Learning distributed representations of RNA and protein sequences and its application for predicting lncRNA-protein interactions

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Abstract
The long noncoding RNAs (lncRNAs) are ubiquitous in organisms and play crucial role in a variety of biological processes and complex diseases. Emerging evidences suggest that lncRNAs interact with corresponding proteins to perform their regulatory functions. Therefore, identifying interacting lncRNA-protein pairs is the first step in understanding the function and mechanism of lncRNA. Since it is time-consuming and expensive to determine lncRNA-protein interactions by high-throughput experiments, more robust and accurate computational methods need to be developed. In this study, we developed a new sequence distributed representation learning based method for potential lncRNA-Protein Interactions Prediction, named LPI-Pred, which is inspired by the similarity between natural language and biological sequences. More specifically, lncRNA and protein sequences were divided into k-mer segmentation, which can be regarded as “word” in natural language processing. Then, we trained out the RNA2vec and Pro2vec model using word2vec and human genome-wide lncRNA and protein sequences to mine distribution representation of RNA and protein. Then, the dimension of complex features is reduced by using feature selection based on Gini information impurity measure. Finally, these discriminative features are used to train a Random Forest classifier to predict lncRNA-protein interactions. Five-fold cross-validation was adopted to evaluate the performance of LPI-Pred on three benchmark datasets, including RPI369, RPI488 and RPI2241. The results demonstrate that LPI-Pred can be a useful tool to provide reliable guidance for biological research.

1. Introduction
The emerging recognition of RNA is that any transcripts, regardless of protein coding potential, can have intrinsic functions [1]. One kind of this transcripts that are no less than 200 nucleotides, known as long non-coding RNA (lncRNA). Existing studies demonstrate that only less than 2% of the human genome can be translated into proteins, whereas more than 80% of it has biochemical functions [2,3]. Furthermore, more than 70% of ncRNA are long ncRNA [4], which means there is massive of precious information lncRNAs contained awaiting our effective mining. The lncRNA often act through functions by binding to partner proteins, and play critical roles in gene regulation, splicing, translation, chromatin modification and poly-adenylation [5–8]. Moreover, emerging evidences have revealed that various complex diseases have strong correlation with lncRNAs, such as Alzheimer [9], lung cancer [10] and cardiovascular diseases [11]. Therefore, the basis for understanding the functions of lncRNA is to identify lncRNA-protein interactions. It's inefficient to examine a large number of under-researched lncRNAs and proteins though wet experiments.

Due to the time-consuming and laborious of high throughput experiments, such as CLIP-seq, RIP-seq and fRIP-seq [12], several computational lncRNA-protein interaction prediction methods have been put forward in recent years, which can be used as guide tools for biological experiments. These methods can be divided into two categories. The first kind of methods mainly based on sequence information, structural information, evolutionary knowledge or physicochemical properties to exploit discriminative features of lncRNA and protein. For instance, Muppirala et al. proposed RPSSeq, which adopted k-mer composition to encode RNA and protein sequences and trained support vector machine (SVM) and Random Forest (RF) model to identify interactions [13]. Suresh et al. used sequence information and structure information to build a SVM predictor to predict novel protein-RNA interactions, named PRI-Pred [14]. Bellucci et al. developed catPA-
PID by using the physicochemical properties of nucleotide and polypeptide, include secondary structure, Van der Waals propensities and hydrogen bonding, to evaluate the interaction propensities, and they further applied this model to predicted protein interactions in the Xist network [15,16]. Lu et al. scored RNA-protein pair by using matrix multiplication and Fisher’s linear discriminant. More recently, Yi et al. presented a deep learning framework RPI-SAN, using stacked autoencoder to extract high-level hidden feature from sequence, then they trained RF classifier and ensemble strategy to robustly and accurately predict ncRNA-protein interactions [17]. These methods suggested that the sequence carried enough information for prediction tasks.

There is another category of methods in this domain, which considered the known interactions between IncRNA and protein. Yun et al. considered the relatedness of heterogeneous objects path-constrained, introduced a method using HeteSim measure to compute the relatedness score, called PLPHS [18]. Zhang et al. using graph regularized nonnegative matrix factorization to discover unknown interacted pairs based on the hypothesis that similar lncRNAs (proteins) have similar corresponding proteins (lncRNAs) [19]. Shen et al. proposed LPI-KTASLP to infer IncRNA-protein interactions with kernel target alignment and semi-supervised link prediction model using multivariate information [20]. Zhang et al. combined multiple sequence-based features and IncRNA-IncRNA similarities and protein-protein similarities, which is calculated by using RNA sequences and protein sequences and known IncRNA-protein interactions [21]. But these kind methods have limitations when predicting new samples, especially those never appeared in the similarity matrices.

This paper aims to develop a new sequence distributed representation learning based method for novel IncRNA-Protein Interactions Prediction, named LPI-Pred, which is inspired by the similarity between biological sequences and natural languages [22]. More specifically, IncRNA and protein sequences were divided into k-mer segmentation, which can be regard as “word” in natural language processing. Furthermore, we trained the RNA2vec and Pro2vec model using skip-gram word embedding model and Human genome-wide IncRNA and protein sequences for IncRNA and protein, respectively. The aforementioned train sequences data are provided by the GENCODE project (release v29) [23]. And then, we measured the importance of features via Gini information impurity, and select top-50 feature as final discriminative features. Finally, these features are used to train RF predictor. We evaluated our model on three benchmark datasets under five-fold cross-validation, including RNA-protein interaction datasets, RPI369 and RPI1807, and IncRNA-protein interaction dataset, RPI488, using six widely used evaluation indicators in machine learning field. And we compared our model with other state-of-the-art models such as RPS2Seq [13], IncPro [24], and RPI-SAN [17]. The rigorous experimental results prove the validity and reliability of our method.

2. Materials and methodology

2.1. Datasets exploration

In practice, three benchmark datasets, including RPI369 [13], RPI1807 [14] and RPI488 [25] were selected to execute our evaluation. The first two are RNA-protein interactions datasets, while the third is IncRNA-protein interactions dataset. The RPI369 dataset is a non-redundant data set, which is generated from RPIDB [26], and only have non-ribosomal complexes (e.g., mRNA, miRNA, tRNA). The dataset RPI369 contains 332 RNA sequences, 338 protein sequences and 369 positive interaction pairs. In the same work, the authors also constructed another dataset RPI2241, which is larger than RPI369 but is strongly biased to ribosomal RNA-protein interactions. That’s why we’re not inclined to adopt it. The RPI1807 also is a non-redundant data set of RNA-protein interactions complexes, generated by parsing the RPIDB and Nucleic Acid Database (NDB) [24]. There are 1078 RNA sequences and 1807 protein sequences in RPI1807, consisting 1807 pairs positive samples and 1436 pairs negative samples. The RPI488 is a IncRNA-protein interactions dataset, contains 245 negative IncRNA-protein pairs, 243 interacted IncRNA-protein pairs. The number of IncRNA and protein in this dataset are 25 and 247, respectively. The details of these three benchmark datasets are listed in Table 1 as below:

2.2. k-mer segmentation

In this section, we will introduce the feature representation scheme used in this study, which is aims to fully exploit the hidden high-level feature from the sequence information. For a given IncRNA or protein sequence, k-mer composition is used to split them into subsequences, which can be considered as “word” in the fellow step. Scan a sequence from beginning to the end, one nucleic acid once time. For a given sequence of length L, we will obtain L – k + 1 k-mers, and the count of possible k-mer are 4^k for RNA (A, C, G, U) and 20^k for protein (Ala, Gly, Val, Ile, Leu, Phe, Pro, Tyr, Met, Thr, Ser, His, Asn, Glu, Lys, Arg, Lys, Asp, Glu, Cys), different from common usage, we do not use the 7-letter reduce alphabet, which reduced 20 amino acids into 7 groups based on their similarity of dipole moments and side chain volume. We set the k to 4 for IncRNA and set k to 3 for protein, which are two commonly accepted empirical parameters [13,17,25,27]. The process of splitting nucleic acids sequence and amino acids sequences into k-mers shown in Fig. 1.

2.3. Distribution representation of IncRNA and protein sequences

And then, we using the genome-wide human IncRNA and protein sequences to train a word embedding model, named RNA2vec and pro2vec, respectively. The training data provided by the GENCODE project and their goal of this project is to identify and classify all gene features in the human and mouse genomes with high accuracy based on biological evidence, and to release these annotations [23,28]. We use the skip-gram [29,30] word representation model to learn distribution representation of RNA and protein sequences. In nature, the model is a neural network with projection layer for learning word representation. The structure of skip-gram is shown in Fig. 2 below.

For a given sequence (w_1, w_2, . . . , w_{k-1}), the goal of training model is to maximize the mean log probability:

\[
\max 1 - \frac{1}{N} \sum_{n=1}^{N} \sum_{1 \leq c \leq m \leq c} \log P(w_{n+m} | w_n)
\]

(1)

c stands for the distance to the central word; the log probability distribution can be defined as follow:

\[
\log P(w_0 | w) = \log \frac{e^{v_0^T v_w}}{\sum_{w} e^{v_w^T v_w}}
\]

(2)

where the v_0 and v_w are the input and output vector of word w, respectively. W is the size of training IncRNA or protein training lexicon.

In natural language processing, the word embedding model has achieved great success [31,32], it has also made progress in computational biology [33–35]. In this work, we regard each k-mer as a word and a sequence as a sentence, and then learning the distribution representation by using skip-gram word2vec model. The procedure for training RNA2vec and pro2vec is shown as Fig. 3.
The parameters of the model are \textit{min\_count} = 1, \textit{size} = 300, \textit{window} = 5, \textit{iter} = 10, \textit{batch\_words} = 100. Where the \textit{size} represents the dimensions of output word vector, and \textit{window} stands for maximum distance between the current and predicted word within a sentence, \textit{iter} is the count of iterations (epochs) over the corpus, \textit{batch\_words} is the target size (in words) for batches of examples passed to worker threads. When the \textit{min\_count} (means minimum word frequency) is set too high, the model only counts high-frequency words, which is not conducive to learning discriminative word vectors from sequence representation. Other parameters are default. Inspired by the additivity of word embedding [30], we represented a given sequence by summing all its \textit{k}-mer word embed-

### Table 1

| Datasets     | # of RNAs | # of proteins | Positive samples | Negative samples | References |
|--------------|-----------|---------------|------------------|------------------|-----------|
| RPI369       | 332       | 338           | 369              | 369              | [13]      |
| RPI1807      | 1078      | 1807          | 1807             | 1807             | [14]      |
| RPI488       | 25        | 247           | 243              | 245              | [25]      |

Fig. 1. Procedure of splitting RNA nucleotides and protein amino acids sequences into smaller \textit{k}-mers.

Fig. 2. The skip-gram word embedding model. Lnc2vec and pro2vec model were trained by using this model and genome-wide human lncRNA and protein sequences. Skip-gram is trained by predicting words surrounding the central word, after training, the weights matrix \textit{W} of the hidden layer is obtained, that is word vectors.

The parameters of the model are \textit{min\_count} = 1, \textit{size} = 300, \textit{window} = 5, \textit{iter} = 10, \textit{batch\_words} = 100. Where the \textit{size} represents the dimensions of output word vector, and \textit{window} stands for maximum distance between the current and predicted word within a sentence, \textit{iter} is the count of iterations (epochs) over the corpus, \textit{batch\_words} is the target size (in words) for batches of examples passed to worker threads. When the \textit{min\_count} (means minimum word frequency) is set too high, the model only counts high-frequency words, which is not conducive to learning discriminative word vectors from sequence representation. Other parameters are default. Inspired by the additivity of word embedding [30], we represented a given sequence by summing all its \textit{k}-mer word embed-

![Diagram](image-url)
2.4. Gini information impurity-based feature selection

A data set often has hundreds of previous features. How to choose the features that have the greatest impact on the results, so as to reduce the number of features when building the model. There are many such methods, for instance, principal component analysis, Lasso [36,37], mRMR [38] and so on. However, here we are going to introduce the use of Random Forest to feature screening based on Gini information impurity.

Assuming that there are \( m \) features \( f_1, f_2, \ldots, f_m \), we can calculate the Variable Importance Measures \( \text{VIM} \) for each feature \( f_i \), that is, the average change of node splitting impurity in all RF decision trees by \( f_i \) feature. The Gini index \( GI \) can be defined as:

\[
GI_i = \sum_{k=1}^{K} \sum_{l=1}^{k} p_k p_l^2 = \sum_{k=1}^{K} p_k^2
\]

where the \( K \) means there are \( k \) categories, and \( p_k \) indicates the proportion of categories \( k \) in \( i_{th} \) node. The \( \text{VIM} \) of feature \( f_i \) in \( i_{th} \) node can be computed from the variation of \( GI \) before and after branching of \( j_{th} \) node:

\[
\text{VIM}^{\text{Gini}} = GI_i - GI_j - GI_l
\]

Among them, \( GI_r \) and \( GI_l \) respectively represent the \( GI \) of the right and left nodes after branching. Suppose there are \( N \) decision trees, so:

\[
\text{VIM}^{\text{Gini}}_j = \sum_{n=1}^{N} \text{VIM}^{\text{Gini}}_n
\]

Finally, all the obtained importance scores can be normalized by:

\[
\text{VIM}_i = \frac{\text{VIM}_i}{\sum_{j=1}^{N} \text{VIM}_j}
\]

Here, we selected the most important top-50 features as final feature.

2.5. Training an LPI-Pred model

The selected top-50 feature would be used to train an LPI-Pred model for predicting potential IncRNA-protein interactions on test data set. In summary, the procedure for training an LPI-Pred is shown in Fig. 4:

- Using human genome-wide IncRNA and protein sequences as corpus, segment them into k-mers as the words.
- Using word2vec model to train out RNA2vec and pro2vec for IncRNA and protein sequence distribution representation.
- Obtaining the word embedding of the protein and ncRNA sequences in the benchmark RNA-protein interaction datasets.
- Select top-50 features based on feature importance to train Random Forest predictor.

2.6. Performance evaluation metrics

In this study, we proposed a novel IncRNA-protein interactions prediction model LPI-Pred, based on sequence distributed representation learning and Gini information impurity measure. The common metrics and five-fold cross-validation are used to evaluate the performance of LPI-Pred. Divided all data into five equal sub-set. For each training, one-fold set data is taken as test data, the rest four-fold are taken as training data. Take the mean performance metrics of five training as final performance. There is no overlap between train data and test data, and this is unbiased comparison. The metrics used in performance evaluation including accuracy (Acc), Sensitivity (Sens), Specificity (Spec), Precision (Pre) and Matthews Correlation Coefficient (MCC). Certainly, and the area under the curve (AUC) of the Receiver Operating Characteristic (ROC) curve are also adopted to evaluate the performance. These metrics can be defined as:

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

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

\[
\text{Specificity} = \frac{TN}{TN + FP}
\]

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

where \(TN, TP\) indicates the correctly predicted negative and positive samples number, \(FN, FP\) represents the false wrongly predicted negative and positive samples number.

3. Results and discussion

In this study, we proposed a novel lncRNA-protein interactions prediction model LPI-Pred, based on sequence distributed representation learning and Gini information impurity measure. In this section, we designed the following experiments to verify the performance of the model. First, we compared the effects of different sequence coding schemes on lncRNA-protein interaction dataset, and the effect of feature selection. Second, we did a performance comparison with different individual predictors. And then, we verify LPI-Pred’s ability to predict lncRNA-protein interactions and compared with other state-of-the-art methods. Finally, we apply our model to lncRNA-protein interactions network construction.

3.1. Comparison between different sequences encoding strategies

We applied a new RNA and protein sequences encoding method in this work, using skip-gram distribution representation model. In order to verify the effectiveness of this sequence numerical coding scheme, we first compare it with the widely used \(k\)-mer frequency on three benchmark datasets. The comparison results are shown in Table 2.

In all three gold standard datasets, the selected word embedding feature, obtained through RNA2vec and pro2vec model, have improved performance compared to \(k\)-mer method. This can prove that distribution representation word vector is effectiveness for biological sequences encoding, for RNA and protein. It can achieve and even exceed the performance of \(k\)-mer, which is very widely used in biological sequence representation. The comparison between LPI-Pred (using RNA2vec and pro2vec with feature selection) and LPI-Pred without feature selection demonstrate the necessity of feature selection.

3.2. Comparison with individual predictors

To verify the effect of RF classifier separately, we compared RF and other machine learning models including SVM (with RBF kernel), Logistic Regression (LR), under same set of features and the same experimental conditions. These models were trained with default parameters. The results are shown in Table 3:

Several Random Forest-based methods have achieved remarkable performance on many issues in the field of computational biology. We trained LPI-Pred based on random forest classifier.
As shown in the comparison results in the above table, LPI-Pred outperformed all other classifiers using same feature set and under same experimental conditions.

3.3. Evaluation of LPI-Pred’s capability to predict lncRNA-protein interactions

Furthermore, we compared our model with other state-of-the-art methods including RPISeq [13], lncPro [24], and RPI-SAN [17] to evaluate the predictive ability to lncRNA-protein interactions of LPI-Pred. The RPISeq and lncPro use only sequence information, which is similar to LPI-Pred. More recently, the RPI-SAN use deep learning model, based on sequence information and evolutionary information to predict novel ncRNA-protein interactions. We follow same performance evaluation measurements. The comparison details are shown as below Table 4.

**Table 4**

Comparing five-fold cross-validation performance of LPI-Pred and other state-of-the-art methods on three gold standard datasets.

| Datasets | Methods   | Acc (%) | Sens (%) | Spec (%) | Pre (%) | MCC (%) | AUC     |
|----------|-----------|---------|----------|----------|---------|---------|---------|
| RPI369   | RPISeq    | 70.4    | 70.5     | 70.2     | 70.7    | 40.9    | 0.767   |
|          | IncPro    | 70.4    | 70.8     | 69.6     | 71.3    | 40.9    | 0.740   |
|          | LPI-Pred  | 73.06   | 75.32    | 71.14    | 72.64   | 46.67   | 0.802   |
| RPI1807  | RPISeq    | 97.3    | 96.8     | 98.4     | 96.0    | 94.6    | 0.996   |
|          | IncPro    | 96.9    | 96.5     | 98.1     | 95.3    | 93.8    | 0.994   |
|          | RPI-SAN   | 96.1    | 93.6     | 99.9     | 91.4    | 92.4    | 0.999   |
|          | LPI-Pred  | 97.10   | 97.89    | 96.14    | 96.91   | 94.13   | 0.994   |
| RPI488   | RPISeq    | 88.0    | 92.6     | 82.2     | 93.2    | 76.2    | 0.903   |
|          | IncPro    | 87.0    | 90.0     | 82.7     | 91.0    | 74.0    | 0.901   |
|          | RPI-SAN   | 89.7    | 94.3     | 83.7     | 95.2    | 79.3    | 0.920   |
|          | LPI-Pred  | 89.92   | 82.75    | 96.72    | 96.32   | 80.59   | 0.911   |

The boldface indicates this measure performance is the best among the compared methods for individual dataset.

As shown in the comparison results in the above table, LPI-Pred outperformed all other classifiers using same feature set and under same experimental conditions.

**Table 2**

Comparing the five-fold cross-validation performance of k-mer and word embedding with and without feature selection on three gold standard datasets.

| Datasets | feature | Acc (%) | Sens (%) | Spec (%) | Pre (%) | MCC (%) |
|----------|---------|---------|----------|----------|---------|---------|
| RPI369   | k-mer   | 68.71   | 67.29    | 70.30    | 69.88   | 37.74   |
|          | embedding without feature selection | 71.97 | 70.27 | 73.76 | 73.19 | 44.24   |
|          | embedding with feature selection   | 73.06 | 75.32 | 71.14 | 72.64 | 46.67   |
| RPI488   | k-mer   | 89.29   | 83.17    | 95.17    | 94.33   | 79.09   |
|          | embedding without feature selection | 87.64 | 83.17 | 91.93 | 90.82 | 75.52   |
|          | embedding with feature selection   | 89.92 | 82.75 | 96.72 | 96.32 | 80.59   |
| RPI1807  | k-mer   | 96.88   | 98.44    | 94.96    | 96.04   | 93.72   |
|          | embedding without feature selection | 96.73 | 97.90 | 95.28 | 96.28 | 93.37   |
|          | embedding with feature selection   | 97.10 | 97.89 | 96.14 | 96.91 | 94.13   |

The boldface indicates this measure performance is the best among the compared methods for individual dataset.

4. Conclusion

The lncRNA-protein interactions play numerous roles in life activities, cellular function and disease. The first step in studying its function and mechanism is to identify interacting lncRNA-protein pairs. In this study, we present a novel lncRNA-protein interaction prediction model named LPI-Pred. First, we trained distribution representation model, RNA2vec and pro2vec, by using skip-gram word embedding model and human genome-wide lncRNA and protein sequences. Then, we convert the lncRNA and protein sequences into a fixed-length vector using this word embedding model. Then, we perform feature selection to further enhance the expressiveness of features and reduce the complexity of model training.
protein sequence into word vector using the model trained above. The Gini impurity-based feature selection is used to obtain discriminative features. Then we training LPI-Pred to predict lncRNA-protein interactions. We compared the performance of different feature representations and predictors, and we also compared LPI-pred with other state-of-the-art methods. The rigorous evaluation experimental results show the effectiveness and robustness of our model.

Inspired by the similarity between biological sequences and natural language sentences, we divided sequence into k-mer, which can be considered as “words” in biological language. The experimental proved this feature extraction scheme works well. However, rethinking of the procedure of RNA2vec and pro2vec, we recognize that k-mer may not be the best way to sequence word segmentation. More bio-semantic sequence segmentation should be explored in the future.

Author contributions
H-C. Y and Z-H. Y conceived the algorithm, carried out analyses, prepared the data sets, carried out experiments, and wrote the manuscript; L. C, Z, T-H. J and X. L designed, performed and analyzed experiments and wrote the manuscript; All authors read and approved the final manuscript.

Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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