Abstract

Many real-world complex networks actually have a bipartite nature: their nodes may be separated into two classes, the links being between nodes of different classes only. Despite this, and despite the fact that many ad-hoc tools have been designed for the study of special cases, very few exist to analyse (describe, extract relevant information) such networks in a systematic way. We propose here an extension of the most basic notions used nowadays to analyse classical complex networks to the bipartite case. To achieve this, we introduce a set of simple statistics, which we discuss by comparing their values on a representative set of real-world networks and on their random versions.

INTRODUCTION

A bipartite graph is a triplet $G = (T, \bot, E)$ where $T$ is the set of top nodes, $\bot$ is the set of bottom nodes, and $E \subseteq T \times \bot$ is the set of links. The difference with classical graphs lies in the fact that the nodes are in two disjoint sets, and that the links always are between a node of one set and a node of the other. In other words, there cannot be any link between two nodes in the same set.

Many real-world complex networks may be modeled naturally by a bipartite graph. Let us cite for example the actors-movies network, where each actor is linked to the movies he/she played in [70], [51], authoring networks, where the authors are linked to the paper they signed [53], [54], occurrence networks, where the words occurring in a book are linked to the sentences of the book they appear in [30], company board networks, where the board members are linked to the companies they lead [61], [19], [7], and peer-to-peer exchange networks in which peers are linked to the data they provide/search [31], [68], [35], [34]. These networks are often called affiliation networks, or two-mode networks.

Although there is nowadays a significant amount of notions and tools to analyse classical networks, there is still a lack of such results fitting the needs for analysing affiliation networks. In such cases, one generally has to transform the affiliation network into a classical one and/or to introduce ad-hoc notions. In the first case, there is an important loss of information, as well as other problems that we detail below (Section III). In the second case, there is a lack of rigor and generality, which makes the relevance of the obtained results difficult to evaluate.

The aim of this contribution is to provide a set of simple statistics which will make it possible and easy to analyse real-world affiliation networks (or at least make the first step towards this goal) while keeping their bipartite nature.

To achieve this, we will first present an overview of the basic notions and methodologies used in the analysis of classical (as opposed to affiliation) networks. We will then show how people usually transform
bipartite networks into classical networks in order to be able to analyse them with the tools designed for this case. This will lead us to a description of the state of the art, then of the methodology used in this paper. Finally, we will present and evaluate the statistics we propose for the analysis of affiliation networks.

Before entering in the core of this contribution, let us notice that we only deal here with simple\footnote{This means that we do not allow loops (links from a node to itself) nor multiple links between two given nodes. This is classical in complex networks studies: loops are managed separately, if some occur, and multiple links are generally encoded as link weights, or simply ignored.} undirected, unweighted, static networks. Considering directed, weighted, and/or dynamic networks is out of the scope of this paper; we will discuss this further in the conclusion. Moreover, in all the cases we will consider here (and in most real-world cases), the graph has a giant connected component, \textit{i.e.} there exists a path in the graph from almost any node to any other. In the following, we will make our statistics on the whole graph everywhere this makes sense, but we will restrict ourselves to the largest connected component where this is necessary (namely for distance computations). Again, this is classical in the literature and has no significant impact on our results.

I. Classical notions.

Let us consider a (classical) graph $G = (V, E)$, where $V$ is the set of nodes and $E \subseteq V \times V$ is the set of links. We will denote by $N(v) = \{u \in V, \ (u, v) \in E\}$ the \textit{neighbourhood} of a node $v$, the elements of $N(v)$ being the \textit{neighbours} of $v$. The number of nodes in $N(v)$ is the \textit{degree} of $v$: $d^o(v) = |N(v)|$.

The most basic statistics describing such a graph are its size $n = |V|$, its number of links $m = |E|$, and its average degree $k = \frac{2m}{n}$. Its density $\delta(G) = \frac{2m}{n(n-1)}$, \textit{i.e.} the number of existing links divided by the number of possible links, also is an important notion. It is nothing but the probability that two randomly chosen (distinct) nodes are linked together.

Going further, one may define the distance between two nodes in the graph as the minimal number of links one has to follow to go from one node to the other. Note that this only make sense if there is a path between the two nodes, \textit{i.e.} if they are in the same connected component. As explained above, in all the paper, we will only consider distances between the nodes in the largest connected component (and we will give its size). Then, the average distance of the graph, $d(G)$, is nothing but the average of the distances for all pairs of nodes in the largest connected component.

The statistics described above are the ones we will call the \textit{basic} statistics. The next one is not so classical. It is the degree distribution, \textit{i.e.} for all integer $i$ the fraction $p_i$ of nodes of degree $i$. In other words, it is the probability that a randomly chosen node has degree $i$. One may also observe the correlations between degrees, defined as the average degree of the neighbours of nodes of degree $i$, for each integer $i$. Other notions concerning degrees have been studied, like assortativity \cite{55} for instance, but we do not detail this here.

The last kind of statistics we will discuss here aims at capturing a notion of overlap: it measures the probability that two nodes are linked together, provided they have a neighbour in common. In other words, it is the probability that any two neighbours of any node are linked together. This may be done using two slightly different notions, both called \textit{clustering coefficient}, among which there often is a confusion in the literature\footnote{Some authors make a difference by calling the first notion \textit{clustering coefficient} and the second one \textit{transitivity ratio}, but we prefer to follow the most classical conventions of complex network studies here.}. Both will be useful in the following therefore we discuss them precisely here.

The first one computes the probability, for any given node chosen at random, that two neighbours of this node are linked together. It therefore relies on the notion of clustering coefficient for any node $v$ of degree at least 2, defined by:

$$cc_o(v) = \frac{|E_{N(v)}|}{\frac{|N(v)||N(v)|-1}{2}} = \frac{2|E_{N(v)}|}{d^o(v)(d^o(v)-1)}$$
where $E_{N(v)} = E \cap (N(v) \times N(v))$ is the set of links between neighbours of $v$. In other words, $cc_\bullet (v)$ is the probability that two neighbours of $v$ are linked together. Notice that it is nothing but the density of the neighbourhood of $v$, and in this sense it captures the local density. The clustering coefficient of the graph itself is the average of this value for all the nodes:

$$cc_\bullet (G) = \frac{\sum_{v \in V} cc_\bullet (v)}{|\{v \in V, \ \delta^0 (v) \geq 2\}|}.$$ 

One may define directly another notion of clustering coefficient of $G$ as a whole as follows:

$$cc_\vee (G) = \frac{3N_\Delta}{N_\vee}$$

where $N_\Delta$ denotes the number of triangles, i.e. sets of three nodes with three links in $G$, and $N_\vee$ denotes the number of connected triples, i.e. sets of three nodes with at least two links, in $G$. This notion of clustering is slightly different from the previous one since it gives the probability, when one chooses two links with one extremity in common, that the two other extremities are linked together.

Both notions have their own drawbacks and advantages. The first one has the advantage of giving a value for each node, which makes it possible to observe the distribution of this value and the correlations between this value and the degree, for instance. It however has the drawback of reducing the role of high degree nodes. Moreover, importantly, these definitions capture slightly different notions, which may both be relevant depending on the context. We will therefore use both notions in the following. This is why we introduced two different notations, namely $cc_\bullet$ and $cc_\vee$, which emphasises the fact that one is centered on nodes and the other is centered on pairs of links with one extremity in common.

One may consider many other statistics to describe complex networks. Let us cite for instance centrality measures, various decompositions, and notions capturing the ability of each node to spread information in the network. See [69], [2], [56], [14], [16] for surveys from different perspectives. We will not consider here such statistics. Instead, we will focus on the most simple ones, described above, because they play a central role in recent complex network studies as we will explain in the next section.

II. CLASSICAL COMPLEX NETWORKS.

Many complex networks have been studied in the literature, ranging from technological networks (power grids, internet) to social ones (collaboration networks, economical relations), or from biological ones (protein interactions, brain topology) to linguistic ones (cooccurrence networks, synonymy networks). See [69], [2], [56], [14], [16] and references therein for detailed examples. It appeared recently [70], [2], [56], [14] that most of these large real-world complex networks have several nontrivial properties in common. This section is devoted to an overview and a discussion of these properties (based on the definitions given in previous section), on which the rest of the paper will rely. We will use the same notations as in Section I.

We are concerned here with large complex networks, which means that $n$ is large. In most real-world cases, it appeared that $m$ is of the same order of magnitude as $n$, i.e. the average degree $k$ is small compared to $n$. Therefore, the density generally is very small: $\delta (G) = \frac{kn}{n(n-1)} \sim \frac{k}{n}$, which is close to 0 since $n$ is much larger than $k$ in general. We will always suppose we are in this case in the following.

It is now a well known fact that the average distance in real-world complex networks is in general very small (small-world effect), even in very large ones, see for instance [45], [70]. This is actually true in most graphs, since a small amount of randomness is sufficient to ensure this, see for instance [70], [41], [42], [9], [24]. This property, though it may have important consequences and should be taken into account, should therefore not be considered as a significant property of a given network (see Section V).

Another issue which received recently much attention, see for instance [26], [4], is the fact that the degree distribution of most real-world complex networks is highly heterogeneous, often well fitted by a power law: $p_k \sim k^{-\alpha}$ for an exponent $\alpha$ generally between 2 and 3.5. This means that, despite most
nodes have a low degree, there exists nodes with a very high degree. This implies in general that the average degree is not a significant property, bringing much less information than the exponent $\alpha$ which is a measurement of the heterogeneity of degrees.

If one samples a random network with the same size (i.e. as many nodes and links) as a given real-world one, thus with the same density, then the obtained degree distribution is qualitatively different: it follows a Poisson law. This means that the heterogeneous degree distribution is not a trivial property, in the sense that it makes real-world complex networks very different from most graphs (of which a random graph is typical). The degree correlations and other properties on degrees, however, behave differently depending on the complex network under concern.

Going further, the clustering coefficients (according to both definitions) are quite large in most real-world complex networks: despite most pairs of nodes are not linked together (the density is very low), if two nodes have a neighbour in common then they are linked together with a probability significantly higher than 0 (the local density is high). However, the clustering coefficient distributions, their correlations with degrees, and other properties related to clustering, behave differently depending on the complex network under concern.

If, as above, one samples a random graph with the same size as an original one then the two definitions of clustering coefficients are equivalent and equal to the density. The clustering coefficients therefore are very low in this case. If one samples a random graph with the same number of nodes and the very same degree distribution then the clustering coefficients still are very small, close to 0 [56]. Clustering coefficients therefore capture a property of networks which is not a trivial consequence of their degree distribution.

Finally, it was observed that the vast majority of large real-world complex networks have a very low density, a small average distance, a highly heterogeneous degree distribution and high clustering coefficients. These two last properties make them very different from random graphs (both purely random and random with prescribed degree distribution). More subtle properties may be studied, but until now no other one appeared to be a general feature of most real-world complex network. The properties described here therefore serve as a basis for the analysis of real-world complex networks, and so we will focus on them in the following. Our aim will be to define and discuss their equivalent for affiliation networks / bipartite graphs.

### III. Projection.

Let us now consider a large affiliation network modeled as a bipartite graph $G = (\top, \bot, E)$. The $\bot$-projection of $G$ is the graph $G_\bot = (\bot, E_\bot)$ in which two nodes (of $\bot$) are linked together if they have at least one neighbour in common (in $\top$) in $G$: $E_\bot = \{ (u, v), \exists x \in \top : (u, x) \in E \text{ and } (v, x) \in E \}$. The $\top$-projection $G_\top$ is defined dually. See Figure 1 for an example.

![Fig. 1. An example of bipartite graph (center), together with its $\top$-projection (left) and its $\bot$-projection (right).](image)

In order to be able to use the many notions defined on classical networks, and to compare a particular network to others, one generally transforms an affiliation network into its $\bot$-projection, often called the

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6We consider here a network chosen uniformly at random among the ones having this size, using typically the Erdős and Rényi model [9], [24].

7We consider here a network chosen uniformly at random among the ones having this number of nodes and this degree distribution, using typically the configuration model [8], [9], [46], [47], [67].
one-mode version of the network. This was typically done for the affiliation networks we presented in the introduction: the actors-movies network is transformed into its \( \perp \)-projection where two actors are linked together if they acted together in a movie [70]; the authoring networks are transformed into their \( \perp \)-projections, \textit{i.e.} coauthoring networks where two authors are linked together if they signed a paper together [53], [54], [51]; the occurrence networks are transformed into their \( \perp \)-projections, \textit{i.e.} cooccurrence networks where two words are linked if they appear in the same sentence [30]; the company board networks are transformed into their \( \perp \)-projections where two persons are linked together if they are member of a same board [61], [19], [7]; and the peer-to-peer exchange networks are transformed into their \( \perp \)-projections where two data are linked together if they are provided/searched by a same peer [31], [68], [35], [34].

This approach is of course relevant since the projections under study make sense, and also encode relevant information. Moreover, this allows the study of affiliation networks using the powerful tools and notions provided for classical networks. We however argue that in most cases there would be a significant gain in considering the bipartite version of the data. The main reasons are as follows.

- Most importantly, there is much information in the bipartite structure which may disappear after projection. For instance, the fact that two actors played in many movies together, and the size of these movies, brings much information which in not available in the projection, in which they are simply linked together. This loss of information is particularly clear when one notices that there are many bipartite graphs which lead to the same projection (while each bipartite graph has only one \( \top \)- and one \( \perp \)-projection), see [37], [36]. The fact that much important information is encoded in the bipartite structure is a central point which we will illustrate all along this paper.

- Notice that each top node of degree \( d \) induces \( \frac{d(d-1)}{2} \) links in the \( \perp \)-projection, and conversely. This induces an inflation of the number of links when one goes from a bipartite graph to its projection, see Table I. In our examples, this is particularly true for peer-to-peer: the number of links reaches more than 10 billions in the \( \perp \)-projection, which needs more than 80 GigaBytes of central memory to be stored using classical (compact) encodings (while the original affiliation network needs less than 500 MegaBytes). This is a typical case in which the huge number of links induced by the projection is responsible for limitations on the computations we are able to handle on the graph in practice.

|                        | actors-movies | authoring | occurrences | peer-to-peer   |
|------------------------|---------------|-----------|-------------|----------------|
| Number of links in \( G \) | 1,470,418     | 45,904    | 183,363     | 55,829,392     |
| Number of links in \( G_\perp \) | 15,038,083   | 29,552    | 392,066     | 10,142,780,673 |
| Number of links in \( G_\top \) | 20,490,112   | 134,492   | 51,405,275  | 1,085,217,140  |

\textbf{TABLE I} \textbf{NUMBER OF LINKS IN AFFILIATION NETWORKS AND THEIR PROJECTIONS, FOR THE FOUR EXAMPLES WE WILL DESCRIBE IN SECTION V.}

- Finally, some properties of the projection may be due to the projection process rather than the underlying data itself. For instance, it is shown in [51], [37], [36] that when considering the projection of a random bipartite graph, one observes high clustering coefficients. Therefore, high clustering coefficients in projections may not be viewed as significant properties: they are consequences of the bipartite nature of the underlying affiliation network. Likewise, the projection may lead to very dense networks, even if the bipartite version is not dense; this is particularly the case here for the \( \top \)-projection of occurrences.

One way to avoid some of these problems is to use a \textit{weighted} projection. For instance, the weight of a link \((u, v)\) between two bottom nodes in the weighted \( \perp \)-projection may be defined as the number of (top) neighbours \( u \) and \( v \) have in common in the bipartite graph. Other definitions may be considered: each top node may contribute to each link it induces in the \( \perp \)-projection in a way that decreases with its degree, for instance. In all cases, and despite such an approach is relevant and promising, one still looses a significant amount of information, and one transforms the problem of analysing a bipartite structure
into the problem of analysing a weighted one, which is not easier. Indeed, despite the fact that important progress has recently been done in this direction [5], [6], [49], much remains to be done to be able to analyse precisely the structure of weighted networks.

Our aim is this paper is to provide an alternative to the projection approach, leading to a better understanding of affiliation networks. It must however be clear that (weighted) projection approaches also lead to significant insight, and we consider that the two approaches should be used as complementary means to understand in details the properties of affiliation networks.

IV. STATE OF THE ART.

Affiliation networks have been studied in an amazingly wide variety of context. Let us cite for instance company boards [61], [19], [7], [51], sport teams [10], [58], movie actors [70], [51], human sexual relations [25], [44], attendance to political events [29], financial networks [18], [20], [32], [71], recommendation networks [60], theatre performances [1], [64], politic ativism [5], [61], [44], student course registrations [38], word cooccurrences [22], [65], file sharing [39], [31], [68], [35], [34], and scientific authoring [62], [48], [53], [54], [50].

These studies are made in disciplines as various as social sciences, computer science, linguistics and physics, which makes the literature very rich. In all these contexts, scientists face affiliation networks which they try to analyse, with various motivations and tools. They all have one feature in common: they insist on the fact that the bipartite nature of their data plays an important role, and should be taken into account. They also emphasise the lack of notions and tools for doing so.

Because of this lack of relevant notions and tools, most authors have no choice but to consider the most relevant projection of their affiliation network. This leads for instance to studies of interlocks between companies, see for instance [61], [19], studies of coauthoring networks, see for instance [53], [54], [50], or studies of exchanges between peers in peer-to-peer systems, see for instance [31], [68], [35], [34].

Many authors realise that this approach is not sufficient, and try to use the bipartite nature of their data. This is generally done by combining the use of projections and the use of basic bipartite statistics, mostly degrees. For instance, one studies the coauthoring relations (typically a projection) and the distributions of the number of papers signed by authors and of the number of authors of papers (i.e. the bipartite degree distributions, see Section IV) [50]. Authors may also consider weighted projections, see for instance [7], [48], [34], [35], [39], [50], which has advantages and drawbacks, as discussed above, Section III.

Going further, some authors introduce bipartite notions designed for the case under study. This is often implicit and restricted to very basic properties, like the case of degree distributions cited above (which essentially capture the size of events and the number of events in which persons or objects are involved, in most cases). But some authors introduce more subtle notions, like notions of overlap [10], clustering [13], [61], [44], centrality measures [28], degree correlations [59], and others [71], [25], [18], [60], [39], [13], [61], [44]. Most of these notions are ad hoc and specific to the case under study, but some of them actually are very general or may be generalised. One of our central aims here is to give a complete and unified framework for the most general of these notions. We will cite appropriate references when the notions we will discuss have already been considered previously.

As already said, a different and interesting approach is developed in [51], [37], [36]. The authors study the expected properties of the projections given the properties (namely the degree distributions) of the underlying bipartite graph. They show in particular that the expected clustering coefficient in the projections is large, and give an efficient estimation formula; this means that a high clustering coefficient in a projection may be seen as a consequence of the underlying bipartite structure rather than a specific property of the network. Conversely, if the clustering coefficient of the projection is different from the expected one, it means that the underlying bipartite structure has nontrivial properties responsible for it. These properties should therefore be further analysed. Our aim here is to propose notions and tools for such an analysis. This approach has been used with profit in several cases, for instance [51], [52], [19], [64].
Finally, a significant effort has already been made to achieve the goal we have here, or similar goals: some studies propose general approaches for the analysis of affiliation networks. This is for instance the case of [28], focused on centrality measures, of [17], which proposes to consider both projections and compare them, and of [10], which studies in depth the notion of overlap.

Let us cite in particular [13], which has the very same aim as we have here, but makes quite different choices. For instance, the authors deal with visualisation, whereas we do not here. On the other hand, they consider rather small networks, while we are particularly interested in large ones. They use a matrix approach, which is very powerful but not suited for large cases. This also leads them to consider some properties, like (betweenness) centrality for instance, which can hardly be computed on very large networks. Finally, they do not use the comparison with random graphs, central to our contribution (see Section V), which probably reflects the fact that this method was not as usual in 1997 as it is now. It is interesting to see that, although the initially claimed aim is very similar, the final contributions are very different.

Other researchers propose formalisms suited for the analysis of affiliation networks, often based on a generalisation of well known models. Let us cite Galois lattices [62], correspondence analysis [40], [27], extensions of blockmodels [12], [23] and p* models [63], [29], [1] and a particularly original approach based on boolean algebra [11].

Therefore, there already exists quite an impressive amount of work on affiliation networks, and on methods for their analysis. However, we observe that many of the approaches proposed previously, though very relevant, are hardly applicable to large networks, typically networks with several hundreds of thousands nodes. Moreover, they often rely on quite complex notions and formalisms, which are difficult to handle for people only interested in analysing a given network. Finally, none of them consists in a generalisation of the notions nowadays widely used to analyse classical (as opposed to affiliation) networks, outlined in Section I.

We propose here such a contribution. We design simple notions and methods to analyse very large affiliation networks, which could be used as a first step in particular studies. These methods may then be extended to fit the details of particular cases, and we explain how to do so. Moreover, they are not only extensions of classical notions; we go further by proposing new notions designed specifically for the bipartite case. Our approach may also be applied to smaller networks, as long as they are not too small (typically thousands of nodes).

As explained above, the topic has a deep interdisciplinary nature. In order to make our techniques usable by a wide audience, we give a didactic presentation and we focus on basic notions. Let us insist however on the fact that this presentation is rigorous and formal, and, as will appear all along the paper, the results are sufficient to bring a significant amount of information on a given network.

Finally, we insist on the fact that analysing properly and in details a given network is a difficult task, which may be handled using different methods. There is no unique way to obtain relevant information and results in such cases. Moreover, much resides in the interpretations made from the outputs of these approaches. All the ones we have cited above, and the one we propose here, should therefore be seen as complementary rather than concurrent.

Let us conclude this section by noticing that, because of the wide dispersion of contributions due to the interdisciplinary nature of the topic (and the fact that it received continuous attention since several decades), we certainly missed some references. We however expect that the ones we have cited span well the contributions on the topic.

V. METHODOLOGY AND DATA.

The methodology we follow to develop tools for the analysis of complex networks of various kinds, mainly used since the end of the 90’s, is the one sketched in Section I. It relies on the introduction of statistical parameters aimed at capturing a given feature of networks under concern, an then on the comparison of the behaviours of real-world networks concerning these parameters as compared to random
ones. The underlying principle is that a parameter which behaves similarly on real-world and random networks is just a property of most networks (of which random networks are representatives) and so, though it may play an important role, it should not be considered as surprising and meaningful concerning the description of the real-world network. Instead, one generally looks for properties which make real-world networks different from most networks.

Our contribution here relies on this methodology. Namely, we will define statistical parameters aimed at capturing properties of bipartite graphs, and then evaluate the relevance of these parameters by comparing their values on random bipartite graphs and on real-world affiliation networks.

Just like one considers purely random graphs and random graphs with prescribed degree distributions in the case of classical networks, we will use both purely random bipartite graphs and random bipartite graphs with prescribed degree distributions. Such graphs are constructed easily by extending the classical case, see for instance [51], [37]. We provide a program generating such graphs at [33]. Note that these models (both the classical and bipartite versions) generate graphs that are not necessarily simple: they may contain some loops and multiple links. There are however very few such links, and simply removing them generally has no impact on the results. This is what is generally done in the literature, and we will follow this convention here: in our context, it cannot have a significant impact.

Notice also that the properties of random graphs may be formally evaluated, see for instance [51], [36]. We will however focus on practical aspects here and leave these evaluations for further work, see Section X.

In order to complete our comparison between random and real-world cases, we also need a set of real-world affiliation networks. We chose the following four instances, which correspond to the examples given in the introduction and have the advantage of spanning well the variety of cases met in practice:

- the actors-movies network as obtained from the Internet Movie Data Base [21] in 2005, concerning $n_{\bot} = 127,823$ actors and $n_{\top} = 383,640$ movies, with $m = 1,470,418$ links;
- an authoring network obtained from the online arXiv preprint repository [3], with $n_{\top} = 19,885$ papers, $n_{\bot} = 16,400$ authors, and $m = 45,904$ links;
- an occurrence graph obtained from a version of the Bible [66] which contains $n_{\bot} = 9,264$ words and $n_{\top} = 13,587$ sentences with $m = 183,363$ links;
- a peer-to-peer exchange network obtained by registering all the exchanges processed by a large server during 48 hours [35], [34], leading to $n_{\top} = 1,986,588$ peers, $n_{\bot} = 5,380,546$ data, and $m = 55,829,392$ links;

We provide these data, together with the programs computing the statistics described in this paper, at [43]. The key point here is that this dataset spans quite well the variety of context in which large affiliation networks appear, as well as the variety of data sizes.

Let us insist on the fact that our aim here is not to derive conclusions on these particular networks: we only use them as real-world instances to illustrate the use of our results and to discuss their generality. This is why we do not detail more the way they are gathered and their relevance to any study. This is discussed in the corresponding references and is out of the scope of this paper.

VI. BASIC BIPARTITE STATISTICS.

The basic statistics on bipartite graphs are direct extensions of the ones on classical graphs. One just has to be careful with the fact that some classical properties give birth to twin bipartite properties while others must be redefined.

Let us consider a bipartite graph $G = (\top, \bot, E)$. We denote by $n_{\top} = |\top|$ and $n_{\bot} = |\bot|$ the numbers of top and bottom nodes, respectively. We denote by $m = |E|$ the number of links in the network. This leads to a top average degree $k_{\top} = \frac{m}{n_{\top}}$ and a bottom one $k_{\bot} = \frac{m}{n_{\bot}}$. One may obtain the average degree in the

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8One may also use the methods described in [67] to obtain directly simple (connected) graphs, but this is more intricate, and unnecessary in our context.
graph \( G' = (\top \cup \bot, E) \) as \( k = \frac{2m}{n^R_n \bot + n^\bot_n \top} = \frac{n^\top n\bot k^\top + n^\bot n\top k^\bot}{n^\top n\bot} \). Finally, we obtain the bipartite density \( \delta(G) = \frac{m}{n^\top n\bot} \), i.e. the fraction of existing links with respect to possible ones. Note that this is different from the density of \( G' \): \( \delta(G') = \frac{2m}{(n^\top + n\bot)(n^\top + n\bot - 1)} \), which is much lower.

Concerning the average distance (again, we restrict distance computations\(^5\) to the largest connected component, which contains the vast majority of nodes, see Table II), there is no crucial difference except that one may be interested by the average distance between top nodes and between bottom nodes, \( d^\top \) and \( d^\bot \). These values may be significantly different but one may expect that they are very close since a path between two top (resp. bottom) nodes is nothing but a path between bottom (resp. top) nodes with two additional links. Notice that there is no simple way to derive the average distance \( d \) in \( G' \) from the bipartite statistics \( d^\bot \) and \( d^\top \).

The values obtained for each of these basic properties on our four examples, together with values obtained for random bipartite networks with the same size, are given in Table II. It appears clearly that our examples may be considered as large networks with small average degrees, compared to their size. The density therefore is small. Moreover, the average distance is also small. These basic properties are very similar to what is observed on classical networks: both classical and affiliation large real-world complex networks are sparse and have a small average distance, and in both contexts this is also true on random graphs.

### Table II

|                  | actors-movies | authoring | occurrences | peer-to-peer |
|------------------|---------------|-----------|-------------|--------------|
|                  | real random   | real random | real random | real random  |
| \( n^\top \)     | 127,823 idem | 19,885 idem | 13,587 idem | 1,986,588 idem |
| \( n^\bot \)     | 383,640 idem | 16,400 idem | 9,264 idem  | 5,380,546 idem |
| \( m \)          | 1,470,418 idem | 45,904 idem | 183,363 idem | 55,829,392 idem |
| \( k^\top \)     | 11.5 idem    | 2.3 idem   | 13.5 idem   | 28.1 idem    |
| \( k^\bot \)     | 3.8 idem    | 2.8 idem   | 19.8 idem   | 10.4 idem    |
| \( \delta \)     | 0.000030 idem | 0.00014 idem | 0.0015 idem | 0.0000052 idem |
| \( \top \) giant | 124,414 125,944 | 16,209 18,512 | 13,579 13,587 | 1,986,343 1,426,978 |
| \( \bot \) giant | 374,511 381,431 | 11,654 14,607 | 9,246 9,264 | 5,380,507 5,054,689 |
| \( d^\top \)     | 6.8 5.3     | 13.1 9.3   | 3.1 3.0     | 5.3 5.0     |
| \( d^\bot \)     | 7.3 5.8     | 13.9 9.9   | 3.8 3.7     | 5.4 4.9     |
| \( d \)          | 7.2 5.8     | 13.5 9.6   | 3.4 3.2     | 5.3 4.9     |

### VII. Bipartite statistics on degrees.

The notion of degree distribution has an immediate extension to the bipartite case. We denote by \( \bot_i \) the fraction of nodes in \( \bot \) having degree \( i \) and by \( \top_i \) the fraction of nodes in \( \top \) having degree \( i \), and then call \((\bot_i)_{i \geq 0}\) the bottom degree distribution and \((\top_i)_{i \geq 0}\) the top one.

The top and bottom degree distributions of our four examples are given in Figure 2. One may observe on these plots that the bottom degree distributions are very heterogeneous and well fitted by power laws (of various exponents). This is true in particular for the occurrences graph, which is a well known fact for a long time [72]: the frequency of occurrences of words in a text generally follows a particular kind of power law, named Zipf law. Instead, the shape of the top degree distribution depends on the case under

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\(^5\)Distance computations are expensive; the exact value cannot be computed in a reasonable amount of time for data of the size we consider here. Instead, we approximate the average by computing the average distance from a subset of the nodes to all the others, this subset being large enough to ensure that increasing it does not improve our estimation anymore, which is a classical method. All other computations are exact.
concern: whereas it is well fitted by a power law in the peer-to-peer and actors-movies cases, it is far from a power law in the authoring and occurrences cases. This is due to the fact that papers have a limited number of authors (none has one hundred authors for instance), and likewise sentences have a limited number of words. Moreover, the number of very short sentences also is not huge. In these two cases, one can hardly conclude that the top degrees are very heterogeneous.

We finally conclude that, even if heterogeneity is present on at least one side of an affiliation network, this is not generally true for both sides. This separates real-world affiliation networks into two distinct classes, which should be taken into account in practice. This also confirms that considering the bipartite statistics brings significant information as compared to the projections, which exhibit power law degree distributions in all cases.

Let us now compare these real-world statistics with random graphs. If one generates purely random bipartite graphs of the same size as the ones considered here, the (⊤ and ⊥) degree distributions are Poisson laws. Therefore, the heterogeneity of some degree distributions is not present, and even in the cases where the distributions are not very heterogeneous they do not fit the random case. We will therefore compare in the following our real-world affiliation networks to random bipartite graphs with the same size and the same (top and bottom) degree distributions.

The next natural step is to observe possible correlations between top and bottom degrees. In order to do this, we plot in Figure 3 the average degree of neighbours of nodes as a function of their degree, both for top and bottom nodes, separately. In other words, for each integer \(i\) we plot the average degree of all nodes which are neighbours of a node of degree \(i\). We plot the same values obtained for random graphs of the same size and same degree distributions.

In the cases of authors-movies and peer-to-peer, the plots for the random cases are close to horizontal lines, showing that there are no correlations between a node degree and the average degree of its neighbours: this last value is independent of the node degree. In both cases, however, the real-world network displays nontrivial correlations. In the case of actors-movies, for instance, the average degree of neighbours of bottom nodes (the lower-left corner plot in Figure 3) decreases with the node degree. In other words, if an actor plays in many movies then he/she tends to play in smaller movies (in terms of the number of involved actors). Such nontrivial observations may be made on the other plots for actors-movies and peer-to-peer as well.

In the cases of authoring and occurrences, the plots for the random graphs are nontrivial: they grow for the top statistics, and are far from smooth for the bottom ones. Here again, the real-world cases exhibit significantly different behaviours, at least for the top statistics, thus demonstrating that these behaviours are nontrivial and related to intrinsic properties of the underlying networks. Detailing this however is
such an information is not available in the authoring affiliation network. To distinguish between different behaviours. For instance, they make it possible to say if a given author to be linked to only one node \( u \) in \( \perp \), with \( N(u) = N(N(v)) \), or when \( v \) is linked to \( |N(N(v))| \) nodes in \( \perp \), each being linked to only one other node in \( \top \). Of course, intermediate cases may occur, and the actual situation may be observed by plotting the correlations between the degree of nodes \( v \), i.e. \( |N(v)| \), and their number of distance 2 neighbours, \( |N(N(N(v)))| \). These statistics therefore offer a way to study how node degrees in the projection appear, and to distinguish between different behaviours. For instance, they make it possible to say if a given author has many coauthors because he/she writes many papers or if he/she writes papers with many authors. Such an information is not available in the projection of the authoring affiliation network.

The plots in Figure 4 show that, as one may have guessed, the number of distance 2 neighbours of a node grows with its degree; more precisely, it generally grows as a power of the degree (the plots follow straight lines in log-log scale), and actually almost linearly. This is in conformance with the intuition that the number of distance 2 neighbours should be close to the degree of the node times the average degree of its neighbours. In the random cases, this leads to very straight plots (except in the top plot of occurrences). The real-world plots are quite close to the random ones, with a few notable exceptions: the slope of the plot is significantly different for the top plot of peer-to-peer, the real-world plots often are significantly below the random ones for large degrees, and they are in general slightly lower than the random ones even for small degrees. This means that there is some redundancy in the neighbourhoods: whereas in random cases the number of distance 2 neighbours is close to the sum of the degrees of the direct neighbours, in real-world cases the direct neighbours have many neighbours in common and so the number of distance 2 neighbours is significantly lower. This is an important feature of real-world complex networks, that we will deepen in the next sections.
VIII. BIPARTITE CLUSTERING AND OVERLAP.

Whereas there were quite direct extensions of the basic statistics and the ones on degrees to the bipartite case, the notion of clustering coefficient does not make any sense in itself in this context. Indeed, it relies on the enumeration of the triangles in the graphs, and there can be no triangle in a bipartite graph. We will therefore have to discuss the features captured by the classical clustering coefficients in order to propose bipartite extensions.

Both definitions of classical clustering coefficients capture the fact that when two nodes have something in common (one neighbour) then they are linked together with a probability much higher than two randomly chosen nodes. Conversely, they capture the fact that when two nodes are linked together then they probably have neighbours in common. In other words, they capture correlations between neighbourhoods. We will use this point of view here and define a first notion of clustering coefficient defined for pairs of nodes (in the same set $\top$ or $\bot$):

$$\text{cc}_\bullet(u, v) = \frac{|N(u) \cap N(v)|}{|N(u) \cup N(v)|}.$$  

This is the most direct generalisation of the classical notion, and it was already suggested in [13], and explicitly used in [35] in the context of peer-to-peer exchange analysis. It captures the overlap between neighbourhoods of nodes: if $u$ and $v$ have no neighbour in common then $\text{cc}_\bullet(u, v) = 0$. If they have the same neighbourhood, then $\text{cc}_\bullet(u, v) = 1$. And if their neighbourhoods partially overlap then the value is in between, closer to 1 when the overlap is large compared to their degrees. See Figure 5 for an illustration.

This definition however has several drawbacks. The first one is the fact that it defines a value for pairs of nodes. One may want to capture the tendency of one particular node to have its neighbourhood included in the ones of other nodes. To achieve this, one may simply define the clustering coefficient of one node as the average of its clustering coefficients with other nodes. We however do not include in this averaging the pairs for which the overlap is empty: most nodes have disjoint neighbourhood, which does not bring information. Like in the classical case, we want to measure the implication of the fact of having one neighbour in common on the rest of the neighbourhoods. We finally obtain:

$$\text{cc}_\bullet(u) = \frac{\sum_{v \in N(N(u))} \text{cc}_\bullet(u, v)}{|N(N(u))|}.$$  

One may then observe the distribution of these values, their correlations with degrees, etc. One may also

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10 As a consequence, the obtained value will never be 0, but it may be very small. Notice also that the clustering coefficient is not defined for nodes $v$ such that $N(N(v)) = \emptyset$ (recall that, by definition, $v \notin N(N(v))$).
define the clustering coefficient of the top (resp. bottom) nodes, denoted by $cc_\top(u,v)$ (resp. $cc_\bot(u,v)$) as the average of this value over top (resp. bottom) nodes. The average over the all graph, denoted by $cc_\star(G)$, can then be obtained easily: $cc_\star(G) = \frac{n_\top cc_\top + n_\bot cc_\bot}{n_\top + n_\bot}$. We will discuss the obtained values below, see Table III.

The notion of clustering coefficient discussed until now is an extension of the first classical one. It captures the fact that a node which has a neighbour in common with another node generally has a significant portion of neighbours in common with it. There is another way to capture this, similar to the second definition of classical clustering coefficient, is to measure the probability that, given four nodes with three links, they actually are connected with four links (all the possible bipartite ones):

$$cc_{\mathcal{N}}(G) = \frac{2N_\times}{N_N}$$

where $N_\times$ is the number of quadruplets of nodes with four links in $G$, and $N_N$ is the number of quadruplets of nodes with at least three. This extension of the second notion of classical clustering coefficient was already proposed in [61] in the context of company board networks. We will discuss the obtained values below, see Table III.

The two notions above generalise the classical definitions of clustering coefficients. Capturing the overlap between neighbours may however need more precision. Suppose that degrees are heterogeneous in the network, as it is often the case (Section VII), and consider two nodes $u$ and $v$. If one of these nodes has a high degree and the other has not, then $cc_\star(u,v)$ will necessarily be small. This will be true even if one of the neighbourhoods is entirely included in the other. One may however want to capture this, which can be done using the following definition:

$$cc_{\mathcal{S}}(u,v) = \frac{|N(u) \cap N(v)|}{\min (|N(u)|, |N(v)|)}.$$ 

One may define dually:

$$cc_{\mathcal{T}}(u,v) = \frac{|N(u) \cap N(v)|}{\max (|N(u)|, |N(v)|)}.$$ 

See Figure 5 for an illustration. These two notions, called min- and max-clustering, were introduced first in [35]. The first one emphasises on the fact that small neighbourhoods may intersect significantly large ones; it is equal to 1 whenever one of the neighbourhoods is included in the other. The second one emphasises on the fact that neighbourhoods (both small or large ones) may overlap very significantly: it is 1 only when the two neighbourhoods are the same and it tends to decreases rapidly if the degree of one of the involved nodes increases. It captures the fact that nodes with similar degrees have high neighbourhood overlaps.

![Fig. 5. Examples of bipartite clustering coefficients, and interpretations. Left: a case in which $cc_\star(u,v) = \frac{2}{3} = 0.333 \cdots$ is quite small, despite the fact that $u$ and $v$ have two neighbours in common, due to the fact that the union of their neighbours is quite large; on the contrary, $cc_{\mathcal{S}}(u,v) = \frac{1}{2} = 0.666 \cdots$ is quite large, revealing that one of the neighbourhoods is almost included in the other; the value of $cc_{\mathcal{T}}(u,v) = \frac{1}{3} = 0.4$ indicates that this may be due to the fact that one of the nodes has a high degree. The situation is different in the case at the center: all clustering coefficients are quite high (resp. 0.5, 0.666 \cdots, and 0.666 \cdots), indicating that there is not only an important overlap, but that this overlap concerns a significant part of each neighbourhoods (and thus the two nodes have similar degrees). On the right, the two nodes have a small clustering coefficient $cc_{\mathcal{S}}(u,v) = \frac{1}{3} = 0.25$, and the fact that the value of $cc_{\mathcal{T}}(u,v) = \frac{1}{3} = 0.4$ remains quite small indicates that this is not due to the fact that one of the two nodes has a very high degree compared to the other one. If ones considers larger degree nodes, then the difference between small and high values is clearer, but the figure would be unreadable.

With these definitions, one may define $cc_\star(v)$, $cc_\star(T)$, $cc_\star(\bot)$, $cc_\star(G)$, $cc_\star(v)$, $cc_\star(T)$, $cc_\star(\bot)$, and
cc\(_{G}(G)\) in a way similar to the one used above for cc\(_*(v)\), cc\(_*(\top)\), cc\(_*(\bot)\), and cc\(_*(G)\). The distributions and various correlations may then be observed.

|                  | actors-movies | authoring | occurrences | peer-to-peer |
|------------------|---------------|-----------|-------------|--------------|
|                  | real          | random    | real        | random       | real        | random       |
| cc\(_*(\top)\)   | 0.046         | 0.064     | 0.27        | 0.29         | 0.066       | 0.066        |
| cc\(_*(\bot)\)   | 0.20          | 0.36      | 0.25        | 0.31         | 0.038       | 0.065        |
| cc\(_N(G)\)      | 0.00024       | 0.0082    | 0.048       | 0.079        | 0.0012      | 0.053        |
| cc\(_G(\top)\)   | 0.23          | 0.24      | 0.20        | 0.56         | 0.19        | 0.61         |
| cc\(_G(\bot)\)   | 0.79          | 0.81      | 0.61        | 0.73         | 0.64        | 0.70         |
| cc\(_G(\top)\)   | 0.062         | 0.087     | 0.097       | 0.36         | 0.097       | 0.34         |
| cc\(_G(\bot)\)   | 0.21          | 0.37      | 0.041       | 0.33         | 0.069       | 0.26         |

TABLE III
BIPARTITE CLUSTERING STATISTICS ON OUR FOUR EXAMPLES AND ON RANDOM BIPARTITE GRAPHS WITH THE SAME SIZE AND SAME DEGREE DISTRIBUTIONS.

We give in Table III the values obtained for our four examples together with the values obtained for random bipartite graphs with same size and degree distributions (the values for purely random bipartite graphs are similar). It appears clearly that the notions we introduced capture different kinds of overlaps between neighbourhoods. However, except for cc\(_N(G)\), the obtained values are not very different on random graphs and on real-world networks. This indicates that these statistics do not capture a very significant feature of real-world complex networks, which will discuss this further below. Instead, the obtained values for cc\(_N(G)\) is significantly larger on real-world networks than on random graphs, which shows that it captures more relevant information.

![Cumulative distributions of clustering coefficients](image)

We show in Figure 6 the cumulative distributions of cc\(_*(v)\), cc\(_G(v)\), and cc\(_G(v)\) for our four examples, i.e. for each value x on the horizontal axis the ratio of all the nodes having a value lower than x for these statistics. Before entering in the discussion of these plots, notice that, by definition, we have cc\(_*_{(v)}\) ≤ cc\(_G_{(v)}\) ≤ cc\(_G_{(v)}\) for any v. Therefore, the lower plots in each case of Figure 6 is the one of cc\(_*_{(v)}\), the upper is the one for cc\(_*_{(v)}\) and the one for cc\(_G_{(v)}\) is in between.

More interesting, the plots exhibit quite different behaviours. In several cases (in particular top of actors-movies, occurrences and peer-to-peer, as well as bottom of occurrences and peer-to-peer) the plots for cc\(_G_{(v)}\) and cc\(_*_{(v)}\) grow very rapidly and are close to 1 almost immediately. This means that the values
of these statistics are very small, almost 0, for most nodes: in these cases, the neighbours of nodes have a small intersection, compared to the union of their neighbourhoods. However, in several cases, the plots for \( cc_v \) grow much less quickly, and remain lower than 1 for a long time. In several cases, it is even significantly lower than 1 by the end of the plot, meaning that for an important number of nodes the value of \( cc_v \) is equal to 1: almost 10% in the case of top of actors-movies, almost 20% in the cases of top authoring and bottom of peer-to-peer, and more than 40% in the case of bottom of occurrences. This means that, despite overlaps are in general small compared to their possible value, the neighbourhoods of many low-degree nodes significantly or even completely overlap with other nodes neighbours.

Other cases display a very different behaviour: in both top and bottom plots of authoring, and in bottom of actors-movies, it appears clearly that a significant number of nodes have a large value for \( cc_v \), \( cc_v \) and \( cc_v \). This means that node neighbours overlap significantly, and that this is not only a consequence of the fact that low degree nodes have their neighbourhoods included in the ones of other nodes.

![Figure 7](image.png)

**Fig. 7.** Cumulative distributions of the \( cc \) clustering coefficient in our four real-world affiliation networks, and in random bipartite graphs of the same size and same degree distributions. First row: for top nodes. Second row: for bottom nodes.

Again, our aim here is not to discuss in detail the specificities of each case, but to give evidence of the fact that these statistics have nontrivial behaviours and capture significant information. It is clear from the discussion above that the three notions of clustering captured by \( cc_v \), \( cc_v \) and \( cc_v \) are different, and give complementary insight on the underlying network properties. One may however be surprised by the fact that \( cc_v \) often is very small, which we deepen now by comparing its behaviours on real-world cases and on random ones, see Figure 711.

In these plots, it appears clearly that, except in the case of bottom of actors-movies, the plots of the real-world values and of the random ones are quite similar. This means that, concerning the values of \( cc_v \), real-world graphs are not drastically different from random ones (they however have slightly higher values of \( cc_v \) in most cases). In other words, this statistics does not capture very significant information, according to the methodology described in Section V. This is true by construction for random graphs, and the plots above show that this is mostly true for real-world networks also, which was not obvious.

Similar conclusions follow from the study of \( cc_v \), but the study of \( cc_v \) leads to the opposite conclusion: an important number of nodes have their neighbourhood included in the one of other (large degree) nodes, as already discussed, which happens much more rarely in random graphs. We do not detail

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11 For clarity and to avoid long discussions on specific behaviours, which is out of our scope here, we only compare the real-world and the random behaviours of \( cc_v \) (not of the two other notions of clustering coefficients).
these results here, since they do not fit in the scope of this paper. Instead, we will propose a new statistics in the next section that has several advantages on the clustering coefficients discussed here and does not have their drawbacks.

Before turning to this other statistics, let us observe the correlations between node degrees and their clustering coefficient. Again, for clarity and to maintain the paper within a reasonable length, we focus on $cc(v)$ and its comparison with the random case. See Figure 8.

![Fig. 8. Correlations of the $cc(v)$ clustering coefficient with node degrees in our four examples, and in random bipartite graphs with the same size and degree distributions. First row: for top nodes. Second row: for bottom nodes.](image)

The values for the random graphs are below the ones for the real-world cases (or they coincide at some points), in all plots. This shows that the value of $cc(v)$ are larger in real-world cases than in random ones, but the difference is small, which confirms the observations above. More interestingly, it appears clearly that in most cases $cc(v)$ decreases as a power of the degree of $v$ (straight line in log-log scale). In other words, the clustering coefficient of low degree nodes is quite large, but the one of large degree nodes is very small, like in random graphs.

**IX. The Notion of Redundancy.**

In the previous section, we discussed several ways to extend the classical notions of clustering coefficient to the bipartite case. One may wonder if the bipartite nature of the networks under concern may lead to new, specific notions, just like we observed concerning degrees in Section VII. Moreover, one may want to capture the notion of overlap concerning one particular node; in previous section, this was only possible by averaging the value obtained for a possibly large number of pairs of nodes. This section answers this: it is devoted to a new notion aimed at capturing overlap in bipartite networks, in a node-centered fashion.

First notice that neighbourhood overlaps correspond to links which are obtained in several ways during the projection, and that these links cannot be distinguished one from another in the projection. They also reveal the fact that, among all the links induced by a node of a bipartite graph in the projection, many (and possibly all) may actually be induced by others too. In other words, if we remove this node in the bipartite graph then the projection may be only slightly changed (or even not at all). This can be captured by the following parameter, which we call the redundancy coefficient of $v$:

$$
rc(v) = \frac{|\{\{u, w\} \subseteq N(v), \exists v' \neq v, (v', u) \in E \text{ and } (v', w) \in E\}|}{N(v)(|N(v)|-1)}.
$$

In other words, the redundancy coefficient of $v$ is the fraction of pairs of neighbours of $v$ linked to another node than $v$. In the projection, these nodes would be linked together even if $v$ were not there, see Figure 9.
this is why we call this the redundancy. If it is equal to 1 then the projection would be exactly the same without \( v \); if it is 0 it means that none of its neighbours would be linked together in the projection.

![Fig. 9. Example of redundancy computation. From left to right: a bipartite graph, its \( \perp \)-projection, and the \( \perp \)-projection obtained if the node \( A \) is first removed. Only two links disappear, leading to \( \text{rc}(A) = \frac{4}{6} = 0.666 \ldots \).]

Again, we can derive from this definition the ones of \( \text{rc}(\top) \), \( \text{rc}(\perp) \) and \( \text{rc}(G) \), as well as distributions and correlations. We give in Table IV the values obtained for our four examples and for comparable random graphs. It appears clearly from these values that, except in the case of occurrences, the redundancy coefficient is much larger in real-world networks than in random graphs, and that it actually is very large: in peer-to-peer, for instance, on average half the pairs of peers that have a common interest for a given data also have a common interest for another data. These values are much larger than the ones for the clustering coefficients in the previous section, see Table III and the difference they make between random graphs and real-world networks is much more significant. To this regard, it may be considered as a better generalisation of clustering coefficients in classical networks (as opposed to affiliation ones) than the bipartite clustering coefficients defined in Section VIII.

The case of occurrences is different: the projections on both sides are very dense, which is very particular as already noticed. The redundancy coefficient therefore is huge, but this is not because of a property of how the neighbourhoods overlap: this is a direct consequence of the high density of the projections. In such a case, the redundancy coefficient is meaningless, and we will therefore not discuss this case any further in this section; simply notice that the redundancy coefficient has similar behaviours in such graphs and in their random equivalent.

|                | actors-movies | authoring | occurrences | peer-to-peer |
|----------------|---------------|-----------|-------------|--------------|
| \( \text{rc}(\top) \) | 0.26          | 0.38      | 0.80        | 0.31         |
| \( \text{rc}(\perp) \) | 0.25          | 0.33      | 0.83        | 0.50         |
| Real           | Random        | Real      | Random      | Real         |
|                | 0.014         | 0.0016    | 0.74        | 0.011        |
|                | 0.011         | 0.00037   | 0.75        | 0.069        |

**TABLE IV**

The redundancy coefficient for our four examples and for random bipartite graphs with the same size and same degree distributions.

We show in Figure 10 the distributions of \( \text{rc}(v) \) for our four examples together with plots for comparable random graphs. These plots confirm that the redundancy coefficient captures a property that makes real-world complex networks different from random ones: in all the cases except occurrences, the value of this coefficient in random graphs is almost 0 for all nodes (both top and bottom); instead, in real-world networks it is significantly larger, and equal to 1 for a large portion of the nodes. This last fact is not surprising since \( \text{cc}_\top(v) = 1 \) implies \( \text{rc}(v) = 1 \) for all nodes \( v \).

However, the redundancy coefficient has a much wider range of values than \( \text{cc}_\top(v) \), which generally is close to 0 or 1, see Figure 6. Moreover, the redundancy coefficient captures a different property: in the case of actors-movies, for instance, it does not only mean that a significant number of movies have a cast that is a sub-cast of another movie (as captured by \( \text{cc}_\top(v) \)), but that when two actors act together in a movie then there often exists (at least) another movie in which they also act together. Both are interesting, and complementary, but the redundancy coefficient certainly captures a more general feature.

Let us now observe the correlations between node redundancy coefficient and their degree, plotted in
Figure 10. Cumulative distributions of the redundancy coefficient in our four real-world affiliation networks, and in random bipartite graphs of the same size and same degree distributions. First row: for top nodes. Second row: for bottom nodes.

Figure 11. In these plots, except for occurrences, the plots for the random graphs coincide with the x-axis, which confirms that the values of node redundancy in random graphs are very small, independently of node degrees. Real-world cases, on the contrary, exhibit nontrivial behaviours. In most cases, the redundancy decreases with the degree, which is not surprising since the number of links needed in the projection so that the redundancy of a node is large grows with the square of its degree. However, the redundancy remains large even for quite large degrees: it is close to 0.15 for nodes of degree 30 for top nodes in actors-movies, for instance, meaning that among the 435 possible pairs of neighbours of these nodes, on average 65 are linked to another top node in common. This has a very low probability in random graphs. Likewise, one may notice that some very high degree nodes have a very large redundancy coefficient in several cases, which also is a significant information.

One may push further the study of the redundancy, for instance by counting how many nodes have an overlap with a given one, and so may be responsible for its high redundancy. This is nothing but the degree of the node in the appropriate projection, which emphasises once again that our approach may be combined with the one based on projection with benefit, as argued in Section III.
X. Conclusion and perspectives.

The core contribution of this paper is a set of rigorous and coherent statistical properties usable as a basis for the analysis of large real-world affiliation networks. These statistics go from the very basics (size, distances, etc) to subtle ones (typically various clustering coefficients and their correlations with degrees). Let us insist on the fact that we do not only extend classical notions to the bipartite case, but also develop new notions which make sense only in this context. Moreover, the proposed approach avoids projection of affiliation networks into classical ones, which makes it possible to grab much richer information. We hope that this unified framework and discussion will help significantly people involved in analysis of such networks.

A first conclusion driven from the computation of these statistics on four representative real-world examples is that, just like real-world classical complex networks, they have nontrivial properties in common which make them very different from random networks. In particular, there is a high heterogeneity between degrees of nodes of at least one kind, and there are significant overlaps between neighbourhoods. Concerning this last property, we show that immediate extensions of the classical notions of clustering coefficients are not sufficient to make the difference between real-world networks and random graphs; we propose the notion of redundancy as a relevant alternative. Overall, these facts are strikingly close to what is met in classical networks and should play a similar role. Conversely, we observed many properties which behave differently depending on the affiliation network under concern, which may be used to describe a particular instance in more details.

Notice that these contributions do not only concern the affiliation networks themselves, but also their projection: keeping the bipartite nature of the data makes it possible to obtain more precise information on the projection itself. For instance, statistics on degrees make it possible to separate high degree nodes in the projection into two distinct classes: the ones which are linked to many nodes in the affiliation network, and the ones linked to nodes of high degree in the affiliation network. This kind of analysis could be deepened using clustering and redundancy notions.

Going further, one may use the notions we introduced here to define new relevant statistics on classical networks. Indeed, any (classical) graph $G = (V, E)$ may be seen as a bipartite graph $G' = (V, V, E)$ where the links are between two copies of $V$. The statistics we studied here may then be computed on this bipartite graph, leading to new insight on the original graph $G$.

There are many directions to improve and continue the work presented here. Among them, let us cite the analytic study of the parameters we propose, which can in particular be done using the techniques in [57]. One might prove in this way the expected behaviour of these parameters and deepen their understanding. Another direction is the development of models of affiliation networks capturing the properties met in practice. Just like it is the case for classical networks, much can be done concerning degrees, but very little is known concerning the modeling of clustering and redundancy. Finally, applying these results to practical cases and giving precise interpretations of their meanings in these different contexts would probably help in designing other relevant notions.

Let us conclude by noticing that large complex network analysis is only at its beginning, though much has been done in recent years on classical networks. However, most real-world networks are directed, weighted, labelled, hybrid, and/or evolve during time. Some work has recently been done concerning weighted networks [5], [6], [49], and we propose here a contribution concerning affiliation networks. However, there is still much to do to be able to analyse efficiently these various kinds of networks. Extending the notions we discussed here to the case of multipartite graphs (nodes are in several disjoint sets, with links between nodes in different sets only) would be a step further in this direction.

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