An Atlas of Genetic Correlations across Human Diseases and Traits

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Abstract

Identifying genetic correlations between complex traits and diseases can provide useful etiological insights and help prioritize likely causal relationships. The major challenges preventing estimation of genetic correlation from genome-wide association study (GWAS) data with current methods are the lack of availability of individual genotype data and widespread sample overlap among meta-analyses. We circumvent these difficulties by introducing a technique for estimating genetic correlation that requires only GWAS summary statistics and is not biased by sample overlap. We use our method to estimate 300 genetic correlations among 25 traits, totaling more than 1.5 million unique phenotype measurements. Our results include genetic correlations between anorexia nervosa and schizophrenia/ body mass index and associations between educational attainment and several diseases. These results highlight the power of a polygenic modeling framework, since there currently are no genome-wide significant SNPs for anorexia nervosa and only three for educational attainment.

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Introduction

Understanding the complex relationships between human behaviours, traits and diseases is a fundamental goal of epidemiology. In the absence of randomized controlled trials and longitudinal studies, many disease risk factors are identified on the basis of population cross-sectional correlations of variables at a single time point. Such approaches can be biased by confounding and reverse causation, leading to spurious associations [1, 2]. Genetics can help elucidate cause or effect, since inherited genetic effects cannot be subject to reverse causation and are biased by a smaller list of confounders.

The first methods for testing for genetic overlap were family studies [3–7]. The disadvantage of these methods is the requirement to measure all traits on the same individuals, which scales poorly to studies of multiple traits, especially traits that are difficult or costly to measure (e.g., low-prevalence diseases). Genome-wide association studies (GWAS) produce effect-size estimates for specific genetic variants, so it is possible to test for shared genetics by looking for correlations in effect-sizes across traits, which does not require measuring multiple traits per individual.

A widely-used technique for testing for relationships between phenotypes using GWAS data is Mendelian randomization (MR) [1, 2], which is the specialization to genetics of instrumental variables [8]. MR is effective for traits where significant associations account for a substantial fraction of heritability [9, 10]. For many complex traits, heritability is distributed over thousands of variants with small effects, and the proportion of heritability accounted for by significantly associated variants at current sample sizes is small [11]. For such traits, MR suffers from low power and weak instrument bias [8,12].

A complementary approach is to estimate genetic correlation, a quantity that includes the effects of all SNPs, including those that do not reach genome-wide significance (Methods). Genetic correlation is also meaningful for pairs of diseases, in which case it can be interpreted as the genetic analogue of comorbidity. The two main existing techniques for estimating genetic correlation from GWAS data are restricted maximum likelihood (REML) [13–18] and polygenic scores [19,20]. These methods have only been applied to a few traits, because they require individual genotype data, which are difficult to obtain due to informed consent limitations.

In response to these limitations, we have developed a technique for estimating genetic correlation using only GWAS summary statistics that is not biased by sample overlap. Our method is based on LD Score regression [21] and is computationally very fast. We apply this method to data from 25 GWAS and report genetic correlations for 300 pairs of phenotypes, demonstrating shared genetic bases for many complex diseases and traits.

Results

Overview of Methods

The method presented here for estimating genetic correlation from summary statistics relies on the fact that the GWAS effect-size estimate for a given SNP incorporates the effects of all SNPs in linkage disequilibrium (LD) with that SNP [21,22]. For a polygenic trait, SNPs with high LD will have higher $\chi^2$ statistics on average than SNPs with low LD [21]. A similar relationship holds if we replace $\chi^2$ statistics for a single study with the product of $z$-scores from two studies of traits with non-zero genetic correlation. Precisely, under a polygenic model [13,15], the expected value of
$z_{1j} z_{2j}$ is

$$
\mathbb{E}[z_{1j} z_{2j}] = \frac{\sqrt{N_1 N_2} \rho_g}{M} \ell_j + \frac{\rho N_s}{\sqrt{N_1 N_2}},
$$

(1)

where $N_i$ is the sample size for study $i$, $\rho_g$ is genetic covariance (defined in Methods), $\ell_j$ is LD Score [21], $N_s$ is the number of individuals included in both studies, and $\rho$ is the phenotypic correlation among the $N_s$ overlapping samples. We derive this equation in the Supplementary Note. If study 1 and study 2 are the same study, then Equation 1 reduces to the single-trait result from [21], because genetic covariance between a trait and itself is heritability, and $\chi^2 = z^2$. As a consequence of equation 1, we can estimate genetic covariance using the slope from the regression of $z_{1j} z_{2j}$ on LD Score, which is computationally very fast (Methods). If there is sample overlap, it will only affect the intercept from this regression (the term $\rho N_s/\sqrt{N_1 N_2}$) and not the slope, so the estimates of genetic correlation will not be biased by sample overlap. Similarly, population stratification will alter the intercept but have minimal impact on the slope, for the same reasons that population stratification has minimal impact on the slope from single-trait LD Score regression [21]. If we know the amount of sample overlap and phenotypic correlation in advance (i.e., the true value of $\rho N_s/\sqrt{N_1 N_2}$), we can constrain the intercept to this value, which reduces the standard error. We refer to this estimator as constrained intercept LD Score regression. Normalizing genetic covariance by the heritabilities yields genetic correlation: $r_g := \rho_g/\sqrt{h_1^2 h_2^2}$, where $h_i^2$ denotes the SNP-heritability [13] from study $i$. Genetic correlation ranges between $-1$ and $1$. Similar results hold if one or both studies is a case/control study, in which case genetic covariance is on the observed scale. There is no distinction between observed and liability scale genetic correlation for case/control traits, so we can talk about genetic correlation between a case/control trait and a quantitative trait and genetic correlation between pairs of case/control traits without difficulties (Supplementary Note).

**Simulations**

We performed a series of simulations to evaluate the robustness of the model to potential confounders such as sample overlap and model misspecification, and to verify the accuracy of the standard error estimates. Details of our simulation setup are provided in the methods. Table 1 shows LD Score regression estimates and standard errors from 1,000 simulations of quantitative traits. For each simulation replicate, we generated two phenotypes for each of 2,062 individuals in our sample by drawing effect sizes approximately 600,000 SNPs on chromosome 2 from a bivariate normal distribution. We then computed summary statistics for both phenotypes and estimated heritability and genetic correlation with LD Score regression. The summary statistics were generated from completely overlapping samples. Results are shown in Table 1. These simulations confirm that LD Score regression yields accurate estimates of the true genetic correlation and that the standard errors match the standard deviation across simulations. Thus, LD Score regression is not biased by sample overlap, in contrast to estimation of genetic correlation via polygenic risk scores, which is biased in the presence of sample overlap [20]. We also evaluated simulations with one quantitative trait and one case/control study and show that LD Score regression can be applied to binary traits and is not biased by oversampling of cases (Table S1).

Estimates of heritability and genetic covariance can be biased if the underlying model of genetic architecture is misspecified, e.g., if variance explained is correlated with LD Score or MAF [21, 23]. Because genetic correlation is estimated as a ratio, it is more robust: biases that affect the
| Parameter | Truth | Estimate | SD  | SE  |
|-----------|-------|----------|-----|-----|
| $h^2$     | 0.58  | 0.58     | 0.072 | 0.075 |
| $\rho_g$  | 0.29  | 0.29     | 0.057 | 0.058 |
| $r_g$     | 0.50  | 0.49     | 0.079 | 0.073 |

Table 1: Simulations with complete sample overlap. Truth shows the true parameter values. Estimate shows the average LD Score regression estimate across 1000 simulations. SD shows the standard deviation of the estimates across 1000 simulations, and SE shows the mean LD Score regression SE across 1000 simulations. Further details of the simulation setup are given in the Methods.

Replication of Psychiatric Cross-Disorder Results

As technical validation, we replicated the estimates of genetic correlations among psychiatric disorders obtained with individual genotypes and REML in [16], by applying LD Score regression to summary statistics from the same data [24]. These summary statistics were generated from non-overlapping samples, so we applied LD Score regression using both unconstrained and constrained intercepts (Methods). Results from these analyses are shown in Figure 1. As expected, the results from LD Score regression were similar to the results from REML. LD Score regression with constrained intercept gave standard errors that were only slightly larger than those from REML, while the standard errors from LD Score regression with intercept were substantially larger, especially for traits with small sample sizes (e.g., ADHD, ASD).

Application to Summary Statistics From 25 Phenotypes

We used cross-trait LD Score regression to estimate genetic correlations among 25 phenotypes (URLs, Methods). Genetic correlation estimates for all 300 pairwise combinations of the 25 traits are shown in Figure 2. For clarity of presentation, the 25 phenotypes were restricted to contain only one phenotype from each cluster of closely related phenotypes (Methods). Genetic correlations among the educational, anthropometric, smoking, and insulin-related phenotypes that were excluded from Figure 2 are shown in Table S4 and Figures S1, S2 and S3, respectively. References and sample sizes are shown in Table S3.

For the majority of pairs of traits in Figure 2, no GWAS-based genetic correlation estimate has been reported; however, many associations have been described informally based on the observation of overlap among genome-wide significant loci. Examples of genetic correlations that are consistent with overlap among top loci include the correlations between plasma lipids and cardiovascular disease [10]; age at onset of menarche and obesity [25]; type 2 diabetes, obesity, fasting glucose, plasma lipids and cardiovascular disease [26]; birth weight, adult height and type 2 diabetes [27,28]; birth length, adult height and infant head circumference [29, 30]; and childhood obesity and adult obesity [29]. For many of these pairs of traits, we can reject the null hypothesis of zero genetic correlation with overwhelming statistical significance (e.g., $p < 10^{-20}$ for age at onset of menarche and obesity).
Figure 1: Replication of Psychiatric Cross-Disorder Results. This plot compares LD Score regression estimates of genetic correlation using the summary statistics from [24] to estimates obtained from REML with the same data [16]. The horizontal axis indicates pairs of phenotypes, and the vertical axis indicates genetic correlation. Error bars are standard errors. Green is REML; orange is LD Score with intercept and white is LD Score with constrained intercept. The estimates of genetic correlation among psychiatric phenotypes in figure 2 use larger sample sizes; this analysis is intended as a technical validation. Abbreviations: ADHD = attention deficit disorder; ASD = autism spectrum disorder; BPD = bipolar disorder; MDD = major depressive disorder; SCZ = schizophrenia.

The first section of table 2 lists genetic correlation results that are consistent with epidemiological associations, but, as far as we are aware, have not previously been reported using genetic data.
Figure 2: Genetic Correlations among 25 GWAS. Blue represents positive genetic correlations; red represents negative. Larger squares correspond to more significant p-values. Genetic correlations that are different from zero at 1% FDR are shown as full-sized squares. Genetic correlations that are significantly different from zero after Bonferroni correction for the 300 tests in this figure have an asterisk. We show results that do not pass multiple testing correction as smaller squares in order to avoid whiting out positive controls where the estimate points in the expected direction, but does not achieve statistical significance due to small sample size. This multiple testing correction is conservative, since the tests are not independent.

The estimates of the genetic correlation between age at onset of menarche and adult height [31], cardiovascular disease [32] and type 2 diabetes [32,33] are consistent with the epidemiological associations. The estimate of a negative genetic correlation between anorexia nervosa and obesity (and
Table 2: Genetic correlation estimates, standard errors and p-values for selected pairs of traits. Results are grouped into genetic correlations that are new genetic results, but are consistent with established epidemiological associations (“Epidemiological”), genetic correlations that are new both to genetics and epidemiology (“New/Nonzero”) and interesting null results (“New/Low”). The p-values are uncorrected p-values. Results that pass multiple testing correction for the 300 tests in Figure 2 at 1% FDR have a single asterisk; results that pass Bonferroni correction have two asterisks. We present some genetic correlations that agree with epidemiological associations but that do not pass multiple testing correction in these data.

a similar genetic correlation with BMI) suggests that the same genetic factors influence normal variation in BMI as well as dysregulated BMI in psychiatric illness. This result is consistent with the observation that BMI GWAS findings implicate neuronal, rather than metabolic, cell-types and epigenetic marks [34,35]. The negative genetic correlation between adult height and coronary artery disease agrees with a replicated epidemiological association [36–38]. We observe several significant associations with the educational attainment phenotypes from Rietveld et al. [39]: we estimate a statistically significant negative genetic correlation between college and Alzheimer’s disease, which agrees with epidemiological results [40, 41]. The positive genetic correlation between college and bipolar disorder is consistent with previous epidemiological reports [42,43]. The estimate of a negative genetic correlation between smoking and college is consistent with the observed differences in smoking rates as a function of educational attainment [44].

The second section of table 2 lists three results that are, to the best of our knowledge, new both
to genetics and epidemiology. One, we find a positive genetic correlation between anorexia nervosa
and schizophrenia. Comorbidity between eating and psychotic disorders has not been thoroughly
investigated in the psychiatric literature [45, 46], and this result raises the possibility of similarity
between these classes of disease. Two, we estimate a negative genetic correlation between ulcerative
colitis (UC) and childhood obesity. The relationship between premorbid BMI and ulcerative colitis is
not well-understood; exploring this relationship may be a fruitful direction for further investigation.
Three, we estimate a positive genetic correlation between autism spectrum disorder (ASD) and
educational attainment, which itself has very high genetic correlation with IQ [39,47,48]. The ASD
summary statistics were generated using a case-pseudocontrol study design, so this result cannot be
explained by the tendency for the parents of children who receive a diagnosis of ASD to be better
educated than the general population [49]. The distribution of IQ among individuals with ASD has
lower mean than the general population, but with heavy tails [50] (i.e., an excess of individuals
with low and high IQ). There is evidence that the genetic architectures of high IQ and low IQ ASD
are dissimilar [51].

The third section of table 2 lists interesting examples where the genetic correlation is close to
zero with small standard error. The low genetic correlation between schizophrenia and rheumatoid
arthritis is interesting because schizophrenia has been observed to be protective for rheumatoid
arthritis [52]. The low genetic correlation between schizophrenia and smoking is notable because of
the high prevalence of smoking among individuals with schizophrenia [53]. The low genetic corre-
lation between schizophrenia and plasma lipid levels contrasts with a previous report of pleiotropy
between schizophrenia and triglycerides [54]. Pleiotropy (unsigned) is different from genetic correla-
tion (signed; see Methods); however, this observation from Andreassen, et al. [54] could be explained
by the sensitivity of the method used to the properties of a few regions with strong LD, rather than
trait biology (Figure S5). We estimate near-zero genetic correlation between Alzheimer’s disease
and schizophrenia. The genetic correlations between Alzheimers disease and the other psychiatric
traits (anorexia nervosa, bipolar, major depression, ASD) are also close to zero, but with larger
standard errors, due to smaller sample sizes. This suggests that the genetic basis of Alzheimer’s
disease is distinct from psychiatric conditions. Last, we estimate near zero genetic correlation be-
tween rheumatoid arthritis (RA) and both Crohn’s disease (CD) and UC. Although these diseases
share many associated loci [55,56], there appears to be no directional trend: some RA risk alleles
are also risk alleles for UC and CD, but many RA risk alleles are protective for UC and CD [55],
yielding near-zero genetic correlation. This example highlights the distinction between pleiotropy
and genetic correlation (Methods).

Finally, the estimates of genetic correlations among metabolic traits are consistent with the esti-
mates obtained using REML in Vattikuti et al. [17] (Supplementary Table S4), and are directionally
consistent with the recent Mendelian randomization results from Wuertz et al. [57]. The estimate
of 0.57 (0.074) for the genetic correlation between CD and UC is consistent with the estimate of
0.62 (0.042) from Chen et al. [18].

Discussion

We have described a new method for estimating genetic correlation from GWAS summary statistics,
which we applied to a dataset of GWAS summary statistics consisting of 25 traits and more than
1.5 million unique phenotype measurements. We reported several new findings that would have
been difficult or impossible to obtain with existing methods, including a positive genetic correla-
tion between anorexia nervosa and schizophrenia. Our method replicated many previously-reported GWAS-based genetic correlations, and confirmed observations of overlap among genome-wide significant SNPs, MR results and epidemiological associations.

This method is an advance for several reasons: it does not require individual genotypes, genome-wide significant SNPs or LD-pruning (which loses information if causal SNPs are in LD). Our method is not biased by sample overlap and is computationally fast. Furthermore, our approach does not require measuring multiple traits on the same individuals, so it scales easily to studies of thousands of pairs of traits. These advantages allow us to estimate genetic correlation for many more pairs of phenotypes than was possible with existing methods.

The challenges in interpreting genetic correlation are similar to the challenges in MR. We highlight two difficulties. First, genetic correlation is immune to environmental confounding, but is subject to genetic confounding, analogous to confounding by pleiotropy in MR. For example, the genetic correlation between HDL and CAD in Figure 2 could result from a causal effect $\text{HDL} \rightarrow \text{CAD}$, but could also be mediated by triglycerides (TG) \cite{10,58}, represented graphically \cite{59} as $\text{HDL} \leftarrow G \rightarrow \text{TG} \rightarrow \text{CAD}$, where $G$ is the set of genetic variants with effects on both HDL and TG. Extending genetic correlation to multiple genetically correlated phenotypes is an important direction for future work. Second, although genetic correlation estimates are not biased by oversampling of cases, they are affected by other forms of selection bias, such as misclassification \cite{16}.

We note several limitations of LD Score regression as an estimator of genetic correlation. First, LD Score regression requires larger sample sizes than methods that use individual genotypes in order to achieve equivalent standard error. Second, LD Score regression is not currently applicable to samples from recently-admixed populations. Third, methods built from polygenic models, such as LD Score regression and REML, are most effective when applied to traits with polygenic genetic architectures. For traits where significant SNPs account for a sizable proportion of heritability, analyzing only these SNPs can be more powerful. Developing methods that make optimal use of both large-effect SNPs and diffuse polygenic signal is a direction for future research.

Despite these limitations, we believe that the LD Score regression estimator of genetic correlation will be a useful addition to the epidemiological toolbox, since it allows for rapid screening for correlations among a diverse set of traits, without the need for measuring multiple traits on the same individuals or genome-wide significant SNPs.
Methods

Definition of Genetic Covariance and Correlation

All definitions refer to narrow-sense heritabilities and genetic covariances. Let $S$ denote a set of $M$ SNPs, let $X$ denote a vector of additively (0-1-2) coded genotypes for the SNPs in $S$, and let $y_1$ and $y_2$ denote phenotypes. Define $\beta := \arg\max_{\alpha \in \mathbb{R}^M} \text{Cor} [y_1, X\alpha]$, where the maximization is performed in the population (i.e., in the infinite data limit). Let $\gamma$ denote the corresponding vector for $y_2$. This is a projection, so $\beta$ is unique modulo SNPs in perfect LD. Define $h_S^2$, the heritability explained by SNPs in $S$, as $h_S^2(y_1) := \sum_j \beta_j^2$ and $\rho_S(y_1, y_2)$, the genetic covariance among SNPs in $S$, as $\rho_S(y_1, y_2) := \sum_{j \in S} \beta_j \gamma_j$. The genetic correlation among SNPs in $S$ is $r_S(y_1, y_2) := \rho_S(y_1, y_2)/\sqrt{h_S^2(y_1)h_S^2(y_2)}$, which lies in $[-1,1]$. Following [13], we use subscript $g$ (as in $h_g^2$, $\rho_g$, $r_g$) when the set of SNPs is genotyped and imputed SNPs in GWAS.

SNP genetic correlation ($r_g$) is different from family study genetic correlation. In a family study, the relationship matrix captures information about all genetic variation, not just common SNPs. As a result, family studies estimate the total genetic correlation ($S$ equals all variants). Unlike the relationship between SNP-heritability [13] and total heritability, for which $h_g^2 \leq h^2$, no similar relationship holds between SNP genetic correlation and total genetic correlation. If $\beta$ and $\gamma$ are more strongly correlated among common variants than rare variants, then the total genetic correlation will be less than the SNP genetic correlation.

Genetic correlation is (asymptotically) proportional to Mendelian randomization estimates. If we use a genetic instrument $g_i := \sum_{j \in S} X_{ij} \beta_j$ to estimate the effect $b_{12}$ of $y_1$ on $y_2$, the 2SLS estimate is $\hat{b}_{2SLS} := g^T y_2 / g^T y_1$ [8]. The expectations of the numerator and denominator are $E[g^T y_2] = \rho_S(y_1, y_2)$ and $E[g^T y_1] = h_S^2(y_1)$. Thus, $\text{plim}_{N \to \infty} \hat{b}_{2SLS} = r_S(y_2, y_1) \sqrt{h_S^2(y_1)/h_S^2(y_2)}$. If we use the same set $S$ of SNPs to estimate $b_{12}$ and $b_{21}$ (e.g., if $S$ is the set of all common SNPs, as in the genetic correlation analyses in this paper), then this procedure is symmetric in $y_1$ and $y_2$.

Genetic correlation is different from pleiotropy. Two traits have a pleiotropic relationship if many variants affect both. Genetic correlation is a stronger condition than pleiotropy: to exhibit genetic correlation, the directions of effect must also be consistently aligned.

Cross-Trait LD Score Regression

We estimate genetic covariance by regressing $z_{1j} z_{2j}$ against $\ell_j \sqrt{N_{1j} N_{2j}}$, (where $N_{ij}$ is the sample size for SNP $j$ in study $i$) then multiplying the resulting slope by $M$, the number of SNPs in the reference panel with MAF between 5% and 50% (technically, this is an estimate of $\rho_{5\%-50\%}$, see the Supplementary Note).

If we know the amount of sample overlap ahead of time, we can reduce the standard error by constraining the intercept with the --constrain-intercept flag in ldsc. This works even if there is nonzero sample overlap, in which case the intercept should be constrained to $N_s \rho / \sqrt{N_1 N_2}$.

Regression Weights

For heritability estimation, we use the regression weights from [21]. If effect sizes for both phenotypes are drawn from a bivariate normal distribution, then the optimal regression weights for genetic
covariance estimation are
\[
\text{Var}[z_j z_j | \ell_j] = \left( \frac{N_1 h_1^2 \ell_j}{M} + 1 \right) \left( \frac{N_2 h_2^2 \ell_j}{M} + 1 \right) + \left( \frac{\sqrt{N_1 N_2} \rho_g \ell_j + \rho N_s}{\sqrt{N_1 N_2}} \right)^2
\]  
(2)
(Supplementary Note). This quantity depends on several parameters \((h_1^2, h_2^2, \rho_g, \rho, N_s)\) which are not known a priori, so it is necessary to estimate them from the data. We compute the weights in two steps:

1. The first regression is weighted using heritabilities from the single-trait LD Score regressions, \(\rho N_s = 0\), and \(\rho_g\) estimated as \(\hat{\rho}_g := (\bar{\ell} \sqrt{N_1 N_2})^{-1} \sum_j z_{1j} z_{2j}\).

2. The second regression is weighted using the estimates of \(\rho N_s\) and \(\rho_g\) from step 1. The genetic covariance estimate that we report is the estimate from the second regression.

Linear regression with weights estimated from the data is called feasible generalized least squares (FGLS). FGLS has the same limiting distribution as WLS with optimal weights, so WLS \(p\)-values are valid for FGLS [8]. We multiply the heteroskedasticity weights by \(1/\ell_j\) (where \(\ell_j\) is LD Score with sum over regression SNPs) in order to downweight SNPs that are overcounted. This is a heuristic: the optimal approach is to rotate the data so that it is de-correlated, but this rotation matrix is difficult to compute.

Assessment of Statistical Significance via Block Jackknife

Summary statistics for SNPs in LD are correlated, so the OLS standard error will be biased downwards. We estimate a heteroskedasticity-and-correlation-robust standard error with a block jackknife over blocks of adjacent SNPs. This is the same procedure used in [21], and gives accurate standard errors in simulations (Table 1). We obtain a standard error for the genetic correlation by using a ratio block jackknife over SNPs. The default setting in ldsc is 200 blocks per genome, which can be adjusted with the \(--\text{num-blocks}\) flag.

Computational Complexity

Let \(N\) denote sample size and \(M\) the number of SNPs. The computational complexity of the steps involved in LD Score regression are as follows:

1. Computing summary statistics takes \(\Theta(MN)\) time.
2. Computing LD Scores takes \(\Theta(MN)\) time, though the \(N\) for computing LD Scores need not be large. We use the \(N = 378\) Europeans from 1000 Genomes.
3. LD Score regression takes \(\Theta(M)\) time and space.

For a user who has already computed summary statistics and downloads LD Scores from our website (URLs), the computational cost of LD Score regression is \(\Theta(M)\) time and space. For comparison, REML takes time \(\Theta(MN^2)\) for computing the GRM and \(\Theta(N^3)\) time for maximizing the likelihood.

Practically, estimating LD Scores takes roughly an hour parallelized over chromosomes, and LD Score regression takes about 15 seconds per pair of phenotypes on a 2014 MacBook Air with 1.7 GhZ Intel Core i7 processor.
**Simulations**

We simulated quantitative traits under an infinitesimal model in 2062 controls from a Swedish study. To simulate the standard scenario where many causal SNPs are not genotyped, we simulated phenotypes by drawing casual SNPs from 622,146 best-guess imputed 1000 Genomes SNPs on chromosome 2, then retained only the 90,980 HM3 SNPs with MAF above 5% for LD Score regression. We used in-sample LD Scores for LD Score regression.

**Summary Statistic Datasets**

We selected traits for inclusion in the main text via the following procedure:

1. Begin with all publicly available non-sex-stratified European-only summary statistics.
2. Remove studies that do not provide signed summary statistics.
3. Remove studies not imputed to at least HapMap 2.
4. Remove studies that include heritable covariates.
5. Remove all traits with heritability $z$-score below 4. Genetic correlation estimates for traits with heritability $z$-score below 4 are generally too noisy to interpret.
6. Prune clusters of correlated phenotypes (e.g., obesity classes 1-3) by picking the trait from each cluster with the highest heritability heritability $z$-score.

We then applied the following filters (implemented in the script `sumstats_to_chisq.py` included with `ldsc`):

1. For studies that provide a measure of imputation quality, filter to INFO above 0.9.
2. For studies that provide sample MAF, filter to sample MAF above 1%.
3. In order to restrict to well-imputed SNPs in studies that do not provide a measure of imputation quality, filter to HapMap3 [60] SNPs with 1000 Genomes EUR MAF above 5%, which tend to be well-imputed in most studies. This step should be skipped if INFO scores are available for all studies.
4. If sample size varies from SNP to SNP, remove SNPs with effective sample size less than 0.67 times the 90th percentile of sample size.
5. Remove indels and structural variants.
6. Remove strand-ambiguous SNPs.
7. Remove SNPs whose alleles do not match the alleles in 1000 Genomes.
8. Because the presence of outliers can increase the regression standard error, we also removed SNPs with extremely large effect sizes ($\chi^2 > 80$, as in [21]).

Genomic control (GC) correction at any stage biases the heritability and genetic covariance estimates downwards (see the Supplementary Note of [21]). The biases in the numerator and denominator of genetic correlation cancel exactly, so genetic correlation is not biased by GC correction. A majority of the studies analyzed in this paper used GC correction, so we do not report genetic covariance and heritability.

Data on Alzheimer’s disease were obtained from the following source:
International Genomics of Alzheimer’s Project (IGAP) is a large two-stage study based upon genome-wide association studies (GWAS) on individuals of European ancestry. In stage 1, IGAP used genotyped and imputed data on 7,055,881 single nucleotide polymorphisms (SNPs) to meta-analyze four previously-published GWAS datasets consisting of 17,008 Alzheimer’s disease cases and 37,154 controls (The European Alzheimer’s Disease Initiative, EADI; the Alzheimer Disease Genetics Consortium, ADGC; The Cohorts for Heart and Aging Research in Genomic Epidemiology consortium, CHARGE; The Genetic and Environmental Risk in AD consortium, GERAD). In stage 2, 11,632 SNPs were genotyped and tested for association in an independent set of 8,572 Alzheimer’s disease cases and 11,312 controls. Finally, a meta-analysis was performed combining results from stages 1 and 2.

We only used stage 1 data for LD Score regression.
URLs

1. ldsc software:
   github.com/bulik/ldsc
2. This paper:
   github.com/bulik/gencor.tex
3. PGC (psychiatric) summary statistics:
   www.med.unc.edu/pgc/downloads
4. GIANT (anthropometric) summary statistics:
   www.broadinstitute.org/collaboration/giant/index.php/GIANT_consortium_data_files
5. EGG (Early Growth Genetics) summary statistics:
   www.egg-consortium.org/
6. MAGIC (insulin, glucose) summary statistics:
   www.magicinvestigators.org/downloads/
7. CARDIoGRAM (coronary artery disease) summary statistics:
   www.cardiogramplusc4d.org
8. DIAGRAM (T2D) summary statistics:
   www.diagram-consortium.org
9. Rheumatoid arthritis summary statistics:
   www.broadinstitute.org/ftp/pub/rheumatoid_arthritis/Stahl_et_al_2010NG/
10. IGAP (Alzheimers) summary statistics:
    www.pasteur-lille.fr/en/recherche/u744/igap/igap_download.php
11. IIBDGC (inflammatory bowel disease) summary statistics:
    www.ibdgenetics.org/downloads.html
    We used a newer version of these data with 1000 Genomes imputation.
12. Plasma lipid summary statistics:
    www.broadinstitute.org/mpg/pubs/lipids2010/
13. SSGAC (educational attainment) summary statistics:
    www.ssgac.org/
14. Beans:
    www.barismo.com
    www.bluebottlecoffee.com
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Author Contributions

MJD provided reagents. BMN and ALP provided reagents. CL, ER, VA, JP and FD aided in the interpretation of results. JP and FD provided data on age at onset of menarche. The caffeine molecule is responsible for all that is good about this manuscript. BBS and HKF are responsible for the rest. All authors revised and approved the final manuscript.

Competing Financial Interests

Unfortunately, we have no financial conflicts of interest to declare.
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Supplementary Note

1.1 Quantitative Traits

Suppose we sample two cohorts with sample sizes $N_1$ and $N_2$. We measure phenotype 1 in cohort 1 and phenotype 2 in cohort 2. We model phenotype vectors for each cohort as $y_1 = Y\beta + \delta$, and $y_2 = Z\gamma + \epsilon$, where $Y$ and $Z$ are matrices of genotypes with columns standardized to mean zero and variance one, with dimensions $N_1 \times M$ and $N_2 \times M$, respectively; $\beta$ and $\gamma$ are vectors of per-standardized genotype effect sizes, and $\delta$ and $\epsilon$ are vectors of residuals, representing environmental effects and non-additive genetic effects. In this model, $Y$ and $Z$ are unobserved matrices of all SNPs, including SNPs that are not genotyped.

We treat all of $Y,Z,\beta,\gamma,\delta$ and $\epsilon$ as random. We model all of these as independent, except for $\beta,\gamma$, $\delta,\epsilon$. Suppose that $(\beta,\gamma)$ has mean zero and covariance matrix

$$\text{Var}[(\beta,\gamma)] = \frac{1}{M} \left( \begin{array}{cc} h_1^2 I & \rho_g I \\ \rho_g I & h_2^2 I \end{array} \right),$$

and $(\delta,\epsilon)$ has mean zero and covariance matrix

$$\text{Var}[(\delta,\epsilon)] = \left( \begin{array}{cc} (1-h_1^2) I & \rho_e I \\ \rho_e I & (1-h_2^2) I \end{array} \right).$$

Let $\rho := \rho_g + \rho_e$. Vectors of genotypes for each individual are drawn i.i.d. from a distribution with covariance matrix $r$ (i.e., $r$ is an LD matrix with $r_{jk} = \mathbb{E}[Y_{ij}Y_{ik}]$). There are $N_s$ individuals who are included in both studies.

Lemma 1. Under this model, the expected genetic covariance (as defined in methods) between phenotypes is $\rho_g$, justifying our use of the notation $\rho_g$.

Proof. Let $X$ denote an $1 \times M$ vector of standardized genotypes for an arbitrary individual. Under the model, the additive genetic component of phenotype 1 for this individual is $\sum_j X_j \beta_j$, and the additive genetic component of phenotype 1 for this individual is $\sum_j X_j \gamma_j$. Thus, the genetic

1We ignore the distinction between normalizing and centering in the population and in the sample, since this introduces only $O(1/N)$ error.

2The assumption that all $\beta$ is drawn with equal variance for all SNPs hides an implicit assumption that rare SNPs have larger per-allele effect sizes than common SNPs. As discussed in the simulations section of the main text and in our earlier work [21], LD Score regression is robust to moderate violations of this assumption, though it may break down in extreme cases, e.g., if all causal variants are rare. In situations where a different model for Var[$\beta$] is more appropriate, all proofs in this note go through with LD Score replaced by weighted LD Scores, $t_j = \sum_k \text{Var}[\beta_j] r_{jk}^2$. 


covariance between phenotype 1 and phenotype 2 is

\[
\text{Cov} \left[ \sum_j X_j \beta_j, \sum_j X_j \gamma_j \right] = \mathbb{E} \left[ \left( \sum_j X_j \beta_j \right) \left( \sum_j X_j \gamma_j \right) \right]
\]

\[
= \sum_j \sum_k \mathbb{E} [X_j X_k \beta_j \gamma_k]
\]

\[
= \sum_j \mathbb{E} [X_j^2 \beta_j \gamma_j]
\]

\[
= \sum_j \mathbb{E} [X_j^2] \mathbb{E} [\beta_j \gamma_j]
\]

\[
= \rho_g.
\]

We compute linear regression z-scores \( z_{1j} := Y_j^T y_1 / \sqrt{N_1} \) and \( z_{2j} := Y_j^T y_2 / \sqrt{N_2} \) for genotyped SNPs \( j \) (where \( Y_j \) and \( Z_j \) denote the \( j^{th} \) columns of \( Y \) and \( Z \)).

**Definition 1.** The LD Score of a variant \( j \) is \( \ell_j := \sum_k r_{jk}^2 \), where the sum is taken over all other variants \( k \).

**Proposition 1.** Let \( j \) denote a genotyped SNP. Under the model described above,

\[
\mathbb{E} [z_{1j} z_{2j}] = \frac{\sqrt{N_1 N_2} \rho_g}{M} \ell_j + \frac{N_s \rho}{\sqrt{N_1 N_2}}.
\]

**Proof.** By the law of total expectation,

\[
\mathbb{E} [z_{1j} z_{2j}] = \mathbb{E} [\mathbb{E} [z_{1j} z_{2j} | Y, Z]]
\]

First we compute the inner expectation from Equation 4, with \( Z \) and \( Y \) fixed.

\[
\mathbb{E} [z_{1j} z_{2j} | Y, Z] = \frac{1}{\sqrt{N_1 N_2}} \mathbb{E} [Y_j^T y_1 y_2^T Z_j]
\]

\[
= \frac{1}{\sqrt{N_1 N_2}} Y_j^T \mathbb{E} [(Y \beta + \delta)(Z \gamma + \epsilon)^T] Z_j
\]

\[
= \frac{1}{\sqrt{N_1 N_2}} Y_j^T \left( Y \mathbb{E} [\beta^T \gamma] Z + \mathbb{E} [\delta^T Z \gamma] + \mathbb{E} [\beta^T Y \epsilon] + \mathbb{E} [\delta^T \epsilon] \right) Z_j
\]

\[
= \frac{1}{\sqrt{N_1 N_2}} Y_j^T \left( \rho_g Y_j^T Y Z_j^T Z + \rho_e Y_j^T Z_j \right).
\]

Next, we remove the conditioning on \( Y \) and \( Z \).

\[
\frac{1}{\sqrt{N_1 N_2}} \mathbb{E} [Y_j^T Z_j] = \frac{N_s}{\sqrt{N_1 N_2}},
\]

\( \square \)
and
\[
\frac{1}{\sqrt{N_1 N_2}} \mathbb{E}[Y_j^T Y Z_j^T Z] = \ell_j + \frac{MN_s}{\sqrt{N_1 N_2}}.
\] (7)

Substituting equations 6 and 7 into Equation 5,
\[
\mathbb{E}[z_{1j} z_{2j}] = \frac{\sqrt{N_1 N_2 \rho_g}}{M} \ell_j + \frac{Ns (\rho_g + \rho_e)}{\sqrt{N_1 N_2}}
= \frac{\sqrt{N_1 N_2 \rho_g}}{M} \ell_j + \frac{Ns \rho}{\sqrt{N_1 N_2}}.
\] (8)

If study 1 and study 2 are the same study, then \(N_1 = N_2 = N_s\), \(\rho_g = \rho_g\) and \(\rho = 1\), so Equation 8 reduces to the LD Score regression equation for a single trait from [21].

1.2 Regression Weights

We can improve the efficiency of LD Score regression by weighting by the reciprocal of the conditional variance function (CVF), \(\text{Var}[z_{1j} z_{2j} | \ell_j]\). The CVF is not uniquely determined by the assumptions about the first and second moments of \(\beta\) and \(\gamma\) used to derive Proposition 1. Therefore we derive the CVF for the case where \(z_{1j}\) and \(z_{2j}\) are jointly distributed as bivariate normal\(^3\). From a standard formula for double second moments of the bivariate normal, the CVF is
\[
\text{Var}[z_{1j} z_{2j} | \ell_j] = \text{Var}[z_{1j}] \text{Var}[z_{2j}] + \mathbb{E}[z_{1j} z_{2j}]^2 = \left( \frac{N_1 h_1^2 \ell_j}{M} + 1 \right) \left( \frac{N_2 h_2^2 \ell_j}{M} + 1 \right) + \left( \frac{\sqrt{N_1 N_2 \rho_g}}{M} \ell_j + \frac{\rho N_s}{\sqrt{N_1 N_2}} \right)^2
\] (9)

The terms on the left follow from the fact that \(\text{Var}[z_{ij}] = \chi_{ij}^2\) and \(\mathbb{E}[\chi^2] = Nh_2^2 \ell_j / M + 1\). The term on the right follows from Proposition 1. Note that if \(z_1 = z_2\), this reduces to the expression for the CVF of \(\chi^2\) statistics from [21].

In cases where the normality assumption does not hold, LD Score regression will remain unbiased, but may be inefficient, because the regression weights will be suboptimal. We also apply a heuristic weighting scheme to avoid overcounting SNPs in high-LD regions, described in the methods.

1.3 Liability Threshold Model

In the liability threshold (probit) model [61], binary traits are determined by an unobserved continuous liability \(\psi\). The observed trait is \(y := 1[\psi > \tau]\), where \(\tau\) is the liability threshold. If \(\psi\) is normally distributed, then setting \(\tau := \Phi^{-1}(1 - K)\) (where \(\Phi\) is the standard normal cdf) yields a population prevalence of \(K\).

For phenotypes generated according to the liability threshold model, we can estimate not only the heritability and genetic covariance of the observed phenotype, but also the heritability and genetic covariance of the unobserved liability.

\(^3\)For instance, it is sufficient but not necessary to assume that \(\beta, \gamma, \delta\) and \(\epsilon\) are multivariate normal. More generally, the z-scores will be approximately normal if \(\beta\) and \(\gamma\) are reasonably polygenic. If the distribution of effect sizes is heavy-tailed, e.g., if there are few casual SNPs, then the CVF may be larger.
In the next lemma, we derive population case and control allele frequencies in terms of the heritability of liability when liability is generated following the model for quantitative traits from section 1.1. Since we are only modeling additive effects and are willing to assume Hardy-Weinberg equilibrium, we lose no generality and simplify notation considerably by stating the proofs in terms of haploid genotypes.

We state this lemma in terms of marginal per-allele effect sizes, instead of the per-standardized-genotype effect sizes considered in section 1.1. Here marginal means that these are the effect sizes obtained by univariate regression of phenotype against genotype in the infinite data limit. Haploid standardized genotypes are defined \( X_{ij} := (G_{ij} - p_j) / \sqrt{p_j(1 - p_j)} \), where \( G_{ij} \) is the 0-1 coded genotype. If \( \beta_j \) is the marginal per-standardized-genotype effect and \( \zeta_j \) is the marginal per-allele effect, we have \( X_j \beta_j = G_j \zeta_j \). Thus, setting \( G_{ij} = 1 \) yields \( \zeta_j = \beta_j \sqrt{(1 - p_j)/p_j} \).

**Lemma 2.** Suppose unobserved liabilities \( \psi, \varphi \) for traits \( y_1, y_2 \) with thresholds \( \tau_1, \tau_2 \) corresponding to prevalences \( K_1, K_2 \) are generated according to the mode for quantitative traits from section 1.1, i.e., \( \psi_i = \sum_j X_{ij} \beta_j + \delta \), \( \varphi_i = \sum_j X_{ij} \gamma_j + \epsilon \), with

\[
\text{Var}[(\beta, \gamma)] = \frac{1}{M} \begin{pmatrix} h_1^2 I & \rho_g I \\ \rho_g I & h_2^2 I \end{pmatrix},
\]

and

\[
\text{Var}[(\delta, \epsilon)] = \begin{pmatrix} (1 - h_1^2) I & \rho_e I \\ \rho_e I & (1 - h_2^2) I \end{pmatrix}.
\]

Let \( \zeta_j \) and \( \xi_j \) denote the marginal per-allele effect sizes of SNP \( j \) on \( \psi \) and \( \varphi \). Let

\[
p_{\text{cas,}k,j} := \mathbb{P}[G_{ij} = 1 | y_{ik} = 1]
\]

\[
p_{\text{con,}k,j} := \mathbb{P}[G_{ij} = 1 | y_{ik} = 0]
\]

denote the allele frequencies of SNP \( j \) in cases and controls for phenotype \( k \), where \( y_{ik} \) denotes the value of phenotype \( k \) for individual \( i \) and \( k = 1, 2 \). Then

\[
E[p_{\text{cas,}1,j} - p_{\text{con,}1,j}] = 0,
\]

\[
E[p_{\text{cas,}2,j} - p_{\text{con,}2,j}] = 0,
\]

\[
\text{Var}[p_{\text{cas,}1,j} - p_{\text{con,}1,j}] = \frac{p_j(1 - p_j)\phi(\tau_1)^2 h_1^2}{MK_1^2(1 - K_1)^2} \ell_j,
\]

\[
\text{Var}[p_{\text{cas,}2,j} - p_{\text{con,}2,j}] = \frac{p_j(1 - p_j)\phi(\tau_2)^2 h_2^2}{MK_2^2(1 - K_2)^2} \ell_j,
\]

\[
\text{Cov}[p_{\text{cas,}1,j} - p_{\text{con,}1,j}, p_{\text{cas,}2,j} - p_{\text{con,}2,j}] = \frac{p_j(1 - p_j)\phi(\tau_1)\phi(\tau_2)\rho_g}{MK_1(1 - K_1)K_2(1 - K_2)} \ell_j,
\]

where the expectation is taken over where \( \phi \) is the standard normal density. These results apply to population allele frequencies, not allele frequencies in a finite sample. We deal with ascertained finite samples in the next section.

**Proof.** This proof is accomplished in two steps. First, we compute allele frequencies conditional on the marginal effects on liability. To do this, we reverse the conditional probability using Bayes’ theorem, which reduces the problem to a series of [Taylor approximations to] Gaussian integrals.
Second, we remove the conditioning on the marginal effects on liability in order to express the allele frequencies in terms of \( h_1^2, h_2^2, \rho_g \) and \( \ell_j \). Since liability is just a quantitative trait, we need only apply the LD Score regression equation for quantitative traits.

By Bayes’ rule,

\[
P(G_{ij} = 1 | y_{i1} = 1, \zeta_j) = \frac{P[y_{i1} = 1 | G_{ij} = 1, \zeta_j] P(G_{ij} = 1)}{P[y_{i1} = 1]} = \frac{p_j}{K_1} P[y_{i1} = 1 | G_{ij} = 1, \zeta_j]
\]

\[
= \frac{p_j}{K_1} P[\psi_i > \tau_1 | G_{ij} = 1, \zeta_j].
\]

(10)

The distribution of \( \psi \) given \( G_{ij} \) and \( \zeta_j \) is \( \psi | (G_{ij} = 1, \zeta_j) \sim N(\zeta_j, 1 - \zeta_j^2) \approx N(\zeta_j, 1) \) (where the approximation that the variance equals one holds when the marginal heritability explained by \( j \) is small, which is the typical case in GWAS). Thus \( P[\psi_i > \tau_1 | G_{ij} = 1] \) is simply a Gaussian integral. We approximate this probability with a first-order Taylor expansion around \( \tau_1 \).

\[
P[\psi_i > \tau_1 | G_{ij} = 1, \zeta_j] = 1 - \Phi(\tau_1 - \zeta_j)
\]

\[
\approx K_1 + \phi(\tau_1) \zeta_j,
\]

(11)

Substituting Equation 11 into Equation 10,

\[
P[G_{ij} = 1 | y_{i1} = 1, \zeta_j] = \frac{p_j}{K_1} (K_1 + \phi(\tau_1) \zeta_j).
\]

(12)

A similar argument shows that

\[
P[G_{ij} = 1 | y_{i1} = 0, \zeta_j] = \frac{p_j}{1 - K_1} (1 - K_1 - \phi(\tau_1) \zeta_j).
\]

(13)

Subtracting Equation 13 from Equation 12,

\[
P[G_{ij} = 1 | y_{i1} = 1, \zeta_j] - P[G_{ij} = 1 | y_{i1} = 0, \zeta_j] = p_j \frac{\phi(\tau_1) \zeta_j}{K_1(1 - K_1)}.
\]

(14)

Similar results hold for trait 2, replacing \( \zeta \) with \( \xi \) and subscript 1 with subscript 2.

We have written the probabilities in question in terms of constants and marginal effects on liability. Since liability is simply a quantitative trait, the means, variances, and covariances of the marginal effects on liability are described by the LD Score regression equation for quantitative traits from Proposition 1. Precisely, \( E[\xi_j] = E[\zeta_j] = 0, \ Var[\xi_j] = (1 - p_j) h_1^2 \ell_j / p_j M, \ Var[\zeta_j] = (1 - p_j) h_2^2 \ell_j / p_j M \) and \( \text{Cov}[\zeta_j, \xi_j] = (1 - p_j) \rho_g \ell_j / p_j M \). If we combine these results with Equation 14, we find that

\[
E[p_{\text{cas,1,j}} - p_{\text{con,1,j}}] = 0;
\]

(15)

\[
\text{Var}[p_{\text{cas,1,j}} - p_{\text{con,1,j}}] = \text{Var} \left[ \frac{p_j \phi(\tau_1) \zeta_j}{K_1(1 - K_1)} \right] = \frac{p_j(1 - p_j) \phi(\tau_1)^2 h_1^2 \ell_j}{MK_1^2(1 - K_1)^2}.
\]

(16)
(similarly for trait two), and
\[
\text{Cov}[p_{\text{cas},1j} - p_{\text{con},1j}, p_{\text{cas},2j} - p_{\text{con},2j}] = \text{Cov} \left[ \frac{p_j \phi(\tau_1) \zeta_j}{K_1(1 - K_1)}, \frac{p_j \phi(\tau_2) \xi_j}{K_2(1 - K_2)} \right] = \frac{p_j (1 - p_j) \phi(\tau_1) \phi(\tau_2) \rho_g}{MK_1(1 - K_1)K_2(1 - K_2)} \ell_j. \tag{17}
\]

### 1.4 Ascertained Studies of Liability Threshold Traits

In the next proposition, we derive an LD Score regression equation for ascertained case/control studies.

Let \( P_i \) denote the sample prevalence of \( y_i \) in study \( i \) for \( i = 1, 2 \). We compute z-scores
\[
z_j := \frac{\sqrt{NP(1 - P)}(\hat{p}_{\text{cas}} - \hat{p}_{\text{con}})}{\sqrt{p_j(1 - p_j)}},
\]
where \( \hat{p}_j \) denotes allele frequency in the entire sample\(^4\), \( \hat{p}_{\text{cas}} \) denotes sample case allele frequency and \( \hat{p}_{\text{con}} \) denotes sample control allele frequency.

We emphasize one subtlety before stating the main proposition. The results in this section allow for study \( k \) to select samples based on phenotype \( l \) only if \( k = l \). If study 1 ascertains on phenotype 2 – for example, if all cases in study 1 have \( y_{11} = y_{12} = 1 \) — then \( \hat{p}_{\text{cas},1j} \) will not be an unbiased estimate of \( p_{\text{cas},1j} \). Indeed, in this example, \( \mathbb{E}[\hat{p}_{\text{cas},1j}] = \mathbb{P}[G_{ij} = 1 \mid y_{11} = y_{12} = 1], \) which will not equal \( p_{\text{cas},1j} = \mathbb{P}[G_{ij} = 1 \mid y_{11} = 1] \) unless \( \rho = 1 \) or \( \rho = 0 \). This follows from the fact that the conditionals and marginals of a bivariate normal are equal iff \( \rho = 0 \) or \( \rho = 1 \). We do not derive formulae describing the bias, except to note that the most common scenario, the “healthy controls” model — cases are sampled independently but all controls are controls for both traits — is probably nothing to worry about, so long as cases for both traits are uncommon. In this scenario, \( \mathbb{P}[G_{ij} = 1 \mid y_{11} = 0] \approx \mathbb{P}[G_{ij} = 1 \mid y_{11} = y_{12} = 0] \). Conditioning on \( y_{12} = 0 \) hardly changes the distribution, because \( y_{12} = 0 \) most of the time, anyway.

**Proposition 2.** Under the liability threshold model from lemma 1.3,
\[
\mathbb{E}[z_j z_2] \approx \frac{\sqrt{N_1 N_2 \rho_{g,\text{obs}}}}{M} \ell_j + \sqrt{N_1 N_2 P_1(1 - P_1)P_2(1 - P_2)} \left( \sum_{a,b \in \{\text{cas,con}\}} \frac{N_{a,b} (-1)^{1+I[a=b]}}{N_{a,1}N_{b,2}} \right) \tag{18}
\]
where
\[
\rho_{g,\text{obs}} := \rho_g \left( \frac{\phi(\tau_1) \phi(\tau_2)}{K_1(1 - K_1)K_2(1 - K_2)} \right)
\]
denotes observed scale genetic covariance, \( N_{a,b} \) denotes the number of individuals with phenotype \( a \) in study 1 and \( b \) in study two for \( a, b \in \{\text{cas,con}\} \) (e.g., \( N_{\text{cas,con}} \) is the number of individuals who are a case in study 1 but a control in study 2), \( N_i \) denotes total sample size in study \( i \) and \( N_{a,i} \) for \( a \in \{\text{cas,con}\} \) and \( i = 1, 2 \) denotes the number of individuals with phenotype \( a \) in study \( i \).

\(^4\) Conditional on the marginal effect of \( j \), the expected value of \( \hat{p}_j \) is not equal to \( p_j \) unless \( P = K \) or the marginal effect of \( j \) is zero.
Observe that \( \rho_{g,obs}/\sqrt{\hat{h}^2_{1,obs} \hat{h}^2_{2,obs}} = \rho_g / \sqrt{h^2_1 h^2_2} = r_g \). Put another way, the natural definition for “observed scale genetic correlation” turns out to be the same as regular genetic correlation, because the scale transformation factors in the numerator and denominator cancel. This is convenient: we can compute genetic correlations for binary traits on a sensible scale without having to worry about sample and population prevalences.

**Proof.** The full form of \( z_{1j}z_{2j} \) is

\[
 z_{1j}z_{2j} = \frac{\sqrt{cN_1N_2}(\hat{p}_{cas,1j} - \hat{p}_{con,1j})(\hat{p}_{cas,2j} - \hat{p}_{con,2j})}{\sqrt{p_{1j}(1 - p_{1j})p_{2j}(1 - p_{2j})}},
\]

where \( c := P_1(1 - P_1)P_2(1 - P_2) \). Our strategy for obtaining the expectation is

\[
 \mathbb{E}[z_{1j}z_{2j}] \approx \sqrt{cN_1N_2} \frac{\mathbb{E}[(\hat{p}_{cas,1j} - \hat{p}_{con,1j})(\hat{p}_{cas,2j} - \hat{p}_{con,2j})]}{\mathbb{E}[\sqrt{p_{1j}(1 - p_{1j})p_{2j}(1 - p_{2j})}]}, \tag{19}
\]

\[
 \approx \sqrt{cN_1N_2} \frac{\mathbb{E}[(\hat{p}_{cas,1j} - \hat{p}_{con,1j})(\hat{p}_{cas,2j} - \hat{p}_{con,2j})]}{\mathbb{E}[(\hat{p}_{1j}(1 - p_{1j})p_{2j}(1 - p_{2j})]} \tag{20}
\]

\[
 = \sqrt{cN_1N_2} \frac{\mathbb{E}[(\hat{p}_{cas,1j} - \hat{p}_{con,1j})(\hat{p}_{cas,2j} - \hat{p}_{con,2j}) | \zeta_j, \xi_j]}{\mathbb{E}[(\hat{p}_{1j}(1 - p_{1j})p_{2j}(1 - p_{2j})] | \zeta_j, \xi_j]} \tag{21}
\]

where \( \zeta_j \) and \( \xi_j \) denote the marginal per-allele effects of \( j \). Approximation 19 hides \( \mathcal{O}(1/N) \) error from moving from the expectation of a ratio to a ratio of expectations. Approximation 20 hides \( \mathcal{O}(1/N) \) error from moving from the expectation of a square root to a square root of expectations, and dear reader we admire your perseverance in making it this far. Equality 21 follows from applying the law of total expectation to the numerator and denominator.

First, we compute the numerator. By linearity of expectation,

\[
 \mathbb{E}[(\hat{p}_{cas,1j} - \hat{p}_{con,1j})(\hat{p}_{cas,2j} - \hat{p}_{con,2j}) | \zeta_j, \xi_j] = \mathbb{E}[(\hat{p}_{cas,1j})(\hat{p}_{1j} \hat{p}_{cas,2j} | \zeta_j, \xi_j] + \mathbb{E}[\hat{p}_{con,1j}(\hat{p}_{con,2j} | \zeta_j, \xi_j] \tag{22}
\]

After conditioning on the marginal effects \( \zeta_j \) and \( \xi_j \), the only source of variance in the sample allele frequencies \( \hat{p}_{cas,1}, \hat{p}_{cas,2}, \hat{p}_{con,1}, \hat{p}_{con,2} \) is sampling error. Write \( p_{cas,1j}\hat{p}_{cas,2j} = (p_{cas,1j} + \eta)(p_{cas,2j} + \nu) \), where \( \eta \) and \( \nu \) denote sampling error. If study 1 and study 2 share samples, \( \nu \) and \( \eta \) will be correlated:

\[
 \mathbb{E}[\hat{p}_{cas,1j}\hat{p}_{cas,2j} | \zeta_j, \xi_j] = p_{cas,1j}p_{cas,2j} + \mathbb{E}[\eta\nu] \approx p_{cas,1j}p_{cas,2j} + \frac{N_{cas,1}N_{cas,2}}{N_{cas,1}N_{cas,2}} \tag{23}
\]

\[
 \approx p_{cas,1j}p_{cas,2j} \left(1 + \frac{N_{cas,1}N_{cas,2}}{N_{cas,1}N_{cas,2}} \right) \tag{24}
\]

where approximation 23 is the (bivariate) central limit theorem, and approximation 24 comes from ignoring the difference between \( \sqrt{p_{cas,1j}(1 - p_{cas,1j})p_{cas,2j}(1 - p_{cas,2j})} \) and \( p_j(1 - p_j) \). This step is justified in the derivation of the denominator. Similar relationships hold for the other terms in Equation 22.
If we combine equations 24 and 17, we obtain
\[
\mathbb{E}[(\hat{p}_{\text{cas},1j} - \hat{p}_{\text{con},1j})(\hat{p}_{\text{cas},2j} - \hat{p}_{\text{con},2j})] \approx p_j(1 - p_j) \left( \frac{\phi(\tau_1)\phi(\tau_2)p_\ell_j + \sum_{a,b \in \{\text{cas,con}\}} N_{a,b}(-1)^{1+[a=b]}}{c'M} \right),
\]
where \(c' := K_1(1 - K_1)K_2(1 - K_2)\).

Next, we derive the expectation of the denominator. Conditional on \(\zeta_j\) and \(\xi_j\), \(\hat{p}_{1j}(1 - \hat{p}_{1j})\) is \(P_1p_{\text{cas},1j} + (1 - P_1)p_{\text{con},1j}\) plus \(O(1/N)\) sampling variance. If studies 1 and 2 share samples, the \(O(1/N)\) sampling variance in \(\hat{p}_{1j}(1 - \hat{p}_{1j})\) and \(\hat{p}_{2j}(1 - \hat{p}_{2j})\) will be correlated, but this still only amounts to \(O(N_s/N_1N_2)\) error. If we remove the conditioning on \(\zeta_j\) and \(\xi_j\), then \(P_1p_{\text{cas},1j} + (1 - P_1)p_{\text{con},1j}\) is equal to \(p_j(1 - p_j)\) plus \(O(h_{\text{obs}}^2\rho_{\ell,j}/M)\) error from uncertainty in \(\zeta_j\). The covariance between uncertainty in \(\zeta_j\) and uncertainty in \(\xi_j\) is driven by \(\rho_{g,\text{obs}}\), so the expectation of the denominator is \(\mathbb{E}\left[\sqrt{p_{1j}(1 - \hat{p}_{1j})\hat{p}_{2j}(1 - \hat{p}_{2j})}\right] = p_j(1 - p_j)\left(1 + O(N_s/N_1N_2) + O(h_{\text{obs}}^2\rho_{\ell,j}/M)\right)\). We make the approximation\(^5\)
\[
\mathbb{E}\left[\sqrt{p_{1j}(1 - \hat{p}_{1j})\hat{p}_{2j}(1 - \hat{p}_{2j})}\right] \approx p_j(1 - p_j).
\]
We obtain the desired result by dividing \(\sqrt{cN_1N_2}\) times Equation 25 by Equation 26.

**Corollary 1.** If study 1 is an ascertained study of a binary trait, and study 2 is a non-ascertained quantitative study, then proposition 2 holds, except with genetic covariance on the half-observed scale
\[
\rho_{g,\text{obs}} := \rho_g \left(\frac{\phi(\tau_1)\sqrt{P_1(1 - P_1)}}{K_1(1 - K_1)}\right).
\]

**Corollary 2.** For a single binary trait,
\[
\mathbb{E}[\chi_j^2] = \frac{Nh_{\text{obs}}^2\rho_{\ell,j}}{M} + 1,
\]
where \(h_{\text{obs}}^2 = h^2\phi(\tau)^2P(1 - P)/K^2(1 - K)^2\).

**Proof.** This follows from proposition 2 if we set study 1 equal to study 2 and note that the observed scale genetic covariance between a trait and itself is observed scale heritability. To show that the intercept is one, observe that if study 1 and study 2 are the same, then
\[
\sqrt{cN_1N_2} \left( \sum_{a,b \in \{\text{cas,con}\}} \frac{N_{a,b}(-1)^{1+[a=b]}}{N_{a,1}N_{b,2}} \right) = NP(1 - P) \left(\frac{1}{N_{\text{cas}}} + \frac{1}{N_{\text{con}}}\right)
\]
\[
= NP(1 - P)(N_{\text{cas}} + N_{\text{con}})
\]
\[
= \frac{N_{\text{cas}}N_{\text{con}}}{N_{\text{cas}}N_{\text{con}}}
\]
But \(NP = N_{\text{cas}}\) and \(N(1 - P) = N_{\text{con}}\), so Equation 28 simplifies to 1.\(\square\)

\(^5\)For \(\ell_j = 100\) (roughly the median 1kG LD Score), \(M = 10^7\) and \(\rho_{g,\text{obs}} = 1\), we get \(\rho_{g,\text{obs}}\ell_j/M = 10^{-5}\). A worst-case value for 2\(s\)/2\(i\) might be 2\(s\) = 2\(i\) = 10\(^3\), in which case 2\(s\)/2\(i\) = 10\(^{-3}\). Thus, \(\rho_{g,\text{obs}}\ell_j/M\) and 2\(s\)/2\(i\) will generally be at least 3 orders of magnitude smaller than 1.
1.5 Flavors of Heritability and Genetic Correlation

The heritability parameter estimated by ldsc is subtly different than the heritability parameter \( h^2_g \) estimated by GCTA. If \( g \) denotes the set of all genotyped SNPs in some GWAS, define \( \beta_{GCTA} := \arg\max_{\alpha \in \mathbb{R}^{|g|}} \text{Cor} [y_1, X_g \alpha] \), where \( X_g \) is a random vector of standardized genotypes for SNPs in \( g \). Then the heritability parameter estimated by GCTA is defined

\[
h^2_g := \sum_{j \in g} \beta^2_{GCTA,j}.
\]

Let \( S \) denote the set of SNPs used to compute LD Scores (i.e., \( \ell_j = \sum_{k \in S} r^2_{jk} \)), and let \( \beta_S := \arg\max_{\alpha \in \mathbb{R}^{|S|}} \text{Cor} [y_1, X_S \alpha] \). Generally \( \beta_{S,j} \neq \beta_{GCTA,j} \) unless all SNPs in \( S \setminus g \) are not in LD with SNPs in \( g \). Define

\[
h^2_S := \sum_{j \in S} \beta^2_{S,j}.
\]

Let \( S' \) denote the set of SNPs in \( S \) with MAF above 5%. Define

\[
h^2_{5-50\%} := \sum_{j \in S'} \beta^2_{S,j}.
\]

The default setting in ldsc is to report \( h^2_{5-50\%} \), estimated as the slope from LD Score regression times \( M_{5-50\%} \), the number of SNPs with MAF above 5%.

The reason for this is the following: suppose that \( h^2 \) per SNP is not constant as a function of MAF. Then the slope of LD Score regression will represent some sort of weighted average of the values of \( h^2 \) per SNP, with more weight given to classes of SNPs that are well-represented among the regression SNPs. In a typical GWAS setting, the regression SNPs are mostly common SNPs, so multiplying the slope from LD Score regression by \( M \) (which includes rare SNPs) amounts to extrapolating that \( h^2 \) per SNP among common variants is the same as \( h^2 \) per SNP among rare variants. This extrapolation is particularly risky, because there are many more rare SNPs than common SNPs.

It is probably reasonable to treat \( h^2 \) per SNP as a constant function of MAF for SNPs with MAF above 5%, but we have very little information about \( h^2 \) per SNP for SNPs with MAF below 5%. Therefore we report \( h^2_{5-50\%} \) instead of \( h^2_S \) to avoid excessive extrapolation error. This lower bound can be pushed lower with larger sample sizes and better rare variant coverage, either from sequencing or imputation.

There are two main distinctions between \( h^2_{5-50\%} \) and \( h^2_g \). First, \( h^2_{5-50\%} \) does not include the effects of common SNPs that are not tagged by the set of genotyped SNPs \( g \). Second, the effects of causal 4% SNPs are not counted towards \( h^2_{5-50\%} \). In practice, neither of these distinctions makes a large difference, since most GWAS arrays focus on common variation and manage to assay or tag almost all common variants, which is why we do not emphasize this distinction in the main text.

The relationship between the genetic covariance parameter estimated by LD Score regression and the genetic covariance parameter estimated by GCTA is similar to the relationship between \( h^2_{5-50\%} \) and \( h^2_g \). Choice of \( M \) is not important for genetic correlation, because the factors of \( M \) in the numerator and denominator cancel.
Supplementary Tables

Simulations with one Binary Trait and one Quantitative Trait

| Prevalence | $\hat{h}^2$  | $\hat{h}_{\text{liab}}^2$ | $\hat{r}_g$  |
|------------|--------------|----------------|--------------|
| 0.01       | 0.72 (0.1)   | 0.59 (0.04)    | 0.51 (0.4)   |
| 0.05       | 0.72 (0.12)  | 0.59 (0.07)    | 0.45 (0.17)  |
| 0.2        | 0.72 (0.11)  | 0.6 (0.08)     | 0.46 (0.14)  |
| 0.5        | 0.73 (0.11)  | 0.59 (0.08)    | 0.42 (0.17)  |

Table S1: Simulations with one binary trait and one quantitative trait. The prevalence column describes the population prevalence of the binary trait. We ran 100 simulations for each prevalence. The $\hat{h}^2$ column shows the mean heritability estimate for the quantitative trait. The $\hat{h}_{\text{liab}}^2$ column shows the mean liability-scale heritability estimate for the binary trait. The $\hat{r}_g$ column shows the mean genetic correlation estimate. Standard deviations across 100 simulations in parentheses. The true parameter values were $r_g = 0.46$, $h^2 = 0.7$ for the quantitative trait and $h_{\text{liab}}^2 = 0.6$ for the binary trait. For all simulations, the quantitative trait sample size was 1000, the binary trait sample size was 1000 cases and 1000 controls, and there were 500 overlapping samples. There were 1000 effective independent SNPs. The environmental covariance was 0.2. We simulated case/control ascertainment using simulated LD block genotypes and a rejection sampling model of ascertainment. This is the same strategy used to simulate case/control ascertainment in [21].
Simulations with MAF- and LD-Dependent Genetic Architecture

| LD Score  | $h^2$(5-50%) | $\rho_g$(5-50%) | $r_g$(5-50%) |
|-----------|--------------|-----------------|--------------|
| Truth     | 0.83         | 0.42            | 0.5          |
| HM3       | 0.53 (0.08)  | 0.28 (0.07)     | 0.52 (0.1)   |
| PSG       | 0.36 (0.08)  | 0.18 (0.06)     | 0.5 (0.13)   |
| 30 Bins   | 0.81 (0.12)  | 0.41 (0.08)     | 0.51 (0.09)  |
| 60 Bins   | 0.81 (0.12)  | 0.41 (0.09)     | 0.51 (0.09)  |

Table S2: Simulations with MAF- and LD-dependent genetic architecture. Effect sizes were drawn from normal distributions such that the variance of per-allele effect sizes was uncorrelated with MAF, and variants with LD Score below 100 were fourfold enriched for heritability. Sample size was 2062 with complete overlap between studies; causal SNPs were about 600,000 best-guess imputed 1kG SNPs on chr 2, and the SNPs retained for the LD Score regression were the subset of about 100,000 of these SNPs that were included in HM3. True parameter values are shown in the top line of the table. Estimates are averages across 100 simulations. Standard deviations (in parentheses) are standard deviations across 100 simulations. LD Scores were estimated using in-sample LD and a 1cM window. HM3 means LD Score with sum taken over SNPs in HM3. PSG (per-standardized-genotype) means LD Score with the sum taken over all SNPs in 1kG as in [21]. 30 bins means per-allele LD Score binned on a MAF by LD Score grid with MAF breaks at 0.05, 0.1, 0.2, 0.3 and 0.4 and LD Score breaks at 35, 75, 150 and 400. 60 bins means per-allele LD Score binned on a MAF by LD Score grid with MAF breaks at 0.05, 0.1, 0.15, 0.2, 0.25, 0.3, 0.35, 0.4 and 0.45 and LD Score breaks at 30, 60, 120, 200 and 300. These simulations demonstrate that naive (HM3, PSG) LD Score regression gives correct genetic correlation estimates even when heritability and genetic covariance estimates are biased, so long as genetic correlation does not depend on LD.
Table S3: Sample sizes and references for traits analyzed in the main text.
Genetic Correlation between Educational Attainment Phenotypes

| Phenotype 1       | Phenotype 2                  | $r_g$ | $se$  |
|-------------------|-----------------------------|-------|-------|
| College (Yes/No)  | Years of Education          | 1.00  | 0.014 |

Table S4: Genetic correlation between the two educational attainment phenotypes from Rietveld, et al. [39].
Supplementary Figures

Genetic Correlations among Anthropometric Traits

Figure S1: Genetic correlations among anthropometric traits from studies by the GIANT and EGG consortia. The structure of the figure is the same as Figure 2 in the main text: blue corresponds to positive genetic correlations; red corresponds to negative genetic correlation. Larger squares correspond to more significant p-values. Genetic correlations that are different from zero at 1% FDR are shown as full-sized squares. Genetic correlations that are significantly different from zero at significance level 0.05 after Bonferroni correction are given an asterisk.
Figure S2: Genetic correlations among smoking traits from the Tobacco and Genetics (TAG) consortium. The structure of the figure is the same as Figure 2 in the main text: blue corresponds to positive genetic correlations; red corresponds to negative genetic correlation. Larger squares correspond to more significant p-values. Genetic correlations that are different from zero at 1% FDR are shown as full-sized squares. Genetic correlations that are significantly different from zero at significance level 0.05 after Bonferroni correction are given an asterisk.
Genetic Correlations among Insulin-Related Traits

Figure S3: Genetic correlations among insulin-related traits from studies by the MAGIC consortium. The structure of the figure is the same as Figure 2 in the main text: blue corresponds to positive genetic correlations; red corresponds to negative genetic correlation. Larger squares correspond to more significant p-values. Genetic correlations that are different from zero at 1% FDR are shown as full-sized squares. Genetic correlations that are significantly different from zero at significance level 0.05 after Bonferroni correction are given an asterisk.
Figure S4: This figure compares estimates of genetic correlations among metabolic traits from table 3 of Vattikuti et al. [17] to estimates from LD Score regression. The LD Score regression estimates used much larger sample sizes. Error bars are standard errors.
Schizophrenia — TG Conditional QQ Plot with and without the MHC

Figure S5: At left, we reproduced the conditional QQ plot comparing schizophrenia (SCZ) and triglycerides (TG) from Andreassen et al. [54] using the same data (PGC1 schizophrenia [76] and TG from Teslovich, et al. [71]). Conditional QQ plots show the distribution of p-values for SCZ conditional on the $-\log_{10}(p)$ for TG exceeding different thresholds. The thresholds are indicated by color, as described in the legends. Dark blue corresponds to no threshold, green corresponds to $-\log_{10}(p) > 1$, red corresponds to $-\log_{10}(p) > 2$ and light blue corresponds to $-\log_{10}(p) > 3$. The major histocompatibility complex (MHC, chr6, 25-35 MB) is a genomic region containing SNPs with exceptionally long-range LD and the strongest GWAS association for schizophrenia [62], as well as an association to TG [71]. If we remove the MHC, the signal of enrichment in the conditional QQ plot is substantially attenuated (middle); in particular, the red line falls below the green and blue lines (which correspond to less stringent thresholds for TG). If in addition we remove SNPs with very high LD Scores ($\ell > 200$, roughly the top 15% of SNPs), the signal of enrichment is further attenuated. The most likely explanation for the attenuation is that conditional QQ plots will report pleiotropy if causal SNPs are in LD (even if the causal SNPs for trait 1 are different from the casual SNPs for trait 2), which is more likely to occur in regions with long-range LD.
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