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Methods

Making connections between novel transcription factors and their DNA motifs

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The key components of a transcriptional regulatory network are the connections between *trans*-acting transcription factors and *cis*-acting DNA-binding sites. In spite of several decades of intense research, only a fraction of the estimated ~300 transcription factors in *Escherichia coli* have been linked to some of their binding sites in the genome. In this paper, we present a computational method to connect novel transcription factors and DNA motifs in *E. coli*. Our method uses three types of mutually independent information, two of which are gleaned by comparative analysis of multiple genomes and the third one derived from similarities of transcription-factor–DNA-binding-site interactions. The different types of information are combined to calculate the probability of a given transcription-factor–DNA-motif pair being a true pair. Tested on a study set of transcription factors and their DNA motifs, our method has a prediction accuracy of 59% for the top predictions and 85% for the top three predictions. When applied to 99 novel transcription factors and 70 novel DNA motifs, our method predicted 64 transcription-factor–DNA-motif pairs. Supporting evidence for some of the predicted pairs is presented. Functional annotations are made for 23 novel transcription factors based on the predicted transcription-factor–DNA-motif connections.

[Supplemental material is available online at www.genome.org.]

To a large extent, gene expression programs depend on the recognition of specific promoter sequences (transcription-factor-binding sites) by transcriptional regulatory proteins (transcription factors, TFs). Studies designed to identify these binding sequences and regulatory proteins, and determine the correct connections between them, provide the data necessary to build a model of the transcriptional regulatory network of an organism (Shen-Orr et al. 2002; Bar-Joseph et al. 2003; Liao et al. 2003; Kao et al. 2004). While traditional experimental methods (e.g., electrophoretic mobility shift and nuclease protection assays) have identified only a fraction of the transcription regulatory interactions of *Escherichia coli* (Martinez-Antonio and Collado-Vides 2003), modern high-throughput methods such as chromatin immunoprecipitation coupled with promoter microarrays (ChIP-chip experiments) (Ren et al. 2000) have the potential to rapidly associate many TFs with their cognate binding sites in the genome, and thus provide the genome-scale interaction data necessary to model the *E. coli* regulatory network. However, in order for ChIP-chip to work, the growth conditions under which the TFs are active need to be known before experiments are conducted, and determining these growth conditions will undoubtedly be a challenging task. Furthermore, cloning a large number of epitope-tagged TFs (including many novel factors) remains a labor-intensive process that would likely face unforeseeable technical difficulties. Another approach was described by Segal et al. (2003), who addressed the problem of connecting a group of TFs with their regulated genes in yeast using only microarray expression data. Their method relies on the assumption that TFs are themselves transcriptionally regulated such that their expression profiles correlate with their target genes. This assumption prob-

ably holds true for many TFs, but is violated in cases where other types of regulation are involved, such as post-translational modification and binding of small molecule effectors. In addition, the expression level of some transcription factors may be too low to be reliably detected with current microarray technology.

Fueled by the ever-increasing number of completely sequenced genomes, comparative genomics has proven to be a powerful tool to address a large variety of biological questions (Marcotte et al. 1999; Overbeek et al. 1999; Pellegrini et al. 1999; Korf et al. 2001; McCue et al. 2001, 2002; Rivas and Eddy 2001; Blanchette and Tompa 2002; Rajewsky et al. 2002; Ji et al. 2004). In particular, computational methods such as phylogenetic footprinting have been applied to the *E. coli* genome, allowing the discovery of many novel TF-binding sites (McCue et al. 2001, 2002; Rajewsky et al. 2002). Furthermore, clustering of phylogenetic footprints has yielded inferences on the sets of coregulated genes (regulons), generating DNA motif models for unknown TFs as well as many previously characterized TFs (van Nimwegen et al. 2002; Qin et al. 2003). Complete genomic sequence has also allowed the prediction of the full repertoire of TFs in *E. coli* using standard computational sequence analysis techniques (Perez-Rueda and Collado-Vides 2000; Babu and Teichmann 2003). Thus, computational predictions have already provided substantial data sets for each of the two necessary components of a regulatory network for *E. coli*: the binding sites and the TFs. In this paper, we describe an *in silico* approach that harnesses the power of comparative genomics to make connections between TFs with their cognate binding sites in the *E. coli* genome.

We hypothesize that information concerning the connection of a TF to its cognate DNA motif is carried in the genomes and can be extracted by comparing multiple genomic sequences and available structural data. Specifically, our method takes advantage of three types of information in order to assign a DNA-binding motif to a given TF: (1) a distance constraint between a

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TF and its closest binding site in the genome (D_{\min} information), (2) the phylogenetic correlation between TFs and their regulated genes (PC information), and (3) a binding specificity constraint for TFs having structurally similar DNA-binding domains (FMC information). For a given TF and DNA motif, the three types of information are combined to calculate the probability that such a TF \leftrightarrow DNA-motif pair is a true pair. We demonstrate the method using a study set of known TFs and their cognate DNA-binding motifs, and further apply the method to predict connections between novel TFs and DNA motifs from *E. coli*. In addition, for functionally unannotated TFs, we are able to infer their target cellular processes based on the overrepresented functional categories of their regulons. Our results demonstrate the value of combining heterogeneous data types to solve a challenging computational problem.

Results

Distance constraint between a TF and its closest binding site in the genome, D_{\min} information

Bacterial TFs are often autoregulatory; 55% (58 of 105) of all known *E. coli* TFs in the database RegulonDB are autoregulated (Martinez-Antonio and Collado-Vides 2003). Besides autoregulation, it has been noticed in many cases (Dickson et al. 1975; Eichler et al. 1996; Palacios and Escalante-Semerena 2000; Torres et al. 2003) that TFs and the genes they regulate are near each other in the genome. Both phenomena imply a distance constraint between the TF and its closest binding site in the genome. In the first case, the distance constraint is due to the requirement for autoregulation. In the second case, the distance constraint may be the result of horizontal gene transfer because it would not be advantageous to acquire a new operon conferring a new function if the operon is not expressed correctly (Lawrence 1999; Martinez-Antonio and Collado-Vides 2003). To use this distance constraint information, we introduced the quantity D_{\min} . Two types of D_{\min} s can be considered: D_{\min_self} is the distance between a TF gene and its closest binding site in the genome. D_{\min_cross} is the distance between a TF gene and the closest binding site for a different TF. Owing to the existence of a distance constraint between a TF and its own closest site in the genome, we expect D_{\min_self} to be smaller than D_{\min_cross} for most other TFs. In Figure 1A, we plot the distributions of D_{\min_self} and D_{\min_cross} for the study set of 35 TFs. The mean of the D_{\min_self} distribution is significantly smaller than the mean of the D_{\min_cross} distribution ($p = 4.8 \times 10^{-7}$). Thus, we can use distance constraint information to connect novel TFs with their cognate DNA motifs. For any pair of transcription factor TF_j and DNA motif M_i , we can calculate the probability that they are a true pair given their D_{\min} value D_{\min}^j , $P(TF_j \leftrightarrow M_i | D_{\min}^j \leq x)$ (Supplemental Fig. 4A).

Phylogenetic correlation between TFs and their regulated genes (regulons), PC information

TFs and their regulated genes tend to evolve concurrently (Mironov et al. 1999; Gelfand et al. 2000; Tan et al. 2001). Thus, we can connect TFs and DNA motifs through correlation between their occurrences in a comparative analysis of multiple species. The "regulon" refers to the set of genes directly regulated by a common TF. If the TF is not known, such as in our problem, the "regulon" can also be defined as the set of genes controlled by a common DNA motif belonging to the as yet unknown TF. For a

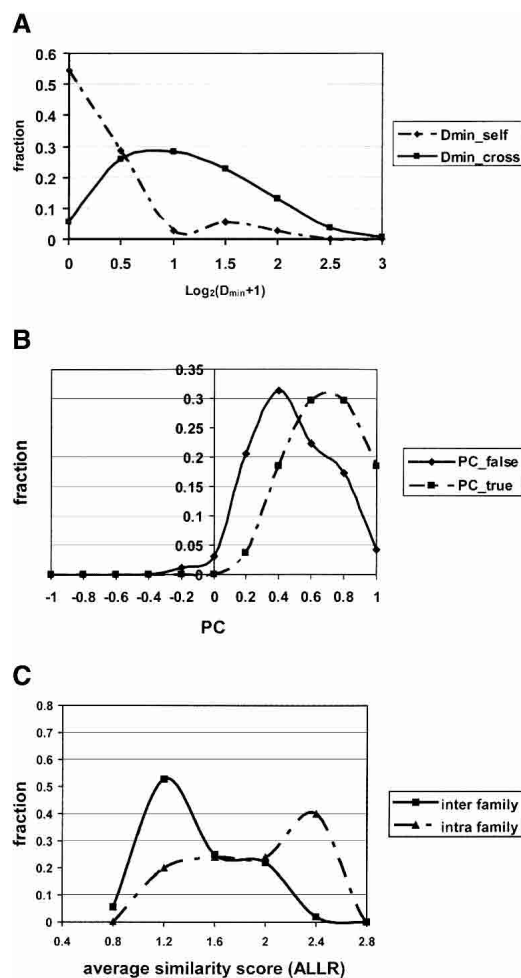


Figure 1. Three types of normalized D_{\min} distributions. D_{\min_self} : the distance between a TF gene and its closest binding site in the genome. D_{\min_cross} : the distance between a TF gene and the closest site for a different TF. (B) Two types of phylogenetic correlation (PC) distributions. PC_true: PC for true TF–DNA-motif pairs. PC_false: PC for false TF–DNA-motif pairs. (C) Distribution of average similarity scores for motifs from the same family and from different families. Motifs were aligned using an ungapped Smith-Waterman algorithm and scored using the ALLR statistic.

given DNA motif M_i and a species G_k , we can build a \overrightarrow{GR}_{ik} vector (see Methods) to represent the putative regulon controlled by the DNA motif M_i in species G_k . Each element of \overrightarrow{GR}_{ik} denotes the probability that a gene is controlled by M_i . An alternative to \overrightarrow{GR}_{ik} is a binary vector whose elements indicate whether or not orthologs in species G_k are controlled by M_i by imposing a binding site cutoff. However, a binary vector contains no information about the number or quality of the binding sites in front of a gene (regulation strength by M_i), whereas the \overrightarrow{GR}_{ik} vector does. In pilot studies, we found that \overrightarrow{GR}_{ik} gave a better performance than binary vectors (data not shown). We then computed the Pearson's correlation coefficient between the *E. coli* \overrightarrow{GR}_{ik} vector ($k = E. coli$) and the \overrightarrow{GR}_{ik} vector for each of the other species considered. This measures the degree of conservation of the *E. coli* regulon determined by M_i in species G_k . When multiple species are con-

sidered, the $\overrightarrow{GR_{ik}}$ vector correlation coefficients form a new vector, $\overrightarrow{RC_i}$, which described the evolutionary changes of the *E. coli* M_i regulon in other species. We then compared the $\overrightarrow{RC_i}$ vector with the $\overrightarrow{TF_j}$ vector that describes the phylogenetic profile of TF_j in the various genomes. We expect that there will be no strong correlation in those species that have lost TF_j . For example, using the TF *trpR*, Table 1 shows the vector for its conservation $\overrightarrow{TF_{trpR}}$ and the vector for its regulon conservation $\overrightarrow{RC_{trpR}}$ in 13 γ -proteobacteria. As can be seen, only those species that have an ortholog to *E. coli* *trpR* (with a 1 in the TF column) also have a large positive value in the $\overrightarrow{RC_{trpR}}$ vector, meaning conservation of the regulon. We can use this phylogenetic correlation information to connect novel TFs with their cognate DNA motifs based on the difference in PC values between true and false TF-DNA-motif pairs ($p = 2.12 \times 10^{-4}$) (Fig. 1B). For any pair of transcription factor TF_j and DNA motif M_i , we can calculate the probability that they are a true pair given their PC^{ij} value $P(TF_j \leftrightarrow M_i | PC^{ij} \geq \gamma)$ (Supplemental Fig. 4B).

Binding specificity constraint for TFs having structurally similar DNA-binding domains, FMC information

TFs that are more similar to one another are expected to bind to sites that are more similar to each other than to dissimilar pairs. For instance, the DNA motifs of many TF family members are often similar (Luscombe et al. 2000; Rigali et al. 2002; Sandelin and Wasserman 2004). Using the study set of 35 TFs, we examined this issue in more detail by studying the relationship of the DNA-binding domain (DBD) of TFs within a family and similarities in their DNA motifs. The database SUPERFAMILY (Madera et al. 2004) contains hidden Markov models (HMMs) for DBDs belonging to different structural families. We used the database to classify the 35 TFs into seven structural families based on the similarity between our query DBDs and the DBD HMMs in the database. For these seven TF families, we calculated the average pairwise similarities of their corresponding DNA motifs. Figure 1C shows the distributions of average pairwise similarity scores for motifs belonging to the same family and motifs from different families. The mean intrafamily pairwise similarity score is significantly larger than the mean interfamily pairwise similarity

Table 1. An example of phylogenetic correlation between the transcription factor *trpR* and its regulon

Species	TF	Regulon conservation
<i>E. coli</i>	1	1.0000
<i>H. influenzae</i>	1	0.3953
<i>P. multocida</i>	1	0.5237
<i>P. aeruginosa</i>	0	-0.0399
<i>P. putida</i>	0	0.0297
<i>P. syringae</i>	0	0.0507
<i>S. oneidensis</i>	0	0.0866
<i>S. typhi</i>	1	0.9721
<i>V. cholerae</i>	1	0.9664
<i>V. parahaemolyticus</i>	1	0.3588
<i>V. vulnificus</i>	1	0.8106
<i>X. campestris</i>	0	-0.0495
<i>Y. pestis</i>	1	0.2105

"1" indicates the existence of a *trpR* ortholog in the given species, "0" otherwise. The degree of regulon conservation is calculated using *E. coli* as the reference species (see Methods for calculation). The phylogenetic correlation is $PC_{trpR} = 0.8016$.

Table 2. Probability increase for true TF-DNA motif pairs using different combinations of information

Type of information	Average P_true/P_avg
No information	1
FMC	2.48
PC	1.33
D_{min}	2.74
FMC- D_{min}	5.38
FMC-PC	3.21
PC- D_{min}	3.49
FMC-PC- D_{min}	6.48

Probabilities calculated using the study set of TFs and their DNA motifs. P_true: the probability of the true TF-DNA motif pair. P_avg: the average probabilities of all TF-DNA motif pairs for a given TF.

score ($p = 0.003$). This result further confirmed the hypothesis that TFs with similar DBDs tend to have similar DNA-binding motifs. To use this binding constraint information, we introduced the quantity FMC (familial motif conservation), which measures the average similarity between a query DNA motif and a family of DNA motifs. FMC serves as an estimate of the membership of a query motif to a family of motifs. For any pair of transcription factor TF_j and DNA motif M_i , we can calculate the FMC between M_i and the DNA motif family of which TF_j 's motif is a member. We can then calculate the probability that they are a true pair given their FMC value FMC^{ij} , $P(TF_j \leftrightarrow M_i | FMC^{ij} \geq z)$ (Supplemental Fig. 4C).

Making TF \rightarrow DNA-motif connections by combining three types of information

Given a set of TFs and DNA motifs, we calculate the probability that a TF \leftrightarrow DNA-motif pair is true for all possible pairs. We can calculate three types of probabilities for each pair based on the three types of information described before: $P(TF_j \leftrightarrow M_i | D_{min}^{ij} \leq x)$, $P(TF_j \leftrightarrow M_i | PC^{ij} \geq \gamma)$, and $P(TF_j \leftrightarrow M_i | FMC^{ij} \geq z)$. Since the different types of information are independent of each other, the joint probability $P(TF_j \leftrightarrow M_i | D_{min}^{ij} \leq x, PC^{ij} \geq \gamma, FMC^{ij} \geq z)$ considering all types of information can be calculated using the three conditional probabilities mentioned above (see Methods). By combining different information, the probabilities of true connections could be raised much higher above the background probabilities of false connections. This increase in signal-to-noise ratio can be illustrated using the P_true/P_avg ratio (Table 2), the probability ratios between the true connections (P_true) and the average of all connections for a given TF (P_avg). On average, the P_true/P_avg ratios were 2.18, 4.03, and 6.48 after one, two, and three types of information were used, respectively.

Assessment of the algorithm using the study set

To test the performance of our algorithm, we assembled a set of 35 well-characterized transcription factors and their corresponding DNA-binding motifs. These 35 TFs regulate a large variety of cellular processes in response to many different stimuli. Based on the number of binding sites, information content, and E-value of the motif models (Supplemental Fig. 5), these 35 DNA motifs are fairly representative of all DNA motifs discovered in *E. coli* so far. The study set of 35 TFs and their DNA motifs was used to evaluate the prediction accuracy of our algorithm in three different ways. First, we use only the study set of 35 motifs and TFs and we determine how accurately we make correct assignments using

different combinations of information. In Figure 2, the cumulative frequency of the true connection ranks was plotted. As can be seen, combining different information improves prediction accuracy. By combining the three types of information, 63% of the true connections were ranked first out of the 35 possible connections for each TF, and 85% of the true connections were ranked in the top three. In comparison, only 3% of the true connections would be ranked first and 9% ranked in the top three if we didn't have any information and made our predictions by random guessing.

In a second test we evaluated our prediction accuracy on the study set after combining the study set with the set of novel TFs and DNA motifs. There are 134 TFs and 105 DNA motifs in this combined set. Now, the probability of success by random guessing is three times smaller than using the study set alone. As illustrated in Figure 3, the prediction accuracy only dropped slightly (59% of the true connections were ranked first, and 81% of the true connections were ranked in the top three), suggesting that our algorithm still works well in a larger search space.

In the third test we did a fivefold cross-validation with the combined study set and novel set, as described for the previous test. For each cross-validation, 28 TF and DNA motif pairs from the study set were used to derive the probability distributions, and the remaining seven TF and DNA motif pairs were used to assess the accuracy. As described above, we combined the test set of seven pairs with the novel set of TFs and DNA motifs when making predictions. The prediction accuracy measurement is the same as described in Figure 1 (fractions of true pairs ranked in the top three). The average prediction accuracy based on the five rounds of cross-validation is similar to that shown in Figure 1: 64.7% of the true pairs ranked as number one, 15.2% of the true pairs ranked as number two, and 10.4% of the true pairs ranked as number three. This demonstrates that the accuracy remains high for known pairs that were not used to determine the three probability distributions.

Predicted connections between novel TFs and DNA motifs

For all 6930 TF-DNA-motif pairs in the novel set, we calculated their probabilities of being a true pair using all three types of information. From these posterior probabilities, we can make two types of predictions. The first type of prediction is the top three DNA motifs for each TF ranked by their probabilities, the top-three-pick (TTP) method. However, since there are more TFs than DNA motifs, this forces DNA motifs to be associated with TFs whose cognate motifs are not present in the set of motifs considered. This also allows two or more TFs to share motifs. Although it is possible for paralogous TFs to have highly similar

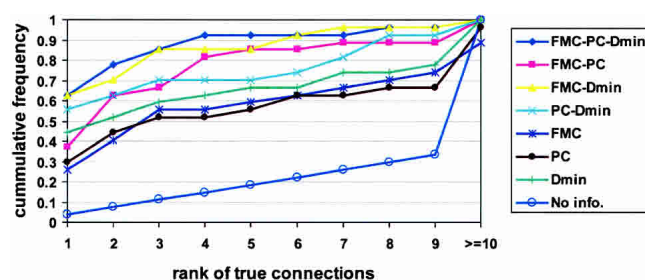


Figure 2. Prediction accuracy on the study set using different combinations of information. Size of search space: 35 TFs \times 35 motifs. The connections are ranked according to $P(TF \leftrightarrow M)$.

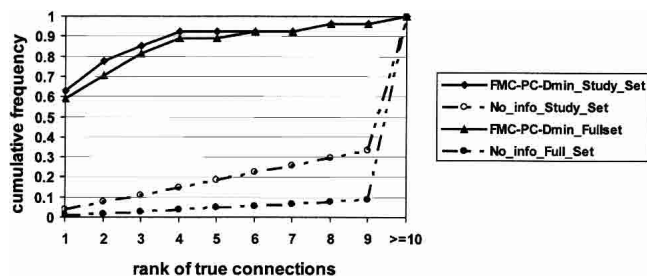


Figure 3. Prediction accuracy on the study set using all three types of information. Size of search space: 134 TFs \times 105 motifs. The connections are ranked according to $P(TF \leftrightarrow M)$.

DBDs and DNA motifs, we expect such cases to be rare. To resolve those issues, we resort to the maximal weighted matching algorithm (MWM) to make a set of unique predictions (Cary and Stormo 1995). Using the table of probabilities as input, the MWM algorithm predicted 64 unique TF-DNA-motif pairs, leaving six motifs with no associated TFs (Supplemental Table 7). Most of the predictions by MWM are ranked in the top three out of the 70 possible motifs that can be associated with each TF, indicating a good overlap between the MWM predictions and the TTP predictions. Specifically, among the MWM predictions, 28 TFs have a predicted DNA motif ranked as number one, 11 TFs have a motif ranked as number two, and 13 TFs have a motif ranked as number three. In total, 81% of the 64 TFs have a predicted DNA motif ranked in the top three.

Assessment of the algorithm on the novel predictions

The three TFs in the novel set with fewer than five known binding sites (*ntrC*, *pdhR*, *xylR*) provided a means to assess the performance of our algorithm. Both *pdhR* and *xylR* were correctly associated with their known DNA motifs, but *ntrC* was left unassociated by the MWM algorithm. However, *ntrC*'s motif was ranked in the top three by the TTP method. This result is very interesting because it provides an independent validation for our algorithm. We also searched the literature for additional supporting evidence for our predictions. Of the 64 predicted TFs, 15 have known target genes discovered via genetic analysis, but no binding sites have been reported. If our method makes correct predictions, there should be high-scoring sites for the predicted motif in the regulatory region for the known target genes. We searched the regulatory regions of these genes for binding sites of the predicted DNA motif for each TF. For nine out of these 15 cases, a binding site (scored above the site cutoff) was identified in the regulatory region of the target gene (Table 3), eight predicted by both the TTP and MWM algorithm and one by the TTP algorithm alone. For all reported sites in Table 3, the probability of occurring by chance is significantly smaller than 0.05. Thus, it is likely that those nine DNA motifs are truly recognized by the predicted TFs.

Cellular target processes of novel transcription factors

Once a TF is connected to a DNA motif, we can infer the target cellular process of the TF by studying the enrichment of functional categories of the regulated genes. We made use of 21 COG (Tatusov et al. 1997) functional categories assigned to the 4279 genes in the *E. coli* K-12 genome (Blattner et al. 1997). The 32 DNA motifs (nine known, 23 novel) whose regulons contain at least one overrepresented COG functional category are listed

Table 3. Predicted TF-DNA motif pairs with known target genes.

TFs	Regulated genes	Predicted DNA motifs	Site scores	$-\log_{10}(P\text{-value})$
ascG	<i>ascFB</i>		10.25	3.40
baeR	<i>mdtABC</i> , <i>yicO</i>		11.32, 9.76	3.83, 3.52
cadC	<i>cadBA</i>		11.48	3.60
dsdC*	<i>dsdXA</i>		7.76	2.43
emrR	<i>emrAB</i>		NA	NA
feaR	<i>tynA</i> , <i>padA</i>		NA	NA
glcC	<i>glcDFGB</i>		9.39	2.93
hydG	<i>hydHG</i>		NA	NA
idnR	<i>idnDOTR</i>		7.59	3.21
lrhA	<i>flhDC</i> , <i>lrhA</i>		6.64, 12.60	2.91, 3.93
mhpR	<i>mhpABCDE</i>		NA	NA
nadR	<i>nadA</i> , <i>nadB</i>		11.22, 5.76	3.93, 2.82
ntrC*	<i>glnALG</i> , <i>glnHPQ</i> , <i>glnK-antB</i>		16.20, 10.52, 11.32	6.13, 4.96, 5.12
pdhR	<i>pdhR-aceEF-tpd</i> , <i>lctPRD</i>		13.53, 11.12	5.32, 4.84
rpiR	<i>rpiB</i>		NA	NA
slyA	<i>hlyE</i>		11.80	4.20

All pairs were predicted by both MWM and TTP except for *dsdC* and *ntrC* (labeled with a star) which are predicted only by TTP. P-value of a site having the observed score was calculated as the tail probability of a normal distribution for all possible scores in the genome.

in Table 4 along with their corresponding TFs. Of the 13 known DNA motifs in the McCue et al. (2002) prediction set, nine have regulons with enriched COG functional categories. All enriched functional categories are consistent with the known cellular process controlled by the associated TFs. We also associated target cellular process(es) for 10 TFs whose DNA motifs were unknown before this study but some knowledge about their target genes were known. Some of the associated processes are consistent with the current knowledge about the TFs. For instance, the TF *baeR* regulates several transmembrane proteins (*mdtABC*) that form a hetero-multimeric drug efflux pump (Nagakubo et al. 2002). This is consistent with the target cellular process predicted by functional category enrichment: cell envelope biogenesis, outer membrane. Another example is the TF *ascG*, which regulates genes involved in carbon source metabolism (Postma et al. 1993). The remaining 13 predictions are for putative TFs without any functional annotation. Thus, our study provided the first description of the cellular processes that these putative TFs may regulate.

Discussion

One of the grand goals for post-genomic research is to understand the organization of the transcriptional regulatory network in an organism. Fundamental requirements for this goal are the delineation of a network's architecture, its wiring diagram, and estimation of the parameters of its components. To do this, one can take a bottom-up approach in which the parts list of TFs and DNA motifs in a genome can be identified first. Subsequently, these two sets of elements can be connected, resulting in a set of primary connections in the genetic network. These primary connections then can serve as a scaffold upon which the entire regulatory network containing higher-order interactions can ultimately be built. Shen-Orr et al. (2002) has taken the first step to identify higher-order connections in the regulatory network of *E. coli* based on a collection of known primary connections. They found three types of network motifs (higher-order connections) that are significantly overrepresented in the *E. coli* network. However, this network is far from complete because of our limited knowledge about the connections between TFs and DNA-binding sites. Thus, identifying primary connections between TFs and DNA-binding sites represents a major bottleneck for modeling transcriptional regulatory networks. In this paper, we have taken the first step to address this problem through computational means.

We hypothesize that information concerning the connection of a TF to its DNA motif is carried in the genome sequences, and we can extract this information by comparing multiple genomic sequences. TFs and their binding sites are often in similar genomic locations (D_{\min} information), and they tend to evolve concurrently with their regulated genes (PC information). We explored both types of information to identify TF-DNA-motif connections. Both D_{\min} and PC information were derived from a comparative analysis of multiple genomes. Using a comparative genomics approach has both advantages and drawbacks. On one hand, it increases our confidence in the predictions as our inference is based on reinforced signals from multiple genomes. On the other hand, this also means the success of our method depends on a wise choice of species. It has been observed that phylogenetic distance, similarity of habitat, genome size, and the number of shared genes are important factors in the selection of species for phylogenetic footprinting (McCue et al. 2002). Since we are also studying transcriptional regulatory systems using comparative genomics, we expect the same factors to be important for species selection. However, we have the additional concern that the species should have enough variation in their regulatory networks to provide our method with the signals necessary to distinguish true DNA-motif \leftrightarrow TF connections from background. Taking both considerations into

Table 4. Enrichment of regulons for genes within COG functional categories

DNA motif ID	TF	COG functional category	$-\log_{10}$ (<i>P</i> -value)
Known TF–DNA motif pairs			
c1373_e	crp	Carbohydrate transport and metabolism; energy production and conversion	10.03, 3.38
c2225_o	fadR	Lipid metabolism	5.03
c594_e	fnr	Energy production and conversion	6.06
c577_e	fruR	Carbohydrate transport and metabolism	8.06
c583_e	lexA	DNA replication, recombination and repair; cell division and chromosome partitioning	10.54, 4.53
c590_e	metJ	Amino acid transport and metabolism	5.06
c591_e	mlc	Energy production and conversion; carbohydrate transport and metabolism	3.42, 3.19
c582_e	purR	Nucleotide transport and metabolism	9.28
c595_e	trpR	Amino acid transport and metabolism	4.27
Novel TF–DNA motif pairs			
c4574	ascG	Carbohydrate transport and metabolism	5.44
c352_o	baeR	Cell envelope biogenesis, outer membrane	3.33
c580_e	cadC	Energy production and conversion; nucleotide transport and metabolism	3.60, 2.65
c489_o	hydG	Transcription	3.33
c6105_o	idnR	Energy production and conversion	5.14
c7434_o	lrhA	Amino acid transport and metabolism	3.12
c645_o	nlp	Nucleotide transport and metabolism; translation, ribosomal structure and biogenesis	4.82, 5.14
c646_o	ntrC	Amino acid transport and metabolism	3.11
c648_o	pdhR	Energy production and conversion	5.55
c647_o	xyIR	Carbohydrate transport and metabolism	4.47
c417_e	yagA	Carbohydrate transport and metabolism	3.25
c419_o	ycfQ	Signal transduction mechanisms	2.74
c428_o	ydcN	Energy production and conversion	4.35
c428_e	ydfH	Energy production and conversion	2.73
c6271_e	ydhB	Energy production and conversion; cell envelope biogenesis, outer membrane	2.91, 3.19
c571_e	yfeR	Lipid metabolism; cell envelope biogenesis, outer membrane	3.51, 3.00
c581_e	yfeT	Nucleotide transport and metabolism	2.77
c477_o	yfhH	Energy production and conversion	2.79
c3037_o	yhjC	Translation, ribosomal structure and biogenesis	9.94
c498_o	yidZ	Translation, ribosomal structure and biogenesis	2.65
c474_o	yjfQ	Translation, ribosomal structure and biogenesis	3.39
c467_o	ynfL	Coenzyme metabolism	3.32
c641_o	yqhC	DNA replication, recombination and repair	3.06

P-values were calculated as the tail probability of the hypergeometric distribution for finding at least *k* genes from a particular COG functional category in a regulon of size *n*. Since we considered 21 functional categories, 2.4×10^{-3} was chosen as the *P*-value cutoff to give an overall significance level of 0.05 for each regulon. Associations of TFs and DNA motifs are made according to the MWM method.

account, we chose the seven species used by McCue et al. (2001, 2002) for discovering novel DNA motifs in *E. coli* and added six more γ -proteobacterial species (Supplemental Table 5). This set of species provides a good balance of conservation and variation for our analyses.

We also took advantage of the observation that TFs from the same structural family tend to have similar DNA motifs. The 248 TFs in *E. coli* fall into 12 families based on the structural similarity of their DBDs. Based on binding data from RegulonDB and DPinteract, DNA motif models can be constructed for 77 TFs in these 12 families. Some of the motif models are less accurate since they are built from a very small number of known sites (<5). As the amount of binding site data increases for various families of TFs, the utility and accuracy of the FMC information will be enhanced.

We have shown the value of combining heterogeneous information to connect novel TFs and DNA motifs. All three types of information provided evidence for particular connections between TFs and DNA motifs. None of these three types of information alone is highly specific and still maintains reasonable sensitivity. But by combining the evidence of each type we were able to achieve good prediction accuracy, as shown in Figures 2 and 3. In addition, the flexibility of our approach allows new and diverse types of information to be incorporated easily because the order of the information is not important. With the availability of multiple finished genomes for many bacterial groups,

our approach can also be applied to other bacterial species with large numbers of neighbors, such as the Gram positive bacterium *Bacillus subtilis* and the environmentally significant bacterium *Shewanella oneidensis*.

Methods

Genomic sequences

All Genomic sequences were downloaded from the NCBI RefSeq database (Pruitt and Maglott 2001). The genomes used in our study are *E. coli* K-12 MG1655, *Haemophilus influenzae* Rd, *Pasteurella multocida*, *Pseudomonas aeruginosa* PAO1, *Pseudomonas putida* KT2440, *Pseudomonas syringae* pv. tomato, *Shewanella oneidensis* MR-1, *Salmonella typhi* CT-18, *Vibrio cholerae* El Tor, *Vibrio parahaemolyticus* RIMD 2210633, *Vibrio vulnificus* CMCP6, *Xanthomonas campestris* pv. *campestris*, and *Yersinia pestis* CO92.

Two gene sequences were deemed orthologous if they satisfied the following three criteria simultaneously (Huynen and Bork 1998; Tan et al. 2001): (1) They were the most similar sequences for each other between the two genomes. (2) Their BLASTP *E*-value was lower than 10^{-10} . (3) Their BLASTP alignment extended to at least 60% of one of the sequences.

Study set transcription factors and their DNA-binding motifs

TF-binding sites obtained through footprinting experiments were extracted from the databases RegulonDB v3.2 (Salgado et al.

2004) and DPinteract (<http://arep.med.harvard.edu/dpinteract/index.html>). After removing redundant sites for the same TF, 35 TFs have at least five experimentally verified binding sites. The average number of binding sites per TF is 20. Weight matrix models for DNA-binding sites were constructed for these 35 TFs using the multiple sequence alignment program CONSENSUS (Hertz and Stormo 1999). In this paper, DNA motifs refer to these weight matrix models. The TFs are araC, arcA, argR, cpxR, crp, deoR, dnaA, fadR, flhD, fnr, fruR, fur, galR, glpR, gntR, hipB, ilvY, lexA, lrp, malT, marA, metJ, metR, mlc, modE, nagC, narL, narP, ompR, phoB, purR, soxS, torR, trpR, and tyrR.

Full sets of TFs and DNA motifs in *E. coli*

Two groups have conducted computational surveys to identify the repertoire of TFs in *E. coli*. Using a combination of sequence homology search and literature mining, Perez-Rueda and Collado-Vides (2000) identified a total of 314 TFs in *E. coli*. By looking for signature protein domains present in TFs, Babu and Teichmann (2003) independently identified a total of 273 TFs in *E. coli*. We took the intersection of the two sets as the set of TFs in *E. coli*, which contains 248 proteins. McCue et al. (2002) conducted a whole-genome phylogenetic footprinting study of *E. coli* and six additional genomes, resulting in thousands of palindromic DNA motif predictions that consist of cross-species site alignments. Clustering a set of statistically significant ($P < 0.05$) motifs from this study yielded predicted regulons (Qin et al. 2003). For this study we used as input a set of 113 motifs (predicted regulons) that resulted from clustering a less stringent set of statistically significant ($P < 0.2$) phylogenetic footprint motifs (L.A. McCue, unpubl.; data available at <http://www.wadsworth.org/resnres/bioinfo/>).

Novel sets of TFs and DNA motifs in *E. coli*

From the full set of 248 TFs and 113 DNA motifs, we identified a set of novel TFs and DNA motifs to make connections. They were selected via the following procedures. For DNA motifs, we removed all motifs containing *E. coli* sites that overlap with verified TF-binding sites for any of the 35 TFs in the study set, reported RNA secondary structures, and intergenic repeats. We ended up with 70 novel DNA motifs. Among them, three motifs contain binding sites for known TFs (ntrC, pdhR, xylR) that are not included in our study set because they have fewer than five verified binding sites. These motifs remained in our novel set as an additional way to evaluate the performance of our algorithm because the TF-DNA-motif associations are known in these cases. For TFs, we removed those with documented sites in RegulonDB and/or DPinteract, including the study set TFs and TFs with known nonpalindromic DNA motifs. Again, ntrC, pdhR, and xylR were retained for performance evaluation. In addition, since the set of DNA motifs was derived from alignments having sites from *E. coli* and at least one more genome that is not *S. typhi* (owing to the close phylogenetic distance between *E. coli* and *S. typhi*, alignments having sites from only these two genomes may not represent functional DNA elements), we removed TFs that only occur in *E. coli* or in *E. coli* and *S. typhi*. This procedure resulted in 99 novel TFs (Supplemental Table 6).

Calculation of normalized D_{\min} (minimal distance)

The study set and novel DNA motifs in this study represent partial regulons; in some cases a DNA motif may contain only one or two binding sites from a given species like *E. coli*. Thus, we needed to more fully delineate the regulons in order to calculate a minimal distance between a TF-encoding gene and a binding site. Specifically, for a given DNA motif M_i , all intertranscription

unit regions in the genome were scored against the weight matrix representing M_i (constructed from clustered sites as described in above). Sites scoring above the mean minus 1.5 standard deviations of the training set scores were regarded as likely binding sites of M_i . Determining the cutoff score for binding sites is a hard problem and inevitably involves some arbitrary decision. Previously, the mean minus one or two standard deviations, or the lowest score of training sequences, have each been used as site cutoffs (Robison et al. 1998; Gelfand et al. 2000; De Wulf et al. 2002). The cutoff used in this study is a compromise that we have found yields many true binding sites without too many false positives (Tan et al. 2001).

For a given transcription factor TF_j , we then calculate a D_{\min}^{ij} by the following procedure. The binding site closest to the translation start of TF_j is used to calculate the unnormalized minimal distance d_{\min}^{ij} , which is the number of genes between the closest binding site and the translation start of TF_j . Each d_{\min}^{ij} is divided by $\langle d_{\min}^i \rangle$ to give the normalized D_{\min}^{ij} , where $\langle d_{\min}^i \rangle$ denotes the mean of all d_{\min}^{ij} 's between M_i and the set of TFs under consideration. The normalization is needed to account for the difference between motifs (having low information content) that occur very frequently and motifs (having high information content) that occur very infrequently. We calculate a D_{\min}^{ij} for each genome and the final D_{\min}^{ij} is the average of all D_{\min}^{ij} 's for species having an ortholog of TF_j .

Calculation of PC (phylogenetic correlation)

For a given DNA motif M_i and a transcription factor TF_j , we calculate a PC^ij . The two vectors, \vec{TF}_j and \vec{RC}_i , are K -element vectors, where K is the number of species under consideration. Each element k of \vec{TF}_j assumes the value 1 or 0, where 1 denotes that an ortholog of TF_j was detected in species k , and 0 otherwise. Each element k of \vec{RC}_i assumes a value that measures the degree of conservation between the *E. coli* regulon determined by M_i and its counterpart in species k . These values are calculated by using the gene regulation vector, \vec{GR}_{ik} . For species k , the elements of \vec{GR}_{ik} represent orthologs shared between *E. coli* and that species. The value of each element is the probability that the orthologous genes are controlled by the DNA motif M_i . Given a DNA motif M_i and the regulatory region S , the probability that this region is bound by a TF having the motif M_i can be approximated as:

$$P(\text{bound} | S, M_i) \approx c \sum_{j=1}^l e^{s_j} / \sum_{j=1}^g e^{s_j}$$

where l is the length of S and g is the length of the genome, s_j is the score of the sequence word starting at position j , and c is a constant depending on the concentration of the TF in the cell (Heumann et al. 1994; Workman and Stormo 2000). All genes in a transcription unit (TU) are assigned the same probability (of being regulated by M_i), which is calculated using the regulatory region of the TU. Transcription units are predicted using the method described in Salgado et al. (2000). The regulatory region of a TU is the intergenic sequence between the first gene of the TU and the last gene of the upstream TU. The Pearson's correlation coefficients between an *E. coli* \vec{GR}_{ik} vector and \vec{GR}_{ik} vectors for other genomes form the \vec{RC}_i vector, which describes the evolutionary changes of the regulon predicted to be controlled by the motif M_i . The phylogenetic correlation between the transcription factor TF_j and the regulon controlled by M_i , PC^ij , is calculated as the Pearson's correlation coefficient between the \vec{TF}_j vector and the \vec{RC}_i vector.

Calculation of FMC (familial motif conservation)

Given a DNA motif M_i and a transcription factor TF_j belonging to the structural family α , FMC^{ij} is calculated as follows:

$$FMC^{ij} = \frac{1}{n} \sum_{k=1}^n ALLR_{ik}$$

where n is the number of TFs in family α and $ALLR_{ik}$ is the similarity score between motifs i and k . The majority of the DNA motifs have inverted or direct repeats; we can compare them by splitting the full matrices at the center of symmetry. This avoids the problem of allowing insertions/deletions in the alignment when comparing motifs with different length spacer regions between the most conserved positions. A few DNA motifs have asymmetric patterns, for them the full matrices are used. Now matrices can be aligned using an ungapped Smith-Waterman algorithm, modified for profile alignment instead of sequence alignment, which finds the highest scoring local alignment between them. The scoring function for the alignment between a pair of columns from two alignment matrices is the average log likelihood ratio (ALLR) (Wang and Stormo 2003):

$$ALLR = \frac{\sum_{b=A}^T n_{bj} \ln \frac{f_{bi}}{p_b} + \sum_{b=A}^T n_{bi} \ln \frac{f_{bj}}{p_b}}{\sum_{b=A}^T n_{bi} + n_{bj}}$$

where i and j are two columns from the two alignment matrices respectively, n_{bi} and n_{bj} are count vectors for base b , f_{bi} and f_{bj} are frequency vectors for base b , and p_b is the frequency of base b in the background model. The score of aligning two matrices is the sum of scores for all columns in the alignment.

Probability of a TF–DNA-motif connection

The posterior probability of a connection being true given one type of information can be calculated using Bayes' rule:

$$P(TF \leftrightarrow M | I) = \frac{P(I | TF \leftrightarrow M) * P(TF \leftrightarrow M)}{P(I)}$$

where I refers to one of the following three terms: $D_{\min} \leq x$, $PC \geq y$, or $FMC \geq z$. We use a flat prior of $1/N$, where N is the number of TFs or DNA motifs, whichever is smaller. Using Bayes' rule and the assumption of independence of the three types of information, the joint probability of a true $TF \leftrightarrow DNA$ connection considering all types of information simultaneously, $P(TF \leftrightarrow M | D_{\min} \leq x, PC \geq y, FMC \geq z)$, is the product of the following four terms:

$$\frac{P(D_{\min} \leq x | TF \leftrightarrow M)}{P(D_{\min} \leq x)}, \frac{P(PC \geq y | TF \leftrightarrow M)}{P(PC \geq y)}, \frac{P(FMC \geq z | TF \leftrightarrow M)}{P(FMC \geq z)}, P(TF \leftrightarrow M).$$

The first three terms represent the application of the three types of information independently, and the fourth term is the prior probability. Probability distributions calculated based on the study set are used to set the probabilities for the novel set of TFs and DNA motifs.

Maximum weighted matching algorithm

A maximum weighted matching algorithm (MWM) has been used to predict RNA secondary structures (Cary and Stormo 1995;

Tabaska et al. 1998). To apply MWM to our problem, we convert the posterior probability table (containing the probabilities for all possible TF–DNA-motif pairs) into a bipartite graph, only allowing edges between each TF and each DNA motif. The edge weight between TF_j and motif M_i is calculated as follows:

$$w^{ij} = \frac{P(TF_j \leftrightarrow M_i | D_{\min}^{ij} \leq x, PC^{ij} \geq y, FMC^{ij} \geq z) - \langle P \rangle}{\langle P \rangle}$$

where $\langle P \rangle$ is the average of the probabilities of associating TF_j with each of the DNA motifs.

To find the best set of TF–DNA-motif pairs, the algorithm finds the match with the highest total edge weight. MWM is guaranteed to reach the optimal solution (Gabow 1976).

Calculation of P -value for functional category enrichment

Genes having a binding site for a DNA motif, scored above the cutoff, are regarded as members of the regulon. The hypergeometric distribution is used to calculate the P -value of observing the number of genes from a particular COG functional category within a regulon. Specifically, the probability of having at least k genes from a functional category within a regulon of size n is given by:

$$P = 1 - \sum_{i=0}^{k-1} \frac{\binom{M}{i} \binom{N-M}{n-i}}{\binom{N}{n}}$$

where M is the total number of genes in a functional category, and N is the total number of genes in the *E. coli* K-12 genome (4279). Since we tested 21 COG functional categories for each regulon, a significance value of 2.4×10^{-3} is used for each functional category to give an overall significance value of 0.05 for each regulon.

Calculation of P -value for binding sites

Scores of all possible l -mers (l is the length of the DNA motif) in the genome is calculated using the program PATSER. The scores conform to a normal distribution. The tail probability of finding a site having the observed score is calculated by converting the score distribution into a standard normal distribution.

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