We present a novel classification-based algorithm called GeneClass for learning to predict gene regulatory response. Our approach is motivated by the hypothesis that in simple organisms such as *Saccharomyces cerevisiae*, we can learn a decision rule for predicting whether a gene is up- or down-regulated in a particular experiment based on (1) the presence of binding site subsequences (“motifs”) in the gene’s regulatory region and (2) the expression levels of regulators such as transcription factors in the experiment (“parents”). Thus our learning task integrates two qualitatively different data sources: genome-wide cDNA microarray data across multiple perturbation and mutant experiments along with motif profile data from regulatory sequences. Rather than focusing on the regression task of predicting real-valued gene expression measurements, GeneClass performs the classification task of predicting +1 and -1 labels, corresponding to up- and down-regulation beyond the levels of biological and measurement noise in microarray measurements. GeneClass uses the Adaboost learning algorithm with a margin-based generalization of decision trees called alternating decision trees. In computational experiments based on the Gasch *S. cerevisiae* dataset, we show that the GeneClass method predicts up- and down-regulation on held-out experiments with high accuracy. We explore a range of experimental setups related to environmental stress response, and we retrieve important regulators, binding site motifs, and relationships between regulators and binding sites that are known to be associated to specific stress response pathways. Our method thus provides predictive hypotheses, suggests biological experiments, and provides interpretable insight into the structure of genetic regulatory networks.

**Supplementary website:** [http://www.cs.columbia.edu/compbio/geneclass](http://www.cs.columbia.edu/compbio/geneclass)

## I. INTRODUCTION

Understanding the underlying mechanisms of gene transcriptional regulation through analysis of high-throughput genomic data has become an important current research area in computational biology. For simpler model organisms such as *S. cerevisiae*, there have been numerous computational approaches that combine gene expression data from microarray experiments and regulatory sequence data to solve different problems in gene regulation: identification of regulatory elements in non-coding DNA [1,2], discovery of
co-occurrence of regulatory motifs and combinatorial effects of regulatory molecules [13], and organization of genes that appear to be subject to common regulatory control into “regulatory modules” [7, 17]. Most of the recent studies can be placed broadly in one of three categories: statistical approaches, which aim to identify statistically significant regulatory patterns in a dataset [1, 7, 13]; probabilistic approaches, which try to discover structure in the dataset as formalized by probabilistic models (often graphical models or Bayesian networks) [5, 11, 12, 17, 18]; and linear network models, which hope to learn explicit parameterized models for pieces of the regulatory network by fitting to data [2, 20]. While these approaches provide useful exploratory tools that allow the user to generate biological hypotheses about transcriptional regulation, in general, they are not yet adequate for making accurate predictions about which genes will be up- or down-regulated in new or held-out experiments. Since these approaches do not emphasize prediction accuracy, it is difficult to directly compare performance of the different algorithms or decide, based on cross-validation experiments, which approach is most likely to generate plausible biological hypotheses for testing in the lab.

In the current work, we present an algorithm called GeneClass that learns a prediction function for the regulatory response of genes under different experimental conditions. The inputs to our learning algorithm are the gene-specific regulatory sequences – represented by the set of binding site patterns they contain (“motifs”) – and the experiment-specific expression levels of regulators (“parents”). The output is a prediction of the expression state of the regulated gene. Rather than trying to predict a real-valued expression level, we formulate the task as a binary classification problem, that is, we predict only whether the gene is up- or down-regulated. This reduction allows us to exploit modern and effective classification algorithms. GeneClass uses the Adaboost learning algorithm with a margin-based generalization of decision trees called alternating decision trees (ADTs). Boosting, like support vector machines, is a large-margin classification algorithm that performs well for high-dimensional problems. We evaluate the performance of our method by measuring prediction accuracy on held-out microarray experiments, and we achieve very good classification results in this setting. Moreover, we can analyze the learned prediction trees to extract significant features or relationships between features that are associated with accurate generalization rather than just correlations in the training data. In a range of computational experiments for the investigation of environmental stress response in yeast, GeneClass retrieves significant regulators, binding motifs, and motif-regulatory pairs that are known to be associated with specific stress response pathways.

Among recent statistical approaches, the most relevant method related to GeneClass is the REDUCE algorithm of Bussemaker et al. [18] for regulatory element discovery. Given gene expression measurements from a single microarray experiment and the regulatory sequence $S_g$ for each gene $g$ represented on the array, REDUCE proposes a linear model for the dependence of log gene expression $E_g$ on presence of
regulatory subsequences (or “motifs”) \( E_g = C + \sum_{\mu \in S_g} F_{\mu} N_{\mu g} \), where \( N_{\mu g} \) is a count of occurrences of regulatory subsequence \( \mu \) in sequence \( S_g \), and the \( F_{\mu} \) are experiment-specific fit parameters. GeneClass generalizes beyond the conditions of a single experiment by using paired \((\text{motif}_g, \text{parent}_e)\) features, where the parent variable represents over- or under-expression of a regulator (transcription factor, signaling molecule, or protein kinase) in the experiment \( e \), rather than using motif information alone. Note, however, that GeneClass uses classification rather than regression as in REDUCE.

Restriction to a candidate set of potential parents has also been used in the probabilistic model literature, including in the regression-based work of Segal et al. for partitioning target genes into regulatory modules for \textit{S. cerevisiae} [17]. Here, each module is a probabilistic regression tree, where internal nodes of the tree correspond to states of regulators and each leaf node prescribes a normal distribution describing the expression of all the module’s genes given the regulator conditions. The authors provide some validation on new experiments by establishing that the target gene sets of specific modules do have statistically significant overlap with the set of differentially expressed genes; however, they do not focus on making accurate predictions of differential expression as we do here. Our GeneClass method retains the distinction between regulator (“parent”) genes and target (“child”) genes, as well as a model that can capture combinatorial relationships among regulators; however, the margin-based GeneClass trees are very different from probabilistic trees. Unlike in [17], we learn from both expression and sequence data, so that the influence of a regulator is mediated through the presence of a regulatory sequence element. We note that in separate work, Segal et al. [18] present a probabilistic model for combining promoter sequence data and a large amount of expression data to learn transcriptional modules on a genome-wide level in \textit{S. cerevisiae}, but they do not demonstrate how to use this learned model for predictions of regulatory response.

The current work follows up on our original paper introducing the GeneClass algorithm for prediction of regulatory response [10]. Here, we report additional computational experiments and more detailed biological validation for specific environmental stress responses (Section IV.C). Due to space constraints, we omit some algorithmic details and refer the reader to the earlier presentation and to additional results available at the supplementary website: [http://www.cs.columbia.edu/compbio/geneclass](http://www.cs.columbia.edu/compbio/geneclass).

II. LEARNING ALGORITHM

A. Adaboost

The underlying classification algorithm that we use is Adaboost, introduced by Freund and Schapire [15], which works by repeatedly applying a simple learning algorithm, called the weak or base
learner, to different weightings of the same training set. For binary prediction problems, Adaboost learns from a training set that consists of pairs \((x_1, y_1), (x_2, y_2), \ldots, (x_m, y_m)\), where \(x_i\) corresponds to the features of an example and \(y_i \in \{-1, +1\}\) is the binary label to be predicted, and maintains a weighting that assigns a non-negative real value \(w_i\) to each example \((x_i, y_i)\). On iteration \(t\) of the boosting process, the weak learner is applied to the training set with weights \(w_1^t, \ldots, w_m^t\) and produces a prediction rule \(h_t\) that maps \(x\) to \(\{0, 1\}\). The rule \(h_t(x)\) is required to have a small but significant correlation with the labels \(y\) when measured using the current weighting. After the function \(h_t\) is generated, the example weights are changed so that the weak predictions \(h_t(x)\) and the labels \(y\) are decorrelated. The weak learner is then called with the new weights over the training examples and the process repeats. Finally, one takes a linear combination of all the weak prediction rules to obtain a real-valued strong prediction function or prediction score \(F(x)\). The strong prediction rule is given by \(\text{sign}(F(x))\):

\[
F_0(x) \equiv 0 \\
\text{for } t = 1 \ldots T \\
w_i^t = \exp(-y_i F_{t-1}(x_i)) \\
\text{Get } h_t \text{ from weak learner} \\
\alpha_t = \ln \left( \frac{\sum_i: h_t(x_i) = 1, y_i = 1 w_i^t}{\sum_i: h_t(x_i) = 1, y_i = -1 w_i^t} \right) \\
F_{t+1} = F_t + \alpha_t h_t.
\]

One can prove that if the weak rules are all slightly correlated with the label, then the strong rule learned by Adaboost will have a very high correlation with the label – in other words, it will predict the label very accurately. Moreover, one often observes that the test error of the strong rule (percentage of mistakes made on new examples) continues to decrease even after the training error (fraction of mistakes made on the training set) reaches zero. This behavior has been related to the concept of a “margin”, which is simply the value \(yF(x)\). While \(yF(x) > 0\) corresponds to a correct prediction, \(yF(x) > a > 0\) corresponds to a confident correct prediction, and the confidence increases monotonically with \(a\). Our experiments in this paper demonstrate the correlation between large margins and correct predictions on the test set (see Section IV).

B. ADTs for Predicting Regulatory Response

Adaboost is often used with a decision tree learning algorithm as the base learning algorithm. For the problem of predicting regulatory response, we use a form of Adaboost that produces a single tree-based decision rule called an alternating decision tree (ADT). More details on learning ADTs for regulatory response can be found in [10].
Briefly, in our problem setting, we begin with a candidate set of motifs $\mu$ representing known or putative regulatory element sequence patterns and a candidate set of regulators or parents $\pi$. For each (gene,experiment) example in our gene expression dataset, we have two sources of feature information relative to the candidate motifs and candidate parent sets: a vector $N_{\mu g}$ of motif counts of occurrences of patterns $\mu$ in the regulatory sequence of gene $g$, and the vector $\pi_e \in \{-1, 0, 1\}$ of expression states for parent genes $\pi$ in the experiment $e$. The data representation is depicted in Figure 1 (A).

Figure 1 (B) shows a toy example of an ADT that could be produced by Adaboost in our setting. An ADT consists of alternating levels of prediction nodes (ovals) – which contain real-valued contributions to the prediction scores – and splitter nodes (rectangles) – which contain true/false conditions. To obtain the prediction score $F(x)$ for a particular example $x$, we sum the values in all prediction nodes that we can reach along all paths down from the root corresponding to yes/no decisions consisent with $x$.

Splitter nodes in our ADTs depend on decisions based on (motif,parent) pairs. However, instead of splitting on real-valued thresholds of parent expression and integer-valued motif count thresholds, we consider only whether a motif $\mu$ is present or not, and only whether a parent $\pi$ is over-expressed (or under-expressed) in the example. Thus, splitter nodes make boolean decisions based on conditions such as “motif $\mu$ is present and regulator $\pi$ is over-expressed (or under-expressed)”. Paths in the learned ADT correspond to conjunctions (AND operations) of these boolean (motif,parent) conditions. Full details on selection of the candidate motifs and regulators and discretization into up and down states is given in Section III.

FIG. 1: Boosting ADTs for regulatory response prediction. In (A), we show the data presentation for our problem. Every (target gene,experiment) is assigned a label of +1 (up-regulated, in red) or -1 (down-regulated, in green) and represented by the gene’s vector of motif counts (only binary values shown here) and the experiment’s vector of regulator expression states. A toy example of an ADT is shown in (B).

In terms of Adaboost, each prediction node represents a weak prediction rule, and at every boosting iteration, a new splitter node together with its two prediction nodes is introduced. The splitter node can be
attached to any previous prediction node, not only leaf nodes. In general, more important decision rules are
added at early iterations. We use this heuristic to analyze the ADTs and identify the most important factors
in gene regulatory response.

III. METHODS

Dataset: We use the Gasch et al. [4] environmental stress response dataset, consisting of cDNA microarray experiments measuring genomic expression in *S. cerevisiae* in response to diverse environmental transitions. There are a total of 6110 genes and 173 experiments in the dataset, with all measurements given as \( \log_2 \) expression values (fold-change with respect to unstimulated reference expression). We do not perform a zero mean and unit variance normalization over experiments, since we must retain the meaning of the true zero (no fold change).

Motif set: We obtain the 500 bp 5' promoter sequences of all *S. cerevisiae* genes from the Saccharomyces Genome Database (SGD). For each of these sequences, we search for transcription factor (TF) binding sites using the PATCH software licensed by TRANSFAC [19]. The PATCH tool uses a library of known and putative TF binding sites, some of which are represented by position specific scoring matrices and some by consensus patterns, from the TRANSFAC Professional database. A total of 354 binding sites are used after pruning to remove redundant and rare sites.

Parent set: We compile different sets of candidate regulators to study the performance and dependence of our method on the set of regulators. We restrict ourselves to a superset of 475 regulators (consisting of transcription factors, signaling molecules and protein kinases), 466 of which are used in Segal et al. [17] and 9 generic (global) regulators obtained from Lee et al. [9].

Due to computational limitations on the number of (motif,parent) features we could use in training, we select smaller subsets of regulators based on the following selection criteria. We use 13 high-variance regulators that had a standard deviation (in expression over all experiments) above a cutoff of 1.2. The second subset consists of the 9 global regulators that are present in the Lee et al. studies but absent in the candidate list of Segal et al. We also include 30 regulators that are found to be top ranking regulators for the 50 clusters identified in Segal et al. The union of these three lists gives 53 unique regulators.

Target set and label assignment: We discretize expression values of all genes into three levels representing down-regulation (-1), no change (0) and up-regulation (+1) using cutoffs based on the empirical noise distribution around the baseline (0) calculated from the three replicate unstimulated (time=0) heat-shock experiments [4]. We observe that 95% of the samples in this distribution had expression values between +1.3 and −1.3. Thus we use these cutoffs to decide what we define as significantly up-regulated (+1) and
down-regulated (-1) beyond the levels of biological and experimental noise in the microarray measurements.

Note that, although we train only on those (gene,experiment) pairs which discretize to up- or down-regulated states, we can test (make predictions) on every example in a held-out experiment by thresholding on predicted margins. That is, we predict baseline if a prediction has margin below threshold (see Section [IV]).

We reduce our target gene list to a set of 1411 genes which include 469 highly variant genes (standard deviation > 1.2 in expression over all experiments) and 1250 genes that are part of the 17 clusters identified by Gasch et al. [4] using hierarchical clustering (eliminating overlaps).

**Software:** We use the MLJAVA software developed by Freund and Schapire [14] which implements the ADT learning algorithm. We use the text-feature in MLJAVA to take advantage of the sparse motif matrix and use memory more efficiently.

### IV. EXPERIMENTAL RESULTS

#### A. Cross-Validation Experiments

We first perform cross-validation experiments to evaluate classification performance on held-out experiments. We divide the set of 173 microarray experiments into 10 folds, grouping replicate experiments together to avoid bias, and perform 10-fold cross-validation experiments using boosting with ADTs on all 1411 target genes.

We train the ADTs for 400 boosting iterations, during most of which test-loss decreases continuously. We obtain an accuracy of 88.5% on up- and down-regulated examples averaged over 10 folds (test loss of 11.5%), showing that predicting regulatory response is indeed possible in our framework.

To assess the difficulty of the classification task, we also compare to a baseline method, $k$-nearest neighbor classification (kNN), where each test example is classified by a vote of its $k$ nearest neighbors in the training set. For a distance function, we optimize the weighted sum of Euclidean distances for motif and parent vectors, trying values of $k < 20$ and both binary or integer representations of the motif data (see [10]). We obtain minimum test-loss of 31.3% at $k=19$ and with integer motif counts, giving much poorer performance than boosting with ADTs.

Since ADTs output a real-valued prediction score $F(x)$ whose absolute value measures the confidence of the classification, we can predict a baseline label by thresholding on this score, that is, we predict examples to be up- or down-regulated if $F(x) > a$ or $F(x) < -a$ respectively, and to be baseline if $|F(x)| < a$, where $a > 0$. Figure 2(A) shows expression values versus prediction scores for all examples (up, down,
FIG. 2: **True expression values versus prediction scores** $F(x)$. The scatter plot (A) shows a high correlation between prediction scores (x-axis) and true log expression values (y-axis) for genes on held-out experiments. The confusion matrix (B) gives truth and predictions for all genes in the held-out experiments, including those expressed at baseline levels. Examples are binned by assigning a threshold $a = \pm 0.5$ to expression and prediction scores.

and baseline) from the held-out experiments using 10-fold cross-validation. The plot shows a high correlation between expression and prediction, reminiscent of the actual regression task. Assigning thresholds to expression and prediction values binning the examples into up, down and baseline we obtain the confusion matrix in Figure 2 (B).

### B. Extracting features for biological interpretation

We describe below several approaches for extracting important features from the learned ADT model, and we suggest ways to relate these features to the biology of gene regulation.

**Extracting significant features:** We rank motifs, parents and motif-parent pairs by two main methods. The *iteration score (IS)* of a feature is the boosting iteration during which it first appears in the ADT. This ranking scheme appears to be useful in identifying important motifs and motif-parent pairs (restricting to iteration scores $< 50$), since features selected at early rounds tend to be most significant. The *abundance score (AS)* of a regulator in the number of nodes in the final tree that include the regulator as the parent in a motif-parent parent. A regulator with a large abundance score will affect a large number of paths through the ADT and hence affect a large number of target genes. If the state of a regulator is changed through stress response or knockout, its predicted effect on target genes will depend on its abundance in the ADT. Note that presence of a strong motif-parent feature does not necessarily imply a direct binding relationship between parent and motif. Such a pair could represent an indirect regulatory relationship or some other kind of predictive correlation, for example, co-occurrence of the true binding site with the motif corresponding
“In silico” knock-outs: By removing a candidate from the regulator list and retraining the ADT, we can evaluate whether test loss significantly decreases with omission of the parent and identify other weaker regulators that are also correlated with the labels. We investigate in silico knock-outs in the biologically-motivated experiments described in Section IV C.

C. Biological Validation Experiments

We designed the following four training and test sets of selected microarray experiments to study the response to specific types of stress. By comparative analysis of the trees learned from these sets, we find and validate regulators that are associated to regulation programs activated by different stresses. More detailed results can be found on the supplementary website.

Heat-shock and osmolarity stress response: In the first study, we train on heat-shock, osmolarity, heat-shock knockouts, over-expression, amino acid starvation experiments, and we test on stationary phase, simultaneous heat-shock and hypo-osmolarity experiments.

We observe a low test loss of 9.3%, supporting the hypothesis that pathways involved in heat-shock and osmolarity stress appear to be independent and the joint response to both stresses can be predicted easily. This result agrees with the observation by Gasch et al. [4] that these two environmental stresses have nearly additive effects on gene expression of environmental stress response (ESR) genes. The high test accuracy also supports the observation by Gasch et al. [4] that the response as well as parts of the underlying regulatory mechanisms for stationary phase induction (test set) are similar to that of heat-shock (training set).

The top five high scoring parents (based on AS) were USV1, XBP1, PPT1, GIS1 and TPK1. These regulators are known or believed to play specific important roles in each of the training and test set stresses. Segal et al. [17] specifically identify USV1 as an important regulator in stationary phase (test set) and PPT1 to be important in the response to osmolarity stress (training set).

The top ranking motif (based on IS) was the STRE element of MSN2/MSN4, a known regulatory element for a significant number of general stress response genes [4]. The connection of the osmolarity response to the HOG and other MAP kinase pathways is well known. Also, the osmolarity response is strongly related to glycerol metabolism and transport and hence closely associated with gluconeogenesis and glucose metabolism pathways. We find the binding sites of CAT8 (gluconeogenesis), GAL4 (galactose metabolism), MIG1 (glucose metabolism and regulator of osmosensitive glycerol uptake) [8], GCN4 (regulator of HOG pathway and amino acid metabolism), HSF1 (heat-shock factor), CHA4 (amino acid
catabolism), MET31 (methionine biosynthesis) and RAP1 to have high iteration scores; these regulators are all related to the stress conditions in the training set.

FIG. 3: Comparison of expression profiles (173 experiments) of USV1, MSN2, HSF1 and PPT1. The mRNA expression levels of USV1 and PPT1 are informative, with about 50% and 35% of experiments (respectively) showing over 2 fold expression change over wildtype. The expression levels for MSN2 and HSF1 fall mostly in the baseline state, with only about 6% and 5% of experiments (respectively) showing at least 2 fold expression change. While MSN2 and HSF1 are not identified as high scoring parents in the learned trees, their binding sites occur as high scoring motifs.

It is interesting to note that while the presence of binding sites of some very important stress factors like MSN2 and HSF1 (heat shock factor) are identified as significant features (high motif iteration score) in the ADT, the mRNA expression levels of these regulators do not seem to be very predictive. HSF1 does not appear as a parent and MSN2 gets low abundance and iteration scores as a parent, despite their importance as heat-shock and general stress response regulators respectively. Similar results are observed in the modules of [17], where HSF1 is not found in any of the regulation programs and MSN2 is found in 3 of the 50 regulation programs but with low significance. If we compare the expression profiles of HSF1, MSN2, USV1 and PPT1, we find that the mRNA levels of MSN2 and HSF1 have small fluctuations (rarely greater than 2 fold change) and fall mostly within the baseline state, while the expression levels of USV1 or PPT1 show much larger variation over many experiments (see Figure 3). It is known that the activity of MSN2 is regulated by TPK1 (a kinase) via cellular localization. TPK1 is identified as an important parent in the ADT (AS) and is found associated with the MSN2 binding site as a motif-parent pair. Thus in this case, where the activity of a regulator is itself regulated post-transcriptionally, we see a clear advantage of using motif data along with mRNA expression data.

**USV1 “in silico” knockout for heat-shock and osmolarity stress:** Using the same training and test microarrays as in the heat-shock/osmolarity setup, we perform a second study by removing one of the strong
regulators, USV1, from the parent set and retraining the ADT. We get a minor but significant increase in test error from 9.3% to 11%.

TPK1 in the upregulated state along with the MSN2/MSN4 binding site is the top scoring feature (IS). TPK1 is also the top scoring regulator based on abundance.

We also study target genes that change label from correct to incorrect due to the removal of USV1. We reason that since these genes require presence of USV1 in the ADT for correct prediction of their regulatory response, they are highly dependent on USV1 activity and provide good candidates for downstream targets of regulatory pathways involving the knocked out parent. We find that 305 target genes change prediction labels. GO annotation enrichment analysis of these target genes reveal the terms cell wall organization and biogenesis, heat-shock protein activity, galactose, acetyl-CoA and chitin metabolism and tRNA processing and cell-growth. These match many of the terms (namely transcription factors, nuclear transport, cell wall and transport sporulation and cAMP pathway, RNA processing, cell cycle, energy, osmotic stress, protein modification and trafficking, cell differentiation) enriched by analyzing GO annotations of genes that changed significantly in a microarray experiment by [17] with stationary phase induced in a USV1 knockout.

**Peroxide, superoxide stress, and pleiotropic response to diamide:** For the third study, we train on heat-shock, heat-shock knockouts, over-expression, H$_2$O$_2$ wild-type and mutant, menadione, DTT experiments, and we test on diamide experiments. Gasch et al. [4] consider the diamide response to be a composite of responses to the experiments in the training set. We observe a moderate test loss of 16%, suggesting that this pleiotropic response is more complex than the simpler additive responses to heat-shock and osmolarity.

Although USV1, XBP1 and TPK1 are the top three regulators, we see the emergence of an important parent, YAP1. This factor appears to be dominant in the ADTs of only those setups that include redox related stresses, specifically peroxide and superoxide stresses, in the training sets. It is well documented that YAP1 plays a significant regulatory role in oxidation stresses, and this role correlates well with our findings. We hypothesize that USV1 is not very important for response to diamide based on analysis of the fourth setup below, and we attribute its presence in the ADT to the presence of heat-shock experiments in the training set (based on the first setup). We thus simulate a knockout by removing USV1 from the training set and retraining on the training data. Test loss reduces dramatically from 16% to 9.2%, indicating that USV1’s presence in the ADT is detrimental to prediction on diamide response. The ADT for this setup also shows YAP1 associated with its binding motif as an important feature (IS).

**Redox and starvation response:** In this study, we train on DTT and diamide stresses and response to nitrogen depletion and stationary phase induction. We test on diauxic shift and amino acid starvation experiments. We observe a poor test loss of 27.9%. This poor prediction accuracy could mean that regulatory
systems active in experiments in the training set and test set are significantly different. Gasch et al. mention that the starvation responses are quite different from each other and significantly more complex than other stresses (DTT, diamide stress) due to cell-cycle arrest and several secondary effects.

Analysis of the ADT reveals YGL099W (KRE35) as the most abundant regulator. KRE35 also scores among the top 5 candidates in other setups involving redox stresses (such as the third setup above). It could thus be an important regulator for redox related stresses.

We observe that the poor prediction accuracy correlates with the absence of USV1 in the ADT, which is otherwise abundant in the ADTs of all other setups. Since the first three setups show that USV1 is an important regulator for heat-shock response, we add the heat-shock experiments to the training set. As expected, on retraining with this new training set, we get a very significant improvement in prediction accuracy on the same test set (from 27.9% to 16%). This could mean that pathways involved in the heat-shock response are an important component of the more complex response to some starvation responses.

V. DISCUSSION

We have shown that the GeneClass learning algorithm makes accurate predictions of gene regulatory response in yeast over a wide range of experimental conditions. In particular, in experiments related to environmental stress response, examination of the learned GeneClass tree models retrieved important regulators, motifs, and regulator-motif relationships associated with specific stress response pathways. We believe that GeneClass provides a promising new methodology for integrating expression and regulatory sequence data to study transcriptional regulation.

One important next step is to use GeneClass to analyze larger data sets. Since the Gasch dataset that we used here involves only environmental stress response experiments, it is likely that many of the regulatory pathways are not activated and therefore cannot be modeled by analysis of this dataset alone. We hope to extend our studies by incorporating additional and more diverse yeast data sets currently available through resources like NCBI’s Gene Expression Omnibus. At the same time, we plan to make improvements in the computational efficiency of the GeneClass software to allow a significant increase the number of parents so that we can identify the possible roles of many additional regulators. In particular, we plan to use using data structures more appropriate for our pairwise interaction features and weighted sampling to reduce the size of the memory required for holding the training data.

A second potential advance would be a more careful treatment of the raw data. While the ratio data (perturbation/wild type) gives a natural input variable for our analysis, better signal to noise is likely to be achieved by taking into account the expression levels separately. In further work, we plan to use two-color
noise modeling to establish expression-level specific thresholds and thus allow inclusion of more genes whose up- or down-regulated states currently fall within the baseline category. This improvement will allow more training examples and should enable us to accurately predict the response of more subtle target genes.

A third direction for improvement would be to treat parent and child expression levels as continuous (rather than binary) quantities. Similarly, the number of motifs in the regulatory region, rather than merely their presence/absence, could be taken into account. While these refinements could potentially yield more realistic models, it is important that they be represented in a way that is informative for the learning problem and avoids overfitting.

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[1] H. J. Bussemaker, H. Li, and E. D. Siggia. Regulatory element detection using correlation with expression. *Nature Genetics*, 27:167–171, 2001.
[2] P. D’Haeseleer, X. Wen, S. Fuhrman, and R. Somogyi. Linear modeling of mRNA expression levels during CNS development and injury. *Pac. Symp. Biocomp.*, pages 41–52, 1999.
[3] Y. Freund and L. Mason. The alternating decision tree learning algorithm. *Proc. of the Sixteenth International Conf. on Machine Learning*, pages 124–133, 1999.
[4] A. P. Gasch, P. T. Spellman, C. M. Kao, Orna Carmel-Harel, M. B. Eisen, G. Storz, D. Botstein, and P. O. Brown. Genomic expression programs in the response of yeast cells to environmental changes. *Mol. Biol. Cell*, 11:4241–4257, 2000.
[5] A. J. Hartemink, D. K. Gifford, T. S. Jaakkola, and R. A. Young. Using graphical models and genomic expression data to statistically validate models of genetic regulatory networks. *Pac. Symp. Biocomp.*, pages 422–33, 2001.
[6] J. D. Hughes, P. W. Estep, S. Tavazoie, and G. M. Church. Computational identification of cis-regulatory elements associated with groups of functionally related genes in Saccharomyces cerevisiae. *J. Mol. Biol.*, 296(5):1205–14, 2000.
[7] J. Ihmels, G. Friedlander, S. Bergmann, O. Sarig, Y. Ziv, and N. Barkai. Revealing modular organization in the yeast transcriptional network. *Nature Genetics*, 31:370–377, 2002.
[8] C. J. Klein, L. Olsson, and J. Nielsen. Glucose control in Saccharomyces cerevisiae: the role of MIG1 in metabolic functions. *Microbiology*, 144:13–24, 1998.
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[9] T. I. Lee, N. J. Rinaldi, F. Robert, D. T. Odom, Z. Bar-Joseph, G. K. Gerber, N. M. Hannett, C. R. Harbison, C. M. Thompson, I. Simon, J. Zeitlinger, E. G. Jennings, H. L. Murray, D. B. Gordon, B. Ren, J. J. Wyrick, J. Tagne, T. L. Volkert, E. Fraenkel, D. K. Gifford, and R. A. Young. Transcriptional regulatory networks in Saccharomyces cerevisiae. Science, 298:799–804, 2002.

[10] M. Middendorf, A. Kundaje, C. Wiggins, Y. Freund, and C. Leslie. Predicting genetic regulatory response using classification. http://www.cs.columbia.edu/compbio/geneclass, 2004.

[11] D. Pe’er, A. Regev, G. Elidan, and N. Friedman. Inferring subnetworks from perturbed expression profiles. Proc. of the Ninth International Conf. on Intelligent Systems for Molecular Biology, pages 215–224, 2001.

[12] D. Pe’er, V. Regev, and A. Tanay. A fast and robust method to infer and characterize an active regulator set for molecular pathways. Proc. of the Tenth International Conf. on Intelligent Systems for Molecular Biology, pages 258–267, 2002.

[13] Y. Pilpel, P. Sudarsanam, and G. M. Church. Identifying regulatory networks by combinatorial analysis of promoter elements. Nature Genetics, 2:153–159, 2001.

[14] R. E. Schapire and Y. Singer. Boostexter: A boosting-based system for text categorization. Machine Learning, 39(2/3):135–168, 2000.

[15] Robert E. Schapire. The boosting approach to machine learning: An overview. In MSRI Workshop on Nonlinear Estimation and Classification, 2002.

[16] Robert E. Schapire, Yoav Freund, Peter Bartlett, and Wee Sun Lee. Boosting the margin: A new explanation for the effectiveness of voting methods. The Annals of Statistics, 26(5):1651–1686, October 1998.

[17] E. Segal, M. Shapira, A. Regev, D. Pe’er, D. Botstein, D. Koller, and N. Friedman. Module networks: Identifying regulatory modules and their condition specific regulators from gene expression data. Nature Genetics, 34(2):166–176, 2003.

[18] E. Segal, R. Yelensky, and D. Koller. Genome-wide discovery of transcriptional modules from DNA sequence and gene expression. Bioinformatics, 19:273–282, 2003.

[19] E. Wingender, X. Chen, R. Hehl, H. Karas, I. Liebich, V. Matys, T. Meinhardt, M. Prüss, I. Reuter, and F. Schacherer. TRANSFAC: an integrated system for gene expression regulation. Nucleic Acids Research, 28:316–319, 2000.

[20] M. K. Yeung, J. Tegner, and J. J. Collins. Reverse engineering gene networks using singular value decomposition and robust regression. Proc. Natl. Acad. Sci. USA, 99:6163–8, 2002.