

Revealing in-block nestedness: Detection and benchmarking

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(Received 17 January 2018; published 5 June 2018)

As new instances of nested organization—beyond ecological networks—are discovered, scholars are debating the coexistence of two apparently incompatible macroscale architectures: nestedness and modularity. The discussion is far from being solved, mainly for two reasons. First, nestedness and modularity appear to emerge from two contradictory dynamics, cooperation and competition. Second, existing methods to assess the presence of nestedness and modularity are flawed when it comes to the evaluation of concurrently nested and modular structures. In this work, we tackle the latter problem, presenting the concept of *in-block nestedness*, a structural property determining to what extent a network is composed of blocks whose internal connectivity exhibits nestedness. We then put forward a set of optimization methods that allow us to identify such organization successfully, in synthetic and in a large number of real networks. These findings challenge our understanding of the topology of ecological and social systems, calling for new models to explain how such patterns emerge.

DOI: [10.1103/PhysRevE.97.062302](https://doi.org/10.1103/PhysRevE.97.062302)**I. INTRODUCTION**

The identification of macroscale connectivity patterns has been central to the development of network science. Beyond the inherent methodological challenges of this task, ascertaining them is of relevance to the specific disciplines and to the area as a whole, inasmuch they are the outcome of distinct microscopic mechanisms of network formation. It is in this context, i.e., understanding network architecture as an emergent feature, that nestedness and modularity arise as prominent macrostructural signatures to study.

The concept of *nestedness* was first coined in biology to characterize the spatial distribution of biotas in isolated, yet spatially-related, landscapes [1] and later found to describe large families of interspecies cooperative relations [2]. In structural terms, a perfectly nested pattern is such that the set of connections of any given node is a subset of the relationships of larger degree ones [3]; see Fig. 1 (left). Nestedness has imposed itself as a landmark feature in mutualistic interactions, with an emphasis in natural ecosystems, triggering a large amount of research spanning fieldwork [2], modeling [4], and simulation [5]. Beyond natural systems, nestedness emerges as well in social, technical, and economic systems, e.g., industrial relationships [4,6,7], international trade [8], information ecosystems [9], anthropology [10], and knowledge

production [11]. In socioeconomic systems, the epitome of this property in unipartite networks, the emergence of nestedness is originated in agents attempting to maximize their own centrality [12–14].

On the other side, the identification of *modular* patterns in networks stands as one of the hallmarks in the area with prominent precedents in social network analysis [15]. Besides social systems, networks with significant community structure, see Fig. 1 (middle), appear in multiple contexts [16], like biology [17] or cognitive science [18]. It implies the existence of subgroups of nodes, strongly connected within but loosely connected to nodes outside. The identification and analysis of community structure constitutes itself a subarea of network science. It poses challenges with respect to detection algorithms, empirical problems, applications, and conclusions derived [19].

Nestedness and modularity have been often treated as incompatible architectures, since they are thought to emerge from conflicting (respectively, cooperative and competitive) dynamics [20]. Thus, most studies have focused exclusively on either of them. The existence of systems which combine both patterns has been largely overlooked, despite challenging indications in natural [21–26] and social ecosystems [9]. As of now, the proper identification of such compound structures lays beyond the capabilities of state-of-the-art techniques.

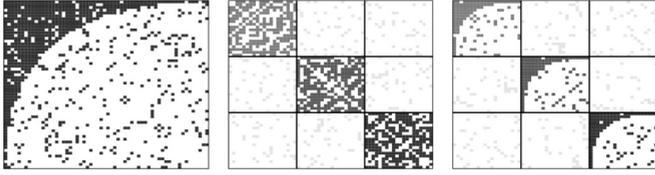


FIG. 1. Left: Example of network with *nested structure*. Rows and columns have been ordered by degree. Center: Example of a network with *block structure*; *blocks* are typically referred to as *communities* and *compartments* in the network science [16] and the ecology [21] literature, respectively. The network exhibits high internal connectivity between nodes of the same block and low connectivity between nodes that belong to different blocks. Right: Example of a network with *IBN structure*. Nodes within blocks exhibit nested structure. Vertical and horizontal lines are a visual guide to show the existing blocks.

In the scarce existing literature we identify two different approaches. The first operates in parallel, measuring modularity and nestedness independently [9,23,27], with the obvious drawback that these properties are treated as emerging unrelatedly. The second approach operates sequentially: After a proper identification of a partition (usually in terms of modularity [28]), it computes the nestedness (usually in terms of node overlap-decreasing fill (NODF) [29]) locally for each block [22,24]. In consequence, modularity takes functional precedence relegating nestedness from a macro- to a mesoscopic pattern. Both approaches overlook that these two network patterns are inherently intertwined and thus cannot be evaluated using *independent* metrics.

To be precise, the presence of modules places hard limits to the extent of nestedness that a network can exhibit [Fig. 2(a)], and, in the other direction, detected communities in a globally nested system lead to aberrant, hardly interpretable modules [22] (see Fig. 2(f) and the related discussion in Sec. III). As is expected, the modularity score is sensitive to the number of communities [Fig. 2(b)] but not to the shape of the nested structure. Overall, these *in-block nested* (IBN) structures are highly undetectable if the network contains few communities and/or the nested structure within the communities is very stylized (Fig. 1, right).

Beyond the methodological challenges, there exists an important epistemological aspect which cannot be overlooked. In most scenarios, the boundaries of the system under consideration are imprecise because the researchers, albeit involuntarily, impose a discretionary observation scale to it. Extending the realm of observation, the network structure can parsimoniously be expected to show a set of loosely interconnected blocks. This is particularly evident in natural ecosystems [30], where, in general, no precise geographic boundaries can be defined, but also in social networks, when it comes to decide which subjects should be included or not in a specific study.

The paper is organized as follows: To overcome the limitations of existing approaches, Sec. II introduces a compact methodological framework that jointly considers both patterns (block—or compartmental—structure and within-block nestedness). Our methodology can unveil the existence of IBN structures, as shown in Sec. III for a suitable benchmark. After discussing the possible resolution limits of our proposal

(Sec. IV), we investigate, in Sec. V, the question of how general (or anecdotal) such property is. We show that a large number of real datasets exhibit in-block nested structures that would have gone undetected under conventional modularity optimization/nestedness detection techniques. Our findings indicate that these previously overlooked structures are in fact common in ecological and social systems. This opens a new direction for the structural analysis of ecological and social systems, discussed in Sec. VI, calling as well for new models to explain how IBN structures emerge.

II. DEFINITION AND QUANTIFICATION OF IN-BLOCK NESTEDNESS

In this section, we develop a proper formulation of the problem of determining to what extent a given network is organized as loosely interconnected *blocks*, each of them internally nested. We begin by defining in a congruent manner both, global nestedness and the new in-block nestedness fitness \mathcal{I} . In particular, maximizing the IBN fitness function \mathcal{I} allows us to unveil the best node partition in terms of IBN structure. We analyze synthetic networks to show that \mathcal{I} -maximization allows us to reconstruct ground-truth IBN structures that would have gone undetected under the widely used modularity optimization. We then proceed to analyze a large set of real networks—originated in the most varied disciplines—to evince that this type of structures is indeed a common occurrence in both uni- and bipartite networks of diverse nature.

A. A nestedness measure

In this section we will present a suitable objective function to detect in-block nested structures in bipartite networks. Later, we indicate how the objective function can be extended to unipartite networks. Consider a bipartite network, describing a relationship between two sets $G = \{s, t, \dots\}$ and $\Gamma = \{\sigma, \tau, \dots\}$ with cardinalities N_r and N_c , respectively. The bipartite network can be represented as a binary adjacency matrix \mathbf{A} whose elements are $A_{s,\tau} = 1$ if a relationship between elements s and τ exists or zero otherwise. In the same spirit as the measure [29] NODF, we introduce the global nestedness fitness $\tilde{\mathcal{N}}$, which measures the amount of global overlap between row and column pairs

$$\tilde{\mathcal{N}} = \frac{2}{N_r + N_c} \left[\sum_{s,t} \frac{O_{s,t}}{k_t(N_r - 1)} \Theta(k_s - k_t) + \sum_{\sigma,\tau} \frac{O_{\sigma,\tau}}{k_\tau(N_c - 1)} \Theta(k_\sigma - k_\tau) \right], \quad (1)$$

where k_i corresponds to the degree of the element i (regardless of whether it belongs to G or Γ); $\Theta(\cdot)$ is the Heaviside step function (such that the only contributing terms are those in which the outer index has larger degree than the inner); and $O_{\cdot,\cdot}$ measures the degree of overlap between row and column pairs as

$$O_{s,t} = \sum_{\nu=1}^{N_r} A_{s\nu} A_{t\nu}, \quad O_{\sigma,\tau} = \sum_{\mu=1}^{N_c} A_{\mu\sigma} A_{\mu\tau}. \quad (2)$$

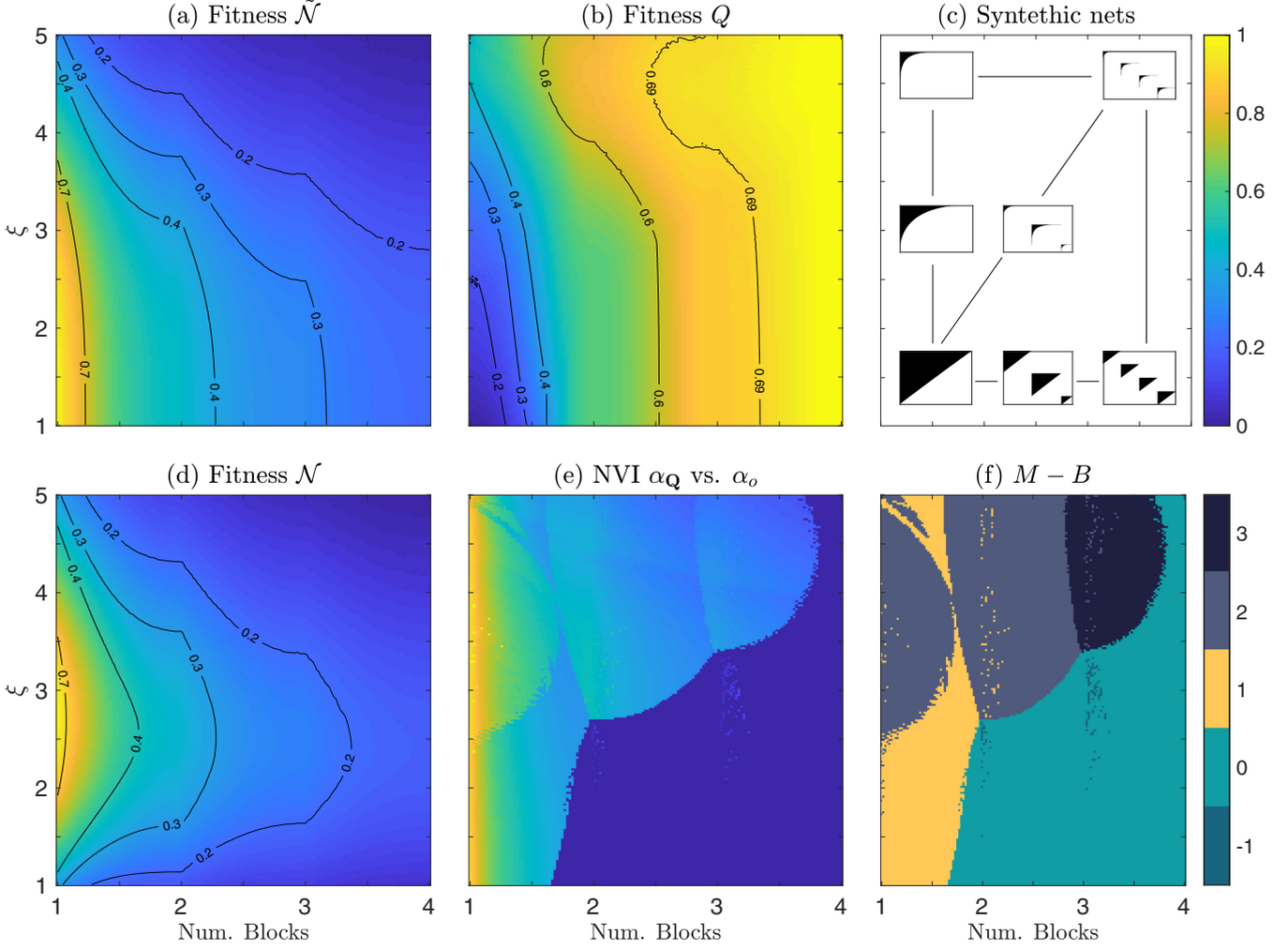


FIG. 2. Behavior of nestedness metrics $\tilde{\mathcal{N}}$ and \mathcal{N} and modularity Q in noiseless IBN synthetic networks. Panel (a) illustrates how global nestedness $\tilde{\mathcal{N}}$ decreases with the number of blocks in a network—regardless of their internal perfectly nested structure. Notice that the shape parameter ξ [see Eq. (5)] also affects $\tilde{\mathcal{N}}$ as a side effect of the strict decreasing condition imposed in the original definition of NODF. Panel (b) illustrates how modularity Q increases with the number of blocks in a network. Panel (c) illustrates the kind of networks generated by the model (see Sec. III A). The vertical axis represents the shape parameter ξ that quantifies the slimness of the nested structures. The horizontal axis stands for the number of modules. In noninteger settings (e.g., 2.5) there is an integer amount of nested blocks of the same size (e.g., 2) and one block of relative size equal to the remaining fraction (e.g., 0.5 of the size of the other communities). Panel (d) shows the global nestedness fitness \mathcal{N} which incorporates a null model, as defined by Eq. (5). Panel (e) shows the normalized variation of information between the modules detected by modularity optimization and the ground-truth (GT) blocks. Panel (f) shows the difference $M - B$ between the number M of modules detected by the modularity optimization and the number B of ground-truth blocks. Note that bottom-right color bar refers to panel (f) only.

It is important to remark that Eq. (1) weighs linearly the contribution of rows and columns to $\tilde{\mathcal{N}}$ (instead of quadratic weights as in NODF [29]). This is preferable when the difference between the number of rows and columns is considerable.

B. Recasting nestedness at the mesoscale level

We now introduce two new elements that allow us to generalize $\tilde{\mathcal{N}}$ to the case where nested structures exist at a mesoscopic scale: a membership variable and a null model. First, we consider that both sets of nodes are partitioned into C disjoint subsets, termed *blocks*. This implies that for each node i , it is possible to define a membership variable α_i . Based

on this, the total size of block ℓ can be obtained as

$$C(\ell) = \sum_{s=1}^{N_r} \delta(\alpha_s, \ell) + \sum_{\sigma=1}^{N_c} \delta(\alpha_\sigma, \ell), \quad (3)$$

where δ is the Kronecker delta. In addition, $C_s = \sum_t \delta(\alpha_s, \alpha_t)$ and $C_\sigma = \sum_\tau \delta(\alpha_\sigma, \alpha_\tau)$ give, respectively, the number of nodes in the block nodes s and σ belong to. The block overlap $O_{\cdot, \cdot}$, now including the membership variable, can be obtained as

$$O_{s,t} = \sum_{v=1}^{N_c} A_{sv} A_{tv} \delta(\alpha_s, \alpha_v),$$

$$O_{\sigma,\tau} = \sum_{u=1}^{N_r} A_{u\sigma} A_{u\tau} \delta(\alpha_u, \alpha_\sigma).$$

The null model gauges the expected overlap between a pair of nodes belonging to a class and aims at compensating the nestedness that can be explained solely by the nodes' degrees. The expected overlap, given two nodes s and t with degrees k_s and k_t , is obtained considering that the neighbors of each node are chosen uniformly at random. In this situation, the probability that both nodes share a common neighbor is simply $(k_s k_t)/N_c^2$ and therefore the expected amount of shared neighbors, i.e., the expected overlap, is given by $\langle O_{s,t} \rangle = k_s k_t / N_c$. The same argument shows that for the columns, $\langle O_{\sigma,\tau} \rangle = k_\sigma k_\tau / N_r$.

We can now introduce the in-block nestedness fitness \mathcal{I} , which quantifies to which extent a network exhibits IBN,

$$\mathcal{I} = \frac{2}{N_r + N_c} \left\{ \sum_{s,t}^{N_r} \left[\frac{O_{s,t} - \langle O_{s,t} \rangle}{k_t (C_s - 1)} \Theta(k_s - k_t) \delta(\alpha_s, \alpha_t) \right] + \sum_{\sigma,\tau}^{N_c} \left[\frac{O_{\sigma,\tau} - \langle O_{\sigma,\tau} \rangle}{k_\tau (C_\sigma - 1)} \Theta(k_\sigma - k_\tau) \delta(\alpha_\sigma, \alpha_\tau) \right] \right\}. \quad (4)$$

In this expression, some normalization factors have disappeared after straightforward simplifications. In the same spirit as in the row and column weighting of $\tilde{\mathcal{N}}$ [Eq. (1)], the per-block nestedness aggregates are weighted by the size of the block (i.e., C_s and C_σ). Notice that, for each pair of row nodes, $O_{s,t}$ only accounts for column nodes within the same block, while $\langle O_{s,t} \rangle$ considers all column nodes regardless of the block they belong to. This implies that, for any pair of rows, \mathcal{I} will be in principle larger when they are assigned to the same block: In this case the difference $O_{s,t} - \langle O_{s,t} \rangle$ has positive contributions. On the other hand, the membership variable α allows us to discard some of the comparisons, assigning row nodes to different communities. In general, an algorithm that correctly maximizes \mathcal{I} will attempt to discard pairs whose contribution is negative to the aggregate. This intuition is equivalent for columns. The balance of such “merge-split” strategy for rows and columns allows an algorithm to identify in-block nested structures by maximizing the objective function \mathcal{I} . In this work, we have adopted a biologically inspired optimization algorithm [31]. However, \mathcal{I} 's formulation—which closely follows that of modularity Q —enables the adoption of many existing heuristics (see Ref. [16] for an extensive review).

Equation (4) is equally valid for unipartite networks, simply imposing that the sets G and Γ are equal. That is, each row node and its column counterpart have the same class. This simplifies the optimization, by reducing the number of unknowns to the number of nodes in the networks.

Noteworthy, the objective function \mathcal{I} reduces to $\tilde{\mathcal{N}}$, corrected by a suitable null model, if one considers a single block ($\alpha_s = \alpha_\sigma = \alpha, \forall s, \sigma$), i.e.,

$$\mathcal{N} = \frac{2}{N_r + N_c} \left[\sum_{s,t}^{N_r} \frac{O_{s,t} - \langle O_{s,t} \rangle}{k_t (N_r - 1)} \Theta(k_s - k_t) + \sum_{\sigma,\tau}^{N_c} \frac{O_{\sigma,\tau} - \langle O_{\sigma,\tau} \rangle}{k_\tau (N_c - 1)} \Theta(k_\sigma - k_\tau) \right].$$

While the difference between NODF and $\tilde{\mathcal{N}}$ is slight—except when $N_r \ll N_c$ or vice versa—the null-model correction in \mathcal{N}

heavily alters the nestedness measure. In particular, note that fully connected nodes do not contribute to \mathcal{N} —as opposed to maximum contribution in the original formulation.

To illustrate this point, it is instructive to consider the case of one single nested block [Num. Blocks = 1 in Fig. 2(a), 2(c) and 2(d)] (note that Fig. 2 is explained in detail in the next section). In this cut, the contrast between noncorrected measure of nestedness [Fig. 2(a)] and corrected [Fig. 2(d)] is very clear: When the perfectly nested network is very dense (bottom-left corner, $\xi < 2.5$), most of the nodes have large expected overlap with the few hubs. Hence, even though the nestedness condition is respected for all the pairs of nodes, the nestedness metric $\tilde{\mathcal{N}}$ only deviates little from its expected value under the null model, resulting in a small value of \mathcal{N} . In other words, the observed level of nestedness $\tilde{\mathcal{N}}$ can be simply explained by the network degree distribution. On the contrary, $\tilde{\mathcal{N}}$ and \mathcal{N} are practically identical for $\xi > 2.5$. In this region, the slimness of the nested structure and the strict decreasing connectivity condition [Heaviside function in Eqs. (1) and (4)] heavily limit the value of nestedness, regardless of the consideration of a null model (\mathcal{N}) or not ($\tilde{\mathcal{N}}$).

III. DETECTION OF IN-BLOCK NESTED STRUCTURES IN SYNTHETIC NETWORKS

In this section, we first introduce a benchmark graph model with planted in-block nested structures (Sec. III A) and then present the results for the Q and \mathcal{I} optimization algorithms' performance in reconstructing such planted structures (Sec. III B).

A. In-block nested structures generator

In a perfectly nested structure, rows (and columns) interact with a subset of the neighbors of the rows (and columns) of larger degree. Correctly ordering its adjacency matrix by row and column degree, it resembles an upper (possibly with some curvature) secondary diagonal matrix. Inspired by the p -norm unit ball equation, we synthetically generate such structures using

$$y = f_n(x) = 1 - (1 - x^{1/\xi})^\xi, \quad (5)$$

where $x \in [0, 1]$ and $\xi \in [1, \infty)$ dictate how stylized the shape of the nested structure is. The adjacency matrix of the nested structure with N_c nodes is constructed tessellating the $[0, 1] \times [0, 1]$ space into N_c^2 squares and then adding a link into each matrix position whose center lies above the curve in Eq. (5). While an approach based on threshold graphs would have been also used, the degree sequences it produces are stochastic, introducing unnecessary fluctuations to the network generation [32].

Such noiseless nested structures are rarely found in real systems. Thus, on top of the previous scheme, we mimic random and uncorrelated noise using a dual-step procedure. In the first stage, we randomly remove links from the perfectly nested structure with probability p . Given a network with E_c edges, pE_c of them will be removed in average. In the second step, the removed edges are randomly distributed across the empty elements of the adjacency matrix. These include initially empty positions [i.e., those lying below the function in Eq. (5)]

and empty positions resulting from the stochastic removal procedure. Parameter value $p = 0$ corresponds to the initial structure and $p = 1$ corresponds to an Erdős-Rényi network with average degree E_c/N_c .

The construction of an adjacency matrix of an ideal in-block nested structure starts off with B (a real-valued number) blocks. Specifically, we build $\lfloor B \rfloor$ blocks of size $\lfloor N_c/B \rfloor$ and another with the remaining $N_c - \lfloor N_c/B \rfloor$ nodes. In the previous, $\lfloor \cdot \rfloor$ stands for the integer part function. In this way, the network produced has some level of heterogeneity (albeit the size of all blocks remains in the same order of magnitude). Then $A_{\mathcal{I}}$ can be constructed repeating the procedure for each block, A^c , and joining them to compose a block diagonal matrix,

$$A_{\mathcal{I}} = \begin{bmatrix} A^{c_1} & A^{o_{11}} & \dots & A^{o_{1B}} \\ A^{o_{21}} & A^{c_2} & \dots & A^{o_{2B}} \\ \vdots & \vdots & \ddots & \vdots \\ A^{o_{B1}} & A^{o_{B2}} & \dots & A^{c_B} \end{bmatrix}, \quad (6)$$

where $A^o = \mathbf{0}$ is a matrix of the required size.

Similarly to the intrablock noise, we reproduce interblock perturbations with an additional dual-step procedure controlled by the parameter $\mu \in [0, 1]$. In this first step, for each block, each link is removed with probability $p_i = \mu(B - 1)/B$. In the second step, those links are distributed at random to connect one node of the original block with a random node of a different block. Probability p_i depends on the number of blocks, since our purpose is that for $\mu = 1$ the amount of links within each block is the same as the amount of links connecting any two distinct blocks. In the limit situation of $p = \mu = 0$ the outcome corresponds to a noiseless in-block nested structure and for $p = \mu = 1$ the outcome corresponds to an Erdős-Rényi network with the same average degree as the germinal noiseless structure, see Appendix A.

The described generative process can be locally implemented in terms of edge probabilities. Along these lines, the probability of having a link between nodes i and j within a block becomes

$$P(A_{ij}^c) = [(1 - p + p p_r)] \Theta[j N_c - f_n(i N_c)] + p_r \{1 - \Theta[j N_c - f_n(i N_c)]\} (1 - p_i), \quad (7)$$

where Θ is the Heaviside function. The term within square brackets is related to the intrablock noise. In the first term, $(1 - p)$ corresponds to the probability of not altering the link. The second, $p p_r$, corresponds to the probability of recovering a link, after removal, in the random dispersion of removed links. These two terms are restricted, by Θ function, to the region where links exist in the noiseless structure. The third term, $p_r = p E_c (N_c - E_c + p E_c)^{-1}$, corresponds to the probability of selecting link A_{ij} in the random distribution of removed links. Eventually, the term $(1 - p_i)$ corresponds to the probability of not removing the link in the process of generating interblock noise.

The probability of a interblock link is

$$P(A_{ij}^o) = \frac{2 E_c p_i}{2(B - 1) N_c^2} = \frac{\mu E_c}{N_c^2 B}. \quad (8)$$

The numerator accounts for the amount of removed links from the blocks related to the off-diagonal block A^o , that

is, compartments i and j . The denominator accounts for the possible places where each of those links can be placed. Note that the “2” on both numerator and denominator explicitly shows that each removed link of compartment A^{c_k} can be reallocated in A^{o_k} or $A^{o_{\bar{k}}}$.

The noisy version given by Eqs. (7) and (8) of the original noiseless in-block nested structure generates a network with equivalent average degree. This is formally proved in Appendix A. An example of these synthetically generated structures is shown in Fig. 2(c) and Fig. 3(c).

B. IBN optimization applied to synthetic networks

In Fig. 2, we unveil the limitations of current techniques to detect IBN structures in noiseless networks (IBN networks) where no links exist between nodes belonging to different blocks. In a more realistic setting, where links can connect nodes that belong to different blocks, such weaknesses become even more apparent (see Fig. 3).

We generated synthetic (unipartite) networks of N_c nodes and E_c edges within blocks, where the level of in-block nestedness and the number of interblock links can be varied in a controlled manner by means of few parameters (see Sec. III A): the number of blocks B , the shape parameter ξ , the in-block nestedness parameter p , and the mixing parameter μ . To allow for heterogeneity in block size, we build $\lfloor B \rfloor$ blocks of size $\lfloor N_c/B \rfloor$ and another with the remaining nodes. ξ determines the density of the network, controlling how stylized the nested structure is. The in-block nestedness parameter p gives the fraction of links that do not respect the notion of perfectly nested organization within a block and the mixing parameter μ measures the fraction of interblock links.

The experiments in Fig. 2 correspond to 3×10^4 networks generated with parameters $p = \mu = 0$, with varying number of blocks and number of edges [see Fig. 2(c) for an illustration of the resulting adjacency matrices]. By construction, all networks have maximum \mathcal{I} , in the range (0.16, 0.90), depending on the shape parameter ξ and number of blocks B . Unsurprisingly, modularity Q increases as the number of block increases [Fig. 2(b)]. We find that in this setting, the modules detected by the modularity-maximization algorithm can be very different from the planted blocks, as measured by the NVI [34,35], Fig. 2(e). The difference between the detected and planted blocks is larger for sparser networks [upper region of Fig. 2(e)]. Figure 2(f) shows the difference between the number M of detected modules by the Q maximization and the number B of planted blocks. The difference is nonzero for a large region of the parameter space. In particular, the modularity optimization algorithm detects more than one module in a network composed of one single block with internal nested structure [see Fig. 2(f), left corner]. This happens because the modularity-optimization algorithm tends to form a module that only contains the nodes with largest degree. Figures 2(e) and 2(f) indicate that modularity optimization is only reliable in the limit of large number of blocks and dense networks (lower-right corner).

The results of Figs. 2(b), 2(e) and 2(f) make clear that measuring modularity and nestedness as two independent network properties is inherently flawed: modularity-optimization algorithms detect more than one module in a network composed of

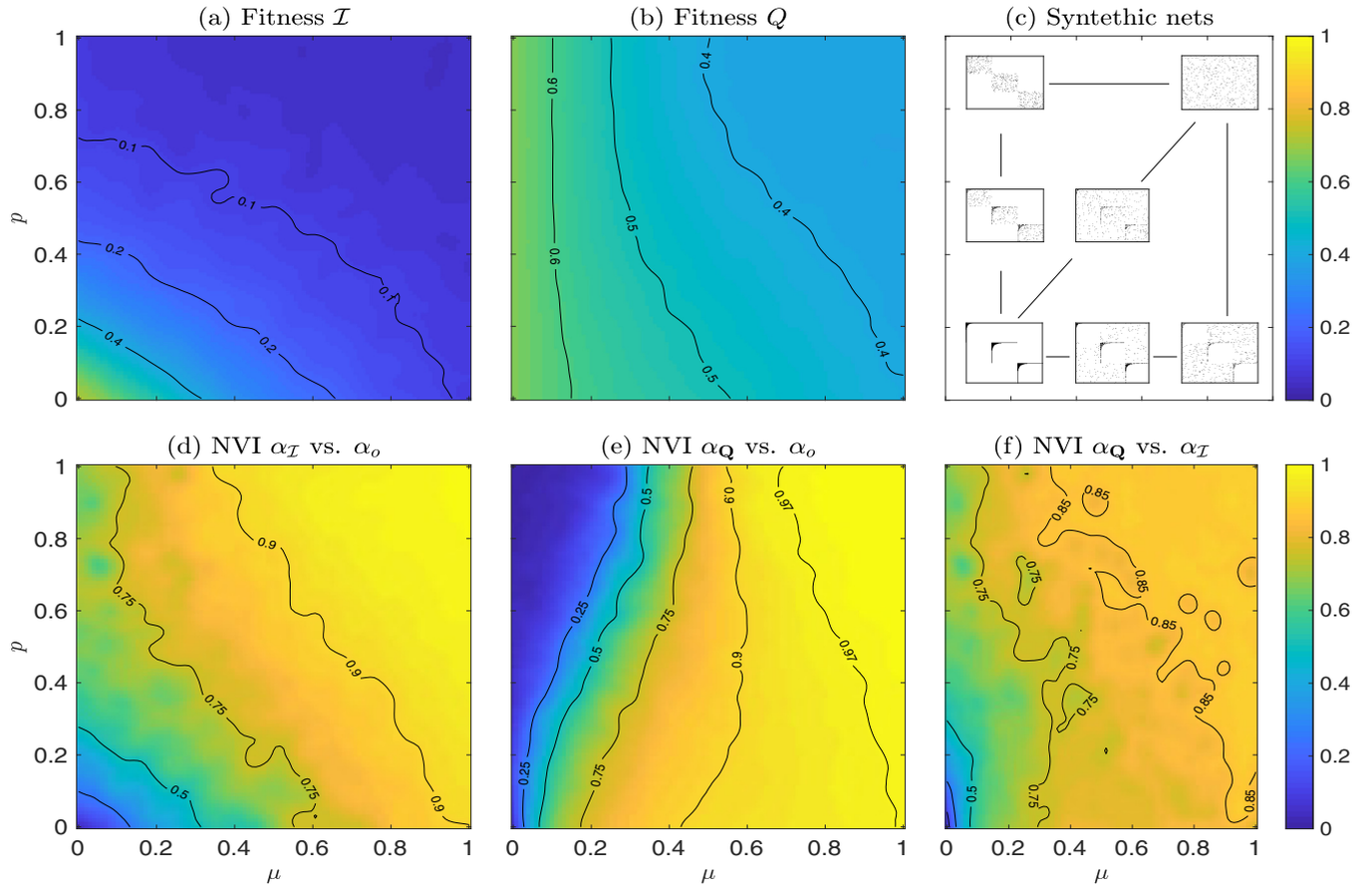


FIG. 3. Results for synthetic IBN networks with $B = 3$ and $\xi = 3$ for varying values of p and μ . Panel (a) displays the value of the fitness function \mathcal{I} as obtained with a biologically inspired optimization algorithm. Similarly, panel (b) reports the value of modularity Q as obtained with the Combo optimization algorithm [33]. Panels (d) and (e) compare the \mathcal{I} -optimized ($\alpha_{\mathcal{I}}$) and the Q -optimized (α_Q) partitions, respectively, to the planted partition (α_0) via the normalized variation of information (NVI). Panel (c) provides a visual intuition of the effect of parameters p and μ on the synthetically generated networks. Panel (f) shows the difference between the \mathcal{I} -optimized ($\alpha_{\mathcal{I}}$) and the Q -optimized (α_Q) partitions as measured by their NVI.

a single nested block, and the NVI between detected modules and planted blocks is in general large. We have verified that the \mathcal{I} -optimization algorithm introduced in this paper overcomes these limitations and is able to correctly recover the planted structure for all the parameter values shown in Fig. 2.

Figure 3 shows the results for an exhaustive exploration of the (p, μ) parameter space over 2600 networks with a fixed $B = 3$ and shape parameter $\xi = 3$ [see Fig. 3(c) for an illustration of the resulting adjacency matrices]. Results on these synthetic networks after a modularity optimization process [Fig. 3(b)] show that Q is only mildly affected by the p parameter (i.e., by the level of IBN). This itself is a consequence of the fact that Q does not consider any particular structure within the blocks, but only their internal density.

Figure 3(a) shows the value of in-block nestedness fitness, \mathcal{I} , after a maximization procedure based on a biologically inspired optimization algorithm (see Methods). Evidently, \mathcal{I} is sensitive to both the modular structure, and the nested organization within, taking a maximum value for $\mu = 0$ and $p = 0$. When we increase the randomness in either dimension the obtained in-block nestedness fitness smoothly decreases, reaching a global minimum when $\mu = 1$ and $p = 1$.

Since it can be assumed that the block structure is known *a priori*, it is possible to quantify how far the detected partition is from the planted one. As before, we resort on NVI to assess the quality of the partitions obtained optimizing \mathcal{I} [Eq. (4)] and the quality of the partitions obtained maximizing Q [33]. Focusing on the similarity of the Q -detected partition with respect to the prescribed one [Fig. 3(e)], we see that modularity recovers the planted partition in the full range of p values and low μ , implying that Q cannot discriminate between structured (nested) and nonstructured blocks. A good approach to detect IBN structures must be sensitive to both parameters p and μ . Remarkably, changes in NVI are closely independent of the parameter p , related to the level of disorder within each block. In contrast, we see that \mathcal{I} optimization allows us to unveil the planted partition for a region along the μ axis, as long as p remains low [Fig. 3(d)]: The presence of internal nestedness compensates the tenuous identity of the blocks, caused by large μ . The parameter region corresponding to low p and large μ is also the region where Q -detected partitions and \mathcal{I} -detected partitions differ the most [Fig. 3(f)]. This points out that the Q -detected partitions are particularly unreliable when there is a clear internal nested structure and there exist a significant number of interblock links.

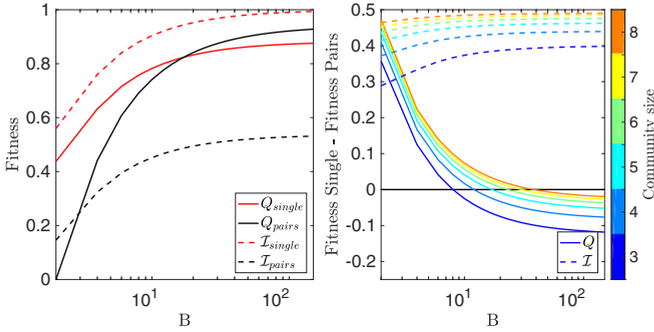


FIG. 4. Left: Q_{single} (solid-red line) and Q_{pairs} (solid-black line) as a function of the number of nested blocks, B (note semilog scale). As in the case of fully connected cliques, Q for this ring of dense subgraphs exhibits a paradoxical behavior, showing a crossover that disregards the *natural* partition favoring a larger-scale one. $\mathcal{I}_{\text{single}}$ (dashed-red line) and $\mathcal{I}_{\text{pairs}}$ (dashed-black line) for the same configuration. Unlike Q , \mathcal{I} uncovers the *natural* partition for all the range of B , i.e., $\mathcal{I}_{\text{single}} > \mathcal{I}_{\text{pairs}}, \forall B$. Furthermore, the ratio $\mathcal{I}_{\text{single}}/\mathcal{I}_{\text{pairs}} \approx 0.5$. Right: Difference between $\mathcal{I}_{\text{single}}$ and $\mathcal{I}_{\text{pairs}}$ (dashed), and between Q_{single} and Q_{pairs} (solid), for different block sizes (note semilog scale). Results show that that the conclusions extracted on the left panel hold as well for different settings.

To see how results vary in terms of the ξ parameter, we present results for $\xi = 4$ in Appendix B. We emphasize that modularity does not recover the planted partition in any parameter configuration, not even at $\mu = 0$, and it is almost insensitive to changes in the parameters of the model, its range is quite narrow, $0.55 < Q < 0.7$. Even further, the value of modularity fluctuates around $Q \approx 0.6$, in remarkable accordance with the predictions in Guimerà *et al.* [36], for sparse graphs, like those obtained with $\xi = 4$.

IV. RESOLUTION LIMIT

Before confronting the proposed methodology to empirical networks, it is convenient to analyze the possible existence of a resolution limit in \mathcal{I} , along the lines of the well-known detectability issues inherent to modularity [37]. Far from anecdotal, Q 's resolution limit implies that a certain detected partition may be at risk of missing important structures at smaller scales—which triggered a wealth of research on the topic [38,39]. To illustrate this situation, Fortunato and Barthelemy propose the simple case of a ring of fully connected (clique) subnetworks, loosely interconnected to the neighboring ones. Intuitively, the optimal partition of such network is the one where the communities correspond to the planted cliques, i.e., Q_{single} . However, increasing the number of communities in the ring yields at some point a different optimal Q_{pairs} , i.e., the one in which pairs of consecutive cliques are considered as single communities. Such behavior, analytically studied in Ref. [37] (and visually evident in Fig. 4, left panel), is ever more paradoxical, provided that *the local structure of each clique has not changed at all*, i.e., the ratio between internal and external links remains constant despite the addition of cliques to the ring.

We consider here a similar setting, where the elements of the ring are not cliques—the optimal structure for Q —but rather perfectly nested blocks; see Appendix C. Figure 4 (left panel)

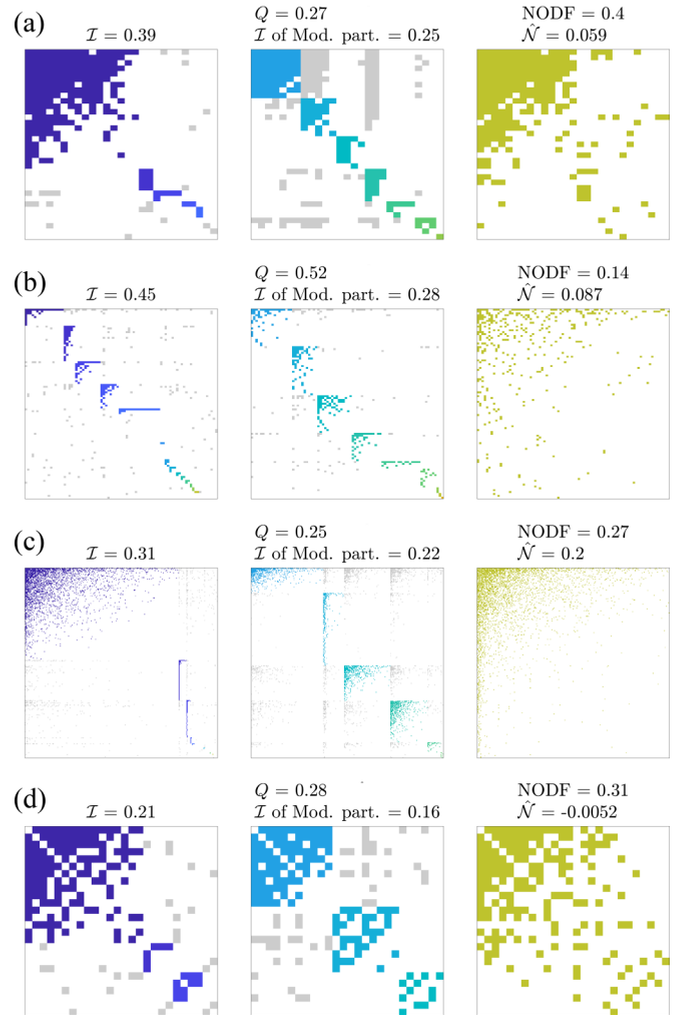


FIG. 5. Interaction matrices of four illustrative systems: (a) a host-parasite competitive bipartite network in the Volga-Kama Nature Reserve; (b) a pollination mutualistic bipartite network in Cordón del Cepo, Chile; (c) an urban bipartite network accounting for citizen visits to city services in Chennai, India; and (d) a unipartite friendship network in a Dutch school class. For all of them, rows and columns have been arranged to highlight different aspects: block membership and degree for \mathcal{I} - (left) and Q -maximization (center) partitions; global degree ranking in the right column.

monitors the evolution of $\mathcal{I}_{\text{single}}$ and $\mathcal{I}_{\text{pairs}}$ against the growth of the ring (expressed as B , the number of nested blocks). This plot shows that there is no evidence of a resolution limit for IBN, because the dependency of $\mathcal{I}_{\text{single}}$ and $\mathcal{I}_{\text{pairs}}$ on the number of blocks (and any other configuration, for that matter) show a parallel behavior, i.e., there will be no crossover between these two quantities. Moreover, the right panel of Fig. 4 confirms this behavior for a few other configurations.

The explanation to this unexpected result becomes apparent by noticing that \mathcal{I} [Eq. (4)] is built up on the sum over normalized, pairwise connectivity overlaps, with the following effect: Once a perfectly nested structure is found, further additions of new nodes to the block will be detrimental for the global score, unless they have a perfectly nested relation with the nodes already identified as members of the block. This is very different from modularity, where the addition of

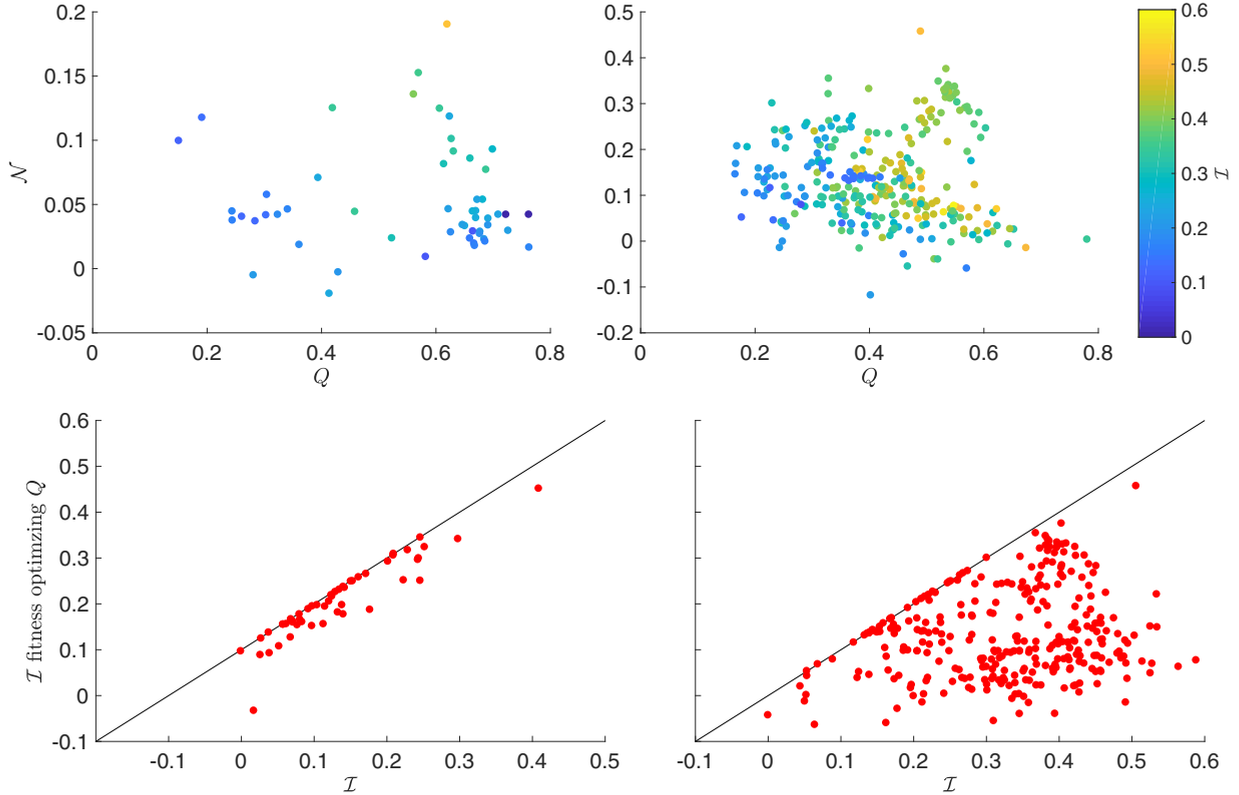


FIG. 6. Top: Scatter plot confronting modularity and nestedness with the \mathcal{I} measure (color coded). Each point represents an ecological, urban, or social network (left panel: unipartite networks; right panel: bipartite networks). Note that bipartite networks have been analyzed under the formulation of Barber’s modularity [42]. Bottom: Comparison of in-block nestedness value obtained optimising modularity or directly \mathcal{I} , as we propose.

nodes to a community always increases the positive term of Q , as long as these new nodes are nonisolated with respect to the community. Appendix C discusses an extension of this idealized setting, showing that, contrary to modularity, the detectability of small blocks increases as we increase the network size. These findings suggest that the study of resolution limits in in-block nested structures is an intriguing problem that deserves a deeper analysis in the future.

V. DETECTION OF IN-BLOCK NESTED STRUCTURES IN REAL DATASETS

The previous sections demonstrate the adequacy and robustness of \mathcal{I} —and the inherent flaws of modularity Q and global nestedness \mathcal{N} —to unveil IBN structures. However, those analysis would be limited to a mere academic exercise in absence of ample (in terms of examples and origin) empirical evidence. To demonstrate the practical aspects of the proposed methodology, we have analyzed a total of 334 networks, including both unipartite (57) and bipartite (277) ones which are known to display some level of nested organization. Most of them (209 bipartite networks) belong to ecology [40]—mostly mutualistic networks—and the rest belong to online platforms (68 bipartite networks) and social networks (57 unipartite networks) [41].

As a visual intuition, Fig. 5 displays the adjacency matrix of four of these networks, where rows and columns have been sorted following different criteria: For left and central columns (\mathcal{I} - and Q -maximizing partitions, respectively), nodes in the

same block are placed together, and they are ranked by degree (within blocks) to make more apparent a possible IBN structure; in the right column, nodes are simply ranked by degree. Figure 5(a) shows such arrangements for a host-parasite network (see A_HP_050 in Ref. [41]). Clearly, the three matrix representations look very different. In this case, \mathcal{I} favors the existence of a large, highly nested block, and a set of smaller clusters with a clear internal organization as well, whereas Q renders several, similarly sized, highly dense modules with no clear internal nested organization. Even though the classical NODF measure hints at some degree of global nestedness, taking into account the null model ($\mathcal{N} = 0.059$) seems to indicate that the nested organization is a simple consequence of the network’s degree distribution. Figure 5(b) shows the results for a pollination mutualistic network (see M_PL_001 in Ref. [41]). The system exhibits a clear IBN structure that cannot be detected through the maximization of modularity. From the results in Figs. 5(a) and 5(b), it is worth remarking that the observation of IBN structures in ecosystems with different types of interactions demands a reconsideration of which patterns should or should not be expected in them. Figure 5(c) shows the results for a urban user-service network (see Chennai in Ref. [41]). We observe again that global nestedness fails to characterize the predominant organization of the system, i.e., an IBN structure. Figure 5(d) shows the results for a unipartite network representing friendship relations in a Dutch school class (see c2 in Ref. [41]). The conclusions of the analysis of this network are similar to the ones in Fig. 5(a).

While Fig. 5 is helpful to get an intuition of how \mathcal{I} , Q , and \mathcal{N} work, these representations may be misleading. For example, the degree-ranked representation in Fig. 5(b) (right) conveys the idea that this particular plant-pollinator network is clearly (and globally) nested—but the other two arrangements are qualitatively convincing as well. For this reason, we have systematically compared the results of \mathcal{N} and Q , on one side, and \mathcal{I} , on the other, for the whole set of real networks mentioned above. In Fig. 6(a), two color-coded scatter plots are shown for uni- (left) and bipartite (right) networks. Strikingly, modularity Q and in-block nestedness \mathcal{I} are not strongly correlated in real datasets. Networks that exhibit small or intermediate values of modularity (compatible with those of a random network [36]) may show high \mathcal{I} , regardless of the \mathcal{N} score. Also, large values of Q —which unsurprisingly display nestedness ≈ 0 —indeed exhibit both large and small \mathcal{I} scores as well. Beyond the uncorrelated behavior between the three descriptors, what surfaces here is the fact that when analyzing data we may be overlooking a relevant pattern—IBN structure—just because two partial views of it (\mathcal{N} and Q taken independently) appear to be nonsignificant.

Further, in Fig. 6(b) we show the value of \mathcal{I} for partitions obtained by maximizing Q confronted to the maximization of \mathcal{I} itself. This plot evidences that modularity optimization may sometimes render partitions which do have some in-block organization (near the diagonal), but most often it is blind to it. This result highlights that using an approach where modularity is maximized, to successively evaluate nestedness within the blocks identified (i.e., the approach in Refs. [22,24]), is not able to unveil the IBN structure in most real-world networks.

VI. CONCLUSIONS

The emergence of structural patterns in complex networks is a consequence of the dynamics that take place on them. While the ultimate goal is to understand how these dynamics operate, this is not feasible until the correct methods to identify those patterns are available. The increasing evidence that nestedness and modularity appear in many empirical contexts, the already abundant hints that they may appear together, and the importance of both to disentangle how they affect—and are affected by—the evolution of a system prompt the need of rethinking the strategy to detect the occurrence of in-block nestedness.

Inspired by the NODF and modularity optimization, in this paper we have developed a methodology to detect in-block nested structures. The objective function at the core of this method naturally embeds a suitable null model to discount the in-block nestedness of the network that can be ascribed to randomness. Beyond the formal correctness of our formulation, demonstrated by means of a suitable benchmark to generate synthetic networks, we have shown that it overcomes the inherent limitations of nestedness and modularity (as independent methods) for this task. In structural terms, our approach can be interpreted as a generalization of the concept of nestedness (as expressed in NODF), reframing it to the mesoscopic network scale. Along this line, the door remains open to further development. For example, slight modifications on our formulation generalize the notion of multiple core-periphery structures, which has been recently

addressed from a different starting point [43]. Other directions may be related to identification techniques (e.g., stochastic block models [44]) or the design of distinct, specialized null models [45,46]. Prominently, an analytical account of the \mathcal{I} 's resolution limit—or lack thereof—is missing, beyond the preliminary intuitions offered in this work.

The analysis of real data has shown that many networks display in-block nested structure, regardless of their NODF and modularity scores. This finding suggests that previous works may have overlooked important features when discussing the organization of real systems and connects with previous works in diverse areas, such as network neuroscience [47,48]. The existence of in-block nested structures affects the debate around population dynamics, in ecology especially, in terms of which patterns maximize survival [5] and why. Our methodological contribution thus uncovers the need for models—beyond host-parasite [49,50]—that explain how networks transition between possible configurations: from modular to combined to purely nested architectures, as suggested by the emergence of collective attention processes [9], or from nested to combined architectures, as one would expect in a growing, but highly structured, system with increasing specialization. Noteworthy, it is not clear whether these processes are reversible, as they may respond to different systemwide adaptive processes.

ACKNOWLEDGMENTS

C.J.T and M.S.M acknowledge financial support of the University of Zürich through the University Research Priority Program on Social Networks.

APPENDIX A: AVERAGE DEGREE OF THE SYNTHETIC NETWORK GENERATION MODEL

The synthetic in-block nested network generator we propose in Sec. III A include two parameters to generate noise. Only as a summary, parameter p controls the intra-compartment noise and parameter μ controls the intercommunity noise. The noisy versions of the initial germinal noiseless in-block nested structure has the same average degree. We first show that adding intracommunity noise does not alter the average number of edges. Under this situation $\mu = 0$ and Eq. (7) becomes

$$P(A_{ij}^c) = [(1-p) + pp_r] \Theta[jN_c - f_n(iN_c)] + p_r \{1 - \Theta[jN_c - f_n(iN_c)]\} \quad (\text{A1})$$

and the expected number of edges within a compartment is $\langle E_c \rangle = \sum_{ij}^{N_c} P(A_{ij}^c)$. Reorganizing common terms of the sums we obtain that

$$\begin{aligned} \langle E_c \rangle &= \sum_{ij}^{N_c} P(A_{ij}^c) \\ &= [(1-p) + pp_r] \sum_{ij}^{N_c} \Theta[jN_c - f_n(iN_c)] \\ &\quad + p_r \sum_{ij}^{N_c} \{1 - \Theta[jN_c - f_n(iN_c)]\}. \end{aligned} \quad (\text{A2})$$

The heaviside step function is one when a link was existing in the original noiseless compartment. Thus, by definition, $E_c = \sum_{ij}^{N_c} \Theta[jN_c - f_n(iN_c)]$, and, consequently, $N_c - E_c = \sum_{ij}^{N_c} \{1 - \Theta[jN_c - f_n(iN_c)]\}$. Substituting, this to Eq. (A2) leads to

$$\begin{aligned} \langle E_c \rangle &= (1 - p + pp_r)E_c + p_r(N_c^2 - E_c) \\ &= E_c - pE + p_r(N_c - E_c + pE_c) = E_c. \end{aligned} \quad (\text{A3})$$

We now consider the case where $\mu \neq 0$. The expected amount of edges within the full in-block nested structure is

$$\begin{aligned} \langle E \rangle &= B \sum_{ij}^{N_c} P(A_{ij}^c) + B(B-1) \sum_{ij}^{N_c} P(A_{ij}^o) \\ &= B(1 - p_i) \sum_{ij}^{N_c} \{[(1 - p) + pp_r]\Theta[jN_c - f_n(iN_c)]\} \\ &\quad + B(1 - p_i) \sum_{ij}^{N_c} \{p_r\Theta[f_n(iN_c) - jN_c]\} \end{aligned}$$

$$\begin{aligned} &+ B(B-1) \sum_{ij}^{N_c} \frac{E_c \mu}{N_c^2 B} \\ &= B(1 - p_i)E_c + B(B-1) \frac{\mu E_c}{N_c^2 B} \sum_{ij}^{N_c} 1 \\ &= B \left[1 - \frac{\mu(B-1)}{B} \right] E_c + (B-1)\mu E_c = BE_c. \end{aligned} \quad (\text{A4})$$

Q.E.D.

APPENDIX B: THE EFFECT OF ξ ON Q AND \mathcal{I} PERFORMANCE

Figure 3 above illustrates the capabilities of Q and \mathcal{I} when confronted with a rich benchmark of synthetic networks, which span three relevant architectures—in-block nested to modular to random. While the main lessons from Fig. 3 hold in general, both Q and \mathcal{I} are quite sensitive to the density of the networks, which our toy model encapsulates under the parameter ξ . Here Fig. 7 presents the same results as Fig. 3 above, with sparser structures ($\xi = 4$).

The sparsity of these synthetic networks may lead to counterintuitive results, such as the fact that, in Fig. 7(e), NVI

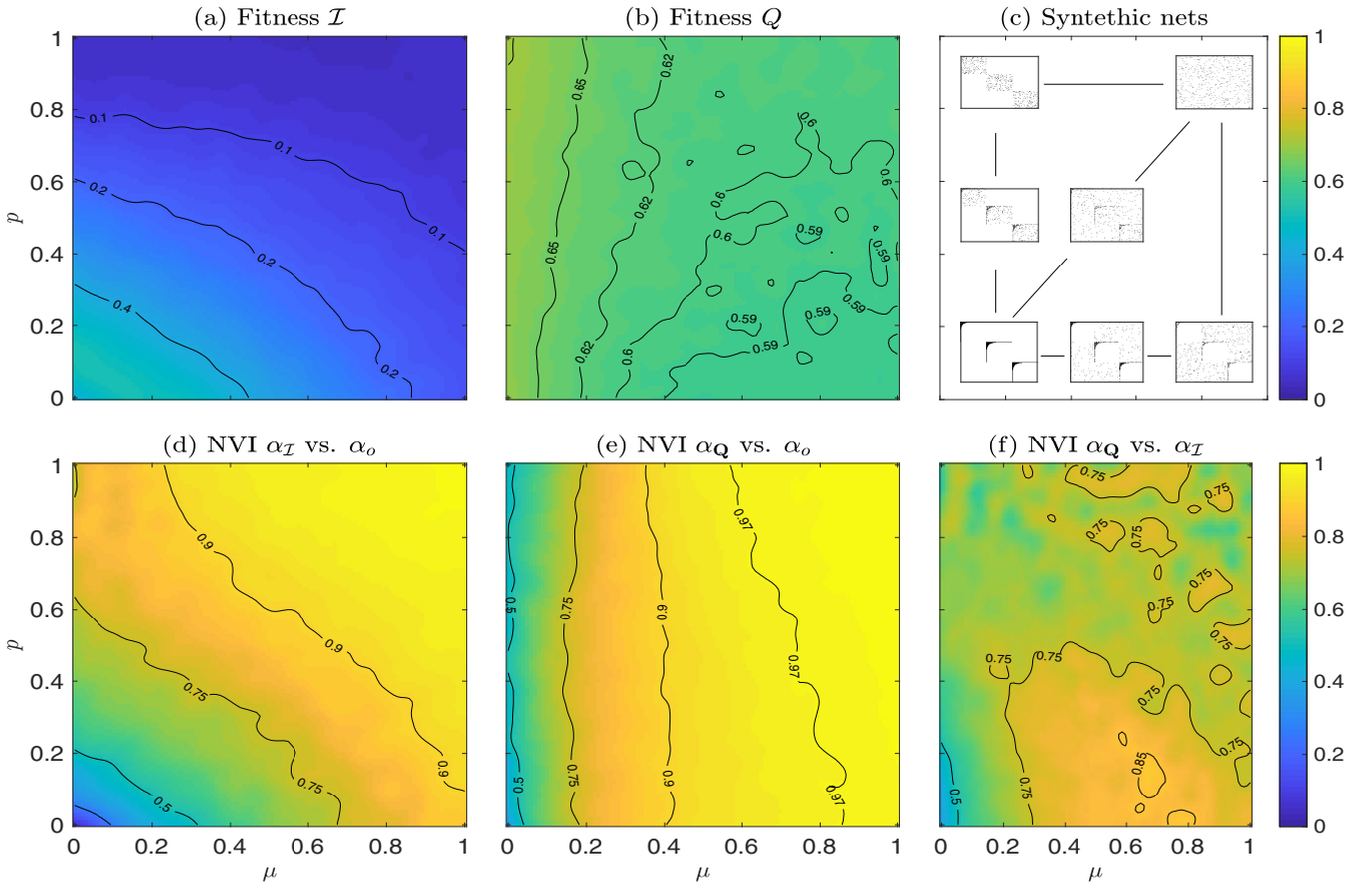


FIG. 7. Results for synthetic IBN networks with $B = 3$ and $\xi = 4$ for varying values of p and μ . Panel (a) displays the value of the fitness function \mathcal{I} as obtained with a biologically inspired optimization algorithm. Similarly, panel (b) reports the value of modularity Q as obtained with the Combo optimization algorithm [33]. Panels (d) and (e) compare the \mathcal{I} -optimized ($\alpha_{\mathcal{I}}$) and the Q -optimized (α_Q) partitions, respectively, to the planted partition (α_0) via the NVI. Panel (c) provides a visual intuition of the effect of parameters p and μ on the synthetically generated networks. Panel (f) shows the difference between the \mathcal{I} -optimized ($\alpha_{\mathcal{I}}$) and the Q -optimized (α_Q) partitions as measured by their NVI.

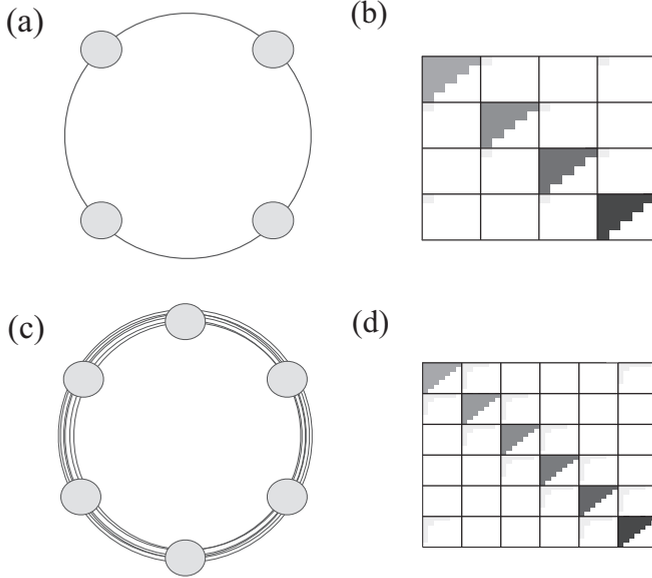


FIG. 8. Panels (a) and (b) represent a ring of in-block nested subgraphs (network and adjacency matrix representation, respectively) connected by a single link. This example, which follows closely Fig. 3 in Ref. [37], is analyzed in Sec. IV. Panels (c) and (d) represent the same ring, although each block is connected by ℓ^* links—the minimum needed to enforce $\mathcal{I}_{\text{pairs}} > \mathcal{I}_{\text{single}}$ at $B = 2$.

values are close to 0.5 instead of ~ 0 in the region $\mu \approx 0$, $p \approx 1$, i.e., the region where a purely modular, non-nested structure is more obvious. The reason for these apparently inconsistent results stems from the sparsity of the network itself, which distorts even the modular partition that seems more natural. For example, in the most extreme case ($p = 1, \mu = 0$), the modularity value that the Combo algorithm [33] delivers ($Q = 0.70$) is larger than the one imposing the planted partition ($Q = 0.66$), thus explaining such high NVI.

APPENDIX C: FURTHER INSIGHTS AROUND Q AND \mathcal{I} 'S RESOLUTION LIMIT

In Sec. IV we have provided a minimal example, along the lines of Ref. [37], which hints at the idea that \mathcal{I} may not be subjected to the inherent limitations of Q regarding the detectability of nested blocks at small scales. Despite the lack of an analytical approach, which we deem necessary in the future, we suggest that such lack of a resolution limit is due to the fact that \mathcal{I} does not depend on a global quantity (like L , the total number of links in the network) but rather is the sum over a relative quantities, i.e., the normalized overlap between pairs of nodes.

Here we want to extend this argument with a striking example that shows that not only \mathcal{I} can retain the *natural*

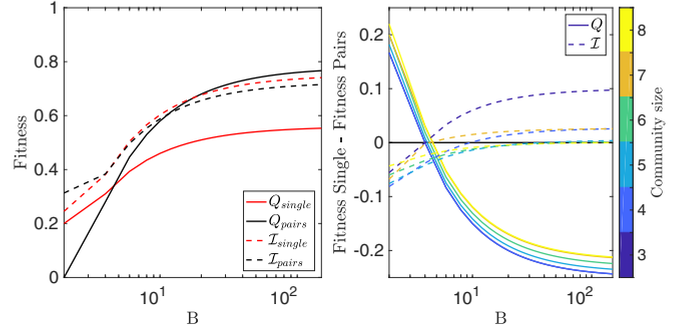


FIG. 9. Left: Q_{single} (solid-red line) and Q_{pairs} (solid-black line) as a function of the number of nested blocks, B . Q for this ring of dense subgraphs—each block connected to its neighbors by ℓ^* links—exhibits the well-known behavior, i.e., showing an earlier crossover (if compared to Fig. 4) that disregards the *natural* partition and favors a larger-scale one. $\mathcal{I}_{\text{single}}$ (dashed-red line) and $\mathcal{I}_{\text{pairs}}$ (dashed-black line) for the same configuration. As prescribed, $\mathcal{I}_{\text{pairs}} > \mathcal{I}_{\text{single}}$ initially. However, as B increases, \mathcal{I} uncovers the *natural* partition, i.e., $\mathcal{I}_{\text{single}} > \mathcal{I}_{\text{pairs}}$. Right: Difference between $\mathcal{I}_{\text{single}}$ and $\mathcal{I}_{\text{pairs}}$ (dashed), and between Q_{single} and Q_{pairs} (solid), for different block sizes. Results show that the conclusions extracted on the left panel hold as well for different settings.

partition as the number of blocks B grows but also it can actually *recover* such natural partition even from the most adverse starting point. To show this, let us consider again a network built from dense subgroups, connected as a ring. Unlike the structure from Figs. 8(a) and 8(b) (which mimics Fig. 3 in Ref. [37]), consider that the blocks in the ring are connected (to the previous and next block) by more than just one link [say, ℓ links, see Fig. 8(c) and 8(d)]. Arguably, the logic of Q 's resolution limit still holds, with the difference that the crossover at which $Q_{\text{pairs}} > Q_{\text{single}}$ will happen for a smaller B : We would observe a shift to the left in Fig. 4. If ℓ is large enough, then $Q_{\text{pairs}} > Q_{\text{single}} \forall B$.

Translating this logic to \mathcal{I} , we perform the following experiment: We start out with a configuration of two nested blocks ($B = 2$) which are connected, in ring shape, with the minimum amount of links ℓ^* , such that ensure, at this level, that $\mathcal{I}_{\text{pairs}} > \mathcal{I}_{\text{single}}$. We then proceed to extend the ring, connecting again each block to its neighbors with the same amount of ℓ^* links. Figure 9 shows that indeed, at $B = 2$ (the minimal ring network), $\mathcal{I}_{\text{pairs}} > \mathcal{I}_{\text{single}}$. However, we observe, as B grows that the difference between $\mathcal{I}_{\text{pairs}}$ and $\mathcal{I}_{\text{single}}$ reduces to eventually reach a point such that \mathcal{I} recovers the most natural partition, i.e., $\mathcal{I}_{\text{single}}$ has higher fitness than $\mathcal{I}_{\text{pairs}}$. This result evidences that, contrary to modularity, the detectability of small blocks increases as we increase the size of the network and allows us to hypothesize that \mathcal{I} may not be susceptible to problems similar to the resolution limit.

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