Evolutionary Implications of the RNA $N^6$-Methyladenosine Methylome in Plants

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Abstract

Epigenetic modifications play important roles in genome evolution and innovation. However, most analyses have focused on the evolutionary role of DNA modifications, and little is understood about the influence of posttranscriptional RNA modifications on genome evolution. To explore the evolutionary significance of RNA modifications, we generated transcriptome-wide profiles of $N^6$-methyladenosine (m$^6$A), the most prevalent internal modification of mRNA, for 13 representative plant species spanning over half a billion years of evolution. These data reveal the evolutionary conservation and divergence of m$^6$A methylomes in plants, uncover the preference of m$^6$A modifications on ancient orthologous genes, and demonstrate less m$^6$A divergence between orthologous gene pairs with earlier evolutionary origins. Further investigation revealed that the evolutionary divergence of m$^6$A modifications is related to sequence variation between homologs from whole-genome duplication and gene family expansion from local-genome duplication. Unexpectedly, a significant negative correlation was found between the retention ratio of m$^6$A modifications and the number of family members. Moreover, the divergence of m$^6$A modifications is accompanied by variation in the expression level and translation efficiency of duplicated genes from whole- and local-genome duplication. Our work reveals new insights into evolutionary patterns of m$^6$A methylomes in plant species and their implications, and provides a resource of plant m$^6$A profiles for further studies of m$^6$A regulation and function in an evolutionary context.

Key words: RNA modification, bioinformatics, epitranscriptome, plant, epigenetic modifications, evolutionary genetics.

Introduction

Epigenetics has informed hereditary and evolutionary theory since Conrad Waddington first defined the term in the 1940s (Waddington 1942; Jablonka and Lamb 2002). To date, epigenetic modifications such as methylation, which regulate many aspects of the genome including chromatin organization, structure, recombination, and gene expression (Kouzarides 2007; Colome-Tatche et al. 2012; Shi et al. 2019; Liang et al. 2020), have been recognized as important contributors to evolutionary novelties in eukaryote genomes (Diez et al. 2014; Vidalis et al. 2016; Wendel et al. 2016; Yi 2017). With the development of high-throughput epitranscriptome sequencing technologies, RNA modifications have also been identified as novel and dynamic regulators of gene expression. RNA $N^6$-methyladenosine (m$^6$A), the most abundant internal RNA modification, can affect many aspects of RNA metabolism (e.g., RNA splicing, localization, translation, stability) and regulates gene expression in many biological processes (Zhao, Roundtree, et al. 2017; Yang et al. 2018; Shen et al. 2019), including the DNA damage response (Xiang et al. 2017). The association of m$^6$A modification with genome integrity and gene expression led to the question of whether the evolution of m$^6$A modification interplays with genome evolution and innovation.

The evolutionary conservation of m$^6$A was revealed through comparative analysis of the m$^6$A methylomes of humans and mice (Dominissini et al. 2012; Song et al. 2021). Through the examination m$^6$A methylomes from humans, chimpanzees, and rhesus macaques, the evolution of m$^6$A was found to occur in parallel with the evolution of the gene sequence and mRNA abundance (Ma et al. 2017). Population genetics analysis showed that a proportion of m$^6$A modifications occurred under positive selection pressure, and were associated with human traits or diseases (Zhang et al. 2020). Because the evolution of genome structures and the composition of m$^6$A functional genes differ widely between humans and plants (Yue et al. 2019), the evolution of plant m$^6$A warrants specific investigation. A single-species m$^6$A analysis uncovered evolutionary conservation of m$^6$A within Arabidopsis (Luo et al. 2014) and an

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association between m^6^A modification and the evolution of duplicate genes in maize (Zea mays) (Miao et al. 2020). However, virtually nothing is known about the global landscape and evolutionary dynamics of m^6^A methylomes in plant species or their potential implications for the evolution of plant genomes.

In this study, we first generated m^6^A profiles for 13 representative plant species spanning over half a billion years of evolution. We then performed comparative analysis of these m^6^A methylomes in order to address topics of conceptual importance to understanding the evolutionary implications of m^6^A methylomes in plant species. Specifically, we: 1) characterized the evolutionary patterns of conservation and diversity of m^6^A methylomes in plants species, 2) dissected the relationship between m^6^A modifications and gene origin and evolution, 3) revealed the evolutionary implications of m^6^A modifications on polyploidization in plant species, and 4) investigated the influence of m^6^A modifications on both gene family expansion and evolution triggered by local-genome duplications.

**Results**

**Transcriptome-Wide Mapping of m^6^A Modifications for 13 Plant Species**

To explore the landscape of m^6^A modifications in the plant kingdom, we constructed and sequenced m^6^A-seq libraries from 12 plant species, namely A. thaliana, Gossypium arboreum, G. hirsutum, Glycine max, Phaseolus vulgaris, Sorghum bicolor, Z. mays, Aegilops tauschii, Triticum dicoccoides, T. aestivum, Oryza sativa, and Phycomitrella patens (supplementary data 1, Supplementary Material online). The sequencing reads are highly accumulated around the stop codon and within 3’-untranslated regions (3’-UTRs) in all m^6^A-seq libraries (supplementary fig. 1, Supplementary Material online). According to the previous study (Miao et al. 2020), we identified m^6^A peaks with high confidence for each species by intersecting peak regions in a pairwise fashion among all three replicates. Regions that overlap in at least two of three replicates were designated as high-confidence m^6^A peak regions (supplementary fig. 2, Supplementary Material online). Strong correlations were observed for the abundance of confident peaks between biological replicates (supplementary fig. 3, Supplementary Material online). After the integrated analysis of these data sets together with previously published m^6^A-seq data sets of four plant species (A. thaliana, O. sativa, Z. mays, and Solanum lycopersicum; supplementary data 2, Supplementary Material online), we obtained transcriptome-wide m^6^A profiles for 13 representative plant species (six dicots, six monocots, and one bryophyte; genome size 135 Mb–17 Gb; number of annotated protein-coding genes 27,420–107,545) representing over half a billion years phylogenetic timescale (fig. 1A). Among these 13 plant species, there were 3,743–31,719 high-confidence m^6^A peaks corresponding to 3,726–30,968 genes (supplementary data 3, Supplementary Material online). For each analyzed species, m^6^A peaks were preferentially enriched around the gene stop codon and 3’-UTR regions (supplementary fig. 4, Supplementary Material online), consistent with previous observations in both animal and plant species (Fu et al. 2014; Liang et al. 2020).

These transcriptome-wide m^6^A profiles provide a new valuable resource for researchers interested in investigating the evolutionary dynamics of m^6^A in plants. The statistical analysis showed that the m^6^A methylation ratio for all genes was significantly negatively correlated with the genome size (Pearson’s r = −0.62, P = 0.02; fig. 1B) and the gene number (Pearson’s r = −0.65, P = 0.02; fig. 1C). To enable interspecies comparison, we identified orthologous genes (OGs) and species-specific genes (SGs) through comparative analysis of protein-coding genes from the 13 species (see Materials and Methods). We observed that the m^6^A methylation ratio for OGs was significantly higher than that for SGs (fig. 1D and supplementary data 4, Supplementary Material online). Of the one-to-one OGs among these species, the number of m^6^A OGs shared by different species were significantly more than expected by chance (supplementary data 5, Supplementary Material online). Gene ontology (GO) enrichment analysis showed that OGs with high-conservation of m^6^A methylation were mainly enriched in protein transport and RNA processing, whereas OGs with species-specific m^6^A methylation were mainly enriched in signal transduction and response to stress (supplementary fig. 5, Supplementary Material online). These results revealed evolutionary dynamics of m^6^A methylation, that should be related to the evolution of gene function. Then, we identified putative functional counterparts of the m^6^A methyltransferases (MT-A70, FIP37, VIRILIZER, and HAKAI), and reader proteins (YTH-domain proteins) in 13 species (supplementary data 6, Supplementary Material online). Statistical analysis showed that the global m^6^A levels for these species were positively correlated with the number of writer and reader genes (supplementary fig. 6, Supplementary Material online); such correlation would be indicative of coevolution between m^6^A methylene and m^6^A functional counterpart genes. Together, these results suggested that species differentiation and genome evolution are accompanied by conservation and variation in the m^6^A methylene.

**Divergence of m^6^A Modification in OGs with Different Evolutionary Origins**

Previous phylostratigraphy analysis showed that OGs between early diverged species were older than those between recently diverged species (Domazet-Loso et al. 2007). Here, we broadly classified OGs with diverse evolutionary origins into six categories: dicot genes orthologous to dicot genes (DDs), dicot genes orthologous to monocot genes (DMs), dicot genes orthologous to Phy. patens genes (DPs), monocot genes orthologous to monocot genes (MMs), monocot genes orthologous to dicot genes (MIDs), and monocot genes orthologous to Phy. patens genes (MPs). Therefore, of the dicot genes, DPs had the oldest evolutionary origins, followed by DMs, and DDs. Likewise, of the monocot genes, MPs had the oldest evolutionary origins, followed by MIDs, and MMs. For both dicot and monocot genes, the mean m^6^A methylation ratio increased with evolutionary age (fig. 2A): DDs ((49.1 ± 4.8)% < DMs ((52.2 ± 4.7)% < DPs...
Fig. 1. Characterization of $m^6$A methylomes in 13 plant species. (A) Overview of genome size, gene number, and transcriptome-wide $m^6$A methylation ratios in 13 plant species. (B) The $m^6$A methylation ratio is negatively correlated with genome size. (C) The $m^6$A methylation ratio is negatively correlated with gene number. (D) Comparison of $m^6$A methylation ratios between OGs and SGs. Statistical analysis conducted using the Student’s t-test. **P<0.001.

Fig. 2. $m^6$A modifications were preferentially maintained in ancient genes. (A) Comparison of $m^6$A methylation ratios between interspecific OGs and intraspecific OGs. DDs, dicot genes orthologous to dicot genes; DMs, dicot genes orthologous to monocot genes; DP, dicot genes orthologous to Physcomitrella patens genes; MMs, monocot genes orthologous to monocot genes; MDs, monocot genes orthologous to dicot genes; MPs, monocot genes orthologous to Phy. patens genes. Statistical analysis conducted using the Student’s t-test. *P<0.05; **P<0.001. (B) The $m^6$A methylation ratios decreased from ancient PSs to young PSs.
these species. Here, the evolutionary rate (we first analyzed the correlation between m6A m eth yl ome c ations throughout plant evolution. To answer this question, evolutionarily ancient genes. divergence and species differentiation time. We used the Supplementary Material online). Then, we analyzed the number of shared m6A OGs among these six categories, and found that the intersections between DDs and DP s were smaller than those between DDs and DMs in dicot species, and the intersections between MM s and MP s were smaller than those between MM s and MDs in monocot species (supplementary fig. 8, Supplementary Material online).

We further investigated the changes in m6A methylation ratios for OGs with different evolutionary origins. These are defined in supplementary figure 9A, Supplementary Material online, a phylor stratigraphic tree of genes built using 34 plants species with clear taxonomic classification. In total, 543,853 OG s were assigned to 29 internodes on the phylor stratigraphic tree (named as phylor stratigraphic stages [PSs]) according to the emergence of their founders in the phylogeny (supplementary fig. 9B, Supplementary Material online). The evolutionary origins of the OGs were identified by tracing each set to its earliest common ancestor and assessing the age of the OGs in the corresponding internodes. OGs at the Chloroplastida internode have the oldest evolutionary origins, and the evolutionary origins of OGs in the subsequent PSs gradually get newer as they advance along the phylor stratigraphic paths. Figure 28 shows six phylor stratigraphic paths for the 13 species. We found that for each of these six phylor stratigraphic paths, the m6A methylation ratio for OGs decreased gradually from old PSs to young PSs (fig. 2B). These results indicated that m6A modifications were maintained in evolutionarily ancient genes.

Evolutionary Dynamics of m6A Modification between OGs

Previous studies demonstrated that ancient genes have been under strong purifying selection because of their fundamental role in plant survival (Guo 2013; Arendsee et al. 2014). This prompted us to ask whether dynamic selection forces may act on plant genomes to shape the diversity of m6A modifications throughout plant evolution. To answer this question, first, we analyzed the correlation between m6A methylome divergence and species differentiation time. We used the Spearman correlation coefficient (SCC) to examine the similarity of m6A abundances between paired OGs in any two given species, and the divergence of m6A methylomes between these two species was measured as 1–SCC. We found that the divergence of m6A methylomes was positively correlated with species differentiation time (fig. 3A). We then investigated the relationship between the evolutionary rate (ω) and the divergence of m6A status for pairs of OGs among these species. Here, the evolutionary rate (ω) was defined as the ratio of nonsynonymous substitutions (K_a)/synonymous sites (K_s). The divergence of m6A status indicated the coexistence or loss of m6A modifications within paired OGs. In order to eliminate the ambiguity resulting from genomic duplication events, we only used single-copy OGs expressed (TPM>1) in seven diploid species: A. thaliana, G. arbo reum, P. vulgaris, Sol. lycopersicum, S. bicolor, Aeg. tauschii, and O. sativa. In the four PSs shared by these seven diploid species (fig. 3B), both the frequency of sequence mutation (supplementary fig. 10, Supplementary Material online) and evolutionary rate (fig. 3C) of OGs increased from older to younger PSs, meaning that OGs originating from old PSs experienced a higher intensity of purifying selection and slower rates of evolution than those originating from young PSs. A similar increase was also observed for the divergence-m6A ratio of OGs originating from Chloroplastida to Spermaphyta PSs (fig. 3C), suggesting that evolutionarily ancient OG pairs had less divergence in m6A methylation than younger OG pairs. Further Pearson correlation analysis revealed a positive correlation between the divergence-m6A ratio and the evolutionary rate of paired OGs (Pearson’s r = 0.59, P = 3.13 × 10^{-9}; fig. 3D). These results together indicate evolutionary constraints on m6A divergence of OG pairs such that the earlier an OG pair originated, the less m6A modification diverged.

Divergence of m6A Modification between Homologous Genes in the Context of Polyploidization in Plants

Polyploidization (whole-genome duplication [WGD]) is a common phenomenon in plants (Otto 2007; Soltis et al. 2015; Cheng et al. 2018) and an essential component of plant genome evolution and innovation (Freeling 2009; Van de Peer et al. 2009). For example, maize and soybean (Gly. max) have undergone recent autopolyploidization and subsequent rediploidization, and upland cotton (G. hirsutum), emmer wheat (T. dicoccon), and common wheat (T. aestivum) have undergone allopolyploidization. Our recent study, moreover, revealed an association between m6A modification and duplicate gene evolution in maize (Miao et al. 2020). By sequencing multiple polyploid plant species with m6A methylomes in this study, we further explored the evolutionary implications of m6A modification in the context of plant genome polyploidization, with a special emphasis on the evolutionary divergence of m6A modifications triggered by autopolyploidization and allopolyploidization.

First, we identified subgenomes of each sequenced polyploid species by analyzing synteny relationships between homologous genes in polyploids and their diploid relatives (e.g., Gly. max vs. P. vulgaris, Z. mays vs. S. bicolor; see Materials and Methods). Then, we classified homologous gene pairs (both partners expressed with TPM>1) into three groups (fig. 4A): IM pairs (both of the paired genes modified with m6A), DM pairs (only one of the paired genes modified with m6A), and NM pairs (none of the paired genes containing m6A peaks). Many sequence variations were found in the duplicated genes after polyploidization (Lockton and Gaut 2005; Freeling and Thomas 2006; Li et al. 2021). Next, we compared the sequence differences in IM, DM, and NM pairs. For both the autopolyploid species and the allopolyploid species, we found less sequence variation in IM than in DM and NM pairs (fig. 4B and supplementary fig. 11, Supplementary Material online). In
addition, both $K_a$ and $K_s$ from IM pairs were significantly lower than from DM pairs. Notably, NM pairs had the largest $K_a$ and $K_s$ values (fig. 4C and supplementary fig. 12, Supplementary Material online). Focusing on 3'-UTR, where m^6A peaks were mainly enriched, we found that sequence identity within IM pairs was significantly higher than that within DM pairs (fig. 4D and supplementary fig. 13, Supplementary Material online). In addition, we compared the length of 3'-UTR sequences between homologous gene pairs. As shown in supplementary figure 14A, Supplementary Material online, the differences of 3'-UTR length between homologous gene pairs were mainly distributed into peaks that span — 500 to 500 nucleotides (zero indicates the same length of two partners of homologous gene pairs). Focusing on the highest part of these peaks, around zero, the peak of IM pairs is significantly higher than that of DM pairs.
(supplementary fig. 14A, Supplementary Material online), meaning less difference of 3'-UTR length between IM pairs compared with DM pairs (supplementary fig. 14B, Supplementary Material online). These results suggest that divergence of m^6A modifications may be related to the occurrence of spontaneous mutations in homologous genes during the evolutionary process postpolyploidization.

In addition to these similarities, we also found differences between autopolyploid and allopolyploid species. As shown in figure 4E, the divergence-m^6A ratio of homologous genes...
(i.e., DM/[IM+DM+NM]) in autopolyplid species (15.2–20.4%) was significantly higher than in allopolyploid species (9.2–11.1%; Wilcoxon test, P < 0.05). This may be the result of a higher frequency of sequence substitutions (Ks and Kt) in homologous genes between autopolyplid species and their diploid relatives than between allopolyploid species and their diploid relatives (fig. 4F). Alternatively, this may be the result of higher m6A methylation ratios for duplicates than for singletens in autopolyplid species (soybean and maize; supplementary fig. 15, Supplementary Material online). That is, the autopolyplidization–rediploidization cycle included the loss of one copy of many duplicate gene pairs (i.e., gene fractionation) (Lockton and Gaut 2005; Freeling and Thomas 2006). The m6A modifications were retained on duplicate genes during the autopolyplidization–rediploidization cycle.

Based on these observations, we inferred that postpolyplidization, sequence variations may be important factors affecting the evolutionary divergence of m6A modifications. The asymmetric retention of m6A modifications between duplicates and singletens during the autopolyplidization–rediploidization cycle may provide for divergence of m6A modification in homologous gene pairs between autopolyplid species and their diploid relatives.

**Number of Family Members Was Correlated with m6A Methylation Ratio during Gene Family Expansion**

Unlike animals, plants cannot move away from unexpected and harsh environments, thus long-term selective pressures leave genetic and genomic signatures in plant enduracnce. In addition to WGD, local-genome duplication, which results in significant expansions of specific gene families, is also an important factor in shaping natural variation for adaptation in plants (Rizzon et al. 2006; Freeling 2009; Wang, Yang, et al. 2019). To pave the way for understanding the evolutionary consequences of m6A modification in the context of gene family expansion, we constructed a phylogenetic tree of local-genome duplication events using genes from 27 diploid species that were reported to have not experienced WGD events since they split from their close relatives (supplementary fig. 16, Supplementary Material online). Our results showed that the gene family either expands or contracts over time at a rate of approximately 0.0011 per gene every million years. This is comparable to the rate previously estimated for Drosophila (0.0012) (Hahn et al. 2007), mammals (0.0016) (Demuth et al. 2006), and plants (0.0014) (Guo 2013). We identified 308 significantly expanded gene families shared by seven angiosperm species with m6A profiling in this study (supplementary data 7, Supplementary Material online). The methylation ratios of these significantly expanded gene families were highly positively correlated (fig. 5A), implying interspecific conservation of evolutionary patterns of m6A modification in the context of gene family expansion. For each angiosperm species, the m6A methylation ratios for genes within significantly expanded gene families were lower than those for genes of the whole genome (fig. 5B). In addition, the m6A methylation ratio increased and the mean number of gene family members decreased along phylogenetic timescale from old PSs to young PSs (fig. 5C). Moreover, the m6A methylation ratio was negatively correlated with the number of gene family members in each PS (supplementary fig. 17, Supplementary Material online) and each species (fig. 5D). These results suggested that elimination of m6A-modified genes during the evolutionary process of gene family expansion may be triggered by local-genome duplications.

The evolutionary correlation between methylation ratio and number of family members was also observed in different types of gene families (e.g., protein kinase [PK], transcription factor [TF], and disease-resistance protein [R]). The PK families are enzymes that can regulate the biological activity of proteins through phosphorylation of specific amino acids. Thirty-eight PK families with more than five family members could be divided into two clusters (low m6A methylation ratio [LMC] and high m6A methylation ratio [HMC]; fig. 5E). The number of family members was significantly larger in LMC than in HMC, and the coefficient of variation in the number of family members was positively correlated with that in the m6A methylation ratio across species (fig. 5E).

Our results were similar in the TF families (supplementary fig. 18, Supplementary Material online). The R families are plant immune receptors that guard the plant against pathogens. Members of these families were identified based on the presence of nucleotide-binding site (NBS) and leucine-rich repeat (LRR) domains. Phylogenetic analysis showed that NBS-LRR genes could be classified as two subfamilies (supplementary fig. 19, Supplementary Material online): Toll/interleukin-1 receptor in the amino-terminal domain (TNL) and coiled-coil motifs in the amino-terminal domain (CNL). Proportions of CNLs were grouped into clades dominated by TNLs, suggesting that homologous recombination or homogenization occurred between some members of the two subfamilies (supplementary fig. 19, Supplementary Material online). Figure 5F depicts the negative correlation between the number of R family members and the m6A methylation ratio. This negative correlation was also present in subfamilies with distinct species compositions (supplementary fig. 20 and data 8, Supplementary Material online).

Together, these results illustrate evolutionary correlation between m6A modification and gene family expansion. That is, ancient gene families have experienced numerous expansion events that led to an increase in the number of family members accompanied by a decrease in m6A methylation ratio for family members, resulting in asymmetric expansion between m6A-modified members and non-m6A members.

**Effects of m6A Modification on Gene Expression and Translational Efficiency in Genomic Duplications**

Previous studies have demonstrated that a variety of gene dosage compensation mechanisms at both the transcriptional and translational levels have evolved to alleviate harmful stoichiometric imbalances caused by genomic duplications (Veitia et al. 2008; Edger and Pires 2009; Song et al. 2020). For example, DNA methylation could rebalance gene dosage after gene duplication by inhibiting the initiation of transcription of duplicate genes (Chang and Liao 2012). Given that the effect of m6A methylation on translational efficiency is multifaceted in its strength and genic location...
**Fig. 5.** Evolutionary consequences of m\(^6\)A modification in the context of gene family expansion. (A) The pairwise correlation of methylation levels of significantly expanded gene families among seven diploid species. (B) Decreased m\(^6\)A methylation ratios for genes within significantly expanded gene families. (C) The m\(^6\)A methylation ratio increased and the mean number of gene family members decreased along the phylogenetic timescale from ancient PSs to young PSs. (D) The m\(^6\)A methylation ratio was negatively correlated with the number of gene family members in each species. (E) For PK families in seven diploid species, the number of family members was significantly larger in LMC than in HMC, and the coefficient of variation in family members was positively correlated with the m\(^6\)A methylation ratios. LMC, clusters with LMC ratio; HMC, clusters with HMC ratio. (F) For R gene families, the number of family members was negatively correlated with the m\(^6\)A methylation ratio. In (B), statistical analysis conducted using the \(\chi^2\) test. **P<0.001. In (E), statistical analysis conducted using the Wilcoxon test. **P<0.001.
(Luo et al. 2020), we wondered what the evolutionary effects of m^6A methylation on gene expression and translational efficiency are in the context of genomic duplications. Integrative analysis of the m^6A-seq and ribosome profiling data sets (Luo et al. 2020) showed that both the expression level and translational efficiency of m^6A-modified genes were significantly higher than those of non-m^6A genes in maize (supplementary fig. 21, Supplementary Material online), therefore, we focused on duplicated genes identified in the maize genome. We identified 7,536 locally duplicated genes from 2,566 gene families. Members of gene families with the largest numbers of members had the lowest m^6A methylation ratio (fig. 6A), average translational efficiency (fig. 6B) and expression level (supplementary fig. 22A, Supplementary Material online). These results indicated that, with gene family expansion, the reduced level of m^6A modification may attenuate the expression level and translational efficiency of gene duplicates caused by local-genome duplication (supplementary figs. 23 and 24, Supplementary Material online). In addition, the translational efficiency of gene duplicates retained after WGD was significantly higher than that of singletons (fig. 6C). Notably, for duplicate genes, the translational efficiency of m^6A-modified genes was significantly higher than that of non-m^6A genes, but this difference was not observed in singletons (fig. 6D), implying biased impact of m^6A modification on translational efficiency in WGD-duplicated genes. In addition, the expression level of m^6A-modified genes was significantly higher than that of non-m^6A genes for both duplicate and singleton genes (supplementary fig. 22B, Supplementary Material online).

Collectively, our results showed that the evolutionary consequences of m^6A modification and the influence on translational efficiency of duplicated genes varied depending on the type of duplication event.

**Discussion**

Although the progress in dissecting the role of m^6A modification in gene regulation is expedited in the plant kingdom (Shen et al. 2019), our understanding of the mechanism of evolution of m^6A modification remains limited. Some biological functions of m^6A modification appear to be evolutionarily conserved in both mammals and plants (Zhao, Roundtree et al. 2017), however, many previous observations of m^6A modification in mammals have been challenged in plant species (Yue et al. 2019). Furthermore, some components of m^6A modification have been present since early plant evolution, but some components were lost during plant evolution and speciation (Liang et al. 2020). Thus, our view of the evolutionary conservation and divergence of m^6A modification across plants species has remained blurry.

In this study, we systematically compared m^6A methylomes among 13 representative plant species, and provided novel insights into the patterns, processes, and influences of m^6A evolution. As an ancient chemical modification, m^6A modification may undertake basic molecular functions such as RNA metabolism and export, and therefore be widespread and conserved to a certain extent in plants. Transcriptome-wide methylation levels differed little between evolutionarily adjacent species (fig. 1). Physcomitrella patens was an outlier, with the smallest genome and the highest methylation level. These results suggest that variation in the m^6A methylome was relatively stable at short evolutionary timescales, whereas at long evolutionary timescales, the methylation levels varied as a result of species differentiation and genome evolution.

Natural selection can drive the evolution of plants through gene sequence mutation and copy number variation. m^6A modifications were mainly enriched in the 3'-UTR gene region such that mutations in this region may affect the evolutionary divergence of the m^6A methylome. In this study, by comparing OG pairs between polyploids and their diploid relatives, we found that m^6A modifications were maintained in ancient genes, and 3'-UTR sequences were more conserved in IM pairs than in DM pairs. We speculated that throughout the long evolution of the plant kingdom, ancient genes have been under strong purifying selection, since ancient genes are mostly fundamental for plant survival, and this resulted in slower evolution and higher conservation of these sequences, which could in turn favor the retention of m^6A modification in ancient OGs. The m^6A modifications in turn affected the evolutionary fate of genes following duplication events by changing gene dosage. Consistent with these ideas, recent studies in mammals have shown that m^6A modifications gained during evolution were subject to positive selection, whereas conserved m^6A peaks were under purifying selection (Ma et al. 2017). Comparative study of m^6A methylomes between humans and mice found that ~40% of the m^6A peaks in the 3'-UTR were subject to purifying selection, and the m^6A sites gained by humans were subject to positive selection and associated with disease (Zhang et al. 2020). These results preliminarily illustrated the conservation of m^6A modification, which contributed the evolutionary conservation and novelty of some genes key to speciation and species differentiation.

Changes in the number of copies of genes can enhance a species’ ability to adapt to change in its environment. Our results indicate that throughout the process of gene family expansion, m^6A modification influenced the gene expression and translational efficiency of family members differently, possibly due to differences in duplication events. Gene families with many copies can be divided into two categories. One includes small-scale local duplication events, which resulted from expansion in short evolutionary time leading to gene dosage proliferation. According to the Gene Balance Hypothesis, duplicates of dosage-sensitive genes tend to be underretained or eliminated in order to maintain proper balance (Freeling 2009). The association between m^6A modification and copy number variation and the influence of m^6A modification on gene expression and translational efficiency may be a mechanism of dosage compensation that has evolved to alleviate harmful dosage imbalances. The other includes WGD events that increased the dosage of all genes simultaneously. The positive contribution of m^6A modification to gene expression and translational efficiency of duplicate genes appears to support the Increased Gene Dosage Hypothesis, which predicts that the WGD and diploidization
cycle yields some highly expressed genes that may be beneficial and in turn result in purifying selection to retain both gene duplicates (Conant and Wolfe 2008).

Based on the results in this study, we propose a conceptual model to explain the evolutionary role of the m6A methylome in plants (fig. 7). The process of plant speciation and genome evolution was accompanied by the divergence of m6A modification in OGs, and m6A modifications were maintained in evolutionarily ancient genes, increasing the diversity of m6A methylomes across plants. In the context of the WGD-diploidization cycle, variations in gene sequences affected the gain and loss of m6A loci, and m6A-modified genes were preferentially retained as duplicates, potentially leading to substantial effects on gene fractionation. In addition, local-genome duplication triggered variation in copy number among family members that was accompanied by the divergence of m6A modification. With gene family expansion, the reduced level of m6A modification attenuated the expression level and translational efficiency of duplicated genes, and this could be a mechanism of evolutionary reaction to gene dosage imbalances caused by local-genome duplication.

**Materials and Methods**

**Plant Material and Growth Conditions**

*Arabidopsis thaliana* (Col-0) was sown in nutrient soil and cultivated in a greenhouse at 22 °C under a 16 h light/8 h dark cycle. On the 14th day after germination, all the above-ground parts of the seedlings of *A. thaliana* were collected and frozen in liquid nitrogen immediately and stored at −80 °C. *Gossypium arboreum* (Shixiya1, AA genome) and *G. hirsutum* (TM-1, AADD genome) were grown at 28 °C, and *P. vulgaris* (G19833) and *Gly. max* (Williams 82) were grown at 20 °C under a 16 h light/8 h dark cycle. The above-ground parts of the seedlings of *G. arboretum*, *G. hirsutum*, *P. vulgaris*, and *Gly. max* were collected on the 18th day after germination. *Sorghum bicolor* (BTx623) and *Z. mays* (B73) were sown in nutrient soil and cultivated in a greenhouse at 25 °C under a 16 h light/8 h dark cycle. *Aegilops tauschii* (AL8/78, DD genome), *T. dicoccoides* (Zavitan, AAB genome), and *T. aestivum* (Chinese Spring, AABBDD genome) were germinated in petri dishes and transplanted into soil under a 16 h light/8 h dark cycle at 22 °C. *Oryza sativa* (Nipponbare) was grown at 28 °C and under a 16 h light/8 h dark cycle. When seedlings of these six monocot species developed three fully expanded leaves, the above-ground parts were collected and frozen at −80 °C. The single shoot tip of *Phy. patens* (Gransden 2004) was taken from the top of the fronds and inserted into BCD medium without ammonium tartrate at 25 °C and under a 16 h light/8 h dark cycle. After 4 weeks, the stems and leaves were mixed together and frozen in liquid nitrogen immediately and stored at −80 °C. For each species, three biological replicates were used for the experiments described below.

![Fig. 6. Evolutionary effects of m6A modification on translational efficiency.](image)

**A** Gene families with more copies had lower m6A methylation levels. **B** Gene families with more copies had lower average translational efficiency. **C** Comparison of translational efficiency between duplicate genes and singletons in the context of a WGD event. **D** Comparison of translational efficiency between m6A-modified genes and non-m6A genes in duplicates and singletons, respectively. In (C) and (D), statistical analyses were conducted using the Student’s t-test. **P < 0.001.
RNA Isolation and High-Throughput m^6^A-Seq

Total RNA was extracted and purified using TRizol reagent (Invitrogen) according to the manufacturer’s instructions, and NanoDrop ND-1000 (NanoDrop) was used to quantify its amount and purity. RNA integrity was assessed using Bioanalyzer 2100 (Agilent, CA) with RIN number > 7.0 and confirmed using denaturing agarose gel electrophoresis. Polyadenylated (PolyA+) mRNA selection was performed using Dynabeads Oligo (dT) 25-61005 (Thermo Fisher) with two rounds of purification from 30 µg total RNA. The poly(A) RNA was fragmented into small pieces using Magnesium RNA Fragmentation Module (NEB, Cat. e6150) under 86 °C for 7 min. The fragmented RNA was then used to construct the input library directly following conventional RNA-seq and the immunoprecipitation (IP) library was enriched using m^6^A-specific antibodies (No. 202003, Synaptic Systems, Germany) in IP buffer (50 mM Tris–HCl, 750 mM NaCl, and 0.5% Igepal CA-630) for 2 h at 4 °C. The Input and IP were reverse-transcribed using SuperScript II Reverse Transcriptase (Invitrogen, Cat. 1896649). U-labeled second-stranded DNAs were synthesized with Escherichia coli DNA polymerase I (NEB, Cat. m0209), RNase H (NEB, Cat. m0297), and dUTP Solution (Thermo Fisher, Cat. R0133).

An A-base was added to the blunt ends of each strand. Adapters containing a T-base overhang were used for ligating the A-tailed fragmented DNA. Single- or dual-index adapters were ligated to the fragments, and size selection was performed with AMPureXP beads. After heat-labile UDG enzyme (NEB, Cat. m0280) treatment of the U-labeled second-stranded DNAs, the ligated products were amplified with PCR under the following conditions: initial denaturation at 95 °C for 3 min; eight cycles of denaturation at 98 °C for 15 s, annealing at 60 °C for 15 s, extension at 72 °C for 30 s, and final extension at 72 °C for 5 min. The average insert size for the final cDNA library was 300 ± 50 bp. Sequencing was carried out on the Illumina NovaSeq 6000 platform (LC-Bio Technology Co., Ltd., Hangzhou, China) with a 2 × 150-bp paired-end strategy (PE150).
Bioinformatic Analysis of m6A-Seq Data

Raw reads from 13 species were preprocessed by trimming the adapter sequences, low-quality bases, and undetermined bases using fastp v0.20.0 (Chen et al. 2018) with default parameters. Clean reads were mapped against their corresponding reference genomes using STAR v2.7.3a (Dobin et al. 2013) with the following settings: outFilterMismatchNmax, 0; outSAMattrHisstart, 0; outFilterMultimapNmax, 1; alignIntronMin, 10; alignIntronMax, an estimated value that covers 99.9% of all intron lengths in the analyzed species. The reference genomes and gene annotations in GFF3 format of A. thaliana, Z. mays, Aeg. tauschii, T. dicocoides, and T. aestivum were obtained from Ensembl Plants (v43) (Howe et al. 2020); G. arboreum from Wang, Wang, et al. (2019); G. hirsutum from CottonGen database (Yu et al. 2014); Sol. lycopersicum from Sol Genomics Network (Lin et al. 2005); P. vulgaris, Gly. max, O. sativa, and S. bicolor from JGI (phytozome v12) (Goodstein et al. 2012) (supplementary data 9, Supplementary Material online). For strand-specific sequencing data, bam files were split into two sets based on samflags: forward strand set (-f 80 [first in pair] and -f 144 [second in pair]), SAMtools v1.9 (Li et al. 2013). Stringtie v2.0 (Pertea et al. 2015) was used to estimate gene expression abundances from input libraries by calculating transcripts per million (TPM) (stringtie -e -j 10; –rf for strand-specific parameter). m6A peak abundance was calculated as (mapped fragments input x Total mapped fragments input)/(mapped fragments input x Total mapped fragments input) using DiffBind package (Ross-Innes et al. 2012). The called peaks with low m6A abundance (<2-fold enrichment) or within lowly expressed genes (TPM<1) were discarded.

GO Enrichment Analysis

The GO annotations were retrieved from Ensembl Plant Biomart (Kinsella et al. 2011), and CottonGen (Yu et al. 2014). GO enrichment analysis was performed using the enricher function from clusterProfiler R-package for the hyper-geometric test (Yu et al. 2012).

Phylostratigraphic Analysis

The phylostratigraphic tree of genes or gene families was constructed using TimeTree (Kumar et al. 2017) according to methods detailed in previous studies (Domazet-Loso et al. 2007; Guo 2013; Lei et al. 2017). To obtain high-quality phylostratigraphic results, we analyzed 34 representative species having clear taxonomic and phylogenetic relationships (Wu et al. 2020). The orthologous genes were assigned to different phylostrata representing different evolutionary ages. If there was an unnamed node in phylostrata, it was renamed using the combination of first letters of the child node names (i.e., AT represented Aeg. tauschii and Triticum, BM represented Brassicaceae and Malvaceae).

Divergence of m6A Methyomes

The species tree and divergence time of internal nodes were obtained from the TimeTree database (http://timetree.org/) using “BUILD A TIMETREE” function (Kumar et al. 2017). The m6A abundance divergence was defined as 1−ρ, where ρ was the Spearman’s correlation coefficient between m6A methylomes in two species.

Identification of Subgeneromes in Species Having Undergone Recent Polyploidyization

The paralogous gene pairs between subgenomes in Z. mays, Gly. max, and T. aestivum, and their syntenic counterparts in sister species S. bicolor/Z. mays and P. vulgaris/Gly. max were obtained from previously published data sets (Zhao, Zhang, et al. 2017; Brohammer et al. 2018; Ramirez-Gonzalez et al. 2018). The subgenomes of G. hirsutum were classified based on chromosomes (A and D subgenomes). The syntenic counterparts in G. arboreum/G. hirsutum (A subgenome), Aeg. tauschii/T. aestivum (D subgenome), and T. dicocoides (A subgenome)/T. aestivum (A subgenome), and T. dicocoides (B subgenome)/T. aestivum (B subgenome) were detected using SynMap within CoGe with a 1:1 quota-align ratio and default parameters (Haug-Baltzell et al. 2017). In maize, genes not included in duplicates formed by WGD were further classified into four duplication types using DupGen_finder (Qiao et al. 2019) with default parameters: tandem duplicates, proximal duplicates, transposed duplicates, and dispersed duplicates.

Analysis of Synonymous and Nonsynonymous Substitutions in OGs

The one-to-one OG pairs from either eight diploid species (A. thaliana, G. arboreum, P. vulgaris, Sol. lycopersicum, S. bicolor, Aeg. tauschii, O. sativa, and Phy. patens) or their sister species (G. arboreum/G. hirsutum, P. vulgaris/Gly. max, S. bicolor/Z. mays, Aeg. tauschii/T. dicocoides/T. aestivum) were extracted from OrthoFinder, and their protein
sequences were aligned using paraAT v2.0 (Zhang et al. 2012). The synonymous and nonsynonymous substitutions, and their resulting $K_s/K_a (\omega)$ values were calculated using KaKs_calculator v2.0 (Wang et al. 2010) with the Model Averaging method.

Analysis of Sequence Divergence
To estimate the sequence divergence between OG pairs, nucleotide sequences for gene pairs were aligned using needle in EMBL analysis tools (Chojnacki et al. 2017) with parameters -gapopen=10 and -gapextend=0.5. The sequence divergence between OG pairs from sister species genomes was calculated by ortholog comparisons. For example, sequence divergence in Z. mays genes was estimated by comparison with their orthologous in S. bicolor. For the aligned sequences of paired genes, gaps in the gene from the species that had not experienced recent polyploidy events were removed. For example, in S. bicolor/Z. mays gene pairs, the gaps in S. bicolor genes were removed. Aligned sequences were further divided into three nonoverlapping parts: 5′-UTR, coding sequence, and 3′-UTR. Only gene pairs with all three parts and each part longer than 100 bp were used in subsequent analysis. Each part was divided into ten bins and the mutation frequency for each bin was calculated using “inconsistent base/base number.” The sequence identity of 3′-UTR was calculated using “identical base/base number.”

Likelihood Analysis of Gene Gain and Loss
The program CAFE (computational analysis of gene family evolution) v4.2.1 (De Bie et al. 2006) was used to analyze the evolution of gene families based on a probabilistic graphical model. To reduce the influence from plant polyploidization events, 27 diploid species were selected to identify expansion and contraction of gene families. The number of genes in gene families was obtained from OrthoFinder results. Gene families with more than 100 gene copies in one or more species were removed from this analysis. The dated species tree was downloaded from the TimeTree database. We set the birth–death parameter to one and the $\omega$ value to 0.01.

Identification of TFs, Kinases, and R Genes
Transcription factors and kinases were obtained by scanning consensus rules and HMM profiles using iTAK v1.7 (Zheng et al. 2016). The identification of R genes was performed using HMMER v3 (Potter et al. 2018) and CD-search (Marchler-Bauer and Bryant 2004). The protein sequences from 13 species were scanned using HMMER and Hidden Markov Model of NB-ARC (Pam accession: PF00931) (El-Gebali et al. 2019). Genes with NB-ARC domain (E-value <1e-5) were further authenticated. The amino-terminal domain (TIR/coiled-coil/other) and carboxyl-terminal domain (LRR) were recognized by CD-search. Only genes with both NBS and LRR domains were defined as R genes.

Sequence Alignment and Estimation of the Phylogenetic Tree
Multiple sequence alignment of R genes was performed using muscle v3.8.1551 (Edgar 2004) with default parameters. The maximum-likelihood phylogenetic tree was then inferred with IQ-Tree v2.0.3 (Minh et al. 2020) with parameters -m VT+F+F9, -bb 1,000, and -T AUTO. All trees were rooted using P25941 from streptomyces as the outgroup.

Calculation of Translational Efficiency
The translational level and corresponding transcriptional level of maize genes were obtained from a previous study (Luo et al. 2020). The translational efficiency of each gene was calculated by “TPM (transcriptional level)/TPM (transcriptional level)” as previously described (Lei et al. 2015; Luo et al. 2020).

Statistical Analysis
The Wilcoxon test, the $\chi^2$ test, and the Student’s t-test were respectively performed using the wilcoxon.test, chisq.test, and t.test functions in R package.

Supplementary Material
Supplementary data are available at Molecular Biology and Evolution online.

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Author Contributions
C.M. and Z.M. conceived the project. T.Z., B.X., and Z.M. performed the data analysis. Y.Q. prepared plant materials. Z.M., T.Z., B.X., and C.M. wrote the article.

Data Availability
The raw-sequencing data generated in this study have been deposited in the Genome Sequence Archive (Wang et al. 2017) in National Genomics Data Center (National Genomics Data Center Members and Partners 2020), Beijing Institute of Genomics (BIG), Chinese Academy of Sciences, under accession number CRA004114 that are publicly accessible at https://bigd.big.ac.cn/gsa. The analysis
pipeline and codes used in this study have been deposited in GitHub (https://github.com/cma2015/m6A_Evolution).

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