Network motifs emerge from interconnections that favor stability

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The microscopic principles organizing dynamic units in complex networks from proteins to power-grid generators — can be understood in terms of network 'motifs': small interconnection patterns that appear much more frequently in real networks than expected in random networks [1, 2]. When considered as subgraphs isolated from a larger network, these motifs are more robust to parameter variations, easier to synchronize than other possible subgraphs, and can provide specific functionalities [3–15]. But one can only isolate these subgraphs by assuming, for example, a significant separation of time scales, and the origin of network motifs and their functionalities when embedded in larger networks remain unclear. Here we show that most motifs emerge from interconnection patterns that best exploit the intrinsic stability characteristics at different scales of interconnection, from simple nodes to whole modules. This functionality suggests an efficient mechanism to stably build complex systems by recursively interconnecting nodes and modules as motifs. We present direct evidence of this mechanism in several biological networks.

In complex natural systems, as biological networks, the particular topologies of network motifs have been shaped by evolution. Evolution and natural selection accumulate *stable* intermediate components (i.e., nodes), which are interconnected to form more complex systems. This modular design principle is observed at many scales, from the motion control architecture of vertebrates to the emotional response of human beings [16–19]. Yet, in general, the interconnection of stable components may result in an unstable system [20]. Thus, it is natural to hypothesize that nature favors interconnections that make it easier to obtain a stable networked system. In this letter, we show that most network motifs in real networks emerge precisely from such consideration. This property can be used for building larger systems by applying it at different scales of interconnection.

To start, we consider a set of N nodes with scalar dynamics of the form

$$\dot{x}_i = f_i(x_i, t) + u_i, \quad y_i = x_i \tag{1}$$

with initial condition $x_i(t_0) = x_{i0}, i = 1, \dots, N$. Here the scalars x_i , u_i and y_i are the state, input and output of node *i*, respectively. The state of a node may represent the expression level of a gene, the concentration of a metabolite, the charge of a capacitor, etc. Vector dynamics are discussed later in the context of modules. The functions $f_i(x_i, t)$, which are typically nonlinear, determine the nodal dynamics.

Nodes interact with each other by interconnecting their inputs $u = \operatorname{col}(u_1, \dots, u_N) \in \mathbb{R}^N$ with their outputs $y = \operatorname{col}(y_1, \dots, y_N) \in \mathbb{R}^N$. Analyzing the stability of networked systems with nonlinear interconnections

$$u = g(y), \quad g : \mathbb{R}^N \to \mathbb{R}^N,$$
 (2)

requires knowledge of the functional form and parameters of the interconnection g(y) and nodal dynamics $f_i(x_i, t)$, which is hard to obtain in most systems (SI-1.7). Linear interconnections do not require such knowledge, enabling us to quantify the contribution of the interconnection to the stability of the networked system without the need of detailed knowledge of the nodal dynamics. More precisely, by considering

$$u = Ay, \tag{3}$$

where $A = (a_{ij}) \in \mathbb{R}^{N \times N}$ is the weighted adjacency matrix of the interconnection network, our analysis requires a single constant per node — its *contraction rate*, defined later on characterizing its intrinsic stability properties. Here $a_{ij} \neq 0$ represents a directed edge from node *i* to node *j*. In general, the linear interconnection (3) can be used to approximate (2) in some working range [5, 7, 21]. Furthermore, diffusive coupling of oscillators and several models of neural networks actually use linear interconnection networks [22–24].

Our standing assumption on the isolated nodes is that they are stable, and we aim to quantify for which interconnections is easier to get a stable networked system. The separate contribution of the isolated nodes and the interconnection to the stability of the networked system is made transparent by using *contraction theory*. Contraction theory is a tool to analyze the stability of dynamic systems based on a differential-geometric viewpoint inspired by fluid mechanics [25], in contrast to Lyapunov stability theory that is based on analogs of mechanical energy. A system is *contracting* if the trajectories associated to any two initial conditions exponentially converge towards each other. More precisely, a dynamic system of the form

$$\dot{x} = f(x,t), \quad x(t_0) = x_0$$
(4)

with state $x \in \mathbb{R}^N$ is contracting with rate $\alpha > 0$ if there exists a vector norm $|\cdot|$ and constant $\beta > 0$ such that for any two initial conditions $x_a, x_b \in \mathbb{R}^N$ their corresponding trajectories $x(x_a, \cdot), x(x_b, \cdot)$ satisfy

$$|x(x_a,t) - x(x_b,t)| \le \beta |x_a - x_b| e^{-\alpha(t-t_0)}, \quad \forall t \ge t_0.$$

Denote by $J(x,t) = \partial_x f(x,t)$ the Jacobian of system (4). Then contraction is equivalent to the existence of a matrix measure μ such that $\mu(J(x,t)) \leq -\alpha$, for all $x \in \mathbb{R}^N$ and $t \geq t_0$ [25, 26]. Any vector norm $|\cdot|$ induces a matrix norm $||\cdot||$ and a matrix measure μ by

$$\|A\|\coloneqq \sup_{|x|=1} |Ax|, \quad \mu(A)\coloneqq \lim_{h\searrow 0} \frac{\|I+hA\|-1}{h}$$

both of which are well defined for any matrix $A \in \mathbb{R}^{N \times N}$. In particular, we prove that

$$\mu_A(A) \coloneqq \min_{\mu \in \mathcal{M}} \mu(A) = \max_{1 \le i \le N} \operatorname{Re}\lambda_i(A), \tag{5}$$

where \mathcal{M} is the set of all matrix measures in $\mathbb{R}^{N \times N}$ and $\lambda_i(A)$ are the eigenvalues of A (Theorem 1 in SI-1). We use the notation μ_A to emphasize that the matrix measure achieving the minimum in (5) depends on the matrix A itself. Recall also that matrix measures are subadditive: $\mu(A_1 + A_2) \leq \mu(A_1) + \mu(A_2)$, for any $A_1, A_2 \in \mathbb{R}^{N \times N}$.

In the case of scalar isolated systems, as in (1) with $u_i = 0$, contraction with rate α_i is equivalent to the condition $J_i(x_i, t) = \partial_{x_i} f(x_i, t) \leq -\alpha_i$ for all $x_i \in \mathbb{R}$ and $t \geq t_0$. The contraction property of isolated nodes might be lost when they are interconnected, so that the networked system is no longer contracting. Indeed, due to the subadditivity of matrix measures, the Jacobian of the networked system (1)-(3) satisfies

$$\mu(J(x,t)) \le \mu(\mathsf{diag}\{J_i\}) + \mu(A) \le \mu(-D_{\alpha}) + \mu(A), \tag{6}$$

where $J_i = J_i(x_i, t)$ and $D_{\alpha} = \text{diag}(\alpha_1, \cdots, \alpha_N)$.

We define $\mu(A)$ as the contraction loss of the interconnection network. Then the inequality (6) indicates that the networked system remains contracting if the effective contraction of the isolated nodes $\mu(-D_{\alpha}) < 0$ dominates the contraction loss $\mu(A)$ due to the interconnection. Consequently, interconnections with small contraction loss best favor stability, since they require smaller contraction rates from the isolated nodes to keep the whole network contracting.

The choice of matrix measure in (6) is a degree of freedom that should be optimized to make $\mu(-D_{\alpha}) + \mu(A)$ as negative as possible. Solving this optimization problem is not trivial, since the matrix measure minimizing the contraction loss $\mu(A)$ might also decrease the effective contraction of the isolated nodes (i.e., making the term $\mu(-D_{\alpha})$ less negative). In Proposition 1 of SI-1, we prove that the optimal matrix measure is given by μ_A defined in (5), provided that the contraction rates of all nodes are equal, or that the off-diagonal entries of A are non-negative (i.e., interactions between nodes are only positive). Choosing the matrix measure (5), the contraction loss of some classes of networks depends only on their topology. For example, Proposition 2a of SI-1.4 shows that acyclic networks have zero contraction loss regardless of their edge-weights (positive or not). This implies that feedforward and bifan interconnections, and their generalizations [27], always have zero contraction loss (SI-1.5). Moreover, non-positive contraction loss for interactions with arbitrary strength requires that reciprocal interactions have opposite signs, and the absence of cycles (feedback loops) of length 3 or more, Proposition 2b in SI-1.4. In particular, the contraction loss of negative feedback between two nodes is at most zero (SI-1.4).

Interestingly, the presence of negative interactions always improves the stability of the networked system by decreasing the contraction loss, in the sense that $\mu(A) \leq \mu(\bar{A})$ for any $\mu \in \mathcal{M}$ and $A = (a_{ij}) \in \mathbb{R}^{N \times N}$, see Lemma 1 in SI-1.3. Here $\bar{A} = (\bar{a}_{ij}) \in \mathbb{R}^{N \times N}$ is defined as $\bar{a}_{ii} = a_{ii}$ and $\bar{a}_{ij} = |a_{ij}|$ if $j \neq i$. Therefore, it is possible to add functionalities requiring negative interactions by replacing a positive interaction by a negative one without decreasing the stability of the networked system. This result motivated us to consider positive interactions only in the rest of the paper because (i) they provide the worst-case analysis of the contraction loss of an interconnection with respect to all possible edge-weights, and (ii) their contraction loss can be optimally computed using μ_A .

We analyzed the contraction loss of all 3- or 4- node subgraphs and identified those with the lowest contraction loss in their *density class*, defined as the set of all subgraphs with the same number of nodes and edges [5]. Those subgraphs with lowest contraction loss best favor the stability of the networked system. Since the precise value of the interactions may change from one system to other, we randomly select them from a uniform distribution on $[0, a_{max}]$ to form an ensemble of 10,000 weighted adjacency matrices with the same connectivity pattern. From this ensemble the *mean contraction loss* $\langle \mu_A \rangle$ is computed, see Fig.1. SI-2 details how to compute analytically and numerically the mean contraction loss of a subgraph. The observed order of the subgraphs according to their mean contraction loss is independent of the interaction strength distribution and the value of $a_{max} > 0$ (SI-3). See also SI-4 for results using a different matrix measure to compute the contraction loss.



FIG. 1. Mean contraction loss of all 3- or 4-node subgraphs. Error bars represent standard deviation. **A.** Mean contraction loss of all subgraphs with 3 and 4 nodes $(a_{max} = 1)$, same color indicating same number of edges. Vertical dashed lines separate groups of subgraphs with the same number of edges (density classes) in increasing order from left to right (2 to 6 edges for 3-node subgraphs, and 3 to 12 edges for 4-node subgraphs). Gray markers show the network motifs reported in [1], circles denoting biological networks (gene transcription, neurons and food webs) and triangles denoting man-made networks (electronic circuits and the WWW). Motif M2 appears in both, gene regulatory networks and some electronic circuits (forward logic chips). **B.** The 12 network motifs reported in [1].

We find that all motifs reported in [1] (except M_3 , M_9 and M_{12} that contain feedback loops of length 3 or more) emerge among the subgraphs with the minimum $\langle \mu_A \rangle$ within their respective density class. In particular, all motifs found in biological networks (marked in circles in Fig.1) have zero contraction loss. Motifs containing feedback loops with length 3 or more do not have the minimum contraction loss within their density class —they do not favor stability— and these motifs also require more accurate tuning of their edge-weights to be stable (SI-1.4). Recall also that feedback provides functionalities associated with performance (like robustness to external disturbances [25]), which do not necessarily favor the stability of the networked system. To further disentangle the relation between network motifs and subgraphs with low contraction loss, we compared the Z-score and *relative* contraction loss of subgraphs in several real networks. As introduced in [1], the Z-score of a subgraph A in a real network quantifies its statistical significance as a motif, and is defined by

$$Z(A) \coloneqq \frac{N_{\mathsf{real}}(A) - \langle N_{\mathsf{rand}}(A) \rangle}{\sigma_{\mathsf{rand}}(A)},$$

where N_{real} is the number of occurrences of subgraph A in the real network, $\langle N_{\text{rand}} \rangle$ the average number of occurrences in an ensemble of its randomizations, and σ_{rand} its standard deviation. A subgraph with a high (low) Z-score is over (under) represented in the real network. The *normalized* Z-score of a subgraph is its Z-score divided by the maximum (in absolute value) Z-score of all subgraphs with the same number of nodes.

We define the *relative contraction loss* of a subgraph A as

$$r(A) \coloneqq \frac{\langle \mu_A(A) \rangle - \mu_{\min}}{\mu_{\max} - \mu_{\min}},$$

where μ_{max} (or μ_{min}) is the maximum (or minimum) mean contraction loss among all subgraphs within the density class of A. The case r(A) = 0 (or r(A) = 1) corresponds to a subgraph with the minimal (or maximal) mean contraction loss among its density class. The relative contraction loss is undefined for all subgraphs with 3-nodes/2-edges or 4-nodes/3edges (i.e., $\mu_{\text{min}} = \mu_{\text{max}} = 0$), since they are acyclic and thus have zero contraction loss (Proposition 2a of SI-1.4). Hence, we discard them from the discussion that follows since subgraphs with zero contraction loss maintain the stability of the networked system.

We compared the relative contraction loss and the normalized Z-score of 3- and 4-node subgraphs in several biological networks, finding that overrepresented subgraphs (e.g., motifs) tend to have low relative contraction loss, see Fig. 2. The phenomenon is stronger for 3-node subgraphs, where underrepresented subgraphs (e.g., anti-motifs) have high relative contraction loss. In other words, subgraphs that favor stability are overrepresented, while 3-node subgraphs which do not favor stability are underrepresented. We did not find this phenomenon in other classes of networks containing feedback motifs with high Z-score (like the electronic circuits shown in SI-6), suggesting that other factors apart of maintaining stability play a central role in their construction.

Next we explore how the small contraction loss property of motifs can be used to build bigger networked systems. Consider a set of N modules (i.e., connected subgraphs) possibly



FIG. 2. Relative contraction loss vs. normalized Z-score. Each marker represents a subgraph of the neuron network of C. *elegans*, the gene transcription networks of Yeast and E. *coli*, and the food-web at Saint Martin Island. Subgraphs with high Z-score tend to have small relative contraction loss. In the case of 3-node subgraphs, under-represented subgraphs (anti-motifs) tend to have high relative contraction loss.

having vector dynamics

$$\dot{x}_i = f_i(x_i, t) + B_i u_i, \quad y_i = C_i x_i, \tag{7}$$

 $i = 1, \dots, N$, where $x_i \in \mathbb{R}^{n_i}$, $u_i \in \mathbb{R}^{m_i}$ and $y_i \in \mathbb{R}^{p_i}$ are the state, input and output vectors of module *i*. The matrices $B_i \in \mathbb{R}^{n_i \times m_i}$ and $C_i \in \mathbb{R}^{p_i \times n_i}$ determine which nodes of the module interact with other modules. The interconnection of modules is again described by equation (3), but the matrix $A \in \mathbb{R}^{(m_1 + \dots + m_N) \times (p_1 + \dots + p_N)}$ is not necessarily square anymore because some modules may have different number of inputs and outputs.

Each isolated module is assumed contracting with rate $\alpha_i > 0$ under measure μ_i . The contraction rate of a module can be calculated using the contraction rate of its internal nodes and their respective interconnection topology A_i . To each module, we associate a *condensed* node, with scalar state and linear dynamics, that inherits the module's contraction rate

$$\dot{z}_i = -\alpha_i z_i + u_i, \quad y_i = z_i. \tag{8}$$

Additionally, we use the interconnection network of the full system to define a *condensed*

weighted adjacency matrix $A_{cond} \in \mathbb{R}^{N \times N}$ as follows:

$$A_{\text{cond}} \coloneqq \begin{bmatrix} \mu_1(M_{11}) & \|M_{12}\|_{1,2} & \cdots & \|M_{1N}\|_{1,N} \\ \|M_{21}\|_{2,1} & \mu_2(M_{22}) & \cdots & \|M_{2N}\|_{2,N} \\ \vdots & \ddots & \vdots \\ \|M_{N1}\|_{N,1} & \|M_{N2}\|_{N,2} & \cdots & \mu_N(M_{NN}) \end{bmatrix},$$
(9)

where $M_{ij} = B_i A_{ij} C_j$, and $A_{ij} \in \mathbb{R}^{m_i \times p_j}$ is the (i, j) block of the original A interconnection network (3), see Fig S1 in supplement. Above $\|\cdot\|_{i,j}$ stands for the induced matrix norm

$$\|M\|_{i,j} \coloneqq \sup_{|x|_i=1} |Mx|_j$$

with $|x|_i = |P_i^{1/2}x|_2$ a weighted Euclidean norm with metric $P_i \in \mathbb{R}^{n_i \times n_i}$ found as the solution to the linear matrix inequality of Theorem 1 in SI-1. When the off-diagonal elements of A_i are non-negative, a diagonal solution to such matrix inequality exists and the metric P_i just assigns different units to different modules (SI-1). Also, in the case when each module has a single input and a single output, A_{cond} takes a particular simple form in which its (i, j)entry is $|\gamma_{ij}A_{ij}|$ if $i \neq j$ and $\gamma_{ij}A_{ij}$ if i = j, with $\gamma_{ij} = B_i^{\mathsf{T}}C_j \in \mathbb{R}$.

In Theorem 2 of SI-1 we prove that if the *condensed networked system* (8)-(9) is contracting, then the original system (7)-(3) is also contracting. This result also holds when instantaneous contraction rates are used for the modules, making the condensed dynamics (8) dependent on the states of the original dynamics (SI-1.6). Hence, the interconnections between modules have minimal contraction loss if they are also network motifs. This suggests a modular design principle to build complex systems, starting by building modules interconnecting nodes as network motifs, and then interconnecting those modules again as network motifs.

To better illustrate this point, consider the feedback interconnection of three 3-node motifs shown in Fig. 3A. Each isolated motif, will be contracting if

$$-\alpha_j \coloneqq -\alpha_{j,\min} + \mu_j(A_j) < 0, \quad j = 1, 2, 3,$$

where $\alpha_{j,\min}$ is the minimum contraction rate of the nodes inside the *j*-th motif, and A_j is its internal interconnection. Indeed, α_j is just the contraction rate of motif *j*. The smaller is the contraction loss of the internal topology, the larger is the contraction inherited by the module. Since the contraction loss of feedforward motifs is zero, in this example each



FIG. 3. Relative contraction loss of motifs at different scales of interconnection. A. Interconnection of motifs as motifs. The original network is condensed using (8) and (9) into the condensed network. Contraction of the condensed network ensures contraction of the original network. B. Original and two consecutive condensations for the Yeast transcription network. Network motifs are recursively found and condensed into a single node, see SI-5 for details. C. Number of motifs vs. relative contraction loss for the original and condensed networks. The Saint Martin food-web network is not shown since it does not contain motifs after a single condensation.

condensed node inherits the minimal contraction rate of its nodes, i.e., $\alpha_j = \alpha_{j,\min}$. To construct the condensed interconnection network, we first note that

$$B_{1} = C_{2}^{\mathsf{T}} = \begin{bmatrix} 0\\0\\1 \end{bmatrix}, B_{2} = C_{3}^{\mathsf{T}} = \begin{bmatrix} 1\\0\\0 \end{bmatrix}, B_{3} = C_{1}^{\mathsf{T}} = \begin{bmatrix} 0\\1\\0 \end{bmatrix}$$

The interconnection of the modules is described by the adjacency matrix of the 3-node feedback interconnection motif $A \in \mathbb{R}^{3\times3}$, whose only nonzero values are A_{12}, A_{23} and A_{31} (see Fig. 3A). Then, it is not surprising that the corresponding A_{cond} obtained using (9) is again the adjacency matrix of a 3-node feedback interconnection. The condensed interconnected system will be contracting if $\alpha_{\min} = \min\{\alpha_1, \alpha_2, \alpha_3\} > \mu_{\text{cond}}(A_{\text{cond}}) = \mu_A(A)$. Additionally, Theorem 2 of SI-1 implies that under such condition the original interconnected system is also contracting. Since the constraints were imposed in the contraction rates only, the details of the node dynamics were not used in the analysis.

The contraction loss of the interconnection plays an important role in the stability of the whole network, because modules inherit a larger contraction rate when their internal interconnection has a smaller contraction loss. And this applies at different scales of interconnection: if the system is recursively condensed, the resulting condensed modules at each step inherit larger contraction rates when the interconnections between the modules in the previous step have smaller contraction loss. In this form, the interconnection of "motifs of motifs" is a recursive and modular network design procedure in which the contraction loss remains minimal at each step of construction of the network. Both humans and nature seem to favor this design principle by recursively interconnecting already designed modules [28, 29].

The idea of "motifs of motifs" was used in [30] to reverse-engineer electronic circuits and coarse-grain a signal-transduction protein network. In contrast, here we aim to check if motifs at different scales still have low relative contraction loss, thus providing direct evidence of a design principle found in real-world complex networks. We used a collection of real networks to test our hypothesis by recursively searching and condensing motifs (details of the method and used networks are found SI-5 and SI-7, respectively). We found that most motifs in the original and condensed regulatory networks of *E. coli* and Yeast have low relative contraction loss, Fig. 3. For the neuron network of *C. elegans* this only happens for 3-node motifs. A closer analysis reveals that most 4-node motifs in the *C. elegans* with high relative contraction loss also have small Z-score, see Fig. 2. In other words, they are not strongly overrepresented.

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