i6mA-Vote: Cross-Species Identification of DNA N6-Methyladenine Sites in Plant Genomes Based on Ensemble Learning With Voting

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DNA N6-Methyladenine (6mA) is a common epigenetic modification, which plays some significant roles in the growth and development of plants. It is crucial to identify 6mA sites for elucidating the functions of 6mA. In this article, a novel model named i6mA-vote is developed to predict 6mA sites of plants. Firstly, DNA sequences were coded into six feature vectors with diverse strategies based on density, physicochemical properties, and position of nucleotides, respectively. To find the best coding strategy, the feature vectors were compared on several machine learning classifiers. The results suggested that the position of nucleotides has a significant positive effect on 6mA sites identification. Thus, the dinucleotide one-hot strategy which can describe position characteristics of nucleotides well was employed to extract DNA features in our method. Secondly, DNA sequences of Rosaceae were divided into a training dataset and a test dataset randomly. Finally, i6mA-vote was constructed by combining five different base-classifiers under a majority voting strategy and trained on the Rosaceae training dataset. The i6mA-vote was evaluated on the task of predicting 6mA sites from the genome of the Rosaceae, Rice, and Arabidopsis separately. In Rosaceae, the performances of i6mA-vote were 0.955 on accuracy (ACC), 0.909 on Matthew correlation coefficients (MCC), 0.955 on sensitivity (SN), and 0.954 on specificity (SP). Those indicators, in the order of ACC, MCC, SN, SP, were 0.882, 0.774, 0.961, and 0.803 on Rice while they were 0.798, 0.617, 0.666, and 0.929 on Arabidopsis. According to the indicators, our method was effectiveness and better than other concerned methods. The results also illustrated that i6mA-vote does not only well in 6mA sites prediction of intraspecies but also interspecies plants. Moreover, it can be seen that the specificity is distinctly lower than the sensitivity in Rice while it is just the opposite in Arabidopsis. It may be resulted from sequence similarity among Rosaceae, Rice and Arabidopsis.

Keywords: N6-methyladenine, plant genomes, cross-species, feature encoding, ensemble learning
INTRODUCTION

DNA N6-methyladenine (6mA) is a methyl modification at the sixth position of the adenine ring, which was discovered by Vanyushin et al. (1968). 6mA is widely found in prokaryotes and eukaryotes (Fu et al., 2015; Greer et al., 2015; Zhang et al., 2015). It is reported that 6mA plays vital roles in DNA replication, repairing nucleotide dislocations, and preventing the invasion of foreign DNA (Wion and Casadesús, 2006). Although 6mA in animal genomes studies have been well studied, those of plants genomes have still known a little, which hampered to explore their functions. To better understand the molecular mechanism of 6mA in plants, it is the first step to determine the 6mA sites accurately.

To detect 6mA sites, several biochemical methods were developed, such as single-molecule real-time sequencing technology (SMRT-seq) (Davis et al., 2013) and restriction endonuclease-based 6mA sequencing (6MA-RE-seq) (Fu et al., 2015). In SMRT-seq, single-nucleotide molecules labeled by different fluorophores were paired with bases of a DNA sequence, and the fluorescence signals were recorded during the process of pairing. The fragment of DNA sequence may be methylated if it showed the continuous same signal during the process of pairing. 6mA-RE-seq explored restriction enzymes to fragment genomic DNA at “CATG” and “GATC” motifs that did not contain 6mA and then retained these motifs containing 6mA. In this way, after end-repair and other operations, the methylated motifs would be enriched in the internal positions of DNA fragments. However, these methods are hard to detect 6mA sites from high-throughput sequences because they are time-consuming and expensive.

Therefore, some machine learning models have been developed to identify 6mA sites in recent years because they are efficient and cheap. At first, iDNA6mA-PseKNC (Feng et al., 2019) was proposed to detect 6mA sites in the mouse genome. In this model, DNA sequences were represented by pseudo-k-tuple nucleotide composition incorporating the physicochemical properties of nucleotides, and then the sequences were classified by a support vector machine (SVM). Subsequently, i6mA-Pred (Chen et al., 2019) trained a novel SVM model to identify 6mA sites in the rice genome based on the chemical properties of nucleotides such as the loop structure, the hydrogen bond, and the amino groups, and the nucleotide frequency of DNA sequences. To avoid overfitting, i6mA-Pred used the maximum correlation maximum distance approach to select the most representative features. Afterward, iN6-methylate (Le, 2019), another novel SVM model, used FastText to generate feature vectors for DNA sequences based on the assumption that a DNA sequence is a word. Unlike previous models, MM-6mAPred (Pian et al., 2019) constructed Markov chains based on DNA sequences with 6mA sites (positive samples) and DNA sequences without 6mA sites (negative samples) in the training dataset. Based on the Markov chains, the positive and negative probabilities of a DNA sequence were calculated separately. It is considered that a sequence contained 6mA site if the ratio of positive probability against negative probability is greater than 1.

To improve the performance of above methods, ensemble learning has been increasingly applied to 6mA sites prediction. In the beginning, iDNA6mA-Rice (Lv et al., 2019), a rice 6mA site classification model based on random forest, encoded DNA sequences via three feature descriptors, namely the k-nucleotide frequency, the mono-nucleotide binary coding, and the natural vector containing the frequency, average position, and second-order central moment of mono-nucleotides. Soon afterward, on the basis of bagging with CART, i6mA-DNCP (Kong and Zhang, 2019) represented rice DNA sequences by two novel feature descriptors: dinucleotide frequency and dinucleotide physicochemical properties. In addition, i6mA-DNCP employed heuristic ideas to select the most representative features. Several months later, i6mA-Fuse (Hasan et al., 2020) was proposed to classify Rosaceae DNA sequences with random forest and linear regression. Subsequently, a random forest-based multi-species 6mA site prediction model 6mA-Finder (Xu et al., 2020) was developed, which contained three modules for mouse, rice, and a general species admixed by mouse and rice DNA sequences, respectively. i6mA-stack (Khanal et al., 2021) developed a two-level stacked ensemble classifier based on linear regression, random forest, support vector machine, and gaussian naive bayes to recognize Rosaceae 6mA sites.

With the development of deep learning, some neural network models were also developed for identifying 6mA sites. For example, iDNA6mA (Tahir et al., 2019) is composed of four layers: two convolution layers which extract features of DNA sequences, a dropout layer which is used to avoid overfitting, and a full-connection layer which performs classification tasks. Subsequently, SNNRice6mA (Yu and Dai, 2019) was improved iDNA6mA by adding a normalization layer and a pooling layer between the convolution layer and the dropout layer, which aimed to reduce redundant features of DNA sequences according to the correlation of the features. i6mA-DNC (Park et al., 2020) is similar with the above two models except it extracted features from nucleotide pairs of DNA sequences rather than from single nucleotides. It is worth noting that the three neural network models mentioned above were all developed for predicting 6mA sites in the rice genome.

Because the previously mentioned models are species-specific, Meta-i6mA (Hasan et al., 2021) was proposed for 6mA site prediction from multiple plants. Although Meta-i6mA has achieved encouraging results in intraspecies, it still has room for improvement in interspecific. To solve this problem, a novel classification model i6mA-vote was developed based on an ensemble learning strategy. In this model, DNA sequences were encoded by nucleotide position-based feature descriptors, and then these sequences were classified by an ensemble classifier integrating random forest, linear discriminant analysis, multi-layer perceptron, stochastic gradient descent, and extreme gradient boosting. The details of i6mA-vote will be introduced in the following sections.
MATERIALS AND METHODS
Framework of i6mA-Vote
In our study, as shown in Figure 1, i6mA-vote was constructed by four steps. Firstly, positive samples of Rosaceae, Rice, and Arabidopsis were derived from MDR (Liu et al., 2019), GEO (Edgar et al., 2002), and MethSMRT (Ye et al., 2017) databases, and negative samples of these plants were downloaded from NCBI. For each plant, the positive and negative samples were filtered by CD-HIT (Li and Godzik, 2006) to reduce high similar samples. Then all samples were divided into three datasets according to organisms for the subsequent experiments. The Rosaceae dataset was split into a training dataset and a test dataset, and datasets for the remaining two species were used as cross-species evaluation datasets. Secondly, to transform DNA sequences into feature vectors, one-hot encoding method was performed on dinucleotides (e.g., AA, AG, ... ) of DNA sequences. Because the known nucleotides can be represented by four symbols (A, G, C, T) and other unknown nucleotides can be denoted by symbol N, in this way, there were twenty-five dinucleotide combinations. Thirdly, an ensemble learning model, named i6mA-vote, was built by integrating random forest (RF), multi-layer perceptron (MLP), stochastic gradient descent (SGD), linear discriminant analysis (LDA), extreme gradient boosting (XGB), based on majority voting strategy. Then all samples were represented by feature vectors and the ensemble learning model was trained on the samples. Finally, to evaluate the performance of the model, i6mA-vote was used to perform simulation a task on test datasets, and its superiority was demonstrated by accuracy, Matthew correlation coefficient, sensitivity, and specificity. In the following sections, the
Our Encoding Strategy

One-hot encoding method for dinucleotides (One-hot2) is based on the one-hot encoding method in natural language processing. The one-hot encoding method compiles a dictionary using the words in the sentences and then encodes each word into a 0-1 vector through this dictionary. The length of the vector is equal to that of the dictionary, and each bit in the vector corresponds to a word in the dictionary. When encoding a word, its corresponding bit is set to 1 in the vector, and the other bits are kept at 0. Similarly, One-hot2 treats DNA sequences as sentences and dinucleotides as words.

A DNA sequence is usually composed of four standard nucleotide symbols: A, C, G, and T. However, sometimes the DNA sequence also include non-standard nucleotide symbol N, which means that the nucleotide was not identified. Accordingly, a DNA sequence may consist of 5 symbols, and it contains 25 possible symbol combinations of dinucleotides like AA, AC, AN.

In our method, the one-hot2 encoded each dinucleotide into a 25-dimensional 0-1 vector. The vector of each dinucleotide is shown in Formula (1).

\[
\begin{align*}
AA &= (1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0) \\
AC &= (0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0) \\
AG &= (0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0) \\
\vdots \\
NT &= (0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0) \\
NN &= (0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0)
\end{align*}
\]

To show how One-hot2 encodes DNA sequences, an example is given below. DNA sequence \( D = ACGTNA \) can be split into five dinucleotides (AC, CG, GT, TN, NA), and then they are replaced with their corresponding one-hot codes. In this way, a vector with the dimension of 125 is generated.

Because the length of the DNA sequences in our datasets are 41bp, the sequences can be spliced into 40 dinucleotides and thus the vectors of these dinucleotides were concatenated into a 1000-dimensional feature vector to describe their primary sequence.

There are three reasons why One-hot2 was chosen: (1) It can solve the problem that classifiers are not good at handling continuous data. In addition, it generates sparse vectors, allowing many machine learning problems to be linearly separated and models more efficient to be stored. (2) It considers the relationship between adjacent nucleotides as it is encoded in dinucleotide. (3) Some studies (Chen et al., 2019; Feng et al., 2019) found position-specific features can better represent sequences containing 6mA sites, and One-hot2 happens to be this kind of method.

The Concerned Encoding Strategies

Density-Based Approach

Accumulated Mono-Nucleotide Frequency (AMNF) represent the frequency of single nucleotides in the subsequence which ranges from the first nucleotide to the current nucleotide of the original sequence. Similarly, Accumulated Di-Nucleotide Frequency (ADNF) (Chen et al., 2017) denotes the nucleotide pairs which appears before current nucleotide. For example, DNA sequence

| Datasets | Number of positive samples | Number of negative samples | Total number |
|----------|---------------------------|---------------------------|--------------|
| DS1      | 29237                     | 29433                     | 58670        |
| DS2      | 7298                      | 7300                      | 14598        |
| DS3      | 153635                    | 153629                    | 307264       |
| DS4      | 31414                     | 31843                     | 63257        |
\( D = \text{ACGTNA} \) can be encoded as \((1, 0.5, 0.33, 0.25, 0.2, 0.33)\) and \((1, 0.5, 0.33, 0.25, 0.2)\) by AMNF and ADNF, respectively.

**Physicochemical-Properties-Based Approach**

Dinucleotide Physical-Chemical Properties (DPCP) and Trinucleotide Physical-Chemical Properties (TPCP) (Manavalan et al., 2019; Wei et al., 2019) replace the DNA sequences with the vectors calculated by Equation (2) using the physicochemical-properties in Supplementary Tables 1,2. In Supplementary Table 1, the columns represent 15 physicochemical properties, and the rows represent 25 dinucleotides. In Supplementary Table 2, the columns represent 11 physicochemical properties, and the rows represent 125 trinucleotides.

\[ x_{\text{PCP}_i} = N_i \times x_{\text{PCP}_{ij}} \]

where \( x = D \) refers to Dinucleotide and \( x = T \) denotes Trinucleotide. When \( x = D \), the values of \( i \) range from 1 to 25, the values of \( j \) range from 1 to 15, \( \text{DPCP}_i \) is the DPCP value of the \( i \)th dinucleotides, \( N_i \) is the count of the \( i \)th dinucleotides in the DNA sequence, and \( \text{DPCP}_{ij} \) is the \( j \)th properties of the \( i \)th dinucleotides; When \( x = T \), the values of \( i \) range from 1 to 125, the values of \( j \) range from 1 to 11, \( \text{TPCP}_i \) is the TPCP value of the \( i \)th trinucleotides, \( N_i \) is the count of the \( i \)th trinucleotides in the DNA sequence, and \( \text{TPCP}_{ij} \) is the \( j \)th properties of the \( i \)th trinucleotides.

**Position-Based Approach**

One-hot encoding method for mononucleotide (One-hot1) is similar to One-hot2, except that its encoding unit is the mononucleotide. It converts a mononucleotide into a one-hot code with a length of five, corresponding to five mononucleotides (A, C, G, T, and N). For instance, the encoded vector of DNA sequence \( D = \text{ACGTNA} \) is \((1,0,0,0,0)\) and \((0,1,0,0,0)\) since the position A is the only one with a value of 1.

**Classifier**

To train a classification model with stable and good performance, five machine learning algorithms was utilized to construct five base-classifiers. Subsequently, majority voting was adopted to integrate these five base-classifiers. Its detailed procedure is illustrated in the following steps.

1. The processed training dataset was inputted into five machine learning algorithms, and five base-classifiers were generated. These five algorithms were random forest (RF), multi-layer perceptor (MLP), stochastic gradient descent (SGD), linear discriminant analysis (LDA), extreme gradient boosting (XGB). Among them, RF refers to one of classifier that utilizes multiple decision trees to train and predict samples. MLP, as a simple neural network, contains three fully connected layers, the input layer, the hidden layer, and the output layer. SGD is a kind of support vector machine model. LDA is a classifier generated according to Bayes’ rule. XGB is also based on trees, but unlike random forests, its trees are regressive, and it also optimizes the algorithm itself, the efficiency and robustness of the algorithm.

2. The five base classifiers were combined into one ensemble classifier by majority voting. That is, when three or more base classifiers judge a sequence to be a positive (or negative) sample, then their combination also treats this sequence as a positive (or negative) sample.

It should be noted that the hyperparameters of the base-learners were optimized by grid search strategy. After manually specifying variation ranges of hyperparameters, this strategy adopted an exhaustive method-like approach to find the best-performing combination from these hyperparameters. In addition, all classifier algorithms in this paper were implemented by sklearn (Hinton, 1989; Belhumeur et al., 1997; Platt, 2000; Breiman, 2001; Bengio and Glorot, 2010; Pedregosa et al., 2011; Kingma and Ba, 2014; He et al., 2015; Chen and Guestrin, 2016).

**Performance Evaluation**

Our model was validated according to accuracy (ACC), Matthew correlation coefficient (MCC), Sensitivity (SN), Specificity (SP) which had been widely adopted in the field of bioinformatics (Huang and Gong, 2020; Liu et al., 2020; Smolarczyk et al., 2020; Wang H. et al., 2020; Wang J. et al., 2020; Shao and Liu, 2021; Zhang et al., 2021). These metrics can be calculated by equations (3) ~ (6).

\[ ACC = \frac{n_{TP} + n_{TN}}{n_{TP} + n_{TN} + n_{FP} + n_{FN}} \]
\[ MCC = \frac{n_{TP} \times n_{TN} - n_{FP} \times n_{FN}}{\sqrt{(n_{TP} + n_{FP}) \times (n_{TP} + n_{FN}) \times (n_{TN} + n_{FP}) \times (n_{TN} + n_{FN})}} \]
\[ SN = \frac{n_{TP}}{n_{TP} + n_{FP}} \]
\[ SP = \frac{n_{TN}}{n_{TN} + n_{FP}} \]

Where \( TP \) and \( TN \) refer to correctly predicted 6mA and non-6mA; \( FP \) and \( FN \) denote incorrectly predicted non-6mA and 6mA; \( n_x \) means the number of \( x \).

**RESULTS AND DISCUSSION**

**DNA Sequence Logos**

To find optimal features of samples, the DNA sequences of samples should be analyzed. Since these sequences were of equal length, they could be analyzed sequence logos (Schneider and Stephens, 1990). Two Sample Logo was employed (Vacic et al., 2006), which calculated the statistical difference between positive and negative samples at specific positions. The logo consists of three parts, the upper and lower parts represent the enriched and depleted nucleotides at specific positions, and the middle part denotes the consistent results of positive and negative samples. The x-axis indicates the position. The length of DNA sequences in our datasets is 41bp, so there are 41 scales on the x-axis. Additionally, as the middle nucleotide is consistent in both positive and negative samples, it is set to the 0th scale. The y-axis represents the amount of information at the position. The higher the symbol in a position, the more

\( \sqrt{ } \)
Figure 2 | Sequence logos of Rosaceae (A), Rice (B), and Arabidopsis (C).

Figures 2A-C are the sequence logos established for Rosaceae, Rice, and Arabidopsis. From the three figures, it can be seen that the sequences have a length of 41bp with “A”s at the center. In addition, “A” enriched at positions -6, -4, -3, 4, 7, 8, 10, 11, 12, “C” enriched at positions -7, -2, 2, 6, 9, “G” enriched at positions -8, -1, 2, 3, 5, 8, and “T” enriched at positions 3. Since these sequences containing 6mA are enriched with some nucleotides at some positions, it is speculated that position-based approaches are more suitable for extracting information from the sequences in our datasets.

Performance Evaluation of Models
To verify the conjecture in the previous section, six methods were chosen to extract the datasets as features and then they were applied to five commonly used well-performing algorithms in sklearn. Since the conjecture is too intuitive and may lead to some significant features being overlooked, not only nucleotide position-based methods are compared, but also density-based and physicochemical property-based methods were also compared.

The experimental results of 5-fold cross-validation are displayed in Table 2. The columns indicate the feature extraction methods which have been introduced in the “Feature Extraction” section. The rows denote classifier algorithms and their evaluation metrics, and they have been briefly described in the “Classifiers” section and the “Performance Evaluation” section.

As can be seen in Table 2, whichever classifier algorithm is selected, the ACCs, SNs, SPs, and MCCs of AMNF, ADNF, DPCP and TPCP are all lower than 0.80, 0.79, 0.83, and 0.60, whereas them of One-hot1 and One-hot2 are all higher than 0.93, 0.93, 0.91, and 0.86. These illustrate that compared with density-based and physicochemical property-based approaches, position-based ways can better express the characteristics contained in DNA sequences in our datasets. XGB performed slightly better with one-hot1 than...
one-hot2. This may be because XGB may lose some valuable information when it was applied on high-dimensional one-hot2 features. Specifically, XGB divides the high-dimensional feature space into many small parts which may be treated as noise. In addition, if the feature descriptor is One-hot1 or One-hot2, all classifiers show good performance, which indicates that all these algorithms are appropriate for this classification task.

Moreover, to judge intuitively whether the above six feature extraction methods were good at distinguishing between positive and negative samples, the tSNE (van der Maaten and Hinton, 2008) technique in sklearn (Pedregosa et al., 2011) was used to project the sample points of these methods from the high-dimensional space into a two-dimensional space. The visualization plots of the projection are shown in Figure 3. It can be seen from Figure 3 that the samples of the two labels are separated by certain dividing lines in Figures 3E, F, while in other subgraphs, the negative sample points are almost covered by the positive ones. These illustrate that One-hot1 and One-hot2 can better discriminate the sample points of the two labels in a high dimensional space than the other four methods.

Through these arguments, the nucleotide position-based methods are indeed more suitable for extracting features from DNA sequences in our datasets, and the assumptions that was made in the previous section are proved to be correct. Therefore, in the subsequent analysis, only One-hot1 and One-hot2 would be considered.

**Comparison of Features**

In the previous section, it has been learned that the position-based approaches express the information contained in our DNA sequences well. However, it is not sure which is the best among One-hot1, One-hot2, and their fusion. Therefore, in this subsection, they are compared. The comparison results are shown in Figure 4.

As can be seen from Figure 4, only when the classifier is XGB, the effect of the other two is slightly better than One-hot2; when the classifier is RF, LDA, MLP, or SGD, One-hot2 is significantly better than One-hot1 and slightly better than the fusion. The reason for this is that when encoding a dinucleotide, some information about the mononucleotide is involved. Therefore, in most cases, One-hot1 is not as good as One-hot2, and their fusion produces some redundant information. Consequently, One-hot2 is the best answer.

**Efficiency of Ensemble Strategy**

Using One-hot2 to extract features and take RF, LDA, MLP, SGD, and XGB as classifiers, five base models can be obtained. As shown in Figure 5, except for some differences between SN and SP of LDA and SGD, SN and SP for the other three classifiers do not differ much, as well as these base models are all with excellent performance, so they were tried to be combined with the majority voting strategy. The integrated results are also shown in Figure 5. It can be found that after voting, except for no enhancement in SP, all the other three metrics improved, which means that after this operation, the performance of the whole classification system has been risen to a higher level.

**Comparison With Other Machine Learning Models**

To evaluate the generalization capability and cross-species identification ability of our model, it was applied to three test datasets, DS2, DS3, and DS4. Moreover, the test results were compared with several other machine learning models to demonstrate the advantages of our model. Table 3 shows the comparative results on Rosaceae, Rice, Arabidopsis. The columns indicate four evaluation indicators that have been introduced in the "Performance Evaluation" section. The rows represent the species and the models applied on these species. The models include Meta-i6mA (Hasan et al., 2021), i6mA-Fuse (Hasan et al., 2020), i6mA-stack (Khanal et al., 2021), i6mA-Pred (Chen et al., 2019), iDNA6mA-Rice (Lv et al., 2019), MM-6mAPred (Pian et al., 2019), and 6mA-Finder (Xu et al., 2020). Among them, i6mA-Fuse consists of two modules, which were trained by the datasets of Fragaria Vesca and Rosa Chinensis, respectively. To
FIGURE 3 | The tSNE scatterplots of AMNF (A), ADNF (B), DPCP (C), TPCP (D), One-hot1 (E), and One-hot2 (F). (Blue and pink dots indicate DNA sequence samples with and without 6mA sites, respectively).

FIGURE 4 | Comparison before and after feature fusion.

FIGURE 5 | Effects of the ensemble strategy.

better distinguish them, i6mA-Fuse_FV and i6mA-Fuse_RC are used instead. The same situation is true for i6mA-stack.

As can be seen from Table 3, when the species is Rosaceae, although our SN and SP values only rank second, our ACC and MCC values are the maximum, suggesting that our model has the best overall performance in Rosaceae. It can be concluded that our model can make cross-species predictions for Rice as all four metrics of our model rank at the top. And it can better find 6mA sites from unknown Rice sequences because our model has the highest SN value. Like Rosaceae, our model predicts 6mA sites well in Arabidopsis, and with the highest SP, our model can better screen out those sequences that do not contain 6mA sites. Considering the comparative results on the three species, our model has better generalization performance and cross-species prediction ability than other methods. This may be because only the best-performing feature descriptor was selected to represent the DNA sequences rather than the fusion of several well-performing features. Thereby, the risk of generating
In the construction process, a hypothesis was put forward. Rosaceae, Rice, and Arabidopsis and achieved good results. In this study, a plant cross-species 6mA site recognition model was constructed by ensemble learning. It has been applied on Rosaceae, Rice, and Arabidopsis and tested on Rosaceae, Rice, and Arabidopsis. The experimental results showed that our model was adept at predicting the 6mA sites in homologous and heterologous species. In addition, it was also found that there might be a strong similarity between Rice sequences and Rosaceae positive sequences, and the similarity between Arabidopsis sequences and Rosaceae positive sequences is weak. The comparison with other models also showed the superiority of our model. In summary, i6mA-vote outperformed other concerned methods in predicting 6mA sites in the plant genomes. Meanwhile, our research also has the limitation that only three plants were considered. Therefore, future studies will focus on the 6mA site formation characteristics of more plants.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://github.com/zhaozhengnan/i6mA-vote/tree/master, github.

**AUTHOR CONTRIBUTIONS**

ZXT improved the model, designed experiments and drafted the manuscript. ZZ proposed the initial idea and implemented the experiments. YL prepared all datasets for experiments. ZT analyzed experimental results. MG revised the manuscript. QL conceived the whole research process and revised the manuscript. GW conceived designed experiments and revised the manuscript. All authors have read and approved the final manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.845835/full#supplementary-material
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