iRNA-m7G: Identifying N7-methylguanosine Sites by Fusing Multiple Features

Wei Chen,1,2 Pengmian Feng,1 Xiaoming Song,2 Hao Lv,3 and Hao Lin3

As an essential post-transcriptional modification, N7-methylguanosine (m7G) regulates nearly every step of the life cycle of mRNA. Accurate identification of the m7G site in the transcriptome will provide insights into its biological functions and mechanisms. Although the m7G-methylated RNA immunoprecipitation sequencing (MeRIP-seq) method has been proposed in this regard, it is still cost-ineffective for detecting the m7G site. Therefore, it is urgent to develop new methods to identify the m7G site. In this work, we developed the first computational predictor called iRNA-m7G to identify m7G sites in the human transcriptome. The feature fusion strategy was used to integrate both sequence- and structure-based features. In the jackknife test, iRNA-m7G obtained an accuracy of 89.88%. The superiority of iRNA-m7G for identifying m7G sites was also demonstrated by comparing with other methods. We hope that iRNA-m7G can become a useful tool to identify m7G sites. A user-friendly web server for iRNA-m7G is freely accessible at http://lin-group.cn/server/iRNA-m7G/.

INTRODUCTION

Besides N1-methyladenosine (m1A), N7-methylguanosine (m7G) is another kind of positively charged RNA modification.1 m7G is added to the 5′ end co-transcriptionally during transcription, and it is essential for efficient gene expression and cell viability.2 It has been found that m7G is required for nearly all phases of the mRNA cycles, such as RNA splicing,3 polyadenylation,3 nuclear export of mRNA,5 translation,6 and so on. Although studies on m7G have been carried out for a long time, the knowledge about its function is still limited. The key step of revealing the functions of m7G is to determine its accurate position in the transcriptome.

By using the mass spectrometry quantification and m7G-methylated RNA immunoprecipitation sequencing (MeRIP-seq) method,7 Zhang et al. not only detected the m7G sites in Homo sapiens and Mus. Musculus but also provided the base resolution m7G sites in human HeLa and HepG2 cells. However, the MeRIP-seq method still has its own limitations,7 and it is cost-ineffective for performing transcriptome-wide detections. Therefore, it is necessary to develop computational methods for identifying m7G sites.

RESULTS AND DISCUSSION

Performance of Each Kind of Feature

We built three models based on the three kinds of features (nucleotide property and frequency [NPF], pseudo nucleotide composition [PseDNC], and secondary structure component [SSC]), and we compared their performances for identifying m7G sites. As indicated in Equations 4 and 5, the PseDNC model is dependent on two parameters, w and λ. Hence, we first optimized the parameters of PseDNC. In general, the greater the λ value is, the more global sequence-order information the model contains. However, a larger λ would reduce the cluster-tolerant capacity so as to lower the cross-validation accuracy due to an overfitting problem. Therefore, the search ranges for

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Correspondence: Wei Chen, Center for Genomics and Computational Biology, School of Life Sciences, North China University of Science and Technology, Tangshan 063000, China.
E-mail: chenweimu@gmail.com
Correspondence: Hao Lin, Key Laboratory for Neuro-Information of Ministry of Education, School of Life Science and Technology, Center for Informational Biology, University of Electronic Science and Technology of China, Chengdu 610054, China.
E-mail: hlin@uestc.edu.cn
w and \( \lambda \) were set in [0, 1] and [1, 10] with a step of 0.1 and 1, respectively. As shown in Figure 1, the PseDNC-based model yielded the best results when \( w = 0.8 \) and \( \lambda = 8 \).

The k-fold cross-validation test method was often used to examine the quality of various predictors.\(^{10}\) For saving computational time, in the current study, the 10-fold cross-validation test was used to evaluate the performance of these models. Their predictive results were reported in Table 1. Among the three models, the NPF-based model obtained the highest accuracy of 89.14%, which is approximately 5% and 14% higher than that of the PseDNC- and SSC-based models, respectively, for identifying m7G sites in the dataset.

To objectively compare their performances, the area under the receiver operating characteristic curve (auROC) of these methods was also calculated. The NPF-based model obtained an auROC of 0.899, higher than the 0.841 and 0.776 obtained by the PseDNC- and SSC-based models, respectively.

### Performance of Fusing Multiple Features

To investigate whether the feature fusion strategy could improve the performance, we built another model by fusing the NPF, PseDNC, and SSC features. The framework of how to build the model is shown in Figure 2. The model thus obtained was then evaluated by using the 10-fold cross-validation test. The detailed results are provided in the last row of Table 1. As indicated in Table 1, the sensitivity (Sn), specificity (Sp), accuracy (Acc), and Mathew’s correlation coefficient (MCC) were all improved compared with those obtained by the NPF-, PseDNC-, and SSC-based models.

To intuitively compare the performance of the models based on different features, their ROC curves from the 10-fold cross-validation test were plotted in Figure 3. The fusion strategy-based model obtained an auROC of 0.946, which is higher than those of the NPF-, PseDNC-, and SSC-based models.

Moreover, to further demonstrate its stability for identifying m7G sites, the fusion strategy-based model was also evaluated by the jackknife test, in which each sample in the training dataset is in turn singled out as an independent test sample, and all the properties are calculated without including the one being identified. In the jackknife test, the fusion strategy-based model obtained an accuracy of 89.88% with the sensitivity of 89.07%, specificity of 90.69%, and MCC of 0.80, which is comparable to those from the 10-fold cross-validation test. These results indicate that the feature fusion strategy is effective and the model is robust for identifying m7G sites.

### Comparison of SVM and Other Classifiers

Since there is no computational method that has been proposed for identifying m7G sites, to demonstrate its effectiveness, we compared the performance of the current SVM-based model with those of the Naive Bayes-, Random Forest-, LogitBoost-, and BayesNet-based models. The Naive Bayes, Random Forest, LogitBoost, and BayesNet were implemented by using WEKA.\(^{11}\) For a fair comparison, all the models were built by using the feature fusion strategy and tested on the same dataset. The 10-fold cross-validation test results of these models are reported in Table 2. As shown in Table 2, the SVM-based model obtained the best results in terms of the four metrics defined in Equation 9. The predictive accuracy of the SVM-based model is 9.7%, 3.3%, 6.1%, and 7.7% higher than those of the Naive Bayes-, Random Forest-, LogitBoost-, and BayesNet-based models, respectively. This result demonstrates that the SVM is more effective than other classification algorithms for identifying m7G sites.

### Conclusions

In this study, we proposed iRNA-m7G, the first computational method to identify m7G sites. In this predictor, the feature fusion strategy was used to represent RNA sequences. Comparative results demonstrated that the feature fusion strategy is much more effective for identifying m7G sites than a single kind of feature.

Moreover, we also compared iRNA-m7G with the other four machine-learning algorithm-based methods, and we found that the SVM-based model achieves the best performance for identifying m7G sites.
For the convenience of the scientific community, a publicly accessible web server called iRNA-m7G that allows the prediction of m7G sites in RNA was established at http://lin-group.cn/server/iRNA-m7G/. We anticipate that iRNA-m7G will become a useful tool for identifying m7G sites. In future works, we will collect more m7G data and use powerful methods such as deep learning to improve the performance of computationally identifying m7G sites.

**MATERIALS AND METHODS**

**Benchmark Datasets**

By using the MeRIP-seq method, Zhang et al. detected 801 base-resolution m7G sites that appeared in human HeLa and HepG2 cells. By mapping these sites to the human genome (hg19), 801 m7G sites containing sequences were obtained. Preliminary tests indicated that the best predictive result was achieved when the sequence length is 41 bp with the m7G site in the center. To build a high-quality dataset, the CD-HIT software with the threshold of 80% was used to remove redundant sequences. Accordingly, we obtained 741 m7G site-containing sequences.

The non-m7G site-containing sequences were obtained by choosing 41-bp-long sequences with the intermediate guanosine not detected as m7G by the MeRIP-seq method. By doing so, a huge number of negative samples is obtained. Since imbalanced datasets affect the performance evaluation of computational methods, to balance the numbers between positive and negative samples in model training, we randomly picked out 741 non-m7G site sequences with the sequence similarity less than 80% to form the negative samples.

**Sequence Representation**

**NPF**

The NPF is an effective sequence-encoding scheme for computationally identifying nucleotide modification sites. According to NPF, the i-th nucleotide $n_i$ in RNA sequence can be represented by a four-dimensional vector $(x_i, y_i, z_i, d_i)$, in which the elements are defined as follows:

$$
x_i = \begin{cases} 
1 & \text{if } n_i \in \{A, G\} \\
0 & \text{otherwise}
\end{cases}$$

$$
y_i = \begin{cases} 
1 & \text{if } n_i \in \{A, U\} \\
0 & \text{otherwise}
\end{cases}$$

$$
z_i = \begin{cases} 
1 & \text{if } n_i \in \{A, C\} \\
0 & \text{otherwise}
\end{cases}$$

(Equation 1)

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**Table 1. Predictive Results for Identifying m7G Sites by Using Different Features**

| Features          | Sn (%) | Sp (%) | Acc (%) | MCC  | auROC |
|-------------------|--------|--------|---------|------|-------|
| NPF               | 88.12  | 90.15  | 89.14   | 0.78 | 0.899 |
| PseDNC            | 81.92  | 87.99  | 84.95   | 0.70 | 0.841 |
| SSC               | 73.11  | 78.71  | 75.91   | 0.52 | 0.776 |
| Fusion            | 88.66  | 90.96  | 89.81   | 0.80 | 0.946 |

Sn, sensitivity; Sp, specificity; Acc, accuracy; MCC, Mathew’s correlation coefficient; auROC, area under the receiver operating characteristic curve; NPF, nucleotide property and frequency; PseDNC, pseudo nucleotide composition; SSC, secondary structure component.

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**Figure 2. Framework of Developing iRNA-m7G**

For an RNA sequence, it is converted into a feature vector by fusing nucleotide property and frequency, pseudo nucleotide composition, and secondary structure component. The support vector machine was used to build the classification model.
Besides the local sequence order information, the global sequence order effect is also important for computationally identifying RNA modification sites. Accordingly, in the current study, the PseDNC was also used to encode the RNA sequences, which can be calculated by using PseKNC and PseKNC-General. Based on PseDNC, the RNA sequence is converted into a discrete vector defined as follows:

$$R = [d_1, d_2, \cdots, d_{16}, d_{18}, \cdots, d_{16} \cdot 1]^{T},$$  \hspace{1cm} (Equation 4)

where

$$d_i = \frac{1}{|N_i|} \sum_{j=1}^{l} f(n_j), \quad f(n_j) = \begin{cases} 1 & \text{if } n_j = n_i; \\ 0 & \text{otherwise.} \end{cases} \hspace{1cm} (Equation 2)$$

where $l$ is the sequence length, and $|N_i|$ is the length of the $i$-th prefix string $\{n_1, n_2, \cdots, n_i\}$ in the sequence.

According to NPF, an RNA sequence with a length of $l$ bp will be encoded by the following vector:

$$R = [x_1, y_1, z_1, d_1, \cdots, x_i, y_i, z_i, d_i, \cdots, x_l, y_l, z_l]^{T}. \hspace{1cm} (Equation 3)$$

**PseDNC**

Besides the local sequence order information, the global sequence order effect is also important for computationally identifying m7G sites by the 10-Fold Cross-Validation Test. Table 2 shows the performance comparison of different classifiers for identifying m7G sites.

| Classifiers     | Sn (%) | Sp (%) | Acc (%) | MCC  |
|-----------------|--------|--------|---------|------|
| Naive Bayes     | 72.47  | 87.85  | 80.16   | 0.61 |
| Random Forest   | 83.27  | 89.88  | 86.57   | 0.73 |
| LogitBoost      | 81.38  | 86.23  | 83.81   | 0.68 |
| BayesNet        | 77.19  | 87.04  | 82.12   | 0.65 |
| SVM             | 88.66  | 90.96  | 89.81   | 0.80 |

Sn, sensitivity; Sp, specificity; Acc, accuracy; MCC, Mathew’s correlation coefficient; SVM, support vector machine.

**SSC**

The formation of RNA modification is affected by RNA structures. Hence, the RNAsfold tool in the ViennaRNA package was used to predict the secondary structure of the RNA sequences in the dataset. For each position in the RNA, the paired nucleotide was represented by a parenthesis (“(” or “)”), while the unpaired one was represented by a dot (“.”). In the current study, we do not distinguish “(“ and “)” and use “(” for both statuses. For a given tri-nucleotide, there are eight (2³) possible structure statuses (i.e., “(((,” “((,” “(,” “.,(,” “(.,” “.,(,” “.,(,” “.)}). Together with the first nucleotide of the tri-nucleotide,
there will be 32 (4 × 8) possible sequence-structure modes denoted as “A-((,” “A-(,” “A-(,” ..., and “U-...” Therefore, by using the sequence-structure mode, an RNA sequence can be represented as follows:

\[ R = \begin{bmatrix} f^A_1 \cdots f^A_8 & f^U_1 \cdots f^U_8 \end{bmatrix}^T. \]  

(Equation 8)

**SVM**

In the current study, the LibSVM package 3.18, which is available at https://www.csie.ntu.edu.tw/~cjlin/libsvm/, was used to perform the classification task. The basic idea of SVM is to transform the input data into a high-dimensional feature space and then determine the optimal separating hyperplane. Because of its better performance, the radial basis kernel function (RBF) was used to obtain the separating hyperplane. The regularization parameter \( C \) and kernel parameter \( \gamma \) of the SVM operation engine were optimized in the ranges of \([2^{-5}, 2^{15}]\) and \([2^{-15}, 2^{-5}]\) with the steps of 2 and 2\(^{-1}\), respectively. The final prediction was made according to the probability obtained by SVM. \(^{22–33}\) If its probability is >0.5, a guanine will be predicted as an m7G site.

**Evaluation Metrics**

In this study, the four metrics, \(^{34–40}\) namely, Sn, Sp, Acc, and MCC, were used to measure the performance of the proposed methods, which are defined as follows:

\[
\begin{align*}
Sn &= 1 - \frac{N^-}{N^+} \quad 0 \leq Sn \leq 1 \\
Sp &= 1 - \frac{N^-}{N} \quad 0 \leq Sp \leq 1 \\
Acc &= 1 - \frac{N^+ + N^-}{N^+ + N^-} \quad 0 \leq Acc \leq 1 \\
MCC &= \frac{1 - \frac{N^+}{N^+} \cdot \frac{N^-}{N^-}}{\sqrt{\left(1 + \frac{N^-}{N^+}\right) \left(1 + \frac{N^+}{N^-}\right)}} \quad -1 \leq MCC \leq 1
\end{align*}
\]

(Equation 9)

where \( N^+ \) represents the m7G site-containing sequence, while \( N^+ \) is the number of m7G site-containing sequences incorrectly predicted to be of false m7G site-containing sequences; \( N^- \) is the total number of false m7G site-containing sequences, while \( N^- \) is the number of the false m7G site-containing sequences incorrectly predicted to be of m7G site-containing sequences.

Moreover, by plotting the sensitivity against (1-specificity) with the varying of the threshold, the ROC curve\(^{41,42}\) was generated to evaluate the performance of the proposed method. The auROC is an indicator of the performance of the method. An auROC value of 0.5 is equivalent to random prediction while an auROC of 1 represents a perfect one.

**AUTHOR CONTRIBUTIONS**

W.C. and H. Lin conceived and designed the study, W.C., P.F., X.S., and H. Lin conducted the experiments. P.F., W.C., and X.S. implemented the algorithms. H. Lv established the web server. W.C., P.F., X.S., H. Lv, and H. Lin performed the analysis and wrote the paper. All authors read and approved the final manuscript.

**CONFLICTS OF INTEREST**

The authors declare no competing interests.

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