Network landscape from a Brownian particle’s perspective

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(Received 22 August 2002; revised manuscript received 13 January 2003; published 21 April 2003)

Given a complex biological or social network, how many clusters should it be decomposed into? We define the distance \(d_{ij}\) from node \(i\) to node \(j\) as the average number of steps a Brownian particle takes to reach \(j\) from \(i\). Node \(j\) is a global attractor of \(i\) if \(d_{ij}=d_{ik}\) for any \(k\) of the graph; it is a local attractor of \(i\) if \(j\in E_i\), the set of nearest neighbors of \(i\) and \(d_{ij}=d_{lk}\) for any \(l\in E_i\). Based on the intuition that each node should have a high probability to be in the same community as its global (local) attractor on the global (local) scale, we present a simple method to uncover a network’s community structure. This method is applied to several real networks and some discussion on its possible extensions is made.

DOI: 10.1103/PhysRevE.67.041908 PACS number(s): 87.10.+e, 89.75.–k, 89.20.–a

A complex networked system, such as an organism’s metabolic network and genetic interaction network, is composed of a large number of interacting agents. The complexity of such systems originates partly from the heterogeneity in their interaction patterns, aspects of which include the small-world [1] and the scale-free properties [2,3] observed in many social, biological, and technological networks [4–6]. Given this high degree of complexity, it is necessary to divide a network into different subgroups to facilitate the understanding of the relationships among different components [7,8].

A complex network could be represented by a graph. Each component of the network is mapped to a vertex (node), and the interaction between two components is signified by an edge between the two corresponding nodes, whose weight is related to the interaction strength. The challenge is to dissect this graph based on its connection pattern. We know that to partition a graph into two equally sized subgroups such that the number of edges in between reaches the absolute minimum is already a \(NP\)-complete problem, a solution is not guaranteed to be found easily; however it is still a well-defined question. On the other hand, the question “How many subgroups should a graph be divided into and how?” is ill posed, as we do not have an objective function to optimize; and we have to rely on heuristic reasoning to proceed.

If we are interested in identifying just one community that is associated with a specified node, the maximum flow method [9] turns out to be efficient. Recently, it is applied to identifying communities of Internet webpages [10]. An community thus uncovered is usually very small; and for this method to work well one needs \textit{a priori} knowledge of the network to select the source and sink nodes properly. Another elegant method is based on the concept of edge betweenness [11]. The degree of betweenness of an edge is defined as the total number of shortest paths between pairs of nodes which pass through it. By removing recursively the current edge with the highest degree of betweenness, one identifies communities of Internet webpages by assigning each edge a length equalling its reciprocal weight. Furthermore, in the sociology literature, there is a relatively long tradition in identifying communities based on the criteria of reachability and shortest distance (see, e.g., Ref. [12]).

In this paper, a method of network community identification is described. It is based on the concept of network Brownian motion: If an intelligent Brownian particle lives in a given network for a long time, what might be its perspective of the network’s landscape? We suggest that, without the need to remove edges from the network, the node-node distances “measured” by this Brownian particle can be used to construct the community structure and to identify the central node of each community. This idea is tested on several social and biological networks and satisfactory results are obtained. Several ways are discussed to extend and improve our method.

Consider a connected network of \(N\) nodes and \(M\) edges. Its node set is denoted by \(V=\{1,\ldots,N\}\) and its connection pattern is specified by the generalized adjacency matrix \(A\). If there is no edge between node \(i\) and node \(j\), \(A_{ij}=0\); if there is an edge in between, \(A_{ij}=A_{ji}>0\) and its value signifies the interaction strength (self-connection is allowed). The set of nearest neighbors of node \(i\) is denoted by \(E_{i}\). A Brownian particle keeps moving on the network, and at each time step it jumps from its present position (say \(i\)) to a nearest-neighbor position \(j\). When no additional knowledge about the network is known, it is natural to assume the following jumping probability \(P_{ij}=A_{ij}/\sum_{j=1}^{N}A_{ij}\) (the corresponding matrix \(P\) is called the transfer matrix). One verifies that at time \(t\gg M\) the probability \(p(k)\) for the Brownian particle to be at any node \(k\) is \textit{nonvanishing} and equals to \(\sum_{i}A_{ki}/\sum_{m,n}A_{mn}\), proportional to the total interaction capacity \(\Sigma_{i}A_{ki}\) of node \(k\).

Define the node-node distance \(d_{ij}\) from \(i\) to \(j\) as the average number of steps needed for the Brownian particle to move from \(i\) through the the network to \(j\). From some simple linear-algebra calculation [13] it is easy to see that

\[
d_{ij} = \sum_{l=1}^{N} \left( \frac{1}{1-B(j)} \right)_{il},
\]

where \(I\) is the \(N\times N\) identity matrix and matrix \(B(j)\) equals to the transfer matrix \(P\) except that \(B_{li}(j)=0\) for any \(l\)
\( \in V \). The distances from all the nodes in \( V \) to node \( j \) can thus be obtained by solving the linear algebraic equation 
\[
(1 - \mathbf{B}(j)) \mathbf{d}_{ij} = \mathbf{1},
\]
where \( \mathbf{B} \) is the adjacency matrix of \( G \). We are mainly interested in sparse networks with \( M = O(N) \); for such networks there exist very efficient algorithms [14,15] to calculate the root of this equation. If node \( j \) has the property that 
\[
d_{ij} < d_{ik} \quad \text{for any } k \in V,
\]
then \( j \) is tagged as a global attractor of node \( i \) (we are intuitively expecting that each node will have a high probability to be in the same subgroup as its local attractor \( j \), since among all the nearest neighboring nodes in \( E_i \), node \( j \) has the shortest “distance” from node \( i \). For simplicity let us just assume this probability to be unity (a possible improvement is discussed later). Thus, we can define a local-attractor-based community (or simply a “L community”) as a set of nodes \( L = \{i_1, \ldots , i_m\} \) such that (1) if node \( i \in L \) and node \( j \) is a local attractor of \( i \), then \( j \in L \), (2) \( i \) is in \( L \), and node \( k \) has \( i \) as its local attractor, then \( k \in L \), and (3) any subset of \( L \) is not a L community. Clearly, two L communities \( L_a \) and \( L_b \) are either identical \( (L_a = L_b) \) or disjoint \( (L_a \cap L_b = \emptyset) \). Based on each node’s local attractor the graph could be decomposed into a set of L communities.

According to the same intuitive argument, on the global scale we expect that each node will have a high probability to be in the same community as its global attractor, and if assume this probability to be unity we can similarly construct the global-attractor-based communities (“G communities”) based on the global-attractor of each node. For small networks, we expect the L- and G-community structures to be identical; while for large networks, each G community may contain several L-communities as its subgroups. A community could be characterized by its size \( N_c \) and an instability index \( I_c \). A node \( i \) in community \( C \) is referred to as unstable if its total direct interaction with nodes in any other community \( C' \), \( \sum_{k \in C'} A_{ik} \), is stronger than its total direct interaction with other nodes in its own community, \( \sum_{k \in C} A_{ik} \cdot I_c \), is the total number of such nodes in each community. We can also identify the center of a community (if it exists) as the node that is the global attractor of itself.

Now we test the above-mentioned simple method on some well-documented networks whose community structures are known. The first example is the social network recorded by Zachary [16]. This network contains 34 nodes and 77 weighted edges, and it was observed to spontaneously fission into two groups of size 16 and 18, respectively [16] [these two groups are marked by two colors in Fig. 1(a)]. The results of our method is shown in Fig. 1(a). Community \( L_1 \) contains 11 elements (node 13 is unstable and has stronger direct interaction with \( L_2 \). \( L_2 \) has 6 elements (node 9 has stronger direct interaction with \( L_1 \), and \( L_3 \) has 17 elements. Nodes 1 (the manager), 3, and 34 (the officer) are the corresponding centers. We find that for this network the G communities coincide with the L communities.

As another example, the scientific collaboration network of Santa Fe Institute [7] is considered. The giant connected component contains 118 nodes and 200 weighted edges, the weights are assigned according to the measure in Ref. [17]. The present method divides the network into six L communities, see Fig. 1(b). All the nodes in community \( L_1 \) (size 14), \( L_2 \) (41), \( L_4 \) (8), \( L_5 \) (26), and \( L_6 \) (17) are locally stable, and one node in \( L_3 \) has stronger direct interaction with community \( L_6 \). Same as the above example, the G-community structure is also identical to the L-community structure. Girvan and Newman divided this network into four major groups by recursively removing edges of highest degree of betweenness [7]; the largest of which was further divided into three subgroups and the second largest was divided into two subgroups. There are still some minor differences between the six subgroups obtained by the present method and those obtained in Ref. [7], which may be attributed to the fact that, in the treatment of Ref. [7] the network was regarded as unweighted.

The method is further tested on a relatively more complicated case, the football match network compiled by Girvan and Newman [7]. It contains 115 nodes and 613 unweighted edges. These 115 teams were distributed into 12 conferences by the game organizers. Based on the connection pattern, the present method divides them into 15 L communities, of which 11 are locally stable: \( L_2 \) (size 9), \( L_3 \) (13), \( L_4 \) (14), \( L_5 \) (10), \( L_6 \) (8), \( L_7 \) (6), \( L_8 \) (7), \( L_9 \) (6), \( L_{10} \) (4), \( L_{11} \) (6), and \( L_{13} \) (size 9). One element of \( L_1 \) (size 9) has stronger interaction with \( L_{10} \), and one element of \( L_{12} \) (size 10) has stronger interaction with \( L_3 \), and all the elements of \( L_{14} \) (size 2) and \( L_{15} \) (size 2) are locally unstable. The G communities of this network are also identical to the L communities. In Fig. 1(c) the community structure of this network is shown, where nodes belonging to each identified community are located together, and the different colors encode the actual 12 conferences [7]. Figure 1(c) indicates that the predicted communities coincide very well with the actual communities. The community structure obtained by the present method is also in very good correspondence with that obtained by Girvan and Newman [7] based on edge betweenness.

The above-studied networks all have relatively small network sizes and the identified G communities coincide with the L communities. Now we apply our method to the protein interaction network (yeast core [18,19]) of baker’s yeast. The giant connected component of this network contains 1471 proteins and 2770 edges (assumed to be unweighted, since the interaction strengths between the proteins are generally undetermined). The present method dissect this giant component into 14 G communities (Table I) and into 69 L communities (11 of them contain one locally unstable node, 15 of them have 2–7 locally unstable nodes, all the others are stable). The relationship between the G and L communities is demonstrated in Fig. 1(d), where proteins are grouped into L communities and those of the same G community have the same color. We see from Fig. 1(d) that if two nodes are in the same L community, they are very probable to be in the same G community. The largest G community \( G_1 \) contains more
than half of the proteins and is centered around nucleoporin YMR047C, which, according to SWISS-PROT description, is “an essential component of nuclear pore complex” and “may be involved in both binding and translocation of the proteins during nucleocytoplasmic transport.” YMR047C interact directly only with 39 other proteins (it is even not the most connected node in the system), but associated with it is a group of 935 proteins as suggested by the present method. The protein interaction network may be evolved to facilitate efficient protein transportation by protein-mediated indirect interactions.

What will happen if the protein YMR047C is removed from the network? The resulting perturbed system has 1463 nodes and 2729 edges, and we find that its L-community structure does not change much. Altogether 72 L communities are identified, and most of them contain more or less the same set of elements as in the unperturbed network. However, there is a dramatic change in the G community structure. There are now 21 G communities (the largest of which has 574 proteins), while G_1 of the original system breaks up into eight smaller G communities. It was revealed that the most highly connected proteins in the cell are the most im-

FIG. 1. (Color) Community structure of some model networks (the nodes of the same L community are spatially grouped together). (a) The karate club network compiled by Zachary [16] (here nodes are colored according to their actual groupings); (b) the scientific collaboration network compiled by Girvan and Newman [7]; (c) the football match network compiled by Girvan and Newman [7] (nodes are colored according to their actual groupings); and (d) the yeast protein interaction network [18,19], here nodes of the same G community are encoded with the same color (open circles denote nodes in G_1).
important for its survival, and mutations in these proteins are usually lethal [21]. Our work suggests that, these highly connected proteins are especially important because they help integrating many small functional modules (L communities) into a larger unit (G community), enabling the cell to perform concerted reactions in response to environment stimuli.

In the above examples, the network studied are all from real world. We have also tested the performance of our method to some artificial networks generated by computer. To compare with the result of Ref. [7], we generated an ensemble of random graphs with 128 vertices. These vertices are divided into four groups of 32 vertices each. Each vertex has on average 16 edges, \( z_{\text{out}} \) of which are linked to vertices of other groups, and the remaining are to vertices within its group; all these edges are drawn randomly and independently in all the other means. Using the method of Girvan and Newman, it was reported [7] that when \( z_{\text{out}} < 6 \) all the vertices could be classified with high probability. Our present method in its simplest form could work perfectly only when \( z_{\text{out}} < 2.5 \). In the artificial network, the vertices are identical to each other in the statistical sense and there is no correlation between the degrees of two neighboring edges. Our method seems not to be the best for such kind of random networks. However, an improved method based on the present work [24] outperforms Ref. [7].

In summary, we have suggested a simple way of grouping a graph of nodes and edges into different subgraphs based on the node-node distance measured by a Brownian particle. The basic idea was applied to several real networked systems and very encouraging results were obtained. The concept of random walking was also used in some recent efforts to facilitate searching on networks (see, e.g., Refs. [22,23]), the present work is an attempt in applying it on identifying network community structure. Some possible extensions of our method are immediately conceivable: First, in the present work we have assumed that a node will be in the same community as its attractor with probability 1. Naturally, we can introduce an “inverse temperature” \( \beta \) and suppose that node \( i \) be in the same community as node \( j \) with probability proportional to \( \exp(-\beta d_{ij}) \). The present work discusses just the zero temperature limit. We believe that the communities identified at zero temperature will persist until the temperature is high enough. Second, we can construct a coarse-grained network by regarding each L community as a single node, and defining the distance from one L community to another as the average node-node distance between nodes in these two communities. The present method can then be applied, and the relationship between different L communities can be better understood. Third, for very large networks, it is impractical to consider the whole network when calculating node-node distance. Actually this is not necessary, since the length of the shortest path between a given node and its attractor should be small. We can therefore focus on a localized region of the network to identify the attractor of a given node.

Furthermore, based on the distance measure of the present paper, we can define a quantity called the dissimilarity index for any two nearest-neighboring nodes. Nearest-neighboring vertices of the same community tend to have small dissimilarity index, while those belonging to different communities tend to have high dissimilarity index. Extensions of the present work will be reported in a forthcoming paper [24].

An interesting task is to use extended versions of the present method to explore the landscape of the Internet’s autonomous system [3] and that of the metabolic network of coli [8,25].

I am grateful to M. Girvan and M. E. J. Newman for sharing data and to Professor R. Lipowsky for support.

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