

# Convolution neural network for the prediction of RNA using heterogeneous network.

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## Abstract

T N6-methyladenosine (m6A), a kind of post-transcriptional alteration, is essential for the stability and control of gene regulations. As a result, identifying m6A is critical for comprehending the functional mechanisms of biological systems. To make the tedious process easier, many machine learning algorithms based on convenient handcraft feature extraction techniques had been presented. Nevertheless, due to poor extracting features, such strategies enhance computing overhead and as a result, reduce the reliability of m6A detection. That research provides a rapid and accurate statistical method for m6A location detection. This suggested approach relies on the CNN, where recovers the much more important aspects from RNA sequences encode by appending as well as nucleotides chemical composition. This proposed approach is tested to state-of-the-art prediction algorithms on different species benchmark datasets. Here on a benchmark dataset of Homo sapiens (*H.sapien*), Mus musculus (*M.musculus*), Saccharomyces cerevisiae (*S.cerevisiae*), as well as Arabidopsis thaliana (*A.thaliana*), the proposed system provides good precision of 93.6 percent, 93.8%, 85.0% and 92.5%, correspondingly.

**Keywords:** Gene expression, Feature extraction, Convolutional neural network, Location detection.

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## Introduction

Many of the moreover 160 RNA alterations, methyladenosine (m6A) is by far the most commonly occurring. It may be found in eukaryotic such as bacteria, bugs, as well as primates. This adenine basis methylation at the 6th position of the nitrogen is referred to as m6A [1-2]. RNA structure dynamical, cellular proliferation and remodeling, RNA localization and destruction, alternative splicing, circadian clocks control, including fundamental microRNA digestion are all linked to the m6A gene [3]. As a result, knowing the biological process's functioning functions is critical. In the latest days, To discover m6A locations, greater designing research such as m6A-seq as well as MeRIP-Seq [4-6] have been used by the editorial assistant who coordinated the evaluation of this work and approved it for publishing. Using parallel processing and antibody-mediated capturing, the m6A-seq gives a transcriptome-wide view of the rat as well as humans' m6A alteration landscapes. MeRIP-Seq, on either hand, detects transcribed that seem to be adenosine methylated targets as well as gives insight into mammal transcriptional modulating the expression [7]. The controlled experiments were wasteful in terms of time and money, including in terms of accurately finding the m6A location. Researchers want to solve the problem of properly and quickly identifying m6A locations, which is now a bottleneck. As a result, developing mathematical algorithms is critical.

The above computer learning-based techniques were primarily focused on the useful constructed characteristics that need

domains expertise for successful predictions of the suggested predictors. Such characteristics are constructed in such a way that knowledge about the patterns in the sequences should be preserved [8].

## Related works

Deep learning-based computationally structures, on either hand, are capable of separating the much more relevant characteristics from sequencing without any need for human interaction, resulting in mathematical computations that are substantially more precise and resilient. Presently, deeply learning basis methods acquires modern outcomes in the fields of natural languages processes, image identification, word detections as well as in the field of mathematical biologist. Deep learning-based algorithms are now achieving outstanding achievements in the fields of natural languages processing, image identification, audio acknowledgment and computational biology.

Inside this context, they present simply and efficient CNN basis framework for the discovery of m6A locations in RNA sequencing to cover the efficiency and computationally complexity limitations in present numerical simulations. pm6A-CNN is how we refer to it. One-hot transcription and nucleotides chemical components are used to describe the incoming RNA sequences [9]. The chemical characteristics of polymorphisms are the most fundamental characteristics of nucleotides is based on functional compounds, hydrogen bonds, including ring system [10-12]. The CNN design can

extract meaningful the much more essential properties from RNA sequence representations, allowing the pm6A-CNN to detect the m6A locations with greater accuracy and reliability. This grid search technique [13,14] is used to find the best hyperparameters for the pm6ACNN. For compatibility with state-of-the-art systems, the achievement of pm6A-CNN has been assessed through the uses of the sub-sampling approach with the parameter values set to 10. Its pm6A-CNN beat current mathematical algorithms as a result of this breakthrough.

## Materials and methods

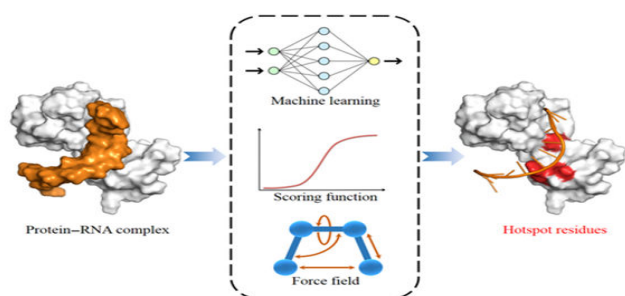
### Proposed methods

Adenines are found in the middle of all 4 benchmarks functions. These affirmative variants include methyladenosine (m6A) spots that have been scientifically confirmed,

S. No	Name of species	Pattern	Length
1	<i>H.sapien</i>	Positive 1120 Negative 1230	42 nt
2	<i>M.musculs</i>	Positive 750 Negative 750	43 nt
3	<i>S.cerevisiae</i>	Positive 1317 Negative 1317	52 nt
4	<i>A.thaliana</i>	Positive 2110 Negative 2110	121 nt

**Table 1.** Datasets of species.

The RNA sequencing is encoded using a mixture of 2 widely utilized encoded techniques; one-hot encoding as well as nucleotides elemental composition (NCP). A is symbolized by (1,0,0,0), C is expressed by (0,1,0,0), G is expressed by (0,0,1,0), whereas U is expressed by (0,0,1,0). (0,0,0,1). Whereas NCP is a 3-dimensional Cartesian coordinates framework representation of each nucleotide in the RNA sequence depending on its chemical properties. C and U are pyrimidines with a single ring, whereas C and U were pyrimidines with two rings. Throughout regards to secondary structures formation, weaker hydrogens bonds exist among A and U, but strong hydrogens bonds exist among C and G. Those 4 nucleotides may be categorized into 3 separate groupings based on these 3 chemicals features, which have been expressed in the 3-dimensional Cartesian coordinates system by giving a value of 1 or 0 (Figure 1).



**Figure 1.** Proposed model.

meanwhile, the negative variants do not even have methyladenosine (m6A) groups. The *H.sapiens* benchmarking database was collected using 1130 positives sequencing as well as 1130 negatives sequencing, which each was 41nt long. The *M.musculus* benchmarking database, with each nucleotide being 41nt long. *M.musculus* has 725 affirmative sequencing plus 725 negative genotypes in its datasets consisting. In 2015, the *S.cerevisiae* benchmarking information was released. It has 1307 affirmative sequencing as well as 1307 negative sequencing, each being 51nt long. It consists of 2100 affirmative and 2100 negative sequencing, which each is 101nt in size.

As illustrated, they present deeply learning-based Architectures that accepts RNA sequencing as an input. A grids research methodology is used to find the best hyperparameters. These hyper-parameter values are shown in Table 1.

These matrices created by combining one hot-encoding plus NCP were connected in series to form a 7-channel vector that represents the RNA string. A CNN model with 2 Conv1D stages and 2 fully connected layers is used to process the resultant vectors. A ReLU activated function follows each Conv1D layer. Furthermore, the initial Conv1D is accompanied by a grouping normalization, with the group's size set at 4. The convolution levels' learned characteristics were sent from dropouts layers with a dropouts rate of 0.5 and subsequently to 2 fully connecting layers. An activated function follows the 1<sup>st</sup> completely linked layers. To reduce the number of features, the strengths as well as biased of the filtration are regularized using the L2 approach. For modeling, the Adam optimizers with a learning rate of 0.001 are used. As a loss function, binary cross-entropy is used. For the maximal amount of training repetitions, a sampling size of 32 is used, with quick terminating depending on validated losses.

$$\text{Conv}(S)_{ij} = \text{ReLU} \left( \sum_{s=0}^{Z-1} \sum_{n=0}^{I-1} W_{sn}^k S_{j+s,n} \right) \quad (1)$$

Conv1D is represented by Equations 1, wherein S is the RNA pattern's inputs, k refers to the filter's number and j indicates the outputs position's index. Each  $W_k$  is convolutional filtering with a  $Z \times I$  weighted matrices, with Z indicating the size of the filter and I indicating the set of input streams.

$$f = w_{d+1} \sum_{k=1}^d m_k w_k z_k \quad (2)$$

Equations 2 depicts the densest layer, with  $w_{d+1}$  denoting an additional bias term,  $mk$  denoting a dropouts operator based on the Bernoulli distributions,  $z_k$  denoting one-dimensional features vectors, as well as  $w_k$  denoting the weighting of  $z_k$  from the preceding stage.

$$\text{ReLU}(x) = \begin{cases} x & \text{if } x > 0 \\ 0 & \text{if } x \leq 0 \end{cases} \quad (3)$$

The ReLU activated function is shown in Equations 3 wherein  $x$  is the inputs. This sigmoid activated function is depicted in Equations 4.

$$\text{Sigmoid}(x) = \frac{1}{1 + e^{-x}} \quad (4)$$

### Performance evaluation

This 10-fold cross-validations approach was used to assess the performances of their proposed approach. This benchmarking information was subsequently split into 10 foldings that were mutually exclusionary. One folding is set aside for order to test the proposed system, another for confirmation and the other folded are set aside for retraining the conceptual approach. This is a continuous procedure that repeats itself 10 times. This averaged consequence of 10 folds had been used to calculate the final estimation of the effectiveness.

S. No	Name of species	Method	ACC	Sn	Sp	Mcc
1	<i>H.sapien</i>	One-hot	0.92	0.81	0.91	0.83
2	<i>M.musculus</i>	Ncp	0.71	0.82	0.82	0.92
3	<i>S.cerevisiae</i>	One hot-Ncp	0.84	0.84	0.96	0.92
4	<i>A.thaliana</i>	One hot-Ncp	0.95	0.95	0.88	0.81

**Table 2.** Representation of sequence methods.

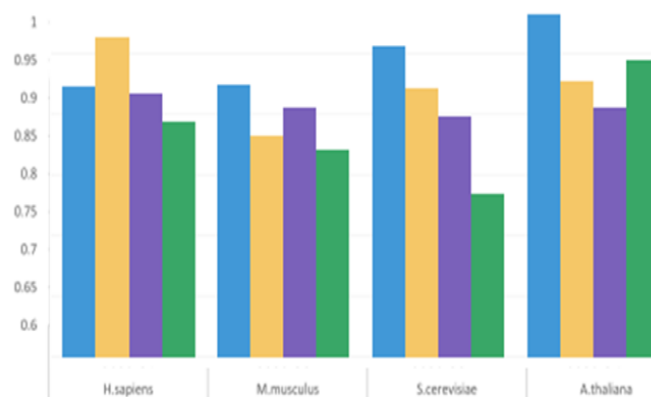
This combined depiction yielded excellent performance in the identifying of m6A locations, as could be observed. Additionally, the AUC of the developed framework together with sample variance mistakes in 10-fold cross-validation utilizing the benchmarking databases of *H.sapienses*, *M.musculuses*, *S.cerevisiaees*, as well as *A.thaliana*. Furthermore, effectiveness was used to evaluate the pm6A-dominance. CNN's By utilizing only the 4<sup>th</sup> benchmarking dataset, the RFathM6A performed better than the M6AMRFS. i.e. *A.thaliana*.

Contrasts of mathematical methods. The '-' symbol indicates that only certain measurements for the necessary instruments just weren't available. Their proposed approach is shown to

S. No	Name of species	ACC	Sn	Sp	Mcc
1	<i>H.sapien</i>	0.9	0.8	0.9	0.8
2	<i>M.musculus</i>	0.7	0.8	0.8	0.9

## Results and Discussion

Using 4 species benchmarking functions, this suggested framework was validated utilizing 10-fold cross-validations. Studies conducted were carried out to determine the efficacy of integrating nucleotides' chemical characteristics. For RNA sequences representation, the very first attempt only employed a one-hot encoding. In experiment 2, just nucleotides chemicals characteristics were employed to model RNA sequences. In experiment 3, the description including both encrypting techniques was combined (Figure 2 and Table 2).

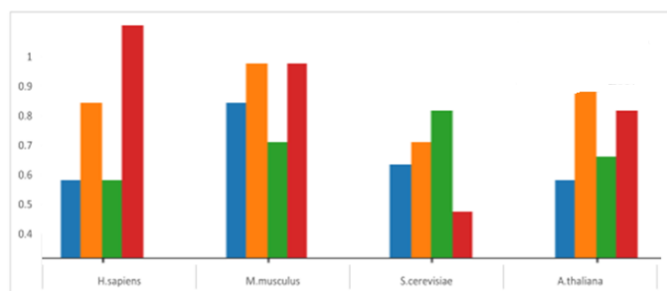


**Figure 2.** The performance measures of the proposed model. Note: ■ One-hot, ■ NCP, ■ One-hot, NCP.

outperform all those other competitive techniques. The proposed methodology outperforms the state of the arting approach iN6-Methyl on the *H.sapienses* and *M.musculuses* benchmarks datasets. The gains in SN, MCC, AUC, and ACC for the *H.sapienseses* benchmarking datasets are 2.5%, 6.5%, 4.3% and 6.2%, correspondingly. The gains in SN, MCC, AUC, and ACC in the *M.musculus* benchmarks dataset were 4.3%, 11.9%, 7.3% and 5.8%, correspondingly. Inside the datasets consisting of *S.cerevisiae*, the prototype system beat iMRM by 7.3%, 7.6%, 7.5%, 15.6% and 7.4%, correspondingly, in all performances indicators ACC, SN, SP, MCC, as well as AUC (Table 3 and Figure 3).

**Table 3.** Proposed system outcomes.

3	<i>S.cerevisiae</i>	0.8	0.8	0.9	0.9
4	<i>A.thaliana</i>	0.9	0.9	0.8	0.8



**Figure 3.** Comparison between the proposed model and existing models. Note: ■ Round 1, ■ Round 2, ■ Round 3, ■ Round 4.

Eventually, the proposed methodology enhanced SN, MCC, ACC, SP with 7.10%, 5.0%, 9.1% and 14.1%, correspondingly, for the *A.thaliana* benchmarks dataset. The overall efficiency of the proposed approach in identifying the m6A location utilizing the mixture of multiple distinct encoded approaches for the representations of RNA sequences is demonstrated by the expected outcomes of the suggested framework in order of all performance benchmarks for all baseline methods.

## Conclusion

Researchers presented effective deeply learning-based Architectures for identifying m6A locations in several organisms in this work. By combining one-hot encoding plus nucleotides chemicals characteristics for the characterization of RNA sequences, the CNN-based prediction identifies far more relevant information. Such mixture aided the predictor's ability to identify m6A locations more effectively and efficiently. Furthermore, it is expected that perhaps the created prediction, in conjunction with both the website, will be an excellent tool for researchers to examine the functioning mechanism of m6A spots.

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