# WHISTLE: a high-accuracy map of the human *N*<sup>6</sup>-methyladenosine (m<sup>6</sup>A) epitranscriptome predicted using a machine learning approach

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# ABSTRACT

 $N^6$ -methyladenosine (m<sup>6</sup>A) is the most prevalent post-transcriptional modification in eukaryotes, and plays a pivotal role in various biological processes, such as splicing, RNA degradation and RNA-protein interaction. We report here a prediction framework WHISTLE for transcriptome-wide m<sup>6</sup>A RNAmethylation site prediction. When tested on six independent datasets, our approach, which integrated 35 additional genomic features besides the conventional sequence features, achieved a major improvement in the accuracy of m<sup>6</sup>A site prediction (average AUC: 0.948 and 0.880 under the full transcript or mature messenger RNA models, respectively) compared to the state-of-the-art computational approaches MethyRNA (AUC: 0.790 and 0.732) and SRAMP (AUC: 0.761 and 0.706). It also outperformed the existing epitranscriptome databases MeT-DB (AUC: 0.798 and 0.744) and RMBase (AUC: 0.786 and 0.736), which were built upon hundreds of epitranscriptome high-throughput sequencing samples. To probe the putative biological processes impacted by changes in an individual m<sup>6</sup>A site, a network-based approach was implemented according to the 'guilt-by-association' principle by integrating RNA methylation profiles, gene expression profiles and protein-protein interaction data. Finally, the WHISTLE web server was built to facilitate the query of our high-accuracy map of the human m<sup>6</sup>A

epitranscriptome, and the server is freely available at: www.xjtlu.edu.cn/biologicalsciences/whistle and http://whistle-epitranscriptome.com.

# INTRODUCTION

Large scale analysis has revealed the abundance of RNA modifications in the human epitranscriptome (1). With the recent advances in the exploration of RNA epigenetics, more than 150 types of RNA modifications have been identified (2). Among them, the most prevalent non-cap modification marker present on eukaryotic messenger RNA (mRNA) and long non-coding RNA,  $N^6$ -methyladenosine  $(m^{6}A)$ , (3) has emerged as an abundant and dynamically regulated modification (4). m<sup>6</sup>A was detected within poly-A RNA for the first time in 1974 (5), and has since been characterized in various eukaryotic species. In the past five decades, various studies have demonstrated the biological significance of m<sup>6</sup>A RNA methylation, which includes roles in the circadian clock (6), regulation of mRNA translation (7), heat shock response (8), microRNA (miRNA) processing (9), DNA damage response (10), RNA-protein interaction (11) and regulation of RNA stability (12). Consequently, the accurate identification of m<sup>6</sup>A locations is critical for the study and understanding of the downstream effects of RNA modification in biology.

To identify the precise location of  $m^6A$  sites on mRNA, the first whole-transcriptome  $m^6A$  profiling technique  $m^6A$ -seq (or MeRIP-seq) was introduced in 2012 (13,14), in which the  $m^6A$  containing RNA fragments is immunoprecipitated, purified and then subjected to further analysis. This technique applies high-throughput sequencing to

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the IP sample enriched with m<sup>6</sup>A-containing mRNA fragments. In contrast to the input control samples, it typically results in the detection of m<sup>6</sup>A containing peaks with around 100 nt resolution using MACS, the exomePeak R/Bioconductor Package and other peak callers (15,16). The precise location of the m<sup>6</sup>A sites may be further narrowed down to base-resolution by searching for the m<sup>6</sup>A motif RRACH within the peaks detected with m<sup>6</sup>A signal. Most existing epitranscriptome databases, such as, MeT-DB and RMBase, rely on this very simple strategy (17,18). A major limitation of this method is that it cannot differentiate between a randomly-occurring RRACH motif and a real m<sup>6</sup>A-containing motif located nearby, i.e. all the RRACH motifs located within an m<sup>6</sup>A peak will be reported as holding an m<sup>6</sup>A site, including the chance occurrences, resulting in false positive predictions. Since m<sup>6</sup>A-seq is currently the most widely used approach for profiling the transcriptome-wide m<sup>6</sup>A, and a very large number of the m<sup>6</sup>A-seq samples have been acquired in different studies, the m<sup>6</sup>A site information extracted from the m<sup>6</sup>A-seq using the motif search strategy essentially dominates the existing epitranscriptome databases. For this reason, it is not surprising that both MeT-DB (426 544 sites) and RMBase (477 452 sites) report a very large number of transcriptome m<sup>6</sup>A sites, many of which may be false positive due to a chance RRACH motif located close to a real m<sup>6</sup>A site (or within an m<sup>6</sup>A peak).

Besides the m<sup>6</sup>A-seq technique, single-based resolution techniques such as the miCLIP (19) and m<sup>6</sup>A-CLIP (20)were also developed. However, these experiments are usually more laborious to perform but still offer limited coverage of the m<sup>6</sup>A epitranscriptome, since the reported RNAmethylation sites are still restricted to the transcripts more readily expressed under a specific cell/tissue condition. Although base-resolution profiling techniques have not been very widely applied in biological studies due to their expense and difficulties, they provide the ground truth of m<sup>6</sup>A site information that is necessary for computational prediction. To date, a large number of RNA-methylation site prediction methods and web servers have been developed based on the information extracted from base-resolution techniques since 2015, including the pseudo nucleotide composition based approach iRNA-Methyl by Chen et al. (21) and physical-chemical properties-based approach pRNAm-PC (22). Subsequently, Zhou et al. employed SRAMP, a random forest machine learning framework, to predict mammalian m<sup>6</sup>A sites using sequence features (23). Many other site predictors have been developed for m<sup>6</sup>A and other RNA modification, such as MethyRNA (24), RNAMethPre (25), RAM-NPPS (26), Target M6A (27), AthMethPre (28), iRNA-PseColl (29), M6APred-EL (30), iMethyl-STTNC (31), iRNA-PseDNC (32), etc. (33–39). These methods have been recently reviewed (40). These site predictors usually take the transcript sequence as the input and report a number of possible m<sup>6</sup>A sites as the output, making them very convenient to use. However, to our knowledge, they are exclusively based on the sequence-derived information-even when the secondary structure or other high level features (41) are used, the information is still directly extracted from sequence without considering other potentially useful genomic features, such as, conservation, transcript type and gene annotation. Although the sequence information probably plays a central role, other genomic features may also be helpful in the prediction of  $m^6A$  sites and thus should be incorporated in the analysis. Additionally, although potentially feasible, none of these approaches have been applied transcriptome-wide to reconstruct the entire  $m^6A$  epitranscriptome, thus limiting their usage in large-scale or high-throughput analysis.

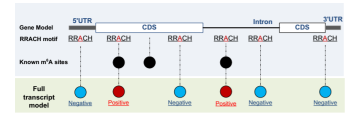
In this project, we proposed a prediction framework, WHISTLE, which stands for <u>who</u>le-transcriptome m<sup>6</sup>A <u>site</u> prediction from multiple <u>genomic</u> features. The framework extracted a comprehensive set of domain knowledge based on various genomic features, and integrated them with conventional sequence-derived features for reconstructing a high-accuracy map of the m<sup>6</sup>A epitranscriptome. The 'guiltby-association' principle was then applied to further annotate the functional relevance of each individual RNAmethylation site by integrating gene expression profiles, RNA methylation profiles and PPI networks.

#### MATERIALS AND METHODS

#### Training and testing data for m<sup>6</sup>A site prediction

The data used for training and benchmarking in m<sup>6</sup>A site prediction includes six single-base resolution m<sup>6</sup>A experiment obtained from five cell types (see Table 1). The baseresolution m<sup>6</sup>A sites in each experiment were downloaded directly from Gene Expression Omnibus (GEO). The two samples (MOLM13 mi-CLIP sample and the A549 m<sup>6</sup>A-CLIP) reported based on the human genome assembly hg18 were lifted using UCSC liftOver tool (https://genome.ucsc. edu/cgi-bin/hgLiftOver). A total of 20 516 and 17 383 m6A sites out of the original 23 480 and 19 683 sites were lifted to hg19, respectively. Both samples have very large number of (>17000) positive sites that can be used for training and testing after liftOver, and the majority (four out of six) baseresolution samples are based on hg19 and thus do not require extra processing step.

In the beginning of the performance evaluation procedure, dataset 6 of the base-resolution data (Table 1) was used as the independent testing data, while the other five datasets were used as the training data. The positive training data (m<sup>6</sup>A sites) was determined as the m<sup>6</sup>A sites under RRACH consensus motifs that have been reproduced in at least two of the five training datasets. The negative training data (non-m<sup>6</sup>A sites) was randomly selected from the nonpositive RRACH adenosines on the full transcripts containing the positive sites (see Figure 1). Initially, the number of randomly selected negative sites was ten times the number of positive sites. Later, the positive-negative ratio was balanced by randomly splitting the negative samples into 10 random subsets. Consequently, 10 training datasets, each with 1:1 positive-to-negative ratio, were constructed using different negative samples. The negative data was also generated similarly on testing data (Dataset 6), i.e. the negative data were randomly selected non-positive m<sup>6</sup>A sites from the m<sup>6</sup>A containing transcripts. The ratio of positive testing data to negative testing data was also kept as 1:10. The testing performances from the 10 independent sessions were averaged.



**Figure 1.** Generation of positive and negative data. The transcriptome  $m^6A$  sites under RRACH consensus motifs that have been reproduced in at least two of the five training datasets were used as positive  $m^6A$  sites. The negative training data (non- $m^6A$  sites) was randomly selected from the non-positive RRACH adenosines on the full transcripts containing the positive sites.

To exclude randomness of testing dataset from the  $m^6A$  site prediction evaluation, we also applied dataset level leave-one out validation over the six base-resolution datasets. In each round of the dataset level validation, one of the six base-resolution datasets was used as the independent testing data, while the remaining five datasets were used as the training data. The same rules of the training and testing data generation were applied as previously described in each individual test. As the training and testing data were all extracted from different independent experiments, there should be no overfitting problem.

### Features for m<sup>6</sup>A site prediction

Sequence-derived features. The sequence-based information around the RRACH motif was encoded using the same method of  $m^6$ Apred (44) and MethyRNA (24), which have been shown to be quite effective and achieved good performance in human and yeast m<sup>6</sup>A site prediction. The sequence feature encodes the nucleotides sequence by three distinct structural chemical properties: ring structures, functional groups and hydrogen bonds. Specifically, adenine and guanine have two ring structures, while cytosine and uracil have only one ring; adenine and cytosine contain the amino group, while guanine and uracil contain the keto group; adenine and uracil can form two hydrogen bonds during hybridization, whereas guanine and cytosine can form three hydrogen bonds. Based on the three structural chemical properties defined above, the *i*-th nucleotide from sequence S can be encoded by a vector  $S_i = (x_i, y_i, z_i)$ :

$$x_{i} = \begin{cases} \lim_{i \in \{A,G\}} \\ \lim_{i \in \{C,U\}} \\ i \in \{C,U\} \end{cases}, y_{i} = \begin{cases} \lim_{i \in \{A,C\}} \\ \lim_{i \in \{G,U\}} \\ i \in \{G,U\} \\ i \in \{C,G\} \end{cases}, z_{i} = \begin{cases} \lim_{i \in \{A,U\}} \\ \lim_{i \in \{C,G\}} \\ i \in \{C,G\} \\ i \in \{C,G$$

Therefore, the A, C, G, U can be encoded as a vector of three features (1,1,1), (0,1,0), (1,0,0) and (0,0,1), respectively. Additionally, a feature of the cumulative nucleotide frequency is calculated for each nucleotide position in the sequence. The density of the *i*-th nucleotide  $d_i$  is defined as the sum of all the instances of the *i*-th nucleotide before the i + 1 position. The nucleotide frequency  $f_i$  is defined by the following formula:  $f_i = d_i/i$ . Using the sequence 'AUG-GACACU' as an example, the cumulative frequency for adenine is 1.00 (1/1), 0.40 (2/5) and 0.43(3/7) at the first, fifth and seven position, respectively; while the frequency for uracil is 0.50 (1/2) and 0.11 (1/9) at the second and ninth respective position.

Genome-derived features. Most existing RNA modification site prediction algorithms use exclusively sequencebased features; however, such features alone may not fully capture the attributes of RNA modification topology. Hence, we generated 35 additional genomic features that may contribute to the prediction. Genomic Features 1–13 are dummy variable features indicating whether the adenosine sites shall fall within the transcript regions that satisfy certain topological properties. All the features in this category are generated by the GenomicFeatures R/Bioconductor package (45) using the transcript annotations hg19 TxDb package. To remove the ambiguity caused by transcript isoforms, only the primary (longest) transcripts of each gene were kept for the extraction of the transcript sub-regions. Genomic Features 14-16 are real valued features defining the relative position of the transcript regions (3'UTR, 5'UTR and whole transcript), i.e. the distance from the adenine to the 5' end divided by the width of the region. The values are also set to zero for sites that do not belong to the region. Genomic features 17–19 represent the length of the transcript region containing the modification site. The values are also set to zero for sites that not belong to the region. Features 20–22 capture the distance from the adenine sites to the 5'end or 3'end of the splicing junctions. Additionally, the distance to the nearest neighboring m<sup>6</sup>A sites in the training data is generated to measure the clustering effect of the m<sup>6</sup>A RNA modification sites. Features 23-26 represent the evolutionary conservation score of the adenosine sites and its flanking regions; two metrics of nucleotide conservation, Phast-Cons score (46) and the fitness consequence scores are used to measure the conservation level of the underlying nucleotide sequence. Features 27 and 28 represent the RNA secondary structures around the adenine site, the RNA secondary structures are predicted using RNAfold from the Vienna RNA package (47). Finally, features 29–35 are the properties of the genes or transcripts containing the m<sup>6</sup>A sites, such as being the miRNA target genes or housekeeping genes. The annotation of miRNA target sites are from miRanda (48) and TargetScan (49). Supplementary Table S1 contains more details about the genomic features we considered in the prediction.

#### Machine learning approach used for m<sup>6</sup>A site prediction

The Support Vector Machine (SVM) is one of the most widely used machine learning algorithms in computational biology. It was previously used for mammalian miRNA target prediction (50), protein kinase-specific phosphorylation sites prediction (51) and mammalian m<sup>6</sup>A modification sites prediction (24,25). In this project, we used an R language interface of LIBSVM (52) to construct the SVMbased m<sup>6</sup>A site predictors. Following previous approaches (21,22), the radial basis function was chosen as the kernel function, and the other parameters were set at the default. Random Forest is another popular machine learning algorithm applied in biology data, and one of the earliest mammalian m<sup>6</sup>A site predictor SRAMP was developed based on the Random Forest approach (23). In this project, we also use Random Forest from the R package randomforest (53) to compare the predictive performance using SVM.

## Performance evaluation of m<sup>6</sup>A site prediction

For both the SVM and random forest classifiers, a 5fold cross-validation was employed on the training datasets for model selection purpose, and the final performance of the predictor was measured on the independent testing dataset. The receiver operating characteristic curve (sensitivity against 1-specificity) was used to measure the prediction performance under different decision thresholds, and the area under the curve (AUC) was calculated as the main performance evaluation metric.

When evaluating the accuracy of  $m^6A$  site information stored in existing epitranscriptome  $m^6A$  site databases MeT-DB Version 2 and RMBase Version 2, the reliability was determined by the number of experiments that support the existence of a specific  $m^6A$  site, based on which the AUC can be calculated. In addition, the sensitivity (*Sn*), specificity (*Sp*) and Matthews correlation coefficient (*MCC*) were calculated to measure the performance of predictor:

$$Sn = \frac{TP}{TP + FN} \tag{2}$$

$$Sp = \frac{TN}{TN + FP} \tag{3}$$

$$MCC = \frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP) \times (TP + FN) \times (TN + FP) \times (TN + FN)}}, (4)$$

where, *TP*, *TN*, *FP* and *FN* represent true positive, true negative, false positive and false negative, respectively. When different methods were compared under AUC, they always use the same positive and negative gold standard dataset, and AUCs were always calculated in the same way. The AUCs of different methods reported in our manuscript are therefore strictly comparable.

#### Estimate the posterior probability of RNA methylation

The existing machine learning approaches usually report the probability of an m<sup>6</sup>A motif to be an actual methylation site under the assumption of equal prior probability, i.e. the prior probability of an m<sup>6</sup>A motif being an m<sup>6</sup>A site is 0.5. However, it is known in practice that the number of m<sup>6</sup>A sites is a lot smaller than the number of m<sup>6</sup>A motifs, so the number of RNA-methylation sites under a specific experimental condition is likely to be significantly over-estimated. To address this bias, a posterior probability of RNA methylation under a specific condition is calculated with:  $q_{M,i} = \frac{\pi_M p_{M,i}}{(1-\pi_M)(1-p_{M,i})+\pi_M p_{M,i}}$ , where,  $\pi_M$  is the prior probability that a transcriptome RRACH motif embraces a true m<sup>6</sup>A site under a specific model, which is calculated empirically from the 6 base-resolution datasets (see Table 1) as the average number of m<sup>6</sup>A sites under a condition divided by the number of occurrences of transcriptome RRACH motifs that are supported by at least one m<sup>6</sup>A record in MeTDB for the mature mRNA model, or RM-Base for the full transcript model. These is also the search space of our predicted m<sup>6</sup>A epitranscriptome.  $p_{M,i}$  is the predicted probability (or likelihood) of the *i*-th site being a real m<sup>6</sup>A site under a specific model M, and  $(1 - p_{M,i})$  is

ID	Cell	Note	Technique	Source
1 2 3 4 5 6	HEK293 HEK293 MOLM13 A549 CD8T HeLa	abacm antibody sysy antibody	mi-CLIP m <sup>6</sup> A-CLIP	(19) (19) (42) (20) (20) (43)

the probability of the opposite being true.  $q_{M,i}$  is a posteriori probability of the *i*-th site being a real m<sup>6</sup>A site under a specific condition. The posterior probability  $q_{M,i}$  is also reported in the WHISTLE database along with the probability  $p_{M,i}$ .

#### **RESULTS AND DISCUSSION**

#### m<sup>6</sup>A site prediction

The predictors on the full-transcript data were established first in which the true  $m^6A$  site and negative sites may be located in both exonic and intronic regions. Because experimental procedures, especially the polyA selection step, may induce bias toward mRNA, we also consider a mature mRNA model, under which, the goal is to predict only exonic  $m^6A$  sites, and thus only the exonic regions are considered.

We show in Supplementary Table S3 that, although the genome-derived features alone are already very effective for predicting m<sup>6</sup>A sites, the best performance is achieved when the sequence features and genomic features are combined. Consequently, our m<sup>6</sup>A site predictor was established based on both the genome-derived features and sequence-derived features.

Feature selection was performed to identify the most effective genomic features for m<sup>6</sup>A site prediction. Here, datasets 1-5 were used as the training data, while dataset 6 was used as the independent testing data. The relative importance of each genome-derived feature in the prediction was firstly assessed with the Perturb method (54) using the R caret package. Next, the N most important features were retained in the prediction analysis, and the prediction performance was evaluated using a 5-fold cross-validation. As shown in Supplementary Figure S1A, the predictor performance under the full transcript model stops increasing after including the top 14 most important genomic features. The top three most critical genomic features under this model are long exon, miRNA target and conservation score. To achieve the most robust performance and to avoid potential overfitting, only the top 14 genomic features were used in the full transcript model for m<sup>6</sup>A site prediction purpose in later analysis. Similarly, the top 19 genome-derived features with the highest importance were selected for the mature mRNA model (see Supplementary Figure S1B). The distance to known m<sup>6</sup>A sites became the most important predictive feature, which demonstrated the clustering effect of m<sup>6</sup>A modification, followed by long exon and conservation under the mature mRNA model.

The performance of the proposed m<sup>6</sup>A predictors was then evaluated using independent datasets and compared

				Performance	e on independent data	set (AUC)		
Model	Method	A549	CD8T	Hela	HEK293 (sysy)	HEK293 (abacm)	MOLM13	Average AUC
Full Transcript	WHISTLE MethyRNA* SRAMP	0.965 0.807 0.856	0.930 0.800 0.841	0.953 0.741 0.762	0.936 0.848 0.883	0.968 0.778 0.838	0.933 0.765 0.759	0.948 0.790 0.761 <sup>#</sup>
Mature mRNA	WHISTLE MethyRNA SRAMP	0.830 0.903 0.751 0.814	0.904 0.734 0.796	0.702 0.894 0.676 0.702	0.885 0.936 0.848 0.869	0.838 0.818 0.698 0.796	0.739 0.823 0.686 0.710	0.701 0.880 0.732 0.706 <sup>#</sup>

Table 2. Performance evaluation of m<sup>6</sup>A site prediction methods

Note: \*The MethyRNA approach uses sequence-derived features with SVM (24), which we reproduced faithfully with the same training data of WHISTLE for comparison.

<sup>#</sup>The SRAMP method was originally trained on A549, CD8T, HEK293 (sysy) and HEK293 (abacm). To avoid overfitting, only Hela and MOLM13 were considered when evaluating its average performance.

Only the  $m^6A$  sites not previously used as training data were considered during performance evaluation, so the training sites and testing sites have no overlap. Please see Supplementary Table S4 for the results when all sites from the independent testing samples were considered.

with competing approaches (Table 2). By combining additional genome-derived features, the performance of our approach was substantially higher in all the tested conditions than MethyRNA and SRAMP, which rely only on information extracted from sequences. WHISTLE achieved AUCs of 0.948 and 0.880 under the full transcript and mature mRNA modes, respectively, representing a major improvement compared to MethyRNA (0.790 and 0.732) and SRAMP (0.761 and 0.706).

## A predicted map of human m<sup>6</sup>A epitranscriptome

With the extensive study of RNA epigenetics, especially the accumulation of large number of m<sup>6</sup>A-seq datasets, the transcriptome-wide distribution of m<sup>6</sup>A sites have been summarized and made available from bioinformatics databases, such as MeT-DB (55) and RMBase (56). MeT-DB is the first transcriptome m<sup>6</sup>A database that provides condition-specific distribution of m<sup>6</sup>A RNA methylation in human and mouse initially, and later in other species as well; while RMBase is a more comprehensive RNA modification database, supporting more species and more RNA modification types. However, as these two databases overwhelmingly rely on m<sup>6</sup>A-seq data, and implemented a data processing pipeline that could not differentiate between true and randomly-occurring m<sup>6</sup>A motif located in close proximity within an m<sup>6</sup>A peak, the information they provide may not be accurate and should be re-assessed.

The reliability of a specific  $m^6A$  site in epitranscriptome databases has been measured by the number of experiments that support the record. This metric will be used when evaluating the accuracy of the two databases. Interestingly, as shown in Table 3, when comparing the two epitranscriptome databases, the exomePeak-based MeT-DB database is slightly more accurate than primarily the MACS-based RMBase database. However, even with hundreds of highthroughput sequencing datasets accumulated, existing epitranscriptome databases are still far less accurate than what we may achieve with machine learning approaches (see Table 2).

We thus performed a whole transcriptome prediction of  $m^6A$  RNA-methylation sites in human to generate a map of human  $m^6A$  epitranscriptome using our proposed WHISTLE approach. Our predicted map is of substantially higher accuracy (average AUC of 0.948 and 0.880) compared with existing epitranscriptome databases MeT-DB (average AUC of 0.798 and 0.744) and RMBase (average AUC of 0.786 and 0.736) when evaluated on independent base-resolution datasets under both full transcript and mature mRNA mode, respectively. Additionally, we calculated a posterior probability of RNA-methylation site under a specific experimental condition. This provided a more empirical evaluation of the methylation status by taking into consideration the prior probability of an m<sup>6</sup>A motif being an m<sup>6</sup>A site, which is estimated from the base-resolution datasets.

Besides CLIP-based approaches, we also tested the accuracy of the proposed method on a high resolution  $m^6A$ seq dataset (57). Although still antibody-based, this  $m^6A$ seq dataset was generated from an improved protocol and achieved near base resolution (58). As shown in Table 4, when antibody-based  $m^6A$ -seq technique is used as the ground truth, WHISTLE still substantially outperformed competing approaches under both the full transcript and mature mRNA models.

## Website interface

An online database has been built to host the predicted human  $m^6A$  epitranscriptome. The individual RNA-methylation sites were then functionally annotated with gene expression data, RNA methylation data and protein–protein interaction data according to the 'guilt-byassociation' principle (detailed in the Supplementary File S2). As is shown in Figure 2, The website supports queries that may be a methylation site, a gene or a specific biological function under the Gene Ontology framework (59). It also supports the download of the original base-resolution datasets (Table 1) used for site prediction and the entire predicted epitranscriptome map with the functional annotations for large-scale analysis.

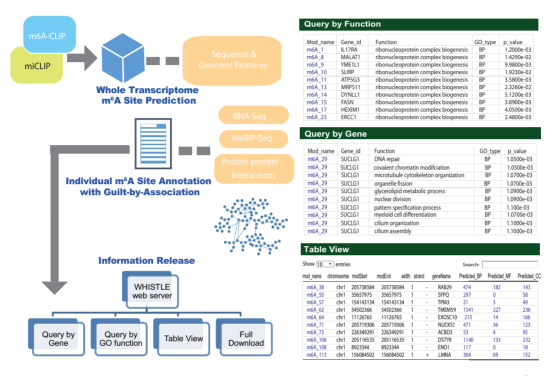
# CONCLUSIONS

Along with recent advances in RNA epigenetics, especially, the development of new techniques for profiling the RNA methylome (60,61), computationally deciphering the epitranscriptome from various omic data presents a major

Table 3.	Performance eva	luation for	bioinformatics	databases
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			Group truth dataset used (AUC)					
Mode	Method	A549	CD8T	Hela	HEK293 (sysy)	HEK293 (abacm)	MOLM13	Average AUC
Full Transcript	RMBase	0.825	0.788	0.832	0.837	0.701	0.733	0.786
	MetDB	0.835	0.802	0.843	0.848	0.719	0.744	0.798
Mature mRNA	RMBase	0.768	0.716	0.790	0.752	0.707	0.682	0.736
	MetDB	0.775	0.730	0.795	0.762	0.716	0.683	0.744

Note: To ensure the results are comparable to Table 2, only the unique sites not previously reported in the training data of predictors were considered. Please see Supplementary Table S4 for the results when all sites from the independent testing samples were considered.



**Figure 2.** WHISTLE website. The WHISTLE website hosts a functionally annotated high-accuracy predicted map of the human  $m^6A$  epitranscriptome. The WHISTLE website supports direct query of RNA-methylation sites with respect to a specific GO function or gene. The  $m^6A$  RNA-methylation sites were predicted from m6A-CLIP data, miCLIP data, sequence features and genome-derived features. And then, the most dynamic RNA-methylation sites were annotated under the Gene Ontology framework using the guilt-by-association principle by integrating gene expression, RNA methylation and protein–protein interaction data. Please Supplementary Figure S2 for the complete data processing pipeline of WHISTLE.

Table 4. Per	formance assessment	using high	resolution m <sup>6</sup>	A-seq data
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	AUC under full transcript model	AUC under mature mRNA model
WHISTLE	0.980	0.904
MethyRNA	0.904	0.826
SRAMP	0.825	0.783
RMBase	0.774	0.758
MeTDB	0.775	0.767

**Note:** The high confidence consensus m6A sites detected in more than two of the total six high resolution m6A-seq experiments (57) were considered. Similar as before, MethyRNA and WHISTLE used the same m6A datasets for training. Only the unique sites not previously reported in the training data were considered here. Please see Supplementary Table S5 for the results when all sites from the independent testing samples were considered.

challenge to the bioinformatics community. In the past few years, sequence-derived features have been widely used for the prediction of RNA modification sites in human (24), mouse (24), other mammals (23,25), yeast (30,62) and other species; and a few major bioinformatics databases, including MeT-DB (18), RMBase (17), m6AVar (63), MOD-OMICS (64) and RNAMDB (65) have been built. These databases address various aspects of the RNA modifications including transcriptome-wide distribution, mechanism pathway, relevance to miRNA and RNA-binding proteins, functional variants, etc., and have greatly benefited researchers in this field.

Here, we constructed a functionally annotated highaccuracy predicted map of human m<sup>6</sup>A epitranscriptome and named it WHISTLE. The most stringent validation strategy was implemented, in which the performance of WHISTLE was assessed on six independent datasets (Tables 2 and 3) and on dataset generated from a different technique (Table 4). By integrating 35 genome-derived features with the conventional sequence-derived features, WHIS-TLE achieved a substantial improvement in accuracy, under both the full transcript model and the mature mRNA model, compared with existing machine learning-based  $m^6A$  predictors and the latest epitranscriptome databases.

It is worth noting that, the prediction performance achieved on the full transcript model (AUC: 0.948) may be significantly over-estimated due to the library preparation (polyA selection) of the miCLIP and m<sup>6</sup>A-CLIP samples used, because they cannot effectively capture the intronic m<sup>6</sup>A sites. The performance achieved on the mature mRNA model (AUC: 0.880) is probably a more realistic estimate.

A web server WHISTLE was built to enable the direct query of predicted RNA-methylation sites, their putative functions and their potential association to other methylation sites or genes, which provides the requisite data for the further epitranscriptome studies in human.

Our work has provided a computational scheme to study the  $m^6A$  epitranscriptome based on multi-omics datasets using machine learning and network-based method. In the future, it can be easily expanded to the study of other RNA modifications, such as  $m^1A$  (66) and Pseudouridine (67), as well as in other species, such as mouse and yeast.

## SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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Author contributions: J.M., R.R., Z.L. and J.P.M. conceived the idea and designed the research; Z.W. constructed the genomic features considered in m<sup>6</sup>A site prediction and processed the raw data; K.C. performed the m<sup>6</sup>A site prediction; Q.Z. and X.W. performed the network-based functional annotation of individual m<sup>6</sup>A sites; K.C. built the website; K.C., Q.Z. and W.Z. drafted the manuscript. All authors read, critically revised and approved the final manuscript. We thank Zoya Farooq at University of Liverpool for her assistance in website building.

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