Functional analysis of cancer gene subtype from co-clustering and classification

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ABSTRACT
Cancer is a heterogeneity genetic disease with huge phenotypic alterations among dissimilar cancers types or even between same cancer types. Recent expansions of genome-wide profiling technologies offer a chance to explore molecular changes variations throughout advancement of cancer. Therefore, various statistical and machine learning algorithms have been designed and developed for the handling and interpretation of high-throughput microarray molecular data. Discovery of molecular subtypes studies have permitted the cancer to be allocated into similar groups that are deliberated to port similar molecular and clinical characteristics. Thus, the main objective of this research is to discover cancer gene subtypes and classify genes to obtain higher accuracy. In particular improved co-clustering algorithm used to discover cancer subtypes. And then supervised infinite feature selection gene selection method was combined with multi class SVM for classification of selected genes and further biological analysis. The analysis on breast cancer and glioblastoma multiforme evidences that top genes involved in cancer and the pathways present in both cancer top genes. The functional analysis is useful in medical and pharmaceutical field for cancer diagnosis and prognosis.

Keywords:
Biological analysis
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1. INTRODUCTION
Abnormalities of cancer genome can be observed through basic researches which have been used to categorize patients with respect to enhance their clinical decision making and implement more efficient treatments. Even though this types of categorization have enhanced the efficiency of treatment of various cancers, but the heterogeneity among the populations still remains as a main challenge. The advancement of DNA microarray technology has permitted an extensive understanding of genes especially in oncology field for start, diagnosis and prognosis of cancers. These various diagnostics are useful for different types of cancer, which lead to individual treatment plans and accurate clinical outcomes estimation [1, 2].

As the initial stage in organizing and investigating high-throughput gene expression datasets is through artificial intelligence in deep machine learning approach by grouping them together (cluster) according to similar biological features (gene) or conditions (samples) conferred on some similarity measures [3-5]. Meanwhile for both features and conditions are typically inadequate with prior knowledge, the clustering process is conducted as an unsupervised process via grouping features and conditions [6]. The conventional clustering is not said to be an ideal method for complicated and heterogeneous cancers. This is because, there are only certain genes in a subset of samples are expressed as a cancer genes in cellular processes among the similar clinical types of cancer in a specific tissue. Hence, it has been found a limitation that a single gene might play role in regulating and participating in numerous clusters and pathways of different conditions [7].

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Therefore, Cheng and Church [8] has been the pioneer in implementation bi-clustering approach on gene expression datasets. Fundamentally, co-clustering [or bi-clustering] simultaneously clusters genes and samples to discover subtypes or subgroups of genes which displays similar patterns within certain conditions subset of experiments [9-11]. On top of that, this application has provided unique opportunities and challenges to classify tumors and discover more tumor subtype mechanisms. In addition to this, the researchers are also trying to identify multiple bi-clusters at a time. For example, statistical method [12], information theory [13], matrix factorization [14], and graph based bi-clustering [15]. Many co-clustering methods were developed by researchers. Cho and Dhillon [16] proposed a minimum sum-squared residue co-clustering (MSSRCC) to identify coherent bi-clusters. Modular Singular Value Decomposition (Mod-SVD) was proposed by Aradhya, Masulli [17] to discover bi-clusters from SVD computation. Huang, Sun [18] developed a modified fuzzy co-clustering (MFCC) while Hussain and Ramazan [19] proposed a method based on co-similarity measure between genes (and conditions).

Therefore, in this research we proposed to improve network assisted co-clustering for the identification of cancer subtypes (iNCIS). Generally this method utilizes gene network prior knowledge to be integrated with gene expression data to obtain bi-clusters.

2. RESEARCH METHOD

2.1. Dataset and Tools

Two cancer microarray datasets were used in this research. They are Breast Cancer (BRCA) and Glioblastoma Multiforme (GBM). In addition, both of these datasets are publicly available at The Cancer Genome Atlas (TCGA) where BRCA obtained from the Network [20] meanwhile GBM is from Verhaak, Hoadley [21]. The cancer datasets were in text file format and had been pre-formatted to be used as an input for the software. These datasets mainly comprise of numerical values; rows representing the genes and columns representing the samples/patients but the classes are unknown. BRCA contains 17814 genes and 547 samples while GBM contains 11861 genes and 202 samples. Co-clustering and classification are performed using MATLAB 2014a, while Feature selection library (FSLib) [22] is used to combine for classification.

2.2. Co-clustering and Validation

In this stage, the first step is to assign weights to each genes using modified PageRank algorithm. Then the beginning of co-clustering (iNCIS) algorithm where the objectives function is improved to minimize the sum-squared residues and optimizes matrix X. The selections of parameters are based on cophenetic correlation coefficient and some of them are default [23, 24]. For the validation, silhouette analysis was done [25]. The larger the silhouette value is better the clustering. Beside this, subnetworks are obtained for a particular gene for both cancer and validated.

2.3. Classification and Validation

The results of number of classes for both datasets were used further for feature selection and classification. Supervised infinite feature selection (SinFS) technique is combining to multiclass support vector machine (mSVM). Hence it is required the number of classes for each genes belongs to which we obtained from the co-clustering method. The genes with highest ranking and lowest threshold p-value are selected for classification further. And then, the selected top ten genes are analyze for functional analysis where the gene is belong to a specific cancer subtypes, biomarkers, oncogenes, transcription factors, tumor suppressor or antigen in addition their pathways involved.

3. RESULTS AND ANALYSIS

This section deeply discussing about the overall results obtained from implementation of co-clustering and classification of two cancer gene expression datasets.

3.1. Cancer Subtypes and Subnetworks of BRCA

From the implementation of BRCA dataset to iNCIS, there are five different cancer subtypes has been identified which are tabulated in Table 1 and visualize in Figure 1 (a) to (e). Gene ABCC8’s subnetwork was produced for these five subtypes that obtained from iNCIS. Subnetwork visualization which shows the expression level and the weight of the gene will be helpful in order to demonstrate the difference of the five subtypes. Primarily, this subnetwork was chosen because it has connected with small number of genes and is easy to present visibly.

From BRCA data set, a small subnetwork (Figure 1) ABCC8 taken as an example to demonstrate the differences of five subtypes. There is total 16 genes presence in this subnetwork. Gene ABCC8 has been highly
expressed in Luminal A and B subtypes, moderately expressed in normal-like subtype and triple-negative and HER2 subtype shows low expression. Besides this, KLK11, KLK13, HDAC5 and RRAD genes are expressed moderate to high level in all subtypes.

### Table 1. Number of samples for BRCA subtypes

| subtype                  | Normal-like | Triple negative/ basal-like | Luminal A | Luminal B | HER2-enriched | TOTAL |
|--------------------------|-------------|----------------------------|-----------|-----------|---------------|-------|
| Normal-like              | 24          | 99                         | 160       | 146       | 118           | 547   |
| Triple negative/ basal-like | 160         | 146                        | 118       | 118       | 99            | 547   |
| Luminal A                | 160         |                            | 146       | 118       |               | 424   |
| Luminal B                | 146         |                            | 118       | 99        |               | 363   |
| HER2-enriched            | 118         |                            | 99        | 547       |               | 764   |
| TOTAL                    | 547         |                            | 424       | 363       | 764           | 1674  |

![Figure 1](image-url)
3.2. Cancer Subtypes and Subnetworks of GBM

In this study, four subtypes have been identified through implantation of GBM in the co-clustering algorithm. Table 2 shows the obtained subtypes and its number of samples. Scientists from TCGA have published the finding of four distinct subtypes of GBM. They are Proneural, Neural, Mesenchymal and Classical subtypes [21, 26].

Table 2. Number of Samples for GBM Subtypes

|          | Proneural | Neural | Mesenchymal | Classical | TOTAL |
|----------|-----------|--------|-------------|-----------|-------|
|          | 51        | 64     | 43          | 44        | 202   |

According to this implementation, gene NPTX1 act as the target gene, and the subnetwork was produced to interpret the relationship among genes. Figure 2 (a) to (d) shows all the four subtypes generated. Figure 2 (a) concluded as Proneural subtypes. This subtype is very common among the young adults which are normally characterised by IDH/TP53 positivity [27, 28]. From iNCIS, 51 samples of Proneural subtype were obtained. In addition, proneural subtype mostly deriving from low-grade gliomas which are associated with better prognosis [27].

![NPTX1 subnetwork in GBM subtypes expression patterns](image)

Figure 2. NPTX1 subnetwork in GBM subtypes expression patterns. Direct connected genes to NPTX1 and genes targeting NPTX1’s downstream are shown. Circle colour shows the gene expression level; circle size is based on gene weight. (a) Proneural; (b) Neural; (c) Mesenchymal; (d) Classical
On average, Figure 3 shows optimal clusters. Hence the identification of five cancer subtypes of BRCA are concluded to be true. On average, Figure 4 shows optimal clusters. Hence the identification of four cancer subtypes of GBM are determined to be true.

3.3. Gene Analysis for BRCA and GBM

Apart from that, for the gene selection process, gene ranking plays an important role. The ranking is given to these genes based on proposed SinfFS-mSVM method. Although a produced gene has obtained highest rank among the gene list, it does not promise that this gene will be selected as one of the best gene.

The top ten (10) ranked genes listed in Table 3 and Table 4 are produced from gene ranking calculation as reported in proposed SinfFS+mSVM method. In addition, genes ranked from this list compared to genes’ $p$-value which is generated from t-test and ANOVA. The $p$-value from ANOVA is calculated for the whole data set of BRCA and GBM. Hence the obtained genes’ $p$-value is compared and selects the lowest $p$-value gene as the threshold value is 0.05 with the highest gene weight score.

### Table 3. Top 10 Genes for BRCA subtypes

|                | Overall | Normal-like | Triple negative | Luminal A | Luminal B | HER2   |
|----------------|---------|-------------|-----------------|-----------|-----------|--------|
| NPY1R          | COL17A1 | BCL11A      | CCL13           | IGF1*     | HOXB13*   |        |
| CEACAM6        | KRT5    | ABCB8*      | EGFR            | JAM2      | MYBL2     |        |
| TFAP2B         | SPRP1   | NAT1        | ESR1*           | LAMA2     | PKMYT1    |        |
| UGT2B11        | ID4     | GRPR        | IL18RAP         | NDN       | CDKN3     |        |
| SCGB2A2        | NFIB    | MLPH*       | LCK             | SLIT2*    | E2F1      |        |
| CBLN2          | KRT17   | CA12        | TBX21*          | RUNX1T1   | PLK1*     |        |
| ROPN1B         | OSR1    | SCUBE2      | PLAGI4A         | JAM3      | AURKA*    |        |
| AREG           | TRIM29  | GFRAL*      | SLAMFI          | BMX       | KIF4B*    |        |
| ROPN1          | EGFR    | ESR1        | PTX3            | CXCL12    | GTSE1     |        |
| PDZK1          | BIRC5   | ERBB4       | BCL11A          | COL14A1   | CX3CR1    |        |

*Red: significant genes; * found particularly in the subtypes

### Table 4. Top 10 genes for GBM subtypes

|                | Proliferative | Neural | Classical | Mesenchymal |
|----------------|---------------|--------|-----------|-------------|
| RPS4Y1         | CHD5          | PTPRJ  | ACTA2     | POLB        |
| LTF            | PGM1          | ACY1*  | LHX1      | CEP76       |
| IL8            | ADARB1        | CEP27  | ASGR1*    | NFATC4      |
| DKK1           | DMD           | CPS1   | AMFR*     | MAP3K14*    |
| EGFR           | OX2T          | PF4    | RPS15A    | ACTA2*      |
| PTX3           | ANGPTL4       | CTNAP2 | CYBA      | DNAJB5      |
| IL13RA2        | CYP7A         | DNAJB5*| CCT6A     | PFARQ       |
| FABP5          | EFI285        | THAP11 | INHA*     | RAB17       |
| CHI3L1         | BCL2L10       | PHKB   | RNGT1     | MYH2        |
| MOXD1          | PCLR          | RPL19  | RGS14*    | TCEG1       |

*Red: significant genes; * found particularly in the subtypes

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The first column of both Table 3 and Table 4 which is named as overall set of gene is produced through comparison of lowest ANOVA \( p \)-value and highest weight score. Whereas, the gene list for each subtype is attained from comparison of lowest \( p \)-value from t-test and highest weight score.

### 3.4. Pathway Analysis for BRCA and GBM

The pathways involved in each subtype of BRCA and GBM investigated. The pathway analysis conducted on gene selected for each subtype as shown Table 5 and 6 for BRCA and GBM respectively. Hence, GeneCards (https://www.genecards.org) and KEGG: Kyoto Encyclopaedia of Genes and Genomes (https://www.genome.jp/kegg/) used to identify the top significant pathways for each subtypes. The pathway analysis was conducted for the selected overall genes on both data sets.

Table 5 and 6, shows the top enriched pathways of the common genes and subtype-specific genes of each subtype from BRCA and GBM correspondingly. The pathways for both data sets have been classified into seven different types, which involve (i) cellular process, (2) metabolism, (3) environmental information processing, (4) nervous system, (5) cancers, (6) immune system, and (7) other organismal systems. The first column in both table shows the pathways obtained based on overall gene from BRCA and GBM data sets.

It can be noticed that the BRCA pathways are relatively different among different subtypes. The pathways for overall genes and samples (in the first column of the Table 5) are related to metabolism and cellular process. Pathways for triple-negative subtype are associated to cancers and metabolism which infers the growth of breast cell tumors. The pathways in subtype Luminal A related to cancers, metabolism and immune systems. Pathways in subtype Luminal B are highly related to cancers and metabolism type. While pathways from HER2 subtype are linked with cellular process and metabolism. Most of these pathways are having linked with breast cancer in breadth and depth.

**Table 5. Top 10 Pathways for BRCA Subtypes**

| Overall | Triple-negative | Luminal A | Luminal B | HER2 |
|---------|----------------|-----------|-----------|------|
| Signaling by GPCR | Development, Leptin signaling via PI3K-dependent pathway | Akt Signaling | ERK Signaling | Regulation of Androgen receptor activity |
| Hematopoietic Stem Cell Differentiation Pathways and Lineage-specific Markers | Integrated Breast Cancer Pathway | NF-kappa B signaling | Focal adhesion | Class I MHC mediated antigen processing and presentation |
| Apoptosis and Autophagy | Type II diabetes mellitus | ERK Signaling | Blood-Brain Barrier and Immune Cell Transmigration | Gene Expression |
| Integrated Breast Cancer Pathway | Metabolism | Hematopoietic Stem Cell Differentiation | Pathways in cancer | Cyclin A/B1 associated events during G2/M transition |
| Synapto genesis induction | Pharmacokinetics | Integrated Breast Cancer Pathway | Endometrial cancer | Cell Cycle, Mitotic |
| PKA-dependent signaling processes | Signaling by GPCR | IL12 signaling mediated by STAT4 | Adipogenesis and Cytokine Signaling in Immune system | Integrated Breast Cancer Pathway |
| PKG signaling processes | Akt Signaling | Immune response IL-23 signaling | Developmental Biology | Gdsti-1/CD106 Signaling |
| Regulation of CFTR activity (norm and CF) and Uricosurics Pathway, Pharmacodynamics | Signaling by Rho GTPases | Akt Signaling | T cell receptor signaling | retrograde transport |
| Activation of cAMP-Dependent PKA | Translational Control | Glucocorticoid receptor regulatory network | G-Beta Gamma Signaling | Developmental Biology |
| Metabolism | CDK-mediated phosphorylation and removal of Cdk6 | p38 MAPK Signaling | Apoptotic cleavage of cellular proteins | Regulation of beta-cell development |
| Environmental information processing | Cellular process | Immune system | Other organismal system |
| Cancer | | | |
| Nervous system | | | |
4. CONCLUSION

From the results, it can be concluded that, there are five subtypes from BRCA and four subtypes from GBM were successfully identified. The iNCIS algorithm is able to produce simple subnetwork to show gene expression in each subtype. By the feature selection and classification, it has been able to prioritize significant genes for each subtype of both data sets which are analyzed for disease prognosis and diagnosis.

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