Logistic Bayesian LASSO for detecting association combining family and case-control data

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Abstract
Because of the limited information from the GAW20 samples when only case-control or trio data are considered, we propose eLBL, an extension of the Logistic Bayesian LASSO (least absolute shrinkage and selection operator) methodology so that both types of data can be analyzed jointly in the hope of obtaining an increased statistical power, especially for detecting association between rare haplotypes and complex diseases. The methodology is further extended to account for familial correlation among the case-control individuals and the trios. A 2-step analysis strategy was taken to first perform a genome-wise single single-nucleotide polymorphism (SNP) search using the Monte Carlo pedigree disequilibrium test (MCPDT) to determine interesting regions for the Adult Treatment Panel (ATP) binary trait. Then eLBL was applied to haplotype blocks covering the flagged SNPs in Step 1. Several significantly associated haplotypes were identified; most are in blocks contained in protein coding genes that appear to be relevant for metabolic syndrome. The results are further substantiated with a Type I error study and by an additional analysis using the triglyceride measurements directly as a quantitative trait.

Background
As next-generation sequencing (NGS) technology becomes more accurate and affordable, many recent studies have focused on assessing associations between common complex diseases and single-nucleotide variants (SNVs), paying particular attentions to those that are rare. Various methods have been proposed, but most can only achieve the identification of candidate genes or regions. To narrow the list of potential causal variants, it would be helpful to investigate haplotype blocks formed by single-nucleotide polymorphisms (SNPs) in regions/genes where associations are suggested but may not necessarily be genome-wide significant. Apart from being able to identify biologically relevant variants, haplotype-based methods can be more powerful than SNV-based methods as multilocus genotypes contain more information than single-locus genotypes, especially when causal loci interact in \textit{cis}, leading to disease etiology \cite{1}. If there are rare causal SNVs in a haplotype block, then rare haplotypes can tag such causal variants, a conclusion based on a simulation study \cite{2}. More importantly, rare haplotypes may be obtained from common SNPs, rendering NGS data unnecessary. The power for detecting rare haplotype associations is further enhanced in a family-based study, as rare associated variants are enriched in families afflicted with the diseases compared to population samples of independent cases and controls of the same size. Currently, numerous methods exist, including a class based on Logistic Bayesian LASSO (LBL) for detecting associations of haplotypes, common or rare, using either case control or family-based data \cite{1, 3}.

The GAW20 Real Data Package provides a good opportunity to apply LBL to identify haplotypes that are associated with metabolic syndrome. Specifically, we consider the ATP binary trait derived from the measurements taken at visit 2 (before drug intervention). Among
the 188 pedigrees in the data set, only 17 contain complete case–parent trios (ie, genotype information for both parents and the child and phenotype status for the child are all available), leading to a total of only 25 such trios. In addition, we extracted 283 cases (ATP = 1) and 475 controls (ATP = 0) with available genotype information. Because the number of trios is extremely small, it is clear that there is insufficient power to detect haplotype association using these data alone. However, their inclusion may enhance detection power compared to when only case-control data are used. Because the current LBL methodology focuses on a single study design, we propose eLBL, an extension of the LBL methodology to combine case–control and case–parent trios data for a joint analysis. Furthermore, because cases, controls, and trios are all extracted from the same set of pedigrees, there are intrinsic correlations. To account for such familial dependency, we have adopted a composite likelihood adjustment approach.

**Methods**

**Extension of the logistic Bayesian LASSO accounting for familial dependency**

Suppose we have \( n = n_1 + n_2 \) individuals, where \( n_1 \) and \( n_2 \) are the numbers of cases and controls, respectively. Let \( Y = (Y_1, Y_2, \ldots, Y_n) \), where \( Y_i \) denotes the affection status of the \( i \)th individual, with case = 1 and control = 0.

Let \( Z = (Z_1, Z_2, \ldots, Z_n) \), where \( Z_i \) is the (unobserved haplotype pair) of individual \( i \), while the observed genotype matrix is denoted by \( G = (G_1, G_2, \ldots, G_n) \), where \( G_i \) is the genotype vector (over a set of SNPs) of the \( i \)th individual. We note that \( G \) contains information about \( Z \), but the mapping is typically many (haplotype pairs) to one (vector of genotypes). The complete data (haplotype) likelihood is

\[
L_c(\varnothing) = \prod_{i=1}^{n_1} P(Z_i | Y_i = 1, \varnothing) \prod_{i=n_1+1}^{n} P(Z_i | Y_i = 0, \varnothing)
\]

where the probabilities are specified, as elaborated below, through a logit link function relating the odds of disease to the haplotypes, and \( \varnothing \) is a vector of parameters including haplotype frequencies and coefficients of the logistic regression model.

Suppose we also have \( m \) trios with each ascertained through the offspring. Let \( Y_{ic} = 1 \) denote that the child is a case. The haplotype configuration of the \( i \)th trio can be written as \( Z_i = (Z_{ij}, Z_{im}, Z_{ic}) \), where the 3 components denote the haplotype pair of the father, mother, and the affected child, respectively. Under the assumption of allelic exchangeability, this is equivalent to \( Z_i = (Z_{ia}, Z_{im}) \), where \( Z_{ia} \) is the untransmitted haplotype pair from the parents. Then, the haplotype-based likelihood for case–parent trios is

\[
L_f(\varnothing) = \prod_{i=1}^{m} P(Z_{ic} | Y_{ic} = 1, \varnothing) P(Z_{ia} | Y_{ic} = 0, \varnothing)
\]

where the probabilities and the parameter vector are specified as in the case–control data. Putting eqs. (1) and (2) together, we obtained the following composite likelihood:

\[
L_c(\varnothing) = L_c(\varnothing) \times L_f(\varnothing)
\]

It is apparent from the description of the GAW20 data given above that our data units are not independent, thus the composite likelihood as specified in eq. (3) is not the correct likelihood based on the observed data. However, owing to the complex relationships among the extracted cases, controls, and trios, it is difficult to formulate the correct likelihood. Fortunately, it is possible to obtain correct inferences based on the misspecified composite likelihood \( L_c(\varnothing) \) through appropriate adjustment [4]. Following the “magnitude adjustment” algorithm [5], we denote \( H(\varnothing) = E[\sum_{j=1}^{K} \ell_j(\varnothing) | \varnothing] \), where \( \ell_j(\varnothing) = \log L_j(\varnothing) \) is the log-composite likelihood, and \( V \) and \( V_0 \) are the first-order and second-order derivatives, respectively. Let \( \lambda_1, \lambda_2, \ldots, \lambda_p \) be the eigenvalues of \( H(\varnothing)^{-1}J(\varnothing) \); based on them we form \( k = p / \sum_{i=1}^{p} \lambda_i \). Then the adjusted log-likelihood

\[
\ell^*(\varnothing) = k \ell_c(\varnothing)
\]

is used for inference, as elaborated in the following paragraph.

To specify the probabilities and elaborate on \( \varnothing \), we assume that, for any given individual in the study with haplotype pair \( Z \), we model the odds of the disease \( \theta_Z = P(Y = 1 | Z) / P(Y = 0 | Z) \) with a logistic model \( \log \theta_Z = \alpha + X_Z \beta \), where \( X_Z \) is the design vector corresponding to haplotype pair \( Z \), coded according to the assumed mode of inheritance (eg, additive, recessive, dominant); \( \beta = (\beta_1, \ldots, \beta_K) \) (part of the collection of parameter vector \( \varnothing \)) is the regression coefficient vector with \( \beta_j \) corresponding to the effect of the \( j \)th variant on the log odds; and \( \alpha \) is the baseline effect (related to the phenocopy rate). Note that if we assume an additive model, then the \( j \)th variant is the \( j \)th haplotype, and the total number of distinct haplotypes is \( K + 1 \). We cast the problem into a Bayesian framework, where the adjusted likelihood in eq. (4) is used for correct posterior inference [5]. The detailed Markov chain Monte Carlo (MCMC) inference procedure follows the original LBL methodology using shrinkage priors to increase power for detecting rare haplotypes [1, 3]; the adjustment factor \( k \) is updated in each MCMC iteration. Convergence of the Markov chain is assured based on commonly used diagnostic tools. The posterior odds over the prior odds, namely the Bayes factor (BF), is used to assess the significance of the \( \beta_j \)s. We have also constructed empirical posterior
credible intervals (CIs) for the odds ratios (ORs). For each haplotype, the OR is essentially the exponential of the corresponding β in the logistic model given above. It is estimated, together with the CIs, from the posterior sample of the β values. Decision on the significance of a haplotype is based on both BF (> 2) and CI (not including the null value 1).

A 2-step analysis strategy
Because the proposed eLBL (extension of the Logistic Bayesian LASSO [least absolute shrinkage and selection operator]) methodology is based on an MCMC procedure to sample from the posterior distribution, it is computationally intensive, and thus not suitable for whole-genome scan. Instead, we adopt the following 2-step strategy. In the first step, we use Monte Carlo pedigree disequilibrium test (MCPDT) [6], a family-based single-SNP association testing method, to scan 654,767 SNPs across the 22 autosomes. We excluded SNPs with low minor allele frequencies (< 1%). MCPDT imputes missing data and takes familial relationships into account; consequently, it is viewed as using all information to the maximum extent possible. In the second step, we formed haplotype blocks around the SNPs selected from Step 1 using haploview [7]. We then applied eLBL to identify haplotype(s) within each block that have a significant influence on the ATP binary trait.

Results
Of the 654,767 SNPs considered, we selected the 10 SNPs with the smallest MCPDT p values for further analysis with eLBL. These 10 SNPs have p values close to \(10^{-4}\) or smaller (Table 1), with the 3 SNPs on chromosome 1 passing the threshold of genome-wide significance at the 5% level after Bonferroni correction. To increase the statistical power for detecting rare variants and to potentially understand the causal mechanism in downstream analysis, we applied eLBL to 9 haplotype blocks that cover the SNPs displayed in Table 1. Note that the first 2 SNPs on chromosome 1 belong to the same haplotype block (block 1; Table 1). The remaining 8 SNPs were placed into 8 separate blocks labeled as blocks 2 to 9 (Table 1).

The results from eLBL, presented in Table 2, show that a number of haplotypes have an estimated CI of OR not including 1 and a BF > 2, providing significant evidence for association between the identified haplotypes and the ATP trait. For instance, haplotype h11101 of block 1, which contains the minor alleles of SNPs rs10915052 and rs1406862 (occupying the first and fourth positions, in bold in Table 2), is seen to have a fairly significant evidence of association with a BF of 15. This haplotype contains the 2 minor alleles strongly suggests that the 2 SNPs may very well interact in cis and play a regulatory role for metabolic syndrome, as the block is not located within the coding region of a gene. As another example, 2 haplotypes in block 7, located within the protein coding gene STARD13, are inferred to be associated with ATP. This haplotype block spans 10 SNPs, with SNP rs8001893 sitting at the last position. For the haplotype containing the minor allele, h1111111111, its effect is protective (OR < 1), which is the opposite of the effect of haplotype h1000000000 (OR > 1). Both haplotypes are rare, especially the risk haplotype (frequency < 0.001). Given its rarity and the large variability in the estimate (reflected in the large upper bound of the CI resulting from small frequency and only moderate sample size), care needs to be taken in the interpretation of the effect size. Nevertheless, the associated gene appears to be relevant for the study of metabolic syndrome. Gene Ontology annotations related to STARD13 include guanosine triphosphatase (GTPase) activator activity and lipid binding. Finally, another protein coding gene, ABCCI, which contains haplotype block 8, is also noteworthy, as it also appears to be pertinent to metabolic syndrome. Among its related pathways are vitamin digestion and absorption, and metabolism.

| Table 1 | Top 10 SNPs with the smallest p values as identified by MCPDT |
| --- | --- |
| SNP | Chr | Position | Allele | MAF | p value | Block |
| rs10915052 | 1 | 30,479,266 | G/A | 0.0104 | 2.59 x 10^{-8} | 1 |
| rs1406862 | 1 | 30,483,442 | T/C | 0.0109 | 3.48 x 10^{-8} | 1 |
| rs16833496 | 1 | 30,509,960 | G/A | 0.0110 | 1.80 x 10^{-7} | 2 |
| rs17068604 | 4 | 57,033,788 | T/C | 0.0597 | 1.22 x 10^{-4} | 3 |
| rs2048091 | 8 | 19,003,646 | A/C | 0.3565 | 5.92 x 10^{-5} | 4 |
| rs12281650 | 11 | 69,360,392 | C/T | 0.2665 | 9.48 x 10^{-5} | 5 |
| rs7943255 | 11 | 124,450,756 | G/A | 0.0544 | 1.23 x 10^{-4} | 6 |
| rs8001893 | 13 | 33,120,261 | C/A | 0.0171 | 6.24 x 10^{-6} | 7 |
| rs35625 | 16 | 16,077,067 | T/C | 0.4123 | 1.15 x 10^{-4} | 8 |
| rs966287 | 18 | 27,580,619 | G/C | 0.3285 | 8.95 x 10^{-5} | 9 |

MAF, minor allele frequency
*Minor allele is listed after the slash

Discussion and conclusions
Motivated by making maximum use of information resulting from the limited sample sizes when only case-control or trio data are considered, we propose eLBL, an extension of the LBL methodology, so that both types of data can be analyzed jointly to increase statistical power. This new approach is further extended to adjust for familial correlations, leading to correct statistical inference using dependent data. Our 2-step analysis strategy was designed to increase statistical power. Indeed, by using all available information, MCPDT identified 3 genome-wide significant SNPs, which disappear when only observed
The results, as expected, are sensitive to the selection of the independent samples, and miss many of the haplotypes identified in Table 2. Similarly, an analysis of 17 independent trios using famLBL [2] reveals that the sample size is too small to obtain interpretable results. With an increase in power, the natural question is whether there is also an increase in Type I error, as eLBL is a new method and has not been studied thoroughly. To answer this question, we performed a limited simulation study wherein data from a null model was simulated. To mimic the family dependent structure and the linkage disequilibrium structure of the real data, we simulated our data using the GAW20 families and the inferred haplotypes with the estimated frequencies from block 8 to preserve linkage disequilibrium. Our results indicate that there is no elevated Type I error. In the contrary, eLBL is seen to be conservative for rare variants. Nevertheless, for haplotypes with frequencies greater than 0.05, the Type I error is as expected. To further substantiate the results from eLBL, we also analyzed the triglyceride level from visit 2 directly as a quantitative trait using a variation of LBL [1], but also accounting for the familial structures in the data. Of the 4 protein coding genes, SRPT2, SLC37A2, STARD13, and ABCCI, identified by eLBL, the quantitative analysis also identified associated haplotypes in blocks contained within these genes. These results, together with the Type I error study and the annotations of the genes, affirm the results from eLBL, leading to our conjecture that the haplotypes identified are potentially either involved in the causal mechanism or playing a regulatory role in metabolic syndrome.

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Availability of data and materials
The data that support the findings of this study are available from the Genetic Analysis Workshop (GAW), but restrictions apply to the availability of these data, which were used under license for the current study. Qualified researchers may request these data directly from GAW.

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Authors’ contributions
XZ, MW, and HZ implemented the algorithms and performed the data analyses. SL conceived the study and supervised the analyses and interpretation. XZ, MW, and SL wrote the manuscript. WS reviewed the manuscript and contributed to the discussion. All authors have read and approved the final manuscript.

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