

# Revising Qualitative Models of Gene Regulation

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**Abstract.** We present an approach to revising qualitative causal models of gene regulation with DNA microarray data. The method combines search through a space of variable orderings with search through a space of parameters on causal links, with weight decay driving the model toward integer values. We illustrate the technique on a model of photosynthesis regulation and associated microarray data. Experiments with synthetic data that varied distance from the target model, noise, and number of training cases suggest the method is robust with respect to these factors. In closing, we suggest directions for future research and discuss related work on inducing causal regulatory models.

## 1 Introduction and Motivation

Like other sciences, biology requires that its models fit available data. However, as the field moves from a focus on isolated processes to system-level behaviors, developing and evaluating models has become increasingly difficult. This challenge has become especially clear with respect to models of gene regulation, which attempt to explain complex interactions in which the expression levels of some genes influence the expression levels of others. A related challenge concerns a shift in the nature of biological data collection from focused experiments, which involve only a few variables, to cDNA microarrays, which measure thousands of expression levels at the same time.

In this paper, we describe an approach that takes advantage of such nonexperimental data to revise existing models of gene regulation. Our method uses these data, combined with knowledge about the domain, to direct search for a model that better explains the observations. We emphasize qualitative causal accounts because biologists typically cast their regulatory models in this form. We focus on model revision, rather than constructing models from scratch, because biologists often have partial models for the systems they study.

We begin with a brief review of molecular biology and biochemistry, including the central notion of gene regulation, then present an existing regulatory model of photosynthesis. After this, we describe our method for using microarray data to improve such models, which combines ideas from learning in neural networks and the notion of minimum description length. Next we report experimental

studies of the method that draws on both biological and synthetic data, along with the results of these experiments. In closing, we suggest directions for future research and discuss related work on inducing causal models of gene regulation.

## 2 Qualitative Causal Models of Gene Regulation

A gene is a fundamental unit of heredity that determines an organism’s physical traits. It is an ordered sequence of nucleotides in deoxyribonucleic acid (DNA) located at a specific position on a chromosome. Genes encode functional products, called proteins, that determine the structure, function, and regulation of an organism’s cells and tissues.

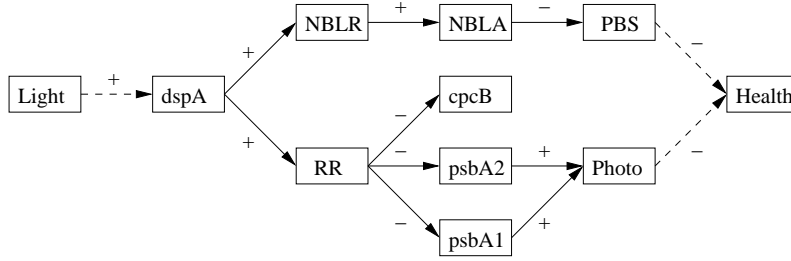
The gene’s nucleotide sequence is used to construct proteins through a multiple stage process. In brief, the enzyme RNA polymerase transcribes each gene into a complementary strand of messenger ribonucleic acid (mRNA) using the DNA as a template. Ribosomes then translate the mRNA into a specific sequence of amino acids forming a protein. Transcription is controlled through the RNA polymerase by transcription factors that let it target specific points on the DNA. The transcription factors may themselves be controlled through signalling cascades that relay signals from cellular or extra-cellular events. Typically, a signalling cascade phosphorylates (or dephosphorylates) a transcription factor, changing its conformation (i.e., physical structure) and its ability to bind to the transcription site. Translation is controlled by many different mechanisms, including repressors binding to mRNA that prevents translation into proteins.

In our work, we focus on revising biological models that relate external cell signals to changes in gene transcription (as measured by mRNA) and, ultimately, phenotype. Specifically, we look at a model of photosynthesis regulation that is intended to explain why *Cyanobacteria* bleaches when exposed to high light conditions and how this protects the organism. This model, shown in Figure 1, was adapted from a model provided by a microbiologist (Grossman et al., 2001).<sup>1</sup> Each node in the model corresponds to an observable or theoretical variable that denotes a measurable stimulus, gene expression level, or physical characteristic. Each link stands for a causal biological process through which one variable influences another. Solid lines in the figure denote internal processes, while dashes indicate processes connected to the environment.

The model states that changes in light level modulate the expression of *dspA*, a protein hypothesized to serve as a sensor. This in turn regulates *NBLR* and *NBLA* expression, which then reduces the number of phycobilisome (PBS) rods that absorb light. The level of PBS is measured photometrically as the organism’s greenness. The reduction in PBS protects the organism’s health by reducing absorption of light, which can be damaging at high levels. The organism’s health under high light conditions can be measured in terms of the culture density.

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<sup>1</sup> The paper describes an initial model for high light response in the *Cyanobacterium Synechococcus*. This model was modified slightly for the *Cyanobacterium* used in our experiments, *Synechocystis* PCC6803, by actions such as replacing *nblS* with its homolog *dspA*.



**Figure 1.** Initial model for photosynthesis regulation of wild type Cyanobacteria.

The sensor *dspA* impacts health through a second pathway by influencing an unknown response regulator *RR*, which in turn down regulates expression of the gene products *psbA1*, *psbA2*, and *cpcB*. The first two positively influence the level of photosynthetic activity (*Photo*) by altering the structure of the photosystem. If left unregulated, this second pathway would also damage the organism in high light conditions.

Although the model incorporates quantitative variables, it is qualitative in that it specifies cause and effect but not the exact numerical form of the relationship. For example, one causal link indicates that increases in *NBLR* will increase *NBLA*, but it does not specify the form of the relationship, nor does it specify any parameters.

The model is both partial and abstract. The biologist who proposed the model made no claim about its completeness and clearly viewed it as a working hypothesis to which additional genes and processes should be added as indicated by new data. Some links are abstract in the sense that they denote entire chains of subprocesses. For example, the link from *dspA* to *NBLR* stands for a signaling pathway, the details of which are not relevant at this level of analysis. The model also includes a theoretical variable *RR*, an unspecified gene (or possibly a set of genes) that acts as an intermediary controller.

### 3 An Approach to Revising Qualitative Causal Models

In this paper, we represent causal models in terms of linear relationships among variables. That is, each quantitative variable  $x(i)$  is represented with an equation of the form

$$x(i) = \sum_{j=1}^{i-1} A(i, j)x(j) + b(i) \quad , \quad (1)$$

where  $A(i, j)$  describes the causal effect of variable  $x(j)$  on  $x(i)$  and  $b(i)$  is an additive constant. The variables in a model are ordered and variable  $x(i)$  can only be influenced by those variables that come before it in the causal ordering.

Using matrix form, we can represent the equations for all  $x(i)$ ,  $i = 1..n$ , as  $\mathbf{x} = \mathbf{Ax} + \mathbf{b}$ . In this formulation,  $A(i, j) = 0$  if  $i \leq j$ , where  $A(i, j)$  denotes the element in row  $i$  and column  $j$  of  $\mathbf{A}$ . This constraint enforces the causal ordering on the variables. A model is completely specified by an ordering of variables in  $\mathbf{x}$  and an assignment of values to all elements of  $\mathbf{A}$  and  $\mathbf{b}$  that satisfy the above constraints. This defines the space of models that our revision method will consider.

However, we still need some way to map an initial biological model onto this notation. If we let  $\mathbf{A}_0$  and  $\mathbf{b}_0$  denote the initial model, then we can transform qualitative models like that in Figure 1 into a matrix  $\mathbf{A}_0$  by setting  $A(i, j) = 1$  if there is a positive link from variable  $j$  to  $i$  in the model,  $A(i, j) = -1$  if the link is negative, and  $A(i, j) = 0$  otherwise. We set the vector  $\mathbf{b}_0$  to zero for all its elements.

Given  $\mathbf{A}_0$ ,  $\mathbf{b}_0$ , and observations on  $\mathbf{x}$ , we learn new values for  $\mathbf{A}$  and  $\mathbf{b}$  by

1. Picking an initial ordering for variables in  $\mathbf{x}$ ;
2. Learning the best real-valued matrix  $\mathbf{A}$  according to a score function that penalizes for differences from  $\mathbf{A}_0$ , and is subject to the ordering constraints;
3. Swapping variables in the ordering and going to step 2 (i.e., performing hill-climbing search in the space of variable orderings), continuing until the score obtained no longer improves; and
4. Transforming the real matrix  $\mathbf{A}$  that has the best score into a discrete version with  $A(i, j) \in \{-1, 0, 1\}$  with a thresholding method.

Step 1 in this revision algorithm determines the starting state of the search. Our approach selects a random ordering that is consistent with the partial ordering implied by the initial model. During step 2, our method invokes an approach to equation revision that transforms the equation  $\mathbf{x} = \mathbf{Ax} + \mathbf{b}$  into a neural network, revises weights in that network, and then transforms the network back into equations in a fashion similar to that described by Saito et al. (2001).

This neural network approach uses a minimum description length (Rissanen, 1989) criterion during training to penalize models that differ from the initial model. For example, suppose  $\mathbf{w}_0$  is the parameter vector of the neural network that corresponds to the initial model. We define our revision task as finding a  $\mathbf{w}$  that lets the network closely replicate the observed data and is also reasonably close to  $\mathbf{w}_0$ . To this end, we consider a communication problem where a sender wishes to transmit a data set to a receiver using a message of the shortest possible length. However, unlike the standard MDL criterion, we assume that the initial model with  $\mathbf{w}_0$  is known to the receiver. Namely, we try to send message length with respect to  $\mathbf{w}_0 - \mathbf{w}$ , rather than with respect to  $\mathbf{w}$ . Since we can avoid encoding parameter values equal to the initial ones, this metric prefers the initial model. The new parameters  $\mathbf{w}_0 - \mathbf{w}$  are regarded as weights of the neural network, and their initial values are set to zero. Then, in order to obtain a learning result that is reasonably close to the initial model, the network is trained with weight decay, using a method called the MDL regularizer (Saito & Nakano, 1997).

When the modeling task includes some unobserved variables, like RR in Figure 1, we cannot directly revise links associated with those variables. To cope with such situations, our method adopts a simple forward-backward estimation based on the initial model. If  $x(i)$  is an unobserved variable, then its value can be estimated in the forward direction by the equation,  $\hat{x}(i)^{(0)} = \sum_j A(i, j)x(j) + b(j)$ . On the other hand, if  $S$  is a set of observed variables linked directly from  $x(i)$ , i.e.,  $S = \{x(k) : k > i \wedge A(k, i) \neq 0\}$ , then for  $x(k) \in S$ , the equation for the backward estimation is  $x(i) = A(k, i)^{-1}(x(k) - \sum_{j \neq i} A(k, j)x(j) - b(k))$ . This lets us estimate the values  $\{\hat{x}(i)^{(1)}, \dots, \hat{x}(i)^{(M)}\}$ , where  $M$  is the number of elements in  $S$ . Finally, our method estimates the value of  $x(i)$  as the average of these values using the equation  $\hat{x}(i) = (M + 1)^{-1} \sum_{m=0}^M \hat{x}(i)^{(m)}$ . One could repeat these two procedures, estimation of the unobserved variables and revision of the parameters, although the current implementation makes only one pass.

As stated above, our method performs gradient search through a space of parameters on causal links, with weight decay driving the model toward integer values. However, the resulting values are not strictly integers. To overcome this problem, in step 4 we employ a simple thresholding method. After sorting the resulting parameter values to predict one variable  $x(i)$ , the system uses two thresholds,  $T_{-1}$  and  $T_{+1}$ , to divide this sorted list into three portions. Parameter value  $A(i, j)$  is set to  $-1$  if  $A(i, j) < T_{-1}$ , to  $+1$  if  $A(i, j) > T_{+1}$ , and to  $0$  otherwise. Note that  $T_{-1} \leq T_{+1}$ , and we can obtain all possible integer lists with computational complexity  $O(N^2)$ , where  $N$  denotes the number of parameters.

Given these integer lists, our method selects the result that minimizes the MDL cost function defined by  $\{0.5 \times s \times \log(MSE)\} + \{r \times \log(N)\}$ , where  $s$  is the number of training samples,  $r$  is the number of revised parameters, and  $MSE$  is the mean squared error on the samples. The first term of the cost function is a code length for transmitting data, derived by assuming Gaussian noise for variables, while the second term is a code length for revision information, i.e., multiplication of the number of revised parameters and the cost of encoding an integer to indicate the parameter that is revised.

## 4 Experimental Studies of the Revision Method

In this section, we describe experimental studies of our revision method. We take a dual approach of evaluating the system using both natural data obtained from microarrays of Cyanobacteria cultures and synthetic data generated from known mathematical models. Natural data lets us evaluate the biological plausibility of changes suggested by our algorithm. However, because we have an extremely limited number of microarrays, it can be difficult to evaluate the reliability of the suggested revisions even if they appear biologically plausible. Therefore, we also used synthetic data to evaluate the robustness and reliability of our approach. Because we can generate synthetic data from a known model, we can measure the sensitivity and reliability of our algorithm in the presence of complicating factors such as errors in the initial model, small sample sizes, and noise.

#### 4.1 Revising the Model of Photosynthesis Regulation

We applied our method to revise the regulatory model of photosynthesis for wild type Cyanobacteria. We have microarray data which includes measurements for approximately 300 genes believed to play a role in photosynthesis. For this analysis, we focus on the genes in the model and do not consider links to other genes. The array data were collected at 0, 30, 60, 120, and 360 minutes after high light conditions were introduced, with four replicated measurements at each time point. We treat both RR and Photo, which represents the structure of the photosystem, as unmeasured variables. We currently treat the data as independent samples and ignore their temporal aspect, along with dependencies among the four replicates.

We implemented our method in the C programming language and conducted all experiments on a 1.3 Ghz Pentium running Linux. Revising the photosynthesis model took 0.02 seconds of CPU time. For each variable, the observed values were normalized to a mean of zero and a standard deviation of one. Figure 2 shows the revised model, which reflects three changes:

1. dropping the link from dspA to RR;
2. connecting Photo to RR instead of psbA1 and psbA2; and
3. changing the sign of the link from PBS to Health from negative to positive.

The first two changes are difficult to explain from a biological perspective. Because dspA is a light sensor, there should be either a direct or indirect path linking it with the genes cpcB, psbA1, or psbA2. Dropping the link disconnects dspA from those genes and removes it as possible cause. Also, the structure of the photosystem (Photo) is believed to depend on at least one of psbA1 or psbA2, and connecting Photo only to RR removes psbA1 and psbA2 as parents.<sup>2</sup>

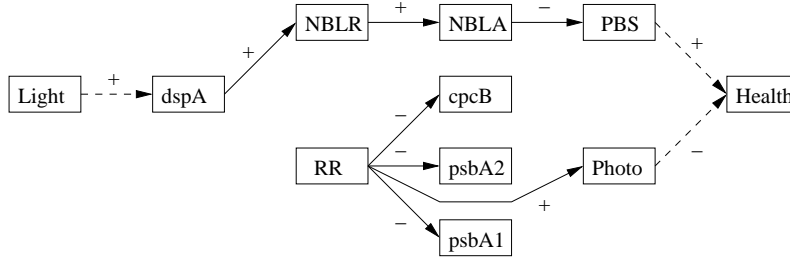
Changing the sign of the link from PBS to Health is more plausible. The initial model was specified for high light conditions in which excessive light levels damage the organism. However, at lower light levels, increased PBS should aid the organism because it is a vital component in energy production. One explanation suggested by the microbiologist is that light levels during the biological experiment may not have been set correctly and were not high enough to reduce health.

#### 4.2 Robustness of the Revision Approach

We evaluated the robustness of our approach by generating synthetic data from a known model and varying factors of interest. Specifically, we varied the number of training samples, the number of errors in the initial model, the observability of variables, and the noise level. We expected each of these factors to influence the behavior of the revision algorithm.

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<sup>2</sup> The genes psbA1 and psbA2 encode variants of the D1 protein, a necessary and central component of the Photosystem II reaction center (Wiklund et al., 2001).



**Figure 2.** Revised model of photosynthesis regulation in Cyanobacteria.

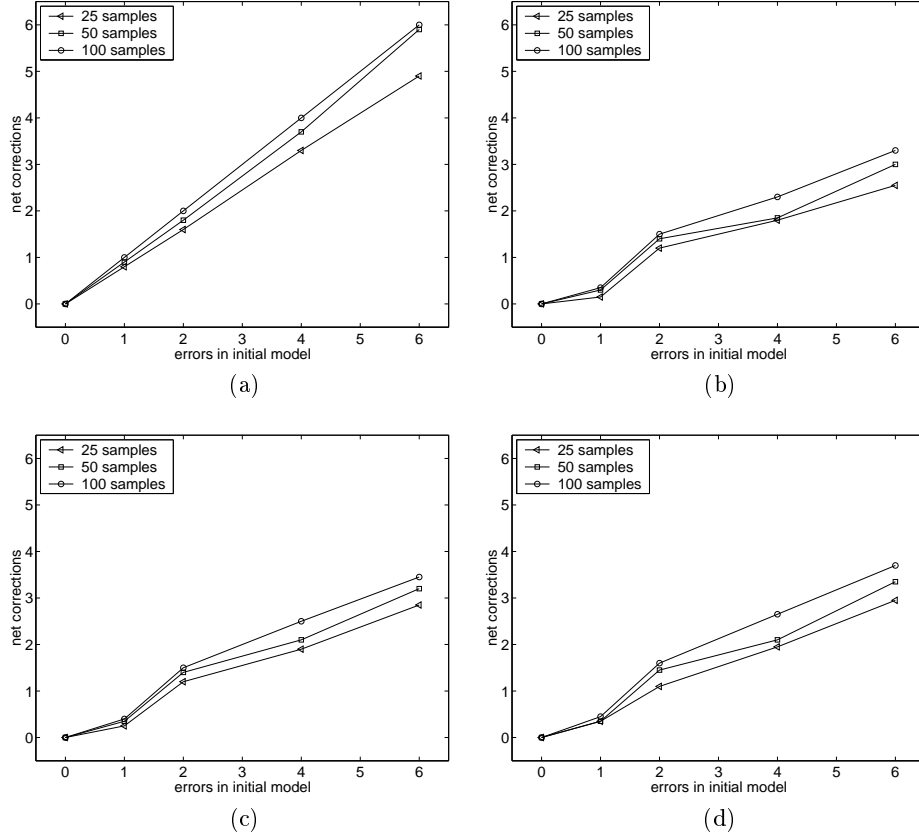
To this end, we generated training data by treating the structure of the model in Figure 1 as the true model. We assumed that each variable was a linear function of its parents with noise added from a random normal distribution. The root causal variable, Light, has no parents and was assigned a random uniform value between 0 and 1. We generated initial models to serve as starting points for revision by randomly adding links to, or deleting links from, the true model in Figure 1. Our dependent measure was the net number of corrections, that is, the number of correct changes minus the number of incorrect changes, suggested by the revision process. For each experimental condition, we generated 20 distinct training sets and averaged the results for this measure.

Figure 3 (a) shows the results from one experimental condition that involved only observable variables and only a small amount of noise ( $\sigma = 0.1$ ). The  $x$  axis in the graph represents the number of errors in the initial model, whereas the  $y$  axis specifies the net number of corrections. The three curves correspond to different size training sets, with the smallest containing only 25 instances and the largest involving 100 observations. In general, the revision method fared quite well, in that it consistently corrected almost all of the errors in the initial model. More data improved this performance, with 100 training cases being enough to give almost perfect results on all 20 runs.

However, other factors can degrade the system’s behavior somewhat. Figure 3 (b) shows the results at the same noise level when the variable RR is unobservable but all others are available. Overall, the net number of corrections decreased substantially compared to the fully observable condition. However, the method still has enough power to recover portions of the true model. Figure 3 (c) and (d) show the system’s behavior with RR unobserved at higher levels of noise, with  $\sigma = 0.2$  and  $\sigma = 0.4$ , respectively. The net number of corrections under these conditions is similar to that when  $\sigma = 0.1$ , which suggests that our approach is robust with respect to noise of this type. Note that  $\sigma = 0.4$  constitutes a rather high noise level in comparison with the range of the variables (e.g., light varies from 0 to 1).

We should also note that the system never suggested changes to the initial model when it was correct (i.e., contained zero errors). This indicates that the revision method is behaving in a conservative manner that is unlikely to make

a good model worse, even in the presence of noise, unobservable variables, and small samples. This in turn suggests that our use of minimum description length is having the desired effect.



**Figure 3.** Average net number of corrections to the initial model for 25, 50, and 100 samples when (a) all variables are observed and  $\sigma = 0.1$ , (b) the variable RR is unobserved and  $\sigma = 0.1$ , (c) RR is unobserved and  $\sigma = 0.2$ , and (d) RR is unobserved and  $\sigma = 0.4$ .

## 5 Directions for Future Research

The results from our experiments on Cyanobacteria data were disappointing, as they were difficult to explain from a biological perspective. However, on synthetic data our system was able to improve incorrect initial models even when there were few training samples, unobserved variables, and noise.

This suggests that our general approach is feasible, but that we may need to address some of the limitations, chosen by design, in the approach. For instance,



we modeled the relationships between genes as a linear function. Although linear models are desirable because they have few parameters, they cannot model combinatorial effects among genes or thresholds in which a gene's expression must be above a certain level before it can affect other genes. The neural network approach to revision is not limited to linear models and we could use a more general form to represent relationships between genes.

We also restricted the genes that could appear in the model to a small subset of those measured by the microarray chips. The complete set of data contains about 300 variables from which we used the 11 variables present in the initial model. Restricting the number of variables involves a tradeoff. Including too many variables for the number of samples makes estimating relationships unreliable because of the multiple hypothesis testing problem (Shaffer, 1995). However, using too few variables increases the likelihood that we may have ignored an important variable from the analysis. Future implementations could minimize this problem by including an operator for adding new genes during the revision process and using domain knowledge to select only the most promising candidates for incorporation into the model.

In addition, we should extend our approach to model revision in various other ways. Since transcriptional gene regulation takes time to occur, future systems should search through an expanded space of models that include time delays on links<sup>3</sup> and feedback cycles. To handle more complex biological processes, it should also represent and revise models with subsystems that have little interaction with each other. Finally, each of these extensions would benefit from incorporation of additional biological knowledge, cast as taxonomies over both genes and regulatory processes, to constrain the search for improved models.

Finally, we must test our approach on both more regulatory models and more microarray data before we can judge its practical value. Our biologist collaborators are collecting additional data on *Cyanobacteria* under more variable conditions, which we predict will provide additional power to our revision method. We also plan to evaluate the technique on additional data sets that we have acquired from other biologists, including ones that involve yeast development and lung cancer.

## 6 Related Research

Although most computational analyses of microarray data rely on clustering to group related genes, we are not the first to focus on inducing causal models of gene regulation. Most research on this topic encodes regulatory models as Bayesian networks with discrete variables (e.g., Friedman et al., 2000; Hartemink, 2002; Ong et al., 2002). Because microarray data are quantitative, this approach often includes a discretization step that may lose important information, whereas our approach deals directly with the observed continuous

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<sup>3</sup> An alternative is to model the regulation between genes with differential equations.

values.<sup>4</sup> These researchers also report methods that construct causal models from scratch, rather than revising an initial model, though some incorporate background knowledge to constrain the search process.

An alternative approach represents hypotheses about gene regulation as linear causal models, which relate continuous variables through a set of linear equations. Such systems evaluate candidate models in terms of their ability to predict constraints among partial correlations, rather than their ability to predict the data directly. Within this framework, some methods (e.g., Saavedra et al., 2001) construct a linear causal model from the ground up, whereas others (e.g., Langley et al., 2002) instead revise an initial model, as in the approach we report here. One advantage of this constraint-based paradigm is that it can infer qualitative models directly, without the need to discretize or fit continuous parameters. In contrast, our technique combines search through a parameter space with weight decay to achieve a similar end.

We should also mention approaches that, although not concerned with gene regulation, also construct causal models in scientific domains. One example comes from Koza et al. (2001), whose method formulates a quantitative model of metabolic processes from synthetic time series about chemical concentrations. Another involves Zupan et al.’s (2001) GENEPATH, which infers a qualitative genetic network to explain phenotypic results from gene knockout experiments. Mahidadia and Compton (2001) report an interactive system for revising qualitative models from experimental results in neuroendocrinology. Finally, our approach to revising scientific models borrows ideas from Saito et al. (2001), who transform an initial quantitative model into a neural network and utilize weight learning to improve its fit to observations.

## 7 Conclusions

In this paper, we characterized the task of discovering a qualitative causal model of gene regulation based on data from DNA microarrays. Rather than attempting to construct the model from scratch, we instead assume an existing model has been provided biologists who want to improve its fit to the data. These models require a causal ordering on variables, links between variables, and signs on these links. We presented an approach to this revision task that combines a hill-climbing search through the space of variable orderings and a gradient descent search for weights on links, with the latter using a weight decay method guided by minimum description length to drive weights to integer values.

We illustrated the method’s behavior on a model of photosynthesis regulation in Cyanobacteria, using microarray data from biological experiments. However, our experimental evaluation also relied on synthetic data, which let us vary systematically the distance between the initial and target models, the amount of training data available, and the noise in these data. We found that the method scaled well on each of these dimensions, which suggests that it may prove a useful

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<sup>4</sup> Imoto et al. (2002) report one way to induce quantitative models of gene regulation within the framework of Bayesian networks.

tool for revising models based on biological data. We noted that our approach has both similarities to, and differences from, other recent techniques for inducing causal models of gene regulation. We must still evaluate the method on other data sets and extend it on various fronts, but our initial experiments on synthetic data have been encouraging.

## Acknowledgements

This work was supported by the NASA Biomolecular Systems Research Program and by NTT Communication Science Laboratories, Nippon Telegraph and Telephone Corporation. We thank Arthur Grossman, Jeff Shrager, and C. J. Tu for the initial model, for microarray data, and for advice on biological plausibility.

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