A multivariate regression approach to association analysis of a quantitative trait network

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ABSTRACT

Motivation: Many complex disease syndromes such as asthma consist of a large number of highly related, rather than independent, clinical phenotypes, raising a new technical challenge in identifying genetic variations associated simultaneously with correlated traits. Although a causal genetic variation may influence a group of highly correlated traits jointly, most of the previous association analyses considered each phenotype separately, or combined results from a set of single-phenotype analyses.

Results: We propose a new statistical framework called graph-guided fused lasso to address this issue in a principled way. Our approach represents the dependency structure among the quantitative traits explicitly as a network, and leverages this trait network to encode structured regularizations in a multivariate regression model over the genotypes and traits, so that the genetic markers that jointly influence subgroups of highly correlated traits can be detected with high sensitivity and specificity. While most of the traditional methods examined each phenotype independently, our approach analyzes all of the traits jointly in a single statistical method to find the genetic markers that perturb a subset of correlated traits jointly rather than a single trait. Using simulated datasets based on the HapMap consortium data and an asthma dataset, we compare the performance of our method with the single-marker analysis, and other sparse regression methods that do not use any structural information in the traits. Our results show that there is a significant advantage in detecting the true causal single nucleotide polymorphisms when we incorporate the correlation pattern in traits using our proposed method.

Availability: Software for GFlasso is available at
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1 INTRODUCTION

Recent advances in high-throughput genotyping technologies have significantly reduced the cost and time of genome-wide screening of individual genetic differences over millions of single nucleotide polymorphism (SNP) marker loci, shedding light to an era of ‘personalized genome’ (The International HapMap Consortium, 2005; Wellcome Trust Case Control Consortium, 2005; Malo et al., 2008). However, in modern studies the patient cohorts are routinely surveyed with a large number of traits (from measures of hundreds of clinical phenotypes to genome-wide profiling of thousands of gene expressions), many of which are correlated among them. For example, in Figure 1, the correlation structure of the 53 clinical traits in the asthma dataset collected as a part of the Severe Asthma Research Program (SARP) (Moore et al., 2007) is represented as a network, with each trait as a node, the interaction between two traits as an edge and the thickness of an edge representing the strength of correlation. Within this network, there exist several subnetworks involving a subset of traits, and furthermore, the large subnetwork on the left-hand side of Figure 1 contains two subgroups of densely connected traits with thick edges. In order to understand how genetic variations in asthma patients affect various asthma-related clinical traits in the presence of such a complex correlation pattern among phenotypes, it is necessary to consider all of the traits jointly and take into account their correlation structure in the association analysis. Although numerous research efforts have been devoted to studying the interaction patterns among many quantitative traits represented as networks (Friedman, 2004; Mehlan et al., 2008) as well as discovering network submodules from such networks (Hu et al., 2005; Segal et al., 2003), this type of network structure has not been exploited in association mapping (Cheung et al., 2005; Stranger et al., 2005). Many of the previous approaches examined one phenotype at a time to localize the SNP markers with a significant association, and combined the results from a set of such single-phenotype association mapping across phenotypes. However, we conjecture that one can detect additional weak associations and at the same time reduce false signals by combining the information across multiple phenotypes under a single statistical framework.

In QTL mapping studies with pedigree data, a number of approaches have been proposed to detect pleiotropic effect of markers on multiple correlated traits by considering the traits jointly. However, these approaches involve only a weak and indirect form of structural information present in the phenotypes. The methods based on multivariate regression with multiple outcomes (Knott and Haley, 2000; Liu et al., 2007; Xu et al., 2005) were concerned with finding genetic loci that influence all of the phenotypes jointly, rather than explicitly taking into account the complex interaction patterns among the phenotypes. A different approach has been proposed that first applies a principal component analysis (PCA) to find the directions in which phenotypes are the most correlated, and then uses a multivariate regression on the projected phenotypes (Mangin et al., 1998; Weller et al., 1996). The transformation via PCA allows relationships between them. Many statistical approaches have been proposed to address various challenges in identifying genetic locus associated with the phenotype from a large set of markers, with the primary focus on problems involving a univariate trait (Li et al., 2007; Malo et al., 2008). However, in modern studies the patient cohorts are routinely surveyed with a large number of traits (from measures of hundreds of clinical phenotypes to genome-wide profiling of thousands of gene expressions), many of which are correlated among them. For example, in Figure 1, the correlation structure of the 53 clinical traits in the asthma dataset collected as a part of the Severe Asthma Research Program (SARP) (Moore et al., 2007) is represented as a network, with each trait as a node, the interaction between two traits as an edge and the thickness of an edge representing the strength of correlation. Within this network, there exist several subnetworks involving a subset of traits, and furthermore, the large subnetwork on the left-hand side of Figure 1 contains two subgroups of densely connected traits with thick edges. In order to understand how genetic variations in asthma patients affect various asthma-related clinical traits in the presence of such a complex correlation pattern among phenotypes, it is necessary to consider all of the traits jointly and take into account their correlation structure in the association analysis. Although numerous research efforts have been devoted to studying the interaction patterns among many quantitative traits represented as networks (Friedman, 2004; Mehlan et al., 2008) as well as discovering network submodules from such networks (Hu et al., 2005; Segal et al., 2003), this type of network structure has not been exploited in association mapping (Cheung et al., 2005; Stranger et al., 2005). Many of the previous approaches examined one phenotype at a time to localize the SNP markers with a significant association, and combined the results from a set of such single-phenotype association mapping across phenotypes. However, we conjecture that one can detect additional weak associations and at the same time reduce false signals by combining the information across multiple phenotypes under a single statistical framework.
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Illustration of association analysis using phenotype correlation graph for asthma dataset.

In this article, we propose a family of methods, called graph-guided fused lasso (GFlasso), that fully incorporates the quantitative-trait network as an explicit representation for correlation structure without applying additional clustering algorithms to phenotypes. Our methods combine multiple phenotypes in a single statistical framework, and analyze them jointly to identify SNPs perturbing a subset of tightly correlated phenotypes instead of combining results from multiple single-phenotype analyses. The proposed methods leverage a dependency graph defined on multiple quantitative traits such as the graph for the asthma-related traits shown in Figure 1, assuming that such a graph structure is available from preprocessing steps or as prior knowledge from previous studies. It is reasonable to assume that when a subset of phenotypes are highly correlated, the densely connected subgraphs over these correlated traits contain variables that are more likely to be synergistically influenced by the same or heavily overlapping subset(s) of SNPs with similar strength than an arbitrary subset of phenotypes.

The proposed approach is based on a multivariate regression formalism with the $L_1$ penalty, commonly known as lasso, that achieves sparsity in the estimated model by setting many of the regression coefficients for irrelevant markers to 0 (Tibshirani, 1996). This property of lasso makes it a natural approach for genome-wide association analysis, where the marker genotypes are treated as the predictors, the phenotype in question is treated as the response, and the (sparse) set of markers having non-zero regression coefficients are interpreted as the markers truly associated with the phenotype. However, when applied to association mapping with multivariate traits, lasso is equivalent to a single-trait analysis that needs to be repeated over every single trait. In other words, for a collection of traits, each trait would be treated as independent of all other traits, and every trait would be regressed on a common set of marker genotypes via its own lasso (Fig. 2A), ignoring the possible coupling among traits. Our innovations in GFlasso that enable a departure from the baseline lasso for a single trait is that, in addition to the lasso penalty, we employ a 'fusion penalty' that fuses regression coefficients across correlated phenotypes, using either unweighted or weighted connectivity of the phenotype graph as a guide. This additional penalty will encourage sharing of information between correlated phenotypes, since the full network contains much richer information about complex interaction patterns than the clusters of phenotypes.

Another disadvantage of this approach is that it relies on a set of single-phenotype statistical tests and combines the results afterwards in order to determine whether a marker is significantly affecting a subgroup of phenotypes, thus requiring a substantial effort in conducting appropriate multiple hypothesis testing. We believe that an approach that considers markers and all of the phenotypes jointly in a single statistical method has the potential to increase the power of detecting weak associations and reduce susceptibility to noise.
A conventional single-trait association via linear regression model denote an $N$ and each of the $K$ $X \beta$ (GwFlasso) that offers a flexible range of stringency of the regression coefficients under GFlasso. The results on two datasets, one simulated from HapMap SNP markers and the other collected from asthma patients, show that our method outperforms competing algorithms in identifying markers that are associated with a correlated subset of phenotypes.

2 LASSO REGRESSION FOR MULTIPLE INDEPENDENT PHENOTYPES

Let $X$ be an $N \times J$ matrix of genotypes for $N$ individuals and $J$ SNPs, where each element $x_{ij}$ of $X$ is assigned 0, 1 or 2 according to the number of minor alleles at the $j$-th locus of the $i$-th individual. Let $Y$ denote an $N \times K$ matrix of $K$ quantitative trait measurements over the same set of individuals. We use $y_{ik}$ to denote the $k$-th column of $Y$. A conventional single-trait association via linear regression model can be applied to this multi-trait setting by fitting the model to $X$ and each of the $K$ traits $y_k$'s separately:

$$y_k = X\beta_k + \epsilon_k, \quad k = 1, \ldots, K, \quad (1)$$

where $\beta_k$ is a $J$-vector of regression coefficients for the $k$-th trait that can be used in a statistical test to detect SNP markers with significant association, and $\epsilon_k$ is a vector of $N$ independent error terms with mean 0 and a constant variance. We center each column of $X$ and $Y$ such that $\sum x_{ik} = 0$ and $\sum y_{ik} = 0$, and consider the model in Equation (1) without an intercept. We obtain the estimates of $B = [\beta_1, \ldots, \beta_K]$ by minimizing the residual sum of squares:

$$B = \arg\min_k \sum_i (y_{ik} - X\beta_k)^T(y_{ik} - X\beta_k). \quad (2)$$

In a typical genome-wide association mapping, one examines a large number of marker loci with the goal of identifying the region associated with the phenotypes and markers in that region. A straightforward application of the linear regression method in Equation (2) to association mapping with large $J$ can cause several problems such as an unstable estimate of regression coefficients and a poor interpretability due to many irrelevant markers with non-zero regression coefficients. Sparse regression methods such as forward stepwise selection (Weisberg, 1980), ridge regression (Hoerl et al., 1975; Malo et al., 2008) and lasso (Tibshirani, 1996) that select a subset of markers with true association have been proposed to handle the situation with large $J$. Forward stepwise selection method iteratively selects one relevant marker at a time while trying to improve the model fit based on Equation (2), but it may not produce an optimal solution because of the greedy nature of the algorithm. Ridge regression has an advantage of performing the selection in a continuous space by penalizing the residual sum of square in Equation (2) with the $L_2$ norm of $\beta_k$'s and shrinking the regression coefficients toward 0, but it does not set the regression coefficients of irrelevant markers to exactly 0. We use lasso that penalizes the residual sum of square with the $L_1$ norm of regression coefficients and has the property of setting regression coefficients with weak association markers exactly to 0, thus offering the advantages of both forward stepwise selection and ridge regression. The lasso estimate of the regression coefficients can be obtained by solving the following:

$$\hat{B}^{\text{lasso}} = \arg\min_k \sum_i (y_{ik} - X\hat{\beta}_k)^T(y_{ik} - X\hat{\beta}_k) + \lambda \sum_j |\hat{\beta}_{kj}|, \quad (3)$$

where $\lambda$ is a regularization parameter that controls the amount of sparsity in the estimated regression coefficients. Setting $\lambda$ to a large value increases the amount of penalization, setting more regression coefficients to 0. Many fast algorithms are available for solving Equation (3) (Efron et al., 2004; Tibshirani, 1996).

Lasso for multiple-trait association mapping in Equation (3) is equivalent to solving a set of $K$ independent regressions for each trait with its own $L_1$ penalty, and does not provide a mechanism to combine information across multiple traits such that the estimates reflect the potential relatedness in the regression coefficients for those correlated traits that are influenced by common SNPs. However, several traits are often highly correlated such as in gene expression of co-regulated genes in eQTL study, and there might be genotype markers that are jointly associated with those correlated traits. Below, we extend the standard lasso and propose a new penalized regression method for detecting markers with pleiotropic effect on correlated quantitative traits.

3 GFLASSO FOR MULTIPLE CORRELATED PHENOTYPES

In order to identify markers that are predictive of multiple phenotypes jointly, we represent the correlation structure over the set of $K$ traits as an edge-weighted graph, and use this graph to guide the estimation process of the regression coefficients within the lasso framework. We assume that we have available a preprocessing step a phenotype correlation graph $G$ consisting of a set of nodes $V$, each representing one of the $K$ traits and a set of edges $E$. In this article, we adopt a simple and commonly used approach for learning such graphs, where we first compute pairwise Pearson correlation coefficients for all pairs of phenotypes using $y_k$'s, and then connect two nodes with an edge if their correlation coefficient is above the given threshold $\rho$. We set the weight of each edge $(m, l) \in E$ to the absolute value of correlation coefficient $|r_{ml}|$, so that the edge weight represents the strength of correlation between the two nodes. This thresholded correlation graph is also known as a relevance network, and has been widely used as a representation of gene interaction networks (Butte et al., 2000; Carter et al., 2004). It is worth pointing out that the choice of methods for obtaining the phenotype network is not a central issue of our method. Other variations of the standard relevance network have been suggested (Zhang and Horvath, 2005), and any of these graphs can also be used within our proposed regression methods. Below, we first introduce $G_0$Flasso that makes use of unweighted graph, and further extend this method to $G_m$Flasso to take into account the full information in the graph including edge weights. Given the correlation graph of phenotypes, it is reasonable to assume that if two traits are highly correlated and connected with an edge in the graph, their variation across individuals might be explained by genetic variations at the same loci, possibly having the same amount of influence on each trait. In $G_0$Flasso, this assumption is expressed as an additional penalty term that fuses two regression...
When used in this setting, GcFlasso looks for associations between traits across the correlated traits given a genotype marker with pleiotropic effect propagates to the regression applied locally to a pair of regression coefficients for neighboring traits. The regression coefficient pair corresponds to the elements of the corresponding matrix B in Equation (4). In GcFlasso, we employ a similar strategy in a multiple-output regression in order to identify pleiotropic effect of a genetic marker and a subgraph of phenotype network rather than a genetic marker and a single phenotype. Unlike other previous approaches for detecting pleiotropic effect that first apply clustering algorithms to learn subgroups of traits and then search for genetic variations that perturb the subgroup, GcFlasso uses the full information on correlation structure in phenotypes available as a graph, where the subgroup information is embedded implicitly within the graph as densely connected subgraphs. Although the fusion penalty in GcFlasso is applied locally to a pair of regression coefficients for neighboring trait pairs in the graph, this fusion effect propagates to the regression coefficients for other traits that are connected to them in the graph. For densely connected nodes, the fusion is effectively applied to all of the members of the subgroup, and the set of non-zero regression coefficients tend to show a block structure with the same values across the correlated traits given a genotype marker with pleiotropic effect on those traits, as we demonstrate in experiments. If the edge connections are sparse within a group of nodes, the corresponding traits are only weakly related and there is little propagation of fusion effect through edges in the subgroup. Thus, GcFlasso incorporates the subgrouping information through the trait correlation graph in a more flexible manner compared to previous approaches.

Now, we present a further generalization of GcFlasso that exploit the full information in the phenotype networks for association mapping. Note that the only structural information used in GcFlasso is the presence or absence of edges between two phenotypes in the graph. GwFlasso is a natural extension of GcFlasso that takes into account the edge weights in graph G in addition to the graph topology. GwFlasso weights each term in the fusion penalty in Equation (4) by the amount of correlation between the two phenotypes being fused, so that the amount of correlation controls the amount of fusion. More generally, GwFlasso weights each term in the fusion constraint in Equation (4) by a monotonically increasing function of the absolute values of correlations, and finds an estimate of the regression coefficients as follows:

\[
\hat{\beta}^{GW} = \arg\min_{\beta} \sum_{x} (y_x - X\beta)^T (y_x - X\beta) + \lambda \sum_{k} \sum_{(m,j) \in E} \sum_{f(r_m)} \sum_{j} \left| \beta_{jm} - \gamma \cdot \text{sign}(r_m) \beta_{kj} \right|.
\]

If the two phenotypes m and l are highly correlated in graph G with a relatively large edge weight, the fusion effect increases between these two phenotypes since the difference between the two corresponding regression coefficients \( \beta_{jm} \) and \( \beta_{jl} \) is penalized more than for other pairs of phenotypes with weaker correlation. In this article, we consider \( f_1(r) = r \) for G1wFlasso and \( f_2(r) = r^2 \) for G2wFlasso. We note that the GcFlasso is a special case of GwFlasso with \( f(r) = 1 \).

The optimization problems in Equations (4) and (5) can be formulated as a quadratic programming as described in Appendix A, and there are many publicly available software packages that efficiently solve such quadratic programming problems. The regularization parameters and can be determined by a cross-validation or a validation set.

4 RESULTS

We compare the results from our proposed methods, GcFlasso, G1wFlasso and G2wFlasso, with the ones from the single-marker analysis as well as multivariate regression methods such as ridge regression and lasso that do not use any structural information in the phenotypes. For ridge regression, lasso and our methods, we selected the regularization parameters using a validation set. We used \([-\log(P\text{-value})]\) for the standard single-marker analysis, and the absolute value of regression coefficients \( |\beta_{jm}| \) for the GcFlasso weights each term in the fusion constraint in Equation (4) by a monotonically increasing function of the absolute values of correlations, and finds an estimate of the regression coefficients as follows:

4.1 Simulation study

We simulated genotype data for 250 individuals based on the HapMap data in the region of 8.79–9.20 M in chromosome 7. The
first 60 individuals of the genotype data came from the parents of the HapMap CEU panel. We generated genotypes for additional 190 individuals by randomly mating the original 60 individuals on the CEU panel. We included only those SNPs with minor allele frequency >0.1. Since our primary goal is to measure the performance of the association methods in the case of multiple correlated phenotypes, we sampled 50 SNPs randomly from the 697 SNPs in the region in order to reduce the correlation among SNPs from the linkage disequilibrium.

Given the simulated genotype, we generated the true associations represented as regression coefficients $\beta$ and phenotype data as follows. We assumed that the number of phenotypes is 10, and that there are three groups of correlated phenotypes of size 3, 3 and 4, respectively, so that the phenotypes in each group form a subnetwork in the correlation graph of phenotypes. For simplicity, we assumed that there were no environmental factors or other genetic effects, and that all of the covariance components in the traits came from the relevant SNPs in the given set of SNPs. We randomly selected three SNPs as affecting all of the phenotypes in the first subnetwork, and four SNPs as influencing each of the remaining two subnetworks. We assumed that there is one additional SNP affecting phenotypes in the first two subnetworks, which corresponds to the case of a SNP perturbing a super network consisting of two subnetworks such as the large subnetwork on the left-hand side of Figure 1. In addition, we assumed one additional SNP affecting all of the phenotypes. We set the effect size of all of the true association SNPs to the same value. Once we set the regression coefficients, we generated the phenotype data with noise distributed as $N(0,1)$, using the simulated genotypes as covariates.

We evaluate the performance of the association methods based on two criteria, sensitivity/specificity and phenotype prediction error. The sensitivity and specificity measure whether the given method can successfully detect the true association SNPs with few false positives. The (1-sensitivity) and (1-specificity) are equivalent to type I error rate and (1-type II error rate), and their plot is widely known as a receiver operating characteristic (ROC) curve.

The phenotype prediction error represents how accurately we can predict the values of phenotypes given the genotypes of new individuals, using the regression coefficients estimated from the previously available genotype and phenotype data. We generate additional dataset of 50 individuals, $y_{new}$ and $X_{new}$, and compute the phenotype prediction error as sum of squared differences between the true values $y_{true}$ and predicted values $y_{pred}$ of the phenotypes, $\sum (y_{true} - y_{pred})^2$. For both criteria for measuring performance, we show results averaged over 50 randomly generated datasets.

In the results shown below, for each dataset of size $N$, we fit lasso and the graph-guided methods using ($N-10$) samples, and use the remaining 30 samples as a validation set for determining the regularization parameters. Once we determine the regularization parameters, we use the entire dataset of size $N$ to estimate the final regression coefficients given the selected regularization parameters.

We apply the various association methods to datasets with varying sample sizes, and show the ROC curves in Figure 3. We used the threshold $\rho = 0.3$ to obtain the phenotype correlation graph and set the effect size to 0.5. The results confirm that lasso is an effective method for detecting true causal SNPs and is affected less by the irrelevant SNPs, compared to the simple-marker analysis, ridge regression and CCA-based regression. When we use the weighted fusion penalty in addition to the lasso penalty as in $G_cFlasso$, $G_1wFlasso$ and $G_2wFlasso$, the performance significantly improves over lasso across all of the samples sizes shown in Figure 3.

In order to see how the effect size affects the performance of the methods for association analysis, we vary the effect size and show the ROC curves in Figure 4, for the threshold $\rho = 0.1$ of the phenotype correlation network and sample size $N = 100$. $G_1Flasso$ and $G_2Flasso$ outperform all of the other methods across all of the effect sizes. Because of the relatively low value of the threshold $\rho = 0.1$, the correlation phenotype contains many edges between a pair of phenotypes that are only weakly correlated. Thus, $G_1Flasso$ that does not distinguish edges for strong correlation from those for weak correlation does not show a consistent performance across different effect size, performing better than lasso at effect size 0.3 but worse than lasso at effect size 1.0. $G_1wFlasso$ and $G_2wFlasso$ have the flexibility to handle different strengths of correlation in the graph, and consistently outperform $G_1Flasso$ as well as the methods that do not consider the structural information in the phenotypes.

In order to examine the effect of the threshold $\rho$ for the phenotype correlation graph on the performance of our methods, we evaluate the $G_1$Flasso methods with $\rho$ at 0.1, 0.3, 0.5 and 0.7, and show the ROC curves in Figure 5. We include the ROC curves for the single-marker analysis, the ridge regression, lasso and the CCA-based method that do not use the thresholded phenotype correlation graph
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Fig. 4. ROC curves for comparison of association analysis methods with varying effect size. Effect size is (A) 0.3; (B) 0.5; (C) 0.8; and (D) 1.0. The sample size is 100, and the threshold $\rho$ for the phenotype correlation graph is 0.1.

Fig. 5. ROC curves for comparison of association analysis methods with different values of threshold ($\rho$) for the phenotype correlation network. (A) $\rho = 0.1$; (B) $\rho = 0.3$; (C) $\rho = 0.5$; and (D) $\rho = 0.7$. The sample size is 100, and the effect size is 0.8.

in each panel of Figure 5 repeatedly for the ease of comparison. We use the sample size $N = 100$ and the effect size 0.8. Regardless of the threshold $\rho$, $G_{c}^{1}$Flasso and $G_{c}^{2}$Flasso outperform all of the other methods or perform at least as well as lasso. As we have seen in Figure 4, $G_{c}$Flasso does not have the flexibility of accommodating edges of varying correlation strength in the phenotype correlation graph, and this negatively affects the performance of $G_{c}$Flasso at the low threshold $\rho = 0.1$ in Figure 5A. As we increase the threshold $\rho_S$ in Figure 5B and C, the phenotype correlation graph include only those edges with significant correlations. Thus, the performance of $G_{c}$Flasso approaches that of $G_{c}^{1}$Flasso and $G_{c}^{2}$Flasso, and the curves of the three methods in the GFlasso family almost entirely overlap. When the threshold is relatively high at $\rho = 0.7$, the number of edges in the graph is close to 0, effectively removing the fusion penalty. As a result, the performance of the graph-guided methods becomes close to lasso. Overall, taking into account the correlation structure in phenotypes improves the detection rate of true causal SNPs. Once the phenotype correlation graph includes the edges that capture strong correlations, including more edges by further lowering the threshold $\rho$ does not significantly affect the performance of $G_{c}^{1}$Flasso and $G_{c}^{2}$Flasso. The same tendency is shown in the prediction errors in Figure 6.

We show an example of a simulated dataset and the estimated association strength in Figure 7, using the sample size $N = 100$ and effect size 0.8. Although lasso is more successful in setting the regression coefficients of irrelevant SNPs to 0 than the ridge regression and the CCA-based method, it still finds many SNPs as having a non-zero association strength. $G_{c}$Flasso, $G_{c}^{1}$Flasso and $G_{c}^{2}$Flasso remove most of those spurious SNPs, and shows a clear block structure in the estimated regression coefficients, with each causal SNP spanning subgroups of correlated phenotypes. Since $G_{c}$Flasso uses only the information on the presence or absence of edges, when edges of weak correlation connect nodes across two true subgraphs, $G_{c}$Flasso is unable to ignore the weak edges and fuses the effect of SNPs on the phenotypes across those two subgraphs. This undesirable property of $G_{c}$Flasso disappears when we incorporate the edge weights in $G_{c}^{1}$Flasso and $G_{c}^{2}$Flasso.

We show the computation time for solving a single optimization problem for lasso, $G_{c}$Flasso and $G_{w}$Flasso in Figure 8 for varying number of SNPs and phenotypes.

4.2 Case study using asthma dataset

We apply our methods to data collected from 543 asthma patients as a part of the Severe Asthma Research Program (SARP). The genotype data were obtained for 34 SNPs within or near IL-4R gene that spans a 40 kb region on chromosome 16. This gene has been previously shown to be implicated in severe asthma (Wenzel et al., 2007). We used the publicly available software PHASE (Li and Stephens, 2003) to impute missing alleles and phase the genotypes. The phenotype data included 53 clinical traits related to severe asthma such as age of onset, family history and severity of various symptoms. The phenotype correlation graph thresholded at 0.7 as
Our goal is to examine whether any of the SNPs in the IL-4R gene are associated with a subnetwork of correlated traits rather than an individual trait. We standardized measurements for each phenotype to have mean 0 and SD 1 so that their values are roughly in the same range across phenotypes.

Figure 9A shows the correlation matrix of the phenotypes after reordering the phenotypes using the agglomerative hierarchical clustering so that highly correlated phenotypes are clustered with a block structure along the diagonal. Using threshold $\rho = 0.7$, we obtain a phenotype correlation graph as shown in Figure 9B, where the white pixel at position $(i, j)$ indicates that the $i$-th and $j$-th phenotypes are connected with an edge in the graph. The graph shows several blocks of white pixels representing densely connected subgraphs. We show the full graph in Figure 1. We present results for the single-marker regression analysis, ridge regression, lasso, $G_1 wFlasso$, and $G_2 wFlasso$ in Figure 9C–H, respectively, where the rows represent phenotypes, and the columns correspond to genotypes, with bright pixels indicating high strength of association. The phenotypes in rows are rearranged according to the ordering given by the agglomerative hierarchical clustering so that each row in Figure 9C–H is aligned with the phenotypes in the correlation matrix in Figure 9A. In the fusion penalty in our proposed methods, we use the edges in Figure 9B obtained at threshold $\rho = 0.7$. The graph obtained at threshold $\rho = 0.7$ seems to capture the previously known dependencies among the clinical traits such as subnetworks corresponding to lung physiology and quality of life. We select the regularization parameters in lasso, $G_1 wFlasso$, $G_2 wFlasso$, and $G_2 Flasso$ using a 5-fold cross validation.

As shown in Figures 9C and E, both the single-marker regression analysis and lasso find a SNP near the top row, known as Q551R, as significantly associated with a block of correlated phenotypes. This subset of traits corresponds to the bottom subnetwork (consisting of baselineFEV1, PredFEFPred, AvgNO, BMI, PostbroPred, BaseFEVPer, PredrugFEVIP, MaxFEVIP, FEV1Diff and PostFEF) that resides within the large subnetwork on the left-hand side of Figure 1, and represents traits related to lung physiology. This Q551R SNP has been previously found associated with severe asthma and its traits for lung physiology (Wenzel et al., 2007), and our results confirm this previous finding. In addition, the results from the single-marker analysis in Figure 9C show that on the downstream of this SNP, there is a set of adjacent SNPs that appears to be in linkage disequilibrium with this SNP and at the same time has generally a high level of association with the
Table 1. Summary of results for the association analysis of the asthma dataset

<table>
<thead>
<tr>
<th>ρ</th>
<th>Number of edges</th>
<th>Number of non-zero βjm's</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lasso</td>
<td>G1Flasso</td>
</tr>
<tr>
<td>0.3</td>
<td>421</td>
<td>125</td>
</tr>
<tr>
<td>0.5</td>
<td>165</td>
<td>125</td>
</tr>
<tr>
<td>0.7</td>
<td>71</td>
<td>125</td>
</tr>
<tr>
<td>0.9</td>
<td>11</td>
<td>125</td>
</tr>
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</table>

same subset of phenotypes. On the other hand, lasso in Figure 9E sets most of the regression coefficients for this block of SNPs in linkage disequilibrium with Q551R to 0 identifying a single SNP as significant. This confirms that lasso is an effective method for finding sparse estimates of the regression coefficients, ignoring most of the irrelevant markers by setting corresponding regression coefficients to 0. The ridge regression as shown in Figure 9D does not have the same property of encouraging sparsity as lasso. In fact, in statistical literature, it is well-known that the ridge regression performs poorly in problems that require a selection of a small number of markers affecting phenotypes.

Since our methods in the GFlasso family include the lasso penalty, the results from G1Flasso, G2Flasso and G3Flasso show the same property of sparsity as lasso in their estimates, as can be seen in Figure 9F–H. In addition, because of the fusion penalty, the regression coefficients estimated by our methods form a block structure, where the regression coefficients for each SNP are set to the same value within each block. Thus, each horizontal bar indicates a SNP influencing a correlated block of phenotypes. It is clear that the horizontal bars in Figure 9F–H are generally aligned with the blocks of highly correlated phenotypes in Figure 9A. This block structure is much weaker in the results from lasso in Figure 9E. For example, Figure 9F–H show that the SNPs rs3024660 and rs3024622 on the downstream of Q551R are associated with the same block of traits as Q551R, generating an interesting new hypothesis that these two SNPs as well as Q551R might be jointly associated with the same subset of clinical traits. These two SNPs were only in a weak linkage disequilibrium with SNP Q551R (r² = 0.48 and 0.012, respectively). This block structure shared by the two SNPs is not obvious in the number of edges are included in the phenotype correlation graph in Table 1. When the threshold is high at the threshold for the correlation graph, and summarize the results of single-marker tests and lasso.

5 DISCUSSION

In this article, we proposed a new family of regression methods called GFlasso that directly incorporates the correlation structure represented as a graph and uses this information to guide the estimation process. Often, we are interested in detecting genetic variations that perturb a sub-module of phenotypes rather than a single phenotype, and GFlasso achieves this through fusion penalty in addition to the lasso penalty that encourages parsimony in the estimated model. Using simulated and asthma datasets, we demonstrated that including richer information on phenotype structure as in G1Flasso and G2Flasso improves the accuracy in detecting true associations.

One of the possible directions for future research is to explore the use of more sophisticated graph-learning algorithms such as graphical Gaussian models within the GFlasso framework instead of simply using a thresholded correlation graph. Another possible extension is to learn the graph structure and the regression coefficients jointly by combining GFlasso with graphical lasso (Friedman et al., 2008) that learns sparse covariance matrix for phenotypes. Also, in this work, we considered only continuous-valued traits, but the method can be extended to include logistic regression for discrete-valued traits. Scalability of the method to a large dataset is another important issue that needs to be addressed in the future research. Since the problem is formulated as a convex optimization, we can directly benefit from advances in convex optimization research to speed up the process of parameter estimation.

Finally, in addition to the structural information in the phenotype, we would like to incorporate the structure in the genome such as linkage disequilibrium structure in order to identify a block of correlated markers influencing a set of correlated phenotypes.

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REFERENCES


APPENDIX A

A1. PARAMETER ESTIMATION

In this section, we describe the procedure for obtaining estimates of the regression coefficients in \( G_k \text{Flasso} \). Since \( G_k \text{Flasso} \) is a special case of \( G_W \text{Flasso} \) with \( f(r) = 1 \), the same procedure can be applied to \( G_k \text{Flasso} \) in a straightforward manner. The optimization problem in Equation (5) is the Lagrangian form of the following optimization problem:

\[
G_k \text{Flasso} : \min_{\beta} \sum_{k} \left( y_k - X_k \beta_k \right)^T (y_k - X_k \beta_k) \tag{6}
\]

subject to

\[
\sum_{j} |\beta_{kj}| \leq \lambda_1 \text{ and } \sum_{j} |\beta_{kj}| \leq \lambda_2 \text{, for } k=1,\ldots,K.
\]

where \( \lambda_1 \) and \( \lambda_2 \) are tuning parameters corresponding to \( \lambda \) and \( \gamma \) in Equation (5).

Since the objective function and constraints in Equation (6) are convex, we can formulate this problem as a quadratic programming (QP) as follows. Let \( \beta_k \) denote a \( (K \times 1) \)-vector that can be obtained by concatenating \( \beta_k^T \)'s such that \( \beta_k = (\beta_k^T, \ldots, \beta_k^T, 0, \ldots, 0)^T \). We represent \( \beta_k^+ = (\beta_k^+)^T \) and \( \beta_k^- = (\beta_k^-)^T \), where \( \beta_k^+ \geq 0 \) and \( \beta_k^- \geq 0 \), and let \( \beta_k^+ \) and \( \beta_k^- \) denote \( (K \times 1) \)-vectors of \( \beta_k^+ \)'s and \( \beta_k^- \)'s, respectively. We define \( \theta_1(m,l) = \theta_1^m(l) - \theta_1^l(m) \) and \( \theta_2(m,l) = \theta_2^m(l) - \theta_2^l(m) \), and let \( \theta_1^+=\theta_1^+, \theta_2^+=\theta_2^+ \), where \( \theta_1^+ \) and \( \theta_2^+ \) are \( (J \times K) \)-matrices for \( e = m, l \). We define \( \theta_1^- \) and \( \theta_2^- \) similarly. Let \( M \) be a \( (J \times K) \)-matrix, or equivalently a \( |E| \times K \)-matrix of \( I \times J \)-submatrices. Each sub-matrix \( B_{k,E} \) of \( M \) for \( e = 1, \ldots, |E| \) and \( k = 1, \ldots, K \) is an identity matrix if \( e = (m,l) \) and \( k = m \). If \( e = (m, l) \) and \( k = l \), \( B_{k,E} \) is set to a diagonal matrix with 1 along the diagonal. Otherwise, \( B_{k,E} \) is set to a matrix of 0's. Let \( R \) be a \( (J \times |E|) \)-matrix of \( I \times J \)-submatrices with length \( J \). Each sub-vector in \( R \) is set to \( f(r_{(m,l)}) \) for \( (m,l) \), which represents a \( J \times 1 \)-vector of 1's. Then, the QP problem for Equation (6) can be written as

\[
\min_{\beta} \sum_{k} \left( y_k - X_k \beta_k \right)^T (y_k - X_k \beta_k) \text{ subject to } \beta_k \geq 0 \text{ and } \beta_k \leq 0 \text{, for } k=1,\ldots,K.
\]

where \( I \) is a \( (K \times (K \times 1)) \)-identity matrix, and \( 1_{(J \times K)} \) is a \( (J \times K) \)-vector of 1's.