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Abstract: Healthcare is a major area of research since few years. Ample amount of biological data getting accumulated daily due to advancement in technologies. Microarray is such technology which captures expressions of thousands of genes at a time. Interactions occur among genes are represented in terms of special network known as Gene Regulatory Network (GRN). It is constructed from Differentially Expressing Genes(DEFs). GRN is a graphical representation containing genes as nodes and regulatory interactions among them as edges. It helps in tracking pathways where usual gene interaction changes leading to malfunctioning of cells and results in illness. Also, now a day's people are diagnosed with new diseases like dengue, swine flu, Nipah, Corona virus infection for which exact molecular pathways are yet to be invented through GRN. Therefore, in this paper, a nature inspired algorithm is used for reconstruction of GRN using differentially expressing genes.

Keywords: Microarray, Genes, Cellular Biology, Gene Regulatory Network, Differentially Expressing Genes

I. INTRODUCTION

Genes contain blue print of living organisms. All cell activities are controlled by synthesis of proteins whose disproportionate share causes malfunctioning in cellular activity. Some gene products known as proteins are required by cells under all growth conditions. Those are called housekeeping genes. These include genes that encode proteins such as DNA polymerase, RNA polymerase, and DNA gyrase. Some gene products are required under specific growth conditions. These include enzymes that synthesize amino acids, break down specific sugars, or respond to a specific environmental condition such as DNA damage [1]. To analyze the insight of biological activities, analysis of gene expressions is necessary. Advanced technology like microarray plays an important role in gene expression

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analysis as it captures expressions of thousands of genes under different conditions simultaneously. Those genes which behave differently under stress conditions are called as Differentially Expressing Genes (DEGs).

Identifying gene interactions is a major challenge in post genomic era. It helps in knowing how cells maintain their form. Though vast amount of biological data getting accumulated day by day, a technique is needed which will successfully model uncertainty lies in gene expressions in terms of GRN.

A. Definition and Concepts

Definition 1: The Gene Regulatory Network is a graph G(E, N), where N represents set of genes and E represents set of regulatory interactions through which genes communicate with each other.

Definition 2: A positive regulation between gene g1 and gene g2 is indicated by a directed edge arising from source gene g1 to the target gene g2 and is denoted as g1 \longrightarrow g2. Gene g1 positively regulates gene g2; iff binding of gene g1 at specific promoter causes gene g2 to express. In this case gene g1 is called activator gene and gene g2 is called target gene.

Definition 3: A negative regulation between gene g1 and gene g2 is indicated by an undirected edge arising from source gene g1, and closed at target gene g2, g1—g2 and is denoted as gene g1 negatively regulates gene g2; iff inactivation of gene g1 at operon site causes gene g2 to express. In this case gene g1 is called inhibitor gene and gene g2 is called target gene. For showcasing GRN, graphical representation is preferred as it is simple and perfect layout to show interaction between genes. Interaction between genes can be shown using any preferred way not necessarily as mentioned in definitions2 and 3. Figure 1 shows sample GRN of budding yeast. Green arrows and red blunt-end ones are activating and inhibiting interactions, respectively. For self-pointed arrows, orange blunt-end indicates self-degradation.

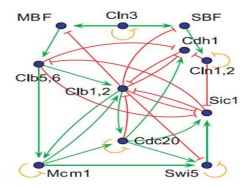


Figure 1. Sample GRN of budding yeast [2]



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In past few years there are many methods proposed in [3][4][5][6][7][8] for inference of GRN but still this research area has a wide scope because of inability to reach to maximum detection of true positive interactions between genes for complex disorders. Based on the chronological order, existing models are classified into two major categories i.e. conventional and non-conventional. A conventional model includes Boolean Network, Model Bayesian Network, Linear Differential non-conventional model includes Neural Network model and Model based on Evolutionary algorithms. Paper[9] gives detailed review of existing mathematical models used for reconstruction of GRN along with database experimental setup used. Disadvantages of some of the important models are given in Figure 2.

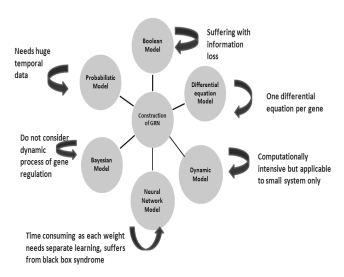


Figure 2. Disadvantage of existing models used for construction of GRN

II. METHODOLGY

Research in bioinformatics demands use of advanced tools for processing huge amount of ambiguous and uncertain biological data. Discovering patterns hidden in the gene expression data across number of samples which are correlated with specific condition has a tremendous opportunity and challenges for functional genomics and proteomics [10][11][12]. Unfortunately, employing any kind of pattern recognition algorithm to such data is hindered by the curse of dimensionality (limited number of samples and to Swarm Intelligence Category have capability to handle enormous data and generate solution from it in simpler way. Errors generated in microarray are more tolerable in SI algorithms than in deterministic algorithms. Errors are treated as contributing factor for population diversity, a desirable property for convergence of SI algorithms [14]. Therefore Ant Colony Optimization based algorithm is proposed which will generate GRN from any number of genes in less time by considering relationship between genes. In 2005, Karaboga [15] gave an interesting idea of artificial ants based algorithm known as Ant colony optimization (ACO) algorithm. Ants are blind, but yet know how to find the shortest distance between the food source and there native place. Ants use pheromones laid by the other ants as footmarks to follow and hence ant reaches the shortest path by using knowledge gained by the other ants and this behavior is imitated in the form of an algorithm that can be used for optimization problems, including gene interaction network optimization [16]. In [16], ACO is used for inference of GRN but author is able to find number of interactions equal to number of genes. It is major drawback because one target gene has many controlling parent which regulates its expressions [17]. Inspired by the foraging activities of ants, ant colony optimization [18] is a class of metaheuristics that provide a generic framework of communication between simple agents (artificial ants), whose task is to construct candidate solutions to the optimization problem under consideration. One type of heuristic that has not been used previously is Ant-Based algorithm. The difference between the Ant Colony Optimization and the Ant-Based algorithm is that in both cases artificial ants maneuver based on the local information and deposited pheromones as they travel but in Ant-Based algorithm cumulative pheromone levels are used to build candidate solution. In Ant Colony Optimization, each ant builds the individual solution and leaves pheromone on the edges which act as guide for remaining ants but in Ant-Based algorithm each ant builds a part of solution and together efforts of all ant gives rise to final solution. We have combine features of both the algorithms and proposed a hybrid approach known Sequential as

very high feature dimensionality) [13]. Algorithms belonging

Algorithm 1 Algorithm for reconstruction of GRN using SDCAA

Initialize pheromone matrix $\tau_{ij}(t2) = [x]_{NXN}$ based on correlation coefficient between genes for first two sample points.

Initialize Tabu list for each gene g_i as $T_i = \{g_i\}$

Initialize Interaction type as $I_{ij} = \{0\}_{NXN}$

Initialize $D_i = \{0\}$ which contains degree of each gene

 $\alpha := 1$ the parameter controlling influence of pheromone on the edge

 $\beta = 2$ the parameter controlling desirability of edge between gene i and j

 $\rho := 0.5$ is pheromone evaporation rate

 $E := [e_{ij}]_{NVM}$ Contains gene expressions of DEGs



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Procedure

1: Initialize one ant at each gene

2: Initialize pheromone between pair of genes using equation 1

3: while stopping criteria not met

4: **for** t = 2 to M **do**

5: for each ant k do

6: Move ant k from gene i to j with probability

$$P_{ij}^{k}(1,t) = \begin{cases} \frac{\left[\tau_{ij}(t)\right]^{\alpha}.\left[\eta_{ij}\right]^{\beta}}{\sum_{\textit{Keallowed}_{k}} [\tau_{ik}(t)]^{\alpha}.\left[\eta_{ik}\right]^{\beta}} & \textit{if jeallowed}_{k} \\ 0 & \textit{otherwise} \end{cases}$$

7: Update pheromone for edges which are selected by ant using expression

$$\tau_{ij}(1,t) \coloneqq (1-\rho)\tau_{ij} + 1/\eta_{ij}$$

8: Update pheromone for the edges which are not selected using

$$\tau_{ij}(1,t) := (1 - \rho)\tau_{ij}$$

9: Update Tabu list T_i , Interaction type I_{ij} and degree vector D_i of each gene

10: end for

11: end while

12: if stopping criteria met then go to step 13 else empty Tabu list and go to step 3

Based on threshold value of pheromone construct adjacency matrix A[i][j]_{NXN} between genes

14: **if** $\tau_{ij} < Th$ **then** A[i][j] = 0

15: else A[i][j] = 1

16: end if

Where M is maximum and m is minimum value of expression for ith gene. Value of pheromone decides the regulatory interaction between genes. Scaling factor 3 is used in order to have large enough differences in pheromone values so that it is easy to select suitable edge.

The algorithm adds new edge between existing gene g1 in GRN and the new gene g2 which has highest probability of getting selected. Due to this, total number of edges at the end is more than number of genes which is advantageous in biological point of view. Amount of pheromone evaporates if edge connecting already added ge ne is not selected in further iteration. By selecting edges having pheromone value above threshold restricted the degree of each node in GRN. The algorithm stops when GRN with maximum deposition of pheromone is generated. The algorithm is compared with existing approaches on the basis of true positive edges matched with benchmark networks. Specificity and sensitivity of reconstructed GRN is also calculated to check the performance of proposed algorithm. GRN is constructed from adjacency matrix generated from Algorithm1 which is given as input to Cytoscape [19]. It is mainly used for graphical display of any kind of biological networks. Solid edges are used to represent positive regulation and dotted edges to represent negative regulation along with label 1 and -1 respectively. In order to compare the result of SDCAA with other existing approaches, GRN is constructed using following datasets.

A.Urilon dataset: It contains 9 significant genes responding to DNA breakdown. This is the most preferred dataset used for

validity of new method. This dataset contains missing values which are first imputed and the GRN is constructed from it. Four different experiments were conducted with different UV light intensities. Using these experiments, expressions of eight major genes, such as uvrD, uvrA, lexA, recA, umuDC, ruvA, polB and uvrY, have been documented as shown in the Figure 3. The displayed relationships express known regulatory interactions between genes. Normal arrow heads denote activation, while diamond-shaped arrow heads denote repression or inhibition.

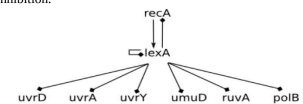


Figure 3. Structure of the SOS DNA repair transcriptional network of E. coli[5].

SOS DNA repair transcriptional network of E.coli using SDCAA algorithm is shown in Figure 4.

III. RESULTS

We have compared GRN modeled using SDCAA with different existing approaches on the basis of known interaction which is shown in Table 1. In this dataset, LexA is a major repressor gene which represses expressions of all other genes.

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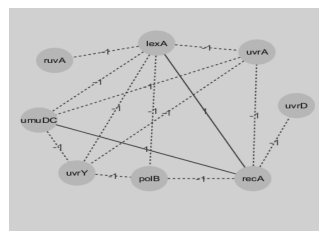


Figure 4 SOS DNA repair GRN of E. coli using SDCAA

In the table Y indicates that known interaction mentioned in first column is correctly predicted using respective method. In methods [20][21][22][23] two genes, uvrY and ruvA were not considered as missing values are there in their expressions for initial time samples. Positive predictive value is calculates using formula as:

$$PPV = \frac{\textit{TP edges}}{\textit{Total number of edges}}$$

False Positive edges are the edges which are incorrectly identified as significant. Method mentioned in [24] reports a less conservative prediction which included all nine true relations but more FP = 7 leading to a lower precision value (PPV = 0.56). Neural network technique is used in [5] which is suffering with black box syndrome. Apart from true edges, list of spurious edges of SOS DNA repair GRN is listed in Table 2. Sensitivity of GRN in Figure 4 using proposed algorithm is 66% and specificity is 33%.

Table 1 Comparative analysis of SDCAA with other methods for SOS GRN

Known interactions	[25]	[26]	[20]	[24]	[21]	[22]	[23]	[5]	[16]	SDCAA
lexA->lexA	Y	Y	Y	N	Y	Y	Y	Y	N	N
lexA->recA	Y	Y	N	Y	Y	Y	Y	Y	Y	Y
recA->lexA	Y	Y	Y	N	Y	Y	Y	N	N	Y
lexA->uvrA	Y	Y	Y	Y	N	Y	Y	Y	Y	Y
lexA->uvrD	N	N	Y	Y	Y	Y	Y	Y	N	N
lexA->uvrY	N	N	-	N	-	-	-	Y	N	Y
lexA->umuD	N	Y	Y	Y	Y	Y	Y	Y	N	N
lexA->ruvA	N	N	-	N	-	-	-	Y	N	Y
lexA->polB	N	N	Y	Y	Y	Y	Y	Y	N	Y
Spurious edges (FP)	5	10	6	7	15	16	11	5	6	6
Precision (PPV)	0.28	0.33	0.50	0.56	0.29	0.30	0.39	0.62	0.25	0.43

Table 2 List of Spurious edges of SOS DNA repair DNA

Sr.No.	Gene 1	Gene 2	Interaction
1.	итиDC	uvrA	negative
2.	итиDC	recA	negative
3.	uvrA	recA	negative
4.	uvrA	uvrY	negative
5.	uvrY	polB	negative
6.	polB	recA	negative

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A.Yeast cell cycle ($\alpha - factor$): It is also called Spellman dataset [21] containing gene expressions of yeast while undergoing cell cycle regulation. Figure 5 shows the standard GRN of yeast cell cycle constructed using GeneNetweaver. GRN is constructed using 10 DEGs.

Figure 6 shows GRN constructed using SDCAA algorithm. Positive interaction is shown using continuous line and negative interaction is shown using dotted line. Total number of edges in the network is 15 out of which 6 are spurious edges which are shown in Table 3.

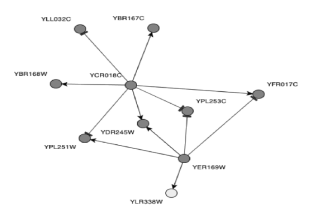


Figure 5. Standard GRN of Yeast from GeneNetweaver

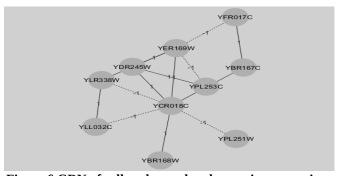


Figure 6 GRN of cell cycle regulated genes in yeast using SDCAA

Table 3 Gene interaction for cell cycle regulated genes in yeast

Gene1	Gene2	Predicted Interaction type	Standard interaction P/N
YCR018C	YBR167C	1	P
YCR018C	YLL032C	-1	P
YCR018C	YBR168W	1	P
YCR018C	YPL251W	-1	N
YCR018C	YDR245W	1	P
YCR018C	YER169W	1	N
YER169W	YLR338W	1	P
YER169W	YFR017C	-1	P
YER169W	YPL253C	-1	N
YER169W	YDR245W	1	P
YFR017C	YBR167C	1	N

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YCR018C	YLR338W	-1	N
YPL253C	YDR245W	1	N
YLL032C	YLR338W	1	N
YFR017C	YER169W	-1	N

IV. DISCUSSION AND CONCLUSION

Sensitivity of SOS and yeast network is 66 % and 60% respectively and specificity is 33% and 50% respectively. SDCAA can build GRN of any size in considerable amount of time. It is flexible and less time consuming which makes it better choice for reconstruction of GRN. Time complexity of SDCAA is O(m*n) where m is number of sample points and n is number of genes.

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