Data and text mining

DeepSec: a deep learning framework for secreted protein discovery in human body fluids

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Abstract

Motivation: Human proteins that are secreted into different body fluids from various cells and tissues can be promising disease indicators. Modern proteomics research empowered by both qualitative and quantitative profiling techniques has made great progress in protein discovery in various human fluids. However, due to the large number of proteins and diverse modifications present in the fluids, as well as the existing technical limits of major proteomics platforms (e.g. mass spectrometry), large discrepancies are often generated from different experimental studies. As a result, a comprehensive proteomics landscape across major human fluids are not well determined.

Results: To bridge this gap, we have developed a deep learning framework, named DeepSec, to identify secreted proteins in 12 types of human body fluids. DeepSec adopts an end-to-end sequence-based approach, where a Convolutional Neural Network is built to learn the abstract sequence features followed by a Bidirectional Gated Recurrent Unit with fully connected layer for protein classification. DeepSec has demonstrated promising performances with average area under the ROC curves of 0.85–0.94 on testing datasets in each type of fluids, which outperforms existing state-of-the-art methods available mostly on blood proteins. As an illustration of how to apply DeepSec in biomarker discovery research, we conducted a case study on kidney cancer by using genomics data from the cancer genome atlas and have identified 104 possible marker proteins.

Availability: DeepSec is available at https://bmbl.bmi.osumc.edu/deepsec/.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Human body fluids, such as blood, saliva and urine, are primary clinical specimens, which hold considerable promises in presenting molecular biomarkers for disease diagnosis and therapeutic monitoring (Anderson, 2010; Lathrop et al., 2003). Since the first study of serum globulin was reported in 1937 (Tiselius, 1937), several large-scale research efforts have been made to profile proteomes in various types of human body fluids, mostly in blood, using different proteomic technologies, such as two-dimensional gel electrophoresis (Margolis and Kenrick, 1969), mass spectrometry (Thomson, 1914) and liquid chromatography (Zhao and Lin, 2014). As a result, a large number of protein species were identified in different body fluids, which have been documented in a large volume of research papers and archived in several community-based proteomic databases such as Human Plasma Proteome Project (Legrain et al., 2011), Plasma Proteome Database (Nanjappa et al., 2014) and Human Plasma PeptideAtlas (Schwenk et al., 2017).

Due to the large complexity of protein content and post-modifications involved in different body fluids, protein identification using conventional proteomics techniques remains a challenging research topic in the past decade. To facilitate this research, several computational attempts have been made to predict secreted proteins in human body fluids, mostly based on machine learning methods, such as Support Vector Machine (SVM) (Cui et al., 2008; Hong et al., 2011; Sun et al., 2015; Wang et al., 2013, 2016b). These
models often used common protein features such as amino acid flexibility index, surface tension and solubility as input and predict secreted proteins associated with a specific body fluid. Although the performances of those methods were promising, featured-based models generally suffered from common limitations such as the blind manual collection of features that might be incomplete and biased and feature selection procedures that need manual intervention. To this end, automatic feature extraction using end-to-end model can dispense with the initial feature selection step and possibly improve the prediction performance.

Deep learning (DL) has been successfully applied in protein research to study new protein functions, structures and interactions (Jain et al., 2021). Among different DL models, Convolutional Neural Network (CNN) has been one of the most frequently used methods and has attained remarkable performances in several classification applications, especially when combined with Gated Recurrent Unit (GRU) (Wilaiprasitporn et al., 2020). GRU, as a new class of Recurrent Neural Networks (RNNs), can effectively solve the vanishing gradient problem by introducing memory cells and a gating mechanism for holding information from the prior inputs in a well-behaved way. Compared to Long Short-Term Memory, GRU has a relatively simple structure, which ensures reduced complexity and faster convergence. Recent successes with GRU on the basis of protein sequences include DeepSig (a model to detect signal peptides of proteins) (Savojardo et al., 2018) and DeepLoc (a model for predicting protein subcellular localization) (Armenteros et al., 2017). The high performances from both studies indicate that protein sequences must carry important characteristics related to protein sorting.

In this study, we present a new DL-framework, named DeepSec, to facilitate body-fluid secreted protein prediction. DeepSec adopts an end-to-end sequence-based approach and employs CNN as a feature extractor and Bidirectional Gated Recurrent Unit (BGRU) with fully connected dense layer as a classifier to predict secreted proteins. We have employed DeepSec on 12 different types of common human body fluids (one model for each body fluid), which include blood, saliva, urine, cerebrospinal fluid, seminal fluid, amniotic fluid, tear fluid, bronchoalveolar lavage fluid, milk, nipple aspirate fluid, pleural effusion and sputum. At last, we demonstrate possible applications of DeepSec in biomarker discovery by a kidney cancer case study.

2 Materials and methods

2.1 Datasets

The positive datasets of DeepSec were collected from our previous work (Huang et al., 2021). For negative dataset generation, we employed a similar method that is proposed by Cui et al. (2008), where the Pfam family annotation was used to select proteins that are potential non-body-fluid-secretory proteins. We chose negative samples from Pfam families (Pfam release 33.1) (Sara et al., 2018) which do not contain any proteins in the positive dataset. According to the size of the positive dataset varies in different fluid type, the selection of negative samples is carried out as follows. For a specific body fluid, if the count of positive-related Pfam families is greater than 30% of the total number of human-related Pfam families, all proteins in the remaining Pfam families were collected as the negative set. In contrast, if the count is <30%, we randomly chose one member from each remaining Pfam families to construct the negative data.

Using the above procedure, we have collected both positive and negative datasets for each of the 12 body fluids (Fig. 1), including 8203 and 2739 proteins in blood, 4072 and 3291 proteins in saliva, 8048 and 5136 proteins in urine, 6260 and 5787 proteins in cerebrospinal fluid, 5376 and 2624 proteins in seminal fluid, 3212 and 3722 proteins in amniotic fluid, 1490 and 4184 proteins in tear fluid, 1117 and 4436 proteins in bronchoalveolar lavage fluid, 2171 and 3887 proteins in milk, 2234 and 4009 proteins in nipple aspirate fluid, 1373 and 4328 proteins in pleural effusion, and 2341 and 3967 proteins in sputum.

![Fig. 1. The distribution of 12 types of body fluids that are analyzed in this study](https://example.com/figure1)

Training DL models generally benefits from datasets with a balanced size. To address the imbalance problem, a random undersampling process is adopted in this study, which is advantageous over the procedure of eliminating proteins from the over-sized larger dataset to match the size of smaller dataset that may lead to information loss. We randomly partitioned the larger dataset into smaller subsets with similar size of the smaller dataset. For a size ratio t between the positive and negative datasets, round(t) subsets are created randomly. Each generated new subset and the original smaller dataset are resampled together to generate independent datasets for model evaluation. At last, bagging algorithm is employed to calculate the overall performance of the model, i.e. the average performance across all sample model denoted as $P = \frac{1}{t} \sum_{i=1}^{t} P_i$ where $P_i$ ($1 \leq i \leq t$) refers to the performance based on one sample set.

Furthermore, the sample space of each body fluid is divided into training, validation and testing datasets according to the proportion of 60%, 20% and 20%, respectively.

2.2 Neural network model

DeepSec takes protein sequences as input and performs a binary classification in terms of secretion into a specific body fluid or not. Figure 2 summarizes the architecture of DeepSec, which comprises three basic components: input sequences, feature extraction and classification.

2.2.1 Input sequence generation

We first create a Position-Specific Score Matrix (PSSM) for each protein sequence to enable subsequent convolution operations. The protein sequence is embodied into a PSSM by position-specific iterative basic local alignment search (PSI-BLAST) (Altschul et al., 1997) against UniRef 90 (released in 2020_01) database with inclusion 0.001 and 3 iterations. For each protein with sequence length L, a PSSM of dimensionality $L \times 20$ is obtained. The columns of PSSM represent the presence of 20 amino acids in each position. We then transform the PSSM described in Wang et al. (2016a) by the Sigmoid function $1/(1 + \exp(-x))$ where $x$ represents a single entry of the PSSM.

Since variable sequence length [from tens to thousands of amino acids (aa) in this case] represents another challenge for building the prediction model, we decide to use a fixed size (1000aa) window to process protein sequences. It has been shown that N-terminus or C-terminus of the sequence carry the most useful signaling information. For instance, N-terminus modifications have a pivotal role in protein regulation and cellular signaling (Varland et al., 2015), and the identity of the C-terminal amino acids has a strong influence on protein expression levels (Weber et al., 2020). Therefore, if a protein’s length exceeds 1000, we will keep 500 aa from N-terminus and C-terminus, respectively, and remove the middle sequence. This method of fixed size window has achieved remarkable performances

![Fig. 2. The architecture of DeepSec](https://example.com/figure2)
in protein localization prediction (Armenteros et al., 2017). In the training dataset, 11.7% of the proteins are truncated using this rule. If a protein sequence is <1000aa, the missing part is filled with 0. As a result, the input representation of each protein is a matrix with size [sequence length (L = 1000) \times size of amino acid vocabulary (c = 20)].

### 2.2.2 Feature extraction

To reveal hidden knowledge from the observed PSSM, a CNN model is trained to extract a complex feature representation of the input sequence. Specifically, the input PSSM matrix is fed into CNN to learn the weight parameters of the convolution filters. The filters are set to calculate the feature map C. The convolution layer outputs the matrix inner product between the input matrix and the filters. The rectified linear unit (ReLU) was applied as the activation function to sparsify the output of the convolution layer:

\[
C_{ij} = \max(0, \sum_{d=-(n-1)/2}^{(n-1)/2} \sum_{e=-(c-1)/2}^{(c-1)/2} X_{i+d,e} \odot W_{d+(n-1)/2,c}^{ij} + b^{ij}) \quad (1)
\]

where \(X\) is the PSSM matrix, \(W^{ij}\) is the weight matrix of the convolution kernels, \(b^{ij}\) is the offset vector, \(C_{ij}\) is the results of feature extraction. Then, \(C_{ij}\) is used as input to feed into the next layer.

### 2.2.3 Classification

For memorizing the residue presence in the C-terminus and N-terminus, respectively, we use the forwards and backwards GRU to capture possible long-range dependencies between the sequence and the prediction. Bidirectional GRU sweeps from both C-terminus to N-terminus and N-terminus to C-terminus, and concatenates the outputs of individual directions before feeding them into the fully connected layer. The hidden states are updated recursively from the convolutional features and the previous value of the hidden states (Fig. 3).

The recurrent calculation at each sequence position \(t\) is denoted as:

\[
\begin{align*}
\bar{h}_t &= (\bar{h}_t, \bar{b}_t) \\
\bar{h}_{t} &= (1 - \xi_t) \odot \bar{h}_{t-1} + \xi_t \odot \bar{y}_t \\
\xi_t &= \sigma(W_c \bar{c}_t + U_c \bar{h}_{t-1} + b_c) \\
\bar{y}_t &= \tanh(W_y \bar{c}_t + U_y \bar{h}_{t-1} + b_y) \\
\bar{r}_t &= \sigma(W_r \bar{c}_t + U_r \bar{h}_{t-1} + b_r)
\end{align*}
\]

where \(\{c\}_{t-1}^L\) is the input features of \(c\) at position \(t\), \(\{b\}_{t-1}^L\) is the hidden states of BGRU, where \(\bar{h}_t\) is a vector of 2-time size of the number of hidden units in the individual direction GRU, \(W_c, W_r, W_y, U_c, U_r, U_y\) and \(U_r\) are the weight metrics and \(b_r, b_c\) and \(b_y\) are the bias. The calculation of \(b_t\) is similar to \(\bar{h}_t\). The tanh function is a non-linear activation function taking the form of \(\tanh(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}} - 1\).

Then, the subsequence classification is performed by a fully connected layer comprised hidden-layers and an output layer. The hidden layer computes a non-linear transformation, defined as follows:

\[
f = \max(0, b_t \cdot \mu + \nu) \quad (7)
\]

where \(\mu\) and \(\nu\) are the weight vector and bias respectively, \(\{b\}_{t-1}^L\) is the learned hidden states of BGRU, ReLU activation is used.

Finally, the output layer computes the probability distribution \(\tilde{y}\), defined as follow:

\[
\hat{y} = \sigma(f \cdot \gamma + \tau) \quad (8)
\]

where \(\gamma\) and \(\tau\) are the weight vector and the bias respectively, and the function \(\sigma\) is the softmax function, denoted as \(\sigma(z) = \frac{e^{z_i}}{\sum_{i=1}^{K} e^{z_i}}\), for \(i = 1, \ldots, K\).

A cross-entropy loss function is used to quantify how ‘far away’ our prediction is from the ground truth. Network parameters are optimized by minimizing the training errors. The backward pass uses the chain rule to back-propagate error signals and computes gradients with respect to all weights throughout the neural network. Given a training set \(\theta = (x_1, y_1), \ldots, (x_n, y_n)\), \(n\) is the number of samples and \(y\) is the true output targets, the cross-entropy loss function is defined as:

**Fig. 2.** The architecture of DeepSec which supports input as PSI-profiles based on protein sequences, feature extraction through CNN, classification based on BGRU with fully connected dense layer, and the outputs as the probability of being secreted protein.

**Fig. 3.** The forwards and backwards GRU capturing possible long-range dependencies between the input sequence and the predicted class.
2.3 Performance assessment

The prediction performance is measured based on both training and validation sets. Specifically, accuracy, Matthew’s correlation coefficient (MCC) and the Area under the ROC Curve (AUC) are applied. Considering the unbalanced cases in this study, we also include additional measures including precision, recall and F-measure. Note that higher values indicate better classification performance for all those measures.

Accuracy represents how many predictions of the classifier are in fact correct, defined as:

\[
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN}
\]  

where \(TP\) and \(TN\) are the true predictions in positives and negatives, respectively, and \(FP\) and \(FN\) are the false predictions in positives and negatives, respectively.

Recall (or sensitivity) shows how many positive examples are correctly identified by the classifier. In this case, this is the percentage of secreted proteins identified as such, defined as:

\[
\text{Recall} = \frac{TP}{TP + FN}
\]

Precision represents the proportion of the correctly predicted positive cases relative to all the predicted positive ones. In this case, this is the percentage of proteins identified as secreted proteins that actually secreted proteins, defined as:

\[
\text{Precision} = \frac{TP}{TP + FP}
\]

F-measure is the harmonic mean of precision and recall, and better reflects the performance of a classifier of unbalanced classes, defined as:

\[
F - \text{measure} = 2 \times \frac{\text{precision} \times \text{recall}}{\text{precision} + \text{recall}}
\]

MCC is a correlation coefficient between the observed and predicted binary classifications, defined as:

\[
\text{MCC} = \frac{(TP \times TN - FP \times FN)}{\sqrt{(TP + FN)(TF + FP)(TN + FP)(TN + FN)}}
\]

2.4 Comparison with other models based on protein features

Since SVM has been previously applied for secreted protein prediction, we first compare DeepSec with SVM models based on reported protein features. In addition, two other state-of-the-art models, including Decision Tree (DT) and Deep Neural Network (DNN), are benchmarked together with DeepSec. To do this, we use the same datasets to evaluate the performance to ensure a fair comparison. We examine features used in the previous studies and also add a few newly reported features, which can be grouped into four categories: (i) sequence properties; (ii) physicochemical properties; (iii) domains/motifs properties; and (iv) structural properties. In total, a total of 1610 protein features are collected (as detailed in Supplementary Table S1). Then, feature selection is performed in a similar workflow as documented in our previous publication (Huang et al., 2021), where \(F\)-test (\(P\) value \(\leq 0.005\)) and false discovery rate (FDR, \(q\) value \(\leq 0.05\)) are employed to rank the features. The top 50 features are selected into the final model for 12 body fluids, respectively (Supplementary Table S2). Feature normalization is performed using Z-score method. Finally, these models are evaluated in terms of accuracy, recall, precision, F-measure, MCC and AUC.

3 Results

3.1 Model performance of DeepSec in 12 types of body fluid

All classification models on 12 body fluids were built and evaluated using Pytorch 1.7.1, running python 3.8 and using Scikit-learn library version 0.23.2. In DeepSec, the input representation of each protein is a \(1000 \times 20\) matrix. Referring to Deeploc (Armenteros et al., 2017), our model has 50 filters at the convolution layer with three different sizes \(w = \{1, 5, 7\}\), which has led to a total of 150 filters and \(1000 \times 150\) feature maps. Next, GRU scanned the sequence using 32 hidden units in each individual direction, leading to a total of \(1000 \times 64\) outputs. The fully connected layer comprised a single hidden-layer with 16 units and an output layer with two units. The parameters were optimized by Adam optimizer with learning rate as 0.0001 and a dropout probability as 0.1 prior to fully connected layer. We chose 0.5 as the prediction threshold, which means that a probability \(\geq 0.5\) indicates a positive class associated with secretion into a specific body fluid. Finally, the performances on 12 body fluids were evaluated based on testing dataset (Table 1) and all datasets (Table 2), respectively.

We have applied DeepSec to screen against all human proteins (20394 unique proteins) in the UniProtKB/Swiss-Prot database (UniProt release 2020.06) in each of body fluids. As shown in Figure 4, DeepSec predicted 12364 proteins (60.6% of the 20394) as blood-secreted proteins, and 9491 (46.5%), 6877 (33.7%) and 6713 (32.9%) proteins to be secreted into urine, cerebrospinal fluid and saliva, respectively.

3.2 DeepSec outperforms feature-based classifiers

We compared the performances between the DeepSec model and aforementioned feature-based models based on average AUC on testing datasets. Here the feature-based DT model, SVM model and DNN model are termed DTf, SVMf and DNNf, respectively. As shown in Figure 5, for all 12 fluid types, DeepSec reported the best performances with average AUC ranged in 0.85-0.94. Especially, DeepSec was 4-17% higher than other models.

3.3 A close-look at the blood secreted protein prediction

In the blood protein case, the positive dataset consists of 8203 proteins while 2739 human proteins were generated in negative dataset. Considering the count ratio \(t\) between positive and negative dataset is round8203/2739 = 3, we repeated 3 times the random sampling processing and obtained 3 positive subsets. In the end, three datasets, each containing 2734 positive samples and 2739 negative samples, were used to train and evaluate DeepSec classifier.

To select the most appropriate model, we compared different model architectures including CNN, BGRU and DeepSec. Each model architecture was evaluated based on testing dataset and all datasets in terms of accuracy, recall, precision, F-measure, MCC and AUC (Table 3). The average ROCs are plotted in Figure 6. Note that DeepSec classifier achieved the highest overall performance on both testing dataset (average AUC: 0.94) and all datasets (average AUC: 0.94), respectively. In the meantime, it also attained the highest average values of accuracy (0.870/0.87), F-measure (0.91/0.87) and MCC (0.69/0.74) on testing dataset and all datasets.

The results showed that the DeepSec model is more effective in capturing sequence information for predicting secreted proteins and reveals relationships between amino acid sequences and secretion status. This is not surprising as BGRU is capable of learning long-term dependencies between sequence and secreted proteins, which may contribute to an improved performance by updating the weight of each hidden state.

4 An application case study on kidney cancer biomarker discovery

To illustrate possible applications of DeepSec, we explored potential kidney cancer biomarkers in blood by using public genomics data and DeepSec prediction. To do that, we first collected gene
expression profiles of 72 paired kidney cancer tissues and adjacent control tissues samples from kidney renal clear cell carcinoma (KIRC) via the cancer genome atlas (TCGA) Data Portal. Each gene-expression dataset covered 19,804 human genes measured using RNA-seq. Note the paired samples helps reduce the impact of individual variability in differential expression analysis. Two-tailed t-test (Liang et al., 2019) was performed to identify genes that have significant differential expression between kidney

| Body fluids                        | Accuracy | Recall  | Precision | F-measure | MCC       | AUC       |
|------------------------------------|----------|---------|-----------|-----------|-----------|-----------|
| Blood                             | 0.871139 | 0.87212 | 0.868200  | 0.910294  | 0.691481  | 0.940572  |
| Saliva                            | 0.824650 | 0.81052 | 0.835108  | 0.797251  | 0.643117  | 0.898319  |
| Urine                             | 0.845857 | 0.80501 | 0.88317   | 0.834207  | 0.692125  | 0.918341  |
| Cerebrospinal fluid               | 0.835470 | 0.66788 | 0.931376  | 0.747156  | 0.637556  | 0.900955  |
| Seminal fluid                     | 0.821891 | 0.83407 | 0.810378  | 0.819841  | 0.644204  | 0.894597  |
| Amniotic fluid                    | 0.828476 | 0.74779 | 0.88995   | 0.790455  | 0.649091  | 0.905148  |
| Tear fluid                        | 0.830080 | 0.57252 | 0.926077  | 0.646611  | 0.545096  | 0.856645  |
| Bronchoalveolar lavage fluid      | 0.859257 | 0.43204 | 0.966373  | 0.551724  | 0.502859  | 0.857458  |
| Milk                              | 0.811124 | 0.67701 | 0.85655   | 0.719173  | 0.580265  | 0.871808  |
| Nipple aspirate fluid             | 0.816795 | 0.56500 | 0.956554  | 0.687654  | 0.594120  | 0.887423  |
| Pleural effusion                  | 0.837814 | 0.57163 | 0.922045  | 0.62887   | 0.530781  | 0.848811  |
| Sputum                            | 0.823529 | 0.82209 | 0.824375  | 0.775122  | 0.633483  | 0.891406  |

Note: The highest scores are in bold, and the lowest scores are underlined.
cancer and control samples. The test statistic $t_k$ for the $k$th feature between case and control is given by

$$t_k = \frac{d_k}{s_k}$$

where $d_k$ and $s_k$ is the mean difference and the standard error of the $k$th feature across $n$ paired samples, denoted as

$$d_k = \frac{1}{n} \sum_{i=1}^{n} d_{ik}$$

$$s_k = \sqrt{\frac{\sum_{i=1}^{n} (d_{ik} - d_k)^2}{(n-1)}}$$

where $d_{ik}$ is the differential between the paired case and control for the $k$th feature of the $i$th sample, given as

$$d_{ik} = X_{1i,k} - X_{2i,k}$$

logFC > 2 and adjusted $P < 0.05$ were used to identified differentially expressed genes between cancer group and normal control. Overall, 138 and 261 genes (Supplementary Tables S3 and S4) were found to be up- and down-regulated in kidney cancer versus control tissues, respectively, hence making them as potential maker candidates in kidney cancer.

Based on all the differentially expressed genes (138 up-regulated and 261 down-regulated) (Fig. 7) in kidney cancer versus control, we applied DeepSec and further inferred that 261 of these biomarkers may be secreted into blood. Since 157 proteins have been included in our positive dataset, the remaining 104 proteins are considered novel marker proteins identified by this prediction model. Despite of the discordance between gene expression and protein

**Table 3. Prediction performance of various model architectures evaluated based on testing dataset and all datasets**

| Measures       | CNN          | BGRU         | DeepSec      | CNN          | BGRU         | DeepSec      |
|----------------|--------------|--------------|--------------|--------------|--------------|--------------|
| Accuracy       | 0.806068     | 0.851455     | 0.871339     | 0.825292     | 0.811404     | 0.871345     |
| Recall         | 0.782884     | 0.911252     | 0.872120     | 0.777778     | 0.931287     | 0.887427     |
| Precision      | 0.875302     | 0.687112     | 0.868200     | 0.872807     | 0.691520     | 0.853263     |
| F-measure      | 0.858212     | 0.904143     | 0.910294     | 0.816577     | 0.831593     | 0.873381     |
| MCC            | 0.587026     | 0.608209     | 0.691481     | 0.653542     | 0.641525     | 0.743075     |
| AUC            | 0.912392     | 0.897955     | 0.940372     | 0.906867     | 0.903960     | 0.941104     |

Note: The highest scores are in bold.
abundance, proteins that show significantly elevated gene expression in kidney cancer tissues versus control can be more promising marker candidates compared to those repressed ones. The detailed prediction results about up- and down-regulated protein markers of kidney cancer in blood are listed in Supplementary Table S5.

5 Discussion

DeepSec represents the first generalized computational model that can predict secreted proteins in multiple human body fluids. Specifically, this end-to-end model is built based on sequence features extracted via CNN followed by bidirectional GRU. Compared to other models based on proteins features, DeepSec has demonstrated improved prediction accuracy and better generalizability (Fig. 5), which indicates the advantage of sequence-based method as compared to feature-based algorithms. In the blood protein case, the model shows reasonably high AUC (0.94/0.94) and high F-measure (0.91/0.87) on testing dataset and all datasets, as compared to other model architectures (Table 3), which also suggests that DeepSec has good representation of the relevant proteins across the whole protein space.

However, there might be some concerns about the limitations of DeepSec. First, due to the lack of clear knowledge about non-body-fluid secretory proteins, this study generates negative dataset based on Pfam family information. It means that the negative datasets may not adequately include the whole space of the non-body-fluid secretory proteins. Indeed, with larger number of positive instances being identified by wet lab experiments, this model can be better tuned, as demonstrated in Cui et al. (2008). Second, since the protein sequences are compressed into a fixed size vector, there is a risk of information loss. Note that 11.7% (1990 proteins) of human proteins have been influenced by the truncation rule. However, when analyzing the physical properties of those proteins, we understand that the probability of the secreted proteins having long anomic acid sequences is very low, which implies a minimal negative impact on the DeepSec’s performance if there is any. Last, since we addressed the imbalance problem by sampling multiple smaller subsets and recalculates the probabilities based on the same small dataset during the training process, there is a certain likelihood that it introduces risks of overfitting while yielding significantly improved performance. However, when evaluating the performance of DeepSec on all datasets (Table 3), it appears confidently that DeepSec doesn’t overly fit the data.

6 Conclusion

In summary, we propose a DL method, DeepSec, to predict secreted proteins in 12 kinds of human body fluids based on protein sequences. To the best of our knowledge, DeepSec is the first system that fully capture the sequence features related to protein secretion automatically using CNN with BGRU architecture. The BGRU network is able to capture possible long-range dependencies between sequence and secreted status of proteins, which contributes to the improved performance. DeepSec is able to predict the secreted protein with higher accuracy than existing state-of-the-art methods. Moreover, DeepSec is useful for discovering novel candidates of blood biomarkers in kidney cancer that have been experimentally verified. Our future effort will focus on including more types of human body fluids into the system and improve the performance toward biomarker discovery in complex human diseases and physiological phenotypes.

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