Exploring Highly Structure Similar Protein Sequence Motifs using SVD with Soft Granular Computing Models

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Abstract- Vital areas in Bioinformatics research is one of the Protein sequence analysis. Protein sequence motifs are determining the structure, function, and activities of the particular protein. The main objective of this paper is to obtain protein sequence motifs which are universally conserved across protein family boundaries. In this research, the input dataset is extremely large. Hence, an efficient technique is demanded. A Rough Granular computing model is created to efficiently extracting protein motif data that transcends protein families. Before apply this model, the very first step of this research is trying to reduce segments. The literature suggests that the Singular Value Decomposition (SVD) computing technique is more suited for reducing segments. After that the reduced segments are followed by applying Rough Granular computing model. The effectiveness of final results effectiveness is tested by several measures. The experimental results suggest that the SVD with Rough Granular computing model generates more number of highly structured motif patterns.


I. INTRODUCTION

A thick relationship between protein sequence and its structure plays a vital role in current bioinformatics research. The biological term ‘sequence motif’ denotes a relatively, functionally or structurally conserved sequence patterns that occur repeatedly in a group of related proteins [12]. These motif patterns may be able to predict the structural or functional area of other proteins, such as enzyme-binding sites, DNA or RNA binding sites, prosthetic attachment sites, or regions involved in binding other small molecules.

PROSITE [1], PRINTS [2], BLOCKS [3], SBASE [4], and PFAM [5] are five popular databases for sequence motifs. There are some commonly used softwares for protein sequence motif discover including MEME [6], Gibbs Sampling [7, 8], Block Maker [9] and some of the latest algorithms include MITRA [10], and Gemoda [11]. These applications, endure a common issue of limiting the size of input dataset. Several protein sequences are required to be input by the user while using these tools.

In this research, protein sequences are converted into segments using sliding window concepts and patterns are extracted from the selected segments. The total sliding sequence segments are trim by Singular Value Decomposition (SVD) [13].These sliding sequence segments are separated into different groups with granular computing models that utilized Fuzzy C-Means, Adaptive Fuzzy C-Means and Rough K-Means clustering algorithms to divide the set of segments into several smaller subsets and then apply K-Means and Rough K-Means algorithm to each subset to discover relevant information. Finally, we merge the information generated by all granules and obtain the final sequence motif information. Various evaluation methods are applied in this study such as structural similarity, Dunn Index (DI) measure, Davis-Bouldin Index (DBI) measure, and HSSP-BLOSUM62 evaluation method. The hybridization of the SVD with Rough Granular computing model generates more number of highly structured motif patterns.

The rest of the paper is organized as follows. Section 2 presents related work in this area of research. Section 3 introduces SVD-Entropy based segment selection process. In section 4, the description of granular computing techniques and clustering algorithms are explained. Experimental setup is explained in section 5. In section 6, experimental results are explained. Section 7 concludes the paper with directions for further enhancement.
II. RELATED WORKS

K-Means clustering algorithm with random initial centroids is utilized by Han et al. [14] to find recurring protein sequence motifs across the boundaries of a protein family. To overcome the inherent problem of K-Means clustering algorithm, Wei et al. proposed an improved K-Means clustering algorithm to obtain initial centroid locations more wisely [15] and the results published by Wei et al. have been improved in their experiment.

Bernard Chen et al. proposed a granular computing model work called FIK model [16, 17] for overcome the high computational cost, which utilizes a Fuzzy C-Means clustering algorithm to divide the whole data space into several smaller subsets and then applies a standard improved K-Means algorithm to each subset to discover relevant information. In FGK model [16, 17] Bernard Chen et al. develop a new greedy K-Means algorithm to further improve secondary structural similarity sequence motifs. In the Greedy K-Means, the best centroids are selected after five runs of K-Means and then K-Means algorithm is executed by considering those centroids. It consumes more time and complexity is also high.

Motif detection from a huge amount of sequences is a challenging task and not all the segments generated are so important. Therefore, Bernard Chen [18] has proposed Super Granular SVM Feature Elimination. In this approach the original dataset is first partitioned using Fuzzy C-Means clustering and then for each partition Greedy K-Means clustering algorithm is been implemented. Then ranking SVM based segment selection is done on each cluster to collect survived sequence segments. The survived segments are then clustered once again using Greedy K-Means to generate motif information. The Super Granular SVM segment selection technique requires more computational time for segment selection process. Here, the computational time includes time taken for Fuzzy Clustering plus time taken for Greedy K-Means clustering before segment selection.

In this paper, SVD Entropy segment selection Technique is applied before clustering, which helps us to reduce computational time. Here, all sequence segments generated by sliding window technique may not yield highly structural similar clusters. Therefore, removing such noisy segments using entropy segment selection [19] helps us to produce clusters with good structural similarity.

III. SEGMENT SELECTION TECHNIQUE

A. SVD Entropy Based Segment Selection Technique

SVD based entropy addresses the problem of selecting the significant segments in the area of protein sequence motif identification [13, 32]. The city block metric is used for calculating the difference between a sequence segment and the centroid of a given sequence cluster. The formula for calculating entropy each sequence segment is given here under.

\[ V_j = \frac{S_j^2}{\sum w S^2_w} \]  

(1)

where \( S_j \) denotes singular values of the segment, \( S^2_w \) denotes eigen values of the segment, \( w \) denotes the window size.

The resulting SVD- Entropy is as follows

\[ E = -\frac{1}{\log(w)} \sum_{j=1}^{w} V_j \log (V_j) \]  

(2)

1) \( E < m + n \), features with high contribution.
2) \( m + n > E > m - n \), features with average contribution.
3) \( E < m - n \), features with negative contribution.

The segments obtained in the first group are said to relevant to our problem. The segments in the second group are said to be neutral and the third group segments will reduce total SVD entropy. In this work, we have selected only those segments which fall under the first category.
Fig. 1 shows SVD Entropy Selection algorithm applied in Fuzzy Granular Model (FGM), Adaptive Fuzzy Granular Model (AFGM) and Rough Granular Model (RGM). The motif information obtained after the segment selection process is said to be more meaningful as well as DBI value considerably decreased after the feature selection process.

IV. GRANULAR COMPUTING TECHNIQUES

A. Fuzzy Granular Model with SVD Entropy

This computation work consists of two phases. Phase one selects significant segments using SVD-Entropy method. Phase two adopts FGM computing technique. The SVD-Entropy has been combined with FGM to identify hidden motif patterns that are available in different protein families. As the dataset is very large, hence the work focuses on segment selection technique to be applied before granular computing which helps us to reduce computational cost. Traditional K-Means [20] and Rough K-Means Clustering algorithms are performed on each information granule generated by FCM. At the final stage, we combine information generated by all granules and obtain final sequence motif information. The Figures 2 and 3 show the structure of FGM using K-Means and FGM using Rough K-Means respectively.

Algorithm : SVD Entropy Based Segment Selection

Input : Sequence segments of N numbers.

Output : Significant protein sequence segments.

Procedure:

Step1: Computation of SVD - Entropy

For i = 1 to N

Calculate singular value decomposition for each sequence segment using (1)

Let K is the number of non zero SVD entries along with window size

For j varies from 1 to K

Apply SVD Entropy using (2)

End For

End For

Step2: Selection of Sequence segments

If (entropy of each sequence segment < threshold value) then

Select those sequence segments for clustering process

Else

Eliminate the segments from clustering process

End If
B. **Fuzzy C-Means**

Fuzzy C-Means (FCM) is a clustering algorithm which allows one segment of data is belongs to one or more clusters. This algorithm is to minimize the following objective function [16]:

\[ J_m = \sum_{i=1}^{N} \sum_{j=1}^{C} u_{ij}^m \| x_i - c_j \|^2, \quad 1 \leq m \leq \infty \]  

(3)

where \( m \), the fuzzification factor, is any real number greater than 1, \( u_{ij} \) is the degree of membership of \( x_i \) in the cluster \( j \), \( x \) is the \( i \)th of \( d \)-dimensional measured data, \( c \) is the \( d \) dimension center of the cluster, and \( \| \cdot \| \) is any norm expressing the similarity between any measured data and the center. Fuzzy partitioning is carried out through an iterative optimization of the objective function shown above, with the update of membership \( u_{ij} \) and the cluster centers \( c_j \) by:

\[ c_j = \frac{\sum_{i=1}^{N} u_{ij}^m \cdot x_i}{\sum_{i=1}^{N} u_{ij}^m} \]  

(4)

where

\[ u_{ij} = \frac{1}{\sum_{k=1}^{C} \left( \frac{\| x_i - c_j \|^2}{\| x_i - c_k \|^2} \right)^{\frac{m-1}{2}}} \]  

(5)

This iteration will stop when \( \max_{0 \leq k \leq \infty} \| U^{(k+1)} - U^{(k)} \| < \delta \) where \( \delta \) is a termination criterion between 0 and 1, whereas \( k \) is the iteration step. This procedure converges to a local minimum or a saddle point of \( J_m \).
The Fuzzy C-Means Clustering algorithm is described as following:

i. Initialize membership function matrix \( U = [u_{ij}] \), and \( U(0) \).

ii. at k step: Calculate the centroid point by the equation (4)

iii. Update \( U^{(k)} \) and \( U^{(k+1)} \) by using equation (5).

iv. if \( |U^{(k+1)} - U^{(k)}| < \varepsilon \) then stop; otherwise return to step 2.

C. Adaptive Fuzzy Granular Model with SVD Entropy

The SVD-Entropy has been combined with AFGM to identify more hidden motif patterns. Traditional K-Means and Rough K-Means Clustering algorithms are performed on each information granule generated by AFCM. At the final stage, we combine information generated by all granules and obtain final sequence motif information. The Figures 4 and 5 show the structure of AFGM with SVD Entropy [21, 22].

![Figure 4. Sketch of AFGM using K-Means Computing Model with SVD Entropy](image)

![Figure 5. Sketch of AFGM using K-Means Computing Model with SVD Entropy](image)

Many of the behavioural problems with standard Fuzzy C-Means algorithm are eliminated when we relax probabilistic constraint imposed on membership function. Further Krishnapuram R and Keller JM [21, 33] have modified the approach for calculating membership values. Equation (6) shows membership calculation.

\[
\sum_{j=1}^{k} \sum_{i=1}^{n} \mu_{ij}(x_i) = n
\]

Here,

- \( \mu_{ij}(x_i) \) is the membership of \( x_i \) in \( j^{th} \) cluster
- \( k \) is the specified number of clusters
- \( n \) is the number of data points

In Adaptive Fuzzy C-Means (AFCM), the total membership quantifiers for all sample points are equal to \( n \). This flexible approach leads to clustering optimization problem, provides a way to improve cluster robustness. Here the algorithm is adaptive; that is membership is based on sample size rather than fixed to upper limit as one in Fuzzy C-Means clustering. The membership values in this method are calculated using Equation (7)
The Adaptive fuzzy clustering algorithm is efficient in handling data with outlier points. It gives very low membership values for outliers since the sum of distances of points in all the clusters involves in membership calculation.

D. Rough Granular Model with SVD Entropy

A set of information granules is built using the Rough Granular Model (RGM) with SVD entropy and then applying K-Means and Rough K-Means Clustering algorithms to obtain the final information. The RGM process is given below in Fig. 6 and Fig. 7 [21, 22].

\begin{equation}
\mu_j(x_i) = \frac{n \left( \frac{1}{d_j} \right)^{m-1}}{\sum_{k=1}^{P} \sum_{z=1}^{D} \left( \frac{1}{d_{kz}} \right)^{m-1}}
\end{equation}

Figure 6. Sketch of RGM using K-Means Computing Model with SVD Entropy

Figure 7. Sketch of RGM using K-Means Computing Model with SVD Entropy

E. Rough Clustering

In rough clustering each cluster has two approximations, a lower and an upper approximation. The lower approximation is a subset of the upper approximation. The members of the lower approximation belong certainly to the cluster; therefore they cannot belong to any other cluster. The data objects in an upper approximation may belong to the cluster. Since their membership is uncertain they must be a member of an upper approximation of at least another cluster.

F. Properties for the Rough Clustering Algorithm

Property 1: a data object can be a member of one lower approximation at most.

Property 2: a data object that is a member of the lower approximation of a cluster is also member of the upper approximation of the same cluster.

Property 3: a data object that does not belong to any lower approximation is member of at least two upper approximations [23].

The Rough K-Means algorithm provides a rough set theoretic flavour to the conventional K-Means algorithm to deal with uncertainty involved in cluster analysis. The Rough K-Means algorithm [24, 25] described as follows:

1. Select initial clusters of n objects into K clusters.
2. Assign each object to the Lower bound (L(x)) or upper bound (U(x)) of cluster/ clusters respectively as: For each object v, let \( d(v, x_i) \) be the distance between itself and the centroid of cluster \( x_i \). The difference between \( d(v, x_i) / d(v, x_j) \), \( 1 \leq i, j \leq k \) is used to determine the membership of v as follows:
   - If \( d(v, x_i) / d(v, x_j) \leq \) thershold, then \( v \in U(x_i) \) & \( v \in U(x_j) \). Furthermore, v will not be a part of any
3. For each cluster $x_i$, re-compute center according to the following equations the weighted combination of the data points in its lower_bound and upper_bound.

$$x_i = \begin{cases} 
  \frac{\sum_{j=1}^{l_i} y_j}{\text{lower}_i} + w_{\text{upper}} \cdot \frac{\sum_{j=l_i+1}^{u_i} y_j}{\text{upper}_i - \text{lower}_i} & \text{if } |\text{lower}_i - \text{upper}_i| \neq 0 \\
  \frac{\sum_{j=l_i}^{u_i} y_j}{\text{lower}_i} & \text{otherwise} 
\end{cases}$$

where $1 \leq j \leq k$. The parameters $w_{\text{lower}}$ and $w_{\text{upper}}$ correspond to the relative importance of lower and upper bounds. If convergence criterion is met, i.e. cluster centers are same to those in previous iteration, then stop; else go to step 2.

**G. K-Means Clustering Algorithm**

Among all clustering algorithms, K-Means clustering algorithm has the advantages of easy interpretation and implementation, high scalability, and low computation complexity. The K-Means clustering take the user input parameter $K$, and partitions a set of $n$ objects into $K$ clusters then iteratively updates the centers until no reassignment of patterns to new cluster centers occurs. In every step, each sample is allocated to its closest cluster center and cluster centers are reevaluated based on current cluster memberships [26].

**V. EXPERIMENTAL SETUP**

**A. Data Set**

The dataset obtained from Protein Sequence Culling Server (PISCES) includes 4946 protein sequences [27]. In this work, we have considered 3000 protein sequences to extract sequence motifs that transcend in protein sequences. The threshold for percentage identity cut-off is set as less than or equal to 25%, resolution cut-off is 0.0 to 2.2, R-factor cut-off is 1.0 and length of each sequence varies from 40 to 10,000. Homology Derived Secondary Structure of Proteins (HSSP) frequency profiles is used to represent each segment [4, 5]. The sliding windows with ten successive residues are generated from protein sequences. Each window represents one sequence segment of ten continuous positions. Around 6, 60,364 sequence segments are generated by sliding window method, from 3000 protein sequences. Each sequence segment is represented by 10 X 20 matrix, where ten rows represent each position of sliding window and 20 columns represent 20 amino acids. Fig. 8 shows sliding window technique. In this sliding window technique we can generate $n$ number of sequence segments (10 X 20 matrices).

Dictionary of Secondary Structure of Proteins (DSSP) assigns secondary structure to eight different classes [28]. These eight structural classes can be reduced to three using reduction method as follows: H, G and I to H (Helices); B and E to E (Sheets); all others to C (Coils) [29].

![Sliding Window techniques with a window size of 10 applied on 3CA8 HSSP file](image)
B. Structural Similarity Measures

A cluster’s average structure is calculated using the following formula:

\[
\sum_{i=1}^{WS} \max\left(\mathcal{P}_{(i,h)}, \mathcal{P}_{(i,b)}, \mathcal{P}_{(i,c)}\right) / WS
\]

where \(WS\) is the window size and \(\mathcal{P}_{(i,h)}\) shows the frequency of occurrence of helix among the segments for the cluster in position \(i\). \(\mathcal{P}_{(i,b)}\) and \(\mathcal{P}_{(i,c)}\) are defined in a similar way. If the structural homology for a cluster exceeds 70\%, the cluster can be considered structurally identical [16]. If the structural homology for the cluster exceeds 60\% and is below 70\%, the cluster can be considered weakly structurally homologous.

C. Distance Measure

The city block metric is more suitable for this field of study since it will consider every position of the frequency profile equally. The city block metric is used for calculating the difference between a sequence segment and the centroid of a given sequence cluster. Han and Baker also chose the city block metric because of complications associated with the use of Euclidean metric for clustering algorithms [14]. The following formula is used to calculate the distance between two sequence segments:

\[
\text{Distance} = \sum_{i=1}^{WS} \sum_{j=1}^{N} |F_x(i, j) - F_c(i, j)|
\]

where \(WS\) is the window size and \(N\) is 20 which represent 20 different amino acids. \(F_x\) \((i, j)\) is the value of the matrix at row \(i\) and column \(j\) used to represent the sequence segment. \(F_c\) \((i, j)\) is the value of the matrix at row \(i\) and column \(j\) used to represent the centroid of a give sequence cluster.

D. Dunn Index Measure

The Dunn Index (DI) also favours clustering with low intra-cluster and high inter-cluster distances, although the compactness of the clusters is assessed in a different way [42]. This index is originally proposed to use at the identification of "compact and well separated clusters". So the result of the clustering has to be small. Thus, based on the Dunn’s index definition, we may conclude that large values of the index indicate the presence of compact and well separated clusters.

The implications of the Dunn index are:

- The considerable amount of time required for its computation.
- The sensitive to the presence of noise in datasets.

Since these are likely to increase the values of \(\text{diam}(c)\).
E. Davis-Bouldin Index (DBI) Measure

The DBI measure [17] is a function of the inter-cluster and intra-cluster distance. A good cluster result should reflect a relatively large inter-cluster distance and a relatively small intra-cluster distance. The DBI measure combines both distance information into one function, which is defined as follows:

$$DBI = \frac{1}{k} \sum_{p=1}^{k} \max_{p \neq q} \left( \frac{d_{\text{intra}}(C_p) + d_{\text{intra}}(C_q)}{d_{\text{inter}}(C_p, C_q)} \right),$$

where

$$d_{\text{intra}}(C_p) = \frac{1}{n_p} \sum_{i=1}^{n_p} \| g_i - g_{pc} \|$$

and

$$d_{\text{inter}}(C_p, C_q) = \| g_{pc} - g_{qc} \|$$

K is the total number of clusters, $d_{\text{intra}}$ and $d_{\text{inter}}$ denote the intra- and inter-cluster distances respectively. $n_p$ is the number of members in the cluster $C_p$. The intra-cluster distance defined as the average of all pairwise distances between the members in cluster $P$ and cluster $P$’s centroid $g_{pc}$. The inter-cluster distance of two clusters is computed by the distance between two clusters’ centroids. The lower DBI value indicates the high quality of the cluster result.

F. HSSP-BLOSUM62 Measure

BLOSUM62 [30] (Fig. 9.) is a scoring matrix based on known alignments of diverse Sequences.

![Figure 9. BLOSUM62 Matrix](image)

By using this matrix, we may access the consistency of the amino acids appearing in the same position of the motif information generated by our method. Because different amino acids appearing in the same position should be close to each other, the corresponding value in the BLOSUM62 matrix will give a positive value. Hence, the measure is defined as the following [31]:

$$\text{If } k = 0: \quad \text{HSSP-BLOSUM62 measure} = 0$$

$$\text{Else if } k > 1: \quad \text{HSSP-BLOSUM62 measure} = \frac{1}{k} \text{BLOSUM62}_{ij}$$

$$\text{If } 0 \% \leq \text{HSSP}_i < 10\%: \quad \text{HSSP-BLOSUM62 measure} = \frac{1}{k} \sum_{i=1}^{k} \text{HSSP}_i \text{BLOSUM62}_{ij}$$

$$\text{Else:} \quad \text{HSSP-BLOSUM62 measure} = \frac{1}{\sum_{i=1}^{k} \sum_{j=1}^{k} \text{HSSP}_i \text{HSSP}_j}$$
G. Parameter Setup

In this work, SVD - Entropy based segment selection is applied and selected around 85% of sequence segments from original data set. Number of clusters has been set to 900. For FCM granular with SVD – Entropy technique, the fuzzification factor is set to 1.15 and number of clusters is equal to ten. This setting produced better results in our specific dataset. In order to separate information granules from FGM results, the membership threshold is set to 18% [32]. The function that decides how many numbers of clusters should be in each information granule is given below:

\[ C_k = \frac{n_k}{\sum_{i=1}^{m} n_i} \times m \]

where \( C_k \) denotes the number of clusters assigned to information granule \( k \), \( n_k \) is the number of members belonging to information granule \( k \), \( m \) is the number of clusters in Fuzzy C-Means. In this technique we are able to indentify 900 clusters.

For Adaptive Fuzzy C-Means, the fuzzification factor is considered as 1.15 and membership threshold is set to 13% [32]. The number of clusters in each granule is decided by the function given below:

\[ C_k = \frac{n_k}{\sum_{i=1}^{m} n_i} \times m \]

where \( C_k \) denotes the number of clusters assigned to information granule \( k \), \( n_k \) is the number of members belonging to information granule \( k \), \( m \) is the number of clusters in Adaptive Fuzzy C-Means. In this technique we are able to indentify 901 clusters.

For Rough K-Means, the epsilon value is considered as 1.001 and the number of clusters in each granule is been decided by the function given below:

\[ C_k = \frac{n_k}{\sum_{i=1}^{m} n_i} \times m \]

where \( C_k \) denotes the number of clusters assigned to information granule \( k \), \( n_k \) is the number of members belonging to information granule \( k \), \( m \) is the number of clusters in Rough K-Means. In this technique we are able to indentify 900 clusters.

VI. EXPERIMENTAL RESULTS

TABLE I. SUMMARY OF THE RESULTS OBTAINED BY THE FCM

<table>
<thead>
<tr>
<th>Granules</th>
<th>Number of Members</th>
<th>Number of Clusters</th>
<th>Data Size (in MB)</th>
</tr>
</thead>
<tbody>
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<tr>
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<td>Original Data Set</td>
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<td>19.20</td>
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</table>

The summary of the results obtained from FCM granular method is shown in Table I. Although the total segment increased from 660364 to 805869, we achieved the goal of reduced data size is to deal with one information granule at a time [22].

The summary of the results obtained from FCM granular method with SVD entropy is shown in Table II. The total number of segments are slight increased, but we achieved the goal of reduced data size is to deal with one information granule at a time.
The summary of the results obtained from AFCM granular method is shown in Table III. Although the total number of members increased from 562745 to 721390, we only deal with one information granule at a time. Therefore, we achieved the goal of reduced space-complexity [22].

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<tr>
<td>Granule 6</td>
<td>125769</td>
<td>170</td>
<td>6.34</td>
</tr>
<tr>
<td>Granule 7</td>
<td>2409</td>
<td>3</td>
<td>0.23</td>
</tr>
<tr>
<td>Granule 8</td>
<td>65409</td>
<td>89</td>
<td>4.14</td>
</tr>
<tr>
<td>Granule 9</td>
<td>2824</td>
<td>4</td>
<td>0.22</td>
</tr>
<tr>
<td>Granule 10</td>
<td>129761</td>
<td>176</td>
<td>6.47</td>
</tr>
<tr>
<td>Total</td>
<td>664609</td>
<td>901</td>
<td>34.93</td>
</tr>
<tr>
<td>Original Data Set</td>
<td>660364</td>
<td>900</td>
<td>19.20</td>
</tr>
</tbody>
</table>
The summary of the results obtained from AFCM granular method with SVD entropy is shown in Table IV. Although the total number of members increased at 686552, we only deal with one information granule at a time. Hence, we achieved the goal of reduced space-complexity with more number of highly structure motif patterns.

The summary of the results obtained from RKM granular method is shown in Table V. The total number of members is exactly same as original data set but identifies more number of hidden highly structure motif patterns.
TABLE VI. SUMMARY OF THE RESULTS OBTAINED BY THE SVD-RKM

<table>
<thead>
<tr>
<th>Granules</th>
<th>Number of Members</th>
<th>Number of Clusters</th>
<th>Data Size (in MB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Granule 1</td>
<td>80341</td>
<td>128</td>
<td>5.55</td>
</tr>
<tr>
<td>Granule 2</td>
<td>21425</td>
<td>34</td>
<td>1.48</td>
</tr>
<tr>
<td>Granule 3</td>
<td>77671</td>
<td>124</td>
<td>5.28</td>
</tr>
<tr>
<td>Granule 4</td>
<td>54727</td>
<td>87</td>
<td>3.89</td>
</tr>
<tr>
<td>Granule 5</td>
<td>43451</td>
<td>69</td>
<td>2.60</td>
</tr>
<tr>
<td>Granule 6</td>
<td>53482</td>
<td>85</td>
<td>3.77</td>
</tr>
<tr>
<td>Granule 7</td>
<td>60673</td>
<td>97</td>
<td>4.13</td>
</tr>
<tr>
<td>Granule 8</td>
<td>45012</td>
<td>72</td>
<td>2.96</td>
</tr>
<tr>
<td>Granule 9</td>
<td>66865</td>
<td>107</td>
<td>4.72</td>
</tr>
<tr>
<td>Granule 10</td>
<td>61623</td>
<td>98</td>
<td>4.17</td>
</tr>
<tr>
<td>Total</td>
<td>565270</td>
<td>901</td>
<td>38.55</td>
</tr>
<tr>
<td>Original Data Set</td>
<td>660364</td>
<td>900</td>
<td>17.70</td>
</tr>
</tbody>
</table>

The summary of the results obtained from RKM granular method with SVD entropy is shown in Table VI. The total number of members is smaller than original data set but identifies more number of hidden highly structure motif patterns.

Figure 10. BLOSUM62 Matrix

Fig. 10 has been interpreted from table VII. From the Fig. 9 we state that the number of strong and weak clusters have been increased in Granular RKM with Rough K-Means technique as well as percentage of sequence segments have also been increased considerably.
Table VII shows the comparative results obtained from different algorithms and granularization methods. From the table VII, we can infer that RGM with Rough K-Means method able to identify more number of hidden motif patterns.

![Comparison of DBI, DI and BLOSUM62 measure values](image)

**Figure 11.** Comparison of DBI, DI and BLOSUM62 measure values
Fig. 11 shows DBI, DI and HSSP-BLOSUM62 measure values obtained from different methods and different granular computing techniques.

![Figure 12. Comparison of Structural Similarity Values](image)

Fig. 12 shows percentage of structural similarity belonging to clusters obtained from different methods and different granular computing techniques. Fig. 11 has been interpreted from table VIII. From the Fig. 11, we state that the number of strong and weak clusters have been increased in RGM with SVD entropy along with Rough K-Means.

![Figure 13. Comparison of DBI, DI and BLOSUM62 measure values](image)

Fig. 13 shows DBI and HSSP-BLOSUM62 measure values obtained from different methods and different granular computing techniques. The low DBI measure and high HSSP-BLOSUM62 values indicate the improvement of the quality of clusters achieved by RGM with SVD entropy along with Rough K-Means technique.


**TABLE VIII. COMPARISON RESULTS OF DIFFERENT ALGORITHMS**

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>After SVD Segment Selection</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of Clusters &gt;70% Structure Similarity</td>
<td>101</td>
</tr>
<tr>
<td>No. of Clusters &gt;60% and &lt;70% Structure Similarity</td>
<td>190</td>
</tr>
<tr>
<td>% of Sequence Segments &gt;70%</td>
<td>11.22</td>
</tr>
<tr>
<td>% of Sequence Segments &gt;60% and &lt;70%</td>
<td>21.11</td>
</tr>
<tr>
<td>DI Measure</td>
<td>0.2206</td>
</tr>
<tr>
<td>Avg. HSSP</td>
<td>0.5361</td>
</tr>
</tbody>
</table>

A. **Sequence Motifs**

Four different motif patterns obtained from RGM with SVD entropy along with Rough K-Means process are shown in tables IX to XII. The following format is used for representation of each sequence motif table. Instead of using the traditional format, in this paper protein logo representation has been used [18].

**TABLE IX. SHEETS-COILS MOTIF**

<table>
<thead>
<tr>
<th>Number of Sequence Segments:171</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural Similarity: 73.1</td>
</tr>
</tbody>
</table>

![Protein Logo Representation]

- **HP**: 0.86, 0.65, 0.72, 0.33, 0.26, 0.24, 0.20, 0.22, 0.26, 0.64
- **Var**: 5, 6, 5, 9, 9, 8, 9, 11, 5
### TABLE X. COILS MOTIF

<table>
<thead>
<tr>
<th>Number of Sequence Segments: 34</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural Similarity: 72.65</td>
</tr>
</tbody>
</table>

### TABLE XI. HELICES MOTIF

<table>
<thead>
<tr>
<th>Number of Sequence Segments: 120</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural Similarity: 73</td>
</tr>
</tbody>
</table>

### TABLE XII. HELICES MOTIF

<table>
<thead>
<tr>
<th>Number of Sequence Segments: 125</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural Similarity: 70.96</td>
</tr>
</tbody>
</table>
• The above tables IX-XII show the number of sequence segments belonging to this motif, percentage of structural similarity. The graph demonstrates the type of amino acid frequently appearing in the given position by amino acid logo. It only shows the amino acid appearing with a frequency higher than 8%. The height of symbols within the stack indicates the relative frequency of each amino or nucleic acid at that position.

• The x-axis label indicates the representative secondary structure (S), the hydrophobicity value (Hyd.) of the position. The hydrophobicity value is calculated from the summation of the frequencies of occurrence of Leu, Pro, Met, Trp, Ala, Val, Phe, and Ile. The variability indicates the number of amino acids with the frequency greater than 5%.

VII. CONCLUSION

In this study, the granular computing models such as FGM, AFGM, RGM and combined these methods with SVD entropy have studied and implemented. The SVD with Rough Granular computing model generates more number of highly structured motif patterns in each granule. Further, the granules obtained in each of the above methods are clustered using K-Means and Rough K-Means. The highly structured clusters are used to construct the motif patterns. The main objective of generating more motif patterns has been achieved with the proposed SVD with rough granular approach and Rough K-Means clustering. It is believed that this SVD entropy with granular strategy make innovative ideas in bioinformatics research.

ACKNOWLEDGMENT

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REFERENCES


