Genomic structural equation modelling provides insights into the multivariate genetic architecture of complex traits

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Genetic correlations estimated from genome-wide association studies (GWASs) reveal pervasive pleiotropy across a wide variety of phenotypes. We introduce genomic structural equation modelling (genomic SEM): a multivariate method for analysing the joint genetic architecture of complex traits. Genomic SEM synthesizes genetic correlations and single-nucleotide polymorphism heritabilities inferred from GWAS summary statistics of individual traits from samples with varying and unknown degrees of overlap. Genomic SEM can be used to model multivariate genetic associations among phenotypes, identify variants with effects on general dimensions of cross-trait liability, calculate more predictive polygenic scores and identify loci that cause divergence between traits. We demonstrate several applications of genomic SEM, including a joint analysis of summary statistics from five psychiatric traits. We identify 27 independent single-nucleotide polymorphisms not previously identified in the contributing univariate GWASs. Polygenic scores from genomic SEM consistently outperform those from univariate GWASs. Genomic SEM is flexible and open ended, and allows for continuous innovation in multivariate genetic analysis.

Genome-wide association studies (GWASs) are rapidly identifying loci affecting multiple social, behavioural and psychiatric phenotypes. Moreover, using cross-trait versions of methods such as genomic-relatedness-based restricted maximum likelihood (GREML) and linkage disequilibrium score regression (LDSC), researchers have identified genetic correlations between diverse traits; for example, age of first birth and risk of smoking, insomnia and psychiatric traits (for example, schizophrenia), major depressive disorder (MDD) and number of children, and educational attainment and cognitive performance. Widespread statistical pleiotropy appears to be the rule rather than the exception across complex traits. Although these findings are currently suggestive of constellations of phenotypes affected by shared sources of genetic liability, existing methods do not permit the causes of the observed genetic correlations to be investigated systematically. Here, we introduce genomic structural equation modelling (genomic SEM)—a new method for modelling the multivariate genetic architecture of constellations of traits and incorporating genetic covariance structure into multivariate GWAS discovery. Genomic SEM is a flexible framework for formally modelling the genetic covariance structure of complex traits using GWAS summary statistics from samples of varying and potentially unknown degrees of overlap, in contrast with existing methods that model phenotypic covariance structure, with specific applications, using raw data. Moreover, genomic SEM allows for the specification and comparison of a range of proposed multivariate genetic architectures, which improves on existing approaches for combining information across genetically correlated traits to aid in discovery.

One powerful feature of genomic SEM is the capability to model shared genetic architecture across phenotypes with factors representing broad genetic liabilities, and compare the fit of different factor structures to the empirical data. When an appropriate model has been identified at the level of the genome-wide covariance structure, the researcher may incorporate individual single-nucleotide polymorphisms (SNPs) into the model to identify variants with effects on general dimensions of cross-trait liability, boost power for discovery, and calculate more valid and predictive polygenic scores (PGSs). Genomic SEM can also evaluate whether the multivariate genetic architecture implied by a specific model is applicable at the level of individual variants using developed estimates of heterogeneity. When certain SNPs only influence a subset of genetically correlated traits, a key assumption of other multivariate approaches is violated. SNPs with high heterogeneity estimates can be flagged as likely to confer disproportionate liability towards individual traits, be removed when constructing polygenic risk scores, or be studied specifically to understand the nature of heterogeneity.

We validate key properties of genomic SEM with a series of simulations, and illustrate the flexibility and utility of genomic SEM with...
analyses of real data. These include a joint analysis of GWAS summary statistics from five genetically correlated psychiatric case-control traits: schizophrenia, bipolar disorder, MDD, post-traumatic stress disorder (PTSD) and anxiety. We model their joint genetic architecture using a general factor of psychopathology ($p$), for which we identify 27 independent SNPs not previously identified in the univariate GWASs, 5 of which can be validated based on separate GWASs. PGs derived using this $p$ factor consistently outperform PGs derived from GWASs of the individual traits in out-of-sample prediction of psychiatric symptoms. Other demonstrations include a multivariate GWAS of neuropsychiatric items, an exploratory factor analysis (EFA) of anthropometric traits and a simultaneous analysis of the unique genetic associations between schizophrenia, bipolar disorder and educational attainment.

Results

Genomic SEM is a two-stage structural equation modelling approach\textsuperscript{12–14}. In stage 1, the empirical genetic covariance matrix and its associated sampling covariance matrix are estimated. The diagonal elements of the sampling covariance matrix are squared standard errors. The off-diagonal elements index the extent to which sampling errors of the estimates are associated, as may be the case when there is sample overlap across GWASs. In stage 2, a SEM is specified and parameters are estimated by minimizing the discrepancy between the model-implied genetic covariance matrix and the empirical covariance matrix obtained in the previous stage. We evaluate fit with the standardized root mean square residual (SRMR), model $\chi^2$, Akaike Information Criterion (AIC) and Comparative Fix Index (CFI; see Methods)\textsuperscript{15–17}. In a set of simulations, we verify key properties of genomic SEM (Methods). We find that genomic SEM produces unbiased parameter estimates when the correct structural model is specified, and that model fit indices consistently favour the correct model over alternative models. In a second set of simulations, we demonstrate that the inclusion of data from overlapping samples does not bias genomic SEM parameter estimates or their standard errors.

Genomic SEM can be employed as a tool for multivariate GWASs based on univariate summary statistics. First, the genetic covariance matrix and its associated sampling covariance matrix are expanded based on univariate summary statistics. First, the empirical genetic covariance matrix is estimated. The genetic covariance matrix and its associated sampling covariance matrix are estimated. The diagonal elements of the sampling covariance matrix are squared standard errors. The off-diagonal elements index the extent to which sampling errors of the estimates are associated, as may be the case when there is sample overlap across GWASs. In stage 2, a SEM is specified and parameters are estimated by minimizing the discrepancy between the model-implied genetic covariance matrix and the empirical covariance matrix obtained in the previous stage. We evaluate fit with the standardized root mean square residual (SRMR), model $\chi^2$, Akaike Information Criterion (AIC) and Comparative Fix Index (CFI; see Methods)\textsuperscript{15–17}. In a set of simulations, we verify key properties of genomic SEM (Methods). We find that genomic SEM produces unbiased parameter estimates when the correct structural model is specified, and that model fit indices consistently favour the correct model over alternative models. In a second set of simulations, we demonstrate that the inclusion of data from overlapping samples does not bias genomic SEM parameter estimates or their standard errors.

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We provide two examples of confirmatory factor analysis (CFA) using genomic SEM. In our first example, we fit a genetic factor model to psychiatric case-control traits. Recent findings indicate that the comorbidity across psychiatric disorders is captured by a general psychopathology factor (that is, the $p$ factor), and this is widely supported based on previous results\textsuperscript{17–21}. We tested for the presence of a single common genetic $p$ factor using genomic SEM with European-only summary statistics for schizophrenia, bipolar disorder, MDD, PTSD and anxiety (Supplementary Table 1 for phenotypes and sample sizes). The model fit was adequate ($\chi^2(5) = 89.55; AIC = 109.50; CFI = 0.848; SRMR = 0.212$). The results indicated that schizophrenia and bipolar disorder loaded the strongest onto the genetic factor (Supplementary Fig. 1)—a pattern of findings that closely replicates previous findings from twin/family studies\textsuperscript{19}.

In a second example, we tested for the presence of a single common genetic factor of neuroticism using summary statistics from 12 item-level indicators from UK Biobank (UKB; Supplementary Table 1), as estimated using the Hail software\textsuperscript{2}. The model fit was good ($\chi^2(54) = 4,884.10; AIC = 4,932.11; CFI = 0.893; SRMR = 0.109$). The results indicated strong positive loadings for all indicators (Supplementary Fig. 2). We used this single common factor model for both neuroticism and the $p$ factor when estimating SNP effects for discovery under the section ‘SNP effects’ below.

EFA of a genetic covariance matrix. We provide two examples of how one might use exploratory methods to guide the specification of more nuanced factor models. In the first example, we submitted the LDSC-derived genetic correlation matrix of the 12 neuroticism items in UKB to EFA (see Supplementary Results). Based on these initial EFA results, follow-up CFAs (Supplementary Fig. 3) were specified using genomic SEM (standardized loadings > 0.4 were retained; Supplementary Table 2). The two-factor solution ($\chi^2(53) = 2,758.18; AIC = 2,808.18; CFI = 0.940; SRMR = 0.077$) and three-factor solution ($\chi^2(51) = 1,879.31; AIC = 1,933.31; CFI = 0.959; SRMR = 0.057$) both provided excellent fit to the data and exceeded the fit of the single, common factor model. Consistent with the superior model fit indices for the two- and three-factor solutions, only 28 and 20 of the 69 Q\textsubscript{SNP} hits from the single common factor model (described in further detail in the ‘SNP effects’ section below) continued to surpass genome-wide significance for the two- and three-factor models, respectively (Supplementary Fig. 4 and Supplementary Table 3). In addition, a GWAS of all HapMap3 SNPs for the two- and three-factor models revealed that the average size of Q\textsubscript{SNP} across all SNPs was largest for the common factor ($\chi^2(1) = 1.68$), followed by the two-factor ($\chi^2(1) = 1.64$) and three-factor models ($\chi^2(1) = 1.51$). Thus, heterogeneity indices of individual SNP effects in the GWAS data agree with model fit indices, with both favouring the three-factor model of neuroticism.

In the second example, EFA was applied to the LDSC-derived genetic correlation matrix for nine anthropometric traits from the Early Growth Genetics and Genetic Investigation of Anthropometric Traits consortia (Supplementary Table 4). EFA results indicated that 2 factors explained 61% of the total genetic variance. Moreover, a heat map of the genetic correlation matrix suggests two primary factors that index overweight and early-life growth phenotypes (Supplementary Fig. 5). A follow-up CFA (Supplementary Fig. 6) within genomic SEM was specified based on the EFA parameter estimates (standardized loadings > 0.25 were retained). The CFA showed good fit to the data ($\chi^2(25) = 12,994.71; AIC = 13,034.71; CFI = 0.962; SRMR = 0.092$). The results indicated highly significant loadings and a small genetic correlation ($r_g$) between the two factors ($r_g = 0.10$; s.e. $= 0.03$; $P < 0.001$). This indicates that early-life physical growth is modestly associated with later life obesity traits via genetic pathways.

Genetic multivariable regression (replicating genome-wide inferred statistics (GWIS)). Nieuwboer et al.\textsuperscript{15} used summary statistics for educational achievement\textsuperscript{1} and both schizophrenia and bipolar disorder\textsuperscript{19} to determine whether genetic correlations with educational achievement are driven by variation specific to either disorder. Educational achievement is genetically correlated with
SNP effects. Common factor models. A powerful application of genomic SEM is to include individual SNP effects in both the genetic covariance matrix and the sampling covariance matrix, to estimate the effect of a given SNP on the latent genetic factor(s). If the summary statistics are composed of M different SNPs, M models are estimated to obtain genome-wide summary statistics for the latent factor. As an example of genomic SEM used for multivariate GWAS, we incorporated SNP effects into the p-factor and neuroticism models presented above. Linkage disequilibrium-independent hits are defined below using a clumping threshold of \( r^2 < 0.1 \) in a 500-kilobase (kb) window, with the exception of a 1-megabase (Mb) window for chromosomes 6 and 8. A total of 128 independent loci were genome-wide significant for the p-factor \((P < 5 \times 10^{-8})\) Supplementary Figs. 8–10 and Figs. 1a and 2a). Of the 128 loci, 27 independent loci were not previously identified in any of the contributing univariate GWASs (Table 1 and Supplementary Table 5). Of these 27 loci, 5 were identified as either genome-wide significant or suggestive of significance \((P < 1 \times 10^{-4})\) in a separate, previously published GWAS of 1 of the 5 traits. A total of 118 loci were genome-wide significant for neuroticism, with 38 loci not identified in the univariate item-level GWASs (Supplementary Table 6 and Figs. 1b and 2b). Plots of item-level effects for individual SNPs revealed high consistency in magnitude and direction for SNPs identified as genome-wide significant for the common factors (Supplementary Fig. 11). Although there is early lift-off in the \( Q-Q \) plots for both common factors, LDSC analyses of the summary statistics produced by genomic SEM indicated that the results were not due to uncontrolled inflation for either the p factor \((\text{intercept} = 0.987; \text{s.e.} = 0.014)\) or neuroticism \((\text{intercept} = 0.997; \text{s.e.} = 0.001)\).

General trends. Mean \( \chi^2 \) statistics were higher for the genomic SEM-derived summary statistics of common factors relative to univariate indicators (Table 1). It is important to note here that, whereas genomic SEM may boost power in many cases, this is not the primary purpose of the method. Rather, it is to identify the relationship between SNPs and observed phenotypes as mediated through a user-specified model and to concurrently evaluate the construct validity of said model. Inspecting the distribution of univariate \( P \) values for the newly identified SNPs for the general factors indicated that these SNPs were generally characterized by relatively low \( P \) values, albeit not low enough to cross the genome-wide significance threshold for any individual phenotype (Supplementary Figs. 12 and 13).

\( Q_{\text{SNP}} \) results. The results revealed 1 and 69 independent \( Q_{\text{SNP}} \) loci for the p-factor and neuroticism, respectively (Fig. 2c,d and Supplementary Fig. 14). For neuroticism, significant \( Q_{\text{SNP}} \) estimates were obtained for SNPs that were highly significant for some traits but not others (Supplementary Table 7 and Supplementary Fig. 15). The association between \( P \) values for SNP effects and \( Q_{\text{SNP}} \) estimates were minimal (Supplementary Fig. 16). Comparing the \( Q_{\text{SNP}} \) estimates for SNPs identified as significant for only the p factor or neuroticism relative to SNPs identified as significant for one of the indicators, but not the common factor, indicated that the latter group of SNPs were characterized, as would be expected, by larger \( Q_{\text{SNP}} \) estimates (that is, greater heterogeneity in individual effects; Supplementary Fig. 17). Intercepts from LDSC analyses of the \( Q_{\text{SNP}} \) statistics also indicated that the results for the heterogeneity index were not attributable to inflation \((p \text{ factor}: \text{intercept} = 0.978; \text{s.e.} = 0.009; \text{neuroticism: intercept} = 0.963; \text{s.e.} = 0.009)\). Slopes from the same LDSC analyses further indicated genetic signal in heterogeneity \((p \text{ factor}: Z = 13.65; P = 6.68 \times 10^{-44}; \text{neuroticism: Z} = 30.23; P = 9.98 \times 10^{-20})\).

Comparison with multi-trait analysis of GWAS (MTAG). Existing multivariate methods use summary statistics of genetically correlated phenotypes to boost power for discovery and prediction for a particular trait\(^{10,20,22}\). Boosting power is only one application of genomic SEM. That said, a genomic SEM common factor GWAS approach has already been shown by an independent research group to perform comparably to existing multivariate approaches for out-of-sample prediction\(^1^\). Moreover, as a flexible modelling framework, genomic SEM may encompass other multivariate approaches. For example, we show mathematically that genomic SEM can be specified to satisfy the same moment conditions as MTAG\(^{11}\) (see Supplementary Methods). Simulation results also revealed near-perfect correspondence from a linear regression in which Z statistics from MTAG were used to predict those from a genomic SEM specified to satisfy the MTAG moment conditions (Supplementary Fig. 18; unstandardized slope = 0.999; intercept = 2.65 \times 10^{-4}).

Performance in empirical data under controlled missingness. We contrast estimates obtained from the common factor model of neuroticism described above with estimates for a GWAS with an imposed missing structure. We first transformed the binary scale neuroticism items into a smaller number of quantitative scores. To do so, we created three parcels of neuroticism items consisting of four items each with scores ranging from 0–4, at which point it is appropriate to treat the parcel as continuous\(^2\). Parcels were constructed based on the same EFA results described above and mirrored the composition of the three-factor model, with the exception that the irritability item was included with parcel 2 so as to have an equal distribution of four items per parcel. Of the 300,000 participants, 100,000 non-overlapping participants were removed from 2 of the 3 parcels for missing data models. The best-powered results (indexed by mean \( \chi^2 \) values) were for genomic SEM of the individual neuroticism items presented above, indicating that construction of composite indices via averaging, although convenient, removes multivariate information that can otherwise be retained with genomic SEM (Supplementary Table 8). Genomic SEM analyses that incorporated supplementary information from parcels containing imposed missing data consistently outperformed GWASs of individual parcels with complete data, and performed nearly as well as analyses of complete data across all three parcels. Thus, the inclusion of summary data from genetically correlated phenotypes in genomic SEM may boost power relative to GWASs of the individual phenotypes.
even when there is high sample overlap and sample sizes are uneven across phenotypes.

**Parcel comparison of $Q_{SNP}$.** Using the three constructed parcels without any missing data, the distribution of $P$ values was compared across SNPs with high ($P < 5 \times 10^{-8}$) and low ($P > 5 \times 10^{-3}$) $Q_{SNP}$ estimates from the item-level genomic SEM analysis of neuroticism for SNPs that were genome-wide significant in at least one of the parcels. These results indicated that, for SNPs with a higher $Q_{SNP}$ for the common factor, there was more discordance of effect sizes among three lower-order factors relative to SNPs that produced lower heterogeneity estimates (Supplementary Fig. 19). The average differences between the highest and lowest $-\log_{10}[P]$ values were 10.56 and 4.96 for high and low $Q_{SNP}$ respectively. This suggests that $Q_{SNP}$ is appropriately indexing discordance in SNP-level effects across genetically correlated indicators.

**Polygenic prediction.** We re-estimated the $p$ factor model using the summary statistics from the schizophrenia and MDD GWASs that did not overlap with the UKB dataset to predict psychiatric symptoms in UKB (see Supplementary Fig. 20 for the phenotypic model). To produce a reliable set of targets for polygenic prediction, and to focus our analyses on construct validation, latent factors of psychiatric symptoms were specified as the out-of-sample targets. We compared the magnitude of out-of-sample prediction for the $p$ factor PGSs predicting the phenotypic $p$ factor and factors of individual psychiatric domains relative to the prediction using PGSs derived from univariate summary statistics (Fig. 3 and Supplementary Table 9). The PGS for the genetic $p$ factor predicted more variance in symptoms of depression, psychotic experiences, mania, anxiety, PTSD and a phenotypic $p$ factor than any univariate PGS. For neuroticism, univariate PGSs were constructed in data from the Generation Scotland study using summary statistics for

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**Fig. 1** Genomic SEM solutions for $p$- and neuroticism-factor models with SNP effect. a, b, Standardized results from using genomic SEM (with WLS estimation) to construct a genetically defined $p$ factor of psychopathology (a) and a genetic neuroticism factor (b) with a lead independent SNP predicting the factors (rs4552973 for the $p$ factor and rs10497655 for neuroticism). Standard errors are shown in parentheses. For a model that was standardized with respect to the outcomes only, the effect of the SNP was $-0.093$ (s.e. $= 0.017$; SNP variance $= 0.252$) for the $p$ factor and $-0.042$ (s.e. $= 0.007$; SNP variance $= 0.432$) for neuroticism. This can be interpreted as the expected standard deviation unit difference in the latent factor per effect allele. The subscript g is used in these path diagrams to denote genetic variables. The $u$ variables reflect the residual variance in the genetic indicators not explained by the common factor. ANX, anxiety; BIP, bipolar disorder; Emb, worry too long after embarrassment; Fed-up, fed-up feelings; Feel, sensitivity/hurt feelings; Irr, irritability; SCZ, schizophrenia.
the 12 neuroticism items, genomic SEM factor of items, 3 neuroticism parcels, genomic SEM factor of parcels and neuroticism sum score. We used PGSSs to predict a sum score composed of the same neuroticism items administered in UKB. We also calculated mean \( \chi^2 \) values for each of these summary statistics, which we used to infer their relative power. Of all of the summary statistics considered, summary statistics derived from a genomic SEM analysis of a common factor of the neuroticism items produced the largest mean \( \chi^2 \) in the summary statistics and predicted the greatest variance in the out-of-sample phenotype (Supplementary Fig. 21). In both cases, the superior performance of genomic SEM analysis of the common factor of items relative to the sum score of the items is probably, in part, a reflection of the fact that the sum score in UKB was created using listwise deletion, resulting in a reduced sample size of 274,008. Conversely, genomic SEM uses all available information from neuroticism items, with sample sizes of ~325,000 each. In more severe cases of sample non-overlap, we would expect even larger power benefits of genomic SEM-derived summary statistics relative to individual items or sum scores. Indeed, in instances of minimal sample overlap, it is not possible to compute sum scores, but genomic SEM can still be used to integrate data across phenotypes.

**Biological annotation.** The biological function of the SNPs related to the \( p \) factor and neuroticism was examined using DEPICT\(^{25} \). Table 1 presents the number of enriched gene sets, prioritized genes, and enriched tissues and cell types across the univariate statistics and common factors (see Supplementary Tables 10–18 for detailed output). Common factors produced more informative results than the individual indicators. As expected, all of the tissue enrichment for the common factors was identified in the nervous system (Supplementary Fig. 22). Neuroticism-prioritized genes indicated a central role of synaptic activity (for example, \( STX1B, NR4A2 \) and \( PCLO \)), including glutamatergic neurotransmission (\( GRM3 \)). The \( p \) factor gene sets were largely characterized by communication between neurons (for example, ‘dendrite development’, ‘dendritic spine’ and ‘abnormal excitatory postsynaptic potential’). Biological annotation of \( Q_{\text{hap}} \) statistics for neuroticism indicated that genes within the 69 loci related to neuroticism, but not through a single factor, include: \( GRIA1 \), a glutamate receptor subunit (involved in signalling in excitatory neurons) that has previously been related to schizophrenia\(^{31} \), chronotype\(^{32} \) and autism\(^{33} \); and \( PCDH17 \), a gene involved in cellular connections in the brain that has been related to intelligence\(^{34} \).

**General guidelines.** When implementing genomic SEM, users should be aware of the limitations and assumptions of the method. First, because genomic SEM is a method for modelling genetic covariance matrices, it relies on the same assumptions as the method used to estimate genetic covariances, and best practices for implementing such a method should be followed. For example, when LDSC is used to construct the genetic covariance matrix, SNPs should not first be pruned for linkage disequilibrium, and summary statistics for different phenotypes should be obtained from ethnically homogeneous samples of similar ancestral backgrounds\(^1 \). With respect to selecting between competing models, users should take into account a variety of both absolute fit indices (for example, SRMR and model \( \chi^2 \)) and relative fit indices (for example, AIC and \( \chi^2 \) difference). We provide general standards for absolute model fit in the Methods. Finally, a formal power analysis should take into account specific characteristics of the summary data, genetic architecture of the phenotypes and model to be specified. This can typically be achieved with simulation. Generally speaking, we would expect power to detect SNP effects on a common genetic factor.
to increase when the constituent univariate GWASs have larger sample sizes, higher heritabilities, higher genetic correlations with one another and lower sample overlap with one another. That said, we still expect some power benefits relative to univariate GWASs when the constituent phenotypes are only moderately heritable and/or moderately genetically correlated and/or sample overlap is high. The choice of included summary statistics, phenotypes and model(s) will of course depend on the researcher’s objectives and the model(s) to be specified.

**Discussion**

Applications of genome-wide methods to data from large-scale population-based samples have uncovered clear evidence of pervasive statistical pleiotropy. Genomic SEM is a method for modelling the multivariate genetic architecture of constellations of genetically correlated traits and incorporating genetic covariance structure into multivariate GWAS discovery. In contrast with methods that model phenotypic rather than genetic covariance structure, and rely on raw data, genomic SEM employs summary GWAS data to model genetic covariance structure. Genomic SEM is computationally efficient, accounts for potentially unknown degrees of sample overlap and allows for flexible specification of covariance structure, such that several broad classes of structured covariance models can be applied. The genomic SEM approach shares benefits of some existing approaches for boosting power by combining information across genetically correlated phenotypes. However, genomic SEM uniquely allows one to compare different hypothesized genetic covariance architectures and to incorporate such architectures into multivariate discovery. Importantly, shared genetic liabilities across phenotypes can be explicitly modelled as factors that may be treated as broad genetic risk factors with equally broad downstream consequences. Multivariate genetic methods have existed for decades in the twin literature, with Martin and Eaves providing a framework for fitting structural equation models of genetic and environmental variance components to multivariate twin data. Using GWAS summary data from unrelated individuals, genomic SEM can be used to estimate multivariate genetic models similar to those from the existing twin literature. Moreover, genomic SEM offers new promise as a method that allows for modelling genetic covariance even among phenotypes for which phenotypic covariance cannot be estimated.

Genomic SEM is not the first method for multivariate GWAS. Other methods, such as MTAG, SHom/SHet, metaUSTAT, min-P and TATES allow researchers to perform multivariate meta-analyses based solely on summary data. The methods can generally be divided into two distinct classes: methods that aggregate test statistics or effect sizes based on a model (genomic SEM, SHom and MTAG) and those that select from the univariate P-values while taking care not to inflate type-I error (min-P, TATES and SHet). As we show with respect to MTAG, models on which existing methods are based may be fit within the genomic SEM framework. We also anticipate that the approaches for selecting the P-values from a set of analyses while maintaining proper type-I error control could be integrated into the genomic SEM framework. For instance, whereas TATES is currently applied to select P-values from

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**Table 1 | Summary of multivariate (genomic SEM) and univariate GWAS results**

| trait | Lead SNPs (P < 5 × 10⁻⁸) | Q₀ᵤᵤ hits | Unique hits | Number of gene sets | Number of prioritized genes | Number of tissues and cells | Mean χ² |
|------|--------------------------|------------|-------------|---------------------|---------------------------|-----------------------------|---------|
| p factor | Genomic SEM (WLS) 128 | 1 (1) | 27 | 71 | 37 | 24 | 1.88 |
| Schizophrenia | 127 | – | 34 (0) | 2 | 25 | 21 | 1.82 |
| Bipolar | 4 | – | 4 (0) | 0 | 0 | 0 | 1.15 |
| MDD | 5 | – | 5 (0) | 0 | 0 | 0 | 1.31 |
| PTSD | 0 | – | 0 (0) | 0 | 0 | 0 | 1.01 |
| Anxiety | 1 | – | 1 (0) | 0 | 0 | 0 | 1.03 |

| Neuroticism | Genomic SEM (WLS) 118 | 69 (5) | 38 | 1 | 19 | 20 | 1.64 |
| Mood | 43 | – | 19 (5) | 0 | 0 | 15 | 1.37 |
| Misery | 31 | – | 6 (4) | 0 | 0 | 0 | 1.32 |
| Irritability | 36 | – | 17 (4) | 0 | 0 | 0 | 1.37 |
| Hurt feelings | 24 | – | 11 (0) | 0 | 0 | 0 | 1.33 |
| Fed-up | 38 | – | 21 (6) | 0 | 0 | 0 | 1.36 |
| Nervous | 41 | – | 25 (12) | 0 | 0 | 0 | 1.36 |
| Worry | 56 | – | 26 (6) | 0 | 13 | 0 | 1.46 |
| Tense | 19 | – | 10 (3) | 0 | 0 | 0 | 1.32 |
| Embarrass | 17 | – | 6 (2) | 0 | 0 | 0 | 1.33 |
| Nerves | 12 | – | 7 (3) | 0 | 0 | 0 | 1.26 |
| Lonely | 6 | – | 4 (3) | 0 | 0 | 0 | 1.19 |
| Guilt | 21 | – | 8 (1) | 0 | 0 | 0 | 1.28 |

*Numbers in parentheses for Q₀ᵤᵤ report how many Q₀ᵤᵤ hits were in linkage disequilibrium with hits identified as significant for the common factor. Unique hits for the common factor refers to lead SNPs that were not in linkage disequilibrium with hits for the individual indicators. Unique hits for the individual indicators refer to hits for the respective indicator that were not in linkage disequilibrium with hits for the common factor. Unique hits for the common factor excluded hits in linkage disequilibrium with Q₀ᵤᵤ hits. For unique hits for indicators, values in parentheses indicate whether any of these hits were identified as significant for Q₀ᵤᵤ. For unique hits for the common factor, hits were excluded that were in linkage disequilibrium with previously reported indicator hits that were removed due to missing values across the other phenotypes. The single Q₀ᵤᵤ hit for WLS estimation of the p factor was significant for both the common factor and schizophrenia. For the common factor and the indicators, independent hits were defined using a pruning window of 5000 kb and r² > 0.1. For chromosomes 6 and 8, an additional pruning filter of 1 Mb and r² > 0.1 was used to account for long-range linkage disequilibrium due to the MHC region and pericentric inversion, respectively. For univariate statistics, we used only the SNPs present across all indicators to facilitate a direct comparison with genomic SEM results.**
In this equation, GW-SEM requires access to phenotypic data, which is a substantial limitation compared to GWAS, which relies on genotype summary statistics. However, GW-SEM is more flexible and user friendly than GWAS, and allows for the combination of items tapping genetically related phenotypes into a single structural model. This question may be of particular interest as the large degrees of phenotypic overlap between psychiatric traits makes it challenging to disentangle the genetic and environmental contributions to each trait individually.

PGSs were constructed using the same set of SNPs for all predictors. $R^2$ on the y-axis indicates the percentage of variance (possible range: 0–100) explained in the outcome unique of covariates. The summary statistics for genomic SEM were estimated using WLS. The genomic SEM-based PGS was derived from a model estimating SNP effects on a common $p$ factor, constructed from schizophrenia, bipolar, MDD, PTSD and anxiety (as in Fig. 1a). To prevent bias, the genomic SEM summary statistics were produced using schizophrenia and MDD GWAS summary statistics that did not include UKB participants. Error bars indicate 95% confidence intervals estimated using the delta method. Phenotypes were constructed for European participants in the UKB for five symptom domains.

Methods

Overview of genomic SEM. Genomic SEM is a two-stage structural equation modelling approach. In the first stage, the empirical genetic covariance matrix and its sampling covariance matrix are estimated. In principle, these matrices may be obtained using a variety of methods for estimating SNP heritabilities, methods for estimating genetic covariances and their joint estimation errors. Here, we use a novel version of LDSC that accounts for potentially unknown degrees of sample overlap by populating the off-diagonal elements of the sampling covariance matrix. The same strengths, as well as assumptions and limitations, that are known to apply to LDSC, apply to its extension used here and to genomic SEM. In stage 2, the user employs a multivariate system of regression and covariance associations involving the genetic components of phenotypes with one another and/or more general latent factors. These associations are represented by parameters that may be fixed or freely estimated, as long as the model is statistically identifiable (for example, the number of freely estimated parameters does not exceed the number of non-redundant elements in the genetic covariance matrix being modelled). A model with genetic parameters $(\Theta)$ is estimated such that the fit function indexing the discrepancy between the model-implied covariance matrix, $\Sigma(\Theta)$, and the empirical covariance matrix, $\Sigma$, estimated in stage 1 is minimized. Model fit is considered good when $\Sigma(\Theta)$ closely approximates $\Sigma$. In the main text, we highlight the results from weighted least squares (WLS) estimation that weights the discrepancy function using the inverse of the diagonal elements of the sampling covariance matrix, and produces model standard errors using the full sampling covariance matrix. In the Supplementary Results, we additionally report the results from an alternative normal theory maximum likelihood estimation method.

Form of structured covariance models. Genomic SEM provides substantial user flexibility with respect to the particular SEM that is specified to produce the model-implied covariance matrix $\Sigma(\Theta)$ that approximates the empirical covariance matrix, $\Sigma$. SEMs can be partitioned into two sets of equations—one describing the measurement model and the other describing the structural model. In the measurement model, the genetic components of $k$ ‘indicant’ phenotypes are described as linear functions of a smaller set of $m$ (continuous) latent variables, $\mathbf{y} = A \mathbf{q} + \mathbf{e}$. In this equation, $\mathbf{y}$ is a $k \times 1$ vector of indicators, $\mathbf{e}$ is a $k \times 1$ vector of residuals, $\mathbf{q} = m \times 1$ vector of latent variables and $A$ is $k \times m$ matrix of factor loadings (that is, regressions relating the latent variables to the set of indicators). In a typical application of genomic SEM, each indicator is a function of exactly one of the latent variables (although this so-called ‘simple structure’ restriction may be relaxed). In a CFA model, only the measurement model is specified, and the set of latent variables are allowed to freely co-vary. Thus, the model-implied covariance matrix of a CFA is $\Sigma(\Theta) = \Sigma(\mathbf{A}) + \Theta$, where $\Sigma(\mathbf{A})$ is an $m \times m$ latent variable covariance matrix and $\Theta$ is a $k \times k$ matrix of covariances among the residuals, $\epsilon$. Typically, $\Theta$ is diagonal, which implies that indicators are mutually independent conditional on the set of latent variables. That constraint may be relaxed such that select pairs of indicators are allowed to co-vary over and above their associations via the latent variable structure (that is, residual covariances are allowed). CFA models are typically used to assess the strength of relations between sets of indicators and their respective underlying latent variables, as well as to assess the fit of a measurement model to data. A well-fitting CFA model implies that the latent variable structure is able to account for the observed covariances among a set of indicator variables.

When a theory aims to explain associations among latent variables, a structural variable may be added to the measurement model to produce a full SEM. The structural model of a SEM relates latent variables to each other via directed regression coefficients. It can be written in matrix notation as $\mathbf{q} = \mathbf{B} \mathbf{z} + \mathbf{e}$, where $\mathbf{B}$ is an $m \times m$ matrix of regression coefficients that relate latent variables to each other, and $\mathbf{z}$ is an $m \times 1$ vector of latent variable residuals. The model-implied covariance matrix of observed variables is $\Sigma(\Theta) = \Sigma(\mathbf{B}) + \Psi(\mathbf{1} - \mathbf{B})^{-1} + \Theta$ where $I$ is an $k \times k$ identity matrix. Thus, in a full SEM, the empirical matrix is represented by a set of parameters that relate observed variables to latent variables, and latent variables to each other in a series of linear equations.

Path diagrams. SEMs can be represented graphically as path diagrams representing regression and covariance relations among variables. In path diagrams, observed variables are represented as squares and unobserved (that is, latent) variables are represented as circles. Regression relationships between variables are represented as one-headed arrows pointing from the independent variable to the dependent variable. Covariance relationships between variables are represented as two-headed arrows linking the two variables. The variance of a variable (that is, the covariance between a variable and itself) is represented as a two-headed arrow connecting the variable to itself. In genomic SEM, we represent the genetic component of each phenotype with a circle, as the genetic variance component is a latent variable that is not directly measured, but is inferred from LDSC (it is the phenotype itself that is observed in the raw data that is used to produce the summary statistics). SNP heritabilities are directly measured, and are therefore represented as squares. When all elements in a SEM are represented in a path diagram, the diagram contains the full system of algebraic equations needed to estimate the full set of SEM parameters, $\Theta$, and to produce the model-implied covariance matrix, $\Sigma(\Theta)$.

![Graphical representation of path diagrams](image-url)
Stage 1 estimation. In stage 1, the empirical genetic covariance matrix (\(S_{\text{LDSC}}\)) and its associated sampling covariance matrix (\(V_{\text{LDSC}}\)) are estimated using our multivariable extension of LDSC. \(S_{\text{LDSC}}\) is a \(k \times k\) symmetric matrix with SNP heritabilities on the diagonal and genetic covariances (\(\sigma_{\text{g,ij}}\)) between phenotypes \(i\) and \(j\) off the diagonal. The genetic covariance between phenotypes \(i\) and \(j\) can be computed as the genetic correlation scaled relative to the total genetic variance of each of the two contributing phenotypes (themselves scaled to unit variances), \(\rho_{\text{g,ij}} = \frac{\sigma_{\text{g,ij}}}{\sqrt{\Sigma_{\text{g,ii}} \Sigma_{\text{g,jj}}}}\). Thus, the genetic covariance matrix of order \(k\) has \(k(k + 1)/2\) non-redundant elements. It can be written as:

\[
S_{\text{LDSC}} = \begin{bmatrix}
\sigma_{\text{g,11}} & \sigma_{\text{g,12}} & \cdots & \sigma_{\text{g,1k}} \\
\sigma_{\text{g,21}} & \sigma_{\text{g,22}} & \cdots & \sigma_{\text{g,2k}} \\
\vdots & \vdots & \ddots & \vdots \\
\sigma_{\text{g,k1}} & \sigma_{\text{g,k2}} & \cdots & \sigma_{\text{g,kk}}
\end{bmatrix}
\]

To produce unbiased standard error estimates and test statistics, we require the sampling covariance matrix, \(V_{\text{LDSC}}\), of the LDSC estimates that is composed of all non-redundant elements in the \(S_{\text{LDSC}}\) matrix. Thus, it is a symmetric matrix of order \(k\), with \(k(k + 1)/2\) non-redundant elements. The diagonal elements of \(V_{\text{LDSC}}\) are sampling variances (that is, squared standard errors of the elements in \(S_{\text{LDSC}}\)). The off-diagonal elements of \(V_{\text{LDSC}}\) are sampling covariances that indicate the extent to which the sampling distributions of the variance and covariance estimates in \(S_{\text{LDSC}}\) co-vary with one another, as would be expected when there is overlap among the samples from which the terms are estimated. This \(V_{\text{LDSC}}\) matrix can be written as:

\[
V_{\text{LDSC}} = \begin{bmatrix}
\text{var}(h_1^2) & \text{cov}(h_1,h_2) & \cdots & \text{cov}(h_1,h_k) \\
\text{cov}(h_2,h_1) & \text{var}(h_2^2) & \cdots & \text{cov}(h_2,h_k) \\
\vdots & \vdots & \ddots & \vdots \\
\text{cov}(h_k,h_1) & \text{cov}(h_k,h_2) & \cdots & \text{var}(h_k^2)
\end{bmatrix}
\]

The diagonal elements of \(V_{\text{LDSC}}\) can be estimated using the jackknife resampling procedure in the bivariate version of LDSC that is currently available from its original developers\(^4\). The LDSC function introduced in the GenomicSEM software package expands the jackknife procedure to the multivariable context to additionally produce sampling covariances (which index dependencies among estimation errors) among the elements of \(S_{\text{LDSC}}\) needed to populate the off-diagonal elements of \(V_{\text{LDSC}}\).

Incorporating individual SNP effects. Several steps are needed to incorporate individual SNP effects into genomic SEM. The first step requires that the inputted genetic covariance matrix be expanded to include covariances between the SNP and each of the phenotypes, \(g_i\) through \(g_m\), by appending a vector of SNP phenotype covariances (\(S_{\text{SNP}}\)) to \(S_{\text{LDSC}}\):

\[
S_{\text{SNP}} = \begin{bmatrix}
\sigma_{\text{SNP}} \\
\sigma_{\text{SNP,1}} & h_1 \\
\sigma_{\text{SNP,2}} & h_2 \\
\vdots & \vdots \\
\sigma_{\text{SNP,kg}} & h_k
\end{bmatrix}
\]

The sampling covariance matrix, \(V_{\text{SNP}}\), associated with this expanded \(S_{\text{SNP}}\) covariance matrix, includes a number of components. One block of this \(V_{\text{SNP}}\) matrix, \(V_{\text{SNP}}^{\text{LDSC}}\), contains the sampling variances and sampling covariances of the latent genetic variances (SNP heritabilities) and genetic covariances, which are obtained from the multivariable LDSC approach introduced above. A second block of the \(V_{\text{SNP}}\) matrix, \(V_{\text{SNP}}^{\text{cov}}\), is composed of the sampling covariance matrix of the SNP effects on the phenotypes. The SNP variance (derived from reference panel data) is treated as fixed, and its sampling variance and sampling covariance with all other terms are fixed to 0 (or to a very small value to facilitate computational tractability). The sampling covariances of the SNP genotype covariances with one another are obtained using cross-trait LDSC intercepts (which represent sampling correlations weighted by sample overlap) after being rescaled relative to the sampling variances of the respective SNP genotype covariances\(^5\). A final block of the \(V_{\text{SNP}}\) matrix represents the sampling covariance of the SNP genotype covariances with the genetic variances and genetic covariances. These are fixed to 0, as sampling variation of the SNP genotype covariance is expected to be independent of the test statistics of all linkage disequilibrium blocks except the one it occupies. Because the sampling variance of the heritabilities and genetic correlations derive from sampling variability in the test statistics within all of the linkage disequilibrium blocks, their sampling covariances with a single SNP effect is expected to approach 0. The \(V_{\text{SNP}}\) matrix can be written in compact form as:

\[
V_{\text{SNP}} = \begin{bmatrix}
V_{\text{SNP}}^{\text{LDSC}} & 0 \\
0 & V_{\text{SNP}}^{\text{cov}}
\end{bmatrix}
\]

Stage 2 estimation. In stage 2, the genetic covariance matrix obtained in the previous stage, \(S\), is used to estimate the parameters in a SEM. In this stage, we allow for both WLS and normal theory maximum likelihood estimators. WLS does not strictly require positive definite \(S\) and \(V\) matrices, but may still benefit from positive definiteness during optimization. Maximum likelihood estimation requires both \(S\) and \(V\) to be positive definite. The GenomicSEM software package therefore smooths \(S\) and \(V\) to the nearest positive definite matrices before stage 2 estimation using the R function nearPD\(^6\).

The fit function minimized in the diagonally weighted version of WLS estimation that is standard in the GenomicSEM software package is the following:

\[
F_{\text{WLS}}(\theta) = \log |\Sigma(\theta)| - |S| + \text{tr}(|S|^{-1} \Sigma(\theta)) - k
\]

where \(\Sigma(\theta)\) is the covariance matrix implied by the set of parameter estimates. Note that, while the formulation of the maximum likelihood fit function does not explicitly include a weight matrix, it is asymptotically equivalent to a more general formulation that is identical to the WLS fit function, with \(\Sigma^j(\Sigma^{-1}(\theta) \otimes \Sigma^{-1}(\theta))D_j\), where \(D_j\) is the duplication matrix of order \(k\), in place of \(D\). Thus, the difference between maximum likelihood and WLS estimation can be construed as a difference in weight matrices only. A comparison between maximum likelihood and WLS results can be found in the Supplementary Results (see also Supplementary Figs. 23–27 and Supplementary Table 19).

WLS estimation more heavily prioritizes reducing misfit in those cells in the \(S\) matrix that are estimated with greater precision. This has the desirable property of potentially decreasing sampling variance of the genomic SEM parameter estimates, which may boost power for SNP discovery and increase polygenic prediction. However, because the precision of cells in the \(S\) matrix is contingent on the sample sizes for the contributing univariate GWASs, WLS may produce a solution that is dominated by the patterns of association involving the most well-powered GWASs, and contain substantial local misfit in cells of \(S\) that are informed by lower-powered GWASs. In other words, WLS relative to maximum likelihood may more heavily prioritize minimizing sampling variance of the parameter estimates in the so-called variance bias tradeoff\(^7\). We expect that this will only occur when the model is overidentified (that is, \(d.f. > 0\)), such that exact fit cannot be obtained, and that divergence in WLS and maximum likelihood estimates will be most pronounced when there is lower sample overlap and the contributing univariate GWASs differ substantially in power. Maximum likelihood estimation may be preferred when the goal is to most evenly weight the contribution of the univariate sample statistics.

Both WLS and maximum likelihood fit functions will produce consistent estimates of the model parameters when the model is true\(^8\). However, the ‘naïve’ standard errors and fit statistic produced in stage 2 estimation will be incorrect, because neither estimator uses the full \(V\) matrix in estimation. Thus, robust corrections must be applied to produce consistent estimates of standard errors and test statistics. The correct sampling covariance matrix of the stage 2, genomic SEM parameter estimates (that is, \(V\)) can be obtained using a sandwich correction\(^9\):

\[
V = (\hat{\Delta}^{-1} \hat{\Sigma} \hat{\Delta}^{-1})^{-1} \hat{\Sigma} \hat{\Delta}^{-1} \hat{\Sigma}^{-1} \hat{\Delta}^{-1}
\]

where \(\hat{\Delta} = \frac{\hat{\Delta}}{\hat{\Delta}}\) is the matrix of model derivatives evaluated at the parameter estimates, \(\hat{\Sigma}\) is the naïve stage 2 weight matrix that takes its form depending on the estimation method used (WLS or maximum likelihood), and \(\hat{\Sigma}\) is the sampling covariance matrix of \(S\) obtained using multivariable LDSC.

It may not always be possible to obtain the full sampling covariance matrix, \(V\). For example, for highly sensitive data, only the matrix \(S\) and the standard errors
of its elements may be available (that is, the diagonal of \( V \)). However, we note that when there is low sample overlap across the GWASs for each phenotype, off-diagonal elements of the sampling covariance matrix are small and pragmatically ignorable. Moreover, in other contexts with complete sample overlap, standard error inflation of the SEM parameters estimated using diagonally weighted versions of WLS has been estimated to be less than 8% without robustness corrections, and 0% with robustness corrections\(^{26}\).

**Standardization and scaling of summary statistics for multivariate GWASs.** Typically, GWAS summary statistics for quantitative phenotypes are not reported in terms of covariances, but are reported as ordinary least squared (OLS) unstandardized regression coefficients, with the phenotypes standardized before analyses (that is, the coefficients are standardized with respect to the outcome, but not the predictor). To transform the partially standardized regression coefficient (\( b_{SNP, 1} \)) of SNP effect on phenotype \( P \) to a covariance model, we multiply by the variance of scores on the SNP. The variance (\( \sigma^2_{SNP} \)) of scores (0, 1, 2) of a biallelic autosomal SNP is estimated as \( 2qg \), assuming Hardy–Weinberg equilibrium, where \( p \) is the minor allele frequency (MAF) and \( q = 1 − MAF \), with the MAF typically obtained from a reference sample. As the latent genetic factors estimated in LDSC are scaled relative to unit-variance-scaled phenotypes (by virtue of the SNP heritability estimates being placed on the diagonal of \( S \)), no further scaling is needed to transform this SNP phenotype covariance into a SNP genotype covariance.

When OLS regression coefficients and standard errors are provided from an analysis in which the phenotype has not been standardized before analyses, or only Z statistics or Pvalues (for which Z statistics can be readily obtained) are provided, the partially standardized regression coefficients and their standard errors can be obtained as:

\[
Z = b_{SNP, 1} \cdot \frac{\sigma_{SNP}}{\sigma_{P}},
\]

where \( b_{SNP, 1} \) is the equal to the regression coefficient for the OLS GWAS of the unstandardized phenotype. These derived partially standardized coefficients are then transformed into covariances by multiplying by the variance of scores on the SNP, as above.

In the case of UKB, where the GWAS summary statistics are available for logistic regression coefficients for liabilities for categorical outcomes (for example, case/control status) on the SNP, the logistic regression coefficients can be transformed into covariances as above, by multiplying by the SNP variances. However, it is appropriate to further transform the coefficients and their standard errors such that they are scaled relative to unit-variance-scaled liabilities. This can be achieved by dividing by \( \sqrt{\sigma_{SNP}^2 \cdot \logit \left( \frac{1}{p} \right)} \) as a logistic regression model implies a residual variance of 2. If GWAS summary statistics are reported for odds ratios, they can be transformed to logistic regression coefficients by taking their natural logarithm. Standard errors for the logistic regression coefficient are obtained as \( s.e. = 1 / (\sigma_{P} \cdot \logit \left( \frac{1}{p} \right)) \), where \( OR \) is the odds ratio. The derived logistic coefficients and their standard errors should further be transformed such that they are scaled relative to unit-variance-scaled liabilities, as above. Note that when the outcomes are categorical, the liability scale heritabilities and genetic covariances from multivariate LDSC (and not what are referred to as the ‘observed scale’ heritabilities and genetic covariances) should be used to populate the \( S \) matrix. This has the desirable property of both modelling the continuous scale of risk in the population and providing estimates that are independent of the observed prevalence of the categorical outcomes.

On occasion, summary statistics will be provided from OLS GWASs of categorical outcomes (for example, case/control status). Such an analysis is sometimes referred to as a linear probability model, as it (incorrectly) assumes that the association between the predictor and the probability of being in the comparison (for example, case) group relative to the reference (for example, control) group is linear. Parameters from the linear probability model are probability model results, and are freely estimated. Residual variances are estimated in step 2 by estimating the variances of \( k \) residual factors defined by the indicators. This provides an estimate of the discrepancy between the model-implied and observed covariance matrices, \( R = S − \Sigma(\theta) \), along with the sampling covariance matrix (\( V_0 \)) of \( R \).

While the discrepancy between model-implied and observed covariance matrices can be computed simply by deriving covariance expectations from the step 1 model and subtracting the observed covariance matrix, such an approach would not provide the corresponding \( V_0 \) matrix necessary for the calculations below. The \( V_d \) matrix is expected to be positive semidefinite and, consequently, have no negative eigenvalues. Therefore, the \( V_d \) matrix has the following eigendecomposition:

\[
V_d = (P_d \cdot P_d') \cdot E \cdot P_d,-N(0, I),
\]

where \( P_d \) is a matrix of principal components (eigenvectors) of \( V_d \), and \( E \) is a corresponding diagonal matrix consisting of non-zero eigenvalues. \( P_d \) reflects the null space of \( V_d \). Projecting \( R \) —a vector of residual covariances estimated from the step 2 model—onto \( P_d \), and adjusting for corresponding eigenvalues, we have:

\[
R = P_d \cdot E^{-1/2} \cdot P_d^{-1} \cdot R \cdot E^{-1/2} \cdot P_d \cdot P_d^{-1}
\]

where \( P_d \) is a matrix of principal components (eigenvectors) of \( V_d \), and \( E \) is a corresponding diagonal matrix consisting of non-zero eigenvalues. \( P_d \) reflects the null space of \( V_d \). Projecting \( R \) —a vector of residual covariances estimated from the step 2 model—onto \( P_d \), and adjusting for corresponding eigenvalues, we have:

\[
R = P_d \cdot E^{-1/2} \cdot P_d^{-1} \cdot R \cdot E^{-1/2} \cdot P_d \cdot P_d^{-1}
\]

This equation produces a test statistic that \( x^2 \) distributed with degrees of freedom (\( r \)) equal to the difference between the number of non-redundant elements (\( k \)) in the empirical covariance matrix (\( S \)) and the number of freely estimated parameters in the proposed model. The CFI is a test of approximate model fit. CFI indexes the extent to which the proposed model fits better than a model that allows all phenotypes to be heritable, but assumes that they are genetically uncorrelated. The \( x^2 \) statistic can be used to calculate CFI by calculating a second \( x^2 \) statistic for a so-called independence model (that is, a model that estimates genetic variances of all phenotypes but assumes all genetic covariances to be zero, such that \( \Sigma(\theta) = 0 \)). CFI is calculated using the formula below\(^{27}\), with \( f = x^2 \) —d.f.:

\[
f = \int (\text{independence model}) - f \text{ (proposed model)} \]

\[
f = \text{independence model})
\]

For the \( x^2 \) value of the independence model, a model is estimated in step 1 that includes only the variance of the indicators and no common factor. In step 2, these variances are fixed and the covariances among the indicators and variances of \( k \) residual factors defined by the indicators are estimated and used to populate the sampling equation above used to calculate the proposed model \( x^2 \). CFI values theoretically range from 0 to 1, with higher values indicating good fit. CFI values of \( \geq 0.90 \) are typically considered acceptable fit, and values of \( \geq 0.95 \) are typically considered good model fit\(^{27}\). When the empirical covariance matrix contains a large number of cells that are very close to 0, CFI values may be low, even when such cells are approximated well by the model.
AIC is a relative fit index that balances fit with parsimony, and can be used to compare models regardless of whether they are nested. AIC is calculated as:

$$AIC = \chi^2 + 2k$$

where $p$ is the number of free parameters in the model. Lower AIC values are considered superior.

SRMR is an index of approximate model fit that is calculated as the standardized root mean squared difference between the model-implied and observed correlations in $\Sigma(\theta)$ and $\Sigma$, respectively. Higher SRMR values indicate a larger discrepancy between $\Sigma(\theta)$ and $\Sigma$. It is positively biased, with larger bias resulting when the contributing univariate GWAS samples are lower powered. SRMR values below 0.10 indicate acceptable fit, values less than 0.05 indicate good fit, and a value of 0 indicates perfect fit.

We recommend that model-fit indices be considered concurrently, as individual indices each have their own strengths and limitations. Model $\chi^2$ is an index of exact fit, with lower values indicating better fit. Model $\chi^2$ may oftentimes be statistically significant, indicating that the model-implied genetic covariance matrix significantly differs from the empirical (unrestricted) genetic covariance matrix, even when the model-implied covariance matrix very closely approximates the empirical genetic covariance matrix. Oftentimes, models that closely (albeit imperfectly) approximate the empirical genetic covariance matrix may be scientifically and inferentially useful. We thus recommend considering CFI and SRMR indices of absolute fit, even when model $\chi^2$ is significant. We also recommend using the SRMR as a relative fit index to compare covariates (that is, different models fit to genetic covariance matrices derived from the exact same summary data for the exact same phenotypes). When models are nested, their respective $\chi^2$ values can be subtracted from one another to calculate a $\chi^2$ difference test, with d.f. equal to the difference in d.f. between the two models. This $\chi^2$ difference test indexes the extent to which the less complex model (that is, the model with fewer $\theta$) approximates the empirical genetic covariance matrix significantly worse than the more complex model (that is, the model with fewer d.f.). If the $\chi^2$ difference test is significant, the more complex model should be chosen. If the $\chi^2$ difference test is not significant, the less complex model should be chosen, as it is more parsimonious and approximates the empirical genetic covariance matrix no worse than the more complex model. Two models are nested when the set of possible model-implied covariance matrices from one model is a subset of the set of possible model-implied covariance matrices of the second model. Nesting can typically be confirmed if the less restrictive model can be derived from the more restrictive model by dropping or fixing parameters. Regardless of whether models are nested, they can be compared on CFI, SRMR and AIC, so long as the same data are being modelled.

**Simulation of factor structure.** To evaluate the ability of genomic SEM to capture the genetic factor structure in the generation of the population, the GCTA package was used to generate 100 sets of 6 independent, 100% heritable phenotypes (orthogonal genotypes) to pair with genotypic data for 39,909 randomly selected, unrelated individuals of European descent from UKB data for 1,209,498 SNPs present in HapMap3. The generating list of causal SNPs was set to 10,000 for all 600 genotypes, with the specific list of causal variants sampled with replacement from the 1,209,498 SNPs. One of the six orthogonal genotypes per set was designated an index of the general genetic factor, and the remaining five were designated indices of specific genetic factors. All of the orthogonal genotypes were scaled to $M=0$ and $\sigma=1$. For the two correlated genotypes were then constructed, each as the weighted linear combination of the general genetic factor and one domain-specific genetic factor. Weights for the general genetic factor were $\lambda g_{30} = 0.70$, 0.60, 0.50, 0.40 and 0.30 for correlated genotypes 1–5, respectively. Weights for the domain-specific factors were $\lambda g_{30} = (1-\lambda g_{30})$. Phenotypes were then each constructed as the weighted linear combination of one of the correlated genotypes and domain-specific environmental factors (randomly sampled from a normal distribution with $M=0$ and $\sigma=1$). Heritabilities for phenotypes 1–5 were set to $h^2 = 35$, 40, 50, 60 and 70%, respectively, such that the weights for the phenotypes were $\lambda g_v$ and the weights for the environmental factors were $\sqrt{(1-h^2)}$. We chose these choices to stabilize the properties of the distributions across simulations at 100 replications with $N=39,000$ each. We expect that with lower SNP $h^2$ values, the same patterns would hold, albeit at larger sample sizes. Each of the 500 phenotypes (100 sets of 5 phenotypes) was then analysed as a univariate GWAS in PLINK to produce univariate GWAS summary statistics. Our multivariable LDSC function was then used to construct 100 sets of 5 × 5 genetic covariance matrices ($\Sigma$) and associated sampling covariance matrices ($\Sigma_{sampl}$) for each simulation.

### Genomic SEM simulations

**Validation of summary-based model fit statistics via simulation.** A generating population with a common factor model defined by 4, 5 or 6 indicators was used to examine the null distribution of the newly derived $\chi^2$ test statistic using a set of 1,000 simulations per model. These simulations did not include individual genotypes, and were simulated solely based on a generating factor structure. For the 6-indicator models, the standardized factor loadings in the generating population were 0.42, 0.64, 0.22, 0.59, 0.19 and 0.64. The four- and five-indicator models specified the same factor loadings, excluding the last, or last two loadings, respectively. The results indicated that the two-step procedure described above produced a test statistic equivalent to the $\chi^2$ statistic calculated by lavaan from the raw data (Supplementary Fig. S1 and Supplementary Table S2). For a $\chi^2$-distributed test statistic, the mean of the null sampling distribution should match the d.f. test statistic. The d.f. test statistic was calculated as $\chi^2 = \sum_{k=1}^{K} \chi^2_{sampl}$, where each $\chi^2_{sampl}$ is the d.f. test statistic computed to approximate a $\chi^2$ distribution with an average approaching the d.f. (Supplementary Fig. S2). Calculated CFI values were also highly consistent with those observed using the CFI statistic provided by lavaan when using raw data (Supplementary Fig. S3 and Supplementary Table S2). Calculated AIC values were not contrasted with those obtained using the lavaan package in R. The software uses a formula that includes a log-likelihood estimate contingent on the provided sample size.

**Null distribution of $Q_{hmm}$.** To verify that the null distribution for $Q_{hmm}$ is $\chi^2$ distributed, a set of simulations specified a generating population in which the direct effects of the SNP on the indicators were entirely mediated through the common factor. Each simulation included 1,000 datasets, with $N=100,000$ completely overlapping participants per dataset. All simulated datasets were analysed using both WLS and maximum likelihood. We examined $Q_{hmm}$ for $F=1$ factor and $k=4$, 5 or 6 phenotypes. Supplementary Table 21 presents descriptive statistics for $Q_{hmm}$. Using a genome-wide significance threshold, in all cases, the false discovery rate for $Q_{hmm}$ was 0, and the power to detect a SNP effect on the common factor was 1. Both WLS and maximum likelihood estimation produced mean values of $Q_{hmm}$ that were approximately equal to the d.f. of the corresponding model. Supplementary Fig. S4 depicts the null sampling distributions of $Q_{hmm}$ estimated using WLS or maximum likelihood. Supplementary Fig. S5 plots $Q_{hmm}$ from these two estimation methods against $\chi^2$ distributions and against one another. These results indicate that both estimation methods produce results that are approximately $\chi^2$ distributed.

**Simulation of partial sample overlap.** To examine the effect of sample overlap on estimates obtained from genomic SEM, the GCTA package was used to generate a 50% heritable, quantitative phenotype with 30,000 causal SNPs. The phenotype was paired with genetic data from 100,000 randomly selected, unrelated individuals of European descent from UKB data for 1,209,498 SNPs. Three sets of 60,000 participants each were created using this same phenotype, with 40,000 participants identified using genomic SEM and 20,000 participants unique to each phenotype (that is, 100,000 total participants). These three subsamples were individually analysed in PLINK to produce univariate GWAS summary statistics. The multivariable LDSC function was then used to construct the genetic covariance and sampling covariance matrix using the three sets of summary statistics, and genomic SEM was used to fit a one-factor model with the SNP predicting the distribution of factor. Two key results were verified at this stage. First, we confirmed that the standardized factor loadings on the common factor were 1 for the identical phenotypes. Second, we verified that the bivariate linkage disequilibrium score intercepts used to account for sample overlap in the sampling covariance matrix were as expected. The equation for the linkage disequilibrium score (LDSC) intercepts below is $N_i = N_s \times (h^2)$, where $N_i$ is the sample overlap, $\rho$ is the LDSC correlation, $N_s$ is the sample size of trait 1, and $N_t$ is the sample size of trait 2. In this simulation, we observed bivariate intercepts of 0.67, which is as expected given sample overlap of 40,000, a phenotypic correlation of 1 and sample sizes of 60,000 (that is, $40,000 \times 1/\sqrt{60,000 \times 60,000} = 0.67$). Finally, estimates
from this multivariate GWAS were compared with estimates from the univariate GWAS in PLINK for the full set of 100,000 participants. If sample overlap is not appropriately accounted for in this example, such that data are incorrectly treated as derived from independent participants (as opposed to 100,000 total participants), we would expect the \( Z \) statistics for the SNP effects from genomic SEM to be upwardly biased relative to those from a univariate GWAS applied directly to the single phenotype in the 100,000 participants. We observed no such bias. A linear regression of \( Z \) statistics from genomic SEM from the 3 overlapping samples of 60,000 participants each predicted univariate GWAS \( Z \) statistics in the complete sample (of 100,000 participants) revealed near-perfect correspondence (unstandardized slope = 1.003; intercept = −0.003).

**MTAG simulation.** To evaluate the relationship between estimates from MTAG and those from a genomic SEM formulation of the MTAG model, we specified a bivariate system of heritable phenotypes, A and u. Phenotype u was constructed separately using the GCTA package, and also specified to be 60% heritable and affected by a random selection of 30,000 HapMap3 SNPs. Phenotype u was constructed separately using the GCTA package, and also specified to be 60% heritable and affected by a random selection of 30,000 HapMap3 SNPs. Both A and u were standardized (\( V_A = V_u = 1.0 \)). Phenotype A was constructed from phenotypes A and u according to the equation \( B = 0.7A + 0.7u \). This procedure resulted in 60% heritabilities for both traits A and B, with a genetic correlation of 0.7 between them. Sample sizes for phenotypes A and B were 25,000 each, with 10,000 participants contributing data for both phenotypes A and B (that is, 40% sample overlap), such that the full dataset was released to 40,000 unique individuals in total. Both MTAG and a genomic SEM model specified to satisfy the same moment conditions as MTAG were used to obtain the MTAG moment conditions were compared. Results indicated near-perfect correspondence from the MTAG and genomic SEM. The moment conditions as MTAG (see Supplementary Methods) were then each run in 60% heritabilities for both traits A and B, with a genetic correlation of 0.7, and each of the individual domains—specified as latent variables. PGS variables were specified to directly predict the latent phenotypes within the model (that is, factor score estimates were not used). To construct PGSs, we removed from both the p-factor and univariate summary statistics the 3 SNPs that were identified as having genome-wide significant \( Q_{PGS} \) estimates for maximum likelihood, along with SNPs that were in linkage disequilibrium with these SNPs using an \( r^2 \) threshold of 0.1 and a 500-kb window. PGSs were constructed using PRSice\(^2\), with linkage disequilibrium clumping set to \( r^2 > 0.25 \) over 250-kb sliding windows. PGSs for the p-factor were based on the WLS summary statistics produced using genomic SEM. We ran PGS analyses using a \( P \) value threshold of 1.0 (that is, we used all available SNPs removed from those removed due to \( Q_{PGS} \) analyses). To maintain comparability, PGSs for the univariate summary statistics were constructed based on the same SNPs with which the PGSs for the p-factor were constructed. In the confirmatory factor models, we included controls for age, sex, genotyping array and 40 principal components of ancestry in conjunction with the PGS predictor.

**Neuroticism.** The raw total on the 12-item neuroticism subtest of the Eysenck Personality Questionnaire-Revised\(^{523}\) (maximum score = 12) was used as the target phenotype for out-of-sample prediction. Both genetic and neuroticism target data were available on 19,876 European participants in the Generation Scotland cohort\(^{10} \). After removing sex and 2016-2017 participants with uncertainty of ancestry before examining out-of-sample prediction. PGSs were constructed using PRSice\(^{2}\), with linkage disequilibrium clumping set to \( r^2 > 0.25 \) over 250-kb sliding windows and using a \( P \) value threshold of 1.0. PGSs for neuroticism were based on the WLS summary statistics produced using genomic SEM. Regression analyses were run using the lme4 function within the coxme package in R with a random intercept to account for nesting of individuals within families.

**Clumping and biological annotation.** Lead SNPs for univariate indicators and the common factors were identified using the clumping algorithm in PLINK\(^{52}\). We defined linkage disequilibrium-independent SNPs using an \( r^2 \) threshold of 0.1 and a 500-kb window using the same 1000 Genomes Phase 3 reference panel used for obtaining MAF. For chromosomes 6 and 8, an additional pruning filter was used of 1 Mb and \( r^2 > 0.1 \) to account for long-range linkage disequilibrium due to the MHC region and pericentric inversion, respectively. Increasing the pruning window further to 4 Mb did not influence our findings on chromosome 6 or 8. The lead SNPs identified using PLINK were entered into DEPCT. Prioritized genes, enriched gene sets and enriched tissues were identified using the standard false discovery rate of 5%.
Code availability
GenomicSEM software is an R package that is available from GitHub at https://github.com/MichelNivard/GenomicSEM. The GenomicSEM R package can be installed directly at https://github.com/MichelNivard/GenomicSEM/wiki. Examples of GenomicSEM code, including code used to produce the results, is provided for each set of analyses at https://github.com/MichelNivard/GenomicSEM/wiki.

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References
1. Lee, S. H. et al. Genetic relationship between five psychiatric disorders estimated from genome-wide SNPs. Nat. Genet. 45, 984–994 (2013).
2. Bush, W. S., Crawford, M. T. & Crawford, D. C. Unravelling the human genome–phenome relationship using phenome-wide association studies. Nat. Rev. Genet. 17, 129–145 (2016).
3. Yang, J., Lee, S. H., Goddard, M. E. & Visscher, P. M. GCTA: a tool for genome-wide complex trait analysis. Am. J. Hum. Genet. 88, 76–82 (2011).
4. ReproGen Consortium et al. An atlas of genetic correlations across human diseases and traits. Nat. Genet. 47, 1236–1241 (2015).
5. Barban, N. et al. Genome-wide analysis identifies 12 loci influencing human reproductive behavior. Nat. Genet. 48, 1462–1472 (2016).
6. Jansen, P. R. et al. Genome-wide analysis of insomnia (N=1,331,010) identifies novel loci and functional pathways. Nat. Genet. 51, 394–403 (2019).
7. Wray, N. R. et al. Genome-wide association analyses identify 44 risk variants and refine the genetic architecture of major depression. Nat. Genet. 50, 668–681 (2018).
8. Okbay, A. et al. Genomic association study identifies 74 loci associated with educational attainment. Nature 533, 539–542 (2016).
9. Verhulst, R., Maes, H. H. & Neale, M. C. GW-SEM: a statistical package to conduct genome-wide structural equation modeling. Behav. Genet. 47, 345–359 (2017).
10. Beaumont, R. N. et al. Genome-wide association study of offspring birth weight in 86,577 women identifies five novel loci and highlights maternal genetic effects that are independent of fetal genetics. Hum. Mol. Genet. 27, 742–756 (2018).
11. Turley, P. et al. Multi-trait analysis of genome-wide association studies using summary statistics. Am. J. Hum. Genet. 90, 229–237 (2012).
12. Cheung, M. W.-L. metaSEM: an R package for meta-analysis using structural equation modeling. Front. Psychol. 5, 1521–1532 (2015).
13. Savalei, V. & Bentler, P. M. A two-stage approach to missing data: theory and application to auxiliary variables. Struct. Equ. Modeling 16, 477–497 (2009).
14. Yuan, K. H. & Bentler, P. M. Robust mean and covariance structure analysis through iteratively reweighted least squares. Psychometrika 65, 43–58 (2000).
15. Browne, M. W. Asymptotically distribution-free methods for the analysis of covariance structures. Br. J. Math. Stat. Psychol. 37, 62–83 (1984).
16. Huedo-Medina, T. B., Sánchez-Meca, J., Marín-Martínez, F. & Botella, J. Assessing heterogeneity in meta-analysis: Q statistic or I² index. Psychol. Methods 11, 193–220 (2006).
17. Aspar, A. & Morfett, T. E. All for one and one for all: mental disorders in one twin, phenotypes in the other twin. Clin. Psychol. Sci. 2, 119–137 (2013).
18. Psychometrica 65, 43–58 (2000).
19. Flora, D. B. & Curran, P. J. An empirical evaluation of alternative methods of estimation for confirmatory factor analysis with ordinal data. Psychol. Methods 9, 466–491 (2004).
20. Baselmans, B. M. et al. Multivariate genome-wide analyses of the well-being spectrum. Nat. Genet. 51, 445–451 (2019).
21. Bates, D., Maechler, M., Davis, T. A., Oehlschlägel, J. & Riedy, M. matrix: Sparse linear algebra. R package version 1.2-12 (2017).
22. Zhu, X. et al. Meta-analysis of correlated traits via summary statistics from linear mixed model association on all-or-none genetic effects that are independent of fetal genetics. Hum. Mol. Genet. 27, 742–756 (2018).
23. Van der Sluis, S., Posthuma, D. & Dolan, C. V. TATES: efficient multivariate genotype–phenotype analysis for genome-wide association studies. PLoS Genet. 9, e1003325 (2013).
24. Allegretti, A. et al. Genomic prediction of cognitive traits in childhood and adolescence. Preprint at biorxiv https://www.biorxiv.org/content/10.1101/418210v1 (2018).
25. Sneller, M. et al. Psychiatric genetics and the structure of psychopathology. Mol. Psychiatry 24, 499–520 (2019).
26. Stockl, J. et al. Mood, anxiety and psychotic phenomena measure a common psychopathological factor. Psychol. Med. 45, 1483–1493 (2015).
27. Seed, C. et al. Hair: An Open-Source Framework for Scalable Genetic Data. Neale Lab http://www.nealelab.is/blog/2017/7/19/scaling-gwas-of-thousands-of-phenotypes-for-large-scale-replication (2017).
28. Nieuwenhoo, H. A., Pool, R., Dolan, C. V., Boomsma, D. I. & Nivard, M. G. GWIS: genome-wide inferred statistics for functions of multiple phenotypes. Am. J. Hum. Genet. 99, 917–927 (2016).
29. Ritveld, C. A. et al. GWAS of 126,559 individuals identifies genetic variants associated with educational attainment. Science 6139, 1467–1471 (2013).
30. Raderfer, D. M. et al. Polygenic dissection of diagnosis and clinical dimensions of bipolar disorder and schizophrenia. Mol. Psychiatry 19, 1017–1024 (2014).
31. Maier, R. M. et al. Improving genetic prediction by leveraging genetic correlations among human diseases and traits. Nat. Commun. 9, 989–993 (2018).
32. Kraus de Suis, S., Posthumus, D. & Dolan, C. V. TATES: efficient multivariate genotype–phenotype analysis for genome-wide association studies. PLoS Genet. 9, e1003325 (2013).
33. Algrenini, A. et al. Genomic prediction of cognitive traits in childhood and adolescence. Preprint at biorxiv https://www.biorxiv.org/content/10.1101/418210v1 (2018).
34. Rieppel, S. et al. A multi-analysis of genome-wide association studies for major depressive disorder. Mol. Psychiatry 18, 497–511 (2013).
35. Rieppel, S. et al. Biological insights from 108 schizophrenia-associated genetic loci. Nature 511, 421–427 (2014).
36. Muthén, L. K. & Muthén, B. O. Mplus: The Comprehensive Modeling Program for Applied Researchers. Version 7.3. https://www.statmodel.com/download/usersguide/MplusUserGuideVer_7.pdf (Muthén & Muthén, 2014).
61. Euesden, J., Lewis, C. M. & O’Reilly, P. F. PRSice: polygenic risk score software. *Bioinformatics* **31**, 1466–1468 (2014).

62. Eysenck, S. B., Eysenck, H. J. & Barrett, P. A revised version of the psychoticism scale. *Pers. Individ. Diff.* **6**, 21–29 (1985).

63. Smith, B. H. et al. Cohort profile: Generation Scotland: Scottish Family Health Study (GS:SFHS). The study, its participants and their potential for genetic research on health and illness. *Int. J. Epidemiol.* **42**, 689–700 (2012).

64. Rossel, Y. lavaan: An R package for structural equation modeling and more. Version 0.5–12 (BETA) http://users.ugent.be/~yrosseel/lavaan/lavaanIntroduction.pdf (2012).

65. Neale, M. C. et al. Cohort profile: Generation Scotland: Scottish Family Health Study (GS:SFHS). The study, its participants and their potential for genetic research on health and illness. *Int. J. Epidemiol.* **42**, 689–700 (2012).

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Author contributions

A.D.G., M.R., H.F.I., M.G.N. and E.M.T.-D. developed the software. A.D.G., M.G.N. and E.M.T.-D. developed the theory underlying genomic SEM. A.D.G., M.R., R.d.V., M.G.N. and E.M.T.-D. developed the techniques and mathematical derivations. A.D.G., T.T.M., M.G.N. and E.M.T.-D. performed the simulation studies. S.J.R., R.E.M. and E.M.T.-D. performed the polygenic prediction analyses. A.D.G., M.G.N. and E.M.T.-D. wrote the manuscript. M.R., S.J.R., T.T.M., W.D.H., A.M.M., I.J.D., R.E.M., P.D.K. and K.P.H. provided feedback and edited the manuscript.

Competing interests

The authors declare no competing interests.

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Software and code

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**Data collection**

No software was used for data collection.

**Data analysis**

- Lead SNPs and GWAS for simulation studies were conducted using PLINK v1.9.
- Simulation of phenotypes with corresponding genotypes was conducting using GCTA v1.91.
- The construction of polygenic scores for out-of-sample prediction was conducted using PRSice v1.25.
- Genomic SEM was run using code we make available for download at https://github.com/MichelNivard/GenomicSEM

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All summary statistics used are available for public download. No raw data was used except for the construction of polygenic scores (PGSs) for out-of-sample prediction in UK Biobank and Generation Scotland. Although we are unable to make this data available ourselves, the data may be requested by others. Supplementary Table 1 provides references for all of the summary statistics used in our analyses. All summary statistics used are available for public use, and links are provide directly below.

For construction of the p-factor, summary statistics for PTSD, Anxiety, Bipolar Disorder, and Major Depressive Disorder were downloaded from: http://www.med.unc.edu/pgc/results-and-downloads. Summary statistics for Schizophrenia were downloaded from: http://walters.pyscm.cf.ac.uk/.

The GWIS summary statistics for Bipolar Disorder and Schizophrenia were also downloaded from the cross-disorder section of the PGC website. The summary statistics for educational achievement are available for download from: https://www.thessgac.org/data.

Summary statistics for item-level indicators of neuroticism were downloaded from: https://docs.google.com/spreadsheets/d/1b3oGI21Ut578cuHttWaZotQcID-mBRpZih87Ms_No/edit?gid=1209628142. We use Round 1 of the Neale's Lab UKB GWAS results.

Summary statistics used for the anthropometric traits example for BMI, waist-hip ratio, hip circumference, waist circumference, and height are available from: https://portals.broadinstitute.org/collaboration/giant/index.php/GIANT_consortium_data_files.

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Behavioural & social sciences study design

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Study description  Our study uses publically available summary statistics to examine a new developed method, Genomic SEM. Genomic SEM can be used to examine any form of structural equation model (e.g., factor analysis, mediation, etc.) using summary statistics. The software provides model fit statistics, and can also be used to produce summary statistics for a latent trait.

Research sample  We use only publically available summary statistics for our analyses. As Genomic SEM relies on ld-score regression (LDSC) to construct genetic covariance matrices, and LDSC requires summary statistics to be within a single ethnic population due to differences in linkage disequilibrium across populations, we use only summary statistics restricted to European populations.

Sampling strategy  This is not applicable as we use previously collected data.

Data collection  This is not applicable as we use previously collected data.

Timing  This is not applicable as we use previously collected data.

Data exclusions  We use only summary statistics from European populations due to the requirements of LDSC, as noted above.

Non-participation  This is not applicable as we use previously collected data.

Randomization  This is not applicable as we use previously collected data.

Reporting for specific materials, systems and methods
Materials & experimental systems

n/a Involved in the study

- Unique biological materials
- Antibodies
- Eukaryotic cell lines
- Palaeontology
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Methods

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- ChIP-seq
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