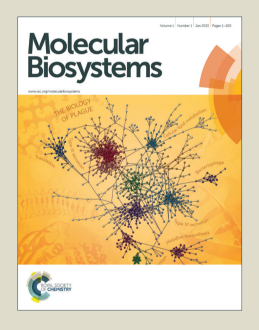
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Network motif frequency vectors reveal evolving metabolic network organisation

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At the systems level many organisms of interest may be described by their patterns of interaction, and as such, are perhaps best characterised via network or graph models. Metabolic networks, in particular, are fundamental to the proper functioning of many important biological processes, and thus, have been widely studied over the past decade or so. Such investigations have revealed a number of shared topological features, such as a short characteristic path-length, large clustering coefficient and hierarchical modular structure. However, the extent to which evolutionary and functional properties of metabolism manifest via this underlying network architecture remains unclear. In this paper, we employ a novel graph embedding technique, based upon low-order network motifs, to compare metabolic network structure for 383 bacterial species categorised according to a number of biological features. In particular, we introduce a new *global significance score* which enables us to quantify important evolutionary relationships that exist between organisms and their physical environments. Using this new approach, we demonstrate a number of significant correlations between environmental factors, such as growth conditions and habitat variability, and network motif structure, providing evidence that organism adaptability leads to increased complexities in the resultant metabolic networks.

Introduction

Many biological systems can be described using the techniques of network science 1,2, which provides a powerful set of tools for analysing the underlying connectivity structures that naturally arise within all living organisms^{3,4}. At the cellular level, networks emerge via interacting proteins, and other macro-molecules, resulting in various biochemical nets, such as gene regulatory networks 5,6, protein-protein interaction networks ⁷ and protein residue networks ^{8,9}. In this regards, the metabolic process in particular plays a fundamental role, providing the building blocks (nucleic acids and amino acids) that enable genes to interact effectively, and thus for the cell to function properly. Moreover, recent evidence suggests that the interaction patterns described by metabolic networks reflect the evolutionary origins of important functional changes ^{10–12}, and thus understanding their topological organisation promises to unravel important features of biological organisation at the systems level.

Metabolic networks have been the focus of a large number of studies (see for example the review by Lacroix *et al.* ¹³ and references therein) and several important structural characteristics have been evidenced. For example, modularity, i.e. the propensity for a network to organise into nearly-independent

structural units, has been shown to be a prevalent feature within metabolic networks ^{14,15}, and has, for example, been related to important biological properties such as robustness ^{16,17} and evolvability ^{10,11,16}. However, metabolic networks are by no means perfectly modular; their inter-module connectivity is relatively high, leading some authors to conclude that these networks are better described as being hierarchically structured ¹⁴, that is metabolic networks may be considered to possess fractal-like properties, such as self-similarity. Indeed, the existence of many small highly integrated units, which then group together to form larger modules and so on, provides one possible explanation for the high level of inter-modular connectivity observed in these networks.

Another popular approach for analysing metabolic networks is provided by *network motifs* ¹⁸, i.e. recurrent, statistically significant subgraphs. Motifs are of particular interest since they are typically associated with certain biological functions, and their relative over-abundance is considered to be an evolutionary result reflecting their "importance" to the organisms involved ^{19,20}. Moreover, they constitute the basic structural units from which complex metabolic networks are formed, and thus provide a simplified framework for probing large-scale topologies. For example, in a recent study Shellman *et al.* ²¹ successfully captured key evolutionary differences between metabolic networks from the six different kingdoms of life, employing network motif analysis. Another example highlighting the considerable potential of such an approach, is provided by the work of Asgari and colleagues ²²,

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in which the authors employ recent advances in the theory of network controllability as a method for improving drugtarget discovery techniques. In particular, they suggest that network motifs provide ideal 'driver' candidates, which can be employed to manoeuvre the system of interest into certain desirable states.

Here, we introduce a novel technique for comparing biological networks of varying size based on local network structure. In particular, we propose a new embedding technique based on low order network motifs. In our approach, each network is mapped to a point in a high-dimensional vector space, the dimension of which depends on the number of motifs considered (n = 212 in our work as we consider all 3 and 4 node motifs²³). By using a suitably defined low-rank approximation, we are able to combine the 212 motif frequency scores into a single network specific measurement, which allows us to compare and contrast networks in terms of just a single parameter. Using this new measure we investigated 383 bacterial metabolic networks with identified growth conditions, as well as a smaller subset of 115 networks classified according to the amount of variability present within their natural habitats, and found a number of significant correlations between network motif structure and fluctuations in environmental conditions.

A new graph embedding approach

Motivated by the prominent role that network motifs have played to date in the analysis of biological networks (see for example the book of Alon³ and references therein), we propose a new, lossy graph embedding technique based on low-order motifs. The proposed technique is lossy in the sense that the original network cannot be recovered from the corresponding vector-space representation. Importantly, such an approach takes a difficult and unwieldy problem, i.e. the analysis of many large, complex biological networks of differing order, and replaces it by one which is 'easier' to manipulate – a plethora of tools and techniques from statistical machine learning ^{24,25} already exist for the analysis of the resultant embedded data.

Motif frequency vectors

Motif frequencies can be used to directly compare different metabolic networks as they provide a 'unique' network signature ²¹. Alternatively, networks can be compared by calculating a feature vector of z-score's, computed in the usual way, i.e.

$$z_{i,j} = \frac{N_j^i - \left\langle N_j^{\text{rand}_i} \right\rangle}{\sigma_i^{\text{rand}_i}},$$

where here, N_j^i denotes the rate of recurrence of the jth motif within the ith network whilst $\langle N_j^{\mathrm{rand}_i} \rangle$ and $\sigma_j^{\mathrm{rand}_i}$ denotes the

mean and standard deviation of the rate of recurrence of the *j*th motif in an ensemble of randomised networks³.

In this way, for each network of interest we can compute a feature vector, \mathbf{z}_i , whose elements are the z-scores of each network motif. For example, if, as in this work, we consider all 3- and 4-node motifs then the result is a vector $\mathbf{z}_i \in \mathbb{R}^{212}$ representing the *i*th network.

Note that it is typically the case that the networks we wish to compare are of varying order and as such we need to take care that network size does not bias any results. To handle this issue one can consider instead of the z-scores defined above, a so-called *significance profile* ²⁶ defined by

$$s_{i,j} = \frac{z_{i,j}}{\sqrt{\sum_k z_{i,k}^2}}.$$

The motif significance profile for the *i*th network, s_i , is simply the normalised vector of z-scores. The motif significance profile allows for direct comparisons between networks of different sizes. This is important due to the fact that motifs in larger networks tend to exhibit larger z-scores than they do in smaller networks 23,26 . Note also, that each entry of the motif significance profile lies in the interval [-1,1].

In the work presented here, we threshold the network significance profiles such that any entries $s_{i,j} < 0$ are set to zero as we are only interested in those motifs that are over represented. Motifs that are under represented are known as antisignificant motifs, or anti-motifs, and although we do not consider them in this study, the approach forwarded here can easily be extended to that case. This results in a matrix

$$S = \left[\mathbf{s}_1, \dots, \mathbf{s}_m\right]^T \geq 0,$$

i.e. a non-negative matrix, whose rows consist of the significance profiles (thresholded) for the m networks under investigation.

To analyse the matrix S we use a matrix decomposition to compute a low-rank approximation of the data²⁵. Since our data is non-negative, it is natural to decompose it using a non-negative matrix factorisation²⁷ (NNMF) (for algorithmic details see the Methods section). Such an approach is akin to a principal component analysis, that reduces the dimension of the problem, thus allowing us to detect important network features. Mathematically, we approximate S as follows:

$$S \approx WH$$
, (1)

where $W \in \mathbb{R}^{m \times k}$ and $H \in \mathbb{R}^{k \times 212}$ are non-negative matrices. Here, k is the rank of the approximation and m the number of networks being considered. Importantly, both the columns of W and the rows of H can be used to reveal important network features 28,29 . Note, that in all of our experiments, the

factorisation in (1) was carried out using k = 3, and W, H were chosen so as to minimise the residual²⁵

$$||S-WH||_F$$
.

Here, $||\cdot||_F$ denotes the Frobenius norm (see the Methods section for further details concerning, for example, the choice of k).

The approach can be concisely summarised into the following three basic steps (see Figure 1 for a schematic description):

Step 1: For each metabolic network compute the significance profile, $\mathbf{s}_i \in \mathbb{R}^{212}$, consisting of the normalised significance scores for each of the 212 three- and four-node motifs.

Step 2: Compute a low-dimensional (k << 212) representation of the thresholded matrix of significance scores, $S = [\mathbf{s}_1, \dots, \mathbf{s}_m]^T$, using a non-negative matrix factorisation

Step 3: Use the columns/rows of W/H to determine important network features.

Global and local motif significance scores

In order to determine the relative importance of the *j*th motif in the *i*th network we construct the following local motif significance score:

$$P(i,j) = s_{i,j} \cdot h_{1,j}. (2)$$

Note that this results in a matrix $P \in \mathbb{R}^{m \times 212}$ (m = 115 or m = 383 here), whose rows encapsulate the network motif structure of each metabolic network, and whose columns provide information pertaining to the relative importance of specific motifs across the network ensemble.

In the experiments in the next section, we derive a *global* significance score for each network by summing the rows of P as follows

$$P_{\text{global}}(i) = \sum_{j} P(i, j) = \sum_{j} s_{i, j} \cdot h_{1, j},$$

$$= \mathbf{s}_{i} \cdot \mathbf{h}_{1}.$$
(3)

As alluded to by the second row in the above, this is equivalent to projecting the significance vector \mathbf{s}_i onto \mathbf{h}_1 , the first row of H. Note that in practice \mathbf{h}_1 is the row of greatest magnitude and thus is likely to provide the optimal single-variable projection of the data²⁵. Moreover, we consider the global significance score in (3) to be a proxy for network complexity, in the sense that a large value indicates the presence of a relatively large number of network motifs, whereas a low value is indicative of a simpler, more tree-like structure.

Environment	Number of Nodes			Number of Edges		
	min	median	max	min	median	max
Obligate (34)	78	273	620	91	340	840
Specialised (5)	442	480	541	566	641	692
Aquatic (4)	541	580	647	700	751	868
Facultative (41)	90	652	809	101	890	1160
Multiple (28)	430	615	800	560	821	1119
Terrestrial (3)	557	689	693	779	944	966
Total (115)	78	541	809	91	730	1160

Table 1 Network statistics for the reaction graphs of 115 bacterial species studied classified according to environmental variability. According to the NCBI³², obligate bacteria have the most constant environment, followed by specialised and aquatic, and then facultative, multiple and terrestrial bacteria in that order.

Environment	Number of Nodes			Number of Edges		
	min	median	max	min	median	max
Aerobic (154)	65	605	892	74	809	1210
Facultative (180)	78	602	816	91	825	1168
Anaerobic (49)	307	488	681	381	645	969
Total (383)	65	581	892	74	789	1210

Table 2 Network statistics for the reaction graphs of the 383 bacterial species studied classified according to species' oxygen requirements. The degree of oxygen required increases in the order anaerobic, facultative and aerobic.

Results and discussion

In this section we present the results of applying the approach described in the previous section to a large cohort of metabolic networks. We begin by giving a brief description of the organisms studied, and details of the network construction process.

Metabolic networks

The metabolic data in this study is the same as used by Takemoto³⁰, and was derived from the KEGG database³¹ on May 20th, 2011. In total, we studied upto 383 bacterial species (see Tables 1 and 2 for an overview of some basic network properties), each being characterised by a number of shared biological features (e.g. environmental variability, oxygen requirements and genome size), using graph theoretical techniques. A complete list of all the bacterial species used in our analysis is provided in the Supplementary Materials[†].

Metabolic processes can be modelled using simple graphs in a number of ways ³³ and it is important to choose an appropriate representation. The most common representation is the *substrate-product* graph whereby nodes and edges correspond to metabolites and reactions, respectively. Note, that a potential caveat of such an approach is that it can lead to the detection of erroneous pathways (see, for example, the discussion in Montañez *et al.* ³⁴). However, since we are not considering

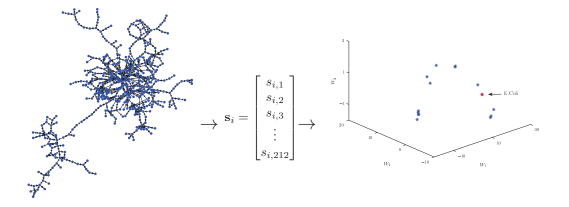


Fig. 1 A schematic illustration of our algorithmic approach in the case of *E. coli*. **Step 1:** by considering all 3- and 4-node motifs, the metabolic network of *E. coli* is mapped into a 212-dimensional Euclidean space. **Step 2:** a low-dimensional representation of the data point is obtained via a non-negative matrix factorisation. **Step 3:** important network features are determined by analysing the resultant low-dimensional representation (not shown).

a path analysis here and for the ease of comparability with related studies, we consider the substrate-product representation in all of our experiments. Moreover, ubiquitous metabolites such as $\rm H_2O$, ATP and NADH were removed from the analysis as they tend not to be involved in higher order functions, and if included, typically lead to physiologically meaningless pathways. Finally, to further simplify the analysis, we consider only the largest connected component for each network. This avoids, for example, issues that arise when constructing randomised networks through the rewiring of metabolic networks consisting of a number of disconnected components. For further details of the network construction the interested reader is referred to the papers of Takemoto and colleagues 11,30,35 .

It is worth noting that, in addition to the modelling issues touched upon above, there are general limitations to any complex network study of metabolism due to noisy and incomplete metabolic maps (e.g. missing/spurious links), the omission of reaction stoichiometry data and incomplete reaction reversibility data. Nevertheless, the approach taken here is standard within the field and provides a global picture of the biochemical systems under investigation.

As an illustration of the approach introduced in the previous section, we carried out two experiments with the aim of testing the hypothesis that organism adaptability is manifested via the network motif structure of the corresponding metabolic networks.

Habitat variability and network motif structure

The first experiment undertaken considered 115 metabolic networks, each being categorised according to their environmental habitat (see Table 1). The organisms can be found in a variety of conditions, ranging from highly specialised (e.g.

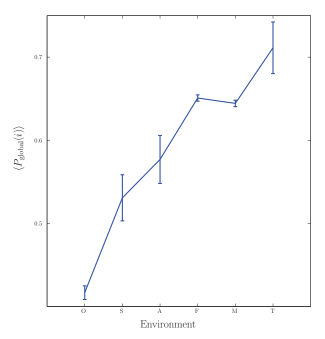


Fig. 2 Relationship between environmental variability and the mean global significance score $\langle P_{\mathrm{global}} \rangle$ for the six bacterial habitats: Obligate, Specialised, AQuatic, Facultative, Multiple and Terrestrial. Vertical bars represent standard errors.

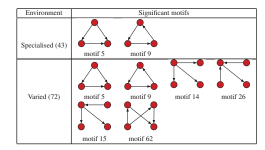


Table 3 Motifs significantly overrepresented in networks pertaining to a specialised and varied environment.

symbiotic bacteria living within a host), to extremely heterogeneous conditions such as soil, and thus have evolved under very different selective pressures.

Figure 2 shows a plot of the mean global significance score, $\langle P_{\mathrm{global}} \rangle$, versus environmental variability for the 115 different bacterial networks. Note that the average here is taken over each of the six environmental classes: obligate, specialised, aquatic, facultative, multiple and terrestrial. Importantly, we found that motif frequency, and thus network complexity, increased significantly with environmental variability. The lowest motif frequency was found for the bacteria within the obligate class, followed by a relatively steep increase to the specialised and aquatic classes, then higher again for the facultative and multiple classes, and then highest for the terrestrial class. The group differences shown in Figure 2 are statistically significant (Kruskal-Wallis test, $p < 10^{-9}$).

This result provides evidence supporting the view that variability in an organisms habitat has important consequences for the topology of the resultant metabolic networks, and is consistent with previous studies ^{10,12,36,37} that have demonstrated important links between the metabolic networks of organisms and their biochemical environments. In the current context, these results can be understood as an evolutionary effect due to the greater uncertainty that accompanies an increasingly fluctuating environment: greater numbers of 3- and 4-node motifs lead to larger numbers of cycles, i.e. closed paths, and thus to increased redundancy in the metabolic network, which in turn promotes greater adaptability and robustness.

The effect of oxygen requirement on network structure

Next, we considered the effect of oxygen requirements on metabolic network structure. We studied 383 bacterial species which were categorised into three groups: 154 aerobes, 180 facultatative aerobes and 49 anaerobes.

Figure 3 shows a plot of the mean global significance scores versus growth conditions for the 383 different bacterial species. Interestingly, we found that networks that have evolved in the presence of oxygen, i.e. aerobes and facultative

aerobes, have a significantly larger number of network motifs. The group differences shown in Figure 3 were found to be significant (Kruskal-Wallis test, $p < 10^{-4}$).

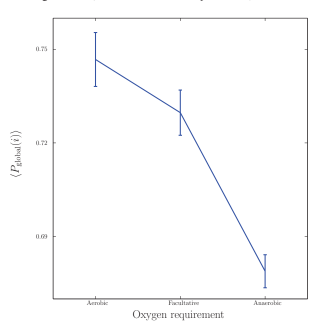


Fig. 3 Relationship between growth conditions, in particular oxygen requirements, and mean global significance score $\langle P_{\rm global} \rangle$. Vertical bars represent standard errors.

The results shown in Figure 3 are in agreement with recent studies (see, for example, the paper by Raymond and Segré ³⁸) demonstrating that bacterial networks that are exposed to oxygen are able to form additional pathways, compared to those that are oxygen deprived. In particular, the study by Raymond and Segrè ³⁸ found that aerobic bacteria have approximately a 1.5 fold increase in the number of reactions and metabolites relative to anaerobic bacteria, resulting in the expansion of metabolic networks evolving in the presence of oxygen, and thus supporting the view of oxygen induced network complexity.

Motifs responsible for the observed differences

To determine the specific motifs driving the observations of the previous section, we considered the quantity $\sum_i P(i,j)$, that is, the column sum of the matrix P defined in Equation (2) – recall that the columns of P contain information specific to individual motifs. Moreover, by restricting the sum above to a particular subgroup of interest (specialised, obligate, multiple, etc.), it is possible to detail the extent to which any particular motif featured within that group. In the following we consider a motif to be significant within a particular group, if the mean

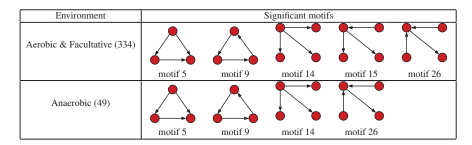


Table 4 Motifs significant to networks with differing oxygen requirements.

local significance score of that motif (restricted to the group of interest) is at least 2 standard deviations greater than the mean score across the entire network ensemble. Note that a list of all 3- and 4-node motifs is provided in the Supplementary Material.

Habitat variability In order to simplify the analysis we considered two groups: specialised (consisting of the obligate, specialised and aquatic classes) and varied (consisting of facultative, multiple and terrestrial classes). The significant motifs are displayed in Table 3. The first thing to note is that motifs 5 and 9, a feed forward loop and closed cycle, respectively, are prominent throughout the entire ensemble of networks, regardless of environmental factors. This is perhaps not too surprising as both of these patterns are considered to play important functional roles in many biological networks. The addition of a feed forward loop to a linear cascade of biochemical reactions, for example, has been hypothesised to accelerate the metabolic process³⁹. Importantly, we found the number of significant motifs to be greatest in those metabolic networks exposed to more variable environments: 2/212 for specialised and 6/212 for varied (see Table 3). Clearly, this represents only a very small percentage of available 3- and 4-node motifs ($\approx 1-3\%$), and so the differences observed in Figure 2 can be attributed to a small set of motifs more or less specific to the different kinds of bacteria.

The increased numbers of network motifs present within the varied class indicates a potentially significant growth in network redundancy within those organisms inhabiting fluctuating environments, and can be considered as further evidence of so-called *functional redundancy mediated robustness* 40 , that is, the observed perseverance of systems level redundancies prevalent in metabolic, as well as more general, cellular networks. More specifically, of the 4 additional significant motifs found in the varied class, motifs 14 and 15 may be considered variants of the single-input motif, motif 62 a bi-parallel fan, and motif 26 a multi-input motif, all of which have been implicated as potential indicators of network redundancy. For example, in the context of metabolism the single-input motif consists of a substrate X that is consumed in multiple reac-

tions, the result of which are the products Y,Z,...; whilst the bi-parallel fan implies the presence of multiple, or compensatory, pathways whose efficiencies may vary according to alterations in environmental conditions. Indeed, these findings are in agreement with a number of recent studies relating genetic robustness and organism adaptability 40,41 , and suggest that bacteria that live in more variable environments typically display a greater abundance of redundant metabolic reactions.

In addition to the topological differences observed between varied and specialised bacteria, we found that the distribution of those metabolites occurring within motif structures present across the entire network ensemble, i.e. motifs 5 and 9, also differed significantly. Figures 4 and 5 shows the mean frequency for metabolites occurring within motif 5 for the 115 metabolic networks, again grouped into the specialised (blue bars) and varied classes (red bars). Note that the frequencies plotted in Figures 4 and 5 have been normalised to remove any bias due to network size (see Methods section for further details), and metabolites are displayed in decreasing order according to the varied class. Figure 4 displays the distribution for those 263 metabolites that occurred at least once within motif 5 across the two classes under consideration. Interestingly, we see that the distribution for the varied class is relatively broad, with a large number of metabolites occurring with a relatively low frequency, whereas the distribution for the specialised class is more akin to a scale-free or power-law distribution, consisting of a small set of relatively high frequency metabolites. Note that similar results were found for motif 9 (see Supplementary Material).

Next we used a Chi-square test (Fisher's exact test, p < 0.01) to explore the differences in proportions of the individual metabolites between the two groups. Figure 5 shows the 47/263 metabolites for which a significant difference in proportions was found, again displayed according to decreasing frequency of the varied class. Metabolites displaying the most significant differences (Fisher's Exact test, $p < 10^{-5}$) included (2R)-2-Hydroxy-3-(phosphonooxy)-propanal, Tetrahydrofolate and Isopentenyl diphosphate, all of which were overrepresented in the specialised group compared to the varied group. Note that the aforementioned overrepre-

sented metabolites are required for biosynthesis of various amino acids, folates and terpenoids and are also responsible for the regulation of carbohydrate metabolism in many bacterial species.

Oxygen requirements Similar to the above, we then investigated which motifs were driving the observed differences between metabolic networks that evolved in the presence or absence of oxygen. Again, for simplicity we divided the bacteria into two separate groups: anaerobic and aerobic (including facultative aerobes). The significant motifs are displayed in Table 4. For aerobic networks 5/212 possible motifs were found to be significant, whilst for the anaerobic networks 4/212 were found to be significant. Again, motifs 5 and 9 were significant across the entire cohort, along with motifs 14 and 26 in this instance. The only motif that differed between the two groups was motif 15, which was specific to the aerobic class. Interestingly, the study by Raymond and Segré³⁸ found that the effects of oxygen exposure on metabolic network structure was most prolific at the periphery of the network, that is, network alterations were largely due to the addition of new reactions and pathways, rather than network rewiring. Thus, the enrichment of motif 15 is a natural consequence, as it acts as a branch point on these newly formed peripheral reactions and pathways.

Figures 6 and 7 shows the distribution of metabolites across motif 5 for the two groups, ordered according to decreasing metabolite frequency for the aerobic class (blue bars). Note, that the aerobic class exhibits a fairly broad distribution, whilst the anaerobic distribution tails off slightly quicker, in a similar but less pronounced manner to that displayed by the specialised bacteria in Figure 4. Figure 7 shows those metabolites that displayed a significant group difference. Interestingly, the majority of metabolites, some 37/52, were found to be overrepresented in the aerobic group compared to the anaerobic group, the most significant of which were Isopentenyl diphophosphate, Fatty acid, trans,trans-Farnesyl diphosphate, Phosphatidylethanolamine, Phosphatidylserine, L-Threonine, L-2-Amino-3-oxobutanoate, Phosphatidylcholine, 2-Acyl-sn-glycero-3-phosphocholine, L-2-Lysophosphatidylethanolamine, 3'.5'-Cyclic (Fisher's Exact test, $p < 10^{-5}$). These metabolites are known to be involved in the biosynthesis of a range of amino acids and secondary metabolites. Again, similar results where found for motif 9 (see Supplementary Material).

Conclusions

In this work, we have introduced a new graph embedding approach for studying large numbers of networks, of possibly differing order, and employed it to investigate the effect of environmental variability on the metabolic network structure

of a large cohort of bacterial species. Using the new technique, we found evidence supporting the view that organisms that evolve in more uncertain environments exhibit more complex metabolic connectivity structures than those evolving under more stable conditions. Note, that the motif based approach forwarded here strongly supports the view that environmental conditions play a pivotal role in shaping the resultant metabolic networks, and is robust in the sense that the patterns described in Figures 2 and 3 are reproducible in both the latest and older, less complete versions of the data 42 (data not shown). This is in contrast to recent studies in which network features that were found to correlate with environmental variability (e.g. modularity) disappeared when tested on newer versions of the data ^{30,43}. These findings suggest that alterations in the motif signature provide a robust indicator of adaptability and evolvability in bacterial metabolic networks.

Methods

Detection of network motifs and the choice of null model

Network motif frequencies were computed using the opensource software $mfinder^{23,44}$. To determine significance, motif frequencies were compared against frequency distributions for some 1000 random graphs, chosen so as to preserve both the in- and out degree, as well as (n-1)-node motifs. Note that the latter condition ensures that the enrichment of n-node motifs is not simply due to the presence of highly significant subgraphs.

Non-negative matrix factorisation

There are many different variants of the non-negative matrix factorisation algorithm 28 . In our work we used the Multiplicative Update algorithm which is included in the MATLAB statistics toolbox*. Starting with initial guesses W^0 , H^0 , typically random matrices, the method computes a rank-k approximation to the data matrix $A \approx WH$ (here $A \in \mathbb{R}^{n \times m}$, $W \in \mathbb{R}^{n \times k}$ and $H \in \mathbb{R}^{k \times m}$) via successive iterations of the equations

$$\begin{split} H^{i+1} &= H.* \frac{W^{iT}A}{W^iH^iH^{iT} + 10^{-9}} \\ W^{i+1} &= W^i.* \frac{AH^{iT}}{W^iH^iH^{iT} + 10^{-9}}. \end{split}$$

Here .* denotes point-wise multiplication.

Note that due to the iterative nature of the scheme, the resulting decomposition can vary depending upon (a) the choice of initial matrices W^0 , H^0 ; (b) the choice of the parameter k, which is usually not obvious *a priori*, and tends to be based on heuristics such as the number of expected clusters in the data,

*http://www.mathworks.com/

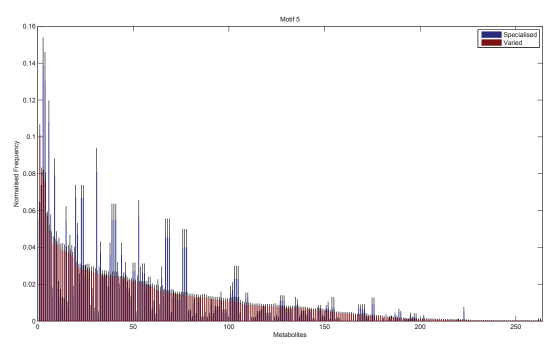


Fig. 4 Mean normalised frequency for the 263 metabolites obtained for the 115 metabolic networks. Blue bars represent the specialised class and red bars represent the varied class. Here, the metabolites are in descending order of the metabolite frequencies for the varied class.

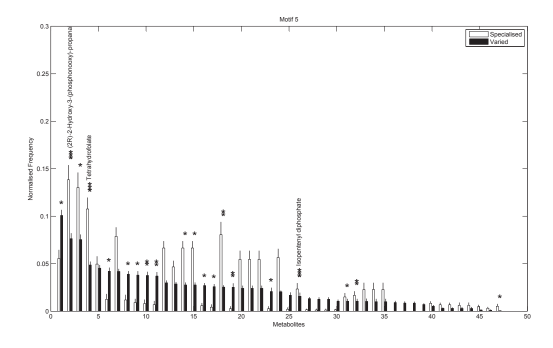


Fig. 5 Mean normalised frequency for the significant metabolites with p < 0.01 (Fisher's Exact test). Vertical bars are standard errors. Asterisks indicate large significant differences between metabolic networks from a specialised and varied environment, where *,**,and *** correspond to p < 0.001, p < 0.0001 and p < 0.00001. Metabolite names are provided for the most significant metabolites.

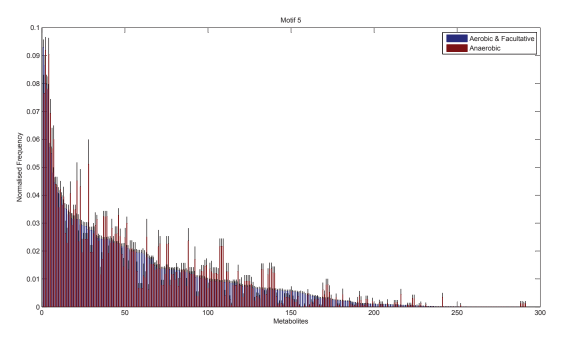


Fig. 6 Mean normalised frequency for the 291 metabolites obtained for the 383 metabolic networks. Blue bars represent the aerobic-facultative class and red bars represent the anaerobic class. Metabolites are displayed in descending order of the metabolite frequencies for the varied class.

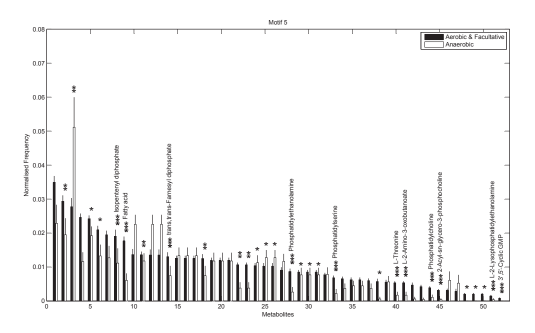


Fig. 7 Mean normalised frequency for the significant metabolites with p < 0.01 (Fisher's Exact test). Vertical bars are standard errors. Asterisks indicate large significant differences between metabolic networks from a specialised and varied environment, where *,**,and *** correspond to p < 0.001, p < 0.0001 and p < 0.00001. Metabolite names are provided for the most significant metabolites.

or a visual inspection of the scree plot. In our experiments, we ran the algorithm with 100 different random initial conditions, choosing as our factorisation the matrices W, H for which the residual $||A-WH||_F$ was minimised, according to the Frobenius norm. Moreover, we repeated the experiments for a range of different k values (up to and including k=25) to test the robustness of our results. We found the effects of varying k to be inappreciable, in the sense that the patterns reported were reproduced for most values of k.

Determining significant metabolites

When considering the differences between the frequency of metabolites occurring in a motif of interest (5 or 9 in our case) care must be taken to eliminate the influence of network size on the analysis. This bias is due to the increased number of motifs exhibited by larger networks which naturally leads to greater frequencies of metabolites. Thus, given a network i and a metabolite j, we denote by $f_{i,j}$ the frequency with which metabolite j appears within the motif of interest, motif q say, for the ith network. Now, in order to remove any bias due to network size we normalise the statistic $f_{i,j}$ by dividing it by the frequency with which motif q appears in network i, which we denote by f_{i,mot_q} . This then leads to the following normalised statistic:

$$\hat{f}_{i,j} = \frac{f_{i,j}}{f_{i,\text{mot}_a}},$$

describing the relative importance of metabolites via their participation within specific motifs.

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