Identifying accurate link predictors based on assortativity of complex networks

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Abstract

Link prediction algorithms in complex networks, such as social networks, biological networks, drug-drug interactions, communication networks, and so on, assign scores to predict potential links between two nodes. Link prediction (LP) enables researchers to learn unknown, new as well as future interactions among the entities being modeled in the complex networks. In addition to measures like degree distribution, clustering coefficient, centrality, etc., another metric to characterize structural properties is network assortativity which measures the tendency of nodes to connect with similar nodes. In this paper, we explore metrics that effectively predict the links based on the assortativity profiles of the complex networks. To this end, we first propose an approach that generates networks of varying assortativity levels and utilize three sets of link prediction models combining the similarity of neighborhoods and preferential attachment. We carry out experiments to study the LP accuracy (measured in terms of area under the precision-recall curve) of the link predictors individually and in combination with other baseline measures. Our analysis shows that link prediction models that explore a large neighborhood around nodes of interest, such as CH2-L2 and CH2-L3, perform consistently for assortative as well as disassortative networks. While common neighbor-based local measures are effective for assortative networks, our proposed combination of common neighbors with node degree is a good choice for the LP metric in disassortative networks. We discuss how this analysis helps achieve the best-parameterized combination of link prediction models and its significance in the context of link prediction from incomplete social and biological network data.

A wide range of real-world problems can be effectively solved by modeling them as complex networks which are represented as a graph having nontrivial topological characteristics compared to random networks\textsuperscript{1}. Such complex networks play a significant role in identifying the significance of nodes and understanding the different connectivity patterns using different algorithms.

Existing algorithms on complex networks can answer different questions such as ranking the nodes based on some characteristics, predicting the structures of different topologies, and showing the flow of node/edge influence within the network among others. Link prediction (LP) models form an important class of such algorithms for complex networks. They are widely used on social networks (like Facebook, Twitter, LinkedIn, YouTube, etc.) as a way for suggesting friends, groups, videos, and any sort of possible group affiliation. In recommendation systems (such as books, movies, music, etc.), LP models were used to improve the similarity measurement of collaborative filtering methods, by exploring the association within user-item interactions to predict user interests and preferences\textsuperscript{2}. For example, they have been used to promote products for people who share the same shopping behavior e.g., on Amazon, or promote movies on YouTube or Netflix and so on. Similarly, in biological systems, the high-throughput methods often detect an incomplete view of the network and exhibit high false-positive and false-negative rates of protein interactions\textsuperscript{3}. LP models play a key role in predicting the possible Protein–Protein interactions in the biological system\textsuperscript{4}. Other applications are collaborative prediction in scientific co-authorship networks\textsuperscript{5}, predicting the spread of epidemic disease, detecting the drug-target interactions for drug discovery\textsuperscript{6,7} and even in bio-inspired networking\textsuperscript{8–11}.

The prediction of the future structure, or more precisely a future edge in a complex network is considered an active and ongoing research area that affects several applications in network science. Apparently, the entry of a new node (or set of nodes) into the network or the creation of new edges (or removing edges) is applicable to many network applications. Therefore, the prediction of future links plays a critical role in understanding the
future structure of the network and hence its effect on the functionality of the entire network or on a particular section of it. It is essential to reveal the mechanism by which a new (or an existing) node makes a connection with another, considering the difference in functionality and structure of the different networks. Hence, link prediction in complex networks has received wide attention using different approaches, such as graph theoretical methods, machine learning methods, probabilistic and statistical methods and so on\[12,13\]. Among them, the graph theoretic approach is considered a favorable way of computing the possible future links in comparison with the other methods, due to its general applicability. It uses the different graph features of the network to score possible future links between two given nodes. These methods consider the similarity between the two nodes as a way of determining the future link(s). Existing similarity-based methods can be classified into three categories: (1) local approaches, (2) global approaches and (3) quasi-local methods. We mainly focus on the local approaches due to their high performance and accuracy. In local similarity-based approaches, the nodes’ local structure (neighborhood) is the key metric used to compare different nodes. Local approaches are faster than the rest of the methods as they only require a scan of one layer of the neighborhood for each comparison. Several methods exist that measure the similarity of two nodes based on their local information.

Due to the complexity exhibited in the structures of complex networks, several algorithms measure either the overall network properties (like the degree distribution, clustering coefficient, hierarchical structures, and so on) or rank a network’s nodes by measuring the node’s influence or contribution in the network (e.g., different centrality measures such as degree, eigenvector, PageRank, betweenness\[14\] and so on\[15\]). Another important metric is the assortativity of the network\[16-18\]—a network property that refers to the preference of the nodes to affiliate/connect with other nodes that share similar features. Such features could refer to the similarity in degree, neighborhood, the existence of shared shortest paths, and so on. Specifically, a network is said to be assortative if its high-degree nodes connect with other high-degree nodes and low-degree nodes connect with other low-degree nodes. For example, in social networks (like Facebook), a node (or a person) has a high chance to make a relationship with others that share the same set of nodes (friends). Conversely, a network is said to be disassortative if its high-degree nodes connect with low-degree nodes. For example, biological networks exhibit such connectivity patterns where nodes with low-degree tend to have a connection with high-degree nodes.

Contributions. In this paper, we make the following contributions. First, we propose three sets of link prediction models based on (1) the similarity, (2) the dissimilarity of neighborhoods, and (3) extended preferential attachment\[19\] models—all using local topologies of the two nodes. We did not consider other path dependent metrics (such as betweenness\[14\], closeness\[19\], average shortest path etc.). Instead, we considered the global and local influence of a node relative to the network as a measure of the node’s participation (or influence) within the network. Second, we propose a model for link prediction for a missing edge \((x, y)\) using a parameterized combination of two different methods and compare their different versions against the assortativity value. In course of the analysis, we employ a combination of common neighbors method and each one of the three prediction models (similarity, dissimilarity, and extended preferential attachment). The common neighbor method is selected by default in the parameterized combination model due to its simplicity, intuitiveness, and performance. Also, it has earlier shown very competitive results in comparison with many complex approaches on real-world networks\[12,13\].

Third, and most importantly, we explore how the accuracy of the link prediction models varies with the assortativity of the networks. To this end, we present an approach that adapts an existing assortative network generation algorithm by Zhou et al.\[21\] to create networks of varying levels of assortativity (and disassortativity). We use this approach to determine the best-parameterized combination of link prediction models. Finally, we carry out extensive experiments on real-world and synthetic networks to evaluate the proposed link prediction models against standard local similarity-based algorithms and well-studied link prediction metrics taken from the literature. We show that similarity-based models perform better in highly assortative networks where a large percentage of edges connect nodes having a similar degree to each other. On the contrary, the dissimilarity-based models perform better in highly disassortative networks where nodes tend to form connections with other nodes that are dissimilar.

Material and methods

Dataset. We employ network data from the following sources to validate the proposed link predictors. (Table 1 shows the essential statistics for each of the selected networks.)

1. Karate\[22,23\] is a network of 34 members of a Karate club that reflects the members’ state of affiliation into groups due to a conflict between administrators and instructors. The dataset was collected and studied by Wayne W. Zachary from 1970 to 1972.
2. Dolphins\[24,25\] is a network that represents the frequent associations among 62 bottlenose dolphins.
3. Polbook is a network of US politics books (as nodes). Edges represent the frequent co-purchasing of books on amazon.com by the same buyer. The network dataset was retrieved from http://www.orgnet.com/.
4. USAir\[26\] is a network of airports (as nodes) and airlines (as edges) that represents the US air transportation system connecting the US around the globe.
5. **Word Adjacency** is a network that represents the existence of either noun-noun, adjective-noun, or adjective-adjective adjacent words in the novel of “David Copperfield”. Nouns and adjectives are represented as nodes, and their adjacency is represented as edges.

6. **Escherichia Coli GRN** is a biological network that reflects the genes and transcription factors of E. Coli and how they interact with each other to regulate the functionality of the organism. Genes and transcription factors represent the nodes and their interactions represent the edges of the network.

7. **Barabasi-Albert** is an algorithm that generates random scale-free networks using a preferential attachment model, i.e., the probability of a new node creating a connection with an existing node is proportional to the number of connections that the existing node has. This would result in new nodes tending/preferring to form connections with highly connected nodes.

8. **Facebook**, **Facebook** are two social networks extracted from Facebook. Here the nodes represent users (friends) and the edges represent the different web-based social interactions (of liking, sharing, or messaging).

9. **ca-sandi_auths** is a collaboration network of 86 scientists at Sandia National Labs.

10. **fb-pages-food** represents Facebook pages of several food companies (collected in 2017) and how they mutually interact among each other.

11. **soc-tribes** is a network acquired from the study conducted on the tribes of central Highlands of New Guinea. It shows the cultural-linguistic groups of that area and what similarities and differences exist among them.

12. **bn-cat-mixed-species_brain_1** is the neural connection networks (connectome) of cortical areas from the brain of cats.

13. **bn-macaque-rhesus_brain_2** represents the connectome that existed in the brain of rhesus macaque monkeys.

14. **bio-celegans** represents the connectome network of the Caenorhabditis elegans.

15. **soc-firm-hi-tech** represents a network of friendships among employees of a small hi-tech computer firm.

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| Name                                           | | | | | | | |
|------------------------------------------------|---|---|------|---|---|------|
| **Bn-macaque-rhesus_brain_2**                  | 91 | 582 | 0.2678 | 0.8601 | 1.8681 | −0.7698 | 3 | 0.1421 |
| **Karate**                                     | 34 | 78  | 0.2557 | 0.5706 | 2.4082 | −0.4756 | 5 | 0.1390 |
| **E. coli**                                    | 1565 | 3742 | 0.0155 | 0.2116 | 3.5791 | −0.3411 | 9 | 0.0031 |
| **Ca-sandi_auths**                             | 86 | 124 | 0.2721 | 0.4149 | 4.8140 | −0.2558 | 11 | 0.0339 |
| **Soc-firm-hi-tech**                           | 33 | 123 | 0.3875 | 0.6705 | 1.7689 | −0.2557 | 2 | 0.2330 |
| **Circuits**                                   | 122 | 304 | 0.0709 | 0.5656 | 1.9744 | −0.2487 | 3 | 0.0412 |
| **Bio-celegans**                               | 758 | 2025 | 0 | 0 | 3.7294 | −0.2233 | 11 | 0.0071 |
| **USAir97**                                    | 332 | 2126 | 0.3964 | 0.6252 | 2.7381 | −0.2079 | 6 | 0.0387 |
| **Word adjacencies**                           | 112 | 425 | 0.1569 | 0.1728 | 2.5356 | −0.1293 | 5 | 0.0684 |
| **Polbooks**                                   | 105 | 441 | 0.3484 | 0.4875 | 3.0788 | −0.1279 | 7 | 0.0808 |
| **Barabasi_albert_graph**                      | 500 | 1491 | 0.0322 | 0.0543 | 3.2335 | −0.0882 | 6 | 0.0120 |
| **Soc-tribes**                                 | 17 | 76  | 0.6131 | 0.6488 | 1.4485 | −0.0792 | 2 | 0.5588 |
| **Dolphins**                                   | 62 | 159 | 0.3088 | 0.2590 | 3.3570 | −0.0436 | 8 | 0.0841 |
| **fb-pages-food**                              | 620 | 2102 | 0.2226 | 0.3309 | 5.0887 | −0.0282 | 17 | 0.0110 |
| **bn-cat-mixed-species_brain_1**               | 65 | 730 | 0.5747 | 0.6614 | 1.6995 | −0.0254 | 3 | 0.3510 |
| **ENZYMES8**                                   | 141 | 133 | 0 | 0 | 4.3333 | 0.0161 | 10 | 0.0135 |
| **Reptilia-tortoise-network-sg**               | 24 | 26  | 0.4390 | 0.2639 | 3.6538 | 0.0162 | 9 | 0.0942 |
| **Reptilia-tortoise-network-mc**               | 15 | 28  | 0.6724 | 0.7094 | 1.6727 | 0.0408 | 3 | 0.2667 |
| **CAG_mat72**                                  | 72 | 750 | 0.6541 | 0.7511 | 2.1291 | 0.0470 | 6 | 0.2934 |
| **Reptilia-tortoise-network-pr**               | 35 | 66  | 0.5045 | 0.4884 | 2.4589 | 0.0583 | 6 | 0.1109 |
| **ENZYMES123**                                 | 135 | 127 | 0 | 0 | 5.7098 | 0.0912 | 12 | 0.0140 |
| **Reptilia-tortoise-network-im**               | 45 | 106 | 0.3644 | 0.4345 | 2.6439 | 0.1181 | 6 | 0.1071 |
| **Aves-weaver-social**                        | 445 | 1335 | 0.5881 | 0.6685 | 4.4699 | 0.2000 | 12 | 0.0135 |
| **Facebook348**                                | 448 | 6384 | 0 | 0 | 3.0253 | 0.2227 | 10 | 0.0638 |
| **Facebook414**                                | 300 | 3386 | 0 | 0 | 3.1918 | 0.3064 | 8 | 0.0755 |
| **Reptilia-tortoise-network-bsr**              | 136 | 374 | 0.3649 | 0.3335 | 3.7357 | 0.3254 | 10 | 0.0407 |
| **Reptilia-tortoise-network-cs**               | 73 | 132 | 0.4158 | 0.3146 | 2.4022 | 0.3934 | 6 | 0.0502 |
| **Reptilia-tortoise-network-fi**               | 787 | 1197 | 0.4199 | 0.2680 | 7.9334 | 0.4766 | 21 | 0.0039 |
| **Bio-SC-TS**                                  | 636 | 3959 | 0.9137 | 0.4712 | 1 | 0.9211 | 1 | 0.0196 |

Table 1. Network properties of 29 networks used in the analysis; **| V**: number of nodes in the network, **| E**: number of edges in the network, **r**: Assortativity coefficient value, **GCC & ACC**: Global and average clustering coefficients, **ASP**: Average shortest path, **d**: diameter, and **D**: graph density. The networks are sorted according to the assortativity level; (0 > r ≥ −1) for disassortative networks, (1 ≥ r > 0) for assortative networks.
(Continued from previous page)

2. Local influence of a node (LI): In this model, the node’s degree is compared to the average degree of the node’s neighbors to reflect the local influence of the node within its neighborhood area. Equation (2) depicts a node $u$’s influence with respect to the average degree of its neighbors, \( I^L_u \).

1. Global influence of a node (GI): In this model, the node’s degree is compared to the average degree of the network to reflect the global influence of the node within the network. Equation (1) depicts node $u$’s influence relative to the average degree of the network, \( I^G_u \).

\[
I^G_u = \frac{|\Gamma_u|}{|V|} \sum_{v \in V} |\Gamma_v| \quad u \in V
\]

\[
I^L_u = \frac{|\Gamma_u|}{\sum_{v \in V} |\Gamma_v|} \quad u \in V
\]
Based on node influence, we present three groups of models (discussed hereafter) to measure the possibility of having a link between two nodes: (1) extended preferential attachment models, (2) dissimilarity models and (3) similarity-based models.

**Extended preferential attachment model.** A well-known phenomenon of rich becomes richer [i.e., preferential attachment, see Eq. (13)] is where low-degree nodes tend to form a connection with highly connected nodes, especially in networks that follow a power law degree distribution. However, nodes may also tend to create a connection with other nodes based on the degree of influence within the network. Future links can be calculated using the same methodology as the preferential attachment, but using the node's influence instead of the node's degree. Herein, the node's influence towards attachment can utilize the node's global influence (\(I^G_u\)) or its local influence (\(I^L_u\)), \(u \in V\). Preferential attachment using global influence (PAGI) and preferential attachment using local influence (PALI) is used for scoring the potential for having a link/edge between \(x, y \in V\) using Eqs. (3) and (4).

\[
S_{x,y}^{PAGI} = I^G_x \ast I^G_y
\]

(3)

\[
S_{x,y}^{PALI} = I^L_x \ast I^L_y
\]

(4)

**Dissimilarity models.** The second model can be viewed as an extension to the preferential attachment where low-degree nodes tend to establish a connection/link with higher-degree nodes. Given this scenario, the potential for having an edge is increased as the difference in power/influence between the two nodes is increased. Herein, the absolute difference of either global influence (GI) or local influence (LI) has been used as a way to measure the possibility of establishing a link between the two given nodes. Therefore, as the difference (or dissimilarity) in power increases, there will be a higher chance of creating a link. Dissimilarity-based attachment using global influence (DAGI), and dissimilarity-based attachment using local influence (DALI) are used for scoring the potential of having a link between \(x, y \in V\) using Eqs. (5) and (6).

\[
S_{x,y}^{DAGI} = |I^G_x - I^G_y|
\]

(5)

\[
S_{x,y}^{DALI} = |I^L_x - I^L_y|
\]

(6)

**Similarity models.** Herein, distance is mostly used to check the degree of similarity of two given items; the high distance value means less similarity and vice versa. Thus, the third model can be viewed as an inverse of the dissimilarity-based attachment using global influence (inDAGI), and dissimilarity-based attachment using local influence (inDALI), see Eqs. (7) and (8).

\[
S_{x,y}^{inDAGI} = \frac{1}{S_{x,y}^{DAGI}} = \frac{1}{|I^G_x - I^G_y|}
\]

(7)

\[
S_{x,y}^{inDALI} = \frac{1}{S_{x,y}^{DALI}} = \frac{1}{|I^L_x - I^L_y|}
\]

(8)

These three different models consider the local and global influence of the nodes (an extension of the node’s degree) in the prediction of a connection between the two nodes. These metrics can also contribute to the local similarity-based metrics (such as common neighbors) in forming the connection. The resultant combined link prediction model that calculates the potential link score is given as follows:

\[
S_{x,y} = \alpha \cdot S_{x,y}^{CN} + (1 - \alpha)S_{x,y}^{Model} \\
Model \in \{PAGI, PALI, DAGI, DALI, inDAGI, inDALI\}
\]

(9)

A parameterized contribution of both of common neighbors and one of the proposed models would provide the final score. The \(\alpha\) parameter ranges in [0.2, 0.4, 0.6, 0.8]. We used the \(\alpha\) value to show the degree of the contribution that each measure has towards better overall link prediction. As the contribution of one measure increases, the contribution of the other will decrease.

**Baseline algorithms.** We compared our proposed models (refer to Section “Proposed models”) with the following local similarity-based algorithms. Local similarity-based approaches use node neighborhoods to measure the similarity of each node with other nodes in the network. Local approaches are faster than non-local approaches and it is highly parallelizable and efficient for dynamic networks. However, all of the following algorithms have a computation complexity of \(O(vk^2)\) except for the preferential attachment which has a computation complexity of \(O(vk)\); \(v\) refers to the number of vertices (nodes) and \(k\) refers to the degree of the node. Most of these methods are well explained in Martinez et. al.13.
1. **Common Neighbors**\(^{42}\) (CN) is the simplest and fundamental local technique. It measures the number of shared neighbors between two nodes \(x, y\). A confirmed hypothesis\(^{43}\) shows that for two distinct nodes, there is a correlation between the number of shared neighbors and the probability of being connected. The formula for \(S_{CN}^{x,y}\) is as follows:

\[
S_{CN}^{x,y} = |\Gamma_x \cap \Gamma_y| 
\]

2. **Adamic-Adar Index**\(^{44}\) (AA) is another variation of common neighbors which measures the similarity between \(x, y\) by logarithmically penalizing the shared neighbors.

\[
S_{AA}^{x,y} = \sum_{z \in \Gamma_x \cap \Gamma_y} \frac{1}{\log |\Gamma_z|} 
\]

3. **Resource Allocation Index**\(^{45}\) (RA) is another variation of both common neighbors and the Adamic-Adar index which models the unit of resources between two unconnected nodes through neighborhood nodes. The number of resource units transmitted from node \(x\) using \(x\)'s neighbors and received by node \(y\) reflects the degree of similarity between \(x, y\).

4. **Preferential attachment**\(^{19}\) (PA) is based on a premise that in a large set of real networks, node degrees tend to follow a power law distribution resulting in scale-free networks. The probability of having an edge between two nodes increases as their degrees increase.

5. **The Jaccard Index**\(^{46}\) (JA) is a widely used similarity measurement that measures the ratio of shared neighbors in the complete set of neighbors for two nodes.

\[
S_{JA}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{|\Gamma_x \cup \Gamma_y|} 
\]

6. **Salton Index**\(^{47}\) (SA) is another related measure to the Jaccard index, which is mostly known as the cosine similarity. In several experiments, the Salton index has been shown to be approximately twice the Jaccard index.

\[
S_{SA}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{\sqrt{|\Gamma_x||\Gamma_y|}} 
\]

7. **Sorensen Index**\(^{48}\) (SI) is a very similar method to the Jaccard index, used to compare the similarity between different ecological community data samples.

\[
S_{SI}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{|\Gamma_x| + |\Gamma_y|} 
\]

8. **Hub Promoted Index**\(^{19}\) (HPI) measures the similarity between \(x, y\) by comparing the ratio of common neighbors of nodes \(x, y\) to the minimum degree of either node.

\[
S_{HPI}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{\min(|\Gamma_x|, |\Gamma_y|)} 
\]

9. **Hub Depressed Index**\(^{19}\) (HDI) measures the similarity between \(x, y\) by comparing the ratio of common neighbors of nodes \(x, y\) to the maximum degree of either node.

\[
S_{HDI}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{\max(|\Gamma_x|, |\Gamma_y|)} 
\]

10. **Local Leicht-Homle-Newman Index**\(^{50}\) (LLHN) is a model where the similarity between \(x, y\) nodes is measured as the ratio of common neighbors of the \(x, y\) nodes to the multiplication of neighbors of the \(x, y\) nodes.

\[
S_{LLHN}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{|\Gamma_x||\Gamma_y|} 
\]

11. **Cannistraci-Alanis-Ravasi-based variation of the resource allocation**\(^{33,51}\) (CAR) is a model, where two nodes are likelier to have a connection if their common neighbors share very strong inner-links, forming so-called “local-community LC”.

\[
S_{CAR}^{x,y} = \frac{|\Gamma_x \cap \Gamma_y|}{|\Gamma_x||\Gamma_y|} 
\]
12. **CH2-L2 Index**\(^{52,53}\) is a link prediction model that assigns a reward for the internal connectivity existing among common neighbors and penalizes outside connectivity. Here \(C_i\) represents the number of neighbors of node \(i\) that exist in \(\hat{W}_x \cap \hat{W}_y\), \(O_i\) represents the number of neighbors of node \(i\) that do not exist in \(\hat{W}_x \cap \hat{W}_y\) nor in \(x\) or \(y\).

\[
S_{CH2-L2}^{x,y} = \sum_{i \in \hat{W}_x \cap \hat{W}_y} \frac{1 + C_i}{1 + O_i}
\]  \(\text{(21)}\)

Here \(C_i\) represents the number of neighbors of node \(i\) that exist in \(\Gamma_x \cap \Gamma_y\), \(O_i\) represents the number of neighbors of node \(i\) that do not exist in \(\Gamma_x \cap \Gamma_y\) nor in \(x\) or \(y\).

13. **CH2-L3 Index**\(^{52,53}\), very similar to CH2-L2 metric, this metric considers all three path lengths (two intermediate nodes \(i, j\)) between the targeted edge \((x, y)\).

\[
S_{CH2-L3}^{x,y} = \sum_{i \in \Gamma_x \cap \Gamma_y} \frac{A_{ij}}{(1 + C_i)(1 + C_j)}
\]  \(\text{(22)}\)

Here \(C_i\) represents the number of links between node \(i\) and all the nodes that exist in the set of intermediate nodes on all 3-hop paths connecting nodes \(x\) and \(y\), \(O_i\) represent the number of links between node \(i\) and all nodes that are not \(x, y\) nor the intermediate nodes on any 3-hop paths connecting \(x\) and \(y\).

**Network assortativity.** Assortativity, or assortative mixing, is a preference for the nodes to attach to other nodes that are similar in some way. Degree-based assortativity coefficient \(r\) of a network is measured as the Pearson correlation coefficient\(^{44,55}\) of the degree between all pairs of linked nodes, ranging from \((-1\) to \(1\)). Positive assortativity indicates a tendency of nodes to connect with other nodes of a similar degree. On the other hand, a negative correlation suggests that connections are more likely to exist between node pairs of dissimilar degrees.

\[
r = \frac{\sum_{ij} (A_{ij} - k_i k_j / 2m) k_i k_j}{\sum_{ij} (k_i \delta_{ij} - k_i k_j / 2m) k_i k_j}
\]  \(\text{(23)}\)

Here, \(A\) is the adjacency matrix of the network, \(k_i, k_j\) is the degree of node \(i, j\) respectively, \(\delta_{ij}\) is Kronecker delta. Equation (23) is an example of a Pearson correlation coefficient where it has covariance in the numerator and a variance in the denominator. Figure 1 shows two networks of assortative and disassortative types.

Figure 1. Level of assortativity for two networks. High-degree nodes are colored dark green while low-degree nodes are colored with a lighter color. Each node is labeled with its degree. (a) Assortative networks \((r = 0.6)\) where high-degree nodes are attached to high-degree nodes and low-degree nodes are attached to low-degree nodes. (b) Disassortative networks \((r = -0.84)\) where high-degree nodes are attached to low-degree nodes.
Generation of networks of varying assortativity. As discussed in Section "Introduction", we extend the Monte Carlo sampling approach presented by Zhou et al.21 to generate assortative and disassortative networks of a given order. While Zhou et al. constrain the degree distribution, we constrain the graph density of the final network. Since the degree distribution affects the assortativity of a network, constraining it may restrict the approach from achieving the necessary level of assortativity (Algorithm 1). The algorithm takes as input the following: the number of nodes in the final generated network \( N \), assortativity level \( \rho \), mode \( m \) (equal to 0, 1 for assortative and disassortative, respectively), initial network order \( n_0 \) and maximum node degree in the final network \( d_{\text{max}} \) with graph density \( D \), and outputs the undirected network \( G \).

As shown in Algorithm 1, the final complete network \( G \) is initialized with \( n_0 \) nodes. Subsequently, new nodes (\( v \)) are added to the network iteratively as follows. In lines 5–7, we calculate \( \hat{E} \) as the difference between the number of edges in \( G \) and the number of edges needed to achieve graph density \( D \). If \( \hat{E} = 0 \) the edges count requirement has already been met; otherwise surplus links are necessary. Specifically, once the surplus edge condition \( \hat{E} = 0 \) is met, the assortative network generation algorithm goes to the next new node. We calculate \( \rho \) as the ratio of the surplus edges to be added and the number of nodes left to be added, i.e., \( \rho = \left\lceil \frac{\hat{E}}{N - |V|} \right\rceil \). We input mode \( m = 0 \) to generate an assortative network, where higher \( \iota \rightarrow |L| \) will make the network increasingly less assortative. Conversely, for disassortative network, \( m \) is set to 1, for which \( \iota \rightarrow 0 \) will make the network increasingly less disassortative (lines 14–19). As depicted in Fig. 2, u-th element in the hash table \( L \) is the absolute difference between the degree of new node \( v \) and node \( u \), such that, the lower the difference, the more similar are nodes \( u \) and \( v \). For every new node \( v \), the idea is to choose a neighbor \( u \) with a (1) low \( L_v \) or (2) high \( L_u \) in order to generate a (1) assortative network or (2) disassortative network, respectively.

We input mode \( m = 0 \) to generate an assortative network, where higher \( \iota \rightarrow |L| \) will make the network increasingly less assortative. Conversely, for disassortative network, \( m \) is set to 1, for which \( \iota \rightarrow 0 \) will make the network increasingly less disassortative (lines 14–19). Specifically, the algorithm controls the extent of assortativity, by sorting \( L \) in the increasing order of \( L_u \) and introducing the assortativity level \( \rho \). The parameter \( \rho \) is necessary to determine \( \iota \) which marks the index of nodes \( u \) in \( L \) that will be candidates for neighbors of \( u \). Finally, in lines 22–27, we select \( k \) neighbors for node \( v (u \in V) \) depending on the choice of mode \( m \), while enforcing the degree distribution, i.e., \( \text{not}([w : w \in V \& \text{degree}(G,w) = \text{degree}(G,u)]) \) \( \leq N \times P_{\text{degree}(G,u)} \). Finally, the network \( G \) is returned as output.
Results

Evaluation criterion. The link prediction models mentioned in the algorithms section measure a similarity score $S_{m,x,y}$ for every missing edge $(x, y)$ in $G^p$, (m refers to the link prediction model in use). The resultant score estimates the possibility of having an edge between nodes $x$, $y$ given its neighbors’ structures. If the similarity-score value $S_{m,x,y}$ equals or surpasses a threshold, then an edge between $x$, $y$ is considered as predicted (or true positive TP) and otherwise rejected (false positive FP).

The range of similarity-score values that resulted from implementing the link prediction models varied even for the same graph. Therefore, to assess the performance of the different link prediction models on the given graph, the AUC (Area Under the receiver operating characteristic Curve) and AUPRC (Area Under Precision-Recall Curve) metrics are used. Given true positive (TP), true negative (TN), and false positive (FP) calculated on the true and predicted link labels, AUC and AUPRC are measured as follows:\(^{(24)}\):

1. **Area under Curve (AUC)** is a metric that measures the extent to which the model is capable of distinguishing between classes. In the context of this paper, AUC measures the probability that a randomly chosen existing edge is given a higher similarity score $S_{m,x,y}$ than a randomly chosen non-existing edge. AUC is measured as follows:

\[
AUC = \frac{n_1 + 0.5n_2}{n}
\]

Here, $n_1$ is the number of times that the missing edge got a higher score than an unconnected edge, $n_2$ is the number of times they are equal, and $n$ is the number of observations done. If AUC = 0.5, then the score is generated from an independent and identical distribution. Thus, an AUC value closer to 1 indicates how much better the link prediction model is when compared to the prediction by chance. Overall, the higher the AUC, the better the model is at assigning the right labels to different classes.

2. **Area under Precision-Recall Curve (AUPRC)** captures the trade-off between precision and recall, where precision is equal to $TP/(TP+FP)$ and recall is $TP/(TP+TN)$. A high AUPRC suggests that the model exhibits both high precision and recall.

Generation of assortative and disassortative networks. Algorithm 1 allows us to generate networks with varying assortativity (by modulating the assortativity level $\rho$). Table 2 shows the mean and standard deviation of the standard topological properties of 100 networks of size 250 nodes along with the level of assortativity and disassortativity and level of density. High assortative networks ($\rho > 0.7$) and less assortative networks ($0.25 < \rho < 0.6$) are generated with $\rho = 0$ and $\rho = 0.2$, respectively. Conversely, high disassortative networks ($\rho < -0.45$) and low disassortative networks ($-0.40 < \rho < -0.25$) are generated with $\rho = 1.0$ and $\rho = 0.8$, respectively. A correlation analysis is conducted between the graph density (D) and network assortativity ($r$) for each newly added node $v$, hash table $L_v$ where $L_u$ is the absolute difference between the degree of new node $v$ and node $u$. The keys in $L$ are arranged in the increasing order of $L_u$. The parameter $i$ marks the offset that controls the level of assortativity.

**Table 2.** Topological properties of a sample of the generated synthetic networks (of size 250) used in the analysis; Mean (and standard deviation of) GCC & ACC: Global and Average Clustering Coefficients, ASP: Average Shortest Path, $r$: Assortativity Coefficient value, $d$: diameter, and D: graph density.

|                  | GCC     | ACC     | ASP         | $r$   | $d$   | D       |
|------------------|---------|---------|-------------|-------|-------|---------|
| **High density** |         |         |             |       |       |         |
| High assortative | 0.67(0.04) | 0.64(0.02) | 2.76(0.23) | 0.74(0.03) | 7.46(1.44) | 0.18(0.01) |
| Less assortative | 0.35(0.02) | 0.29(0.02) | 2.03(0.03) | 0.3 (0.07) | 4(0.01) | 0.19(0.01) |
| Less disassortive | 0.18(0.02) | 0.2(0.02) | 1.86(0.01) | 0.28(0.05) | 3.19(0.39) | 0.19(0) |
| High disassortive | 0.24(0.02) | 0.47(0.04) | 1.81(0.01) | 0.43(0.02) | 2.59(0.49) | 0.19(0.01) |
| **Low density**  |         |         |             |       |       |         |
| High assortative | 0.54(0.01) | 0.17(0.02) | 11.92(0.18) | 0.88(0.01) | 24.97(0.72) | 0.01(0) |
| Less assortative | 0.21(0.03) | 0.15(0.03) | 2.74(0.24) | 0.47(0.05) | 5.57(0.64) | 0.08(0.02) |
| Less disassortive | 0.1(0.01) | 0.15(0.02) | 2.64(0.05) | 0.37(0.01) | 4.82(0.39) | 0.05(0) |
| High disassortive | 0.11(0.01) | 0.15(0.01) | 2.66(0.05) | 0.51(0.03) | 4.73(0.45) | 0.05(0) |

Figure 2. For each newly added node $v$, hash table $L_v$ where $L_u$ is the absolute difference between the degree of new node $v$ and node $u$. The keys in $L$ are arranged in the increasing order of $L_u$. The parameter $i$ marks the offset that controls the level of assortativity.
networks created by the generative algorithm. As shown in Fig. 3, no relationship exists between \( r \), \( D \) on both high density networks (Fig. 3a) and low density networks (Fig. 3b).

In addition to the standard sampling criteria that have been used in the literature for dividing the network edges into training (\( E^T \)) and testing (\( E^P \)), we apply another edge sampling criteria (termed similar degree edges, SDE criteria) to evaluate the performance of the different categories of the link prediction models. We collect the edges that have nodes with the same degree, sorted by the difference of degrees. This results in a descending list of edges, ranked in the decreasing order of score equal to the absolute value of the difference of nodes’ degree, i.e., \(|\Gamma_x - \Gamma_y||\). We have shown in Section “The effect of edge sampling” that the removal of similar degree links results in disassortative training networks. Therefore, this sampling criteria represents a worst-case scenario where the metrics are used to predict edges among similar degree nodes despite being trained on disassortative networks.

Cross validation on the synthetic networks. We implement the following cross-validation strategy. Unless otherwise stated, link prediction results reflect 50 runs (with 25 folds each) on 100 networks each of order 100, 200, 250, 500, and 1000 nodes.

1. For each of the 25 folds, the network set was randomly divided into 80% training and 20% testing set of networks. A network participates in either the training or the testing set.
2. The resultant evaluation values of the different link prediction models were collected and averaged resulting in an average over 25 folds \( \times \) 50 runs.

Prediction on standard networks. Standard networks are divided based on the assortativity coefficient \( r \) into assortative (\( 1 \geq r > 0 \)) and disassortative networks (\( 0 > r \geq -1 \)), see Table 1. Figure 4 shows the accuracy of the different link prediction metrics on assortative and disassortative networks, measured in terms of Area Under the Precision-Recall Curve (AUPRC). Figure 4a and b show the AUPRC for the assortative network group for random and similar degree edge removal, respectively, where local similarity-based metrics (AA, RA, and CH2-L2), relying on common neighborhoods, perform better than other metrics.

In the case of disassortative networks using random sampling of Fig. 4c, we also noticed that several weighted forms of the combined dissimilarity-based model of (CN+DALI, CN+DAGI) perform better than other local similarity models. Most of the standard and combined link prediction models that use local similarity (such as SA, JA, SO, CN+inDAGI, CN+inDALI) perform poorly. In Fig. 4d, we note that (CN+inDAGI) outperformed other metrics, using a similar degree of edge removal. We have not shown the results for preferential attachment using global influence (or PAGI as discussed in Section “Proposed models”). This is because it has a very similar formulation (i.e., the PAGI score is equivalent to the PA score divided by a constant) and yields the same accuracy as the preferential attachment (PA) metric for both standard and synthetic networks. However, we have reported high accuracy for the combination of PAGI with the common neighbor (CN).

Prediction on synthetic networks. Figures 5 and 6 summarise the AUPRC results by implementing the link prediction metrics on the assortative and disassortative, high density (\( D = 0.15 - 0.19 \)) synthetic networks, respectively. (The corresponding AUC results for the synthetic networks have been shown in Supplementary section 1 and section 2. Furthermore, the AUPRC results for the low-density networks (with density \( D = [0.01, 0.05, 0.1] \)) have been shown in Supplementary Section 3).
Assortative networks analysis. Like in the standard assortative networks (discussed in Section “Prediction on standard networks”), the local similarity measures, namely CH2-L2, CH2-L3, Jaccard, HDI, and CAR, once again exhibit high accuracy for both random as well as similar degree edge sampling (see Fig. 5). The superior performance of local similarity-based metrics that rely on shared neighbors suggests that the similar degree nodes in the assortative networks are strongly interconnected.

Disassortative networks analysis. Figure 6 shows the AUPRC scores for the disassortative networks. CH2-L3, DALI, and DAGI outperform other LP metrics for the highly disassortative network, followed by combined dissimilarity-based models (with low CN contribution). Unlike the assortative networks, most similarity-based models (along with combined models) show very low AUPRC performance. Interestingly, in the case of similar degree edges sampling, similarity-based models of (inDALI and inDAGI) show higher AUPRC performance than preferential attachment extension and dissimilarity metrics.

Key observations from the prediction models. First, we find that CH2-L2 and CH2-L3 perform consistently well for assortative and disassortative networks. This is because, unlike local similarity-based LP metrics, these metrics explore larger neighborhoods around the nodes of interest. Second, we report that the combined LP metrics of common neighbors (CN) and inDAGI exhibit an improvement over local similarity-based metrics in standard assortative networks (see Fig. 4b). This shows that a combined influence of local and global neighborhoods can often be a better strategy for standard assortative networks. Also, since similar degree nodes tend to group together in assortative networks, the local similarity-based metrics that rely on common neighbors (such as AA, RA, CN, etc.) can emerge as good choices for LP metrics. Third, for synthetic networks, since nodes in assortative networks tend to have connections with similar degree nodes, the AUPRC of the local similarity-based metrics decreases with network assortativity. We find low (positive) assortativity to be associated with the improved performance of the dissimilarity metrics (DALI, DAGI, etc.) as well as the preferential attachment combined metrics (PAGI, PALI) with CN. This is most evident in case of random sampling, as depicted in Figs. 5a,c and 6a,c. Fourth, we find CN + inDAGI to perform well in case of similar...

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**Figure 4.** Performance of the LP models (i.e., the AUPRC values) implemented on the standard networks using random edges sampling (a, c), descending Similar Degree Edges (b, d). Assortative networks are shown in (a, b), and disassortative networks are shown in (c, d). Refer to Table 1 for details on the assortativity group each standard network belongs to. Red bars refer to standard LP models, blue bars refer to dissimilarity-based metrics along with their extensions and green bars refer to similarity-based metrics along with their extensions.
degree node removal and disassortative networks. A combination of low CN and high inDAGI has proven to be effective, showing the importance of the global influence of nodes (measured by their degree) in determining their network connections. Two nodes seem more likely to be connected if they have similar degrees rather than neighbor-based similarity. Overall, we intuit that the combined models can be particularly useful for predicting connections in real-world networks (namely, biological, social, and technological networks) which are often disassortative in nature.

The effect of edge sampling. The standard random sampling of edges guarantees that the type of edge to be selected is not biased and that no specific global or local feature of the network is targeted. Therefore, we observe that the assortativity coefficient $r$ for the different standard networks maintained a close distance (small error ranges) with future sampling. Figure 7a reports the values of the assortativity coefficient $r$ for each of the given networks such that we removed (20%) of the edges 30 times.

On the other hand, similar degree edges removal (or sampling) guarantees that the edges to be selected are determined by the similarity in degree of the two nodes at the specified edge. It is noteworthy that the two nodes are not necessarily similar in neighbors (as in common neighbors) or other standard local similarity-based metrics. Intuitively and after the removal of similar degree edges, the resultant network would have less assortativity coefficient $r$ value, as can be seen in Figure 7b, where we report $r$ for standard networks after (20%) edges removal of the top similar degree ones, and for 30 times. We observe that the assortativity coefficient $r$ is effected in a descending fashion, and gradually converts the network into disassortative networks. We can assume that the opposite is true such that the removal of dissimilar degree edges will increase the assortativity level.

Overall, the random and similar edge removal techniques are employed to demonstrate two aspects of the link prediction analysis. The random edge removal approach eliminates links without bias. Thus, the prediction occurs on probe networks very similar to the original networks (see Fig. 7a and refer to Section “Formal problem setting” for details on training and probe networks). On the other hand, as depicted in Fig. 7b, the similar edge removal scheme challenges the predictors by altering the assortativity coefficients of the training networks. In
Figure 6. Performance of the LP models (i.e., the average AUPRC values) implemented on the high density, disassortative synthetic networks of size (100, 200, 250, 500, and 1000 nodes), 500 networks each. Table 2 depicts the high and less disassortativity coefficient $r$ ranges, respectively. Red bars refer to standard LP models, blue bars refer to dissimilarity-based metrics along with their extensions and green bars refer to similarity-based metrics along with their extensions.

Figure 7. The average and standard deviation of the assortativity level of each of the standard networks after removing 20% of network edges at random for 30 separate runs. Edge removal of each run is implemented independently of other runs.
other words, the training is carried out on a disassortative network and the metrics are used to predict the possibility of links and degree-similar nodes.

**Exploratory analysis of $\alpha$ and the combined model.** To better understand the impact of ($\alpha$) in the combined model, we reported the results that were collected from different simulations by using $\alpha = [0, 0.05, 0.1, \cdots , 1]$. We have already noticed the effect that $r$ plays along with the two edge sampling approaches (Random, and SDE). Also, we have demonstrated that the combined model is constructed by two terms: common neighbors and either one of the proposed metrics of 1) similarity models (inDAGI, inDALI), 2) preferential attachment extension models (PAGI, PALI) and 3) dissimilarity models (DAGI, DALI). As the assortativity coefficient value $r$ increases, the combined model tends to incorporate similarity metrics, i.e., CN + inDAGI and CN + inDALI, putting high weight on the CN model. Similarly, as the assortativity coefficient value $r$ decreases (to disassortativity level), the combined model tends to incorporate dissimilarity metrics, i.e., (CN + DAGI and CN + DALI), putting high weight on the dissimilarity metrics.

However, in order to find the effect of ($\alpha$) on the combined model at high $r$, we particularly report additional analysis on the best similarity metrics of (CN + inDAGI) by considering a range of discrete values of $\alpha$. If $\alpha = 0$, we consider 100% of the inDAGI value. Likewise, if $\alpha = 1$, then we consider 100% of the CN value, see Fig. 8. We count the average AUPRC value (with its standard deviation) for the 500 networks of size 200 nodes, using high assortative, and less assortative networks with random edges sampling. Also, we count average AUPRC values (with their standard deviation) using similar degree edges. These three models show the best AUPRC performance for CN + inDAGI. We notice that there is a marginal contribution of the (inDAGI) of 10% in the high and less assortative networks (refer to Fig. 8).

**Discussions**

In this paper, we have attempted to identify metrics for link prediction based on network assortativity. As part of this task, we introduced an approach that generates networks of varying assortativity levels and proposed three different models for link prediction that measure the different link properties. These models are local dissimilarity-based models (DAGI, DALI), extended preferential attachment models (PAGI, PALI) and finally similarity models (inDAGI, inDALI). These link prediction models are then combined with the most standard local similarity-based metric of common neighbors to form the weighted combined models. We have also introduced an algorithm to generate assortative and disassortative networks of varying levels. We carry out extensive simulation experiments to demonstrate the contribution of several standard local neighborhood-based

![Image](https://www.nature.com/scientificreports/)
metrics along with the common neighbors in the composition of the accurate link predictors for most cases. Our dissimilarity-based models outperform most of the other models in link prediction. Although there is less association between the assortativity coefficient of the network \( r \) and the link prediction models, we were able to show high prediction accuracy of specific models for assortative and disassortative networks.

This work opens up a few interesting research directions. First, we shall employ the proposed similarity and dissimilarity metrics to predict links in large-scale social and biological networks. In addition to the assortativity levels, this analysis will take into account other node and link labels as well as the directionality of links. Second, Fig. 8 shows the effect of varying the weighing parameter \( \alpha \) on the overall accuracy of link predictions. Going forward, we intend to leverage these findings to infer general rules that will inform the selection of the link prediction metrics contingent on the assortativity and relevant feature information (of the nodes and links). Specifically, the rules will be mined using adaptive optimization algorithms that will learn the right \( \alpha \) that maximizes accuracy for myriad assortativity levels as well as other topological properties of networks. Moreover, we shall delve deeper into the relationship between graph density and network assortativity in Algorithm 1. This will involve finding a range of graph densities for which it is feasible to generate networks of a prespecified assortativity coefficient. This effort will be particularly useful to the community of social and biological network researchers who need to analyze and make inferences from diverse families of partially available network datasets.

**Data availability**

The datasets used, generated, and/or analyzed during the current study along with the associated code are available in the GitHub repository (https://github.com/almusawia2/Identifying-Accurate-Link-Predictors-based-on-Assortativity-of-Complex-Networks/).

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**References**

1. Ben-Naim, E., Frauenfelder, H. & Toroczkai, Z. *Complex Networks* Vol. 650 (Springer, 2004).
2. Chen, H., Li, X. & Huang, Z. Link prediction approach to collaborative filtering. in *Proceedings of the 5th ACM/IEEE-CS Joint Conference on Digital Libraries (JCDL05)*, 141–142 (IEEE, 2005).
3. Qi, Y., Bar-Joseph, Z. & Klein-Seetharaman, J. Evaluation of different biological data and computational classification methods for use in protein interaction prediction. *Proteins Struct. Funct. Bioinform.* 65, 490–500 (2006).
4. Kovacs, I. A. et al. Network-based prediction of protein interactions. *Nat. Commun.* 10, 1–8 (2019).
5. Chuan, P. M. et al. Link prediction in co-authorship networks based on hybrid content similarity metric. *Appl. Intell.* 48, 2470–2486 (2018).
6. Lu, Y., Guo, Y. & Korhonen, A. Link prediction in target-tranger interactions network using similarity indices. *BMC Bioinform.* 18, 1–9 (2017).
7. Abbas, K. et al. Application of network link prediction in drug discovery. *BMC Bioinform.* 22, 1–21 (2021).
8. Ghosh, P. et al. Principles of genomic robustness inspire fault-tolerant wsn topologies: A network science based case study. in *IEEE international conference on Pervasive computing and communications workshops (PERCOM workshops)*, 160–165 (IEEE, 2011).
9. Kamapantula, B. K. et al. Leveraging the robustness of genetic networks: A case study on bio-inspired wireless sensor network topologies. *J. Ambient Intell. Hum. Comput.* 5, 323–339 (2014).
10. Niazi, A., Raj, M., Di Francesco, M., Ghosh, P. & Das, S. K. Deployment of robust wireless sensor networks using gene regulatory networks: An isomorphism-based approach. *Pers. Mob. Comput.* 13, 246–257 (2014).
11. Roy, S., Ghosh, P., Ghosh, N. & Das, S. K. Transcriptional regulatory network topology with applications to bio-inspired network- ing: A survey. *ACM Comput. Surv.* https://doi.org/10.1145/3468266 (2021).
12. Lu, L. & Zhou, T. Link prediction in complex networks: A survey. *Physica A* 390, 1150–1170 (2011).
13. Martinez, V., Berzel, F. & Cubero, J.-C. A survey of link prediction in complex networks. *ACM Comput. Surv. (CSUR)* 49, 1–33 (2016).
14. Brandes, U. A faster algorithm for betweenness centrality. *J. Math. Soc.* 25, 163–177 (2001).
15. Newman, M. *Networks: An Introduction* (Oxford University Press, 2018).
16. Newman, M. E. Mixing patterns in networks. *Phys. Rev. E* 67, 026126 (2003).
17. Noldus, R. & Van Mieghem, P. Assortativity in complex networks. *J. Comput. Netw.* 3, 507–542 (2015).
18. Mayo, M., Abdelzaher, A. & Ghosh, P. Long-range degree correlations in complex networks. *Comput. Soc. Netw.* 2, 1–13 (2015).
19. Barabási, A.-L. & Albert, R. Emergence of scaling in random networks. *Science* 286, 509–512 (1999).
20. Freeman, L. C. Centrality in social networks conceptual clarification. *Soc. Netw.* 1, 215–239 (1978).
21. Zhou, J. et al. Generating an assortative network with a given degree distribution. *Int. J. Bifurc. Chaos* 18, 3495–3502 (2008).
22. Zachary, W. W. An information flow model for conflict and fission in small groups. *J. Anthropol. Res.* 33, 452–473 (1977).
23. Aaron Clauset, E. T. & Sainz, M. The colorado index of complex networks. https://icon.colorado.edu/ (2016).
24. Lusseau, D. et al. The bottlenose dolphin community of doubtful sound features a large proportion of long-lasting associations. *Behav. Ecol. Sociobiol.* 59, 396–405 (2003).
25. Xu, Z. & Harris, R. Exploring the structure of the us interstate passenger air transportation network: A weighted complex network approach. *Geojournal* 73, 87 (2008).
26. Newman, M. E. Finding community structure in networks using the eigenvectors of matrices. *Phys. Rev. E* 74, 036104 (2006).
27. Schaffer, T., Marbach, D. & Floreano, D. Genenetweaver: In silico benchmark generation and performance profiling of network inference methods. *Bioinformatics* 27, 2263–2270 (2011).
28. Albert, R. & Barabási, A.-L. Statistical mechanics of complex networks. *Rev. Mod. Phys.* 74, 47 (2002).
29. Leskovec, J. & Krevl, A. SNAP datasets: Stanford large network dataset collection. http://snap.stanford.edu/data (2014).
30. Rossi, R. & Ahmed, N. The network data repository with interactive graph analytics and visualization. in *Twenty-ninth AAAI Conference on Artificial Intelligence* (2015).
31. Read, K. E. Cultures of the central highlands, New Guinea. *J. Anthropol. Res.* 15, 396–405 (1954).
32. Wang, R. et al. Hierarchical connectome modes and critical state jointly maximize human brain functional diversity. *Phys. Rev. Lett.* 123, 038301 (2019).
33. Cannistraci, C. V., Alanis-Lobato, G. & Ravasi, T. From link-prediction in brain connectomes and protein interactomes to the local-community-paradigm in complex networks. *Sci. Rep.* 3, 1–14 (2013).
34. Varshney, L. R., Chen, B. L., Paniagua, E., Hall, D. H. & Chklovskii, D. B. Structural properties of the *Caenorhabditis elegans* neuronal network. *PLoS Comput. Biol.* 7, e1001066 (2011).
35. Simonis, N. et al. Empirically controlled mapping of the Caenorhabditis elegans protein–protein interactome network. Nat. Methods 6, 47–54 (2009).
36. Milo, R. et al. Superfamilies of evolved and designed networks. Science 303, 1538–1542 (2004).
37. van Dijk, R. E. et al. Cooperative investment in public goods is kin directed in communal nests of social birds. Ecol. Lett. 17, 1141–1148 (2014).
38. Cho, A. et al. Wormnet v3: A network-assisted hypothesis-generating server for Caenorhabditis elegans. Nucleic Acids Res. 42, W76–W82 (2014).
39. Lo, S., Monagan, M. & Wittkopf, A. Strongly connected graph components and computing characteristic polynomials of integer matrices in maple. (2006).
40. Sah, P. et al. Inferring social structure and its drivers from refuge use in the desert tortoise, a relatively solitary species. Behav. Ecol. Sociobiol. 70, 1277–1289 (2016).
41. Ahmad, I., Akhtar, M. U., Noor, S. & Shahnaz, A. Missing link prediction using common neighbor and centrality based parameterized algorithm. Sci. Rep. 10, 1–9 (2020).
42. Liben-Nowell, D. & Kleinberg, J. The link-prediction problem for social networks. J. Am. Soc. Inf. Technol. 58, 1019–1031 (2007).
43. Newman, M. E. Clustering and preferential attachment in growing networks. Phys. Rev. E 64, 025102 (2001).
44. Adamc, L. A. & Adar, E. Friends and neighbors on the web. Soc. Netw. 25, 211–230 (2003).
45. Zhou, T., Lu, L. & Zhang, Y.-C. Predicting missing links via local information. Eur. Phys. J. B 71, 623–630 (2009).
46. Jaccard, P. Etude de la distribution florale dans une portion des alpes et du jura. Bulle. Soc. Vaudoise Sci. Nat. 37, 547–579 (1901).
47. Chowdhury, G. G. Introduction to Modern Information Retrieval (Facet publishing, 2010).
48. Sorensen, T. Method of establishing groups of equal amplitude in plant sociology based on similarity of species content and its application to analyses of the vegetation on danish commons. (1948).
49. Ravasz, E., Somera, A. L., Mongru, D. A., Ólafsdóttir, Z. N. & Barabási, A.-L. Hierarchical organization of modularity in metabolic networks. Science 297, 1551–1555 (2002).
50. Leicht, E. A., Holme, P. & Newman, M. E. Vertex similarity in networks. Phys. Rev. E 73, 026120 (2006).
51. Ghasemian, A. et al. Stacking models for nearly optimal link prediction in complex networks. Proc. Natl. Acad. Sci. 117, 23393–23400 (2020).
52. Muscoloni, A., Abdelhamid, I. & Cannistraci, C. V. Local-community network automata modelling based on length-three-paths for prediction of complex network structures in protein interactomes, food webs and more. BioRxiv 346916 (2018).
53. Zhou, T., Lee, Y.-L. & Wang, G. Experimental analyses on 2-hop-based and 3-hop-based link prediction algorithms. Physica A 564, 125532 (2021).
54. Newman, M. E. Assortative mixing in networks. Phys. Rev. Lett. 89, 208701 (2002).
55. Newman, M. Networks: An introduction (Springer, 2010).
56. Boyd, K., Eng, K. & Page, D. Area under the precision-recall curve: point estimates and confidence intervals. In Joint European conference on machine learning and knowledge discovery in databases, 451–466 (Springer, 2013).
57. Saito, T. & Rehmsmeier, M. The precision-recall plot is more informative than the roc plot when evaluating binary classifiers on imbalanced datasets. PLoS ONE 10, e0118432 (2015).
58. Yang, Y., Lichtenwalter, R. & Chawla, N. Evaluating link prediction methods. Knowl. Inf. Syst. 45, 751–782 (2015).
59. Jiang, M., Chen, Y. & Chen, L. Link prediction in networks with nodes attributes by similarity propagation. http://arxiv.org/abs/1502.04380 (2015).
60. Johnson, S., Torres, J. J., Marro, J. & Munoz, M. A. Entropic origin of disassortativity in complex networks. Phys. Rev. Lett. 104, 108702 (2010).

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Author contributions
A.A. conceived the study, and A.A., S.R., and P.G. developed the methodology. A.A. developed the software and visualization. All authors wrote the paper and reviewed the manuscript.

Competing interests
The authors declare no competing interests.

Additional information

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