Supplementary Figures

Supplementary Figure 1 Region-specific LD heterogeneity of the genome. LD score of each variant is defined as the sum of LD r^2 between the target variant and all variants (including the target variant) within ± 10 Mb distance. For better graphic display, the y-axis is truncated at 1,000.

Supplementary Figure 2 LDAK weights vs. MAF for the 233,588 variants on chromosome 22 in the UK10K-WGS data. For better graphical presentation, shown are the mean weights in 100 MAF bins.

Supplementary Figure 3 GREML-LDMS estimate of h_{WGS}^2 using sequence variants and simulated phenotype based on the UK10K-WGS data. Each column represents the mean estimate from 200 simulations. Error bar is the s.e. of the mean estimate. The true h^2 parameter is 0.8 for the simulated traits (see Online Methods for the 4 simulation scenarios). Variants are stratified based on the distribution of the segment-based mean LD score (see Online Methods for details). 4LD: first, second, third and forth quartiles. 3LD: first, (second + third), and forth quartiles. 2LD: (first + second), and (third + forth) quartiles. In each LD group, variants are further stratified by MAF. 7MAF: variants are stratified into 7 MAF groups, i.e. MAF \leq 0.001, 0.001 < MAF \leq 0.01, 0.01 < MAF \leq 0.1, 0.1 < MAF \leq 0.2 , $0.2 <$ MAF \leq 0.3, $0.3 <$ MAF \leq 0.4 and 0.4 $<$ MAF \leq 0.5. 2MAF: variants are stratified into 2 MAF groups, i.e. MAF \leq 0.01 and 0.01 \leq MAF \leq 0.5.

(a) Common variants (MAF \geq 0.01)

(b) Rare variants $(MAF < 0.01)$

Supplementary Figure 4 Proportion of variation at sequence variants captured by 1KGP imputation in the UK10K-WGS data. Details of the 4 simulation scenarios can be found in Online Methods. The estimates are from GREML-LDMS analysis (4 LD groups with each LD group being further stratified into 7 MAF groups, 28 groups in total) using all the 1KGP-imputed variants after QC (without filtering variants for IMPUTE-INFO). The proportion of variation at sequence variants captured by imputation (i.e. multi-variant tagging) is defined as the estimate of phenotypic variance explained by 1KGPimputed variants summed over all the relevant groups (i.e. 20 groups for common variants and 8 groups for rare variants in the GREML-LDMS analysis) divided by true simulation parameter (i.e. variance explained by causal variants). Plotted value is the mean estimate from 200 simulations. Error bar is the s.e. of the mean estimate. Panel (a): common variants. Panel (b): rare variants. The multi-variant tagging averaged across SNP arrays and simulation scenarios is ~ 0.97 for common variants and ~ 0.68 for rare variants.

(c) Number of variants

Supplementary Figure 5 Estimate of variance explained by 1KGP-imputed variants stratified by MAF and LD for height and BMI. The estimates of variance explained are from the GREML-LDMS analyses of fitting all the 28 genetic components simultaneously in the combined data from 7 GWAS cohorts $(44,126$ unrelated individuals and 17M variants). Segmental LD score increases from the 1st to 4th quartiles (See Online Methods for the LD stratified approach). Error bar represents the standard error. For better display, negative estimates are not shown on the figure, which are available in **Supplementary Table 3**. Panel (c) shows the number of variants in each of the 28 groups.

(a) Correlation of GRM (b) Correlation of the estimate of genetic variance

Supplementary Figure 6 Correlation of GRM (or estimate of variance component) between each pair of the variant groups for the GREML-LDMS analysis in the combined GWAS data set. There are ~17M 1KGP-imputed variants stratified by MAF and segment-based LD score into 28 groups (Online Methods). MAF1: MAF ≤ 0.001; MAF2: 0.001 < MAF ≤ 0.01, MAF3: 0.01 < MAF ≤ 0.1; MAF4: 0.1 $<$ MAF \le 0.2; MAF5: 0.2 $<$ MAF \le 0.3; MAF6: 0.3 $<$ MAF \le 0.4; MAF7: 0.4 $<$ MAF \le 0.5. LD1, LD2, LD3 and LD4 are the four quartiles of the segment based LD score distribution (from low to high), respectively. Panel (a) shows the correlation of the off-diagonal elements of GRM between each pair of groups. Panels (b) shows the correlation of the estimate of genetic variance $(\hat{\sigma}^2_\nu)$ between each pair of groups for height. The correlation of $\hat{\sigma}_{v}^{2}$ is calculated as $cov(\hat{\sigma}_{v(i)}^{2}, \hat{\sigma}_{v(j)}^{2}) / \sqrt{var(\hat{\sigma}_{v(i)}^{2})var(\hat{\sigma}_{v(j)}^{2})}$. The GRMs of different groups are positively correlated, in particular for those in the same LD group (e.g. a maximum correlation of 0.471 between groups MAF6_LD4 and MAF7_LD4), which leads to negative correlation of $\hat{\sigma}_{\nu}^2$. The correlation of $\hat{\sigma}_{\nu}^2$ for BMI is almost identical to that for height and is therefore not shown.

Supplementary Figure 7 Estimate of variance explained by 1KGP-imputed variants stratified by MAF for height and BMI. The estimates of variance explained are from the GREML-MS analyses of fitting all the 7 genetic components simultaneously in the combined data set (44,126 unrelated individuals and 17M variants). Error bar is the standard error. This figure shows that rare variants (MAF \leq 0.01) do explain a significant amount of variance for both height (variants with $0.001 \leq \text{MAF} \leq 0.01$) and BMI (variants with MAF ≤ 0.001), consistent with the results from GREML-LDMS (**Supplementary Table 3 and Supplementary Fig. 5**).

Supplementary Figure 8 Variance explained per variant. The estimate of variance explained per variant is calculated as the $\hat{h}^2_{KGP(i)}/m_i$ where $\hat{h}^2_{KGP(i)}$ is the estimate of variance explained by the variants in *i*-th MAF group (shown on x-axis) from the GREML-LDMS analysis in the combined set data, and m_i is the number of variants in the *i*-th MAF group.

(a) GREML-LDMS estimate of h_{IKGP}^2 (height)

 $^{2}_{\text{KGP}}$ (height) (b) GREML-LDMS estimate of h_{KGP}^{2} (BMI)

Supplementary Figure 9 Variance explained by the first PCs for height and BMI and the GREML-LDMS estimates of h_{IKGP}^2 with different number of PCs estimated from different sets of variants. Shown are the results from the analyses in the combined data set. Shown in panels (a) and (b) are the results from the GREML-LDMS analyses fitting different numbers of PCs as fixed covariates for (a) height and (b) BMI. 10 PCs (HapMap3): GREML-LDMS analysis fitting the first 10 PCs estimated from the common variants on HapMap3. 20 PCs (All): GREML-LDMS analysis fitting the first 20 PCs estimated from all the variants. In panels (c) and (d), plotted are cumulative variance explained by the first *x* PCs, which are the adjusted multiple regression R^2 of height (or BMI) phenotype on the first *x* PCs calculated from using (c) all common variants on HapMap3 or (d) all the variants.

Supplementary Figure 10 Evidence that height- and BMI-associated genetic variants being under natural selection is not driven by population stratification. We re-ran the analyses as presented in **Fig. 4c** and **4d** using slightly different methods. In panel (b), we performed mixed linear model based association analyses using BOLT-LMM¹, association tests of ~17M 1KGP-imputed variants (each fitted as a fixed effect) with ~1.2M common variants on the HapMap3 fitted as random effects to control for population stratification. In panel (b), we calculated the loading of each SNP along first PC (PC computed from all ~17M 1KGP-imputed variants) following the method described in Galinsky et al.², and adjusted the minor allele effect (b_m) by SNP loading using linear regression. The results are almost identical as those shown in **Fig. 4c** and **4d**.

Supplementary Figure 11 Standard error (s.e.) of the estimate from GREML-LDMS is approximately inversely proportional to sample size (*n*). We randomly sampled the 10,000 to 40,000 samples by steps of 5,000 from the combined GWAS data set (*n* = 44,126 unrelated individuals), and repeated the GREML-LDMS analyses for height and BMI using 1KGP-imputed variants as described in Online Methods. The result shows that the s.e. of GREML-LDMS estimate of $h²_g$ (variance explained by all sequence or imputed sequence variants) is approximately 927 / *n*.

Supplementary Figure 12 Estimates of variance explained by 1KGP-imputed variants stratified by MAF. The results are from the GREML-LDMS analyses of fitting 28 genetic components in the combined GWAS data set of 44,126 unrelated individuals for (a) height and (b) BMI. Shown is the sum of estimates in each of the 5 MAF groups. Error bar is the standard error. Assumption A: assuming a normal distribution of the effect sizes corresponding the standardised genotype variables (default assumption for the GREML methods). Assumption B: assuming a normal distribution of the allelic substitution effects. For height, the log likelihood (logL) of the model is -20189.2 (AIC = 40458.4 and BIC = 40564.0) under Assumption A, and is 20189.2 (AIC = 40458.3 and BIC = 40563.9) under Assumption B. For BMI, the logL is -21234.5 (AIC = 42548.9 and BIC = 42654.4) under Assumption A, and -21235.7 (AIC = 42551.3 and BIC = 42656.8) under Assumption B. These results suggest that the two models fit almost equally well with the data for both height and BMI.

Supplementary Figure 13 Simulations with effect sizes of causal variants sampled from the distribution estimated from real data for height. It is shown in Fig. 4c that mean \hat{b}_m is correlated with log10(mean MAF) across 100 MAF bins for both height and BMI, where each MAF bin is the 1% quartile of MAF distribution, and b_m is defined as the effect size of the minor allele of a variant. Using the same data as in Fig. 4c, we show in panel (a) that mean \hat{u}_m is also correlated log10(mean MAF) for height (*r* = 0.44, *P*_{permu} = 1.3e-5) but not for BMI (*r* = -0.03, *P*_{permu} = 0.76), where $u_m = b_m \sqrt{2p(1-p)}$ with p being the MAF and P_{permu} is calculated from 1 million permutations. The relationship between

 (a) (b)

 u_m and MAF violates the assumption of the GREML methods. We therefore performed simulations (based on the UK10K-WGS data) to test the robustness of GREML-LDMS to such a correlation. The variance of u_m , as shown on the y-axis of panel (b), is calculated as $var(u_m) = var(\hat{u}_m) - SE^2(\hat{u}_m)$. The simulation strategy largely follows that in Online Methods with an important difference that we simulated u_m from a normal distribution with mean from that shown on the y-axis of panel (a) and variance from that of panel (b) for height according to its MAF. For the ease of stimulation, we used the absolute value of var (u_m) if the estimate of var (u_m) is negative. Shown on panel (c) are the mean estimates from 200 simulations. Error bar is the s.e. of the mean estimate.

(c) Proportion of variants with $MAF < 0.1$ (d) Adjusted estimate

Supplementary Figure 14 Adjusting the estimate of h_{KGP}^2 in each LD group by the proportion of low-MAF variants in the group. The estimates are from the GREML-LDMS analyses (28 genetic components) in the combined data set for height and BMI. It is shown in panel (a) that variants in lower LD regions tend to explain a larger proportion of variance, for height in particular. However, there is also an enrichment of low-MAF variants in regions with lower LD as shown in panels (b) and (c). We therefore adjusted \hat{h}^2_{KGP} (and its s.e.) by the proportion of variants with MAF ≤ 0.1 (θ) in each MAF group, using a regression model $y = \alpha + x\beta + e$ with $y = \hat{h}_{1KGP}^2$ (or its s.e.) and $x = \theta - \text{mean}(\theta)$. Shown in panel (d) is the adjusted estimate is calculated as $\hat{h}^2_{KGP} - x\hat{\beta}$ with $\hat{\beta}$ estimated from the regression above.

Supplementary Figure 15 Power comparison between WGS and imputation. Power is measured by the noncentrality parameter (NCP) of a χ^2 test-statistic. Given a fixed budget, NCP(imputation) / NCP(WGS) = r^2n_1/n_2 where n_1 is the sample sizes of a study using SNP genotyping followed imputation, n_2 is the sample size of a WGS study, r^2 is the proportion of variation at a sequence variant tagged by imputed variant(s). Assuming genotyping cost per individual using SNP array is 20 times cheaper than that using WGS, NCP(imputation) / NCP(WGS) = $20r^2$. Error bar is the s.e.m.. Multivariant tagging was quantified as in **Fig. 3**, and single-variant tagging is calibrated as in **Fig. 5**, by imputing variants on Illumina CoreExome array from the UK10K-seq data to 1KGP reference panels.

a) Rare variants

b) Common variants

Supplementary Figure 16 Single-variant tagging of sequence variants by 1KGP-imputed variants based Illumina OmniExpress array vs. that based on Illumina CoreExome array. Shown is the squared correlation $(r²_{\text{max}})$ between a sequence variant from the UK10K-WGS data and the best tagging variant from 1KGP imputation within ±1Mb distance. The 1KGP imputation analyses are based on variants on Illumina OmniExpress and Illumina CoreExome arrays extracted from the UK10K-WGS data (see Online Methods for details about the imputation analyses based on the UK10K-WGS data). There are 10.2% (8.3%) rare variants and 1.1% (0.3%) common variants that are almost not tagged by imputation, i.e. r_{max}^2 < 0.05, based on Illumina CoreExome (OmniExpress) array.

Supplementary Figure 17 MAF distribution of variants in the UK10K-WGS data. After quality controls (Online Methods), there are 17.6M variants in total (9.3M variants with MAF < 0.01).

Supplementary Figure 18 Proportions of variance explained by the simulated causal variants in 7 MAF groups under different simulation scenarios. I) Random: 1,000 causal variants randomly sampled from all the sequence variants; II) More common: 1,000 random and additional 500 common causal variants; III) Rarer: 1,000 random and additional 500 rare causal variants; IV) Rarer & DHS: 1,000 random and additional 500 rare causal variants all sampled from the variants at DHSs. Shown are averages from 200 simulation replicates. The total heritability is 0.8.

Supplementary Figure 19 Principal component analysis (PCA) of ancestry in the combined GWAS data set. The PCA is based on 1KGP-imputed variants that are on HM3, with MAF ≥ 0.01 and imputation $R^2 \ge 0.3$.

Supplementary Tables

Supplementary Table 1 Estimates of heritability using WGS variants under different simulation scenarios based on the UK10K-WGS data.

Random: 1,000 causal variants sampled from the WGS variants at random; **More common:** 1,000 causal variants sampled at random + 500 causal variants with MAF > 0.01; **Rarer:** 1,000 causal variants sampled at random + 500 causal variants with MAF < 0.01; **Rarer & DHSs:** 1,000 causal variants sampled at random from the variants at DHSs + 500 causal variants with MAF < 0.01 sampled from the variants at DHSs; **Est.**: mean estimate averaged from 200 simulations; **s.e.m.**: standard error of the mean.

Supplementary Table 2 Number of SNPs used in the imputation analysis based on the UK10K-WGS data

These are the number of SNPs that are in common between the SNPs arrays and the UK10K data after QC (i.e. UK10K-WGS). The list of SNPs for Affymetrix 6.0, Illumina OmniExpress, and Illumina Omni2.5 were from the ARIC, TwinGene, and HRS data, respectively, after QC. The SNPs for Illumina CoreExome and Affymetrix Axiom were from the strand alignment data files produced and hosted by Will Rayner

(http://www.well.ox.ac.uk/~wrayner/strand/).

Supplementary Table 3 Estimates of variance explained by 1KGP-imputed variants from GREML-LDMS analysis for height and BMI in the combined GWAS data set.

		1st LD quartile		2nd LD quartile		3rd LD quartile		4th LD quartile		Row	
		Est	s.e.	Est	s.e.	Est	s.e.	Est	s.e.	sum	s.e.
Height	2.5×10^{-5} < MAF ≤ 0.001	-0.002	0.014	-0.003	0.011	0.004	0.010	-0.003	0.006	-0.005	0.016
	$0.001 < \text{MAF} \leq 0.01$	0.048	0.013	-0.001	0.011	0.019	0.010	0.018	0.007	0.084	0.019
	$0.01 < \text{MAF} \leq 0.1$	0.031	0.008	0.043	0.007	0.036	0.006	0.021	0.004	0.130	0.011
	$0.1 < \text{MAF} \leq 0.2$	0.029	0.005	0.019	0.004	0.018	0.004	0.014	0.003	0.081	0.008
	$0.2 <$ MAF ≤ 0.3	0.024	0.005	0.028	0.004	0.028	0.004	0.012	0.003	0.092	0.008
	$0.3 < \text{MAF} \leq 0.4$	0.024	0.005	0.029	0.004	0.010	0.004	0.022	0.003	0.085	0.008
	$0.4 < \text{MAF} \leq 0.5$	0.034	0.004	0.019	0.004	0.022	0.004	0.013	0.003	0.088	0.007
	Column sum	0.188	0.019	0.134	0.016	0.137	0.015	0.097	0.009		
	Total sum	0.555	0.023								
	Log likelihood	-20189.2									
	Sample size	43,599									
BMI	2.5×10^{-5} < MAF ≤ 0.001	0.003	0.015	0.015	0.013	0.009	0.011	0.010	0.008	0.038	0.018
	$0.001 < \text{MAF} \leq 0.01$	0.001	0.014	-0.009	0.012	-0.012	0.010	0.006	0.007	-0.014	0.019
	$0.01 < \text{MAF} \leq 0.1$	0.027	0.008	0.008	0.006	0.007	0.005	0.009	0.004	0.051	0.011
	$0.1 < \text{MAF} \leq 0.2$	0.009	0.005	0.020	0.004	0.020	0.004	0.010	0.003	0.059	0.008
	$0.2 < \text{MAF} \leq 0.3$	0.012	0.005	0.014	0.004	0.013	0.004	0.003	0.003	0.042	0.007
	$0.3 < \text{MAF} \leq 0.4$	0.013	0.004	0.010	0.004	0.010	0.003	0.011	0.003	0.044	0.007
	$0.4 < \text{MAF} \leq 0.5$	0.023	0.004	0.015	0.004	0.014	0.003	0.002	0.002	0.054	0.006
	Column sum	0.088	0.020	0.073	0.018	0.061	0.015	0.052	0.010		
	Total sum	0.274	0.025								
	Log likelihood	-21234.5									
	Sample size	43,366									

Note: s.e. for "column sum" or "row sum" refers to standard error of the sum estimate rather than sum of the standard errors.

Supplementary Table 4 Descriptive summary of the cohorts and quality control criteria of the genotype data

Supplementary Note

1. The circumstances under which the estimate of variance explained by all the variants is a (un)biased estimate of heritability

Under an additive genetic model, the quantitative phenotype of an individual *i* can be expressed as

$$
y_i = g_i + e_i \tag{1}
$$

where g_i is the genetic value, $g_i \sim N(0, \sigma_g^2)$ with σ_g^2 being the genetic variance, and e_i is the residual with $e_i \sim N(0, \sigma_e^2)$. The narrow-sense heritability is defined as $h^2 = \sigma_g^2/(\sigma_g^2 + \sigma_e^2)$. If causal variants are known, *g*ⁱ can be expressed as

$$
g_i = \mathbf{z}_i' \mathbf{q}
$$

where $z_i = \{z_{ik}\}\$ is an $m_q \times 1$ vector of the standardized genotype variables of m_q causal variants,

 $z_{ik} = (x_{ik} - 2p_k) / \sqrt{2p_k(1-p_k)}$, x_{ik} is the genotype variable of the *k*-th causal variant (coded as 0, 1 or 2), p_k is the frequency of the coded allele, and **q** is an $m_q \times 1$ vector of the effect sizes of the causal variants with $\mathbf{q} \sim N(0, \mathbf{I}\sigma_g^2/m_q)$. Since z_{ik} is standardized, $E(z_{ik}) = 0$ and var(z_{ik}) = 1. The phenotypic covariance between two individuals based on this model is

$$
cov(y_i, y_j) = cov(g_i, g_j) = cov(\mathbf{z}_i' \mathbf{q}, \mathbf{z}_j' \mathbf{q}) = \mathbf{z}_i' var(\mathbf{q}) \mathbf{z}_j = \sigma_g^2 \mathbf{z}_i' \mathbf{z}_j / m_q
$$
 [3]

If we define $G_{ij} = \mathbf{z}_i' \mathbf{z}_j / m_q$ as the genetic relationship between two individuals at the causal variants³, then $cov(y_i, y_j) = \sigma_g^2 G_{ij}$.

In practice, however, causal variants are largely unknown. We therefore estimate the variance explained using all the variants from whole genome sequencing based on the model below $y_i = g_{v(i)} + e_i$ with $g_{v(i)} = \mathbf{w}'_i$ **u** $[4]$

where $g_{v(i)}$ is the genetic value captured by all the variants with $g_{v(i)} \sim N(0, \sigma_v^2)$, $\mathbf{w}_i = \{w_{ik}\}\$ are the standardized genotype variables of *m* variants, and **u** is an $m \times 1$ vector of the effect sizes of the variants with $\mathbf{u} \sim N(0, \mathbf{I}\sigma_v^2/m)$. The phenotypic covariance between two individuals based on this model is

$$
cov(y_i, y_j) = cov(g_{v(i)}, g_{v(j)}) = cov(\mathbf{w}_i' \mathbf{u}, \mathbf{w}_j' \mathbf{u}) = \mathbf{w}' var(u) \mathbf{w}_j = \sigma_v^2 \mathbf{w}_i' \mathbf{w}_j / m
$$
 [5]

If we define $A_{ij} = \mathbf{w}_i' \mathbf{w}_j / m$ as the genetic relationship between two individuals at all the variants, then $cov(y_i, y_j) = \sigma_v^2 A_{ij}$. The proportion of variance explained by all the variants is defined as $h_v^2 = \sigma_v^2 / (\sigma_v^2 + \sigma_e^2)$.

We have known previously³ that $h_v^2 = h^2$ if A_{ij} is an unbiased predictor of G_{ij} in a sense that

 $E(G_{ij} | A_{ij}) = A_{ij}$, i.e. $h_v^2 = \frac{\text{cov}(G_{ij}, A_{ij})}{\text{cov}(A)}$ $var(A_{ij})$ h^2 . We show below the circumstance under which the estimate of

 h_v^2 is an unbiased or biased estimate of h^2 . Assuming causal variants are a subset of all the variants, we have

$$
cov(G_{ij}, A_{ij}) = cov(\mathbf{z}_{i}^{\prime}\mathbf{z}_{j} / m_{q}, \mathbf{w}_{i}^{\prime}\mathbf{w}_{j} / m) = E(\mathbf{z}_{i}^{\prime}\mathbf{z}_{j} \mathbf{w}_{i}^{\prime}\mathbf{w}_{j}) / (m_{q}m)
$$

\n
$$
= E(\sum_{k}^{m_{q}} z_{ik} z_{jk} \sum_{l}^{m} w_{il} w_{jl}) / (m_{q}m) = E(\sum_{k}^{m_{q}} \sum_{l}^{m} z_{ik} z_{jk} w_{il} w_{jl}) / (m_{q}m)
$$

\n
$$
= \sum_{k}^{m_{q}} \sum_{l}^{m} E(z_{ik} w_{il}) E(z_{jk} w_{jl}) / (m_{q}m) = \sum_{k}^{m_{q}} \sum_{l}^{m} r_{kl}^{2} / (m_{q}m) = \overline{r}_{MQ}^{2}
$$

with \bar{r}_{QM}^2 being the mean linkage disequilibrium (LD) r^2 between the causal variants and all the variants (including the causal variants), and

$$
\operatorname{var}(A_{ij}) = \operatorname{var}(\mathbf{w}_i' \mathbf{w}_j / m) = E(\mathbf{w}_i' \mathbf{w}_j \mathbf{w}_i' \mathbf{w}_j) / m^2
$$

=
$$
\sum_{k}^{m} \sum_{l}^{m} E^2(w_{ik} w_{il}) / m^2 = \sum_{k}^{m} \sum_{l}^{m} r_{kl}^2 / m^2 = \overline{r}_{MM}^2
$$
 [7]

with \bar{r}_{QM}^2 being is the mean LD r^2 between all the variants (including the causal variants), suggesting that $h_v^2 = \frac{\overline{r}_{QM}^2}{\overline{r}_{QM}^2}$ 2 $\overline{r}_{\!\scriptscriptstyle M\!M\!}^{\scriptscriptstyle 2}$ $\frac{2M}{2}h^2$. We therefore can conclude that

1) If causal variants are a random subset of all the variants used in the analysis, $\overline{r}_{MQ}^2 = \overline{r}_{MM}^2$ so that

$$
h_v^2 = h^2.
$$

- 2) Since LD r^2 is a function of allele frequencies between two variants⁴, if there are disproportionally more common or rare causal variants, then \bar{r}_{MQ}^2 will be different from \bar{r}_{MM}^2 , so that $h_v^2 \neq h^2$.
- 3) If causal variants are enriched in genomic regions with lower or higher LD than average, then $\overline{r}_{MQ}^2 \neq \overline{r}_{MM}^2$ so that $h_v^2 \neq h^2$.

It is notable that if there is a difference between h_v^2 and h^2 due to the difference between \bar{r}_{MQ}^2 and \bar{r}_{MM}^2 , such difference (bias) is caused by the difference in models rather than the difference in methods for parameter estimation.

2. The standard error of the estimate of cumulative contribution of variants with MAF $\leq \theta$ **to the genetic variance**

Let L be the variance and covariance matrix of the estimates of genetic components from GREML-LDMS analysis (**L** matrix is available in GCTA-GREML output). The diagonal elements of **L** are $var(\hat{\sigma}_{v(i)}^2)$ (see Online Methods for definition of $\sigma_{v(i)}^2$), and the off-diagonal elements of LD are cov($\hat{\sigma}_{v(i)}^2, \hat{\sigma}_{v(j)}^2$). The cumulative contribution of variants with MAF $\leq \theta$ to the genetic variance is calculated as $\hat{\sigma}_{v}^2(MAF \le \theta) / \hat{\sigma}_{v}^2(MAF \le 0.5)$, where is the sum of estimates of $\hat{\sigma}_{v(i)}^2$ for variants with

MAF < 0.1 and $\hat{\sigma}_{v}^{2}$ (MAF ≤ 0.5) is the sum of all the 28 components. Let $x = \hat{\sigma}_{v}^{2}$ (MAF ≤ θ) and $y = \hat{\sigma}_v^2(MAF \le 0.5)$, the sampling variance of $\hat{\sigma}_v^2(MAF \le \theta) / \hat{\sigma}_v^2(MAF \le 0.5)$ can be calculated from the Delta method⁵ as $var[\frac{x}{x}]$ *y* $\big] \approx \frac{E(x)^2}{E(x)^2}$ $E(y)^2$ var(*x*) $\frac{\text{var}(x)}{E(x)^2} + \frac{\text{var}(y)}{E(y)^2}$ $\frac{\text{var}(y)}{E(y)^2} - \frac{2\text{cov}(x, y)}{E(x)E(y)}$ \int {
ነ $\overline{\mathfrak{l}}$ $\overline{}$ $\left\{ \right.$ \int . In practice, $E(x)$ and $E(y)$ are unknown and is therefore replaced by their estimates $(x \text{ and } y)$. The var (x) , var (y) , and cov (x, y) are

calculated from the sum of the relevant sub-matrix of **L**.

3. Variance explained by population stratification is very small for both height and BMI in our data

In all the analyses described above, we fitted in the model as fixed covariates the first 10 principal components (PCs) calculated from common (MAF > 0.01) imputed variants on HapMap3. We also performed analyses fitting the first 20 PCs or the PCs calculated from all the imputed variants. The results were all very similar (**Supplementary Fig. 9**). This is because the first 20 PCs computed from either all the imputed variants or the common variants on HapMap3 only explained up to 0.8% of variance for height and 0.08% for BMI.

4. GREML-LDMS is robust to the model assumption about the relationship between MAF and effect size

We found strong evidence that variants with MAF < 0.1 explained a larger proportion of variance for height than what we would expect under a neutral model (**Fig. 4a**). We then asked whether or not the estimate could be biased due to our model assumption. If we define $w = (x - 2p)/\sqrt{2p(1-p)}$ with *x* being the genotype variable of a variant (coded as 0, 1 or 2) and *p* being the allele frequency, i.e. *w* is the standardized form *x*, and define *b* and *u* as the regression coefficients of phenotype on *x* and *w*, respectively, then the genetic variance attributable to this variant can be written as $var(xb) = 2p(1-p)b^2$ or var(*wu*) = u^2 , meaning that $u^2 = 2p(1-p)b^2$. By default, the GREML approach³ calculates the genetic relationship using the equation $A_{ij} = \frac{1}{i}$ m_t $(x_{ik} - 2p_k)(x_{jk} - 2p_k)$ *k* $2p_k(1-p_k)$ $\sum_{k}^{m_t} \frac{(x_{ik} - 2p_k)(x_{jk} - 2p_k)}{2n(1 - n)}$ (see Equation 2 in Online

Methods for notations). This model assumes a normal distribution of *u* with constant variance regardless of MAF, which implicitly assumes that the per-allele effect size (*b*) of a rare variant is on average larger than that of a common variant (Assumption A). We therefore re-analysed the data assuming a normal distribution of *b* (Assumption B), which means that variance explained for a rare variant is on average smaller than that for a common variant. Assumption B is equivalent to calculating the genetic relationship using the following equation^{6,7},

$$
A_{ij} = \sum_{k}^{m_i} [(x_{ik} - 2p_k)(x_{jk} - 2p_k)] / \sum_{k}^{m_i} [2p_k(1 - p_k)]
$$
. We found that the GREML-LDMS estimates based on these two assumptions were remarkably consistent (Supplementary Fig. 12), demonstrating the robustness of the GREML-LDMS method to different assumptions about the distribution of effect sizes. In addition, we observed from the genome-wide association analysis a correlation between effect size and MAF, and showed by simulations (mimicking the observed genetic architecture for height) that the GREML-LDMS was also robust to such a correlation (Supplementary Fig. 13).

5. An approximate method to estimate $h^2{}_{\text{WGS}}$ from $h^2{}_{\text{1KGP}}$

We quantified by simulations based on 5 different types of SNP arrays under four different scenarios that on average \sim 97% and \sim 68% of variation at common and rare variants, respectively, can be captured by KGP imputation (**Supplementary Fig. 4**). This implies that

 $h^2 \approx h_{1KGP(rare)}^2/0.68 + h_{1KGP(common)}^2/0.97$, where $h_{1KGP(rare)}^2$ and $h_{1KGP(common)}^2$ denote h_{1KGP}^2 for rare and common variants, respectively. Note that this is still likely to be an underestimate of heritability because complicated structure variation and extremely rare variation are likely to be less well captured by 1KGP imputation than those used in our simulation. Using this approximate method, we quantified the lower limit of heritability of 0.61 (s.e. $= 0.045$) for height and 0.29 (s.e. $= 0.047$) for BMI.

6. Other possible sources of missing heritability

It has been suggested that epistasis and GxE can explain a substantial amount of variance for allelespecific expression traits⁸. However, the extent to which epistasis and GxE contribute to complex trait variation in general remains unclear. The missing heritability problem refers to the gap between additive genetic variance estimated from family studies and that from population-based studies (e.g. GWAS), so that non-additive genetic variation is irrelevant unless h^2 is overestimated in family-based studies due to the confounding with non-additive genetic variation. However, such confounding is negligible if the total amount of variance attributed to non-additive genetic variation is small. In fact, a recent study⁹ shows that on average across a number of quantitative traits dominance variation explains \leq 4% of phenotypic variance. Quantitative genetics theory also predicts that epistasis and other higher order interaction terms are unlikely to explain a substantial amount of variance for complex traits¹⁰. Therefore, non-additive genetic variation is unlikely to contribute substantially to the missing heritability. GxE can be relevant only if there is a strong component of GxE variance and h^2 is estimated from family-based samples in a single homogenous environment where GWAS is performed in samples from multiple heterogeneous environments.

7. Enrichment of variance explained in lower-LD regions is driven by MAF

Our results also seem to suggest that variants in genomic regions with lower LD tend to explain a larger proportion of variance than those in regions with higher LD, 18.8% vs. 9.7% ($P_{\text{difference}} = 3.0 \times 10^{-5}$) for height and 8.8% vs. 5.2% ($P_{difference} = 0.12$) for BMI (**Supplementary Table 3**). This, however, is

likely due to the enrichment of lower-MAF variants in regions with lower LD (**Supplementary Fig. 14**).

8. The Morrison et al. WGS study

Morrison et al.¹¹ applied the GREML method in a WGS data set of 962 individuals and concluded that the majority of the heritability of high-density lipoprotein cholesterol level can be attributable to common variation. The s.e. of \hat{h}^2_{WGS} for common variants from Morrison et al. is 0.14, much smaller than that expected from theory¹² given the sample size. The reported s.e. is consistent with the variance of the off-diagonal elements of the genetic relationship matrix (GRM) being 1.1×10^{-4} for common variants, \sim 7 times larger than what we observed in the UK10K data (1.6 \times 10⁻⁵). This suggests that the variance of GRM in Morrison et al. was highly inflated, possibly due to the adjustment that they made to the GRM (the adjustment is under strong assumptions³ and actually not necessary for WGS data) and/or due to cryptic relatedness. If all the 962 individuals are unrelated and there is no adjustment to the GRM, the s.e. of \hat{h}_{WGS}^2 for common variants should be ~ 0.37 rather than 0.14. Such inflation in the variance of estimated genetic relatedness could potentially bias their results from simulations and their conclusion about variance explained by common variation from the analysis of real data.

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