Population genetic differentiation of height and body mass index across Europe

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Across-differences in the mean values for complex traits are common1–8, but the reasons for these differences are unknown. Here we find that many independent loci contribute to population genetic differences in height and body mass index (BMI) in 9,416 individuals across 14 European countries. Using discovery data on over 250,000 individuals and unbiased effect size estimates from 17,500 sibling pairs, we estimate that 24% (95% credible interval (CI) = 9%, 41%) and 8% (95% CI = 4%, 16%) of the captured additive genetic variance for height and BMI, respectively, reflect population genetic differences. Population genetic divergence differed significantly from that in a null model (height, $P < 3.94 \times 10^{-9}$; BMI, $P < 5.95 \times 10^{-4}$), and we find an among-population genetic correlation for tall and slender individuals ($r = -0.80$, 95% CI = $-0.95$, −0.60), consistent with correlated selection for both phenotypes. Observed differences in height among populations reflected the predicted genetic means ($r = 0.51$; $P < 0.001$), but environmental differences across Europe masked genetic differentiation for BMI ($P < 0.58$).

Many of the phenotypes that vary within human populations are complex, in that they are determined by alleles at multiple loci and many non-genetic factors9–15. Therefore, it is reasonable to assume that regional differences in such traits have a complex basis16–18. Understanding these regional differences requires knowledge of the relative roles of environmental and genetic effects, which can be gained through estimating the amount of population genetic variance in phenotype and by determining the amount of observed differences that are explained by population genetic effects19. Thus far, these estimates have yet to be made outside of laboratory study populations20, and experimental designs for human populations have been lacking because of confounding from genetic and environmental effects.

At least 135 million European citizens are obese21, resulting in major direct and indirect health and economic costs22,23. Regional differences across Europe in height and susceptibility to weight gain, as defined by BMI, are well documented2,24–26, but the reasons for these differences are not well understood. Height and BMI are extensively studied, and there is strong evidence that a large number of genetic polymorphisms influence both traits, with a considerable proportion of genetic variance captured by common SNPs24–28. For height, there is empirical evidence for selection at height-associated SNP loci within Europe26,31 and between European populations and the rest of the world32. However, the true extent of population genetic differentiation is unlikely to have been captured or well represented by the limited number of ascertained loci examined thus far. For BMI, it is not known whether genetic differentiation exists, and, for both traits, the extent to which common loci contribute to population genetic variance or the observed regional phenotypic differences remains unknown (although see ref. 26).

Here we estimate cumulative population genetic differentiation for height and BMI captured by multiple unlinked loci across 9,416 Europeans from 14 countries, using population genetic analyses (Supplementary Methods and Supplementary Figs. 1 and 2). We performed genome-wide association study (GWAS) meta-analyses on data from recent studies33,34, selecting independent loci (linkage disequilibrium (LD) $r^2 < 0.1$ and >1 Mb apart35) associated with either trait in a European-ancestry sample (~250,000 individuals for height and ~350,000 individuals for BMI). We reestimated the effects of each SNP in a within-family study design, which is unbiased by population stratification, and used these effect sizes to create a genetic predictor for both phenotypes (also termed a ‘profile’ or ‘polygenic score’)36.

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The proportion of variance in the genetic predictor attributable to population differences was estimated in a Bayesian mixed-effects model, alongside the co-differentiation of the phenotypes and the predicted means for each nation (Online Methods). Using theory and simulation study, we show that, whereas within-family effect sizes are unbiased, population genetic analyses conducted using loci ascertained from a standard GWAS can be biased if population stratification is not fully accounted for (Online Methods, Supplementary Figs. 3–5 and Supplementary Note). There is no certainty that population stratification is completely controlled for in large-scale meta-analyses, and we thus repeated our analysis using (i) non-ascertained, unlinked (LD $r^2 < 0.1$ and >1 Mb apart), common (minor allele frequency (MAF) >1%) HapMap 3 loci (~40,000 SNPs) and (ii) unlinked (LD $r^2 < 0.1$ and >1 Mb apart), common (MAF >1%) HapMap 3 loci selected on the basis of their within-family association with each phenotype (~40,000 SNPs for both traits). This analysis provides an unbiased, genome-wide estimate, representing a lower limit of population genetic differentiation at common, unlinked loci.

The maximum proportion of variance in a polygenic predictor attributable to population genetic differences was 24% (95% CI = 9%, 41%) and 8% (95% CI = 4%, 16%) for height and BMI, respectively, using 2,660 SNPs for height and 11,919 SNPs for BMI. For height, the largest proportion of population-level variance was captured by SNPs of low P value in the meta-analysis (Supplementary Fig. 6). For BMI, the continual addition of SNPs increased the proportion of population-level variance captured (Supplementary Fig. 6). For both traits, the among-population variation was greater in predictors that explained greater phenotypic variance (Supplementary Fig. 7). Our results were confirmed using the non-ascertained independent, genome-wide loci (height: 8.6%, 95% CI = 3%, 15.7%; BMI: 2.8%, 95% CI = 1.1%, 5.3%) and the set of independent, genome-wide loci selected on the basis of their within-family association (height: 11.9%, 95% CI = 4.5%, 21.8%; BMI: 8%, 95% CI = 3.4%, 14.7%). The lower among-population variance captured using non-ascertained loci reflects reduced prediction accuracy, likely due to the addition of a large number of loci with no detectable association. Subsequent results are presented using the predictor for each trait that captured the greatest amount of population- and individual-level variance (comprising 2,660 SNPs for height and 11,919 SNPs for BMI); however, the results remained the same irrespective of the SNPs selected (Supplementary Fig. 8). The predicted population genetic means for the traits are shown in Figure 1 alongside the observed values, estimated from an independent set of recently published data, accounting for trends over time. Among-population differences in allele frequency are expected to create genetic differences in height, such that people from the Netherlands are on average 1 cm taller than those from Italy, and genetic differences in BMI, such that, on average, the BMIs for people from Italy and Denmark differ by 0.2 units (Figs. 1 and 2).

Genetic differences among populations may occur by random, chance processes or through natural selection in the evolutionary past. We thus compared our estimates to the values from a null quantitative genetic model of multivariate population differentiation. We found strong evidence that the divergence of each trait was greater than expected under the neutral model (Fig. 2). The overall level of neutral genetic differentiation was small for both height (1.2%; 95% CI = 0.01%, 1.78%) and BMI (1.9%, 95% CI = 0.48%, 2.97), reflecting the average $F_{ST}$ (a measure of population differentiation due to genetic structure; Supplementary Note) of the SNP sets between the populations of 1% for height and 1.2% for BMI. Our results were confirmed using non-ascertained independent, genome-wide loci (height, $P = 3.29 \times 10^{-6}$; BMI, $P = 0.018$) and independent, genome-wide loci selected on the basis of their within-family association (height, $P = 2.67 \times 10^{-6}$; BMI, $P = 8.35 \times 10^{-5}$). We therefore reject the null model, and our results suggest that population genetic differentiation across these 14 European countries for height and BMI has been driven by selection on standing genetic variation across geographical regions in the evolutionary past.

The significant departure from a neutral model occurs because, on average, the common loci comprising the genetic predictor are differentiated in a direction that is consistent with the direction of
their effects on each trait, which in turn creates differences among countries in a genetic predictor. Our interpretation does not depend on LD, as we used only independent loci, nor does it depend on the SNPs used, as we generated our null model estimates from the same set of SNPs that we used to create the genetic predictor. Loss of LD, genotype-environment interactions and genetic heterogeneity can change the relationships between SNPs and underlying causal variants across countries, but such changes would reduce the likelihood of detecting population genetic differentiation. In this study, we minimized these effects by using populations of European ancestry throughout.

We estimated the population genetic co-differentiation of height and BMI to ask whether selection has acted on both traits independently. We found a negative correlation between the population genetic means of −0.80 (95% CI = −0.95, −0.60; Fig. 2). This finding was consistent across predictors comprising non-ascertained genome-wide loci (−0.77, 95% CI = −0.94, −0.55) and independent genome-wide loci selected on the basis of their within-family association (−0.89, 95% CI = −0.97, −0.77). These results imply that selection has acted on common loci to increase height while reducing BMI and vice versa, and a genetic predisposition for tall stature at the population level was associated with a genetic predisposition for slenderness (low BMI). As height and BMI are nearly uncorrelated at the individual level (correlation among genetic profile scores within populations r = −0.016, 95% CI = −0.041, 0.001), selection for one trait should not elicit a response in the other. Our results suggest that selection has acted on both phenotypes, although, as some genes affect both phenotypes, we cannot rule out differentiation in one trait having been mediated by selection for the other. The population genetic co-divergence shown here is inconsistent with random genetic drift because the expectation with drift is that the among-population genetic correlation will equal the within-population correlation (77, 49).

We tested whether the observed phenotypic differences across the 14 European countries reflect genetic differentiation at common loci or whether current environmental differences among countries (in diet, economy, climate, etc.) mask the population genetic

![Figure 2](image1.png)  
**Figure 2** Predicted genetic differentiation compared to the expectation under genetic drift for height and BMI across 14 European nations. (a, b) The mean predicted genetic differentiation (blue) and differentiation under the null model representing genetic drift (gray) are shown for 14 European nations, with 95% credible intervals, for height (a) and BMI (b). ISO 2 country codes indicate each nation. The average P value of differentiation from the null expectation is shown in the figure. (c) The pattern of population-level co-differentiation for height and BMI across 14 European nations. The negative population genetic co-differentiation for these traits of −0.80 (95% CI = −0.95, −0.60) is represented by the blue ellipse.

![Figure 3](image2.png)  
**Figure 3** Association between observed population means and predicted genetic population means for height and BMI across 14 European nations. (a, b) Predicted population genetic means are plotted against observed population means for height (a) and BMI (b). P values give the significance of the multivariate Pearson product-moment correlation between the predicted population genetic means and the observed population means for both traits. For height, the correlation (r = 0.51, 95% CI = 0.39, 0.61) was greater than that expected under the null model (r = 0.03, 95% CI = −0.21, 0.17). For BMI, the correlation (r = −0.010, 95% CI = −0.19, 0.01) was not significantly different from the null expectation (r = −0.08, 95% CI = −0.24, 0.15).
Supplementary Figs. 9 and 10), a pattern expected on the basis of our theory (Supplementary Figs. 11 and 12) and supported by the fact that the proportion of population-level variance was greater in a predictor explaining greater phenotypic variance. We found no evidence for significant among-population differentiation at any SNP across the genome, implying that many loci of small effect combine across the genome to create the detected genome-wide population genetic differentiation. Annotation of the 500 SNPs contributing the most to each trait with genes suggests that population genetic variation across the 14 countries for height and BMI is underlain by the combined effects of multiple pathways, with an overlap in the genes involved (Supplementary Fig. 13).

Finally, we examined population genetic differentiation in height and BMI on both a local and worldwide scale. We found no evidence for population genetic differentiation across six northern Italian villages10 (Supplementary Fig. 14). We then examined population differentiation for height and BMI in the Human Genetic Diversity Panel (HGDP), as used in a previous study32 (Supplementary Fig. 15). For both phenotypes, we found evidence to reject the null hypothesis that population genetic differentiation reflects neutrality (Supplementary Fig. 15), extending previous work that reported no significant differentiation for BMI using a limited number of loci32. Additionally, we found no evidence for population genetic co-differentiation of height and BMI, implying a European-specific pattern of selection.

The conclusions of our study are fourfold: (i) many common loci combine in a consistent manner to create population genetic differences for height and BMI; (ii) population genetic differentiation for height and BMI does not reflect a pattern expected under neutrality, and selection has thus driven the differences observed; (iii) population genetic divergences for height and BMI are correlated within Europe, with this correlation reaching a greater level than expected under neutrality, implying that the height-associated loci under selection are enriched for loci with effects that reduce BMI; and (iv) selection-driven population genetic differences for height reflect the phenotypic patterns we see across Europe, whereas, for BMI, environmental factors are masking the population genetic differences. Although genotype-environment effects and rare variants will have a role in shaping population genetic differentiation, the focus of our approach is on estimating the amount of population-level variance that is ‘tagged’ by a specific set of common SNP markers. As additional genetic variation is captured for both traits, it is likely that power will increase to fully capture the among-population genetic effects. The theoretical and analysis framework builds upon previous approaches26,32, is entirely general and can be applied to estimate the role of commonly varying loci in shaping population differences for any set of phenotypes.

METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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ONLINE METHODS

Genetic differentiation of height and BMI across Europe. The theoretical basis of our analysis framework builds upon those from refs. 32,47,51–53 and is outlined in the Supplementary Note. We apply this framework to examine population genetic differentiation in height and BMI across Europe. An overview of the analysis steps is provided in Supplementary Figure 1.

Imputation. All of the cohorts used in this study were independently imputed to the 1000 Genomes Project reference panel, using identical quality control procedures on the initial data sets of per-SNP missing data rate <0.02, MAF >0.01, per-individual missing data rate <0.05 and Hardy-Weinberg disequilibrium P <0.0001. Imputation for the majority of the cohorts was performed in two stages. First, the target data were assigned to haplotypes using HAPI-UR. Second, IMPUTE2 (ref. 55) was used to impute the haplotypes to the 1000 Genomes Project reference panel (release 1, version 3). We then selected SNPs that were present across all data sets at an imputation information score of >0.8. The imputation for the Netherlands cohort was identical, except that SHAPEIT2 (ref. 57) was used for haplotyping. We performed these same quality control steps again after combining the data from the different cohorts, including comparisons of allele frequencies across populations.

Selection of SNPs for genomic profiling across Europe. We performed GWAS meta-analyses on data from recent studies33–34 to select independent loci (r² <0.1 and >1 Mb apart) using the PLINK clumping procedure35 that were associated with the traits in a large sample of individuals (~250,000 for height and ~350,000 for BMI) of European ancestry. We excluded cohorts overlapping with our within-family and prediction samples. For height, we excluded the TWINGENE study, the TwinsUK study, the QIMR sample, the Framingham Heart Study sample and the Netherlands Twin Register. For BMI, cohort-level summary statistics were not available for all samples, and we could only exclude the QIMR sample.

Within-family estimation of SNP effects. We reestimated the SNP effects at these loci in a within-family sibling pair data set (Supplementary Table 1) using the QFAM procedure in PLINK described in equation (3.1) in the Supplementary Note.

Population genetic analyses conducted using ascertainment loci from a standard GWAS can be biased if population stratification is not fully accounted for (Supplementary Note). Thus, to confirm our results, we also selected a non-ascertained genome-wide set of unlinked (LD r² <0.1 and >1 Mb apart) that passed quality control in both the within-family and prediction samples. We estimated the effects of these SNPs again using our within-family sibling pair data set with the QFAM procedure in PLINK described in equation (3.1) in the Supplementary Note. Additionally, we used the clumping procedure in PLINK to select genome-wide unlinked (LD r² <0.1 and >1 Mb apart), common (MAF >1%) HapMap 3 loci for height and BMI (~40,000 SNP loci) that passed quality control in both the within-family and prediction samples. We estimated the drift profile and trait profile scores of the phenotype and genetic predictor, respectively, in equation (6.1) to equation (6.4) in the Supplementary Note.

Genomic profile scoring in a collection of European genomic data. We used the within-family effect sizes to create genetic predictors for individuals across a collection of European genomics data. All data were imputed as described above, and details of the cohort are provided in Supplementary Table 1.

From the POPRES study, we selected individuals from France, Portugal, Spain, Italy and Switzerland whose grandparents were born in the same country as the sampled individuals. From the Estonian and Finnish cohorts, we selected 1,000 individuals at random who were included in all analyses. For the Netherlands cohort, we selected 1,000 control individuals from the MinE ALS study. These individuals were healthy controls born in the Netherlands, whose grandparents and parents were also born in the Netherlands. From the Psychiatric Genomic Consortium and the Wellcome Trust Case Control Consortium 2, we used control individuals from Bulgaria, Ireland, Norway, Denmark, Sweden and the UK (Supplementary Table 1). The POPRES data set used in this manuscript was obtained from the database of Genotypes and Phenotypes (dbGaP) through accession phs000145.v4.p2.

We used the genetic predictors for height and BMI as response variables in a bivariate Bayesian mixed-effects model, which is outlined in equation (2.2) of the Supplementary Note. This model was estimated using the R package MCMCglmm 39, with uninformative inverse Wishart priors, a burn-in period of 7,000 iterations, a sampling interval of 10 iterations and a total of 17,000 iterations, providing 1,000 posterior estimates. This provided estimates of the population genetic (co)variance of height and BMI, the residual (individual-level) (co)variance and the best linear unbiased predictors of the means of each nation along with the 95% credible intervals. For each set of SNPs, the genetic predictor was standardized to a z score to allow comparison across SNP sets and comparison with the null model.

Comparison with a null model. At each stage of the analysis, the estimates obtained were compared to the expectations under random genetic drift, using a quantitative genetic framework for studying population differentiation that is outlined in the Supplementary Note.

In brief, the within-family regression coefficients were randomized across SNPs 1,000 times, and 1,000 genetic predictors were created in the European prediction sample. By keeping the effect sizes consistent but attributing these effects across SNPs at random, the genetic predictors generated reflect the action of genetic drift. Second, each set of genetic predictors was standardized to a z score and used as a response variable in the Bayesian mixed-effects model outlined in equation (2.2) of the Supplementary Note. This provided 1,000 estimates of the population genetic variance and population means under drift; these values are displayed in the figures as the estimates from the neutral model. Third, the sample covariance matrix of these 1,000 estimates was calculated, which provided an estimate of the expected population-level covariance in phenotype under drift. We then used a Mahalanobis distance statistic to provide a measure of the relative deviation of our predicted population-level means from their multivariate theoretical expectations under drift (Supplementary Note). This calculation provided the z² test statistic used to compare our predicted estimates to the expected values under drift. As both the drift profile and trait profile scores were transformed to a z score, this comparison was on the same standard deviation scale.

European phenotypic data. Height and BMI measures for males for each of the 14 European countries were taken from recently published estimates25,37. For height, measures were available from 1860 to the present day, and, for BMI, measures were available from 1980 to the present day. Values for both phenotypes were adjusted for trends over time before estimating the population means within a mixed-effects model. The model-generated estimates of these means were then compared to the predicted genetic means as described in equation (5.1) to equation (5.4) in the Supplementary Note.

Transforming population means onto the observed scale. For graphical presentation of the results, each genetic predictor zg created for each individual i for each trait m, was approximately transformed back to the observed scale as outlined in equation (6.1) in the Supplementary Note.

We used an independent sample of population data to determine the amount of phenotypic variance explained by the genetic predictors created from different sets of SNPs, as measured by cov(zg, zp), where zg and zp are the z scores of the phenotype and genetic predictor, respectively, in equation (6.1) of the Supplementary Note. Individuals within the Health and Retirement Study (HRS) (Supplementary Table 1) were unrelated, and phenotypic values were adjusted by the first 20 principal components of the SNPs used in the predictor, to account for any population stratification before estimating the within-population covariance. The standard deviation for height and BMI measured in the sample of 17,500 quasi-independent sibling pairs was estimated accounting for sex differences. The HRS data set was obtained from dbGaP through accession phs000428.v1.p1.

Testing the contribution of each SNP to the pattern of population differentiation. As described in equation (4.4) in the Supplementary Note, we estimated a z² value for each SNP representing its contribution to the pattern of population genetic differentiation. We tested for the association between a SNP’s contribution to differentiation and its MAF and expected contribution to phenotypic variation estimated as 2pqβ², where p is the frequency of the minor allele, q is the frequency of the major allele and β is the within-family effect size.
Positional and functional annotation of SNPs. Our aim was to describe the positional genetic annotation and gene ontology (GO) categories for the height- and BMI-associated SNPs that contributed the most to the genome-wide pattern of population genetic variance. We selected the 500 height- and BMI-associated SNPs that contributed the most to differentiation. First, on the basis of the genomic position of these SNPs, we assigned them to genes and simply estimated the number of overlapping genes involved in the genetic differentiation of height and BMI. Second, SNPs were assigned to the following positional genetic annotation categories: 3’ UTR variant, 5’ UTR variant, intronic variant, noncoding transcription variant, variant 1–1,000 bp downstream of a gene, variant 1–1,000 bp upstream of a gene, missense transcription variant, synonymous transcription variant and noncoding exonic variant. Finally, SNPs were assigned Ensembl gene identities and then GO terms. The top GO annotations are listed in Supplementary Table 2.

We then conducted statistical testing. As a baseline, we used the top 10,000 SNPs for height and BMI. We first compared the number of SNPs among the 500 SNPs contributing the most to differentiation in each genic category, to count data from the top 10,000 SNPs. This provided a list of genic categories potentially enriched for highly differentiated SNPs. We used Fisher’s exact tests (hypergeometric) with Bonferroni P-value correction. Second, we compared the number of SNPs among the 500 SNPs contributing the most to differentiation in each GO term, to count data from the top 10,000 SNPs. Again, we used Fisher’s exact tests with Bonferroni P-value correction. From this analysis, we selected a list of the top 20 functional categories potentially enriched for each trait. We assigned P values to our comparisons of count data but used these only as a guide to select the top categories, rather than as a definitive test of enrichment. All annotation was conducted using the R library biomaRt from Bioconductor.

Genomic profile scoring worldwide. We repeated our analyses for height and BMI using data from the HGDP as analyzed in ref. 32. We imputed the data following our protocol outlined above.

Simulation study using real genotype data. In addition to the simulations described in the Supplementary Note, we used the common, independent HapMap 3 SNPs from the 17,300 sibling pairs in the main empirical analyses (Supplementary Table 1) as the basis for a series of simulations. We randomly assigned 5,000 of the independent loci across the genome to be causal variants, with their effects sampled from a normal distribution with a mean of 0 and a variance of 1. The heritability of the trait was simulated to be 90%, and a phenotype was created as $y = \sum x_i b_i + \epsilon$, where $x_i$ is the indicator value of locus $k$ (0, 1, 2), $b_i$ is the effect size of the locus and $\epsilon$ is the residual variance, with $\epsilon = N(0, 1 - h^2)$, where $h^2$ is the heritability. Fifty simulation replications were conducted.

For each simulation replicate, we randomly selected 16,000 sibling pairs to create an estimation set, leaving 1,500 pairs as a prediction set. We then tested the effects of each SNP on the phenotype in the estimation set in three ways. First, we used a within-family sibling pair analysis implemented using the QFAM procedure in PLINK as described in equation (3.1) in the Supplementary Note. Second, we selected one member of a sibling pair at random to create an unrelated set of individuals and then estimated SNP effects using ordinary least-squares regression (the standard GWAS approach) without any control for population stratification. Finally, we repeated the GWAS estimation and controlled for the first 20 principal components estimated from the sibling pair sample.

We then used these three sets of estimates to create three different profile scores in the prediction set. We followed a recent approach to partition variance in a predictor in sibling pairs into genetic, environmental, common genetic and common environmental terms51. Variance attributable to common genetic or common environmental terms indicates population stratification bias in the effect size estimates, enabling us to demonstrate that our within-family estimates are unbiased by population stratification.

Additionally, we used the 3 sets of effect size estimates to create 3 different profile scores in the European prediction data of 9,416 individuals used in the main empirical analyses. Our European prediction data were projected onto the first principal component estimated in the within-family sample, and two groups of individuals were then created based on the upper and lower quartiles of the distribution of the projected principal component. The first principal component has been shown to reflect population stratification within genotype samples, and we thus stratified the independent prediction sample by the predominant axis of potential bias in the discovery sample. If there is no population stratification bias in estimates of SNP effects, then there should be no significant differentiation between the groups corresponding to the upper and lower quartiles with respect to the mean for the genetic predictor. We therefore compared the estimates obtained from the three predictors to those generated when a predictor was created using the true simulated effects and to those generated when a predictor was created using randomly allocated effect sizes under our null model. To test for ascertainment biases, we selected the top 100 and top 500 SNPs identified by a GWAS that did not control for population stratification, created 2 predictors containing these SNPs—one from the GWAS effect size estimates and one from the within-family estimates—and then compared the mean differences between the groups corresponding to the upper and lower quartiles. This procedure identifies SNPs from a GWAS and then tests for population differentiation as in previous approaches.

Finally, we repeated the 50 simulations described above, but we created a genotype-environment correlation. We used the same effect sizes and phenotype as in the simulation described above, but we added a z score of the standardized individual-level eigenvalue of the first principal component estimated in the within-family data. Thus, our phenotype was described as $y = \sum x_i b_i + 0.2PC_1 + \epsilon$, where PC_1 is the z score–standardized value for each individual at the first principal component. This creates a correlation between the phenotype and the first principal component of ~0.2 across simulations, representing a phenotypic difference that aligns to the major axis of genetic differentiation. Note that, in this scenario, there is no selection because causal variants are allocated at random across the genome such that, on average, their frequencies across simulations are not expected to differ in a manner that will create a consistent directional difference in profile score along any axis of population stratification. We repeated the estimation and prediction in our independent European prediction sample to test for a directional deviation in mean profile scores between the groups corresponding to the upper and lower quartiles. Under this scenario, ascertainment biases are expected to be large because the SNPs identified by a GWAS that does not control for population stratification should correspond to the ones with the strongest genotype-environment correlation, creating a predictor with mean differences across the leading principal component. However, across the genome, the expectation is that the direction of SNP differentiation should not align with the direction of the effect size and the direction of the phenotypic differentiation, provided that the predictors are created from SNP effect sizes that are unbiased by population stratification.

Code availability. Full computer code for the analysis and the values used to produce the figures are available from the lead author upon request.