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# Partial Context Similarity of Gene/Proteins in Leukemia Using Context Rank Based Hierarchical Clustering Algorithm

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Article Info	ABSTRACT		
Article history:	In this paper we proposed a method which avoids the choice of natural		
Received Dec 31, 2014	language processing tools such as pos taggers and parsers reduce the processing overhead. Moreover, we suggest a structure to immediately create		
Revised Feb 24, 2015	a large-scale corpus annotated along with disease names, which can be		
Accepted Mar 16, 2015	applied to train our probabilistic model. In this proposed work context rank based hierarchical clustering method is applied on different datasets namely colon, Leukemia, MLL medical diseases. Optimal rule filtering algorithm is applied on these datasets to remove unwanted special characters for gene/protein identification. Finally, experimental results show that proposed method outperformed existing methods in terms of time and clusters space.		
Keyword:			
Biomedical			
Clustering			
Gene/protein			
Machine learning			
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## 1. INTRODUCTION

Life science studies are characterized by the construction of large and heterogeneous patterns of biological study, including protein or gene series. Therefore, a number of methods based upon text-mining have been used to improve the identify protein and genes names in medical texts. Text mining has been defined as the discovery by computer of recent, previously unknown, data by automatically extracting data from different written resources. Machine learning means the development and study of systems that could learn from data. This is actually a technique of teaching computers in order to make and enhance behaviors based on some data. Machine learning is a huge field with hundreds of algorithms for addressing different issues. Machine learning provides challenging problems in terms of algorithmic approach, data representation, computational effectiveness, and quality of the resulting program. Biomedical data along with its updates are saved in natural language style. Due to the enhanced amount of biomedical sources, it is becoming more and more challenging to find useful and relevant information regarding a specific topic. All research inventions come and enter the repository at high-rate, making the strategy of finding out and disseminating quality information a very difficult task. Manual assessment of such large amount of data will probably be very difficult and time-consuming. The issue is further magnified by the consumption of large evaluation measures, and datasets that contain essentially different annotation formats and task definitions.

Medical text documents continuously hide valuable structured data. For example, a collection of newspaper content will contain details on the location of the head-quarters of various entities. If we need to find the position of the head-quarters of, say Microsoft we could try and utilize conventional data retrieval techniques for discovering documents that contain the answer on the present query. An application of systems biology is to uncover the bio-processes underlying the patterns of a cell. Relationships within genes encode most of this data and are occasionally discovered and symbolized as key products. Understanding these relationships is an extremely challenging issue as even the simplest organisms contain variety genes that interact in complex combinations to deal with ecological circumstances. Another complicating element is current high throughput technique designed to determine the activity level of genes is extremely noisy [8]. As there exists very few well understood genetic activities, unsupervised clustering is a common first step to understand these data.

The clustering procedure is a basic tool to organize a collection of objects within a metric space into a set of smaller partitions called clusters. By using clusters, the representation of the object pool can be made easier and the computation expense of data management can be reduced. The created clusters can be used to introduce rules of top levels describing the common characteristics of data objects. In the case of grammar induction structures, the rules of grammar are stated on word classifications as the words within the same category are transformed similarly. If word categories are known, grammar principles might be explored in a better way.

Nearest neighbor is a machine learning method introduced in the literature that often learns by comparing each individual new case to prior examples. Machine learning is definitely an area of artificial intelligence focusing on the development of approaches which permit computers to learn. More clearly, machine learning is a method for generating computer programs for the evaluation of datasets. Instance based learning, of which nearest neighbor is a subset, is a branch of machine learning techniques; other branches include: rule based genetic algorithms, ANN and support-vector-machines.

In the whole nearest neighbor algorithm, all tuples are generally saved in memory during data training. When a new query instance is accepted the memory is searched to find the instance that suits the query instance most closely. Nearest neighbor will then infer that the concept label of the query instance is similar as the notion label of the most similar instance stored in memory.

Noise present in data is a significant challenge avoiding machine learners away from being more quality, or applicable to the large selection of domains. Noise is an incorrect attribute or model value information which can be a effect of errors in manual data entry, compilation, measurement or corruption of data. If the potential for noise is certainly not recognized, this can lead to machine learning algorithms fitting the noise. Fitting the noise happens when the machine learner learns the noisy data as if were not noisy information. Noise will often make instances in memory oppose one another.

#### 2. RESEARCH METHOD

Following are the limitations of the related work discussed in this section. Eliminate the Non-Functional Characters

- Apply Heuristic Policies to Remove Non-Functional Symbols
- Remove and replace the following symbols with gaps: #â€e? \$&\*ó@|~!\
- Remove the subsequent characters if they are followed by a space: ;: .,
- Eliminate the following pairs of brackets if the open bracket is preceded by a space and the closed bracket is followed by a space: [] ()
- Eliminate the single quotation symbol if it is associated with by a space or if it is preceded by a space.
- Remove s and t if they are associated with by a space
- Eliminate slash / if it is associated with by a space.

Our proposed work overcomes all these limitations. We take three biomedical disease datasets offline to extract hidden patterns using feature extraction and hierarchical clustering approaches. Each dataset is preprocessed to remove non-functional characters to identify disease names by using gene/protein database. Hierarchical methods for supervised and unsupervised datamining give multilevel indexing of data. It can be relevant for several applications associated to data extraction, patterns retrieval and data organization.

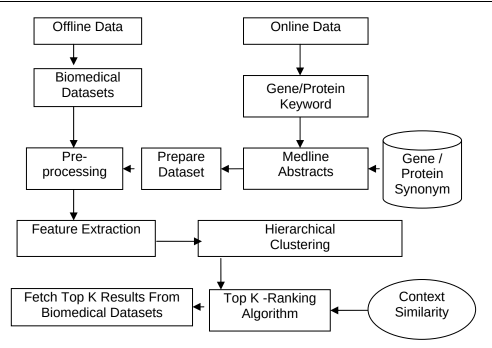


Figure 1. Proposed method for eliminating the Non-Functional Characters.

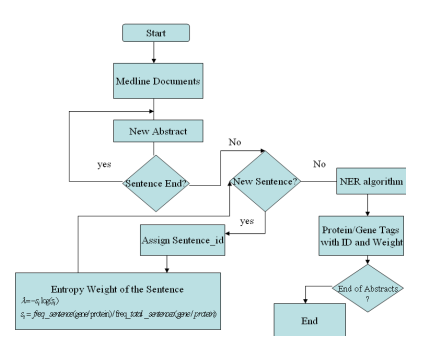


Figure 2. Proposed method flow chart for eliminating the Non-Functional Characters

#### **Hierarchical Clustering Algorithm:**

Input : Name entity Gene/Protein tags Tgp using NER approach,Gene/Protein DB, Probability P, Classes
Positive pos, Negative neg, Tokenset Tk, Sentenceset Sen .
Read k, Threshold, Entropy weight;
Output: Quality k- abstracts.
Tgp=Get(Name \_Entity\_Gene/ Protein\_Tags)
for each tg in Tgp
For each in Tk
Calculate tag probability
List.add(tg)

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List.add() count=count+1 end end. For each token t in Tk For each sen in Sentenceset If((t \_Sen)&& (t Tgp)&&( >getProb(t)) List Data ← Sentence\_id,token, Pmid, Entropy\_weight,Synonyms,Data,Title,PositiveClass Else

List Data ← Sentence\_id,token, Pmid, Entropy\_weight,Synonyms,Data,Title,NegativeClass End

End

For each pair of objects in Data

Calculate distance between two objects as

$$D(c1_i, c2_j) = (1 - \mathbf{r}_{ij}) * 0.5$$
  
$$r_{ij} = (\sum_{i=0}^d (c1_{ij} - \overline{c1_i})(c1_{ji} - \overline{c1_j})) / \sqrt{\sum_{i=1}^d (c1_{ij} - \overline{c1_i})^2 \sum_{i=1}^d (c1_{ji} - \overline{c1_j})}$$

6.

a. Start with the disjoint clustering that have level as 0 and sequence\_number m = 0.

b. Rank the pairs from smallest distance (similarities in common) to the maximal distance.

c. Calculate and count pairs, say n pairs.

If  $n \ge 0$ 

do,

c.1 Explore the median as root hierarchical node.

c.2 Split the pairs as left and right side branches based on the median.

c.3 Explore the smallest unlike pair of clusters in the leftside and rightside current clustering, say pair rs, ls according to  $d[(rs),(ls)] = \min r[(i),(j)]$  in which the minimum value is taken over all pairs of clusters in the current clustering.

c.4 If leftside and rightside have atleast one similar object. In this case merge it collectively in one cluster, and look up smallest value over all pairs of clusters in the currentclustering.

Else

c.5 Find the maximal dissimilar pair of clusters in the leftside and rightside current clustering, say pair rs, ls according to  $d[(rs),(ls)] = \max r[(i),(j)]$  in which the m value is taken over all pairs of clusters in the current clustering.

d. Increment the sequence number: m = m + 1. (In both left and right sides) Merge clusters (r) and (s) into a single-cluster to form the subsequent cluster m. Place the level of this cluster to L(m) = r[(r),(s)]

e. Revise the tree, T, by eliminating the nodes corresponding to clusters (p) and (q) and adding a node corresponding to the newly composed cluster. The neighborhood betweenthe new cluster, denoted (p,q) and old cluster (m) is stated in this way:

 $d[(m), (p,q)] = \min r[(m),(p)], d[(m),(q)].$ 

If d < 0

Then

#### Minimum Variance:

The distance between two clusters is defined as the increase in the sum of squared errors (SSE) when the two clusters are merged. The SSE for a given cluster  $C_i$  is given as:

$$SSE_i = \sum_{\mathbf{x}_j \in C_i} \|\mathbf{x}_j - \boldsymbol{\mu}_{C_i}\|^2$$

and the SSE for a clustering  $C = \{C_1, \dots, C_m\}$ , is given as:

$$SSE = \sum_{i=1}^{m} SSE_i = \sum_{i=1}^{m} \sum_{\mathbf{x}_j \in C_i} \|\mathbf{x}_j - \boldsymbol{\mu}_{C_i}\|^2$$

When we merge  $C_i$  and  $C_j$  into  $C_{ij}$ , the change in the SSE involves these three clusters, and is given as:

$$\Delta SSE_{ij} = SSE_{ij} - SSE_i - SSE_j$$

Plugging in into the equation above, after simplification, we thus obtain the distance between the two clusters as:

$$\delta(C_x, C_y) = \Delta SSE_{xy} = \left(\frac{|C_x| \cdot |C_y|}{|C_x| + |C_y|}\right) \|\boldsymbol{\mu}_{C_x} - \boldsymbol{\mu}_{C_y}\|^2$$

f. If all objects are in one cluster, stop. Else, go to step b.
Algorithm2:
Input : Hierarchical clusters from top to bottom
Output: Top K Disease Results.
6.1 For each cluster in Cluster-set
6.1.1 t1=gene/protein search keyword.

6.1.2 For each synonym in the cluster

#### t2=synonym.

Find context similarity between t1 and t2.

Context Similarity Score:

End for

 $\sum_{t1 \in cluster} Cos(t1, t2) / \prod_{i=1}^{m} size of (cluster_i)$ t2∈keyword

6.2 Sort <t1,t2> according to context similarity score.

6.3 Get abstracts from biomedical databases according to tag pair score.

Table 1. The Performance of			
Variable	Speed (rpm)	Power (kW)	
х	10	8.6	
у	15	12.4	
Z	20	15.3	

#### 3. RESULTS AND ANALYSIS

Input		×
?	Enter file name Leukemia OK Cancel	

Figure 3. Loading leukemia disease data

```
Partial Context Simiarity of Gene/Proteins in leukemia:
Context Simiarity %5.3f===>0.2526455026455026
```

```
<=== U19107_rna1_at ===> synonyms are ZNF127 (ZNF127) gene
        Context Simiarity %5.3f===>0.3436507936507936
<=== U19142_at ===> synonyms are GAGE1 G antigen 1 (GAGE-1)
        Context Simiarity %5.3f===>0.4829059829059829
<=== U19180_at ===> synonyms are BAGE B melanoma antigen
        Context Simiarity %5.3f===>0.4363929146537842
<=== U19261 at ===> synonyms are Epstein-Barr virus-induced protein mRNA
        Context Simiarity %5.3f===>0.2578347578347578
<=== U19345_at ===> synonyms are AR1 protein (AR) mRNA
        Context Simiarity %5.3f===>0.43915343915343913
<=== U19487_at ===> synonyms are Prostaglandin E2 receptor mRNA
        Context Simiarity %5.3f===>0.26296296296296295
<=== U19517_at ===> synonyms are (apoargC) long mRNA
        Context Simiarity %5.3f===>0.38791423001949316
<=== U19523_at ===> synonyms are GCH1 GTP cyclohydrolase 1 (dopa-responsive dystonia) {alternative
products}
        Context Simiarity %5.3f===>0.41629629629629633
<=== U19718 at ===> synonyms are MFAP2 Microfibrillar-associated protein 2
        Context Simiarity %5.3f===>0.3785004516711834
<=== U19796_at ===> synonyms are Melanoma antigen p15 mRNA
        Context Simiarity %5.3f===>0.43407407407407406
```

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<=== U19878 at ===> synonyms are Transmembrane protein mRNA Context Simiarity %5.3f===>0.43304843304843305 <=== U19906 at ===> synonyms are VASOPRESSIN V1A RECEPTOR Context Simiarity %5.3f====>0.0 <=== U19948\_at ===> synonyms are Protein disulfide isomerase (PDIp) mRNA Context Simiarity %5.3f===>0.2578347578347578 <=== U19977\_at ===> synonyms are Preprocarboxypeptidase A2 (proCPA2) mRNA Context Simiarity %5.3f===>0.42407407407407405 <=== U20158 at ===> synonyms are 76 kDa tyrosine phosphoprotein SLP-76 mRNA Context Simiarity %5.3f===>0.42328042328042326 = U20230\_at ===> synonyms are "GB DEF = Guanyl cyclase C gene, partial cds" <=== U20240\_at ===> synonyms are "CEBPG CCAAT/enhancer binding protein (C/EBP), gamma" Context Simiarity %5.3f===>0.4199860237596087 <=== U20285 at ===> synonyms are Gps1 (GPS1) mRNA Context Simiarity %5.3f===>0.0 <=== U20325\_at ===> synonyms are Cocaine and amphetamine regulated transcript CART (hCART) mRNA Context Simiarity %5.3f===>0.41816009557945044 <=== U20350 at ===> synonyms are CMKRL1 Chemokine receptor-like 1 Context Simiarity %5.3f===>0.38078703703703703 <=== U20362 at ===> synonyms are Tg737 mRNA Context Simiarity %5.3f===>0.4037037037037037037 <=== U20391\_rna6\_at ===> synonyms are Folate receptor (FOLR1) gene Context Simiarity %5.3f===>0.32936507936507936 <=== U20428\_at ===> synonyms are SNC19 mRNA sequence Context Simiarity %5.3f===>0.0 <=== U20530\_at ===> synonyms are GB DEF = Bone phosphoprotein spp-24 precursor mRNA Context Simiarity %5.3f===>0.37703703703703706 **Correlation Distance Metric:** Correlation Distances: 0.5246662304909925 Correlation Distances: 0.5619362422999764 Correlation Distances: 0.6513947712224407 Correlation Distances: 0.48759512587181975 Correlation Distances:0.5319049159237761 Correlation Distances: 0.5246662304909925 Correlation Distances: 0.5619362422999764 Correlation Distances: 0.6513947712224407 Correlation Distances: 0.5319049159237761 Correlation Distances: 0.5246662304909925 Correlation Distances: 0.5619362422999764 Correlation Distances: 0.6513947712224407 Correlation Distances: 0.5319049159237761 Correlation Distances: 0.5619362422999764 Correlation Distances: 0.6221864849879517 Correlation Distances: 0.6058234775837336 === Clustering stats for training data === **Clustered Instances** 11 (92%) 0 1 (8%) 1 === ACCURACY DETAILS=== TOTAL GENE DETECTION ACCURACY 12 100 % ERROR RATE OF PROPOSED ALGORITHM 0 0 % Correlation Efficiency 1 Total Number of Instances 12

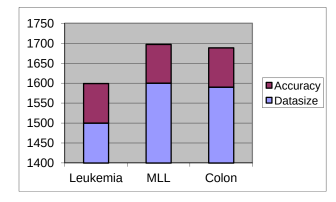


Figure 4. Comparision between datasize and accuracy in different datasets

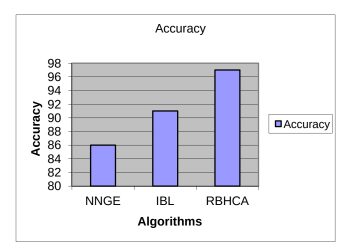


Figure 5. Comparison between proposed and traditional algorithms for leukemia dataset

### 4. CONCLUSION

In this proposed work context rank based hierarchical clustering method is applied on different datasets namely colon, Leukemia, MLL medical diseases. Optimal rule filtering algorithm is applied on these datasets to remove unwanted special characters for gene/protein identification. This work overcomes some of the limitations in the literature such as : noise elimination in medical datasets, robustness, high disease prediction rate, high quality cluster result with less search space and high true positive rate. Finally, experimental results show that proposed method outperformed well in terms of time and clusters search space are concerned. In future this work can be extended to implement similar disease clusters on online medical documents like medline, pubmed etc.

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