Better estimation of SNP heritability from summary statistics provides a new understanding of the genetic architecture of complex traits

Doug Speed\textsuperscript{1,2,*} and David J Balding\textsuperscript{2,3}

*Corresponding author: doug@aias.au.dk

\textsuperscript{1}Aarhus Institute of Advanced Studies (AIAS), Aarhus University, Denmark.
\textsuperscript{2}UCL Genetics Institute, University College London, United Kingdom.
\textsuperscript{3}Melbourne Integrative Genomics, School of BioSciences and School of Mathematics & Statistics, University of Melbourne, Australia.

LD Score Regression (LDSC) has been widely applied to the results of genome-wide association studies. However, its estimates of SNP heritability are derived from an unrealistic model in which each SNP is expected to contribute equal heritability. As a consequence, LDSC tends to over-estimate confounding bias, under-estimate the total phenotypic variation explained by SNPs, and provide misleading estimates of the heritability enrichment of SNP categories. Therefore, we present SumHer, software for estimating SNP heritability from summary statistics using more realistic heritability models. After demonstrating its superiority over LDSC, we apply SumHer to the results of 24 large-scale association studies (average sample size 121 000). First we show that these studies have tended to substantially over-correct for confounding, and as a result the number of genome-wide significant loci has under-reported by about 20\%. Next we estimate enrichment for 24 categories of SNPs defined by functional annotations. A previous study using LDSC reported that conserved regions were 13-fold enriched, and found a further twelve categories with above 2-fold enrichment. By contrast, our analysis using SumHer finds that conserved regions are only 1.6-fold (SD 0.06) enriched, and that no category has enrichment above 1.7-fold. SumHer provides an improved understanding of the genetic architecture of complex traits, which enables more efficient analysis of future genetic data.
Results

SumHer

SumHer has the same four aims as LDSC:1–3 we outline them here, with methodological details provided in Online Methods. Suppose we are provided with summary statistics from a GWAS where each of \( m \) SNPs has been tested individually for association with a particular trait. Suppose also that we have access to a well-matched reference panel, from which we can reliably estimate \( \nu_j^2 \), the squared correlation between SNPs \( j \) and \( l \). Let \( S_1, S_2, \ldots, S_m \) denote the \( \chi^2(1) \) test statistics from single-SNP analysis; the first aim is to estimate the average inflation of these test statistics due to confounding. Let \( h_j^2 \) denote the heritability directly contributed by SNP \( j \); the second aim is to estimate \( h_{SNP}^2 = \sum_{j=1}^m h_j^2 \), the SNP heritability of the trait.9 Let \( C \) index a category of SNPs; the third aim is to estimate \((\sum_{j\in C} h_j^2)/h_{SNP}^2\), the proportion of SNP heritability contributed by SNPs in \( C \) (we can then estimate the enrichment of the category by dividing its estimated proportion of SNP heritability by its expected proportion). Finally, if we are also provided with summary statistics from a second trait, the fourth aim is to estimate the correlation between SNP effect sizes for the two traits.10

In order to achieve these four aims, we must specify a heritability model, which describes how \( h_j^2 \) is expected to vary across the genome. Suppose this heritability model takes the form \( \mathbb{E}[h_j^2] \propto q_j \). The main difference between LDSC and SumHer is that SumHer allows for any heritability model (i.e., the user can specify arbitrary \( q_j \)), whereas LDSC assumes all \( q_j \) are the same. We recommend using SumHer with the “LDAK Model”: \( q_j = [f_j(1 - f_j)]^{0.75} w_j \), where \( f_j \) is the minor allele frequency (MAF) of SNP \( j \) and \( w_j \) is a weighting based on local levels of LD.5,6 In this model, a SNP with high MAF is expected to contribute more heritability than one with low MAF, while a SNP in a region of low LD is expected to contribute more than one in a region of high LD. By contrast, LDSC estimates are obtained by setting \( q_j = 1 \), which corresponds to the assumption that all SNPs are expected to contribute equally.1 We refer to this as the “GCTA Model” as this is a core assumption of the software GCTA.5,9

A second difference between LDSC and SumHer is how they estimate confounding bias. In a GWAS with no confounding, \( \mathbb{E}[S_j] = 1 + n \nu_j^2 \), where \( n \) is the sample size and \( \nu_j^2 = h_j^2 + \sum\text{near } j \nu_j^2 h_j^2 \) is the heritability tagged by SNP \( j \) (a working definition of “near” is within 1 Mb1). Both LDSC and SumHer estimate the deviation of test statistics from their expected values assuming no confounding. LDSC uses the model \( \mathbb{E}[S_j] \approx 1 + A + n \nu_j^2 \), where \( A \) indicates the average amount each test statistic is inflated additively due to confounding (LDSC then reports 1 + \( A \), which it refers to as “the intercept”). By contrast, we recommend using the model \( \mathbb{E}[S_j] \approx C(1 + n \nu_j^2) \), where \( C \) reflects how much each test statistic is inflated multiplicatively. There are two reasons why we prefer our approach. Firstly, it is standard practice to correct test statistics by scaling (i.e., divide each by \( C \));10 in theory, one could instead shift test statistics (i.e., subtract \( A \) from each), but this would result in some negative values. Secondly, the SumHer model for estimating bias accommodates test statistics that have been subjected to genomic control.11 Although genomic control is intended to reduce bias, we find that it is often the biggest source of bias in GWAS results and a major hindrance when using summary statistics to interrogate genetic architecture (see below).

In total we use six versions of SumHer, which differ according to their assumed heritability model and allowance for confounding bias. **LDSC-Zero** assumes the GCTA Model and that there is no bias \( (A = 0, C = 1) \); this is equivalent to using the LDSC software1 with the option --intercept-h2 1. **LDSC** assumes the GCTA Model and allows for additive bias \( (A \) free to vary, \( C = 1) \); this is equivalent to using the LDSC software1 with default options. **SumHer-Zero** assumes the LDAK Model and that there is no bias \( (A = 0, C = 1) \); this is our recommended version when estimating \( h_{SNP}^2 \) or enrichments and confident that confounding is negligible. **SumHer-GC** assumes the LDAK Model and allows for multiplicative bias \( (A = 0, C \) free to vary)\); this is our recommended version when estimating confounding or genetic correlations, or when estimating \( h_{SNP}^2 \) or enrichments and it is likely that test statistics are biased due to population structure or relatedness, or were obtained using genomic control or mixed-model association analysis (see below). **Hybrid-Zero** assumes the heritability model

\[
q_j = (1 - p) \times 1/m + p \times [f_j(1 - f_j)]^{0.75} w_j/Q'
\]

where \( Q' = \sum_j [f_j(1 - f_j)]^{0.75} w_j \), (1)

and that there is no bias \( (A = 0, C = 1) \), while **Hybrid-GC** assumes the same heritability model but allows for multiplicative bias \( (A = 0, C \) free to vary). Model (1) is a linear combination of the GCTA and LDAK models, where \( p \) indicates the weight assigned to
Estimates can depend sensitively on the heritability model. (a) We generate 500 phenotypes with $h^2_{SNP} = 0.5$ for each of three heritability models, GCTA, LDAK and Schoech et al.\textsuperscript{13} (see main text for details of each model). Bars report average estimates of $h^2_{SNP}$ from LDSC-Zero, LDSC, SumHer-Zero and SumHer-GC. (b) For the same phenotypes, bars now report average estimates of the enrichment of heritability in conserved regions from LDSC-Zero and SumHer-Zero (true enrichment is 1). For each method, the three bars correspond to estimating enrichment using a 2-part, 25-part or 53-part model (see main text). (c) For each of the three heritability models, we now generate 500 additional pairs of phenotypes with genetic correlation 0.5. Bars report average estimates of genetic correlation from LDSC-Zero, LDSC, SumHer-Zero and SumHer-GC. In all plots, vertical line segments mark 95% confidence intervals for the average estimates.

Figure 1: Estimates can depend sensitively on the heritability model. (a) We generate 500 phenotypes with $h^2_{SNP} = 0.5$ for each of three heritability models, GCTA, LDAK and Schoech et al.\textsuperscript{13} (see main text for details of each model). Bars report average estimates of $h^2_{SNP}$ from LDSC-Zero, LDSC, SumHer-Zero and SumHer-GC. (b) For the same phenotypes, bars now report average estimates of the enrichment of heritability in conserved regions from LDSC-Zero and SumHer-Zero (true enrichment is 1). For each method, the three bars correspond to estimating enrichment using a 2-part, 25-part or 53-part model (see main text). (c) For each of the three heritability models, we now generate 500 additional pairs of phenotypes with genetic correlation 0.5. Bars report average estimates of genetic correlation from LDSC-Zero, LDSC, SumHer-Zero and SumHer-GC. In all plots, vertical line segments mark 95% confidence intervals for the average estimates.

For all analyses, our reference panel is 8,850 unrelated Caucasian individuals from the Health and Retirement Study\textsuperscript{12} (HRS). When estimating enrichments of SNP categories, we use the 24 functional annotations used by Finucane et al.,\textsuperscript{3} which include coding, conserved, enhancer and promoter regions (see Supplementary Table 1 for a full list). When analyzing real data, we exclude SNPs within the major histocompatibility complex (Chromosome 6: 25–34 Mb), as well as SNPs which individually explain $>1\%$ of phenotypic variation, and SNPs in LD with these (within 1 cM and $r^2_{ij} > 0.1$).

Simulated phenotypes

We first demonstrate the importance of choosing an appropriate heritability model via simulations. For this we use 7,548 unrelated Caucasian individuals from the three control cohorts of the Wellcome Trust Case Control Consortium (WTCCC), recorded for 3,280,768 common SNPs (MAF $> 0.01$). We first generate 1,000 phenotypes each with 2,000 causal SNPs and $h^2_{SNP} = 0.5$; for half the phenotypes, we sample causal SNP effect sizes according to the GCTA Model, for the other half, according to the LDAK Model. We then analyze each phenotype using LDSC-Zero, LDSC, SumHer-Zero and SumHer-GC.

Figure 1a shows that, as expected, accurate estimates of $h^2_{SNP}$ are returned when phenotypes are analyzed assuming the matching heritability model (i.e., when GCTA phenotypes are analyzed using LDSC-Zero or when LDAK phenotypes are analyzed using SumHer-Zero), but that using a different heritability model can result in very poor estimates; SumHer-Zero tends to over-estimate $h^2_{SNP}$ by about 20% when applied to GCTA phenotypes, while LDSC-Zero tends to under-estimate $h^2_{SNP}$ by about 50% when applied to LDAK phenotypes. Supplementary Figure 1a shows LDSC correctly infers that there is no confounding when used on GCTA phenotypes (average intercept 1.000, SD 0.0003), and therefore its estimates of $h^2_{SNP}$ closely match those from LDSC-Zero. However, when used on LDAK phenotypes, LDSC wrongly infers that much of the causal signal is in fact confounding (average intercept 1.033, SD 0.0003), and as a result, its estimates of $h^2_{SNP}$ are on average about half those from LDSC-Zero and about 75% lower than the true value.

Figure 1b reports the estimated enrichment of SNPs in conserved regions. As causal SNPs were picked at random from across the
genome, the true enrichment is 1. Again, we see that assuming the correct heritability model produces reliable estimates, but assuming the wrong model can lead to misleading conclusions. In particular, we find that when LDSC-Zero is used to analyze LDAK phenotypes, it infers that conserved regions are at least 3-fold enriched for heritability. We chose to focus on conserved regions as this was the category that, by applying LDSC to real data, Finucane et al.\textsuperscript{3} found to be most enriched. We have previously shown that the LDAK Model better reflects real data than the GCTA Model\textsuperscript{5} (and provide further evidence below), and thus our simulations suggest that a substantial portion of the enrichment observed by Finucane et al. is an artifact of misspecifying the heritability model.

Figure 1b also examines how estimates of enrichment are affected by the way the genome is divided into SNP categories (Supplementary Fig. 1c provides a zoomed-in version). The simplest approach is to use a 2-part model, in this case partitioning SNPs into those inside and those outside conserved regions. Next we use a 25-part model, dividing the genome into the 24 functional categories (of which conserved regions are one), plus a category containing all SNPs. Finally, we use a 53-part model, constructed by adding 28 “buffer regions”, the approach recommended by Finucane et al. We find that when the correct heritability model is assumed, results appear insensitive to the approach used. However, when the wrong model is assumed, the three approaches can give substantially different estimates.

As well as the GCTA and LDAK phenotypes, for each analysis we additionally generate phenotypes according to a heritability model recently proposed by Schoech et al.\textsuperscript{13} We formally define this model in the Discussion, however, loosely speaking, it is intermediate between the GCTA and LDAK models. This is reflected by the estimates of $h^2_{SNP}$ in Figure 1a: on average those from LDSC-Zero are about 5% too low, while those from SumHer-Zero are about 10% too high. We also consider changing the reference panel. While we prefer using the HRS dataset, as its larger sample size enables more accurate estimation of $r^2_{jl}$, Supplementary Figure 2 shows that estimates are similar if we instead use the 404 non-Finnish Europeans from the 1000 Genomes Project.\textsuperscript{14}

Real phenotypes

We now show that the choice of heritability model is also important when analyzing real data. We use “25 raw GWAS” (18 binary traits, 7 quantitative, average sample size 9 700; see Supplementary Table 2), for which we have individual-level genotype and phenotype data from either the WTCCC\textsuperscript{15} or the eMERGE Network.\textsuperscript{16} The 13 WTCCC GWAS all examine diseases (e.g., Bipolar Disorder, Ischaemic Stroke and Parkinson’s Disease), while the 12 eMERGE GWAS consider a mixture of diseases (e.g., age-related macular degeneration, heart failure and peripheral artery disease) and clinical measurements (e.g., blood pressure, height and lipid levels). After imputation, strict quality control, and excluding SNPs not present in our reference panel, the WTCCC data contain on average 2.3 M SNPs, while the eMERGE data contain 2 972 162 SNPs. When performing single-SNP analysis, we include as covariates sex and ten principal components (five calculated from the data, five derived from the 1000 Genomes Project\textsuperscript{14}). As we have access to raw data, we can use REML to estimate how much of the total phenotypic variance explained by SNPs is inflation due to population structure or relatedness.\textsuperscript{17,18} We estimate that on average 3.6% of the variance explained is inflation (range -0.9% to 6.8%), indicating that confounding due to population structure and relatedness is modest (Supplementary Fig. 3).

Figure 2a and Supplementary Table 2 show that across the 25 traits, estimates of $h^2_{SNP}$ from SumHer-Zero are on average 2.0 times larger (SD 0.05) than those from LDSC-Zero. Figure 2b and Supplementary Table 3 report estimates of enrichment for the 24 functional categories, averaged across the 25 traits. First we estimate enrichment using LDSC-Zero with a 53-part model (the approach
Figure 2: Importance of the heritability model for the 25 raw GWAS. (a) Estimates of $h^2_{SNP}$ from LDSC-Zero (x axis) and SumHer-Zero (y axis). Colors distinguish between the 13 WTCCC, the 5 binary eMERGE and 7 quantitative eMERGE traits (black denotes the 25-trait average). Horizontal and vertical line segments mark 95% confidence intervals. Estimates for binary traits are on the observed scale. (b) Average estimates of enrichment for the 24 functional categories from LDSC-Zero (using a 53-part model) and SumHer-Zero (using a 25-part model). Colors indicate significant enrichments ($P < 0.05$) from one or both methods. (c) Estimates of genetic correlations between pairs of eMERGE traits using LDSC (x axis) and SumHer-GC (y axis). Colors indicate significant correlations ($P < 0.05$) from one or both methods.

taken by Finucane et al.\(^3\)), then using SumHer-Zero with a 24-part model (our recommended approach). While there is reasonable agreement between which categories SumHer-Zero and LDSC-Zero declare to be significant, their estimates of enrichment are very different. Notably, LDSC-Zero estimates that conserved regions on average contribute 28% (SD 4) of SNP heritability (corresponding to 11-fold enrichment), whereas SumHer-Zero estimates that they contribute only 5.1% (SD 0.7) of SNP heritability (1.6-fold enrichment). Supplementary Figure 4 compares estimates of enrichment from 2-part, 25-part and 53-part models. When we use LDSC-Zero, we find substantial differences between the results of the three approaches; in particular, not one of the 24 estimates from the 2-part model is consistent ($P > 0.05/24$) with either of the corresponding estimates from the 25-way or 53-way models. By contrast, when we use SumHer-Zero, there is much stronger concordance between the three sets of results; for example, all 24 (19 out of 24) of the 2-part model estimates are consistent with those from the 25-way (53-way) model.

Figure 2c and Supplementary Table 4 report genetic correlations between pairs of traits. In general, it is only possible to get a meaningful estimate when both traits have substantial $h^2_{SNP}$, so we restrict to the 18 traits for which both LDSC-Zero and SumHer-Zero find significant $h^2_{SNP}$ ($P < 0.05/25$). As predicted by our earlier simulations, we find good concordance between the genetic correlations reported by LDSC and SumHer-GC, but that the latter produces more precise estimates: on average the SumHer-GC estimates have SD about half that of the LDSC estimates, and SumHer-GC finds 18 pairs of traits with significant correlation ($P < 0.05$), whereas LDSC finds only 4.

Comparing heritability models

Previously, we compared different heritability models based on the likelihood from REML analysis; we showed that across 42 human traits (average sample size 7,400), log likelihood was on average 9.8 higher if we assumed the LDAK Model rather than the GCTA Model.\(^5\) Those 42 traits included the 13 WTCCC traits we study here. If we repeat that analysis using the 12 eMERGE traits (average sample size 13,000), we find that log likelihood is on average 9.8 higher if we assumed the LDAK Model rather than the GCTA Model. In Supplementary Table 6, we show that it remains possible to compare models based on likelihood if only summary statistics are available. However, an alternative, and easier to visualize, method is to fit both the GCTA and LDAK Models simultaneously and allow the data to decide the relative weighting of each. Specifically, we use Hybrid-Zero, with the focus on estimating $p$ in Model (1), the “proportion of LDAK” in the heritability model. Figure 3a demonstrates proof of principle: we see that Hybrid-Zero correctly estimates $p = 0$ when
applied to GCTA phenotypes, $p = 1$ for LDAK phenotypes, and $p = 0.5$ for “Hybrid Phenotypes” (each created by summing a GCTA and an LDAK phenotype). We then apply Hybrid-Zero to our 25 raw GWAS (Figure 3b and Supplementary Table 6), finding on average $p = 1.03$ (SD 0.02), indicating that the data overwhelmingly support the LDAK Model over the GCTA Model.

**Population structure and relatedness**

Until recently, it was standard to estimate confounding bias via the genomic inflation factor\(^{11}\) (GIF). However, the GIF tends to over-estimate bias, because it makes the assumption that all observed inflation of test statistics is due to confounding.\(^{19}\) LDSC provides a method for estimating confounding bias which appreciates that substantial inflation can instead be due to causal variation.\(^{1}\) However, our above simulations indicate that LDSC also tends to over-estimate bias, due to the poor fit of the GCTA heritability model. This is supported by our analysis of the 25 raw GWAS (Supplementary Table 2), for which we performed very careful quality control and verified using REML that confounding due to population structure and relatedness is modest (Supplementary Fig. 3). LDSC finds substantial bias; its average estimate of the intercept ($1 + A$) is 1.031 (SD 0.002), which is similar to the average GIF, 1.035. By contrast, SumHer-GC finds that bias is slight; its average estimate of the scaling factor ($C$) is 1.001 (SD 0.002).

We now construct GWAS when there is substantial confounding. For each of the 13 WTCCC GWAS, we replace 2504 of the controls (on average 67%) with 2504 individuals from POBI\(^{20}\) (People of the British Isles); this generates population structure because, although both WTCCC and POBI individuals were recruited from the UK, the latter predominately came from isolated, rural regions (Supplementary Figure 5). Supplementary Table 7 reports estimates of confounding bias and $h^2_{\text{SNP}}$ for each GWAS. SumHer-GC now finds substantial bias; its average estimate of the scaling factor is 1.049 (SD 0.003). For comparison, the average GIF is 1.098, while the average LDSC intercept is 1.088 (SD 0.002). SumHer-Zero estimates of $h^2_{\text{SNP}}$ after switching controls are on average 95% (SD 4) higher than SumHer-Zero estimates prior to switching, demonstrating how population structure can lead to substantial inflation when estimating SNP heritability. However, SumHer-GC estimates of $h^2_{\text{SNP}}$ after switching controls are on average only 4% (SD 7) higher, consistent with SumHer-GC being able to reliably estimate SNP heritability despite the population structure.

**Genomic control and mixed-model association analysis**

Although its use is declining, it remains that the majority of published GWAS have performed genomic control (divided test statistics by the GIF) at least once in their analyses.\(^{11}\) As the GIF tends to over-estimate confounding,\(^{19}\) genomic control tends to produce negatively
Figure 4: **Correcting for genomic control.** For the 25 raw GWAS, each plot compares estimates from SumHer using raw test statistics (x-axis) with those using test statistics subjected to genomic control (y-axis). Horizontal and vertical line segments mark 95% confidence intervals. (a) Estimates of $h^2_{SNP}$ using SumHer-Zero both pre and post genomic control. (b) Estimates of $h^2_{SNP}$ using SumHer-Zero pre genomic control and SumHer-GC post genomic control. (c) Estimates of average enrichment for the 24 functional categories using SumHer-Zero both pre and post genomic control. (d) Estimates of average enrichment for the 24 functional categories using SumHer-Zero pre genomic control and SumHer-GC post genomic control.

biased test statistics. For this reason, we recommend using SumHer-GC to estimate $h^2_{SNP}$ and enrichment, instead of SumHer-Zero, if it is likely that genomic control has been applied. By way of demonstration, we perform genomic control for each of the 25 raw GWAS. Figure 4a shows that if we continue to use SumHer-Zero, estimates of $h^2_{SNP}$ are centered on zero. However, if we instead use SumHer-GC, estimates of $h^2_{SNP}$ are in general consistent with estimates from SumHer-Zero prior to genomic control (Figure 4b); an exception is stroke, indicative of the LDAK Model being sub-optimal for this trait (Supplementary Figure 6). Similarly, Figures 4c & d show that if genomic control has been performed, it is beneficial to use SumHer-GC instead of SumHer-Zero when estimating enrichments.

SumHer, like LDSC, is designed to be used with test statistics from classical regression (see Online Methods). However, with the development of software such as Fast-LMM, GCTA-LOCO and Bolt-LMM, a popular alternative is for GWAS to use mixed-model association analysis. Supplementary Figures 6 & 7 show that when estimating $h^2_{SNP}$ and enrichment, the impact of mixed-model analysis is similar to, albeit less severe than, that of genomic control. For example, SumHer-Zero estimates of $h^2_{SNP}$ based on mixed-model test statistics are on average about half those based on classical test statistics. However, as with genomic control, reliable estimates can be obtained by using SumHer-GC instead of SumHer-Zero.

**Analysis of published GWAS results**

For our main analysis, we use “24 summary GWAS” (12 binary traits, 12 quantitative, average sample size 121,000; see Table 1). These are GWAS for which we do not have individual-level data, but have downloaded summary statistics from previously-published analyses. After excluding SNPs not present in our reference panel, on average there are summary statistics for 2.2 M SNPs per trait. All of the GWAS either performed genomic control or used mixed-model association analysis, so in general we use SumHer-GC. However, we also repeat all analyses using SumHer-Zero restricted to the 11 traits that did not use mixed-model analysis and for which the impact of genomic control was lowest (Supplementary Table 8).

Figure 3c reports estimates of $p$, the proportion of LDAK in the heritability model, obtained using Hybrid-GC. Across the 24 traits, we estimate $p = 0.91$ (SD 0.01), indicating that again the data strongly support the LDAK Model over the GCTA Model, although not as strongly as for the 25 raw GWAS (see Discussion). Supplementary Table 9 shows that we reach the same conclusion if we instead use Hybrid-Zero restricted to the 11 traits least impacted by genomic control ($p = 0.90$; SD 0.01), or if we compare the GCTA and LDAK Models based on likelihood (the log likelihood is on average 103 higher under the LDAK Model than the GCTA Model).

**Confounding**

Table 1 reports the LDSC intercept \((1 + A)\) and the SumHer-GC scaling factor \((C)\) for each trait. LDSC finds that test statistics are on
Table 1: Estimates of confounding bias for the 24 summary GWAS. Columns 2 & 3 report the average sample size ($n$) and genomic inflation factor (GIF) for each trait. Columns 4-7 report estimates of $h^2_{SNP}$ and confounding from both LDSC and SumHer-GC (LDSC measures confounding via the intercept $1 + A$, while SumHer-GC uses the scaling factor $C$). Columns 8-11 report the number of significant loci based on the published test statistics, then after correction via genomic control, LDSC and SumHer-GC (dividing test statistics by the GIF, $1 + A$ and $C$, respectively). When estimating $h^2_{SNP}$, LDSC requires uncorrected test statistics from classical regression. However the test statistics from all 24 GWAS were calculated either using genomic control or mixed-model association analysis; † indicates the 11 traits for which classical regression was used and the impact of genomic control was lowest.
average inflated by 5.7% (SD 0.2), implying that the GWAS tended to under-correct for confounding. By contrast, SumHer-GC finds that test statistics are on average deflated by 7.4% (SD 0.3), indicating that the studies tended to over-correct. We note that the four GWAS with lowest $C$ (0.56 for body mass index, 0.74 for HDL and LDL cholesterol, and 0.73 for triglyceride levels), are all meta-analyses that used genomic control both before and after combining results across cohorts.\textsuperscript{35,38} Table 1 also reports the number of independent loci with $P < 5 \times 10^{-8}$. Without adjustment (i.e., using the published test statistics), there are on average 73 loci per trait. If we correct using LDSC, this number is reduced to 51, but if we correct using SumHer-GC, it increases to 91. For these counts, we defined two significant SNPs as dependent if they are within 1 cM and have $r^2_{jl} > 0.05$, but we find that results are very similar if instead we increase the window size to 3 cM, or use $r^2_{jl} > 0.2$ (Supplementary Table 10).

**Functional enrichments**

Figure 5a and Supplementary Table 11 report estimates of enrichment for the 24 functional categories, averaged across the 24 traits. We again see striking differences between the estimates from LDSC (using a 53-part model) and those from SumHer-GC (using a 25-part model). For example, LDSC estimates of enrichment range from -1.5 to 7.8, whereas SumHer-GC estimates range from 0.80 to 1.7. Supplementary Table 12 shows that large differences remain if we instead estimate enrichment using LDSC-Zero and SumHer-Zero restricted to the 11 traits least impacted by genomic control. Based on the results from SumHer-GC, we conclude that conserved regions\textsuperscript{44,45} and transcription start sites\textsuperscript{46} are most enriched for heritability; they are estimated to contribute 5.8% (SD 0.2) and 3.0% (SD 0.2) of $h^2_{SNP}$, respectively, 1.6 (SD 0.06) and 1.7 (SD 0.08) times higher than their expected contributions under the LADAK Model. Repressed regions\textsuperscript{46} are the only category significantly depleted; although they are estimated to contribute 36% (SD 0.4) of $h^2_{SNP}$, this is 0.80 (SD 0.01) times lower than expected. Further results are provided in Supplementary Figure 8, which confirm that LDSC estimates are similar if we change the SNP sets (LDSC recommends that the reference panel contains as many SNPs as possible, but that only HapMap\textsuperscript{34} SNPs with MAF > 0.05 are used when performing the regression\textsuperscript{1,3}), or if we use the 75-part model used by Gazal \textit{et al.}\textsuperscript{48}

**Genetic correlations**

Supplementary Figure 9 and Supplementary Table 13 provide estimates of genetic correlations for the 276 pairs of traits. As expected, there is strong concordance between estimates from LDSC and SumHer-GC, but the SumHer-GC estimates are more precise; for example, across the 41 pairs of traits that both methods find to be significantly correlated ($P < 0.05/276$), the SD of SumHer-GC estimates is on average a third lower than the SD of LDSC estimates.
Figure 6: Prediction of five quantitative traits. For each trait, we use data from the 24 summary GWAS to construct Bayesian polygenic risk scores (PRS) corresponding to four heritability models: GCTA, Enriched-GCTA, LDAK and Enriched-LDAK (see main text for details of each model). (a) The prior distribution of $\nu_j^2$, the heritability tagged by SNP $j$, corresponding to each heritability model. (b) Prediction accuracy, measured as correlation between observed and predicted phenotypes in the (independent) eMERGE data, for the Classical PRS (effect sizes are frequentist estimates from single-SNP analysis) and for each of the four Bayesian PRS (effect sizes are posterior means). Vertical line segments mark 95% confidence intervals.

Improving the efficiency of future analyses

Figure 5b shows that there is strong concordance between the average estimates of enrichment obtained from the 12 binary traits and those from the 12 quantitative traits. This suggests broad similarities between the genetic architectures of different traits, which in turn implies that it should be possible to use information from existing GWAS to improve the efficiency of future analyses. As a demonstration, we consider prediction using polygenic risk scores (PRS). We focus on body mass index, height, HDL & LDL cholesterol and triglyceride levels, as for these five traits we can train prediction models using the 24 summary GWAS, then measure how well these perform on the independent eMERGE data.

To construct a PRS, we need estimates of SNP effect sizes. The current standard is to use estimates from single-SNP analysis (“Classical PRS”). However, in Online Methods, we explain how, given a heritability model, we can obtain a prior distribution for $\nu_j^2$, the heritability tagged by SNP $j$, then calculate a “Bayesian PRS” using the posterior mean effect sizes. For each trait, we construct four Bayesian PRS corresponding to four heritability models. First we use the GCTA heritability model ($q_j = 1$). Next we use the “Enriched GCTA Model”, obtained by scaling the $q_j$ based on the (53-part) estimates of enrichment (e.g., if a category was estimated to have 2-fold enrichment, then the SNPs it contains would have average $q_j = 2$). We similarly construct PRS based on the LDAK Model, then the “Enriched LDAK Model”, where the $q_j$ are scaled according to the SumHer-GC (25-part) estimates of enrichment.

Figure 6a compares the four prior distributions for $\nu_j^2$. We see the two priors derived from the GCTA Model are more diffuse than the two from the LDAK Model. As explained in the Online Methods, this is because the GCTA Model predicts that $\nu_j^2$ scales with local levels of LD, which vary considerably across the genome, whereas the LDAK Model predicts that $\nu_j^2$ scales with local MAFs, which vary less. Figure 6b and Supplementary Table 14 report the performance of each PRS, measured as correlation between predicted and observed phenotypes for the eMERGE individuals. Averaged across the five traits, the Bayesian PRS constructed from the GCTA and Enriched GCTA Model are, respectively, 1.1% (SD 2.0) worse and 2.3% (SD 2.0) better than the Classical PRS, whereas the Bayesian PRS constructed from the LDAK and Enriched LDAK Model are, respectively, 5.8% (SD 2.0) and 7.5% (SD 2.0) better. The fact that the PRS based on the Enriched GCTA Model outperforms the PRS based on the GCTA Model, is because introducing partitions relaxes the unrealistic assumption that $\nu_j^2$ scales with local levels of LD; however, the fact that the PRS based on the LDAK Model outperforms the PRS based on the Enriched GCTA Model, reflects that it is better to start with a more realistic model, than try to fix a sub-optimal model by partitioning.\(^5\)
Discussion

We have presented SumHer, software for estimating confounding bias, SNP heritability, enrichments of heritability and genetic correlations from GWAS results. While the aims of SumHer are the same as those of LDSC, the key difference is that SumHer allows the user to specify the heritability model. If SumHer is run using the GCTA Model, its estimates will match those from LDSC. However, we instead recommend using the LDAK Model, which we have shown better reflects real data, and therefore produces more accurate estimates. We have analyzed GWAS results for tens of traits, showing that the impact of using an improved heritability model is often substantial, and overall provides a very different description of the genetic architecture of complex traits than has to date been obtained from LDSC analyses.

While the GCTA Model has been used almost exclusively when estimating SNP heritability from summary statistics, alternative models have been used when analyzing individual-level data. A recent submission by Schoech et al., which has three authors in common with the original LDSC publication, uses the following model: $q_j = [f_j(1 - f_j)]^{0.62}(1 - 0.3LLD_j)$, where $LLD_j$ is a local measure of tagging (obtained by computing LD Scores, binning by MAF, quantile normalizing, then truncating). Like the LDAK Model, the model of Schoech et al. assumes that a SNP with high MAF contributes more heritability than one with low MAF, and that a SNP in a region of low LD contributes more than one in a region of high LD. Schoech et al. claim that this model is more realistic than both the GCTA Model and the LDAK Model. However, while they compared their model with the GCTA Model on real data (comparing the two models according to REML likelihood), their only comparison with the LDAK Model was using phenotypes simulated according to their heritability model. Supplementary Table 5 compares the model of Schoech et al. to the GCTA and LDAK Models for the 25 raw GWAS. While we agree with Schoech et al. that their model is superior to the GCTA Model (on average, its log likelihood is 9.4 higher), we find it to be inferior to the LDAK Model (on average, its log likelihood is 9.5 lower). Moreover, Figure 1 shows that even if the model of Schoech et al. does accurately reflect the genetic architecture of complex traits, it would remain the case that LDSC is producing inaccurate estimates of confounding bias, $h^2_{SNP}$ and enrichment.

We note that, despite remaining superior to the GCTA Model, the LDAK Model does not perform as well for the 24 summary GWAS (average LDAK Proportion 0.91) as for the 25 raw GWAS (average 1.02). One possible reason for this is that for the 24 summary GWAS we had to rely on the quality control performed by the original analysts, which was generally much less strict than we would recommend for heritability analysis. For example, when analyzing the 25 raw GWAS, we restricted to SNPs with imputation information score $> 0.99$, whereas for the 24 summary GWAS, the most common thresholds were 0.5 and 0.8. There will be a correlation between the heritability of a SNP and its genotyping certainty (a SNP genotyped with error will tag less causal variation than were it perfectly typed), and similarly, there will be a correlation between genotyping certainty and local levels of LD (low-LD regions tend to contain more low-MAF SNPs, which are often hard to genotype reliably, while imputation is easier in high-LD regions). Therefore, including lower-certainty SNPs in a GWAS will generate correlation between the heritability of each SNP and levels of LD, which will result in traits appearing more “GCTA-like”. Alternatively, the results for the 24 summary traits might simply reflect that the LDAK Model is not perfect, and will fit some traits better than others. Therefore, we encourage readers to find ways to improve the LDAK Model, either generally or on a per-trait level, which they can do using the tools provided by SumHer for testing and comparing different heritability models on large-scale GWAS data.

Two practical advantages of SumHer over LDSC are evidenced by the results for height in Table 1 (for these we used summary statistics from the most recent GIANT Consortium meta-analysis, which has average sample size 245 000). First we focus on estimates of $h^2_{SNP}$. Accurate estimates of $h^2_{SNP}$ are important not only because they improve our understanding of genetic architecture, but also because they are now being incorporated in software for analyzing complex traits (e.g., the mixed-model association software Bolt-LMM and the prediction software LDpred). Multiple studies have estimated that common SNPs explain at least 40% of the variation in height, this indicates that the SumHer-GC estimate of $h^2_{SNP}$ (0.38, SD 0.03) is closer to the truth than the LDSC estimate (0.16, SD 0.01). Complementary to estimates of $h^2_{SNP}$ are estimates of confounding bias. These are used both to assess the quality of a GWAS and when correcting test statistics. LDSC estimates the intercept ($1 + A$) to be 1.73 (SD 0.05), implying that there is substantial bias and that strong correction is required; were we to divide test statistics by 1.73, the number of significant loci drops by over half.
By contrast, SumHer-GC estimates the scaling factor \((C)\) to be 1.02 (SD 0.03), indicating that confounding is slight and that almost no correction of test statistics is needed (dividing the test statistics by 1.02 reduces the number of significant loci by only 3%).

The most striking differences between LDSC and SumHer are observed when estimating heritability enrichments, as highlighted in Figure 5a. Whereas analyses using LDSC have found that heritability is highly focused in specific genomic regions, SumHer instead shows that heritability is spread far more diffusely across the genome, supporting an omnigenic view of genetic architecture. While the realization that complex traits are even more complicated than previously thought is daunting, as our prediction example demonstrates, it is only by properly understanding their complexity that we can develop more efficient tools for analyzing genetic data.

URLs

LDAK, http://www.ldak.org/; LDSC, http://www.github.com/bulik/ldsc; 24 functional annotations, https://data.broadinstitute.org/alkesgroup/LDSCORE/baseline_bedfiles.tgz.

Methods

Methods, including statements of data availability and any associated accession codes and references, are available in the online version of the paper.

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Author contributions

D.S. performed the analysis, D.S. and D.J.B. wrote the manuscript.
Competing financial interests

The authors declare no competing financial interests.

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Online Methods

The Supplementary Note provides step-by-step code for using SumHer to estimate SNP heritability, confounding bias, heritability enrichments and genetic correlations from summary statistics.

Estimating SNP heritability. Suppose that we have summary statistics from a GWAS on $n$ individuals and $m$ SNPs; let $S_j$ denote the $\chi^2(1)$ test statistic from regressing the phenotype on $X_j$, the vector of additively-coded genotypes for SNP $j$, and let $n_j \leq n$ denote the number of individuals used in this regression (note that in the main text, for simplicity, we assumed $n_j = n$). If $S_j$ was obtained using classical (i.e., least-squares) linear regression, then

$$\mathbb{E}[S_j] \approx 1 + n_j v_j^2 \quad \text{with} \quad v_j^2 = h_j^2 + \sum_{l \neq j} \text{Cor}(X_j, X_l)^2 h_l^2,$$

(2)

where $v_j^2$ is the total amount of heritability tagged by SNP $j$. In the main text, we referred to $h_j^2$ as the heritability “directly contributed” by SNP $j$, to emphasize that while a causal variant can contribute to multiple $v_j^2$ (i.e., be tagged by multiple SNPs), it can only contribute to one $h_j^2$. More formally, the $h_j^2$ represent a partitioning of $h_{\text{SNP}}^2$, the total heritability tagged by the $m$ SNPs genotyped by the GWAS; this formal definition appreciates that a causal variant need not be typed to contribute towards $h_{\text{SNP}}^2$, provided it is tagged by one or more SNPs that have been typed (in which case its heritability will be shared across the $h_j^2$ of the tagging SNPs, even though none of these “directly contribute” this heritability). If there is no population structure or cryptic relatedness, then $\text{Cor}(X_j, X_i)^2$ will be negligible for distant SNPs, while for local SNPs, an unbiased estimate of $\text{Cor}(X_j, X_i)^2$ is

$$v_{ji}^2 = \text{Cor}(X'_j, X'_i)^2 - \frac{1 - \text{Cor}(X'_j, X'_i)^2}{n' - 2},$$
where $X_j'$ is the vector of SNP $j$ genotypes for the $n'$ individuals in a reference panel (for accurate estimates of $r_{jkl}$, these individuals should have similar ancestry to those used in the GWAS). Therefore, in place of Equation (2) we use

$$\mathbb{E}[S_j] \approx 1 + n_j v_j^2 \quad \text{with} \quad v_j^2 = h_j^2 + \sum_{i \in N_j} r_{j1}^2 h_i^2,$$

(3)

where the set $N_j$ indexes those SNPs “near” SNP $j$; a working definition of near is two SNPs within 1 cM (Supplementary Figure 2). Finally, given a heritability model defined as $\mathbb{E}[h_j^2] \propto q_j$, we replace $h_j^2$ by its expected value $q_j h_{\text{SNP}}^2/Q$, where $h_{\text{SNP}}^2 = \sum h_j^2$ and $Q = \sum q_j$, resulting in

$$\mathbb{E}[S_j] \approx 1 + u_j h_{\text{SNP}}^2 \quad \text{with} \quad u_j = (q_j + \sum_{i \in N_j} q_i r_{j1}^2) n_j/Q,$$

(4)

which allows us to estimate $h_{\text{SNP}}^2$ by regressing $(S_j - 1)$ on $u_j$. To account for correlated datapoints and heteroscedasticity we use weighted least squares regression. Specifically, if $D$ is a diagonal matrix whose non-zero entries are the regression weights, then the estimate of $h_{\text{SNP}}^2$ would be $(a^T D a)^{-1} a^T D (S - 1)$, where $a$ and $S$ are vectors containing the $m$ values for $u_j$ and $S_j$, respectively. Following LDSC, we use $1/D_{jj} = (\sum_{i \in N_j} r_{j1}^2)(1 + u_j h_{\text{SNP}}^2)$. Since $h_{\text{SNP}}^2$ is unknown, we proceed iteratively starting at $h_{\text{SNP}}^2 = 0$, then successively updating $D_{jj}$ and $h_{\text{SNP}}^2$ until convergence. Again following LDSC, we estimate standard errors via block jackknifing (by default we use 200 blocks).

Comparing heritability models. The (weighted) log likelihood is

$$L(S_j|h_{\text{SNP}}^2, D) = -\frac{d}{2} (2\pi \gamma/d + 1) \quad \text{with} \quad d = \sum_j D_{jj} \quad \text{and} \quad \gamma = \sum_j D_{jj} (S_j - 1 - u_j h_{\text{SNP}}^2)^2.$$

(5)

To evaluate $L(S_j|h_{\text{SNP}}^2, D)$ requires values for $h_{\text{SNP}}^2$ and $D$. While the natural choice is to use the values after the final iteration, in order to compare different heritability models based on $L(S_j|h_{\text{SNP}}^2, D)$, we must use the same $D$ for each (else models leading to lower $D_{jj}$ would have an unfair advantage). Therefore, SumHer reports $L(S_j|h_{\text{SNP}}^2, D^0)$, where $h_{\text{SNP}}^2$ is the final estimate of $h_{\text{SNP}}^2$, but $D^0$ is the initial weight matrix (obtained by setting $1/D_{jj} = \sum_{i \in N_j} r_{j1}^2$); Supplementary Table 6 shows that for the 25 raw GWAS, comparisons of heritability models based on $L(S_j|h_{\text{SNP}}^2, D^0)$ align closely with those based on REML likelihood.

Estimating confounding bias. We recommend replacing Equation (3) with $\mathbb{E}[S_j] \approx C(1 + u_j h_{\text{SNP}}^2)$, where $C$ denotes the multiplicative inflation of test statistics due to confounding; we can then estimate $C$ and $Ch_{\text{SNP}}^2$ by jointly regressing $S_j$ on 1 and $u_j$. To instead copy LDSC, which considers the additive inflation of test statistics, we replace Equation (4) with $\mathbb{E}[S_j] \approx 1 + A + u_j h_{\text{SNP}}^2$, then estimate $A$ and $h_{\text{SNP}}^2$ by jointly regressing $(S_j - 1)$ on 1 and $u_j$.

Estimating enrichments. Suppose we have $K$ categories; let $I_{jk} \in \{0, 1\}$ indicate whether SNP $j$ belongs to Category $k$. We wish to estimate $h_{\text{Cat}}^2 = \sum_k I_{jk} h_{\text{Part}}^2$, the heritability contributed by SNPs in Category $k$. We now use the heritability model

$$\mathbb{E}[h_j^2] = q_j \sum_k I_{jk} h_{\text{Part}}^2/Q_k \quad \text{with} \quad Q_k = \sum_j q_j I_{jk},$$

(6)

where $q_j h_{\text{Part}}^2/Q_k$ indicates the contribution of Category $k$ to the expected heritability of SNP $j$, and $h_{\text{SNP}}^2 = h_{\text{Part}1}^2 + \ldots + h_{\text{Part}K}^2$. We consider two scenarios: in Scenario 1, the $K$ categories partition the genome ($\sum_k I_{jk} = 1$); in Scenario 2, the first $K - 1$ categories correspond to annotations, and the $K$th is the base category containing all SNPs ($I_{jk} = 1$). Using Equation (6) ensures that when there is no enrichment, $\mathbb{E}[h_j^2] = q_j h_{\text{SNP}}^2/Q$. To appreciate why, consider that for Scenario 1, $h_{\text{Part}}^2 = h_{\text{Cat}}^2$, so that when no categories are enriched, $h_{\text{Part}}^2 = Q_k h_{\text{SNP}}^2/Q$; for Scenario 2, $h_{\text{Part}1}^2, \ldots, h_{\text{Part}K-1}^2$ indicate how much each annotation increases the SNP heritability from $h_{\text{Part}}^2$, so that when there is no enrichment, $h_{\text{Part}1}^2 = \ldots = h_{\text{Part}K-1}^2 = 0$ and $h_{\text{Part}}^2 = h_{\text{SNP}}^2$. Now when we replace $h_j^2$ in Equation (3) by its expected value, we obtain

$$\mathbb{E}[S_j] \approx 1 + \sum_k u_{jk} h_{\text{Part}}^2 \quad \text{with} \quad u_{jk} = (q_j I_{jk} + \sum_{l \in N_j} q_l I_{lk} r_{j1}^2) n_j/Q_k,$$

(7)

and therefore we can estimate $h_{\text{Part}1}^2, \ldots, h_{\text{Part}K}^2$ by jointly regressing $(S_j - 1)$ on $u_{j1}, \ldots, u_{jk}$. Given these, our estimate of $h_{\text{Cat}}^2$ is $\sum_k (\sum_j I_{jk} q_j I_{jk}) h_{\text{Part}k}^2/Q_k$, which we then divide by $Q_k h_{\text{SNP}}^2/Q$ to get an estimate of the enrichment of Category $k$. 

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Estimating genetic correlation. Suppose we have summary statistics from two GWAS. Instead of \( \chi^2(1) \) test statistics, we now use (signed) \( Z \)-statistics. Let \( Z_{A_j} \) and \( Z_{B_j} \) denote the two \( Z \)-statistics for SNP \( j \), computed using \( n_{A_j} \) and \( n_{B_j} \) individuals, respectively, of which \( n_{C_j} \) were common to both GWAS (if the two GWAS were independent, \( n_{C_j} = 0 \)). We assume

\[
E[Z_{A_j}Z_{B_j}] \approx \frac{c_{AB}n_{B_j}}{n_{A_j}n_{B_j}} + u'_j h_{AB}^2 \quad \text{with} \quad u'_j = (q_j + \sum_{l \in N_j} q_l r_{jl}^2) / \sqrt{n_{A_j}n_{B_j}} / Q,
\]

where \( c_{AB} \) is the phenotypic correlation between the two traits and \( h_{AB}^2 \) is their genetic covariance. This equation matches that used by LDSC,\(^2\) except we have replaced \( r_{jl}^2 / m \) by \( q_j r_{jl}^2 / Q \). By regressing \( Z_{A_j}Z_{B_j} \) on \( u'_j \), we obtain an estimate of \( h_{AB}^2 \), which we then divide by estimates of \( \sqrt{h_{SNP}^2} \) for each trait to produce an estimate of their genetic correlation.

Regions of extreme LD and large-effect SNPs. Estimates of SNP heritability can be unduly affected by regions of extreme LD and by SNPs with disproportionately large effect size.\(^4,5\) Therefore, for all analyses, we exclude SNPs within the major histocompatibility complex (Chromosome 6: 25.34 Mb), as well as SNPs which individually explain >1% of phenotypic variation \( (S_j > n_j / 99) \), and SNPs in LD with these (within 1 cM and \( r_{jl}^2 > 0.1 \)); the latter resulted in the exclusion of SNPs for 8 of the 25 raw traits and for 6 of the 24 summary traits (Supplementary Table 15).

Binary phenotypes. So far, we have implicitly assumed the trait is quantitative. If instead it is binary (i.e., for case-control GW AS), there are two considerations. Firstly, heritability estimates now correspond to the “observed scale;” SumHer will convert them to the “liability scale” if provided with the prevalence and ascertainment.\(^51,52\) Secondly, it is likely the \( p \)-values came from (classical) logistic regression instead of linear regression; however, Supplementary Figure 10 shows that, because linear and logistic \( p \)-values closely match for SNPs with small or moderate effect, this tends to have limited impact.

Polygenic risk scoring. To predict phenotypes, we construct PRS of the form \( \sum_j \beta_j X'_j \), where the vector \( X'_j \) contains standardized genotypes for SNP \( j \) (obtained by centering \( X_j \), then scaling to have variance 1). Without loss of generality, we assume phenotypes have also been standardized, in which case the estimate of \( \beta_j \) from classical linear regression is \( \rho_j \), the correlation observed between SNP \( j \) and the phenotype, and has variance \( (1 - \rho_j^2) / n \). Note that given summary statistics, we can recover \( \rho_j \) by appreciating that the Wald test \( Z \)-statistic is \( \rho_j / \sqrt{(1 - \rho_j^2) / n} \). For the Classical PRS, our estimate of \( \beta_j \) is \( \rho_j \). For the Bayesian PRS, we must specify a prior distribution for \( \beta_j \). Given a heritability model, we use \( \beta_j \sim \mathcal{N}(0, \sigma_j^2) \), where \( \sigma_j^2 = (q_j + \sum_{l \in N_j} q_l r_{jl}^2) h_{SNP}^2 / Q \); this prior is motivated by recognizing that for the “true PRS”, \( \beta_j = \nu_j \), the heritability tagged by SNP \( j \). We then estimate \( \beta_j \) by its posterior mean, a shrunken version of the classical estimate. To calculate this, we approximate the likelihood distribution by \( \mathcal{N}(\rho_j, (1 - \rho_j^2) / n) \),\(^53\) then the posterior mean equals \( \tilde{\rho}_j \sigma_j^2 / (\sigma_j^2 + (1 - \rho_j^2) / n_j) \). Two technicalities. Firstly, to construct each Bayesian PRS requires a value for \( h_{SNP}^2 \), so we used the corresponding estimate from LDSC-Zero or SumHer-Zero (for Enriched GCTA we used the 53-part model, for Enriched LDAK we used the 25-part model). Supplementary Table 14 confirms that the ranking of PRS remains the same if we instead agnostically set \( h_{SNP}^2 = 0.5 \). Secondly, for the Bayesian PRS, we performed clumping\(^54\) (thinned SNPs to ensure no pair within 1 cM had \( r_{jl}^2 > 0.05 \), whereas for the Classical PRS we did not; this was because we found clumping benefited all Bayesian PRS, but was detrimental to the Classical PRS. Supplementary Table 14 shows that the ranking of Bayesian PRS remains the same if we do not clump, but then they are inferior to the Classical PRS.

Choosing a heritability model. Although the heritability model is defined in terms of \( h_j^2 \), as is clear from Equation (3), its fit depends only on how well it models \( \nu_j^2 = h_j^2 + \sum_{l \in N_j} r_{jl}^2 h_l^2 \). Therefore, when specifying \( q_j \), the focus should be on accurately describing how \( \nu_j^2 \) is expected to vary across the genome. LDSC assumes \( E[h_j^2] \) is constant \( (q_j = 1) \), and therefore that \( \nu_j^2 \) correlates with local levels of LD; we refer to this as the GCTA Model\(^5\) because the same assumption is made by GCTA, software for estimating SNP heritability via REML.\(^55\) The GCTA Model is widely used in statistical genetics. For example, it is implicitly assumed by any regression method where SNPs are standardized and the same penalty function / prior distribution is applied to each, and likewise by any simulation study where causal SNPs are picked at random and their standardized effect sizes are sampled from a common distribution. However, as we have shown, the GCTA Model poorly reflects real data.\(^5\) Even for traits where a correlation is observed between \( \nu_j^2 \) and \( S_j \) (those with \( p < 1 \) in Figures 3b & 3c), the correlation is substantially weaker than predicted by the GCTA Model (hence \( p >> 0 \)), and as we discussed above, might be a consequence of including lower-certainty SNPs in the GWAS. We currently recommend using the LDAK.
Model. Originally, this took the form \( q_j = w_j \), where the weights \( w_j \) are calculated based on the assumption that \( \mathbb{E}[v_j^2] \) is constant. We recently updated it to \( q_j = w_j [f_j (1 - f_j)]^{0.75} r_j \), reflecting that \( v_j^2 \) tends to depend on MAF, \( f_j \), and genotype certainty, \( r_j \) (the latter is not relevant in this study as either \( r_j \approx 1 \) or was unknown), and we recognize that further improvements will be possible.

Quality control and association analysis. To prepare the 13 WTCCC GWAS, we used our previously described protocol. In summary, after excluding apparent population outliers, samples with extreme missingness or heterozygosity and SNPs with MAF < 0.01, call rate < 0.95 or Hardy-Weinberg \( P < 1 \times 10^{-6} \), we phased using SHAPEIT, then imputed using IMPUTE2 with the 1000 Genomes Project Phase 3 (2014) reference panel. When merging case and control datasets, we converted genotype probabilities to hard calls using a certainty threshold of 0.95, then retained only autosomal SNPs that in all cohorts had MAF \( \geq 0.01 \) and info score \( \geq 0.99 \) (using IMPUTE2 \( r_2 \_t y p e 2 \) for directly genotyped SNPs). Finally, we thinned individuals, so no pair remained with estimated relatedness \( > 0.05 \). We performed the same quality control for the our reference panel, the Health and Retirement Study. The emerge data were provided post-imputation; this was also performed using SHAPEIT and IMPUTE2, but used the 1000 Genomes Project Phase 2 reference panel. We converted genotype probabilities to hard calls using a certainty threshold of 0.95, then retained only biallelic SNPs with MAF \( \geq 0.01 \), call rate \( \geq 0.95 \), info score \( \geq 0.99 \) and whose genomic position matched that in the 1000 Genomes Project. Finally, we excluded individuals ancestrally inconsistent \( (P < 0.05) \) with non-Finnish Europeans from the 1000 Genomes Project (see Supplementary Figure 11) and those whose ethnicity was reported as “Hispanic or Latino”, then filtered until no pair remained with estimated relatedness \( > 0.05 \) (which left 25,875 individuals).

For the 25 raw GWAS, we performed the association analysis using linear regression (regardless of whether the trait was quantitative or binary), including as covariates sex and ten principal components (five derived from the reference panel, five from the 1000 Genomes Project). For the 24 summary GWAS, we used publicly-available summary statistics (Supplementary Table 8). For each SNP, SumHer requires the two alleles, the \( \chi^2(1) \) test statistic, \( n_j \) and, when estimating genetic correlations, the direction of effect (relative to the first allele). Given summary statistics, we generally used all SNPs in common with our reference panel (and with consistent alleles), however, for the five GWAS which provided info scores, we also excluded SNPs with score \( < 0.95 \). Per-SNP sample sizes were only available for eight of the summary GWAS, so for the remainder, we set \( n_j = n \), the total sample size.

Run times. Given summary statistics and a reference panel, a SumHer analysis has two steps: the first generates a “tagfile”, which contains \( q_j + \sum_{l \in N_j} q_l v_{jl}^2 \) for each SNP; the second performs the regression. The latter is trivial, and typically finishes within minutes. The time to compute the tagfile depends mainly on the size of the reference panel; using our preferred reference panel (8,850 individuals) takes \(~1 \text{ day on a single CPU, whereas using the non-Finnish Europeans from the 1000 Genomes Project}^{14} \) (404 individuals) takes a couple of hours. While we recommend users compute tagfiles from scratch, starting with the subset of SNPs common to both the GWAS and reference panel, we alternatively provide a pre-computed tagfile, which should suffice when the GWAS coverage is high.