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# Conservation of DNA Regulatory Motifs and Discovery of New Motifs in Microbial Genomes

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Regulatory motifs can be found by local multiple alignment of upstream regions from coregulated sets of genes, or regulons. We searched for regulatory motifs using the program AlignACE together with a set of filters that helped us choose the motifs most likely to be biologically relevant in 17 complete microbial genomes. We searched the upstream regions of potentially coregulated genes grouped by three methods: (1) genes that make up functional pathways; (2) genes homologous to regulons from a well-studied species (*Escherichia coli*); and (3) groups of genes derived from conserved operons. This last group is based on the observation that genes making up homologous regulons in different species are often assorted into coregulated operons in different combinations. This allows partial reconstruction of regulons by looking at operon structure across several species. Unlike other methods for predicting regulons, this method does not depend on the availability of experimental data other than the genome sequence and the locations of genes. New, statistically significant motifs were found in the genome sequence of each organism using each grouping method. The most significant new motif was found upstream of genes in the methane-metabolism functional group in *Methanobacterium thermoautotrophicum*. We found that at least 27% of the known *E. coli* DNA-regulatory motifs are conserved in one or more distantly related eubacteria. We also observed significant motifs that differed from the *E. coli* motif in other organisms upstream of sets of genes homologous to known *E. coli* regulons, including Crp, LexA, and ArcA in *Bacillus subtilis*; four anaerobic regulons in *Archaeoglobus fulgidus* (NarL, NarP, Fnr, and ModE); and the PhoB, PurR, RpoH, and FhlA regulons in other archaeobacterial species. We also used motif conservation to aid in finding new motifs by grouping upstream regions from closely related bacteria, thus increasing the number of instances of the motif in the sequence to be aligned. For example, by grouping upstream sequences from three archaeobacterial species, we found a conserved motif that may regulate ferrous ion transport that was not found in individual genomes. Discovery of conserved motifs becomes easier as the number of closely related genome sequences increases.

Motif-finding algorithms can be used to discover alignments of sites, which correspond to transcriptional regulatory motifs in upstream regions of genes. These motifs are often binding sites for DNA-binding proteins. Several different algorithms have been used previously for motif finding, including Gibbs sampling (Lawrence et al. 1993; Liu et al. 1995), MEME (Bailey and Elkan 1995; Grundy et al. 1996), ClustalW (Thompson et al. 1994), MACAW (Schuler et al. 1991), and algorithms based on computational analysis of oligonucleotide frequencies (van Helden et al. 1998).

AlignACE (Roth et al. 1998) is a local alignment program based on the Gibbs-sampling algorithm, optimized for DNA sequence alignment. AlignACE chooses statistically significant alignments in the input sequence. However, regulatory signals often have relatively weak alignments when compared to other common genomic elements found by AlignACE, such as ribosome-binding sites and repetitive elements. Additional filters have been developed to select the motifs found by AlignACE that are the most likely to corre-

spond to biologically relevant motifs (Roth et al. 1998; Hughes et al. 2000). Together with these filters, AlignACE has been used for motif-finding in *Saccharomyces cerevisiae*, using expression data from microarrays (Roth et al. 1998; Tavazoie et al. 1999) as well as metabolic functional group categories (Hughes et al. 2000). Known *S. cerevisiae* motifs were computationally identified in these studies, as well as new motifs.

Computationally identifying upstream-regulatory motifs in bacterial genomes by AlignACE is complicated by the presence of operons. It is difficult to locate the regulatory region for a gene found within an operon, since the promoter for that operon can lie several genes upstream, and it is difficult to predict which gene is at the head of the operon. In addition, there are fewer instances of most regulatory motifs in a bacterial genome than in the *S. cerevisiae* genome, as there is usually only one instance of a regulatory motif per operon instead of one instance per gene. It is easier to discover a motif that is found in more copies in the genome. However, one can increase the number of instances of a conserved regulatory motif by pooling together upstream sequence from orthologous genes in

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closely related organisms, assuming the motif is conserved across these organisms.

A similar method was employed recently by Gelfand et al. (2000) to predict transcriptional regulatory sites in archaeal genomes by comparative genomics for four specific groups of genes. In phylogenetic footprinting analysis, noncoding sequences from several organisms are aligned to identify conserved regions (Tagle et al. 1988; Duret and Bucher 1997; Hardison et al. 1997; Wasserman and Fickett 1998). In addition, genomic comparisons of regulatory sites have been done between *Escherichia coli* and *Haemophilus influenzae* for several known motifs by using search algorithms employing position weight matrices (Robison 1997; Mironov et al. 1999).

Microarray data, as well as at least five additional methods based on comparative genomics might be used to obtain functionally linked sets of genes that are good candidates for coregulation. Pellegrini et al. (1999) describe a method for predicting functional linkages between nonhomologous genes based on the assumption that proteins that function together in the cell are likely to evolve in a correlated fashion. Marcotte et al. (1999) describe a method based on protein fusions. We obtained potentially coregulated sets of bacterial genes for motif finding by three different methods. We used groups of genes that make up functional pathways, genes homologous to the members of regulons from *E. coli*, and groups of genes derived from conserved operons. Genes with similar expression profiles will be used for motif finding in bacteria as more microbial expression data becomes available.

By aligning the upstream regions of potentially coregulated sets of genes, we were able to find many known bacterial regulatory motifs, and to predict new regulatory motifs in 17 bacterial genomes. In addition, by pooling together upstream sequences from orthologous genes in closely related organisms, we were able to find conserved motifs with only a few instances in each genome. We also analyzed conservation between organisms of motifs in orthologous sets of upstream regions.

## RESULTS

### Potentially Coregulated Groups of Genes

We searched in each species for orthologs to the *E. coli* genes known to be regulated by 55 DNA-binding proteins ([http://arep.med.harvard.edu/ecoli\\_matrices/](http://arep.med.harvard.edu/ecoli_matrices/); Robison et al. 1998; see Methods). This results in 55 potentially coregulated groups in each of 17 genomes. These groups allow us to assess our motif-finding techniques in *E. coli*, as well as to look at conservation of *E. coli* DNA motifs in other organisms and to identify potential new mechanisms for regulating the same cellular process in more distantly related organisms.

The second method we used to predict coregulated groups of genes is based on analysis of conserved operons. Functional couplings between genes can be inferred from conserved spatial proximity of gene pairs on a chromosome (Dandekar et al. 1998; Overbeek et al. 1998,1999). Two genes that are spatially separated on the chromosome in one organism, but have homologs which are spatially close in two or more other species, are good candidates for being coregulated. We can discover regulatory motifs by aligning the upstream regions of sets of genes that were predicted to be coregulated in this manner. We used 343 groups of genes from the WIT database (<http://wit.mcs.anl.gov/WIT2/>) in each of 17 complete bacterial genomes. These groups, containing between 0 and 54 genes in a particular organism, were constructed by searching for pairs of genes contained in conserved operons in >30 complete or partial genomes (Overbeek et al. 1999). In *E. coli*, most of these gene clusters contain parts of known metabolic systems, and at least 11 of these clusters are large groups that correspond to substantial pieces of known pathways (including the purine and arginine biosynthesis pathways).

We also used 68 different functional group categories in each of 17 bacterial species from the KEGG database (<http://www.genome.ad.jp/kegg/>). These groups are based on a compilation of experimental data on metabolic pathways (Ogata et al. 1999).

### Motif Discovery Strategy

We used the AlignACE program (Roth et al. 1998) together with a set of filters based on known properties of binding sites for DNA regulatory proteins to find motifs in the upstream regions of potentially coregulated sets of genes. Our method for selecting upstream-regulatory regions to be searched in genomes containing operons is described in Methods. In addition to aligning groups of upstream regions from a single organism, we also combined sequence from orthologous genes in closely related organisms (see Methods for a listing of groups).

Running AlignACE on all of these groups of genes in each of 17 organisms, as well as in sets of closely related organisms, results in 104,282 motifs. To select significant motifs that are the most likely to be functional, regulatory motifs, we calculated several indices for each matrix: MAP score, site specificity score ( $S_{site}$ ), positional bias, AT content, and palindromicity. The MAP score is a measure used by the AlignACE program to judge alignments sampled during the course of the algorithm, based on the over-representation of the motif in the input sequence (Liu et al. 1995).  $S_{site}$  is a measure of how specific a motif is for the sequence in which it was aligned, compared to the genome as a whole. The positional bias is a measure of the tendency of the top-scoring motif instances in the genome to be

unevenly distributed relative to the start codon of the closest gene (Hughes et al. 2000). A large fraction (almost half) of the footprinted *E. coli* motifs are palindromic. To identify palindromic motifs, we used the CompareACE program (Hughes et al. 2000) to compare a motif with its reverse complement. These indices are described in more detail in Methods.

Table 1 shows how many motifs score above various cutoffs in the values for these indices. The AlignACE output files, as well as the values for these indices for all of the motifs, are available on our website ([http://arep.med.harvard.edu/microbial\\_motifs](http://arep.med.harvard.edu/microbial_motifs)).

## Controls

By searching the upstream regions of genes making up the known regulons in *E. coli* with AlignACE, we can determine what fraction of the known *E. coli* DNA-binding motifs can be found, and how these known motifs score in terms of the parameters that we use to rank motifs. We used the 32 *E. coli* footprinted regulons in our database with between 5 and 100 known binding sites (Robison et al. 1998) as controls. Twenty-six of these regulons have known regulatory motifs that can be found by AlignACE (81%). The six motifs that are not found by AlignACE have low or dispersed information content (Ihf, Hns, Lrp, Fis, OmpR, and SoxS; [http://arep.med.harvard.edu/ecoli\\_matrices](http://arep.med.harvard.edu/ecoli_matrices)). Table 1 shows that by using our highest cutoffs we can

eliminate the majority of false positives, but we also only find the true positives with the strongest motifs. Without using any additional cutoffs with AlignACE, the false positive rate is very high (95%). However, by restricting our analysis to the motifs with the lowest  $S_{site}$  and those that are palindromic, we are able to make high-confidence predictions. Optimization of the choice of cutoffs for a certain false positive rate could also be performed using automatic classification schemes, such as decision trees and linear discriminant functions. Here we applied very stringent cutoffs based on simple criteria obtained from the biological properties of promoters. Table 2 lists the number of motifs from all of the AlignACE runs in all organisms that score above each cutoff.

Figure 1 shows the MAP scores and  $S_{site}$  values for the positive controls (green triangles), as well as all of the motifs found by AlignACE with MAP scores greater than 5.0 (30,252 motifs from all AlignACE runs in three kinds of gene groupings in 17 organisms). Some of the controls are plotted more than once because the motif was found multiple times. Motifs in the upper right hand corner of the plot (nonspecific motifs with good alignments) tend to be either repetitive elements (i.e., *E. coli* BIME elements) or common elements in the genome such as Shine–Dalgarno sequences. The fraction of motifs corresponding to known controls is highest in the upper left corner of the plot. Motifs

with MAP score > 10.0 and  $S_{site} < 10^{-25}$  are listed in Table 3a and 3b. Pooling together upstream regions from orthologs in several closely related organisms in order to increase the number of instances of a motif improves the ability to discriminate known conserved motifs (magenta diamonds). These conserved motifs tend to have higher MAP scores and lower  $S_{site}$  than the same motif found in sequence from *E. coli* alone (green triangles).

The most useful parameter for discriminating the known motifs in the *E. coli* regulon controls from the rest of the motifs found by AlignACE is  $S_{site}$  (see Fig. 1). Palindromicity is also a useful parameter. Almost half of the known motifs are palindromic, but only ~5% of the 104,282 motifs found in our analysis are palindromic. Thus, selecting for palindromic motifs does increase greatly the

**Table 1. Number of Controls Scoring Above Different Cutoffs**

MAP	Cutoffs <sup>a</sup>		Motifs from <i>E. coli</i> controls <sup>b</sup>	True positives <sup>c</sup>	Additional real motifs <sup>d</sup>	False positives <sup>e</sup>
	$S_{site}$	additional				
0	1	(no cutoffs)	591	26 (81%)	5	95%
5	1		299	22 (69%)	5	91%
10	1		152	22 (69%)	3	84%
5	1e-10		92	20 (63%)	4	74%
10	1e-10		56	20 (63%)	3	59%
10	1e-15		20	15 (47%)	1	20%
10	1e-20		9	9 (28%)	0	0%
10	1e-25		3	3 (9%)	0	0%
10	1e-10	pal	10	8 (25%)	2	0%
10	1e-10	pal, AT < 80	10	8 (25%)	2	0%

<sup>a</sup>Motifs where more than half of the aligned sites originate from a single input sequence were not included to eliminate repetitive elements. Minimum MAP score and maximum site specificity score ( $S_{site}$ ) are indicated. “pal” means the motif is palindromic (CompareACE score to its reverse complement is >0.7). AT is the %AT content.

<sup>b</sup>Total number of different motifs found within each *E. coli* regulon by aligning the upstream regions of known regulons in *E. coli*. Motifs found in the upstream regions of each regulon were clustered by similarity, and the total number of distinct motifs summed over all 32 regulons is given here.

<sup>c</sup>Percentage of 32 *E. coli* regulons with 5–100 known sites where the known regulatory motif can be found.

<sup>d</sup>Additional known motifs found due to overlap of regulons (i.e., the ArcA motif found upstream of the genes making up the Fnr regulon).

<sup>e</sup>Percentage of all motifs above this cutoff found in the upstream regions of footprinted regulons in *E. coli*, which are different from a motif known to regulate this groups of genes (CompareACE score greater than 0.7).

**Table 2.** Number of Motifs Scoring Above Different Cutoffs

Cutoffs <sup>a</sup>			Number of motifs found in all organisms <sup>b</sup>					
			WIT <sup>c</sup>		KEGG <sup>d</sup>	Regulons <sup>e</sup>		total
MAP	$S_{site}$	additional	indiv. <sup>f</sup>	pooled <sup>g</sup>	indiv. <sup>f</sup>	indiv. <sup>f</sup>	pooled <sup>g</sup>	
0	1	(no cutoffs)	22,134	32,760	21,677	9967	17,744	104,282
5	1		4924	4228	13,416	4696	2988	30,252
10	1		2706	1829	8484	2830	1480	17,329
5	1e-10		606	252	2035	794	219	3906
10	1e-10		316	125	1231	485	147	2304
10	1e-15		63	20	321	139	50	593
10	1e-20		7	4	90	58	24	183
10	1e-25		0	2	23	18	16	59
10	1e-10	pal	18	27	128	74	53	300
10	1e-10	pal, AT < 80	7	23	44	50	42	166

<sup>a</sup>See Table 1.<sup>b</sup>May contain multiple instances of the same motif (motifs are not clustered by similarity).<sup>c</sup>Groups derived from conserved gene adjacencies from the WIT database (Overbeek et al., 1999).<sup>d</sup>Groups derived from KEGG metabolic pathways (Ogata et al., 1999).<sup>e</sup>Groups derived from *E. coli* footprinted regulons.<sup>f</sup>AlignACE runs in individual organisms.<sup>g</sup>AlignACE runs on groups of closely related organisms pooled together.

fraction of biologically relevant motifs. The MAP score was not as useful as  $S_{site}$  for distinguishing known motifs in the *E. coli* regulon controls from the rest of the motifs, since many nonspecific chromosomal features have high MAP scores (i.e., Shine–Dalgarno sequences and repetitive elements). The positional bias statistic was also not useful in distinguishing the *E. coli* regulon controls, because most known motifs do not have significant values for our positional bias parameter (data not shown). However, there are many other motifs found in this study that do have significant positional bias. Most of these likely correspond to locationally conserved features such as the ribosome binding site (Shine–Dalgarno sequence), which is always located close to the start codon (4–13-bp upstream of ATG).

### Conservation of Known *E. coli* Motifs in Other Bacteria

In organisms other than *E. coli*, running AlignACE on upstream regions of orthologs of members of *E. coli* footprinted regulons allows for study of the conservation of *E. coli* regulatory motifs, as well as identification of other possible mechanisms for regulating the same cellular process. For each of the 34 *E. coli* regulons with more than five members, Figure 2 shows in which organisms the *E. coli* motif is conserved, and in which organisms there is a new and significant motif in the upstream regions of the homologous regulon.

A new motif can indicate either a different mechanism for regulating a similar cellular process, or divergence of binding site residues in a conserved DNA-binding protein. In *Bacillus subtilis*, there is no Crp pro-

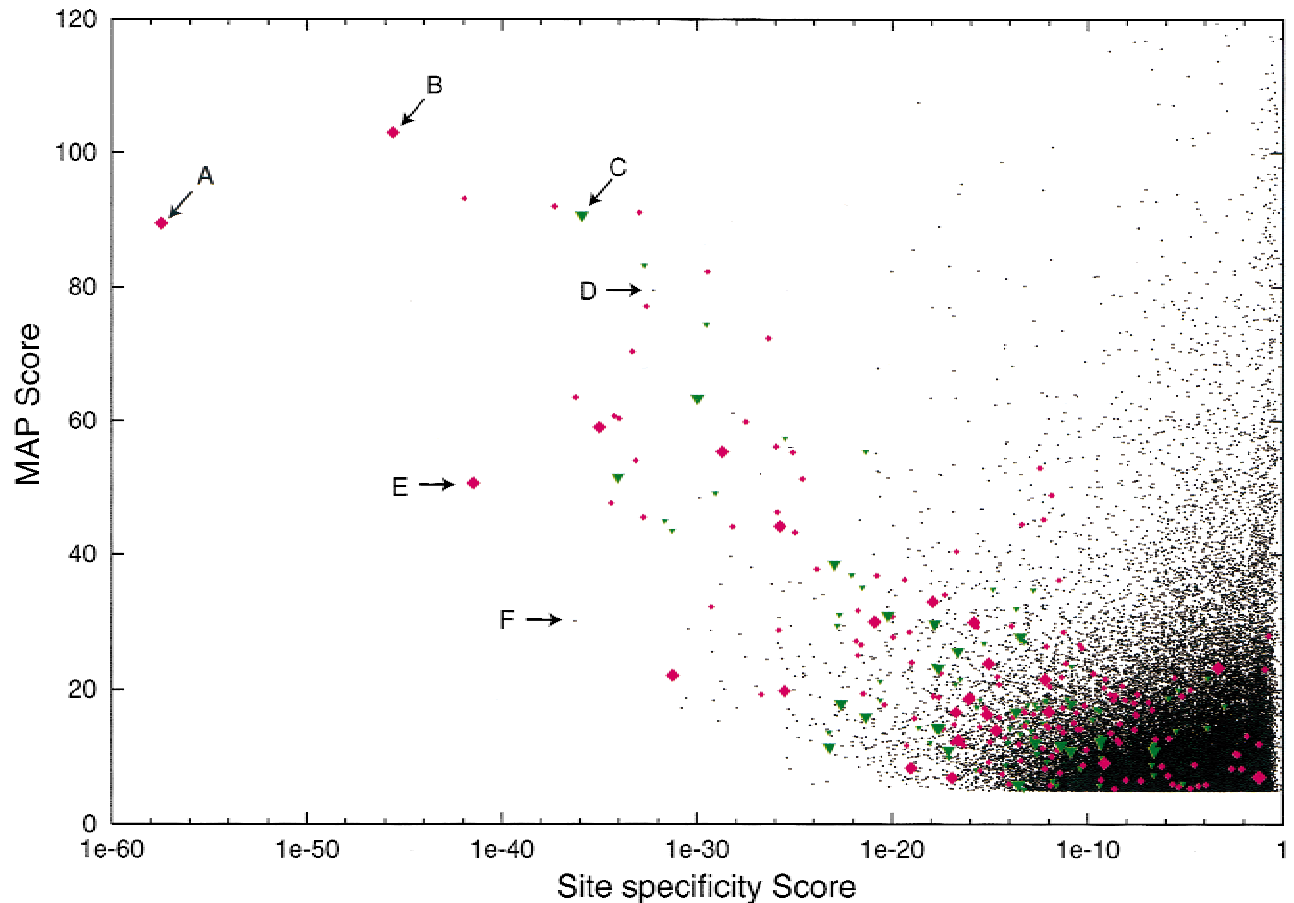
tein; instead, CcpA regulates carbon metabolism by a different mechanism (Henkin 1996). We detect the CcpA binding motif in our analysis. An example of divergence of binding site residues is LexA in *B. subtilis*. *H. influenzae* and *E. coli* have very similar LexA binding motifs. However, gram-positive bacteria, including *B. subtilis* and *Mycobacterium tuberculosis*, have a completely different binding motif for LexA (Lovett et al. 1993; Cheo et al. 1991). The LexA protein is conserved but its binding-site residues are mutated. Thus *B. subtilis* LexA binds to a completely different motif, which is found using our strategy.

Five new motifs were identified in *B. subtilis*, and ten were identified in archaeobacte-

rial species (*Methanococcus janaschii*, *Pyrococcus horokoshii*, *Methanobacterium thermoautotrophicum*, and *Archaeoglobus fulgidus*). These motifs (Fig. 2, yellow) are listed in Tables 3a and 3c. In *B. subtilis*, the CcpA motif is found in the Crp, AraC, and PhoB categories. The motif found in the *B. subtilis* ArcA category is similar to the *E. coli* –Crp motif. Therefore this could simply be a variation on the Fnr motif, which is closely related to the Crp motif in *E. coli*. The Fnr and ArcA regulons overlap significantly.

In *A. fulgidus*, there are new motifs in the categories for NarL, NarP, Fnr, and ModE. In *E. coli*, all four of these DNA-binding proteins regulate overlapping regulons related to anaerobic metabolism. The motifs from the NarL, NarP, and Fnr categories are very similar to each other (pairwise CompareACE scores < 0.7). However, the motif that is found in the ModE category is different. Thus, we predict that these two new motifs control anaerobic metabolism in *A. fulgidus* (see Tables 3a and 3c).

Combining upstream sequences from orthologous genes in closely related organisms can help in the discovery of conserved motifs with few instances in each genome. A known *E. coli* motif (MetR) is not found when AlignACE is run on the upstream regions of the members of the MetR regulon in *E. coli* alone because there are too few instances of the motif in the *E. coli* genome. However, when *E. coli* and *B. subtilis* upstream regions are pooled together, the MetR motif can be found because it is also present in *B. subtilis*. Eleven additional *E. coli* motifs can be identified in *B. subtilis* and/or *H. influenzae* using this method. Even for the



**Figure 1** All of the motifs with MAP scores  $>5.0$  (30,252 motifs) from the AlignACE runs in all 17 organisms are presented here (black points). The controls (*E. coli* motifs found in alignments of upstream regions of genes with  $>5$  footprints and which are known to constitute a regulon) are plotted with colored symbols: (green triangles) *E. coli* motifs found in alignments of upstream regions of genes making up the regulon in *E. coli* only; (magenta diamonds) conserved *E. coli* motifs found in alignment of sequence from *E. coli* together with sequence from *B. subtilis*, *H. influenzae*, or both. Several of the footprinted motifs are represented several times on this plot, as we perform two AlignACE runs on every group of genes, one for each of two different kinds of operon prediction. Also, a very strong motif can occasionally show up more than once in the same AlignACE run despite iterative masking in AlignACE. Therefore, for each control, we have represented the motif instance with the lowest value for  $S_{site}$  with a large colored symbol, and all other instances of the same motif with a small colored symbol. Several of the most specific motifs are labeled: (A) PurR conserved in *E. coli* and *H. influenzae*; (B) LexA conserved in *E. coli* and *H. influenzae*; (C) LexA in *E. coli*; (D) T-box in *B. subtilis*; (E) ArgR conserved in *E. coli*, *H. influenzae*, and *B. subtilis*; (F) new motif found in the methane metabolism functional group in *M. thermoautotrophicum*. See text and Table 3 for details.

motifs that occur frequently enough to be found in a single genome, pooling together sequence from closely related organisms increases the number of instances of the motif. This improves the values of the MAP and  $S_{site}$  parameters in the alignments, making the motif easier to identify (see Fig. 1).

### New Motifs

Table 3 shows the most specific motifs, as well as the top palindromic motifs, that result from AlignACE runs. Some of these motifs are known, and some of these correspond to potential new regulatory motifs. Only motifs scoring above stringent cutoffs are presented here; alignments and parameters for all motifs found in this analysis are available ([http://arep.med.harvard.edu/microbial\\_motifs](http://arep.med.harvard.edu/microbial_motifs)).

### Specific Motifs Found by Aligning Sequence from Individual Organisms

Table 3a displays the most specific motifs found in AlignACE runs on sequence from single organisms. Out of 53,778 motifs, the 41 motifs with  $S_{site} < 1e-25$  were clustered according to their similarity using pairwise CompareACE scores (see Methods), resulting in 18 motif clusters. The member from each cluster with the lowest  $S_{site}$  value is shown in the Table 3. Three of these clusters correspond to known *E. coli* motifs (the LexA, Crp, and PurR control sets), and one corresponds to a known *B. subtilis* motif (the T-box). The T-box is a known regulatory motif found in the functional group made up of aminoacyl-tRNA synthetases in gram-positive bacteria (Henkin et al. 1992).

The other 14 clusters correspond to new motifs.

**Table 3.** Top Motifs Found by AlignACE<sup>a</sup>

(a) Most Specific Motifs Found by Aligning Sequence from Individual Organisms

Known <sup>b</sup>	Category <sup>c</sup>	Organism <sup>d</sup>	$S_{site}$	MAP	Motif logo <sup>e</sup>
—	RpoH, 00680, 00630, 00790	TH	5.6e-37	30.1	ATATAAA TI I
LexA	LexA	EC	1.2e-36	90.7	CTG-ATA A A CAG
Crp	Crp, 00500, 00052, 00010	EC	9.2e-35	51.6	GTGA T CACA A
T-box	00970	BS	5.7e-33	79.6	AGGG-GG A C G
—	00071	AG	1.6e-31	17.3	AA A A A A
—	00310, 00061, 00120	AG	2.7e-31	18.8	TTTAA T A A
PurR	PurR	EC	1.1e-30	63.4	G-AA-CG-TTGC G
—	NarL, Fnr, 00380	AG	7.2e-30	18.5	A A TTA A T A
—	PurR	PH	1.7e-29	35.5	TTT A T A T A A
—	00970	BS	6.5e-29	61.2	GGGACG A G
—	00310, 00290	AG	5.1e-29	39.7	AAAAATTT x
—	00360	EC	1.6e-28	28.2	A GA A AAA A AA
—	modE	TH	4.0e-27	24.5	T A T A A T T A
—	00650	AG	1.6e-26	30.7	AAA A AAAAA
—	PurR, 00640	AG	4.3e-26	14.4	TTT A A AA A A
—	fhlA	TH	4.5e-26	20.0	T T T T A A A A
—	00630	TH	6.2e-26	8.3	AT A A T A A A
—	00071 (AG); 00640 (TH)	AG, TH	8.9e-26	17.3	AT T A A A A T T

(b) Most Specific Motifs Found by Aligning Sequence from Two or More Species

Known <sup>b</sup>	Category <sup>c</sup>	Organism <sup>d</sup>	$S_{site}$	MAP	Motif Logo <sup>e</sup>
PurR	PurR, GalR	EC, HI	3.7e-58	89.5	A G-AAACG-TT C
LexA	LexA	EC, HI	5.1e-38	92.0	CTG T A T A CAG
TrpR	TrpR	EC, HI	5.8e-32	22.1	G A T A T A TAG
—	103	TH, AG	5.1e-30	28.7	TTAGG T C TAAA
Crp	Crp	EC, HI	6.7e-29	44.1	GTGA T CACA T
Fur	Fur	EC, BS	3.2e-26	19.8	A T A TGA AA T T T A T
ArgR	ArgR	EC, HI, BS	8.6e-26	55.3	G A T A A A A T C A

(Continued)

The motif with the lowest value for  $S_{site}$  ( $S_{site} = 5.6e - 37$ ) for which we are not aware of any documentation in the literature is an AT-rich motif found in the methane metabolism functional group (00680) of the methane-producing archaeobacteria *M. thermoautotrophicum*. Another very specific member of this motif cluster ( $S_{site} = 3.5e - 31$ ) shows up in the closely related folate biosynthesis functional group (00790) in *M. thermoau-*

*trophicum*. Therefore this motif could regulate metabolism of one-carbon units in this organism. A similar motif also shows up with a slightly higher  $S_{site}$  in the glyoxylate and dicarboxylate metabolism functional group (00630) and the group corresponding to the orthologs of the RpoH (heat shock) regulon in *M. thermoautotrophicum*. The relationship between these two groups and metabolism of one-carbon units is not

**Table 3.** (Continued)

(c) Top Palindromic Motifs Found in Sequence from Individual Organisms

Known <sup>b</sup>	Category <sup>c</sup>	Organism <sup>d</sup>	$S_{site}$	MAP	Motif logo <sup>e</sup>
Crp	Crp, 00010, 00051, 00052, 00380, 00500, 00530, 00910, 02060 (EC); 00240, 00710 (HI)	EC, HI	9.2e-35	51.6	
LexA	LexA	EC	2.1e-33	83.3	
PurR	PurR, 00230, 001 (EC); PurR, 00230 (HI)	EC, HI	1.1e-30	63.4	
ArgR	ArgR, 054, 00220	EC	1.1e-23	38.5	
Fnr	Fnr, NarL (EC); NarL (HI)	EC, HI	2.5e-23	18.0	
CcpA, GalR	GalR (EC); Crp, AraC, PhoB, 00010, 00051, 00052, 00500, 02060 (BS)	BS, EC	4.5e-22	16.1	
Cheo	LexA	BS	6.5e-22	11.5	
Crp	Fnr, ArcA	BS	4.9e-21	27.0	
—	PurR	PH	5.7e-20	15.4	
—	156	PH	4.0e-19	20.2	
—	NarP	AG	3.2e-18	14.9	
TyrR	TyrR	EC	2.3e-18	23.3	
FruR	FruR, 00010	EC	2.6e-18	14.5	
TrpR	TrpR	EC	3.8e-16	12.4	
—	FhlA, 00230	MJ	1.2e-15	14.0	
—	255	MG	7.3e-15	16.7	
—	PhoB (PH); 00290, 00310 (AG); 00020 (BS)	PH, AG, BS	9.0e-15	11.5	
—	00330	MP	1.3e-14	18.5	
Crp	CytR	EC	2.7e-13	21.7	
—	ModE	AG	5.3e-13	11.6	
—	00330	BS	6.6e-13	32.0	
—	00260	MT	2.4e-12	41.6	
—	00240	HI	8.3e-12	14.2	
—	00061	BS	8.4e-12	13.3	
—	00260, 00251	CY	1.3e-11	25.8	
—	00950	HI	2.5e-11	10.3	
—	00910	MJ	2.9e-11	11.0	
—	00230	EC	7.3e-11	18.0	
—	033	MJ	9.6e-11	11.8	
—	199, 00061	CT	9.9e-11	23.2	

(Continued)

**Table 3.** (Continued)

(d) Top Palindromic Motifs Found by Aligning Sequence from Two or More Species

Known <sup>b</sup>	Category <sup>c</sup>	Organism <sup>d</sup>	$S_{site}$	MAP	Motif logo <sup>e</sup>
PurR	PurR, 001	EC, HI	3.7e-58	89.5	<b>A G AAACG TT C</b>
ArgR	ArgR, 054	EC, HI, BS	3.7e-42	50.6	<b>GAAT AT CA</b>
LexA	LexA	EC, HI	5.1e-38	92.0	<b>CTG ATACAG</b>
TrpR	TrpR	EC, HI	5.8e-32	22.1	<b>G AATA T A TAG</b>
—	103	TH, AG, PH	5.1e-30	28.7	<b>TTAGG CTAAA</b>
Crp	Crp	EC, HI	6.7e-29	44.1	<b>GTGA TACA</b>
GalR	GalR	EC, HI, BS	1.8e-26	44.2	<b>G AA CG TT CA</b>
Fur	Fur, 101	EC, BS	3.4e-22	19.4	<b>A T A TG AA TT</b>
TyrR	TyrR	EC, HI, BS	1.0e-19	24.1	<b>TGTAAA TACA</b>
—	OxyR	EC, HI, BS	1.9e-19	17.4	<b>AT AT T AT</b>
ModE	ModE	EC, HI	2.5e-17	12.6	<b>T T A TA ATA</b>
HipB	HipB	EC, HI	4.3e-17	16.1	<b>T TCG A A AA</b>
ArcA	ArcA, Fnr	EC, HI	5.9e-17	10.6	<b>TGTGAA TA</b>
—	190	EC, BS	1.1e-14	11.3	<b>A AC TA ATATAG</b>
—	DnaA	EC, BS	4.7e-12	14.8	<b>A CAAT TGTG</b>
Fnr	Fnr, NarP, narL	EC, HI, BS	5.2e-12	19.1	<b>TGA T A CAA</b>
—	199	MJ, PH	1.2e-11	24.7	<b>G GG TT AA CC</b>
—	073	EC, HI	4.2e-11	16.6	<b>CG ACAAT</b>
—	009	EC, BS	6.4e-11	16.8	<b>AA ATTG</b>

<sup>a</sup>The top motifs were clustered according to their pairwise CompareACE scores, and the most specific member of each cluster is listed here (see Methods).

<sup>b</sup>If this motif has been previously documented, its name is given.

<sup>c</sup>The potentially coregulated group of genes in which this motif was found by AlignACE. For motifs found in categories derived from *E. coli* footprinted regulons, the name of the *E. coli* DNA-binding protein that regulates that regulon is listed. For motifs found in categories derived from conserved operons and KEGG functional group categories, either a 3-digit or 5-digit number is listed, respectively. These correspond to the following groups:

KEGG functional group categories (84 total groups):

00010	Glycolysis/gluconeogenesis	00360	Phenylalanine metabolism
00020	Citrate cycle (TCA)	00380	Tryptophan metabolism
00051	Fructose and mannose metabolism	00500	Starch and sucrose metabolism
00052	Galactose metabolism	00530	Aminosugars metabolism
00061	Fatty acid biosynthesis	00630	Glyoxylate and dicarboxylate metabolism
00071	Fatty acid metabolism	00640	Propanoate metabolism
00120	Bile acid biosynthesis	00650	Butanoate metabolism
00220	Urea cycle and amino group metabolism	00680	Methane metabolism
00230	Purine metabolism	00710	Carbon fixation
00240	Pyrimidine metabolism	00790	Folate biosynthesis
00251	Glutamate metabolism	00910	Nitrogen metabolism
00260	Glycine, serine, and threonine metabolism	00950	Alkaloid biosynthesis I
00290	Valine, leucine, and isoleucine biosynthesis	00970	Aminoacyl-tRNA biosynthesis
00310	Lysine degradation	02060	Phosphotransferase system (PTS)
00330	Arginine and proline metabolism		

Groups derived from conserved operons in WIT (343 total groups):

001	Purine biosynthesis	103	Ferrous ion transporter
009	Nucleotide biosynthesis	156	HYP operon
033	Pyruvate synthase	190	Ribonucleoside diphosphate
054	Arginine biosynthesis	199	Transcription
073	Fatty acid biosynthesis	255	Heat shock
101	Iron transport		

<sup>d</sup>See Methods for abbreviations.

<sup>e</sup>The Motif Logo (Schneider and Stevens, 1990). The height of the stack of letters is proportional to the information content, and the relative frequency of each base is given by its relative height. A sequence span containing the ten strongest positions of the logo is displayed.

	EC	HI	BS	IMT	IRP	HP	BB	TP	CT	ICY	MP	MGAA	IMJ	PH	TH	AG	
Crp	77*	93*	78*	43*	12	19	14	14*	7	33*	16	14	16	15	23	16	19*
NarL	41*	30*	20*	17*	8	4	1	3	5	11*	2	1	20	6*	7	6	25*
PhoB	52*	6*	34	37*	3	3*	7	3	2	33*	11	9	8*	10	13	11	9
NarP	29*	23*	10*	6*	7	2	1	3	2	7*	2	0	13	3	3	3	15
PurR	28*	13*	29*	22*	3	13	3	3	6	27	2	2	22	19	19	18	20*
Fnr	26*	16*	21	16	3	4	1	1	4	8*	0	1	13*	7	6*	6	14
FruR	22*	11	18	7*	3	8	11	6	5	10	10	8	5	5	9	4	4
RpoN	21*	3	22*	11	4	8*	3*	4*	4*	10	5*	3*	16*	14*	14	16	8
LexA	20*	0*	12*	12*	9	8	8	10	7	13*	9	8	5	3	4*	8	3*
RpoH	19*	18*	20	19	17*	18	19	16	15	30	12	11	19	10	9*	13	14
CoxR	18*	8*	24*	23*	14*	9*	5	8	1	18*	0	0	4*	0	3	2	4
AraC	15*	6*	20*	16*	10*	7	2	1	5	11*	3	1	8	5	6	8	9
Fur	13*	5*	10*	13*	10	2*	1	0	2*	5*	0	0	4*	1	2	2	1*
PhiA	14*	1	1*	6	0	4*	0	0*	1	6	0*	0*	4*	10	11	10	4*
GlpR	13*	12*	12*	6	3	1	3	3	4	5*	2	4	3	2	2	4*	4
ArgR	12*	5*	14*	10*	0	2	2	0	0	10	3	0	11	12	9	9	12
FinCD	12*	0	9	2	0	8	7	8	8	0	0	0	7	2	0	0	2
TyrR	11*	6*	8*	0	0	1	0	3	5	1	1	0	0	2	1	2	1
ModE	11*	10*	9	11	0	7	0	0	0	8	0	0	10	5	8	8*	13*
GalP	10*	9*	13*	4	1	1	2	5	2*	3	3	2	1	5	6	4*	3
NagC	10*	4	14*	4	2	2	4	1	2	4	1	1	3	3	3	2	2
AraC	10*	1	13	1	0	1	1	2	1	2	1	0	0	0	1	0	1
SoxS	10*	4	8	5	1	2	2	2	4	3	2	1	2	0	3	2	0
TrpR	9*	8*	6	7	0	6*	0	3	5*	6	0	0	5	6	2	5	4
MalT	9*	2	10	8*	0	3	4	4	2	6	3	3	3	5	8*	2	2
CytR	8*	4	9*	5	0	3	1	3	0	2	4	4	1	2	2	2	5
Fis	8*	8*	5	2	7	7	5	3	6	6	0	0	8	4	4	3	3
PdhR	7*	3	5	4	3	0	0	0	3	3	2	2	2	0	5*	1	3
DnaA	6*	6*	3*	3*	6*	5*	2*	4*	5*	5*	1*	2*	3*	1	4*	0	1
MetJ	6*	5*	2	1	0	0	0	1	1	1	0	1	2	1	3	1	3
NtrC	6*	2	18*	5	4*	4*	3*	4*	3*	4	5	3	10*	4	3	8	4
FadR	6*	3*	5	14	1	1	2	1	0	1*	1	0	2	0*	3	2	14*
MetR	6*	4*	4	4	2	1	1	2	2	2	1	2	3	3	3	3	5
RnaS	6*	0	5*	0	0	1	1	0	0	1	0	0	1	0	1	0	0

**Figure 2** This figure summarizes the results in each organism from AlignACE runs on groups derived from *E. coli* footprinted regulons. The 34 known *E. coli* DNA-binding proteins that regulate at least five genes in *E. coli* are listed in the left column. The numbers indicate how many regulon members are present in that organism. (\*) Ortholog to the *E. coli* DNA-binding protein was found. (Red or pink square) Motif similar to the known *E. coli* motif (CompareACE score >0.7) is conserved in this organism and was found by AlignACE with MAP score >5 and  $S_{site} < 1e - 10$ . (Red square) Motif was found by aligning sequence from this organism only; (pink square) motif was found by aligning sequence from this organism together with sequence from *E. coli* (>30% of the aligned instances of the motif were from this organism). (Yellow square) New and significant motif found in sequence from this organism that is not similar to the *E. coli* motif. Yellow squares correspond to the motifs described in Tables 3a and 3c that were found in groups derived from *E. coli* regulons (i.e., these motifs score better than one of two cutoffs: (1)  $S_{site} < 1e - 25$ , MAP score >10; (2)  $S_{site} < 1e - 10$ , MAP score >10, palindromicity >0.7, % AT <80%). (White square) No new motif that scores better than these cutoffs, and the *E. coli* motif is not conserved in this organism.

clear. This motif resembles the central AT-rich core (ATATAAxxTT) of the known archaeal heat-shock promoter (Thompson and Daniels 1998; Gelfand et al. 2000). It has a CompareACE score of 0.72 when compared to this known heat shock motif (consensus TTCTATATAAxxTTTCG). Another new motif in this list is the motif discussed earlier which we predict to regulate anaerobic metabolism in *A. fulgidus* (see Fig. 2). It is not clear why this motif is also found in the tryptophan metabolism functional group (00380) in *A. fulgidus*. Another very specific motif is the AT-rich motif found in the PurR (purine biosynthesis) group in *Pyrococcus horokoshii*. This motif shows simi-

larity to the purine biosynthesis motif found by Gelfand et al. (2000) in three *Pyrococcus* genomes, except that the highly conserved C and G positions are not present.

Many of these 14 new clusters contain AT-rich archaeobacterial motifs that are very specific to the groups in which they are found. There are no known *E. coli* motifs with AT content greater than 80%. However, not much is known about regulatory mechanisms in the archaeobacteria, so these could be functional regulatory elements. These are not AT-rich genomes (*A. fulgidus* and *M. thermoautotrophicum* both have 51% AT content).

#### Specific Motifs Found by Combining Sequence from Closely Related Organisms

Of the 50,504 motifs found by aligning sequence from closely related organisms together, there are 18 motifs with  $S_{site} < 1e - 25$ , which fall into seven distinct clusters (Table 3b). Six of these clusters correspond to known *E. coli* motifs conserved in either *H. influenzae* or *B. subtilis* (PurR, LexA, TrpR, Crp, Fur, and ArgR). Many of these conserved motifs show up with lower values for  $S_{site}$  and higher MAP scores than when sequence from only one organism containing the motif is aligned, because a stronger alignment can be obtained when more instances of the motif are present. Therefore, more known *E. coli* motifs are present in Table 3b than in Table 3a, which contains alignments of sequence from *E. coli* only.

The only motif in this list that has not been documented previously is found in a group derived from conserved operons (group 103) in *A. fulgidus*, *M. thermoautotrophicum*, and *P. horokoshii*. In *A. fulgidus*, the motif is found upstream of two genes with homology to ferrous ion transporters; in *M. thermoautotrophicum*, the motif is found upstream of a ferrous ion transporter and a gene with homology to the iron repressor; and in *P. horokoshii*, the motif is found upstream of a ferrous ion transporter (see Table 3d). This motif is highly palindromic (consensus TTAGG-x4-CCTAA). This pattern of two palindromic halvesites separated by a short linker sequence is common among the binding sites for known bacterial regulatory DNA-binding proteins. Our prediction is that this motif is a binding site for a protein regulating iron transport in these archaeobacteria. Since the motif is found upstream of a putative iron repressor in *M. thermoautotrophicum*, it is possible that this putative repressor (MTH214) is the regulatory protein that binds to these sites, and that it is autoregulatory.

#### Top Palindromic Motifs Found by Aligning Sequence from Individual Organisms

Palindromicity is a parameter that is strongly correlated with regulatory function. Only about 5% of all of

the motifs found by AlignACE in this study were palindromic, whereas almost half of all of the known motifs are palindromic. By considering palindromic motifs separately, we can increase the  $S_{site}$  cutoff and retain a high rate of true positives (see Table 1). A selection of these motifs is shown in Table 3c. All 101 palindromic motifs with MAP >10,  $S_{site} > 1e - 10$ , and AT-content <80% were clustered into 30 clusters. One representative from each cluster is shown here. If we do not impose the AT-content cutoff, there are over twice as many motifs present (220 motifs). Ten of these 30 clusters correspond to known *E. coli* or *H. influenzae* motifs, and two correspond to known *B. subtilis* motifs. The known *E. coli* motifs scoring above this cutoff are Crp, LexA, ArgR, Fnr, TyrR, FruR, TrpR, and GalR. The known *B. subtilis* motifs are the Cheo motif (the *B. subtilis* SOS box), and the CcpA motif. In *B. subtilis* there is also a variant of the Fnr motif that resembles the *E. coli* Crp motif. ArgR and PurR are also found above this cutoff in groups derived from conserved operons (groups 054 and 001, which contain parts of the purine and arginine biosynthesis pathways, respectively). Of the 22,134 motifs found by running AlignACE on the upstream regions of all 343 gene groups predicted from conserved operons, the ArgR and PurR motifs in *E. coli* are among the most specific palindromic motifs.

Eighteen of the 30 motif clusters in Table 3c show no similarity to known motifs. Thirty percent of these motifs are also AT rich (70%–80% AT content). Three of these motifs are found in groups derived from conserved operons. These are group 156 (Hyp operon genes) in *P. horokoshii*, group 255 (heat shock genes) in *M. genitalium*, and group 033 (pyruvate synthase genes) in *M. janaschii*. The presence of regulatory motifs upstream of the genes making up these groups lends additional support to the hypothesis that these are indeed functionally coupled groups of genes. Since this method for predicting regulons is based purely on the chromosomal gene order across genomes, the high-scoring motifs found in these groups are not biased toward well-studied organisms. In contrast, the other kinds of groups of potentially coregulated genes that we used in this study (groups from metabolic pathways and groups based on footprinted *E. coli* regulons) both originate from experimental information determined in the well-studied organisms, so the high-scoring motifs from these two methods are largely from *E. coli* and *B. subtilis*.

The motif found upstream of group 255 (heat shock genes) in *M. genitalium* does not resemble the CIRCE (Controlling Inverted Repeat of Chaperone Expression) motif, which is known to regulate several heat shock-related genes in a wide variety of organisms, including *M. genitalium*, through the binding of the repressor HrcA (Naberhaus 1999). Since HrcA is the

only transcription factor known to be involved in regulating heat shock genes in *M. genitalium*, it is not clear what transcription factor could bind to our predicted motif. This motif could represent a false positive since it is also not found in the closely related *Mycoplasma pneumoniae*. A motif resembling the CIRCE element is found upstream of this group of heat shock genes in *M. genitalium*; however, it did not score below our stringent specificity score cutoff ( $S_{site} = 2.9e - 7$ ).

#### Top palindromic motifs found by combining sequence from closely related organisms

Using this same cutoff ( $S_{site} < 1e - 10$ , MAP >10, AT content < 80%) on the motifs obtained from aligning upstream sequence from orthologous genes in two or more closely related organisms together, we obtain 65 motifs that reduce to 19 clusters (Table 3d). Of these 19 clusters, 12 correspond to known *E. coli* motifs conserved in either *H. influenzae* or *B. subtilis* (PurR, ArgR, LexA, TrpR, Crp, GalR, Fur, TyrR, ModE, HipB, ArcA, and Fnr). Again, these conserved motifs show up with lower  $S_{site}$  and higher MAP scores when they are aligned in multiple organisms containing the same motif because there are more instances of the motif to align.

Of the seven clusters corresponding to new motifs, one is the ferrous ion-transport motif in *A. fulgidus*, *M. thermoautotrophicum*, and *P. horokoshii* described above. Four of the remaining six clusters correspond to conserved motifs found in groups derived from conserved operons. These are group 190 (ribonucleoside diphosphate) in *E. coli* and *B. subtilis*, group 199 (transcription) in *M. janaschii* and *P. horokoshii*, group 073 (fatty acid biosynthesis) in *E. coli* and *H. influenzae*, and group 009 (nucleotide biosynthesis) in *E. coli* and *B. subtilis*.

## DISCUSSION

We found many significant new motifs in 17 bacterial genomes. Known regulons in *E. coli* were used as controls to calibrate the significance of these motifs. More motifs were found in larger genomes with more complex regulation. Some of the highest-scoring new motifs are found in archaeobacteria, for which there is relatively little experimental data because of difficulties in performing experiments in these organisms. New, significant motifs include two motifs potentially regulating anaerobic metabolism in *A. fulgidus*, a motif potentially regulating methane metabolism in *M. thermoautotrophicum*, and a palindromic motif regulating ferrous ion transport that is conserved in *M. thermoautotrophicum*, *A. fulgidus*, and *P. horokoshii*.

We also identified a number of motifs that are conserved in several eubacterial species. At least 22% of the known *E. coli* DNA regulatory motifs are conserved in *H. influenzae*. In the more distantly related organism *B.*

*subtilis*, at least 15% of the known *E. coli* motifs are conserved. In even more distantly related organisms, motifs can differ considerably. We found cases in which there are different but significant motifs upstream of genes homologous to known *E. coli* regulons, including Crp, LexA, and ArcA in *B. subtilis*; four anaerobic regulons in *A. fulgidus* (NarL, NarP, Fnr, and ModE); and PhoB, PurR, RpoH, and FhlA in other archaeobacterial species. This can indicate that the organisms have evolved different methods for regulating the same cellular processes, or it can indicate parallel mutations in the DNA-binding protein and the motif that it recognizes.

The three methods of predicting regulons that we use here are based on different sources of biological information. The gene groups obtained from homologs to members of known *E. coli* regulons and the groups based on functional pathways in each organism are based on the prior body of knowledge from biological experiments. Therefore, many known motifs are found in these groups. In contrast, the groups derived from conserved operons in other organisms are not based on any biological information other than the positions of genes within the genomic sequence. Therefore, these groups are less biased towards motifs that were already known. However, some known motifs are found in these groups as some of the known regulons can be reconstructed in this manner (i.e., ArgR and PurR). By our criteria for ranking motifs, the PurR and ArgR motifs are two of the top 10 motifs to come out of the AlignACE runs on the groups derived from conserved operons, which lends credibility to the use of this method for predicting regulons and their regulatory motifs.

The usefulness of motif finding upstream of potential regulons depends strongly on how good the regulon predictions are. The groups of genes that were constructed based on conserved operons in other organisms were limited in several ways. The first limitation is that only top reciprocal FASTA hits were considered orthologs (Overbeek et al. 1999), which is a restrictive ortholog definition. When groups of orthologs are constructed across many organisms, and one or more organism contains close paralogs, actual orthologs will be missed or split into separate ortholog groups. This ortholog definition also causes problems in the case of multidomain protein fusions, which can be orthologous to multiple nonhomologous proteins in another species. Thus, using a method that treats multidomain protein fusions as multiple separate entities and employs a looser ortholog definition will increase the size of the functionally related gene clusters that can be predicted. Including multidomain protein fusions as separate entities will also extend the usefulness of this approach to include eukaryotic genomes. The second limitation in the way the groups were constructed is

that only tandemly oriented genes were considered to be potentially functionally related (Overbeek et al. 1999). However, divergently transcribed genes are also good candidates for coregulation in prokaryotes (Robison 1997), as well as in eukaryotes (Smith and Zhang 1998). Thus, a looser operon definition that also includes divergently transcribed genes should increase the usefulness of this approach for predicting coregulated sets of genes.

These methods will continue to become more powerful as the amount of available sequence data increases. Overbeek et al. (1998) suggest that the number of ortholog pairs contained in conserved operons increases roughly as the square of the number of organisms included in the analysis. This method of predicting regulons based on conserved operons will work best when using sequence from a variety of distantly related organisms, including representatives from all families of bacteria. A conserved operon is more likely to represent a functional coupling when it has been conserved across a large evolutionary distance. However, motif finding in bacteria using these predicted regulons will be most effective when there are many closely related genomic sequences available. As we have seen, regulatory motifs conserved more often in closely related genomes. With more bacterial genomes, upstream sequences from large groups of closely related bacteria can be pooled together, increasing the number of instances of conserved motifs.

The three methods that we use separately for predicting coregulated sets of genes can be combined to obtain larger and more complete groups. Groups obtained by these three methods can also be combined with groups of genes experimentally observed to have similar expression profiles, as well as groups obtained by methods such as that of Pellegrini et al. (1999), assuming that proteins that function together in the cell are likely to evolve in a correlated fashion. Combining different methods for predicting coregulated sets of genes, together with alignment of upstream regions of these groups using AlignACE, should be a very powerful method for predicting functionally related and coregulated sets of genes, discovering the upstream regulatory motifs controlling these groups, and generating hypotheses concerning gene regulation.

The biological significance of some of the motifs presented here should be verified experimentally, including determination of factors binding to these motifs. Predictions for which DNA-binding protein might be interacting with the motif can be obtained by computational methods, such as finding which predicted DNA-binding proteins have the motif in their upstream region (assuming autoregulation), and searching for a member of a known DNA-binding protein family that is linked to the regulon via a conserved operon in another organism. The regulon prediction

and motif discovery methods described here should be an increasingly powerful addition to the current array of tools used to elucidate connectivities in bacterial regulatory networks.

## METHODS

### Organisms

AlignACE runs were performed on upstream regions from the following 17 bacterial organisms (abbreviations used in the tables and figures are given in parentheses): *A. fulgidus* (AG), *Aquifex aeolicus* (AA), *Borrelia burgdorferi* (BB), *B. subtilis* (BS), *Chlamydia trachomatis* (CT), *E. coli* K12 (EC), *H. influenzae* (HI), *Helicobacter pylori* (HP), *Mycoplasma genitalium* (MG), *M. janaschii* (MJ), *Mycoplasma pneumoniae* (MP), *M. thermoautotrophicum* (TH), *M. tuberculosis* (MT), *P. horokoshii* (PH), *Rickettsia prowazekii* (RP), *Synechocystis sp.* (CY), and *Treponema pallidum* (TP). For AlignACE runs on groups of genes derived from conserved operons, sequence from the following 12 groupings of closely related organisms were pooled together: EC and HI; EC and BS; EC, HI, and BS; BS and MT; EC and MT; AG and TH; AG, TH, and MJ; AG, TH, Mji, and PH; MJ and PH; MG and MP; CY and CT; and TP and BB. For AlignACE runs on upstream regions of groups of genes derived from *E. coli*-footprinted regulons, 16 groups are constructed by pooling *E. coli* sequence together with sequence from each organism separately, including distantly related organisms. Two additional groups were also used: EC, HI, and BS; and BS and MT.

### Identification of Orthologs

In each organism, we searched for orthologs to the members of the footprinted-*E. coli* regulons ([http://arep.med.harvard.edu/ecoli\\_matrices](http://arep.med.harvard.edu/ecoli_matrices)), as well as for orthologs to the *E. coli* DNA-binding proteins controlling these regulons. To identify potential orthologs, we performed reciprocal BLAST searches between *E. coli* and each of 16 other completely sequenced organisms. To not discount closely related paralogs from our analysis, we allowed up to five potential orthologs for each gene to be included. It is desirable to include upstream regions from several potential orthologs in the alignments because AlignACE can tolerate some superfluous sequence, and we want to make sure that the correct ortholog is included; however, if too much extra sequence is added, the real motif will not be found.

To find potential orthologs in genome  $G_b$  for a gene  $x_a$  in genome  $G_a$ , we performed a BLAST search over all genes in genome  $G_b$  using  $x_a$  as a query. We identified the raw BLAST score of the top hit in genome  $G_b$ , and selected up to five hits in genome  $G_b$  with raw score >70% of this value. For each of

these genes  $x_{bi}$  in genome  $G_b$ , we performed a BLAST search over all genes in genome  $G_a$ . If the original gene  $x_a$  turned up with a raw score >70% of the value of the top BLAST hit in genome  $G_a$ , then  $x_a$  and  $x_{bi}$  are potential orthologs.

### Identification of Upstream Regulatory Regions

If a gene lies within an operon, its promoter and regulatory region could lie several genes upstream. It is difficult to predict the first gene in an operon, especially in less well-studied organisms. To ensure that the sequence we align contains the regulatory intergenic region, we must include several possible intergenic regions. However, if we add too many extra intergenic sequences, the regulatory motif will not be found by AlignACE because there will be too much noise.

Our definition of an operon is two or more tandem genes separated by less than a certain cutoff distance. We used two different operon cutoff distances (100 and 300 bp). For each operon defined in this manner, we recorded the entire sequence of all of the intergenic segments of length greater than 20 bp between the gene of interest and the operon head, as well as 300 bp upstream of the operon head (see Fig. 3). We ran AlignACE twice for each group of potentially coregulated genes: once using a loose distance cutoff in the operon definition (300 bp) to ensure inclusion of the correct upstream region, and once using a more conservative distance cutoff (100 bp) to reduce inclusion of extraneous intergenic regions.

To increase the signal to noise ratio, we took no more than 300 bp of sequence upstream of the operon head because the overwhelming majority of the binding sites for DNA-binding proteins in bacteria are found within the first 300 bp upstream of the start codon. In cases where there is a site further upstream, there is usually also a site close to the promoter (Gralla and Collado-Vides 1996). We looked only at noncoding regions, because most binding sites for DNA-binding proteins are found within noncoding regions. Thus, we excluded regulatory sites downstream of start codons, as well as rare sites far upstream of the start codon.

### AlignACE Runs

The AlignACE (Roth et al. 1998), ScanACE, and CompareACE (Hughes et al. 2000) programs are available at <http://arep.med.harvard.edu/mrnadata/mrnasoft.htm>. Default AlignACE parameters were used, except that the expected number of sites was set to five, and the fractional GC content was set to the proper value for each genome. A total of 14,863 AlignACE runs took two months of CPU time on eight 300–400 MHz Pentium II processors.

Each motif found in this analysis was compared to the 55 footprinted *E. coli* motifs (Robison et al. 1998) using the CompareACE program (Hughes et al. 2000). CompareACE



**Figure 3** Example of upstream region prediction. This shows the predicted upstream region for gene *E* taking up to 300-bp of sequence upstream of the predicted operon head, and using two different cutoffs for operon prediction (300 and 100 bp). First the algorithm checks the length of the upstream region for the gene in question. If an intergenic region is shorter than the distance cutoff, then the entire intergenic region is stored for motif finding and the next intergenic region further upstream is considered as well. This continues until an intergenic region is encountered that is either divergently transcribed, or longer than the distance cutoff.

finds the best alignment between two motifs and calculates the correlation coefficient between the two position-specific scoring matrices. Two motifs with a CompareACE score >0.7 were considered to be similar. This cutoff was determined by looking at sequence logos and alignments for motifs similar to the known *E. coli* motifs, but containing slightly different sets of aligned sites. Most variants of the same motif had CompareACE scores > 0.7 when compared to the original motif alignment. Only 0.8% of 100,000 randomly selected pairs of motifs have CompareACE similarity scores > 0.7.

The CompareACE program was also used to compare and cluster new motifs coming out of the analysis. A matrix of all pairwise CompareACE scores was calculated, and then motifs were clustered using a simple joining algorithm (Hartigan 1975). The member of each cluster with the best value for  $S_{site}$  is shown in Table 3.

## Parameters Used in Motif Analysis

### Site Specificity Score ( $S_{site}$ )

$S_{site}$  is a measure of how specific a motif is for the sequence in which it was aligned. This statistic is similar to the specificity score described by Hughes et al. (2000), but modified to better accommodate bacterial motifs, which are located upstream of operons rather than individual genes and are often found in multiple copies within a single regulatory region.

For each motif, we constructed a position-specific weight matrix and searched for additional instances of the motif in the whole genome. We used the Berg and von Hippel weight matrix (Berg and von Hippel 1987, 1988; Berg 1988a, 1988b), implemented in the ScanACE program (Hughes et al. 2000) to perform this search, and saved the best 200 sites. Of these top 200 sites, we calculated the number of sites that are located within the upstream regions used to align the motif. Then we calculated the probability of obtaining this number of sites or more within this particular subset of the genomic sequence by chance using the hypergeometric distribution:

$$S_{site} = \sum_{i=x}^{\min(s_1, s_2)} \frac{\binom{s_1}{i} \binom{N-s_1}{s_2-i}}{\binom{N}{s_2}}$$

$N$  is the total number of possible sites (the total number of base pairs in the genome(s) considered),  $s_1$  is the number of ScanACE hits considered (here we are using the 200 highest-scoring hits in the genome),  $s_2$  is the total number of possible sites in the set of upstream regions used to align the motif (equal to the number of base pairs in the sequence input to AlignACE), and  $x$  is number of the top 200 ScanACE hits that fall within the upstream regions input to AlignACE.

### AT Content

Many of the motifs that were found by AlignACE, including motifs with low values of  $S_{site}$ , are AT rich (>90% AT content). However, no known matrices for *E. coli* DNA-binding proteins have AT content greater than 80% (Robison et al. 1998). We are also not aware of any known motifs in other organisms, including archaeobacterial genomes (Gelfand et al. 2000), with AT content > 80%. Thus, we explored the use of this measure to exclude AT-rich motifs that do not resemble known regulatory motifs. However, many high-scoring motifs in less well-studied organisms, including archaeobacteria, are AT rich. Since we know very little about binding sites for DNA regula-

tory proteins in these bacteria, we did not exclude the most specific AT-rich motifs.

### Palindromicity

To select palindromic motifs for further analysis, we used the CompareACE program to compare a motif with its reverse complement. We used the same CompareACE cutoff score for comparing motifs to one another (0.7).

### Additional Cutoffs used in Selecting Interesting Motifs

To exclude repetitive elements from our analysis, we excluded motifs in which more than half of the aligned sites came from a single upstream region. In the AlignACE runs combining sequence from several closely related organisms, we only looked at those motifs where < 70% of the aligned instances of the motif came from a single organism, in order to limit our analysis to conserved motifs.

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