A Sequence-Motif Based Approach to Protein Function Prediction via Deep-CNN Architecture

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Abstract: The challenge of determining protein functions, inferred from the study of protein sub-sequences, is a complex problem. Also, a little literature is evident in this regard, while a broad coverage of the literature shows a bias in the existing approaches for the full-length protein sequences. In this paper, a CNN-based architecture is introduced to detect motif information from the sub-sequence and predict its function. Later, functional inference for sub-sequences is used to facilitate the functional annotation of the full-length protein sequence. The results for the proposed approach demonstrate a great future ahead for further exploration of sub-sequence based protein studies. Comparisons with the ProtVecGen-Plus – a (multi-segment + LSTM) approach – demonstrate, an improvement of +1.24% and +4.66% for the biological process (BP) and molecular function (MF) ontologies, respectively. Next, the proposed method outperformed the hybrid ProtVecGen-Plus + MLDA by a margin of +3.45% for the MF dataset, while ranked second for the BP dataset. Overall, the proposed method produced better results for significantly large protein sequences (having sequence length > 500 amino acids).

1 INTRODUCTION

The study of the role of proteins in (i) the disease Pathobiology, (ii) the examination of meta-genomes, and (iii) the discovery of therapeutic targets, are important tasks that require deep knowledge about the functions of proteins. In this regard, the functional knowledge acquisition about proteins is well supported by the computational approaches that are fast and economical, though, still needing a good amount of effort to compete with the evolving dynamics of proteins – only less than 1% of proteins have reviewed annotations.¹ The recent trend to infer protein function(s) show a biasness of the existing works for protein sequences (Jiang et al., 2016), (Kumari et al., 2019), (Radivojac et al., 2013), (Fa et al., 2018), (Kulmanov and Hoehndorf, 2020), (Makrodimitris et al., 2019), (Ranjan et al., 2019), (Ranjan et al., 2021) – mostly due to their large and cheap availability, though works based on protein structures (Yang et al., 2015), (Gligorijević et al., 2021), protein interaction network (Kulmanov et al., 2018), and others (You et al., 2018) are also available.

Protein sequences encode vital patterns, which are formed due to interactions among amino acids that in turn fold into proteins’ sub-structures, for example, binding sites, to perform the function. This justifies the necessity for the sub-sequence based approaches, while the existing approaches are primarily focused on full-length protein sequences (Cao et al., 2017), (Kulmanov et al., 2018), (Kulmanov and Hoehndorf, 2020) which makes the function prediction a little less effective. There exist only a few notable works (Ranjan et al., 2019), (Ranjan et al., 2021) that have demonstrated the utility of a sub-sequence-based methodology. In (Ranjan et al., 2019), the proposed solution is a (multi-segmentation + LSTM) based framework. The other work (Ranjan et al., 2021) is an ensemble (multi-segmentation + tf-idf + MLDA) method. Both works involve utilizing predicted function(s) for protein sub-sequences to infer the function(s) of the full-length protein sequence.

Convolutional neural networks (CNNs) have recently gained popularity as a strong alternative to recurrent neural networks (RNNs), automating feature representations for biological sequences, and for a variety of tasks such as function prediction (Kulmanov...
et al., 2018), (Kulmanov and Hoehndorf, 2020), drug- target prediction (Öztürk et al., 2018), (Öztürk et al., 2019), etc. In, (Kulmanov et al., 2018) and (Kulmanov and Hoehndorf, 2020), they applied CNNs for the complete protein sequences to infer protein function(s). This paper proposes a framework that uses a deep CNN-based architecture to first infer the function(s) of protein sub-sequences and then uses the inferred function(s) for protein sub-sequences to determine the function(s) of the full-length protein sequence. The proposed CNN-based architecture extracts motif information from the sub-sequence, and uses it to predict the GO-term(s) for the protein sub-sequence.

The evaluations of the proposed framework conducted for two independent datasets – biological process (BP) and molecular function (MF) sub-ontologies – demonstrated a significant effort of the proposed framework. The overall improvements with respect to the similar multi-segment based ProtVecGen-Plus (Ranjan et al., 2019), based on RNNs, i.e., LSTM network, demonstrated improvements of: +1.24% for the BP dataset and +4.66% for the MF dataset. Further, when compared to the hybrid, ProtVecGen-Plus + MLDA (Ranjan et al., 2019) method, the proposed work produces improvement of +3.45% for the MF dataset, while ranked second for the BP dataset. The proposed method showed better results for handling the longer protein sequences (having sequence length > 500 amino-acids).

Following is the organization of the paper. Section 2 is an elaboration of the dataset used for the experiments and the proposed methods. Following this is a Section 3 for the results discussion. Lastly, the Section 4 is a conclusion.

2 DATASETS AND METHODS

Here, we will discuss the experimental datasets, the steps for the segmented dataset construction, and the proposed method.

2.1 Datasets

Experiments were conducted for two datasets, – corresponding to the biological process (BP) and molecular function (MF) sub-ontologies as defined by the Gene Ontology (GO) (Ashburner et al., 2000). These datasets were created by downloading reviewed protein sequences and their mapped functional annotation from the UniProtKB/SwissProt (2017) repository (Consortium, 2015). The functional annotation is a unique identifier, known as a GO-term, that indicates the distinct protein function. The biological process dataset has 58,310 protein sequences and 295 unique GO-terms. The other dataset, molecular function, has 43,218 protein sequences and 135 unique GO-terms. For each GO-terms, the least number of protein sequences is taken as 200. Only proteins with a sequence length higher than 40 and lower than 2,000 were chosen for this study.

2.2 Steps to Construct the Protein Sub-Sequence Dataset

The proposed framework to infer the protein function(s) is based on the protein sub-sequences, hence, the steps taken to construct the segmented dataset are explained next. Let the training dataset be denoted as $S = \{s_i, Y_i\}_{i=1}^n$; where, $s_i$ and $Y_i$ denote the $i^{th}$ protein sequence and the corresponding GO-term(s), respectively. As shown in Figure 1, for both the BP and MF sub-ontologies, large number of protein sequences have lengths around 200 to 300. So, the maximum length for the protein sub-sequences is set to 200, with a gap of 60 amino acids between two consecutive sub-sequences.

1. For each protein sample pair, $(s_i, Y_i);\ i \in \{1, 2, ..., n\}$, a protein sequence $s_i$ is split to generate a set of protein sub-sequences of size 200. Zero-padding is done for the short protein sub-sequences.

2. The output labels for each sub-sequences are assumed equivalent to the parent protein sequence.
2.3 Proposed Method

The paper introduces a two-step framework for inferring protein function: (i) a deep stacked CNN-based architecture is used to first infer function(s) of protein sub-sequences, and (ii) the inferred function(s) for protein sub-sequences are used to determine function(s) of the full-length protein sequence. A discussion on the input sequence representation and the proposed architecture are given next.

2.3.1 One-Hot Encoding Based Input Sequence Representation

The protein sequences are pre-processed and represented as a string of amino acids, where amino acids are represented via the one-hot encoding scheme. The dimension for the one-hot encoding scheme is taken as 21. The first dimension is used to indicate the rare amino acids (O, U, X), whereas the remaining 20 dimensions correspond to the well-known 20 amino acids.

Table 1: Hyper-parameters configurations with different CNN blocks.

<table>
<thead>
<tr>
<th>S. No.</th>
<th>CNN Block</th>
<th>Filter-Size</th>
<th>Filters Count</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CNN Block 1</td>
<td>1 x 5</td>
<td>96</td>
<td>–</td>
</tr>
<tr>
<td>2</td>
<td>CNN Block 2</td>
<td>1 x 7</td>
<td>96</td>
<td>–</td>
</tr>
<tr>
<td>3</td>
<td>CNN Block 3</td>
<td>1 x 9</td>
<td>96</td>
<td>–</td>
</tr>
<tr>
<td>4</td>
<td>CNN Block 4</td>
<td>1 x (F_i)</td>
<td>128</td>
<td>(F_i) denote filter size in range 6, 9, 12, ..., (3N + 3).</td>
</tr>
</tbody>
</table>
### 2.3.2 Stacked CNN-Based Architecture

The proposed architecture employs stacked layers of Convolutional Neural Networks (CNNs) blocks to process protein sub-sequences character-by-character (where amino acids represent a character). The most notable feature of CNNs is their ability to capture local dependencies – between amino acids – through the use of trainable filters that aid in the transformation of protein sequence into a viable representation. The complete architecture is shown in Figure 2. There are two consecutive layers of two CNN blocks (CNN Block 1 and CNN Block 2) that are further divided into a set of sub-networks before being combined down the architecture. The components of both CNN blocks and sub-networks are discussed next:

### 2.1) CNN Block: The CNN block as shown in Figure 2 has following layers:

1. **ID-CNN Layer:** The purpose of the 1-dimensional CNN layer is to learn local dependencies between the amino acids along the sequence. Here, the hyper-parameters that are customized for different CNN blocks include the number of filters and filter-size. A more detailed discussion on these hyper-parameters with different CNN blocks is given in Table 1. Let \( I_f[n] \) denote the output feature map after applying the convolution operation, and the equation can be seen as:

\[
I_f[n] = x[n] * h[n] = \sum_{k=-\infty}^{\infty} x[k] * h[n-k] \tag{1}
\]

where,
- \( h[n] \) is the kernel
- \( x[n] \) is the input feature
- \( * \) denotes the convolution operation.

2. **Batch-Normalization Layer:** This layer acts as a regularizer that controls the biasness of the model – utilizing the statistics of the mini-batch (Ioffe and Szegedy, 2015).

Let \( I_f \) denote an input feature-map corresponding to the \( f^{th} \) filter, where \( 1 \leq f \leq 128 \) (given in Table 1), then the output of batch-normalization layer for the \( f^{th} \) input feature-map, denoted as \( I'_f \), is defined as:

\[
I'_f = \gamma_f \left( \frac{I_f - m[I_f]}{\sqrt{\text{var}[I_f]}} \right) + \beta_f \tag{2}
\]

where,
- \( I_f \) is the \( f^{th} \) feature map,
- \( m[I_f] \) is the mean of the \( f^{th} \) feature-map,
- \( \text{var}[I_f] \) is the variance for the \( f^{th} \) feature-map,
- \( \gamma_f \) and \( \beta_f \) are two learning parameters which control \( m[I_f] \) and \( \text{var}[I_f] \), respectively.

3. **LeakyRelu Layer:** This layer transforms the output of the previous layer in the range as given in Equation 3 and saves the unit from being non-functional (Maas et al., 2013).

\[
f(I_f) = \begin{cases} 
\alpha I_f, & \text{if } I_f < 0 \\
I_f, & \text{otherwise} 
\end{cases} \tag{3}
\]

where,
- \( \alpha \) is the constant taken as 0.2.
- \( I_f \) is the given input to the Leaky Relu layer.

### 2.2) Components of Sub-Network: Each sub-network is composed of layers as follows:

1. A consecutive layers of two CNN blocks (represented as CNN Block3\(N\) and CNN Block4\(N\)). Here \( N \) represents the number of sub-networks. Different-sized filters with the CNN Block4\(N\) are used to extract motifs of different sizes as given in Table 1.

2. Next, the MaxPooling layer is employed to extract features that emphasize the important motifs present in protein sub-sequences. This also helps to prevent the over-fitting of the model by reducing the feature maps.

\[
I'_f = \frac{(I_f - k)}{s + 1} \tag{4}
\]

where,
- \( I'_f \) is the output feature vector.
- \( I_f \) is the input feature vector.
- \( k \) is the kernel size
- \( s \) is the stride.

3. The last layer of each sub-network is the flatten layer to reduce the output to a 1-dimensional vector.

The combined output from each sub-networks, obtained using the concatenate layer, is then passed to the dropout layer (with dropout probability = 0.4). Finally, the output layer with sigmoid activation is used as a classification layer.

Importantly, a residual connection (shown with dashed line) is added between the CNN blocks for efficient training of deep neural architecture, as shown in Figure 2. This provides a significant improvement in the network’s ability to overcome vanishing gradients. The hyper-parameters for the proposed architecture are shown in Table 2.

### 2.3.3 Final Prediction for the Full-Length Protein Sequence

The mean of inferred protein functions obtained for all the protein sub-sequences is computed to get the...
3 RESULTS AND DISCUSSION

We have trained this model using Keras built on the Tensor-flow architecture as the backend. The datasets discussed in Section 2.1 were split into training (0.75%) and testing (0.25%) datasets. For monitoring the training, we have used check-pointers, early-stopping criteria along with 10% of training data as validation data.

3.1 Evaluation Metrics

Here, let \( Y_i = \{y_{i1}, y_{i2}, \ldots \} \) denotes the actual GO-terms and \( P_i = \{p_{i1}, p_{i2}, \ldots \} \) denotes the predicted GO-terms for protein sequence \( S_i; i \in \{1, 2, \ldots, n\} \). The metrics were defined as:

1. Average Recall: Recall catches the true prediction made by the model over all the predicted true samples.

\[
Re_{avg} = \frac{\sum_{i=0}^{n} Y_i \cap P_i}{Y_i} \quad (5)
\]

2. Average Precision: Precision catches the true prediction made by the model over all the actual true samples.

\[
Pr_{avg} = \frac{\sum_{i=0}^{m} Y_i \cap P_i}{P_i} \quad (6)
\]

3. Average F1-Score: F1-Score balance both the precision & recall and return the value lowest between the recall and precision.

\[
F1_{avg} = \frac{\sum_{i=0}^{n} 2|Y_i \cap P_i|}{|Y_i| + |P_i|} \quad (7)
\]

4. Consistency factor: This is based on variance which emphasizes a method’s overall generality with regard to protein sequences of various lengths. This is defined as follows:

\[
\text{Consistency factor} = \sqrt{\frac{1}{4} \sum (f1 - f_{1ri})^2} \quad (8)
\]

where, the average \( f1 - \text{score} \) for the test samples in the sequence length range \( ri \) is \( f_{1ri} \), while \( f1 \) is the overall \( f1 - \text{score} \) for the test dataset. A low value indicated high consistency and vice versa.

3.2 Baseline Comparison Methods

A lot of work has been done in the past to predict protein function using GO and amino acid sequences. Notable works used for the fair comparison includes:

3.2.1 MLDA (Wang et al., 2016)

MLDA stands for Multi-Label Linear Discriminant Analysis. This is based on the complete protein sequence that uses \( tf-idf \) features, further reduced in dimension using the Multi-label LDA (MLDA) approach, as the input representation for the protein sequence. To reduce the features, MLDA project the input feature to some other feature space.

3.2.2 ProtVecGen-Plus (Ranjan et al., 2019)

This work of ours, is the first to present the deep-learning-based method that exploits the protein sub-sequences to infer functional annotation(s) for the full-length protein sequence. To infer protein functions, multiple LSTM-based network architectures are used, each trained with different-sized protein subsequences (i.e., 100, 120, 140).

3.2.3 ProtVecGen-Ensemble (Ranjan et al., 2021)

Another sub-sequence-based method, this time employing the \( tf-idf + \) MLDA technique. However, this also entails discarding a few protein sub-sequences that have been found to be less informative and relying on the remaining sub-sequences to infer annotation(s) for the full-length protein sequence.

3.2.4 ProtVecGen-Plus + MLDA (Ranjan et al., 2019)

This is an ensemble of standard machine learning and deep learning methods. The results of the MLDA approach are combined with the results of deep learning-based ProtVecGen-Plus (Ranjan et al., 2019). This method showed great potential for predicting protein functions.
Table 3: Biological Process Dataset: Classification report with respect to different number of sub-networks with the proposed approach on protein sub-sequences (N stands for number of sub-networks).

<table>
<thead>
<tr>
<th>S. No.</th>
<th>N</th>
<th># parameters (Millions)</th>
<th>( P_{\text{avg}} ) (%)</th>
<th>( R_{\text{avg}} ) (%)</th>
<th>( F_{1\text{avg}} ) (%)</th>
<th>( CF )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>6</td>
<td>≈ 1.79</td>
<td>92.75</td>
<td>33.53</td>
<td>34.44</td>
<td>8.101</td>
</tr>
<tr>
<td>2.</td>
<td>7</td>
<td>≈ 2.21</td>
<td>93.17</td>
<td>32.44</td>
<td>33.37</td>
<td>8.707</td>
</tr>
<tr>
<td>3.</td>
<td>8</td>
<td>≈ 2.66</td>
<td>92.30</td>
<td>33.84</td>
<td>34.73</td>
<td>8.960</td>
</tr>
<tr>
<td>4.</td>
<td>9</td>
<td>≈ 3.15</td>
<td>91.99</td>
<td>35.48</td>
<td>36.33</td>
<td>9.232</td>
</tr>
</tbody>
</table>

Table 4: Molecular Function Dataset: Classification report with respect to different number of sub-networks with the proposed approach on protein sub-sequences (N stands for number of sub-networks).

<table>
<thead>
<tr>
<th>S. No.</th>
<th>N</th>
<th># parameters (Millions)</th>
<th>( P_{\text{avg}} ) (%)</th>
<th>( R_{\text{avg}} ) (%)</th>
<th>( F_{1\text{avg}} ) (%)</th>
<th>( CF )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>6</td>
<td>≈ 1.67</td>
<td>94.34</td>
<td>45.40</td>
<td>s</td>
<td>5.722</td>
</tr>
<tr>
<td>2.</td>
<td>7</td>
<td>≈ 2.07</td>
<td>93.24</td>
<td>50.82</td>
<td>51.35</td>
<td>5.544</td>
</tr>
<tr>
<td>3.</td>
<td>8</td>
<td>≈ 2.50</td>
<td>94.35</td>
<td>49.58</td>
<td>50.35</td>
<td>5.938</td>
</tr>
<tr>
<td>4.</td>
<td>9</td>
<td>≈ 2.97</td>
<td>93.68</td>
<td>49.84</td>
<td>50.84</td>
<td>5.481</td>
</tr>
</tbody>
</table>

3.3 Study the Effect of Number of sub-Networks

In this sub-section, the effect of number of sub-networks (denoted as N), considering N as 6, 7, 8, 9, with the proposed architecture is studied for both the datasets. The experiments are conducted for two different cases:

1. Full-length Protein Sequence: The proposed architecture is trained and evaluated based on the full-length protein sequences. The truncation of protein sequences larger than 500 amino acids are done.

2. Protein Sub-sequence: This represents the complete proposed framework that is based on the protein sub-sequences.

This sort of study will allow to understand advantages with the sub-sequence based method over the methods that are based on the full-length sequence model. The results for both cases are reported in Table 3 (for the BP dataset) and Table 4 (for the MF dataset).

On increasing the number of sub-networks, with respect to various performance metrics, including \( P_{\text{avg}}, R_{\text{avg}}, \) and \( F_{1\text{avg}} \), a general increase is observed. Further, increasing the number of sub-networks also helps improve the consistency factor, a lower value of consistency factor indicate a better generalized behavior of the model towards protein sequences of different lengths. For the sub-sequence based framework, the best \( F_{1\text{avg}} \) reported are 55.89% (for the BP dataset) and 70.57% (for the MF dataset), with the nine sub-networks. These experimental observations stand true for full-length sequences as well.

Importantly, the results as shown in the Table 3 and the Table 4, clearly indicate that the proposed sub-sequence based framework is superior. The performances obtained for the full-length sequence model are notably worse when compared to the sub-sequence based framework, and this observation applies regardless of choice of the sub-networks. Moreover, the full-length sequence model tends to favor a particular size of protein sequence more as quantified with the consistency factor. With respect to the best \( F_{1\text{avg}} \), in comparison to the full-length sequence model, the sub-sequence based framework is able to produce an improvements of 19.56% for the BP dataset and 19.73% for the MF dataset.

The poor performance of the full-length sequence model can be attributed primarily to the model’s inability to efficiently retain the useful information. This is because, the useful information is masked by the presence of too much not useful information, especially for the case involving long-sized protein sequences.
Table 5: Comparison between the state-of-art approach and proposed model (CF = consistency factor).

<table>
<thead>
<tr>
<th>Dataset —&gt;</th>
<th>Biological Process</th>
<th>Molecular Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>S. No.</td>
<td>Pravg (%)</td>
<td>Reavg (%)</td>
</tr>
<tr>
<td>1. MLDA (Wang et al., 2016)</td>
<td>52.61</td>
<td>49.42</td>
</tr>
<tr>
<td>2. ProtVecGen-Plus (Ranjan et al., 2019)</td>
<td>56.65</td>
<td>56.42</td>
</tr>
<tr>
<td>3. ProtVecGen-Ensemble (Ranjan et al., 2021)</td>
<td>58.59</td>
<td>56.09</td>
</tr>
<tr>
<td>4. ProtVecGen-Plus + MLDA (Ranjan et al., 2019)</td>
<td>58.80</td>
<td><strong>58.19</strong></td>
</tr>
<tr>
<td>5. Proposed model</td>
<td><strong>59.70</strong></td>
<td>57.32</td>
</tr>
</tbody>
</table>

3.4 Overall Comparison with State-of-the-Art Approaches

In this section, the proposed model is compared with the existing state-of-the-art literature works, that include: (i) Multi-label LDA (MLDA) (Wang et al., 2016), (ii) ProtVecGen-Plus (Ranjan et al., 2019), (iii) ProtVecGen-Ensemble (Ranjan et al., 2021), and (iv) hybrid approach ProtVec-Plus + MLDA (Ranjan et al., 2019). The observed performance metrics for each of the methods are shown in Table 5 for both the BP and MF datasets.

3.4.1 [object Promise]

For the BP dataset, the proposed approach easily betters the results with the MLDA (Wang et al., 2016), ProtVecGen-Plus (Ranjan et al., 2019) and ProtVecGen-Ensemble (Ranjan et al., 2021) approaches, the respective absolute enhancement in $F_1$avg being +6.62%, +1.24% and +0.55%, as shown in Table 5. A similar trend is seen for the MF dataset as well, with the proposed approach showing an improvement of +12.66%, +4.66%, and +5.10% over the MLDA (Wang et al., 2016), ProtVecGen-Plus (Ranjan et al., 2019), ProtVecGen-Ensemble (Ranjan et al., 2021), respectively. The other metrics, $Pr_{avg}$ and $Re_{avg}$ follow this behavior as well.

In comparison to the ProtVecGen-Plus + MLDA (Ranjan et al., 2019), the proposed methods stood second for the BP dataset, while comfortably outperforming for the MF dataset. The increase in the $F_1$avg for the MF dataset is +3.45%.

3.4.2 [object Promise]

Hereby, the consistency factor gives an indication about the model’s behavior to perform for protein sequences of various lengths. For the BP dataset, the consistency factor of the proposed model is reduced by 6.49, 1.21, 0.58, and 0.81 units with respect to MLDA (Wang et al., 2016), ProtVecGen-Plus (Ranjan et al., 2019), ProtVecGen-Ensemble (Ranjan et al., 2021) and ProtVecGen-Plus + MLDA (Ranjan et al., 2019), respectively. For MF, the proposed model stood next to the ProtVecGen-Ensemble (Ranjan et al., 2021) ($consistency\ factor = 3.279$), while reducing the consistency factor by 4.35, 0.67, and 0.96 units over the MLDA, ProtVecGen-Plus, and ProtVecGen-Plus + MLDA respectively.

![Figure 3: Biological Process: Length-wise performances of protein sequences.](image)
4 CONCLUSION

In this work, a sub-sequence based method for protein function prediction is introduced. The proposed method takes benefits from information collected for multiple sequence motifs – captured using the CNN network – to determine the function for each sub-sequence. Later, the functional inference for subsequences are used to facilitate the functional annotation of full-length protein sequence. Overall, the proposed method showed great potential, especially for long protein sequences. The research focused on protein sub-sequence is still an open research area, and remarkably, can be great asset to improve the protein studies. Future work will focus on merging additional features and putting different deep learning models to the test.

REFERENCES


