

Skeleton and Fractal Scaling in Complex Networks

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We find that the fractal scaling in a class of scale-free networks originates from the underlying tree structure called a skeleton, a special type of spanning tree based on the edge betweenness centrality. The fractal skeleton has the property of the critical branching tree. The original fractal networks are viewed as a fractal skeleton dressed with local shortcuts. An *in silico* model with both the fractal scaling and the scale-invariance properties is also constructed. The framework of fractal networks is useful in understanding the utility and the redundancy in networked systems.

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Emerging unifying concepts such as small-world property [1], scale-free behavior [2], and hierarchical modularity [3] now constitute our basic understanding of the organization of complex networked systems, which appear in such diverse examples as the World Wide Web, social networks, and biochemical reaction networks inside cells. The small-world property refers to the one that the average separation $\langle D \rangle$ between pairs of vertices in the network scales at most logarithmically in the total number of vertices N in the system, $\langle D \rangle \sim \ln N$. Scale-free behavior means the lack of characteristic scales in the number of links k a vertex has, called the degree, manifesting itself in the form of a power-law degree distribution $p_d(k) \sim k^{-\gamma}$ for large k . The recent discovery of fractal scaling and topological self-similarity in the World Wide Web and metabolic networks [4], however, raised a new perspective on our view of such networked systems. Fractal scaling stands for the power-law relation between the minimum number of boxes N_B needed to cover the entire network and the size of the boxes ℓ_B ,

$$N_B(\ell_B) \sim \ell_B^{-d_B}, \quad (1)$$

with a finite fractal dimension d_B [5]. Self-similarity here refers to the scale invariance of the degree distribution under coarse graining with different box sizes (length scales) as well as under the iterative application of coarse graining (the network renormalization) [4,6]. It has been observed, however, that not all networks are fractal and most of the random network models proposed yet are not fractal, either. This poses a fundamental question on the origin of the fractal scaling observed in real-world networks [4,7–9]. In this Letter, we show that the fractal property of the network can be understood from its underlying tree structure.

As highly entangled as a network looks, a more simple structure is embedded underneath it, that is, the spanning tree. A spanning tree is a tree composed of $N - 1$ edges in a way that they connect all the N vertices in the network. Of particular significance is the so-called skeleton [10] of a network. The skeleton is a particular spanning tree, which

is formed by the edges with highest betweenness centralities [11,12] or loads [13]. The remaining edges in the system are called shortcuts, which contribute to forming loops. The skeleton of a scale-free network is also scale-free but with different γ . Since the betweenness centrality is related to the amount of information flow along a given edge, the skeleton can be considered as the “communication kernel” of the network [10]. If a network is organized in a modular way, as it is believed to be for the World Wide Web and biological systems, the intermodular connections offer communication channels across the modules, thus gaining high-betweenness centralities. By construction, the skeleton is composed preferentially of such high-betweenness intermodular connections, which will preserve the modular structure while greatly simplifying the complexity. Furthermore, if the modular structure is distinct enough, i.e., there is a rather clear-cut separation between modules, we can expect that even a random spanning tree can capture the modular structure. Thus, by looking at the properties of its spanning trees, we can visualize more easily the topological organization of the network.

With the underlying skeleton and random spanning tree, here we perform the fractal scaling analysis by measuring $N_B(\ell_B)$ for several real-world networks and network models [14]. Comparison of the fractal scalings in each original network with the corresponding spanning trees reveals distinct patterns according to the presence or the absence of fractality in the network. For the fractal networks, such as the World Wide Web [Fig. 1(a)] [15], the metabolic network of *Escherichia coli* [Fig. 1(b)] [16], and the protein interaction network of *Homo sapiens* [Fig. 1(c)] [17], the numbers of boxes needed to cover the original network and its skeleton are almost the same. Moreover, the random spanning tree, while possessing a different statistics of N_B , shows nevertheless the same fractal dimension d_B . This is surprising, because in the World Wide Web, for example, more than $2N$ edges (shortcuts) are added onto the skeleton (the average degree of the World Wide Web is 6.7), which is a tremendous number in the graph-theoretical sense and

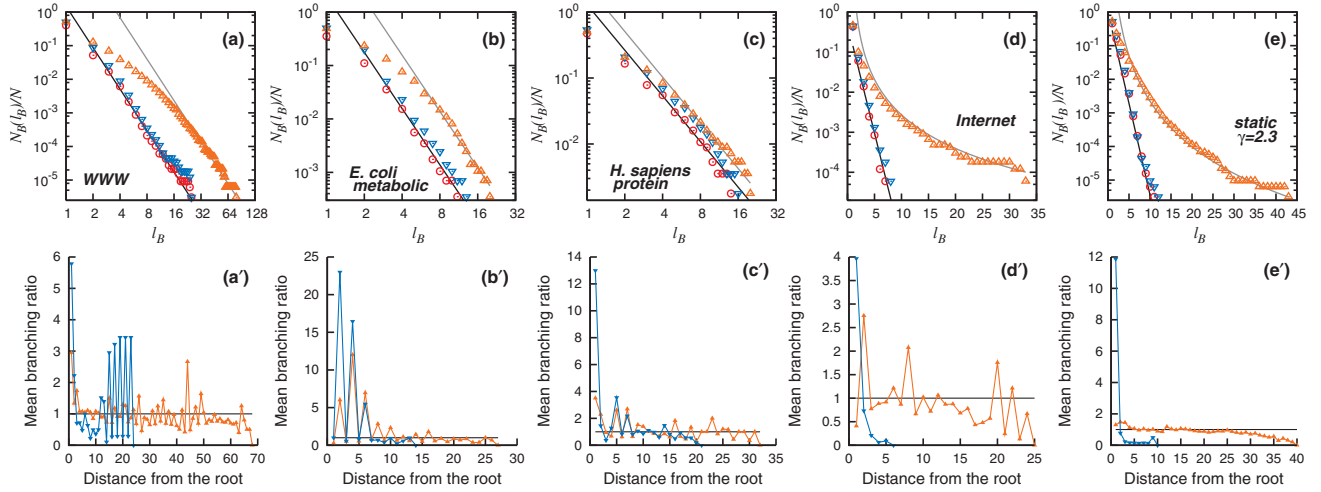


FIG. 1 (color). (a)–(e) Box counting analysis of original networks (\circ , red) and their skeleton (∇ , blue) and random spanning tree (\triangle , orange). Shown are cases for (a) the World Wide Web, (b) the metabolic network of *Escherichia coli*, and (c) the protein interaction network of *Homo sapiens*, (d) the Internet at the autonomous systems level as of the year 2004, and (e) the static model network with $\gamma = 2.3$ and $\langle k \rangle = 4$. In (a)–(c), we also show two guidelines for the fractal scaling of the original network (black) and its random spanning tree (gray). The slope of each guideline is (a) -4.1 , (b) -3.5 , and (c) -2.3 . In (d)–(e), the black line is a fit to the exponential function and the gray one is to a power law. (a')–(e') Branching analysis. The mean branching number as a function of distance from the root for the skeleton (∇ , blue) and the random spanning tree (\triangle , orange) of the networks in (a)–(e). For (a')–(c'), both the skeleton and the random spanning tree fulfill the criticality condition $\langle m \rangle = 1$ (horizontal line) as the distance from the root increases, while for (d')–(e'), the mean branching number of the skeleton decays to zero with no plateau at $\langle m \rangle = 1$.

by no means a minute perturbation. Such a robustness of fractal scaling in the World Wide Web shows that, even though the network is far from being a tree, the shortcuts are distributed in a way that they preserve the fractality and modularity. In other words, shortcuts are present mainly inside modules and the connections between different modules are made largely through the skeleton. This topological structure can be measured by the fraction of intra-branch shortcuts among the total number of shortcuts, a branch being the subtree connected to the most connected vertex. We find that the ratio is 0.78, 0.33, and 0.45 for Figs. 1(a)–1(c), respectively. On the other hand, other networks exhibit different features in the fractal scaling analysis. For example, in the Internet autonomous systems [Fig. 1(d)] or the static model with $\gamma = 2.3$ and $\langle k \rangle = 4$ [Fig. 1(e)], the box counting number of the original network and the skeleton decays with ℓ_B much faster than that of the random spanning tree, and the fraction of intra-branch shortcuts is small: 0.087 for Fig. 1(d) and 0.015 for Fig. 1(e). Also for social networks, such as the actor network and the collaboration network, the $N_B(\ell_B)$ curves are appreciably different from those of the skeletons, implying that the global topology of the social network is highly interwoven on a large scale to form a more compact structure [18].

The scaling behavior of the box counting relation Eq. (1) in Figs. 1(a)–1(c) for the original network and its skeleton suggests that the fractal property of the network originates from that of the skeleton. In addition, we argue here that the criticality in the topology of the skeleton is required for

a network to be a fractal: The tree structures, such as the skeleton and the random spanning tree, may be seen as generated through a multiplicative branching process starting from a root vertex [19]. At each branching step, each vertex born in the previous step generates m offsprings with probability b_m . The criticality condition means the average branching number

$$\langle m \rangle \equiv \sum_{m=0}^{\infty} m b_m = 1. \quad (2)$$

Thus, the branching tree grows perpetually with offspring neither flourishing nor dying out. In this case, when $b_m \sim m^{-\gamma}$, the number of vertices s in the tree scales with its linear size t in a power-law form as $s \sim t^z$, with $z = (\gamma - 1)/(\gamma - 2)$ for $2 < \gamma < 3$ and $z = 2$ for $\gamma > 3$ [19,20], and the tree structure is fractal with fractal dimension $d_B = z$. Such a critical branching tree is similar in the topological characteristics to the homogeneous scale-free tree network proposed in Ref. [21]. To check the validity of our suggestion, we examine if the criticality condition is fulfilled for the skeleton and the random spanning trees of the four real-world networks and the static model in Figs. 1(a')–1(e'). Indeed, for the fractal networks [Figs. 1(a')–1(c')], both the skeleton and the random spanning tree fulfill the criticality condition, even though in reality, the dynamic origins of their formations may well be more complicated than the pure branching dynamics. In our analysis, the root is taken as the most connected vertex in the tree. On the other hand, for nonfractal networks, the mean number of branches of

the skeleton decays to zero rapidly as the distance from the root increases [see Figs. 1(d') and 1(e')]. A similar behavior is observed for the actor network as well [18]. Thus, the actor network is not a fractal. However, the random spanning tree satisfies the criticality condition in all cases, suggesting the generic fractal structure of this kind of trees as shown in Ref. [22]. In short, a fractal network contains a fractal skeleton underneath it, which is perturbed by local shortcuts, thus preserving its fractal property.

Incorporating all the findings so far, we set up a fractal network model. The model is based on the multiplicative branching tree [19]. We first introduce an exponent γ for the branching probability $b_m \sim m^{-\gamma}$ ($m \geq 1$), designed to produce the desired power-law degree distribution $p_d(k) \sim k^{-\gamma}$, with $\gamma > 2$. The branching probability is properly normalized to be critical; i.e., it satisfies Eq. (2). Next a parameter p is introduced to control the number of shortcuts added and, hence, the mean degree of the network. One final parameter q accounts for the relative frequency of the local and the global shortcuts. The construction of the model network proceeds as follows: (i) A tree is grown by the multiplicative branching rule with branching probability b_m . (ii) After the branching process, every vertex increases its degree by a factor p and attempts to make shortcuts to its local neighbors. (iii) For each successful shortcut in (ii), with probability q , we replace it by reconnecting it to a randomly chosen vertex, not restricted to its local neighbors. In the latter case, we choose the vertex with a weight in proportion to its degree in the branching tree, so as to maintain the same power-law scaling of the degree distribution. This rule is schematized in Fig. 2(e) [18]. Figure 2(a) is an illustrative example of a branching tree of size $N = 164$ and $\gamma = 2.3$. The vertices of the tree are colored according to which box they belong to in a particular box-covering with $\ell_B = 2$. For such a tree, even a simple graph drawing algorithm, such as Pajek [23], can capture its inherent hierarchical structure. Figure 2(b) shows the fractal network structure with the addition of local shortcuts, where we use $p = 0.5$ and $q = 0$. The hierarchical modular structure presented in the tree network [Fig. 2(a)] persists. On the other hand, the network with $q = 1$ shown in Fig. 2(c), in which the same number of shortcuts as in Fig. 2(b) are attached, does not retain the modularity. Such an absence of modularity can be readily seen in Fig. 2(d), a different layout of the same network as Fig. 2(c) generated by Pajek in an unsupervised manner. Consequently, the fractal property is preserved in the case of $q = 0$, whereas it is not for $q = 1$, as is clearly revealed by the box counting analysis in Fig. 2(f).

It is noteworthy that there exists an important distinction between the present model and other scale-free trees such as the Barabási-Albert tree [2] and the geometrically growing scale-free tree [24]. Such models are not fractal, because they do not fulfill the criticality condition. Indeed, their mean branching rate decreases to zero monotonically

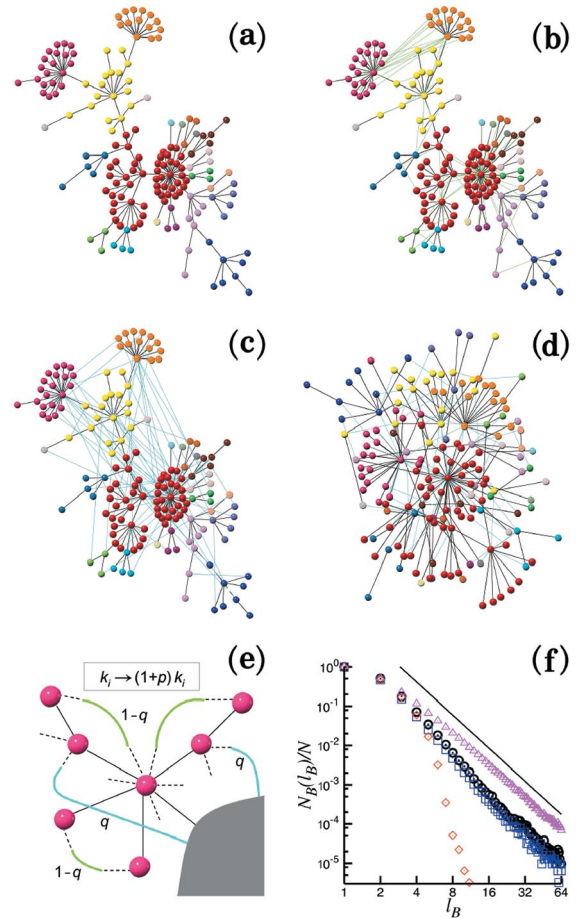


FIG. 2 (color). Fractal network model. (a) Uncorrelated scale-free tree with the degree exponent $\gamma = 2.3$ and the number of vertices $N = 164$. It is grown by the multiplicative branching rule [19], with the branching probability $b_m \sim m^{-\gamma}$. (b) Fractal model network created by adding local shortcuts (green) to the tree in (a). (c) A nonfractal model network created by adding random shortcuts (blue) allowing global connection. (d) The generic configuration of the network in (c) generated by Pajek, and the absence of inherent modular structure. In (b)–(d), the color of each vertex is that of the corresponding vertex in (a). (e) Schematic illustration of the growth rule of the fractal network model. The shaded region indicates the rest of the network generated so far. (f) The fractal scaling analysis of networks with larger size $N = 311,043$, constructed in the same manner as those in (a)–(c). Squares (blue) correspond to the fractal network model in (b) with $p = 0.5$, circles (black) to its skeleton, and the diamonds (red) to the nonfractal network model in (c) with $q = 1$. The solid line is the guideline with slope ≈ -2.8 .

as the branching proceeds, without a plateau at $\langle m \rangle \approx 1$ [18]. Such a type of tree network was classified as “causal” trees in Ref. [25]. Note also that the fractal trees are not small worlds. However, with a small number of global shortcuts, e.g., $q = 0.01$ in our model with $N \sim 3 \times 10^5$, the network turns into a small world. This is seen in the mean box mass versus ℓ_B plot in the cluster growing method analysis [18].

The fractal network model is self-similar. To check it, we perform the coarse graining through the box counting method by replacing each box with a single supernode and connecting them if any of their member vertices are connected [4]. We find that the degree distribution of the fractal network model is invariant under the coarse graining by the boxes with different sizes. In addition, the degree distribution is invariant under successive applications of the coarse-graining transformation [18]. It is interesting to note that some networks are self-similar, that is, exhibit the scale-invariant degree distribution, yet are not fractal. A typical example of such networks is the Internet [18]. So fractality and self-similarity do not always imply each other in complex networks.

The framework of a fractal network is helpful to understand, for example, the utility and the redundancy in the metabolic networks from a purely topological aspect. The high flux backbone in the metabolic network of *E. coli* obtained through the flux balance analysis was shown to be composed of many branches with few interbranch connections and to merge into the biomass reaction [26]. Obviously, its topological shape resembles the branching tree skeleton rooted from a vertex with the largest number of connections if the direction in edge is ignored. On the other hand, recent *in silico* flux analysis [27] has shown that the metabolic network of *E. coli* contains a high density of backup reactions (redundancies) for a given condition. Such a reaction is barely used in the normal condition but takes up a high flux when a certain reaction on the backbone is blocked. When the simultaneous blockade of such a reaction pair blocks biomass production, they are called synthetic lethal [28]. Most synthetic lethal reactions are located very close to each other, being apart in three reaction steps or less along the metabolic network. Also they are mostly in the same functional category. Thus, the reactions with high (low) flux in a wild type can be regarded as the edges on the skeleton (shortcuts) in the framework of the fractal network.

The critical branching tree that can be found in various phenomena, such as earthquake processes, population and biological dynamics, epidemics, social cascades, etc., also appears in the skeleton of fractal networks, as we found here. While the evolving process of the fractal networks would be complex and diverse depending on the specific systems, the underlying structure, the skeleton, has the topology of the critical branching tree. Identifying such a simple structure underneath is a step forward towards further studies on renormalization and universality in complex networks.

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