Revisiting the General Concept of Network Centralities: A Propose for Centrality Analysis in Network Science

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Abstract

Background: In network science, although different types of centrality measures have been introduced to determine important nodes of networks, a consensus pipeline to select and implement the best tailored measure for each complex network is still an open field. In the present study, we examine the node centrality profiles of protein-protein interaction networks (PPINs) in order to detect which measure is succeeding to predict influential proteins. We study and demonstrate the effects of inherent topological features and network reconstruction approaches on the centrality measure values.

Results: PPINs were used to compare a large number of well-known centrality measures. Unsupervised machine learning approaches, including principal component analysis (PCA) and clustering methods, were applied to find out how these measures are similar in terms of characterizing and assorting network influential constituents. We found that the principle components of the network centralities and the contribution level of them demonstrated a network-dependent significancy of these measures. We show that some centralities namely Latora, Decay, Lin, Freeman, Diffusion, Residual and Average had a high level of information in comparison with other measures in all PPINs. Finally, using clustering analysis, we restated that the determination of important nodes within a network depends on its topology.

Conclusions: Using PCA and identifying the contribution proportion of the variables, i.e., centrality measures in principal components, is a prerequisite step of network analysis in order to infer any functional consequences, e.g., essentiality of a node. Our conclusion is based on the signal and noise modeling using PCA and the similarity distance between clusters. Also, an interesting strong correlation between silhouette criterion and contribution value was found which corroborates our results.

Keywords:

Network science, Centrality analysis, Protein-Protein Interaction Network (PPIN), Clustering, Principal Components Analysis (PCA)

Introduction

Network modelling of diverse biological complex processes is a pervasive approach in contemporary biological researches [1]. This application of modelling is divided into two parts; moving from elements and their relationships toward a whole system and getting back from a whole into important and local structures among the system [2, 3]. For instance, researchers experimentally or computationally collect information about a set of proteins and the interactions between them to reconstruct a protein-protein interactions network (PPIN). Then, they define an importance or an influence concept such as degree centrality (hub) in order to identify influential structures among the given PPINs which contains individual nodes, edges and subgraphs such as cliques or pathways. Not only in the biological networks but also in all types of networks such as social or literary, finding center of the network is a chief question which is called centrality analysis [4, 5]. By this analysis, the influential parts of the networks which are useful to predict the behavior of the whole network could be determined. Any change of these parts could be followed through the system and may disturb the whole [6]. Various centrality metrics or measures have been defined during last forty years, mostly in the context of social network analyses [5, 7]. Whilst basic concepts of the graph theory, including number of links, distances, eigenvalues and local structures, are represented in almost all centrality analyses, some of these measures are very popular and mostly used in diverse context using specific tools [8]. There is much research in biological areas which has studied the correlation of the lethality and essentiality with some centrality measures [4, 9-16]. Additionally, these measures have been applied in many social and epidemiological research studies, examples include predicting the details of information controlling or disease spreading within a specific network in order to delineate how to effectively implement target marketing or preventive healthcare [7, 17-21].

However, the selection of the appropriate metric for given networks is still an open question. Which one is better in translating center of the real networks? Do all of them independently highlight the central network elements and encompass independent information or are they correlated? Are computations of all these measures meaningful in all different network models or do they depend on the network architecture and the logic of the network modeling?

In 2004, Koschützki and Schreiber compared five centrality values in two biological networks and showed different patterns of correlations between centralities. They generally concluded that all Degree, Eccentrecity, Closeness, random walk Betweenness and Bonacich's Eigenvector centralities should be considered and could be useful in various applications without explaining any preference among them [22]. Two years later, they showed the independence behavior of centrality measures in a PPIN using 3D parallel coordinates, orbit-based and hierarchy-based comparison [23]. Valente et al. examined the correlation

between the symmetric and directed versions of four measures which are commonly used by the network analysts. By comparing 58 different social networks, they concluded that network data collection methods change the correlation between the measures and these measures show distinct trends [24]. Batool and Niazi also studied three social, ecological and biological neural networks and they concluded the correlation between Closeness-Eccentricity and Degree-Eigenvector and insignificant pattern of Betweenness. They also demonstrated that Eigenvector and Eccentricity measures are better to identify influential nodes [25]. In 2015, Cong Li et al. further investigated the question of correlation between centrality measures and introduced a modified centrality measure called *m*th-order degree mass. They observed a strong linear correlation between the Degree, Betweenness and Leverage centrality indices within both real and random networks [26].

Similar in methodology but different in networks, all these studies attempted to quantify correlations between different well-known centrality measures. In contrast, in this study, we develop a formal methodology (besides the correlation analysis) for centrality comparison and usage within network analysis. We comprehensively compared 27 distinct centrality measures applied to 14 small to large biological and random networks. All biological networks are PPIN among same set of proteins which are reconstructed using a variety of computational and experimental methods. We demonstrate how the ranking of nodes (or edges) using these measures depends on the network structure (topology) and why this network concept i.e. centrality requires renewed attention.

Materials and methods

The workflow of this study is schematically presented in Fig. 1. Our workflow starts by constructing and retrieving networks, followed by network global analysis. The centrality analysis and comparing the centrality measures using machine learning methods were the next main steps.

Reconstruction of the networks

In this study, a UniProtKB reviewed dataset [27] was used to retrieve proteins in *Saccharomyces cerevisiae* (6721 proteins). UniProtKB accessions were converted to STRING using the STRINGdb R package, which resulted in 6603 protein identifiers (3rdSep2016). Interactions among proteins were extracted based on the STRING IDs . In the 2017 edition of the STRING database the results of these interactions are structured in a way to provide maximum coverage; this is achieved by including indirect and predicted interactions on the top of the set. Each result is assigned a score which represents to what extent the interactions are biological, meaningful, specific and reproducible according to the supporting evidence. The STRING database is comprised of channels that are known as "evidence channels", which are formed according to the origin and type of evidence [28]. In this study, 13 evidence channels indicating PPIN of yeast are present: co-expression, co-expression-transferred, co-occurrence, database, database-transferred, experiments, experiments-transferred, fusion, homology, neighborhood-transferred, textmining, textmining-transferred and combined-score.

For the purpose of comparison with real network behavior, a null model network was generated. The null network is the Erdős–Rényi model [29] and was generated using the igraph R package [30]. The generated null network was created with a size similar to the yeast reconstructed PPIN in order to have a more fair comparison.

Fundamental network concepts analysis

To understand the network structure, we reviewed various network features using several R packages [31-33]. First of all, we inquired the score distribution of different channels and whether there is any significant correlation among the scores for each PPI. Then the networks were made readable in the R environment. The network density, clustering coefficient, network heterogeneity, and network centralization properties of the network were calculated. The number of connected components and graph diameter for each network were also computed. Then, the power-law distribution was determined by computing α and r values.

Centrality analysis

For this research study, we are only considering undirected, loop-free connected graphs according to the PPIN topology. For centrality analysis, the following 27 centrality measures were selected: Average Distance [34], Barycenter [35], Closeness (Freeman) [7], Closeness (Latora) [36], Residual closeness [37], ClusterRank [38], Decay [39], Diffusion degree [40], Density of Maximum Neighborhood Component (DMNC) [41], Geodesic K-Path [42, 43], Katz [44, 45], Laplacian [46], Leverage [47], Lin [48], Lobby [49], Markov [50], Maximum Neighborhood Component (MNC) [41], Radiality [12], Eigenvector [51], Subgraph scores [52], Shortest-Paths betweenness [7], Eccentricity [53], Degree, Kleinberg's authority scores [54], Kleinberg's hub scores [54], Harary graph [53] and Information [55]. All these measures can be calculated for undirected networks in a reasonable time. These measures were established using the centiserver [8], igraph [30] and sna [56] R packages. We assorted the centrality measures into five distinct parts including Distance-, Degree-, Eigen-, Neighborhood-based and miscellaneous groups depend on their logic and formulas (Table 1).

Unsupervised machine learning analysis

We used principal components analysis (PCA) [57] as a key step to understanding the influence of a centrality measure within a network. PCA was performed on normalized computed centrality measures. In order to visualize the result of the PCA and clustering analysis, we applied some R packages [58-60]. After normalization, we examined whether the centrality measures in all networks can be clustered by performing the clustering tendency procedure. This required calculating the Hopkins' statistic values and using VAT (Visual Assessment of cluster Tendency) plot. We applied the clustering validation measures to get the best number of clusters using some other R packages [61, 62]. This will provide silhouette scores in different kinds of clustering methods such as hierarchical, k-means, and PAM (Partitioning Around Medoids). Finally, distance metrics could be formed using the Pearson correlation in order to find any relationships between two centrality measures in each PPIN. To be able to compare the clustering results in the various PPINs, the Jaccard similarity index [63] was used relying on the similarity metrics of the clustering results [64].

Results

Evaluation of network global properties and centrality analysis

By importing the same set of protein names, the 13 different PPINs were extracted from the STRING database using different predictive channels. Note that the PPI scores derived from the neighborhood channel of yeast were all zero. The distribution of STRING links and edge scores are presented for each channel (Fig. 2). As shown in the figure, all these channels independently identify an interaction for each protein pair with specified score or weight. Therefore, we have 13 dissimilar network prediction methods and the following results are independent from each other. The independency between evidence channels is also shown in Fig. 3 by a pairwise scatterplot and Pearson's r correlation coefficient. The scores are not correlated and most of the coefficients are around the zero.

In the following, we consider the evidence channel scores as edge weights in each PPIN. So, along with the Erdos-Renvi null network, the 14 networks were utilized in order to establish an examination of centrality measures. Note that the giant component of each network was calculated to compute corresponding measures. To make sense of the networks and their giant component's structure, some fundamental network concepts were computed as shown in Table 2. The homology, fusion, co-occurrence and database networks contain high numbers of unconnected components. Except the homology network which has the smallest giant component, the density of all networks are between 0.01-0.05, as was expected as real network are typically sparse. The network diameter of the fusion, co-occurrence, database and coexpression are one order of magnitude greater than the others. All of the PPINs except homology network have high r correlation values to the power-law distribution with the diverse alpha coefficients. The high value of the average clustering coefficients of the database and homology indicates the modular structure of these networks. Most of the PPINs have the high value of the heterogeneity and network centralization compared with the null network. The degree distribution and clustering coefficients for the networks are also plotted in Fig. 4 and Fig. 5 respectively. Except the homology network, all the degree distribution are left-skewed similar to the scale-free networks. The scale-free property of each network was visualized (See Supplementary file 1).

In the next step, the 27 centrality measures of nodes were computed in all of the 14 networks (Data is available upon request). As shown in Fig. 6, beside the observed correlation between centrality measures in both combined-score and Erdos-Renyi networks, their distribution and the level of associations showed the vast diversity among all five centrality groups especially in Distance-, Neighborhood-based and Miscellaneous groups. The Eccentricity and Leverage centralities which comes from Distance- and Degree-based groups respectively showed the independent shape of distribution compared to the others. This pattern is repeated in all PPINs to some extent (See Supplementary file 2). However, the multimodal distributions

of centrality measures are presented in the random network but not in the real networks. Also, the value of the associations on average are interestingly higher in the null network than the PPINs.

Dimension reduction and clustering analysis

In the next step, PCA-based dimensionality reduction was used not only to minimize information loss, but to reveal which centrality measures contain the most relevant information and it can effectively identify important or influential nodes in networks. As illustrated in Fig. 7, the profile of the distance to the center of the plot and their directions are mostly similar except for the homology which is similar to the random network. The contribution values of each centrality measure are shown according to rank order in Table 3, based on their corresponding principal components. A similar pattern of the contribution of centrality measures could be observed in all real networks even in homology networks compared with the random null network (See Supplementary file 3). The Closeness centrality Latora is the main contributor of the principal components on average and it showed that it contains more information compared with the other centrality measures in PPINs. In contrast, other well-known centralities i.e. Betweenness and Eccentricity revealed a low contribution value in all PPINs similar to the null network and lower than random threshold. On the contrary, the Degree displayed moderate levels of contribution in all real networks whilst it is the fourth rank of random network contributors. Although, the overall patterns of contributions are similar, all PPINs exhibited this special fingerprint of centrality importance.

Finally, by performing unsupervised categorization, we aim to cluster centrality measures found on the computed values in PPINs. First, by performing the clustering tendency procedure, we find that this data set is clusterable. The distance metrics formed with Pearson correlation coefficients using the normalized centrality measures can be seen in Fig. 8. Then, we validate the measures to obtain the silhouette scores for different clustering methods such as hierarchical, k-means, and PAM (Supplementary files 4 & 5). Considering these scores, the hierarchical clustering algorithm was applied to categorize the standardized centrality measure results in each PPIN. The output of applying the clustering algorithms and the corresponding number of clusters is shown in Table 4. Using the hierarchical algorithm based on Ward's method [65], which relies on determining the adequate number of clusters, the centrality measures are clustered in each PPINs (Fig. 9). The number of clusters, the distance between centralities and the centrality composition in all 13 PPINs displayed independent outcomes and inconstant behavior of each centrality to rank nodes within networks. For better visualization, Table 5 displays the pairwise Jaccard similarity index values for each network pair. The lowest values are related to the homology, neighborhood-transferred and co-occurrence PPINs and among the genome context prediction methods, fusion PPIN is more associated to the others. The high similarity between co-expression and co-expression-transferred was expected but

the similar centrality clustering of the database with both aforementioned PPINs and combined-score with textmining-transferred are remarkable.

Interestingly, it is also illustrated that silhouette scores (calculated from the centralities in each cluster) are related to the contribution value of the corresponding centrality measure. Where there is a high silhouette value, a high contribution value is observed, however, a high contribution value will not always mean a high silhouette value. Fig. 10 shows the relationship between the silhouette scores and contribution values of each centrality measure in the combined-score PPIN. In panel A, each color represents a cluster of centrality measures. As shown in the figure, Latora, Radiality, Residual, Decay, Lin, Leverage, Freeman and Barycenter centralities have been gathered together in the same cluster where the corresponding silhouette scores are all at a high level except the Leverage's score, while the average silhouette score is around 0.66 in that cluster. On the other hand in panel B of Figure 8, we can find the Leverage's contribution value that is under the threshold line and placed in the group with the least amount of contribution. Centralities named Lobby index, ClusterRank, Laplacian, MNC, Degree, Markov, Diffusion degree, Kleinberg's hub, Eigen vector, Authority score, Katz centralities fell in the same group where all the silhouette scores are more than the average equal to 0.61 and in the same way, their corresponding contribution values are at high levels, too. On the other hand, we observe that Shortest path Betweenness (which is in a separated cluster) and Geodesic k path, Subgraph and DMNC (which are all in one cluster) had silhouette values lower than the average 0.03. The silhouette values influence the contribution values, which are the lowest. In all other PPINs, the same relationship between silhouette scores and contribution values was observed as shown in Supplementary files 3 & 5.

Discussion

Network biology is a major part of the relatively young biological research approach called systems biology. Topological analysis of biological networks is a basic approach for understanding complex behaviors of a cell, which can be classified into three categories, which are collective behaviors, subnetwork behaviors and individual behaviors (prioritizing of important nodes by centrality index). It consists of various computational methods and algorithms in order to infer biological conclusions such as biomarker discovery or drug design and repurposing which are mostly at the stage of introduction or development. However, there is no suitable benchmark for network biologists and this could result in inconsistent and non-reproducible outcomes. Therefore, reconsidering the computational methods towards recommending a standard protocol is needed and inevitable.

One of the unsolved problems in network biology and PPIN analysis is sorting the proteins with respect to their influence based on the centrality-lethality rule [66]. Commonly, high Degree value of a protein in PPIN, i.e., a hub is translated to the extent of its corresponding impact on biological functions [67]. However, dependency of Degree and vitality role of proteins has been riddled with some controversy. In a pioneering work, the authors claimed that the high value of degree centrality in the yeast PPIN is likely to be the main protein for the yeast survival [66]. In another study, this rule was re-examined in three distinct PPINs of three species which confirmed the essentiality of central proteins for survival [11]. Similar results were reported for gene co-expression networks of three different species [68] and for metabolic network of Escherichia coli [69, 70]. Ernesto Estrada generalized this rule to six other centrality measures. They showed that the subgraph centrality is the best to find important proteins, and using these measures performed significantly better than a random selection [13]. However, He and Zhang proposed that the relationship between hub nodes and essentiality is not related to network architecture. This finding is explained with the involvement of probability theory [67]. Also, regarding modular structure of PPIN, Joy et al. concluded that Betweenness centrality is more likely to be essential than Degree [71]. The representative power of Betweenness as a topological characteristic was also mentioned in mammalian transcriptional regulatory networks which was clearly correlated to Degree [72]. Recently, it has been shown that existing hub nodes in a network do not have a direct relationship with prognostic genes across cancer types [73].

Some works revisited the centrality notion and focused on repairing this rule by adding new definitions and measures [74]. Tew and Li demonstrated functional centrality and showed that it correlates more strongly than pure topological centrality [75]. Recently, Peng et al. introduced localization-specific centrality and claimed that their measure is more likely to be essential in different species [76]. Khuri and

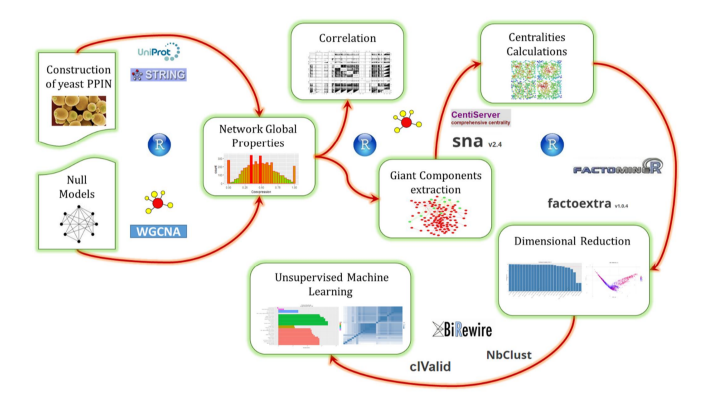
Wuchty introduced minimum dominating sets of PPIN which are enriched by essential proteins. They described that there is a positive correlation between degree and lethality [10]. One of the proposed solutions is to apply functional methods in this context according to the type of biological networks to be analyzed.

In this study, we worked on updated versions of PPINs which were built on different approaches and as it is shown, while the approaches have different properties, all of them were undirected in a same biological category. In addition to improve our network set compared to previous studies, a large number of centrality measures was used to explore the proximity and preference of the measures. We demonstrated that a similar profile of centrality ranking in PPINs i.e. Latora, Barycenter, Diffusion degree, Freeman, Residual, Average, Radiality centralities plays an important role in detecting the center of PPIN models. We inferred that the rationale and logic of networking in each network dictates which centrality indices should be taken. Also, our results demonstrated the relationship between contribution value derived from PCA and silhouette width as a cluster validity index. We are currently working on producing several built-in functions for various networks to recognize the most appropriate network-dependent centrality measure.

However, from this research, we first reasserted that the architecture and global properties of a network can impact on the central component detection and that the center of network would be differed according to network inherent topology. However, the main step before applying centrality measures for biological inference is identifying their ability to demonstrate the distinction between nodes or edges. In other words, we consider whether a centrality measure has enough information to be used for ranking the network components. Similar to other data mining studies, data reduction and low-dimensional projection help to extract interesting features i.e. centralities and corresponding relationships. Thus, in order to quantify connectivity in biological networks, we recommended that before calculating any centrality measures, an analysis of principal components of these measures should be carried out. The analysis of principal components is necessary to find out which measures have the highest contribution values, i.e., which measures carry the most important information.

Figure legends

Fig. 1. A workflow for studying the centrality measures. This follows the construction of the yeast PPIN relying on different kinds of evidence channels as well as the generation of a null network. The workflow contains a comparison of the behavior of several centrality measures using machine learning methods such as principal components analysis and clustering procedures.



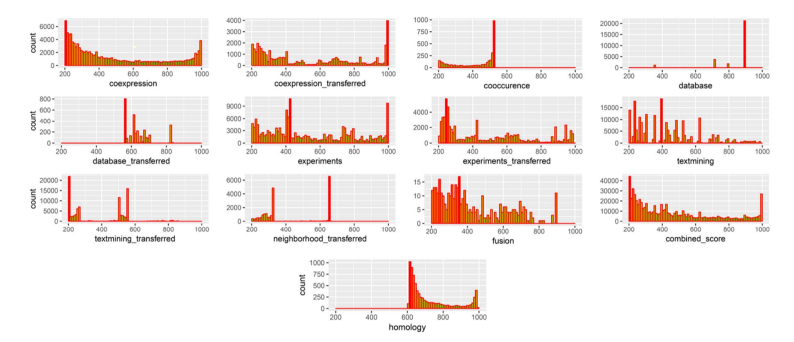


Fig. 2. A graphical representation of the distribution of the evidence channel scores presented in STRING database for the yeast PPIN. The color spectra from green to red indicates low to high values.

Fig. 3. Pairwise scatterplot between the evidence channel scores. The Pearson's r correlation coefficients are also indicated between the evidence channels. The distributions of scores in each evidence are presented at the diameters of the figure.

neighb_tr	fusion	cooccur.	homol.	coexp.	coexp_tr	experim.	xperim_t	data.	data_tr	text.	text_tr	mb_sco	
0 - 0 - 0 -	0.05	0.04	-0.01	0.01	0.27	-0.05	0.12	0.04	-0.01	-0.05	0.24	0.15	neigup_tr
		0.07	0	0	0.01	0	0	0.02	0	0.01	0.02	0.02	TUSION
	177		0.5	0.02	0.02	0.02	0.03	0.04	0.01	0.07	0.06	0.04	cooccur.
	L.:			0.03	0	0.04	0.04	0.03	0.03	0.14	0.08	0.05	nomol.
	35				0.4	-0.03	0.08	0.12	-0.01	-0.04	-0.06	0.37	coexp.
		1				0.03	0.38	0.22	0.02	-0.02	0.22	0.36	uTdxaoo
		1				[0.09	0.14	0.04	0.08	-0.07	0.52	r experim.
			Ē					0.2	0.12	0.02	0.28	0.33	Tuuedx
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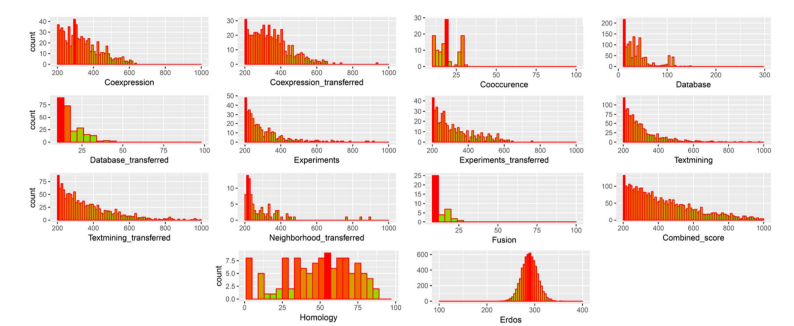
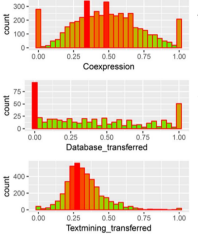
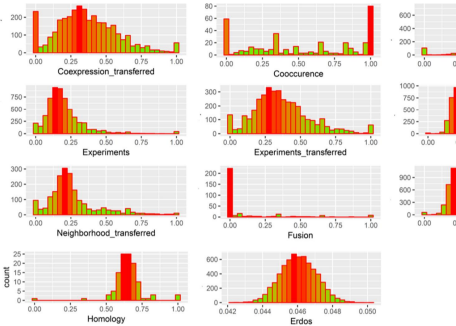
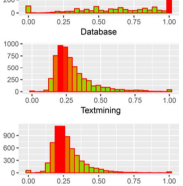


Fig. 4. A graphical representation of the degree distributions in each reconstructed PPIN and the generated null network. The color spectra from green to red indicates low to high values.

Fig. 5. A graphical representation of the clustering coefficient distribution in each reconstructed PPIN and the generated null network. The color spectra from green to red indicates low to high values.

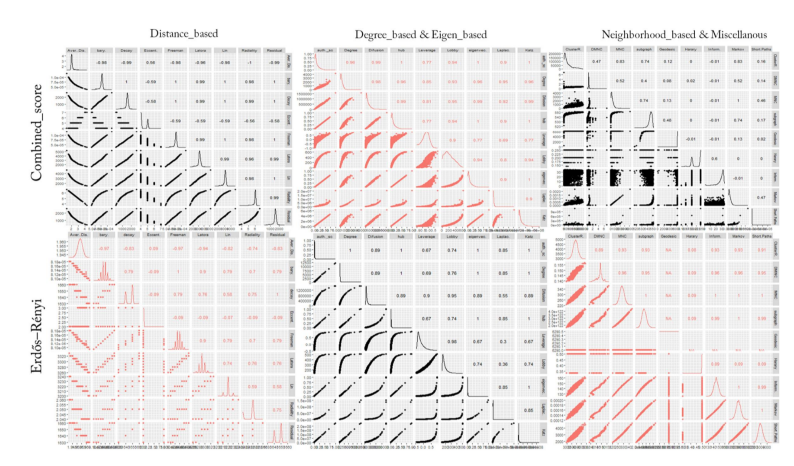






Combined_score

Fig. 6. Pairwise scatterplot between the centrality measures. This figure contains combined-score PPIN and the null network. In this figure, the r Pearson correlation coefficients between centralities beside the centralities distribution are also presented in both networks. For better representation, the scatterplot was divided into three parts corresponding to Table 1 groups. For all networks please refer to the supplementary file 2.



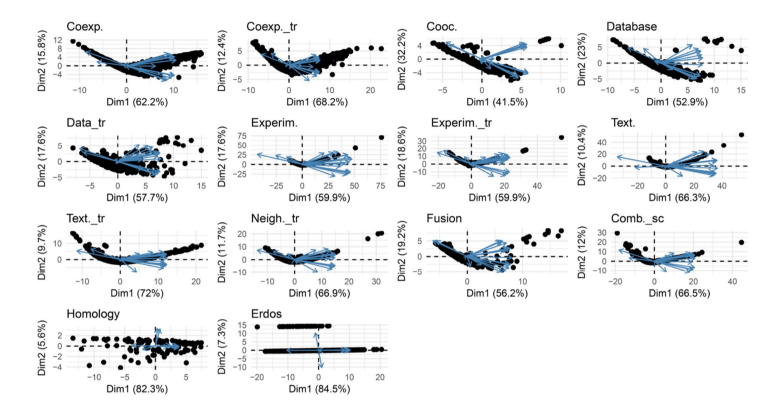


Fig. 7. Biplot representation of the centrality measures in each network. The first two dimensions of multivariate data which can be visualized graphically using PCA. In each plot, nodes are shown as points and centrality measures as vectors.

Fig. 8. Distance metrics. The dissimilarities between centrality measures in each PPIN are shown using colored distance measures found on r Pearson correlation coefficients. Colors from blue to red represents distances which vary from 0 to 2.

Coexpression	Coexpression_transferred	Cooccurence	Database
Database_transferred	Experiments	Experiments_transferred	Textmining
Textmining_transferred	Neighborhood_transferred	Fusion	Combined_score
Homology			

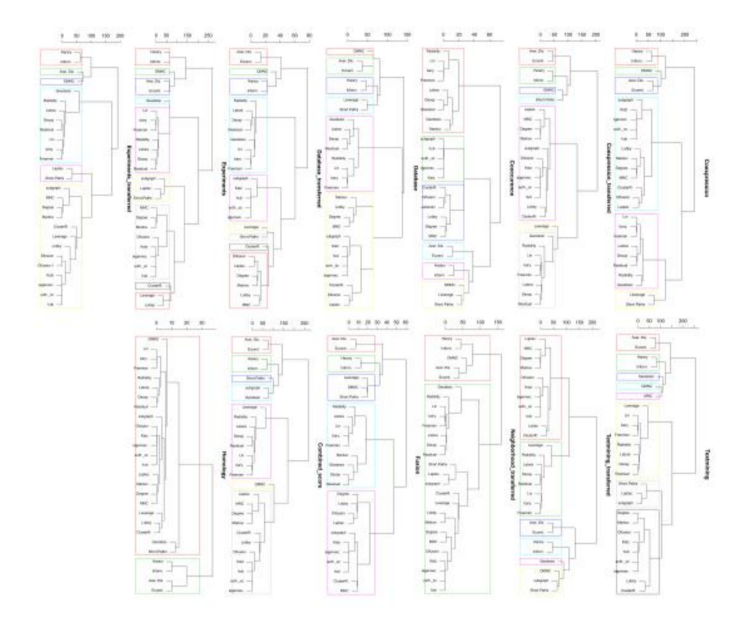


Fig. 9. The clustering dendrograms. In each dendrogram, the colored boxes show ensued clusters of centrality measures in each PPIN based on predefined distance threshold.

Fig. 10. (A) Clustering silhouette plot of the combined-score PPIN. The colors represent the six clusters of the centrality measures in this PPIN. The average silhouette width was equaled to 0.49. (B) Contribution values of centrality measures according to their corresponding principal components in this PPIN. The number of principal components stand on the network architecture was equal to 3. The dashed line indicates the random threshold of contribution. (C) Line plot between silhouette and contribution values. R value shows the result of regression coefficient analysis and p value has been computed found on Pearson correlation test.

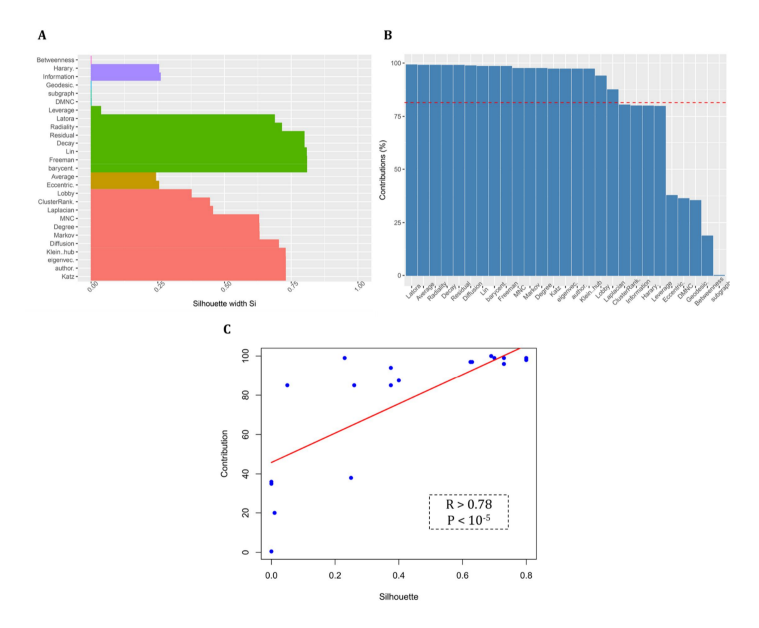


Table legends

Table 1.Centrality measures. These measures represented in five groups depend on their logic and formulas.

Distance_based	Degree-based	Eigen-based	Neighborhood-based	Miscellanous
Average Distance	Authority_score	Eigenvector centralities	ClusterRank	Geodesic K-Path Centrality
Barycenter	Degree Centrality	Katz Centrality (Katz Status Index)	Density of Maximum Neighborhood Component (DMNC)	Harary Graph Centrality
Closeness Centrality (Freeman)	Diffusion Degree	Laplacian Centrality	Maximum Neighborhood Component (MNC)	Information Centrality
Closeness centrality (Latora)	Kleinberg's hub centrality scores		Subgraph centrality scores	Markov Centrality
Decay Centrality	Leverage Centrality			Shortest-Paths Betweenness Centrality
Eccentricity of the vertices	Lobby Index (Centrality)			
Lin Centrality				
Radiality Centrality				
Residual Closeness				
Centrality				

Network Centralizatio n	0.34	0.06	0.05	0.06	0.41	0.06	0.26	0.77	0.30	0.11	0.35	0.56	0.62	0.01
Heterogeneit y	0.47	1.02	1.00	0.86	1.04	1.07	1.39	1.59	1.09	1.56	0.92	1.54	06.0	0.06
Average Clustering Heterogeneit Coefficent y	0.64	0.47	0.07	0.36	0.26	0.69	0.35	0.35	0.32	0.44	0.30	0.19	0.27	0.05
	0.00	0.61	0.91	69.0	0.72	0.56	0.79	0.81	0.75	0.82	0.66	0.82	0.63	0.01
α value r value (Power Law) (Power Law)	0.01	1.07	1.76	1.23	0.91	1.18	0.96	1.03	0.88	1.08	1.01	1.22	0.76	_1.095
Diamete r	'n	22	26	6	9	20	8	6	6	15	v	9	7	3
Density	0.43	0.02	0.01	0.02	0.04	0.01	0.03	0.02	0.04	0.01	0.02	0.01	0.05	0.05
Edges/Giant Component	2813	1653	789	2930	71236	26574	168364	163403	364816	195643	379341	211612	911413	911413
Nodes/giant Edges/Giant component Component	115	425	437	583	2004	2058	3607	3870	4207	5254	5896	6024	6292	6292
Properties Nodes Edges Connected s r s s s	648	209	222	10	1	100	4	1	1	27	1	2	2	1
Edges	7545	3275	1408	2975	71236	27766	168368	3870 163403	364816	5310 195676	379341	211613	911414	6292 911413
Nodes	2479	1221	1187	622	2004	2496	3614	3870	4207	5310	5896	6026	6294	6292
Networks Properties	Homology	Cooccurence	Fusion	Database_transferred	Neighborhood_transferred	Database	Coexpression_transferred	Experiments_transferred	Textmining_transferred	Coexpression	Textmining	Experiments	Combined_score	Erdős_Rényi

Table 2. Network global properties of all PPINs and the null network.

Table 3. Ranking of the contribution values based on the PCA for each network. The red to green highlighted cells represent the top to bottom ranked centrality measures in each network. The underlined ranking values are the centrality measure's contribution value which are less than random threshold of contribution.

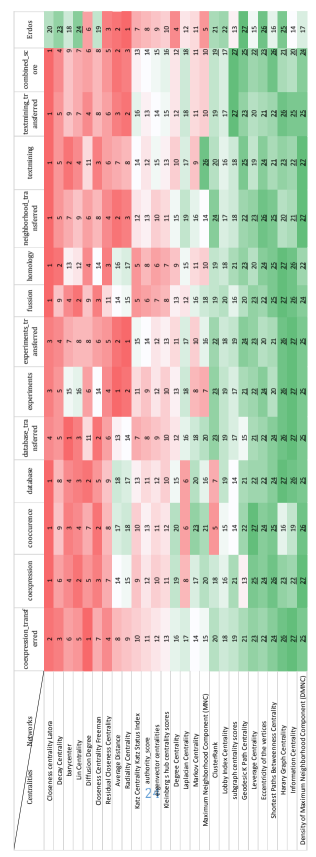
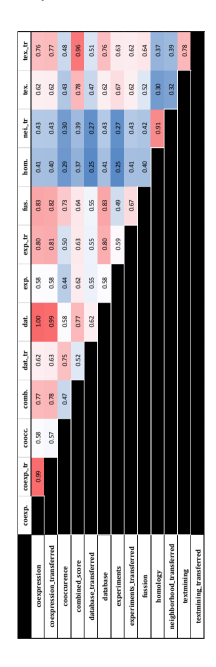


Table 4. Clustering information values for PPINs. The Hopkin's statistics threshold for clusterability is 0.05.

Network	Hopkins Statistic	Number of Clusters	Silhouette Average Value	
Coexpression	0.25	6	0.36	
Coexpression_transferred	0.21	7	0.33	
Cooccurence	0.18	6	0.55	
Database	0.24	6	0.33	
Database_transferred	0.20	9	0.32	
Experiments	0.21	9	0.31	
Experiments_transferred	0.16	6	0.43	
Textmining	0.24	8	0.28	
Textmining_transferred	0.20	6	0.35	
Neighborhood_transferred	0.26	2	0.39	
Fussion	0.16	5	0.48	
Combined_score	0.30	7	0.27	
Homology	0.23	2	0.46	

Table 5. Jaccard index coefficient values for PPINs. The values represent how similar the networks are, in terms of their clustering results. Where a value of 1.00 indicates an exact match, and values equal to 0 show dissimilarity.



Supplementary data

Supplementary file 1: Fitted power law distribution. The degree distribution of each network has been compared to the power law distribution in order to visualize the scale free property in the structure of each network.

Supplementary file 2: Scatterplots between groups of centralities. Each panel indicates scatterplots between centralities groups of two networks.

Supplementary file 3: Contribution values of centralities in each network. These values were computed based on the principal components. The red line shows the threshold used for identifying effective centralities.

Supplementary file 4: Clustering properties results. These properties include connectivity, Dunn and Silhouette scores. These scores suggest the sufficient clustering method by a specific number of clusters.

Supplementary file 5: Clusters silhouette plots. Each color represents a cluster and each bar with specific color indicates a centrality.

Supplementary file 6: Optimal number of clusters. The suitable number of clusters for hierarchical clustering method was computed using the average silhouette values.

Supplementary file 7: Visual assessment of cluster tendency plots. Each rectangular represents the clusters of the calculated results of the centrality measures.

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