Predicting Missing Links via Local Information

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Abstract. Missing link prediction of networks is of both theoretical interest and practical significance in modern science. In this paper, we empirically investigate a simple framework of link prediction on the basis of node similarity. We compare nine well-known local similarity measures on six real networks. The results indicate that the simplest measure, namely common neighbors, has the best overall performance, and the Adamic-Adar index performs the second best. A new similarity measure, motivated by the resource allocation process taking place on networks, is proposed and shown to have higher prediction accuracy than common neighbors. It is found that many links are assigned same scores if only the information of the nearest neighbors is used. We therefore design another new measure exploited information of the next nearest neighbors, which can remarkably enhance the prediction accuracy.

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1 Introduction

Many social, biological, and information systems can be properly described as networks with nodes representing individuals or organizations and edges mimicking the interactions among them. The study of complex networks has attracted increasing attention and become an common focus of many branches of science. Many efforts have been made to understand the evolution of networks [1, 2], the relations between topologies and functions [3,4], and the network characteristics [5]. Very recently, a fresh question is raised [6], that is, how to predict missing links of networks? For some networks, especially the biological networks such as protein-protein interaction networks, metabolic networks and food webs, the discovery of links/interactions costs much in the laboratory or the field, and thus the
current knowledge of those networks is substantially incomplete [7,8]. Instead of blindly checking all possible interactions, to predict in advance based on the interactions known already and focus on those links most likely to exist can sharply reduce the experimental costs if the predictions are accurate enough. For some others like the friendship networks in web society [9], very likely but not yet existing links can be suggested to the relevant users as recommendations of promising friendships, which can help users finding new friends and thus enhance their loyalties to web sites.

Majority of the previous works on missing link prediction have used some external information besides the network topology [10]. Graven et al. [11] predicted the semantic relationships of the world wide web with the help of web content. Popescul and Ungar [12] designed a regression model to predict citations made in scientific literature based on not only the citation graph, but also the authorship, journal information and content. Taskar et al. [13] applied the relational Markov network algorithm to predict missing links in a network of web pages and a social network, in which the well-defined attributes of each node are exploited. O’Madadhain et al. [14] constructed local conditional probability models for link prediction, based on both structural features and nodes’ attributes. The usage of external information can somewhat enhance the algorithmic accuracy, however, the content and attribute information are not available for general networks, thus the applications of above algorithms are strongly limited. Goldberg and Roth [15] exploited the neighborhood cohesiveness property of small-world networks to assess confidence for individual protein-protein interactions. Liben-Nowell and Kleinberg [16] empirically investigated the similarity-based prediction algorithms for large scientific collaboration networks. Clauset et al. [17] designed a prediction algorithm based on the inherent hierarchical organization of social and biological networks.

These mentioned works are practically successful in dealing with specific networks, however, thus far, a comprehensive picture about the dependence of algorithmic performance on network topology is lacking. The reason is twofold: (i) the works from engineering and biological communities have not yet caught up with the current state of development in characterizing the topologies of complex networks [9], while (ii) the physics community has not paid enough attention to the link prediction problem. Accordingly, dozens of important issues are still less explored. For example, one may concern about how to choose a suitable algorithm given some structural descriptions of a network, such as small-world phenomenon [18], degree heterogeneity [19], mixing pattern [20], community structure [21], and so on. In the opposite viewpoint, comparison of the performances of some prediction algorithms may reveal some structural information of the networks. It is just like the community structure has significantly effect on the network synchronizability [22], while the synchronizing process can be properly used to reveal the underlying community structure [23]. In addition, the algorithms only based on local information are generally fast but of lower accuracy, while the ones making use of the knowledge of
global topology are of higher accuracy yet higher computational complexity \cite{10}. Can we find a good tradeoff that provides high quality predictions while requires light computation?

In this paper, we empirically investigate a simple framework of link prediction on the basis of node similarity. Although the framework is simple, it opens a rich space for exploration since the design of similarity measures is challenging and can be related to very complicated physics dynamics and mathematical theory, such as random walk \cite{24} and counting problem of spanning trees \cite{25}. Here we concentrate on local-information-based similarities. We compare nine well-known local measures on six real networks, and the results indicate that the simplest measure, namely common neighbors, has the best overall performance, which is in accordance with the empirical results reported in Ref. \cite{16}. Motivated by the resource allocation process in transportation networks, we next propose a new similarity measure, which performs obviously better than common neighbors, which requires no more information and computational time. Furthermore, it is found that many links get same scores under local similarity measures, just like the degeneracy of energy level. We therefore design a new measure using the information of the next nearest neighbors, which can break the “degenerate states” thus remarkably enhance the algorithmic accuracy. Finally, we outline some future interests on this issue.

2 Method

Considering an undirected simple network $G(V, E)$, where $V$ is the set of nodes and $E$ is the set of links. The multiple links and self-connections are not allowed. For each pair of nodes, $x, y \in V$, every algorithm referred in this paper assigns a score as $s_{xy}$. Since $G$ is undirected, the score is symmetry, namely $s_{xy} = s_{yx}$. This score can be viewed as a measure of similarity between nodes $x$ and $y$, hereinafter, we do not distinguish similarity and score. All the nonexistent links are sorted in decreasing order according to their scores, and the links in the top are most likely to exist.

To test the algorithmic accuracy, the observed links, $E$, is randomly divided into two parts: the training set, $E^T$, is treated as known information, while the probe set, $E^P$, is used for testing and no information in probe set is allowed to be used for prediction. Clearly, $E = E^T \cup E^P$ and $E^T \cap E^P = \emptyset$. In this paper, the training set always contains 90% of links, and the remaining 10% of links constitute the probe set. We use a standard metric, area under the receiver operating characteristic (ROC) curve \cite{26}, to quantify the accuracy of prediction algorithms. In the present case, this metric can be interpreted as the probability that a randomly chosen missing link (a link in $E^P$) is given a higher score than a randomly chosen nonexistent link (a link in $U \setminus E$, where $U$ denotes the universal set). In the implementation, among $n$ times of independent comparisons, if there are $n'$ times the missing link having higher score and $n''$ times the missing link and nonexistent link having the same score, we define the ac-
curacy as $n' + 0.5n''$. If all the scores are generated from an independent and identical distribution, the accuracy should be about 0.5. Therefore, the degree to which the accuracy exceeds 0.5 indicates how much better the algorithm performs than pure chance.

### 3 Data

In this paper, we consider six representative networks drawn from disparate fields: (i) PPI.— A protein-protein interaction network containing 2617 proteins and 11855 interactions [27]. Although this network is not well connected (it contains 92 components), most of nodes belong to the giant component, whose size is 2375. (ii) NS.— A network of coauthorships between scientists who are themselves publishing on the topic of networks [28]. The network contains 1589 scientists, and 128 of which are isolated. Here we do not consider those isolated nodes. The connectivity of NS is not good, actually, NS is consisted of 268 connected components, and the size of the largest connected component is only 379. (iii) Grid.— An electrical power grid of western US [18], with nodes representing generators, transformers and substations, and edges corresponding to the high voltage transmission lines between them. (iv) PB.— A network of the US political blogs [29]. The original links are directed, here we treat them as undirected links. (v) INT.— The router-level topology of the Internet, which is collected by the Rocketfuel Project [30]. (vi) USAir.— the network of US air transportation system, which contains 332 airports and 2126 airlines [31].

**Table 1.** The basic topological features of six example networks. $N$ and $M$ are the total numbers of nodes and links, respectively. $N_c$ denotes the size of the giant component, for example, the entry $2375/92$ in the first line means that the network has 92 components and the giant component consists of 2375 nodes. $e$ is the network efficiency [32], defined as $e = \frac{2}{N(N-1)} \sum_{x,y \in V, x \neq y} d_{xy}^{-1}$, where $d_{xy}$ is the shortest distance between $x$ and $y$, and $d_{xy} = +\infty$ if $x$ and $y$ are in two different components. $C$ and $r$ are clustering coefficient [18] and assortative coefficient [20], respectively. Nodes with degree 1 are excluded from the calculation of clustering coefficient. $H$ is the degree heterogeneity, defined as $H = \frac{\langle k^2 \rangle}{\langle k \rangle^2}$, where $\langle k \rangle$ denotes the average degree.

| Nets   | $N$  | $M$  | $N_c$  | $e$  | $C$  | $r$  | $H$  |
|--------|------|------|--------|------|------|------|------|
| PPI    | 2617 | 11855| 2375/92| 0.180| 0.387| 0.461| 3.73 |
| NS     | 1461 | 2742 | 379/268| 0.016| 0.878| 0.462| 1.85 |
| Grid   | 4941 | 6594 | 4941/1 | 0.063| 0.107| 0.003| 1.45 |
| PB     | 1224 | 19090| 1222/2 | 0.397| 0.361| -0.079| 3.13 |
| INT    | 5022 | 6258 | 5022/1 | 0.167| 0.033| -0.138| 5.50 |
| USAir  | 332  | 2126 | 332/1  | 0.406| 0.749| -0.208| 3.46 |

Table 1 summarizes the basic topological features of those networks. Brief definitions of the monitored topological measures can be found in the table caption, for more details, please see the review articles [1, 2, 3, 4, 5]. We here give a few remarks for the numbers which may be unexpected to some readers: (i) It is well known that in the protein-protein interaction networks, links between highly connected proteins are systematically suppressed, while those between highly-connected and low-connected pairs...
are favored \cite{33}. That is to say, the assortative coefficient should be negative for PPI (for example, as reported in Ref. \cite{20}, the Yeast PPI network has an assortative coefficient -0.156), however, in the present network, the assortative coefficient is very positive, as 0.461. It is because the data set used here \cite{27} is determined from functional interactions and not from physical interactions. More detailed discussion can be found in Ref. \cite{34}. (ii) The extremely large clustering coefficient of NS dues to the specific constructing rule of collaboration networks, namely all the participants in an act are fully connected. Relevant discussion can be found in Appendix B of Ref. \cite{35}.

4 Comparison of Nine Similarity Measures Based on Local Information

In this section, we compare prediction accuracies of nine similarity measures. All these measures are based on the local structural information contained in the testing set. We first give a brief introduction of each measure as follows.

(i) **Common Neighbors.**— For a node $x$, let $\Gamma(x)$ denote the set of neighbors of $x$. In common sense, two nodes, $x$ and $y$, are more likely to have a link if they have many common neighbors. The simplest measure of this neighborhood overlap is the directed count, namely

$$s_{xy} = |\Gamma(x) \cap \Gamma(y)|. \quad (1)$$

(ii) **Salton Index.**— Salton index \cite{36} is defined as

$$s_{xy} = \frac{|\Gamma(x) \cap \Gamma(y)|}{\sqrt{k(x) \times k(y)}}, \quad (2)$$

where $k(x) = |\Gamma(x)|$ denotes the degree of $x$. Salton index is also called cosine similarity in the literature.

| Algorithms | PPI | NS | Grid | PB | INT | USAir |
|------------|-----|----|------|----|-----|-------|
| CN         | 0.889 | 0.933 | 0.590 | 0.925 | 0.559 | 0.937 |
| Salton     | 0.869 | 0.911 | 0.585 | 0.874 | 0.552 | 0.898 |
| Jaccard    | 0.888 | 0.933 | 0.590 | 0.882 | 0.559 | 0.901 |
| Sørensen   | 0.888 | 0.933 | 0.590 | 0.881 | 0.559 | 0.902 |
| HPI        | 0.868 | 0.911 | 0.585 | 0.852 | 0.552 | 0.857 |
| HDI        | 0.888 | 0.933 | 0.590 | 0.877 | 0.559 | 0.895 |
| LHN        | 0.866 | 0.911 | 0.585 | 0.772 | 0.552 | 0.758 |
| PA         | 0.828 | 0.623 | 0.446 | 0.907 | 0.464 | 0.886 |
| AA         | 0.888 | 0.932 | 0.590 | 0.922 | 0.559 | 0.925 |
| ZLZ-I      | 0.890 | 0.933 | 0.590 | 0.931 | 0.559 | 0.955 |
| ZLZ-II     | 0.939 | 0.938 | 0.639 | 0.936 | 0.632 | 0.900 |

Table 2. Accuracies of algorithms, measured by the area under the ROC curve. Each number is obtained by averaging over 10 implementations with independently random divisions of testing set and probe set. The abbreviations, CN, Salton, Jaccard, Sørensen, HPI, HDI, LHN, PA, and AA, stand for Common Neighbors, Salton Index, Jaccard Index, Sørensen Index, Hub Prompted Index, Hub Depressed Index, Leicht-Holme-Newman Index, Preferential Attachment, and Adamic-Adar Index, respectively. The entries corresponding to the highest accuracies among these nine measures are emphasized by black. ZLZ-I and ZLZ-II stand for the similarity measures proposed in Section 5 and Section 6 respectively, where ZLZ is the abbreviation of the authors’ names.
(iii) **Jaccard Index.**— This index was proposed by Jaccard [37] over a hundred years ago, which is defined as

\[ s_{xy} = \frac{|\Gamma(x) \cap \Gamma(y)|}{|\Gamma(x) \cup \Gamma(y)|}. \]  

(3)

(iv) **Sørensen Index.**— This index is mainly used for ecological community data [38], which is defined as

\[ s_{xy} = \frac{2 \times |\Gamma(x) \cap \Gamma(y)|}{k(x) + k(y)}. \]  

(4)

(v) **Hub Prompted Index.**— This index is proposed for quantifying the topological overlap of pairs of substrates in metabolic networks [39], defined as

\[ s_{xy} = \frac{|\Gamma(x) \cap \Gamma(y)|}{\min\{k(x), k(y)\}}. \]  

(5)

Under this measure, the links adjacent to hubs (here, the term “hub” stands for node with very large degree) are probably assigned high scores since the denominator is determined by the lower degree only.

(vi) **Hub Depressed Index.**— Analogously to the above index, we consider a measure with opposite effect on hubs for comparison, which is defined as

\[ s_{xy} = \frac{|\Gamma(x) \cap \Gamma(y)|}{\max\{k(x), k(y)\}}. \]  

(6)

(vii) **Leicht-Holme-Newman Index.**— This index gives high similarity to node pairs that have many common neighbors compared not to the maximum number possible but to the expected number of such neighbors [40]. It is defined as

\[ s_{xy} = \frac{|\Gamma(x) \cap \Gamma(y)|}{k(x) \times k(y)}, \]  

(7)

where the denominator, \(k(x) \times k(y)\), is proportional to the expected number of common neighbors of nodes \(x\) and \(y\) in the configuration model [41].

(viii) **Preferential Attachment.**— The mechanism of preferential attachment can be used to generate evolving scale-free networks (i.e., networks with power-law degree distributions), where the probability that a new link is connected to the node \(x\) is proportional to \(k(x)\) [19]. Similar mechanism can also lead to scale-free networks without growth [42], where at each time step, an old link is removed and a new link is generated. The probability this new link is connecting \(x\) and \(y\) is proportional to \(k(x) \times k(y)\). Motivated by this mechanism, a corresponding similarity index can be defined as

\[ s_{xy} = k(x) \times k(y), \]  

(8)

which has already been suggested as a proximity measure [43], as well as been used to quantify the functional significance of links subject to various network-based dynamics, such as percolation [44], synchronization [45] and transportation [46]. Note that, this index requires less information than all others, namely it does not need to know the neighborhood of each node. As a consequence, it also has the minimal computational complexity.

(ix) **Adamic-Adar Index.**— This index refines the simply counting of common neighbors by assigning the lower-connected neighbors more weights [47], which is defined as:

\[ s_{xy} = \sum_{z \in \Gamma(x) \cap \Gamma(y)} \frac{1}{\log k(z)}. \]  

(9)

We present the algorithmic accuracies for the six example networks in Table 2, with those entries corresponding to the highest accuracies being emphasized by black. To our surprise, the simplest measure, common neighbors,
performs the best. This result is in accordance with the one reported in Ref. [16] for social collaboration networks. Besides CN, Adamic-Adar index performs the next best for its accuracies are always close to the best one, while others, such as Jaccard index, Sørensen index and HDI, perform far worse in the cases for PB and USAir.

Note that, the first seven measures, from CN to LHN, only different in denominators. If all the nodes have pretty much the same degrees, corresponding to a very small $H$, then the difference among those measures becomes insignificant. In addition, for a given network, if its clustering coefficient is very small, whether two nodes have common neighbors plays the most important role, while the denominator is less important. In a word, remarkable difference among those seven measures can be found only if the monitored network simultaneously has large clustering coefficient and large degree heterogeneity, such as PPI, PB and USAir. As shown in Table 2, the performances of those seven algorithms on PB and USAir are obviously different, but for PPI, they are more or less the same. A possible reason is that PPI is a very assortative network (i.e., $r = 0.461$), and thus two nodes of a link tend to have similar degrees, which reduces the difference in denominators.

The preferential attachment has the worst overall performance, however, we are interested in it for it requires the minimal information. One may intuitively think that PA will give good predictions for assortative networks, while performs badly for disassortative networks. However, no obvious correlation between assortative coefficient and algorithmic accuracy based on PA can be found from our numerical results. The reason is twofold. Firstly, links between pairs of large-degree nodes contribute positively to the assortative coefficient and are assigned high scores by PA, while links between pairs of small-degree nodes also contribute positively to the assortative coefficient but are disfavored by PA. Actually, assortative coefficient is an integrated measure involving many ingredients, and there is no simple relation between this measure and the performance of PA. Secondly, assortative coefficient itself is very sensitive to the degree sequence, and a network of higher degree heterogeneity tends to be disassortative [18]. Therefore, this single parameter can not reflect the detailed linking patterns of networks. Clearly, if the large-degree nodes are very densely connected to each other, and the small-degree nodes are rarely connected to each other, PA will perform relatively good. The former relates to the so-called rich-club phenomenon [49], and we have checked that PB and USAir exhibit obvious rich-club phenomenon with respect to their randomized versions (we followed the method proposed by Colizza et al. [50], and they have already demonstrated that the air transportation network shows the presence of rich-club phenomenon). In addition, in USAir, more than 40% of nodes are very small local airports, with degrees no more than 3. A local airport usually connects to a nearby central airport and a very few hubs, the direct links between two local airports are rarely found. This topological feature is also favored by PA. As shown in Table 2, PA gives relatively good predictions for PB and USAir, in accordance with the above discussion.
Note that, all the other eight measures will automatically assign zero score to the pair of nodes located in different components. Therefore, PA performs badly when the network is consisted of many components. This is the very reason why PA gives very bad predictions for NS, although NS has clear rich-club phenomenon. We also note that, PA performs even worse than pure chance for the Internet at router level and the power grid. In these two networks, the nodes have well-defined positions and the links are physical lines. Actually, geography plays a significant role and the links with very long geographical distances are rare (the empirical analysis of spatial dependence of links in the Internet can be found in Ref. [51], and the absence of clustering-degree correlation in the router-level Internet and power grid can be considered as an indicator of a strong geographical constraint [52]). PA can not take into account the effect of geographical localization at all. As local centers, the large-degree nodes have longer geographical distances to each other than average, correspondingly, they also have less probability to directly connect to each other. Actually, these two networks exhibit the anti-rich-club phenomenon, that is, the link density among very-large-degree nodes are even lower than the randomized versions. This anti-rich-club effect leads to the bad performance of PA. In contrast, although USAir has well-defined geographical positions of nodes, its links are not physical. Empirical data demonstrated that the air transportation networks show an inverse relation between clustering coefficient and degree [53], and the number of airline flights is not sensitive to the geographical distance within the range of about 2000 kilometers [54]. As a final remark, comparing Eq. (7) and Eq. (8), LHN is, to some extent, inverse to PA, therefore when PA performs badly, LHN will give relatively good predictions, and vice versa.

5 Similarity Measure Based on Resource Allocation

Except PA, all the others introduced in the last section are neighborhood-based measures. Although they are simple and mathematically graceful, they are not tightly related to any physical processes. In this section, motivated by the resource allocation process taking place in networks [55], we propose a new similarity measure, which has remarkably higher accuracy than all the measures mentioned in Section 4.

Considering a pair of nodes, $x$ and $y$, which are not directly connected. The node $x$ can send some resource to $y$, with their common neighbors playing the role of transmitters. In the simplest case, we assume that each transmitter has a unit of resource, and will averagely distribute it to all its neighbors. Define the amount of resource $y$ received the similarity between $x$ and $y$, one has:

$$s_{xy} = \sum_{z \in \Gamma(x) \cap \Gamma(y)} \frac{1}{k(z)}$$  \hspace{1cm} (10)

Clearly, this measure has a similar form as the Adamic-Adar index, and is also symmetry, namely $s_{xy} = s_{yx}$.

We denote this similarity index as ZLZ-I with ZLZ the abbreviation of the authors’ names. The algorithmic accuracies on the six example networks are presented in Table 2. Compared with all the nine measures introduced
in Section 4, ZLZ-I performs the best. Especially for the networks (i.e., PB and USAir) with large clustering coefficient, high degree heterogeneity and absence of strongly assortative linking pattern, ZLZ-I largely outperforms all other measures. It is observed that, ZLZ-I exhibits particularly good performance on USAir. The reason may be that the resource allocation process is originally proposed to explain the nonlinear correlation between transporta-
tion capacity and connectivity of each airport [53, 56, 57].

ZLZ-I can be easily extended to the case of directed networks. Assuming a unit of resource is located in \( x \), which will be equally send to all \( x \)'s neighbors, each of which will equally distribute the received resource one step further to all its neighbors. The amount of resource a node \( y \) received can be considered as the importance of \( y \) in \( x \)'s sense, denoted as

\[
s_{xy} = \frac{1}{k(x)} \sum_{z \in \Gamma(x) \cap \Gamma(y)} \frac{1}{k(z)}.
\]

(11)

In this case, \( s_{xy} \neq s_{yx} \). This asymmetry measure has already been applied in a personal recommendation algorithm of bipartite user-object networks, and been demonstrated to be much more accurate than the simple counting of common neighbors [53, 56].

6 Improving Algorithmic Accuracy by

Breaking the Degenerate States

The neighborhood-based measures require only the information of the nearest neighbors, therefore have very low computational complexity. However, the information usually seems insufficient, and the probability that two node pairs are assigned the same score is high. That is to say, the neighborhood-based similarity is less distinguishable. If we consider the score assigned to a node pair as its energy, then many node pairs crowd into a very few energy levels. Taking INT as an example, there are more than \( 10^7 \) node pairs, 99.59% of which are assigned zero score by CN. For all the node pairs having scores higher than 0, 91.11% of which are assigned score 1, and 4.48% are assigned score 2. Using a little bit more information involving the next nearest neighbors may break the “degenerate states” and make the scores more distinguishable. Denoting \( A \) the adjacent matrix, where \( A_{xy} = 1 \) if \( x \) and \( y \) are directly connected, and \( A_{xy} = 0 \) otherwise. Obviously, \( (A^2)_{xy} \) is the number of common neighbors of nodes \( x \) and \( y \), which is also equal to the number of different paths with length 2 connecting \( x \) and \( y \). And if \( x \) and \( y \) are not directly connected (this is the case we are interested in), \( (A^3)_{xy} \) is equal to the number of different paths with length 3 connecting \( x \) and \( y \). The information contained in \( A^3 \) can be used to break the degenerate states, and thus we define a new measure as

\[
S = A^2 + \epsilon A^3,
\]

(12)

where \( S \) denotes the similarity matrix and \( \epsilon \) is a free parameter. Clearly, this measure degenerates to CN when \( \epsilon = 0 \). Here, the information in \( A^3 \) is only used to break the degenerate states, therefore \( \epsilon \) should be a very small number close to zero (of course, given a network, one can tune \( \epsilon \) to find its optimal value corresponding to the highest accuracy, however, this optimal value is different for different networks, and a parameter-dependent measure
is less practical in dealing with huge-size networks since the tuning process may take too long time). In the real implementation, we directly count the number of different paths with length 3, which is much faster than the matrix multiplication, and thus Eq. (12) is also based on local calculation.

The algorithmic accuracies on the six example networks are presented in Table 2, where this measure is denoted by ZLZ-II and the parameter is fixed as $\epsilon = 10^{-3}$. It is happy to see that the accuracy, except for USAir, can be largely enhanced by ZLZ-II. In USAir, the large-degree nodes are densely connected and share many common neighbors. Some links among large-degree nodes are removed into the probe set. Even without the contribution of $\epsilon A^3$, those links are assigned very high scores, thus the additional item, $\epsilon A^3$, changes little of their relative positions. Considering two small local airports, $x$ and $y$, which are connected to their local central airports, $x'$ and $y'$. Of course, many hubs are common neighbors of $x'$ and $y'$, and $x'$ and $y'$ may be directly connected. If the link $(x, x')$ is removed, the similarities between $x$ and other nodes are all zero for both CN and ZLZ-II. If $(x, x')$ exists, by ZLZ-II, the similarities $s_{xy'}$ (by $x$-$x'$-$y'$-$y'$), $s_{xy}$ (by $x$-$x'$-$y'$-$y'$), and $s_{xh}$ where $h$ stands for hub node (by $x$-$x'$-$hub$-$hub$) are positive due to the contributions of paths with length 3. There are many links connecting small local airports and local centers, some of which are removed, and the others are kept in the testing set. According to the above discussion, the removed links have lower score than the nonexistent links due to the additional item $\epsilon A^3$. In a word, the very specific structure of USAir (the hierarchical organization consisted of hubs, local centers and small local airports) makes the ZLZ-II worse than the simple CN. In this specific case, we can break the degenerate states in the opposite direction by setting $\epsilon$ being equal to $-10^3$, which lead to an accuracy 0.945, obviously higher than the one by CN, 0.937.

7 Conclusion and Discussion

In this paper, we empirically compared some link prediction algorithms based on node similarities. All the similarity measures discussed here, including the two newly proposed ones, can be obtained by local calculations. Numerical results on the nine well-known measures indicate that: (i) The simplest measure, common neighbors, performs the best, and the Adamic-Adar index is the second; (ii) Remarkable difference among these measures, excluding the Adamic-Adar index and the preferential attachment, can be observed only if the monitored network is with large clustering coefficient, high degree heterogeneity, and absence of strongly assortative linking pattern; (iii) The preferential attachment performs relatively good if the monitored network has the rich-club phenomenon.

We proposed a new measure motivated by the resource allocation process, which is equivalent to the one-step random walk starting from the common neighbors. This measure has a similar form as the Adamic-Adar index, but performs much better. We guess it is particularly suitable for link prediction of transportation networks, whose validity needs further evidence from more empirical results.
We here strongly recommend this measure to relevant applications and theoretical analyses, not only for its good performance, but also for its simplicity and grace.

Furthermore, we found that many links are assigned the same scores based on the local measures using the information of the nearest neighbors only. Exploitation of some additional information of the next nearest neighbors can therefore break the degenerate states and enhance the algorithmic accuracy. In real applications, the algorithms based on global calculations may be less efficient for they require long time and/or large memory, while the algorithms only exploited very local information may be less effective for their low accuracies. A properly designed algorithm can provide a good tradeoff just like the one presented in this paper. A similar idea has also been adopted in the network-based traffic dynamics, where the information of the next nearest neighbors can sharply enhance the traffic efficiency compared with the case in which only the information of the nearest neighbors is known [60].

Although the framework adopted here is very simple, it opens a rich space for investigation since in principle, all algorithms can be embedded into this framework differing only in the similarity measures. Besides ones discussed in this paper, a number of similarities are based on the global structural information, such as the average commute time of random walk [24], the number of spanning trees embedding a given node pair [25], the pseudoinverse of the Laplacian matrix [61], and so on. Some other similarity measures are even more complicated, depending on parameters. These include the Katz index [62] and its variant [63], the PageRank index [64], and so on. These measures should give better predictions than the local ones, however, the calculation of such measures, including determination of the optimal parameters for specific networks, is of high complexity, and thus infeasible for huge-size networks. Anyway, up to now, we lack systematic comparison and clear understanding of the performances of these measures, which is set as our future works.

Empirical analysis on more real networks as well as more known and newly proposed similarity measures is very valuable for building up knowledge and experience, and we can expect a clear picture of this issue can be completed by putting together of many fragments from respective empirical studies. However, the empirical results may be not clear at all since many unknown and uncontrollable ingredients are always mixed together in real networks. An alternative way is to build artificial network models with controllable topological features, and to compare the prediction algorithms on these models.

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