Modeling Causality for Pairs of Phenotypes in System Genetics

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ABSTRACT Current efforts in systems genetics have focused on the development of statistical approaches that aim to disentangle causal relationships among molecular phenotypes in segregating populations. Reverse engineering of transcriptional networks plays a key role in the understanding of gene regulation. However, transcriptional regulation is only one possible mechanism, as methylation, phosphorylation, direct protein–protein interaction, transcription factor binding, etc., can also contribute to gene regulation. These additional modes of regulation can be interpreted as unobserved variables in the transcriptional gene network and can potentially affect its reconstruction accuracy. We develop tests of causal direction for a pair of phenotypes that may be embedded in a more complicated but unobserved network by extending Vuong’s selection tests for misspecified models. Our tests provide a significance level, which is unavailable for the widely used AIC and BIC criteria. We evaluate the performance of our tests against the AIC, BIC, and a recently published causality inference test in simulation studies. We compare the precision of causal calls using biologically validated causal relationships extracted from a database of 247 knockout experiments in yeast. Our model selection tests are more precise, showing greatly reduced false-positive rates compared to the alternative approaches. In practice, this is a useful feature since follow-up studies tend to be time consuming and expensive and, hence, it is important for the experimentalist to have causal predictions with low false-positive rates.

A key objective of biomedical research is to unravel the biochemical mechanisms underlying complex disease traits. Integration of genetic information with genomic, proteomic, and metabolomic data has been used to infer causal relationships among phenotypes (Schadt et al. 2005; Li et al. 2006; Kulp and Jagalur 2006; Chen et al. 2007; Zhu et al. 2004, 2007, 2008; Aten et al. 2008; Liu et al. 2008; Chaibub Neto et al. 2008, 2009; Winrow et al. 2009; Millstein et al. 2009). Current approaches for causal inference in systems genetics can be classified into whole network scoring methods (Zhu et al. 2004, 2007, 2008; Li et al. 2006; Liu et al. 2008; Chaibub Neto et al. 2008, 2010; Winrow et al. 2009; Hageman et al. 2011) or pairwise methods, which focus on the inference of causal relationships among pairs of phenotypes (Schadt et al. 2005; Li et al. 2006; Kulp and Jagalur 2006; Chen et al. 2007; Aten et al. 2008; Millstein et al. 2009; Li et al. 2010; Duarte and Zeng 2011). In this article we develop a pairwise approach for causal inference among pairs of phenotypes.

Two key assumptions for causal inference in systems genetics are genetic variation preceding phenotypic variation and Mendelian randomization of alleles in unlinked loci. These conditions together, which provide temporal order and eliminate confounding of other factors, justify causal claims between QTL and phenotypes. Causal inference among phenotypes is justified by conditional independence relations under Markov properties (Li et al. 2006; Chaibub Neto et al. 2010).

Given a pair of phenotypes, \( Y_1 \) and \( Y_2 \), that co-map to the same quantitative trait locus, \( Q \), our objective is to learn which of the four distinct models, \( M_1, M_2, M_3, \) and \( M_4 \), depicted in Figure 1, is the best representation for the true relation between \( Y_1 \) and \( Y_2 \). Models \( M_1, M_2, M_3, \) and \( M_4 \) represent, respectively, the causal, reactive, independence, and full models as collapsed versions of more complex regulatory networks.
For instance, when the data are transcriptional and one gene is upstream of other genes, the regulation of the upstream gene may affect those downstream, even when the regulation takes place via post-transcriptional mechanisms and, hence, is mediated by unobserved variables. Transcriptional networks should be interpreted as collapsed versions of more complicated networks, where the presence of an arrow from a QTL to a phenotype or from one phenotype to another simply means that there is a directional influence of one node on another (that is, there is at least one path in the network where the node in the tail of the arrow is upstream of the node in the head). Supporting Information, Figure S1 shows a few examples of networks and their collapsed versions. Our goal in this article is to infer the causal direction between two nodes, and the term “causal” should be interpreted as causal direction, meaning either direct or indirect causal relations.

In this article, we propose novel causal model selection hypothesis tests and compare their performance to the AIC and BIC model selection criteria and to a causality inference test (CIT) proposed by Millstein et al. (2009). AIC (Akaike 1974) and BIC (Schwarz 1978) are widely used penalized likelihood criteria performing model selection among models of different sizes. Overparameterized models tend to overfit the data and, when comparing models with different dimension, it is necessary to counterbalance model fit and model parsimony by adding a penalty term that depends on the number of parameters. CIT is an intersection-union test, in which a number of equivalence and conditional F tests are conservatively combined in a single test. P-values are computed for models M1 and M2 in Figure 1, but not for the M3 or M4 models, and the decision rule for model calling goes as follows: (1) call M1 if the M1 P-value is less than a significance threshold α and the M2 P-value is greater than α; (2) call M2 if it is the other way around; (3) call M1 if both P-values are greater than α; and (4) make a “no call” if both P-values are less than α. M1 and M2 call actually means that the model is not M1 or M2 and could correspond to an M3 or M4 model. Note that the CIT makes a no call when both M1 and M2 models are simultaneously significant.

Our causal model selection tests (CMSTs) adapt and extend Vuong’s (1989) and Clarke’s (2007) tests to the comparison of four models. Vuong’s model selection test is a formal parametric hypothesis test devised to quantify the uncertainty associated with a model selection criterion, comparing two models based on their (penalized) likelihood scores. It uses the (penalized) log-likelihood ratio scaled by its standard error as a test statistic and tests the null hypothesis that both models are equally close to the true data generating process. While the (penalized) log-likelihood scores can determine only whether, for example, model A fits the data better than model B, Vuong’s test goes one step further and attaches a P-value to the scaled contrast of (penalized) log-likelihood scores. In this way it can interrogate whether the better fit of model A compared to model B is statistically significant. Vuong’s test tends to be conservative and low powered. Clarke (2007) proposed a nonparametric version that achieves an increase in power at the expense of higher miss-calling error rates by using the median rather than the mean of (penalized) log-likelihood ratio.

We propose three distinct versions of causal model selection tests: (1) the parametric CMST test, which corresponds to an intersection-union test of six separate Vuong’s tests; (2) the nonparametric CMST test, constructed as an intersection-union test of six of Clarke’s tests; and (3) the joint-parametric CMST test, which mimics an intersection-union test and is derived from the joint distribution of Vuong’s test statistics. These CMST tests inherit from Vuong’s test the property that none of the models being compared need be correct. That is, the true model may describe a more complicated network, including unobserved factors. Our approach simply selects the wrong model that is closest to the (unknown) true model.

Methods

Vuong’s model selection test

The Kullback–Leibler Information Criterion (KLIC) (Kullback 1959) measures the closeness of a probability model to the true distribution of data. Sawa (1978) showed that the KLIC orders approximate models by comparing the expected value of the log likelihood under the true model. Vuong (1989) used this result to develop an empirical test of two models based on the sample mean of the log-likelihood ratio scaled by its sample standard error.

Formally, \( f(y|x; \theta) : \theta \in \Theta \) represents a parametric family of conditional models and

\[
\text{KLIC}(h^0; f) = E^0[\log h^0(y|x)] - E^0[\log f(y|x; \theta_0)] = \int y \left( \log h^0(y|x) - \log f(y|x; \theta_0) \right) h^0(x) \, dx,
\]

(1)

where \( E^0 \) represents the expectation with respect to the true joint distribution \( h^0(y, x) = h^0(y|x)h^0(x) \), and \( \theta_0 \) is the parameter value that minimizes the KLIC distance from \( f \) to the true model (Sawa 1978). Note that \( f \) need not belong to the same parametric family as \( h^0 \).

A model \( f_1(y|x; \theta_1) \), denoted \( f_1 \), is a better approximation to the true model \( \theta_0 \) if and only if \( \text{KLIC}(h^0; f_1) < \text{KLIC}(h^0; f_2) \), or alternatively, \( E^0[\log f_1] > E^0[\log f_2] \) (Sawa 1978). Vuong’s model selection test is based on the latter criterion and the null and alternative hypotheses are defined as

\[
H_0 : E^0[LR_{12}] = 0, \quad H_1 : E^0[LR_{12}] > 0, \quad H_2 : E^0[LR_{12}] < 0,
\]

(2)

where \( LR_{12} = \log f_1 - \log f_2 \). The null hypothesis is \( f_1 \) and \( f_2 \) are equally close to the true distribution. The alternative
hypothesis $H_1$ means that $f_1$ is better than $f_2$ and conversely for the alternative $H_2$.

The quantity $E[LR_{12}]$ is unknown, but under fairly general conditions the sample mean and variance of $LR_{12} = \log \hat{f}_{1i} - \log \hat{f}_{2i}$ converge almost surely to $E[LR_{12}]$ and $\text{Var}[LR_{12}] = \sigma_{12}$, where $\hat{f}_{1i} = f_1(y_j | x_i; \hat{\theta}_1)$ and $\hat{\theta}_1$ is the maximum-likelihood estimate of $\theta_1$ (Vuong 1989). Let $LR_{12} = \sum_{i=1}^{n} LR_{12i}$, then, under $H_0$,

$$n^{-1/2} LR_{12}/\sqrt{\sigma_{12}} \rightarrow d N(0,1).$$ (3)

Under $H_1$ this test statistic converges almost surely to $\infty$, whereas, under $H_2$, it converges to $-\infty$ (Vuong 1989).

Vuong’s test is based on the unadjusted log-likelihood ratio statistic. However, competing models may have different dimensions, requiring a complexity penalty. The penalized log-likelihood ratio is given by $LR_{12} = LR_{12} - D_{12}$, where the penalty $D_{12}$ is the difference of the number of parameters between models 1 and 2 (for AIC) or this value rescaled by $(\log n)/2$ (for BIC). Because the penalty term is of smaller size than $n^{1/2}$, the adjusted log-likelihood ratio accounting for different model dimensions

$$Z_{12} = n^{-1/2} LR_{12}/\sqrt{\sigma_{12}}$$ (4)

has the same asymptotic properties as $n^{-1/2} LR_{12}/\sqrt{\sigma_{12}}$ (Vuong 1989).

The $P$-value of Vuong’s test is given by $p_{12} = P(Z_{12} \geq z_{12}) = 1 - \Phi(z_{12})$, where $\Phi()$ represents the cumulative density function of a standard normal variable (Vuong 1989). Note that since $Z_{12} = -Z_{12}$, $p_{21} = 1 - \Phi(z_{21}) = \Phi(z_{12})$, so that $p_{12} + p_{21} = 1$. This property of the Vuong’s test ensures that the $P$-values of the intersection-union tests cannot be simultaneously significant.

Figure S2 illustrates how Vuong’s test trades a reduction in false positives against a reduction in statistical power. In our applications we need to account for both nested and nonnested models. While the presented test corresponds to Vuong’s test for strictly nonnested models, it is also valid for nested models when we adopt penalized likelihood scores (see File S1, for further details).

**Clarke’s model selection paired sign test**

The model selection paired sign test (Clarke 2007) is a nonparametric alternative to Vuong’s test, testing the null hypothesis that the median log-likelihood ratio is 0. Clarke’s test statistic, $T_{12}$, is a sign test on $LR_{12}$. Under the null hypothesis that the median log-likelihood ratio is zero, $T_{12}$ has a binomial distribution, and the $P$-value for comparing models 1 and 2 is

$$p_{12} = P(T_{12} \geq t_{12}) = \sum_{k=t_{12}}^{n} C_n^k 2^{-n},$$ (5)

with $C_n^k = n!/(k!(n-k)!$. The $P$-values for $T_{12}$ and $T_{21}$ do not add to 1 since the statistics are discrete, $p_{12} + p_{21} = 1 + C_n^0 2^{-n}$. Nonetheless, the $C_n^0 2^{-n}$ term decreases to 0 as $n$ increases, and, in practice, $p_{12} + p_{21} \approx 1$ even for moderate sample sizes. We adjust this test using the AIC or BIC penalty $D_{12}$,

$$T_{12} = \sum_{i=1}^{n} \mathbb{I}\{LR_{12i} \geq n^{-1}D_{12} > 0\},$$ (6)

to account for the varying dimensionality of the models.

**Causal model selection tests**

The four models $M_1$, $M_2$, $M_3$, and $M_4$ (Figure 1) are used to derive intersection-union tests based on the application of six separate Vuong (or Clarke) tests comparing, namely, $f_1 \times f_2, f_1 \times f_3, f_1 \times f_4, f_2 \times f_3, f_2 \times f_4$, and $f_3 \times f_4$. Sun et al. (2007) implicitly used intersection unions of Vuong’s tests to select among three nonnested models. Here, we present three distinct versions of the CMST: (1) parametric, (2) nonparametric, and (3) joint-parametric CMST tests. We implement the tests with penalized log likelihoods, but state results for log likelihoods.

Here we focus on model $M_1$ and $P$-value $p_1$, with analogous results and notation for the other three models. Starting with the parametric version, we test the null $H_0$: model $M_1$ is no closer to the true model than $M_2$, $M_3$, or $M_4$, against the alternative $H_1$: $M_1$ is closer to the true model than $M_2$, $M_3$, and $M_4$. More explicitly, we test,
Table 1 Model selection tests for models $M_1$, $M_2$, $M_3$, and $M_4$

| $H_0$ | Null distribution | $P$-value |
|-------|------------------|-----------|
| $H_0^{M1}$ | $Z_1 = (Z_{12}, Z_{13}, Z_{14}) \sim N_3(0, \rho_1)$ | $p_1 = P(Z_{12} \geq w_1, Z_{13} \geq w_1, Z_{14} \geq w_1)$ |
| $H_0^{M2}$ | $Z_2 = (Z_{21}, Z_{23}, Z_{24}) \sim N_3(0, \rho_2)$ | $p_2 = P(Z_{21} \geq w_2, Z_{23} \geq w_2, Z_{24} \geq w_2)$ |
| $H_0^{M3}$ | $Z_3 = (Z_{31}, Z_{32}, Z_{34}) \sim N_3(0, \rho_3)$ | $p_3 = P(Z_{31} \geq w_3, Z_{32} \geq w_3, Z_{34} \geq w_3)$ |
| $H_0^{M4}$ | $Z_4 = (Z_{41}, Z_{42}, Z_{43}) \sim N_3(0, \rho_4)$ | $p_4 = P(Z_{41} \geq w_4, Z_{42} \geq w_4, Z_{43} \geq w_4)$ |

Here $w_k = \min(p_k)$ for $k = 1, \ldots, 4$, and $\rho_k$ is defined in analogy with $\rho_1$ in Equation 10.

For any $\alpha$, if $p_1 \leq \alpha$, then $\min(p_2, p_3, p_4) \geq 1 - \alpha$. Therefore, the proposed CMST test has at most one significant model $P$-value at a time, in contrast to the CIT approach.

The joint-parametric CMST test corresponds to an intersection union of Clarke’s tests, exactly analogous to the parametric version. Because in practice $p_1 \geq p_2 \approx 1$ for Clarke’s test, the nonparametric CMST test also does not allow the detection of more than one significant model $P$-value.

Simple application of separate Vuong tests overlooks the dependency among the test statistics. A multivariate extension, the joint parametric CMST CMST, can be developed to address this caveat. For model $M_1$, and under the same general regularity conditions of Vuong (1989), the sample covariance of $LR_{12}$ and $LR_{13}$, $\sigma_{12,13}$, converges almost surely to $\text{Cov}_{\theta}[LR_{12}, LR_{13}] = \sigma_{12,13}$ (and similarly for all other covariance terms). Therefore, the sample covariance matrix, $\Sigma_1$, converges almost surely to $\Sigma_1$. From the multivariate central limit and Slutsky’s theorems (Shao 2003), if

$$
\left( \begin{array}{c} E^0[LR_{12}] \\ E^0[LR_{13}] \\ E^0[LR_{14}] \end{array} \right) = \left( \begin{array}{c} 0 \\ 0 \\ 0 \end{array} \right)
$$

then $Z_1 = \text{diag}(\Sigma_1)^{-\frac{1}{2}} \mathbf{L}_1 / \sqrt{n} \rightarrow N_3(0, \rho_1)$, where $\mathbf{L}_1 = (LR_{12}, LR_{13}, LR_{14})^T$ and $\rho_1 = \text{diag}(\Sigma_1)^{-\frac{1}{2}} \Sigma_1 \text{diag}(\Sigma_1)^{-\frac{1}{2}}$ is the correlation matrix $\rho_1$.

The condition in (9) is the worst case of the more general null hypothesis that $M_1$ is not better than at least one of $M_2$, $M_3$, or $M_4$, or

$$
H_0 : \min\{E^0[LR_{12}], E^0[LR_{13}], E^0[LR_{14}]\} \leq 0.
$$

We test this against the alternative that $M_1$ is better than all three, or

$$
H_1 : \min\{E^0[LR_{12}], E^0[LR_{13}], E^0[LR_{14}]\} > 0,
$$

using the statistic $W_1 = \min\{Z_1\}$, with $P$-value

$$
P(W_1 \geq w_1) = P(\min\{Z_{12}, Z_{13}, Z_{14}\} \geq w_1) = P(Z_{12} \geq w_1, Z_{13} \geq w_1, Z_{14} \geq w_1).
$$

The joint-parametric CMST test with $W_1$ follows the spirit of an intersection union test while accounting for dependency among test statistics. Table 1 depicts the joint CMST tests for all models.

The CMST tests are implemented in the R/qtlhot package available at CRAN. Although not explicitly stated in the notation, the pairwise models can easily account for additive and interactive covariates, and our code already implements this feature. When using this package please cite this article.

**Simulation studies**

We conducted two simulation studies. In the first “pilot study,” we focus on performance comparison of the AIC, BIC, CIT, and CMST methods with data generated from simple causal models. The goal is to understand the behavior of our methods in diverse settings. In the second “large-scale study,” we perform a simulation experiment, with data generated from causal models emulating QTL hotspot patterns. The goal is to understand the impact of multiple testing on the performance of our causality tests.

The pilot simulation study has data generated from models A to E in Figure 2. We conducted 10 simulation studies, generating data from the five models described above under sample sizes 112 (the size of our real data example) and 1000. For each model, we simulated 1000 backcrosses. We chose simulation parameters to ensure that 99% of the $R^2$ coefficients between phenotypes and QTL ranged between 0.08 and 0.32 for the simulations based on sample size of 112 subjects and between 0.01 to 0.20 for the simulations based on 1000 subjects (see File S2, File S3, and File S4 for details). We evaluated the method’s performance using statistical power, miss-calling error rate, and precision. These quantities were computed as,

$$
\text{Power} = \frac{TP}{N}, \quad \text{Miss-calling error} = \frac{FP}{N},
$$

$$
\text{Precision} = \frac{TP}{TP+FP}
$$

where $N$ is the total number of tests, and TP (true positives) and FP (false positives) are defined according to Table 2, which depicts possible calls against simulated models and tabulates whether a specific call correctly represents the
causal relationship between the phenotypes in the model from which the data were generated.

In the large-scale simulation study we investigate the empirical FDR (1 minus the precision) and power levels achieved by the CMST tests using the Benjamini and Hochberg (1995) and the Benjamini and Yekutieli (2001) FDR control procedures (denoted, respectively, by BH and BY), as well as no multiple testing correction. We simulate data from the models in Figure 3, which emulate eQTL hotspot patterns, i.e., genomic regions to which hundreds or thousands of traits co-map (West et al. 2007). In each simulation we generated 1000 distinct backcrosses with phenotypic data on 5001 traits on 112 individuals. We simulated unequally spaced markers for model F, but equally spaced markers for G, with Q1 and Q set 1 cM apart. Because we fit almost three million hypothesis tests in this simulation study, we did not include the CIT tests in this investigation, restricting our attention to the computationally more efficient CMST tests. The details for our choice of simulation parameters and QTL mapping are presented in File S2, File S3, and File S4. A frequent goal in eQTL hotspots studies is to determine a master regulator, i.e., a transcript that regulates the transcription of the other traits mapping to the hotspot. A promising strategy toward this end is to test the cis traits (i.e., transcripts physically located close to the QTL hotspot) against all other co-mapping traits. Our simulations evaluate the performance of the CMST tests in this setting.

**Results**

**Pilot simulation study results**

Figure 4 depicts the power, miss-calling error rate, and precision of each of the methods based on the simulation results of all five models in Figure 2. The results of the AIC and BIC approaches are constant across all significance levels since these approaches do not provide a measure of statistical significance. For those methods, we simply fit the models to the data and select the model with the best (smallest) score.

Overall, the AIC, BIC, and CIT showed high power, high miss-calling error rates, and low precision. The CMST methods, on the other hand, showed lower power, lower miss-calling error rates, and higher precision. The non-parametric CMST tended to be more powerful but less precise than the other CMST approaches. As expected, for sample size 1000, all methods showed an increase in power and precision and decrease in miss-calling error rate.

Figure S3, Figure S4, Figure S5, Figure S6, and Figure S7 show the simulation results data for each one of models A to E, when sample size is 112. Figure S8, Figure S9, Figure S10, Figure S11, and Figure S12 show the same results for sample size 1000. Some of the simulated models were intrinsically more challenging than others. For instance, in the absence of latent variables the causal and independence relations can often be correctly inferred by all methods (see the results for models A and D in Figure S3, Figure S6, Figure S8, and Figure S11). However, the presence of hidden variables in models B and E tend to complicate matters. Nonetheless, although the AIC, BIC, and CIT methods tend to detect false positives at high rates in these complicated situations, the CMST tests tend to forfeit making calls and tend to detect fewer false positives (see Figure S4, Figure S7, Figure S9, and Figure S12). Model C is particularly challenging (Figure S5 and Figure S10), showing the highest false-positive detection rates among all models.

In genetical genomics experiments we often restrict our attention to the analysis of cis-genes against trans-genes. In this special case it is reasonable to expect the pleiotropic

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**Table 2 True and false positives**

| CMST  | Model A | Model B | Model C | Model D | Model E |
|-------|---------|---------|---------|---------|---------|
| $M_1$ | TP      | TP      | FP      | FP      | FP      |
| $M_2$ | FP      | FP      | FP      | FP      | FP      |
| $M_3$ | FP      | FP      | FP      | FP      | FP      |
| $M_4$ | FP      | FP      | TP      | FP      | TP      |

| CIT   | Model A | Model B | Model C | Model D | Model E |
|-------|---------|---------|---------|---------|---------|
| $M_1$ | TP      | TP      | FP      | FP      | FP      |
| $M_2$ | FP      | FP      | FP      | FP      | FP      |
| $M_3$ | FP      | FP      | TP      | FP      | TP      |

Each entry $i,j$ represents whether the call on row $i$ is a true positive (TP) or as false positive (FP), when the data are generated from the model on column $j$. For instance, when data are generated from models $A$ or $B$, a $M_3$ call represents a true positive, whereas a $M_2$, $M_3$, or $M_4$ call represents a false positive for the AIC, BIC, and CMSTs approaches (for the CIT a $M_2$ or $M_3$ call represents false positive). Note that a $M_3$ call is considered a true positive for model $C$ (in addition to model $E$) because it corresponds to model $M_2^*$ on Figure 1 and, hence, is distribution equivalent to model $M_3$. Please note too that because the CIT provides $P$-values for only the $M_1$ and $M_2$ calls, but not for the $M_3$ and $M_4$ calls, and its output is $M_1$, $M_2$, or $M_3$ we classify a $M_3$ call as a true positive for models $C$, $D$, and $E$. Observe that by doing so we are actually giving an unfair advantage for the CIT approach, since when the data are generated from, say, model $E$, the CIT needs only to discard models $M_1$ and $M_2$ as nonsignificant to detect a “true positive.” The AIC, BIC, and CMST approaches, on the other hand, need to discard models $M_1$, $M_2$, and $M_3$ as nonsignificant and accept model $M_3$ as significant.
causal relationship depicted in model C to be much less frequent than the relationships shown in models A, B, D, and E, so that the performance statistics shown in Figure 4 might be negatively affected to an unnecessary degree by the simulation results from model C.

To investigate the performance of methods in the cis-against trans-case, we present in Figure 5 the simulation results based on models A, B, D, and E only. Comparison of Figures 4 and 5 show an overall improvement in power, decrease in miss-calling rates and increase in precision.

In the analysis of trans- against trans-genes there is no a priori reason to discard the relationship depicted in model C, and more false positives should be expected. The CMST approaches, specially the joint parametric and parametric CMST methods, tend to detect a much smaller number of false positives than the AIC, BIC, and CIT approaches, as shown in Figure S5 and Figure S10.

Large-scale simulation study results

With the possible exception of the nonparametric version, the previous simulation study suggests that the CMST tests can be quite conservative. Therefore, it is reasonable to ask whether multiple testing correction is really necessary to achieve reasonable false discovery rates (FDR).

Figure 6 presents the observed FDR and power using uncorrected, BH corrected, and BY corrected P-values for the simulations based on model G. Figure 6, top, shows that, except for the AIC-based nonparametric CMST, the observed FDRs were considerably lower than the P-value cutoff, suggesting that multiple testing adjustment is not necessary for the CMST tests. Furthermore, comparison of the bottom panels shows that the BH and BY adjustments leads to a reduction in power (specially for the BY adjustment) for the joint and parametric tests at the expense of small drop in FDR levels (that were already low without any correction). For the nonparametric tests, on the other hand, BH corrections leads to bigger drops in FDR (specially for the AIC based test) and smaller drops in power. The BY correction appears too conservative even for the nonparametric tests. The results for model F are similar (Figure S13).

Yeast data analysis and biologically validated predictions

We analyzed a budding yeast genetic genomics data set derived from a cross of a standard laboratory strain and a wild isolate from a California vineyard (Brem and Kruglyak 2005).
The data consist of expression measurements on 5740 transcripts measured on 112 segregant strains with dense genotype data on 2956 markers. Processing of the expression measurements raw data was performed as described in Brem and Kruglyak (2005), with an additional step of converting the processed measurements to normal scores. We performed QTL analysis using Haley–Knott regression (Haley and Knott 1992) with the R/qtl software (Broman et al. 2003). We used Haldane’s map function, genotype error rate of 0.0001, and set the maximum distance between positions at which

Figure 5 Power (A and D), miss-calling error rate (B and E), and precision (C and F) restricted to the cis- vs. trans-cases. The x-axis represents the significance levels used for computing the results. The results were computed using only the simulated models A, B, D, and E in Figure 2, since the pleiotropic causal relationship depicted in model C is expected to be much less frequent than the others when testing cis- vs. trans-case. (A–C) The simulations based on sample size 112; (D–F) the results for sample size 1000. Dashed and solid curves represent, respectively, AIC- and BIC-based methods. Green: parametric CMST. Red: nonparametric CMST. Blue: joint-parametric CMST. Black: AIC and BIC. Orange: CIT. The shaded line on B and E corresponds to the $\alpha$ levels.

Figure 6 Observed FDR and power for the simulations based on model G. The x-axis represents the P-value cutoffs used for computing the results. Dashed and solid curves represent, respectively, AIC- and BIC-based methods. Green: parametric CMST. Red: nonparametric CMST. Blue: joint-parametric CMST. Black: AIC and BIC. The shaded line in the top corresponds to the $\alpha$ levels.
genotype probabilities were calculated to 2 cM. We adopted a permutation LOD threshold (Churchill and Doerge 1994) of 3.48, controlling the genome-wide error rate of falsely detecting a QTL at a significance level of 5%.

To evaluate the precision of the causal predictions made by the methods we used validated causal relationships extracted from a database of 247 knock-out experiments in yeast (Hughes et al. 2000; Zhu et al. 2008). In each of these experiments, one gene was knocked out, and the expression levels of the remainder genes in control and knocked-out strains were interrogated for differential expression. The set of differentially expressed genes form the knock-out signature (ko-signature) of the knocked-out gene (ko-gene) and show direct evidence of a causal effect of the ko-gene on the ko-signature genes. The yeast data cross and knocked-out data analyzed in this section are available in the R/qtlyeast package at GITHUB (https://github.com/byandell/qtlyeast).

To use this information, we: (i) determined which of the 247 ko-genes also showed a significant eQTL in our data set; (ii) for each one of the ko-genes showing significant linkages, we determined which other genes in our data set also co-mapped to the same QTL of the ko-gene, generating, in this way, a list of putative targets of the ko-gene; (iii) for each of the ko-gene/putative targets list, we applied all methods using the ko-gene as the Y1 phenotype, the putative target genes as the Y2 phenotypes, and the ko-gene QTL as the causal anchor; (iv) for the AIC- and BIC-based nonparametric CMST tests we adjusted the P-values according to the Benjamini and Hochberg FDR control procedure; and (v) for each method we determined the “validated precision,” computed as the ratio of true positives by the sum of true and false positives, where a true positive is defined as an inferred causal relationship where the target gene belongs to the ko-signature of the ko-gene, and a false positive is given by an inferred causal relation where the target gene does not belong to the ko-signature.

In total 135 of the ko-genes showed a significant QTL, generating 135 putative target lists. A gene belonged to the putative target list of a ko-gene when its 1.5 LOD support interval (Lander and Botstein 1989; Dupuis and Siegmund 1999; Manichaikul et al. 2006) contained the location of the ko-gene QTL. The number of genes in each of the putative target lists varied from list to list, but in total we tested 31,975 “ko-gene/putative target gene” relationships.

Figure 7 presents the number of inferred true positives, number of inferred false positives, and the prediction precision across varying target significance levels for each one of the methods. The CIT, BIC, and AIC had a higher number of true positives than the CMST approaches, with the AIC-based CMST methods having less power than the BIC-based CMST methods. However, the CIT, BIC, and AIC also inferred the highest numbers of false positives (Figure 7B) and showed low prediction precisions (Figure 7C). From Figure 7C we see that the CMST tests show substantially higher precision rates across all target significance levels compared to the AIC, BIC, and CIT methods. Among the CMST approaches, the joint-parametric CMST tended to show the highest precision, followed by the nonparametric and parametric CMST tests.

The results presented in Figure 7 were computed using all 135 ko-genes. However, in light of our simulation results, which suggest that the analysis of cis- against trans-genes is usually easier than the analysis of trans- against trans-genes, we investigated the results restricting ourselves to ko-genes with significant cis-QTL. Only 28 of the 135 ko-genes were cis-traits, but, nonetheless, were responsible for 2947 of the total 31,975 “ko-gene/putative target gene” relationships. Figure 8 presents the results restricted to the cis-ko-genes. All methods show improvement in precision, corroborating our simulation results. Once again, the CMST tests showed higher precision than the CIT, AIC, and BIC.

Discussion

In this article, we proposed three novel hypothesis tests that adapt and extend Vuong’s and Clarke’s model selection tests to the comparison of four models, spanning the full range of possible causal relationships among a pair of phenotypes. Our CMST tests scale well to large genome wide analyses because they are fully analytical and avoid computationally expensive permutation or resampling strategies.

Another useful property of the CMST tests, inherited from Vuong’s test, is their ability to perform model selection among misspecified models. That is, the correct model need not be one of the models under consideration. Accounting for the misspecification of the models is key. In general, any two phenotypes of interest are embedded in a complex network and are affected by many other phenotypes not considered in the grossly simplified (and thus misspecified) pairwise models.

Overall, our simulations and real data analysis show that the CMST tests are better at controlling miss-calling error rates and tend to outperform the AIC, BIC, and CIT methods in terms of statistical precision. However, they do so at the expense of a decrease in statistical power. While an ideal method would have high precision and power, in practice there is always a trade-off between these quantities. Whether a more powerful and less precise, or a less powerful and more precise, method is more adequate depends on the biologist’s research goals and resources. For instance, if the goal is to generate a rank-ordered list of promising candidates genes that might causally affect a phenotype of interest and the biologist can easily validate several genes, a larger list generated by more powerful and less precise methods might be more appealing. However, in general, follow-up studies tend to be time consuming and expensive, and only a few candidates can be studied in detail. A long list of putative causal traits is not useful if most are false positives. High power to detect causal relations alone is not enough. A more precise method that conservatively identifies candidates with high confidence can be more appealing (see also Chen et al. 2007).
Further, the exploratory goal is often to identify causal agents without attempting to reconstruct entire pathways. Therefore, much information about the larger networks in which the tested pairs of traits reside is unknown and generally unknowable and contributes to the large unexplained variation that in turn results in low power. Our method accurately reflects this difficulty to detect causal relationships in the presence of noisy high-throughput data and poorly understood networks.

Interestingly, our data analysis and simulations also suggest that the analysis of cis-against trans-gene pairs is less prone to detect false positives than the analysis of trans-against trans-gene pairs. Our simulations suggest that model selection approaches have difficulty ordering the phenotypes when the QTL effect reaches the truly reactive gene by two or more distinct paths, only one of which is mediated by the truly causal gene (see Figure S1C, for an example).

When we test causal relationships among gene expression phenotypes, the true relationships might not be a direct result of transcriptional regulation. For instance, the true causal regulation might be due to methylation, phosphorylation, direct protein–protein interaction, transcription factor binding, etc. Margolin and Califano (2007) have pointed out the limitations of causal inference at the transcriptional level, where molecular phenotypes at other layers of regulation might represent latent variables. Model $M_4$ (see Figure 1) can account for these latent variables and can test this scenario explicitly.

Furthermore, as pointed out by Li et al. (2010), causal inference depends on the detection of subtle patterns in the correlation between traits. Hence, it can be challenging even when the true causal relations take place at the transcriptional level. The authors point out that reliable causal inference in genome-wide linkage and association studies require large sample sizes and would benefit from: (i) incorporating prior information via Bayesian reasoning; (ii) adjusting for experimental factors, such as sex and age, that might induce correlations not explained the the causal relations; and (iii) considering a richer set of models than the four models accounted in this article.

The CMST tests represent a step in the direction of reliable causal inference in three accounts. First, they tend to be precise, declining to make calls in situations where alternative approaches usually deliver a flood of false-positive calls. Second, the CMST tests can adjust for experimental factors by modeling them as additive and interactive covariates. Third, the CMST tests can be applied to nonnested models of different dimensions and can be readily extended to incorporate a larger number of models by implementing intersection-union tests on a larger number of Vuong’s tests. For the joint-parametric test a higher-dimensional null distribution is required.

FDR control for the CMST approaches is a challenging problem as our tests violate the key assumption, made by FDR control procedures, that the distribution of the $P$-values under the null hypothesis are uniformly distributed (Benjamini and Hochberg 1995; Storey and Tibshirani 2003). Recall that the CMST $P$-values are computed as the maximum across other $P$-values, and the maximum of multiple uniform random variables no longer follows a uniform distribution. Additionally, the CMST tests are usually not independent since we often test the same cis-trait against several trans-traits, so that the additional assumption of independent test statistics made by the original Benjamini–Hochberg procedure does not hold. The Benjamini–Yekutieli (BY) procedure relaxes the independent test statistics assumption, and we explore both these corrections in our simulations. Our results suggest that BH and BY multiple testing correction should not be performed for the joint and the parametric CMST tests, as the FDR levels are lower than the nominal level without any correction and are too conservative with severe reduction in statistical power with the application of
BH and BY control. The nonparametric CMST tests, on the other hand, seemed to benefit from BH correction, showing slight decrease in power with concomitant decrease in FDR, in spite of the nonparametric CMST tests being based on discrete test statistics and the BH procedure being developed to handle *P*-values from continuous statistics. Inspection of the *P*-value distributions (see Figure S14, Figure S15, Figure S16, and Figure S17) suggests that the smaller *P*-values of the nonparametric tests, relative to the other approaches, are the reason for the higher power achieved by the BH corrected nonparametric tests. The BY procedure, on the other hand, tended to be too conservative even for the nonparametric CMST tests.

The CMST approach is currently implemented for inbred line crosses. Extension to outbred populations involving mixed effects models is yet to be done. Although in this article we focused on mRNA expression traits, the CMST tests can be applied to any sort of heritable phenotype, including clinical phenotypes and other “omic” molecular phenotypes.

The higher statistical precision and computational efficiency achieved by our fully analytical hypothesis tests will help biologists to perform large-scale screening of causal relations, providing a conservative rank-ordered list of promising candidate genes for further investigations.

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Modeling Causality for Pairs of Phenotypes in System Genetics

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Figure S 1 Network models and their collapsed versions. The collapsed networks (bottom panels) represent simplified versions of the true networks (top panels), where nodes other than $Q$, $Y_1$, and $Y_2$ are ignored, even though they still represent the correct causal flow among these three nodes in the true network. Consider, for example, network $c$ and its collapsed version $c'$. The path $Q \rightarrow Y_3 \rightarrow Y_1$ in $c$ is collapsed to $Q \rightarrow Y_1$ in $c'$. The paths $Y_1 \rightarrow Y_5 \rightarrow Y_2$ and $Y_1 \rightarrow Y_6 \rightarrow Y_2$ in $c$ are collapsed to $Y_1 \rightarrow Y_2$ in $c'$. The path $Q \rightarrow Y_3 \rightarrow Y_4 \rightarrow Y_7 \rightarrow Y_2$ in $c$ is collapsed to $Q \rightarrow Y_2$ in $c'$. 
Figure S2  Model selection via log-likelihood ratio versus Vuong’s test.

Figure S2 illustrates how Vuong’s test works. We generated 1,000 data-sets from the model $X \rightarrow Y_1 \rightarrow Y_2$ and applied Vuong’s test to the comparison of models $M_1 : X \rightarrow Y_1 \rightarrow Y_2$ against $M_2 : X \rightarrow Y_2 \rightarrow Y_1$. The top panels present 3D scatter plots of the test statistics $Z_{12}$ against the $R^2$ values of the regression of $Y_1$ on $X$, $R^2(Y_1, X)$, and the $R^2$ values of the regression of $Y_2$ on $X$, $R^2(Y_2, X)$. The data points are color coded as blue, red and grey, representing, respectively, $M_1$, $M_2$ and “no calls”. Note that because model $M_1$ corresponds to the true model, we have that the a $M_1$ call is always correct, whereas
a $M_2$ call is always incorrect in this example. Therefore, blue and red points represent, respectively, correct and incorrect calls. The bottom panels follow the same color coding and show the projections of the 3D scatter plots into the $R^2(Y_1, X)$ by $R^2(Y_2, X)$ plane.

The left panels of Figure S2 show the model selection results based on the log-likelihood ratio (LR) criterium, where positive $\hat{L}_{R_{12}}$ values support $M_1$ and negative $\hat{L}_{R_{12}}$ values support $M_2$ (note that we actually use the $Z_{12}$ test statistics, instead of $\hat{L}_{R_{12}}$ statistics, but the results are equivalent). Because we generate the data from model $X \rightarrow Y_1 \rightarrow Y_2$, it will usually be the case that $X$ explains a greater proportion of the variability of $Y_1$ than of $Y_2$. In other words, $R^2(Y_1, X)$ will tend to be higher than $R^2(Y_2, X)$. However, some of the data-sets show the opposite trend due to random noise on the data. The bottom left panel shows that the log-likelihood criterium tends to make incorrect calls when $R^2(Y_1, X) < R^2(Y_2, X)$.

The right panels of Figure S2 show the model selection results derived from Vuong’s test. Now we see that most of the incorrect calls made by the log-likelihood criterium (red points) are not significant (grey points) according to Vuong’s test, that requires that $Z_{12} \leq -1.64$ or $Z_{12} \geq 1.64$ for statistical significance at a 5% level. The drawback is the reduction in power to detect the correct calls, since not only red dots are replaced by grey dots, but many of the blue dots are turned into grey, as well. These figures illustrate how Vuong’s test trade an increase in precision for a reduction in statistical power to detect true positives.
A technical note on Vuong’s test

Vuong (1989) fully characterized the asymptotic distribution of the log-likelihood ratio statistic under the most general conditions. He showed that the form of the asymptotic distribution of the log-likelihood ratio depends on whether the models are observationally identical or not. Two models are observationally identical if their probability densities are the same, when evaluated at the respective pseudo-true parameter values, i.e., $f_1(y \mid x; \theta_{1*}) = f_2(y \mid x; \theta_{2*})$ for almost all $(y, x)$, where the pseudo-true parameter values, $\theta_{k*}$, corresponds to the parameter value that minimizes the Kullback-Leibler distance from the true model (Sawa 1978).

Explicitly, Vuong showed (Theorem 3.3 on page 313) that under very general conditions:

1. If $f_1(y \mid x; \theta_{1*}) = f_2(y \mid x; \theta_{2*})$, then $2LR_{12}(\hat{\theta}_1, \hat{\theta}_2)$ converges in distribution to a weighted sum of chi-square distributions.

2. If $f_1(y \mid x; \theta_{1*}) \neq f_2(y \mid x; \theta_{2*})$, then

$$
\frac{1}{\sqrt{n}} \left( LR_{12}(\hat{\theta}_1, \hat{\theta}_2) - E^0 \left[ \log \frac{f_1(y \mid x; \theta_{1*})}{f_2(y \mid x; \theta_{2*})} \right] \right) \rightarrow^d N(0, \sigma_{12,12})
$$

Because of this interesting asymptotic behavior Vuong had to proposed 3 distinct model selection tests: one for strictly non-nested models, that are always not observationally identical; another for overlapping models that might or might not be observationally identical; and a third for nested models, that are always observationally identical. (Nested models are always observationally identical because the nested model cannot be better
than the full model and both models are equally close to the true model if and only if they are the same.)

In our applications, models $M_1$, $M_2$ and $M_3$ are not nested on each other, but are nested on models $M'_1$, $M'_3$ and $M'_4$, respectively (Figure 1 in the main text). Hence, our model selection tests consider pairs of models that are either non-nested or nested. In the Methods section we presented Vuong’s test for not observationally identical models, that is suitable for the comparison of strictly non-nested models ($M_1 \times M_2$, $M_1 \times M_3$ and $M_2 \times M_3$).

We point out, however, that even though we perform model selection tests between nested models ($M_1 \times M_4$, $M_2 \times M_4$ and $M_3 \times M_4$) we don’t need to use Vuong’s test for nested models because our test statistics are based on penalized log-likelihoods instead of log-likelihoods, and our penalized models are not observationally identical for nested models too. In other words, even though $f_1(y \mid x; \theta_{1*}) = f_4(y \mid x; \theta_{4*})$ when model 1 is nested in model 4, we have that $f_1(y \mid x; \theta_{1*}) - p_1 \neq f_4(y \mid x; \theta_{4*}) - p_4$ since the penalty $p_1$ is smaller than $p_4$. Therefore, we can simply use Vuong’s test for not observationally identical models in this case too.

On a technical note, we point out that Vuong’s Theorem 3.3 still holds when we replace the log-likelihood ratio by the penalized log-likelihood ratio. The demonstration mimics Vuong’s original proof presented on page 327. We just need to replace the log-likelihoods by penalized log-likelihoods in the Taylor expansion of the log-likelihoods around the maximum likelihood estimates.
File S2

Simulation studies

Here we provide further details on the simulation studies presented in the main text.

File S3

Pilot simulation study

We conducted a total of 10 simulation studies, generating data from the five models described in Figure 2 in the main text using sample sizes 112 and 1,000 (the choice 112 was motivated by the sample size in our real data example). For each model, we simulated 1,000 backcrosses composed with 3 chromosomes of length 100cM containing 101 unequally spaced markers per chromosome. For each one of the simulated backcrosses, the additive and dominance genetic effects were sampled, respectively, from the $U[-0.75, 0.75]$ and $U[0, 0.75]$ distributions, where $U[a, b]$ represents the uniform distribution on the interval $[a, b]$. Residual error rates were sampled from $U[0.5, 1.5]$, and the phenotype to phenotype regression coefficients in Figures 2 A, B and C were sampled from $U[-1, 1]$. The hidden-variable to phenotype regression coefficients on Figures 2 B and E were sampled from $U[-1, 1]$ and $U[0.5, 1]$, respectively. This choice of parameters ensured that approximately 99% of the $R^2$ coefficients between phenotypes and QTL ranged between 0.08 and 0.32 for the simulations based on sample size of 112 subjects (see Figure SI.2a, and the axis scales on Figures S3-S7) and between 0.01 to 0.20 for the simulations based on 1,000 subjects (see Figure SI.2b, and the axis scales on Figures S8-S12).

The backcross simulations and the QTL mapping analyses were performed using the R/qtl software (Broman et al. 2003). We performed Haley-Knott regression (Haley and Knott 1992) and adopted Haldane’s map function, genotype error rate of 0.0001, and set...
the maximum distance between positions at which genotype probabilities were calculated to 2cM. We used a permutation LOD threshold (Churchill and Doerge 1994) of 2.24 for the QTL mapping analysis, aiming to control the genome wide error rate of falsely detecting a QTL at a 5% rate.

Often times the phenotypes map to nearby but not precisely the same QTL, and we need to decide which QTL to use as the causal anchor. When testing expression traits against clinical traits, Millstein et al. (2009) and Schadt et al. (2005) suggest using the clinical trait QTL as the anchor. We adopt a different approach. When the phenotypes map to distinct regions that are less than 2cM apart we determine the QTL position using both phenotypes, jointly, as follows. For each pair of phenotypes \((Y_1, Y_2)\) we perform unconditional mapping analysis for \(Y_1\) and \(Y_2\) and conditional mapping analysis for \(Y_2\) given \(Y_1\). Let \(LOD_1\) represent a LOD score for the mapping analysis of \(Y_1\), and \(LOD_{2|1}\) for the mapping analysis of \(Y_2\) given \(Y_1\). Since

\[
\log_{10} \left\{ \frac{f(y_1, y_2 | q)}{f(y_1, y_2)} \right\} = \log_{10} \left\{ \frac{f(y_2 | y_1, q)}{f(y_2 | y_1)} \right\} + \log_{10} \left\{ \frac{f(y_1 | q)}{f(y_1)} \right\},
\]

we compute the joint LOD score of \((Y_1, Y_2)\) as \(LOD_{1,2} = LOD_1 + LOD_{2|1}\) (or equivalently as \(LOD_{1,2} = LOD_2 + LOD_{1|2}\)). We determine the peak QTL position, \(\lambda\), using the \(LOD_{1,2}\) scores profile and assign the QTL to \(Y_1\) and \(Y_2\) if \(LOD_1\) and \(LOD_2\) are greater than the mapping threshold at the \(\lambda\) position. Figure SI1 illustrates our approach. When both phenotypes co-map to more than one QTL we select the QTL with the highest joint mapping peak.
Figure S1.1  We simulated data from a model $Q \rightarrow Y_1 \rightarrow Y_2$, with a QTL, $Q$, at 50cM. The blue and red curves show the (unconditional) LOD profiles of phenotypes $Y_1$ and $Y_2$, respectively. The black curve depicts the joint LOD curve, and the peak QTL position $\lambda$ is given by the black vertical line. Instead of having to perform an arbitrary choice between the QTLs given by the red and blue vertical lines we use the QTL given by the black line. The dashed line shows the QTL mapping threshold.
We performed two separate simulation studies generating data from the models in Figure 5 in the main text. In model $F$, $Y_1$ plays the role of a master regulator cis trait, and all other traits map in trans to QTL hotspot QTL $Q$ because of the causal effect of $Y_1$. In model $G$, $Y_1$ plays the role of a cis trait mapping to a QTL closely linked to $Q$, and, therefore, causally independent of the trans traits in the hotspot.

In each simulation study we generated 1,000 distinct backcrosses with genetic data composed of 3 chromosomes of length 100cM containing 101 markers per chromosome, and phenotypic data on 5,001 traits on 112 individuals. We simulated unequally spaced markers for model $F$, but equally spaced markers for $G$, with $Q_1$ and $Q$ set 1cM apart. The additive and dominance genetic effects of $Q$ on $Y_1$ were sampled, respectively, from the $U[0; 5; 1]$ and $U[0; 0; 5]$ distributions. Residual error rates were sampled from $U[0; 5; 1; 5]$, and the coefficients of the regressions of $Y_k$ on $Y_1$ were sampled from $U[0; 5; 1]$. Figure SI.3 shows the overall $R^2$ distributions. QTL mapping was performed as in the pilot study, but here we used the QTL for trait $Y_1$ as a causal anchor.

For each simulated data set we tested $Y_1$ against all other phenotypes $Y_k$, $k = 2, \ldots, 5001$, that share the QTL with $Y_1$, so that the number of hypothesis tests varied from simulation to simulation. Figure SI.4 shows the distribution of the number of tests per simulation study. In total we performed 1,656,261 tests for the simulations with model $F$, and 1,286,243 tests for the simulations with model $G$.

The empirical FDR (that corresponds to one minus the precision) was computed as the ratio of the number of FPs by the sum of the number of FPs and TPs across all tests. The empirical power was computed as before. For model $F$, a FP is defined as any
statistically significant $M_2$, $M_3$, or $M_4$ call, and a TP is given by a significant $M_1$ call. For model $G$, on the other hand, a FP corresponds to any statistically significant $M_1$, $M_2$, or $M_4$ call, and a TP is given by a significant $M_3$ call. For the evaluations without multiple testing correction, a call $M_k$ was statistically significant if the respective p-value, $p_k$, was smaller than a fixed significance level $\alpha$.

Multiple testing correction procedures based on the control of family wise error rates tend to be very conservative, and are not generally advisable (Benjamini and Hochberg 1995). Here, we investigate the performances of the Benjamini and Hochberg (1995) and Benjamini and Yekutieli (2001) FDR control procedures (denoted, respectively, by BH and BY for now on). The BH and BY adjusted p-values were computed based on the p-values across all simulations pooled together, separately by model call (e.g., for the model $F$ simulations, we pool together all 1,656,261 $M_1$ p-values and apply the BH adjusted for this set of p-values, and similarly for the $M_2$, $M_3$ and $M_4$ p-values), and then compute the FDR and power empirical estimates using the adjusted p-values.
Figure SI.2 Overall distribution of the $R^2$ statistics across all simulated models in Figure 2. Panels a and b present the $R^2$ statistics for sample sizes 112 and 1,000, respectively.
Figure S3  Simulation results for Model A in Figure 2 and sample size 112. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For this model, blue dots represent true positives. Red, green and black dots represent false positives for the AIC, BIC and CMST methods. Red and yellow dots represent false positives for the CIT.
Figure S4  Simulation results for Model B in Figure 2 and sample size 112. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For this model, blue dots represent true positives. Red, green and black dots represent false positives for the AIC, BIC and CMST methods. Red and yellow dots represent false positives for the CIT.
Figure S5  Simulation results for Model C in Figure 2 and sample size 112. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For the AIC, BIC and CMST methods, black dots represent true positives, and blue, red and green dots represent false positives. For the CIT test, yellow dots represent true positives and blue and red dots show false positives.
Figure S6  Simulation results for Model D in Figure 2 and sample size 112. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For the AIC, BIC and CMST methods, green dots represent true positives, and blue, red and black dots represent false positives. For the CIT test, yellow dots represent true positives and blue and red dots show false positives.
Figure S7  Simulation results for Model E in Figure 2 and sample size 112. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For the AIC, BIC and CMST methods, black dots represent true positives, and blue, red and green dots represent false positives. For the CIT test, yellow dots represent true positives and blue and red dots show false positives.
Figure S8  Simulation results for Model A in Figure 2 and sample size 1,000. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For this model, blue dots represent true positives. Red, green and black dots represent false positives for the AIC, BIC and CMST methods. Red and yellow dots represent false positives for the CIT.
Figure S9  Simulation results for Model B in Figure 2 and sample size 1,000. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For this model, blue dots represent true positives. Red, green and black dots represent false positives for the AIC, BIC and CMST methods. Red and yellow dots represent false positives for the CIT.
Figure S10  Simulation results for Model C in Figure 2 and sample size 1,000. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For the AIC, BIC and CMST methods, black dots represent true positives, and blue, red and green dots represent false positives. For the CIT test, yellow dots represent true positives and blue and red dots show false positives.
Figure S11  Simulation results for Model D in Figure 2 and sample size 1,000. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For the AIC, BIC and CMST methods, green dots represent true positives, and blue, red and black dots represent false positives. For the CIT test, yellow dots represent true positives and blue and red dots show false positives.
Figure S12  Simulation results for Model E in Figure 2 and sample size 1,000. Blue, red, green and black dots represent, respectively, $M_1$, $M_2$, $M_3$ and $M_4$ calls. Yellow dots (CIT plot only) represent $M_i$ calls. Grey dots show the “no calls”. Results were computed using significance level 0.05. For the AIC, BIC and CMST methods, black dots represent true positives, and blue, red and green dots represent false positives. For the CIT test, yellow dots represent true positives and blue and red dots show false positives.
Figure S13  Observed FDR and power for the simulations based on model $F$. The x-axis represents the p-value cutoffs used for computing the results. Dashed and full curves represent, respectively, AIC- and BIC-based methods. Green: parametric CMST. Red: non-parametric CMST. Blue: joint-parametric CMST. Black: AIC and BIC. The grey line in the top panels corresponds to the $\alpha$ levels.
Figure SI.3 Overall $R^2$ statistics distributions for the large scale simulation study. The left and right panels show the distribution for the cis-trait and trans-trait, respectively.
Figure SI.4 For each model F and G we performed 1,000 separate simulations, and tested $Y_1$ against all other phenotypes $Y_k$, $k = 2, \ldots, 5001$, that shared the QTL with $Y_1$, at each simulation. The panels show the distribution of the number of tests, i.e, the number of trans-traits that co-mapped to $Y_1$, per simulation study. In total, we performed 1,656,261 tests across the 1,000 simulations with model F, and 1,286,243 tests across the simulations with model G.
Figure S14  Uncorrected p-value distributions for the BIC-based CMST tests with data simulated from model $F$ in Figure 5. Results based on 1,656,261 tests. For these simulations, the $M_1$ call is the correct one, hence the skewed distribution towards small p-values at the left panels. The skewness towards larger p-values for the $M_2$, $M_3$, and $M_4$ calls follows from the fact that whenever a p-value for one model is smaller than $\alpha$, then the p-values for the other three models are greater than $1 - \alpha$. Note the larger frequency of small $M_1$ p-values for the non-parametric CMST test (bottom left panel - the discrete nature of the histogram is a consequence of the test statistic being discrete for the non-parametric test).
Figure S15  Uncorrected p-value distributions for the AIC-based CMST tests with data simulated from model $F$ in Figure 5. Results based on 1,656,261 tests. For these simulations, the $M_1$ call is the correct one, hence the skewed distribution towards small p-values at the left panels. The skewness towards larger p-values for the $M_2$, $M_3$, and $M_4$ calls follows from the fact that whenever a p-value for one model is smaller than $\alpha$, then the p-values for the other three models are greater than $1 - \alpha$. Note the larger frequency of small $M_1$ p-values for the non-parametric CMST test (bottom left panel - the discrete nature of the histogram is a consequence of the test statistic being discrete for the non-parametric test).
Figure S16 Uncorrected p-value distributions for the BIC-based CMST tests with data simulated from model $G$ in Figure 5. Results based on 1,286,243 tests. For these simulations, the $M_3$ call is the correct one, hence the skewed distribution towards small p-values at the $M_3$ panels. The skewness towards larger p-values for the $M_1$, $M_2$, and $M_4$ calls follows from the fact that whenever a p-value for one model is smaller than $\alpha$, then the p-values for the other three models are greater than $1 - \alpha$. Note the larger frequency of small $M_3$ p-values for the non-parametric CMST test (bottom left panel - the discrete nature of the histogram is a consequence of the test statistic being discrete for the non-parametric test).
Figure S17 Uncorrected p-value distributions for the AIC-based CMST tests with data simulated from model G in Figure 5. Results based on 1,286,243 tests. For these simulations, the $M_3$ call is the correct one, hence the skewed distribution towards small p-values at the $M_3$ panels. The skewness towards larger p-values for the $M_1$, $M_2$, and $M_4$ calls follows from the fact that whenever a p-value for one model is smaller than $\alpha$, then the p-values for the other three models are greater than $1 - \alpha$. Note the larger frequency of small $M_3$ p-values for the non-parametric CMST test (bottom left panel - the discrete nature of the histogram is a consequence of the test statistic being discrete for the non-parametric test).