

PREDICTING TRIADIC CLOSURE IN NETWORKS USING COMMUNICABILITY DISTANCE FUNCTIONS*

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Abstract. We propose a communication-driven mechanism for predicting triadic closure in complex networks. It is mathematically formulated on the basis of communicability distance functions that account for the quality of communication between nodes in the network. We study 25 real-world networks and show that the proposed method correctly predicts 20% of triadic closures in these networks, in contrast to the 7.6% predicted by a random mechanism. We also show that the communication-driven method outperforms the random mechanism in explaining the clustering coefficient, average path length, and average communicability. The new method also displays some interesting features with regards to optimizing communication in networks.

Key words. network analysis, triangles, triadic closure, communicability distance, adjacency matrix, matrix functions

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1. Introduction. Complex networks are ubiquitous in many real-world scenarios, ranging from the biomolecular—those representing gene transcription, protein interactions, and metabolic reactions—to the social and infrastructural organization of modern society [12, 16, 44]. Mathematically, these networks are represented by graphs, where the nodes represent the entities of the system and the edges represent the “relations” among those entities. The accumulation of a mountain of empirical evidence has left little doubt that, in general, real-world networks are very different from those based on the uniform model $G(n, p)$ in many structural and functional aspects [16]. In particular, it is well documented that real-world networks are significantly more “clustered” than one would expect from the previously mentioned uniform model [44]. The degree of “clustering” is usually quantified in network theory through the use of the *clustering coefficient* (see [53]). This accounts for the ratio of the number of triangles to the number of open triads, i.e., subgraphs of the type $i - j - k$. The fact that triangles are abundant in real-world networks has long been appreciated—for example, in 1922, when Simmel [52] theorized that people with common friends are more likely to create friendships. This *friendship transitivity* definitively implies a social mechanism for triadic closure in social networks which may then be applied to explain the evolution of triangle closures [30]. This Simmelian principle of triadic closure due to friendship transitivity assumes that individuals can benefit from cooperative relations, and this may induce individuals to choose new acquaintances from among their friends’ friends.

The high degree of transitivity is not a unique feature of social networks; indeed, it is a common characteristic of many other types of networks such as biomolecular,

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cellular, ecological, infrastructural, and technological (see [16] and references therein). It is natural to assume that analogous cooperative principles to the one proposed by Simmel for social networks could be applied to find mechanisms that explain triadic closure in these other types of networks. Although intuitive, this simple idea has some fundamental drawbacks. First, it is not always true that pairs of nodes benefit from cooperative relations, and therefore the Simmelian principle is useless in such situations. Second, it is evident that not every pair of nodes separated by two edges participates in a triangle in a real-world network. Thus, some kind of selective process has been taking place, closing some of the triads in a network and leaving many others open.

The goal of this paper is to propose a general mechanism to account for such a selective process of triadic closure in networks. We propose a strategy for predicting triadic closure based on the idea that triadic closure is a communication-driven process. This paradigm is formulated on the basis of communicability distance functions that account for the quality of communication between pairs of nodes using a mechanism accounting for both local and long-range interactions. We start with an overview of related work. All the mathematical concepts we use are introduced in section 3 in order to make the paper self-contained. Sections 4 and 5 are devoted to the introduction of the new method for predicting triadic closure. We finish with a presentation and discussion of the results.

2. Related work. Triadic closure, loosely defined as the process in which an edge is added to a triad to form a triangle, has long been considered a fundamental mechanism of social networks' evolution. The theoretical basis of this mechanism is due to Simmel [52], and one of the pioneering studies to use this principle to predict triadic closure in social networks was published by Krackhardt and Handcock [30].

When considering undirected networks, the main focus of triadic closure models has been to create simple mechanisms that provide insight into how (social) networks grow and generate their main topological characteristics. A simple model of network growth based on triadic closure has been proposed by Bianconi et al. [5]. They show that the evolution of networks based on such simple mechanisms *naturally leads to the emergence of community structure, together with fat-tailed distributions of node degree and high clustering coefficients*. Similar results by Klimek and Thurner [27] suggest that triadic closure can be identified as one of the fundamental dynamical principles in social multiplex network formation. These two works use triadic closure mechanisms based on the random selection of the nodes which will be involved in the triangles.

In the case of directed graphs an exhaustive computational analysis was performed by Leskovec et al. [31]. They consider several strategies to model how a node u selects a node v , two steps from it, to form a triangle. The basic strategy is for u to randomly select a node v from all the nodes at distance two. An alternative strategy is to assume that u first selects a neighbor node w according to some mechanism, and then w selects a neighbor v according to some (possibly different) mechanism. The edge (u, v) is then formed, and the triangle Δ_{uvw} is closed. The selection of a neighbor w for u (or v for w) has been carried out using the following techniques: (i) uniformly at random; (ii) proportional to degree of w raised to a power; (iii) proportional to the number of friends that u and w have in common; (iv) proportional to the time passed since w last created an edge raised to a power; (v) proportional to the product of the number of common friends of u and w multiplied by the last activity time, all raised to a power.

The quantitative predictions made by Leskovec et al. are summarized in Table 1, where we report the percentage of correct prediction of triadic closure in four

TABLE 1

Illustration of the percentage of correct prediction of triadic closure in online social networks by the random-random selection of nodes and the best of all predictions made by Leskovec et al. [31].

Network	% correct triadic closure	
	random-random	best
Flickr	13.6	16.9
Delicious	11.7	18.2
Answers	6.8	16.4
Linkedin	16.0	21.4

networks, using a random-random selection of the nodes, and the best percentage of improvement over the log-likelihood of picking a random node two hops away (baseline) reported by the authors. The network LinkedIn is the only one of the four which is undirected.

In a more recent paper, Lou et al. [32] have developed a method that adds sociological information to the network structure in order to predict triadic closure in a Twitter network. Their approach uses information about (i) geographic distance, i.e., whether users have a higher probability of following each other when they are located in the same region; (ii) homophily, i.e., whether similar users tend to follow each other; (iii) implicit network, i.e., how the following network on Twitter correlates with other implicit networks, such as the retweet and reply network; and (iv) social balance, i.e., whether the reciprocal relationship network on Twitter satisfies social balance theory and to what extent. When this nontopological information is added, the developed method outperforms other structure-only approaches in the prediction of triadic closures. A similar approach, which uses demographic information instead, has been developed by Huang et al. [24]. They have used a large microblogging network as the source of their study, which reveals how user demographics and network topology influence the process of triadic closure. Their experimental results on the microblogging data show the efficiency of the proposed model for the prediction of triadic closure formation.

Here we will not account for extratopological information, i.e., information apart from that provided by the topological structure of the network. Thus, our current work is more in the spirit of that of Leskovec et al. [31] with the difference that the networks we study are undirected.

3. Mathematical preliminaries. A graph $\Gamma = (V, E)$ is defined by a set of n nodes (vertices) V and a set of m edges $E = \{(u, v) | u, v \in V\}$ between the nodes. An edge is said to be *incident* to a vertex u if there exists a node $v \neq u$ such that either $(u, v) \in E$ or $(v, u) \in E$. The *degree* of a vertex u , denoted by d_u , is the number of edges incident to u in Γ . The graph is said to be *undirected* if the edges are formed by unordered pairs of vertices. A *walk* of length k in Γ is a set of nodes $u_1, u_2, \dots, u_k, u_{k+1}$ such that for all $1 \leq l \leq k$, $(u_l, u_{l+1}) \in E$. A *closed walk* is a walk for which $u_1 = u_{k+1}$. A *path* is a walk with no repeated nodes. A closed walk of length 3 is called a *triangle*. We will call *triad* every triplet of nodes u, v , and w such that $(u, v), (v, w) \in E$ but $(u, w) \notin E$. Hence a triad is a triangle missing one edge. We shall call this missing edge a *potential edge*. A graph is *connected* if there is a path joining u and v for every $u, v \in V$. A graph with unweighted edges, no edges from a node to itself, and no multiple edges is said to be *simple*.

Let $A = (a_{uv}) \in \mathbb{R}^{n \times n}$ be the *adjacency matrix* of the graph. It is worth noting that for undirected, simple, and connected networks the associated adjacency matrix

is symmetric, binary, hollow (i.e., has zeros on the main diagonal), and irreducible (see [23]), and its entries are

$$a_{uv} = \begin{cases} 1 & \text{if } (u, v) \in E, \\ 0 & \text{otherwise} \end{cases} \quad \forall u, v \in V.$$

It is possible to define several distance measures on networks. The most common is the *shortest-path* (or *geodesic*) *distance* between two nodes $u, v \in V$, which is defined as the length of the shortest path connecting these nodes. We will write $d(u, v)$ to denote the geodesic distance between u and v , and hence the *average path length* [16, 44, 45], the average of the shortest-path distances in the graph, is given by

$$\bar{\ell} = \frac{1}{2m} \sum_{u, v \in V} d(u, v).$$

Another useful measure for characterizing the structure of networks is the so-called *local clustering coefficient* of a node u [53], which quantifies the degree of transitivity of local relations in a network and is defined as

$$C_u = \frac{2t_u}{d_u(d_u - 1)},$$

where t_u is the the number of triangles in which node u participates. Taking the mean of these values as u varies among all the nodes in Γ gives the *clustering coefficient* of the network,

$$\bar{C} = \frac{1}{n} \sum_{u=1}^n C_u.$$

An important quantity to be considered when studying communication processes in networks is the *communicability function* [14, 17, 15], which is defined as

$$G_{uv} = (e^A)_{uv} = \sum_{k=0}^{\infty} \frac{(A^k)_{uv}}{k!} = \sum_{k=1}^n e^{\lambda_k} \mathbf{q}_k(u) \mathbf{q}_k(v) \quad \forall u, v \in V,$$

where $A = Q\Lambda Q^T$ is the spectral decomposition of the adjacency matrix (see [23]), with Λ a diagonal matrix containing the eigenvalues of A and $Q = [\mathbf{q}_1, \dots, \mathbf{q}_n]$ an orthogonal matrix containing the associated eigenvectors.

Communicability counts the total number of walks starting at node u and ending at node v , weighting their length by a factor $\frac{1}{k!}$, thus considering shorter walks more influential than longer ones. The G_{uu} terms of the communicability function, which are usually called *subgraph centralities* of the nodes, characterize the degree of participation of a node in all subgraphs of the network, giving more weight to the smallest ones. Here we will use the *average communicability* as a way to characterize the quality of the communication taking place in the network as a whole:

$$\bar{G} = \frac{1}{n(n-1)} \sum_{u \neq v} G_{uv}.$$

The communicability function can be used to quantify the quality of communication between nodes in a network. When two nodes u and v are exchanging information,

the quality of their communication depends on two factors: how much information departing from a source node reaches its target (G_{uv}), and how much information departing from the node returns to it without ending at its destination (G_{uu}). That is, the quality of communication increases with the amount of information that departs from the originator and arrives at its destination, and decreases with the amount of information which is wasted due to the fact that the information returns to its source without being delivered to the target. In [18] the *communicability distance* is defined as

$$(3.1) \quad \xi_{uv} = \sqrt{G_{uu} + G_{vv} - 2G_{uv}}.$$

It is a Euclidean distance between the nodes u and v in Γ (see [18, 19]). From its definition, it is clear that ξ_{uv} characterizes the quality of the communication taking place between nodes u and v .

4. Communicability distances and triad closure. We start by considering the square of the communicability distance defined in (3.1) for a pair of nodes uv in a connected graph. This distance characterizes communication quality between nodes u and v by assuming that the information departing from node u travels to node v (and viceversa) by taking a series of one-hop steps between the nodes in any of the walks that connect them. From (3.1), it is clear that the smaller the value of ξ_{uv}^2 , the better nodes uv are at exchanging information. The communicability distance is dependent on e^A , where A is the adjacency matrix of a simple graph. If we consider u and v such that $a_{uv} = 1$, then we are assuming that these two nodes are attracted to each other. If instead we were to consider that these nodes repel each other, we would use e^{-A} .

If the (squared) communicability distance between two pairs of nodes uv and pq satisfies $\xi_{uv}^2 < \xi_{pq}^2$, then we say that the attraction between the pair uv is stronger than that between the pair pq in the corresponding network.

Now, consider a triad u, w, v , where $(u, w) \in E$, $(w, v) \in E$, but $(u, v) \notin E$. Because $a_{uw} = 1$ and $a_{wv} = 1$ we can infer that there are attractive *forces* between u and w and between w and v . A simple metaphoric way to represent such attractive forces between pairs of nodes is to suppose that they have opposite charges which attract to each other. For instance, we can consider either of the following schemes for the previous example: $u^+ - w^- - v^+$, $u^- - w^+ - v^-$. Notice that considering a particle spin, as is usually done in sociophysical models of opinion dynamics, also works here as an appropriate metaphor (see, for instance, [51]). Observe that there are two types of interactions between the nodes u and v . First, due to the attractions between u and w and w and v , the node v “feels” an attractive force from u , which is transmitted through the edges of the network. On the other hand, due to the fact that both u and v have the same charge, they experience some repulsion from each other, which takes place in a “through-space” fashion (which we will clarify later). We can expect the link (u, v) to be created if the through-edge attractive force between the nodes u and v is larger than the through-space repulsive force between them.

In order to understand the nature of the interactions described in the previous paragraph, we consider a molecular system as a model example. In this case there is a communication between pairs of atoms which occurs through the covalent bonds of the molecule. This kind of interaction takes place through the edges (covalent bonds) and is analogous to the attractive forces we have previously described. Hereafter we will refer to this interaction as the *through-edges communicability* (TEC). If two non-covalently bonded atoms are close in space, they can interact with each other through

noncovalent interactions, for example, by hydrophobic, polarity, or electrostatic forces. These interactions are analogous to our through-space repulsion, and we will refer to them as *long-range communicability* (LRC). In a social network, TEC is present when information is transmitted from one individual to another in the network by using the social ties that define the edges of the graph. On the other hand, LRC is realized by the direct influence of an individual on another through any source of social signaling.

Note that although the shortest-path distance between every pair of nodes in a triangle equals one, every pair of nodes in it is connected by a pair of adjacent edges through the third vertex. A natural way to account for all the pairs of nodes connected by pairs of adjacent edges is to consider the number of walks of length two between the pairs of nodes. We can then transform a graph, accordingly. Let $\Gamma = (V, E)$ be a simple and connected graph, and let $W_2(\Gamma) = (V, E')$ be the graph with the same set of nodes as Γ but whose edges are weighted by the number of walks of length two between every pair of (not necessarily distinct) nodes in Γ . More precisely, if $\mu_2(u, v)$ is the number of walks of length two between nodes i and j , then the adjacency matrix \tilde{A} of $W_2(\Gamma)$ is

$$(\tilde{a})_{uv} = \begin{cases} \mu_2(u, v), & u \neq v, \\ \mu_2(u, u) = d_u, & u = v. \end{cases}$$

Remark 1. Clearly $\tilde{A} = A^2$, and so we do not need to explicitly construct the graph $W_2(\Gamma)$, since we can simply work with the square of the adjacency matrix of the graph Γ .

Note that two nodes are connected in $W_2(\Gamma)$ if they have the same charge, and so connected nodes in $W_2(\Gamma)$ repel each other. Consequently the repulsive communicability between a given pair of nodes in Γ is given by $\tilde{G}_{uv} = (e^{-\tilde{A}})_{uv} = (e^{-A^2})_{uv}$.

A communicability distance based on \tilde{G}_{uv} accounts for the quality of LRC between pairs of nodes separated by two adjacent edges, i.e., pairs of nodes feeling mutual repulsion in Γ . We can define a communicability distance function by

$$(4.1) \quad \eta_{uv} = \sqrt{\tilde{G}_{uu} + \tilde{G}_{vv} - 2\tilde{G}_{uv}}.$$

A large value of η_{uv} indicates that there is a weak repulsion between nodes u and v . The proof that η_{uv} is a Euclidean distance between the nodes u and v follows the same lines as in [18, 19] and is omitted.

Remark 2. The graph $W_2(\Gamma)$ is not always connected, and so the function η_{uv} is defined only for pairs of nodes which are in the same connected component of the graph. Elsewhere η_{uv} is set to infinity.

Before continuing, consider the following example. The tree illustrated on the left in Figure 1 can be transformed by adding an edge which closes any of the three nonequivalent existing triads of the graph, i.e., by adding the edge $(2, 3)$, $(2, 4)$, or $(1, 5)$. The resulting unicyclic graphs are illustrated on the right of Figure 1. In Table 2 we report the values of ξ_{uv}^2 and η_{uv}^2 for each of the three triads. Now assume that we have information indicating that the process giving rise to the closure of the 1, 2, 3-triad is favored over the other two. We cannot know a priori for any particular system how the attractive and repulsive forces scale. In real physical systems such terms are scaled by minimizing the global energy of the system. Here we simply consider the weighted difference between the two terms, $\alpha\xi_{uv}^2 - \beta\eta_{uv}^2$. We will propose a method to determine the values of the empirical parameter α and β in a given network a little later. For this example it can be verified that, for instance,

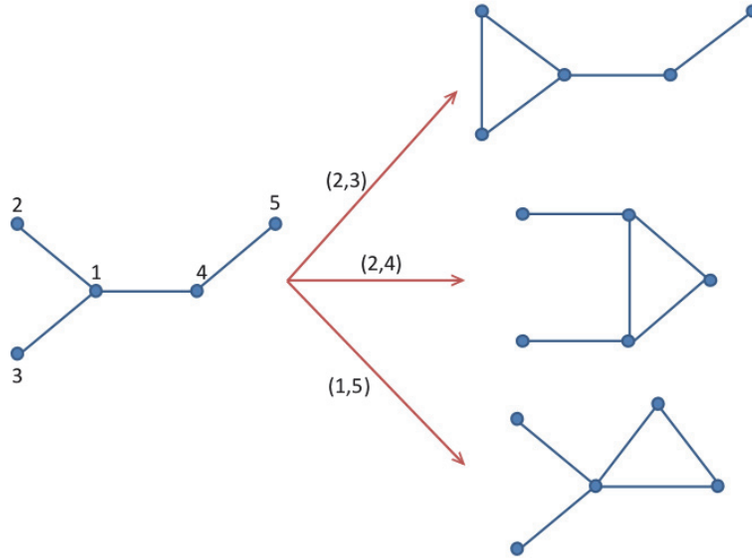


FIG. 1. Example of evolution of a tree after one edge is added to close a triangle.

$\xi_{uv}^2 - 1.5\eta_{uv}^2$ produces a negative value only for the pair (2,3) (see Table 2). This weighted difference between ξ_{uv}^2 and η_{uv}^2 corresponds to the case in which the attractive forces between the corresponding nodes outweigh the magnitude of the repulsive ones. As noted previously, a large value of η_{uv}^2 indicates a small repulsion between the corresponding nodes, and here we have multiplied η_{uv}^2 by a coefficient $\beta > 1$, which further reduces the repulsive forces.

Now suppose instead that we have information indicating that the process giving rise to the closure of the 1, 4, 5-triad is favored over the other two. In this case it can be verified (see Table 2) that the weighted difference $-0.5\xi_{uv}^2 + 1.5\eta_{uv}^2$ is negative only for the pair (1,5). Here, we have considered that the attractive forces between the nodes make a negative contribution to the creation of an edge closing the triad. This may correspond to the situation in which the links (u, w) and (w, v) are both very weak, i.e., friendship ties between the corresponding individuals are not too strong. We further weaken those relations by multiplying ξ_{uv}^2 by a coefficient $\alpha < 0$. At the same time, by multiplying η_{uv}^2 by a coefficient $\beta < 0$ we have assumed that the repulsive factor does not play a major role in determining whether the new edge is created or not. Indeed, in this way η_{uv}^2 is transformed into an attraction term. In the charged-particles analogy this corresponds to a situation in which the charges between the corresponding nodes are very weak and there is no repulsion between those nodes separated by two adjacent edges.

Finally, suppose that we have information indicating that the process giving rise to the closure of the 1, 2, 4-triad is favored over the other two. In this case it can be verified (see Table 2) that the weighted difference $\xi_{uv}^2 + \eta_{uv}^2$ reaches the smallest value for (2,4). The values of the weighted differences for the three triad closure processes are positive, but that corresponding to the closure of the 1, 2, 4-triad is the lowest among the three. In this case, triadic closure is dominated by attractive forces only. The term $\alpha\xi_{uv}^2$ with $\alpha > 0$ indicates the normal attractive forces between the corresponding pair of nodes, while $\beta\eta_{uv}^2$ with $\beta < 0$ is transformed into an attractive term.

TABLE 2
 Values of weighted sum of ξ^2 and η^2 for the potential edges considered in Figure 1.

Pair	ξ_{uv}^2	η_{uv}^2	$\xi_{uv}^2 - 1.5\eta_{uv}^2$	$-0.5\xi_{uv}^2 + 1.5\eta_{uv}^2$	$\xi_{uv}^2 + \eta_{uv}^2$
2,3	2.000	2.000	-1.000	2.000	4.000
1,5	3.184	0.960	1.744	-0.152	4.144
2,4	2.545	1.312	0.577	0.696	3.857

A case we have not considered here is if $\alpha < 0$ and $\beta > 0$ when both terms represent repulsive forces between nodes. In this case $\alpha\xi_{uv}^2 - \beta\eta_{uv}^2 < 0$ for all u, v , and the order in which the triads will be closed is determined by the magnitudes of α and β . In such a repulsive system there are no attractive forces to fuel the creation of new edges. Consequently, the creation of the new edges to close triads is controlled by factors such as their similarities or complementarity in their functions which do not depend particularly on the communicability between nodes. In this case, predictions of triad closure made on the basis of communicability distances are not expected to differ significantly from those made by a random closure of the triads.

In summary, we can use the function

$$(4.2) \quad \Delta_{uv}(\alpha, \beta) := \alpha\xi_{uv}^2 - \beta\eta_{uv}^2 \quad \forall u, v \in V$$

to determine which triad is closed in the network.

In order to predict which triads will close in a given network it is necessary to know the values of α and β . We now propose a method that allows us to estimate these empirical parameters and consequently to determine which triads will close in a given network.

5. Proposed method. In order to predict the triadic closure in a network based on $\Delta_{uv}(\alpha, \beta)$ we develop a procedure to find the values of the empirical parameters α and β which best predict the triadic closure in a network from which we have a priori removed all the triangles. That is, if we take a network Γ , we first detect all its existing triangles. We then transform Γ into a triangle-free graph Γ' by removing one and only one of the edges forming each triangle. The deleted edges are selected uniformly at random from the three edges forming each triangle. As this procedure is likely to be repeated a large number of times (see below for details), the chance that each of the three edges is selected at least once is very high. We keep a list of all these removed edges which we call R . It may happen that two triangles T_1 and T_2 share an edge e . If we select e when considering T_1 , then, when it is time to select an edge in T_2 , we pick an edge which may or may not coincide with e . If it does, we do not add it to the list. It may also happen that T_2 consists of e and two other edges, one of which has also already been removed because it was in common with a third triangle. In such cases, we do not remove the last connection remaining in T_2 , since it could disconnect the network.

We can also create a list, P , of all the pairs of nodes which form triads in Γ but were not part of any triangle. Finally, we create the list $L = R \cup P$. Because we have removed one edge from each triangle, the nodes in R are now separated by two adjacent edges in Γ' , similarly to the pairs of nodes in P . Our task is to select appropriate values of the empirical parameters α and β that differentiate as much as possible the pairs of nodes in R from those in P . We do this by using a nonincreasing ranking of all the pairs of nodes in L according to $\Delta_{uv}(\alpha, \beta)$. We have previously predicted that the triadic closure process should be controlled by the smallest values of $\Delta_{uv}(\alpha, \beta)$ (see the example in Figure 1). Thus, we expect that a nonincreasing

ranking of the values of $\Delta_{uv}(\alpha, \beta)$ contains most of the elements of R at the top of the ranking and those of P at the bottom.

In order to quantify the percentage of triangles that were correctly predicted, we proceed as follows. We first rank the entries of L in nonincreasing order. We select the top r entries of $L = R \cup P$, where r is the cardinality of R . Then, we count the number r_p of entries in this top r which are elements of R . These entries correspond to those pairs of nodes which were originally part of the triangles of Γ . That is, r_p represents the number of correct predictions made by the current method. We call the (percentage) ratio of r_p to r the percentage of **detected**.

5.1. Datasets and computational methods. We now give some computational details on how we implemented these calculations to find the optimal values of α and β for a selection of networks.

We study 25 networks representing complex systems from a wide variety of environments, such as social, ecological, biomolecular, technological, infrastructural, and informational. A brief description of all these networks is given in the appendix.

In order to find the optimal values of the empirical parameters α and β for these networks, we proceed as follows. We calculate all the values of α and β in the interval $I = [-2.1, 2.1]$ with a step length of 0.1. This interval I has been determined empirically as smaller intervals led to worse results and larger ones did not improve the results. Then, for each combination of α and β in $\Delta_{uv}(\alpha, \beta)$, we rank all the elements of L in nonincreasing order and find the percentage of **detected**. The optimal values of α and β for this particular network are those that produce the highest percentage of **detected**. These computations were repeated 100 times.

The effectiveness of the proposed method is tested by considering a simple null model constructed as follows. We randomly order the edges in L , select the top r pairs of nodes, and count how many of them were in the set R . With this information we compute the percentage of correct predictions made by a random ordering of the pairs of nodes (**rand**). Similar values of the percentages of **detected** and **rand** indicate that the ranking produced by the function $\Delta_{uv}(\alpha, \beta)$ does not differ significantly from a random ordering of the pairs of nodes and consequently is not a good one, while larger differences between the percentages of **detected** and **rand** indicate good performance of the proposed method.

Before starting with the detailed analysis of these 25 datasets we consider the possibility of fixing one of the parameters (α or β) and letting the other vary in the bounded interval $[-2.1, 2.1]$. To do this we set $\alpha = 1$ and let β vary. This seems reasonable, since this choice allows us to tune the disturbance caused by the repulsion in the values of Δ_{uv} . However, the results obtained for 10 of the studied networks discouraged us from proceeding with this approach. On average the use of the two parameters α and β makes predictions of triadic closure which are 7% higher than those using only one parameter, with maximum differences of up to 20% for one network (results not shown here). Thus, we will use the more general approach of calibrating both empirical parameters.

5.2. Bounds for communicability distance functions. Although in our experiments we use the exact values of the communicability distance functions in order to obtain the values of Δ_{uv} , we now give some bounds for ξ_{uv} and η_{uv} , which can be used in the computations when working on extremely large networks. It is clear from the definitions given in (3.1), (4.1), and (4.2) that for large matrices these values may be too costly to compute. To avoid the computation of the matrix exponential, we derive bounds for ξ_{uv}^2 and η_{uv}^2 (and therefore for $\Delta_{uv}(\alpha, \beta)$) by means of a Gauss–Radau

quadrature rule. In order to make the present paper self-contained, we summarize the approach used as described in [2, 1, 21] before giving these bounds.

It is well known that the problem of computing bilinear expressions of the form $\mathbf{u}^T f(A) \mathbf{v}$ can be reduced to the approximation of a Riemann–Stieltjes integral with respect to a certain measure using quadrature rules. Indeed, in a series of papers, Golub and collaborators use one step of the symmetric Lanczos iteration to give bounds on the entries of $f(A)$ based on Gauss-type quadrature rules when f is a *strictly completely monotonic* (s.c.m.) function on an interval \mathcal{I} containing the spectrum of A . Recall that a function is s.c.m. on \mathcal{I} if $f^{(2k)}(x) > 0$ and $f^{(2k+1)}(x) < 0$ for all $x \in \mathcal{I}$ and for all $k \geq 0$. Since $g(x) = e^x$ is not s.c.m., we need to work on $f(x) = e^{-x}$ to derive bounds on the quantities of interest here.

The key result that allows us to easily compute a priori bounds using Gauss-type quadrature rules is that we can use the element in position (1, 1) of the matrix $f(J_{p+1})$ (see Theorem 6.6 in [21]), where

$$J_{p+1} = \begin{pmatrix} \omega_1 & \gamma_1 & & & & \\ \gamma_1 & \omega_2 & \gamma_2 & & & \\ & & \ddots & \ddots & \ddots & \\ & & & \gamma_{p-1} & \omega_p & \gamma_p \\ & & & & \gamma_p & \omega_{p+1} \end{pmatrix}$$

is a tridiagonal matrix whose eigenvalues are the nodes of the quadrature rule, and the rule's weights are given by the squares of the first entries of J_{p+1} 's normalized eigenvectors.

Our results are summarized in the following theorems.

THEOREM 5.1. *Let A be the adjacency matrix of an unweighted and undirected network. Then*

$$(5.1) \quad \Phi\left(b, \omega_1 + \frac{\gamma_1^2}{\omega_1 - b}\right) \leq \frac{(\xi_{uv})^2}{2} \leq \Phi\left(a, \omega_1 + \frac{\gamma_1^2}{\omega_1 - a}\right),$$

where

$$(5.2) \quad \Phi(x, y) = \frac{c(e^{-x} - e^{-y}) + xe^{-y} - ye^{-x}}{x - y}, \quad c = \omega_1,$$

$$\begin{cases} \omega_1 = a_{uv}, \\ \gamma_1 = \left[\frac{d_u + d_v}{2} - \omega_1 - A_{uv}^2\right]^{\frac{1}{2}}, \end{cases}$$

and $[a, b]$ is an interval containing the spectrum of $-A$.

Remark 3. If $(u, v) \notin E$, the bounds simplify considerably. Indeed, in this case $\omega_1 = 0$, and hence

$$\frac{b^2 e^{\frac{\gamma_1^2}{b}} + \gamma_1^2 e^{-b}}{b^2 + \gamma_1^2} \leq \frac{(\xi_{uv})^2}{2} \leq \frac{a^2 e^{\frac{\gamma_1^2}{a}} + \gamma_1^2 e^{-a}}{a^2 + \gamma_1^2}.$$

Before proceeding with the proof of the result, note that $(\xi_{uv})^2$ can be written as

$$(\xi_{uv})^2 = (\mathbf{e}_u - \mathbf{e}_v)^T (e^A) (\mathbf{e}_u - \mathbf{e}_v),$$

where \mathbf{e}_u and \mathbf{e}_v are the u th and v th vectors of the canonical basis, respectively.

Proof. Using the Lagrange interpolation formula for the evaluation of matrix functions [22], one can easily show [1] that

$$\mathbf{e}_1^T(e^{-C})\mathbf{e}_1 = \frac{c_{11}(e^{-\mu_1} - e^{-\mu_2}) + \mu_1 e^{-\mu_2} - \mu_2 e^{\mu_1}}{\mu_1 - \mu_2},$$

where μ_1, μ_2 are the distinct eigenvalues of the matrix C .

We now want to build explicitly the matrix $J_2 = \begin{pmatrix} \omega_1 & \gamma_1 \\ \gamma_1 & \omega_2 \end{pmatrix}$ in such a way that $\tau_1 = a$ or $\tau_1 = b$ is a prescribed eigenvalue. The values of ω_1 and γ_1 are derived explicitly by applying one step of the Lanczos iteration to the matrix $-A$ with starting vectors $\mathbf{x}_{-1} = \mathbf{0}$ and $\mathbf{x}_0 = (\mathbf{e}_u - \mathbf{e}_v)/\sqrt{2}$.

Note that if $\gamma_1 = 0$, we simply take $\omega_2 = \tau_1$, and the matrix J_2 is diagonal with eigenvalues $\mu_1 = \omega_1$ and $\mu_2 = \tau_1$. Thus, let us assume $\gamma_1 \neq 0$. Using the three-term recurrence for orthogonal polynomials

$$\gamma_j p_j(\lambda) = (\lambda - \omega_j) p_{j-1}(\lambda) - \gamma_{j-1} p_{j-2}(\lambda), \quad j = 1, 2, \dots, p,$$

with $p_{-1}(\lambda) \equiv 0, p_0(\lambda) \equiv 1$, we find that $\omega_2 = \tau_1 - \frac{\gamma_1}{p_1(\tau_1)}$. Using the same recurrence, we also find that $p_1(\tau_1) = (\tau_1 - \omega_1)/\gamma_1 \neq 0$, since the zeros of orthogonal polynomials satisfying the three-term recurrence are distinct and lie in the interior of \mathcal{I} (see [21, Theorem 2.14]).

The matrix

$$J_2 = \begin{pmatrix} \omega_1 & \gamma_1 \\ \gamma_1 & \tau_1 - \frac{\gamma_1^2}{\tau_1 - \omega_1} \end{pmatrix}$$

has (distinct) eigenvalues $\mu_1 = \tau_1$ and $\mu_2 = \omega_1 + \frac{\gamma_1^2}{\omega_1 - \tau_1}$. The result then follows by applying Theorems 6.4 and 6.6 from [21]. \square

Similar bounds can be computed for η_{uv}^2 , and are described in the following theorem, whose proof matches that of Theorem 5.1.

THEOREM 5.2. *Let A be the adjacency matrix of an unweighted and undirected network. Then*

$$\Phi\left(\tilde{b}, \tilde{\omega}_1 + \frac{\tilde{\gamma}_1^2}{\tilde{\omega}_1 - \tilde{b}}\right) \leq \frac{(\eta_{uv})^2}{2} \leq \Phi\left(\tilde{a}, \tilde{\omega}_1 + \frac{\tilde{\gamma}_1^2}{\tilde{\omega}_1 - \tilde{a}}\right),$$

where Φ is defined as in (5.2) with $c = \tilde{\omega}_1, \tilde{\mathcal{I}} = [\tilde{a}, \tilde{b}]$ is an interval containing the spectrum of A^2 , and

$$\begin{cases} \tilde{\omega}_1 = \gamma_1^2 + \omega_1, \\ \tilde{\gamma}_1 = \left[\frac{1}{2} \sum_{w=1}^n (A_{uw}^2 - A_{wv}^2) - \tilde{\omega}_1^2\right]^{\frac{1}{2}} \end{cases}$$

with ω_1 and γ_1 as in Theorem 5.1.

Remark 4. Since the behavior of the eigenvalues of A is known (see [40]), we may take $\tilde{a} = 0$ and $\tilde{b} = a^2$ as the square of the approximation to the largest eigenvalue of A . For these values, the bounds simplify to

$$\Phi\left(a^2, \tilde{\omega}_1 + \frac{\tilde{\gamma}_1^2}{\tilde{\omega}_1 - a^2}\right) \leq \frac{(\eta_{uv})^2}{2} \leq \Phi\left(0, \tilde{\omega}_1 + \frac{\tilde{\gamma}_1^2}{\tilde{\omega}_1}\right) = \frac{\tilde{\omega}_1^2 e^{-\frac{\tilde{\omega}_1^2 + \tilde{\gamma}_1^2}{\tilde{\omega}_1}} + \tilde{\gamma}_1^2}{\tilde{\omega}_1^2 + \tilde{\gamma}_1^2}.$$

Combining the results described in the previous theorems, one easily gets bounds for the values of $\frac{\Delta_{uv}(\gamma, \beta)}{2}$. Indeed, the computation is straightforward, since the new bounds are linear combinations of the previous ones. For example, if we have $\gamma \geq 0$ and $\beta \leq 0$, we get as lower bound for $\frac{\Delta_{uv}(\gamma, \beta)}{2}$

$$\gamma\Phi\left(b, \omega_1 + \frac{\gamma_1^2}{\omega_1 - b}\right) + \beta\Phi\left(\tilde{a}, \tilde{\omega}_1 + \frac{\tilde{\gamma}_1^2}{\tilde{\omega}_1 - \tilde{a}}\right),$$

and as upper bound

$$\gamma\Phi\left(a, \omega_1 + \frac{\gamma_1^2}{\omega_1 - a}\right) + \beta\Phi\left(\tilde{b}, \tilde{\omega}_1 + \frac{\tilde{\gamma}_1^2}{\tilde{\omega}_1 - \tilde{b}}\right),$$

where ω_1 , γ_1 , $\tilde{\omega}_1$, and $\tilde{\gamma}_1$ depend on the choice of u and v .

6. Modeling results and discussion.

6.1. Predicting and interpreting triadic closure. The first series of results refers to the finding of the optimal values of α and β for the studied networks and the comparison of the percentage of triadic closures correctly predicted by the proposed method in comparison with the random process. The results of the tests are listed in Table 3. The columns $\langle\alpha^*\rangle$ and $\langle\beta^*\rangle$ contain the average best values for the parameters, where the average is taken over the 100 iterations we run. The results show that on average the method based on the communicability distance functions (`detect`) correctly predicts 20% of the triad closures in the real-world networks studied. In seven cases this percentage of correct prediction reaches values larger than 25%. In contrast, the random closure of triads identifies 7.6% of the real triangles existing in those networks.

We can now gain some insights about the processes that have governed the triad closure in the studied networks. Recall that, in general, the triadic closure process consists of two different means of transmission of information, namely, the TEC and the LRC. If we refer to the nature of the two kinds of transmission in the order TEC-LRC we can have any of the following four scenarios:

- $\alpha > 0, \beta < 0$: the triads close by means of attractive-attractive interactions.
- $\alpha > 0, \beta > 0$: the triads close by means of attractive-repulsive interactions.
- $\alpha < 0, \beta < 0$: the triads close by means of repulsive-attractive interactions.
- $\alpha < 0, \beta > 0$: the triads close by means of repulsive-repulsive interactions.

In Table 3 we have arranged the values of $\langle\alpha^*\rangle$ and $\langle\beta^*\rangle$ to correspond to these four classes. For instance, the networks Sawmill, Social3, Matheoremethod, Galesburg, Prison, Zachary, and Colorado (all social networks), Grassland and Bridge Brook (food webs), and Transc_yeast (a gene transcription network) close their triads following a scheme of attractive-attractive interactions. The three social networks of High Tech, Drugs, and Geom, as well as the networks of USAir97 (air transportation network), neurons (neural network), Ythan1 (a food web), and the Internet at Autonomous System level, all belong to the class of networks in which triads are closed by an attraction-repulsion mechanism. The only network with a repulsion-attraction triad closure mechanism is the social network of High School, while there are seven networks closing triads with a repulsion-repulsion mechanism (three protein-protein interaction (PPI) networks, two food webs, one animal social network, and the Roget thesaurus).

TABLE 3
Results of the proposed method for predicting triad closure in 25 complex networks.

Network	r	detected	rand	$\langle\alpha^*\rangle$	$\langle\beta^*\rangle$
Sawmill	18	27%	10%	1.906	-1.25
social3	32	24%	11%	1.164	-1.258
Matheoremetho	19	25%	10%	1.196	-0.574
Grassland	30	25%	9%	1.833	-1.203
Galesburg	29	23%	11%	0.902	-0.648
Prison	58	30%	12%	0.294	-1.492
Zachary	45	42%	10%	1.696	-0.392
BridgeBrook	774	13%	3%	1.977	-1.046
Colorado	17	20%	1%	0.754	-0.044
Transc_yeast	72	4%	1%	0.221	-0.544
USAir97	12181	45%	18%	1.452	0.63
High tech	77	31%	16%	0.198	0.288
Drugs	3598	27%	16%	0.526	1.048
Neurons	2808	16%	8%	0.526	0.978
Geom	12325	12%	6%	0.14	1.149
Ythan1	507	10%	4%	0.248	0.492
Internet	2331	26%	0%	0.1	1.842
High School	199	28%	18%	-0.654	-0.434
Dolphins	95	24%	13%	-0.364	0.586
ScotchBroom	358	31%	4%	-0.372	0.660
StMartin	278	16%	11%	-0.232	0.335
PIN_Ecoli	478	10%	5%	-1.025	0.137
PIN_Yeast	3530	13%	4%	-1.53	1.842
PIN_Human	1047	5%	2%	-0.203	0.291
Roget	1550	7%	6%	-0.305	0.008

The group of networks with attractive-attractive interactions consists of 63% of all the social networks studied here. Among them we find a communication network within a small enterprise: a sawmill, where all employees were asked to indicate the frequency with which they discussed work matters with each of their colleagues on a five-point scale ranging from less than once a week to several times a day. Two employees were linked in the communication network if they rated their contact as a three or higher. This is a cooperative process in which both advisers and advisees cooperate to share the information needed for improving their skills and knowledge. Thus, closing the potential triangles in order to enhance the communication between the individuals involved seems a very reasonable mechanism. The other social networks included in this class all share a common characteristic. In the networks Social3 (a network of social contacts among college students participating in a leadership course), Galesburg (a network of friendship among physicians), and Matheoremetho (a network of friendship among school superintendents), the participants in the respective studies were asked the following questions:

- Choose the three members they wished to include in a committee.
- Nominate three doctors with whom they would choose to discuss medical matters.
- Name their best friends among the chief school administrators in Allegheny County.

In the first two cases it is very clear that the participants were asked to nominate individuals with whom they would easily cooperate, e.g., members of a committee or colleagues with whom to discuss medical matters. The third resembles a general kind of friendship relation, but the question was formulated in the context of analyzing the diffusion of a new mathematical method among high schools in the county. Thus, se-

lecting your best friends among other chief school administrators also means selecting those with whom you would easily cooperate in technical matters. These facts may explain the kind of attraction-attraction interaction which dictates the main mechanism for closing the triads in these networks. Transmission of information through the edges as well as a direct long-range interaction between peers both benefit the cooperative atmosphere needed for performing the tasks for which these networks are created.

In the class of networks in which triads have been closed by attraction-repulsion mechanisms, we find networks of very different natures, and it is difficult to extend the previous analysis to all these networks. This group includes a social network in a small high-tech computer firm which sells, installs, and maintains computer systems, where individuals were asked “Who do you consider to be a personal friend?” It could be speculated that a mechanism of the type based on Simmelian principles dominates here. That is, if $A - B - C$ is a triad and the two pairs $A - B$ and $B - C$ have strong social relations, it is natural to think that there is not a strong repulsion between A and C and they can create a new social tie. The friendship network among boys in a high school, which is the only one showing repulsion-attraction mechanisms, was created by asking the pupils “What fellows here in school do you go around with most often?” The triads here are formed when the relations between the pairs $A - B$ and $B - C$ are not strong enough to tie A and C together. If the pairs $A - B$ and $B - C$ have some strong relation, i.e., if they are dating, a link between A and C could be seen as offensive to the already established couples. The final class of networks, that characterized by repulsion-repulsion interactions, does not contain any human social network. The three PPIs included in this study are characterized by this type of triad closure mechanism, together with two food webs, an animal social network, and a thesaurus. The repulsion-repulsion mechanism is characterized by weak through-edge transmission of information and weak long-range interaction between pairs of nodes separated by two adjacent edges. Thus, it is expected that the triad closure is controlled by nontopological factors, such as similarities or complementarities among the nodes. This is a plausible explanation for the case of the PPI networks where triads of proteins may form triangles due to their functional similarities. We notice that, as expected, the percentages of correct prediction of triad closure in this group are the smallest of the four groups. That is, the difference between the predictions made by the proposed method and the random one in this group is 8.7%, in contrast with 15.5% for the attraction-attraction, 14.1% for the attraction-repulsion, and 10% for the only network with repulsion-attraction mechanisms.

6.2. Network evolution under triadic closure. Finally, in this section we use the results of the proposed method for modeling the triadic closure evolution in a given network. Although we can model the future evolution of a network from its current state, we prefer to consider a network in an early stage of its evolution and to predict how it has evolved toward its current structure. This method allows us to contrast the predictions made by the current method with some control parameters obtained for the real-world network in its current state. For this experiment we selected the network of injecting drug users (Drugs) for which we consider the clustering coefficient, the average path length, and the average communicability of the actual network. In order to perform these experiments, we select 50% of the total number of triangles existing in the network, and we remove one edge from each of them. Edges are selected uniformly at random among those present in the corresponding triangle. As before, let L be the list of edges obtained from the union of the potential edges and of those we removed. The values for α^* and β^* are those determined empirically using the calibration method already described (cf. Table 3).

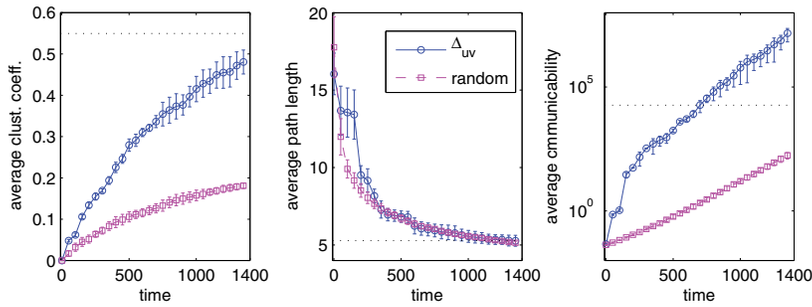


FIG. 2. Illustration of the evolution of the clustering coefficient, average path length, and average communicability for the network of injecting drug users (Drugs) versus the number of links added using the function (Δ_{uv}) and at random (see the text for explanations).

The iteration process goes as follows. We select the potential edge realizing the smallest value for $\Delta_{uv}(\alpha^*, \beta^*)$, and we add this edge to the network. Then we compute the values of the parameters of interest: the average clustering coefficient, the average path length, and the average communicability. Finally, the values for $\Delta_{uv}(\alpha^*, \beta^*)$ are recomputed using the new adjacency matrix, obtained by the addition of the selected potential edge. Here every addition of an edge is considered as a time step.

This iteration is run as many times as the number of edges we have removed. That is, if we removed r edges, we consider a discrete time evolution for $0 \leq t \leq r$. We then repeat this experiment 10 times, taking the average and standard deviations of the corresponding parameter. In order to compare the results, we simulate the same process by adding such edges uniformly at random.

The results of this experiment are illustrated in Figure 2, where we plot the values for the parameters of interest (with the corresponding error bar) versus time. The horizontal dotted line represents the actual value of the property for the original real-world network. As can be seen in Figure 2, the proposed method outperforms the random one for predicting the clustering coefficient of the network. The current value of \bar{C} for this network is 0.549, while the one predicted by Δ_{uv} is 0.486, which contrasts with that of 0.183 obtained by the random method. We remark here that this is the most important parameter to be considered in this experiment as it is the one which accounts more directly for the ratio of triangles to paths of length two in the network. Both methods predict the average path length of the network very well, returning values very close to the actual one ($\bar{\ell} = 5.287$). In addition, the proposed method increases the average communicability of the network more significantly than the random triadic closure. This feature is important when one is interested in maximizing the total average communicability of a network, which is equivalent to increasing the quality of communication among the nodes in the network.

7. Conclusions. The prediction of triadic closure is a very important and far from trivial problem in network theory. The fact that most real-world complex networks have more triangles than random counterparts makes the problem interesting per se. In addition, there is a large amount of evidence that shows that triadic closure in (social) networks is an important driver for other important structural characteristics of networks, such as degree distributions, clustering, and community structure. In this work, we introduce a triad closure mechanism based on communicability distances among pairs of nodes in a network. Our results show acceptable levels of predictability

and interpretability of the potential mechanisms controlling triad closure in real-world networks. Finally, we would like to remark on some other potentialities of the current approach. In a recent paper, Cozzo et al. [11] have considered the structure of triadic relations in multiplex networks. A multiplex is a complex structure formed by several layers representing the same series of nodes which are connected differently in each layer [6, 26]. Those layers may be interconnected by a certain coupling factor which allows the representation of the system as a supra-adjacency matrix [4]. By using this matrix representation of a multiplex the communicability function has been obtained and studied for social and technological multiplexes [13]. Consequently, the current approach can be extended to consider communicability distances among the nodes in multiplexes and in this way to apply the current methodology to detecting triadic closure in these structures.

Appendix.

In this section we give a brief description of the networks used for the tests throughout the paper. For each network in the dataset, Table 4 contains the number of nodes (n), the number of existing triangles in the network (r), and the number of open triads ($|P|$).

TABLE 4

Dataset: n is the number of nodes in the network, r is the number of existing triangles, and $|P|$ is the number of open triads.

Name	n	r	$ P $
Matheoremethod	30	19	175
Galesburg	31	29	224
High tech	33	77	390
Zachary	34	45	393
Sawmill	36	18	165
social3	37	32	299
StMartin	44	278	1732
Dolphins	62	95	638
Prison	67	58	430
High School	69	199	874
BridgeBrook	75	774	9829
Grassland	75	30	427
Ythan1	134	507	9019
ScotchBroom	154	358	4094
PIN_Ecoli	230	478	7803
Neurons	280	2808	33973
USAir97	332	12181	55646
Colorado	324	17	1273
Drugs	616	3598	18533
Transc_yeast	662	72	13069
Roget	994	1550	30116
PIN_Yeast	2224	3530	92882
PIN_Human	2783	1047	85617
Internet	3015	2331	462232
Geom	3621	12325	127794

Brain networks:

- **Neurons:** Neuronal synaptic network of the nematode *C. elegans*. Includes all data except muscle cells and uses all synaptic connections [54, 41].

Ecological networks:

- **BridgeBrook:** Pelagic species from the largest of a set of 50 New York Adirondack lake food webs [47].

- Grassland: All vascular plants and all insects and trophic interactions found inside stems of plants collected from 24 sites distributed within England and Wales [36].
- ScotchBroom: Trophic interactions between the herbivores, parasitoids, predators, and pathogens associated with broom, *Cytisus scoparius*, collected in Silwood Park, Berkshire, England, UK [37].
- StMartin: Birds and predators and arthropod prey of Anolis lizards on the island of St. Martin, which is located in the northern Lesser Antilles [35].
- Ythan1: Mostly birds, fishes, invertebrates, and metazoan parasites in a Scottish Estuary [25].

Informational networks:

- Roget: Vocabulary network of words related by their definitions in *Roget's Thesaurus of English Words and Phrases*. Two words are connected if one is used in the definition of the other [49].

PPI networks:

- PIN_Ecoli: Protein-protein interaction network in *Escherichia coli* [8].
- PIN_Human: Protein-protein interaction network in human [50].
- PIN_Yeast: Protein-protein interaction network in *S. cerevisiae* (yeast) [7, 38].

Social and economic networks:

- Colorado: The risk network of persons with HIV infection during its early epidemic phase in Colorado Springs, (Colorado, USA) using analysis of community wide HIV/AIDS contact tracing records (sexual and injecting drugs partners) from 1985–1999 [48].
- Dolphins: Social network of frequent association between 62 bottlenose dolphins living in the waters off New Zealand [33].
- Drugs: Social network of injecting drug users (IDUs) that have shared a needle in the last six months [43].
- Galesburg: Friendship ties among 31 physicians [10, 28, 46].
- Geom: Collaboration network of scientists in the field of Computational Geometry [3].
- High School: Network of relations in a high school. The students chose the three members they wanted to have in a committee [56].
- High tech: Friendship ties among the employees in a small high-tech computer firm which sells, installs, and maintains computer systems [29, 46].
- Matheoremethd: This network concerns the diffusion of a new mathematics method in the 1950s. It traces the diffusion of the modern mathematical method among school systems that combine elementary and secondary programs in Allegheny County (Pennsylvania, USA.) [9, 46].
- Prison: Social network of inmates in prison who chose “What fellows on the tier are you closest friends with?” [34].
- Sawmill: Social communication network within a sawmill, where employees were asked to indicate the frequency with which they discussed work matters with each of their colleagues [39, 46].
- social3: Social network among college students in a course about leadership. The students chose which three members they wanted to have in a committee [56].
- Zachary: Social network of friendship among the members of a karate club [55].

Technological networks:

- Internet: The Internet at the autonomous system (AS) level as of September, 1997, and of April, 1998 [20].

- USAir97: Airport transportation network between airports in the U.S.A. in 1997 [3].

Transcription networks:

- Transc_yeast: Direct transcriptional regulation between genes in *Saccharomyces cerevisiae* [41, 42].

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