



GENE ONTOLOGY BASED FUNCTIONAL ANALYSIS AND GRAPH THEORY FOR PARTITIONING GENE INTERACTION NETWORKS

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ABSTRACT

Exploring gene-disease associations improves the understanding of the underlying cause of the disease, which leads to further improvements in the diagnosis and treatment. Since genes that belong to the same topological, functional or disease module has an increased tendency of being involved in the same disease or phenotype, cluster analysis is the efficient approach to identify functionally similar genes. The aim of this work is to identify biologically relevant gene clusters using graph theory, which is an essential and influential scientific tool for modeling and exploring interconnected groups. The current work exhibits a computationally efficient algorithm for improving the performance of community detection in graphs using edge pruning techniques. The algorithm does not demand a priori judgment on the size of communities; it helps in automatic detection of gene communities with better performance. The optimized and streamlined approach is applied on cancer dataset and is compared and validated with standard clustering solutions using different validation measures.

KEYWORDS : Community detection, Gene Ontology, Similarity Matrix, Node Connection Score, Silhouette Index, Modularity



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INTRODUCTION

With the development of high-throughput devices and highly efficient technologies, the volume of biological data is increasing with a great pace. Traditional methods for gene mapping such as linkage analysis and association studies are used for identifying disease-associated genes¹. But these methods only associate a disease with chromosomal regions that usually contain tens or even hundreds of genes². Hence it is essential to develop efficient computational methods to identify the groups of strongly associated genes from a long list of candidate genes. Several methods have been proposed to address this problem by identifying, ranking and grouping candidate set of genes. Guilt-by-direct-association³ principle is used to facilitate the prioritization of candidate genes in many problems where disease susceptibility genes are ranked based on their relevance to the genes that are already known to be associated with the disease genes. Even though different clustering solutions are widely used for pattern extraction and for logically grouping homogeneous genes, clustering techniques with the ability to visualize is more commonly used by experts. This increases the understanding of the connections between datasets within groups and makes it simpler to comprehend the results of the clusters. Graph-based models represent complex interactions where data objects/observations are represented as nodes and relationships between observations as edges. The most significant and expressive feature of graphs in demonstrating real-time system is the community structure. The goal of the current approach is to discover communities or clusters by partitioning the vertices in a large semantic similarity gene network into different clusters based on neighborhood similarity. The algorithm uses gene ontology information to identify the functionally accurate clusters. The biological significance of these newly identified clusters is evaluated using gene-gene interactions.

Related Work

Different clustering solutions were developed so far to address the problem of grouping functionally similar genes⁴³⁻⁴⁵. Graph based clustering is the most realistic approach to demonstrate the biological reality. Numerous studies focus on topological characteristics of gene networks for extracting homogeneous groups. Jonsson *et al.*^{4,5} performed cluster analysis using PPI networks to conclude that disease genes belong to communities with a higher degree of connectivity and act as a key central node. Even though Cai⁶ infer similar findings in their studies, they could observe an inverse relationship between disease genes and clustering coefficient. Rahmani⁷ used network properties to predict disease genes in PPI networks. Two parameters were considered for analysis in this approach; the functional context of the proteins and the rank of target proteins with respect to the selected proteins set using ANOVA analysis. Zhang *et al.*⁸ studied online community detection for large, complex networks by proposing a node split based on the degree of the nodes for creating new clusters. The method has been applied to a set of real world network datasets and was compared with Agglomerative⁹ and Eigen vector

approach¹⁰ for community detection. Fast greedy algorithm¹¹ uses greedy method while walktrap algorithm¹² uses a random walk process to find the distance between two nodes. Both approaches used hierarchical agglomerative clustering for deriving meaningful communities. To maximize the density of the clusters obtained, Monte Carlo algorithm was proposed by Spirin *et al.*¹³ where the algorithm demands the cluster size as input parameter for detecting the protein complexes. Highly connected sub graph algorithm is used in several studies for deriving meaningful sub graphs from a connected network by Hartuvetal.¹⁴. A sub graph with n nodes are said to be a highly connected sub graph if more than $n/2$ edges must be removed in order to disconnect it. Przulj *et al.*¹⁵ used this approach in determining the protein complexes in PPI network. The algorithm partitions the graph by finding a minimum cut in the graph. A cut in a graph is a partition of nodes into two non-overlapping sets. Recursive process is initiated until highly connected components are obtained. Krogan *et al.*¹⁶ and Enright *et al.*¹⁷ used Markov clustering (MCL) algorithm for complex detection in PPI network. An adjacency matrix is computed for a graph $G = (V, E)$, $A|V| \times |V| = \{a_{ij}\}$ where $a_{ij} = 1$ if and only if v_i and v_j are neighbors is a matrix. High and low flow regions are identified and the process converges towards high flow regions of sub graphs known as protein complexes separated by no flow regions. Clique percolation (Adamcsek *et al.*)¹⁸, spectral methods (Bu *et al.*)¹⁹, graph flow simulation (Pereira-Leal *et al.*)²⁰, edge-betweenness clustering (Dunn *et al.*)²¹ etc are the additional graph clustering techniques applied to explore sub graphs from a network. Most of the research effort on building knowledge based gene clustering algorithm is focused on integrating gene ontology's for extracting the semantic similarity between genes. Graph based gene ontology structures were used by Cheng *et al.*²² for inferring the degree of relationship between genes. The similarity metric is then applied to a community detection algorithm for finding the group of genes that are related by their biological significance and then integrated with an expression-based metric to perform co-cluster analysis for performance improvement. Dotan-Cohen *et al.*²³ integrated semantic similarity measures for deriving clusters using hierarchical clustering approach. The knowledge of bimolecular functions engraved in the GO was used for cutting the dendrogram into meaningful clusters. Graph clustering algorithms make extensive requests on computational assets and demands cluster size as input parameter which increases the computational complexity and performance of the system. The present research proposes an Edge Reduction and Community Detection Algorithm (ERCD) that improves the performance of the graph clustering by finding out the highly connected edges in the graph. The objective is to partition the nodes of a graph based on a threshold value and to create the optimum clusters from the dataset automatically into k independent groups that maximizes the intragroup similarities and minimizes the intergroup similarities. The method uses beta-PERT distribution for modeling the distributions of variables and finding a threshold value which represents the upper bound to segregate the strong and weak connections between the nodes. The similarity threshold

value for each node analytically guarantees the connectivity of each node with adjacent nodes and creates more accurate groupings/clusters than the existing clustering methods.

MATERIALS AND METHODS

Gene Ontology (GO) is a major initiative in bioinformatics that includes complete and ample knowledge about genes and gene-products in a structured and controlled format. They are symbolized as directed acyclic graphs (DAGs) in which the nodes represents the terms and the edges represents the semantic relations. GO is subdivided into three non-overlapping ontology's, Molecular Function (MF), Biological Process (BP) and Cellular Component (CC). Genes and gene products are connected to GO terms using annotations which are linked to a source file which can be a computational, database evidence or a literature reference (Gene Ontology Consortium. 2004²⁴; Smith *et al.*²⁵). GO is extensively used in data analysis and broadly incorporated into various data analysis platforms and applications. GO annotations help in explaining the similarity of a group of genes using different measures. Recent studies on gene ontology using gene annotations enlighten the functioning of complex biological systems. Functional similarity between proteins based on GO helps in extrapolation of disease genes²⁶. Vafaei *et al.*²⁷ introduced semantic similarity measures for predicting gene functional associations. Arnaud *et al.*²⁸ conducted study on ontology to look up genes with similar functionality or

location within the cell. Rhee *et al.*²⁹ and Lovering *et al.*³⁰ did a study to infer the location or function of genes that are over- or under-expressed using GO annotations. De Bodt *et al.*³¹ evaluated the protein-protein interactions (PPI) utilizing the cellular component ontology and inferred that proteins are likely to interact if they are in the same location. Since Gene Ontology represents highly enlightening and comprehensive annotations of genes products and due to its ability to quickly compute pairwise similarity between gene annotations, we used GO annotations as a measure to compute the similarity between genes in the current work.

Edge reduction and community detection (ERCD)-Process Flow

Here we propose a graph based clustering approach that uses the weight of the edges to explore sub graphs of functionally similar genes from Semantic Similarity Network. The process of building a gene semantic similarity network and extracting communities are detailed in Figure 1. ERCD algorithm tries to partition a weighted undirected graph G into homogeneous sub groups or communities such that nodes in the same group are thickly associated, having numerous edge "within" the community than edges linking nodes "between" different communities by

- Cutting or removing edges of each node which are weakly connected based on statistical approach.
- Detecting dense sub graphs by grouping nodes with maximum neighborhood connectivity

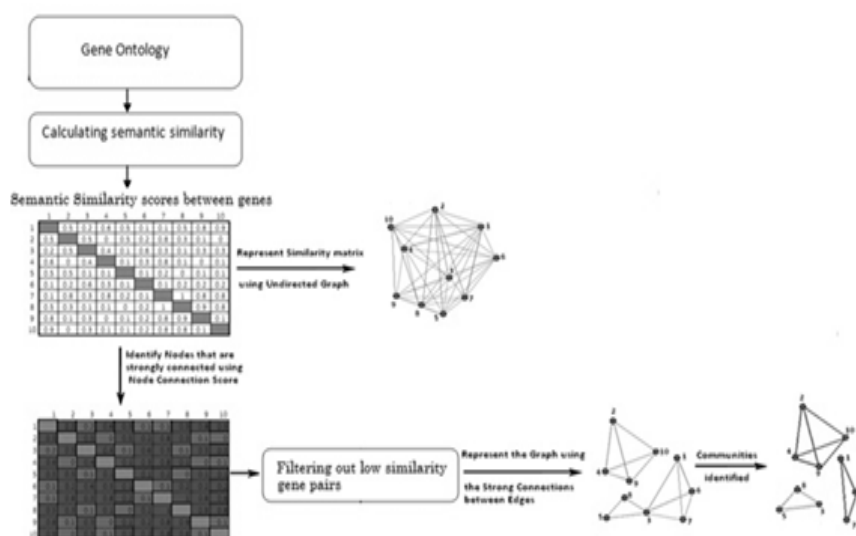


Figure 1
ERCD Process Flow

Components of the ERCD process includes a semantic similarity matrix constructed based on the knowledge source used, the matrix that is transformed into a graph with genes as nodes and edge weights as the association between nodes, optimum semantic graph constructed based on strong associations only and the community extraction process from the optimum graph. The process is detailed in the following steps

Step 1 - Select a Measure of Similarity

There are various measures to express similarity between pairs of objects. If the attribute values are discrete, the similarities between two nodes are determined by examining each of the attributes and counting the number of attribute values they have in common. For continuous attributes, different well known measurements are utilized to compute the separation between two data objects like Euclidean, Manhattan, Murkowski, Canberra, correlation etc. (Pandit *et al.*)³².

Semantic Similarity between GO terms

Usually the analysis happens at the gene level and each gene is associated with multiple GO terms. Different approaches are used to calculate the similarity based on GO terms associated with it. Information Content (IC) based approach is one of the typical methods used to

measure semantic similarity of genes. IC-based similarity measures mainly depend on the factors like frequency of the GO terms and their closest common ancestor term. Equation (1) defines the method for calculating the frequency of a term t ;

$$p(t) = \frac{n_t}{N} \quad | \quad t' \in \{t, \text{children of } t\} \quad (1)$$

where n is the number of term t and N is the total number of terms in GO corpus. IC of a GO term is calculated by negative logarithmic probability of the term, defined in equation (2):

$$IC(t) = -\log(p(t)) \quad (2)$$

Since multiple parents for each concept are allowed in GO allows multiple parents, it is possible to have two terms with common parents with multiple paths. IC based methods finds the similarity of GO terms based on their common ancestor term's IC. Such a common

ancestor term is also known as Most Informative Information Ancestor (MICA). There are four IC based similarity measures that are commonly used, proposed by Resnik³³, Jiang et al.³⁴, Lin and D.³⁵ and Schlicker et al.³⁶.

Resnik Method

$$sim_{Resnik}(t_1, t_2) = IC(MICA) \quad (3)$$

Lin Method

$$sim_{Lin}(t_1, t_2) = \frac{2IC(MICA)}{IC(t_1) + IC(t_2)} \quad (4)$$

Rel Method

This is a combination of Resnik's method and Lin's method. It uses the following equation to find out the similarity.

$$sim_{Rel}(t_1, t_2) = \frac{2IC(MICA)(1-p(MICA))}{IC(t_1) + IC(t_2)} \quad (5)$$

Jiang Method

$$sim_{Jiang}(t_1, t_2) = 1 - \min(1, IC(t_1) + IC(t_2) - 2IC(MICA)) \quad (6)$$

At the end of this process, an $N \times N$ similarity matrix is formed where entries in the matrix quantify the similarity of node pairs.

Step2 – Computation of the Threshold value, Node Connection Score

Any Similarity matrix can be visualized as a network or as a graph. A graph is said to have community structure if it comprises of subsets of genes, with high cohesion within the subset having many edges connecting nodes of the same subset but few edges lying between subsets³⁷. Finding communities within a graph is an efficient way to identify groups of related nodes. It is expensive if we construct the graph for all pairs of similarity index/distances, since for that we have $(N * (N-1))/2$ edges. The performance of the process can be improved by extracting the optimum connections from the graph. For that we followed PERT modeling, which is based on beta distribution and is useful in real-world analysis because it explains the degree of randomness

of the dataset. Typically real dataset follows skewed distribution and beta distribution can handle a variety of skewness in data, both positive and negative efficiently³⁸. An optimum threshold value, NCS is computed for identifying the optimum edge weights/node connections. The method uses three estimation parameters, minimum distance, maximum distance and average path length of each node. For each node 'i', a_i denotes the shortest path of all connected nodes of node i; b_i denotes the largest path of all connected nodes of node i; m_i denotes average path of all connected nodes of node i, NCS is computed using the weighted average mean of all the three parameters for each node shown in equation (7). NCS for each observation reflects the degree of closeness of the nodes in terms of their attribute values

$$NCS_i = \frac{a_i + 4m_i + b_i}{6} \quad (7)$$

Step 3 – Partitioning the Semantic Similarity Graph into Communities

To produce reliable clusters, this approach prunes the similarity matrix, removing edges weights whose similarity indices are less than NCS. Edge weights with higher values represent a high degree of similarity between two nodes. The reduced Similarity matrix determines the intra cluster edges that represent the thickly associated components in the dataset. From the

reduced similarity matrix, vertices are ordered based on the length of total connections on the nodes with each of the adjacent nodes. Pick the nodes from the sorted list and find the intersection of node connection with each of the node. Based on the confidence level set, remove or keep the nodes in the community list. Sub groups or communities are generated from the community list. Each community is distinguished by setting different colors for clusters from 1 to k.

Implementation Of The Proposed Algorithm

The algorithm is implemented in R (<http://www.r-project.org>), a widely used open source platform for data analysis. Table 1 shows the notations and definitions used in ERCD for obtaining natural

communities/clusters. Table 2 represents the pseudo code of the proposed algorithm and Table 3 represents the procedure for building the optimum graph from semantic similarity network using GO terms.

Table 1
Notations used in the Algorithm

D	Gene Dataset
N	Number of genes in the gene set
S	N x N similarity matrix, each cell quantifies the similarity between pair of genes
G	(V, E) be a weighted matrix. The set of nodes/genes and the set of edges of a graph G are denoted by V (G) and E (G), respectively
V(G)	{v1. . . vN} is the node set
E(G)	{e _{ij} } is the edge set with an edge e _{ij} connects vertices v _i and v _j if they are adjacent or neighbors
W _{ij}	The positive weight of an edge (i, j) representing the amount of similarity between genes, gene _i and gene _j .

Table 2
ERCD Algorithm

Input: D=Geneset, T1 = Confidence level
Output: k sub graphs or communities G₁... G_k such that G_i = (V_i, E_i), where V_i ∩ V_j = ∅ ∀ i ≠ j
 Step 1: Graph G = *Generate_OptimumGraph* (D)
 Step 2: For each node in Graph G, *NodeConnected_List*[i] = *getconnectednodes*(G_i), i = 1 to N
 Step 3: Sort the nodes of the Graph by descending order of their degree
 Step 4: Select the *NodeConnected_List* of the first node (V₁) from the sorted list to form communities
 Step 5: Compute confidence level of each node, V_{k=1 to N} with the sorted *NodeConnected_List* using (V₁∩V_k)
 Step 6: If confidence level > T₁; Put all the nodes into a community and remove node from the initial list of nodes
 Else create a new community and add the nodes to the new list.
 Step 7: Repeat Step 4 to 6 until the list is empty

Table 3
Generate Optimum Graph from Semantic Similarity Network

Generate_OptimumGraph(D)
 # Identify strongly connected nodes using Node Connection Score #
 Input: D= Geneset
 Output: Optimum Weighted Graph, G
Begin
 1. Compute the semantic similarity matrix S based on the biological knowledge source
 2. For each row in S, i = 1 to N compute
 i. Edge with minimum value, Z_{iMin} = Minimum (S_{ij}) ∀ i ≠ j, j = 1 to N
 ii. Edge with Maximum Value, Z_{iMax} = Maximum (S_{ij}) ∀ i ≠ j, j = 1 to N
 iii. Average Path Length, $Z_{iAverage} = \frac{\sum_{i \neq j} d(i,j)}{n(n-1)}$
 iv. Similarity threshold value, $Z_{iNCS} = \frac{Z_{iMin} + Z_{iMax} + 4 * Z_{iAverage}}{6}$
 3. # Remove edges which has reduced weight or similarity from Similarity matrix #
 For each row in S, i = 1 to N
 If (S_{ij} < Z_{iNCS}) ∀ i ≠ j, j = 1 to N Delete S_{ij}
 4. Create Graph, G (V, E) using the reduced Similarity Matrix S, which contains only the connections with strong association between genes.
End

Experiment and Result Analysis

The proposed approach is evaluated using a set of 93 cancer genes from Illumina Inc., a research center focusing on the analysis of genetic variation and biological function. Information Content based Resnik approach is employed to find out the semantic similarity between the genes from the cancer gene list. Figure 2 represents the community structure formed using the proposed methodology. The performance of clustering results is validated using three main components based on the study conducted by Jiang *et al.*³⁹, the precise judgment of the cluster quality, performance evaluation

based on ground truth and evaluation of the reliability of the communities formed. Cluster quality can be measured in terms of homogeneity of data objects within the cluster and the separation of data objects between different clusters. Assessment based on ground truth involves comparison with respect to domain knowledge such as protein interactions or known gene functions relating to the clusters. Checking the PPI interactions of genes in each cluster reveals the network similarity. The final aspect of validation gives emphasis on the reliability of the clusters.

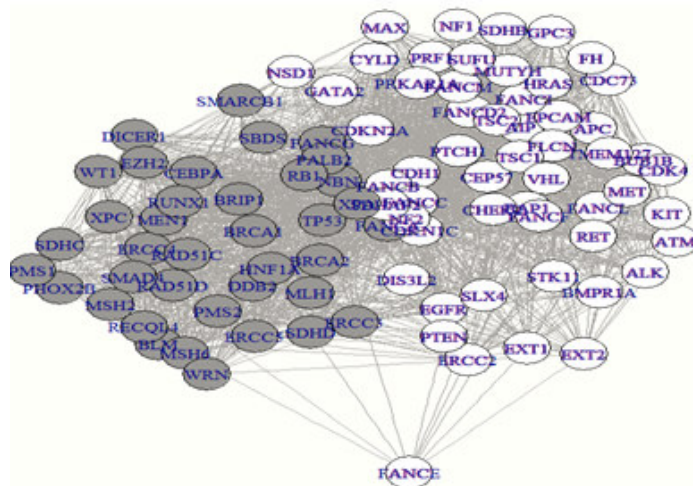


Figure 2
Automatic detection of two communities using ERCD Algorithm

Validation using Modularity measure

Modularity is one of the commonly used metric to evaluate the quality of network’s partition into sub graphs called communities (Newman et al⁴⁰; Newman and M. E.⁴¹). Graphs with a high value of modularity

indicate a good community structure where nodes within groups are strongly connected whereas nodes in different group are sparsely connected. Modularity is defined as

$$Q = \frac{1}{2m} \sum_{ij} [A_{ij} - \frac{k_i \cdot k_j}{2m}] \delta_{c_i c_j} \tag{8}$$

where m is the total number of edges in the network. A_{ij} represents the element of the adjacency matrix with row i and column j, k_i is the degree of node i, c_i is the label of the community to which node i is assigned and $\delta(x, y)$ is 1 if $x=y$ and 0 otherwise. The performance of the proposed approach is compared with three different benchmark community detection algorithms like fast greedy, walktrap and leading eigenvector. Table 4 shows the modularity value and the size of clusters formed when different community detection algorithms

are applied on the same cancer dataset. Modularity of new approach, ERCD is higher when compared to other algorithms and the result (Figure 2) clearly distinguishes the communities formed where as no clear distinction can be inferred from the outputs plotted using other algorithms (Figure 3). Identifying the significant edges, 1844 from a total edge count of 3796 significantly improves the performance of the proposed algorithm and provides an interactive exploration of the clustering results.

Table 4
Comparison of Modularity measure using different community detection algorithms

Algorithm	Modularity	Community Sizes	Edge count
ERCD Algorithm	0.1799317	55, 38	1844
Fast greedy Algorithm	0.05338913	31, 7, 3, 34, 18	3796
Walktrap Algorithm	0.04921651	28, 54, 11	3796
Leading eigenvector Algorithm	0.04985941	48, 38, 7	3796

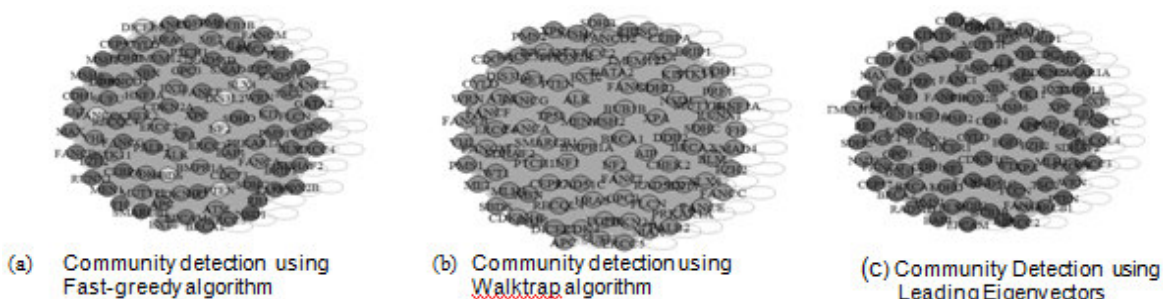


Figure 3
Communities formed when running the three algorithms Fast-greedy (a) Walktrap (b) and Leading Eigenvectors (c) on the cancer dataset

Validation of Clustered genes based on conformity with gene-gene interactions

Several measures have been identified to find the degree of similarity between clusters formed and the ground truth using indices like Rand, Jaccard coefficient,

Minkowski etc. If N denotes the number of data objects and C denotes a set of k clusters $\{C_1, C_2, \dots, C_k\}$, an $N \times N$ binary matrix B can be constructed where $B_{ij} = 1$ if object pairs D_i and D_j belong to the same cluster and $B_{ij} = 0$ otherwise. Similarly, a ground truth matrix $M = \{M_1,$

M_2, \dots, M_s can be constructed to represent the ground truth. Here we extracted the gene interactions of 93 cancer genes from NCBI's database (<http://www.ncbi.nlm.nih.gov/gene>) and checked the conformity of the clusters formed with the gene interactions. Total of 314 interactions were identified which has association with the genes in the training list. The underlying hypothesis is that the number of predicted clusters matching the gene interactions should

be maximum for better clustering solutions. Accuracy is computed by taking the ratio of sum of the count of object pairs showing value "1" in both matrix B and M for each clusters to the total number of gene interactions. Table 5 and 6 lists the gene interactions in the each cluster. We got an accuracy of 78.34% for the clusters derived using ERCD which shows the conformity of the clustering solutions with the ground truth.

Table 5
Gene Interactions of cluster 1 genes that are associated with the genes in the training set.
Total of 88 interactions were detected in cluster 1.

Genes	Count of interactions	Genes	Count of interactions	Genes	Count of interactions	Genes	Count of interactions
AIP	2	FANCB	3	HRAS	1	EGFR	7
EGFR		FANCC		NF1		AIP	
RET		FANCL		MET	2	BMPR1A	
APC	1	FANCM		CDH1		CDH1	
BUB1B		FANCC	5	EGFR		MET	
ATM	5	FANCD2		NF1	1	PRKAR1A	
CHEK2		FANCE		HRAS		PYCH1	
FANCD2		FANCF		NF2	1	RET	
FANCI		FANCL		TSC1		EXT1	2
STK11		FANCM		PRKAR1A	2	EXT2	
VHL		FANCD2	5	CDK4		PYCH1	
BAP1	1	ATM		EGFR		EXT2	1
PTEN		FANCC		PTCH1	2	EXT1	
BUB1B	1	FANCE		EGFR		FANCL	6
APC		FANCI		EXT1		FANCB	
CDH1	3	FANCL		PTEN	3	FANCC	
EGFR		FANCE	4	BAP1		FANCD2	
MET		FANCC		CDH1		FANCF	
PTEN		FANCD2		STK11		FANCI	
CDK4	3	FANCF		RET	2	FANCM	
CDKN1C		FANCM		AIP		FANCM	5
CDKN2A		FANCF	4	EGFR		FANCB	
PRKAR1A		FANCC		STK11	2	FANCC	
CDKN1C	1	FANCE		ATM		FANCE	
CDK4		FANCL		PTEN		FANCF	
CDKN2A	2	FANCM		TSC1	2	FANCL	
CDK4		FANCI	3	NF2		VHL	3
VHL		ATM		TSC2		ATM	
CHEK2	2	FANCD2		TSC2	1	CDKN2A	
ATM		FANCL		TSC1		CHEK2	
VHL						Grand Total	88

Table 6
Gene Interactions of cluster 2 genes that are associated with the genes in the training set.
Total of 158 interactions were detected in cluster 2.

Genes	Count of interactions	Genes	Count of interactions	Genes	Count of interactions	Genes	Count of interactions	Genes	Count of interactions
BLM	11	BRIP1	5	NBN	6	MLH1	8	MSH6	8
BRCA1		BLM		BLM		BLM		BLM	
BRIP1		BRCA1		BRCA1		BRCA1		BRCA1	
FANCA		MLH1		MLH1		BRIP1		MLH1	
FANCG		PMS1		MSH2		MSH2		MSH2	
MLH1		PMS2		MSH6		MSH6		NBN	
MSH2		CEBPA	4	WRN		NBN		PMS1	
MSH6		HNF1A		PALB2	2	PMS1		PMS2	
NBN		RB1		BRCA1		PMS2		TP53	
RAD51D		RUNX1		BRCA2		XPA	4	FANCG	4
TP53		SMAD4		PMS1	5	DOB2		BLM	
WRN		DDB2	3	BRCA2		ERCC4		BRCA2	
BRCA1	15	EZH2		BRIP1		MSH2		ERCC4	
BLM		XPA		MLH1		XPC		FANCA	
BRCA2		XPC		MSH2		WRN	3	HNF1A	1
BRIP1		ERCC3	3	MSH6		BLM		CEBPA	
EZH2		TP53		PMS2	6	NBN		MEN1	2
FANCA		ERCC4	4	BRCA1		TP53		TP53	
MLH1		FANCA		BRCA2		WT1	3	SMARCB1	5
MSH2		FANCG		BRIP1		EZH2		BRCA1	
NBN		MSH2		MLH1		MEN1		RB1	
PALB2		FANCG		MSH2		TP53		RUNX1	
PMS2		MSH2		MSH6		XPC	5	XPC	
PMS2		XPA		RAD51C	1	DOB2		Grand Total	158
RB1		ERCC5	3	RAD51D		ERCC3			
SMAD4		ERCC3	3	RAD51D	2	SMARCB1			
SMARCB1		EZH2	3	BLM		TP53			
TP53		BRCA1		RAD51C		XPA			
BRCA2	6	DOB2		RB1	4	TP53	11		
BRCA1		WT1		BRCA1		BLM			
FANCG		MSH2	11	CEBPA		BRCA1			
PALB2		BLM		SMARCB1		BRCA2			
PMS1		BRCA1		TP53		ERCC3			
PMS2		ERCC4		RECQL4	1	MEN1			
TP53		FANCA		TP53		MSH2			
FANCA	5	MLH1		RUNX1	2	MSH6			
BLM		MSH6		CEBPA		RB1			
BRCA1		NBN		SMARCB1		RECQL4			
ERCC4		PMS1		SMAD4	2	SMARCB1			
FANCG		PMS2		BRCA1		WRN			
MSH2		TP53		CEBPA		WT1			
		XPA				XPC			

Validation based on Silhouette Index

Quality of the clustering is also measured using the internal validation index, Silhouette Index⁴². For a given

$$s = \frac{b-a}{\max(a,b)} \quad (9)$$

where a is the average distance between a sample and all other data points in the same cluster and b is the average distance between a sample and all other data points in the neighboring clusters. A higher Silhouette score relates to a model with better defined clusters. K-means clustering is applied on the same dataset by setting $k = 2$ to 10 to extract the optimum Silhouette

community or cluster, G_i ($i = 1 \dots k$), silhouette width is a quality measure that indicates the membership of a sample in a cluster. It's defined as

index. Silhouette index is maximized in the second cluster solution with an average silhouette width of 0.64 shown in Figure 4. The automatic detection of two clusters using our approach shows a better Silhouette index of 0.754 which shows the performance of the new clustering solution.

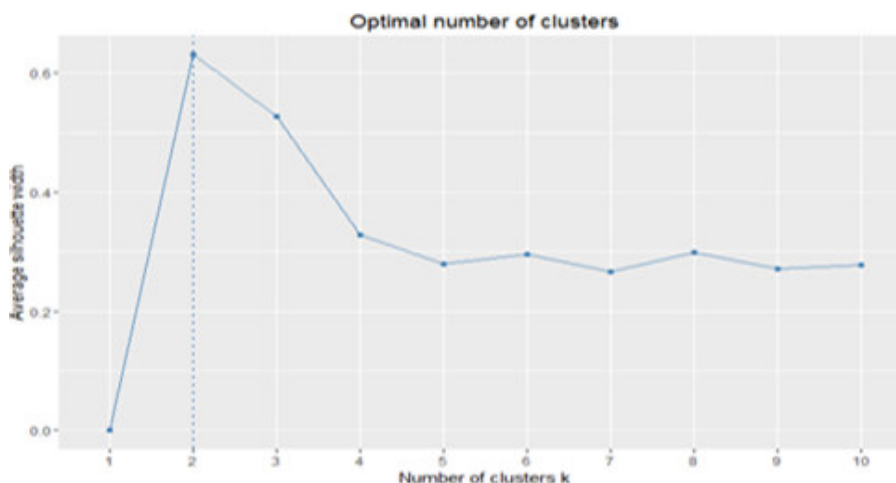


Figure 4
Average Silhouette Width of cancer dataset for different cluster sizes

CONCLUSION AND FUTURE WORK

Many networks in biological system, including gene networks are found to divide naturally into communities or clusters. Discovering and characterizing this community structure is one of the major challenges in the study of graph/networked systems. The reliance of many clustering solutions depends on the prior knowledge of cluster or community size, which degrades the performances of the system. Edge Reduction and Community Detection Algorithm overcomes this by exploring the communities automatically using the edge reduction approach which gives a true set of distinguished clusters. Results from ERCD is compared with existing community detection algorithms like Fast greedy, Walk trap, Leading eigenvector and K-means clustering to demonstrate the data clustering capability. Different validation measures are used to check the performance of the system. The proposed algorithm exhibits excellent performance, with an added advantage of low computational complexity that enables one to analyze large systems. As an enhancement we propose to use integrated knowledge to extract functionally similar genes that contribute to a single

phenotype or disease type from the set of promising candidate genes.

Abbreviations

GO	Gene Ontology
PIP	Protein-Protein Interactions
ERCD	Edge Reduction and Community Detection
MF	Molecular Function
BP	Biological Process
CC	Cellular Component
DAG	Directed Acyclic Graphs
NCS	Node Connection Score

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CONFLICT OF INTEREST

Conflict of interest declared none.

REFERENCES

1. Botstein D, Risch N. Discovering genotypes underlying human phenotypes: past successes for mendelian disease, future approaches for complex disease. *Nature genetics*. 2003 Mar 1;33:228-37.
2. Manolio TA, Collins FS, Cox NJ, Goldstein DB, Hindorf LA, Hunter DJ, McCarthy MI, Ramos EM, Cardon LR, Chakravarti A, Cho JH. Finding the missing heritability of complex diseases. *Nature*. 2009 Oct 8;461(7265):747-53.
3. Aravind L. Guilt by association: contextual information in genome analysis. *Genome Research*. 2000 Aug 1;10(8):1074-7.
4. Jonsson PF, Bates PA. Global topological features of cancer proteins in the human interactome. *Bioinformatics*. 2006 Sep 15;22(18):2291-7.
5. Jonsson PF, Cavanna T, Zicha D, Bates PA. Cluster analysis of networks generated through homology: automatic identification of important protein communities involved in cancer metastasis. *BMC bioinformatics*. 2006 Jan 6;7(1):2.
6. Cai JJ, Borenstein E, Petrov DA. Broker genes in human disease. *Genome biology and evolution*. 2010 Jan 1;2:815-25.
7. Rahmani H, Blockeel H, Bender A. Predicting genes involved in human cancer using network contextual information. *Journal of Integrative Bioinformatics (JIB)*. 2012 Mar 1;9(1):44-71.
8. Pan G, Zhang W, Wu Z, Li S. Online community detection for large complex networks. *PLoS one*. 2014 Jul 25;9(7):e102799.
9. Clauset A, Newman ME, Moore C. Finding community structure in very large networks. *Physical review E*. 2004 Dec 6;70(6):066111.
10. Newman ME. Modularity and community structure in networks. *Proceedings of the national academy of sciences*. 2006 Jun 6;103(23):8577-82.
11. Newman ME. Fast algorithm for detecting community structure in networks. *Physical review E*. 2004 Jun 18;69(6):066133.
12. Pons P, Latapy M. Computing communities in large networks using random walks. In *International Symposium on Computer and Information Sciences 2005 Oct 26 (pp. 284-293)*. Springer Berlin Heidelberg.
13. Spirin V, Mirny LA. Protein complexes and functional modules in molecular networks. *Proceedings of the National Academy of Sciences*. 2003 Oct 14;100(21):12123-8.
14. Hartuv E, Shamir R. A clustering algorithm based on graph connectivity. *Information processing letters*. 2000 Dec 31;76(4-6):175-81.
15. Pržulj N, Wigle DA, Jurisica I. Functional topology in a network of protein interactions. *Bioinformatics*. 2004 Feb 12;20(3):340-8.
16. Krogan NJ, Cagney G, Yu H, Zhong G, Guo X, Ignatchenko A, Li J, Pu S, Datta N, Tikuisis AP, Punna T. Global landscape of protein complexes in the yeast *Saccharomyces cerevisiae*. *Nature*. 2006 Mar 30;440(7084):637-43.
17. Enright AJ, Van Dongen S, Ouzounis CA. An efficient algorithm for large-scale detection of protein families. *Nucleic acids research*. 2002 Apr 1;30(7):1575-84.
18. Adamcsek B, Palla G, Farkas IJ, Derényi I, Vicsek T. CFinder: locating cliques and overlapping modules in biological networks. *Bioinformatics*. 2006 Apr 15;22(8):1021-3.
19. Bu D, Zhao Y, Cai L, Xue H, Zhu X, Lu H, Zhang J, Sun S, Ling L, Zhang N, Li G. Topological structure analysis of the protein-protein interaction network in budding yeast. *Nucleic acids research*. 2003 May 1;31(9):2443-50.
20. Pereira-Leal JB, Enright AJ, Ouzounis CA. Detection of functional modules from protein interaction networks. *PROTEINS: Structure, Function, and Bioinformatics*. 2004 Jan 1;54(1):49-57.
21. Dunn R, Dudbridge F, Sanderson CM. The use of edge-betweenness clustering to investigate biological function in protein interaction networks. *BMC bioinformatics*. 2005 Mar 1;6(1):39.
22. Cheng J, Cline M, Martin J, Finkelstein D, Awad T, Kulp D, Siani-Rose MA. A knowledge-based clustering algorithm driven by gene ontology. *Journal of biopharmaceutical statistics*. 2004 Dec 29;14(3):687-700.
23. Dotan-Cohen D, Kasif S, Melkman AA. Seeing the forest for the trees: using the gene ontology to restructure hierarchical clustering. *Bioinformatics*. 2009 Jul 15;25(14):1789-95.
24. Gene Ontology Consortium. The Gene Ontology (GO) database and informatics resource. *Nucleic acids research*. 2004 Jan 1;32(suppl 1):D258-61.
25. Smith B, Ashburner M, Rosse C, Bard J, Bug W, Ceusters W, Goldberg LJ, Eilbeck K, Ireland A, Mungall CJ, Leontis N. The OBO Foundry: coordinated evolution of ontologies to support biomedical data integration. *Nature biotechnology*. 2007 Nov 1;25(11):1251-5.
26. Jiang R, Gan M, He P. Constructing a gene semantic similarity network for the inference of disease genes. *BMC systems biology*. 2011 Dec 14;5(2):S2.
27. Vafae F, Rosu D, Broackes-Carter F, Jurisica I. Novel semantic similarity measure improves an integrative approach to predicting gene functional associations. *BMC systems biology*. 2013 Mar 14;7(1):22.
28. Arnaud MB, Costanzo MC, Shah P, Skrzypek MS, Sherlock G. Gene Ontology and the annotation of pathogen genomes: the case of *Candida albicans*. *Trends in microbiology*. 2009 Jul 31;17(7):295-303.
29. Rhee SY, Wood V, Dolinski K, Draghici S. Use and misuse of the gene ontology annotations. *Nature Reviews Genetics*. 2008 Jul 1;9(7):509-15.
30. Lovering RC, Dimmer EC, Talmud PJ. Improvements to cardiovascular gene ontology. *Atherosclerosis*. 2009 Jul 31;205(1):9-14.
31. De Bodt S, Proost S, Vandepoele K, Rouzé P, Van de Peer Y. Predicting protein-protein interactions in *Arabidopsis thaliana* through integration of orthology, gene ontology and co-

- expression. BMC genomics. 2009 Jun 29;10(1):288.
32. Pandit S, Gupta S. A comparative study on distance measuring approaches for clustering. International Journal of Research in Computer Science. 2011 Jan 1;2(1):29.
 33. Resnik P. Semantic similarity in a taxonomy: An information-based measure and its application to problems of ambiguity in natural language. J. Artif. Intell. Res.(JAIR). 1999 Jul 11;11:95-130.
 34. Jiang JJ, Conrath DW. Semantic similarity based on corpus statistics and lexical taxonomy. arXiv preprint cmp-lg/9709008. 1997 Sep 20.
 35. Lin D. An information-theoretic definition of similarity. In:ICML 1998 Jul 24 (Vol. 98, No. 1998, pp. 296-304).
 36. Schlicker A, Domingues FS, Rahnenführer J, Lengauer T. A new measure for functional similarity of gene products based on Gene Ontology. BMC bioinformatics. 2006 Jun 15;7(1):302.
 37. Girvan M, Newman ME. Community structure in social and biological networks. Proceedings of the national academy of sciences. 2002 Jun 11;99(12):7821-6.
 38. Moitra SD. Skewness and the beta distribution. Journal of the Operational Research Society. 1990 Oct 1;41(10):953-61.
 39. Jiang D, Tang C, Zhang A. Cluster analysis for gene expression data: A survey. IEEE Transactions on knowledge and data engineering. 2004 Nov;16(11):1370-86.
 40. Newman ME. J, Girvan M (2004) Finding and evaluating community structure in networks. Physical Review E.;69(2):026113.
 41. Newman ME. Modularity and community structure in networks. Proceedings of the national academy of sciences. 2006 Jun 6;103(23):8577-82.
 42. Rousseeuw PJ. Silhouettes: a graphical aid to the interpretation and validation of cluster analysis. Journal of computational and applied mathematics. 1987 Nov 1;20:53-65.
 43. Ashok S, Judy MV. A novel iterative partitioning approach for building prime clusters. International Journal of Advanced Intelligence Paradigms. 2015;7(3-4):313-25.
 44. MALLICK PK, MISHRA D, PATANAIK S, SHAW K. A novel supervised gene clustering approach by mining interdependent gene patterns. International Journal of Pharma and Bio Sciences.;7.
 45. Sonumol NS, Uma VR, Ashok S, Judy MV. Community detection in multidimensional genomic dataset. International Journal of Artificial Intelligence™. 2015 Sep 22;13(2):109-17.