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Reconstruction of Networks with Direct and Indirect Genetic Effects

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ABSTRACT Genetic variance of a phenotypic trait can originate from direct genetic effects, or from indirect effects, *i.e.*, through genetic effects on other traits, affecting the trait of interest. This distinction is often of great importance, for example, when trying to improve crop yield and simultaneously control plant height. As suggested by Sewall Wright, assessing contributions of direct and indirect effects requires knowledge of (1) the presence or absence of direct genetic effects on each trait, and (2) the functional relationships between the traits. Because experimental validation of such relationships is often unfeasible, it is increasingly common to reconstruct them using causal inference methods. However, most current methods require all genetic variance to be explained by a small number of quantitative trait loci (QTL) with fixed effects. Only a few authors have considered the "missing heritability" case, where contributions of many undetectable QTL are modeled with random effects. Usually, these are treated as nuisance terms that need to be eliminated by taking residuals from a multi-trait mixed model (MTM). But fitting such an MTM is challenging, and it is impossible to infer the presence of direct genetic effects. Here, we propose an alternative strategy, where genetic effects are formally included in the graph. This has important advantages: (1) genetic effects can be directly incorporated in causal inference, implemented via our PCgen algorithm, which can analyze many more traits; and (2) we can test the existence of direct genetic effects, and improve the orientation of edges between traits. Finally, we show that reconstruction is much more accurate if individual plant or plot data are used, instead of genotypic means. We have implemented the PCgen-algorithm in the R-package pcgen.

KEYWORDS Structural equation models; multivariate mixed models; causal inference

TO attain higher genetic gains, modern plant and animal breeders increasingly scale up their programs via the implementation of genomic prediction technologies (Cooper *et al.* 2014). Most genomic prediction applications are based on linear mixed- or Bayesian models that predict the phenotype for the target trait (yield) as a function of a multivariate distribution for single nucleotide polymorphism (SNP) effects. In these models, the physiological mechanisms and traits that modulate the genotypic response to the environment over time

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are modeled implicitly via the SNP effects directly affecting the target trait (Calus and Veerkamp 2011; Zhou and Stephens 2014). The availability of high throughput phenotyping technologies has enabled breeders to characterize additional traits, and to monitor growth and development during the season. This opens new opportunities in breeding strategies, in which better-adapted genotypes result from combining loci that regulate complementary physiological mechanisms. This kind of breeding strategy is called physiological breeding (Reynolds and Langridge 2016).

In physiological breeding, prediction accuracy for the target trait potentially benefits from a joint model for all underlying traits. This is partly because of the physiological knowledge that can be incorporated, but also because the use of genetically correlated traits with sufficiently large heritability increases accuracy (Thompson and Meyer 1986;

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van Eeuwijk *et al.* 2019). Often, however, a realistic model should account for at least some of the causal relations between traits, which is difficult with the regression models that are used in most genomic prediction literature. Structural equation models (SEMs), proposed by Wright (1921), and extended with random genetic effects in Gianola and Sorensen (2004), are a promising approach to deal with this problem (Rosa *et al.* 2011). In SEMs, each trait is modeled explicitly as a function of the other traits and a noise term. Therefore, SEMs are a useful tool to identify which are the key traits that could be selection targets, or be incorporated in multi-trait genomic prediction models to improve the prediction accuracy for the target trait.

A first advantage of SEMs, compared to regression models, is that one can predict the behavior of the system when one or more of the structural equations are modified by some kind of intervention. Figure 1 shows an illustration of an intervention. For example, a question that could be of interest to plant breeders is: which would be the contribution of a trait (say, radiation use efficiency) to yield, if flowering time is fixed for all genotypes at a particular value?

Second, SEMs make possible to distinguish between direct and indirect effects of one trait on another, and, similarly, between direct and indirect genetic effects. For example, let plant height (trait \mathbf{Y}_1) be modeled as $\mathbf{Y}_1 = \mathbf{G}_1 + \mathbf{E}_1$, *i.e.*, as the sum of a random genetic and residual effect, where all terms are $n \times 1$ vectors, containing the values for a population of nindividuals. Suppose plant height has a linear effect on yield (\mathbf{Y}_2), with additional random effects \mathbf{G}_2 and \mathbf{E}_2 :

$$\mathbf{Y}_2 = \lambda \mathbf{Y}_1 + \mathbf{G}_2 + \mathbf{E}_2 = (\lambda \mathbf{G}_1 + \mathbf{G}_2) + \lambda \mathbf{E}_1 + \mathbf{E}_2,$$

where λ is the path (or structural) coefficient associated with the effect of \mathbf{Y}_1 on \mathbf{Y}_2 . On the one hand, we have the *direct* genetic effects \mathbf{G}_1 and \mathbf{G}_2 ; on the other, we have the *total* (or marginal) genetic effects $\mathbf{U}_1 = \mathbf{G}_1$ and $\mathbf{U}_2 = \lambda \mathbf{G}_1 + \mathbf{G}_2$, the indirect effects being $\mathbf{U}_1 - \mathbf{G}_1 = 0$ and $\mathbf{U}_2 - \mathbf{G}_2 = \lambda \mathbf{G}_1$. Similarly, we can distinguish between the 2×2 matrices Σ_G , containing the (co) variances of \mathbf{G}_1 and \mathbf{G}_2 , and V_G , with the (co) variances of \mathbf{U}_1 and \mathbf{U}_2 . The latter is the matrix of genetic (co) variances, appearing in the usual MTM (multi-trait mixed model) for \mathbf{Y}_1 and \mathbf{Y}_2 ; here, it is a function of Σ_G and λ .

Knowledge of the direct genetic effects is often of great interest to breeders (Valente *et al.* 2013, 2015). However, routine use of these models is currently difficult for two reasons. First, for a given SEM, not all parameters may be identifiable, *i.e.*, because of overparameterization, different sets of parameter values can lead to the same model, making estimation infeasible. Gianola and Sorensen (2004) provided criteria for identifiability, and suggested putting constraints on some parameters, although automatic generation of interpretable and meaningful constraints remains difficult, especially in high-dimensional settings.

A second (and more fundamental) obstacle for the use of SEMs with genetic effects is that the underlying structure is often unknown. In such cases, causal inference methods (Spirtes *et al.* 2001; Pearl 2009; Maathuis and Nandy

2016) can be used, which reconstruct causal models that are, in some sense, most compatible with the observed data. Most causal inference methods, however, require independent samples, and cannot account for genetic relatedness. For this reason, genotypic differences are most often modeled using a small number of quantitative trait loci (QTL) with fixed effects (Chaibub Neto et al. 2008, 2013; Scutari et al. 2014), but when part of the genetic variance is not explained by QTL (missing heritability), the use of random genetic effects seems inevitable. Only a few works have studied reconstruction in the presence of such effects. Valente et al. (2010) and Töpner et al. (2017) proposed to perform causal inference after subtracting genomic predictions obtained from an MTM. Similarly, Gao and Cui (2015) applied the PC algorithm (Spirtes et al. 2001) to the residuals of multi-single nucleotide polymorphism (SNP) models. The difficulty with these approaches is that the MTM is limited to small numbers of traits, and that the existence of direct genetic effects cannot be tested. For example, if the causal graph among three traits is $Y_1 \rightarrow Y_2 \leftarrow Y_3$, and there are direct genetic effects on Y_1 and Y_3 , then the absence of a direct genetic effect on Y_2 cannot be inferred from MTM residuals.

Inspired by these problems, we define a framework in which direct genetic effects are part of the causal graph, and a single node G represents all direct genetic effects. For each trait Y_i , an arrow $G \rightarrow Y_i$ is present if and only if the direct genetic effect on Y_i is nonzero, *i.e.*, if the *j*th diagonal element of Σ_G is positive. See Figure 3 below for an example. Although our causal interpretation of genetic effects is not new (Stephens 2013; Valente et al. 2013, 2015), this work appears to be the first to formalize it. In particular, we show that the Markov property holds for the graph extended with genetic effects (Theorem 1 below). Informally speaking, this means that there is a one-to-one correspondence between edges in the causal graph and conditional dependencies in the distribution of the traits and genetic effects. This means that edges (either between two traits, or between a trait and G) can be inferred from multi-trait data. Consequently, while some of the covariances between direct genetic effects (contained in Σ_G) may still be unidentifiable, we can at least identify which rows and columns in Σ_G are zero.

Building on the Markov-property, we propose the PCgen algorithm. PCgen stands for PC with genetic effects, and is an adaptation of the general PC-algorithm (named after its inventors Peter Spirtes and Clark Glymour). Briefly, PCgen assesses the existence of a direct genetic effect on a given trait by testing whether its genetic variance is zero, conditional on various sets of other traits. For the existence of an edge between traits Y_1 and Y_2 , we test whether, in a bivariate MTM, the residual covariance between Y_1 and Y_2 is zero, again conditional on sets of other traits. Under the usual assumptions of independent errors, recursiveness, and faithfulness, we show that PCgen can recover the underlying partially directed graph (Theorem 2). Because fitting an MTM for all traits simultaneously is no longer necessary, PCgen can handle a considerably larger number of traits.

While our approach is generally applicable to any species and relatedness matrix, our implementation of PCgen is currently limited to the specific (but important) case of populations where observations on genetically identical replicates are available, assuming independent genetic effects (i.e., as in the classical estimation of broad-sense heritability). This is partly for pragmatic reasons (e.g., lower computational requirements), and partly for statistical reasons. In particular, successful reconstruction requires sufficient power in the tests for direct genetic effects $(G \rightarrow Y_i)$, and those for the between traits relations $(Y_i \rightarrow Y_k)$. Given the availability of replicates, this power is likely to be highest when the original observations are used, instead of genotypic means and a marker-based genetic relatedness matrix (GRM), modeling additive effects (Kruijer et al. 2015). Although mixed models with both replicates and a GRM may further increase power, the increase is often modest, and statistical inference can become biased under model misspecification (e.g., when the GRM models additive effects, and the true architecture is partly epistatic; see Kruijer 2016). By contrast, using only replicates, unbiased estimation of broad-sense heritability is always possible, regardless of the population structure and genetic architecture. The downside is that the contributions of different types of genetic effects cannot be distinguished. On the positive side, PCgen appears to be the first algorithm that can infer the presence of direct genetic effects based on phenotypic data alone.

Our approach is related to that of Stephens (2013), who inferred the sets of traits being affected directly and indirectly by a given locus, assuming unrelated individuals and using only summary statistics. Here, we instead consider sums of individual locus effects, for possibly related individuals. Moreover, PCgen also aims to reconstruct structural relations among the traits themselves, and can deal with larger numbers of traits.

The paper is organized as follows. After introducing SEMs with genetic effects, we define their graphical structure, and, from this perspective, review existing approaches. We then describe the general form of the PCgen-algorithm for estimation of the graphical structure, followed by various proposals for the required conditional independence tests. Next, we test PCgen performance in data simulated with both statistical and crop-growth models, and analyze a maize and a rice dataset. Finally, we state several results regarding PCgen's statistical properties. Supplemental Material, Table S1 provides an overview of the notation, and Appendix A.1 contains the necessary graph-theoretic definitions. Figure 2 provides a graphical summary of our theory and methodology.

Materials and Methods

Structural equation models

To introduce structural models, we first consider a simple linear SEM without genetic effects:

$$\mathbf{y}_{\mathbf{i}} = \mathbf{x}_{\mathbf{i}}B + \mathbf{y}_{\mathbf{i}}\Lambda + \mathbf{e}_{\mathbf{i}},\tag{1}$$

where \mathbf{y}_i is a $1 \times p$ vector of phenotypic values for p traits measured on the *i*th individual, \mathbf{e}_i is a vector of random errors, and Λ is a $p \times p$ matrix of structural coefficients. The $q \times p$ matrix $B = [\beta^{(1)} \cdots \beta^{(p)}]$ contains intercepts and trait-specific fixed effects of (exogenous) covariates, whose values are contained in the $1 \times q$ vector \mathbf{x}_i .

To write Equation 1 in matrix-form, we define the $n \times q$ design matrix **X** with rows \mathbf{x}_i . Similarly, we define $n \times p$ matrices $\mathbf{Y} = [\mathbf{Y}_1 \cdots \mathbf{Y}_p]$ and $\mathbf{E} = [\mathbf{E}_1 \cdots \mathbf{E}_p]$, with rows \mathbf{y}_i and \mathbf{e}_i , and columns \mathbf{Y}_j and \mathbf{E}_j . We can then write

$$\mathbf{Y} = \mathbf{X}\mathbf{B} + \mathbf{Y}\Lambda + \mathbf{E}.$$
 (2)

 Λ has zeros on the diagonal, and defines a directed graph $\mathcal{G}_{\mathcal{V}}$ over the traits Y_1, \ldots, Y_p , containing the edge $Y_j \rightarrow Y_k$ if and only if the (j,k)th entry of Λ is nonzero. The columns in Equation 2 correspond to *p* linear structural equations, one for each trait. These are determined by the path coefficients, the nonzero elements in Λ . For example, in Figure 1, if $\mathbf{X} = \mathbf{1}_n$ is the $n \times 1$ vector of ones and $B = [\mu_1 \, \mu_2 \, \mu_3]$, the third trait has values $\mathbf{Y}_3 = \mu_3 \mathbf{1}_n + \lambda_{13} \mathbf{Y}_1 + \lambda_{23} \mathbf{Y}_2 + \mathbf{E}_3$. The equality sign here should be understood as an assignment, *i.e.*, \mathbf{Y}_3 is determined by the values of Y_1 and Y_2 (its parents in the graph $\mathcal{G}_{\mathcal{V}}$) and an error. If the directed graph does not contain any cycle (*i.e.*, a directed path from a trait to itself), it is a directed acyclic graph (DAG), and the SEM is said to be recursive. In the notation, we will distinguish between the nodes Y_1, \ldots, Y_p in the graph $\mathcal{G}_{\mathcal{Y}}$ (normal type), and the random vectors $\mathbf{Y}_1, \ldots, \mathbf{Y}_p$ that these nodes represent (bold face).

As mentioned above, SEMs can be used to predict the effects of *interventions*, which mathematically correspond to changes in the structural equations. For example, suppose that, in Figure 1, Y_1 , Y_2 , and Y_3 are the expression levels of three genes, and Y_4 is plant height. Then, after forcing Y_2 to be zero (*e.g.*, by knocking out the gene), the total effect of Y_1 on Y_4 changes from $(\lambda_{13}\lambda_{34} + \lambda_{12}\lambda_{23}\lambda_{34})$ to $\lambda_{13}\lambda_{34}$ (File S7.3 and File S7.4 provide other examples, involving genomic prediction). More generally, the new joint distribution of $\mathbf{Y}_1, \ldots, \mathbf{Y}_p$ after an intervention can be obtained from the manipulation or truncated factorization theorem (Pearl 2009), *without* observations from the new distribution. For the consequences for genomic prediction, see Valente *et al.* (2013) and the *Discussion* section below.

GSEM: structural equation models with genetic effects

Gianola and Sorensen (2004) extended model (1) with random genetic effects \mathbf{g}_i : for individuals i = 1, ..., n, it is then assumed that

$$\mathbf{y}_{\mathbf{i}} = \mathbf{x}_{\mathbf{i}}B + \mathbf{y}_{\mathbf{i}}\Lambda + \mathbf{g}_{\mathbf{i}} + \mathbf{e}_{\mathbf{i}},\tag{3}$$

where the $1 \times p$ vectors, \mathbf{g}_i , contain the direct genetic effects for individuals i = 1, ..., n. We will refer to model (3) as a linear genetic structural equation model (GSEM). While the



Figure 1 An example of a linear SEM. The SEM can be represented by a graph (middle), which is defined by the nonzero elements of Λ , the matrix containing the path coefficients (top). The total effect of Y_1 on Y_4 can be obtained by summing the contributions of the directed paths $Y_1 \rightarrow Y_3 \rightarrow Y_4$ and $Y_1 \rightarrow Y_2 \rightarrow Y_3 \rightarrow Y_4$, where each contribution is the product of the corresponding path coefficients. After the intervention $Y_2 := 0$ (bottom), the effect changes from $(\lambda_{13}\lambda_{34} + \lambda_{12}\lambda_{23}\lambda_{34})$ to $\lambda_{13}\lambda_{34}$.

genetic effects introduce relatedness between individuals, there is no form of social interaction [as in Moore *et al.* (1997) and Bijma (2014)]. Each \mathbf{g}_{i}^{t} follows a $N(0, \Sigma_{G})$ distribution, where Σ_{G} is a $p \times p$ matrix of genetic variances and covariances. The vectors \mathbf{g}_{i} are independent of the \mathbf{e}_{i} 's, but not independent among themselves. Defining a $n \times p$ matrix $\mathbf{G} = [\mathbf{G}_{1} \cdots \mathbf{G}_{p}]$, with rows \mathbf{g}_{i} and columns \mathbf{G}_{j} $(j = 1, \dots, p)$, we can extend Equation 2 as follows:

$$\mathbf{Y} = \begin{bmatrix} \mathbf{Y}_1 \cdots \mathbf{Y}_p \end{bmatrix} = \mathbf{X}B + \mathbf{Y}\Lambda + \mathbf{G} + \mathbf{E}.$$
 (4)

Each vector G_j contains the direct genetic effects on the *j*th trait. We make the following assumptions about the GSEM defined by (4), which are summarized in Figure 2:

- All traits are measured in the same experiment: the rows y_i of Y may be either observations at plot or plant level or genotypic means across plots or plants, but the observations should always come from the same experiment. In addition, the residual errors originate from biological variation, *i.e.*, measurement errors are negligible [this in contrast to related work on Mendelian randomization (Hemani *et al.* 2017)].
- 2. *Recursiveness:* the graph $\mathcal{G}_{\mathcal{Y}}$ defined by Λ is a DAG. Consequently, there should be no feedback loops.
- 3. *Causal sufficiency:* the covariance matrix Σ_E of the error vectors \mathbf{e}_i is diagonal, *i.e.*, there are no latent variables. This means that all nonzero (nongenetic) correlations between traits must be the consequence of causal relations between the traits. We also assume the diagonal elements of Σ_E to be strictly positive.
- 4. *Genetic relatedness among individuals:* **G** is independent of **E**, and has a matrix-variate normal distribution with row-covariance *K* and column covariance Σ_G , where *K* is a $n \times n$

relatedness matrix, which we describe in more detail below (see the section *Genetic relatedness*). Equivalent to this, the $np \times 1$ vector $vec(\mathbf{G}) = (\mathbf{G}_1^t, \dots, \mathbf{G}_p^t)^t$ is multivariate normal with covariance $\Sigma_G \otimes K$, where *vec* denotes the operation of creating a column vector by stacking the columns of a matrix. Consequently, each \mathbf{G}_j is multivariate normal with covariance $\sigma_{G_j}^2 K$, where the variances $\sigma_{G_j}^2$ form the diagonal of Σ_G . Using the same notation, we can write that \mathbf{E} is matrix-variate normal with row-covariance I_n and column covariance Σ_E , and that $vec(\mathbf{E}) \sim N(0, \Sigma_E \otimes I_n)$.

5. No collinear genetic effects: the diagonal elements of Σ_G do not need to be strictly positive, but, for all nonzero elements, the corresponding correlation should not be 1 or -1.

Assumptions 1-4 were also made in related work on structural models with random genetic effects (Valente et al. 2010; Töpner et al. 2017), and 1–3 are commonly made for structural models without such effects. Assumption 1 is implicit in the GSEM model (4) itself, as it is assumed that the structural equations propagate all errors to traits further down in the graph. Network reconstruction with traits from different experiments would rely completely on the genetic effects, requiring Σ_G to be diagonal, which is a rather unrealistic assumption (see the Discussion, section Data from different experiments). A small amount of measurement error does not seem to pose problems for our PCgen algorithm. Larger amounts of measurement error will decrease power, which can, however, be avoided by increasing the number of genotypes or replicates (see Table S4, discussed below). Assumption 1 does not require traits to be measured at the same time. In particular, it is possible to include the same trait measured at different time-points, which, of course, puts constraints on causality. Such constraints can, in principle, be incorporated in our model, just as other biological constraints (see e.g., Peters et al. 2017), although we will not explore this here. What is also implicit in Equation 4 is that all causal relations between traits are linear. Our PCgen algorithm relies on this rather heavily, and we discuss the consequences of nonlinearity in the Results below (specifically, in the APSIM simulations, and the example just before the Discussion). In specific cases, it may be possible to obtain linearity by certain transformations of the data, but this requires prior knowledge that is typically unavailable. In the *Discussion*, we suggest various directions of future work to deal with nonlinear relationships, as well as non-Gaussian errors. In any case, as long as the other assumptions hold, the core of our framework (the graphical representation of genetic effects with a single node G, and the Markov property in Theorem 1 below) is still valid for nonlinear GSEMs.

Assumption 2 (no cycles) is essential given the type of data considered here, as the reconstruction of feedback loops requires time-course data (Peters *et al.* 2017), typically with high-resolution. Without such data (or only a few time-points), it is impossible to verify this assumption, but Maathuis *et al.* (2010) provide examples of interventions in yeast data, where cycles are likely to exist, but structural models still outperform nonstructural models.



Figure 2 Graphical summary of the theory and methodology. The Markov property on the right (green; for the residuals) is well known from the literature, while the Markov property on the left (blue, for the conditional distributions of Y_1, \ldots, Y_p, G) is established in Theorem 1. Table S1 contains an overview of the notation, and Appendix A.1 provides the necessary graph-theoretic definitions.

Assumption 3 (no latent variables) is important for orientation of the edges, and has been studied in detail by many authors. In particular, Spirtes *et al.* (2001) and Colombo *et al.* (2012) proposed the FCI and RFCI algorithms, which are extensions of the PC-algorithm, and allow for latent variables. These algorithms could be extended with genetic effects, as we do here for the PC-algorithm (see the *Discussion*). Apart from nonlinear trait-to-trait relations, the APSIM simulations below also contain latent variables.

As in related work (Valente *et al.* 2010; Töpner *et al.* 2017), as well as in much of the literature on multi-trait genomic prediction and genome-wide association studies (GWAS), the relatedness matrix K is the same for all traits (Assumption 4). This may not hold if traits have very different genetic architectures, but seems a good approximation if most of the underlying QTL are small. Large QTL may be added as additional fixed effects.

Assumption 5 implies that, for each pair of traits with direct genetic effects, these effects should not be the result of exactly the same set of QTL, with exactly the same effect sizes. This seems a reasonable assumption whenever the underlying biological structures or processes are really different; see the section *Dealing with derived traits* in the *Discussion*. Of course, reconstruction of direct genetic effects will be more difficult under strong correlations, similar to, for example, the reduced power in GWAS when two causal loci are in strong LD.

Finally, there are a few additional assumptions which are required for the PCgen-algorithm, and are not essential for the definition of GSEM; see the overview in Figure 2 and the section *Statistical properties of PCgen* in the *Results*. In particular, we require the *faithfulness* assumptions defined by expressions 9 and 10 below, and assumptions about the conditional distributions. Appendices A.5 and A.6 provide additional examples and results regarding faithfulness.

Graphical representation of GSEM: extending $\mathcal{G}_{\mathcal{Y}}$ with genetic effects

In contrast to previous work, we will explicitly take into account the possibility that there are no direct genetic effects on some of the traits. In this case, the corresponding rows and columns in Σ_G are zero. Following the notation of Stephens (2013), we use $D \subseteq \{1, \ldots, p\}$ to denote the index set of traits with direct genetic effects, and write $\Sigma_G[D, D]$ for the submatrix with rows and columns restricted to *D*. From Assumption 5 above, it follows that $\Sigma_G[D,D]$ is nonsingular, *i.e.*, there can be no perfect correlations between direct genetic effects.

We graphically represent model (4) by a graph \mathcal{G} with nodes Y_1, \ldots, Y_p , and a node G, which represent, respectively, $\mathbf{Y}_1, \ldots, \mathbf{Y}_p$ and the matrix $\mathbf{G} = [\mathbf{G}_1 \cdots \mathbf{G}_p]$. \mathcal{G} contains an edge $Y_j \rightarrow Y_k$ if the (j, k)th entry of Λ is nonzero, and an edge $G \rightarrow Y_j$ if \mathbf{G}_j is nonzero with probability 1, *i.e.*, if $\sigma_{G,j}^2 > 0$. See Figure 3 for an example. In words, \mathcal{G} is defined as the original graph $\mathcal{G}_{\mathcal{Y}}$ over the traits, extended with the node G and arrows $G \rightarrow Y_j$ for traits with a direct genetic effect, *i.e.*, for all $j \in D$. Consequently, our main objective of reconstructing trait-to-trait relationships and direct genetic effects translates as reconstructing \mathcal{G} .

As for the \mathbf{Y}_j 's, we distinguish between the node *G* in the graph (normal type) and the random matrix **G** it represents (bold face). **G** is represented by a *single* node *G*, instead of multiple nodes G_1, \ldots, G_p . This choice is related to our assumption that *K* is the same for all traits; see File S7.1 for a motivating example. The orientation of any edge between *G* and Y_j is restricted to $G \rightarrow Y_j$, because the opposite orientation would be biologically nonsensical. Because of our assumption that $\mathcal{G}_{\mathcal{Y}}$ is a DAG, it follows that \mathcal{G} is a DAG as well, as a cycle would require at least one edge pointing into *G*.

We emphasize that \mathcal{G} is just a mathematical object and not a complete visualization of all model terms and their distribution, as is common in the SEM literature. In particular, \mathcal{G} does not contain nodes for the residual errors, path coefficients, or information about the off-diagonal elements of Σ_G . While in general, Σ_G is not entirely identifiable (Gianola and Sorensen 2004), we will see that \mathcal{G} is identifiable in terms of its skeleton (the undirected graph obtained when removing the arrowheads) and some of the orientations. The skeleton is generally not equal to the conditional independence graph, which is the undirected graph associated with the inverse covariance or precision matrix (Spirtes *et al.* 2001; Kalisch and Bühlmann 2007). See File S6.2 for an example.

Direct and indirect genetic effects

As pointed out by various authors (Gianola and Sorensen 2004; Valente *et al.* 2010, 2013; Töpner *et al.* 2017), the genetic variance of a trait is driven not only by its direct genetic effect (\mathbf{g}_i), but also by direct genetic effects on traits affecting it, *i.e.*, its parents in the graph $\mathcal{G}_{\mathcal{Y}}$. Assuming that the inverse $\Gamma = (I-\Lambda)^{-1}$ exists, it follows from Equation 3 that

$$\mathbf{y}_{\mathbf{i}} = \mathbf{x}_{\mathbf{i}}B\Gamma + \mathbf{g}_{\mathbf{i}}\Gamma + \mathbf{e}_{\mathbf{i}}\Gamma = \mathbf{x}_{\mathbf{i}}B\Gamma + \mathbf{u}_{\mathbf{i}} + \mathbf{e}_{\mathbf{i}}\Gamma \sim N(\mathbf{x}_{\mathbf{i}}(B\Gamma), \Gamma^{t}\Sigma_{G}\Gamma + \Gamma^{t}\Sigma_{E}\Gamma) = N(\mathbf{x}_{\mathbf{i}}(B\Gamma), V_{G} + V_{E}),$$
(5)

where the $1 \times p$ vector $\mathbf{u}_i = \mathbf{g}_i \Gamma$ contains the *total* genetic effects for the *i*th individual. The $n \times 1$ vector $\mathbf{U}_j = \mathbf{G}\gamma_j$ contains the total genetic effects for the *j*th trait, where γ_j is defined as the *j*th column of Γ . The vector of *indirect* genetic effects is the difference $\mathbf{U}_j - \mathbf{G}_j$. In Figure 3 for example, $\mathbf{G}_3 = (0, \dots, 0)^t$ and $\mathbf{U}_3 = \lambda_{13}\mathbf{G}_1 + \lambda_{23}\mathbf{G}_2$.



Figure 3 An example of a graph G representing a genetic structural equation model (GSEM), with path-coefficients λ_{13} and λ_{23} . There is no direct genetic effect on Y_3 , and therefore no edge $G \rightarrow Y_3$.

Likewise, we can distinguish between the contribution of direct and indirect genetic effects to the genetic covariance. The (j, k)th element of $V_G = \Gamma^t \Sigma_G \Gamma$ in Equation 5 is the *total* genetic covariance between Y_j and Y_k . This is what is usually meant with genetic covariance. Most often, this is different from the covariance between the direct genetic effects G_i and **G**_k, given by $\Sigma_G[j,k]$. Indeed, $\Sigma_G[j,k]$ affects the total genetic covariance, but the latter is also driven by causal relationships between traits, as defined by $\Gamma = (I - \Lambda)^{-1}$. If no such relationship exist, then Λ contains only zeros, and $V_G = \Sigma_G$. In general, however, these matrices are different, and, depending on the structure of the graph and the path coefficients, the correlation $\sum_{G}[j,k]/\sqrt{\sum_{G}[j,j]\sum_{G}[k,k]}$ may be much larger than $V_G[j,k]/\sqrt{V_G[j,j]}V_G[k,k]$, or vice versa. For example, given direct effects G_1 and G_2 with equal variance and correlation 0.9, and an effect $Y_1 \rightarrow Y_2$ of size -1, the total genetic correlation is -0.22. Regarding the diagonal of V_G , we note that traits without a direct genetic effect may still have positive genetic variance.

Genetic relatedness: The genetic relatedness matrix *K* introduced in Assumption 4 determines the covariance between the rows of **G**. In principle, our approach allows for any type of GRM, but, for simplicity, we focus on the following types. In all cases, *K* has dimension $n \times n$.

- 1. $K = ZZ^t$, *Z* being the $n \times m$ incidence matrix assigning n = mr plants (or plots) to *m* genotypes, in a balanced design with *r* replicates for each genotype. This *K* is obtained when each genotype has an independent effect, as in the classical estimation of broad-sense heritability (or repeatability). Since no marker information is included, the model cannot be used directly for genomic prediction, but we will see that, for the reconstruction of *G* (using the training genotypes), it has considerable computational and statistical advantages.
- 2. Given only a single individual per genotype (or genotypic means), we assume K = A, A being a $(n \times n)$ GRM estimated from a dense set of markers, assuming additive infinitesimal effects.
- 3. Given both *r* replicates of *m* genotypes and a GRM *A* of dimension $m \times m$, we assume that $K = ZAZ^t$. In absence of nonadditive effects, this covariance structure uses all available information. However, for computational reasons it is usually easier to work with either the replicates

or with genotypic means and the GRM *A*. We further explore this issue in the simulations below and in the *Discussion*.

The balance required when $K = ZZ^t$ is necessary in Theorems 5 and 6 below, but is not a general requirement for our models, nor for the PCgen algorithm.

The joint distribution implied by the GSEM: The sum $\mathbf{G} + \mathbf{E}$ does not, in general, have a matrix-variate normal distribution, but from our Assumption 4, it still follows that $vec(\mathbf{G} + \mathbf{E})$ is multivariate normal with covariance $\Sigma_G \otimes K + \Sigma_E \otimes I_n$. We can rewrite Equation 4 as

$$\mathbf{Y} = \mathbf{X}B\Gamma + \mathbf{G}\Gamma + \mathbf{E}\Gamma = \mathbf{X}B\Gamma + \mathbf{U} + \mathbf{E}\Gamma, \tag{6}$$

where $\mathbf{U} = \mathbf{G}\Gamma$ is the $n \times p$ matrix of total genetic effects, with columns \mathbf{U}_{j} . Equation 5 now generalizes to

$$\operatorname{vec}(\mathbf{Y}) \sim N\left(\operatorname{vec}\left(\mathbf{X}\tilde{B}\right), V_G \otimes K + V_E \otimes I_n\right),$$
 (7)

where $B = B\Gamma$ is the matrix of fixed effects transformed by Γ . This is a common model for multi-trait GWAS and genomic prediction [see, among others, Korte *et al.* (2012), Stephens (2013), and Zhou and Stephens (2014)]. In those works, however, V_G and V_E are arbitrary covariance matrices, whereas here they are modeled as functions of Σ_G , Σ_E , and $\Gamma = (I-\Lambda)^{-1}$.

Under Assumption 3 (Σ_E diagonal), Σ_G , Σ_E , and Λ together have, at most, p(p+1)/2 + p + p(p-1)/2 = p(p+1) parameters, as many as V_G and V_E together. This suggests that Σ_G , Σ_E , and Λ might be identifiable from the distribution in Equation 7. In Appendix A.2 we show how Σ_G , Σ_E , and Λ can be obtained from V_G and V_E . Apart from Assumption 3, this requires knowledge of the graph, and the faithfulness Assumptions (9)–(10) given below. Equations 17 and 18 in Appendix A.2 can be used to derive estimates $\hat{\Sigma}_G$, $\hat{\Sigma}_E$, and $\hat{\Lambda}$ from estimates \hat{V}_G and \hat{V}_E , although the development of good estimators of Σ_G , Σ_E , and Λ is beyond the scope of this work. Such estimators should account for the structure of the GSEM, as defined by Λ and Σ_G , and might rely on alternative restrictions on the parameters (instead of diagonal Σ_E); see Gianola and Sorensen (2004).

Using the results of Spirtes *et al.* (2001) (p. 371), it turns out that Γ can be written directly in terms of sums of products of path coefficients (see Appendix A.3). Consequently, there is no need to invert $(I - \Lambda)$, although it still holds that $\Gamma = (I - \Lambda)^{-1}$, provided the inverse exists. Recalling that γ_j is the *j*th column of Γ , we can express the *j*th trait as

$$\mathbf{Y}_{\mathbf{j}} = \mathbf{X}B\boldsymbol{\gamma}_{j} + \mathbf{G}\boldsymbol{\gamma}_{j} + \mathbf{E}\boldsymbol{\gamma}_{j} = \mathbf{X}B\boldsymbol{\gamma}_{j} + \mathbf{U}_{\mathbf{j}} + \mathbf{E}\boldsymbol{\gamma}_{j}, \qquad (8)$$

which is Equation 6 restricted to the *j*th column. Similarly, for any nonempty index-set $S \subset \{1, ..., p\}$, the $n \times |S|$ matrix \mathbf{Y}_{S} of traits in *S* (*i.e.*, **Y** with columns restricted to *S*) equals $\mathbf{X}B\Gamma_{S} + \mathbf{G}\Gamma_{S} + \mathbf{E}\Gamma_{S}$, where Γ_{S} is the $p \times |S|$ matrix with columns γ_j $(j \in S)$. Equation (26) (Appendix A.7) provides an expression for the covariance of $vec(\mathbf{Y}_S)$. For the corresponding nodes in the graph, we write $Y_S = \{Y_m : m \in S\}$.

Causal inference without genetic effects: So far, we have assumed that \mathcal{G} is known, in which case estimation of Λ , Σ_G , and Σ_E is usually of interest. In this work, however, we focus on the reconstruction of an unknown \mathcal{G} , based on observations from a GSEM given in Equation 4. We will do this with the PCgen algorithm introduced below, but will first review the necessary concepts, as well as existing methods. Appendix A.1 contains a more detailed introduction.

Suppose for the moment we have observations generated by an acyclic SEM without latent variables, and without genetic effects. From the pioneering work of Judea Pearl and others in the 1980s, it is known that we can recover the skeleton of the DAG and some of the orientations, *i.e.*, those given by the *v*-structures. A *v*-structure is any triple of nodes Y_j, Y_k, Y_l , such that $Y_j \rightarrow Y_k \leftarrow Y_l$, without an edge between Y_j and Y_l . All DAGs with the same skeleton and *v*-structures form an equivalence class, which can be represented by a completed partially directed acyclic graph (CPDAG). DAGs from the same equivalence class cannot be distinguished using observational data, at least not under the assumptions we make here. For the reconstruction of the CPDAG, constraintbased and score-based methods have been developed (for an overview, see Peters *et al.* 2017).

Here, we focus on constraint-based methods, which rely on the equivalence of conditional independence (a property of the distribution) and directed separation (d-separation; a property of the graph). An important result is that an edge $Y_i - Y_k$ is missing in the skeleton of the DAG if and only if Y_i and Y_k are d-separated by at least one (possibly empty) set of nodes, Y_S . Such Y_S is called a *separating set* for Y_i and Y_k . Given the equivalence of d-separation and conditional independence, this means that we can infer the presence of the edge $Y_i - Y_k$ in the skeleton by testing $\mathbf{Y}_i \perp \mathbf{Y}_k | \mathbf{Y}_s$ for all \mathbf{Y}_s . The PC- and related algorithms therefore start with a fully connected undirected graph, and remove the edge $Y_i - Y_k$ whenever Y_i and Y_k are found to be conditionally independent for some Y_S . While the first constraint-based algorithms such as IC (Pearl et al. 1991) exhaustively tested all possible subsets, the PC-algorithm (Spirtes et al. 2001) can often greatly reduce the number of subsets to be considered. Although this is not essential for the equivalence of d-separation and conditional independence, most constraint-based algorithms assume that observations be indendently and identically distributed, and structural equations with additional random effects are usually not considered.

Existing approaches for estimating $\mathcal{G}_{\mathcal{Y}}$ *, given genetic effects:* To deal with the dependence introduced by the genetic effects, Valente *et al.* (2010) and Töpner *et al.* (2017) proposed to predict the total genetic effects (*i.e.*, the term **U** in Equation 6), and perform causal inference on the residuals. These methods are flexible, in the sense that any genomic

prediction method can be used, and combined with any causal inference method. A disadvantage, however, is that the presence of direct genetic effects cannot be tested. Suppose, for example, that $G \rightarrow Y_1 \rightarrow Y_2 \rightarrow Y_3$, and we subtract the total genetic effects. Then, given only the residuals, we can never know if part of the genetic variance of Y_2 was due to a direct effect $G \rightarrow Y_2$. Another disadvantage is that fewer of the between-trait edges can be oriented. Technically, this is because, in the CPDAG (showing which orientations can be recovered from data), typically more edges are undirected; see Appendix A.1 for more details. In the preceding example, the CPDAG associated with \mathcal{G} is $G \rightarrow Y_1 \rightarrow Y_2 \rightarrow Y_3$, *i.e.*, all orientations can be recovered. By contrast, the CPDAG associated with $\mathcal{G}_{\mathcal{Y}}$ is $Y_1 - Y_2 - Y_3$, and we only know that the orientation is *not* $Y_1 \rightarrow Y_2 \leftarrow Y_3$ (see Figure 2).

To use the causal information associated with genetic effects, Töpner *et al.* (2017) estimated "genomic networks", based on the predictions themselves. These, however, seem to require additional assumptions, which are not required for the residual networks (in particular, diagonal Σ_G). Moreover, it seems difficult to relate edges in such a network to direct genetic effects (see the section *Data from different experiments* in the *Discussion*, and File S7.2). In summary, residual and genomic networks only estimate the (CPDAG associated with the) subgraph $\mathcal{G}_{\mathcal{Y}}$ of traitto-trait relations, instead of the complete graph \mathcal{G} .

Another disadvantage is that, without specific models putting restrictions on V_G and V_E , the MTM (7) can only be fitted for a handful of traits (Zhou and Stephens 2014) for statistical as well as computational reasons. For example, Zwiernik et al. (2017) showed that, for general Gaussian covariance models, (residual) ML-estimation behaves like a convex optimization problem only when $n \ge 14p$. Similar problems are likely to occur for Bayesian approaches. The problem with fitting the MTM to data from the GSEM model (Equation 4) is that one cannot exploit the possible sparseness of G. Even for sparse graphs with few direct genetic effects, the matrices $V_G = \Gamma^t \Sigma_G \Gamma$ and $V_E = \Gamma^t \Sigma_E \Gamma$ may still be dense, requiring a total of p(p+1) parameters. To overcome these limitations, we explicitly consider the presence or absence of direct genetic effects to be part of the causal structure, and develop PCgen, a causal inference approach directly on \mathcal{G} .

The PCgen algorithm: The main idea behind PCgen is that the PC-algorithm is applicable to any system in which d-separation and conditional independence are equivalent, and where conditional independence can be tested. We first describe the algorithm and propose the various independence tests; the equivalence is addressed in Theorems 1 and 2 below. If we define $Y_{p+1} := G$, and temporarily rename the node *G* as Y_{p+1} , PCgen is essentially the PC-algorithm applied to Y_1, \ldots, Y_{p+1} :

1. Skeleton-stage. Start with the fully connected undirected graph over $\{Y_1, \ldots, Y_{p+1}\}$, and an empty list of separation sets. Then, test the conditional independence between all pairs Y_i and Y_k , given subsets of other variables Y_s .

Whenever a p-value is larger than the prespecified significance threshold α , update the skeleton by removing the edge $Y_j - Y_k$, and add Y_S to the list of separation sets for Y_j and Y_k . This is done for conditioning sets of increasing size, starting with the empty set ($S = \emptyset$; marginal independence between \mathbf{Y}_j and \mathbf{Y}_k). Only consider S that, in the current skeleton, are adjacent to Y_j or Y_k .

2. **Orientation-stage.** Apply the orientation rules given in File S1 (R1–R3 in Algorithm 1) to the skeleton and separating sets found in the skeleton-stage. For example, if the skeleton is $Y_1 - Y_2 - Y_3$, and $\{Y_2\}$ is *not* a separating set for Y_1 and Y_3 , the skeleton is oriented $Y_1 \rightarrow Y_2 \leftarrow Y_3$; otherwise, neither of the two edges can be oriented.

In order to obtain PCgen, we need to make a few refinements to these steps. First, in the skeleton stage, we need to specify how to test conditional independence statements. Clearly, independence between two traits requires a different test than independence between a trait (Y_i) and G (*i.e.*, Y_{p+1}), in particular because the latter is not directly observed. Second, we need to modify the orientation rules, in order to avoid edges pointing into G. The usual rules give the correct orientations when given perfect conditional independence information, but statistical errors in the tests may lead to edges of the form $Y_i \rightarrow G$. Third, statistical errors can also make the output of PC(gen) order-dependent, *i.e.*, putting the columns (traits) in a different order may lead to a different reconstruction. We therefore adopt the PC-stable algorithm of Colombo and Maathuis (2014), who proposed to perform all operations in the skeleton- and orientation-stage list-wise (details given in File S1). Apart from eliminating the order-dependence, this has the advantage that all conditional independence tests of a given size |S| = s can be performed in parallel.

In summary, PCgen is the PC-stable algorithm with: (1) specific conditional independence tests (described below); and (2) modified orientation rules, in order to avoid edges pointing into *G* (File S1.2). As in the original PC-algorithm, the number of type-I and type-II errors occurring in the tests is determined by the choice of the significance threshold α , which is discussed in section *Assessing uncertainty* below and in the *Discussion*.

Skeleton stage: conditional independence tests. We can distinguish between the following types of conditional independence statements in the skeleton stage:

$$Y_{j} \bot G | Y_{S},$$
 (A)

$$\mathbf{Y}_{\mathbf{i}} \mathbf{\bot} \mathbf{Y}_{\mathbf{k}} | \{ \mathbf{G}, \mathbf{Y}_{\mathbf{S}} \}, \tag{B}$$

$$\mathbf{Y}_{\mathbf{j}} \perp \mathbf{Y}_{\mathbf{k}} | \mathbf{Y}_{\mathbf{S}},$$
 (C)

where $j,k \in \{1,\ldots,p\}$ $(j \neq k)$ and $S \subseteq \{1,\ldots,p\} \setminus \{j,k\}$ [or, in statement (A), $S \subseteq \{1,\ldots,p\} \setminus \{j\}$]. In words, (A) means that the trait \mathbf{Y}_j is independent of all genetic effects (**G**),

conditional on the traits $\mathbf{Y}_{\mathbf{m}}$ ($m \in S$). If *S* is the empty set, this is understood as marginal independence of $\mathbf{Y}_{\mathbf{j}}$ and \mathbf{G} . Similarly, (B) and (C) express conditional independence of traits $\mathbf{Y}_{\mathbf{j}}$ and $\mathbf{Y}_{\mathbf{k}}$ given \mathbf{G} and $\mathbf{Y}_{\mathbf{S}}$, or given $\mathbf{Y}_{\mathbf{S}}$ alone.

We now propose statistical tests for statements (A) and (B), which rely on the linearity of our GSEM, as well as some additional assumptions, which we discuss in more detail below (section *Statistical properties of PCgen*, and Figure 2). Statement (C) can be tested using standard partial correlations and Fisher's z-transform. However, as we show in File S6, this test is redundant, since for any set Y_S that d-separates Y_j and Y_k , the set $Y_S \cup \{G\}$ will also d-separate them. We therefore skip any test for $Y_j \bot Y_k | Y_S$, and instead test the corresponding statement including G, *i.e.*, $Y_j \bot Y_k | \{Y_S, G\}$.

Testing $\mathbf{Y}_j \perp \mathbf{G} | \mathbf{Y}_{\mathbf{S}}$: Our test for statement (A) is based on the intuition that \mathbf{Y}_j is independent of $\mathbf{G} = [\mathbf{G}_1 \cdots \mathbf{G}_p]$ given $\mathbf{Y}_{\mathbf{S}}$, whenever there is no direct genetic effect on \mathbf{Y}_j (*i.e.*, $\mathbf{G}_j = 0$), and all directed paths from *G* to Y_j are blocked by the set $Y_S = \{Y_m : m \in S\}$. In particular, if *S* is the empty set, there should not be any directed path from *G* to Y_j . Because directed paths from *G* to Y_j , will generally introduce some genetic variance in \mathbf{Y}_j , the idea is to test whether there is significant genetic variance in the conditional distribution of \mathbf{Y}_j given $\mathbf{Y}_{\mathbf{S}} = \tilde{y}_S$. This is done as follows:

- 1. When $K = ZZ^t$, we use the classical F-test in a one-way ANOVA, with **X** and \tilde{y}_S as covariates. Technically, this is an ANCOVA (analysis of covariance), where the treatment factor genotype is tested conditional on the covariates being in the model.
- 2. For other *K*, one can use a likelihood ratio test (LRT). The asymptotic distribution under the null-hypothesis is a mixture of a point mass at zero and a chi square.

In both cases, it is assumed that the conditional distribution of $\mathbf{Y}_{\mathbf{j}}$ given $\mathbf{Y}_{\mathbf{S}} = \tilde{y}_{S}$ is that of a single-trait mixed model, the mean being a linear regression over the conditioning traits. This assumption is made mathematically precise below in Equations 12 and 14.

Testing $Y_i \perp Y_k | \{G, Y_s\}$: For statement (B), we mostly use the residual covariance (RC) test, which is based on the conditional distribution of \boldsymbol{Y}_j and \boldsymbol{Y}_k given the observed $\mathbf{Y}_{\mathbf{S}} = \tilde{y}_{S}$. It is assumed that this distribution is that of a bivariate MTM, again with the mean being a linear regression over the conditioning traits; see Equations 13 and 14 below. Assuming the bivariate MTM, we test whether the residual covariance is zero, using the LRT described in File S1.3. The underlying idea is that a nonzero residual covariance must be the consequence of an edge $Y_i \rightarrow Y_k$ or $Y_k \rightarrow Y_i$, because of the assumed normality and causal sufficiency. On the other hand, a nonzero genetic covariance may also be due to covariance between direct genetic effects on these variables, or due to a genetic effect on a common ancestor. The RC-test therefore compares the full bivariate mixed model with the submodel with diagonal residual covariance, while accounting for all genetic (co)variances. The RC-test is not to be confused with a test for zero genetic covariance. The latter is often useful for data exploration, but has no role in PCgen (although in File S1.4 we describe a LRT test that is implemented in our software).

An alternative to the RC-test is the RG-test [Residuals of GBLUP, i.e., the best linear unbiased prediction of the genetic effects]. Fitting the MTM (Equation 7), we obtain the BLUP \mathbf{U}^* of the total genetic effects $\mathbf{U} = \mathbf{G}\Gamma$, and the BLUE \tilde{B}^* of the fixed effects. We then test the significance of partial correlations among residuals, *i.e.*, the columns of $\mathbf{Y} - \mathbf{U}^* - \mathbf{X}\tilde{B}^*$. When U^{*} is close enough to U, it follows from Equations 6 and 7 that the covariance of $vec(\mathbf{Y} - \mathbf{U}^*)$ is approximately $(\Gamma^t \Sigma_E \Gamma) \otimes I_n$, *i.e.*, that of independent samples, without any genetic relatedness. This approach is very similar to the work of Valente et al. (2010) and Töpner et al. (2017), who instead took a fully Bayesian approach to predict U. In either case, the performance of the RG-test critically depends on the prediction error $(\mathbf{U}^* - \mathbf{U})$. As mentioned before, fitting an MTM is usually challenging for >5-10 traits; we therefore also consider residuals of single-trait GBLUP, as an approximation.

PCres: reconstructing only trait-to-trait relationships: Testing only conditional independencies of the form (B), one can reconstruct the graph $\mathcal{G}_{\mathcal{Y}}$ of trait-to-trait relations (see the green boxes in Figure 2). Moreover, if this is done with the RG-test, the algorithm is very similar to the residual approaches of Valente *et al.* (2010) and Töpner *et al.* (2017). Staying within the context of the PC-algorithm, and using residuals from GBLUP, we will call this PCres. As for the RG-test in PCgen, PCres can be based on residuals of either single or multi-trait mixed models.

Software: In our R-package pcgen, we implemented PCgen for the case $K = ZZ^t$. PCres is implemented for $K = ZZ^t$, K = A, as well as $K = ZAZ^t$. Moreover, PCres can be based on either residuals of the full MTM (Equation 7) (only for small numbers of traits), or from univariate models (the default). Tables 1 and 2 in File S2 provide a complete overview of the options, with the required R-commands. The package is freely available at https://cran.r-project.org/web/ packages/pcgen/index.html. pcgen is built on the pcalg package (Hauser and Bühlmann 2012; Kalisch *et al.* 2012), in which we modified the orientation rules and the default conditional independence test.

Assessing uncertainty: The PC-algorithm is asymptotically correct, in the sense that the underlying CPDAG is recovered if conditional independence can be tested without error (Spirtes *et al.* 2001). In Theorem 2 below, we provide a similar consistency result for PCgen. In practice however, type-I or type-II errors are likely to occur, leading to incorrect edges in the graph. Depending on the significance level α used in each test, there may be more type-I errors (large α) or rather more type-II errors (small α). Reliable control of the (expected) false positive rate, or total number of false positives, remains challenging; see the *Discussion (Assessing uncertainty*). We will therefore just consider the *P*-values as they

are, and analyze the real datasets for different significance thresholds. Following Kalisch and Bühlmann (2007) and Kalisch *et al.* (2012), we report, for each remaining edge, the largest *P*-value found across all conditioning sets for which the edge was tested.

Extensions of PCgen: File S3 describes several extensions of PCgen, which are partly implemented in our software. Among others, the causal graph \mathcal{G} and PCgen could be extended with fixed effect QTL, and PCgen can be sped up by starting with a skeleton obtained from PCres ("prior screening"). As in the pcalg-package, it is possible to restrict the maximum size of the conditioning sets, also to improve computation time.

Data availability

The authors state that all data necessary for confirming the conclusions presented in the article are represented fully within the article. The maize and rice data used below can be accessed at https://doi.org/10.15454/IASSTN and https://doi.org/10.6084/m9.figshare.7964357.v1, respectively. Supplemental materials are available at figshare: https://doi.org/10.6084/m9.figshare.11635392.

Results

Simulations with randomly drawn graphs

To compare the different algorithms, we simulated random GSEMs by randomly sampling the sets *D* (defining the traits with direct genetic effects) and the covariance matrices Σ_G , combined with randomly drawn DAGs over the traits ($\mathcal{G}_{\mathcal{Y}}$). Traits were simulated for an existing population of 256 maize hybrids (Millet *et al.* 2016). Two replicates of each genotype were simulated. Given the 256 × 256 additive relatedness matrix *A* based on 50k SNPs, genetic effects were simulated such that $vec(\mathbf{G}) \sim \Sigma_G \otimes (ZAZ^t)$ (*i.e.*, the vector of genetic effects for all the replicates, and all traits). File S4.1 provides further details, such as the magnitude of genetic (co) variances. We focus here on the comparison of:

- 1. PCgen based on the replicates, assuming $K = ZZ^t$ (*i.e.*, ignoring *A*). By default, we apply the prior-screening with PCres.
- 2. PCres (replicates): PCres based on residuals from univariate GBLUP, again using only the replicates.
- 3. PCres (means): PCres based on residuals from multivariate GBLUP, using genotypic means and the relatedness matrix *A* that was used to simulate the data.

Table S2 provides results for variations on these algorithms, including PCgen without prior screening. In all simulations, the significance threshold was $\alpha = 0.01$. The effect of sample size, and the trade-off between power and false positives as function of α , was already investigated by Kalisch and Bühlmann (2007) for the standard PC-algorithm, and is likely to be similar for PCgen.

We separately evaluated the reconstruction of $\mathcal{G}_{\mathcal{Y}}$ and the edges $G \rightarrow Y_j$, as the latter is only possible with PCgen. To

assess the difference between estimated and true skeleton of $\mathcal{G}_{\mathcal{Y}}$, we considered the true positive rate (TPR), the true discovery rate (TDR), and the false positive rate (FPR). Additionally, we used the Structural Hamming Distance (SHD), which also takes into account the orientation of the edges. File S4.2 provides definitions of these criteria. Reconstruction of $G \rightarrow Y_j$ is assessed only in terms of TPR, TDR, and FPR, as these edges can have only one orientation.

Simulation results: We first performed simulations with p = 4 traits (scenario 1), with each potential edge between traits occurring in the true graph with probability $p_t = 1/3$. Hence, for any given trait, the expected number of adjacent traits was $(p-1)p_t = 1$. The edges $G \rightarrow Y_j$ were included in the true graph with probability $p_g = 1/2$. In a related set of simulations (scenario 2), p_t was increased to 0.5, giving denser graphs. In both scenarios, PCgen reconstructed the edges $G \rightarrow Y_i$ with little error, the average TPR being ~0.97 and FPR \sim 0.03 (Table 1). In the first scenario, about onethird of the actual edges between traits was not detected with PCgen (TPR ≈ 0.65 , *i.e.*, the proportion of true edges that was discovered). At the same time, the number of false edges in $\mathcal{G}_{\mathcal{V}}$ was very low, which is also reflected in high TDR values (the proportion of edges in the reconstruction that is true). In scenario 2, the TPR, FPR, and TDR all increased. Hence, for denser graphs, more of the true edges were found, at the expense of a somewhat higher number of false edges.

PCres (replicates) outperformed PCres (means), in spite of the use of univariate GBLUP, and ignoring the actual relatedness matrix. Hence, the information contained in the replicates appears much more important than the precise form of the relatedness matrix, or unbiased estimation of genetic correlations. The performance of PCres strongly depends on the prediction error of the GBLUP, and, in line with the results of Kruijer et al. (2015), this error appeared lowest when using the replicates. The use of both the replicates and the markerbased GRM (*i.e.*, assuming $K = ZAZ^t$, as the data were generated), further improved performance, but only slightly (Table S1, PCres-uni-RA). Unsurprisingly, the MTM required for PCres (means) was computationally more demanding, and could often not be obtained for more than four traits. Motivated by this computational advantage, and the statistical advantages mentioned in the Discussion, all analyses in the remainder will consider only PCgen and PCres based on replicates.

For trait-to-trait relations, PCgen and PCres (replicates) had very similar performance in terms of TPR, TDR, and FPR. However, PCgen substantially improved the orientation of these edges, as shown by the reduced SHD. This is a consequence of the additional edges $G \rightarrow Y_j$ in the underlying graph: because of the fixed orientation of these edges, this generally increases the number of v-structures, and, hence, the number of between-trait edges $Y_j - Y_k$ that can be oriented. See again the example in Figure 2.

To assess performance in higher dimensions, we simulated data sets with p = 20 traits, $p_g = 0.3$, and $p_t = 0.1$ (scenario 3),

Table 1 Performance of PCgen and residuals-based approaches, averaged over 500 simulated datasets per scenario.

	$\mathcal{G}_{\mathcal{Y}}$				$G \rightarrow Y_j$		
	TPR	FPR	TDR	SHD	TPR	FPR	TDR
Scenario 1 ($p = 4$)	$(p_t = 1/3)$				$(p_q = 0.5)$		
PCgen	0.647	0.006	0.981	0.442	0.982	0.026	0.995
PCres (replicates)	0.650	0.039	0.908	1.410			
PCres (means)	0.521	0.390	0.438	3.174			
Scenario 2 ($p = 4$)	$(p_t = 0.5)$				$(p_q = 0.5)$		
PCgen	0.804	0.033	0.986	1.246	0.976	0.031	0.995
PCres (replicates)	0.819	0.073	0.939	2.320			
PCres (means)	0.672	0.364	0.659	3.628			
Scenario 3 ($p = 20$)	$(p_t = 0.1)$				$(p_q = 0.3)$		
PCgen	0.895	0.002	0.985	6.806	0.969	0.018	0.991
PCres (replicates)	0.911	0.004	0.961	9.874			
Scenario 4 ($p = 100$)	$(p_t = 0.01)$				$(p_q = 0.1)$		
PCgen	0.959	0.001	0.942	27.288	0.976	0.022	0.943
PCres (replicates)	0.962	0.001	0.940	38.410			

SE for the TPR, FPR, and TDR were between 0.001 and 0.015. SE for the SHD were ~0.06 (scenarios 1 and 2), 0.18 (scenario 3), and 0.28 (scenario 4). For the performance of other variants of PCgen and PCres in scenarios 1 and 2, see Table S2. In scenario 4, we used PCgen with the RG-test (PCgen-RG-uni); in the other scenarios we used the RC-test, with prior screening (PCgen-RC-screening). All acronyms are explained in Table 1 in File S2. PCres (replicates) and PCres (means) refer to PCres-uni-R and PC-multi-A.

and with p = 100, $p_g = 0.1$, and $p_t = 0.01$ (scenario 4). Both scenarios consider sparse graphs; denser graphs can be analyzed as well, but, for p > 20–30, require several hours, or even days, unless the size of the conditioning sets is restricted, or our implementation of PCgen would be parallelized. Here, we limited the size of conditioning sets to three (scenario 3) and two (scenario 4). As in the first two scenarios, PCgen achieved a strong reduction in SHD, and reliable reconstruction of the direct genetic effects (Table 1).

To assess the effect of thresholding the size of conditioning sets, we simulated 200 datasets with p = 10 traits and a relatively dense graph ($p_g = 0.4$ and $p_t = 4/9$), and used PCgen with various thresholds (Table S3). The restricted maximum size means that a certain number of conditional independence tests is skipped, which may lead to extra false positives. However, the thresholding is only done in PCgen itself and not in the prior screening with PCres (which is much faster, and already removes most false edges). Consequently, thresholding had very little effect on the reconstruction of trait-to-trait relations ($\mathcal{G}_{\mathcal{Y}}$), but did lead to a higher FPR in the reconstruction of the direct genetic effects (0.07 without thresholding, 0.08 with m = 3, and 0.48 with m = 1). Also, the accuracy in the orientations of $\mathcal{G}_{\mathcal{Y}}$ slightly decreased (SHD increasing from 15.9 to 16.2).

In another set of simulations, we explored the effect of measurement error. As expected, increasing amounts of measurement error decreased the power to detect between-trait edges as well as direct genetic effects (Table S4). However, the loss in power could largely be compensated by increasing the number of replicates, or the number of genotypes. The latter was most effective for between-trait edges, while increased replication gave the highest power for the edges $G \rightarrow Y_i$.

In our final set of simulations (Table S5), we explored the effect of strong correlations in Σ_G , *i.e.*, when Assumption 5 is close to being violated. We simulated an example with two traits whose direct genetic effects had unit variance, and increasing covariance (0, 0.5, and 0.95). The corresponding

TPR values for the genetic effects were respectively 0.94, 0.85, and 0.60. Consequently, even in the presence of strong correlations, PCgen still had some power to detect direct genetic effects.

Simulations using a crop-growth model

We also simulated data using the popular crop growth model APSIM-wheat (Keating et al. 2003; Holzworth et al. 2014). Compared to the preceding simulations, this represents a more challenging scenario, as several of the underlying assumptions are violated. In particular, the data-generating process introduces nonlinearities and latent variables. We simulated 12 traits for an existing wheat population of 199 genotypes, with three replicates each. The traits included seven primary traits, four secondary traits, and yield (Y). File S5 provides further details, and trait acronyms are given in Table S6. Traits were simulated by running a discrete dynamic model from the beginning (t = 0) to the end (t = T)of the growing season. Motivated by the fact that some trait measurements are destructive, observations are taken only at t = T. Figure S1A shows the summary graph, defining the causal effects from one time-step to the next (Peters et al. 2017). We note that the summary graph does not directly describe the distribution of the traits at t = T (obtained by marginalizing over previous time points), which can be represented by an ancestral graph (Richardson and Spirtes, 2002). As such graphs are outside the scope of this work, we investigate the extent to which we can reconstruct the summary graph, given observations taken at t = T. There are direct genetic effects on all of the primary traits, which have heritability 0.9. The genetic effects originate from 300 trait-specific QTL, with randomly drawn effect sizes. There are no direct genetic effects on secondary traits and yield.

Compared to the simulations above, it turned out to be much harder to detect the absence of direct genetic effects: in the PCgen reconstruction, all 12 traits had such effects (Figure S1B; highest P-value: 1.7×10^{-4}). These false positives seemed to be a consequence of the nonlinearities in the data-generating distribution, which are not accounted for in our tests. The reconstructed trait-to-trait relations were mostly correct, except for the missing edge $GN \rightarrow Y$, and one incorrect orientation $(Y \rightarrow GW)$. PCres made the same errors (Figure S1C), with an additional false arrow (MGS - SP). The standard PC-stable algorithm applied to all traits and QTL led to many more errors (Figure S1D), such as the false edge between GW and RUE, the missing edge $TFI \rightarrow FT$, and some incorrect orientations. These errors occurred because, for various traits Y_i, many QTL-effects were removed from the graph, *i.e.*, for some set of traits Y_S , the conditional independence $Y_i \perp QTL | Y_s$ was mistakenly accepted. This, in turn, led to problems in the remaining tests, where part of the genetic variance was not taken into account. We emphasize that all 300 QTL were available to the PC-algorithm, and no other markers were provided. Hence, the poor performance in this case is really a consequence of the small effects, rather than the difficulty of QTL detection.

Two case-studies

We now use PCgen to analyze real data from four field trials and one experiment in a phenotyping platform. In all network reconstructions, we used a significance threshold of $\alpha = 0.01$. Reconstructions with $\alpha = 0.001$ are shown in Figures S2 and S4. Figure S5 and Table S9 contain *P*-values for the remaining edges. In all datasets, we removed traits that were derived as sums or ratios of other traits, rather than being directly measured. In particular, the maize data do not contain grain number, which was defined as the ratio of yield over grain weight. We return to this issue in the *Discussion*.

Maize: First, we analyze the field trials described by Millet *et al.* (2016, 2019), with phenotypic data for 254 hybrids of maize (*Zea mays*). We consider a subset of four trials, representing four (out of a total of five) different environmental scenarios described in Millet *et al.* (2016). See Table S8 for an overview. The scenarios were derived from physiological knowledge, crop-growth models, and environmental sensors in the fields. Scenarios were defined as a combination of well-watered or water-deficient conditions (WW *vs.* WD) and temperature. The latter was classified as "Cool" (average maximum and night temperature below respectively 33 and 20°), "Hot" (above 33 and 20°) or "Hot (days)" (maximum temperature above 33, night temperature below 20). Most trials included seven traits:

- three height traits, *i.e.*, tassel height (*TH*), ear height (*EH*), and plant height (*PH*); the latter is missing in the Ner12R trial.
- two flowering time traits: anthesis (*A*) and silking (*Sk*), which are male and female flowering, respectively.
- two yield-related traits: grain weight (*GW*) and yield (*Y*).

Table S7 provides an overview of trait acronyms. Each trial was laid out as an alpha-lattice design, with either two or three

In the PCgen reconstruction, all traits have direct genetic effects, and traits mostly cluster according to their biological category (height, flowering, and yield-related), especially in the WW scenarios (Figure 4, A and B). In the Ner13W and Ner12R trials (Figure 4, B and C), there are edges between yield and respectively tassel- and plant-height, but these (conditional) dependencies are weak and disappear in the reconstruction with $\alpha = 0.001$ (Figure S2). Much stronger are the edges between yield and the flowering traits in the water-deficit trials (Figure 4, C and D); the corresponding conditional independence tests gave highly significant Pvalues for all of the considered conditioning sets (Table S9). By contrast, in the trial without heat or drought stress (Kar12W), the Y - Sk and Y - A edges were already removed in the test conditioning only on the genetic effects; Figure S3 provides an illustration. The relation between yield and delay in silking in maize is well known [see e.g., Borrás et al. (2007) and Araus et al. (2012)]. In the most stressed environment (Bol12R), there is an additional edge between plant height and silking. This may relate to the fact that the timing of anthesis determines the number of phytomeres (number of internodes and leaves) that a plant will generate, which, in turn, affects plant height (McMaster et al. 2005). The strong correlation between anthesis (A) and silking (Sk) may explain the presence of the edge PH - Sk (rather than PH - A).

Finally, apart from the Bol12R trial, there is never an edge between *Y* and *GW*, which seems due to the choice of the genetic material (giving little variation in grain weight) and the design of the trials (targeting stress around flowering time, rather than the grain filling period). See Millet *et al.* (2016) for further details. For all trials, the structure of the graphs is such that none of the between-trait edges can be oriented (technically, this is due to a lack of v-structures). However, for some of these edges, physiological knowledge clearly suggests a certain orientation, in particular for Sk - Y and GW - Y.

The trials also illustrate the difference between the total genetic covariance (V_G) and the covariance among direct genetic effects, as defined by Σ_G . For most pairs of traits, the total genetic correlation (ρ_g) was between 0.3 and 0.9 (Table S10). The (total) genetic correlation between yield and silking was strongly negative in both WD trials (-0.44 and -0.61), and, in the Bol12R trial, also for yield and anthesis (-0.43). In all trials, genetic correlation with *GW* was negative for most traits, but not always significant. In the Kar12W trial for example, we found $\rho_g = -0.010$ for *GW* and *PH*, and $\rho_g = -0.435$ for *GW* and *Sk* (silking). In both cases, the two traits are d-separated in the graph (conditioning on $\{G\}$), but only for *Sk* is the genetic covariance significant ($p = 1.31 \times 10^{-9}$).



Figure 4 Estimated networks with $\alpha = 0.01$, for four of the DROPS field trials. Trait categories are flowering (orange), height (blue), and yield (brown). Each trial represents a different environmental scenario, arising from well-watered (WW) or water-deficit (WD) conditions, and different temperatures (see text). (A) Kar12W, (WW, Cool). (B) Ner13W (WW, Hot). (C) Ner12R (WD, Hot (days)). (D) Bol12R (WD, Hot). Trait acronyms are given in Table S7.

As we have seen in the examples following Equation (5), the existence of an edge between two traits in the graph does not necessarily imply a strong genetic correlation. In other words, having a shared genetic basis is not the same thing as the presence of a causal effect, found after conditioning on the genetic effects and other traits. In the Ner12R trial, for example, there is no edge between yield and grain weight, but a significant genetic correlation, whereas in the Bol12R trial it is the other way round.

Rice: Next we analyze 25 traits measured on 274 indica genotypes of rice (Oryza sativa) under water deficit, reported by Kadam et al. (2017). Three replicates of each genotype were phenotyped in a randomized complete block design, and block was included as a covariate in all conditional independence tests. Tests were restricted to conditioning sets of, at most, four traits. A first run of PCgen produced several inconsistencies in the genetic effects, *i.e.*, traits with significantly positive heritability but without a partially directed path coming from the node G. We therefore applied the correction described in File S3, adding edges $G \rightarrow Y_i$ for all traits with this inconsistency, and then re-ran PCgen. The final reconstruction is given in Figure 5, where traits are grouped into 3 shoot morphological traits (blue), 1 physiological trait (rose), 13 root morphological traits (green), 5 root anatomical traits (gray), and 3 dry matter traits (orange).

After correcting the inconsistencies, there were nine traits without a direct genetic effect. Five of these (MRL, ART, RL2530,

RL3035, and RL35) were completely isolated in the graph, without edges connecting to any other trait. All of these traits are related to either root length, or to the length of thicker roots, which contribute to drought adaptation under field conditions (Uga et al. 2013). However, as the experiment was done in pots, roots were constrained in their exploration range and therefore genotypic differences in root length would not translate into differential access to water and biomass (Poorter et al. 2012). Four other traits (TRWD, RW, RL0510, and RV) had at least one adjacent trait in the graph, but no direct genetic effect. At a lower significance level ($\alpha = 0.001$, Figure S4), direct genetic effects disappeared also for cumulative water transpiration (CWT), and for three root anatomical traits (RD, CD, and SD). For RV (root volume), a direct genetic effect was only present with $\alpha = 0.001$, which was an artifact of the way the initial consistencies were resolved.

Traits related to root surface area (SA), root volume (RV), and roots with small diameter class (RL005, RL1015) had direct genetic effects, and were connected among each other. As expected, traits related to root volume and area influenced root weight and total root weight density (RW, TRWD). In the reconstruction with $\alpha = 0.001$, cumulative water transpired (CWT) was affected by stem and leaf weight (SW, LW), and by RL0510, in agreement with the physiological knowledge that water transpiration is influenced by water demand (related to the above-ground biomass) and water supply (related to the roots' water uptake capacity). The corresponding edges were also present in the reconstruction with $\alpha = 0.01$, where,



Figure 5 PCgen-reconstruction for the rice data from Kadam *et al.* (2017), with $\alpha = 0.01$. Five traits (MRL, ART, RL2530, RL3035, and RL35) are not shown, as they were completely isolated in the graph, without any connections to other traits or *G*. Trait acronyms are given in Table S11.

however, they could not be oriented because of the denser network (in particular, the presence of $G \rightarrow CWT$). Root anatomical traits (LMXD, SD, CD, and RD) appeared as a separate module, not related to the plant water dynamics, suggesting that root anatomy had a smaller impact on water uptake compared with root biomass.

Statistical properties of PCgen

We now investigate a number of statistical issues: the assumptions required for asymptotic consistency of PCgen, the assumptions required for faithfulness, and properties of the conditional independence tests. Readers primarily interested in the application of PCgen could skip this section and continue with the *Discussion*. Proofs of Theorems 1–6 are given in Appendix A.

Consistency: Asymptotic consistency holds if, for increasing sample size, the probability of finding the correct network converges to 1. Correct in this context means that we recover the CPDAG that contains the underlying DAG. Consistency of the PC-algorithm was shown by Spirtes *et al.* (2001) (for low dimensions) and Kalisch and Bühlmann (2007) (for high dimensions). These authors distinguished between consistency of the oracle version of PC, where conditional independence information is available without error, and the sample version, where conditional independence is

obtained from statistical tests. For PCgen we will focus on the oracle version and consistency of the skeleton, leaving the sample version and the correctness of the orientations for future work.

As for the standard PC-algorithm, consistency of PCgen requires equivalence between conditional independence and d-separation in the graph. Part of this is the Markov property, which states that d-separation of two nodes in the graph, given a set of other nodes, implies conditional independence of the corresponding random variables. The converse (conditional independence implying d-separation) is known as faithfulness. The following result provides the Markov property for SEM with genetic effects. The proof (Appendix A.9) is a straightforward adaptation of Pearl's proof for general SEMs (Pearl 2009).

Theorem 1 Suppose we have a GSEM as defined by Equation (4), with a graph \mathcal{G} as defined in the *Materials and Methods*, and satisfying Assumptions 1–4. Then, the global Markov condition holds for \mathcal{G} and the joint distribution of $\mathbf{G}, \mathbf{Y}_1, \ldots, \mathbf{Y}_p$. In particular, d-separation of Y_j and G given Y_S implies $\mathbf{Y}_j \parallel \mathbf{G} \mid \{\mathbf{Y}_S\}$, and d-separation of Y_j and Y_k given $\{Y_S, G\}$ implies $\mathbf{Y}_j \parallel \mathbf{Y}_k \mid \{\mathbf{Y}_S, \mathbf{G}\}$, for all traits \mathbf{Y}_j and \mathbf{Y}_k and subsets \mathbf{Y}_S .

If we now assume faithfulness, the preceding result directly gives the equivalence between conditional independence and d-separation. This, in turn, implies that PCgen will recover the correct skeleton: **Theorem 2** Let $dsep(\mathcal{G})$ denote d-separation in the graph \mathcal{G} . Suppose we have a GSEM as in Theorem 1, and we make the additional assumptions of faithfulness:

$$\mathbf{Y}_{\mathbf{j}} \perp \mathbf{Y}_{\mathbf{k}} | \{ \mathbf{Y}_{\mathbf{S}}, \mathbf{G} \} \Rightarrow Y_{j} dsep(\mathcal{G}) Y_{k} | \{ Y_{S}, \mathbf{G} \}$$
(9)

$$\mathbf{Y}_{\mathbf{j}} \bot \mathbf{G} | \{ \mathbf{Y}_{\mathbf{S}} \} \Rightarrow Y_{j} dsep(\mathcal{G}) \ G | \{ Y_{S} \}, \tag{10}$$

for all traits Y_j and Y_k and subsets Y_s . The oracle version of PCgen then gives the correct skeleton.

Faithfulness: For our first faithfulness condition (expression 9) to hold, it suffices to have faithfulness for the graph without genetic effects:

Theorem 3 Let $P_{\mathbf{Y}|\mathbf{U}}$ denote the joint distribution of $\mathbf{Y}_1, \ldots, \mathbf{Y}_{\mathbf{p}}$ conditional on $\mathbf{U} = \mathbf{G}\Gamma$, the matrix of total genetic effects. Then $\mathbf{Y}_j \perp \mathbf{Y}_k | \{\mathbf{Y}_S, \mathbf{G}\}$ is equivalent with $\mathbf{Y}_j \perp \mathbf{P}_{\mathbf{Y}|\mathbf{U}} \mathbf{Y}_k$, and $Y_j dsep(\mathcal{G}) Y_k | \{Y_S, G\}$ is equivalent with $Y_j dsep(\mathcal{G}_{\mathcal{Y}}) Y_k | Y_S$. Therefore, (9) holds if

$$\mathbf{Y}_{\mathbf{j}} \mathbf{I}_{P_{\mathbf{y}_{iii}}} \mathbf{Y}_{\mathbf{k}} | \{ \mathbf{Y}_{\mathbf{S}} \} \Rightarrow Y_{j} \, dsep(\mathcal{G}_{\mathcal{Y}}) \, Y_{k} | Y_{S}. \tag{11}$$

Consequently, we can rephrase (9) in terms of a faithfulness assumption for the analogous SEM without genetic effects. A necessary (but not sufficient) condition for this is that contributions from different paths do not cancel out (Appendix A.5).

The second faithfulness statement (10) involves d-separation of Y_j and G, and requires that the genetic effects are not collinear. If, for example, we have $\mathbf{Y}_3 = \mathbf{Y}_1 + \mathbf{Y}_2 + \mathbf{E}_3$, with $\mathbf{Y}_1 = \mathbf{G}_1 + \mathbf{E}_1$, $\mathbf{Y}_2 = \mathbf{G}_2 + \mathbf{E}_2$, and $\mathbf{G}_2 = -\mathbf{G}_1 = \mathbf{G}$, it follows that $\mathbf{Y}_3 = \mathbf{E}_1 + \mathbf{E}_2 + \mathbf{E}_3$. Consequently, because $\mathbf{G}_3 = (0, \dots, 0)^t$, we find that \mathbf{Y}_3 and $\mathbf{G} = [\mathbf{G}_1 \ \mathbf{G}_2 \ \mathbf{G}_3]$ are marginally independent, but, in the graph \mathcal{G} , the nodes Y_j and G are not d-separated by the empty set, as there are directed paths $G \rightarrow Y_2 \rightarrow Y_3$ and $G \rightarrow Y_1 \rightarrow Y_3$. Conversely, if \mathbf{G}_1 and \mathbf{G}_2 are not perfectly correlated, this violation of faithfulness cannot occur. The following theorem shows that marginal independence always implies d-separation. We conjecture (but could not prove) that (10) also holds for nonempty conditioning sets.

Theorem 4 Suppose we have a GSEM satisfying Assumptions 1–5, and faithfulness for the graph without genetic effects, given by (11). Then implication (10) holds for $S = \emptyset$, *i.e.*, marginal independence implies d-separation of Y_i and G.

Hence, faithfulness involving Y_j and G requires (at least) absence of collinearities between genetic effects, as well as faithfulness for the corresponding SEM without genetic effects.

Properties of the tests: Theorem 2 provides consistency of the oracle version of PCgen, where conditional independence information is available without error. Proving consistency of the sample version is challenging for two reasons. First, the assumptions made for our conditional independence tests may not always hold, introducing approximation errors. Second, even without these errors, the probabilities of type-I and type-II errors still need to converge to zero with increasing sample size. This is well known for the PC-algorithm with

independent Gaussian data (Kalisch and Bühlmann 2007), but more difficult to establish in the presence of genetic effects. Here, we address the first issue, leaving the second for future work.

Our tests for conditional independence statements (A) and (B) (*i.e.*, $Y_j \bot G | \{Y_S\}$ and $Y_j \bot Y_k | \{Y_S, G\}$) rely on the conditional distributions of, respectively, Y_j and $vec([Y_j, Y_k])$, given the observed Y_S :

$$\begin{split} \mathbf{Y_{j}} | \textit{vec}(\mathbf{Y_{S}}) &= \textit{vec}(\tilde{y}_{S}) \ \sim \ N\Big(\mu_{j|S}, \Sigma_{j|S}\Big) \\ & \Big(\begin{array}{c} \mathbf{Y_{j}} \\ \mathbf{Y_{k}} \end{array} \Big) | \textit{vec}(\mathbf{Y_{S}}) &= \textit{vec}(\tilde{y}_{S}) \ \sim \ N\Big(\left(\begin{array}{c} \mu_{j|S} \\ \mu_{k|S} \end{array} \right), \Sigma_{jk|S} \Big). \end{split}$$

The normality of these distributions directly follows from the assumed normality of the genetic and residual effects. We made the following assumptions about the form of their covariance and mean:

1. The covariance matrix $\Sigma_{j|S}$ is that of a single-trait mixed model with the same relatedness matrix *K* assumed in the GSEM, *i.e.*,

$$\Sigma_{j|S} = \sigma_G^2(j|S)K + \sigma_E^2(j|S)I_n, \qquad (12)$$

for some variance components $\sigma_G^2(j|S)$ and $\sigma_E^2(j|S)$.

2. The covariance matrix $\Sigma_{jk|S}$ is that of a bivariate MTM, again with the same *K* assumed in the GSEM:

$$\Sigma_{jk|S} = V_G(jk|S) \otimes K + V_E(jk|S) \otimes I_n,$$
(13)

for some 2 × 2 matrices $V_G(jk|S)$ and $V_E(jk|S)$.

The conditional means μ_{j|S} and μ_{k|S} are linear regressions over the conditioning traits:

$$\mathbf{X}B\boldsymbol{\gamma}_{j} + \tilde{\boldsymbol{y}}_{S}\boldsymbol{\beta}_{S}^{(j)}, \quad \mathbf{X}B\boldsymbol{\gamma}_{k} + \tilde{\boldsymbol{y}}_{S}\boldsymbol{\beta}_{S}^{(k)}, \tag{14}$$

where $XB\gamma_j$ is the marginal mean of Y_j (see Equation 8), and $\beta_S^{(j)}$ and $\beta_S^{(k)}$ are $|S| \times 1$ vectors of regression coefficients.

In the following theorems we show that when $K = ZZ^t$, the assumptions in Equations 12 and 13 always hold, *i.e.*, they directly follow from our GSEM model.

Theorem 5 When $K = ZZ^t$, the distribution of $vec([\mathbf{Y}_j, \mathbf{Y}_k])|vec(\mathbf{Y}_s)$ has covariance of the form given by Equation 13, *i.e.*, that of a bivariate MTM. Moreover, under faithfulness condition (9), the residual covariance in the MTM is zero if and only if $\mathbf{Y}_j \perp \mathbf{Y}_k | \{\mathbf{Y}_s, \mathbf{G}\}$

Theorem 6 Suppose we have a GSEM as described in Theorem 1, with $K = ZZ^t$. Then, the covariance of $\mathbf{Y}_j | \mathbf{Y}_S$ is of the form $\sigma_G^2(j|S)K + \sigma_E^2(j|S)I_n$, for any conditioning set S. Moreover, assuming the faithfulness condition (10) and $\Sigma_G[D, D]$ of full rank (Assumption 5), $\sigma_G^2(j) = \sigma_G^2(j|\emptyset)$ is zero if and only if $\mathbf{Y}_j \perp \mathbf{G}$.

Apart from the covariance structure, these theorems address the correctness of our tests. In particular, Theorem 5 shows that the residual covariance in the distribution of $\{Y_i, Y_k\}|Y_S$ is indeed the right quantity to test statement

(B). Similarly, the genetic variance in the conditional distribution of $\mathbf{Y}_{j}|\mathbf{Y}_{S}$ is the relevant thing for testing (A). This appears to be true for any conditioning set *S*, although (in Theorem 6) we could prove it only for the empty conditioning set, because faithfulness is required (which we also established only for $S = \emptyset$; see Theorem 4).

The situation is different for the assumption in Equation 14, regarding the conditional means: even when $K = ZZ^t$, it holds for certain conditioning sets and not for others. We illustrate this with the following example. Suppose that $\mathbf{Y}_1 = \mathbf{G}_1 + \mathbf{E}_1$ and $\mathbf{Y}_2 = \lambda \mathbf{Y}_1 + \mathbf{E}_2$, with independent vectors $\mathbf{G}_1 \sim N(0, \sigma_{G,1}^2 K)$, $\mathbf{E}_1 \sim N(\sigma_{E,1}^2 I_n)$, and $\mathbf{E}_2 \sim N(\sigma_{E,2}^2 I_n)$. Then, the graph \mathcal{G} is given by $G \rightarrow Y_1 \rightarrow Y_2$. There is no edge $G \rightarrow Y_2$, although this is not essential for the example. The distributions are given by

$$\begin{split} \mathbf{Y}_{1} &\sim N(0, \Sigma_{1}) = N\Big(0, \sigma_{G,1}^{2}K + \sigma_{E,1}^{2}I_{n}\Big), \\ \mathbf{Y}_{2} &\sim N(0, \Sigma_{2}) = N\Big(0, \lambda^{2}\sigma_{G,1}^{2}K + \Big(\lambda^{2}\sigma_{E,1}^{2} + \sigma_{E,2}^{2}\Big)I_{n}\Big), \\ Cov(\mathbf{Y}_{1}, \mathbf{Y}_{2}) = \Sigma_{12} = \lambda\Big(\sigma_{G,1}^{2}K + \sigma_{E,1}^{2}I_{n}\Big) = \lambda\Sigma_{1}. \end{split}$$

The conditional mean of \mathbf{Y}_2 given $\mathbf{Y}_1 = y_1$ is $\mu_{2|1} = \Sigma_{12} \Sigma_1^{-1} y_1 = \lambda y_1$. As expected given the graph, the conditional mean is a simple linear regression on Y_1 . However, the conditional mean of \mathbf{Y}_1 given $\mathbf{Y}_2 = y_2$, equals

$$\mu_{1|2} = \lambda \Sigma_1 \left(\lambda^2 \Sigma_1 + \sigma_E^2(2) I_n \right)^{-1} y_2$$

which is a linear transformation, but not a multiple of y_2 . In summary, our models for $Y_i | Y_s$ and $\{Y_i, Y_k\} | Y_s$ are sometimes misspecified in terms of the mean, although still correct in terms of covariance, provided $K = ZZ^t$ (Theorems 5 and 6). Despite the approximation error occurring sometimes for the conditional means, our tests still seem to perform reasonably, as shown in the simulations above. The assumption in (14) is more problematic if relations between traits are nonlinear. Suppose, for example, that, for each individual *i*, $\mathbf{Y}_2[i] := (\mathbf{Y}_1[i])^2$ and $\mathbf{Y}_1 := \mathbf{G}_1 \sim N(0, ZZ^t)$, where, for the sake of argument, we assume absence of residual errors. Then, the factor genotype will generally be significant in the ANCOVA with Y_1 as covariate. For example, there could be two replicates of three genotypes, with genetic effects (-1, -1, 0, 0, 1, 1). Then, clearly, there is some unexplained genetic variance when regressing $\mathbf{Y}_2 = (1, 1, 0, 0, 1, 1)^t$ on \mathbf{Y}_1 .

Finally, we briefly discuss how the approximation could be improved. In general, the conditional mean is a function of the genetic and residual covariances between \mathbf{Y}_j and \mathbf{Y}_S . In Appendix A.7 (Equation 23) we derive that $\mathbf{Y}_j | vec(\mathbf{Y}_S) = vec(\tilde{y}_S)$ has mean $\mu_{j|S} = \mathbf{X}B\gamma_j + \sum_{j,S}\sum_S^{-1}vec(\tilde{y}_S - \mathbf{X}B\Gamma_S)$. Defining $\eta_{j|S} = 0$ for $S = \emptyset$, we can write $\mu_{j|S} = \mathbf{X}B\gamma_j + \eta_{j|S}$. Consequently, our approximation of the conditional mean models $\eta_{j|S}$ as a linear regression on \tilde{y}_S . This approximation could probably be improved if we have good estimates of $\hat{\Sigma}_{j,S}$ and $\hat{\Sigma}_S^{-1}$, and set $\hat{\eta}_{j|S} = \hat{\Sigma}_{j,S} \hat{\Sigma}_S^{-1} vec(\tilde{y}_S - \mathbf{X}\hat{B}\Gamma_S)$. Such estimates, however, require fitting an MTM for |S| + 1 traits, which for large *S* is statistically and computationally challenging, unless pairwise or other approximations are applied (Furlotte and Eskin 2015; Joo *et al.* 2016). Moreover, it seems unclear how $\hat{\eta}_{i|s}$ should be incorporated in the tests.

Discussion

Causal inference for data with random genetic effects is challenging because of the covariance between these effects, and because the usual assumption of independent observations is violated. To address these problems we have proposed a model where random genetic effects are part of the causal graph, rather than a nuisance factor that first needs to be eliminated. The resulting distributions and graphs were shown to satisfy the Markov property. This led us to develop the PCgen algorithm, which tests conditional independence between traits in the presence of genetic effects, and also conditional independence between traits and genetic effects. We showed that the presence of a direct genetic effect can be tested, just like the direct (fixed) effect of a QTL can be tested. This is, of course, relative to the observed traits, *i.e.*, for any effect $G \rightarrow Y_j$.

In the linear simulations as well as in the rice data, our tests could indeed identify the absence of many direct genetic effects. By contrast, in the APSIM simulations and maize data all traits had such effects. In the latter case, this could be for biological reasons, *i.e.*, the genetic variance of each trait might really be "unique" to some degree. However, the APSIM results showed that nonlinearities could increase the false positive rate in the edges $G \rightarrow Y_j$, which may be avoided in future versions with better conditional independence tests. Such tests might also allow for non-Gaussian data.

In our simulations, PCgen also improved the reconstruction of between-trait relations. Part of this improvement is due to phenotypic information on replicates, reducing the number of errors in the tests. Another part is due to the improved orientation, which is a consequence of the additional edges $G \rightarrow Y_j$. Compared to previous algorithms, PCgen also appeared to be computationally more efficient: depending on the sparseness of the network, it can analyze ~10–50 traits on a single core, and many more if we limit the maximum size of the conditioning sets, or would parallelize the conditional independence tests.

As for the original PC-algorithm, PCgen is most efficient for sparse graphs, *i.e.*, when each trait is connected to only a few other traits, and when there are few direct genetic effects. But, even when this is not the case, PCgen still has an advantage over existing approaches: by incorporating the genetic effects in the PC-algorithm, we do not need to fit an MTM for all traits simultaneously, but only for bivariate models. Our approach also makes genetic network reconstruction feasible with just two traits, and in the absence of QTL or even no genotypic data at all.

As any causal inference method, PCgen only suggests causal models that are, in some sense, compatible with the data, and cannot validate the existence of a functional relationship, which is possible only through additional experiments. Because of the required assumptions, the identifiability issues and the uncertainty in the estimated networks, it may be better to speak of an algorithm for causal "exploration" than for causal "discovery". At the same time, analysis of one trait conditional on other traits (*e.g.*, yield given plant height) is a common and natural thing to do (Stephens 2013). From that perspective, PCgen could be seen simply as a tool that performs such analyses systematically, combines them, and visualizes the results. PCgen results for different significance levels could then be reported alongside other "descriptive" statistics like heritability estimates and genetic correlations, suggesting functional hypotheses that are of interest in future research.

Dealing with derived traits

We analyzed the maize and rice datasets using the traits as they were measured, without adding "derived" traits defined by ratios, sums, or differences of the original traits. Because such derived traits are not measured themselves, there is no error associated with them, apart from "copies" of errors in the original trait. For example, if there is much variation in leaf weight but almost none in the total weight of a plant, the derived trait "leaf weight ratio" will be essentially a copy of the original leaf weight trait. This can violate our assumption of faithfulness, and lead to errors in the reconstruction; see Figure 7 in Appendix A for an example. Sometimes, derived traits are biologically highly relevant. It may then be desirable to include them in the analysis, and omit some of the original traits. Alternatively, derived traits may be added after running PCgen. For example, we may extend the reconstructed graph with the node $Y_3 := Y_1 + Y_2$ and edges $Y_1 \rightarrow Y_3$ and $Y_2 \rightarrow Y_3$, provided that this makes sense biologically.

Data from different experiments

We assumed traits to be measured on the same individuals in the same experiment, with residuals errors arising from biological variation (Assumption 1). In certain applications, this assumption can indeed be restrictive, but seems to be inevitable. Suppose traits were measured in different experiments, or residual errors would only come from measurement errors. Then there would be no propagation of residual errors, and the reconstruction would rely completely on how the genetic effects are propagated through the network. The GSEM model (Equation 4) would need to be replaced by $\mathbf{Y} = \mathbf{X}B + \mathbf{U} + \mathbf{E}$ and $\mathbf{U} = \mathbf{U}\Lambda + \tilde{\mathbf{U}}$, where $\tilde{\mathbf{U}}$ in a sense models direct genetic effects, reminiscent of the genomic networks of Töpner et al. (2017). However, if data are actually generated by Equation (4), these networks provide only partial information about the direct genetic effects, even without any type-I or type-II error in the tests (see File S7.2). Moreover, the use of the PC-algorithm would require the columns of \tilde{U} to be independent, which appears to be a rather unrealistic assumption.

Biologically, the genomic networks have a different interpretation: for example, we would assume that the genetic component in high blood pressure causes some cardiovascular disease, rather than high blood pressure itself. The alternative model (**Y** = **X***B* + **U** + **E**) implies that the observed traits have diagonal residual covariance, instead of the matrix $\Gamma^t \Sigma_E \Gamma$ obtained under Assumption 1 (see Equation 5). However, the latter matrix turned out to be essential for network reconstruction (see *e.g.*, Theorems 5–6 above). This is why, without Assumption 1, we would need to rely completely on the genetic effects.

A relevant alternative approach here is that of invariance causal prediction (Peters *et al.* 2016), which infers causal effects that are consistent across several experiments, but still requires all traits to be measured in each experiment (as well as low genotype-by-environment interaction).

Replicates vs. means

In principle, PCgen allows for any type of genetic relatedness. We have however focused on the case of independent genetic effects, for the following reasons:

- 1. Performance under model misspecification: different types of genetic effects could, in theory, be represented by introducing multiple genetic nodes, with conditional independence tests that can distinguish between these effects. But this seems difficult in practice due to the computational requirements and lack of statistical power (Blair et al. 2012; Uhler et al. 2013; Kruijer 2016). For this reason, it seems, previous work on network reconstruction used genotypic means and an additive GRM. For the analysis of a single trait, however, Kruijer (2016) showed that broad-sense heritability estimates (obtained with $K = ZZ^t$) capture any type of genetic effect, while a model assuming only additive effects can produce strongly biased heritability estimates, if the actual genetic effects are, for example, partly epistatic. It seems plausible that this robustness extends to the multivariate models considered here, for example, when direct genetic effects are driven by different sets of QTL, leading to trait-specific relatedness matrices.
- 2. Higher power: estimates of (total) genetic variance based on replicates are typically more accurate than markerbased estimates based on genotypic means (Kruijer *et al.* 2015; Visscher and Goddard 2015), and the use of replicates is therefore also likely to improve hypothesis testing. For the reconstruction of trait-to-trait relations with PCres, our simulations indeed suggest that replicates give more power. Mixed models with both replicates and a GRM might further increase power if the true architecture is really additive (Kruijer *et al.* 2015), but also these models lead to biased inference if the actual architecture is different (Kruijer 2016).
- 3. When $K = ZZ^t$, the conditional independence statement considered in the RC-test is completely equivalent with $Y_j \perp Y_k | \{Y_s, G\}$ (Theorem 5), while for other *K* it is not, and an alternative test might be required.

Apart from these statistical issues, there is also a computational advantage: the test for $Y_j \perp G | Y_s$ can be based on standard ANCOVA, which is many times faster than the

LRT for a mixed model. Also the tests for $\mathbf{Y}_{j} \perp \mathbf{Y}_{k} | \{\mathbf{Y}_{S}, \mathbf{G}\}$ are faster when $K = ZZ^{t}$.

Finally, we have not investigated the performance of PCgen for unbalanced designs, but it seems likely that small unbalancedness has only a minor effect. A more fundamental challenge seems to be the presence of incomplete blocks or spatial trends (Flaxman *et al.* 2015; Rodríguez-Álvarez *et al.* 2018).

Assessing uncertainty

If one mistakenly rejects the null-hypothesis of conditional independence (type-I error), PCgen leaves the corresponding edge, although it may still be removed at a later stage, with a different conditioning set. If the null-hypothesis is mistakenly not rejected (type-II error), a true edge is removed, and will not be recovered. Moreover, it may affect the remaining tests, since d-separation of Y_j and Y_k is only tested given conditioning sets contained in $adj(Y_j)$ or $adj(Y_k)$, where the adjacency sets are defined relative to the current skeleton. This is correct in the oracle version, but in the sample version of PC(gen), $adj(Y_j)$, or $adj(Y_k)$ may become smaller than the corresponding adjacency sets in the true graph, and the algorithm may therefore not perform an essential independence test. See Colombo and Maathuis (2014) for examples.

Consequently, assessing uncertainty for constraint-based algorithms is difficult, and cannot be achieved by just applying some multiple testing correction to the *P*-values. To obtain bounds on the expected number of false edges in the skeleton, several authors have used stability selection (Meinshausen and Bühlmann 2010; Stekhoven *et al.* 2012) or other sample-splitting techniques (Töpner *et al.* 2017), but these are often too stringent and require an additional exchangeability assumption (Bühlmann *et al.* 2013; Meinshausen *et al.* 2016). Moreover, these approaches do not provide a level of confidence for specific edges. For example, an edge present in 60% of all subsamples may appear to be present in the true graph with a probability of 0.6, but there is no justification for such a statement.

Alternatively, uncertainty may be assessed using Bayesian methods, which are, however, computationally very demanding and outside the scope of this work. Moreover, despite the recent progress in Bayesian asymptotics (Ghosal and van der Vaart 2017), there seem to be no results regarding the correct coverage of posteriors in these models.

Genomic prediction

PCgen can select traits with direct genetic effects, which are the most relevant in genomic selection. More generally, the usefulness of structural models for genomic selection depends on whether there is an interest in some kind of intervention (Valente *et al.* 2013, 2015). Informally speaking, an intervention is an external manipulation that forces some of the traits to have a particular distribution. For example, with a so-called hard intervention on the *j*th trait, Y_j is forced to a constant level *c*, *e.g.*, c = 0, when Y_j is the expression of a gene that is knocked out. The manipulation or truncated factorization theorem (Spirtes *et al.* 2001; Pearl 2009) can then predict the joint distribution of the system after the intervention:

$$p_{\mathbf{Y}_{j}:=c}(\mathbf{G},\mathbf{Y}_{-j}) = p(\mathbf{G}) \prod_{j' \neq j} p(\mathbf{Y}_{j'}|pa(\mathbf{Y}_{j'}), \mathbf{G}_{j'}),$$
(15)

where $pa(\mathbf{Y}_{j'})$ is the set of parents of $Y_{j'}$. This is generally different from the distribution

$$p(\mathbf{G}, \mathbf{Y}_{-\mathbf{j}} | \mathbf{Y}_{\mathbf{j}} = c), \tag{16}$$

obtained from conditioning on $Y_j = c$, prior to the intervention [see *e.g.*, Peters *et al.* (2017)]. In other words, conditioning is not the same as doing (intervening). In a simulated example (File S7), we show that the use of Equation 15 can indeed greatly improve accuracy after an intervention, compared to standard methods ignoring the underlying structure. When, however, the intervention is on a root node, Equations 15 and 16 are the same (see again Peters *et al.* 2017, p. 110).

The example in File S7.4 is special in the sense that PCgen could correctly infer the complete graph, for most of the simulated datasets. A technical obstacle for a more general use of our networks in genomic prediction is the identifiability issue mentioned in the introduction. PCgen (and the PC-algorithm in general) typically outputs a partially directed graph, several DAGs being compatible with this graph. This is particularly problematic for edges between the traits in *D* (traits with direct genetic effects). For traits with only indirect genetic effects, it is possible to estimate how much of the genetic variance originates from a particular trait in *D*, the result being independent of the chosen DAG. This would first require estimates of the (total) genetic covariance among traits in *D*, obtained either by fitting an MTM, or by an approximation (as in Furlotte and Eskin 2015).

In absence of interventions on the traits, we can think of genomic prediction in terms of an intervention on the node G. Because the latter is a root node by definition, standard genomic prediction methods can, in principle, have optimal performance (Valente *et al.* 2013). More specifically, genomic prediction usually involves a regression of a target trait on a number of markers, having either fixed or random effects. In either case, it is only the total effect of each underlying locus on the target trait that matters, not through which other traits this effect passes.

Optimal prediction accuracy, however, requires that the regression model contains the true distribution (or a good approximation), and a sufficiently accurate estimate of this distribution. We therefore believe that structural models may sometimes be an appealing alternative, especially if the underlying model is highly nonlinear, or when prior physiological knowledge can be incorporated. The extent to which this can really improve accuracy remains to be investigated.

Open questions and extensions

Although we have shown the Markov property for our model, and studied consistency of PCgen, there are a number of open questions left for future work. First, it may be possible to construct better tests, especially for nonlinear structural models and non-Gaussian error distributions. The recent work of Pfister et al. (2018) seems particularly relevant here. A second issue is the consistency of the orientations: while we have shown PCgen's consistency in reconstructing the skeleton, we did not address this for the final CPDAG. This is well known for the PC-algorithm without genetic effects (Spirtes et al. 2001; Kalisch and Bühlmann 2007), but more difficult to establish here, as the class of CPDAGs needs to be restricted to those without errors pointing to G. More generally, orientation constraints seem to be of interest for trait-to-trait relationships as well, e.g., one may require that, if there is an edge, the expression of a gene can only affect a metabolite and not the other way round. To the best of our knowledge, current methodology and theory has considered only the forced absence/presence of an edge, leaving the orientation to the algorithm [The pcalg-package (Kalisch et al. 2012) has the addBgKnowledge option to add orientations ('background knowledge'), in the estimated CPDAG. This is however only done after running PC or a related algorithm, and is only allowed if compatible with the CPDAG]. A final question for future work is whether Theorems 4 and 6 hold for general conditioning sets.

Apart from these open questions, we believe that the idea of explicitly modeling direct genetic effects can be applied more generally. In particular, we hope that the ideas developed here provide a first step toward the more ambitious goal of modeling multiple traits through time, simultaneously for many environments. A first generalization would be to replace the PC-algorithm with other constraint-based algorithms, in particular FCI and RFCI (Spirtes et al. 2001; Colombo et al. 2012). These have the advantage that the causal sufficiency assumption (no latent variables) can be dropped or considerably weakened. The presence or absence of direct genetic effects could also be incorporated in (empirical) Bayesian approaches for genetic network reconstruction, or in invariant causal prediction (Peters et al. 2016). It might also be possible to extend the approach of Stephens (2013), and focus only on the detection of traits with direct genetic effects. Another application of GSEM might be as covariance models in multi-trait GWAS, as an alternative to unstructured (Zhou and Stephens 2014) or low-rank (Millet et al. 2016) models. Finally, the concept of direct and indirect genetic effects may be useful in deeplearning models for high-dimensional phenotypes, observed on genetically diverse individuals.

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APPENDIX A. Faithfulness, conditional distributions, and proofs of Theorems 1-6

A.1. Overview of graph theoretic definitions

Definitions of, for example, d-separation and CPDAGs can be found in many books and articles on graphical models and causal inference (see for example, Lauritzen 1996; Spirtes *et al.* 2001; Kalisch and Bühlmann 2007; Pearl 2009). The following summary was inspired by Shipley (2016) and Maathuis (2014).

- Given different nodes Y_j and Y_k , a path from Y_j to Y_k is a sequence of edges connecting Y_j and Y_k . When all edges are directed and pointing toward the same node, we have a directed path. A path that is not directed is an undirected or nondirected path.
- A cycle is a path from Y_j to Y_k with an additional edge between Y_j and Y_k . A *directed cycle* is a directed path from Y_j to Y_k together with a directed edge $Y_k \rightarrow Y_j$.
- A *directed acyclic graph* (DAG) is a directed graph without any directed cycle. When a graph underlying a SEM is a DAG, the SEM is said to be *recursive*.
- $pa(Y_j)$ is the set of nodes Y_k for which there is a directed edge $Y_k \rightarrow Y_j$; in this case, Y_j is a *child* of Y_k , and Y_k is a *parent* of Y_j . The nodes Y_j and Y_k are *adjacent* if there is an edge $Y_k \rightarrow Y_j$, $Y_j \rightarrow Y_k$, or $Y_k - Y_j$.
- If in a DAG \mathcal{G} there is a directed path from Y_i to Y_k , then Y_k is a descendant of Y_i , and Y_i is an ancestor of Y_k .
- In a DAG with nodes Y_1, \ldots, Y_p , it is always possible to relabel the nodes, such that for each node $Y_j, k < j$ for all parents Y_k in the set $pa(Y_j)$. Such a relabeling is known as a *topological ordering* of the DAG. Using this ordering, the root nodes (without any parents) have the lowest labels, the sink nodes (without any children) have the highest labels, and the matrix of path coefficients Λ is upper triangular. Formally, a topological ordering is a function $\pi : \{1, \ldots, p\} \rightarrow \{1, \ldots, p\}$, such that the preceding property holds. A topological ordering always exists, and does not need to be unique (Peters *et al.* 2017).
- If, for a given path, two directed edges point into the same node, the latter is a *collider*. For example, given the DAG $A \rightarrow C \leftarrow B$, *C* is a collider on the (only) path between *A* and *B*. In all other cases ($A \leftarrow C \rightarrow B$, $A \rightarrow C \rightarrow B$ and $A \leftarrow C \leftarrow B$), *C* is a *noncollider*. Several different paths can pass through a node, and being a (non)collider is always relative to the path.
- In a DAG, a *v*-structure or immorality is a collection of three nodes (say *A*, *B*, and *C*), such that there are directed edges $A \rightarrow B$ and $C \rightarrow B$, but no edge between *A* and *C*. In this case *B* is an *unshielded* collider. If there is an edge between *A* and *C*, it is a *shielded* collider. Similarly, in an undirected graph, *A*, *B*, and *C* form an *unshielded* triple if there are edges A B and C B, but no edge A C.

The skeleton of a (partially) directed graph is the undirected graph obtained after removing all arrowheads.

Given a directed graph *G*, two nodes *A* and *B*, and a (possibly empty) subset of nodes *S* not containing *A* and *B*, a path between *A* and *B* is *blocked* by *S* if at least one of the following two conditions holds: (i) there exists a collider on the path which is not in *S*, and also none of its descendants are in *S*; (ii) there exists a noncollider on the path that is in *S*.

Nodes A and B are *d*-separated by a set S if S blocks all paths from A to B.

Given disjoint sets *U*, *V*, and *S* (*U* and *V* should be nonempty), *U* and *V* are *d*-separated by *S* if *S* blocks all paths from Y_j to Y_k , for all nodes $Y_j \in U$ and $Y_k \in V$.

Two DAGs are equivalent if they have the same skeleton and the same v-structures.

An equivalence class of DAGs is a set containing all DAGs that are equivalent to one another. For example, given a skeleton A - B - C, there is one equivalence class containing the three DAGs $A \rightarrow B \rightarrow C$, $C \rightarrow B \rightarrow A$, and $A \leftarrow B \rightarrow C$, and one equivalence class with only one DAG $(A \rightarrow B \leftarrow C)$. Any DAG in the class can be used to represent the class. But instead of picking an arbitrary DAG, an equivalence class can also be represented by a *completed partially directed acyclic graph* (CPDAG). A *partially directed acyclic graph* (PDAG) is "a graph where some edges are directed and some are undirected and one cannot trace a cycle by following the direction of directed edges and any direction for undirected edges" (Kalisch and Bühlmann 2007). A PDAG is a CPDAG if (a) every directed edge in the PDAG exists in all DAGs in the equivalence class it represents (b) for every undirected edge A - B in the PDAG, the equivalence class contains at least one DAG with $A \rightarrow B$, and at least one with $B \rightarrow A$. Chickering (2002) showed that CPDAGs represent equivalence classes uniquely.

A.2. Identifiability

In a general GSEM, the parameters in Σ_G , Σ_E , and Λ are not identifiable, which was pointed out by Gianola and Sorensen (2004). However, when we know a topological ordering of the graph (defined above in Appendix A.1), and Assumption 3 (diagonal Σ_E) and faithfulness assumptions (9) and (10) hold, it appears that Σ_G , Σ_E , and Λ are identifiable from the joint distribution of $\mathbf{Y}_1, \ldots, \mathbf{Y}_p$ (see Equation 7). Our approach relies on the L^tDL decomposition of V_E , and is probably known, although we could not find it in the literature. Neither could we find a proof, but the decomposition seemed valid in all examples we considered. It works as follows:

- 1. Relabel the nodes (traits) Y_1, \ldots, Y_p according to a topological ordering. Then, both Λ and $\Gamma = (I \Lambda)^{-1}$ are upper triangular. Recall that Λ has zeros on the diagonal. Γ always has ones on the diagonal, *i.e.*, it is *unit* upper-triangular.
- 2. Now, we use the fact that every positive definite matrix *A* can be decomposed as $A = LDL^t = L^tDL$, with a diagonal matrix *D* and unit upper-triangular *L* [see *e.g.*, Petersen *et al.* (2008), section 5.7]. We apply this result to V_E :

$$V_E = L^t D L, \tag{17}$$

and set Σ_E equal to *D*, and Γ equal to *L*. Let $\Lambda = I - \Gamma^{-1}$.

3. Finally, using V_G and L we obtain Σ_G :

$$\Sigma_G = (LL^t)^{-1} (LV_G L^t) (LL^t)^{-1}.$$
(18)

For example, consider the graph $Y_3 \rightarrow Y_2 \rightarrow Y_1$, with path coefficients equal to one and unit error variances (for simplicity, we ignore the genetic effects in this example). We need to relabel the graph such that $Y_1 \rightarrow Y_2 \rightarrow Y_3$. After relabeling, we have

$$\Lambda = \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 0 & 0 & 0 \end{pmatrix}, \quad \Gamma = (I - \Lambda)^{-1} = \begin{pmatrix} 1 & 1 & 1 \\ 0 & 1 & 1 \\ 0 & 0 & 1 \end{pmatrix}, \quad V_E = \Gamma^t \Sigma_E \Gamma = \begin{pmatrix} 1 & 1 & 1 \\ 1 & 2 & 2 \\ 1 & 2 & 3 \end{pmatrix},$$

as Σ_E is the identity matrix. In R, the L^tDL decomposition can be computed as follows:

b <- matrix(c(1,1,1,1,2,2,1,2,3), ncol = 3) #b <- b[c(3,2,1), c(3,2,1)] v <- Cholesky(Matrix(b, sparse = T), LDL = T, perm = F)@x Gamma <- matrix(0,3,3) Gamma[lower.tri(Gamma, diag = T)] <- v D <- diag(diag(Gamma)) diag(Gamma) <- 1 Gamma <- t(Gamma) Lambda <- diag(3) - (solve(Gamma))</pre>

We emphasize that the topological ordering is crucial. Without the relabeling (*e.g.*, uncomment the second line in the above R-code), a different Γ is obtained (also after interchanging the first and third row). Although in this example the topological ordering is unique, there may in general be multiple valid orderings; *e.g.*, Y_1, Y_3, Y_2 and Y_3, Y_1, Y_2 in case the graph is $Y_1 \rightarrow Y_2 \leftarrow Y_3$. Based on all investigated examples, we conjecture (but could not prove) that these orderings lead to the same parameter estimates.

A.3. The matrix Γ expressed as a function of path coefficients

Let $\mathcal{G}_{\mathcal{Y}}$ denote the DAG over the nodes Y_1, \ldots, Y_p , with edges defined by Λ . For each $j \in \{1, \ldots, p\}$, let V_j denote the union of the set $\{Y_j\}$ and the set of root traits (*i.e.*, those without parents in $\mathcal{G}_{\mathcal{Y}}$) for which there is a directed path toward Y_j . For all $j, k \in \{1, \ldots, p\}$, let Π_{jk} denote the set of all directed paths from Y_j to Y_k . For k = j, Π_{jj} contains only the empty path from Y_j to itself. For any directed path π from Y_j to Y_k , let $L(\pi)$ denote the product of the corresponding path coefficients as given by Λ ; for the empty path we define $L(\pi) = 1$.

Using these definitions, we can decompose the variance of a trait into contributions from different ancestors, as well as its own error variance. To this end, we follow Spirtes *et al.* (2001) and define the $p \times 1$ column vector γ_j with elements (l = 1, ..., p)

$$\gamma_{j,l} = \begin{cases} \sum_{\pi \in \Pi_{lj}} L(\pi) & \text{if } Y_l \in V_j \\ 0 & \text{otherwise.} \end{cases}$$
(19)

A.4. The covariance between Y_i and Y_k as function of path coefficients

Since $\mathbf{Y}_{j} = \mathbf{X}B\gamma_{j} + \mathbf{G}\gamma_{j} + \mathbf{E}\gamma_{j}$ (Equation 8 in the main text), the covariance between the $n \times 1$ vectors \mathbf{Y}_{j} and \mathbf{Y}_{k} can be written in terms of γ_{j} and γ_{k} :

$$Cov(\mathbf{Y}_{\mathbf{j}}, \mathbf{Y}_{\mathbf{k}}^{t}) = E[(\mathbf{Y}_{\mathbf{j}} - \mathbf{X}B\gamma_{j})(\mathbf{Y}_{\mathbf{k}} - \mathbf{X}B\gamma_{k})^{t}] = (\gamma_{j}^{t}\Sigma_{G}\gamma_{k})K + (\gamma_{j}^{t}\Sigma_{E}\gamma_{k})I_{n},$$
(20)

for all $j, k \in \{1, ..., p\}$. Consequently, we can express the genetic and residual covariance between traits in terms of quadratic forms, involving Σ_G , Σ_E , and the path coefficients. As a special case of Equation 20, it follows that, without random genetic effects,

$$Cov(\mathbf{Y}[i,j],\mathbf{Y}[i,k]) = oldsymbol{\gamma}_i^t \Sigma_E oldsymbol{\gamma}_k$$

is the covariance between the *j*th and *k*th trait, for each individual *i*. See also Spirtes *et al.* (2001) (Lemma 3.1.6), or Lynch and Walsh (1998) (Appendix 2). Using standard expressions for multivariate Gaussian distributions, this implies that when $Y_j = XB\gamma_i + E\gamma_j$,

$$Cov(\mathbf{Y}[i,j],\mathbf{Y}[i,k]|\mathbf{Y}[i,S]) = \gamma_j^t \Sigma_E \gamma_k - \left(\gamma_j^t \Sigma_E \Gamma_S\right) \left(\Gamma_S^t \Sigma_E \Gamma_S\right)^{-1} \left(\Gamma_S^t \Sigma_E \gamma_k\right).$$
(21)

A.5. The path coefficients condition

It is well known that faithfulness is violated when contributions from different paths cancel out. For example, in the SEM defined by $Y_1 \rightarrow Y_2$, $Y_1 \rightarrow Y_3$, and $Y_2 \rightarrow Y_3$, with respective path coefficients 1, 1, and -1, Y_1 and Y_3 are marginally independent but not d-separated. Conversely, when faithfulness holds, we know that such cancellations cannot occur, and that the sum in Equation 19 is never zero, *i.e.*, $\gamma_{i,l} = 0$ only for $Y_l \notin V_j$. We will refer to this as the path coefficients condition.

A.6. The path coefficients condition and faithfulness

The path coefficients condition is a necessary but not a sufficient condition for faithfulness. First, faithfulness can also be violated when contributions from different paths cancel out when summing over a subset of (rather than all) the directed paths; see Example 2.10 in Peters (2012). Second, it is not only the contributions of directed paths that should not cancel out, but also those of treks. A trek between Y_j and Y_k is any path between these nodes without a collider (Spirtes *et al.* 2001). Every trek consists of two directed paths, starting at the source of the trek, and going toward Y_k and Y_k . One of these can be the empty path; hence, each directed path is also a trek. Figure 6 provides an example where contributions from different treks cancel out, leading to nonfaithfulness.

Another necessary condition for faithfulness is that all error variances are strictly positive. Figure 7 provides an example of nonfaithfulness due to a zero error variance. An extended version of the path coefficients condition (involving sums over subset of treks) together with strictly positive error variances may be sufficient for faithfulness, but we could not find such a result in the literature. However, from Equation 21 it follows that, for a Gaussian linear SEM (again without genetic effects), faithfulness is equivalent with

$$\gamma_j^t \Sigma_E \gamma_k - \left(\gamma_j^t \Sigma_E \Gamma_S\right) \left(\Gamma_S^t \Sigma_E \Gamma_S\right)^{-1} \left(\Gamma_S^t \Sigma_E \gamma_k\right) = 0 \Rightarrow Y_j \text{ and } Y_k \text{ d-separated by } Y_S.$$
(22)

A.7. Conditional means and covariances

Using the notation [,S] to select the columns corresponding to S, and $[S_1, S_2]$ to select both rows and columns, it follows from Equation 7 that $\mathbf{Y}_i | vec(\mathbf{Y}_S) = vec(\tilde{y}_S)$ is multivariate normal with mean and covariance

$$\mu_{j|S} = (\mathbf{X}B\Gamma)[,j] + \Sigma_{j,S}\Sigma_{S}^{-1}vec(\tilde{y}_{S} - (\mathbf{X}B\Gamma)[,S]) = \mathbf{X}B\gamma_{j} + \Sigma_{j,S}\Sigma_{S}^{-1}vec(\tilde{y}_{S} - \mathbf{X}B\Gamma_{S}),$$
(23)

$$\Sigma_{j|S} = \Sigma_j - \Sigma_{j,S} \Sigma_S^{-1} \Sigma_{j,S}^t, \tag{24}$$

where

$$\Sigma_{j,S} = \left(\Gamma^{t}\Sigma_{G}\Gamma\right)[j,S] \otimes K + \left(\Gamma^{t}\Sigma_{E}\Gamma\right)[j,S] \otimes I_{n} = \left(\gamma_{j}^{t}\Sigma_{G}\Gamma_{S}\right) \otimes K + \left(\gamma_{j}^{t}\Sigma_{E}\Gamma_{S}\right) \otimes I_{n},\tag{25}$$

$$\Sigma_S = \left(\Gamma_S^t \Sigma_G \Gamma_S\right) \otimes K + \left(\Gamma_S^t \Sigma_E \Gamma_S\right) \otimes I_n,\tag{26}$$

$$\Sigma_{j} = \left(\Gamma^{t}\Sigma_{G}\Gamma\right)[j,j]K + \left(\Gamma^{t}\Sigma_{E}\Gamma\right)[j,j]I_{n} = \left(\gamma_{j}^{t}\Sigma_{G}\gamma_{j}\right)K + \left(\gamma_{j}^{t}\Sigma_{E}\gamma_{j}\right)I_{n}.$$
(27)

The matrices Σ_j , Σ_s , and $\Sigma_{j,s}$ are the variance-covariance matrix of $vec(\mathbf{Y}_j) = \mathbf{Y}_j$ and $vec(\mathbf{Y}_s)$, respectively, and the covariance between \mathbf{Y}_j and $vec(\mathbf{Y}_s)$. From Equation 7 in the main text we also obtain the conditional distribution

$$\operatorname{vec}([\mathbf{Y}_{\mathbf{j}} \ \mathbf{Y}_{\mathbf{k}}])|\operatorname{vec}(\mathbf{Y}_{\mathbf{S}}) = \operatorname{vec}(\tilde{y}_{S}) \sim N\left(\begin{pmatrix} \mu_{j|S} \\ \mu_{k|S} \end{pmatrix}, \Sigma_{jk|S} \right) = N\left(\begin{pmatrix} \mu_{j|S} \\ \mu_{k|S} \end{pmatrix}, \Sigma_{jk} - \Sigma_{jk,S}\Sigma_{S}^{-1}\Sigma_{jk,S}^{t}\right),$$
(28)



Figure 6 An example of a SEM (without genetic effects) where faithfulness does not hold, because the contributions to the covariance from the treks $Y_3 \leftarrow Y_1 \rightarrow Y_4$ and $Y_3 \leftarrow Y_2 \rightarrow Y_4$ cancel out. If Y_1 and Y_2 are Gaussian with equal error variances, it follows that for every individual *i*, $Cov(\mathbf{Y}_3[i], \mathbf{Y}_4[i]) = Cov(\mathbf{Y}_1[i] + \mathbf{F}_2[i] + \mathbf{F}_3[i], \mathbf{Y}_1[i] - \mathbf{Y}_2[i] + \mathbf{E}_4[i]) = Cov(\mathbf{Y}_1[i] - \mathbf{Y}_2[i]) = 0$. Consequently, \mathbf{Y}_3 and \mathbf{Y}_4 are marginally independent, but not d-separated by the empty set.

where $\mu_{j|S}$ and $\mu_{k|S}$ are as in Equation 23, and Σ_{jk} is the $2n \times 2n$ block matrix with diagonal blocks Σ_j and Σ_k (defined as in Equation 27), and off-diagonal blocks $(\gamma_j^t \Sigma_G \gamma_k)K + (\gamma_j^t \Sigma_E \gamma_k)I_n$. Similarly, given the $p \times 2$ matrix Γ_{jk} with columns γ_j and γ_k , it follows that

$$\Sigma_{jk,S} = \left(\Gamma_{jk}^t \Sigma_G \Gamma_S\right) \otimes K + \left(\Gamma_{jk}^t \Sigma_E \Gamma_S\right) \otimes I_n$$

is the $2n \times |S|n$ covariance between $vec([\mathbf{Y}_{i} \mathbf{Y}_{k}])$ and $vec(\mathbf{Y}_{S})$.

A.8. Covariance structure of the conditional distributions

When $K = ZZ^t$ is block-diagonal, with *m* blocks of ones of dimension $r \times r$ on the diagonal, then for any positive constants *c* and *d*,

$$(cI_n + dK)^{-1} = c^{-1}I_n - \frac{1}{c^2(1/d + r/c)}K$$

Hence, the inverse of $(cI_n + dK)$ is again a linear combination of I_n and K. This follows from the Woodbury identity (Petersen *et al.* 2008; Golub and Van Loan 2012)

$$(A + CBC^{t})^{-1} = A^{-1} - A^{-1}C(B^{-1} + C^{t}A^{-1}C)^{-1}C^{t}A^{-1},$$
(29)

with $A = cI_n$, $B = dI_m$, and C = Z. In addition we have $Z^tZ = rI_m$, and therefore $K^2 = rK$. Consequently, any product of matrices of the form $(cI_n + dK)$ or their inverse is a linear combination of I_n and K. Combining Equations 25, 26 and 27, and using that $(A \otimes B)(C \otimes D) = AC \otimes BD$ for any matrices A, B, C, D of appropriate dimensions, it follows that when $K = ZZ^t$, $\Sigma_{j|S}$ in Equation 24 is of the form $\sigma_G^2(j|S)K + \sigma_E^2(j|S)I_n$, for some numbers $\sigma_G^2(j|S)$ and $\sigma_E^2(j|S)$. Similarly, it follows that $\Sigma_{jk|S}$ (Equation 28) is of the form

$$V_G(jk|S) \otimes K + V_E(jk|S) \otimes I_n$$

for some 2 × 2 matrices $V_G(jk|S)$ and $V_E(jk|S)$.

A.9. Proofs of Theorems 1 and 2

Pearl (2009) (p. 51) showed that, under quite general assumptions, structural equation models satisfy the global Markov property, which means that d-separation in the graph implies conditional independence. It turns out that, in our case, the required assumption of independent errors applies to the *p* error variables and not to *G*. The intuition behind this is that *G* is not just an additional error node, but part of the causal graph, and we can always distinguish between residual (co)variance and genetic (co)variance. We now give the proof of Theorem 1, which only requires minor modifications of the proof given by Pearl for the case without the genetic effects.

Let \mathcal{G}_E denote the graph, obtained by extending \mathcal{G} with the error variables, *i.e.*, for traits j = 1, ..., p we add the node E_j and an edge $E_j \rightarrow Y_j$. We first show that the local Markov property holds for \mathcal{G}_E , *i.e.*, for any variable $Z \in \{G, Y_1, ..., Y_p, E_1, ..., E_p\}, Z$ is conditionally independent of its nondescendants given its parents. This is obvious for $Z \in \{G, E_1, ..., E_p\}$; we now consider Y_j . In \mathcal{G}_E , the set of parents of Y_j is $pa(Y_j) \cup \{E_j\}$, where $pa(Y_j)$ contains G if $j \in D$. By construction, Y_j is entirely determined by



Figure 7 An example of a SEM where faithfulness does not hold, because the variance of the error variables E_2 is zero. The random vectors \mathbf{Y}_1 and \mathbf{Y}_4 are conditionally independent given \mathbf{Y}_3 , but, in the graph, the nodes Y_1 and Y_4 are not d-separated by Y_3 .

 $pa(Y_j) \cup \{E_j\}$, and constant conditional on these variables. Consequently, given $pa(Y_j) \cup \{E_j\}$, it is independent of any E_k $(k \neq j)$, and of any Y_k that it a nondescendant of Y_j [note that if $G \notin pa(Y_j)$, Y_j is indeed conditionally independent of any nondescendant; if $G \in pa(Y_j)$, G cannot be the nondescendant because it is already in the conditioning set]. Therefore, the local Markov property holds for \mathcal{G}_E . By the lemma below, we find that also the Markov factorization property holds for \mathcal{G}_E , since for any distribution having a density it is equivalent with the local and global Markov properties. Given the Markov factorization property for \mathcal{G}_E , and the fact that $f(e_1, \ldots, e_p) = \prod_{j=1}^p f_j(e_j)$, we can just integrate out the e_j , and obtain the Markov factorization property for \mathcal{G} . This concludes the proof of Theorem 1.

Given the result of Theorem 1 and the assumed faithfulness, Theorem 2 now follows directly from the consistency for the general PC-algorithm (Spirtes *et al.* 2001); see the first part of the proof of their Theorem 5.1 (p. 407).

Markov properties: The following lemma is taken from Lauritzen (1996) (p. 51), and reformulated with somewhat less general conditions, which, however, suffice for our purpose.

Lemma. Let *P* be the joint distribution of random variables (Y_1, \ldots, Y_p) , having a density *f*, and let \mathcal{H} be a DAG on these variables. The following properties are equivalent:

1. The Markov factorization property: given the parents pa_i of each x_i , the joint density (f) can be decomposed as

$$f(\mathbf{y}_1,\ldots,\mathbf{y}_p)=\prod_{j=1}^p f_j(\mathbf{y}_j|p\mathbf{a}_j),$$

where the f_j are the conditional densities.

2. The local Markov property: any variable is conditionally independent of its nondescendants, given its parents.

3. The global Markov property: for all disjoint sets $U, V, S \subset \{Y_1, \ldots, Y_p\}$, d-separation of U and V by S in the graph \mathcal{H} implies conditional independence of U and V given S. In contrast to U and V, the conditioning set S may be empty here. A definition of d-separation is given in Appendix A.1.

A.10. Proof of Theorems 3 and 5: We first prove Theorem 3, by showing the equivalence of the left- and right-hand sides of (9) and (11) in the main text. The d-separation statements on the right-hand sides are equivalent, as *G* can never be a (or descendant of a) collider. Also the left-hand sides $(Y_j \perp Y_k | \{Y_s, G\} \text{ and } Y_j \perp_{P_{vin}} Y_k | \{Y_s\})$ are equivalent, since

$$p_{\mathbf{Y}|\mathbf{U}}(y_i, y_k|y_S) = p(y_i, y_k|y_S, G\Gamma) = p(y_i|y_S, G\Gamma)p(y_k|y_S, G\Gamma) = p_{\mathbf{Y}|\mathbf{U}}(y_i|y_S)p_{\mathbf{Y}|\mathbf{U}}(y_k|y_S).$$

For Theorem 5 we make the additional assumption that $K = ZZ^t$, $Z = I_m \otimes (1, ..., 1)^t$ being the $mr \times m$ design matrix for r replicates of m genotypes in a balanced design (with mr = n). The first part of Theorem 5 then follows from the results in Appendix A.8. For the second part, we first recall the equivalence of $\mathbf{Y}_j \perp \mathbf{Y}_k | \{\mathbf{Y}_S, \mathbf{G}\}$ and $\mathbf{Y}_j \perp_{P_{\mathbf{Y}|U}} \mathbf{Y}_k | \{\mathbf{Y}_S\}$. Because of the Gaussianity and the assumed faithfulness, the latter conditional independence is equivalent with

$$\gamma_j^t \Sigma_E \gamma_k - \left(\gamma_j^t \Sigma_E \Gamma_S\right) \left(\Gamma_S^t \Sigma_E \Gamma_S\right)^{-1} \left(\Gamma_S^t \Sigma_E \gamma_k\right) = 0, \tag{30}$$

where we used Equation 21.

Next, we consider the conditional distribution of $vec([\mathbf{Y}_j \ \mathbf{Y}_k])|vec(\mathbf{Y}_S) = vec(\tilde{y}_S)$ given in Equation 28, whose covariance is the $2n \times 2n$ block matrix $\Sigma_{jk} - \Sigma_{jk,S} \Sigma_S^{-1} \Sigma_{jk,S}^t$. All its four $n \times n$ blocks are a linear combinations of K and I_n , and it suffices to show that the coefficient of I_n in the off-diagonal blocks is zero if and only if Equation 30 holds. We recall from Equation 26 that

$$\Sigma_{S} = \left(\Gamma^{t}\Sigma_{G}\Gamma\right)[S,S] \otimes K + \left(\Gamma^{t}\Sigma_{E}\Gamma\right)[S,S] \otimes I_{n} = \left(\Gamma_{S}^{t}\Sigma_{G}\Gamma_{S}\right) \otimes K + \left(\Gamma_{S}^{t}\Sigma_{E}\Gamma_{S}\right) \otimes I_{n}$$

Using the Woodbury identity (Equation 29) with $A = V_E \otimes I_n$, $B = V_G \otimes I_m$, and $C = I_p \otimes Z$, it follows that, for any positivedefinite $p \times p$ matrices V_G and V_E , we have

$$(V_G \otimes K + V_E \otimes I_n)^{-1} = (V_E^{-1} \otimes I_n) - (V_E^{-1} (V_G^{-1} r V_E^{-1})^{-1} V_E^{-1}) \otimes K.$$
(31)

Setting $V_G = \Gamma_S^t \Sigma_G \Gamma_S$, $V_E = \Gamma_S^t \Sigma_E \Gamma_S$, and $R = V_E^{-1} (V_G^{-1} r V_E^{-1})^{-1} V_E^{-1}$, it follows that

$$\Sigma_{S}^{-1} = \left(\Gamma_{S}^{t}\Sigma_{E}\Gamma_{S}\right)^{-1} \otimes I_{n} - R \otimes K.$$
(32)

Combining this with the expressions for Σ_{jk} and $\Sigma_{jk,S}$ given in Appendix A.7, we find that $\Sigma_{jk} - \Sigma_{jk,S} \Sigma_S^{-1} \Sigma_{jk,S}^t$ has off-diagonal blocks

$$\left(\gamma_{j}^{t}\Sigma_{G}\gamma_{k}\right)\otimes K+\left(\gamma_{j}^{t}\Sigma_{E}\gamma_{k}\right)\otimes I_{n}-\left(\left(\gamma_{j}^{t}\Sigma_{E}\Gamma_{S}\right)\otimes I_{n}+T_{j}\otimes K\right)\left(\left(\Gamma_{S}^{t}\Sigma_{E}\Gamma_{S}\right)^{-1}\otimes I_{n}-R\otimes K\right)\left(\Gamma_{S}^{t}\Sigma_{E}\gamma_{k}\otimes I_{n}+T_{k}^{t}\otimes K\right),$$

for $T_j = \gamma_j \Sigma_G \Gamma_S$ and $T_k = \gamma_k \Sigma_G \Gamma_S$. Finally, working out the products in the last display [using that $K^2 = (ZZ^t)^2 = rK$], we find that all terms involving a Kronecker product with I_n correspond exactly to the left-hand side of Equation 30. Consequently, the residual covariance in the distribution $\{\mathbf{Y}_j, \mathbf{Y}_k\}|\mathbf{Y}_S = \tilde{y}_S$ is zero if and only if $\mathbf{Y}_j \perp \mathbf{Y}_k | \{\mathbf{Y}_S, \mathbf{G}\}$.

A.11. Proof of Theorem 4

To obtain faithfulness for $S = \emptyset$, we need to prove that $\mathbf{Y}_j \perp \mathbf{G}$ implies d-separation of Y_j and G in the graph \mathcal{G} . Because the conditioning set is empty, it suffices to show that there are no directed paths from G to Y_j , where we can assume that $j \notin D$ (otherwise \mathbf{G}_j would be nonzero, and because of the noncollinearity, \mathbf{Y}_j and \mathbf{G} would not be independent). Because of the assumed Gaussianity, the independence of \mathbf{Y}_j and \mathbf{G} implies that

$$Cov(\mathbf{Y}_{\mathbf{j}}^{t}, \mathbf{G}) = Cov(\gamma_{\mathbf{j}}^{t}\mathbf{G}^{t}, \mathbf{G}) = \operatorname{trace}(K)(\gamma_{\mathbf{j}}^{t}\Sigma_{G}) = (0, \dots, 0),$$
(33)

where we used that $vec(\mathbf{G}) \sim N(0, \Sigma_G \otimes K)$, and, therefore, $E(\mathbf{G}[i, j]\mathbf{G}[i, k]) = \Sigma_G[j, k]K[i, i]$, for all $i \in \{1, ..., n\}$ and $j, k \in \{1, ..., p\}$. Since trace(K) is strictly positive and the submatrix $\Sigma_G[D, D]$ has full rank, Equation 33 implies that $\gamma_{j,l} = 0$ for all $l \in D$. Finally, we use that the assumed faithfulness implies the path coefficients condition (see Appendices A.3–A.6). Consequently, it follows from $\gamma_{j,l} = 0$ that there is no directed path from Y_l to Y_j . Since this is the case for all $l \in D$, there can neither be a directed path from G to Y_j .

A.12. Proof of Theorem 6

Assuming $K = ZZ^t$, the first part of the theorem follows from the results in Appendix A.8. For the second part, we use that \mathbf{Y}_j has genetic variance $\sigma_j^2(G) = \gamma_j^t \Sigma_G \gamma_j$ (see Equation 20). Because for traits without a direct genetic effect, rows and columns in Σ_G are zero, we can rewrite this as $\gamma_j^t[D]\Sigma_G[D,D]\gamma_j[D]$. Hence, $\sigma_j^2(G) = 0$ is equivalent with $\gamma_{j,l} = 0$ for all $l \in D$, where we used that $\Sigma_G[D,D]$ is of full rank (which is a consequence of Assumption 5). Using the arguments from the proof of Theorem 4 and the assumed faithfulness, it follows that this is equivalent with independence of \mathbf{Y}_j and \mathbf{G} .