Latino 272 populations, but the heterogeneity test showed that the difference is not significant 273  $(I^2 = 0.49 \text{ and } 0.50, \text{ respectively})$ . We observed that the concordance correlation 274 coefficient across gene sets was 0.63(0.51 - 0.68) between Latino and European 275 (Fig 4(k)-(n), S11 Table). Compared to the original LDSC-SEG, cov-LDSC-SEG 276 appeared to have increased statistical power in detecting tissue type specific enrichment 277 in the African American and Latino population (S12 Fig, S14 Fig, S15 Fig, S16 Fig). 278

# Discussion

As we expand genetic studies to explore admixed populations around the world, 280 extending statistical genetics methods to make inferences within admixed populations is 281 crucial. This is particularly true for methods based on summary statistics, which are 282 dependent on the use of LD scores that we showed to be problematic in admixed 283 populations. In this study, we confirmed that LDSC that was originally designed for 284 homogenous populations, should not be applied to admixed populations. We introduced 285 cov-LDSC which regresses out global PCs on individual genotypes during the LD score 286 calculation, and showed it can yield less biased LD scores, heritability estimates and its 287

enrichment, such as trait-relevant cell and tissue type enrichments, in homogenous and <sup>288</sup> admixed populations. <sup>289</sup>

Although our work provides a novel, efficient approach to estimate genetic 290 heritability and to identify trait-relevant cell and tissue types using summary statistics 291 in admixed populations, it has a few limitations. First, covariates included in the 292 summary statistics should match the covariates included in the covariate-adjusted LD 293 score calculations (S1 Appendix). To demonstrate this, we simulated the phenotypes 294 using real genotypes included in the SIGMA cohort. We performed cov-LDSC to 295 measure total heritability and its enrichment with varied number of PCs included in 296 summary statistics and in LD score calculation. As the differences between the number 297 of PCs included in the summary statistics and LD score calculation increase, we 298 observed an increase in bias of the total heritability estimation (S17 Fig) and a loss in 299 power when detecting tissue-specific enrichment (S18 Fig). Second,  $h_a^2$  estimates and 300 their enrichment in admixed populations are more sensitive to potentially unmatched 301 LD reference panels. Unmatched reference panels are likely to produce biased 302 estimates [42,43] and under-powered enrichment analysis (S12 Table, S14 Fig, S15 Fig, 303 S16 Fig). We examined the performance of using an out-of-sample reference panel in 304 admixed populations (See S2 Appendix) and caution that when using 1000 Genomes or 305 any out-of-sample reference panels for a specific admixed cohort, users should ensure 306 that the demographic histories are shared between the reference and the study cohort. 307 Large sequencing projects such as TOPMed [44] that include large numbers (N > 1,000)308 of admixed samples can potentially serve as out-of-sample LD reference panels, although 309 further investigations are needed to study their properties. We therefore advise to 310 compute in-sample LD scores from the full or a random subset of data (N > 1,000)311 used to generate the admixed GWAS summary statistics when possible. For tissue and 312 cell type-specific analyses, this means one needs to compute covariate-adjusted LD 313 scores for the genome annotations that were derived from the publicly available gene 314 expression data. We have released open-source software implementing our approach 315 based on all genome annotations derived previously (**URLs**). We strongly encourage 316 cohorts to release their summary statistics and in-sample covariate-adjusted LD scores 317 at the same time to facilitate future studies. Third, when applying cov-LDSC to 318 imputed variants, particularly those with lower imputation accuracy (INFO < 0.99), we 319

caution that the heritability estimates and its enrichment can be influenced by an 320 imperfect imputation reference panel, especially in Latino populations [45, 46]. To limit 321 the bias in varying genotyping array and imputation quality in studied admixed cohorts, 322 we recommend restricting the heritability analyses to common HapMap3 variants. Any 323 extension to a larger set of genetic variants, especially across different cohorts should be 324 performed with caution. Fourth, when we evaluated the performance of cov-LDSC in 325 case-control studies, we assumed no presence of binary covariates with strong effects 326 and demonstrated that cov-LDSC can yield robust  $h_q^2$  estimates. However, it has been 327 shown that LDSC can provide biased estimates in the presence of extreme ascertainment 328 for dichotomous phenotypes [47]. Adapting cov-LDSC into case-control studies under 329 strong binary effects remains a potential avenue for future work. Fifth, recent studies 330 have shown that heritability estimates can be sensitive to the choice of the LD- and 331 frequency-dependent heritability model [8, 11, 13, 48]. Since our approach can flexibly 332 add annotations to estimate heritability under the model that is best supported by the 333 data, we believe it provides a good foundation for addressing the question of how to 334 incorporate ancestry-dependent frequencies in the LD-dependent annotation in the 335 future (Methods). Sixth, summary statistics derived from linear mixed models cannot 336 currently be used for cov-LDSC analysis (S19 Fig). This is due to the fact that, just as 337 the LD needs to be adjusted for the same covariates included in the summary statistics 338 (S1 Appendix), it also needs to be corrected appropriately for the random effect. We 339 leave efficient computation of random effect-adjusted LD score to future work. 340

Despite these limitations, in comparison with other methods, such as those based on 341 restricted maximum likelihood estimation (REML) [17,19] with an admixture-aware 342 GRM [26], for estimating  $h_g^2$  in heterogeneous or admixed populations, cov-LDSC has a 343 number of attractive properties. First, covariate-adjusted in-sample LD scores can be 344 obtained with a subset of samples, enabling analysis of much larger cohorts than was 345 previously possible. Second, LD scores only need to be calculated once per cohort; this 346 is particularly useful in large cohorts such as 23 and Me and UK Biobank [49], where 347 multiple phenotypes have been collected per individual and per-trait heritability and its 348 enrichment can be estimated based on the same LD scores. Third, as a generalized form 349 of LDSC, it is robust to population stratification and cryptic relatedness in both 350 homogenous and admixed populations. Fourth, similar to the original LDSC methods, 351 cov-LDSC can be extended to perform analyses such as estimating genetic correlations, partitioning  $h_g^2$  by functional annotations, identifying disease-relevant tissue and cell types and multi-trait analysis [6,9,50,51].

By applying cov-LDSC to approximately 344,000 individuals from European, 355 African American, and Latin American ancestry, we observed evidence of heritability 356 differences across different populations. Differences in environmental exposures and 357 biological mechanisms can both contribute to the observed differences in genetic 358 heritability across trans-ethnic populations. These differences highlight the importance 359 of studying diverse populations In particular, the differences in biological mechanisms 360 may lead to mechanistic insights about the phenotype. One strategy to do this, which 361 we explored by extending cov-LDSC, is to partition heritability by different cell type-362 and tissue-specific annotations to dissect the genetic architecture in admixed 363 populations. Our results demonstrated that although there are some cases of nominal 364 heterogeneity across populations among tested tissue-types, most of the tissue-specific 365 enrichments are consistent among the populations studied here. This is consistent with 366 the previous findings that show strong correspondence in functional and cell type 367 enrichment between Europeans and Asians [52, 53]. Seeing the same tissue-type for a 368 single trait emerge in multiple populations can give us more confidence that this tissue 369 may account for polygenic heritability. Larger sample sizes are needed to increase the 370 power of our current analyses and to enhance our understanding of how genetic variants 371 that are responsible for heritable phenotypic variability differ among populations. 372

As the number of admixed and other diverse GWAS and biobank data become readily available [1,44,54], our approach provides a powerful way to study admixed populations.

## Materials and methods

### Mathematical framework of cov-LDSC

Details of the mathematical derivation of cov-LDSC are presented in S1 Appendix. 378 Briefly, in the standard polygenic model on which LDSC is based,  $x_1, \ldots, x_N$  are the 379 length-M genotype vectors for the N individuals, where M is the number of SNPs. We 380

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model the phenotypes  $y_i$ 

$$y_i = x_i \beta + \epsilon_i,\tag{1}$$

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where  $\epsilon_1, ..., \epsilon_N \stackrel{iid}{\sim} \mathcal{N}(0, \sigma_e^2)$  and  $\beta \in \mathbb{R}^M$  is a vector of per-normalized-genotype effect sizes, which we model as random with mean zero. In standard LDSC, the variance of  $\beta_j$ ,  $\operatorname{Var}(\beta_j)$ , is the per-SNP heritability of SNP j, that is, the total SNP-heritability  $h_g^2$ divided by the total number of SNPs M ( $h_g^2/M$ ). In stratified LD score regression the variance of  $\beta_j$  depends on a set of genome annotations.

Let  $\chi_j^2$  denote the chi-square statistic for the  $j^{th}$  SNP, approximately equal to  $(X_j^T Y)^2/N$ , where  $X_j = (x_{1j}, ..., x_{Nj})^T$  and  $Y = (y_1, ..., y_N)^T$ . The main equation on which LDSC is based is: 389

$$\mathbb{E}[\chi_j^2] \approx 1 + Na + \frac{Nh_g^2}{M} \ell(j), \tag{2}$$

where a is a constant that reflects population structure and other sources of confounding, and the LD score,  $\ell(j)$ , is:

$$\ell(j) = \sum R_{jk}^2$$

 $R_{jk}^{2}$  is the correlation between SNPs j and k in the underlying population. A new derivation for this equation is given in S1 Appendix. We estimate the total SNP-heritability  $h_{g}^{2}$  via weighted regression of  $\chi_{j}^{2}$  on our estimates of  $\ell(j)$ , evaluating significance with a block jackknife across SNPs [6].

In the absence of covariates, the LD scores can be estimated from an external reference panel such as 1000 Genomes, as long as the correlation structure in the reference panel matches the correlation structure of the sample. In most homogeneous populations, we can also assume that the true underlying correlation is negligible outside of a 1-cM window.

In the presence of covariates, we let C denote the  $N \times K$  matrix of covariates, each column centered to mean zero, and let  $c_i$  be the *i*-th row of C. Equation (1) can then be replaced with

$$y_i = x_i\beta + c_i\beta_{cov} + \epsilon_i,\tag{3}$$

where  $\beta_{cov}$  is a vector of effect sizes of covariates. We can project the covariates out of 402

this equation by multiplying by  $P = I - C(C^T C)^{-1} C^T$  on the left to get

$$\widetilde{Y} = \widetilde{X}\beta + \widetilde{\epsilon},\tag{4}$$

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where  $\tilde{Y} = PY$ ,  $\tilde{X} = PX$  and  $\tilde{\epsilon} = P\epsilon$  (if the covariates are genotype principal components, then  $P = I - CC^T$ ). Under this model, an equation identical to Equation (2) can be derived, but where both summary statistics and LD are adjusted for the same covariates (see S1 Appendix).

If X is a homogeneous population, then the covariate-adjusted LD will be similar to the non-covariate-adjusted LD and well-approximated by a reference panel. However, if X is the genotype matrix from an admixed or heterogeneous population and the covariates include PCs, then the covariate-adjusted LD is no longer well-approximated by either non-covariate-adjusted LD or by a reference panel. Thus, in cov-LDSC, we compute LD scores directly from the covariate-adjusted in-sample genotypes or a random subsample thereof. We call them the covariate-adjusted LD scores.

Using genotype data to compute LD scores means that the model being fit is based 415 on the joint effects of a sparser set of SNPs, e.g. the genotyped SNPs, than when 416 sequence data is used to compute LD scores. For estimating total SNP-heritability, this 417 means that cov-LDSC estimates the same estimand as GCTA  $(h_a^2)$  and not the usual 418 estimand of LDSC  $(h_{\text{common}}^2; \text{ see below})$ . For partitioned heritability, the density of 419 reference panel SNPs can be important because the joint effect of a SNP in an 420 annotation can include the tagged effect of an untyped SNP that is not in the 421 annotation, deflating estimates of enrichment. Thus, we recommend using cov-LDSC 422 only on annotations made of large contiguous regions, such as gene sets. Moreover, we 423 urge caution when interpreting quantitative estimates of heritability enrichment. Here, 424 we look at the significance of the conditional enrichment (i.e., regression coefficient) of 425 gene sets for our tissue-specific analysis (see below).

#### Window size and number of PCs in LD score calculations

In addition to computing LD from the covariate-adjusted genotypes, we also investigate 428 the appropriate window size for estimating LD scores. To do this, we examine the effect 429 of varying the genomic window size for both simulated and real data sets. We determine 430

that LD score estimates were robust to the choice of window size if the increase in the 431 mean LD score estimates was less than 1% per cM beyond a given window. Using this 432 criterion, we use window sizes of 5-cM and 20-cM for the simulated and real genotypes, 433 respectively (S13 Table, S2 Table, S5 Table). We also calculate the squared correlations 434 between LD score estimates using the chosen window size and other LD score estimates 435 with window sizes larger than the chosen window. The Pearson squared correlations 436 were greater than 0.99 in all cases (S14 Table, S15 Table, S16 Table) indicating the LD 437 score estimates were robust at the chosen window sizes. 438

Similarly, to determine the number of PCs needed to be included in the GWAS 439 association tests and cov-LDSC calculations, we examine the effect of varying the 440 genomic window size using different numbers of PCs. The number of PCs that needed 441 to be included for covariate adjustment depended on the population structure for 442 different datasets. 443

## Genotype simulations

We evaluate the performance of LDSC and cov-LDSC with simulated phenotypes and 445 both simulated and real genotypes. For the simulated genotypes, we used msprime [21] 446 version 0.6.1 to simulate population structure with mutation rate  $2 \times 10^{-8}$  and 447 recombination maps from the HapMap Project [55]. We adapt the demographic model 448 from Mexican migration history [56] for Latinos and the out of Africa model [57] for 449 African Americans using parameters that were previously inferred from the 1000 450 Genomes Project [20]. We assume the admixture event happened approximately 500 451 years and 200 years ago for Latino and African American populations, respectively. We 452 set different admixture proportions to reflect different admixed populations. In each 453 population, we simulate 10,000 individuals after removing second degree related 454 samples (kinship> 0.125) using KING [58]. 455

# Slim Initiative in Genomic Medicine for the Americas (SIGMA)456Type 2 Diabetes (T2D) cohort457

8,214 Mexican and other Latin American samples were genotyped with Illumina 458 HumanOmni2.5 array. We further filter the genotyped data to be MAF > 5% and 459

remove SNPs in high LD regions. After QC, a total of 8, 214 individuals and 943, 244 SNPs remain. We estimate the in-sample LD score with a 20-cM window and 10 PCs in all scenarios.

We use these genotypes for simulations. We also analyze three phenotypes from the SIGMA cohort: height, BMI, and type 2 diabetes (T2D). For T2D, we assume a reported prevalence in Mexico of 0.144 [16]. For each phenotype, we include age, sex, and the first 10 PCs as fixed effects in the association analyses.

#### Phenotype simulations

We simulate phenotypes with two different polygenic genetic architectures, given by 468 GCTA [17] and the baseline model [6], respectively. In the GCTA model, all variants 469 are equally likely to be causal independent of their functional or minor allele frequency 470 (MAF) structure, and the standardized causal effect size variance is constant, i.e. 471  $\operatorname{Var}(\beta_i) = h_a^2/M$ . In contrast, the baseline model incorporates functionally dependent 472 architectures. Briefly, it includes 53 overlapping genome-wide functional annotations 473 (e.g. coding, conserved, regulatory). It models  $\operatorname{Var}(\beta_j) = \sum_C \alpha_c(j) \tau_c$  where  $\alpha_c(j)$  is the 474 value of annotation  $\alpha_c$  at variant j and  $\tau_c$  represents the per-variant contribution, of 475 one unit of the annotation  $\alpha_c$ , to heritability. We generate all causal variants among 476 common observed variants with MAF > 5% ( $\sim 40,000$  SNPs in simulated genotypes 477 and 943,244 SNPs in the SIGMA cohort). To represent environmental stratification, 478 similar to previously described [5], we add  $0.2 \times$  standardized first principal component 479 to the standardized phenotypes. 480

We simulate both quantitative and case-control traits with both GCTA and baseline model genetic architectures, using both simulated and real genotypes, varying the number of causal variants, the true heritability, and environmental stratification. For case-control simulations, we adopt a liability threshold model with disease prevalence 0.1. We obtain 5,000 cases and 5,000 controls for each simulation scenario.

To obtain summary statistics for the simulated traits, we apply single-variant linear 486 models for quantitative traits and logistic models for binary trait both with 10 PCs as 487 covariates in association analyses using PLINK2 [23].

#### 23andMe cohort

All participants were drawn from the customer base of 23andMe, Inc., a direct to 490 consumer genetics company. Participants provided informed consent and participated in 491 the research online, under a protocol approved by the external AAHRPP-accredited 492 IRB, Ethical & Independent Review Services (www.eandireview.com). Samples from 493 23 and Me are then chosen from consented individuals who were genotyped successfully 494 on an Illumina Infinium Global Screening Array ( $\sim 640,000$  SNPs) supplemented with 495  $\sim 50,000$  SNPs of custom content. We restrict participants to those who have 496 European, African American, or Latino ancestry determined through an analysis of 497 local ancestry [59]. 498

To compute LD scores, we use both genotyped and imputed SNPs. We filter 499 genotyped variants with a genotype call rate < 90%, non-zero self-chain score, strong 500 evidence of Hardy Weinberg disequilibrium  $(p > 10^{-20}$  to accommodate large sample 501 sizes included for detecting deviations), and failing a parent-offspring transmission test. 502 For imputed variants, we use a reference panel that combined the May 2015 release of 503 the 1000 Genomes Phase 3 haplotypes [20] with the UK10K imputation reference 504 panel [60]. Imputed dosages are rounded to the nearest integer (0, 1, 2) for downstream 505 analysis. We filter variants with imputation r-squared  $\leq 0.9$ . We also filter genotyped and imputed variants for batch effects (if an F-test from an ANOVA of the SNP dosages 507 against a factor dividing genotyping date into 20 roughly equal-sized buckets has a 508 p-value less than  $10^{-50}$ ) and sex dependent effects (if the r-squared of the SNP is 509 greater than 0.01 after fitting a linear regression against the gender). To minimize 510 rounding inaccuracies, we prioritize genotyped SNPs over imputed SNPs in the merged 511 SNP set. We restrict the merged SNP set to HapMap3 variants with MAF  $\geq 0.05$ . We 512 measure LD scores in a subset of African Americans (61, 021) and Latinos (9, 990) on 513 chromosome 2 with different window sizes from 1-cM to 50-cM (S5 Table) and squared 514 correlation between different window sizes (S16 Table). We compute all LD scores with 515 a 20-cM window. 516

In genome-wide association analyses, for each population, we choose a maximal set of unrelated individuals for each analysis using a segmental identity-by-descent (IBD) estimation algorithm [61]. We define individuals to be related if they share more than 519

#### 700-cM IBD.

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We perform association tests using linear regression model for quantitative traits and logistic regression model for binary traits assuming additive allelic effects. We include covariates for age, sex and the top 10 PCs to account for residual population structure. We list details of phenotypes and genotypes in S4 Table.

## Heritability estimation

We calculate in-sample LD scores using both a non-stratified LD score [5] model and 526 the baseline model [6]. In simulated phenotypes generated with the GCTA model, we 527 use non-stratified LDSC to estimate heritability. In simulated phenotypes generated 528 using the baseline model, we use LDSC-baseline to estimate heritability. We use the 53 529 non-frequency dependent annotations included in the baseline model to estimate  $h_q^2$  in 530 the 23 and Me research database and the SIGMA cohort real phenotypes. We recognize 531 that recent studies have shown that genetic heritability can be sensitive to the choice of 532 LD-dependent heritability model [8, 11, 13]. However, understanding the LD- and 533 MAF-dependence of complex trait genetic architecture is an important but complex 534 endeavor potentially requiring both modeling of local ancestry as well as large 535 sequenced reference panels that are currently unavailable. We thus leave this complexity 536 for future work. 537

# $h_g^2$ versus $h_{common}^2$

The quantity  $(h_a^2)$  we reported in the main analysis is defined as heritability tagged by 539 HapMap3 variants with MAF  $\geq 5\%$ , including tagged causal effects of both 540 low-frequency and common variants. This quantity is different from  $h_{\text{common}}^2$ , the 541 heritability casually explained by all common SNPs excluding tagged causal effects of 542 low-frequency variants, reported in the original LDSC [5]. In Europeans and other 543 homogeneous populations, it is possible to estimate  $h_{\text{common}}^2$ , since reference panels, 544 such as 1000 Genomes Project [20], are available which include > 99% of the SNPs with 545 frequency > 1%. However, in-sample sequence data is usually not available for an 546 admixed GWAS cohort, and so cov-LDSC can only include genotyped SNPs in the 547 reference panel, and thus can only estimate the heritability tagged by a given set of 548

genotyped SNPs. In order to compare the same quantity across cohorts, we use common 549 HapMap3 SNPs (MAF  $\geq 5\%$ ) for in-sample LD reference panel calculation, since most 550 of them should be well imputed for a genome-wide genotyping array. To quantify the 551 difference between  $h_q^2$  and  $h_{common}^2$ , we pre-phase the genotype data in the SIGMA 552 cohort using SHAPEIT2 [62]. We use IMPUTE2 [63] to impute genotypes at untyped 553 genetic variants using the 1000 Genomes Project Phase 3 [20] dataset as a reference 554 panel. We merge genotyped SNPs and all well imputed (INFO> 0.99) SNPs (> 6.9555 million) in the SIGMA cohort as a reference panel and reported  $h_{\text{common}}^2$ , to 556 approximate what the estimate of  $h_{\text{common}}^2$  would have been with a sequenced reference 557 panel (S17 Table). 558

### Tissue type specific analyses

We generate the  $\tau$  for 53 baseline annotations with 40% of annotations with non-zero  $\tau$ 560 and 60% of annotations with zero  $\tau$ . We then generate different regression coefficients  $\tau$ 561 for limbic system in gene sets defined in Franke et al [64, 65] with different enrichment. 562 We scale all the  $\tau$  to make the total  $h_q^2 = 0.5$ . For each variant j, the variance of  $\beta_j$  is 563 the sum of the of all the categories that the variant is in  $(\operatorname{Var}(\beta_j) = \tau_c)$ . We randomly 564 draw j from a normal distribution with mean zero and variance  $\sum_{c:i \in C_c} \tau_c$  to simulate 565 the phenotypes. We run 1,000 simulations for each enrichment set (ranging from no 566  $(1\times)$  enrichment to 2.5× enrichment). We annotate the genes with the same set of 567 tissue specific expressed genes identified previously [9] using the Genotype–Tissue 568 Expression (GTEx) project [66] and a public dataset made available by the Franke 569 lab [64, 65]. We calculate within-sample stratified cov-LD scores with a 20-cM window 570 and 10 PCs in the 23andMe cohort for each of these 205 gene sets and 53 baseline 571 annotations. We obtain regression coefficients  $\hat{\tau}_c$  from the model and normalize them as 572

$$\tau_c^* = \frac{M_{h_g^2} \cdot sd_c}{h_g^2} \hat{\tau_c},$$

where  $M_{h_g^2}$  is the number of SNPs used to calculate  $h_g^2$  and  $sd_c$  is the standard deviation (sd) of annotation  $a_c$  [8]. We interpret  $\tau_c^*$  as the proportional change of averaged per-SNP heritability by one sd increase in value of the annotation of each cell type, conditional on other 53 non-cell type specific baseline annotations. We calculate a

one-tailed p-value for each coefficient where the null hypothesis is that the coefficient is non-positive [9]. All the significant enrichments are reported with false discovery rate  $< 5\% \ (-\log_{10}(p) > 2.75)$ . We perform fixed-effect inverse variance weighting meta-analysis using  $\tau_c^*$  and normalized standard error across populations.

## Software Availability

An open-source software implementation of covariate-adjusted LD score regression is publicly available (see Web Resources).

## Web Resources

cov-LDSC software and tutorials, https://github.com/immunogenomics/cov-ldsc $$	585
msprime, https://pypi.python.org/pypi/msprime;	586
GCTA, http://cnsgenomics.com/software/gcta/;	587
BOLT-LMM, v2.3.4, https://data.broadinstitute.org/alkesgroup/BOLT-LMM/;	588
LDSC, https://github.com/bulik/ldsc/;	589
PLINK2, https://www.cog-genomics.org/plink2;	590
$REAP \ v1.2, \ http://faculty.washington.edu/tathornt/software/REAP/download.html;$	591
ADMIXTURE v1.3.0,	592
http://www.genetics.ucla.edu/software/admixture/download.html;	593

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## Supporting information

S1 Table. Mean of LD scores with varying window sizes for populations 791 included in the 1000 Genomes project. AMR (N = 347) represents Admixed 792 American and EUR represent European populations (N = 503). 10 PCs are included in 793 all cov-LDSC estimates. 794

S2 Table. Mean of LD scores with varying window sizes for the SIGMA 795 cohort using LDSC and cov-LDSC. 10 PCs are included in all cov-LDSC 796 estimates. 797

S3 Table. Genomic inflation factor  $(\lambda_{gc})$ , mean chi-square statistics, estimated  $h_g^2$  and intercept under different simulation scenarios using the SIGMA cohort as described in Fig 2 and S10 Fig. Each estimate represents the mean  $h_g^2$  estimates from 100 simulations of 10,000 unrelated individuals. s.e. represents for standard error.

 S4 Table.
 Sample sizes (N) and number of SNPs (M) used in LD
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 calculation and heritability estimation of seven selected traits in the
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 23andMe cohort.
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S5 Table. mean of LD scores with varying window sizes for the 23 and Me cohort using LDSC and cov-LDSC. 10 PCs are included in all cov-LDSC estimates.

S6 Table. Heritability estimates of three quantitative and five binary809traits included in 23andMe and SIGMA cohorts using different LD models.810Stratified LD model uses genome-wide functional information from all SNPs and811explicitly models LD based on 53 functional annotations.812

S7 Table. Heritability estimates, mean chi-square statistics and genomic control inflation factor ( $\lambda_{gc}$ ) of three quantitative and four binary traits included in 23andMe using LDSC and cov-LDSC. cov-LDSC reports the

stratified LD model that uses genome-wide functional information from all SNPs and	816
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S8 Table. Pairwise heritability comparison for seven traits reported in the	818
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S10 Table. Tissue and type specific analysis on three traits in the	823
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S13 Table. Mean of LD scores with varying window sizes for the	833
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in all cov-LDSC estimates.	835
S14 Table. Pearson r-squared of LD scores with different window sizes	836
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genotypes. 10 PCs are included in all cov-LDSC estimates.	838
S15 Table. Pearson r-squared of LD scores with different window sizes	839
when using cov-LDSC in the SIGMA cohort. 10 PCs are included in all	840
cov-LDSC estimates.	841

S16 Table. Pearson r-squared of LD scores with different window sizes when using cov-LDSC in the 23andMe cohort. 10 PCs are included in all cov-LDSC estimates.

S17 Table. Difference between  $h_{common}^2$  and  $h_g^2$  in the SIGMA cohort for height, body mass index (BMI) and type 2 diabetes (T2D).

S1 Fig. LD score estimates with varying window size in populations from the 1000 Genomes project. LD score estimates with varying window size using unadjusted LDSC (orange) and cov-LDSC (blue) with 10 PCs with varying window size in both Europeans (N = 503, dashed line) and Admixed Americans (N = 347, solid line) from the 1000 Genomes Project. The x-axis shows the genomic window size used for estimating LD scores measured in centimorgan (cM). The y-axis shows the mean LD score estimates.

**S2** Fig. LD score estimates with varying window size and number of PCs in Admixed Americans included in the 1000 Genomes project. LD score estimates (y-axis) using different numbers of PCs at different window sizes (x-axis).

S3 Fig. Estimates of heritability  $(h_g^2)$  under different simulation scenarios 857 using the simulated genotypes reflecting a Latino population. LDSC (orange) 858 underestimated  $h_q^2$  and cov-LDSC (blue) yielded robust estimates under all settings. 859 Each boxplot represents the mean LD score estimate from 100 simulations of 10,000 860 unrelated individuals. For cov-LDSC, a window size of 5-cM with 10 PCs are used in all 861 scenarios. For LDSC, a window size of 5-cM are used in all scenarios. A true polygenic 862 quantitative trait with  $h_q^2 = 0.4$  is assumed for scenarios (a), (b) and (d). 1% causal 863 variants are assumed for (a) and (c) - (d). (b)-(d) assumed a dataset with an admixture 864 proportion of 50% from two different ancestral populations. (a)  $h_q^2$  estimation with 865 varying admixed proportions (x-axis) from two ancestral populations. (b)  $h_q^2$  estimation 866 with varying proportions of causal variants (0.01% - 50%). (c)  $h_q^2$  estimation with 867 varying heritability (0.05, 0.1, 0.2, 0.3, 0.4 and 0.5). (d)  $h_q^2$  estimation when an 868 environmental stratification component aligned with the first PC of the genotype data 869 is included in the phenotype simulation. 870

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S4 Fig. Estimates of heritability  $(h_g^2)$  in simulated genotypes reflecting an 871 African American population. LDSC (orange) underestimated and cov-LDSC 872 (blue) yielded less biased  $h_g^2$  estimates with varying admixed proportions (x-axis). Each 873 boxplot represents the mean LD score estimate from 100 simulations of 10,000 874 unrelated African American individuals. For cov-LDSC, a window size of 5-cM with 10 875 PCs are used in all scenarios. For LDSC, a window size of 5-cM are used in all scenarios. 876 A true polygenic quantitative trait with 1% causal variants and a true  $h_q^2 = 0.4$  is 877 assumed for scenarios. 878

S5 Fig. Estimates of heritability  $(h_g^2)$  in case-control phenotypes under 879 different simulation scenarios using the simulated genotypes reflecting a 880 **Latino population.**  $h_a^2$  estimation in a phylogenetic binary trait with assumed 881 prevalence of 0.1. 50,000 unrelated individuals are simulated in total. Each scenario has 882 5,000 cases and 5,000 controls.  $h_q^2$  estimation (a) with varying admixed proportions 883 (x-axis) from two ancestral populations; (b) with varying proportions of causal variants 884 (0.01% - 50%); (c) with varying heritability (0.05, 0.1, 0.2, 0.3, 0.4 and 0.5); and (d) 885 when an environmental stratification component aligned with the first PC of the 886 genotype data is included in the phenotype simulation. For cov-LDSC, a window size of 887 5-cM with 10 PCs are used in all scenarios. For LDSC, a window size of 5-cM are used 888 in all scenarios. 889

 S6 Fig. ADMIXTURE analysis (K = 5) of individuals included in the
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 SIGMA cohort and the 1000 Genomes Project. Each individual is represented
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 as a thin vertical bar. The colors can be interpreted as different ancestries. AFR
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 represents African; AMR represents Admixed American; EAS represents East Asian;
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 EUR represents European and SAS represents South Asian.
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# **S7 Fig. LD score estimates with varying window size in the SIGMA cohort.** LD score estimates using LDSC (orange) and cov-LDSC (blue) with varying window size in the SIGMA cohort (N = 8, 214). The x-axis shows the genomic window size used for estimating LD scores measured in centimorgan (cM). The y-axis shows the mean LD score estimates. For cov-LDSC, 10 PCs are used in all scenarios.

**S8 Fig. LD score estimates with varying window size and number of PCs in the SIGMA cohort.** LD score estimates (y-axis) using different number of PCs at different window sizes (x-axis).

S9 Fig. Estimates of heritability  $(h_g^2)$  with varying window sizes used in LD score estimation in the SIGMA cohort. cov-LDSC (blue) with 10 PCs and varying window size used to obtain LD score. We assumed a true  $h_g^2$  of 0.4 and 1% causal variant in each simulation. 100 replicates are used for each window size.

# S10 Fig. Intercept of estimated $h_g^2$ under different simulation scenarios using the SIGMA cohort as described in Figure 2. LDSC (orange)

underestimated  $h_g^2$  and cov-LDSC (blue) yielded less biased  $h_g^2$  estimates under all 909 settings. Each boxplot represents the mean LD score estimate from 100 simulations of 910 8,124 individuals included in the SIGMA project. For cov-LDSC, a window size of 911 20-cM with 10 PCs are used in all scenarios. For LDSC, a window size of 20-cM are 912 used in all scenarios. A true polygenic quantitative trait with  $h_q^2 = 0.4$  is assumed for 913 scenarios (a), (c) and (d). 1% causal variants are assumed for scenarios (b)-(d). (a) 914 Intercept with varying numbers of causal variants (0.01% - 50%). (b) Intercept with 915 varying heritability (0, 0.05, 0.1, 0.2, 0.3, 0.4 and 0.5). (c) Intercept with the presence of 916 an environmental stratification component aligned with the first PC of the genotype 917 data is included in the phenotype simulation. (d) Intercept when using a subset of total 918 samples and using admixed American samples included in the 1000 Genomes Project. 919

S11 Fig. Estimates of heritability  $(h_g^2)$  in simulated genotypes using LD scores estimated with varying sample sizes. cov-LDSC (blue) is used with varying sample sizes used to obtain LD scores. A random subset of 1%, 5%, 10% and 50% of the total samples (N = 10,000) in the simulated genotypes are used to calculate in-sample LD scores and then to obtain  $h_g^2$  estimates. LD scores are also obtained using independent genotypes (N = 1,000) using the perfect matching demographic model.

S12 Fig. Simulation results assessing type I error and power for LDSC 926 and cov-LDSC. We simulate a polygenic trait with  $h_g^2 = 0.5$ . LDSC (orange) shows 927 less power compared to cov-LDSC (blue) in detecting tissue. Each point shows the 928

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proportion of simulations (1,000 for each point) in which a null hypothesis of no tissue enrichment is rejected (Pr(rejected at P ; 0.05)), as a function of the z-score of total SNP heritability.

S13 Fig. LD score estimates with varying window size in populations from  $^{932}$ 23andMe. LD score estimates using unadjusted LDSC (orange) and cov-LDSC (blue)  $^{933}$ with 10 PCs with varying window size in both African Americans (N = 46, 844, dashed  $^{934}$ line) and Latinos (N = 161, 894, solid line) from the 23andMe cohort. The x-axis shows  $^{935}$ the genomic window size used for estimating LD scores measured in centimorgan (cM).  $^{936}$ The y-axis shows the mean LD score estimates.  $^{937}$ 

Tissue and cell type specific analysis with summary statistics in S14 Fig. 938 23 and Me Latinos using in-sample original LD and in-sample cov-LD for 939 **BMI.** The left panel (a) shows the tissue and cell type specific analysis using original 940 LDSC with in-sample LD scores; while the right panel (b) shows the tissue and cell type 941 specific analysis using cov-LDSC with in-sample cov-LD scores for BMI in 23andMe 942 cohort. The label on the top right in each plot indicates the number of significant tissue 943 type enrichments for each analysis. We observed no difference between LDSC and 944 cov-LDSC in European populations. In contrast, we observed more enrichment in and 945 around sets of genes that are specifically expressed in tissue- and cell-types using 946 cov-LDSC in Latinos and African Americans. 947

Tissue and cell type specific analysis with summary statistics in S15 Fig. 948 23andMe Latinos using in-sample original LD and in-sample cov-LD for 949 height. The left panel (a) shows the tissue and cell type specific analysis using original 950 LDSC with in-sample LD scores; while the right panel (b) shows the tissue and cell type 951 specific analysis using cov-LDSC with in-sample cov-LD scores for height in 23andMe 952 cohort. The label on the top right in each plot indicates the number of significant tissue 953 type enrichments for each analysis. We observed no difference between LDSC and 954 cov-LDSC in European populations. In contrast, we observed modest increased 955 enrichment using cov-LDSC in Latinos and African Americans. 956

Tissue and cell type specific analysis with summary statistics in S16 Fig. 957 23 and Me Latinos using in-sample original LD and in-sample cov-LD for 958 morning person. The left panel (a) shows the tissue and cell type specific analysis 959 using original LDSC with in-sample LD scores; while the right panel (b) shows the tissue 960 and cell type specific analysis using cov-LDSC with in-sample cov-LD scores for morning 961 person in 23andMe cohort. The label on the top right in each plot indicates the number 962 of significant tissue type enrichments for each analysis. We observed no difference 963 between LDSC and cov-LDSC in European populations. In contrast, we observed 964 modest increased enrichment using cov-LDSC in Latinos and African Americans. 965

S17 Fig. Heritability estimate with different number of PCs for GWAS 966 association test and LD score adjustment. We simulated the phenotypes on the 967 SIGMA cohort using additive model assuming 1% causal SNPs with. We performed 968 univariate cov-LDSC to measure heritability. We varied number of PCs included in 969 summary statistics and varied number of PCs used in cov-LDSC. The x-axis shows the 970 number of PCs included in the cov-LDSC calculation and the y-axis shows the number 971 of PCs included in the summary statistics calculation within the same sample. Numbers 972 in each cell represent the mean estimates from 100 replications. The color (from white 973 to red) represents the statistical difference between the estimated and the truth 974 (measured in  $-\log 10(P)$ ). A red cell indicates the  $h_q^2$  estimate is significantly different 975 from the truth. 976

S18 Fig. Type I error in tissue-type-specific enrichment when different 977 number of PCs are used to generate summary statistics and LD scores. We 978 generated 1,000 simulations for scenarios where there are different number of PCs (2, 5, 5)979 10, 20 and 50) included when calculating LD scores and generating summary statistics 980 (10 PCs) in the cell and tissue-specific enrichment analysis. We simulated a polygenic 981 trait with  $h_a^2 = 0.5$ . Each bar shows the proportion of simulations in which a null 982 hypothesis of no tissue enrichment is rejected ( $Pr(rejected \ at \ P \ 0.05)$ ), as a function of 983 the z-score of total SNP heritability. The horizontal red line indicates P = 0.05. 984

S19 Fig. LDSC and cov-LDSC with summary statistics derived from linear mixed models. Estimation of heritability (truth  $h_g^2 = 0.4$ ) using LDSC and cov-LDSC with 10 (blue) and 50 (green) PCs and a window size of 20-cM. Each boxplot represents the mean LD score estimate from 100 simulations of genotypes from the 8, 124 individuals included in the SIGMA cohort. All summary statistics are derived from linear mixed models with genetic relationship matrix (GRM) only or GRM with 10 genome-wide PCs using GEMMA [67].

S20 Fig. Results of multiple-tissue analysis for body mass index (BMI), height and type 2 diabetes (T2D) in the SIGMA cohort. Each point represents a tissue type from either the GTEx data set or the Franke lab data [64,65]. From left to right, (a)-(d) show multiple-tissue analysis for BMI when using LDSC and cov-LDSC with in-sample and out-of-sample LD reference panels. (e-h) show multiple-tissue analysis for height (e-h) when using LDSC and cov-LDSC with in-sample and out-of-sample LD reference panels. (i-l) show multiple-tissue analysis for T2D when using LDSC and cov-LDSC with in-sample and out-of-sample LD reference panels. (990

S21 Fig. Enrichment analysis using in-sample and out-of-sample LD 1000 reference panel. We simulated a polygenic trait with  $h_a^2 = 0.5$ . Similar power was 1001 obtained when using in-sample (obtained from the SIGMA cohort, turquoise) and 1002 out-of-sample (obtained from 1000 Genomes Admixed American (AMR) samples, red) 1003 reference panel. In both cases, type I error (at no (1x) enrichment) are well controlled. 1004 Each bar shows the proportion of simulations (1,000 for each point) in which a null 1005 hypothesis of no tissue enrichment is rejected ( $Pr(rejected at P \mid 0.05)$ ), as a function of 1006 the z-score of total SNP heritability. 1007

S22 Fig. Principal component analysis (PCA) of the SIGMA samples.1008Samples included in the SIGMA cohort projected onto the first two principal1009components using SNP weights precomputed from samples in the 1000 Genomes Phase 31010project using SNP weights. AFR represents Africans (green); AMR represents Admixed1011Americans (orange); EAS represents East Asians (yellow); EUR represents Europeans1012(blue); SAS represents South Asians (pink) and SIGMA samples are presented in gray.1013

S23 Fig. Tissue and cell type specific analysis with summary statistics in 1014 23andMe Latinos using in-sample cov-LD and out-of-sample cov-LD obtained using 1000G AMR samples. In sample LD is obtained in 23andMe1016Latinos with 20-cM window size and 10PCs. We observed cell type enrichments in both1017BMI and height using in-sample cov-LD. However, when we used out of sample 1000G1018AMR cov-LD with 20cM window size and 10PCs, we observed no cell type enrichments1019in either BMI and height.1020

S24 Fig. Principal component analysis (PCA) of the 23andMe samples.1021Samples included in the 23andMe cohort projected onto the first two principal1022components using SNP weights precomputed from samples in the 1000 Genomes Phase 31023project using SNPweights. AFR represents Africans (green); AMR represents Admixed1024Americans (red); EAS represents East Asians; EUR represents Europeans (blue); SAS1025represents South Asians (brown) and the 23andMe samples are presented in gray.1026

S1 Appendix. Mathematical framework of cov-LDSC

#### S2 Appendix. In-sample versus out-of-sample LD

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## S1 Appendix. Mathematical framework of cov-LDSC

Here, we will first provide a derivation of standard LD score regression that differs 1030 somewhat from published derivations, and in particular gives a mathematical 1031 interpretation for the value of the intercept. Then we will extend this derivation to 1032 cov-LDSC. 1033

#### S.1 Review of LD score regression without covariates

#### S.1.1 Summary statistics without covariates

We begin by describing the input data to LD score regression, which is the output of a standard GWAS.

In a standard GWAS of a quantitative trait, a marginal linear model is fit for each 1038 SNP *j*. Let *Y* denote the  $N \times 1$  vector of phenotypes and  $X_j$  denote the  $N \times 1$  vector 1039 of genotypes for SNP *j*, centered to mean zero. In the absence of covariates, we 1040 typically fit the model 1041

$$Y = X_j \beta_j^{(marg)} + \epsilon^{(marg)} \tag{1}$$

where  $\beta_j^{(marg)}$  is the marginal effect size of SNP j and  $\epsilon^{(marg)} \sim N(0, \sigma^2_{(marg)}I)$ .

The F-statistic, which at GWAS sample sizes is approximately chi-square distributed 1043 under the null and often referred to as the chi-square statistic, is equal to 1044

$$\chi_j^2 = \left(\hat{\beta}_j^{(marg)}\right)^2 / \hat{s}_j^2 \tag{2}$$

where

$$\hat{\beta}_j^{(marg)} = \frac{X_j^T Y}{X_j^T X_j}$$

and

$$\hat{s}_j^2 = \frac{\hat{\sigma}_{(marg)}^2}{X_j^T X_j},$$

where  $\hat{\sigma}^2_{(marg)}$  is an estimate of  $\sigma^2_{(marg)}$  that, if  $\hat{\beta}^{(marg)}_j$  is small, satisfies

$$^{2}_{(marg)} \approx \frac{1}{N} Y^{T} Y.$$

We will assume that  $\beta_j^{(marg)}$  and its estimate  $\hat{\beta}_j^{(marg)}$  are indeed small, so that this is a 1045 valid approximation.

Let 
$$V(X_j) = X_j^T X_j / N$$
 and  $V(Y) = Y^T Y / N$  be the empirical variances of  $X_j$  and 1047  
Y, and let  $\tilde{X}_j = X_j / \sqrt{V(X_j)}$ , and  $\tilde{Y} = Y / \sqrt{V(Y)}$  be  $X_j$  and Y, normalized to 1048

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empirical variance one. Note that when  $X_j$  and Y are random, so are  $V(X_j), V(Y), \tilde{X}_j$ , 1049 and  $\tilde{Y}$ . Note also that  $\tilde{X}_j^T \tilde{X}_j = \tilde{Y}^T \tilde{Y} = N$ , deterministically. We can now simplify the 1050 expression for  $\chi_j^2$ : 1051

$$\chi_j^2 \approx \frac{1}{N} (\tilde{X}_j^T \tilde{Y})^2 \tag{3}$$

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We will assume that we have as input  $\chi_j^2$  for a genome-wide set of SNPs j.

#### S.1.2 The polygenic model

In LD score regression, we take these chi-square statistics as input, and we derive their <sup>1054</sup> expectation under a standard polygenic model. Specifically, instead of the marginal <sup>1055</sup> model used in GWAS, LD score regression is based on a joint model with random SNP <sup>1056</sup> effect sizes: <sup>1057</sup>

$$Y = X\beta + \epsilon \tag{4}$$

where Y is the phenotype vector,  $X = (X_1 \dots X_M)$  is the  $N \times M$  genotype matrix,  $\epsilon \sim \mathcal{N}(0, \sigma_{\epsilon}^2 I)$ , and  $\beta$  is the  $M \times 1$  vector of joint effect sizes. Let  $\tilde{\beta}_j = \beta_j \sqrt{V(X_j)}$ , and note that  $X\beta = \tilde{X}\tilde{\beta}$ . We will model  $\tilde{\beta}_j$  as random with mean zero, independent of each other and of  $\epsilon$ . Here, we will perform derivations in which  $Var(\tilde{\beta}_j) = \sigma_{\tilde{\beta}}^2$ ; these derivations extend easily to the case in which  $Var(\tilde{\beta}_j)$  depends on functional annotations. We don't specify a distribution for  $\tilde{\beta}$ .

In LD score regression, we derive the expectation of  $\chi_j^2$  under this polygenic model, 1064 and we use the resulting equation to estimate parameters such as  $\sigma_{\tilde{\beta}}^2$ . Because X is not 1065 observed, we ultimately treat it as random. Here, we will derive  $E[\chi_j^2]$  by first deriving 1066  $E[\chi_j^2|X]$  and then using the law of total expectation to remove the conditioning on X. 1067

#### **S.1.3** Deriving the expression for $E[\chi_i^2|X]$

Before deriving the expression for  $E[\chi_j^2|X]$ , we will first derive the expected empirical variance of Y, where the variance is over the random individuals in our GWAS and the

expectation is over random  $\beta$  and  $\epsilon$ , conditional on X.

$$\begin{split} E[V(Y)|X] &= \frac{1}{N} E\left[ \left( X\beta + \epsilon \right)^T \left( X\beta + \epsilon \right) |X \right] \\ &= \frac{1}{N} E\left[ \left( \tilde{X}\tilde{\beta} + \epsilon \right)^T \left( \tilde{X}\tilde{\beta} + \epsilon \right) |X \right] \\ &= \frac{1}{N} E\left[ \tilde{\beta}^T \tilde{X}^T \tilde{X} \tilde{\beta} |X \right] + \frac{1}{N} E\left[ \epsilon^T \epsilon \right] \\ &= \frac{1}{N} \sum_{j,k} E\left[ \tilde{\beta}_j (\tilde{X}^T \tilde{X})_{j,k} \tilde{\beta}_k |X \right] + \sigma_\epsilon^2 \\ &= \frac{1}{N} \sum_{j \neq k} E\left[ \tilde{\beta}_j \right] E\left[ \tilde{\beta}_k \right] (\tilde{X}^T \tilde{X})_{j,k} + \frac{1}{N} \sum_j E\left[ \tilde{\beta}_j^2 \right] (\tilde{X}^T \tilde{X})_{j,j} + \sigma_\epsilon^2 \\ &= 0 + \frac{1}{N} \sum_j \sigma_{\tilde{\beta}}^2 (\tilde{X}^T \tilde{X})_{j,j} + \sigma_\epsilon^2 \\ &= M \sigma_{\tilde{\beta}}^2 + \sigma_\epsilon^2 \end{split}$$

We will let  $h_g^2$  denote  $M\sigma_{\tilde{\beta}}^2/E[V(Y)|X]$ , noting that definitions of heritability depend on the model on which they are based, and so  $h_g^2$  as used here is a different value than in a model in which  $\beta$  is fixed.

It will also be useful to have

$$E\left[\left(\tilde{X}_{j}^{T}\epsilon\right)^{2}|X\right] = E\left[\tilde{X}_{j}^{T}\epsilon\epsilon^{T}\tilde{X}_{j}|X\right]$$
$$= \tilde{X}_{j}^{T}E\left[\epsilon\epsilon^{T}\right]\tilde{X}_{j}$$
$$= \sigma_{\epsilon}^{2}\tilde{X}_{j}^{T}\tilde{X}_{j}$$
$$= N\sigma_{\epsilon}^{2}$$

We can now derive the expected chi-square statistic:

$$\begin{split} E[\chi_j^2|X] &= E\left[\frac{1}{N}\left(\tilde{X}_j^T\tilde{Y}\right)^2|X\right] \\ &= E\left[\frac{1}{NV(Y)}\left(\tilde{X}_j^T\left(X\beta + \epsilon\right)\right)^2|X\right] \\ &\approx \frac{1}{NE[V(Y)|X]}E\left[\left(\tilde{X}_j^T\left(X\beta + \epsilon\right)\right)^2|X\right] \\ &= \frac{1}{NE[V(Y)|X]}E\left[\left(\tilde{X}_j^T\left(\tilde{X}\tilde{\beta} + \epsilon\right)\right)^2|X\right] \\ &= \frac{1}{NE[V(Y)|X]}E\left[\left(\sum_k \tilde{X}_j^T\tilde{X}_k\tilde{\beta}_k + \tilde{X}_j^T\epsilon\right)^2|X\right] \\ &= \frac{N}{E[V(Y)|X]}\sum_k \left(\frac{\tilde{X}_j^T\tilde{X}_k}{N}\right)^2 E[\tilde{\beta}_k^2] + \frac{1}{NE[V(Y)|X]}E\left[\left(\tilde{X}_j^T\epsilon\right)^2|X\right] \\ &= \frac{N\sigma_{\tilde{\beta}}^2}{E[V(Y)|X]}\sum_k \left(\frac{\tilde{X}_j^T\tilde{X}_k}{N}\right)^2 + \frac{\sigma_{\epsilon}^2}{E[V(Y)|X]} \\ &= \frac{N\sigma_{\tilde{\beta}}^2}{E[V(Y)|X]}\sum_k \left(\left(\frac{\tilde{X}_j^T\tilde{X}_k}{N}\right)^2 - \frac{1}{N}\right) + \frac{M\sigma_{\tilde{\beta}}^2}{E[V(Y)|X]} + \frac{\sigma_{\epsilon}^2}{E[V(Y)|X]} \\ &= N\frac{h_g^2}{M}\sum_k \left(\left(\frac{\tilde{X}_j^T\tilde{X}_k}{N}\right)^2 - \frac{1}{N}\right) + 1 \end{split}$$

#### **S.1.4** Removing the conditioning on X

When analyzing summary statistics, we do not have access to the true value of X, and 1073 so we need to compute the expectation of  $\chi_i^2$  treating X as random and integrating it 1074 out. To do this, we use the law of total expectation, and so the relevant quantity is 1075 . We would like our method to be applicable in the most general  $E\left|\left(\frac{\tilde{X}_{j}^{T}\tilde{X}_{k}}{N}\right)\right.$ 1076 circumstances, and so we do not want to assume a particular distribution on X, or even 1077 that its rows are drawn i.i.d. from some distribution. Instead, we will let  $W_j$  denote the 1078 set of SNPs in an LD window around j, and we will make three assumptions that will 1079 allow us to complete our derivation: 1080

1. There is a c such that for  $k \notin W_j$ , we have  $E\left[\left(\frac{\tilde{X}_j^T \tilde{X}_k}{N}\right)^2\right] \approx c$ , and the

approximation is good enough that  $N \frac{h_g^2}{M} \sum_{k \notin W_j} \left( E\left[ \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 \right] - c \right)$  is 1082 negligible. If there is no structure or relatedness in our samples (and if N is high 1083 enough that the difference between standardization in the population and in our 1084

sample is negligible), then c can be shown to be 1/N.

2. For  $k \in W_j$ , there is a value  $R_{jk}$  satisfying  $R_{jk} \approx E\left[\left(\frac{\tilde{X}_j^T \tilde{X}_k}{N}\right)^2\right] - c$ , where the 1086 approximation is good enough that  $N\frac{h_g^2}{M}\sum_{k\in W_j}\left(E\left[\left(\frac{\tilde{X}_j^T \tilde{X}_k}{N}\right)^2\right] - c - R_{jk}^2\right)$  is 1087 negligible. Note that if the rows of X are drawn i.i.d. from some distribution and 1088  $R_{jk}$  is the correlation between SNPs j and k in this underlying distribution, and if 1089  $|W_j|$  is small compared to M, then this condition in satisfied. 1090

We can now apply the law of total expectation to complete the derivation:

$$\begin{split} E[\chi_j^2] &\approx N \frac{h_g^2}{M} \sum_k \left( E\left[ \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 \right] - \frac{1}{N} \right) + 1 \\ &= N \frac{h_g^2}{M} \sum_k \left( E\left[ \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 \right] - c \right) + N \frac{h_g^2}{M} \sum_k \left( c - \frac{1}{N} \right) + 1 \\ &\approx N \frac{h_g^2}{M} \sum_{k \in W_j} \left( E\left[ \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 \right] - c \right) + N h_g^2 \left( c - \frac{1}{N} \right) + 1 \\ &\approx N \frac{h_g^2}{M} \sum_{k \in W_j} R_{jk}^2 + N h_g^2 \left( c - \frac{1}{N} \right) + 1 \\ &= N \frac{h_g^2}{M} \sum_{k \in W_j} R_{jk}^2 + N a + 1, \end{split}$$

where  $a = h_q^2(c - 1/N)$ . Letting

$$\ell_j = \sum_{k \in W_j} R_{jk}^2$$

denote the LD score of SNP j, we obtain the main LD score regression equation:

$$E[\chi_j^2] \approx N \frac{h_g^2}{M} \ell_j + Na + 1.$$
(5)

We typically estimate  $\ell_j$  using a reference panel, and we estimate  $h_g^2$  via weighted 1092 regression of  $\chi_i^2$  on  $\ell(j)$ , evaluating significance with block jackknife across SNPs. 1093

### S.2 LD score regression in the presence of covariates

We will now discuss LD score regression for a quantitative trait, in the presence of 1095 covariates. For a treatment of LD score regression for case-control traits with covariates, 1096 see [Weissbrod et al. 2018 AJHG]. 1097

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#### S.2.1 Summary statistics

Let C denote an  $N \times K$  matrix of covariates, each column centered to mean zero. In a 1099 GWAS of a quantitative trait with covariates, we typically fit the model 1100

$$Y = X_j \beta_{SNP,j}^{(marg)} + C \beta_{cov,j}^{(marg)} + \epsilon_j^{(marg)}$$
(6)

where  $\beta_{SNP,j}^{(marg)}$  is the marginal effect size of SNP j and  $\beta_{cov,j}^{(marg)}$  is the effect size vector of 1101 the covariates.

The chi-square statistic is equal to

$$\chi_j^2 = \left(\hat{\beta}_{SNP,j}^{(marg)}\right)^2 / \hat{s}_j^2,\tag{7}$$

where  $\hat{\beta}_{SNP,j}^{(marg)}$  is the least-squares estimate of  $\beta_{SNP,j}^{(marg)}$ , and

$$\hat{s}_j^2 = \hat{\sigma}_{(marg)}^2 (A_j^T A_j)^{-1}{}_{11},$$

where  $A_j$  is the design matrix, given by  $A_j = (X_j \ C)$ , where  $(A_j^T A_j)^{-1}_{11}$  denotes the upper left entry of the matrix  $(A_j^T A_j)^{-1}$ , and where  $\hat{\sigma}^2_{(marg),j}$  is again an estimate of  $\sigma^2_{(marg),j}$ .

Let  $P = I - C(C^T C)^{-1} C^T$ . By the Frisch-Waugh-Lovell theorem, we have

$$\hat{\beta}_{SNP,j}^{(marg)} = \frac{(PX_j)^T PY}{(PX_j)^T PX_j},$$

and by block matrix inversion, we have

$$(A_j^T A_j)_{11}^{-1} = \frac{1}{(PX_j)^T (PX_j)}.$$

Again assuming that the effect size  $\beta_{SNP,j}^{(marg)}$  is small, we have

$$\hat{\sigma}^2_{(marg)} \approx \frac{1}{N} (PY)^T PY.$$

Let 
$$V(PX_j) = ((PX_j)^T PX_j)/N$$
 and  $V(PY) = (PY)^T PY/N$ , and let

$$\tilde{X}_j = PX_j/\sqrt{V(PX_j)}$$
, and  $\tilde{Y} = PY/\sqrt{V(PY)}$ . Then, we can rewrite:  
 $\chi_j^2 \approx \frac{1}{N} \left(\tilde{X}_j^T \tilde{Y}\right)^2$ 
(8)

#### **S.2.2** Deriving the expression for $E[\chi_j^2|X]$

In cov-LDSC, we assume that there are covariates in our GWAS model (Eq (1)) and we include the same set of covariates in the polygenic model that we would like to fit:

$$Y = X\beta + C\beta_{cov} + \epsilon, \tag{9}$$

where Y, X,  $\beta$ , C, and  $\epsilon$  are as before. Note that under this polygenic model,

$$PY = PX\beta + P\epsilon.$$

Let  $\tilde{\beta}_j = \beta_j \sqrt{V(X_j)}$ . Note that  $PX\beta = \tilde{X}\tilde{\beta}$ . We will model  $\tilde{\beta}_j$  as random with

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mean zero and variance  $\sigma_{\tilde{\beta}}^2$ . Now we have

$$\begin{split} E[V(PY)|X] &= \frac{1}{N} E[(PY)^T PY|X] \\ &= \frac{1}{N} E\left[ (PX\beta + P\epsilon)^T (PX\beta + P\epsilon) |X \right] \\ &= \frac{1}{N} E\left[ (\tilde{X}\tilde{\beta} + P\epsilon)^T (\tilde{X}\tilde{\beta} + P\epsilon) |X \right] \\ &= \frac{1}{N} E[\tilde{\beta}^T \tilde{X}^T \tilde{X}\tilde{\beta} |X] + \frac{1}{N} E[(\epsilon^T P^T P\epsilon] \\ &= \frac{1}{N} \sum_{j,k} E\left[ \tilde{\beta}_j (\tilde{X}^T \tilde{X})_{j,k} \tilde{\beta}_k |X \right] + \frac{1}{N} \sum_{j,k} E\left[ \epsilon_j \left( P^T P \right)_{j,k} \epsilon_k \right] \\ &= \frac{1}{N} \sum_{j \neq k} E\left[ \tilde{\beta}_j \right] E\left[ \tilde{\beta}_k \right] (\tilde{X}^T \tilde{X})_{j,k} + \frac{1}{N} \sum_j E\left[ \tilde{\beta}_j^2 \right] (\tilde{X}^T \tilde{X})_{j,j} \\ &+ \frac{1}{N} \sum_{j \neq k} E\left[ \epsilon_j \right] E\left[ \tilde{\epsilon}_k \right] (P^T P)_{j,k} + \frac{1}{N} \sum_j E\left[ \epsilon_j^2 \right] (P^T P)_{j,j} \\ &= 0 + \frac{1}{N} \sum_j \sigma_{\tilde{\beta}}^2 (\tilde{X}^T \tilde{X})_{j,j} + \sigma_{\epsilon}^2 + 0 + \frac{1}{N} \sum_j \sigma_{\epsilon}^2 (P^T P)_{j,j} \\ &= M \sigma_{\tilde{\beta}}^2 + \sigma_{\epsilon}^2 \frac{N - K}{N} \end{split}$$

where K is the rank of C. If K is small compared to N, as is typical of most GWAS, then we can say that

$$E[V(PY)|X] \approx M\sigma_{\tilde{\beta}}^2 + \sigma_{\epsilon}^2.$$

We will let  $h_g^2$  denote  $M\sigma_{\tilde{\beta}}^2/E[V(PY)|X]$ . It will again be convenient to have

$$\begin{split} E\left[(\tilde{X}_{j}^{T}P\epsilon)^{2}|X\right] &= E\left[\left(\frac{1}{\sqrt{V(PX_{j})}}X_{j}^{T}P^{T}P\epsilon\right)^{2}|X\right] \\ &= E\left[\left(\frac{1}{\sqrt{V(PX_{j})}}X_{j}^{T}P^{T}\epsilon\right)^{2}|X\right] \\ &= E\left[\left(\tilde{X}_{j}^{T}\epsilon\right)^{2}|X\right] \\ &= E\left[\left(\tilde{X}_{j}^{T}\epsilon\right)^{2}|X\right] \\ &= \tilde{X}_{j}^{T}E\left[\epsilon\epsilon^{T}\right]\tilde{X}_{j} \\ &= \sigma_{\epsilon}^{2}\tilde{X}_{j}^{T}\tilde{X}_{j} \\ &= N\sigma_{\epsilon}^{2}. \end{split}$$

Now we have:

$$\begin{split} E[\chi_j^2|X] &\approx \frac{1}{N} E\left[ \left( \tilde{X}_j^T \tilde{Y} \right)^2 |X \right] \\ &= E\left[ \frac{1}{NV(PY)} \left( \tilde{X}_j^T PY \right)^2 |X \right] \\ &\approx \frac{1}{NE\left[ V(PY) |X \right]} E\left[ \left( \tilde{X}_j^T (PX\beta + P\epsilon) \right)^2 |X \right] \\ &= \frac{1}{NE\left[ V(PY) |X \right]} E\left[ \left( \tilde{X}_j^T (\tilde{X}\tilde{\beta} + P\epsilon) \right)^2 |X \right] \\ &= \frac{1}{NE\left[ V(PY) |X \right]} \sum_k (\tilde{X}_j^T \tilde{X}_k)^2 E\left[ \tilde{\beta}_k^2 \right] + \frac{1}{NE\left[ V(PY) |X \right]} E\left[ (\tilde{X}_j^T P\epsilon)^2 |X \right] \\ &= \frac{N\sigma_{\beta}^2}{E\left[ V(PY) |X \right]} \sum_k \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 + \frac{\sigma_{\epsilon}^2}{E\left[ V(PY) |X \right]} \\ &= \frac{N\sigma_{\beta}^2}{E\left[ V(PY) |X \right]} \sum_k \left( \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 - \frac{1}{N} \right) + \frac{M\sigma_{\beta}^2}{E\left[ V(PY) |X \right]} + \frac{\sigma_{\epsilon}^2}{E\left[ V(PY) |X \right]} \\ &\approx \frac{Nh_g^2}{M} \sum_k \left( \left( \frac{\tilde{X}_j^T \tilde{X}_k}{N} \right)^2 - \frac{1}{N} \right) + 1 \end{split}$$

#### **S.2.3** Removing the conditioning on X

We will make the same two assumptions as for LD score regression without covariates. 1115

- 1. There is a c such that for  $k \notin W_j$ , we have  $E\left(\frac{X_j^T X_k}{N}\right)^2 \approx c$ . One way to formalize the notion that C captures all structure in X is that c = 1/N in this case.
- 2. For  $k \in W_j$ , we have access, for example from a reference panel, to an estimate  $R_{jk}$  satisfying  $R_{jk} \approx E\left(\frac{X_j^T X_k}{N}\right)^2 c$ . When X contains admixture or other marked for a seturated from a reference panel may not suffice. In that marked case, we can set  $R_{jk}$  to be  $\left(\frac{\tilde{X}_j^T \tilde{X}_k}{N}\right)^2$ , or an estimate of that quantity from a marked from a reference if window size is 30 cM, marked for the GWAS. We note also that even if window size is 30 cM, marked for the still only approximately 1% of the genome, and so  $|W_j|$  is still small marked for M.

With these assumptions satisfied, the rest of the derivation is identical to the case <sup>1125</sup> without covariates. <sup>1126</sup>

#### S2 Appendix. In-sample versus out-of-sample LD

To test the reliability of using an out-of-sample reference LD panel for cov-LDSC 1128 applications, we first examined the performance of out-of-sample LD scores obtained 1129 from 1,000 samples with a perfectly matching demographic history in the simulated 1130 genotypes. cov-LDSC yielded less biased estimates when using 1,000 samples in an 1131 out-of-sample reference panel with a perfectly matching population structure (S11 Fig). 1132 Next, we tested the accuracy of heritability estimates and type I error of enrichment 1133 analysis when using 1000 Genomes Project [20] Admixed American (AMR) samples to 1134 obtain out-of-sample LD scores. When using the AMR panel as a reference panel for 1135 the SIGMA cohort, we observed a less biased  $h_a^2$  estimate (P = 0.33, Fig 2(d)), 1136 However, as we decreased the number of samples included in the subsampling, the 1137 cov-LDSC regression intercepts deviated further from one (S10 Fig(d)). This is 1138 probably due to attenuation bias from noisily estimated LD scores at N < 1,000. We 1139 observed similar tissue type specific enrichment results for BMI, height and T2D (S20 1140 Fig). We further assessed the power and biases of using 1000 Genomes AMR samples as 1141 an external reference panel when applying it in the SIGMA cohort for tissue type 1142 specific analysis via simulation. We observed well calibrated type I error and similar 1143 power compared to in-sample LD reference panel (S21 Fig). This suggested that the 1144 AMR panel included in the 1000 Genomes Project has similar demographic history 1145 compared to the SIGMA cohort (S6 Fig, S22 Fig). 1146

Next, we explored the feasibility of applying 1000 Genomes AMR samples in 1147 heritability estimation and its enrichment analyses in the 23andMe cohort. We obtained 1148 stratified LD scores using 1000 Genomes AMR samples (N = 347) and applied it on 1149 summary statistics obtained from 23 and Me. In contrast to the SIGMA cohort, we 1150 discovered total heritability estimates are significantly different from those estimated 1151 using in-sample LD scores (S12 Table) and discovered no significant tissue type 1152 enrichment (S23 Fig). This suggested that 1000 Genome AMR samples might have 1153 different demographic history compared to 23 and Me samples (S24 Fig). 1154

We therefore caution that when using 1000 Genomes or any out-of-sample reference <sup>1155</sup> panels for a specific admixed cohort, users should ensure that the demographic histories <sup>1156</sup> are shared between the reference and the study cohort. We highly recommend <sup>1157</sup>

computing in-sample LD scores on a randomly chosen subset of at least 1,000 individuals <sup>1158</sup> from a GWAS. We also strongly encourage cohorts to release their summary statistics <sup>1159</sup> and in-sample covariate-adjusted LD scores at the same time to facilitate future studies. <sup>1160</sup>

# Figure and table legends

Fig 1. Overview of the covariate-adjusted LD score regression. (a) As input, cov-LDSC takes raw genotypes of collected GWAS samples and their global principal components. (b) cov-LDSC regresses out the ancestral components based on global principal components from the LD score calculation and corrects for long-range admixture LD. Black and red lines indicate estimates before and after covariate adjustment respectively (c) Adjusted heritability estimation based on GWAS association statistics (measured by  $\chi^2$ ) and covariate-adjusted LD scores. (d) Estimation of heritability enrichment in tissue-specific gene sets.



> Fig 2. Estimates of heritability  $(h_g^2)$  under different simulation scenarios using the SIGMA cohort. LDSC (orange) underestimated  $h_g^2$  and cov-LDSC (blue) yielded robust  $h_g^2$  estimates under all settings. Each boxplot represents the mean LD score estimate from 100 simulated phenotypes using the genotypes of 8,214 unrelated individuals from the SIGMA cohort. We used a window size of 20-cM in both LDSC and cov-LDSC, and 10 PCs were included in cov-LDSC in all scenarios. A true polygenic quantitative trait with  $h_g^2 = 0.4$  is assumed for scenarios (a), (c) and (d) and 1% causal variants are assumed for scenarios (b)-(d). (a)  $h_g^2$  estimation with varying proportions of causal variants (0.01% - 30%). (b)  $h_g^2$  estimation with varying heritabilities (0, 0.05, 0.1, 0.2, 0.3, 0.4 and 0.5). (c)  $h_g^2$  estimation when an environmental stratification component aligned with the first PC of the genotype data was included in the phenotype simulation. (d)  $h_g^2$  estimation when using a subset of the cohort to obtain LD score estimates and using out-of-sample LD score estimates obtained from Admixed Americans included in the 1000 Genomes Project [20].



Fig 3. Estimates of heritability  $(h_g^2)$  of three quantitative and four dichotomous traits in two admixed populations in the 23andMe research cohort. For seven selected non-disease phenotypes (body mass index (BMI), height, age at menarche, left handedness, morning person, motion sickness and nearsightedness) in the 23andMe cohort, we reported their estimated genetic heritability and intercepts (and their standard errors) using the baseline model. LD scores were calculated using 134,999, 161,894, 46,844 individuals from 23andMe European, Latino and African American individuals respectively. For each trait, we reported the sample size in obtained summary statistics used in cov-LDSC. For BMI and height, we also reported the  $h_q^2$  estimates from the SIGMA cohort.

Trait	Heritability (s.e.)	Sample size	Intercept (s.e.)
		8,124	1.02 (0.01)
BMI		125,465	1.02 (0.01)
	H	130,866	1.11 (0.02)
		40,454	1.00 (0.01)
		8,124	1.07 (0.01)
hoight	<b>⊢</b> •∣	125,465	1.07 (0.03)
neight		130,866	1.13 (0.03)
	1 <del>- 1</del>	40,454	1.00 (0.01)
		95.663	1.02 (0.01)
age at menarche		17.679	1.04 (0.01)
_		12,419	1.00 (0.01)
		121 271	1 01 (0 01)
left handedness		94 786	1.01 (0.01)
		42,328	0.99 (0.01)
		94.015	1.02 (0.01)
morning person		100.409	1.03 (0.01)
		29,966	1.00 (0.01)
		102 281	1 03 (0 02)
motion sickness		17 894	1 02 (0.02)
		13,491	1.00 (0.01)
		117.059	1 04 (0 02)
nearsightedness	Hel	117,258	1.04 (0.02)
near signteuriess		22 581	1.02 (0.01)
		22,301	1.01 (0.01)
	0.0 0.2 0.4 0. Heritablilty (standard error)	6	
23andMe African Ame	ricans 🖝 23andMe Latinos 🔶 23andMe Europeans 🛖 SIC	-ima	

## Fig 4. Results of multiple-tissue analysis for height, BMI and morning

**person.** Each point represents a tissue type from either the GTEx data set or the Franke lab data set as defined in Finucane et al [9]. From top to bottom, (a)-(d) show multiple-tissue analysis for BMI in the cross-population meta-analysis and in Europeans, Latinos and African Americans respectively. (e) shows the scatter plot of the estimated per-standardized-annotation effect size  $\tau^*$ , which represents the proportional change of averaged per-SNP heritability for one standard deviation increase in value of the annotation of each cell type, conditional on other 53 non-cell type specific baseline annotations, in the three populations for all tested tissue types (**Methods**). The x-axis shows the  $\tau^*$  in European populations and the y-axis shows either  $\tau^*$  in Latinos (blue) or African Americans (orange). We reported the slope and p-value when we regress Latinos (blue) and African Americans (orange)  $\tau^*$  on Europeans  $\tau^*$  for all tissue types. Error bars indicate standard errors of  $\tau^*$ . Similarly, the results are shown in (f)-(j) for height and (k)-(n) for morning person. The significance threshold in plots (a)-(d), (f-i) and (k-m) is defined by the FDR < 5% cutoff,  $-\log_{10}(p) = 2.75$ . Numerical results are reported in S10 Table.



Table 1. Heritability estimates of height, BMI and type 2 diabetes using different estimation methods. Reported values are estimates of  $h_g^2$  (with standard deviations in brackets) from LDSC using a 20-cM window, cov-LDSC using a 20-cM window and 10 PCs, and GCTA using REAP [26] to obtain the genetic relationship matrix with adjustment by 10 PCs. The final column provides reported  $h_g^2$  estimates in European populations from various studies [12, 24, 25].

Phenotype	LDSC	cov-LDSC	GCTA	Public
	(baseline)	(baseline)	(REAP)	
Height	0.159(0.037)	$0.379\ (0.079)$	0.450(0.042)	0.450-0.685 [12,24]
BMI	0.113(0.030)	0.248(0.061)	0.235(0.041)	0.246-0.270 [24]
T2D	$0.121 \ (0.035)$	0.263(0.073)	0.376(0.046)	0.139-0.414 [24,25]