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Bipartite Structure of all Complex Networks
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Abstract

The analysis and modelling of various complex networks has received much attention in the last few years. Some such networks display a natural bipartite structure: two kinds of nodes coexist with links only between nodes of different kinds. This bipartite structure has not been deeply studied until now, mainly because it appeared to be specific to only a few complex networks. However, we show here that all complex networks can be viewed as bipartite structures sharing some important statistics, like degree distributions. The basic properties of complex networks can be viewed as consequences of this underlying bipartite structure. This leads us to propose the first simple and intuitive model for complex networks which captures the main properties met in practice.

Keywords: graphs, interconnection networks, modelling, bipartite graphs.

Context.

In a random network [7, 13] with \( n \) nodes, each of the \( \frac{n(n-1)}{2} \) possible links exists with a given probability \( p \). In other words, a random network is constructed from \( n \) nodes by choosing \( m = p \cdot \frac{n(n-1)}{2} \) links at random. Until recently, this model was the only one available for the study of complex networks.

However, it has been shown recently [2, 11, 25, 29, 31] that this model does not capture some of the main features of complex networks observed in practice. In particular the clustering (probability that two nodes are connected, given that they are both connected to a same third) and the degree distribution (probability that a randomly chosen node has \( k \) links, for each \( k \)) are inaccurate. Real-world complex networkshave a high clustering and a heavy-tailed degree distribution (degrees variety can be spread over several orders of magnitudes), while random networks of comparable size have a clustering equal to \( p \) (which is small when the average degree is small) and a Poisson-shaped degree distribution (all nodes nearly have a typical degree). In many contexts, these properties have a strong influence on phenomena of interest [4, 21, 24, 27]. For surveys on the variety of complex networks and associated phenomena that have been studied, see [2, 8, 11, 25, 28, 29].
Various models have recently been proposed to capture the main properties met in practice. The first one [31] provides highly clusterized networks but the obtained degree distributions is still Poisson-shaped. Another important step was the introduction of the preferential attachment principle [1, 12]: nodes are added one by one and link to preexisting nodes with a probability depending on the degree of these nodes. As explained in [1, 2, 6], this reflects the way a large variety of real-world networks are indeed constructed. The networks obtained have a power law degree distribution, but their clustering is much lower than in real-world networks.

Many other attempts have been made to produce networks having all the properties we have cited [2, 12, 26, 31]. However, none of them give at the same time an intuitive, realistic and simple interpretation of the causes of the observed properties. They either rely on artificial construction processes, produce networks which do not have all the desired properties, or are very difficult to analyze.

In this communication, we show that all complex networks have an underlying bipartite structure. This makes it possible to view their main properties as consequences of this underlying structure. This also leads to two very efficient models for the generation of complex networks having realistic properties. Moreover, these models can be studied both experimentally and analytically, and they give an interpretation of how these properties are induced by real-world construction processes. As we will discuss, all these aspects make these models interesting alternatives to those previously proposed.

Throughout our presentation, we will use a representative set of complex networks which have received much attention and span quite well the variety of context in which complex networks appear. The set consists of a protein network [9, 18], a map of the core of the Internet at router level [15, 16], the map of a large Web site (considered as undirected) [3, 9], the actors’ co-starring relation [10, 31], the co-occurrence relation of words [14] in the sentences of [30], and a co-authoring relation between scientists [5, 22, 23, 26]. We will refer to these networks as Proteins, Internet, Web, Actors, Cooccurrence, and Coauthoring respectively. They are precisely defined and studied in the cited references.

**Underlying bipartite structure.**

A bipartite network is a triple \( G = (\top, \bot, E) \) where \( \top \) and \( \bot \) are two disjoint sets of nodes, respectively the top and bottom nodes, and \( E \subseteq \top \times \bot \) is the set of links of the network. The difference with classical (unipartite) networks lies in the fact that links exist only between top nodes and bottom nodes.

Two degree distributions can naturally be associated to such a network, namely the top degree distribution: \( \top_k = \frac{|\{t \in \top : d^\top(t) = k\}|}{|\top|} \) and the bottom degree distribution: \( \bot_k = \frac{|\{t \in \bot : d^\bot(t) = k\}|}{|\bot|} \). These two distributions play a central role in the following.

Some complex networks display a natural bipartite structure. For instance, one can view Actors (two actors are linked if they are part of a same cast) as a bipartite network.
where \( \top \) is the set of movies, \( \bot \) is the set of actors, and each actor is linked to the movies he/she played in. Coauthoring can also be viewed this way with \( \top \) being the set of papers and \( \bot \) being the set of authors, each author being linked to the papers he/she (co-)signed. Likewise, in Cooccurrence one can link each sentence to the words it contains.

Given a bipartite network \( G = (\top, \bot, E) \), one can easily obtain its unipartite version defined as \( G' = (\bot, E') \) where \( \{u, v\} \in E' \) if \( u \) and \( v \) are both connected to a same (top) node in \( G \). See Figure 1. From the bipartite versions of Actors, Coauthoring and Cooccurrence networks, one can then recover their original (unipartite) versions. In this unipartite version of the network, each top node induces a clique (complete subnetwork) between the bottom nodes to which it is linked.

![Figure 1: A bipartite network and its unipartite version. Notice that the link \{B, C\} is obtained twice since B and C have two neighbors in common in the bipartite network.](image)

However, most complex networks do not display a natural bipartite structure. For instance there is no immediate way to see Internet, Web or Proteins as bipartite networks. We can however decompose these as follows. For each link \( \{u, v\} \) we compute the largest clique it belongs to. Notice that this clique may contain only two nodes. Moreover, if there are several largest cliques for the same link, we pick one at random. We then construct a bipartite network in which the top nodes are the obtained cliques (without repetition of the ones we may obtain several times), and the bottom nodes are the nodes of the network itself. A clique and a node are linked together if the node belongs to the clique in the original network.

In the case of Figure 1 we obtain several cliques of size 2 (namely \{C, E\} and \{D, F\}), and we have to choose at random between \{A, B, C\} and \{B, C, D\} when considering the link \{B, C\}. However, these two cliques are obtained from other links, and we finally obtain a unique decomposition which is nothing but the bipartite network on the left of the figure.

Notice that the computation of the largest clique containing a given link \( \{u, v\} \) may be very expensive (it is NP-complete). However, some heuristics make it possible to compute it if the network is not too large. In our case, we use the following remarks. Let us denote the sets of neighbors of a node and a link by \( N(u) = \{v \in V | \{u, v\} \in E\} \) and \( N(u, v) = N(u) \cap N(v) \) respectively. First notice that a largest clique containing \( \{u, v\} \) in \( G' \) is also a largest clique containing \( \{u, v\} \) in the sub-network of \( G' \) induced by \( N(u, v) \cup \{u, v\} \). Moreover, if we denote by \( \mathcal{C} \) the largest clique in the sub-network of \( G' \) induced by \( N(u, v) \), then \( \mathcal{C} \cup \{u, v\} \) is the clique we are looking for. We observed that,
in real-world complex networks, the sub-networks induced by $N(u, v)$ for all links $\{u, v\}$ are in general very dense and very small, which is due to the high clustering and to the power law degree distribution, respectively. This makes it possible to compute the largest clique containing $\{u, v\}$ very efficiently. For more details, see [17].

This decomposition scheme gives a bipartite version of any classical network. It is designed to produce (artificial) bipartite networks with the same kind of properties as the natural ones, in order to make them suitable for the modelling issues below. To this respect, it is important that we pick only one clique for each link, since we want the number of top nodes to be in the same order of magnitude as the size of the original network (if we take for example all the maximal cliques but not just one then we can obtain an exponential number of top nodes). Likewise, it is crucial to choose large cliques whenever we can since in real-world bipartite networks, a non-trivial number of large cliques appear. This is why we chose to take maximal cliques. As shown in Figure 2, the degree distributions of the obtained artificial bipartite networks are very similar to the distributions of the natural bipartite ones: bottom degree distributions fit very well power laws, while top degree distributions are of two kinds, either Poisson shaped or heavy tailed.

This last point indicates that the distribution of cliques sizes (i.e. the top degree distributions) may vary qualitatively between networks. This is true both for natural bipartite networks and for the ones obtained using our decomposition scheme.

Figure 2: Top and bottom degree distributions for the natural bipartite versions of Actors, Cooccurrence, and Coauthoring, and for the bipartite version of Internet, Web, and Proteins obtained with the decomposition scheme.

Since all these networks have similar underlying bipartite structure, one may wonder if their properties (like clustering) can be viewed as consequences of this structure. We
explore this idea below.

**Two bipartite models.**

The first natural way to derive a model from our observations is to sample uniformly random bipartite networks with prescribed (top and bottom) degree distributions. As explained in [26], this can be achieved as follows: first generate both top and bottom nodes and assign to each a degree drawn from the given distributions\(^1\), then create for each node as many connection-points as its degree, and finally link top and bottom connection-points randomly.

The properties of this random bipartite model are studied in [17, 26], where it is shown, both analytically and empirically, that the obtained networks have the desired properties (mainly high clustering, and power law degree distribution). Therefore, these properties can be viewed as consequences of the underlying bipartite structure.

Notice that the degree of a bottom node is *not* its degree in the unipartite version of the bipartite network (see for instance Figure 1), but the number of cliques (top nodes) it belongs to (it is linked to). Therefore, it is not trivial that the random bipartite model captures well the original degree distribution. Likewise, the high clustering of the obtained network may be seen as a consequence of the fact that there are cliques of exactly the same sizes (distributed according to the top degree distribution), and it is indeed true. However, it was not clear until now that the presence of cliques is sufficient to capture this, and, more important, no method was known to sample realistic networks with both high clustering and power law degree distribution.

The random bipartite model assumes that two distributions, for both top and bottom degrees, are explicitly given. We can also use the preferential attachment principle to define them implicitly. Indeed, as already noticed, the bottom degree distributions generally follow a power law. This leads to a growing model defined as follows. At each step, a new top node is added and its degree \(d\) is sampled from a prescribed (top) distribution (which qualitatively varies between networks). Then, for each of the \(d\) links of the new top node, either a new bottom node is added (with probability \(\lambda\)) or we pick one among the preexisting ones using preferential attachment (with probability \(1-\lambda\)). The parameter \(\lambda\) is the *overlap ratio*, defined as 1 minus the ratio between the number of nodes and the sum of the sizes of the cliques: \(\lambda = 1 - \frac{n}{\sum_{v \in \tilde{T}} d^\theta(v)}\). We use it to compute the ratio of new bottom nodes to which a new top node is connected, which ensures that the obtained network has this overlap ratio. It can be computed both on real-world bipartite networks and on ones obtained using the decomposition scheme, which gives values greater than 0.5 in general (see Table 1).

\(^1\)Notice that this may lead to incompatible top and bottom degree sequences, in the sense that the sum of top degrees is different from the sum of bottom degrees. In such a case, one can simply solve the problem by sampling again the degree of a randomly chosen top and bottom node, until the distributions are compatible.
By definition, at each step the bipartite network has the required degree distributions and the same overlap ratio as the original network. Notice that this construction process is very similar to the one observed in some real-world cases. For instance, *Actors* is built exactly this way: when a new movie is produced (which corresponds to the addition of a top node), it is linked to actors according to their popularity, and to some new actors, playing in a movie for the first time.

The degree distribution and the clustering of the networks obtained with both models can be compared with the actual networks’ degree distribution and clustering, and with the performances of the main models currently used. This is summarized in Table 1 (clustering and other) and in Figure 3 (degree distributions). In several cases, the simulations fit the real-world values surprisingly well. In all the cases, our models give results much more accurate than the commonly used models, considering all the features together.

There are however differences between the values obtained from the bipartite models and the real-world networks. They are consequences of the following fact. In the original bipartite networks (both natural ones and the ones obtained from the decomposition), many top nodes have a large neighborhood intersection. In other words, the overlap between cliques is large, and more precisely, if two cliques have one node in common then they certainly have many. This behavior can be viewed as a kind of *bipartite clustering*, which is not captured by the bipartite models: the random linking implies that most cliques have only one node in common, if any. This is responsible both for the inaccuracy of the models concerning some clusterings and for the irregularities one can observe on some distributions.

Figure 3: The original degree distribution of our six examples, together with the ones of the unipartite networks obtained from the random bipartite model and with the preferential attachment bipartite model.

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<table>
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<tr>
<th></th>
<th>Internet</th>
<th>Web</th>
<th>Actors</th>
<th>Co-auth</th>
<th>Co-occur</th>
<th>Protein</th>
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<td>1090108</td>
<td>15038083</td>
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<tr>
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<td>2.5</td>
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<td>2.4</td>
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<td>269236</td>
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<td>19885</td>
<td>13588</td>
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<td>0.643</td>
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<tr>
<td>$C$</td>
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<td>0.466</td>
<td>0.785</td>
<td>0.638</td>
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<td>0.0002</td>
<td>0.009</td>
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<td>0.0005</td>
<td>0.0015</td>
<td>0.003</td>
<td>0.028</td>
<td>0</td>
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<tr>
<td>$C_{ws}$</td>
<td>0.171</td>
<td>0.461</td>
<td>0.74 (*)</td>
<td>0.523 (*)</td>
<td>0.74 (*)</td>
<td>0.06 (*)</td>
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<td>$C_{rb}$</td>
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<td>0.663</td>
<td>0.767</td>
<td>0.542</td>
<td>0.831</td>
<td>0.187</td>
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<tr>
<td>$C_{gb}$</td>
<td>0.65</td>
<td>0.708</td>
<td>0.793</td>
<td>0.632</td>
<td>0.768</td>
<td>0.244</td>
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<tr>
<td>$d$</td>
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<td>3.98</td>
<td>2.6</td>
<td>5.45</td>
</tr>
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</table>

Table 1: The main statistics for the complex networks we use in this paper. For each network, we give its number of vertices $n$, its number of links $m$, the value of the exponent $\alpha$ of its power law degree distribution, its clustering coefficient $C$, and its average distance $d$. We also give the number of cliques $nc$ in the corresponding decomposition, as well as the overlap ratio $\lambda$. Moreover, we give the values of these parameters for typical graphs with the same number of vertices and links obtained with commonly used models and the bipartite models: the purely random model [13] ($C_{rd}$ and $d_{rd}$), the random model with prescribed degree distribution [19, 20] ($C_{dd}$ and $d_{dd}$), the Albert and Barabási model [3] ($C_{ab}$ and $d_{ab}$), Watts and Strogatz model [31] ($C_{ws}$ and $d_{ws}$), the random bipartite model with prescribed degree distributions ($C_{rb}$ and $d_{rb}$), and the growing one with preferential attachment ($C_{gb}$ and $d_{gb}$). Notice that purely random graph and Watts and Strogatz ones have a Poisson degree distribution, which makes the $\alpha$ exponent irrelevant in these cases, and makes them irrelevant for realistic modelling of complex networks. Moreover, in the cases pointed by a star (*), the real clustering coefficient is too large to be obtained with the Watts and Strogatz model. Therefore we used in these cases the parameters inducing the maximal clustering, which yields very large average distances.
For example, in the case of Internet, we noticed the presence of a sub-network of only 94 nodes which contains all the 494 cliques of size 14 and more. This makes this sub-network very dense. On the other hand, in the unipartite versions of random bipartite networks, these large cliques are disseminated all over the network which brings two artifacts: there are a lot of nodes having a degree between 14 and 29 which explains the bump on degree distribution (a similar phenomenon can be observed on Web), and the clustering is drastically increased.

Finally, let us insist on the fact that our aim is to produce networks fitting qualitatively the properties of the original ones, which is achieved in all the cases. Moreover, the results we obtain are much more accurate than the ones obtained with commonly used models.

**Conclusion.**

In this communication, we proposed a scheme to decompose any complex network into a bipartite one. This allowed us to study and compare the statistics of both natural and artificial bipartite networks, which revealed surprising similarities. In particular, the bottom degree follows a power law, and there are large cliques in real-world networks.

The properties of various complex networks, like clustering and degree distributions, are also properties of unipartite versions of random bipartite networks. Therefore, they can be seen as consequences of a hidden underlying bipartite structure.

This leads us to propose both the random bipartite model and the growing bipartite model with preferential attachment as two general models for complex networks. The choice between these two models depends on one’s aim. For instance, the second should be preferred if the growth of the network is a key issue, whereas the first should be used in situations where one knows exactly the desired degree distributions.

The obtained networks fit very well the properties of real-world networks, using only their general bipartite structure. The bipartite models therefore provide an interesting alternative to those previously proposed. Of course, real-world networks have other properties not captured by the bipartite models (we discussed the case of the overlap between cliques). There is still a lot to do in this direction, and certainly extensions of these models can be defined to improve their accuracy (we currently work in this direction).

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**References**


