

Topological units of environmental signal processing in the transcriptional regulatory network of *Escherichia coli*

G. Balázsi^{†‡}, A.-L. Barabási[§], and Z. N. Oltvai^{†¶}

[†]Department of Pathology, Northwestern University, Chicago, IL 60611; [§]Department of Physics and Center for Complex Networks Research, University of Notre Dame, Notre Dame, IN 46556; and [¶]Department of Pathology, University of Pittsburgh, Pittsburgh, PA 15261

Edited by Charles R. Cantor, Sequenom, Inc., San Diego, CA, and approved April 12, 2005 (received for review January 21, 2005)

Recent evidence indicates that potential interactions within metabolic, protein–protein interaction, and transcriptional regulatory networks are used differentially according to the environmental conditions in which a cell exists. However, the topological units underlying such differential utilization are not understood. Here we use the transcriptional regulatory network of *Escherichia coli* to identify such units, called *origons*, representing regulatory subnetworks that originate at a distinct class of sensor transcription factors. Using microarray data, we find that specific environmental signals affect mRNA expression levels significantly only within the *origons* responsible for their detection and processing. We also show that small regulatory interaction patterns, called *subgraphs* and *motifs*, occupy distinct positions in and between *origons*, offering insights into their dynamical role in information processing. The identified features are likely to represent a general framework for environmental signal processing in prokaryotes.

cellular networks | regulation | transcription

Transcriptional regulatory (TR) networks govern cellular life by initiating and mediating gene expression in response to environmental and intracellular cues that result in the execution of cellular programs such as metabolic adjustments, sporulation, or cell division. The nodes of this network are transcription units (genes or operons) together with their protein products, whereas the links connecting them correspond to TR interactions mediated by transcription factor (TF) proteins. The genome-scale identification of TF-binding sites, their binding specificities, and their condition-dependent utilization (1–7) results in increasingly comprehensive data sets amenable for analysis of TR-network topology and function.

Previous studies analyzing the TR-network topology of the prokaryote *Escherichia coli* and the eukaryote *Saccharomyces cerevisiae* have demonstrated that their TR networks share several characteristics such as the exponential distribution of in-degree connectivity, the scale-free distribution of out-degree connectivity, and the very low number of feedback circuits except for self-regulation (8–11). In addition, the same small-scale connectivity patterns [e.g., the feed-forward loop (FFL) and bifan motif] are overrepresented in both TR networks (12–15), suggesting that their topology has evolved to accomplish similar tasks in various organisms (14). Recent studies of motif dynamics (16–17) generated the first insights into their information-processing capabilities, although the position of motifs within TR networks and their aggregation into larger topological structures (10) may modify their dynamic behavior.

Despite these advances, there is a clear need to decipher the system-level organization of dynamic TR-network utilization (5) triggered by a various environmental and intracellular cues. Here, based on the inherent directionality of TR interactions in *E. coli* (11, 12), we identify topological units of environmental signal processing (called *origons*) as TR subnetworks originating at a distinct class of TFs. Using microarray data, we demonstrate that environmental signals affect significantly only the *origons* rooted at sensor TFs

specialized for their detection. We also show that small-scale regulatory interaction patterns (subgraphs and motifs) occupy distinct positions in and between *origons* that, together with their filtering properties, are suggestive of their role in information processing. Taken together these results suggest that *E. coli* uses specific topological units of its TR network to detect the elementary components (modes) of complex environmental signals and subsequently develop a response by reassembling these elementary modes near the output layer of the network.

Methods

Databases and Software. The publicly available data on the TR network of *E. coli* MG1655 (www.weizmann.ac.il/mcb/UriAlon/Network_motifs_in_coli/ColiNet-1.1) (12), based predominantly on the RegulonDB database (18), was used for our work. The network originally contained 423 operons and 578 regulatory interactions, reduced to 418 operons and 519 links by the removal of all operons that have only autoregulatory links.

We downloaded microarray data from two public sources: A Systematic Annotation Package for Community Analysis of Genomes (ASAP) at the University of Wisconsin (Madison) (<https://asap.ahabs.wisc.edu/annotation/php/ASAP1.htm>) and the Oklahoma University Microarray Core Facility database (<http://chase.ou.edu/macro/runs.php>). The ASAP database contained 91 experiments on *E. coli* MG1655 transposon insertion mutants (as of June 2004), 50 using cDNA microarrays and 41 using Affymetrix (Santa Clara, CA) oligoarray platforms and 41 aerobic-shift experiments on *E. coli* K-12 MG1655 using Affymetrix oligoarray platforms; the Oklahoma University database contained 104 experiments performed on *E. coli* cells in various conditions.

For network representation we used the program PAJEK (<http://vlado.fmf.uni-lj.si/pub/networks/pajek>). For the rest of our programming (as described below), we used MATLAB (Mathworks, Natick, MA), C, and PERL.

Microarray Data Assembly. Within the three microarray experiment sets, we considered the following as control: in the 50 cDNA microarray experiments, 11 experiments designated as “wild-type, standard-growth conditions”; in the 41 Affymetrix experiments, five experiments described as “wild-type, standard-growth conditions”; and in the 41 aerobic shift experiments, three designated as “wild-type aerobic.” To construct log ratios, we first calculated the average estimated transcript copy number (ETCN) within the control experiments for each of the three experimental data sets. Next, we divided the ETCNs from all other experiments by the

This paper was submitted directly (Track II) to the PNAS office.

Abbreviations: TR, transcriptional regulatory; TF, transcription factor; FFL, feed-forward loop; DIV, divergence; CNV, convergence; CAS, cascade; SRI, single regulatory interaction.

[‡]Present address: Applied Biodynamics Laboratory, Department of Biomedical Engineering, Boston University, Boston, MA 02215.

[¶]To whom correspondence should be addressed. E-mail: oltvai@pitt.edu.

© 2005 by The National Academy of Sciences of the USA

control ETCNs and calculated the base-10 logarithm of the resulting values. Finally, we renormalized the resulting log ratios such that the average of all log ratios for each experiment became zero. The Oklahoma University data set already contained log ratios, and thus these data were only renormalized to yield average log ratios equal to 0.

We assembled expression tables from the four different data sets after renormalizing the data again such that the mean of all gene-expression profiles became 0, and the standard deviation of all expression values within each set of experiments became 1. The rows in the obtained tables corresponded to *E. coli* genes, and the columns corresponded to various experiments. Because the *E. coli* TR network used here consists of 855 genes distributed among 418 operons, we reduced the expression tables to rows corresponding only to these 855 genes. The resulting tables contained normalized expression values (log ratios) $LR_{r,c}$ in $N_r = 855$ rows and N_c columns, corresponding to N_c microarray experiments.

Node–Node Correlation. Because the rows were normalized to zero mean, we calculated the node–node cross-correlation coefficients $cor_{i,j}$ as

$$cor_{i,j} = \left\langle \frac{\sum_{c=1}^{N_c} LR_{r,c} LR_{r',c}}{\sigma_r \sigma_{r'}} \right\rangle_{r \in I, r' \in J}, \quad [1]$$

where the averages are taken over all genes belonging to operons i and j , and the standard deviation of the gene from row r is

$$\sigma_r = \sqrt{\frac{\sum_{c=1}^{N_c} LR_{r,c}^2}{N_c}}. \quad [2]$$

Node–Signal Covariance. We calculated the node–signal covariance between node i and signal S_c with N_c binary values as

$$cov_i = \left\langle \sum_{c=1}^{N_c} LR_{r,c} S_c \right\rangle_{r \in I}, \quad [3]$$

where the averages are taken over all genes belonging to operon I .

Double z Scores. We consider an origon to be significantly affected by the external signal if it is significantly enriched in signal-affected nodes compared to the same number of nodes chosen randomly.

First, to identify the nodes significantly affected by a certain signal, we calculate $cov_{NS}(n)$ for all $n = 1, 2, \dots, N$ nodes within the network. Then we calculate a z score $z(n)$ for each individual node, defined as

$$z_{NS}(n) = \frac{cov_{NS}(n) - \mu_{NS}}{\sigma_{NS}}, \quad [4]$$

where μ_{NS} and σ_{NS} are the mean and standard deviation of $cov_{NS}(n)$, respectively.

To determine origons significantly affected by the external signal, we calculate the average z score $\mu_O = \langle z_{NS} \rangle_O$ over all N_O nodes within an origon. Then we repeatedly select N_O nodes at random within the *E. coli* TR network to estimate the mean, μ_R and standard deviation, σ_R of the quantity $\langle z_{NS} \rangle_R$. Finally, we calculate the double z score, Z_{NS} , for the origon as

$$Z_{NS} = \frac{\mu_O - \mu_R}{\sigma_R}. \quad [5]$$

Modeling the Dynamics of Small Subgraphs. To simulate subgraph dynamics, we used the built-in delay differential equation solver *dde23* from MATLAB using mass-action kinetic modeling and quasiequilibrium approximation for reactions occurring at much faster time scales than the others. First, we modeled the alteration of sensor TF-binding activity by a small metabolite. Next, we modeled RNA polymerase binding and transcription initiation, elongation, and termination. Finally, we modeled translation initiation, elongation, and termination. In the simulations, we used generic constants found in the literature, and we varied them over reasonable ranges to test the robustness of our conclusions. Because of space limitations, we describe the details of the modeling in the supporting information.

Supporting Information. For more information, see *Supporting Appendices 1–3* and *Tables 2–5*, which are published as supporting information on the PNAS web site.

Results

Directionality of Regulatory Interactions Defines TR Subnetworks.

The *E. coli* TR network is defined by its nodes (operons and their protein products) and the links connecting them (TR interactions mediated by binding of TFs to the promoter regions of operons) (8–12). Because all links are unidirectional (12), and the network contains no cycles other than autoregulatory loops (8), the 418 nodes can be arranged hierarchically into five regulatory layers as described previously (11) (Fig. 1A). We define layers as the set of nodes for which the longest path connecting them to the input layer has one, two, three, or at most four links. The layers reflect the flow of information from 76 input nodes (representing sensor TFs that are not regulated transcriptionally by any other TF) to 312 output nodes (representing non-TF proteins) (12, 19).

Besides TFs in the input layer, most TFs from intermediate regulatory layers (such as AraC, ArcA, etc.) also mediate environmental signaling into the TR network, because their conformation and activity are affected by specific changes in the environment (e.g., metabolite availability, oxygen pressure, etc.). However, the expression of sensor TFs from lower layers can also be transcriptionally regulated by other TFs, in contrast to the operons in the input layer that can only be regulated by their own protein products. Therefore, we differentiate topological inputs (operons from layer 0) from sensory inputs (TFs with environment-dependent activity).

Because of link directionality and the sparseness of the *E. coli* TR network (8, 12, 19) (see the supporting information for a description of the network's topological characteristics), TFs regulate only a limited number of operons (nodes). Thus, the set of operons regulated directly or indirectly by a given TF form a transcriptional subnetwork, rooted at the given TF. All such subnetworks are parts of other subnetworks except for the ones originating at the 76 topological input nodes, to which we refer as origons (see *Discussion*) and label them according to their input node (Fig. 1A). Because the input layer contains 76 operons, there are a total of 76 origons in the *E. coli* TR network, each affected by different environmental signal(s) (see the supporting information for their detailed description). An actual example of an *E. coli* origon is shown in Fig. 1B.

If an environmental signal affects the activity or expression level of a single sensor TF, only the expression of the genes within its origon should be affected transcriptionally, the perturbation gradually percolating toward the output layer (Fig. 1A). In turn, when several sensors are affected simultaneously by complex external signals, signal processing may involve propagation through isolated origons or signal combination by connected origons. To examine their connectedness, we redrew the TR network with origons as nodes. Two nodes in this origon network are connected if they share at least one node (operon). As shown in Fig. 1C, 45 of the 76 origons form a single connected component, indicating that a perturbation

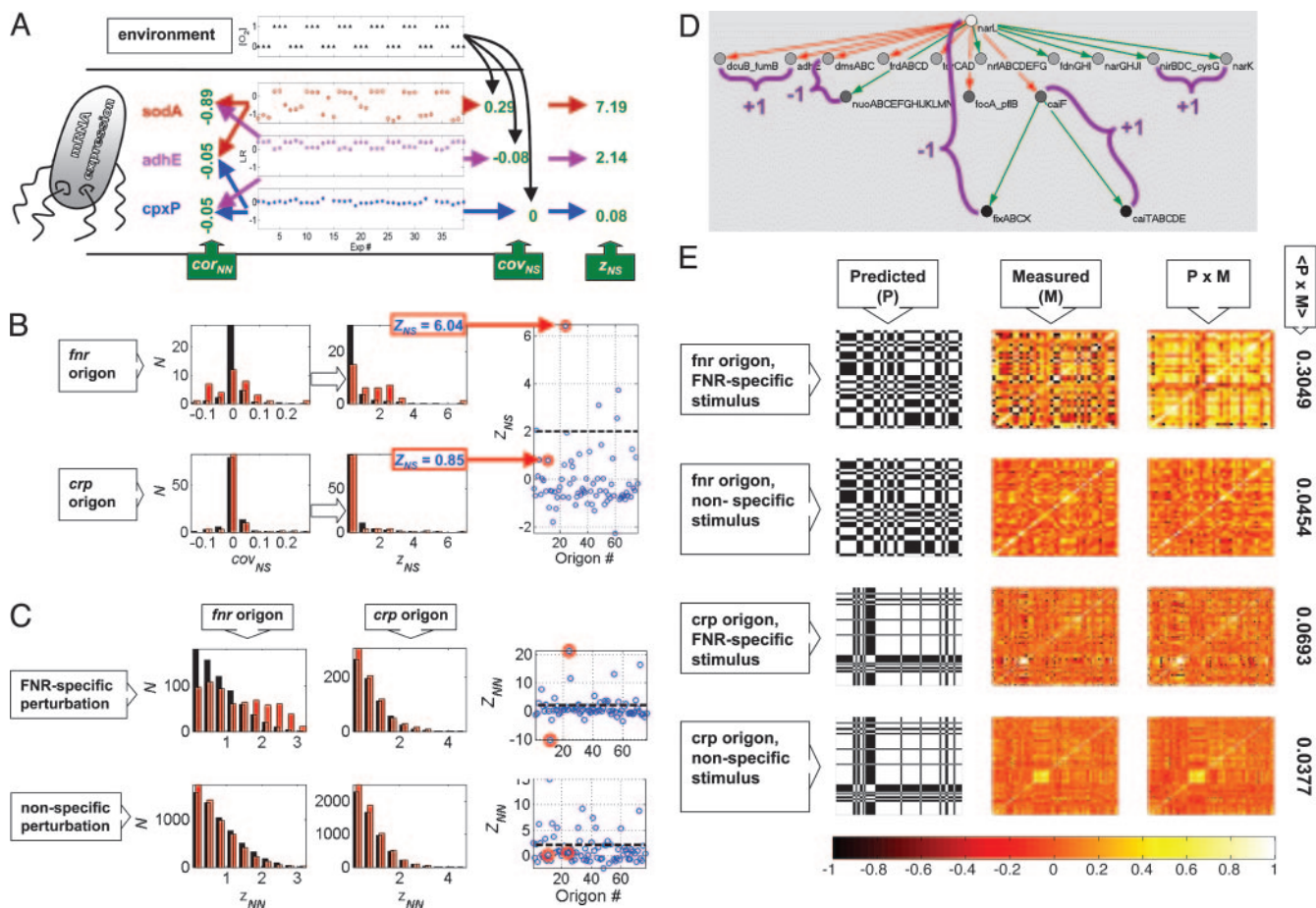


Fig. 2. Validation of the origon concept by microarray data. (A) Definition of node–signal covariance (cov_{NS}) and node–node cross-correlation (cor_{NN}). The quantity cov_{NS} characterizes the impact of the changes in environmental oxygen concentration (39 binary values of $[O_2]$ in the top graph) on the intracellular mRNA expression [log ratios (LRs): vertical axes in the bottom three graphs] of three representative single-gene nodes (*sodA*, *adhE*, and *cpxP*). The z_{NS} measures how different the cov_{NS} value of a node is from other nodes in the network. The quantity cor_{NN} characterizes the similarity between the mRNA expression profiles of the different genes. (B) Histograms of cov_{NS} (Left) within the *crp* and *fnr* origins (red bars) versus equal number of nodes chosen randomly (black bars) for FNR-specific (aerobic-shift) perturbation. The corresponding z_{NS} histograms are shown (Center), as are the double z scores, Z_{NS} , for all origins (Right) (the Z_{NS} values for the *fnr* and *crp* origins are circled). The dashed black line corresponds to a cutoff of $Z_{NS} = 2$. (C) Histograms of z_{NN} within the *crp* and *fnr* origins (red bars) versus equal number of nodes chosen randomly (black bars) for FNR-specific (aerobic-shift) and nonspecific perturbation. The Z_{NN} values for all origins are shown (Right) (*fnr* and *crp* origins are circled, and the dashed black line corresponds to a cutoff of $Z_{NN} = 2$). (D) The predicted binary node–node correlation (within the *narL* origon) between two nodes is equal to the product of link types along any path connecting the two nodes in a nondirectional version of the origon. Link types are considered as follows: +1, activating; –1, repressing; 0, dual. (E) Predictions are only indicative of measured node–node cross-correlations in the *fnr* origon for FNR-specific stimulus. Along the horizontal and vertical axes are operons within the *crp* or *fnr* origon so that the diagonal contains values calculated for an operon with itself. The agreement between predicted binary (Left) and measured (Center) cross-correlations is calculated as the product of the predictions and measurements (Right) and the corresponding averages ($P \times M$). All values in the right column are between –1 and 1, and correct predictions result in positive values, as indicated by the color scale shown at the bottom.

suggesting that origons are meaningful topological units of the TR network, dynamically used for environmental signal processing.

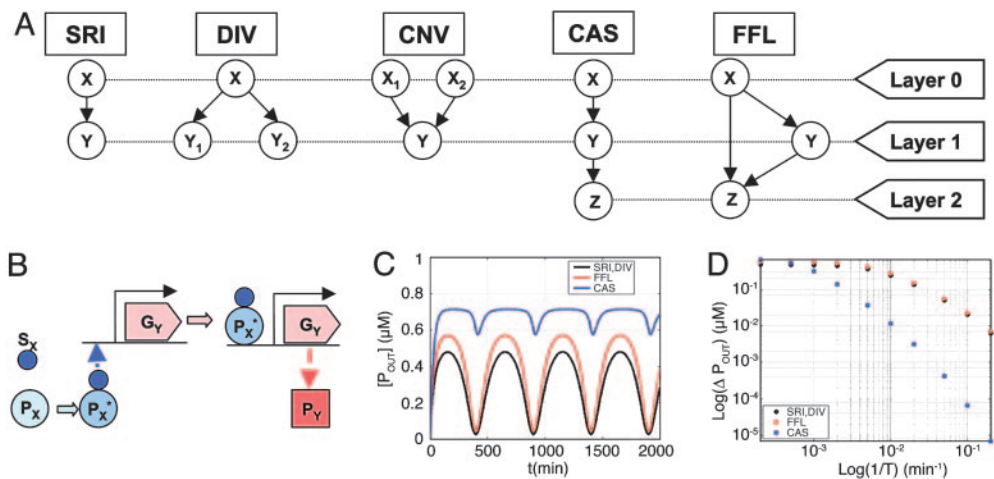
Consistency Between Known TR Interactions and Microarray Data.

After investigating how mRNA expression levels in the *E. coli* TR network reflect external changes, we studied how internal variations in the mRNA expression of various nodes relate to each other. To this end, we calculated node–node correlations, i.e., the cross-correlation between the expression profiles of two nodes (cor_{NN} , Fig. 2A). Given two nodes, we measured the deviation of their correlation from the cor_{NN} of randomly chosen node pairs by z scores (z_{NN}) again. We compared the z_{NN} values of node pairs located within the *crp* and *fnr* origins to z_{NN} values of node pairs chosen randomly from the network. For an FNR-specific stimulus, the number of outlying z_{NN} values is much higher for nodes within the *fnr* origon than for nodes chosen randomly or within the *crp* origon (Fig. 2C). We again used double z scores to statistically

characterize the observed differences and obtained for FNR-specific perturbation $Z = 21.5$ for nodes within the *fnr* origon, compared to $Z = -9.2$ for nodes within the *crp* origon. For non-FNR-specific perturbations (Fig. 2C), both double z scores were < 2 ($Z = 0.56$ for the *fnr* origon and $Z = 0.16$ for the *crp* origon), indicating that node–node correlations were not significantly affected in either of the two origins. Extending the analysis to all origins reveals that 17 origins (*fnr*, *yhdG*, *fis*, *purR*, *fruR*, *argR*, *lrp*, *cysB*, *uidR*, *nlpD*, *rpoS*, *narL*, *rhaSR*, *mlc*, *adiA*, *adiY*, *nhaR*, *soxR*, *glcC*, *birA*, *murA*) are characterized by significant ($Z > 2$) node–node correlations. Besides the known anoxic regulators FNR and NarL, this list contains origins rooted at redox sensors (SoxR), σ factors (RpoS), or TFs initiating reduced amino acid and nucleotide synthesis and transport in conditions of stress (PurR, ArgR, CysB, Lrp) (25).

We also examined whether the measured cor_{NN} values agree with the ones predicted by the type of interaction (activation or repres-

Fig. 3. Filtering properties and dynamics of small subgraphs. (A) Directed small subgraphs: SRI, DIV, CNV, CAS, and FFL are shown. The nodes are labeled by X, Y, and Z progressively, depending on their distance from the input layer of the subgraph (layer 0). (B) Schematic illustration of SRIs, the basic building unit of all three-node subgraphs, reflecting the mechanism considered in our modeling. The signal S_X activates the input TF, P_X . Once activated, P_X binds to the operator region of gene Y and allows binding of RNA polymerase P_R to its promoter region. The polymerase–operator complex initiates transcription, resulting in the synthesis of an mRNA molecule R_Y after a delay τ_R , which, bound by a ribosome P_P , is translated into the output protein P_Y after a delay τ_P . (C) Time courses of output protein levels after a periodic perturbation for SRIs and DIVs (black line), CASs (blue line), and FFLs (red line). The amplitude of fluctuations at the output of the FFL is nearly the same as at the output of the SRI, although it is substantially reduced because of the stronger filtering properties of CAS. (D) Frequency response of SRI and DIV (black circles), CAS (blue squares), and FFL (red triangles) showing the amplitude of fluctuations at the output of subgraphs (vertical axis) versus the frequency of the signal applied to their input (horizontal axis). All subgraphs are low-pass filters, but CAS has a much stronger filtering effect than either the SRI (DIV) or FFL.



sion) between a TF and its target. We associated a binary value (1 or -1) with each node within an origon depending on how its expression level is expected to change when the expression of the root node is altered. We defined the predicted binary cross-correlation between two nodes as the product of the values associated with them. Dual interactions, in which TFs can be both activating or repressing, were not considered. Fig. 2D illustrates such predictions for the narL origon. Fig. 2E illustrates the predicted binary cross-correlations between every pair of nodes in the *crp* and *fnr* origons (P , left column of plots), the measured cov_{NN} values between the same pair of nodes (M , center column of plots), and the quality of predictions estimated as the product of the predicted and measured gene–gene cross-correlations ($P \times M$, right column of plots). Increasingly lighter colors in the plots correspond to increasingly higher values, clearly illustrating that for FNR-specific stimulus, the predicted and measured cov_{NN} values agree substantially more within the *fnr* origon than within the *crp* origon. To quantitatively characterize this difference, we calculated averages over the plots in Fig. 2E Right, obtaining the values of $\langle(P \times M)\rangle$, from top to bottom) 0.3049, 0.0454, 0.0693, and, 0.0377, respectively. These values indicate again that cross-correlations are very noisy and have little predictive value of actual transcriptional regulation for nonspecific perturbations applied to both the *fnr* and *crp* origon. In contrast, FNR-specific perturbation applied to the *fnr* origon results in a 10-fold increase in the quality of the predictions.

Motifs and Subgraphs Occupy Distinct Positions Within Origons and Filter Environmental Signals Differently. The effect of many environmental signals is initiated through altered sensor TF activity, followed by the dynamical propagation of the perturbation to lower layers, eventually altering the expression of all genes within an origon. This propagation takes place through small motifs (12), or

subgraphs (15), that connect subsequent regulatory layers to each other. To elucidate the type and information-processing function of subgraphs encountered by the propagating signal, we next investigated the position and abundance of three-node subgraphs in the whole TR network as well as with respect to individual origons. Fig. 3A shows all three-node subgraphs found within the *E. coli* TR network: divergence (DIV), convergence (CNV), cascade (CAS), and FFL, all of which [composed of single regulatory interactions (SRIs)], connect two or three consecutive layers of the TR network. To examine the relative abundance of all three-node subgraphs present in the *E. coli* TR network, we used a randomization protocol (13) preserving all five layers and keeping the network acyclic (see the supporting information). Our analysis confirms that FFL subgraphs are significantly more abundant than expected (13). However, the abundance of DIV and CAS also deviates from the random expectation, with DIV occurring more frequently, whereas CAS is less frequent than expected (Table 1 and supporting information).

Based on the randomization protocol described above, we expected to find 5.07 ± 4.44 CNV subgraphs within individual origons, whereas the probability of finding no CNV subgraph in any origons is 0.11. It is surprising that CNV is completely absent from any of the individual *E. coli* origons. Therefore, the apparent role of CNV subgraphs is to combine perturbations propagating in different origons. We also found that the majority of origons are trees (Fig. 1C), containing only DIV and CAS subgraphs (Fig. 9). However, seven origons also contain FFL subgraphs in addition to their backbone tree structure (Figs. 12–18). Thus, origons can be classified into those with tree structure and those with FFL-tree structure (Fig. 1C) (see the supporting information for details). An example of an origon with tree structure is shown in Fig. 2C, and an origon with FFL-tree structure [containing aggregated FFL subgraphs (10)] is shown in Fig. 1B.

Table 1. Abundance of three-node subgraphs in the TR network of *E. coli* and the randomized version of this network

Subgraph	CNV	DIV	CAS	FFL
Abundance in real network	227	4777	160	42
Abundance in randomized network	231.92 ± 8.05	4339.3 ± 132	186.69 ± 7.08	9.50 ± 4.17
z score	0.61	3.31	3.77	7.79

The corresponding z scores (13) (listed in the bottom row) indicate the deviation of the values for the real network from those for the randomized network. The corresponding distributions are shown in Fig. 6.

For free-living bacteria such as *E. coli*, environmental signals (e.g., available carbon sources, pH, or oxygen) change frequently. To understand how the effect of fluctuating environmental changes propagate within individual origons, we developed dynamical models of the elementary topological units (subgraphs) connecting subsequent layers (Fig. 3A) by using a generic mass-action kinetic model (26) and taking into account the time delay required for mRNA and protein synthesis (27, 28). We applied perturbation to each of the subgraphs by altering the activity of their input node through the periodically changing concentration of a signaling molecule (see the supporting information for details) and investigated their response amplitude through the concentration of the protein product of their output node (Fig. 3B). As shown in Fig. 3C and D, the response amplitude of SRIs (Fig. 3A) decreases with increasing frequency of the input signal. Therefore, SRIs are low-pass filters, as described previously (29), i.e., they allow slow fluctuations to pass through but filter out fast signal fluctuations. Because SRIs are the basic building blocks of all three-node subgraphs, they all possess filtering properties to some degree. When keeping all parameters used in the simulation of the various subgraphs identical, we find that DIV (composed of two parallel SRIs) has an identical filtering effect to SRI. In contrast, the effect of the two subsequent SRIs comprising a CAS is combined, resulting in the strongest signal filtering. Finally, in FFLs with an AND-type promoter logic (30, 31), the strong filtering effect is again substantially reduced (Fig. 3C and D) because of the extra link directly connecting their input and output nodes. Thus, in contrast to what was suggested previously (12), FFLs might have the role to diminish the strong filtering effects of CASs and/or combine two related signals from two sensory inputs (see *Discussion*).

Discussion

Our analysis indicates that in *E. coli*, distinct transcriptional subnetworks (called *origons*) are responsible for environmental perturbation processing. In relation to existing concepts of multigene regulatory structures, origons are more complex entities than modulons but less complex than stimulons (32). Specifically, all nodes within a modulon must be controlled directly by a common, “pleiotropic” regulator, whereas in the origon they can be controlled indirectly by percolation of altered transcriptional levels through the network. On the other hand, origons are subnetworks originating at a single TF, whereas stimulons include all nodes affected by an environmental signal and are composed of all the origons rooted at TFs sensitive to the signal.

Based on their complexity, environmental perturbations can be classified as elementary (change of a single factor on a constant environmental background) or complex (simultaneous changes of two or more environmental factors).

The transcriptional response of the cell will depend on the type of perturbation, the structure of the affected origon, and its interconnectivity with other origons. If an elementary perturbation is highly specific to one (or few) origon(s), it will affect only operons within them. On the other hand, complex perturbations involving related signals (e.g., two different sugars) typically affect two or more sensory proteins within an origon, resulting in a combined response within individual origons. In this case, processing of the incoming signals is mediated by CAS and/or FFL subgraphs that combine related signals within the same origon. Finally, complex perturbations involving unrelated stimuli (e.g., oxygen and one sugar) are often processed independently and then combined by overlapping origons through CNVs. In addition to signal combination by sensor TFs, the target promoters of FFLs and CNVs use combinatorial logic (30) (with a dependence on signal intensity, promoter strength, etc.) that needs to be determined experimentally (16, 33).

The origon concept suggests that at the transcriptional regulatory level cells perceive their environment by first dissecting complex external signals into elementary perturbations (or modes) processed by individual origons and then developing a response by reassembling the elementary modes near the output of the TR network. Decomposition into such elementary modes is common in signal analysis (Fourier components). Thus, the origon concept may also explain why dimensionality-reduction techniques such as singular-value decomposition (34–36) or network-component analysis (37) have been successful in uncovering biologically significant information. Because the origon concept is intimately related to the directed nature of the TR network, the identification of similar topological units in undirected (protein–protein interaction) and reversibly directed (metabolic) networks will require alternative approaches. Yet, to fully understand the response to external stimuli in these networks, one cannot disregard the origon-specific directed flow of environmental signals characterizing transcriptional regulation.

We thank Q. K. Beg for compiling Table 4 for supporting information; R. Dobrin, E. Ravasz, and A. Vázquez for discussion; and J. J. Collins for comments on the manuscript. Research at the University of Notre Dame, Northwestern University, and University of Pittsburgh was supported by the U.S. Department of Energy, National Institutes of Health, and National Science Foundation.

- Ren, B., Robert, F., Wyrick, J. J., Aparicio, O., Jennings, E. G., Simon, I., Zeitlinger, J., Schreiber, J., Hannett, N., Kanin, E., et al. (2000) *Science* **290**, 2306–2309.
- Lieb, J. D., Liu, X., Botstein, D. & Brown, P. O. (2001) *Nat. Genet.* **28**, 327–334.
- Zeitlinger, J., Simon, I., Harbison, C. T., Hannett, N. M., Volkert, T. L., Fink, G. R. & Young, R. A. (2003) *Cell* **113**, 395–404.
- Harbison, C. T., Gordon, D. B., Lee, T. I., Rinaldi, N. J., Macisaac, K. D., Danford, T. W., Hannett, N. M., Tagne, J. B., Reynolds, D. B., Yoo, J., et al. (2004) *Nature* **431**, 99–104.
- Luscombe, N. M., Babu, M. M., Yu, H., Snyder, M., Teichmann, S. A. & Gerstein, M. (2004) *Nature* **431**, 308–312.
- Pritsker, M., Liu, Y. C., Beer, M. A. & Tavazoie, S. (2004) *Genome Res.* **14**, 99–108.
- Mukherjee, S., Berger, M. F., Jona, G., Wang, X. S., Muzzey, D., Snyder, M., Young, R. A. & Bulyk, M. L. (2004) *Nat. Genet.* **36**, 1331–1339.
- Thieffry, D., Huerta, A. M., Perez-Rueda, E. & Collado-Vides, J. (1998) *BioEssays* **20**, 433–440.
- Guelzim, N., Bottani, S., Bourguin, P. & Kepes, F. (2002) *Nat. Genet.* **31**, 60–63.
- Dobrin, R., Beg, Q. K., Barabási, A.-L. & Oltvai, Z. N. (2004) *BMC Bioinformatics* **5**, 10.
- Ma, H. W., Buer, J. & Zeng, A. P. (2004) *BMC Bioinformatics* **5**, 199.
- Shen-Orr, S., Milo, R., Mangan, S. & Alon, U. (2002) *Nat. Genet.* **31**, 64–68.
- Milo, R., Shen-Orr, S. S., Itzkovitz, S., Kashtan, N. & Alon, U. (2002) *Science* **298**, 824–827.
- Conant, G. C. & Wagner, A. (2004) *Nat. Genet.* **34**, 264–242.
- Vázquez, A., Dobrin, R., Sergi, D., Eckmann, J.-P., Oltvai, Z. N. & Barabási, A.-L. (2004) *Proc. Natl. Acad. Sci. USA* **101**, 17940–17945.
- Mangan, S., Zaslaver, A. & Alon, U. (2003) *J. Mol. Biol.* **334**, 197–204.
- Setty, Y., Mayo, A. E., Surette, M. G. & Alon, U. (2003) *Proc. Natl. Acad. Sci. USA* **100**, 7702–7707.
- Salgado, H., Gama-Castro, S., Martínez-Antonio, A., Díaz-Peredo, E., Sánchez-Solano, F., Peralta-Gil, M., García-Alonso, D., Jimenez-Jacinto, V., Santos-Zavaleta, A., Bonavides-Martínez, C., et al. (2004) *Nucleic Acids Res.* **32**, D303–D306.
- Martínez-Antonio, A. & Collado-Vides, J. (2003) *Curr. Opin. Microbiol.* **6**, 482–489.
- Kiley, P. J. & Beinert, H. (1998) *FEMS Microbiol. Rev.* **22**, 341–352.
- Crack, J., Green, J. & Thomson, A. J. (2004) *J. Biol. Chem.* **279**, 9278–9286.
- Allen, T. E., Herrgard, M. J., Liu, M., Qiu, Y., Glasner, J. D., Blattner, F. R. & Palsson, B. O. (2003) *J. Bacteriol.* **185**, 6392–6399.
- Chang, D. E., Smalley, D. J. & Conway, T. (2002) *Mol. Microbiol.* **45**, 289–306.
- Middendorf, M., Ziv, E. & Wiggins, C. H. (2005) *Proc. Natl. Acad. Sci. USA* **102**, 3192–3197.
- Neidhardt, F. C., Curtiss, F., Ingraham, J. L., Lin, E. C. C., Low, K. B., Magasanik, B., Reznikoff, W. S., Riley, M., Schaechter, M. & Umberger, H. E. (1996) *Escherichia coli and Salmonella typhimurium Cellular and Molecular Biology* (Am. Soc. Microbiol., Washington, DC), 2nd Ed.
- Bolouri, H. & Davidson, E. H. (2002) *BioEssays* **24**, 1118–1129.
- Santillan, M. & Mackey, M. C. (2001) *Proc. Natl. Acad. Sci. USA* **98**, 1364–1369.
- Santillan, M. & Mackey, M. C. (2004) *Biophys. J.* **86**, 1282–1292.
- Simpson, M. L., Cox, C. D. & Saylor, G. S. (2004) *J. Theor. Biol.* **229**, 383–394.
- Buchler, N. E., Gerland, U. & Hwa, T. (2003) *Proc. Natl. Acad. Sci. USA* **100**, 5136–5141.
- Mangan, S. & Alon, U. (2003) *Proc. Natl. Acad. Sci. USA* **100**, 11980–11985.
- Neidhardt, F. C., Ingraham, J. & Schaechter, M. (1990) *Physiology of the Bacterial Cell: A Molecular Approach* (Sinauer, Sunderland, MA).
- Kalir, S. & Alon, U. (2004) *Cell* **117**, 713–720.
- Alter, O., Brown, P. O. & Botstein, D. (2000) *Proc. Natl. Acad. Sci. USA* **97**, 10101–10106.
- Holter, N. S., Mitra, M., Maritan, A., Cieplak, M., Banavar, J. R. & Federoff, N. V. (2000) *Proc. Natl. Acad. Sci. USA* **97**, 8409–8414.
- Yeung, M. K., Tegner, J. & Collins, J. J. (2002) *Proc. Natl. Acad. Sci. USA* **99**, 6163–6168.
- Liao, J. C., Boscolo, R., Yang, Y. L., Tran, L. M., Sabatti, C. & Roychowdhury, V. (2003) *Proc. Natl. Acad. Sci. USA* **100**, 15522–15527.