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## Deciphering the connectivity structure of biological networks using MixNet

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### Abstract

**Background:** As biological networks often show complex topological features, mathematical methods are required to extract meaningful information. Clustering methods are useful in this setting, as they allow the summary of the network's topology into a small number of relevant classes. Different strategies are possible for clustering, and in this article we focus on a model-based strategy that aims at clustering nodes based on their connectivity profiles.

**Results:** We present MixNet, the first publicly available computer software that analyzes biological networks using mixture models. We apply this method to various networks such as the *E. coli* transcriptional regulatory network, the macaque cortex network, a foodweb network and the *Buchnera aphidicola* metabolic network. This method is also compared with other approaches such as module identification or hierarchical clustering.

**Conclusion:** We show how MixNet can be used to extract meaningful biological information, and to give a summary of the networks topology that highlights important biological features. This approach is powerful as MixNet is adaptive to the network under study, and finds structural information without any a priori on the structure that is investigated. This makes MixNet a very powerful tool to summarize and decipher the connectivity structure of biological networks.

### Background

With the increasing power of high throughput technologies and storage capacities, it is now possible to explore datasets which are in the form of complex networks. Many scientific fields are concerned by these major advances,

such as physics, social sciences, and molecular biology [1,2]. One characteristics of interest when studying complex networks is their topology or the way particles, proteins or social agents interact [1]. More generally, studying the topology is crucial to understand the organization of

networks, as structure often affects function. Since networks show complex structural patterns, one common task is to find an appropriate way to summarize their structure. Many indicators have been proposed for this purpose: the degree distribution [3], the clustering coefficient [2,4], and the small world property [1] are among the most popular. However since summarizing a topology using those indicators gives a crude view of the networks topology, another research direction has been to gather nodes that behave similarly from the point of view of a user defined criterion [5-7].

Clustering methods that have been proposed are mainly focused on community detection, *i.e.* they aim at finding groups of nodes that are highly intra-connected and poorly inter-connected [8]. Hierarchical versions of these methods are also available [5]. However, when performing exploratory data analysis, it may be difficult to search for a particular structure. Real networks may not show community structure for instance, or may be characterized by various connectivity patterns among which community is only one feature.

Model-based clustering is a powerful alternative to those methods, as the model underlying the algorithm allows the blind search of connectivity structure without any *a priori* [7,9,10]. The basics of this strategy is to consider that nodes are spread among an unknown number of connectivity classes which are unknown themselves. Many names have been proposed for this model, and in the following, it will be denoted by MixNet, which is equivalent to the Block Clustering model [9].

When using MixNet one central question is the estimation of the parameters, and the associated optimization method. Bayesian strategies have been proposed, but they are limited as they can handle networks with hundreds of nodes only [9]. Heuristics have also been proposed for this problem [10]. In this work, we present the MixNet software program which is the first publicly available software that fits mixture models on large networks using non Bayesian maximum likelihood estimation. The statistical developments associated with this software have been published elsewhere [7], and our algorithm uses a variational approach that has been developed in the context of graphical models [11]. Here we consider the application of MixNet to different biological networks such as regulatory, cortex, foodweb and metabolic networks. We show how flexible the method is, how it summarizes the connectivity structure of a complex network, and how this summary can be used to understand topology-based biological features.

## Results

### Brief recall of MixNet principles

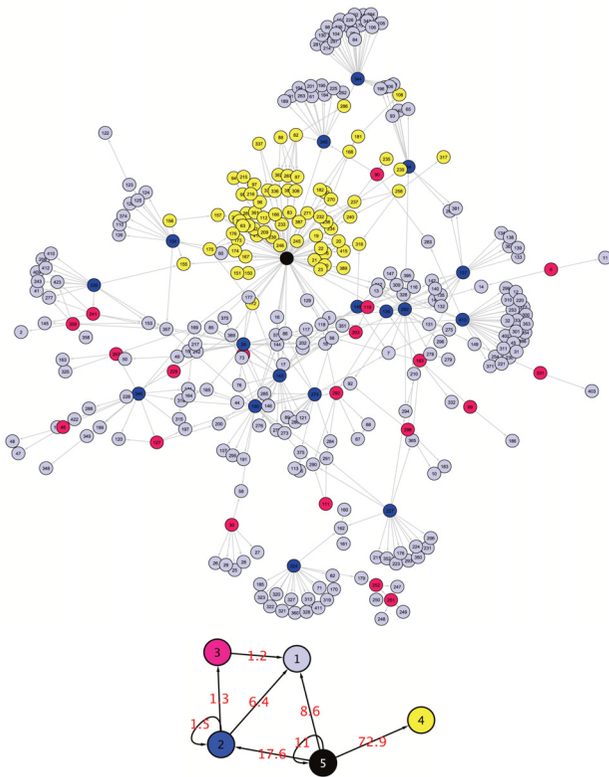
In this first paragraph we briefly recall the principle of mixture models when applied to random graphs.

This is a general setting that has been developed extensively from the statistical point of view [7,9,10].

The network is modeled as a random graph with  $X$  representing its connectivity matrix, such that  $X_{ij} = 1$  if nodes  $i$  and  $j$  are connected and 0 otherwise. In this article, we consider directed networks, such that  $X_{ij}$  may be different from  $X_{ji}$ . The idea of MixNet is to consider that nodes can be spread into  $Q$  connectivity classes which are hidden, with  $Q$  being unknown as well. Then we consider that there exists a sequence of hidden label variables  $Z$  such that  $Z_{iq} = 1$  if node  $i$  belongs to class  $q$ . The parameters of this model are  $\alpha$ , the proportion of each group, and  $\pi$  the connectivity of the groups, such that  $\pi_{q\ell}$  represents the probability for a node of group  $q$  to be connected to a node from group  $\ell$  (given in percentage in the sequel). To this extend,  $\pi$  is a summary of the connectivity of the original network, at the group level. MixNet results can be displayed in two ways. The first intuitive representation is to map the MixNet classes on the nodes of the network as in Figure 1. However, this view may not be informative when too many nodes/colors are present. The second way is to give a graphical representation of the connectivity matrix  $\pi$  which provides a synthetic view of the intensity and direction of connexions between and within MixNet classes (Fig. 1, Table 1). Then the purpose is to interpret such a summary, and our work aims at showing how biological information can be extracted from MixNet results. A classical difficulty when using clustering techniques is to determine how many clusters there are. The advantage of model-based clustering is that it gives a framework for deriving theoretical criteria for model selection. However, our point is that since there is no "true" number of clusters, it may be valuable to study the results given with different configurations. To this extend, we will use two criteria in this article. The first one is called the Integrated Classification Likelihood (ICL [7]), it is based on a penalization of the likelihood of the model. The second one is called the "adaptive strategy". Its principle is to study the increase of the likelihood according to the dimension of the model, and to select the number of clusters for which this increase is less significant [12]. These criteria are briefly described in the Method section.

### A meta-regulation diagram in the TRN of *E. Coli*

Transcriptional regulatory networks (TRN) constitute one important example of biological networks that are studied from the structural point of view. Nodes of the network correspond to operons which are linked if one operon encodes a transcription factor that directly regulates



**Figure 1**  
**E. Coli TRN with 5 MixNet classes with proportions.**  
 $\hat{\alpha}_1 = 65.49$ ,  $\hat{\alpha}_2 = 5.18$ ,  $\hat{\alpha}_3 = 7.92$ ,  $\hat{\alpha}_4 = 21.10$ ,  $\hat{\alpha}_5 = 0.30$

another operon. Such networks have been shown to share some important properties, such as a relative sparseness, a very low number of feed back circuits, and a hierarchical organization [13]. Thus grouping operons based on their connectivity structure appears essential to understand the wiring diagram of such complex networks. In this paragraph, we consider the connex component of the the E. Coli TRN [14].

**Table 1: Connectivity matrix for E. Coli TRN with 5 classes. The probabilities of connexion are given in percentage, and probabilities lower than 1% are not displayed.**

	MixNet Classes				
	1	2	3	4	5
1	.	.	.	.	.
2	6.40	1.50	1.34	.	.
3	1.21	.	.	.	.
4	.	.	.	.	.
5	8.64	17.65	.	72.87	11.01
alpha	65.49	5.18	7.92	21.10	0.30

*Summarizing regulatory structure: the MixNet representation*

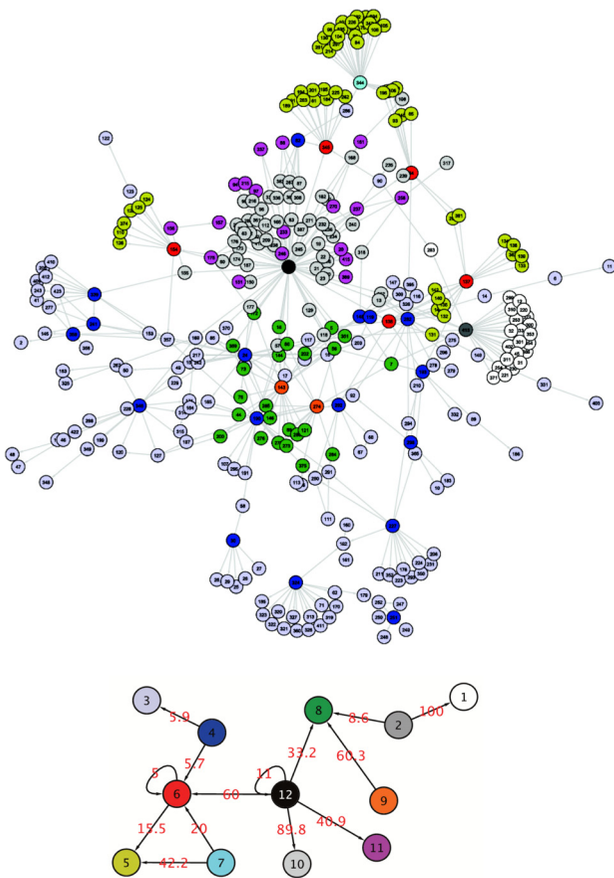
The clustering results with 5 classes (given by the ICL criterion) gives a rough picture of the network's structure. The connectivity matrix  $\pi$  of the TRN is characterized by (i) empty rows and (ii) small diagonal elements (Table 1): (i) means that some groups are made of strictly regulated operons (nodes that receive edges only), and (ii) that there is no community structure, i.e. there is no group which is heavily intra-connected and poorly inter-connected. This result is coherent with the structure of regulatory circuits which form cascades of regulations without feedback [13], meaning that nodes do not share modularity patterns in this regulatory network. Figure 1 indicates that the majority of operons are regulated by very few nodes. At this resolution level, the network is summarized into regulated operons (groups 1 and 4), which receive edges only. These two groups are distinguished based on their regulatory elements: operons of group 4 are regulated by crp only (which makes its own group), whereas operons of group 1 are regulated by many cross-talking elements (group 2, 3, and 5).

*Meta Motifs of regulation*

It has been shown that some motifs like the popular Feed Forward Loop constituted a core structure of the E. Coli regulatory network [14]. When looking at Figure 1, it appears that MixNet exhibits the same global structures at the group level. Groups 5 and 4 form a Single Input Module (SIM), i.e. one TF regulating other operons that do not communicate ( $\pi_{4,4} < 1\%$ ). Similarly, groups 2-3-1 and 2-5-1 form a "meta" Feed-Forward loop. In both cases the effector group is group 1, and groups 2 and 3 can be viewed as information relays.

*Getting a more detailed picture*

The adaptive strategy selects 12 groups which highlight the hierarchical structure of the regulation wiring diagram (Figure 2). The majority of nodes are strictly regulated operons (groups 1, 3, 5, 8, 10), whereas regulators are clustered into small groups that are distinguished based on their connectivity patterns and on their targets. For example yhdG\_fis (group 2) regulates nodes of groups 1 and 8, operons of group 9 (fnr, narL) regulate operons of group 8. MixNet can also be used to detect operons that act as global TF from the connectivity point of view. For instance, rpo operons are clustered in "regulatory" classes (operon rpoE\_rseABC forms group 7 on its own). This result is not surprising though, as rpo operons are involved in the  $\sigma$  unit of the RNA polymerase. More generally, beyond groups that are made of unique major regulatory elements, MixNet gather "regulatory-like" elements together. For instance, group 4 is made of both global TF and  $\sigma$  factors (Table 2).



**Figure 2**  
**E. Coli TRN with MixNet 12 classes with proportions.**  
 $\hat{\alpha}_1 = 6.66, \hat{\alpha}_2 = 0.30, \hat{\alpha}_3 = 37.10, \hat{\alpha}_4 = 5.35, \hat{\alpha}_5 = 16.61,$   
 $\hat{\alpha}_6 = 1.52, \hat{\alpha}_7 = 0.30, \hat{\alpha}_8 = 8.59, \hat{\alpha}_9 = 0.61, \hat{\alpha}_{10} = 16.84,$   
 $\hat{\alpha}_{11} = 5.81, \hat{\alpha}_{12} = 0.30$

Meta motifs are also present in this representation: a Meta Feed Forward Loop (5-6-7) and Single Input Modules (12-10, 12-11, 12-8, 2-8 and 2-1). Their formation is due to groups 12 and 2 which are made of one operon only (crp and yhdG\_fis respectively). Another meta motif is the Dense Overlapping Regulon (DOR motif, groups 4-3). A DOR motif is formed when a set of operons are each regulated by a combination of a set of input transcription factors.

**Discovering Hub families in the macaque cortex network**  
 The dataset consists in cortical regions connected by inter-regional pathways in the Macaque Cortex [15]. As brain function is based on inter-regional connexions, studying the way cortical regions interact may offer new perspectives in the comprehension of information flows within the brain. It appears that particular brain regions may play

**Table 2: Repartition of the E. Coli TRN in MixNet classes.**

Operon	class id	out degree	in degree
yhdG_fis	2	26	0
arcA†	4	20	1
argR	4	6	0
cytR	4	7	0
fadR	4	5	0
FruR	4	7	0
himA†‡	4	21	0
hns†	4	7	1
Irp†	4	14	0
marRAB	4	5	1
metJ	4	4	0
nlpD_rpoS*	4	14	0
ompR_envZ†	4	6	1
oxyR†	4	4	0
purR†	4	16	0
rob†	4	12	0
rpoN*	4	13	0
soxS†	4	6	1
cpxAR†	6	9	1
flhDC	6	7	3
fliAZY*	6	12	2
fur†	6	9	1
rpoH*	6	10	4
rpoE_rseABC*	7	24	0
fnr†	9	22	0
narL†	9	13	0
crp†	12	72	0

List of operons which correspond to regulatory operons in the Coli regulation network with  $Q = 12$  groups. † Global TF from [35]. Note that flhDC is a master compound regulator for motility and chemotaxis, and has not yet been reported to regulate other TFs [36]. ‡himA is the  $\alpha$ -subunit of the Integration Host Factor. \* for  $\sigma$ -factors.

different roles: some regions can be at the "center" of a particular part of the network, meaning that a lot of information will pass through them, whereas other parts of the network may be more "peripheral". Consequently, identifying central zones would be important, as their lesion may compromise the integrity of the whole network.

From a topological view, finding those "hubs" as focused much attention, with a popular definition based on degree. However, there exists many ways for a node to be a hub, and degree is only one criteria. As there is no formal definition of what a hub is, there are many different hubs (provincial and central). This is why multi-criteria strategies were developed to find nodes that can be called "hubs" [15]. From a methodological point of view, this approach seems to be limited as the resulting hubs will be criteria-dependent. The gain of MixNet is that the model





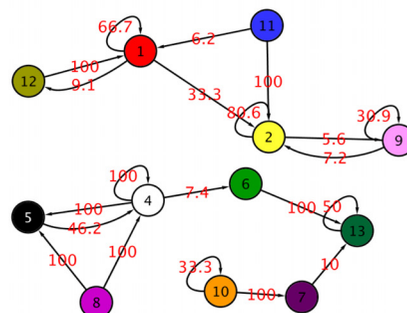
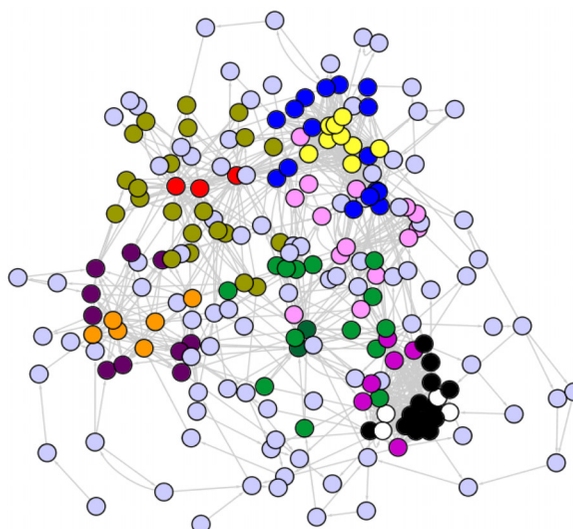


**Table 4: Average degree for the metabolic network with 13 MixNet classes.**

Mixnet Class	alpha	Ave. In Deg	Ave. Out Deg
1	1.4	24.33	7
2	4.1	24.88	8.88
3	45.4	1.52	1.40
4	1.8	16	17.67
5	6.0	10.33	2.58
6	6.3	2.78	3.14
7	4.6	6.80	1.40
8	2.8	1.16	19.67
9	6.4	5.14	6.29
10	2.8	2.66	12.33
11	7.4	1.81	10.56
12	10.1	1.54	4
13	0.9	18.50	3.50

strate to produce phosphate by class 1, which is also produced by all reactions of class 11. The distinction between producers and consumers can also be seen with the average in/out degree of each class (Table 4). It is important to note that the presence of phosphate here is not the sub-product of the transformation of ATP in ADP or other cofactor transformation. Interestingly in *B. aphidicola*, the use of phosphate as substrate occur in degradation of purines whose the products may lead to the synthesis of several other important metabolites as the chorismate, key compound in the synthesis of amino acids. A similar pattern can be found with CO<sub>2</sub> that is used by reactions used by reactions of class 13, and with reactions that use/produce protons (Table 5). If we go to further details, sugars also structure the network (class 9), with reactions are span among the pentose phosphate pathway and the glycolysis.

In component 2, we observe a very strong structure which is due to the use of glutamate, with irreversible reactions that produce glutamate from glutamine (class 8), and reactions that use glutamate (classes 5 and 4). Interestingly these consumer reactions are split because of their different reversibility despite their strong probability of connexion ( $\pi_{4,5} = 100\%$ ). Reactions of class 4 are all reversible and are involved in the metabolism of 3 Amino Acids (Isoleucine, Leucine and Valine) with a common EC number (2.6.1.42), whereas reactions of class 5 are strictly irreversible (83% of which being with EC numbers 2.6.1 and 6.3.2). The glutamate is a key compound in the synthesis of amino acids and thus plays a very important role in the symbiotic function of *B. aphidicola*. Consequently, MixNet enables to emphasize the the central role of the glutamate in the network.



**Figure 6**  
**Metabolic network with 13 MixNet classes with proportions.**  $\hat{\alpha}_1 = 1.4, \hat{\alpha}_2 = 4.1, \hat{\alpha}_3 = 45.4, \hat{\alpha}_4 = 1.8, \hat{\alpha}_5 = 6.0, \hat{\alpha}_6 = 6.3, \hat{\alpha}_7 = 4.6, \hat{\alpha}_8 = 2.8, \hat{\alpha}_9 = 6.4, \hat{\alpha}_{10} = 2.8, \hat{\alpha}_{11} = 7.4, \hat{\alpha}_{12} = 10.1, \hat{\alpha}_{13} = 0.9.$

**Discussion and conclusion**

In this work we show how MixNet can be used to study biological network by providing an accurate summary of the main topological features that structure the network. We explored networks that show very diverse structures: the transcription and the foodweb networks are sparse and globally structured by hubs, whereas the cortex and the metabolic network are dense with some hubs and some strongly connected components. Interestingly MixNet is adaptive to each structure, and catches very diverse features like hubs, hub families, connecting classes, cliques, and local hierarchies. This makes this tool very flexible, and very powerful to detect many features within the same network, whereas oriented clustering techniques like module identification will search for specific features only, even if these features are not in the network. Overall, the graphical representation of a network is

**Table 5: Reactions of the Buchnera metabolic network that involve protons.**

MixNet class 7	
substrate(s)	product(s)
proton+cpd-602	→ cpd-1086
proton+super-oxide	→ hydrogen-peroxide+oxygen-molecule
proton+hydroxy-methyl-butenyl-dip	→ delta(3)-isopentenyl-pp
proton+hydroxy-methyl-butenyl-dip	→ cpd-4211
proton+3-dehydro-shikimate	→ shikimate
proton+2,3-dihydrodipicolinate	→ delta1-piperideine-2-6-dicarboxylate
proton+2-amino-3-oxo-4-phosphonoxybutyrate	→ l-amino-propan-2-one-3-phosphate+carbon-dioxide
proton+2-aceto-lactate	→ diol-isovalerate
proton+methylene-thf	→ 5-methyl-thf
proton+l-aspartate-semialdehyde	→ homo-ser

MixNet class 10	
substrate(s)	product(s)
erythrose-4p	→ proton+erythronate-4p
2-d-threo-hydroxy-3-carboxy-isocaproate	→ proton+cpd-7100
cpd-296	↔ proton+lipoic-acid
proton+oxygen-molecule	↔ proton
sirohydrochlorin+fe+2	→ proton+siroheme
glc-6-p	→ proton+d-6-p-glucono-delta-lactone

a challenging task, and MixNet provides a global view of the network and emphasizes the key elements that make the topology. Summarizing nodes into a small number of meta-nodes linked by meta-edges gives a representation that constitutes a clear synthesis of the network topology.

Here we presented how MixNet parameters can reveal interesting features from the biological point of view. This emphasizes that MixNet is not only a computer software, but also a powerful model that can be used to simulate networks, or as a reference model under which theoretical statistics can be derived. This approach has already been demonstrated in network motifs analysis [25].

Note that the topology of a network is only one structural information that can be used to understand networks functions. It is worth being noted that the incorporation of edge direction improves the interpretability of the results, as the topology itself only constitutes a crude information. Moreover, many networks also have informations on edges: transcription regulatory networks have labeled edges (Activator/Repressor), and metabolic network have stoichiometry which reflects compounds flow in the network. A future research direction will be to use this additional information [26].

**Methods**

**Data description**

The transcription regulatory network has been downloaded from U. Alon web page [27]. We use only the connex component of the 1.1 version of the network, which

is made of 328 nodes with 456 interactions. The food web network has been provided by A. Clauset, and is made of 86 nodes and 113 edges. The cortex network is made of 47 nodes and 505 interactions. It is available in the supplementary material of [15]. The metabolic network was build by the pathway-tools software [28] from the genomic annotations provided by the MAGE annotation platform [29]. The genome of *B. aphidicola* is quite well annotated since it can be considered as a subset of the intensively curated genome of *Escherichia coli*. Consequently, the construction of the *B. aphidicola* metabolic network is supposed to be meaningful from the biological point of view. Overall the network is made of one connex component with 946 edges and 218 nodes.

**Model Selection**

In this paragraph we explain briefly the model selection procedure employed to select the number of clusters. The first criterion ICL is a particular penalized likelihood criterion: it is used to make a trade-off between a reasonable number of parameters and a good quality of fit of the data. In addition to the traditional BIC, ICL also considers the quality of the partition, meaning that it will select a number of clusters for which the classes are well separated (with low entropy). Consequently, ICL is based on the penalization of the complete-data log likelihood of the model, that accounts for the observed X and the missing data Z. The number of classes is selected such that:

$$\hat{Q} = \arg \max_Q \{ \log \mathcal{L}(X, Z) - \text{pen}(Q) \},$$

with  $\text{pen}(Q)$  a penalty that depends on the number of nodes in the network, as well as on the number of parameters in the model [7].

The second method we employ is based on the geometrical behavior of the incomplete-data likelihood when the number of classes increases. It is an adaptive method that has been successfully employed in diverse contexts [12,30,31]. The principle of this method is to calculate the second derivative of the likelihood, and to select the number of classes for which this derivative exceeds a threshold, which is set to 0.5 in practice. This method is close to the L-curve method [12].

### The MixNet software

All the presented algorithms are implemented into the MixNet software package which is written in ANSI C++ and includes Fortran 77 subroutines from the ARPACK [32] library. Optional post-treatment programs written in Perl are also included in the package. Compilation and installation are compliant with the GNU standard procedure. The library is freely available on the MixNet webpage [33]. Online documentation and man pages are also available. MixNet is licensed under the GNU [34] General Public License.

The complexity of the algorithm is proportional to the number of edges of the network (sparse storage format), and  $\mathcal{O}(n^2Q^2)$  in time (where  $n$  stands for the number of nodes and  $Q$  for the number of clusters). If MixNet is run for 1 to  $Q$  clusters, the overall complexity is then  $\mathcal{O}(n^2Q^3)$ . We present the speeds of execution of MixNet on the webpage [33]. Our experience is that on the studied networks, the execution speeds were similar to the simulated annealing method.

### Competing interests

The authors declare that they have no competing interests.

### Authors' contributions

FP wrote the manuscript and conducted the analysis, VM developed the MixNet software program, LC created and did the analysis of the metabolic network, JJD and SR supervised the study.

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### References

1. Strogatz S: **Exploring complex networks.** *Nature* 2001, **410**:268-276.
2. Newman M, Watts D, Strogatz S: **Random graph models of social networks.** *PNAS* 2002, **99**:2566-2572.
3. Barabási A, Albert R: **Emergence of scaling in random networks.** *Science* 1999, **286**:509-512.
4. Albert R, Barabási A: **Statistical mechanics of complex networks.** *R Modern Physics* 2002, **74**:47-97.
5. Girvan M, Newman M: **Community structure in social and biological networks.** *PNAS* 2002, **99**(12):7821-7826.
6. Radicchi F, Castellano C, Cecconi D, Loreto V, Parisi D: **Defining and identifying communities in networks.** *PNAS* 2004, **101**(9):2658-2663.
7. Daudin J, Picard F, Robin S: **A mixture model for random graphs.** *Stat and Computing* 2008, **18**:173-183.
8. Guimera R, Amaral LN: **Functional cartography of complex metabolic networks.** *Nature* 2005, **433**:895-900.
9. Nowicki K, Snijders T: **Estimation and prediction for stochastic blockstructures.** *JASA* 2001, **96**(455):1077-1087.
10. Newman M, Leicht E: **Mixture models and exploratory analysis in networks.** *PNAS* 2007, **104**(23):9564-9569.
11. Jordan M, Ghahramani Z, Jaakkola T, Saul L: **An Introduction to Variational Methods for Graphical Models.** *Mach Learn* 1999, **37**(2):183-233.
12. Lavielle M: **Using penalized contrasts for the change-point problem.** *Signal Processing* 2005, **85**(8):1501-1510.
13. Balazsi G, Barabasi AL, Oltvai Z: **Topological units of environmental signal processing in the transcriptional network of Escherichia Coli.** *PNAS* 2005, **102**(22):7841-7846.
14. Shen-Orr S, Milo R, Mangan S, Alon U: **Network motifs in the transcriptional regulation network of Escherichia coli.** *Nature genetics* 2002, **31**:64-68.
15. Sporns O, Honey C, Kötter R: **Identification and classification of hubs in brain networks.** *PLoS ONE* 2007, **2**:e1049.
16. Dunne J, Williams R, Martinez N: **Food-web structure and network theory: the role of connectance and size.** *PNAS* 2002, **99**(20):12917-12922.
17. Clauset A, Moore C, Newman M: **Hierarchical structure and the prediction of missing links in networks.** *Nature* 2008, **453**:98-101.
18. Dawah H, Hawkins B, Claridge M: **Structure of the parasitoid communities of grass-feeding chalcid wasps.** *Journal of animal ecology* 1995, **64**:708-720.
19. Martinez N, Hawkins B, Dawah H, Feifarek B: **Effects of sampling effort on characterization of food-web structure.** *Ecology* 1999, **80**(3):1044-1055.
20. Buchner P: *Endosymbiosis of animals with plant microorganisms* John Wiley & Sons, Inc., New York, NY; 1965.
21. Shigenobu S, Watanabe H, Hattori M, Sakaki Y, Ishikawa H: **Genome sequence of the endocellular bacterial symbiont of aphids Buchnera sp. APS.** *Nature* 2000, **407**(6800):81-6.
22. Caspi R, Foerster H, Fulcher C, Kaipa P, Krümmenacker M, Latendresse M, Paley S, Rhee S, Shearer A, Tissier C, Walk T, Zhang P, Karp P: **The MetaCyc Database of metabolic pathways and enzymes and the BioCyc collection of Pathway/Genome Databases.** *Nucleic Acids Res* 2008, **36**:D623-D631.
23. Arita M: **The metabolic world of Escherichia coli is not small.** *Proc Natl Acad Sci USA* 2004, **101**(6):1543-1547.
24. Handorf T, Christian N, Ebenhoh O, Kahn D: **An environmental perspective on metabolism.** *J Theor Biol* 2008, **252**(3):530-537.
25. Picard F, Daudin JJ, Koskas M, Schbath S, Robin S: **Assessing the exceptionality of network motifs.** *J Comput Biol* 2008, **15**:1-20.
26. Mariadassou M, Robin S: **Uncovering latent structure in valued graphs: a variational approach.** *In Tech Rep 10 SSB*; 2007.
27. U. Alon webpage [[http://www.weizmann.ac.il/mcb/UriAlon/Network\\_motifs\\_in\\_coli/ColiNet-LLI](http://www.weizmann.ac.il/mcb/UriAlon/Network_motifs_in_coli/ColiNet-LLI)]
28. Karp PD, Paley S, Romero P: **The Pathway Tools software.** *Bioinformatics* 2002, **18**(Suppl 1):S225-S232.

29. Vallenet D, et al.: **MaGe: a microbial genome annotation system supported by synteny results.** *Nucleic Acids Res* 2006, **34**:53-65.
30. Antoniadis A, Bigot J, von Sachs R: **A multiscale approach for statistical characterization of functional images.** *Journal of Computational and Graphical Statistics* 2008 in press.
31. Picard F, Robin S, Lavielle M, Vaisse C, Daudin JJ: **A statistical approach for CGH microarray data analysis.** *BMC Bioinformatics* 2005, **6**:27.
32. **ARPACK** [<http://www.caam.rice.edu/software/ARPACK/>]
33. **MixNet webpage** [<http://pbil.univ-lyon1.fr/software/mixnet>]
34. **GNU** [<http://www.gnu.org/licenses/>]
35. Martinez-Antonio A, Collado-Vides J: **Identifying global regulators in transcriptional regulatory networks in bacteria.** *Curr Opin Microbiol* 2003, **6(5)**:482-489.
36. Martinez-Antonio A, Jangra S, Thieffry D: **Functional organization of Escherichia Coli transcriptional regulatory network.** *J Mol Biol* 2008, **381**:238-247.

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