MoAIMS: efficient software for detection of enriched regions of MeRIP-Seq

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Abstract

Background: Methylated RNA immunoprecipitation sequencing (MeRIP-Seq) is a popular sequencing method for studying RNA modifications and, in particular, for N6-methyladenosine (m6A), the most abundant RNA methylation modification found in various species. The detection of enriched regions is a main challenge of MeRIP-Seq analysis, however current tools either require a long time or do not fully utilize features of RNA sequencing such as strand information which could cause ambiguous calling. On the other hand, with more attention on the treatment experiments of MeRIP-Seq, biologists need intuitive evaluation on the treatment effect from comparison. Therefore, efficient and user-friendly software that can solve these tasks must be developed.

Results: We developed a software named “model-based analysis and inference of MeRIP-Seq (MoAIMS)” to detect enriched regions of MeRIP-Seq and infer signal proportion based on a mixture negative-binomial model. MoAIMS is designed for transcriptome immunoprecipitation sequencing experiments; therefore, it is compatible with different RNA sequencing protocols. MoAIMS offers excellent processing speed and competitive performance when compared with other tools. When MoAIMS is applied to studies of m6A, the detected enriched regions contain known biological features of m6A. Furthermore, signal proportion inferred from MoAIMS for m6A treatment datasets (perturbation of m6A methyltransferases) showed a decreasing trend that is consistent with experimental observations, suggesting that the signal proportion can be used as an intuitive indicator of treatment effect.

Conclusions: MoAIMS is efficient and easy-to-use software implemented in R. MoAIMS can not only detect enriched regions of MeRIP-Seq efficiently but also provide intuitive evaluation on treatment effect for MeRIP-Seq treatment datasets.

Keywords: RNA modification, N6-methyladenosine, Negative binomial model, Treatment effect

Background

RNA modification, represented by the epitranscriptome [1], refers to biochemical modifications of RNAs that are involved in functional regulations such as translation efficiency and mRNA stability without a change in the RNA sequence. Over 100 types of RNA modifications have been reported [2]. Among them, researchers have recently focused on certain abundant modifications such as N6-methyladenosine (m6A) [3], N1-methyladenosine (m1A) [4], and 5-methylcytidine (m5C) [5].

With the fast growth of next-generation sequencing (NGS), scientists can study RNA modifications at a whole-transcriptome scale. Methylated RNA immunoprecipitation sequencing (MeRIP-Seq) is a type of NGS technology for studying RNA modifications and is particularly widely used to detect m6A, a modification found in various species including human, mouse, and zebrafish [6–8]. In MeRIP-Seq, an antibody specific to a certain type of RNA
modification (such as m6A or m1A) is used to immunoprecipitate RNA; it is similar to another popular sequencing technology, i.e., ChIP-Seq (Chromatin immunoprecipitation sequencing) [9], which is used in studies of transcription factor binding. However, based on the inherent features of DNA and RNA, there is some difference between MeRIP-Seq and ChIP-Seq data. First, the distribution of ChIP-Seq read counts is relatively uniform while that of MeRIP-Seq is more variable owing to transcript abundance so that MeRIP-Seq requires an input RNA-Seq sample as a control. Second, transcript abundance affects the duplication rate, which must be considered in preprocessing MeRIP-Seq data. Third, because RNA sequencing can store strand information, which provides more accurate transcriptome profiling by strand-specific sequencing can store strand information, which provides a mixture negative-binomial (NB) model. It is compatible with different RNA sequencing protocols including paired/single-end and non-strand/strand-specific sequencing. Our results demonstrated the excellent processing speed (it only takes several minutes to finish analysis of one dataset) and competitive performance of MoAIMS compared with other tools. When MoAIMS is applied to studies of m6A, the detected enriched regions contain known biological features of m6A. Furthermore, MoAIMS can provide an intuitive indicator of treatment effect for treatment experiments. The signal proportion inferred from MoAIMS for m6A treatment datasets (perturbation of m6A methyltransferases) showed a decreasing trend, consistent with experimental observations. Finally, functional analysis on the m6A perturbation datasets reveals the interplay between m6A and histone modification. In conclusion, we developed efficient and user-friendly software for MeRIP-seq analysis.

Implementation

A MeRIP-Seq dataset consists of one immunoprecipitation (IP) sample and one input sample (used as control). MoAIMS takes aligned IP and input bams as input. Aligned bams are generated from pre-processing as shown in the workflow of MeRIP-Seq analysis (Fig. 1). In the pre-processing, reads are aligned to a target genome by transcriptome-based aligners such as STAR [15], Tophat [16], and HISAT [17]. Only uniquely mapped reads are kept. Then, reads are sorted and marked for duplication using PicardTools [18] or samtools [19]. Given the RNA sequencing protocol (single-end or paired-end, strand-specific or not) and a target genome annotation (in GTF format), MoAIMS is ready for analysis. Typically, MoAIMS requires several minutes to complete the analysis of one MeRIP-Seq dataset. The primary outputs of MoAIMS contain enriched regions (in BED12 format), goodness of fitting (GOF) plot (Fig. 2), and a summary table of the fitted models (Table 1). The source code and the user’s manual are available at https://github.com/rreybeyb/MoAIMS.

In the analysis performed by MoAIMS, it firstly obtains transcriptome bins by concatenating all exons for the expressed genes. Then, it uses featureCounts for counting reads in the bins. Subsequently, it models the distribution of the bin counts by a mixture NB distribution and detects the enriched regions. The details are described as follows.

Read counts of bins

Counting reads in bins was performed for the transcriptome of expressed genes because unexpressed genes provide little information for signal detection. The default threshold for expressed genes is 0.5 TPM (transcripts per million). All exons for the expressed genes were concatenated and split into bins with size 200 bp (default setting). Subsequently, featureCounts [20] was used for counting reads in the bins. The parameters used in featureCounts include the following: requireBothEndsMapped=TRUE (for paired-end sequencing), read2pos=5, ignoreDup=T, allowMultiOverlap=T.

Model construction

A negative-binomial mixture model

Our software implements and extends the statistical framework proposed by MOSAiCS [21], which is used to detect ChIP-Seq enriched regions and cannot be directly applied to MeRIP-Seq data because it is designed for processing DNA Sequencing and models the bin counts
on the whole-genome scale. The statistical framework assumes that the observed bin counts of an IP sample follows a mixture negative-binomial model composed of a background component and a signal component that are unobserved. Let \( Z \) represent the components, where \( Z \in \{0, 1\} \) (0 for the background component and 1 for the signal component) and \( Y_j \) is the observed read count of the \( j \)th bin; therefore, the mixture model can be written as Equation (1),

\[
P(Y_j) = (1 - \pi_s)P(Y_j|Z_j = 0, \Theta_B) + \pi_s P(Y_j|Z_j = 1, \Theta_S),
\]

of which \( \pi_s \) is the signal proportion (\( \pi_s \in [0, 1] \)), equal to \( \pi(Z_j = 1) \), and \( (1 - \pi_s) \) is equal to \( \pi(Z_j = 0) \); \( \Theta_B \) and \( \Theta_S \) are parameters of background and signal distribution respectively.

When the bin is from the background component, the read count follows the distribution \( NB(a, \frac{a}{a+\mu_j}) \), with \( a \) the size parameter and \( \frac{a}{a+\mu_j} \) the probability parameter of the NB distribution. When the bin is from the signal component, the read count can be represented as \( Y_j = N_j + S_j + k \) (one-signal, named 1S mode), where \( N_j \) is the count from a non-specific background following \( NB(a, \frac{a}{a+\mu_j}) \), \( S_j \) is the count from an actual enrichment following \( NB(b_j, \frac{b_j}{c+1}) \) (\( c = \frac{b_j}{\mu} \), \( \mu \) is the mean), and \( k \) is the minimal read count required for the signal component. Thus, the distribution of the signal component is a convolution of negative binomials. Details of the distributions are provided in the Supplementary. Additionally, our software implements the mixture NB model of the signal component (two-signal, named 2S mode) from MOSAiCS considering the complexity of the signal component, where \( S_j \) is the count following the distribution \( \pi_{s1}NB(b_1, \frac{b_1}{c+1}) + (1 - \pi_{s1})NB(b_2, \frac{b_2}{c+1}) \), with \( \pi_{s1} (\pi_{s1} \in [0, 1]) \) the first signal proportion.

In summary, the parameters of NB to be estimated in the model are represented as \( \Theta = \{\Theta_B, \Theta_{S1}, \Theta_{S2}\} \), where \( \Theta_B = (a, \mu_j) \) for the background component, \( \Theta_{S1} = (b, c) \) for the signal component in 1S mode and \( \Theta_{S2} = (b_1, c_1, b_2, c_2) \) for the signal component in 2S mode.

**Parameters estimation**

First, we estimated the parameters of the background component, \( \Theta_B = (a, \mu_j) \). \( \mu_j \) is estimated by regression using the input bin count data. A simple illustrative figure for the regression process is shown in Figure S1. The detailed explanation is described as follows.

Each IP bin count \( Y_j \) has a corresponding input bin count \( X_i \). For the bins from the background component, it is assumed that \( \{Y_j\}_{i=1}^{T} \) with the same input bin count from the same distribution; thus, \( \{Y_j\} \) are grouped by the input bin count to \( S_i = \{Y_j|X_j = x_i\} \) \( x_i \) is the group value equal to available and unique bin count value, i.e. 0, 1, 2, ..., for input sample and \( i \) is the group index. For \( Y_j \in S_i \) it follows that \( NB(a, \frac{a}{a+\mu_j}) \). Subsequently, regression is performed with \( x_i \) as the predictor variable and \( \mu_j \) (equal to \( E(S_i) \), the median value of \( Y_j \in S_i \)) as the
Fig. 2 Examples of goodness of fitting (GOF) plots for a human and a mouse dataset. X-axis is bin count and Y-axis is frequency. Real data, simulation data of 1S (one-signal) mode, and simulation data of 2S (two-signal) mode, are plotted in black, red, and blue lines, respectively.

response variable. MOSAICS uses the weighted robust fitting of linear model (RLM) [22] for regression with the function \( \log(\mu_i) = \beta_0 + \beta_1 \log(x_i) \), of which \( \beta_0 \) and \( \beta_1 \) are the coefficients. However, in some cases of RNA sequencing, we found that the generalized additive model (GAM) [23] can provide better fitting as shown in Figure S1. GAM uses a sum of unspecified smooth functions \( \sum_{s=1}^{G} f_s(v_s) \) to replace the linear form \( \sum_{s=1}^{G} \beta_s v_s \) in the generalized linear model where \( v \) is predictor variable and \( G \) is the number of predictor variables. Here, we used only one predictor variable, that is, the input bin count. Therefore, when using GAM, \( \mu_i \) can be estimated by \( \log(\mu_i) = \beta_0 + f(\log(x_i)|\beta) \), where \( f \) is represented using smoothing splines and \( \beta \) is a vector of coefficients for the spline term with length of 9 as default. We implemented GAM using R package mgcv [24] and set the restricted maximum likelihood [25] as the method for estimating the smoothing parameters. To optimize the model, MoAIMS implements both RLM and GAM and subsequently uses that with a lower BIC (Bayesian Information Criterion) [26]. BIC scores were calculated in the general method by \( r \ln(T) - 2 \ln(\hat{L}) \), where \( r \) is the number of parameters, \( T \) the number of bins, and \( \hat{L} \) the maximum likelihood.

### Table 1

An example of the model summary table

| Dataset   | \( \pi_i \) | BIC_1S | BIC_2S | optim_k | optim_reg |
|-----------|-------------|--------|--------|---------|-----------|
| WT_rep1   | 0.138       | 1679168| 1678590| 2       | rlm       |
| WT_rep2   | 0.11        | 1212063| 1212005| 2       | rlm       |

The columns represent dataset names, signal proportion, BIC values for 1S (one-signal) mode, BIC values for 2S (two-signal) mode, optimized \( k \), and optimized regression methods.
The size parameter $a$ is estimated by $\hat{a} = \sum_i n_i \hat{a}_i / \sum_i n_i$, where $\hat{a}_i = [E(S_i)]^2 / [\text{Var}(S_i) - E(S_i)]$ (the expectation is calculated using median value; the variation is calculated using the median absolute deviation) and $n_i$ is the number of bins.

After estimating the parameters of the background component, expectation maximization (EM) algorithm [27] was applied to estimate the parameters of the signal component in 1S mode, $\Theta_{1S} = (b, c, \pi_s, \pi_b)$, is estimated in the maximization step with optimized $k$ value rather than based on a pre-defined $k$ value in MOSAiCS. For the parameters $b$ and $c$, the method of moments is used as MOSAiCS. The details of modified EM process for 1S mode are provided in the Supplementary. We performed the EM process to estimate the parameters of the signal component in 2S mode, $\Theta_{2S} = (b_1, c_1, b_2, c_2)$, and $\pi_{s1}$ unchanged as MOSAiCS.

Model design for MeRIP-Seq analysis

The modification and extension of the statistical framework proposed by MoSAiCS is aimed to make our software more suitable for MeRIP-Seq analysis. This statistical framework is based on the negative-binomial distribution that is capable of modeling the variance of gene expression. We validated it by plotting the residuals between IP signal and estimated background corresponding to the gene expression. As Figure S1 shows, IP signal increases as the gene expression increases.

The modification and extension involved three aspects. First, we used log-transformation in estimating the background means instead of power-transformation in MOSAiCS because log-transformation is more commonly used in RNA sequencing analysis [28], and this can simplify the parameter tuning required in power transformation. Second, we set $k$, the minimum count in the signal regions, flexible instead of pre-defined in MOSAiCS. Because $k$ may depend on the library size and signal-to-background ratio of the experiments [29], we set $k$ flexible and optimized in the model fitting. With the optimized $k$, the signal proportion ($\pi_s$) was estimated by EM rather than based on a pre-defined $k$ value in MOSAiCS. Third, in addition to the RLM used by MOSAiCS in estimating background means, we applied GAM for regression to obtain better fitting for some cases of RNA sequencing data, as shown in Figure S1. An example of summary table of the fitted models is shown as Table 1 that provides signal proportion, BIC values for 1S (one-signal) mode, BIC values for 2S (two-signal) mode, optimized $\hat{k}$, and optimized regression methods.

Detection of enriched regions

The enriched regions were decided under the threshold of the false discovery rate (FDR), which was calculated as in [29, 30]. In this study, false discovery means a genomic region that is claimed to be significant when it is not. For a set $\mathcal{M}$ of $m$ enriched regions that satisfies a defined cut-off (default is 0.05), the estimated FDR is equal to $\left(1 / m \right) \sum_{i \in \mathcal{M}} \hat{p}(Z = 0 | Y_i)$, where $\hat{p}(Z = 0 | Y_i)$ is equal to $\frac{(1-\hat{\pi}_{b1}) \hat{p}_{0j} + (1-\hat{\pi}_{s1}) \hat{p}_{1j}}{\hat{p}_{0j} + (1-\hat{\pi}_{s1}) \hat{p}_{1j}}$ for the 1S mode and $\frac{(1-\hat{\pi}_{b2}) \hat{p}_{0j} + (1-\hat{\pi}_{s2}) \hat{p}_{1j}}{\hat{p}_{0j} + (1-\hat{\pi}_{s2}) \hat{p}_{1j}}$ for the 2S mode with $\hat{p}_{0j}$ and $\hat{p}_{1j}$ as the post probability for the $j$th bin from the background component and the signal component respectively. Finally, the enriched regions were merged and output in the BED12 format with the highest bin count of merged regions as the score, which can be used as a filter to obtain higher confident signal region candidates.

Goodness of fitting (GOF)

To display the goodness of fitting (GOF), the simulations are performed using the estimated parameters. For the simulation of the 1S mode, $m$ background bins and $n$ signal bins were randomly sampled according to $\pi_s$, where $m + n = T$. The background read count of $T$ bins were generated from the background distribution $\text{NB}(a, \frac{a}{a+b})$, for the 1S mode and $\text{NB}(a, \frac{a}{a+b+c})$ for the 2S mode. Subsequently, for $n$ signal bins, the read count was composed of the background read count, the count sampled from the signal distribution $\text{NB}(1, \frac{c}{c+1})$, and the minimal count $k$. For the simulation of the 2S mode, $m$ background bins, $n_1$ first-signal bins, and $n_2$ second-signal bins were randomly sampled according to $\pi_s$ and $\pi_{s1}$, where $m + n_1 + n_2 = T$. The background read count of $T$ bins were generated from the background distribution $\text{NB}(a, \frac{a}{a+b+c})$, for the signal bins, the read count was composed of the background read count, the count sampled from the corresponding signal distribution $\text{NB}(1, \frac{c}{c+1})$ or $\text{NB}(2, \frac{c_2}{c_2+1})$, and the minimal count $k$. Figure 2 gives an example of GOF plot.

Results

Comparison with other tools

Detection of m6A-enriched regions

We performed analysis on two m6A MeRIP-Seq studies. One is from mouse embryonic stem cell [31] that uses the single-end and strand-specific sequencing protocol. The mouse datasets include the wild type and knock-out of Mettl3 (an m6A methyltransferase), of which each has two biological replicates. The other is from human A549 cell line [32] that uses the paired-end and strand-specific sequencing protocol. The human datasets contain negative control (shGFP) and perturbation of three types of m6A methyltransferases including Mettl14, Mettl3, and WTAP, of which each has two replicates. Table S1 summarized the information of datasets. Raw fastq files were retrieved from Gene Expression Omnibus [33] with accession numbers GSE52662 and GSE54365. Reads were aligned to human (hg19) and mouse (mm10).
Three commonly-used tools for comparison are MACS (version MACS2), exomePeak (v2.13.2) and MeT-Peak (v1.0.0). Duplication-removed bam files were used as input for the three tools. For MACS, we specified parameters 

```
--nomodel --extsize=100 --keep-dup=all -g
286,000,000 (for human)/221,000,000 (for mouse).
```

We genome using STAR (version 2.6.0c, default setting) [15] with annotation files of GENCODE (human release 19 and mouse release M19) [34]. Only uniquely mapped reads were kept. The sorted (by coordinates) and duplication-marked bam files were generated by Picard (version 2.18.1) and subsequently used as input for MoAIMS.
Table 2 Features of MoAIMS compared with other tools

| Features                        | MoAIMS | exomePeak | MeTPeak | MACS |
|---------------------------------|--------|-----------|---------|------|
| Exome-based                     | Y      | Y         | Y       | N    |
| Strand-specific/Paired-end      | Y      | N         | N       | N    |
| Time-consuming                  | N      | Y         | Y       | N    |
| Inference of signal proportion  | Y      | N         | N       | N    |
| Visualization of model fitting  | Y      | N         | N       | N    |
| Output in BED12 format          | Y      | Y         | Y       | N    |
| Support for differential methylation analysis | N | Y         | Y       | N    |

kept the peaks called by MACS overlapped with exonic regions for comparison. For exomePeak and MeTPeak, we used the default setting.

First, we compared the m6A-enriched regions called by MoAIMS with MACS, exomePeak, and MeTPeak. We verified to what extent the enriched regions called by the four tools agree with each other using BEDTools [35]. To obtain higher confident regions, we chose the enriched regions (FDR ≤ 0.05) called by MoAIMS with score ≥ 10. Table S2 shows the results for the mouse wild-type datasets. Each cell of the table represents the percentage of enriched regions of tools in the columns detected by tools in the rows; the number in bracket is the number of enriched regions called by each tool. It is indicated that our enriched regions are overlapped more with MACS and exomePeak. Additionally, MeTPeak called relatively less peaks and, in some cases, could miss enriched regions, as shown in Figure S1.

Subsequently, we verified the occurrence of the DRACH motif [36], a classic m6A motif where D = A, G, or U; R = A or G; and H = A, C, or U, in the top-5000 enriched regions. The ranking scheme for MACS, exomePeak, and MeTPeak is fold change. For MoAIMS, the ranking scheme of fold change (FC) and score are both used for comparison. Sequences of length 200 bp were extracted around the summits of the enriched regions. For MACS, we used the summits it provided; for MoAIMS, exomePeak, and MeTPeak, the summits were defined as the positions with the highest read coverage. Because we had the strand-specific sequencing data, we only counted the motifs that occurred in the expressed genes with coverages (for MACS, only motifs with coverages were counted). Figure 3 compares the percentage of motif occurrence in the decreasing peak ranks for a wild-type mouse dataset (comparisons are also conducted for the other untreated datasets shown in Figure S1). The results indicated that our software achieved comparable performance to the other three tools.

![Fig. 5 Example of detection of strand-specific enriched regions. The plot is generated using IGV [38], showing the enriched region called by MACS, exomePeak, MeTPeak, and MoAIMS in the first four tracks. The following tracks are coverage and aligned reads (strand orientation is colored) for the IP and input sample, respectively, and the genome annotation.](image-url)
Table 3 Performance on the time cost

| Dataset          | MoAIMS | exomePeak | MeTPeak |
|------------------|--------|-----------|---------|
| Human shGFP_rep1 | 14.1   | 141.0     | 176.4   |
| Mouse WT_rep1    | 10.6   | 110.4     | 143.4   |

shGFP_rep1 is one human negative control dataset. WT_rep1 is one wild-type mouse dataset. The units of time is minute.

Next, we are interested to know to what extent the m6A miCLIP sites agree with the MeRIP-Seq enriched regions. We collected miCLIP-Seq data of human A549 cell line from [37], which maps m6A sites at single-base resolution. We counted the number of regions containing miCLIP sites in the top-5000 enriched regions detected by the four tools (the ranking scheme is the same as that for counting motif occurrence). Figure 4 shows that our software with score ranking has the most number of regions with m6A miCLIP sites in the decreasing peak ranks (comparisons were also conducted for the other human dataset provided in Figure S1). To determine whether the number was affected by the length of the enriched regions, we compared the length of the top-5000 enriched regions between the tools, as shown in Table S3. The result shows that compared with MeTPeak, which ranks second with regard to consistency with miCLIP sites, MoAIMS can detect more regions with m6A miCLIP sites under the similar resolution.

Features of MoAIMS

MoAIMS is efficient software with appealing features, as shown in Table 2. Thus, we performed comparison analysis with regard to those features. First, because our software is compatible with general RNA sequencing protocols in counting reads, we investigated how the methods of counting reads affected the detection of enriched regions for pair-end RNA sequencing. The comparison was conducted for the human shGFP (negative control) datasets among exome-based callers: MoAIMS, exomePeak, and MeTPeak. Table S4 lists the number of enriched regions detected by these three tools using pair-end reads and first-in-pair reads, separately. The result indicates that exomePeak and MeTPeak differ in the method of counting paired-end reads, while the difference is limited for our software.

Next, our software is a strand-aware caller; thus, it can avoid calling ambiguous regions that are overlapped with other regions on different strands. Figure 5 shows an example of how our software called strand-specific enriched regions. As shown in the figure, a protein-coding gene Mtmr10 and an antisense gene RP23-84M17.2 are partially overlapped. The coverage track in red (colored by strand) indicates the signal in Mtmr10, not the antisense gene. For this case, exomePeak and MeTPeak have callings on both genes, but MoAIMS can avoid the ambiguous callings.

Finally, our software offers excellent processing speed compared with exome-based callers exomePeak and MeTPeak, which require approximately 2 hours to analyze one dataset (MeTPeak needs even more time because it applies HMM). Table 3 lists the time cost for a human and a mouse dataset, indicating that our software is competitive as it only requires several minutes and can yield comparable performance.

Fig. 6 Position profile of m6A-enriched regions for a wild-type mouse dataset. X-axis is the relative position coordinates and Y-axis is the mean coverage of the enriched regions. The plot is generated using RCAS [39].
**Application on feature and functional analysis of m6A**

m6A is characterized by its location preference close to three prime untranslated regions (3' UTRs); thus, we verified the position preference of the enriched regions (with score $\geq 10$) called by MoAIMS. For the wild-type mouse datasets, as shown in Figure 6 and S7(a), the enriched regions exhibit location bias near 3' UTRs, which is consistent with the results of the original study [31]. For the human negative control datasets, we observed that enriched regions appeared near 5' UTRs, as shown in Figures S7(b) and (c), which agrees with the findings of the original study [32] regarding methylated m6A at transcription start sites.

Because our software infers the signal proportion from the mixture NB model, we assumed that this value can reflect the treatment effect; for example, the knocking-down/out of methyltransferases (such as WTAP, METTL3, or METTL14) can cause decreased signal proportion. For the mouse datasets, as shown in Fig. 7, Mettl3 knock-out exhibits a clear decreasing trend for signal proportion, which agrees with the findings of a recent study [40] that include a discussion on the m6A methyltransferase treatment experiments and the effect of treatment in this dataset. For the human datasets, as shown in Figure S1, WTAP shows a relatively clear effect after perturbation, while Mettl3 and Mettl14 shows less effect. This trend is consistent with the original study [32], in which the authors observed the necessity of WTAP for m6A methylation, while perturbation of Mettl3 and Mettl14 exhibited milder effects in decreasing methylation level. These results suggest that the signal proportion can be used as an intuitive indicator of the m6A treatment effect, which can facilitate biologists' evaluation on the treatment experiments.

Finally, we conducted a functional analysis on the genes affected by the perturbation of methyltransferases. We performed gene ontology (GO) analysis by RCAS [39] on genes with lost m6A-enriched regions. The loss of m6A-enriched regions is defined as a state from being detected in all the replicates of the wild type to being undetected in all the replicates of the treated type. The GO results of enriched biological process (BP) terms are shown in Fig. 8. For the mouse datasets of the wild type and Mettl3 knock-out, the enriched BP terms are related to planar polarity and polarity, thus suggesting that the loss of m6A affects the development of embryo cells. For the human datasets of negative control and WTAP perturbation, the enriched BP terms are related to histone methylation and acetylation, which also appeared in the term list for mouse. This observation agrees with that of [41] regarding m6A's function in destabilizing transcripts that encode histone modification enzymes.

**Discussion**

MoAIMS is an efficient and user-friendly software for the analysis of MeRIP-Seq. Nonetheless, improvements are still required. First, MoAIMS currently supports only the analysis of single samples. For replicate samples, although enriched regions common in all the replicates can be easily extracted using our software, a joint statistical model can be developed as an alternative that considers the variance among replicates. Next, apart from the NB
distribution, other statistical distributions are worth being tested owing to the wide diversity of RNA sequencing data. For example, Poisson–Tweedie has been proposed for studying differential expressed genes as it is a more general family of count data distributions that can fit RNA sequencing data under situations of heavy tail or zero inflation [42]. Additionally, the double Poisson distribution has been applied to manage under-dispersion RNA sequencing data [43]. Last but not least, because our software can provide user-friendly outputs for downstream analysis, it is feasible to integrate MeRIP-Seq datasets with other biological data for a comprehensive functional analysis, especially for MeRIP-Seq-treatment experiments.

**Conclusion**

We developed MoAIMS, which is an efficient and user-friendly software for analysis of MeRIP-Seq. MoAIMS is compatible with general RNA sequencing protocols, achieves excellent speed and competitive performance, and provides user-friendly outputs for downstream analysis. When MoAIMS was applied to studies of m6A, m6A’s known biological features and its interplay with histone
The authors declare that they have no competing interests.

**Competing interests**

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**Availability and requirements**

- **Project name:** MoAIMS
- **Project home page:** https://github.com/rreybeyb/MoAIMS
- **Operating systems:** Linux, Mac OS, Windows
- **Programming language:** R
- **Other requirements:** R version 3.4.0 or higher
- **License:** GNU GPL
- **Any restrictions to use by non-academics:** None

**Supplementary information**

**Supplementary information** accompanies this paper at https://doi.org/10.1186/s12859-020-3430-0.

**Additional file 1:** Supplementary materials for "MoAIMS: efficient software for detection of enriched regions of MeRIP-Seq".

**Abbreviations**

3’ UTRs: three prime untranslated regions; BIC: bayesian information criterion; BP: biological process; ChIP-Seq: Chromatin immunoprecipitation sequencing; EM: expectation maximization; FDR: false discovery rate; GAM: generalized additive model; GO: gene ontology; GOF: goodness of fitting; HMM: hidden markov model; IP: immunoprecipitation; m1A: N1-methyladenosine; m5C: 5-methylcytidine; m6A: N6-methyladenosine; MeRIP-Seq: Methylated RNA immunoprecipitation sequencing; NB: negative-binomial; NGS: next-generation sequencing; RLM: robust fitting of linear models; TPM: transcripts per million.

**Acknowledgements**

The authors thank Dr. Chao Zeng for valuable comments.

**Author's contributions**

YZ conceived and MH supervised this study, YZ implemented the methods, performed the experiments and wrote the draft. MH revised the manuscript critically. YZ and MH contributed to the construction of methods and analysis/interpretation of the data. All authors read and approved the final manuscript.

**Funding**

Publication costs are funded by Waseda University (basic research budget). This study was supported by the Ministry of Education, Culture, Sports, Science and Technology (KAKENHI) [grant numbers JP17H01318, and JP16H05879, JP16H01318, and JP16H02484 to MH]. The funding bodies did not play any role in the design of the study or collection, analysis and interpretation of data or in writing the manuscript.

**Availability of data and materials**

The datasets and materials can be downloaded from https://github.com/rreybeyb/MoAIMS.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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Received: 22 November 2019 Accepted: 24 February 2020

Published online: 14 March 2020

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