Disease biomarker identification from gene network modules for metastasized breast cancer

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Advancement in science has tended to improve treatment of fatal diseases such as cancer. A major concern in the area is the spread of cancerous cells, technically referred to as metastasis into other organs beyond the primary organ. Treatment in such a stage of cancer is extremely difficult and usually palliative only. In this study, we focus on finding gene-gene network modules which are functionally similar in nature in the case of breast cancer. These modules extracted during the disease progression stages are analyzed using p-value and their associated pathways. We also explore interesting patterns associated with the causal genes, viz., SCGB1D2, MET, CYP1B1 and MMP9 in terms of expression similarity and pathway contexts. We analyze the genes involved in both the stages—non metastasis and metastasis and change in their expression values, their associated pathways and roles as the disease progresses from one stage to another. We discover three additional pathways viz., Glycerophospholipid metabolism, h-Efp pathway and CARM1 and Regulation of Estrogen Receptor, which can be related to the metastasis phase of breast cancer. These new pathways can be further explored to identify their relevance during the progression of the disease.

A normal cell follows a well-known path of growth, division and death although there are exceptions. Some cells do not die. They keep on dividing again and again, leading to abnormal growth. Such condition of cells is called cancerous, and it corresponds to a class of diseases known as cancer. Unrestricted growth of cells leads to the formation of lumps known as tumors, which can interfere in activities of bodily systems such as the nervous system or the circulatory system. These lumps of cells can be benign or malignant. Benign tumors do not disturb the normal functioning of the body. They are still usually controlled by genes controlling apoptosis. On the other hand, malignant tumor cells invade organs, thereby disrupting their normal functioning. They can also move throughout the body using the blood or the lymphatic system as the channel and invade other parts. Such spread of the disease is referred to as the metastatic stage. When cancerous cells metastasize, it becomes very difficult to treat them. The current characterization of this crucial stage of the disease is incomplete and as a result, proper diagnosis is often not feasible. Certain studies such as⁴⁻⁵ have analyzed differentially expressed genes for metastatic stages of lymphoma, lung cancer and leukaemia. Veer et al.⁴ and Wang et al.⁵ have identified 70 genes that may be associated with the metastatic stage of breast cancer. Chuang et al.⁶ use a protein-protein interaction network to determine the markers in the metastasis stage. All these studies focus on extracting individual disease markers responsible for disease progression. However, studies have shown that these genes may be associated with a variety of functions in the body, and interplay among the genes may lead to disorders that finally lead to cancer. Therefore, it would be beneficial to study the disease progression from the pathway point of view. This is because a biological pathway involves a number of molecules simultaneously, and these molecules have to work in coordination to perform normal cellular activities. A slight change in any of the molecules may lead others in the chain to behave abnormally, resulting in disease diagnosis.

Treating cancer for the non-metastatic stage is possible by drugs or by chemotherapy, but if the disease has spread to other organs, it becomes very difficult to diagnose on time and accordingly provide treatment. In this paper, we use the biological similarity between genes to identify functionally related modules. Gene expression data only consider the preparatory information available during the experiments. Analyzing the modules using only a part of the laboratory information would not be justifiable as genes are known to be functionally correlated in nature. Therefore, we use both topological and functional properties of genes to find biologically significant

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modules. Finding modules with high coherence is very essential in our case, as the main target of this work is to analyze the different characteristics of genes including the causal genes of breast cancer from different perspective. Such an analysis incorporating evidence from published literature can be used by biologists and other researchers to carry forward their work on these genes and their associated features.

This work provides a comprehensive study of biomarkers associated with breast cancer. In this paper, we make the following contributions.

• We propose an effective gene-gene network module extraction technique from micro-array gene expression data. We use both the topological and functional characteristics of the gene networks to find highly enriched modules.
• We analyze members of these modules from different perspectives, e.g., expression patterns and pathways.
• We identify relevant pathways by analyzing modules for both the stages. Such pathways provide an insight into the role of each gene in the body and can be further extended to identify its role during disease progression.
• We discover certain interesting characteristics of causal genes such as MET and WARS.
• We find three more pathways to be associated with the metastasis stage of the disease. These pathways can be further explored to find their role in progression of the disease.

### Experimental Results

We implemented our network construction and module extraction method in MATLAB running on an HP Z 800 workstation with two 2.4 GHz Intel(R) Xeon(R) processors and 12 GB RAM, using the Windows7 operating system. The semantic similarity for a given pair of genes is found using the GOSemSim package\(^7\) in R.

| Modules | SST = 0.3 | SST = 0.5 | SST = 0.7 | SST = 0.3 | SST = 0.5 | SST = 0.7 | SST = 0.3 | SST = 0.5 | SST = 0.7 |
|---------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
|         | CCT = 0.3 | CCT = 0.3 | CCT = 0.3 | CCT = 0.5 | CCT = 0.5 | CCT = 0.5 | CCT = 0.3 | CCT = 0.5 | CCT = 0.7 |
| M1      | 5.41E-6   | 2.25E-5   | 5.41E-6   | 4.14E-6   | 5.41E-6   | 5.10E-7   | 5.41E-6   | 2.25E-5   | 5.41E-6   |
| M2      | 3.41E-5   | 3.41E-5   | 4.14E-6   | 5.41E-6   | 3.41E-5   | 7.14E-7   | 3.41E-5   | 1.56E-5   | 3.41E-5   |
| M3      | 6.25E-5   | 1.68E-4   | 3.41E-5   | 3.41E-5   | 2.25E-5   | 8.41E-6   | 1.68E-4   | 2.25E-5   | 2.25E-5   |

Table 1. p–value of top 3 modules obtained using different thresholds in metastasis stage.

Figure 1. Chemokine signalling pathway (source: http://www.kegg.jp/kegg/kegg1.html).
| Module No.: Members | Non-metastasis modules | Pathway associated gene names | p-value |
|---------------------|------------------------|-------------------------------|---------|
| 1: SRGN, MX1, GBP1, PLEK, PDE4B, SLA, IL32, CXCL9, RUNX3, IFH44L, CD52, LRMP, TRBV19, CTSS, PDE4B, CCL5, CXCL10, CCND2, CYP1B1, WARS, MMP9, PFKP, TAP1, ARHGAP4, SLC2A3 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 5.38E-6 |
| 2: CCL5, CCND2, WARS, LAG3 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 3.08E-5 |
| 3: CCL5, CCND2, WARS, IGK@, IGLV5-45 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 1.68E-4 |
| 4: CCL5, CCND2, WARS, IGLV3-19 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 1.68E-4 |
| 5: SRGN, TRAV, TRAC, | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 5.10E-7 |
| 2: WARS, PDYN, NUCB1, GUSBP3 | tryptophan metabolism, Aminoacyl-tRNA biosynthesis | WARS | 7.14E-7 |
| 3: WARS, ESR1, NUCB1, ASC1 | tryptophan metabolism, Aminoacyl-tRNA biosynthesis | WARS | 8.41E-6 |
| 4: CCL5, CCND2, WARS, SRGN, TRBV19 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 9.27E-6 |
| 5: CCL5, WARS, LCAT, MFGE8 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5, LCAT | 1.52E-5 |

| Metastasis modules | Pathways | Pathway associated gene names | p-value |
|--------------------|----------|-------------------------------|---------|
| 1: CCL5, CCND2, WARS, SRGN, TRAC, | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 5.10E-7 |
| 2: WARS, PDYN, NUCB1, GUSBP3 | tryptophan metabolism, Aminoacyl-tRNA biosynthesis | WARS | 7.14E-7 |
| 3: WARS, ESR1, NUCB1, ASC1 | tryptophan metabolism, Aminoacyl-tRNA biosynthesis | WARS | 8.41E-6 |
| 4: CCL5, CCND2, WARS, SRGN, TRBV19 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCL5, CCND2 | 9.27E-6 |
| 5: CCL5, WARS, LCAT, MFGE8 | cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5, LCAT | 1.52E-5 |

| Pathways | p-value |
|----------|---------|
| antigen processing and presentation | CTSS |
| purine metabolism | PDE4B |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway, RIG-1-like receptor signalling pathway. | CXCL10 |
| p53 signalling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway, cyclins and cell cycle regulation. | CCND2 |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |
| antigen processing and presentation | SLC2A3 |
| lipid metabolism | LCAT |
| steroid hormone biosynthesis | WARS |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| purine metabolism | WARS |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| purine metabolism | WARS |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| purine metabolism | WARS |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| purine metabolism | WARS |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |
| cytokine-cytokine receptor interaction, chemokine signalling pathway, NOD-like receptor signalling pathway, cytosolic DNA-sensing pathway, Toll-like receptor signalling pathway. | CCL5 |
| purine metabolism | WARS |
| purine metabolism | WARS |
| antigen processing and presentation | TAP1 |

Table 2. Non-Metastasis and Metastasis modules along with pathway information and p-value.
user. CCT takes care of the topological structure of the network and SST is concerned with the biological similarity among the nodes involved in the network. In order to choose the optimal value of the parameter, we compute the p-value for each module at different thresholds. This is because p-value gives the probability for a set of genes to be enriched with the same functional group. The p-value for a module M enriched with functional group F is given as

\[ p-value = 1 - \sum_{i=0}^{q-1} \frac{|F|}{|V|} \left( \frac{|V| - |F|}{|M| - i} \right) \]

where \( M \) contains \( q \) genes in \( F \) and entire network contains \(|V|\) genes.

To choose the best set of parameters, we compute the p-value of each module obtained at CCT values of 0.3, 0.5 and 0.7. The set of SST values is set to be 0.3, 0.5 and 0.7 for each run of CCT. The p-values of top three modules obtained at different threshold values are given Table 1.

In Table 1, we see that two of the three top modules show p-value of \( 5.10E-7 \) and \( 7.14E-7 \) at SST value of 0.7 with CCT value set to 0.5, while for the rest of the modules, it can be seen that the best p-value obtained is in the range of \( E-6 \).

Therefore, we can say that the optimal parameter set is achieved at CCT = 0.5 and SST = 0.7. This is supported by the fact that lower p-value signifies modules which are more biologically similar in nature among themselves and thus the ultimate purpose of module extraction is satisfied.

**Pathway identification from module members.** We have carried out an extensive analysis of top five modules in terms of p-value in both non-metastasis and metastasis stages. We have identified the pathways to which each member gene belongs to in a given module. This is done using the DAVID tool\(^8\). The members of the modules along with the pathways they associate with are given in Table 2.

In Table 2, we see that three pathways viz., Glycerophospholipid metabolism, h-Efp pathway and CARM1 and Regulation of Estrogen Receptor are associated with genes found in modules in the metastasis stage. These three new pathways can be further explored to find how they are responsible for spreading the disease to other organs.

As explained in Proposition 2, we suggest that genes (other than the disease genes) found in a disease associated module also contribute to the disease. Let us consider the example of the top module in the metastasis stage. This module has CCL5, CCND2 (a disease gene), WARS, SRGN and TRAC as its member elements. In Table 2, we see the different pathways associated with its members. We provide pathways for only those genes which are found in literature. We see that the disease gene CCND2 is related to \( p53 \) signaling pathway, Wnt signalling pathway, focal adhesion, Jak-STAT signalling pathway and others. Among its member genes, we find that CCL5 is involved in cytokine-cytokine receptor interaction, chemokine signalling pathway etc. Looking into the pathway structure of chemokine signalling pathway given in the KEGG database\(^9\), we see that these two pathways are linked to the Jak-STAT signalling pathway, which is shown to be linked to the disease gene pathway. Figure 1 shows the chemokine-signalling pathway. The WARS gene belongs to the tryptophan metabolism pathway, which aids in glycolysis as seen in Fig. 2. The glycolysis mechanism is indirectly regulated by the Wnt signalling pathway\(^10\). This signalling pathway also corresponds to the disease gene pathway. Slight perturbation in any of these pathways may affect the changes to be carry forwarded, thus leading to certain disorders. Therefore, we can say that member genes of a disease module also contribute to the disease.

**Interpreting utility of common genes in both stages.** A Venn diagram representation of the common genes found among the modules in both the stages is shown in Fig. 3. We see that genes CCL5, CCND2 and WARS are found among all the five modules in non-metastasis stage. However, in metastasis stage, Fig. 3(b), we see that these three genes occur simultaneously in modules M1 and M4 only. Among the rest of the modules, only the WARS gene is present as a common element. The other two genes are not associated with three of the modules in the metastasis stage. This may be due to its low semantic value with the seed node for the respective modules. In addition to the disease gene, two other genes are found to be strongly related to the breast cancer disease. CCND2, which is the causal gene found in both stages is known to have a higher invasive ability. Certain results also suggest that overexpression of this gene in carcinom cells has an enhanced effect on in vivo aggressive growth pattern\(^11\). CCL5/CCR3 signalling actively encourages metastasis by polarization of CD4+ T cells, for luminal breast cancer\(^12\). WARS, popularly known as Tryptophanyl-tRNA synthetase corresponds to the aminoacyl-tRNA synthetase family. They are involved in RNA transcription, protein synthesis and in angiogenic signalling pathways\(^13\). Overexpression of tRNA synthetase promotes migratory movements of carcinogenic cells\(^14\). This may be one of the reasons why WARS is present in all modules of both the stages.

**Common pathways among modules and change in expression value of associated genes.** We also analyze the common pathways associated with the common genes in both the stages and observe the changes in their expression values as the disease progresses from the non-metastatic to the metastatic stage. This is shown in Table 3. Among all the genes which are common to the two stages, they show a decrease in their expression value in the two stages.

The role of these common pathways in terms of the disease is highlighted here.
• **Cytokine-cytokine pathway**: Cytokines are released whenever there is some infection or inflammation so as to hamper the tumor’s development. Carcinogens can also respond to the host with cytokines so as to promote growth and spreading of the disease to other organs.

• **Chemokine signaling pathway**: Expression values of chemokines are changed in many forms of malignancies and eventually lead to disturbed chemokine signalling pathway.

• **p53 signaling pathway**: p53 loss can disrupt metastasis related pathways. However, transcriptionally defective TP53 mutants can promote metastasis stage of cancer.

• **Wnt signaling pathway**: Activity of the Wnt/β-catenin signalling pathway plays a significant role during the development of breast cancer.

• **Tryptophan metabolism**: Higher expression of enzymes involved in tryptophan degradation is known to be associated with several forms of cancer such as lung cancer, breast cancer and melanoma.

• **Toll-like receptor signaling pathway**: Toll like receptor signaling in tumor cells play a role in aggressive behavior of tumor by excessive secretion of cytokines/chemokines.

• **Cytosolic DNA sensing pathway**: The role of this pathway is yet not clear in context of the disease, but since it is found in both the stages, we can say that it plays a significant role in causing the disease.

![Figure 2. Tryptophan metabolism (source:http://www.kegg.jp/kegg/kegg1.html).](image1)

![Figure 3. Common genes found among the modules in both the stages.](image2)

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Scientific Reports | 7: 1072 | DOI:10.1038/s41598-017-00996-x
Proposition 1. A causal gene \( g_i \in m_{nm} \), the \( i \)th non-metastasis module may not participate in coherent module formation at metastatic stage.

Explanation: Assume that a gene \( g_i \in m_{nm} \) also \( \in m_{nm} \), the \( j \)th module in metastatic stage. It is evident from the literature and also from our experimental study that during stage transition from non-metastasis to metastasis, often a gene undergoes significant variation (i.e., fall or rise) in expression or semantic similarity values. Such a gene with significantly varied expression or semantic similarity values may not remain coherent with other module forming genes, and hence may not satisfy the \( CCT \) and \( SST \) cutoffs, and may not participate in module formation during the metastasis stage. It contradicts our assumption and hence the proof.

We analyzed the semantic similarity values among these five genes and found that XBPI, MET and CYP1B1 are probable candidates for module formation. However, these could not be members of the modules in the metastasis stage probably due to their low semantic similarity score with the seed nodes.

Analyzing genes based on expression values. Apart from analyzing expression pattern of only the common genes found among the modules across the stages, we also tried to find if genes involved during module formation are coherent among themselves or not. We show the expression patterns of genes in the top three modules obtained during both the stages in Figs 4 and 5. In these figures, we see that participating genes demonstrate high coherence across the samples.

We then observe expression patterns of genes associated with the disease as given in GeneCard. These genes are referred to as causal genes. The average expression levels of these genes across the two stages are analyzed and we find that CYP1B1 and MMP9 show an increasing trend in their average expression values from non-metastasis to metastasis stage. This is clearly different from the normal trend of expression values during progression of diseases. The other three genes XBPI, SCGB1D2 and MET show a decreasing trend during the progression phase. We plot the expression patterns of these three genes across the two stages in Fig. 6. Two of the genes SCGB1D2 and MET show normal trend as seen among the rest of the genes. However, XBPI shows a peculiar trend in both stages. The expression values of this gene shows very low variations across all the 286 samples. This may be due to its inherent nature, as it acts as a transcription factor which regulates the gene expression levels for the immune system and in other cellular responses. This may be one of the reasons of its low variation during disease progression as the immune system prompts the body to respond equally during the non-metastasis and metastasis stages.

Discussion
Cancer is a complex disease. Its distinguishing characteristics can best be represented in the form of a modular structure, technically referred to as a cancer hallmark network. Earlier studies have revealed six types of hallmarks associated with cancer: (i) cancerous cells that promote their own growth, (ii) they thwart inhibitory signals that would harm them, (iii) they do not take part in from programmed cell death, (iv) they promote growth and functioning of blood vessels which provide nutrition to the cancerous cells, (v) they have a tendency to keep on multiplying and (vi) they move out to other organs thereby spreading the disease. Four new characteristic features have been discovered to be associated with cancerous cells. These were added to the hallmarks of cancer: (vii) associated pathways show abnormality, (viii) cancer cells remain hidden from the immune system surveillance, (ix) abnormalities occur in the number of chromosomes in the cancerous cells and
(x) there is inflammation in the affected tissue. For a cancerous cell to survive, divide and move out, it needs to have functional capabilities acquired through activation of different hallmarks during different times of cancerous progression. An extensive study of the signalling network along with the tumor-genome sequencing output has illustrated the importance of metamorphing in case of tumor suppressor genes and their role in cancer development and progression over stages. Mutation in these genes results in genome instability and chromosome amplification. Such a situation gives rise to heterogeneity in the tumor cells or the cancerous cells. This heterogeneity leads to loss of robustness among the disease biomarkers. It poses a major hurdle in cancer biomarker discovery as the contemporary methods of discovering biomarkers based on gene expression profiles are inadequate. Identification of robust cancer biomarkers has recently gained importance in the research community. Methods such as MSS (Multiple Survival Screening) and ensembling methods work by characterizing cancer samples into various classes, depending on their gene expression profiles or signalling network structure. Once they are narrowed down to sub-classes, one can easily predict the associated biomarkers depending on their class characteristics.

The module extraction method proposed here is based on semantic similarity among the genes. This method extracts biologically similar modules as suggested by their p-values. Our module extraction method is used over the metastatic and non-metastatic stages of the breast cancer gene expression dataset. We have analyzed members obtained from the top five modules in terms of expression similarity and pathway point of view.

Certain interesting behaviors of the causal genes which are members such as SCGB1D2, MET, CYP1B1 and MMP9 are also studied. We have also suggested reasons for their peculiar behavior.

We have also identified reasons why certain genes are found in modules of only one stage. Apart from this, we give justification for how genes in a module coordinate with others to achieve the function associated with the module. We also justify why certain genes such as WARS are present in all identified modules in both the stages.

In addition to this, we analyze the expression values of the common genes in both stages of the disease and also speculate how these genes play role in causing the disease considering the pathway.

Lastly, we also find three new pathways namely Glycerophospholipid metabolism, h-Eff pathway and CARM1 and Regulation of Estrogen Receptor to be associated with the metastasis stage of the disease. Understanding these pathways will give better insight into the progression of the disease.

Materials and Method

Besides skin cancer, breast cancer is the most commonly diagnosed disease among women in the United States. About 40,610 women in the U.S. are expected to die in 2017 from breast cancer. If the disease is detected while still at the in-situ stage, the chance of survival is close to 100%. However, if the disease is left untreated, it usually spreads to other parts of the body (metastasize), and the first place it usually spreads to is the lymph nodes in the axilla area. Metastatic breast cancer cells differ from the primary ones in properties such as receptor status. The cells often develop resistance to several lines of previous treatment and acquire special properties that permit them to metastasize to distant sites. Metastatic breast cancer can be treated, sometimes...
for many years, but it cannot be cured. Women in this stage of the disease have less than 16% chances of survival. Therefore, it is very crucial to study the progression of the disease to this stage. We make one such effort here, and the findings of our work can be used by the biologists to better interpret the causes of the disease, to aid on-time treatment.

**Dataset Used.** The study of progression of metastasis cancer from the non-metastasis stage requires a thorough understanding of the genes which get mutated during the disease progression, how their expression values change from time to time, what other changes are brought in during the progression, etc. In order to get a vivid picture of the disease progression, we perform our experiments on the GSE 20304 dataset on breast cancer relapse free survival, obtained from ref. 32. It comprises of expression values of 286 patients. Among the 286 samples, 106 samples are from the metastatic stage and the rest are from the non-metastatic stage.

**Preprocessing and Sample selection.** The dataset consists of 22,273 genes with expression values. In order to scale the values obtained over the large number of samples, we use log2 transformation. We then use variance measure to decide upon the genes which are actively involved across all the 286 samples. We tune variance threshold $V_{th}$ in the range 0.9–1.5, and accordingly we obtain seven different samples of genes, each with 7900, 7000, 6129, 5292, 4529, 3903 and 3356 genes respectively as given in Table 4.

From our observation, we find the best suitable range to be 1.1–1.3. We carry out our module formation process using genes at this range. Modules obtained in this range are analyzed, and we find slightly higher number of modules at $V_{th} = 1.1$ than at $V_{th} = 1.2$ or $V_{th} = 1.3$. This may be a result of the extra 872 genes at this sample stage, which are later removed in the other two stages due to its low variance as compared to $V_{th}$. For our extensive analysis purpose, we use module results obtained at $V_{th} = 1.2$, as it is the mid-point of the two values. Apart from being the average of the two values, it also shows higher number of common modules when compared with the modules obtained at both the ends. This threshold value is also supported by Wang et al. in their work, where they use it to get the maximum variance of genes among the samples for the dataset.

**Table 4.** Various gene samples drawn at different $V_{th}$ threshold.

| $V_{th}$ | Number of genes |
|---------|----------------|
| 0.9     | 7900           |
| 1.0     | 7000           |
| 1.1     | 6129           |
| 1.2     | 5292           |
| 1.3     | 4529           |
| 1.4     | 3903           |
| 1.5     | 3356           |
Network construction and module extraction. The process of network construction begins with 5292 genes. Two different networks are constructed, one for non-metastatic stage and the other for the metastatic stage. These two networks are obtained based on the patient’s characteristics. The network A(i,j) is constructed using the Pearson’s correlation coefficient between the genes, PrC(i,j), where

\[
A_{i,j} = \begin{cases} 
1 & \text{if } PrC(i,j) \geq 0.5 \\
0 & \text{otherwise.} 
\end{cases}
\] (2)

We choose 0.5 as the threshold for Pearson coefficient because we want to extract subnetworks or modules with balanced contributions from both gene expression and semantic similarity. Once the networks, i.e., A_non for non-metastatic stage and A_m for metastatic stage are ready, we start with a seed growing technique to find modules. Seed selection for modules is based on the clustering coefficient of each gene in the network. The choice of seed is based on the node with the maximum clustering coefficient, such that this is greater than an input threshold value for CCT. In order to grow the seed, we use a second criterion which is the semantic similarity between gene pairs so as to find their relation with each other. We choose semantic similarity between gene pairs in order to get functionally relevant modules. The more biologically relevant a module is, the more will be its correspondence to the disease. This statement is supported by Proposition 2.

**Proposition 2.** If for a given disease \(D_p\) the causal gene \(g_i \in m_i\), i.e., \(i^{th}\) network module of high biological significance, then any other gene \(g_j \in m_i\) will also have a correspondence with \(D_p\).

**Explanation:** A pair of genes \((g_i, g_j)\) will be members of a module, \(m_i\), of high biological significance (i.e., with very low p-value) iff \(PrC(g_i, g_j) \geq 0.5\) and \(SS(g_i, g_j) \geq SST\). So, any other member gene of \(m_i\) will share highly similar functionality with gene \(g_i\) or \(g_j\). Therefore, since \(g_i\) corresponds to a disease \(D_p\), other member genes, say \(g_j \in m_i\) will also correspond to \(D_p\), and hence the proof.

This proposition can be better understood with the help of an example given in Section Pathway identification from module members. In order to get the best semantic similarity between a pair of genes, we use Wang’s semantic similarity, which is known to be the best among all the existing measures. However, we have restricted ourselves to only those genes during expansion whose semantic similarity value is at least equal to SST, set by the user. The parameter is set experimentally as discussed in the Section Parameter tuning for p-value computation. We now define some terms which are used during the module extraction process.

**Definition 1 (Semantically connected).** Two genes \(g_i\) and \(g_j\) are said to be semantically connected iff \(PrC(g_i, g_j) \geq CCT\) and \(SS(g_i, g_j) \geq SST\), where CCT and SST are user-defined thresholds.

**Definition 2 (Clustering coefficient).** For an undirected graph, clustering coefficient of a node \(v_i\) (representing a gene, say \(g_i\)) is the ratio of the number of links the node has among its neighbors, \(N_i\), to the total number of possible links for that node. Mathematically,

\[
CC(v_i) = \frac{2 \times N_i}{d_i \times (d_i - 1)}
\] (3)

where \(d_i\) is the degree of node \(v_i\).

**Definition 3 (Seed node).** A node \(v_i\) is said to be a seed node if \(\forall v_j, CC(v_i) > CC(v_j)\), where \(v_j \in \{V - v_i\}\) and \(V\) is the set of nodes.

**Definition 4 (Module).** A module is a set of nodes/genes which are semantically connected among each other.

The steps for module extraction is given in Algorithm 1.

**Lemma 1.** Two genes \((g_i, g_j)\) are functionally coherent if they have high expression and semantic similarity.

**Proof.** This proof is trivial because a pair of genes sharing high expression and semantic similarity show similar trend and are more functionally coherent.

**Lemma 2.** Nodes included by our method in a module have high functional coherence.

**Proof.** A gene \(g_i\) can be a member of module, \(m_i\), iff \(Expression(g_i, g_j) \geq 0.5\) and \(SS(g_i, g_j) \geq SST\), where SST is the user-defined threshold for Semantic similarity. So, for any two member genes of a module, the expression similarity and semantic similarity between them is always high. Hence, the nodes in a module have high functional coherence.

**Complexity Analysis.** The module extraction technique involves selection of seed nodes based on the clustering coefficient of the nodes. This involves identifying neighbors of each node, which takes \(O(n)\) time in the worst case. To identify the number of links present among the neighbor sets takes another \(O(n)\) time. Thus, the seed selection process takes \(O(n) \times O(n) \equiv O(n^2)\) time. The module expansion process starts from the node with
Algorithm 1: Algorithm for module extraction from gene gene network

Input: A = (V, E) (Gene network); CCT (Clustering coefficient threshold); SST (Semantic similarity threshold); NSS (Semantic similarity score matrix)

Output: Modules = (C1, C2, …, CN) (a set of N modules)

1. Initialize clusterExpNode = V, Modules = NULL;
2. while |clusterExpNode| > 0 do
3. choose vn ∈ clusterExpNode such that ∀ vi ∈ clusterExpNode, CC(vn) ≥ CC(vi) and CC(vn) ≥ CCT;
4. partialCluster = partialCluster ∪ vn;
5. while vn exists do
6. choose another vn from N_reachable if and only if vn ∈ partialCluster such that NSS(vn, vn) ≥ SST
7. clusterExpNode = clusterExpNode − vn;
8. N_reachable = (N_reachable ∪ N_reachable) choose next vn;
9. end
10. Mark partialCluster as C_vn, only when |partialCluster| ≥ 3;
11. Modules = Modules ∪ C_vn;
12. count ++;
13. end
14. Return Modules;

the highest clustering coefficient. Sorting the nodes in terms of decreasing clustering coefficient requires \( O(n \log n) \) time and then comparing the top most node with user defined CCT requires \( O(1) \) time. Expansion of modules requires \( O(n^2) \) as it requires computing the neighbor set again for the seed node, which takes \( O(n) \) time and then finding its corresponding semantic similarity, which takes at most \( O(n) \) time. This semantic similarity value has to be compared with the SST value defined by the user, which takes \( O(1) \) time. Therefore, the overall time complexity for module extraction is \( O(n^2) + O(n \log n) + O(1) + O(n^2) + O(1) = O(n^3) \).

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