Learning co-evolution information with natural language processing for protein folding problem*

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Co-evolution information is crucial for building protein sequence descriptors. Its computation is traditionally based on Multiple Sequence Alignments. Applications of MSA are limited, for instance, it fails for sequences with shallow alignment. In this work, we investigate pretrained language models as a potential alternative to MSA. Specifically, our main focus is fold classification problem. We examine several state-of-the-art methods and modify them by replacing MSA-based feature-generation parts with pre-trained LMs. We compare language models and MSA in their ability to extract sequence evolution information.

Keywords: protein fold classification; co-evolution, feature generation; transformers; BERT;

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1 Introduction

- Protein properties are determined by its shape, which is specified by its amino acid sequence
- [13]. Therefore it is important to learn how to analyze this sequence. Any analysis[...] begins by
- constructing protein sequence descriptors. Their essential part is a co-evolution information.
- This information is obtained by searching for homologues of the protein in large databases
- and further computing multiple sequence alignment (MSA) on it. This technique requires
- significant computational efforts, moreover, it does not guarantee precise results since it fully
- relies on finite databases. Besides, this method fails for sequences with shallow alignment
- Co-evolution-based methods assume that a mutation of one amino acid leads to mutations of
- others. Therefore, given an amino acid sequence it is natural to look for other similar sequences.
- That is performed with MSA [5]. This method fails when we are dealing with the sequences
- that did not evolve a lot. They have shallow alignments. There are only a few similar sequences 12
- in the database. Therefore, information extracted from these alignments is insignificant. 13
- Computing MSA is a heavy task. Alignment databases are finite. If for a given sequence 14
- there are no matches in the database, MSA-based methods fail to produce reliable results. 15 In this work, we analyse pre-trained language models (LMs) [8] as a potential alternative
- to traditional MSA approaches. It is assumed that protein sequences are not random [4], that

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there is logic in amino acids structure. Therefore, a set of all amino acids could be seen as a language with complicated inner rules. With the invention of BERT [6] it has become possible to learn the structure of any abstract language. It is natural to assume that by learning amino acids structure, BERT will be able to implicitly learn co-evolution information.

Our main aim is to study the efficiency of pre-trained LMs in application to the fold classification problem. To start with, we consider the state-of-the-art method DeepSF [9]. It solves the protein fold classification problem. By replacing its MSA-based feature-generation part with a pre-trained LM, we study how NLP approach affects learning the fold-related information. The whole framework is schematically represented in Figure 1. Further, we perform the same with other fold classification algorithms [7,14] as well as with end-to-end protein structure prediction methods [10,15].

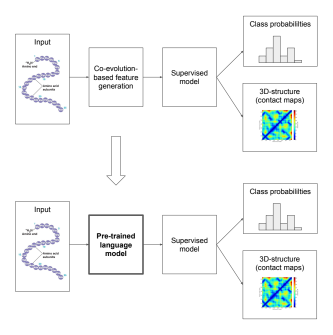


Figure 1 The main framework

2 Problem statement

30 2.1 General description

Proteins that share similarities in their structure are divided into classes – folds [12]. In this
work we investigate a fold classification problem.

We use the following notation: Y is a set of all known classes, A is a set of all proteins with known folds and $f: A \to Y$ is a mapping between proteins and their classes.

Suppose, $x \notin A$ is a protein with unknown fold. We aim to build a model capable of classifying x as an element of one of the present classes, extending f on the set $A \cup \{x\}$.

37 2.2 Feature generation

- In the fold classification problem x_i represents i-th protein description in the D-dimensional space. However, initially each protein is described by its amino acid sequence. Let Σ be a 20-letter alphabet of acid sequences.
- We are given a set of $A = \{a_i \in \Sigma^+\}_{i=1}^n$. Let $g_{\theta,D} : \Sigma^+ \to \mathbb{R}^D$ be an embedding of the space of acid sequences into D-dimensional set of descriptors. These embeddings are parameterized by θ .
- We separate A into training and validation set. Training set is used to formulate fold classification problem. Validation set is used to formulate feature generation problem.
- $A = A_{train} \cup A_{val}$
- Protein sequence descriptors are generated using $g_{\theta,D}$ as $x_i = g_{\theta,D}(a_i)$

48 2.3 Fold classification problem

- Suppose there is a given set of pairs $S = \{(x_i, Y_i)\}_{i=1}^n$ where $x_i \in \mathbb{R}^D$ denotes the sequence descriptor of i-th protein and $Y_i \in \mathbb{Y} = \{(1, 0, ..., 0), (0, 1, 0, ..., 0)....(0, ..., 0, 1)\}$ denotes the i-th protein known class, $|\mathbb{Y}| = m$
- We separate S into training and validation set. Training set is used to formulate fold classification problem.
- $S = S_{train} \cup S_{val}$

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Let $W = \{(w_1, ..., w_m) | \forall i \in \{1...m\} w_i \in \mathbb{R}^D\}$ be a space of parameters for classification models. Let g_w be a model parameterized by w. We use e_k to denote a vector with 1 on k-th position, with zeros on all other positions.

$$e_k = (0, \dots, 0, 1, 0, \dots, 0)$$

$$\uparrow$$

$$k$$

We work under the following assumption:

- $P_w(Y = e_k | x) = \frac{\exp(x^T w_k)}{\sum_j \exp(x^t w_j)}$
- Let $p_{w,x}(y)$ be discrete density of Y, $L_{Y,x}(w)$ be likelihood function of Y.
- $p_{w,x}(y) = \prod_{d=1}^{m} (P_w(Y = e_d|x))^{y_d},$
- $L_{Y,x}(w) = \prod_{i=1}^{n} p_{w,x}(Y_i)$
- Given S_{train} , classification problem is a maximization of likelihood on training set:
- $L_{train}(w) = L_{Y_{train}, x_{train}}(w) \to \max_{w \in W}$

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$_{56}$ 2.4 Feature generation problem

- For fixed θ , D and descriptors x_i let's denote $\hat{w} = \underset{w \in W}{\arg \max} L_{Y_{train}, x_{train}}(w)$.
- Feature generation problem is a maximization of likelihood function on validation set:

$$L_{Y_{val},x_{val}}(\hat{w}) \to \max_{\theta,D}$$

70 2.5 Overview

The problem of determining proteins' class is separated into two parts: first, we encode amino acid sequences as *D*-dimensional descriptors, then these descriptors are used to train a prediction model.

4 3 Computational experiment

75 **3.1** Data

We use SCOP2 [1,2] dataset. It contains information about proteins whose structure is already known. All the proteins in the database have classes assigned to them (folds). At present, there are 1517 known folds [2]. The number of proteins contained in each fold is less than 50 for most of the folds. Detailed histogram is represented in Figures 2, 3.

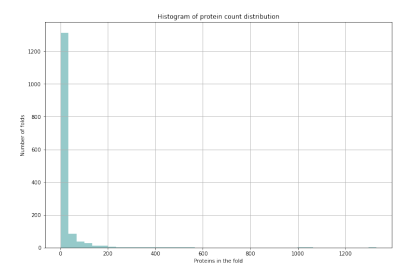
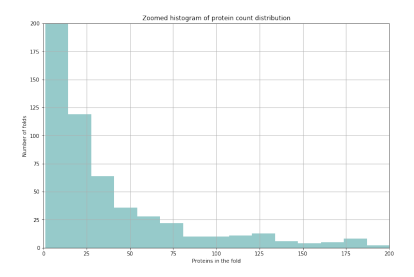
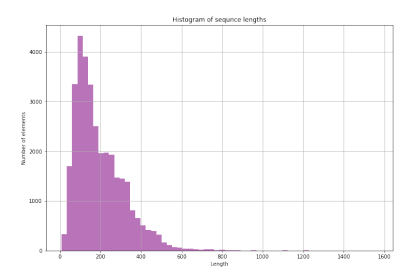


Figure 2 Protein count histogram

In SCOP2 proteins are represented by their amino acid sequences. Each sequence is stored as a string in the 20 letter alphabet and an ID. The length of the string varies mostly between 25 and 500 from protein to another. The length diagram is represented on Figure 2. Search by ID allows database users to extract information about target proteins, including their fold classes.



 ${\bf Figure~3~Protein~count~zoomed~histogram}$



 ${\bf Figure~4~Sequence~length~histogram}$

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$_{5}$ 3.2 Baseline solution

Our solution is a modification of the DeepSF [9] method. The designers of DeepSF use PSI-BLAST [3] to generate features from MSA as well as SCRATCH [11] to extract secondary structure (3 classes) and solvent accessibility (2 classes). In the core of this classification method PSI-BLAST is used during the first stage of prediction [11].

Hoe et al. then use these features as an input to the deep convolutional neural network. The depth of CNN used in DeepSF is 10. The output of the model is a vector of fold probabilities.
Fold with the maximal probability is accepted as a target protein fold.

93 3.3 Proposed solution

DeepSF feature generation is entirely based on MSA. The main point of this work is to suggest another approach to feature generation which is independent of MSA.

We apply BERT to the language of amino acids. We take the representation of acid sequences BERT creates as a new set of features. Then we train DeepSF model on these features.

It shall be noted that we do not reject co-evolution information in our research. Instead of

directly applying MSA we let BERT learn information about amino acid language. We suppose that the model learns evolution information implicitly.

101 3.4 Evaluation

We use standard cross-validation procedure to compute accuracy of our model. We then compare it to the accuracy score of DeepSF method based on MSA feature generation as well as several other state-of-the-art techniques.

105 4 Experiment result

Though the data is available for 1517 classes, during the experiments we approached the problem slowly. First, we solved a classification problem for 2 classes, then for 10 classes.

108 4.1 First experiment

In this experiment we took two balanced classes. They contain 66 and 80 elements respectively.
The distributions of their sequence length are similar. They are displayed on figure 5. We
used a model that consists of two essential layers - 30-max-pooling followed by a linear layer.
Using this architecture we achieved 93% score on validation data. The architecture and training
curves are represented on figures 6 and 7.

114 4.2 Second experiment

We decided to use exactly the same architecture for 10-fold classification problem. Our experiments show that increasing the k hyper-parameter in k-max-pooling layer improves the quality

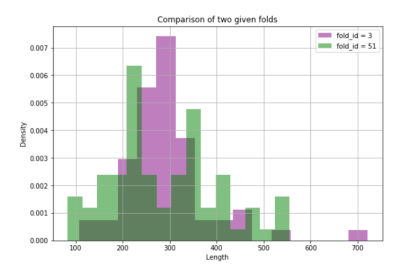


Figure 5 Sequence lengths of two folds

Layer (type:depth-idx)	Output Shape	Param #
K_max_pooling_1d: 1-1	[32, 4096, 30]	
ReLU: 1-2	[32, 4096, 30]	==
-Flatten: 1-3	[32, 122880]	
-Linear: 1-4	[32, 2]	245,762
Total params: 245,762 Trainable params: 245,762		
Trainable params: 245,762 Non-trainable params: 0 Total mult-adds (M): 7.86		
Trainable params: 245,762 Non-trainable params: 0		
Trainable params: 245,762 Non-trainable params: 0 Total mult-adds (M): 7.86		

Figure 6 Experiment 1 architecture

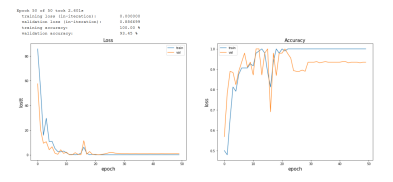


Figure 7 Experiment 1 training curve

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of the model. We chose k=60. It allowed us to reach 95% accuracy on validation and 93% accuracy on test samples. The training curve is represented on figure 8.

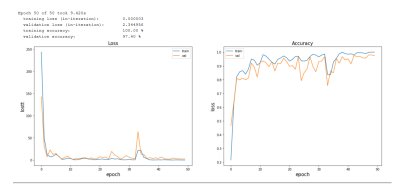


Figure 8 Experiment 2 training curve

119 5 Results

Conducted experiments show that BERT managed to capture the structure of the protein sequence language. We managed to solve 10-fold classification problem with 93% accuracy.

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