Knowledge Discovery in Protein Sequence Analysis Using Hierarchical Clustering Method

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ABSTRACT

Introduction: New Data Mining techniques play a very important role in the large growing biological databases. The clustering technique is an unsupervised method in data mining. Hierarchical Clustering techniques are useful to represent relationships between protein families.

Objective: Bioinformatics urges the need of discovering knowledge in the vast area of molecular biology by using data mining as the core. Data Mining aims to discover hidden data from a large volume of data.

Method: This paper discusses the hierarchical clustering technique of data mining on protein sequence datasets to identify genes that are consistent, easy to implement, finding and grouping the number of clusters pattern recognition. However, the valuable data is sometimes not useful but the knowledge hidden in that valuable data is meaningful.

Result: The distance is used to determine how closely two organisms are related, whereas the dendrogram shows a graphical representation of the distance calculated between the clusters.

Conclusion: The hierarchical clustering will help the biologist to judge which genes were clustered rightfully by viewing the tree structure, e the dendrogram. Therefore, the main aim is to unfold the knowledge in the vast field of bioinformatics by using information technologies as the key.

Key Words: Clustering, Phylogenetic tree, Sequence, Hierarchical clustering, Pattern, Dendrogram

INTRODUCTION

Bioinformatics and data mining provide exciting research challenges for computational science. Data Mining is the process of knowledge discovery of various patterns from a sea of data whereas Bioinformatics involves the storage, analysis, and construction of information from biological data in the form of sequences, pathways, and gene expression. New Data Mining techniques play a very important role in the large growing biological databases. Many times Data Mining is also known as Knowledge Discovery, which means searching a large volume of data to discover patterns and new trends that go beyond simple analysis.¹

The Clustering technique is an unsupervised method in data mining.² It helps to improve the accuracy of the data. A phylogenetic tree can be build-using Hierarchical clustering for known corresponding protein sequences or DNA sequences. A Phylogenetic tree is useful for biologists in solving both scientific and practical problems. The problems would be stated as an evolution of complex features of the species, prediction about fossils, the evolution of diversity, etc.

Hierarchical Clustering techniques are useful to represent relationships between protein families. There are two types of hierarchical clustering methods Agglomerative (bottom-up) and Divisive clustering (top-down). Using agglomerative clustering the very first single cluster is chosen and further in recursive fashion two or more relative clusters keep on merging. Divisive clustering starts with all data points merged in one cluster and further splits into most relative clusters recursively. Cluster analysis is a method for finding similar data objects present in the data.³ This method divides the section into sets of clusters in such a way that two objects a picked, which resembles similar from the same cluster, whereas clusters, which belong to different clusters, are not similar.

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METHODOLOGY

The process of data mining is part of many steps, which require repetition and refinement to achieve accuracy. Finding out new methods of Data Mining will act as a key factor in the vast mounting of biological data. It is a key point to state that the whole process of Data Mining covers a gathering of many techniques [Fig. 1], which involves finding the pattern from a large set of sequences, applying a hierarchical clustering algorithm and discovering knowledge through the result, which is a phylogenetic tree.

The phylogenetic tree is compared against the problem if it is solved with a single phylogenetic tree then there is no recursive procedure to be followed else the data is changed and the hierarchical clustering steps are executed to get the right phylogenetic tree.4 The data is collected from a distance matrix [Fig. 2] on which the hierarchical clustering method is applied.5

RESULTS

The closest clusters were displayed through the hierarchical clustering algorithm as follows:

The closest Clusters are:
- C, DE:: 45
- A, B:: 20
- AB, CDE:: 72
- D, E:: 30

The distance is used to determine how closely two organisms are related. The line that connects all other lines is the representation of the common ancestor. Fig.3 describes the dendrogram generated from the distance matrix.

DISCUSSION

A dendrogram is a cluster tree, which is used to represent sequence data providing links to various groups. Each group is connected to a similar group and ends up forming a tree-like structure.

The above figure shows the results of a hierarchical cluster analysis of the sequence. The horizontal axis shows the distance between each cluster, in which four clusters were identified with an optimal number of 2–4 nodes in each cluster. Cluster 1 is a combination of A and B, which are the closest cluster with a distance of 20. The next closest found is D and E with a distance of 30. The common factors for C and (D and E) are grouped in Cluster 3, whereas Cluster 4 is a combination of (A and B) and (C and (D and E)). This information, which is divided amongst the nodes, has a huge impact on the selection of the closest cluster.
CONCLUSION

The advantage of hierarchical clustering helps the biologist to judge which genes were clustered rightfully by viewing the tree structure, the dendrogram. It also enables us to predict how the genes will change in the future. Phylogenetics provides a more accurate description of patterns based on sequence data.

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