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Mapping complex traits using Random Forests
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Abstract

Random Forest is a prediction technique based on growing trees on bootstrap samples of data, in conjunction with a random selection of explanatory variables to define the best split at each node. In the case of a quantitative outcome, the tree predictor takes on a numerical value. We applied Random Forest to the first replicate of the Genetic Analysis Workshop 13 simulated data set, with the sibling pairs as our units of analysis and identity by descent (IBD) at selected loci as our explanatory variables. With the knowledge of the true model, we performed two sets of analyses on three phenotypes: HDL, triglycerides, and glucose. The goal was to approach the mapping of complex traits from a multivariate perspective. The first set of analyses mimics a candidate gene approach with a high proportion of true genes among the predictors while the second set represents a genome scan analysis using microsatellite markers. Random Forest was able to identify a few of the major genes influencing the phenotypes, such as baseline HDL and triglycerides, but failed to identify the major genes regulating baseline glucose levels.

Background

Trees are nonparametric prediction models. In the context of genetic linkage analysis, Zhang et al. [1] used single trees to identify markers where mean identity-by-descent (IBD) sharing in sib pairs is predictive of the affection status of the pair. Breiman [2] and others have reported that significant improvements in prediction accuracy are achieved by using a collection of trees, called a random forest. We applied Random Forest to the first replicate of the Genetic Analysis Workshop 13 simulated data set, with the sibling pairs as our units of analysis. We used the IBD scores (mean IBD or probability of sharing two alleles IBD) at various loci on the genome to predict the absolute difference between phenotypic values in each pair of sib-
We focused our efforts on baseline levels of the following selection of traits: fasting glucose levels, fasting triglyceride level, and HDL level. Baseline values for each individual were estimated using the first visit measurement for glucose and triglyceride levels, because those traits increase with age. HDL level is stable over time and we used the mean HDL level over all visits to get a more precise estimate. These variables were stratified by ever having smoked and gender and then adjusted for covariates using the SAS [5] software program PROC REG. The covariates used to adjust the final variables were body mass index (BMI) (mean over all visits), baseline grams of alcohol per day, smoking amount (baseline number of cigarettes per day), and baseline age. In adjusting for smoking, both linear and quadratic terms were included. Plots of the adjusted trait values against the covariates showed no remaining trend. We then computed the absolute difference between the values of these adjusted variables for each sibling pair. A fourth phenotype was considered: the first principal component, as calculated by PROC PRINCOMP in SAS, of the absolute sib-pair differences of the adjusted glucose, adjusted triglycerides, and adjusted HDL levels. The first principal component explains 38% of the variance of the three phenotype differences.

A Random Forest is created by growing trees, without pruning, on bootstrap samples of the data, selecting the best split at each node among a random selection of the explanatory variables. For quantitative outcomes the forest is made of regression trees, where the tree predictor is the mean value of the training set observations in each terminal leaf. The Random Forest predictor is computed by averaging the tree predictors over trees for which the given observation was "out-of-the bag", i.e., not included in the bootstrap sample used to build the tree. The mean squared generalization error or predictive error (PE) is used to assess the predictive accuracy. Because the prediction for an observation is based on trees grown without the observation, an idea akin to cross-validation, the estimated PEs are unbiased. The importance of a variable is quantified by calculating the PE increase resulting from randomly permuting the values of a particular variable among the out-of-bag observations for each tree. Shuffling the values of variables of high importance should increase the PE, while the PE should not be affected by permuting values of variables of low importance. Random Forest regression was performed using the program RRFOREST [6] with the following parameter settings: the number of variables selected at each node was set to a value where empirical evidence indicated that the importance index estimates had converged. That value was 5000 in the candidate gene analysis and 10,000 in the genome scan.

The set of candidate genes consisted of all quantitative trait loci (QTLs) affecting clinical phenotypes (b12 to b38 and s3 to s12) and four random locations on each of chromosomes 2, 6, 10, and 16, which contain no QTL [7]. The 399 marker loci were included in the genome-scan analysis. Since IBD is measured at marker loci instead of directly at the QTLs in the genome-scan analysis, a definition of a region surrounding marker loci with high importance index values is needed to determine whether an important marker locus is close enough to a QTL to be called a true positive. We defined an importance peak interval as a region surrounding one or more consecutive markers with importance greater than 0.05%. Each peak interval originates at a marker that is a local maximum of the importance index and extends to the left and right over 10 cM or to the first marker with importance lower than 0.05%, whichever is furthest away. Overlapping peak intervals were collapsed into a single peak interval.

We combined the results for the separate mean IBD and $Z_2$ analyses by ranking the peak intervals from the two analyses in decreasing order of the maximum importance reached within the interval, counting overlapping peak intervals only once. The success of that combined analysis was evaluated by determining whether each peak interval contained genes influencing the particular trait. Candidate genes were ranked similarly.

**Results**

The importance of the mean IBD at candidate genes for predicting phenotype absolute differences is reported in Figure 1. The same measure at the genome scan markers is reported in Figure 2. For adjusted HDL level difference, genes b12 and b20, which both contribute to the HDL phenotype, have the highest importance in the candidate gene analysis. Mean IBD at the first three markers on chromosome 9, close to b12, at a random location on chromosome 12, and at two locations on chromosome 17 linked to genes b19, b20, b18, and b29 have the highest importance in the genome-scan analysis. Figure 3 shows the importance values when using $Z_2$ and both mean IBD and $Z_2$ (joint analysis) in the genome-scan analysis. While the region around b12 on chromosome 9 is identified by all choices of IBD measures, the chromosome 17 region is common only to the mean IBD and joint analyses. The $Z_2$ analysis points to a chromosome 17 locus near s11 that is not seen in the other two analyses. In general, we observed more consistency between the mean IBD and joint analyses than between those two types of analysis and the $Z_2$ analysis. For adjusted triglyceride level difference, the mean IBD at candidate genes b17 and b14 is the highest. A marker on chromosome 1 linked to b17 is detected in the genome scan analysis, as well as a false-positive signal on chromosome 17. For adjusted glucose level difference, the candidate genes most predictive of the phenotype
were random locations on even numbered chromosomes. Similarly, the locations with the highest importance in the genome scan are unlinked to any susceptibility gene. In the analysis of the first principal component of the adjusted HDL, triglyceride, and glucose level differences, the mean IBD at gene b12 stands out as the most important in both analyses. The second highest importance peak in the genome scan analysis on chromosome 19 does not contain any QTL.

The ranking of the importance peaks from combining the mean IBD and $Z_2^2$ separate analyses in Table 1 shows QTLs among the top hits with both candidate genes and genome scan markers for all phenotypes except glucose level. A few genes influencing a phenotype indirectly were also detected. The candidate gene analysis yielded fewer false positives, as expected.

**Discussion**

Although Random Forest was able to identify several genes in the candidate-gene approach and several chromosomal regions harboring susceptibility genes in the genome-scan approach, the findings were not consistent across different phenotypes. In addition, the ranking of genes identified did not typically follow the ranking of effect sizes used in the simulation. The predictive power was in general very low, with PE in excess of 98% of the variance of the absolute difference. However, that observation does not imply failure in the context of a linkage analysis. By analogy, the Haseman and Elston [8] regression of squared difference against IBD sharing in sibs may give a very significant result at a disease locus while at the same time explain little of the total variance of the regressed variable. A measure of the variability of the importance index would be more relevant to assess the significance of observed importance levels.

Since the selected phenotypes are correlated, some genes also have an indirect influence on phenotypes other than the one they directly affect. Genes affecting phenotypes correlated to the one analyzed do figure among the loci with highest importance in Table 1 and could represent indirect effects. However, we did not expect the genes influencing the slope with age to show elevated importance for baseline levels.

We elected to use the baseline value of the phenotypes considered because a larger number of genes were
Figure 2
Importance of genome scan markers
Importance measured by percent increase in PE. Variables are the mean IBD at the 399 genome scan markers. Only the variables with non-zero importance are shown. Random forests are 10,000 trees. A sample of 100 variables is considered at each split.

Figure 3
Importance of genome scan markers using $Z_2$ and both mean IBD and $Z_2$ for HDL level
Importance of $Z_2$ at the 399 genome scan markers (A). Importance of mean IBD and $Z_2$ at the 399 genome scan markers (798 variables) (B). Only the variables with non-zero importance are shown. Random forests are 10,000 trees. A sample of 100 variables is considered at each split.
affecting baseline value than the slope of the phenotype with age. Since we did not identify any baseline genes for glucose levels in our first round of analyses, we subsequently applied the Random Forest method to the slope of the regression of glucose level against age within individual after some groups reported extremely strong signals near the s3 gene on chromosome 5. We observed importance levels much larger than those reported in Figures 1 and 2 (increases in PE exceeding 5% at b22, a locus tightly linked to s3, and around 3% at s3 itself, but no other candidate above 0.5%, under the candidate gene approach, and exceeding 0.8% at the closest marker to s3 with no other peak reaching 0.3% in the genome scan analysis, using either mean IBD or Z). This confirms that Random Forest can identify a major locus when one is present.

It is encouraging that the ability to detect genes was not limited to the candidate gene setting which had relatively few random locations unlinked to each of the three phenotypes. In the genome-scan approach where we simultaneously considered all genome scan markers, Random Forest was again able to select several genomic regions of interest without substantially increasing the number of false-positive signals among variables with the largest importance, compared to the candidate-gene approach.

Table 1: QTLs at candidate loci and in peak intervals with highest importance

| Rank | Candidate Genes Analysis | Genome Scan Analysis |
|------|--------------------------|----------------------|
|      | HDL Glucose Triglycerides | HDL Glucose Triglycerides |
|      | 1st Principal Component  | 1st Principal Component |
| 1    | b12 FP b17 b12           | b12 FP FP b19,b20     |
| 2    | b20 FP b14 b20           | b19,b20,b18 b30      |
| 3    | b13 FP b12 FP            | b13 b12              |
| 4    | s3 FP b27 FP             | s3 b14              |
| 5    | s4 FP b17 FP             | FP b17              |
| 6    | FP FP b19 FP            | FP b14              |
| 7    | b19 FP b33 FP           | FP b29              |
| 8    | FP b23 b25 FP           | FP b29              |
| 9    | FP b22 FP b13           | FP FP              |
| 10   | s7 FP s7 FP             | b24,b35             |
| 11   | b24 FP FP s7           | FP FP              |
| 12   | FP s4 FP s3           | b20 FP              |

List of genes influencing a trait directly or indirectly through a correlated trait. For peak intervals (genome scan), all genes directly affecting a trait are listed. A Genes influencing only a correlated trait. FP, false positive (for genome scan, peak intervals containing no gene influencing the trait).

References
1. Zhang H, Tsai C-P, Yu C-Y, Bonney G: Tree-based linkage and association analyses of asthma. Genet Epidemiol 2001, 21(suppl):S317-S322.
2. Breiman L: Random forests. Machine Learning 2001, 45:5-32.
3. Mukhopadhyay N, Almasy L, Schroeder M, Mulvihill WP, Weeks DE: Mega2, a data-handling program for facilitating genetic link-

age and association analyses [abstract]. Am J Hum Genet 1999, 65:A436.
4. Kruglyak L, Daly MJ, Reeve-Daly MP, Lander ES: Parametric and nonparametric linkage analysis: a unified multipoint approach. Am J Hum Genet 1996, 58:1347-1363.
5. SAS Institute Inc: SAS Language Reference, Version 6, 1st ed. Cary, NC, SAS Institute Inc. 1990.
6. Random Forests for Regression Version 1.0 [http://www.stat.berkeley.edu/users/breiman/rf.html]
7. Daw EW, Morrison J, Zhou Z, Thomas DC: Genetic Analysis Workshop 13: simulated longitudinal data on families for a system of oligogenic traits. BMC Genetics 2003, 4(suppl 1):S3.
8. Haseman JK, Elston RC: The investigation of linkage between a quantitative trait and a marker locus. Behav Genet 1972, 2:3-19.