A comprehensive comparison and analysis of computational predictors for RNA N6-methyladenosine sites of *Saccharomyces cerevisiae*

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Abstract

N6-methyladenosine (m6A) modification, as one of the commonest post-transcription modifications in RNAs, has been reported to be highly related to many biological processes. Over the past decade, several tools for m6A sites prediction of *Saccharomyces cerevisiae* have been developed and are freely available online. However, the quality of predictions by these tools is difficult to quantify and compare. In this study, an independent dataset M6Atest6540 was compiled to systematically evaluate nine publicly available m6A prediction tools for *S. cerevisiae*. The experimental results indicate that RAM-ESVM achieved the best performance on M6Atest6540; however, most models performed substantially worse than their performances reported in the original papers. The benchmark dataset Met2614, which was used as the training dataset for the nine methods, were further analyzed by using a position bias index. The results demonstrated the significantly different bias of dataset Met2614 compared with the RNA segments around m6A sites recorded in RMBase. Moreover, newMet2614 was collected by randomly selecting RNA segments from non-redundant data recorded in RMBase, and three different kinds of features were extracted. The performances of the models built on Met2614 and newMet2614 with the features were compared, which shows the better generalization of models built on newMet2614. Our results also indicate the position-specific propensity-based features outperform other features, although they are also easily over-fitted on a biased dataset.

Key words: N6-methyladenosine sites; computational predictor; dataset bias; position-specific propensity; web servers
**Introduction**

N6-methyladenosine (m6A) modification is the first internal mRNA modification discovered [1, 2]. It is considered as one of the commonest post-transcription modifications in RNAs and mRNA modification discovered [1, 2]. It is considered as one of the most widely utilized model organisms in biotechnology worldwide, and 13 computational methods [33, 34, 36, 38, 39, 42, 44, 45, 47, 48, 50–52] have been developed to predict its m6A sites. In this study, we focused on these methods. These models were summarized in Table 1; we noticed all the models were built on a training dataset Met2614 [33] which was based on the pioneering work of Schwartz et al. [57] and contains only 1307 positive examples. However, the number of m6A sites of S. cerevisiae collected in a RNA modification database (RMBase [58]) is over 60,000, of which 23,581 m6A sites are centered with m6A modification was associated with lots of diseases such as thyroid tumor [21], prostate cancer [22], breast cancer [23–25], pancreatic cancer [26, 27], leukemia [28], etc. Obviously, the identification of m6A sites would be of great benefit for cell biology and disease mechanism research.

Various types of methods have been developed to identify m6A sites. Experimental approaches such as two-dimensional thin layer chromatography [29], high performance liquid chromatography [30] and next-generation sequencing techniques (e.g. m6A-seq [31] and MeRIP-Seq [32]) have been used to identify m6A sites in mRNAs. However, these experimental methods are too costly and time-consuming to perform genome-wide analysis. These limitations of experimental methods may be overcome by developing computational methods for m6A sites identification.

Over the past few years, a variety of computational methods [33–56] have been developed for predicting m6A sites in different species of RNAs, such as Saccharomyces cerevisiae, Homo sapiens, Mus musculus and Arabidopsis thaliana. Saccharomyces cerevisiae is one of the most widely utilized model organisms in biotechnology worldwide, and 13 computational methods [33, 34, 36, 38, 39, 42, 44, 45, 47, 48, 50–52] have been developed to predict its m6A sites. In this study, we focused on these methods. These models were summarized in Table 1; we noticed all the models were built on a training dataset Met2614 [33] which was based on the pioneering work of Schwartz et al. [57] and contains only 1307 positive examples. However, the number of m6A sites of S. cerevisiae collected in a RNA modification database (RMBase [58]) is over 60,000, of which 23,581 m6A sites are centered with GAC pattern.

Our original idea is to select a representative subset to validate the generalization of the available methods built on a small dataset Met2614. We first carefully collected an independent test set and carried out an unbiased evaluation of these predictors that have been publicly available as open access web

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Training dataset</th>
<th>Independent test set</th>
<th>Feature representation</th>
<th>Classifier</th>
<th>Cross validation</th>
<th>Web server</th>
<th>Ref</th>
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<tr>
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<td>Pseudo dinucleotide composition (PseDNC)</td>
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<td>10-fold and</td>
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<td>[33]</td>
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<td>Met950^</td>
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<td>SVM</td>
<td>Jackknife</td>
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<td>SVM</td>
<td>10-fold</td>
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<td>Met414^</td>
<td>Enhanced nucleic acid composition</td>
<td>RF</td>
<td>5-fold and 10-fold</td>
<td><a href="http://www.bioinfogo.org/bermp">http://www.bioinfogo.org/bermp</a></td>
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<td>Dinucleotide binary encoding; local position-specific dinucleotide frequency</td>
<td>XGBoost</td>
<td>10-fold and</td>
<td><a href="http://server.malab.cn/">http://server.malab.cn/</a></td>
<td>[52]</td>
</tr>
</tbody>
</table>

^The datasets are derived from Met2614, so all the datasets are not overlapped with M6Atest6540.

LR, logistic regression; SVM, support vector machine; RF, random forest; BGRU, bidirectional Gated Recurrent Unit; XGBoost, eXtreme Gradient Boosting.
portals. As shown in Table 1, totally 11 predictors have web services as reported in their original papers; however, two of them, TargetM6A [42] and M6A-HPCS [39], are not open-access now. Therefore, we tested and compared the rest nine m6A prediction webservers in this study. An independent test set, M6Atest6540 including 3270 positive examples, is collected from RMBase [58]. Our comparative results indicate that RAM-ESVM achieved the best performance on M6Atest6540; however, the generalization and robustness of all nine methods are not good compared with the performances on Met2614. The RNA segments in Met2614 were further compared with the segments around m6A sites recorded in RMBase, and the results indicate the bias of Met2614 compared with the segments recorded in RMBase.

To further analyze if the bias of Met2614 affects the generalization of the models, we built new datasets by randomly selecting 1307 nonredundant positive examples from RMBase, which was called newMet2614. Then, three different kinds of features were extracted from the RNA segments, namely nucleotide composition-based features, position-specific propensity-based features and physical–chemical properties-based features. With these features, the performances of the models built on newMet2614 were compared with the models built on Met2614. The results indicate that generalization of the models based on newMet2614 is better than the models based on Met2614. Our results also indicate the position-specific propensity-based features outperform other features; however, they are also easily over-fitted on a biased dataset.

Materials and methods

Datasets

The dataset Met2614 [33] contains total 1307 positive RNA segments with experimentally verified m6A sites at the center, which were collected from Schwartz et al.’s work [57]. In order to fairly compare the predictive performance of different web tools, we constructed an independent dataset, M6Atest6540, which contains 3270 experimentally verified m6A sites and other remaining non-m6A sites. Note that benchmark dataset Met2614 and M6Atest6540 are mutually exclusive.

The following seven steps were conducted to create the benchmark dataset M6Atest6540 (Figure 1): (i) collecting all RNA segments of 51-nt nucleotides with exactly the same RGAC consensus motif at the center position by sliding a flexible window along each RNA sequences transcribed from the S. cerevisiae genome. This set of segments obtained here is called T1. (ii) Collecting all RNA segments whose center positions were recorded in RMBase [58], this set of segments is called P1 because all examples are positive. Then P1 was removed from T1 to obtain the set of segments N1, which contains all the negative examples. (iii) A subset of P1 with ‘SupportNum’ > 15 and ‘Region’ == ‘cds’ was selected to increase the reliability of positive examples. ‘SupportNum’ is a term in RMBase to describe the number of supporting experiments or studies for that position, and ‘Region’ is another term to describe the region the nucleotide located. This subset is called P2. (iv) CD-HIT [59] was used to remove the redundancy [60] of all the positive examples of P2 and Met2614 with the sequence identity cutoff set as 75%. A total of 3270 positive examples were obtained from this step. (v) The negative examples of Met2614 were removed from N1 to get a new negative dataset N2. (vi) A total of 3270 negative examples were randomly selected from N2. (vii) The 3270 positive samples and 3270 negative samples were assembled into the benchmark dataset M6Atest6540. All the RNA segments of M6Atest6540, P1 and N1 can be found in Table S1, S2, S3, respectively.

Web-accessible prediction tools

As shown in Table 1, there are totally 13 methods for predicting m6A sites of S. cerevisiae, and 11 of them provide web servers. Currently, only 9 of these 11 web servers are open-access for users to predict m6A sites, namely iRNA-Methyl [33], m6AProf [34], pRNAm-PC [36], RAM-ESVM [44], RAM-NPPS [45], M6APred-EL [47], iRNA(m6A)-PseDNC [50], BERMP [51] and M6AMRFS [52]. Table 1 lists the basic information of these methods. As shown in Table 1, the validation and test datasets are all based on Met2614. Notably, the outputs of the servers pRNAm-PC and RAM-ESVM are color labeled, which makes it a little bit hard to collect the results.

Features extraction for machine learning-based predictors

In the process of training a model, it is crucial to extract informative, discriminative and independent features for converting the RNA sequence into the numeric vector. Among the past studies, many features have been extracted to interpret and keep the sequence information [44, 61–66]. In this study, three groups of features were gathered to analyze the possible reasons that caused the weak generalization of the web tools and to study the effects of benchmark dataset bias on the performances of different features for m6A sites prediction. The three groups of features are nucleotide composition-based features, position-specific propensity-based features and physical–chemical properties-based features (Table 2). The detailed descriptions of all the features can be found in Supplementary Data.

Support vector machine

The basic idea of the support vector machine (SVM) is to determine the optimal separating hyperplane that can correctly
divide the training data set and have the largest interval for linearly separable data \([67, 68]\). In terms of the nonlinearly separable data, SVM maps the original data to high-dimensional space by using the specific kernel, which transforms the problem into linear one. Kernel functions include linear kernels, polynomial kernels, Gaussian kernels, sigmoid kernels and so forth. Among them, Gaussian kernels, also called Radial Basis Function (RBF), are most commonly used \([69–72]\), which can map data to infinite dimensions. There are two indispensable parameters for SVM models with RBF, which are \(C\) and \(\gamma\). The former indicates the tolerance of the model to the error, and the latter implicitly determines the distribution of the data after mapping to the new feature space. As shown in Table 1, most of the models were trained by using SVM.

**Performance evaluation parameters**

According to the previous related studies \([73, 74]\), Sensitivity (SN), Specificity (SP), Accuracy (ACC) and Matthews correlation coefficient (MCC) are the mostly used performance evaluation parameters for the computational predictors, which are defined as follows:

\[
\begin{align*}
SN &= \frac{TP}{TP + FN} \\
SP &= \frac{TN}{TN + FP} \\
ACC &= \frac{TP + TN}{TP + TN + FP + FN} \\
MCC &= \frac{(TP \times TN) - (FP \times FN)}{\sqrt{(TP + FN) \times (TP + FP) \times (TN + FP) \times (TN + FN)}}
\end{align*}
\]

where, \(TP, TN, FP\) and \(FN\) represent the number of true positive, true negative, false positive and false negative, respectively. SN and SP show the ability of the model to correctly predict positive samples and negative samples, respectively. Furthermore, it is worth noting that ACC and MCC are the most important ones of the aforementioned four metrics, because the former reflects the overall accuracy of the predictor, while the latter signified the overall stability.

**Position bias index**

Here, we introduced a position bias index (PBI) to evaluate the bias at a specified position between two groups of sequences.

\[
PBI(k) = \sum_{i=1}^{t} (P_{i,k}^{1} - P_{i,k}^{2})^2
\]

where \(k\) is the specified sequence position, \(t\) is the number of nucleotide or amino acid types and \(P_{i,k}^{g} (g \in \{1, 2\})\) is the probability of the \(i\)th type residue or nucleotide at the specified position \(k\) of the sequence group \(g\), which was defined as follows:

\[
P_{i,k}^{g} = N_{i,k}^{g}/N_{g}
\]

where, \(N_{g} (N_{1} = m, N_{2} = n)\) is the number of sequences in group \(g\) and \(N_{i,k}^{g}\) is the number of \(i\)th type residue at the specified position \(k\) of the sequence group \(g\).

Generally speaking, the bigger the PBI is, the more biased the corresponding position is. However, the bias is also determined by the distribution of PBIs at the specified position.

**Results and Discussion**

**Comparative results on the independent dataset M6Atest6540**

Our original objective of this study is to conduct a fair comparison of existing prediction methods with the help of an independent dataset. We first built the independent dataset M6Atest6540 as described in the Material and Methods section. Then, we compared the nine existing prediction methods with
available web servers, namely, M6APred-EL [47], RAM-NPPS [45], RAM-ESVM [44], pRNAm-PC [36], iRNA-Methyl [33], BERMP [51], M6AMRFS [52], iRNA(m6A)-PseDNC [50] and m6Apred [34]. The predictive results of these methods on the independent dataset are shown in Table 3 and Figure 2, and the predictive labels of these methods for the RNA segments in M6Atest6540 are shown in Table S1.

As shown in Table 3, RAM-ESVM achieved the best performance among the nine tested predictors, with the highest ACC of 61.90% and MCC of 0.238. pRNAm-PC and iRNA-Methyl are the 2nd and 3rd best performers, which achieved ACC of 60.78%, 60.75% and MCC of 0.217, 0.215, respectively. Besides, BERMP achieved the highest specificity of 86.76%, and iRNA(m6A)-PseDNC achieved the highest sensitivity of 83.85%.

Moreover, we also tried to plot the receiver operating characteristics (ROCs) curves for all the predictive results of the nine servers; however, only six of them outputted scores or probabilities. So, we only plotted the ROC curves (Figure 3) for the six methods. The area under the curves are 0.652, 0.649, 0.638, 0.577, 0.574 and 0.533 for M6APred-EL, iRNA-Methyl, BERMP, M6AMRFS, m6Apred and iRNA(m6A)-PseDNC, respectively, which is basically in line with the ACCs in Table 3, except M6APred-EL.

In addition, we compared the performances of these models on their original training dataset and the independent dataset M6Atest6540 to evaluate the generalization of these models. Figure 2 shows the comparison results. MCCs of these models on the training dataset Met2614 and the independent test set M6Atest6540 were shown in Figure 2A, and it is surprising that the generalization of most models is not good, especially those models with high MCCs. If the difference of MCCs between the training dataset and the independent dataset is considered as the parameter to evaluate the generalization, the three best performing models are iRNA-Methyl, pRNAm-PC and BERMP, with differences of MCCs of 0.075, 0.183 and 0.245, respectively. Because MCC is an integral parameter for model evaluation, the sensitivities and specificities of models on the training dataset and the independent dataset were also compared to further explore the cause of the poor generalization. Figure 2C and D show that, for all models except iRNA(m6A)-PseDNC, the low sensitivity on the independent dataset is the major cause of the poor generalization. For iRNA(m6A)-PseDNC, the low specificity on the independent dataset is the main reason. Compared with training datasets used in other methods, the training dataset of iRNA(m6A)-PseDNC contains different negative examples, which may explain the different causes of poor generalization of iRNA(m6A)-PseDNC.

Analysis of the bias of the training dataset Met2614

Given that Met2614 is an early small dataset which was collected from Schwartz et al.’s work [57], we analyzed if the bias between the positive and negative samples in Met2614 is significantly different from the bias between the positive and negative samples recorded in RMBase. By using the FBI defined in the Material and Methods section, the position bias of the training dataset Met2614 was analyzed. As shown in Figure 4, three positions of the dataset Met2614, 22, 24 and 30, are substantially biased. Intuitively, the two positions, 24 and 30, are more seriously
biased. To show if the bias is statistically significant, we focused on these two positions.

PBIs were first calculated based on all the positive and negative examples in the whole S. cerevisiae genome. As mentioned in the Material and Methods section, all the positive and negative examples were generated during generation of the dataset M6Atest6540. As shown in Figure 4, the PBIs at positions 24 and 30 of Met2614 are substantially higher than the corresponding PBIs for the whole genome.

Furthermore, 5000 different datasets were created by random selection of 1307 positive examples and 1307 negative examples from the genome, respectively. The PBIs at each position of the 5000 datasets were calculated, and the distributions of PBIs at positions 24 and 30 are shown in Figure S1. The distributions were fitted by the Weibull distribution. The statistics and parameters of the distributions for positions 24 and 30 are summarized in Table 4. The cumulative distribution function of these two distributions are showed in Figure S2. Based on the fitted distribution, the P-values of PBIs at positions 24 and 30 of Met2614 were calculated to be 1.652E-16 and 1.432E-158, respectively, which proved the bias of the dataset Met2614.

The data were further analyzed to explore the possible causes of the bias. Firstly, the nucleotide composition of the positive examples of Met2614 were compared with that of the positive examples recorded in RMBase [58] at positions 24 and 30. Figure 5A shows that the nucleotide composition of positive examples of Met2614 is substantially different from that of the positive examples recorded in the genome at these two positions, while Figure 5B shows that the nucleotide composition of the negative examples of Met2614 is similar to that of the negative examples recorded in the genome at these two positions. Thus, the bias is more likely to be caused by the positive examples of Met2614.

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### Table 4. The statistics and parameters of the PBI distribution at positions 24 and 30 of the length 51 sequence segment

<table>
<thead>
<tr>
<th>Position</th>
<th>Mean of PBI</th>
<th>Std of PBI</th>
<th>Scale parameter</th>
<th>Shape parameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>24</td>
<td>7.5199E-04</td>
<td>0.0010</td>
<td>0.0006</td>
<td>0.6827</td>
</tr>
<tr>
<td>30</td>
<td>0.0012</td>
<td>0.0010</td>
<td>0.0013</td>
<td>1.2678</td>
</tr>
</tbody>
</table>

*aThese are the two parameters for Weibull distribution.
Performances of the models built on Met2614 and newMet2614 with different features

To further explore if the bias of Met2614 affects the generalization of the models, we collected a new dataset, newMet2614, by randomly selecting positive examples from a non-redundant dataset based on RMBase. The newMet2614 also contains 1,307 positive and negative examples, respectively. Three groups of features as shown in Table 2 were extracted from the RNA segments of both Met2614 and newMet2614. With each feature, the models were built on Met2614 and newMet2614 by using SVM, respectively. The two parameters, KernelScale and BoxConstraint, of the MATLAB function FITCSVM were selected by a grid search according to the results of the leave-one-out cross validation. The range of KernelScale is from $2^{-10}$ to $2^6$, and the range of BoxConstraint is from $2^{-5}$ to $2^{15}$. Note that all the works for newMet2614, including the generation of newMet2614, were repeated three times. Figure 6 shows the cross validation results of the models on Met2614 and newMet2614, respectively. Figure 7 shows the predictive results of the models on dataset M6Atest6540. For the models based on newMet2614, it indicates that the performances on M6Atest6540 are better than the cross validation results on newMet2614; in other words, the ACCs and MCCs on M6Atest6540 are higher than the cross validation results (Figures 6C and D, 7C and D and Table S4). These models show good performances on the positive examples of M6Atest6540; in other words, the sensitivities on M6Atest6540 are higher than the cross validation results (Figures 6A and 7A and Table S4). This is normal because the positive examples in M6Atest6540 are typical for their experimental 'SupportNum' $> 15$ (see Datasets). For the models based on Met2614, especially the models built with position-specific propensity-based features (Features 1–21 and 44), the performances on M6Atest6540 are substantially worse than the cross validation.
results on Met2614 (Figures 6C and D, 7C and 7D and Table S5), especially for sensitivities (Figures 6A and 7A and Table S5). Intuitively, the models built with position-specific propensity-based features are easily affected by the position bias of Met2614, which is in line with our results.

In addition, we compared the performances of the models based on newMet2614 with the models based on Met2614 on the independent test set M6Atest6540. Figure 7C and D shows that the models built on newMet2614 outperform the models based on Met2614; in other words, the ACCs and MCCs of the models based on newMet2614 are higher than the corresponding values of the models based on Met2614; these values are also higher than the corresponding values of the nine previous developed models. Although the models based on newMet2614 show better generalization than the models based on Met2614, it is better to use a large dataset to build the models to predict RNA m6A of S. cerevisiae.

Moreover, the ratio of m6A sites and non-m6A sites in both Met2614 and M6Atest6540 is set to 1.0, which creates the balanced datasets; however, there are more non-m6A sites in the genome than the m6A sites realistically. In other words, the datasets for this classification problem should be imbalanced. Random under-sampling is one of the ways to solve the imbalanced learning problem; however, the important information of negative examples may be lost, and the model may be easily to predict more examples as positive, such that the sensitivity will be overestimated. So, other methods such as informed under-sampling, cost-sensitive learning and kernel-based method could be used to deal this problem.

**Conclusion**

In this study, a benchmark dataset M6Atest6540 was compiled to evaluate and compare the performances of nine available methods for predicting m6A sites. Most of the methods performed substantially worse on the independent dataset M6Atest6540 than on the training dataset Met2614. Note that the Met2614 is a benchmark dataset used for most method building, and this dataset is a small dataset based on the early work of Schwartz et al. [57]. Further analysis shows that Met2614 is a biased dataset compared with the examples recorded in RMBase. In addition, the impact of the dataset bias on the generalization and robustness of models built on different features was further analyzed. The position-specific propensity-based features were found to be easily over-fitted on a biased dataset. Due to the more data we can obtain for RMBase, it is better to use a large benchmark to build RNA m6A sites prediction models of S. cerevisiae. And other kinds of modification [75, 76] would be studied in the future work.

**Key Points**

- We comprehensively review a variety of existing computational methods for the prediction of RNA m6A sites of S. cerevisiae and conduct a comparative study of available web servers.
- Benchmarking results demonstrate the substantially worse generalization of most of the nine available m6A prediction servers compared with the performances reported in the original papers.
- A FBI was introduced to analyze the possible biased positions between two sets of sequences. And the statistical analysis demonstrates that the widely used dataset Met2614 is significantly biased compared with the examples recorded in RMBase.
- The performances on M6Atest6540 indicate that the models based on newMet2614 that was randomly selected from RMBase outperform the models based on Met2614.

**Supplementary Data**

Supplementary data are available online at https://academic.oup.com/bfg.

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**References**


