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The mystery of missing heritability: Genetic interactions create phantom heritability

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Contributed by Eric S. Lander, December 5, 2011 (sent for review October 9, 2011)

Human genetics has been haunted by the mystery of “missing heritability” of common traits. Although studies have discovered >1,200 variants associated with common diseases and traits, these variants typically appear to explain only a minority of the heritability. The proportion of heritability explained by a set of variants is the ratio of (i) the heritability due to these variants (numerator), estimated directly from their observed effects, to (ii) the total heritability (denominator), inferred indirectly from population data. The prevailing view has been that the explanation for missing heritability lies in the numerator—that is, as-yet undiscovered variants. While many variants surely remain to be found, we show here that a substantial proportion of missing heritability could arise from overestimation of the denominator, creating “phantom heritability.” Specifically, (i) estimates of total heritability implicitly assume the trait involves no genetic interactions (epistasis) among loci; (ii) this assumption is not justified, because models with interactions are also consistent with observable data; and (iii) under such models, the total heritability may be much smaller and thus the proportion of heritability explained much larger. For example, 80% of the currently missing heritability for Crohn’s disease could be due to genetic interactions, if the disease involves interaction among three pathways. In short, missing heritability need not directly correspond to missing variants, because current estimates of total heritability may be significantly inflated by genetic interactions. Finally, we describe a method for estimating heritability from isolated populations that is not inflated by genetic interactions.

genome-wide association studies | statistical genetics

A continuing mystery in human genetics is the so-called missing heritability of common traits. Genome-wide association studies (GWAS) have led to the identification of >1,200 loci harboring genetic variants associated with >165 common human diseases and traits, revealing previously unknown roles for scores of biological pathways (1–3). However, early GWAS were puzzling because they appeared to explain only a small proportion of the “heritability” of the traits. With larger GWAS, the proportion of heritability apparently explained has grown (to 20–30% in some well-studied cases and >50% in a few), but, for most traits, the majority of the heritability remains unexplained (1).

This is our first in a series of papers exploring the explanations for missing heritability. Geneticists define the proportion of (narrow-sense) heritability of a trait explained by a set of known genetic variants to be the ratio explained = h² known/h² all, where (i) the numerator h² known is the proportion of the phenotypic variance explained by the additive effects of known variants and (ii) the denominator h² all is the proportion of the phenotypic variance attributable to the additive effects of all variants, including those not yet discovered. The numerator can be calculated directly from the measured effects of the variants, but the denominator must be inferred indirectly from population data.

The prevailing view among human geneticists has been that the explanation for missing heritability lies in the numerator, that is, in as-yet undiscovered variants. While many variants surely remain to be found, we show here that a substantial proportion of missing heritability could arise from overestimation of the denominator, creating “phantom heritability.” Specifically, (i) estimates of total heritability implicitly assume the trait involves no genetic interactions (epistasis) among loci; (ii) this assumption is not justified, because models with interactions are also consistent with observable data; and (iii) under such models, the total heritability may be much smaller and thus the proportion of heritability explained much larger. For example, 80% of the currently missing heritability for Crohn’s disease could be due to genetic interactions, if the disease involves interaction among three pathways. In short, missing heritability need not directly correspond to missing variants, because current estimates of total heritability may be significantly inflated by genetic interactions. Finally, we describe a method for estimating heritability from isolated populations that is not inflated by genetic interactions.

quantitative genetics

The potential magnitude of phantom heritability can be illustrated by considering Crohn’s disease, for which GWAS have so far identified 71 risk associated loci (13). Under the usual assumption that the disease arises from a strictly additive genetic architecture, these loci explain only 21.5% of the estimated heritability. However, if Crohn’s disease instead follows an LP(k) model, the phantom heritability is 62.8%, thus genetic interactions could account for 80% of the currently missing heritability.

To avoid being fooled by phantom heritability, one might hope to be able to recognize when traits involve genetic interactions, for example, based on population data (such as phenotypic correlations among close relatives) or genetic data (such as pairwise tests...
of epistasis). We show, however, that this task may be difficult. For the case of Crohn’s disease above, detecting the genetic interactions may require sample sizes in the range of 500,000.

In short, genetic interactions may greatly inflate the apparent heritability without being readily detectable by standard methods. Thus, current estimates of missing heritability are not meaningful, because they ignore genetic interactions.

Finally, we present a method to estimate $h^2_{\text{adj}}$ that is consistent not only for additive traits but for any genetic architecture. The method involves the study of isolated populations. It may provide a path forward for accurately measuring explained and missing heritability.

Extensive mathematical details and extensions are provided in SI Appendix. A Matlab software package used for the mathematical calculations is available at http://www.broadinstitute.org/mpg/hc.

**Results**

**Quantitative and Disease Traits.** Quantitative traits are assumed to depend on genotype $G$ and environment $E$, according to a function $P = \Psi(G,E)$. Here, $G = (g_1, g_2, \ldots, g_n)$ is the diploid genotype at $n$ biallelic variant sites across the genome, $g_i$ is the number of copies (0, 1, or 2) of a designated allele at the $i$th site, and $f_i$ is the frequency of the designated allele. The variant sites are assumed to be in linkage equilibrium. The environment $E$ may involve both a “shared” environment that is shared among pairs of relatives and a “unique” environment that is specific to each individual, which includes stochastic noise.

Disease traits are given by a binary function $\Delta(G,E)$ that is assumed to arise from a liability threshold model. Specifically, there is an underlying (and unobserved) quantitative trait $P = \Psi(G,E)$, called a “liability.” For a specified threshold $\tau$, individuals are affected ($\Delta = 1$) if $\Psi(G,E) \leq \tau$ and unaffected ($\Delta = 0$) otherwise. (This condition is often equivalently defined as $\Delta = 1$ if $\Psi(G,E) > \tau$.)

For convenience, we assume throughout that $P$ has been normalized to have mean 0 and variance 1. With $\text{Var}(P) = 1$, the amount of variance explained by a factor is equal to the proportion of variance explained.

**Broad-Sense vs. Narrow-Sense Heritability.** Heritability is measured in two ways: broad-sense heritability $H^2$ and narrow-sense heritability $h^2$.

Broad-sense heritability $H^2$ measures the full contribution of genes. It is defined as $H^2 = V_G / \text{Var}(P)$, where $V_G$ is the total variance due to genes. Specifically, $V_G = \text{Var}(P) - \text{Var}(P|G)$, where $\text{Var}(P|G)$ is the phenotypic variance between genetically identical individuals. $H^2$ is the relevant quantity for clinical risk assessment, because it measures our ultimate ability to predict phenotype from genotype.

By contrast, narrow-sense (or additive) heritability $h^2$ is meant to capture the “additive” contribution of genes to the trait: It is the maximum variance that can be explained by a linear combination of the allele counts, $g_i$. Although $h^2$ is a less intuitive concept, it is routinely used to measure progress toward explaining the genetic basis of a trait because one can readily calculate the contribution of individual loci to $h^2$, as described below.

**Explained, Missing, and Phantom Heritability.** We next define “explained” and “missing” heritability, focusing on narrow-sense heritability $h^2$. Let $h^2_{\text{known}}$ (or $h^2_{\text{known}}$) denote the proportion of the phenotypic variance explained by a set $S$ of known variants, and $h^2_{\text{adj}}$ (or $h^2_{\text{adj}}$) denote the proportion of the phenotypic variance explained by all variants that affect the trait. For the variants in $S$, the proportion of “explained heritability” is $\pi_{\text{explained}} = h^2_{\text{known}} / h^2_{\text{adj}}$ and “missing heritability” is $\pi_{\text{missing}} = 1 - \pi_{\text{explained}}$. When all trait-associated variants have been found, $\pi_{\text{missing}} = 0$.

Human geneticists typically use a “bottom-up” approach to estimate the numerator and a “top-down” approach to estimate the denominator.

**Bottom-up.** The numerator $h^2_{\text{known}}$ is straightforward to estimate, based on the effects of the individual variants. The variance explained by the $i$th variant is $V_i = 2f_i(1 - f_i)\beta^2$, where $f_i$ is the frequency and $\beta$ is the additive effect of the locus (defined as the regression coefficient of the phenotype $P$ on the single-locus genotype $g_i$). Under linkage equilibrium, the variance explained by a set $S$ of variants is the sum over the individual loci: $V_{\text{known}} = \sum_{i \in S} V_i$. Because $\text{Var}(P) = 1$, we have $h^2_{\text{known}} = V_{\text{known}}$ and $h^2_{\text{adj}} = \sum_{i \in S} V_i$. We can thus estimate $h^2_{\text{known}} = \sum 2f_i(1 - f_i)\beta^2$ from the allele frequencies and effect sizes estimated in a genome-wide association study.

**Top-down.** The problem comes in estimating the denominator $h^2_{\text{adj}}$. Because not all variants are known, human geneticists must infer their total contributions indirectly, typically via a top-down quantity based on phenotypic correlations in a population. We refer to such quantities as “apparent heritability” and denote them by such symbols as $h^2_{\text{pop}}$.

Missing heritability is then estimated by assuming that $h^2_{\text{adj}} = h^2_{\text{pop}}$ and obtaining an estimate of $h^2_{\text{pop}}$. The problem is that $h^2_{\text{adj}}$ and $h^2_{\text{pop}}$ are not guaranteed to be equal unless the trait is strictly additive, that is, involves neither gene–gene (G-G) nor gene–environment (G-E) interactions. For traits with genetic interactions, $h^2_{\text{pop}}$ may significantly exceed $h^2_{\text{adj}}$. If so, even when all variants have been discovered, the estimate of $\pi_{\text{missing}}$ will not converge to zero with increasing sample size.

The term heritability and the symbol $h^2$ are often used in the literature to refer to the true heritability $H^2$ and to several definitions of apparent heritability $h^2_{\text{pop}}$, despite the fact that these various quantities need not be equal (SI Appendix, section 1.5). We have introduced distinct terminology and notation to avoid confusion about these important differences.

**Assuming Additivity.** We next describe the typical framework for analyzing human traits, noting why the equality $h^2_{\text{adj}} = h^2_{\text{pop}}$ depends on the assumption of additivity. We focus on one measure of apparent heritability, $h^2_{\text{pop}}(ACE)$, which considers additive genetic, common environmental and unique environmental variance components, but discuss alternative measures in SI Appendix, section 1.3.

**Quantitative traits.** A commonly used definition for apparent heritability is $h^2_{\text{pop}}(ACE) = 2r_{MZ} - r_{DZ}$, where $r_{MZ}$ and $r_{DZ}$ are the phenotypic correlations between monozygotic twins and dizygotic twins, respectively (14). (The measure is based on the ACE model of twin studies.) One can show that

$$h^2_{\text{pop}}(ACE) = h^2_{\text{adj}} + W,$$

with

$$W = \sum_{(i,j) \neq (1,0)} \frac{1}{2} \left( 1 - 2^{-(i+j/2)} \right) V_{AD} \geq 0,$$

where $V_{AD}$ denotes the (nonnegative) variances due to all possible $i$th-order additive interactions and $j$th-order dominance interactions among loci (SI Appendix, section 1). The key point is that, if there are any genetic interactions, then $W > 0$, and $h^2_{\text{pop}}(ACE)$ overestimates $h^2_{\text{adj}}$. Unfortunately, there has been no way to estimate $W$ from population data. In most human genetic studies, the “solution” has been simply to make the (usually unstated) assumption that there is no genetic interaction, that is, that $W = 0$. Typically, the studies assume a strictly additive model. (Some studies allow dominance terms at each locus, but they invariably assume additivity across loci, that is, no genetic interactions.)
Assuming a strictly additive model, the genetic architecture takes the form
\[ \Psi(G, E) = \left( a + \sum \beta g_i \right) + \epsilon, \]
with the two terms each being roughly normally distributed with mean 0 and with \( \Psi \) being normalized to have variance 1. Under this model, the variance of the first term is the narrow-sense heritability \( h^2_{\text{add}} \). The environmental noise \( \epsilon \) consists of shared and nonshared environments, with a vector \( \epsilon_R \) denoting the proportion of the environmental variance \( \text{Var}(\epsilon) = 1 - h^2_{\text{add}} \) that is shared between relatives of type \( R \). (For example, \( c_{\text{sib}} \) is the proportion of environment shared among sibs.) We will refer to this additive model as \( A(h^2, c_R) \).

The additive model has many elegant properties. If \( \rho(R) \) denotes the phenotypic correlation between relatives of type \( R \), then
\[ \rho(R) = h^2 \gamma_R + (1 - h^2) c_R. \]
Here, \( \gamma_R \) is the genetic relatedness between relatives of type \( R \) \( (\gamma_R = 1, 1/2, 1/4, \text{ and } 1/8 \text{ for MZ twins, sibs, grandparent–grandchild, and first cousins)} \). The phenotypic correlation is proportional to genetic relatedness under the additive model with no shared environmental variance.

**Disease traits.** Disease traits are traditionally assumed to follow a liability threshold model, where the unseen liability \( \Psi \) follows the additive model above and disease occurs if \( \Psi \leq \tau \). We will refer to this additive disease model as \( A_{\lambda}(h^2, c_R, \mu) \), where \( \mu = \text{Prob}(\Psi \leq \tau) \) denotes the disease prevalence. The model parameters \( (h^2, c_R, \mu) \) completely determine the epidemiologically observable quantities \( (\mu, \lambda_{\text{MZ}}, \lambda_{\text{sib}}, \ldots) \), where \( \mu \) is the disease prevalence and \( \lambda_R \) is the increased risk to relatives of type \( R \) (15). To apply the model to a disease, one fits the model parameters based on observable quantities. (Geneticists often assume that \( c_R = 0 \), in which case the remaining two parameters can be fit based on \( \mu \) and \( \lambda_{\text{MZ}} \).) For genetic variants associated with the disease, one then uses the model to convert an observed increase in disease risk to an inferred additive effect on the liability scale. Heritability calculations are performed not on the observed disease status but on the unseen liability scale. One advantage of using the liability scale is that heritability calculations tend to be robust to uncertainty about disease prevalence. (See SI Appendix, section 2 for details, including the use of both \( \lambda_{\text{MZ}} \) and \( \lambda_{\text{sib}} \) to deal with shared environment.)

**Genetic Interactions Create Phantom Heritability.** What will happen if a geneticist analyzes a trait that involves genetic interactions under the erroneous assumption that it is additive? To explore this question, we introduce a simple and biologically plausible class of models.

**Quantitative traits.** Biological processes often depend on the rate-limiting value among multiple inputs, such as the levels of components of a molecular complex required in stoichiometric ratios, reactants required in a biochemical pathway, or proteins required for transcription of a gene. We thus define a limiting pathway model, in which a trait \( P \) depends on the rate-limiting input from \( k \geq 1 \) biological processes. For simplicity, we will assume that the inputs, \( \Psi_1, \Psi_2, \ldots, \Psi_k \), all follow the standard additive model in Eq. 2 above, each with exactly the same parameters, \( h^2_{\text{pathway}}, \text{ and } c_R \). Apart from the fact that the \( \Psi_i \) are roughly normal, we place no restrictions on the number or allele frequencies of the causal variants.

We define the trait \( LP(k, h^2_{\text{pathway}}, c_R) \) to be the minimum value of the \( \Psi_i \). For a single pathway \( (k = 1) \), the definition reduces to the simple additive model. What happens for \( k > 1 \)? Let us consider a specific example: \( P^* = LP(4, 50\%, c_R) \), with \( c_{\text{sib}} = 50\% \) (yielding shared environmental variance \( V_{\epsilon} = 27\% \)) and \( c_R = 0 \) for other relatives. Suppose that a geneticist analyzes \( P^* \) under the standard (but erroneous) assumption that it is additive. Because we know the true genetic architecture (although the geneticist does not), we can calculate the exact value of all relevant parameters (SI Appendix, section 3). Because we are interested in asymptotic bias, we ignore sampling variation. The geneticist would start by estimating the apparent heritability to be explained. The observed phenotypic correlation among twins is \( r_{\text{MZ}} = 62.4\% \) and \( 35.4\% \), yielding \( h^2_{\text{pop}} = 2(r_{\text{MZ}} - r_{\text{DZ}}) = 54.0\% \). The geneticist would then conduct a genetic study, identify variants associated with the trait, estimate their effect sizes, estimate the heritability \( h^2_{\text{known}} \) explained by the variants, and compare it to the estimated value of \( h^2_{\text{pop}} \). Assuming the sample is so large that all variants are identified (although the geneticist does not know this), \( h^2_{\text{known}} \) will be the true heritability, \( h^2_{\text{all}} = 25.4\% \).

![Fig. 1. Phantom heritability under the limiting pathway model. (A) Quantitative trait model \( LP(k, h^2_{\text{pathway}}, c_R) \). For various parameters, curves show apparent heritability \( h^2_{\text{pop}}(ACE) \) and phantom heritability \( h^2_{\text{phantom}} \). Curves connect points with various values of \( k \) (1, 2, 3, 4, 5, 6, 7, 10, with tip of arrow at \( k = 10 \)), and for specific values of \( h^2_{\text{pathway}}(10, 30, 50, 70, \text{ and } 90\%) \), indicated by color of the curve and \( c_R \) (0% filled circles and arrows, 50% open boxes and arrows). The red asterisk indicates example \( P^* \) referred to in the text. (The quantity \( \text{h}^2_{\text{pop}} \) can exceed 100%, as seen for some models with high heritability and for some real traits. In such cases, we set \( \text{h}^2_{\text{pop}} = 100\% \).) Raw data are in SI Appendix, Table 6. (B) Disease model \( LP_2(k, h^2_{\text{pathway}}, 0\%) \). For various parameters, curves show value of \( \lambda_{\text{MZ}} \) and phantom heritability \( h^2_{\text{phantom}} \). Values of \( k \) and \( h^2_{\text{pathway}} \) are as in A. Values of prevalence \( \mu \) are 0.1% (solid), 1% (dotted), and 10% (dashed). The red asterisk indicates example \( \Delta^* \) referred to in the text. In both A and B, as \( k \) increases, the trait becomes more nonlinear; phantom heritability increases to 50% and beyond. Raw data are in SI Appendix, Table 7.]
Even though all variants have been discovered, they will appear to explain only 47% (=25.4/54.0) of the apparent heritability, $h_{pop}^2$. The remaining 53% is phantom heritability, which will never be explained by additional variants. It is the result of analyzing the data under an erroneous model.

Similar results are obtained for a wide range of parameters. Fig. 1A shows results for $k = 1-10$, $h_{langle q\rangle}^2 = 10-90\%$, and $c_{RE} = 0$ or $50\%$. The phantom heritability grows steadily with $k$. A mathematical theorem (16) implies that $\pi_{phantom} \rightarrow 100\%$ as $k$ grows (SI Appendix, section 3.4).

**Disease traits.** We can similarly define a limiting pathway model for disease traits by applying a threshold to the LP model for quantitative traits. Specifically, we define the disease trait $LP_d(k, h_{langle q\rangle}^2, c, \mu)$ as occurring if and only if $LP(k, h_{langle q\rangle}^2, c, \mu) \leq \tau$, with $\mu$ denoting the disease prevalence. The case $k = 1$ again reduces to the additive model. What happens for $k > 1$?

Again, let us consider a specific case: $\Delta^* = LP_d(3, 50\%, c_{RE}, 1\%)$, with $c_{RE} = 0\%$ for all relatives. Based on the observed relative risks to MZ and DZ twins, a geneticist would calculate that $h_{pop}^2 = 49.0\%$. However, an infinitely large genetic mapping study would yield $h_{known}^2 (=h_{all}^2) = 21.2\%$. Even though all variants had been identified, it would appear to explain only $43.2\% = (21.2/49.0)$ of the apparent heritability $h_{pop}^2$. The remaining $56.8\%$ is phantom heritability. Similar results are obtained for a wide range of parameters. Fig. 1B shows results for $k = 1-10$, $h_{langle q\rangle}^2 = 10-90\%$, and $c = 0.1, 0.1, 0.1, 0.3$.

**Epistasis is Common.** The results show that mistakenly assuming that a trait is additive can seriously distort inferences about missing heritability. From a biological standpoint, there is no a priori reason to expect that traits should be additive. Biology is filled with nonlinearity: The saturation of enzymes with substrate concentration and receptors with ligand concentration yields sigmoid response curves; cooperative binding of proteins gives rise to sharp transitions; the outputs of pathways are constrained by rate-limiting inputs; and genetic networks exhibit bistable states.

Genetic studies in model organisms have long identified specific instances of interacting genes (17). Important examples include synthetic traits (e.g., 18), which occur only when multiple loci or pathways are all disrupted. With the advent of genome-wide mapping in controlled genetic backgrounds in model organisms, studies have been able to reveal that epistasis is pervasive. In the yeast **Saccharomyces cerevisiae**, Brem et al. (19) analyzed as quantitative traits the levels of gene transcripts in segregants of a cross between two strains. For each transcript, they found the strongest quantitative trait locus (QTL) in the cross and then, conditional on the genotype at this locus, identified the strongest remaining QTL. In 67% of cases, these two QTLs demonstrated epistatic interactions. In bacteria, Khan et al. (20) and Chou et al. (21) have recently demonstrated clear epistasis among collections of five mutations that increase growth rate. In mouse and rat, Shao et al. (22) analyzed a panel of chromosome substitution strains, with each strain carrying a different chromosome from a donor strain on a common recipient genetic background. For dozens of quantitative traits, the sum of the effect attributable to the individual donor chromosomes far exceeds (median eightfold) the total effect of the donor genome, indicating strong epistasis. Although genetic interactions are hard to detect in humans (see below), several cases involving variants with large marginal effects have been recently reported in Hirschsprung’s disease, amyloidosis, spondylitis, psoriasis, and type I diabetes (SI Appendix, section 7.1).

Several arguments are sometimes offered in support of the assumption of additivity (e.g., linearity of responses to selection). We discuss the flaws in such reasoning (SI Appendix, section 11).

**Can We Detect Genetic Interactions by Comparisons Across Relatives?** Can a geneticist avoid being fooled by phantom heritability by detecting a priori that a trait involves genetic interactions, based on the phenotypic correlations between close relatives? The task turns out to be difficult even if we restrict attention only to LP models.

**Phenotypic distribution.** The phenotypic distribution of a quantitative trait would not reveal the presence of genetic interactions. The distribution for $LP(k)$ traits with modest values of $k$ (say, $k \leq 10$) is reasonably similar to the normal distribution in the additive model (SI Appendix, section 7.1). Moreover, deviations from perfect normality are common in real traits and are typically resolved by applying a transformation to the distribution.

**Sib correlations.** Phenotypic correlations among sibs would not reveal that a trait involves genetic interactions. For quantitative traits, the correlations $(r_{MZ}, r_{DZ})$ for the LP models above are similar to those seen for real traits: They fit comfortably within the range of values recently reported by Hill et al. (23) for 86 traits (SI Appendix, section 5.1). For disease traits, the relative risks $(\lambda_{MZ}, \lambda_{DZ})$ for various LP models similarly resemble those seen for real traits, for example, those reported for 15 actual diseases by Wray et al. (24) (SI Appendix, section 5.2).

**Correlations among extended relatives.** That sib correlations alone do not distinguish between additive and nonadditive LP models is not surprising: For either model, one can select parameters that largely fit the observed correlations. One might expand the analysis by considering additional relatives. For a trait with no shared environment, the phenotypic correlation between relatives should decrease linearly with genetic relatedness ($r_{ER}$) if the trait is additive (by Eq. 3), but should be concave up if the trait involves genetic interactions. In theory, one could test for genetic interactions by fitting different genetic models to the curve of phenotypic correlations among relatives. In practice, it is difficult to draw strong conclusions from such analysis. First, such tests essentially depend on fitting a handful of values (e.g., correlations for individuals with $r_{ER} = 1, 1/2, 1/4, 1/8$) with limited precision. Second, differences in the degree of shared environmental variance between relative types can substantially alter the shape of the curve (SI Appendix, section 6).

**Examples: Crohn’s disease and schizophrenia.** The problem of discerning genetic architecture from a few parameters can be illustrated by considering alternative models for real diseases.

For Crohn’s disease, current GWAS have identified 71 risk loci. Assuming the disease follows an additive model, these known loci explain $h_{known}^2 = 10.8\%$ of the total phenotypic variance, or $\pi_{explained} = 21.5\%$ of the heritability (assuming $h_{all}^2 = h_{pop}^2 = 50\%$). Alternatively, one can define an LP(3) model that is consistent with the prevalence and sib risks. Under this model, the phantom heritability is $\pi_{phantom} = 62.8\%$. Genetic interactions would account for $80\% = (62.8/(1-0.215))$ of the currently missing heritability. The known variants would account for $\pi_{explained} = 57.5\% = (21.5/(1-0.628))$ of the true heritability $h_{all}^2 = 18.6\%$ (SI Appendix, section 6).

For schizophrenia, Risch (15) presented recurrence risks for various relative types ($r_{ER} = 1, 1/2, 1/4, 1/8$) with limited precision. Second, differences in the degree of shared environmental variance between relative types can substantially alter the shape of the curve (SI Appendix, section 6).

**Can we detect genetic interactions from pairwise epistasis?** Even though it is difficult to detect genetic interactions a priori based on population data such as sib correlations, one might still hope to detect epistasis among variants a posteriori once they have been mapped. Indeed, geneticists have tested for pairwise epistasis between loci, but have found few significant signals. Should failure to detect pairwise epistasis allay our concerns about phantom heritability? Unfortunately, the answer is no.

The reason is that individual interaction effects are expected to be much smaller than linear effects, and the sample size required to detect an effect scales inversely with the square of the effect size. If $n$ loci had equivalent effects, the sample size to
detect the n loci would thus scale with \( n^2 \), whereas the sample size to detect their \( \sim n^2 \) interactions scales with \( n^4 \).

Consider the \( LP(3) \) disease model \( \Delta^* \) discussed above, with phantom heritability of 56.8%. Suppose that we consider two variants with frequency 20% that contribute to different pathways and increase risk by 1.3-fold (which is a large effect relative to those typically seen in GWAS). The sample size required to detect the variants is \( \sim 4,900 \) (with 50% power and genome-wide significance level of \( \alpha = 5 \times 10^{-8} \) in a genome-wide association study with an equal number of cases and controls), whereas the sample size required to detect their pairwise interaction is roughly 450,000 (at 50% power and an appropriate significance level to account for multiple hypothesis testing). A researcher who studied 100,000 samples would likely discover all of the loci but would find little evidence of epistatic interactions. The researcher might conclude that the genetic architecture is additive, although the phantom heritability is actually >50%. In short, the failure to detect epistasis does not rule out the presence of genetic interactions sufficient to cause substantial phantom heritability. (We discuss other ways to potentially detect epistasis in SI Appendix, section 7.5.)

**Consistent Top-Down Estimator of \( h^2_{\text{add}} \).** What we need is a top-down estimator \( h^2_{\text{add}} \) that is consistent not simply for additive traits but for any genetic architecture. Traditional approaches fail because they focus on phenotypic correlations between close relatives; this creates two problems: (i) extensive allele sharing between close relatives makes it difficult to disentangle the effects of genetic interactions; and (ii) differences of shared environment between different relative types make it difficult to disentangle the effects of environment.

We can eliminate these problems by studying nearly unrelated individuals in a population. Specifically, one can (i) identify pairs of individuals whose probability of allele sharing at the causal loci differs slightly from the population average, and (ii) measure how their phenotypic similarity depends on their genotypic similarity.

This goal can be accomplished by studying recent genetically isolated populations (such as Iceland, Finland, the Hutterites, or the Amish), in which one can use dense genotyping to reliably detect large segments shared identical-by-descent (IBD) between individuals (SI Appendix, section 8). We have the following theorem.

**Theorem 1.** Consider a population in which one can detect large segments shared IBD between individuals. Given two individuals \( I_1 \) and \( I_2 \), let \( \kappa_{ij} = \kappa(I_1, I_2) \) denote the proportion of their genomes shared in large IBD segments. Given a trait, let \( \rho(\kappa) \) denote the average phenotypic correlation between pairs of individuals who share proportion \( \kappa \) of their genomes in large IBD blocks. Regardless of the genetic architecture of the trait, the true heritability \( h^2_{\text{add}} \) equals \( h^2_{\text{slope}(\kappa)} = (1 - \rho(\kappa)) \rho'(\kappa) \), where \( \rho'(\kappa) \) is the rate of change of phenotypic correlation around the average sharing level of large IBD segments. Accordingly, \( h^2_{\text{slope}(\kappa)} \) provides a consistent top-down estimator for \( h^2_{\text{add}} \).

The theorem applies to both quantitative traits and disease traits [with heritability measured on the disease (0,1) scale] with individuals sampled from the general population. The proof appears in SI Appendix, section 8, along with a version for individuals ascertained in a case–control study.

To apply this result in practice, one would (i) take a collection of individuals from the population; (ii) for each pair of individuals, calculate the product \( Q \) of the phenotype and the degree \( \kappa \) of IBD sharing; and (iii) estimate \( \rho'(\kappa) \), as the regression coefficient of \( Q \) on \( \kappa \), for pairs with \( \kappa \) in a neighborhood around \( \kappa_{ij} \).

Fig. 2 illustrates the approach on simulated data for the quantitative trait \( \rho^* \) above, where \( h^2_{\text{add}} = 25.4\% \) and \( h^2_{\text{pop}} \) = 54%. With simulated data for 1,000 individuals with IBD sharing similar to that seen in Qatar (25), we estimate \( h^2_{\text{slope}(\kappa)} = 25.8 \pm 8.2\% \), which is very close to the correct value of \( h^2_{\text{add}} = 25.4\% \).

It is instructive to compare our approach with two elegant methods recently introduced by Visscher and colleagues, which inspired our own work. Both methods involve regressing phenotypic correlation on genotypic similarity. The first (26) measures genotypic similarity in terms of IBD within sib pairs—essentially measuring \( \rho'(1/2) \), in our terminology. It eliminates the effects of shared environment by studying a single type of relative, but is confounded by genetic interactions because it studies close relatives (SI Appendix, section 10). The second (27) measures genotypic similarity in terms of identity by state across an SNP catalog for pairs of individuals in a population. As the authors note, the approach is not confounded by genetic interactions, but does not yield a consistent estimator because its sensitivity to causal variants falls with allele frequency. Nonetheless, this method yields a valuable lower bound on \( h^2_{\text{add}} \).

**Discussion.** The main points of this paper are that (i) current estimates of the proportion of heritability explained by known variants (\( \pi_{\text{explained}} \)) implicitly assume that traits involve no genetic interactions; (ii) this assumption is not justified, because many models with interactions are equally consistent with available data; and (iii) under some of these models, the true value of \( \pi_{\text{explained}} \) may be much larger than current estimates. Accordingly, the widely held belief that missing heritability directly reflects the variance due to as-yet undiscovered variants is unjustified. Rather, missing heritability may be due in significant part to genetic interactions.

We focus here on a simple and biologically natural model, the limiting pathway model; it cannot readily be distinguished from an additive model based on population data or tests of pairwise
epistasis, yet entails substantial phantom heritability. Our focus on the LP model is not meant to imply that real traits necessarily follow this particular model; it simply provides an evidence proof that erroneous assumptions may give rise to substantial missing heritability. We discuss more general multiple pathway models (SI Appendix, section 4.4), which also show substantial phantom heritability. (Beyond G–G interactions, we note that G–E interactions can produce additional phantom heritability.)

Importantly, we do not mean to propose that missing heritability is entirely, or even primarily, due to genetic interactions. On the contrary, many more causal variants are likely to exist, and to account for a significant part of the missing heritability. Discovery efforts should continue vigorously.

The case of Crohn’s disease illustrates these points. The currently known loci can explain ~22%, ~58%, or more of the true heritability, depending on whether the disease follows an LP(1), LP(3), or other model. The available data cannot distinguish among the models. This spectacular degree of uncertainty underlines “inference by default,” for example, the frequent conclusion that rare variants must largely cause a disease, because common variants explain “too little” of the heritability. [Notably, a recent study of Crohn’s disease (28) reported that the rare variants explained 10- to 20-fold less of the heritability than the common variants at 56 disease-associated loci.]

Given the dependence of results on genetic architecture, authors reporting proportions of heritability explained or missing should state clearly that the calculations are made under the arbitrary assumption that the trait is additive.

In LP models, phantom heritability increases with the number of pathways. More generally, traits with greater biological complexity may have greater phantom heritability. Current studies are broadly consistent with such a notion: The apparent heritability explained for “simpler” traits such as levels of fetal hemoglobin is greater than for “more complex” traits such as body–mass index or age at menarche (SI Appendix, section 6.3). Such differences may reflect both the number of loci and the genetic interactions underlying the traits.

The fraction of the apparent heritability of human traits due to genetic interactions cannot be inferred from available data, although the perversiveness of epistasis in experimental organisms suggests that the true heritability \( h^2 \) of traits may be much lower than current estimates. (Lower values of \( h^2 \) do not mean that traits “are less genetic” in the popular use of the term, which refers to the total contribution of genes, \( H^2 \). It simply means that additive effects comprise a smaller fraction of \( H^2 \).)

We describe a potential solution to overcome the problem of genetic interactions: Theorem 1 provides a top-down method to remove additive heritability that is consistent regardless of the underlying genetic architecture. In principle, the approach can provide an accurate assessment of heritability, as well as allow detection of the presence of genetic interactions by comparing top-down estimates obtained from different methods. To assess its practical utility, it will be necessary to apply it to appropriate data from isolated populations.

Finally, notwithstanding our focus here, we believe that concerns about missing heritability should not distract from the fundamental goals of medical genetics. Human genetic studies to discover variants associated with common traits should primarily be regarded as the analog to mutant hunts in model organisms, with the primary purpose being to identify the underlying pathways and processes. The key focus should be to study the biological role of the variants discovered so far. The proportion of phenotypic variance explained by a variant in the human population is a notoriously poor predictor of the importance of the gene for biology or medicine. [A classic example is the gene encoding HMGCoA reductase, which explains only a tiny fraction of the variance in cholesterol levels but is a powerful target for cholesterol-lowering drugs (1).] Ultimately, the most important goal for biomedical research is not predicting heritability—that is, predicting personalized patient risk—but understanding pathways underlying disease and using that knowledge to develop strategies for therapy and prevention.

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