Paper

Hierarchical Multi-label Classifier for Gene Ontology Annotation using Multi-head and Multi-end Deep CNN Model

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Gene ontology (GO) annotation is known to be a very complicated multi-label classification task, and the hierarchical multi-label classification (HMC) approaches with local classifiers have been shown to be effective for the task. In a traditional HMC method, a set of hierarchically organized simple local classifiers are usually used, each of which for one hierarchical level separately. In this paper, we propose a novel hierarchical multi-label classifier implementing the whole set of hierarchically organized local classifiers in one deep convolution neural network (CNN) model with multiple heads and multiple ends (MHME). The proposed MHME CNN model consists of three parts: the body part of a deep CNN model shared by different local classifiers for feature extraction and feature mapping; the multi-end part of a set of autoencoders performing feature fusion transforming the input vectors of different local classifiers to feature vectors with the same length so as to share the feature mapping part; and the multi-head part of a set of linear multi-label classifiers. Furthermore, a sophisticated recursive algorithm is designed to train the MHME CNN model to realize the functions of a set of hierarchically organized local classifiers. In this way, by sharing a deep CNN with multiple local classifiers, we are able to construct more powerful local classifiers for each level with limited training samples, and to achieve better classification performance. Experiment results on various benchmark datasets show that the proposed deep CNN based model has better performance than the state-of-the-art traditional models. Moreover, it gives rather good performance even under a transfer learning.

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Keywords: hierarchical multi-label classification, GO annotation, deep learning, autoencoder, feature extraction

1. Introduction

As gene ontology (GO) reveals, individual genes contribute to the biology of an organism at the molecular, cellular and organism levels. GO annotation plays an essential role in many biomedical studies (1). However, with the GO functions continuing to be enriched, the number of GO functions which could be annotated is more than 40 thousands. Therefore, fully experimental protein annotation becomes time-consuming and cost-consuming; automatic prediction with the help of a machine learning method is widely considered to be more practical. Compared to the biological experiment methods, the machine learning method is more economical and convenient.

Automatic GO annotation method is a process to predict the GO functions (classes) as a multi-label classification task (2) by checking the combination of proteins. But it could not be dealt with a normal multi-label method as GO annotation is very complicated, not only the number of GO functions is huge, but also the functions are arranged in a hierarchy:

typically a direct acyclic graph (DAG)⁽³⁾ where each node represents a function. These functions could be divided into different levels according to the grade. It is based on the 'is-a' relationship ⁽⁴⁾. Hierarchical multi-label classification is one kind of the algorithms known to have better performance ⁽⁵⁾, since it is a method ranking labels into a hierarchical structure, and performing classification on each level independently.

Hierarchical multi-label classification (HMC) (6)~(8), is a kind of methods which separate the labels into hierarchical levels and apply the classifiers to different levels, respectively. Multiform HMC methods could be divided into two circumstances, local hierarchical classification approach and global hierarchical classification. Local approach also called topdown approach (9)~(11) trains the hierarchical structure network from the higher level to the lower level. Global approach uses a classification algorithm which learns a global classifier about the whole classes. For example, the authors of Ref. (12) investigated the flat classification algorithm naive Bayes applied in protein function prediction. Flat classification is the simplest approach in HMC methods, but it ignores classrelationship. However, the local approach is used to solve protein function prediction, as it is used augmented versions of dealing with the problem which has hierarchical classes (13).

In Refs (14)~(17), R. Cerri et.al applied the hierarchical multi-label classification to protein function prediction problem with local approach. It is a set of multilayer perceptron (MLP) classifiers trained for each level, separately. The local-

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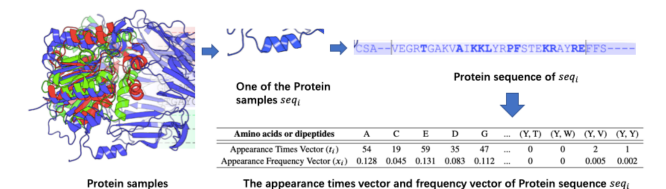


Figure 1: An image of '1-gram + 2-gram' encoding based feature extraction of protein sequences.

based classification model trains MLPs for each level, called a hierarchical multi-label classification with local MLP (HMC-LMLP). In a HMC-LMLP, the predictions with MLP at a given level are used as inputs to the neural network responsible for the prediction in the next level. These methods focused on optimizing model by training it level by level and propagating the positive signal from the previous output. Although the HMC-LMLP achieves a state of the art performance, it is difficult to construct more powerful local classifiers due to the limitation of training samples available in each class level.

Following the basic idea of HMC with local neural network models and the analyzing method of protein sequences, in this paper, we propose a novel hierarchical multi-label classifier implementing the whole set of hierarchically organized local classifiers in one deep convolution neural network (CNN) model with multiple heads and multiple ends (MHME). For this purpose, the proposed MHME CNN model consists of three parts: the body part, the multi-end part, and the multihead part. The body part is a deep CNN model shared by different local classifiers for the feature extraction and feature mapping. The multi-end part is a set of autoencoders performing feature fusion and dimension reduction which transforms the input vectors of different local classifiers to feature vectors with the same length so as to share the feature mapping of deep CNN model. The multi-head part is a set of linear multi-label classifiers realizing the multi-label protein function prediction. Furthermore, we design a sophisticated recursive algorithm to train the MHME CNN model in such way that it performs the functions of a set of hierarchically organized local classifiers. In this way, we are able to design a better hierarchical multi-label classifier by increasing the complexity of the feature mapping model via sharing a deep CNN with multiple local classifiers, and achieve better annotation performance.

On the other hand, protein sequence as a very normal feature has been used in many types of research. Considering that protein sequence consists of 20 different amino acids, which is similar to nature language (18). In this way, the n-gram algorithm utilized to encode proteins is defined as the appearance frequency of n consecutive amino acids (19)(20). The n-gram method generated from natural language processing is now developed in the bioinformatics field. In Ref. (21), the authors investigated lager numbers of protein GO function annotations dealing with convolutional neural networks. Protein sequences have been represented as one-hot encoding in

units of the tripeptide, and a single layer CNN model used as a feature extractor. In this paper, we use a '1-gram + 2-gram' encoding method to obtain an appearance frequency vector as the input of the MHME CNN model.

The rest of our paper is organized as follows. Section 2 introduces the '1-gram + 2-gram' encoding method and formulates local classifiers for each level. Section 3 describes the details of the MHME CNN model and the training algorithm. Section 4 carries out experiments on various benchmark datasets and compares the proposed MHME CNN model with the related existing methods. Finally, Section 5 gives the conclusions.

2. Problem Formulation

In this section, we first introduce the notations and then formulate the problem.

2.1 The Notation and the Problem Suppose we denote the protein instances by $\mathbf{S} = \{seq_1, seq_2, seq_3, ...seq_i | i = N\}$, N is the number of instance. and seq_i represent the protein sequence of instance i. Protein functions are denoted by $\mathbf{C} = \{c_1, c_2, c_3, ...c_j | j = C\}$, C is the species number of functions. Different from sample binary classification and multi-class classification, in hierarchical classification databases used $D = \{(x_i, y_i), 1 \le i \le N\}$ to express the instance in different levels, if x_i is belong to function c_j , then $c_j = 1$, or $c_j = 0$. We use $y_n = \{c_1, c_2, c_3, c_4, ...c_j | j = C\}$ to express the label matrix. At label space, we assume that the number of level of the hierarchy classes is q, expressed as $\mathbf{Y} = \{Y_1, Y_2, Y_3, ...Y_n | n = q\}$. For label in level q is that $Y_q = \{c_{q1}, c_{q2}, ...c_{qn} | c_{qn} \in \mathbf{C}\}$.

2.2 '1-gram + 2-gram' Encoding The encoding method encodes the protein sequences of amino acids seq_i (i = 1, N) into appearance frequency vectors, x_i (i = 1, ..., N), which are column vectors with a length of 420, the number of 20 amino acids and 400 dipeptides.

Figure 1 shows an image of the '1-gram+2-gram' encoding method. For a given protein sequence seq_i , the appearance time vector t_i is first obtained by counting the number of times of each amino acid and dipeptide appearing in the protein sequence seq_i . Then the appearance frequency vector x_i is defined by

$$x_i = \frac{t_i}{\dim(x_i)} \tag{1}$$

where dim $(x_i) = 20^1 + 20^2 = 420$. x_i (i = 1,...,N) is then

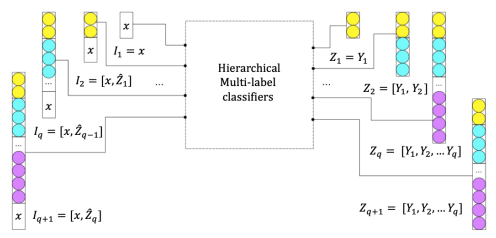


Figure 2: The formulation of HMC with local classifers.

used as an input vector of our MHME CNN model, where *N* is the number of protein samples.

2.3 Local Classifiers for Each Level Figure 2 shows the formulation of HMC with local classifiers. The local classifiers are designed to be hierarchically organized way so as to capture a top-down relationship of labels between different levels. The local classifier for level 1 (classifier 1) has the input vector of $I_1 = x$ and the output vector of $Z_1 = Y_1$. The local classifier for level 2 (classifier 2) has the input vector of $I_2 = [x, \hat{Y}_1] = [x, \hat{Z}_1]$, combined x with the output vector of classifier 1, and the output vector of $Z_2 = [Y_1, Y_2]$. In this way, the local classifier for level n (n = 1, ..., q) (classifier n) will have the input vector of $I_n = [x, \hat{Y}_1, \hat{Y}_2, ..., \hat{Y}_{n-1}] = [x, \hat{Z}_{n-1}],$ combined x with the output vector of classifier n-1, and the output vector of $Z_n = [Y_1, Y_2, ..., Y_n]$. Traditionally, the classifiers 1 to n are implemented by using simple MLP models due to the limitation of training samples in each level.

2.4 An Additional Global Classifier There exists not only top-down relationship of labels between different levels, but also button-up label relationship. In order to capture both top-down relationship and button-up relationship of labels between different levels, in the proposed model, we add another global classifier which has the input vector of $I_{q+1} = [x, \hat{Z}_q]$, combined x with the output vector of the last level classifier q, and the output vector of $Z_{q+1} = [Y_1, Y_2, ..., Y_q]$. †

Traditionally, one constructs q+1 neural network classifiers separately for the above q local classifiers and one global classifier.

$$Z_n = F_n(I_n), \quad n = 1, ..., q + 1$$
 (2)

where $I_1 = x$ and $I_n = [x, \hat{Z}_{n-1}]$. In the next section, we will design a deep CNN model with multiple heads and multiple ends to implement both the q local classifiers and the additional global classifier in one deep neural network, and design a sophisticated recursive algorithm to train the MHME CNN model to perform a set of hierarchically organized powerful classifiers with the limited training samples.

3. The Deep MHME CNN Model

Figure 3 shows the structure of deep CNN model with mul-

tiple ends and multiple heads (MHME) to implement the q+1 hierarchically related classifiers using one deep CNN model. The MHME CNN model consists of three parts: the body part, the multi-end part and the multi-head part, which realizes the q+1 classifiers with sharing the body part, referred to Figure 3.

$$Z_n = f_n\left(\Omega_n^{(3)}, \phi\left(\Omega^{(2)}, A_n\left(\Omega_n^{(1)}, I_n\right)\right)\right),\tag{3}$$

where n=1,...,q+1, $\phi(\Omega^{(2)},\cdot)$, $A_n(\Omega^{(1)}_n,\cdot)$ and $f_n(\Omega^{(3)}_n,\cdot)$ denote the body part, the multi-end part and the multi-head part, and $\Omega^{(2)}$, $\Omega^{(1)}_n$, $\Omega^{(3)}_n$ are the parameters, respectively.

3.1 The Multi-End Part: A Set of Autoencoders The multi-end part corresponds to the inputs of the q+1 hierarchically related classifiers. As can be seen from the description of the previous section, the local classifiers and the additional global classifier have their input vectors with different lengths. In order to share the body part for feature extraction, these input vectors should be transformed to a set of feature vectors with same length at the specific feature space.

$$\begin{cases} a_1 = x \\ a_n = A_n(\Omega_n^{(1)}, I_n), & n = 2, ..., q + 1 \end{cases}$$
 (4)

where a_n is the outputs of multi-end part. For simplicity, we set $\dim(a_n) = \dim(x)$.

Due to no teacher signals for a_n , we apply autoencoders (22), which realizes the transforming of Eq.(4) by minimizing the information loss. As shown in Figure 4, an autoencoder neural network is an unsupervised learning algorithm that applies backpropagation (23), by setting the target values to be equal to the inputs. The general idea of autoencoder is to represent the data through a nonlinear encoder to a hidden layer and use the hidden units as the new feature representations. The major purpose of autoencoder is to learn an approximation to the identity of original protein representation, by encouraging its output to be as similar to its input as possible. Recently, the variants of original autoencoder have drawn increasing attention as an effective feature extractor or descriptor (24). The autoencoder plays two roles: feature fusion and dimension reduction.

3.2 The Body Part: A Deep CNN Model The rect-

 $[\]dagger \hat{Z}_1, \hat{Z}_2, ..., \hat{Z}_q$ are the prediction values of $Z_1, Z_2, ..., Z_q$ by the local classifiers

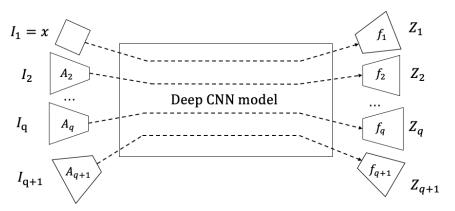


Figure 3: The structure of the multi-head and multi-end deep CNN model.

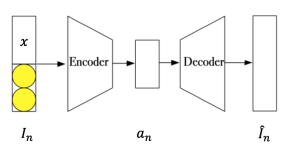


Figure 4: The structure of autoencoder A_n .

angle in the middle of Figure 3 represents the body part, which realizes a feature mapping or feature extraction. CNN outperforms in dealing with the spatial information like graph, text, etc. Recently, it is also shown to be effective in extracting feature from protein sequences in the some related work (21)(25). To demonstrate the ordered arrangement of protein sequences, the proteins are described as the frequency of amino acids and dipeptides. Moreover, the GO terms (predictions of linear classifiers, mentioned in the second section) added in input during the training makes the features very high-dimensional. Compared with the one-layer CNN model used in Ref. (21), a deep CNN model is more reasonable for our purposes of feature extraction and feature mapping. The details of deep CNN model used in our experiments will be described in Subsection 5.2.1.

$$\phi_n = \phi(\Omega^{(2)}, a_n), \quad n = 1, ..., q + 1$$
 (5)

where ϕ_n is the output of the body part.

3.3 The Multi-Head Part: A Set of Linear Classifiers The multi-head part consists of a set of linear multi-label classifiers with input vectors from the outputs of body part. As the linear multi-label classifiers, we simply use a linear network with logistic sigmoid outputs.

$$Z_n = f_n(\Omega_n^{(3)}, \phi_n), \quad n = 1, ..., q + 1$$
 (6)

4. Training Method

The output of hierarchical multi-label classifier Z_{q+1} is calculated recursively by

$$Z_1 = f_1\left(\Omega_1^{(3)}, \phi\left(\Omega^{(2)}, A_1\left(\Omega_1^{(1)}, I_1\right)\right)\right), \ I_1 = x$$

$$Z_{2} = f_{2} \left(\Omega_{2}^{(3)}, \phi \left(\Omega^{(2)}, A_{2} \left(\Omega_{2}^{(1)}, I_{2} \right) \right) \right), \ I_{2} = [x, Z_{1}]$$

$$...$$

$$Z_{q+1} = f_{q+1} \left(\Omega_{q+1}^{(3)}, \phi \left(\Omega^{(2)}, A_{q+1} \left(\Omega_{q+1}^{(1)}, I_{q+1} \right) \right) \right),$$

$$I_{q+1} = [x, Z_{q}]. \quad \cdots \quad \cdots \quad \cdots \quad (7)$$

1) Training of Parameters $\Omega_n^{(1)}$ $\Omega_n^{(1)}$ (n=1,...,q+1) are the parameters of autoencoders. As shown in Figure 4, they are trained in an unsupervised manner by optimizing the mean squared error (MSE) loss function $E_n = \varepsilon_{MSE}(\hat{I}_n, I_n)$. Let z be the input of encoder and \hat{z} be the output of decoder. Then ε_{MSE} is defined by

$$\varepsilon_{MSE}(\hat{z}, z) = \frac{1}{N} \sum_{i=1}^{N} (\hat{z}_i - z_i)^2$$
 (8)

where N is the number of samples.

2) Training of Parameters $\Omega^{(2)}$ and $\Omega_n^{(3)}$ $\Omega^{(2)}$ and $\Omega_n^{(3)}$ (n=1,...,q+1) are the parameters of deep CNN and the linear network with logistic outputs. They are trained in a supervised manner by optimizing the sigmoid cross-entropy (SCE) loss function $E = \sum_{i=1}^n \varepsilon_{SCE}(\hat{Z}_n, Z_n)$. In this formulation, ε_{SCE} is defined by

$$\varepsilon_{SCE}(\hat{z}, z) = -\frac{1}{N} \sum_{i=1}^{N} \sum_{j=1}^{C_n} [z_{ij} \times log(\hat{z}_{ij}) + (1 - z_{ij}) \times log(1 - \hat{z}_{ij})]$$

$$(9)$$

and $C_n = \dim(Z_n)$.

However, as we can see from Eq.(7) that the q+1 classifiers are hierarchically related. A sophisticated training algorithm is needed to train the parameters of $\Omega_n^{(1)}$, $\Omega^{(2)}$ and $\Omega_n^{(3)}$ (n=1,...,q+1). The training algorithm consists of the following three steps.

- Step 1: Pre-training of $\Omega^{(2)}$ and $\Omega_n^{(3)}$ Set $a_n = x$ (n = 1, ..., q + 1). Then train $\Omega^{(2)}$ and $\Omega_n^{(3)}$ based on Eqs. (5), (6) and (9). After the pre-training, we are able to calculate the approximated predictions \hat{Z}_n (n = 1, ..., q + 1) using Eqs. (5) and (6);
- Step 2: Training of $\Omega_n^{(1)}$ Let $I_1 = x$ and $I_n = [x, \hat{Z}_{n-1}]$ (n = 2, ...q + 1). Then train $\Omega_n^{(1)}$ based on Eqs. (4) and (8).
- Step 3: Training of $\Omega^{(2)}$ and $\Omega^{(3)}_n$ Let $I_1 = x$ and $I_n = [x, \hat{Z}_{n-1}]$ (n = 2, ...q + 1). Then

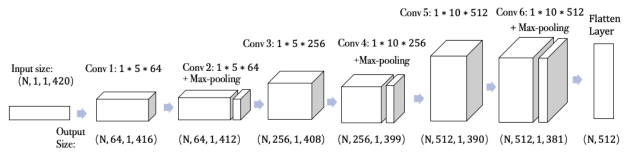


Figure 5: The structure of deep CNN model consisting of 6 convolutional layers and one flatten layer.

train $\Omega^{(2)}$ and $\Omega_n^{(3)}$ based on Eqs. (7) and (9). After the training, we calculate the predictions \hat{Z}_n (n = 1, ..., q + 1) using Eq. (7);

 Step 4: Repeat Steps 2 and 3, until an appropriate stop condition is satisfied.

5. Experiments

In this section, the proposed MHME CNN model is applied to various banchmark protein function datasets, and the results are compared with the state-of-the-art traditional methods.

5.1 Evaluation Metrics The evaluation metrics for multi-label classification include the multi-label precision (MPrecision), the multi-label recall (MRecall), the multi-label F-score (MF-score) (26) and the multi-label G-mean(MG-mean). Considering that the multi-label classification method in our work is a combination of multi-binary classifier for each class, MPrecision and MRecall are the averages for the precious and recall of all classes. They correspond to the micro average of the multi-label precision and the multi-label recall. Let us assume that D is a multi-label dataset, and the sample in |D| is (x_i, y_i) , i = 1...|D|, $y_i \in \{0, 1\}^m$ is the set of labels. Suppose $z_i \in \{0, 1\}^m$ be the set of labels predicted by HMC classifier for sample x_i . The definitions of the evaluation metrics are given by

$$MPrecision = \frac{1}{m} \sum_{i=1}^{|D|} \frac{|y_i \cap z_i|}{|z_i|} \dots (10)$$

$$MRecall = \frac{1}{m} \sum_{i=1}^{|D|} \frac{|y_i \cap z_i|}{|y_i|} \dots (11)$$

$$MF-Score = \frac{2 * MPrecision * MRecall}{MPrecision + MRecall} \dots (12)$$

$$MG-mean = \sqrt{MPrecision * MRecall} \dots (13)$$

In the multi-label classification, considering that the number of labels is imbalanced, so the result of MF-score and MG-means are more important.

5.2 Experiment Settings

5.2.1 Details of the deep CNN model Figure 5 shows the structure of deep CNN model used in the experiments, which has 6 convolutional layers and one flatten layer. Each layer of the CNN model is an 1-d convolution neural network with ReLU activation function. Smaller size filters are used in layers 1 to 3 to extract the micro information of amino acids and dipeptides. And larger size filters are used in layers 4 to 6 so as to learn the ranking rule of amino acids.

The size of filters and the number of channels are shown above each layer. For example, "Conv 3: 1*5*256" denotes that the convolution layer 3 has a filter of size 1*5, and 256 channels. The pooling layers are used in the convolution layers 2, 4, 6, which are max-pooling with pool-size: 1*2. The notation such as (N, 64, 1, 416) describes an input or output of a CNN layer, which denotes that there are N training samples, 64 channels, and each channel with the size of [1, 416]. Finally, the output of deep CNN model is a vector with the size of 1*512. Obviously, the proposed MHME CNN model has a more complicated network compared to the traditional HMC-LMLP model. We are expecting it to have much better performance.

5.2.2 Details on the supervised and unsupervised training algorithms As the deep learning framework, we use Chainer developed by Community, Preferred Networks, Inc (27). The training algorithm described in Section 4 are implemented by using Python programming language. In the experiments, each dataset is divided into three sets: training set, validation set and test set.

The training set is used to train the models. Since the hyper-parameters are not so sensitive for the proposed model training in our experiments, the default values suggested by the deep learning framework, Chainer are used. When supervised leaning of the body part and the multi-head part by using the algorithm described as *Step* 1 and *Step* 3 in Section 4, the initial learning rate used is 0.0075 and training process stops after 120 epochs where the performance is not improved anymore for the validation set. On the other hand, when the unsupervised learning of the autoencoders of the multi-end part by using the algorithm described as the *Step* 2, the initial learning rate used is 0.05 and the learning was stopped after 30 epochs where the performance for validation set stops improving.

The *Step* 2 and *Step* 3 of training process was repeated 2 times where the performance for validation set stopped improving. Finally, the trained models are evaluated on test set. In the experiments, efforts are made to prevent the training of models stuck at a local minimum. The proposed model and the conventional models are compared under a condition of being well-trained. All the results shown are an average of 10 trials, but the standard deviations are ignored since they are rather small in our experiments.

5.3 Comparison Results with the State-of-the-Art Methods Three datasets of protein sequences are used in this comparison, which are from the paper of DeepGO (21). Table 1 describes the details of datasets. For the classes,

Table 1: Datasets used in the paper of DeepGO

Datasets	Training	Validation	Test	Classes	Levels	Classes per level
DeepGO(BP)	20000	9000	10000	932	5	30/180/292/477/932
DeepGO(CC)	20000	8800	10000	439	4	18/126/260/439
DeepGO(MF)	20000	6300	5000	589	5	23/160/261/392/589

Table 2: Comparison results of the proposed MHME-CNN model with the HMC-LMLP, the DeepGO and the simple DCNN models

Datasets	Methods	Evaluation Metrics				
		MPrecision	MRecall	MF-score	MG-mean	
	DeepGO	0.3900	0.3400	0.3600	0.3700	
DeepGO(BP)	simple DCNN	0.4841	0.5001	0.4917	0.4920	
	HMC-LMLP	0.4847	0.5005	0.4921	0.4935	
	MHME-CNN	0.8028	0.5226	0.5315	0.6418	
	DeepGO	0.6600	0.6100	0.6300	0.6400	
DeepGO(CC)	simple DCNN	0.5028	0.5005	0.4959	0.5015	
	HMC-LMLP	0.4973	0.5068	0.5014	0.5017	
	MHME-CNN	0.9149	0.6122	0.6619	0.7456	
	DeepGO	0.6000	0.3800	0.4600	0.5100	
DeepGO(MF)	simple DCNN	0.5005	0.4998	0.4974	0.5001	
	HMC-LMLP	0.4941	0.5010	0.4975	0.4998	
	MHME-CNN	0.5451	0.5125	0.5093	0.5261	

GO functions are divided into three independent groups: the molecular functions (MF), that represent the activities performed by gene products; the cellular component (CC) functions, that describe the locations relative to cellular structures; and the biological process (BP) functions, that describe some 'biological programs' (1).

When applying the proposed MHME CNN model, according to the hierarchy of GO functions, the labels are divided into 5 levels for BP and MF functions, 4 levels for CC functions. Table 1 shows the numbers of classes per level for each dataset. The autoencoder used for each level is an one-layer MLP with a size of $\dim(I_n) \times \dim(x)$. The CNN model used is described in Figure 5.

After training, the MHME CNN model is evaluated on test data. Table 2 shows the experimental results on the three datasets. For comparison, the first row is the results of DeepGO model copied from the DeepGO paper (21). The second row shows the results of simple DCNN (deep CNN) model, whose structure is similar to

$$Z_{q+1} = f_{q+1} \left(\Omega_{q+1}^{(3)}, \phi \left(\Omega^{(2)}, x \right) \right), \tag{14}$$

a deep CNN model without considering the label hierarchical relationship; and the third row is the result of the HMC-LMLP method ⁽¹⁶⁾. The last row is the results of the proposed MHME CNN model.

From the results of Table 2, we can see that the proposed MHME CNN model performs better than the state-of-the-art models. The results of HMC-LMLP stand out when the dataset has more GO terms (BP, 932). Because it is the model trained GO terms level by level, whose result is influenced by GO terms more. The DeepGO or simple DCNN model does well when the dataset has less GO terms (CC, 439; MF, 589). As these two methods are concentrating on the feature extraction of protein sequences. The proposed MHME CNN model improves the performance by taking into account both hierarchy GO training and feature extraction.

5.4 Performance with Transfer Learning The proposed MHME CNN models trained on the three DeepGO datasets in the previous subsection are applied to other

Table 3: UniProt datasets (Number of samples)

Human	Mouse	Thaliana	Cerevisiae
3229	2743	488	488
7194	6055	2149	2149
3024	2756	776	776
	3229 7194	3229 2743 7194 6055	7194 6055 2149

datasets picked up from the UniProt ⁽²⁸⁾. Table 3 shows the details of these datasets. These 4 new datasets are not used in the training and only used for testing.

Table 4 shows the results. The proposed MHME CNN model is compared with the simple DCNN model and the HMC-LMLP model. From Table 4, we can see that the proposed MHME CNN model performs much better than other two methods. Moreover, the simple DCNN model outperforms over the HMC-LMLP model in transfer learning, which also proves that the deep learning based method has better performance in the case of transfer learning. Further studies are needed to show the potential applications of our proposed MHME CNN model based on transfer learning.

6. Conclusions

In this paper, we proposed a multi-head and multi-end (MHME) deep CNN model for GO annotation to improve the performance of complex hierarchical multi-label classification. The proposed MHME CNN model shares one deep CNN model with a set of linear classifiers, in which autoencoders are applied to fusing the features and reducing the dimension of feature vectors. The MHME CNN model is trained level by level to realize a set of local classifiers corresponding to the hierarchy labels, and one global classifier in order to capture the two-way relationship of labels. The simulation results show that our proposed MHME CNN model works well and has much better performance than the state of the art traditional methods. And our pre-built MHME CNN model also achieve a good performance on other GO annotation examples based on transfer learning.

In this paper, we only used the hierarchy GO dataset upto 5 levels. In our future work, we will apply the proposed model

Table 4: Results of the proposed MHME CNN model under transfer learning

Datasets	Functions	Methods	Evaluation Metrics			
			MPrecision	MRecall	MF-score	MG-mean
		simple DCNN	0.5127	0.5081	0.5085	0.5102
	BP	HMC-LMLP	0.4968	0.4989	0.4978	0.4979
		MHME-CNN	0.6967	0.6999	0.6979	0.6983
		simple DCNN	0.5339	0.5308	0.5276	0.5314
Human	CC	HMC-LMLP	0.4929	0.4977	0.4953	0.4957
		MHME-CNN	0.6592	0.6651	0.6613	0.6615
		simple DCNN	0.5249	0.5202	0.5208	0.5225
	MF	HMC-LMLP	0.4952	0.5000	0.4976	0.4977
		MHME-CNN	0.6565	0.6612	0.6587	0.6588
		simple DCNN	0.5119	0.5069	0.5080	0.5093
	BP	HMC-LMLP	0.4977	0.4994	0.4981	0.4985
		MHME-CNN	0.6873	0.6907	0.6887	0.6890
		simple DCNN	0.5273	0.5251	0.5218	0.5252
Mouse	CC	HMC-LMLP	0.4927	0.4977	0.4952	0.4951
		MHME-CNN	0.6601	0.6663	0.6623	0.6626
		simple DCNN	0.5201	0.5167	0.5165	0.5183
	MF	HMC-LMLP	0.4949	0.5000	0.4975	0.4974
		MHME-CNN	0.6452	0.6502	0.6476	0.6477
		simple DCNN	0.5273	0.5162	0.5207	0.5216
	BP	HMC-LMLP	0.4974	0.4989	0.4981	0.4982
		MHME-CNN	0.8793	0.8815	0.8802	0.8803
		simple DCNN	0.5509	0.5498	0.5473	0.5496
Thaliana	CC	HMC-LMLP	0.4940	0.4977	0.4958	0.4959
		MHME-CNN	0.7833	0.7881	0.7849	0.7851
		simple DCNN	0.5538	0.5479	0.5497	0.5507
	MF	HMC-LMLP	0.4958	0.5001	0.4979	0.4980
		MHME-CNN	0.8205	0.8242	0.8222	0.8023
		simple DCNN	0.5265	0.5199	0.5223	0.5231
	BP	HMC-LMLP	0.4973	0.4989	0.4980	0.4981
		MHME-CNN	0.8788	0.8809	0.8797	0.8798
		simple DCNN	0.5518	0.5499	0.5473	0.5498
Cerevisiae	CC	HMC-LMLP	0.4940	0.4976	0.4957	0.4956
		MHME-CNN	0.7822	0.7870	0.7838	0.7840
		simple DCNN	0.5539	0.5510	0.5514	0.5523
	MF	HMC-LMLP	0.4962	0.5000	0.4981	0.4981
		MHME-CNN	0.8187	0.8225	0.8105	0.8206

to GO datasets with more levels (29). Moreover, we will carry out a study on applying other deep learning models, especially transfer learning for protein function predictions.

References

- Gene Ontology Consortium, "The gene ontology (GO) database and informatics resource," *Nucleic acids research*, vol. 32, no. suppl_1, pp. D258–D261,
- (2) H. B. Borges and C. N. Julio, "Multi-label hierarchical classification using a competitive neural network for protein function prediction," in *International Joint Conference on Neural Networks (IJCNN'2012)*. IEEE, 2012, pp. 1–8.
- (3) S. Feng, P. Fu, and W. Zheng, "A hierarchical multi-label classification algorithm for gene function prediction," *Algorithms*, vol. 10, no. 4, 2017.
- (4) F. Wu, J. Zhang, and V. Honavar, "Learning classifiers using hierarchically structured class taxonomies," in *International Symposium on Abstraction, Re*formulation, and Approximation(AAAI'2005). Springer, 2005, pp. 313–320.
- (5) B. Chen and J. Hu, "Hierarchical multi-label classification based on oversampling and hierarchy constraint for gene function prediction," *IEEJ Trans.* on Electical and Electronic Engineering, vol. 7, no. 2, pp. 183–189, 2012.
- (6) X. Zhu and M. Bain, "B-cnn: branch convolutional neural network for hierarchical classification," arXiv preprint arXiv:1709.09890, 2017.
- (7) C. N. Silla and A. A. Freitas, "Novel top-down approaches for hierarchical classification and their application to automatic music genre classification," in 2009 IEEE International Conference on Systems, Man and Cybernetics. IEEE, 2009, pp. 3499–3504.
- (8) L. Zhang, S. K. Shah, and I. A. Kakadiaris, "Hierarchical multi-label classification using fully associative ensemble learning," *Pattern Recognition*, vol. 70, pp. 89–103, 2017.
- (9) R. Cerri and de Carvalho André Carlos PLF, "Hierarchical multilabel classification using top-down label combination and artificial neural networks," in

- 2010 Eleventh Brazilian Symposium on Neural Networks. IEEE, 2010, pp. 253–258
- (10) X.-L. Wang, H. Zhao, and B.-L. Lu, "Enhance top-down method with metaclassification for very large-scale hierarchical classification," in *Proceedings* of 5th International Joint Conference on Natural Language Processing, 2011, pp. 1089–1097.
- (11) A. Mahdi, J. Qin, and G. Crosby, "Deepfeat: a bottom-up and top-down saliency model based on deep features of convolutional neural nets," *IEEE Transactions on Cognitive and Developmental Systems*, 2019.
- (12) C. N. Silla Jr and A. A. Freitas, "A global-model naive bayes approach to the hierarchical prediction of protein functions," in *International Conference on Data Mining (ICDM'2009)*. IEEE, 2009, pp. 992–997.
- (13) C. N. Silla and A. F. Alex, "A survey of hierarchical classification across different application domains," *Data Mining and Knowledge Discovery*, vol. 22, no. 1-2, pp. 31–72, 2011.
- (14) R. Cerri, C. B. Rodrigo, and C. d. C. André, "Hierarchical multi-label classification using local neural networks," *Journal of Computer and System Sciences*, vol. 80, no. 1, pp. 39–56, 2014.
- (15) ——, "Hierarchical classification of gene ontology-based protein functions with neural networks," in *International Joint Conference on Neural Networks* (IJCNN'2015). IEEE, 2015, pp. 1–8.
- (16) R. Cerri, C. B. Rodrigo, C. d. C. André, and Y. Jin, "Reduction strategies for hierarchical multi-label classification in protein function prediction," *BMC bioinformatics*, vol. 17, no. 1, p. 373, 2016.
- (17) J. Wehrmann, R. Cerri, and R. Barros, "Hierarchical multi-label classification networks," in *International Conference on Machine Learning*, 2018, pp. 5225–5234.
- (18) R. Cao, C. Freitas, L. Chan, M. Sun, H. Jiang, and Z. Chen, "ProLanGO: Protein function prediction using neural machine translation based on a recurrent neural network," *Molecules*, vol. 22, no. 10, pp. 1732–1745, 2017.
- (19) J. Zeng, D. Li, Y. Wu, Q. Zou, and X. Liu, "An empirical study of features fusion techniques for protein-protein interaction prediction," *Current Bioinformatics*, vol. 11, no. 1, pp. 4–12, 2016.
- (20) L. Wei, P. Xing, J. Zeng, J. Chen, R. Su, and F. Guo, "Improved prediction

- of protein–protein interactions using novel negative samples, features, and an ensemble classifier," *Artificial Intelligence in Medicine*, vol. 83, pp. 67–74, 2017
- (21) M. Kulmanov, M. A. Khan, and R. Hoehndorf, "Deepgo: predicting protein functions from sequence and interactions using a deep ontology-aware classifier," *Bioinformatics*, vol. 34, no. 4, pp. 660–668, 2017.
- (22) I. Goodfellow, Y. Bengio, and A. Courville, *Deep learning*. MIT press, 2016.
- (23) A. Makhzani and B. J. Frey, "A winner-take-all method for training sparse convolutional autoencoders," NIPS Deep Learning Workshop, 2014.
- (24) —, "Winner-take-all autoencoders," Advances in neural information processing systems, pp. 2791–2799, 2015.
- (25) X. Pan, P. Rijnbeek, J. Yan, and H.-B. Shen, "Prediction of rna-protein sequence and structure binding preferences using deep convolutional and recurrent neural networks," *BMC genomics*, vol. 19, no. 1, p. 511, 2018.
- (26) G. Tsoumakas and K. Ioannis, "Multi-label classification: An overview"," International Journal of Data Warehousing and Mining (IJDWM), vol. 3, no. 3, pp. 1–13, 2007.
- (27) S. Tokui, K. Oono, S. Hido, and J. Clayton, "Chainer: a next-generation open source framework for deep learning," in *Proceedings of workshop on ma*chine learning systems (LearningSys) in the twenty-ninth annual conference on neural information processing systems (NIPS), vol. 5, 2015, pp. 1–6.
- (28) UniProt. [Online]. Available: https://www.uniprot.org/uniprot//
- (29) F. K. Nakano, M. Lietaert, and C. Vens, "Machine learning for discovering missing or wrong protein function annotations," *BMC bioinformatics*, vol. 20, no. 1, p. 485, 2019.

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