

DEEPre: sequence-based enzyme EC number prediction by deep learning

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ABSTRACT

Motivation: Annotation of enzyme function has a broad range of applications, such as metagenomics, industrial biotechnology, and diagnosis of enzyme deficiency-caused diseases. However, the time and resource required make it prohibitively expensive to experimentally determine the function of every enzyme. Therefore, computational enzyme function prediction has become increasingly important. In this paper, we develop such an approach, determining the enzyme function by predicting the Enzyme Commission number.

Results: We propose an end-to-end feature selection and classification model training approach, as well as an automatic and robust feature dimensionality uniformization method, DEEPre, in the field of enzyme function prediction. Instead of extracting manually-crafted features from enzyme sequences, our model takes the raw sequence encoding as inputs, extracting convolutional and sequential features from the raw encoding based on the classification result to directly improve the prediction performance. The thorough cross-fold validation experiments conducted on two large-scale datasets show that DEEPre improves the prediction performance over the previous state-of-the-art methods. In addition, our server outperforms five other servers in determining the main class of enzymes on a separate low-homology dataset. Two case studies demonstrate DEEPre's ability to capture the functional difference of enzyme isoforms.

Availability: The server could be accessed freely at <http://www.cbr.c.kaust.edu.sa/DEEPre>.

1 INTRODUCTION

Enzymes, an essential kind of proteins in the human body, catalyzing reactions *in vivo*, play a vital role in regulating biological processes. Annotation of enzyme function has a broad range of applications, such as metagenomics, industrial biotechnology, and diagnosis of enzyme deficiency-caused diseases. The dysfunction of certain enzymes would

cause serious metabolic diseases. For example, the deficiency of alpha-galactosidase, which hydrolyses the terminal alpha-galactosyl moieties from glycolipids and glycoproteins, would cause the Fabry disease, resulting in full body pain, kidney insufficiency, and cardiac complications (Hoffmann *et al.*, 2007). The deficiency of DNA repair enzymes, which recognize and correct the physical damage in DNA, can cause the accumulation of mutations, which may further lead to various cancers (Wood *et al.*, 2001). To investigate the causation of such diseases, an indispensable step of finding a way to cure them, it is crucial to understand the function of the related enzymes first. The most straightforward and accurate way of doing such investigation is through experimental techniques, such as enzymatic assays (Goddard and Reymond, 2004). However, conducting experiments requires significant amount of time and expert efforts, which may not cope with the rapid increase in the number of new enzymes. In this context, computational methods emerged to assist biologists in determining enzyme function and guiding the direction of setting up the validating experiments.

According to SWISS-PROT (Bairoch and Apweiler, 2000) (released on September 7, 2016), among the 539,566 manually annotated proteins, 258,733 proteins are enzymes. Such a large number of enzymes are usually classified using the Enzyme Commission (EC) system (Cornish-Bowden, 2014), the most well-known numerical enzyme classification scheme, which specifies the function of an enzyme by four digits. This classification system has a tree structure. After the root of the tree, there are two main nodes, standing for enzyme and non-enzyme proteins, respectively. The enzyme main node extends out six successor nodes, corresponding to the six main enzyme classes: (1) oxidoreductases, (2) transferases, (3) hydrolases, (4) lyases, (5) isomerases, and (6) ligases, represented by the first digit. Each main class node further extends out several subclass nodes, specifying the enzyme's subclasses, represented by the second digit. With the same logic, the third digit indicates the enzyme's sub-subclasses and the fourth digit denotes the sub-sub-subclasses. Take type II restriction enzyme, which is annotated as EC 3.1.21.4, as an example, the "3" denotes that it is a hydrolase; the "1" indicates that it acts on ester bonds; the "21" shows that it is an endodeoxyribonuclease producing 5-phosphomonoesters; and the "4" suggests that it is a Type II site-specific deoxyribonuclease. By predicting the EC numbers precisely, computational methods can annotate the function of enzymes. It should also be noted that a substantial number of enzymes

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annotated with some reactions in databases such as UniProt or Brenda do not have EC numbers associated, which is out of the scope of this study.

A number of computational methods have already been proposed to determine the enzyme function by predicting enzyme EC numbers. There have been three main research directions of this problem since (des Jardins *et al.*, 1997), who used machine learning methodologies and sequence information to investigate the problem for the first time. Firstly, because it is commonly believed that structures determine function, some researches, such as (Dobson and Doig, 2005; Roy *et al.*, 2012; Nagao *et al.*, 2014; Yang *et al.*, 2015; Zhang *et al.*, 2017), focused on predicting the enzyme function by predicting the structure of the enzyme first. After obtaining the structure, they scanned the database or the library, whose entries' EC numbers have already been determined and validated by experiments, and assigned the EC number of the template with the most similar structure to the query. However, structure prediction is still relatively immature and time-consuming. Besides, since both the structure prediction step and the EC number prediction step would cause errors, the accumulated error would have a negative effect on the final prediction result. Secondly, the common assumption that enzymes with high sequence similarity tend to have similar functionality leads to a number of studies utilizing sequence similarity (Tian *et al.*, 2004; Arakaki *et al.*, 2009; Yu *et al.*, 2009; Quester and Schomburg, 2011; Kumar and Skolnick, 2012). Although this category of methods is widely used in practice, they are unable to make a prediction when encountering a sequence without significant homologies in the current databases. Thirdly, extracting features from the sequence and classifying the enzyme using machine learning algorithms is the most extensively studied direction (Chou and Elrod, 2003; Cai *et al.*, 2003, 2004; Chou, 2005; Cai and Chou, 2005; Cai *et al.*, 2005; Shen and Chou, 2007; Huang *et al.*, 2007; Lu *et al.*, 2007; Zhou *et al.*, 2007; Lee *et al.*, 2008; Nasibov and Kandemir-Cavas, 2009; Qiu *et al.*, 2009, 2010; Wang *et al.*, 2010, 2011; De Ferrari *et al.*, 2012; Kumar and Choudhary, 2012; De Ferrari *et al.*, 2012; Volpato *et al.*, 2013; Sharif *et al.*, 2015; Zou and Xiao, 2016; Li *et al.*, 2016). Although this direction has already been studied for over 15 years with a number of softwares and servers available, few of them combines the procedure of feature extraction and classification optimization together. Instead, previous studies rely heavily on manually-crafted features, and consider feature extraction and classification as two separate problems. In spite of the success of such methods, with the rapid expansion of the known enzyme sequences, such manually-designed features are very likely to be a suboptimal feature representation which may be unsustainable in the omic era.

In addition to those difficulties, another issue in the protein general function prediction field is the feature dimensionality nonuniformity problem, which usually lies in the sequence-length-dependent features, such as PSSM (position-specific scoring matrix). For example, in this paper, the dimensionality of PSSM can range from 50 by 20 to 5000 by 20, according to the corresponding sequence length. The feature uniformity requirement of mainstream classifiers has pushed out three strategies to this problem. Firstly, avoiding using the sequence-length-dependent features is the most straightforward solution to the problem. Although this approach can work under some certain circumstances, it eliminates the possibility of taking advantage of some powerful representation, such as PSSM, which can provide evolutional information. The second solution is to manually derive sequence-length-independent features from the sequence-length-dependent features (Chen *et al.*, 2013, 2014, 2016). Pse-AAC (pseudo amino acid composition) and Pse-PSSM are typical examples of this category, which have been proved successful in a number of applications (Chou, 2009, 2011). The third solution is to systematically generate sequence-length-independent features, such as string kernels (Leslie *et al.*, 2002, 2004; Rätsch *et al.*, 2005; Wang *et al.*, 2014; Dai *et al.*, 2017), which, however, do not consider the classification problem when extracting features. Despite the previous success of these

three strategies, they still heavily depend on either manually-designed or pre-defined features, which are most likely to be suboptimal. To take full advantage of the bursting of data in recent years, a more robust, automatic framework to extract problem-specific sequence-length-independent features from the sequence-length-dependent ones for dealing with the dimensionality problem is desired.

To conquer the aforementioned limitations, which are homology requirement, feature design and feature dimensionality nonuniformity, here we propose a novel level-by-level prediction approach based on deep learning, by only utilizing the sequence information. The enzyme sequence is represented by two kinds of raw encoding, sequence-length-dependent encoding, such as raw sequence one-hot encoding and PSSM, and sequence-length-independent encoding, such as functional domain encoding. Those two kinds of raw representations are combined into a deep learning model with a novel architecture to perform dimensionality uniformization, feature selection and classification model training simultaneously. This paper makes the following contributions: (1) We propose a framework for hierarchical EC number prediction, the idea of which can also be applied to hierarchical classification of protein general function. (2) To solve the feature dimensionality nonuniformity problem, we propose a robust, automatic framework based on deep learning to extract problem-specific sequence-length-independent features from the sequence-length-dependent ones. (3) We propose a sequence-based enzyme EC number predictor, DEEPred, which is based on the above two frameworks. (4) Two case studies demonstrate our tool's ability of performing functionality prediction of different enzyme isoforms caused by alternative splicing. (5) We investigate the importance of local information in determining the functionality of an enzyme.

2 RELATED WORK

In this section, we introduce four representative methods for enzyme function prediction, followed by a brief overview of deep learning and hierarchical classification.

2.1 EzyPred

EzyPred (Shen and Chou, 2007) is a 3-level EC number predictor, which predicts whether an input protein sequence is an enzyme, and its main class and subclass if it is. It uses two features, pseudo position-specific scoring matrix (Pse-PSSM) and functional domain (FunD) encoding. Pse-PSSM is developed from the pseudo amino acid, a highly innovative manually designed feature which has already been proved successful in a number of problems (Chou, 2009; Hayat and Khan, 2012). It encodes the PSSM of proteins with different lengths using a uniform length matrix, which not only preserves the average score of the amino acid residues in the whole sequence that were changed to a certain type of amino acid during the evolution process but also avoids the complete loss of the sequence order information. Functional domain encoding captures the local functional domain information, which could be referred to Section 3.2.5. With these two features, EzyPred uses optimized evidence-theoretic k -nearest neighbor (OET-KNN) as the classifier, which is an improved version of k -nearest neighbor (KNN). By considering not only the label of the k nearest neighbors of the input query data point but also the distance of the neighbors to the query data as the supporting evidence, OET-KNN alleviates the problem of the original version of KNN for being too sensitive to noise. Although having been developed for ten years, EzyPred still remains as one of the state-of-the-art methods in predicting enzyme function. Its server is easy-to-use with a user-friendly interface as well.

2.2 SVM-Prot

SVM-Prot was proposed in 2004 (Cai *et al.*, 2003, 2004) and updated in 2016 (Li *et al.*, 2016). It can not only predict enzyme functional families but also non-enzyme functional families. It represents the protein sequence using 13 properties, including amino acid composition, polarity, hydrophobicity, surface tension, charge, normalized Van der Waals volume, polarizability, secondary structure, solvent accessibility, molecular weight, solubility, number of hydrogen bond donors in side chain, and number of hydrogen bond acceptors in side chain. Employing composition, translation, and distribution (CTD) to encode each of the above properties, SVM-Prot can make prediction irrespective of sequence similarity. Specifically, composition specifies the fraction of amino acids with a particular property; translation specifies the transition percentage of one amino acid with particular property to another amino acid with different properties; distribution specifies the distribution of amino acids with certain property within the first 25%, 50%, 75%, 100% of the sequence. The original version used SVM as the classifier, while the 2016 update made two more classifiers, KNN and probabilistic neural networks, available.

2.3 COFACTOR

COFACTOR (Roy *et al.*, 2012; Zhang *et al.*, 2017) is a structure-based protein function annotation web-server. In terms of EC number prediction, for an input structural model, which can be obtained either by experiments or computational modeling, it threads the structure against the template library, whose entries' annotation has already been validated by experiments, to identify the template enzyme with the most similar folds and functional sites. Obtaining the template and assuming that structures determine function, the server assigns the EC number of the template enzyme to the query, with the confidence being evaluated by a function considering both the global similarity and the local similarity. In addition to enzyme function prediction, the server can predict the Gene Ontology terms and protein-ligand binding interactions as well. COFACTOR has been proved successful in protein-ligand binding interaction prediction in the CASP9 competition (Moult *et al.*, 2011).

2.4 EFICAz

EFICAz (Tian *et al.*, 2004; Arakaki *et al.*, 2009; Kumar and Skolnick, 2012) is an EC number prediction server using combined approaches. In addition to using the sequence similarity, it also incorporates the PROSITE and PFAM database information. The original version consists of four components: (1) pairwise sequence comparison-based enzyme function inference, (2) conservation controlled hidden Markov model (HMM) iterative procedure for enzyme family classification-based functionally discriminating residue identification, (3) multiple PFAM-based functionally discriminating residue recognition, and (4) multiple PROSITE pattern recognition. Those four components work independently, determining the final prediction by voting. In the later updates in 2009 and 2012, two more components, multiple PFAM family-based SVM evaluation and conservation controlled HMM iterative procedure for enzyme family classification-based SVM evaluation, and larger databases were added. Although it is unable to make EC number annotation if the query sequence has no homology, this server works pretty well in practice with completely four digits assigned.

2.5 Deep Learning and Hierarchical Classification

Since (Krizhevsky *et al.*, 2012), deep learning has become an extremely popular machine learning method. Its two main architectures, convolutional neural network (CNN) and recurrent neural network (RNN), have made a profound contribution to many bioinformatic problems, such as genetic analysis (Xiong *et al.*, 2015), sequence binding specificity prediction (Alipanahi *et al.*, 2015), and cryo-EM image processing (Wang

et al., 2016a). Instead of being a pure classifier that depends on the manually-designed features such as SVM, CNN is considered as an end-to-end wrapper classifier, being able to perform feature extraction based on the classification result and improve the performance in a virtuous circle. As a complement to CNN's capability of capturing significant features from a two-dimensional or three-dimensional matrix, RNN has the potential of encoding long term interactions within the input sequence, which is usually a one-dimensional vector, such as the encoding of English words. In our paper, we combined the advantages of CNN and RNN, using CNN to conduct feature extraction and dimensionality compression starting from the raw two-dimensional encoding matrices, and using RNN to extract the sequential, long-term interactions within the input sequence.

A classification problem with a tree structure in the label space, such as the enzyme function prediction problem discussed in this paper, is often regarded as a hierarchical classification problem. Because this kind of problems can be regarded as multi-label classification and multi-class classification at the same time, the solutions to the problem can be classified into three categories based on different angles to the problem (Silla and Freitas, 2011), namely, flat classification approach, local classifier approach, and global classifier approach. According to the property of our problem, we chose the local classifier approach, which constructs one classifier for each internal node, to be the overall strategy.

3 METHODS

3.1 Datasets

We adopt three datasets in this paper. The first dataset is a widely used one from (Shen and Chou, 2007), constructed from the ENZYME database (released on 01-May-2007), with 40% sequence similarity cutoff. More details of that dataset could be referred to (Shen and Chou, 2007). This dataset is denoted as the *KNN* dataset in the rest of the paper.

Following the same procedure of constructing the *KNN* dataset, we constructed a larger dataset using up-to-date databases. The steps of constructing the dataset are as follows:

1. The SWISS-PROT (released on September 7, 2016) database was separated into enzymes and non-enzymes based on the annotation.
2. To guarantee the uniqueness and correctness, enzyme sequences with more than one set of EC numbers or incomplete EC number annotation were excluded.
3. To avoid fragment data, enzyme sequences annotated with "fragment" or with less than 50 amino acids were excluded. Enzyme sequences with more than 5000 amino acids were also excluded.
4. To remove redundancy bias, we used CD-HIT (Fu *et al.*, 2012) with 40% similarity threshold to sift upon the raw dataset, resulting in 22,168 low-homology enzyme sequences.
5. To construct the non-enzyme part, 22,168 non-enzyme protein sequences were randomly collected from the SWISS-PROT (released on September 7, 2016) non-enzyme part, which were also subject to the 2, 3, 4 steps.

This larger dataset would be referred to as the *NEW* dataset in the rest of this paper.

Other than *KNN* and *NEW*, which will be used as the benchmark to evaluate the proposed method based on cross-fold validation, it is also important to test the generalization power of the proposed method. This can be done by training the model on one dataset, and testing it on an independent and non-overlapping dataset, to avoid being overfitted on a particular dataset. Thus, the third dataset, the benchmark dataset from (Roy *et al.*, 2012), is used for cross-dataset validation. This

non-homologous dataset was collected from PDB, satisfying two requirements: (1) the pair-wise sequence similarity within the dataset is below 30%, and (2) there is no self-BLAST hit within the dataset to ensure that there are no enzymes that are homologous to each other in this set (Roy *et al.*, 2012). All enzymes in this dataset have experimentally determined 3D structures. To avoid overlaps between the training and testing datasets, sequences contained in both our training dataset and this dataset were removed, which reduced the size of the dataset from 318 to 284. This benchmark dataset would be referred to as the *COFACTOR* dataset in the following. Table 1 summarizes the three datasets.

Table 1. Dataset Summary. The *KNN* dataset and *NEW* dataset are used for cross-fold validation. The *COFACTOR* dataset is used for cross-dataset validation.

Dataset	<i>KNN</i> Dataset	<i>NEW</i> Dataset	<i>COFACTOR</i> Dataset
Source	(Shen and Chou, 2007)	Self-constructed	(Roy <i>et al.</i> , 2012)
Enzymes	9,832	22,168	284
Non-enzymes	9,850	22,168	-

3.2 Sequence Representation

The deep learning framework explained in Section 3.3 eliminates the necessity of performing manual dimensionality uniformization and building complex, manually-designed features, which are unlikely to sustain the increasing amount and complexity of data, by conducting feature reconstruction and classifier training simultaneously. Therefore, we use the following raw features, constructed from the input sequence directly, to represent the sequences. Based on their dimensionality, they can be classified into two categories, sequence-length-dependent features and sequence-length-independent features. The first four features described below belong to the former while the last one belongs to the latter.

3.2.1 Sequence One-hot Encoding To preserve the original sequence information, we use one-hot encoding as the first raw representation of the input sequence. This encoding uses one 1 and nineteen 0s to represent each amino acid. For example, A is encoded as $(1 \ 0_1 \ \dots \ 0_{19})$, while C is encoded as $(0_1 \ 1 \ 0_2 \ \dots \ 0_{19})$. For each input protein sequence, the one-hot encoding would produce an L by 20 matrix, where L represents the sequence length, with each row representing a specific spot and each column representing the appearance of a certain amino acid. For those sequences with undetermined amino acid at a particular spot, a vector with 20 0s is used to represent that special position.

3.2.2 Position Specific Scoring Matrix To provide the evolutional information to the training model, we deploy PSSM as the second sequence representation, which was obtained through PSI-BLAST (Altschul *et al.*, 1997) from BLAST+ (Camacho *et al.*, 2009) with three iterations, E-value being 0.002, against SWISS-PROT (released on May 11, 2016).

3.2.3 Solvent Accessibility Solvent accessibility describes the openness of a local region. Because such information is unavailable directly from the database, we use DeepCNF (Wang *et al.*, 2016b) to predict it. Taking the protein sequence as the input, DeepCNF outputs the possibilities of each amino acid of the sequence being in the state of buried, medium or exposed, respectively. The three states are defined by two solvent accessibility thresholds. Buried is defined as less than 10%; exposed is defined as more than 40%; and medium is defined within the range of 10% and 40%. This encoding produces an L by 3 matrix. More details could be referred to (Wang *et al.*, 2016b).

3.2.4 Secondary Structure One-hot Encoding An amino acid could be in one of the three main secondary structure states, alpha-helix, beta-sheet, and random coil, which indicate the protein’s local folding information. Similar to solvent accessibility, we take advantage of DeepCNF (Wang *et al.*, 2016b) to predict the secondary structure of a given sequence, whose result is an L by 3 matrix, each row of which shows the possibility of the amino acid folding into alpha-helix, beta-sheet or random coil, respectively. The details could be referred to (Wang *et al.*, 2016b).

3.2.5 Functional Domain Usually, a protein sequence contains one or several functional domains, which provide distinct functional and evolutional information. Pfam (Finn *et al.*, 2016) is a collection of such functional domains, each represented by an HMM. Searching against the database and encoding in the following way generates the functional domain encoding used in our model.

1. Pfam has its default searching engine as HMMER (Eddy, 2011). For each protein sequence, we use HMMER, with the inclusion E-value threshold as 0.01, to search against Pfam (Pfam 30.0 Released on July, 1 2016), which contains 16,306 entries.
2. We employ a 16,306-dimensional vector to encode the searching result. If the i -th entry in the database is reported as hit, 1 appears on the corresponding position of the vector, otherwise it is 0.

As a result, the functional domain encoding of a protein sequence would be:

$$\mathbf{F}_{FuncD} = [I_1 \ I_2 \ \dots \ I_i \ \dots \ I_{16,306}] \quad (1)$$

where

$$I_i = \begin{cases} 1, & \text{the } i\text{-th entry in Pfam reported as hit,} \\ 0, & \text{otherwise.} \end{cases} \quad (2)$$

3.3 Classification Model

The enzyme function prediction problem has a tree-structured label space, which makes it a typical hierarchical classification problem. To solve this kind of problems, we propose a level-by-level prediction framework, building a model for each internal label node. The model contains two main components, namely, the problem-specific feature extractor, which is able to perform dimensionality uniformity and feature extraction, and the classifier. Such a novel, end-to-end model can perform feature selection and classifier training simultaneously in a virtuous circle, making it more likely to achieve high performance.

3.3.1 Level-by-level Strategy As have been discussed in the hierarchical classification part, because of the relative small size (22,168 data points are assigned to 58 classes until the second digit) and, even worse, the extreme imbalance property (for example, the *NEW* dataset contains 22,168 sequences belonging to non-enzyme while only 10 sequences belonging to subclass 1.20) of the data, we choose the local classifier approach for this problem. Particularly, the level-by-level prediction strategy is used. That is, given a sequence, the trained model would firstly predict whether it is an enzyme or not. If it is an enzyme, the model will further predict the first digit, which indicates its main class. Knowing the main class, our algorithm will choose the trained model for that specific main class and further predict the second digit, that is, the subclass. Corresponding to the label hierarchy, we build one model for determining whether the input is an enzyme, one model to determine the first digit, six models to determine the second digit. This prediction strategy could be referred to Figure 1(A).

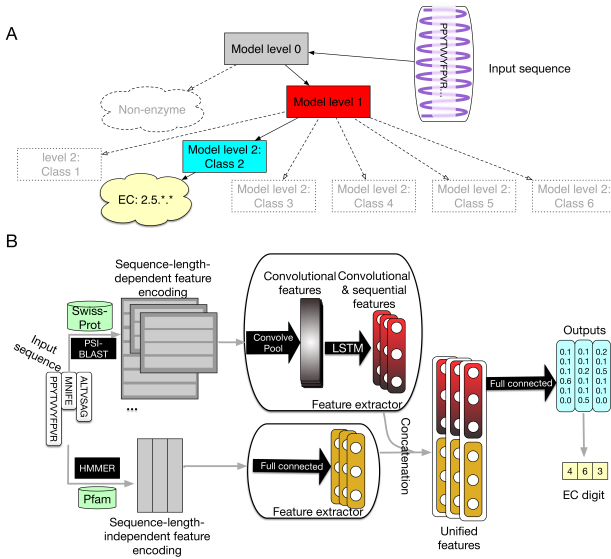


Fig. 1. (A) Strategy for predicting detailed function. Following the structure of the EC number system, we use this level-by-level classification approach. (B) Overview of the model. We use the convolutional neural network component to extract convolutional features and the recurrent neural network component to extract sequential features from each sequence-length-dependent raw feature encoding, followed by a fully-connected component, which concatenates all extracted features together, serving as the classifier. Here we show the procedure of predicting the main class digit of three enzymes with different sequence lengths.

3.3.2 Deep Neural Network Model For each level of prediction, we build the end-to-end model based on several deep neural network components. In terms of the sequence-length-dependent features, such as PSSM, we build a feature extractor exploiting the convolutional neural network component to extract convolutional features from the input map and, after that, a recurrent neural network component, comprised of long short-term memory (LSTM) cells, to extract sequential features from the output of the previous component. As for the sequence-length-independent feature, i.e., the functional domain encoding, which is a vector, we use a fully connected component to perform dimensionality reduction and feature extraction. We employ a fully connected component to combine those different pieces of information together, followed by a softmax layer for classification. The structure of the model could be referred to Figure 1(B). More details of the model structure could be referred to Section S1. It should be noted that the model we evaluate throughout the paper except for Section 4.4 uses only three input features, sequence one-hot encoding, PSSM and functional domain encoding. We encode local features, i.e., secondary structure and solvent accessibility, only in Section 4.4 to evaluate the importance of local information. During training, the training error is back-propagated to each component. The error would guide the convolutional neural network component and recurrent neural network component to perform an end-to-end feature selection, weighing more on the features which would improve the final performance while weighing less on unimportant features automatically. At the same time, the weights of other components would be adjusted simultaneously to adopt the change. Such coupling effect of feature extraction and classifier training optimizes the performance dramatically.

The high complexity and flexibility of the proposed model bring high risks of overfitting. We adopt several methods to alleviate the problem. The first method is weight decay, which is a well-known method to handle the overfitting issue. The deep neural network model is likely to reproduce the detail of noise, which is usually non-smooth, by the usage of extreme

weights. Modifying the objective function by adding an L-2 norm term of weights, we can reduce the probability of ending up with extreme weights and thus mitigate the overfitting issue. The second method is dropout (Srivastava *et al.*, 2014). The key idea of this technique is to randomly drop nodes during training so as to prevent them from co-adapting too much and reduce the model complexity while preserving its power. The third method is batch normalization (Ioffe and Szegedy, 2015). This approach extends the idea of normalization, which is usually done during data pre-processing. In fact, because of the weight and parameter adjustment, the input of the internal layers of the model is possible to be too large or too small, known as “internal covariate shift”, which makes the pre-processing normalization meaningless. To conquer the issue, in addition to normalizing the data before inputting them in the model, we also normalize the input of each internal layer. In addition to the advantage of mitigating the overfitting problem, this manipulation would also reduce the strong dependency of knowledge-intensive initialization when training the model and allow a larger learning rate when tuning the model.

We choose adaptive moment estimation (Adam) as the optimizer (Kingma and Ba, 2014), which is an improved version of stochastic gradient descent, to minimize the weighted cross entropy loss. In this way, our method could handle the class imbalance issue by re-scaling predictions of each class by its weight. Instead of setting the learning rate as a hyper-parameter manually as in stochastic gradient descent and momentum, this method computes the adaptive learning rate of each individual parameter by estimating the first and second movement of the gradients at the cost of computational time and memory. Essentially, this optimizer combines the advantage of RMSprop (Tieleman and Hinton, 2012), which computes the adaptive learning rate during each step, and momentum, which reduces the oscillation problem of stochastic gradient descent by making the weight update considering both the gradient and the update of the previous step.

When training the second-digit prediction models, we adopt an idea that is similar to transfer learning. Since the limited number of data is further divided into six parts corresponding to the six main classes, the amount of data belonging to each main class is insufficient to produce a model with the ability to extract features and being generalized well. To solve this problem, we pre-train the convolutional neural network component and the recurrent neural network component by using all the training data. Then for training each second-digit prediction model, we fix the parameters of those components and only fine tune those fully connected components using the specific subset of the training data.

In practice, we use TensorFlow (Abadi, 2016) as the framework to construct the deep neural network. With two Pascal Titan X cards, it takes around 4 hours to obtain a well-trained model. In Section S2, we provide details on setting the model parameters.

4 RESULTS AND DISCUSSION

4.1 Evaluation Criteria

For the enzyme or non-enzyme prediction, since it is a binary classification problem, we use accuracy, Cohne's Kappa Score (Viera and Garrett, 2005), precision, recall and F_1 score to evaluate the classifiers' performance. For other predictions, since they are multi-class classification problems, we use accuracy, Cohen's Kappa Score, Macro-precision, Macro-recall and Macro- F_1 score to evaluate the classifiers' performance, whose definitions are in Section S3.

4.2 Compared Methods

For the cross-fold validation, in which training and testing are based on different parts within the same dataset, we compare our method with five other methods, including two state-of-the-art methods, EzyPred (Shen

and Chou, 2007) and SVMProt (Li *et al.*, 2016), and three baseline methods. One of the baseline methods uses SVM with the raw features used in our model; another baseline method uses SVM with Pse-PSSM; and the last baseline method uses the traditional neural network with our raw features. Due to the unchangeable database of EFICAz (Kumar and Skolnick, 2012) and COFACTOR (Zhang *et al.*, 2017), we do not include them in the cross-fold validation comparison. However, we perform cross-dataset validation, where the training and testing are performed on different datasets, to compare our method with EzyPred, SVM-Prot, COFACTOR, and EFICAz.

4.3 Cross-fold Validation

Here we report the 5-fold cross validation results, which are shown in Figure 2. Our method almost always outperforms the other methods in both the *KNN* dataset and the *NEW* dataset across the five criteria and across the three hierarchical levels of prediction. As for the *NEW* dataset, DEEPre outperforms the other five methods consistently in level 0 and level 1 prediction across the five criteria. As for the level 2 prediction, the only criterion that DEEPre does not improve over the existing methods is the Macro-Precision, which is an unweighted average of precision of each label. The appearance of very small classes (for example, subclass 1.20 only has 10 enzymes) in the second level prediction might be the reason for this result. In terms of the *KNN* dataset, although the smaller dataset makes the improvement of DEEPre over the other methods in level 0 prediction less significant, it still significantly outperforms the other methods in level 1 and level 2 classification.

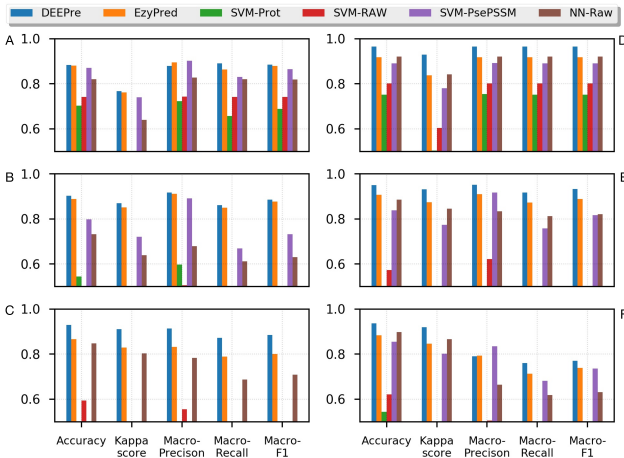


Fig. 2. Cross-fold validation results. (A) Performance comparison of level 0 prediction (predicting whether the input is an enzyme or not) on the *KNN* dataset. (B) Performance comparison of level 1 prediction (predicting the input enzyme's main class) on the *KNN* dataset. (C) Performance comparison of level 2 prediction (predicting the input enzyme's subclass given the main class) on the *KNN* dataset. (D) Performance comparison of level 0 prediction on the *NEW* dataset. (E) Performance comparison of level 1 prediction on the *NEW* dataset. (F) Performance comparison of level 2 prediction on the *NEW* dataset.

4.4 Feature Importance Analysis

It is believed that both global features and local features determine the function of a protein. For detailed function, local information would weigh even more in determining it. The features extracted by the convolutional component and the recurrent component from PSSM and sequence raw encoding could be considered as global features while the functional domain encoding would be considered as a local feature. We remove the

three input raw encoding one by one and show the comparison of their performance on the *NEW* dataset. The comparison is shown in Figure 3(A). It is clear that as the level goes deeper, the importance of functional domain is evidently increasing, which demonstrates the well-recognized hypothesis. To further prove it, we design another experiment, in which we input more local feature encodings, including secondary structure and solvent accessibility, into our model. Details of this experiment could be referred to Section S4. Figure 3(B) shows the performance comparison of this model and the previous model in level 2 prediction. It is clear that the additional local features further improve the performance of our model, with accuracy improved by 1.8% while Macro-precision, Macro-recall and Macro-F1 score improved by at least 11%.

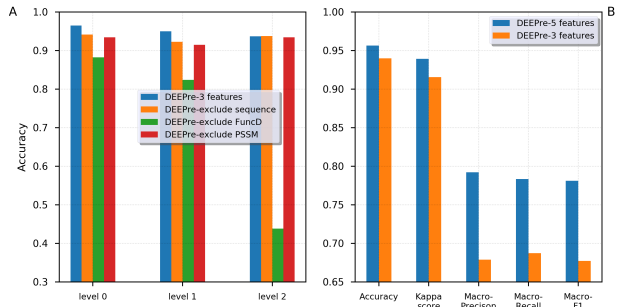


Fig. 3. (A) Feature contribution investigation considering sequence one-hot encoding (sequence), PSSM, and functional domain (FuncD). (B) The performance change of the model before and after we input more local feature encoding. Macro-precision, Macro-recall and Macro-F1 score are improved by at least 11% by inputting solvent accessibility and secondary structure information.

4.5 Cross-dataset Validation

In this experiment, we directly compare the performance of different servers in predicting the first digit and the second digit of an enzyme. We use the *COFACTOR* benchmark dataset, which is proved to be a difficult dataset in the enzyme function prediction field (Roy *et al.*, 2012), as the test dataset. Firstly, we eliminate the sequences in the *COFACTOR* benchmark data which overlap with the DEEPre's training database (*NEW*) by 40% sequence similarity filtering, reducing the data size from 318 to 284, to ensure that there is no bias in the DEEPre's results. Then we input the remaining sequences to each server manually and collect the prediction results. For *COFACTOR*, since it is quite time-consuming to run the server, about 4 hours to obtain the result for one query, we report the results from the original paper. As shown in Figure 4, for the first-digit prediction, DEEPre outperforms the other servers consistently across the five criteria, improving the accuracy by at least 6% over the other servers, including *COFACTOR*. This is significant because *COFACTOR* requires 3D structures of enzymes whereas DEEPre only requires the sequence information. On the other hand, we should admit that we have changed the original *COFACTOR* dataset to some extent by reducing the overlap between the training and testing sets, which might explain some of the performance difference between *COFACTOR* and DEEPre. We should also notice that all of those five servers have different training datasets, but those training datasets highly overlap with each other and each method was optimized on its corresponding dataset. In addition, although we removed overlapping enzymes from our training set and the *COFACTOR* test set, there may still be homologs of the enzymes in *COFACTOR* in our training set. However, DEEPre is a sequence-based statistical method, which explores the statistical properties of training data and does not benefit from knowing

enzyme structures. Therefore, although the performance of DEEPre on *COFACTOR* may still be biased, the influence is not as much as that by nearest neighbor-based methods. We also perform comparison of different servers' performance on the second-digit prediction (Section S5). The results show that DEEPre and EFICAz both perform well on the second-digit prediction on the *COFACTOR* dataset. It is worth noting that EC numbers have regular corrections, such as deletions and transfers. We check all the corrections that are related to the test enzymes in the *COFACTOR* dataset and find that none of them influences the comparison reported here.

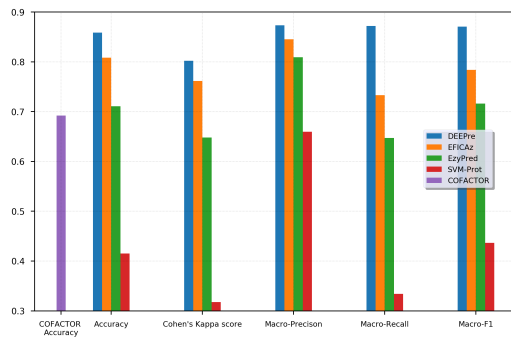


Fig. 4. The performance comparison of different servers on predicting the main class of the *COFACTOR* dataset. DEEPre improves the prediction accuracy over the other servers by at least 6%.

4.6 Third-digit and Fourth-digit Prediction

Using the same framework described above, we are also able to predict the enzyme's third digit, which represents its sub-subclass, on the *NEW* dataset. The accuracy across all the sub-subclasses is 0.9415; the Kappa score is 0.8918; the macro-precision is 0.8942; the macro-recall is 0.8578; and the macro-F1 score is 0.8665. Regarding the fourth-digit prediction, more data are needed to perform normal machine learning training-and-testing procedure. For example, within the sub-subclass 1.1.1 in the *NEW* dataset, there are 188 classes. Each of those classes has less than 40 enzyme sequences, with 175 classes having less than 10 enzyme sequences. Using the current dataset with such distribution would lead to unreliable results.

4.7 Case Study

Glutaminase is a phosphate-activated enzyme, which catalyzes the first step of glutaminolysis, hydrolysing glutamine into glutamate (Curthoys and Watford, 1995). The alternative splicing of its messenger RNA results in its three isoforms, with isoform 1 and isoform 3 being capable of catalyzing while isoform 2 lacking the catalytic activity (Li *et al.*, 2017). To validate our model's ability to distinguish the different functionality of different isoforms, we obtained the sequences of the three Glutaminase isoforms from the UniProt and put them into our model. Our model predicted that isoform 1 and isoform 3 of Glutaminase were hydrolases acting on carbon-nitrogen bonds, being consistent with the experimental results. Our model also recognized isoform 2 as non-enzyme, which is consistent with the experimental result as well.

Aurora kinases B is a key enzyme regulating chromosomal segregation during mitosis, ensuring correct chromosome alignment and segregation as well as chromatin-induced microtubule stabilization and spindle assembly (Carmena *et al.*, 2009). Over-expression of it is possible to cause unequal distribution of genetic information, resulting in aneuploid cells, which may become cancerous (Sorrentino *et al.*, 2005). Aurora kinases B has five isoforms resulted from alternative splicing. Four of them

have roughly equal length with high similarity, while isoform 3, having high expression in the metastatic liver with no expression in the normal liver, is only half of the length of the "canonical" isoform (142 amino acids v.s. 344 amino acids). Despite its much shorter length, the isoform does not lose its functionality. To further validate our model's ability of handling isoforms' functionality prediction, we collected the sequence of the five isoforms from the database and put them into our model. Our model's result is consistent with the experimental results. Particularly, our model predicted the functionality of the isoform 3 successfully, despite its sequence's large difference from the "canonical" sequence.

The detailed performance comparison of different servers on these two case studies could be referred to Section S6. Among the five compared methods, only our method and EzyPred produced correct predictions for both cases.

5 CONCLUSION

In this paper, we proposed a novel end-to-end feature extraction and classifier training method for enzyme function prediction. The method proposed in this paper would force the model to learn to extract features by itself and adapt the parameters of the classifier simultaneously so that it can improve the performance in a virtuous circle. The thorough experiments conducted on two datasets demonstrate the high performance of our method in both a smaller dataset from 10 years ago and a larger dataset constructed half a year ago. The cross-dataset validation experiment proves the performance of our model in handling sequences with no close homologs. Although it is just a starting point, the user-friendly server, DEEPre, will provide users a good guess of enzyme function and help them set up downstream experiments. Since DEEPre predicts a score for each candidate value of a certain EC digit, it can be potentially used to detect the enzyme promiscuity (Carbonell and Faulon, 2010; Mellor *et al.*, 2016), which means that some enzymes show multiple activities by either accepting multiple substrates or catalyzing multiple reactions. Our webserver provides the predicted scores for all candidate EC values. In addition to providing the server in the enzyme function prediction field, we believe the idea proposed in this paper can be quite helpful in handling the feature length nonuniformity problem and the dataset evolution in a wide spectrum of computational biology problems.

Among the global features, the most important one is the functional domain (Figure 3(A)). This sequence-length-independent feature cannot be replaced by sequence-length-dependent features. Nevertheless, the global and local features explored in this paper provide complementary information and together provide improved performance (Figure 3(B)), in spite of leading to the higher dimensionality of the predictor.

A large number of protein function prediction problems are hierarchical classification problems, such as GO term (Camon *et al.*, 2004), transporter classification (Saier *et al.*, 2016), and GPCR hierarchy (Davies *et al.*, 2007). The high extensibility and flexibility of our level-by-level prediction framework make it possible to adopt our framework in those problems. Furthermore, the robust, automatic framework based on deep learning to extract problem-specific sequence-length-independent features from the sequence-length-dependent features can also be extended to other features in addition to the features mentioned in this paper.

There are two directions of the future work. First, more robust methods for the fourth-digit prediction are needed. The increasing number of enzymes that have experimentally validated functions, as well as the advance in method development for learning from imbalanced-data and small samples (Maadooliat *et al.*, 2016), provide potential solutions to the problem. Second, instead of predicting the EC numbers for enzymes, it is practically useful to predict enzymatic reactions of the enzymes. The use of reaction fingerprints, for instance, could be one viable solution for this

(Segler and Waller, 2017). Another possible solution is through the use of descriptors of the reaction centers as in (Rahman *et al.*, 2014).

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