A STUDY AND ANALYSIS OF CLUSTERING ALGORITHMS ON HIV-1 INFECTION MICROARRAY DATASET FOR FINDING CLUSTER WISE COMMON GENES

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Abstract: Data mining refers to collecting or mining knowledge from large amounts of data. It is used in various medical applications like tumor clustering, protein structure prediction, gene selection, cancer classification based on microarray data, clustering of gene expression data, statistical model of protein-protein interaction etc. The analyzing the clustering algorithms phase consist of four clustering algorithms namely K-means, Fuzzy c-means, Hierarchical algorithm and Partitioning Around Medoids(PAM) on HIV – 1 infection effect on macrophages in vitro time course microarray data set. The clustering algorithms are validated using validation measures and based on internal validation measures such as Dunn index, Dunn index 2, Calinski-Harabasz index and Average Silhouette width, the best clustering algorithm out of 4 is to be identified and finally the proposed research work is also to find common genes present in each cluster produced by the four clustering algorithms.

Keywords: Data mining, Microarray, Preprocessing, Clustering algorithm, Finding common genes cluster wise

I. INTRODUCTION

Data mining is often defined as finding hidden information or extracting meaningful information from large database. The aim of data mining is to discover structure inside unstructured data, extract meaning from noisy data, discover patterns in apparently random data, and use all this information to better understand trends, patterns, correlation in existing data [1]. Clustering is task of grouping a set of objects in such way that objects in the same group called a cluster. Clustering is unsupervised learning problem so as every problem it deals with finding structure in a collection of unlabelled data. This model are sometimes called descriptive model. Clustering is one of the most common untested data mining methods that explore the hidden structures embedded in a dataset. Data Mining is the process of automatic discovery of novel and understandable models and patterns from large amounts of data [2]. Bioinformatics can be defined as the application of computer technology to the management of biological information. It is the science of storing, extracting, organizing, analyzing, interpreting and utilizing information from biological sequence and molecules. The primary goal of bioinformatics is to increase the understanding of biological processes. Bioinformatics is the science of managing, mining, and interpreting information from observations of biological processes [3].

Various genome projects have contributed to an exponential growth in DNA and protein sequence databases, and advance in high-throughput technology such as microarrays and mass spectrometry have further created the fields of functional genomics and proteomics, in which one can monitor quantitatively the presence of multiple gene, proteins, metabolites and compounds in a given biological state [4]. Microarray is used for gene expression analysis. It comprises of a tiny membrane or glass slide having sample of many regularly arranged genes. Microarray analysis can detect thousands of genes in a small sample along with the expression of those genes. Microarray Datasets are often characterized by high-dimensions and small samples [5].

The paper organized as follows: section 1 describes the introduction on data mining, section 2 describes the literature review, section 3 describes the methodology of the proposed work, section 4 describes result and discussion and finally the paper is concluded in section 5.

II. LITERATURE REVIEW

analysis, Gene expression datasets of four different cancers are chosen such as Breast Cancer, Leukemia, Lung Cancer and Carcinoma Cancer. Cluster gram is used for automatically clusters genes and samples based on gene expression patterns. Proposed algorithm is compared with rough K-means and K-means algorithms. Comparison based on the elapsed time of three clustering algorithms. K-means take less elapsed time compare to other two algorithms. Fuzzy soft Rough K-means has more internal process but presents better results in less time. Number of iterations for convergence rate is reduced with our proposed algorithm (9, 7, 11, and 3 for Leukemia, lung, carcinoma and breast cancer). These two are used for validation measures namely Xie-Beni index and DB-index. In DB index has less value for the proposed algorithm compare to other two clustering algorithms for all datasets (0.0673, 0.0154, 0.1064 and 0.1135 for Leukemia, lung, carcinoma and breast cancer). The Xie-Beni also less with our proposed algorithm (0.3573, 0.0874, 0.2482 and 0.1274 for Leukemia, lung, carcinoma and breast cancer).

Porkodi Rajendran, Deepika Thangavel, 2016 [7] has presented a work to identify gene enrichment analysis of clustered genes from asthma microarray dataset using k-means clustering algorithm. It follows two strategies that are clustering gene expression profile using K- Means clustering algorithm and conducting Gene Enrichment Analysis using Gene Ontology (GO) to identify the enriched GO terms in each cluster. The microarray dataset contains 41,000 genes and out of that only 9,425 genes. After processing it produces the 3 sets of clusters such as four clusters set, six clusters set and eight clusters set. In the first cluster set with 4 clusters have the number of genes 1720, 2636, 2458 and 2611 respectively. Similarly, the other two cluster sets numbers of genes are identified. The gene enrichment analysis is conducted for each cluster in the cluster set based on top most significant molecular functions GO terms enriched in each cluster.

Yanchi Liu, Zhongmou Li, et al. 2010 [8] has of the investigated the validation properties of a suite for 11 existing internal clustering validation measures (RMSSTD, RS, T, CH, ID, S, DB, XB, SD and S-Dbw) for hard clustering in five different aspects namely monotonicity, noise, density, subclusters and skewed distributions. From the experiment results, most of the existing measures have certain limitations in different application scenarios. Finally, the experiment results show that S-Dbw is the only internal validation measure which performs well in all five aspects, while other measures have certain limitations in different application scenarios, mainly in aspects of noise and sub-clusters.

Sarah M. Kim, Matthew I. Pena et al. 2016 [9] has evaluated the ability of three separate clustering methods like hierarchical, k-means, and k-medoids along with three pair-wise distance measures such as Levenshtein, Jaccard, and n-gram to expertly group lysine, isoleucine, and 3-hydroxypropionic acid (3-HP) biosynthesis pathways. From the experimental results it analyze that average Silhouette width was also lower for 3-HP (0.12) than for those for lysine (0.55) and isoleucine (0.22). For Silhouette width and to a lesser extent Dunn index, clustering using Jaccard distance had the lowest values across the three pathways. Levenshtein distance showed the most stability, with values above 0.9 across all clustering methods. Isoleucine clusters using n-gram distance showed the most stability, with values close to 0.9. For 3-HP clusters, there was no one distance measure that resulted in high stability values; And conclude that though there was no one combination of clustering methods and distances that performed well across all three example biosynthesis pathways, some clustering methods and distance measures namely hierarchical clustering and Levenshtein distance resulted in clusters that matched the expected external. This suggests that an automated method for grouping pathways is possible because it gives in the form of directed approach.

Eréndira Rendón, et al. 2011 [10] has presented a comparison between two clustering validity index approaches that are internal and external. For analyses it uses four external indexes (F-measure, NMIMeasure, Entropy, Purity) and six internal indexes (BIC, CH, DB, SIL, NIVA and DUNN). Totally 13 datasets were used, which were clustered using the K-means and Bissection K-means algorithms. For Bissection K-means algorithm using internal indexes, correctly classified group numbers are obtained as 86% of the time, and in 51.9% when external indexes were used. When clusters of the K-means algorithm were clustered using internal indexes as 76.9% of accuracy was obtained and 61.5% with external indexes. From which the internal indexes are more specific in real group number determination than external indexes, or at least with the used datasets.

Satya Chaitanya Sripada, M.Sreenivasa Rao, 2011 [11] has compared the for K-means and Fuzzy C means clustering using the validity measure as Purity and Entropy. It is medical data for evaluating external measures. For the medical data sets the Fuzzy C means and K-means algorithms were framed and the clusters were generated. The values show that the Fuzzy C Means has high value of purity and low value of entropy. This means a good clustering. The K-means has lower value of purity and high value of entropy compared to Fuzzy C Means. The Fuzzy C means clustering is more suited for medical data sets when compared to K means.

III. METHODOLOGY

The framework of the proposed research work is shown in Fig.1. The framework consists of four phases; Pre-processing, Analyzing K-means, fuzzy c-means, Hierarchical algorithm and PAM using HIV-1 infection effect on macrophages in vitro time course microarray data set. The results of clustering algorithms are validated using
certain validation measures and based on internal validation measures such as Dunn index, Dunn index 2, Calinski-Harabasz index and Average Silhouette width, the best clustering algorithm out of 4 is to be identified. Finally the proposed research is also to find common genes present in each cluster produced by the four clustering algorithms.

**Fig.1 Methodology of the Proposed Work**

A. **Dataset**

The HIV-1 Infection dataset is downloaded from GEO Database Browser and used for the proposed work. The dataset is GEO series with 12625 probe IDs and 16 samples with their corresponding information.

B. **Preprocessing**

The preprocessing is required to remove noisy data or irrelevant data from the dataset. In addition to that, there are some other preprocessing steps are to be considered based on the type and nature of database. As handling of microarray dataset in R is concerned, there are two steps are important one is converting GEO dataset into Expression set object and the other is to convert expression values in the dataset to log2 normalized form. Then the Proposed research work has done two other preprocessing tasks which are filtering and standardization. Filtering step to remove genes which are expressed at low levels or show only small changes in expression. Expression values of genes are standardized to have a mean value of zero and a standard deviation of one.

C. **Analyzing clustering Algorithms**

This section describes the basic idea behind in k-means, fuzzy c-means, hierarchical clustering and PAM.

D. **K-means algorithm**

The algorithm assigns each individual data point to one of the clusters in a random fashion. The underlying idea of the algorithm is that a good cluster is the one which contains the smallest possible with-in cluster variation of all observations in relation to each other [12]. The most common way to define this variation is using the squared Euclidean distance. K-Means clustering is to decide on the
numbers of clusters to use. There numbers of clusters to decide which options is the most useful, applicable or interpretable solution.

E. Fuzzy- c-means algorithm
A major problem in applying the FCM method for clustering microarray data is the choice of the fuzziness parameter $m$. We show that the commonly used value $m = 4$ is not appropriate for some data sets, and that optimal values for $m$ vary widely from one data set to another. By setting threshold levels for the membership values, genes which are tightly associated to a given cluster can be selected. Using a brain cancer data set as an example, we show that this selection increases the overall biological significance of the genes within the cluster [13].

F. Hierarchical algorithm
Hierarchical of clusters is often graphically represented by a dendrogram A dendrogram captures the process by which a hierarchical clustering is generated by showing the order in which clusters are merged (bottom-up view) or clusters are split (top-down view). One of the attractions of hierarchical techniques is that they correspond to taxonomies that are very common in the biological sciences. Instead, any desired number of clusters can be obtained by ‘cutting’ the dendrogram at the proper level. Also, hierarchical techniques are sometimes thought to produce better quality clusters [14].

G. PAM
The PAM (Partioning Around Medoids) algorithm, also called the K-medoids algorithm, represents a cluster by a medoid. Initially, the number of desired clusters is input and a random set of k items is taken to be the set of medoids. Then at each step, all items from the input dataset that are not currently medoids are examined one by one to see if they should be medoids. That is, the algorithm determines whether there is an item that should replace one of the existing medoids [15].

H. Validating the cluster Results
The evaluation of the performance of the above four algorithm are done using cluster validation measures. In general, clustering validation can be categorized into two classes, external clustering validation and internal clustering validation. The Proposed research work uses four widely used internal cluster validation measures to validating the results of clustering algorithms.

I. Dunn index
The Dunn index defines the ratio between the minimal intracluster distances to maximal intercluster distance [16].

$$\text{Dunn index} = \frac{d_{\text{min}}}{d_{\text{max}}} \quad (1)$$

Where $d_{\text{min}}$ denotes the smallest distance between two objects from different clusters and $d_{\text{max}}$ denotes the largest distance of two objects from the same cluster. The Dunn index is limited to the interval [0,∞] and should be maximized.

J. Calinski-Harabasz Index
Calinski-Harabasz Index is the math formula to the measure [17].

$$\frac{SS_B}{SS_W} \times \frac{N-K}{K-1} \quad (2)$$

Where $K$ is the number of clusters and $N$ is the total number of observations (data points), $SS_W$ is the overall within-cluster variance equivalent to the total within sum of squares calculated above), $SS_B$ is the overall between-cluster variance.

K. Average Silhouette width
The silhouette width is calculated by averaging all the silhouette values for each pathway, where the silhouette value is calculated using the following function [18].

$$s(i) = \frac{b(i) - a(i)}{\max(a(i), b(i))} \quad (3)$$

Observations with a large $s(i)$ (almost 1) are very well clustered, a small $s(i)$ (around 0) means that the observation between two clusters, and observations with a negative $s(i)$ are probably placed in the wrong cluster.

L. Finding of common genes- cluster wise
The proposed research work takes clustering results produced by all the specified algorithm as input and identify the common genes present in different clusters of the four clustering algorithm. The four clustering algorithms produced 4 clusters uniformly c1, c2, c3 and c4. Then the common genes present in cluster 1 of all algorithms are identified using the below intersection function.

$$\text{CommonGene} = \sum_{c_1=1}^{4} (\text{FCM}_{c_1} \cap \text{KM}_{c_1} \cap \text{HC}_{c_1} \cap \text{PAM}_{c_1})$$

Similarly the common genes present in cluster 2, cluster3 and cluster 4 are identified.

IV. RESULTS AND DISCUSSION

A. Result of Preprocessing
The experimental dataset consists of 12625 genes in which 3956 genes are excluded through filtering and 2871 genes are excluded through standardization function.

B. Results of clustering algorithms
The K-means algorithm, Fuzzy c-means Algorithm, Hierarchical algorithm and PAM successfully produced 4 clusters and the number of genes in each cluster is represented in Table 1.
Table 1. Clusters by clustering algorithms

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Cluster 1</th>
<th>Cluster 2</th>
<th>Cluster 3</th>
<th>Cluster 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>K-Means</td>
<td>1356</td>
<td>1131</td>
<td>1326</td>
<td>1985</td>
</tr>
<tr>
<td>Fuzzy c-means</td>
<td>1970</td>
<td>815</td>
<td>1481</td>
<td>1532</td>
</tr>
<tr>
<td>Hierarchical</td>
<td>2205</td>
<td>1347</td>
<td>1724</td>
<td>522</td>
</tr>
<tr>
<td>PAM</td>
<td>2379</td>
<td>1261</td>
<td>1081</td>
<td>1077</td>
</tr>
</tbody>
</table>

C. Result of k-means Algorithm

In Fig. 2, cluster 1 shows the overall picture on how the genes present in this cluster are expressed. It is observed that the genes in this cluster uniformly expressed between -1 and +1 expression pattern. Similarly cluster 2, cluster 3, and cluster 4 has followed the gene expression patterns between -1 and +1, 0 and 1 and -2 & +1 respectively.

The 2D representation of k-means clusters solution is depicted in Fig. 3.

D. Result of Hierarchical Algorithm

The genes have been organized into a tree, leaves represent genes and length of the paths between leaves represents distances between genes. Similar genes lie within same subtrees. The number of cluster to be formed is set as four and ward method. The Hierarchical clustering results are shown in dendrogram in Fig. 6.

The 2D representation of Hierarchical cluster solutions is depicted in Fig. 7.
E. Result of PAM algorithm

The PAM clustering algorithm has produced four clusters based on Average Silhouette Width. The Cluster 1 grouped 2379 genes of average silhouette width 0.14. Similarly Cluster 2, Cluster 3 and Cluster 4 grouped 1261 genes of average silhouette width 0.11, 1081 genes of average silhouette width 0.09 and 1077 genes of average silhouette width 0.08 respectively as shown in Fig. 8.

The 2D representation of partitioning around medoids cluster solutions is depicted in Fig. 9.

F. Result of cluster Validation

The proposed research work has validated the performance of four clustering algorithms using internal cluster validation measures such as Dunn index, Dunn index 2, Calinski-Harabasz index and Average Silhouette width. The result of these validation measures are represented in Table 2 and the same is depicted using bar chart in Fig. 10 (a), (b) & (c).

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>K-means algorithm</th>
<th>Fuzzy c-means algorithm</th>
<th>Hierarchical algorithm</th>
<th>Partitioning Around Medoids</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dunn index</td>
<td>0.115987</td>
<td>0.04933466</td>
<td>0.1270628</td>
<td>0.127322</td>
</tr>
<tr>
<td>Dunn index 2</td>
<td>1.064114</td>
<td>1.108705</td>
<td>0.986741</td>
<td>1.037815</td>
</tr>
<tr>
<td>Calinski-Harabasz index</td>
<td>789.841</td>
<td>794.0111</td>
<td>589.629</td>
<td>755.5447</td>
</tr>
<tr>
<td>Average Silhouette width</td>
<td>0.1179472</td>
<td>0.1234762</td>
<td>0.0641567</td>
<td>0.1109011</td>
</tr>
</tbody>
</table>

The experiment results are validated using internal validation measures which performs well in all four aspects, while other measures have certain limitations in different clustering algorithms. The best resulted clusters, of the experimental dataset are compared based on compared the high values of Dunn index, Dunn index 2, Calinski-Harabasz index and Average Silhouette width values which are 0.1270628, 1.108705, 794.0111, and 0.1234762 respectively. From the above inference identified that fuzzy c-means clustering algorithm gives better clustering accuracy then other three clustering algorithms.
**Finding of common genes-cluster wise**

The common genes present in each cluster of all the four clustering algorithms have been identified using basic intersection function and the result has been tabulated in Table 3.

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>C1</th>
<th>C2</th>
<th>C3</th>
<th>C4</th>
<th>No. of Common genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>K-means</td>
<td>1356</td>
<td>1131</td>
<td>1326</td>
<td>1985</td>
<td>22</td>
</tr>
<tr>
<td>Fuzzy c-means</td>
<td>1970</td>
<td>815</td>
<td>1481</td>
<td>1532</td>
<td>0</td>
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<td>2205</td>
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<td>15</td>
</tr>
<tr>
<td>PAM</td>
<td>2379</td>
<td>1261</td>
<td>1081</td>
<td>1077</td>
<td>0</td>
</tr>
</tbody>
</table>

The experimental results of this phase has identified 22 genes as common genes present in cluster 1 of K-means, fuzzy c-means, hierarchical and PAM with its size 1356, 1970, 2205 and 2379 respectively. Similarly 15 genes has been identified as common genes present in cluster 3 of K-means, fuzzy c-means, hierarchical and PAM with its size 1326, 1481, 1724 and 1081 respectively. The research work has also found that there is no common genes present in cluster 2 and cluster 4 of all the four algorithms. The cluster wise common genes found have been represented in Table 4.
Table 4. Common genes in cluster 1 and cluster 3

<table>
<thead>
<tr>
<th>Cluster Number</th>
<th>No. of genes</th>
<th>Common genes in cluster 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cluster 1</td>
<td>22</td>
<td>AGPAT1, AMFR, BSG, CSPG4, DMD, GRPR, KCNMB1, LONRF1, MAP3K11, MDH2, MTHFR, MYO1C, OBSL1, PARVB, PLS1, RIN1, SLC29A2, TIMP2, TRPM2, TUBB2B, VGF, XIST.</td>
</tr>
<tr>
<td>Cluster 3</td>
<td>15</td>
<td>AHSA1, CSF2RB, DYRK2, FAM53B, FTO, HNRNPA0, MGAT2, NOP2, PARP4, PIK3C2A, PPP1R8, RFNG, RPS6KA3, SNAPC5, USP.</td>
</tr>
</tbody>
</table>

V. CONCLUSION AND FUTURE WORK

Data mining in Bioinformatics can be defined as the application of computer technology to the management of biological information. The primary goal of bioinformatics is to increase the understanding of biological processes. The proposed research work consists of four main phases namely preprocessing phase, Analyzing Clustering algorithms phase, validation Phase and Finding of common genes-cluster wise have been successfully implemented in R. The analysing the clustering algorithms phase applied four clustering algorithms namely K-means, Fuzzy c-means, Hierarchical algorithm and Partitioning Around Medoids on HIV – 1 infection effect on macrophages in vitro time course microarray data set. The results of the above clustering algorithms have been validated using certain validation measures and based on these measures; it is found that Fuzzy C-Means clustering algorithm gives better clustering accuracy than the other three clustering algorithms. Finally, the proposed research work have also found common genes present in each cluster of four algorithms.

In future, the research may be extended by considering hybrid approach to improve the clustering accuracy.

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