Epitranscriptomic RNA Methylation in Plant Development and Abiotic Stress Responses

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Recent advances in methylated RNA immunoprecipitation followed by sequencing and mass spectrometry have revealed widespread chemical modifications on mRNAs. Methylation of RNA bases such as N6-methyladenosine (m6A) and 5-methylcytidine (m5C) is the most prevalent mRNA modifications found in eukaryotes. In recent years, cellular factors introducing, interpreting, and deleting specific methylation marks on mRNAs, designated as “writers (methyltransferase),” “readers (RNA-binding protein),” and “erasers (demethylase),” respectively, have been identified in plants and animals. An emerging body of evidence shows that methylation on mRNAs affects diverse aspects of RNA metabolism, including stability, splicing, nucleus-to-cytoplasm export, alternative polyadenylation, and translation. Although our understanding for roles of writers, readers, and erasers in plants is far behind that for their animal counterparts, accumulating reports clearly demonstrate that these factors are essential for plant growth and abiotic stress responses. This review emphasizes the crucial roles of epitranscriptomic modifications of RNAs in new layer of gene expression regulation during the growth and response of plants to abiotic stresses.

Keywords: abiotic stress, epitranscriptome, RNA metabolism, RNA methylation, RNA modification

INTRODUCTION

Epigenetic regulation of gene expression via DNA methylation and histone modifications is an important strategy for living organisms to achieve fine-tuned regulation of developmental processes or responses to environmental cues. Similar to DNA methylation in epigenetic regulation, posttranscriptional RNA modifications are emerging as important “epitranscriptomic” regulatory networks in recent years (Saletore et al., 2012; Meyer and Jaffrey, 2014). Over 150 different chemical modifications on mRNAs, tRNAs, and rRNAs are currently known for all kingdoms of life (Cantara et al., 2010; Boccaletto et al., 2018). Among diverse modifications found on mRNAs, N6-methyladenosine (m6A) is the most prevalent modification in both plants and animals (Liu and Pan, 2016; Covelo-Molares et al., 2018). Recent advances in methylation RNA sequencing (Met RNA-seq) and deep RNA sequencing have revealed transcriptome-wide m6A methylation patterns in plants as well as in animals (Luo et al., 2014; Wang et al., 2015a; Cui et al., 2017). These modifications within mRNAs can affect multiple steps of transcript’s fate, including splicing (Haussmann et al., 2016; Xiao et al., 2016), nucleus-to-cytoplasm export (Zheng et al., 2013), RNA turnover (Du et al., 2016; Mauer et al., 2017; Wei et al., 2018), and translation (Meyer et al., 2015; Wang et al., 2015b; Choi et al., 2016).
The level and status of RNA methylation in cells depend on two crucial proteins: RNA methyltransferase (MT) designated as “writer” and RNA demethylase (DMT) designated as “eraser” (Figure 1). In addition to these two essential proteins required for the addition and removal of methyl groups on RNAs, a third protein designated as “reader” is involved in the recognition and processing of methylated RNAs (reviewed in Meyer and Jaffrey, 2014). In animals, genes encoding m^6A writer (Ping et al., 2014; Schwartz et al., 2014), reader (Luo and Tong, 2014; Xu et al., 2014), and eraser (Jia et al., 2011; Zheng et al., 2013) proteins have been identified and characterized (Table 1). Notably, mutants lacking specific m^6A writer, reader, or eraser have displayed abnormal development and altered response to hypoxia and high temperatures, suggesting crucial roles of RNA methylation in animal development and adaptation to changing environmental cues (reviewed in Meyer and Jaffrey, 2014; Yue et al., 2015; Hsu et al., 2017).

Although these recent studies clearly point to the importance of RNA methylation and essential roles of writers, readers, and erasers in the development of animals, functions of these proteins in plants are just beginning to be uncovered. *Arabidopsis* contains functional orthologs of m^6A writer complex components, erasers, and reader proteins, some of which have been found to play essential roles in normal plant development (Bodi et al., 2012; Shen et al., 2016; Růžička et al., 2017; Arribas-Hernández et al., 2018; Scutenaire et al., 2018; Wei et al., 2018). All these aforementioned studies have emphasized the essential roles of RNA methylation in plant development. However, the identity and functions of most writers, readers, and erasers in plants are currently unclear. In this review, we systematically identified potential m^6A writers, readers, and erasers in *Arabidopsis* and rice (*Oryza sativa*) by comparing sequence homology to animal counterparts. We also reviewed multiple functions and potential significance of m^6A RNA methylation in the development and response of plants to diverse abiotic stresses.

### DIVERSE MODIFICATIONS FOR EUKARYOTIC RNAs

Over 150 different internal modifications on RNAs have been identified (Cantara et al., 2010; Boccaletto et al., 2018), with different degree, topology, and kinds of modifications between mRNAs, tRNAs, and rRNAs. For instance, approximately 17% of total nucleotides in tRNAs are modified, whereas only 2% of nucleotides in rRNAs are modified (Jackman and Alfonzo, 2013). Among diverse modifications identified for tRNAs and rRNAs, 2’-O-ribose methylation and pseudouridilation of rRNAs and 5-methylcytosine (m^5C) and 1-methylguanidine (m^1G) of tRNAs are the most abundant (Chou et al., 2017). Despite emerging roles of mRNA modifications in its processing and function, mRNA is less densely modified compared to tRNAs and rRNAs (Gilbert et al., 2016). Only a handful of different modifications have been identified so far in mRNAs, with N6-methyladenosine (m^6A) being the most abundant (Liu and Pan, 2016; Covelo-Molares et al., 2018). These methylations of bases can influence the structure of RNAs by increasing its hydrophobicity and disrupting the canonical Watson-Crick base pairing (Oerum et al., 2017; Väre et al., 2017).

Importantly, all organisms have evolved to cluster methylation marks in functionally critical positions rather than randomly distributing them along RNA molecules. Most of these modified bases in rRNAs are located at the interface between ribosomal large and small subunits corresponding to P-site and A-site (Sharma and Lafontaine, 2015; Sloan et al., 2017). Wobble positions 34 and 37 of the anticodon loop in tRNAs are the most frequently and diversely modified (Väre et al., 2017). These conserved modification patterns reflect the essential role of RNA methylation in ribosome structure and biogenesis, codon recognition and decoding, and translation initiation or elongation (Jackman and Alfonzo, 2013; Chou et al., 2017; Sloan et al., 2017; Väre et al., 2017). Similar to tRNAs and rRNAs, mRNAs are also methylated in specific regions. For instance, m^6A maps preferentially to the transcription start site, the stop codon, and the 3’ UTR (Dominissini et al., 2012; Luo et al., 2014; Meyer and Jaffrey, 2014), while m^5C is predominantly found in 3’ UTR and coding regions (Squires et al., 2012; David et al., 2017). Several studies have shown that m^5A methylation is frequently found in the start codon and the first splicing

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**Figure 1** Roles and structural characteristics of m^6A RNA methylation-related proteins. (A) Cellular factors introducing, deleting, and interpreting m^6A marks are methyltransferase (“writers”), demethylase (“eraser”), and RNA-binding protein (“reader”), respectively. (B) The writer complex consists of five components: MTA/B (methyltransferase A/B), FIP37 (FKBP12 interacting protein 37), VIR (Virilizer), HAKAI (for “destruction” in Japanese, a c-Cb1-like protein), erasers belong to AlkB-homology (ALKBH) family proteins, and readers are YT512-B homology domain (YTHD) proteins. Numbers at the C terminus indicate the number of amino acid residues in each Arabidopsis protein. MT_A70, methyltransferase, A70; FE2OG_OXY, Fe^2+ - 2-oxoglutarate dioxygenase domain; WTAP, WT1-associated protein.
site which influences translation (Dominissini et al., 2016; Safra et al., 2017). Clearly, the degree, topology, and non-random distribution of RNA modifications are crucial for its specific cellular functions.

WRITERS, ERASERS, AND READERS INVOLVED IN m^6A RNA METHYLATION AND RECOGNITION

Writers
Genes encoding m^6A writer complexes have been identified and characterized firstly in animals. Several proteins including methyltransferase-like 3 (METTL3) and METTL14, Wilms’ tumor 1-associating protein (WTAP), and Vir like m^6A methyltransferase-associated (VIRMA; KIAA1429) are known to form multicomponent m^6A writer complexes in animals (Shah and Clancy, 1992; Ping et al., 2014; Schwartz et al., 2014; Table 1).

Methyltransferase-like 3 is the principal enzyme exerting methyltransferase activity, while METTL14 has a supporting role forming a METTL3-METTL14 heterodimer (Sledz and Jinek, 2016; Wang et al., 2016). After the identification of METTL3 in mammals as a homolog of yeast methyltransferase IME4 (Shah and Clancy, 1992), its orthologs were identified in different species including Arabidopsis and Drosophila. At present, Arabidopsis orthologs of animal m^6A writer components have been identified, including MTA (ortholog of METTL3) and MTB (ortholog of METTL14).

TABLE 1
| Type  | Gene name | Arabidopsis gene ID | Target RNA | Function | Rice ortholog | Animal counterpart |
|-------|-----------|---------------------|------------|----------|---------------|-------------------|
| Writers | MTA | At4g10760 | m^6A | Embryo development | LOC_Os02g45110 | METTL3 |
| Writers | MTB | At4g09980 | m^6A | Embryo development | LOC_Os01g16180 | METTL14 |
| Writers | RR37 | At3g54170 | m^6A | Development | LOC_Os06g27970 | WTAP |
| Writers | VIRILIZER | At3g05680 | m^6A | Development | LOC_Os03g35340 | VIRMA |
| Writers | HAKAI | At3g01160 | m^6A | Development | LOC_Os10g35190 | |
| Writers | TRM4A | At4g40000 | m^5C | | LOC_Os08g37780 | |
| Writers | TRM4B | At2g22400 | m^5C | Root development | LOC_Os09g29630 | |
| Readers | YTH01 (ECT1) | At1g09810 | m^6A m^1A | | LOC_Os01g22630 | YTHDF1 |
| Readers | YTH02 (ECT9) | At1g27960 | | | LOC_Os08g12760 | YTHDF2 |
| Readers | YTH03 (CPSF30) | At1g30460 | | | LOC_Os06g46400 | YTHDF3 |
| Readers | YTH04 (ECT7) | At1g48110 | | | LOC_Os03g20180 | YTHDC1 |
| Readers | YTH05 (ECT4) | At1g55500 | | Development | LOC_Os03g53670 | YTHDC2 |
| Readers | YTH06 (ECT8) | At1g79270 | | | LOC_Os01g48790 | |
| Readers | YTH07 (ECT1) | At3g03950 | | | LOC_Os04g51940 | |
| Readers | YTH08 (ECT5) | At3g13060 | | | LOC_Os08g44200 | |
| Readers | YTH09 (ECT2) | At3g13460 | | Trichome branching | LOC_Os07g07490 | |
| Readers | YTH10 (ECT6) | At3g17330 | | | LOC_Os04g51950 | |
| Readers | YTH11 | At4g11970 | | | | |
| Readers | YTH12 (ECT10) | At5g58190 | | Trichome branching | LOC_Os05g04000 | |
| Readers | YTH13 (ECT3) | At5g61020 | | Trichome branching | LOC_Os05g01520 | |
| Erasers | ALKBH1A | At1g11780 | | | LOC_Os03g60190 | ALKBH1 |
| Erasers | ALKBH1B | At3g14140 | | | LOC_Os11g29690 | |
| Erasers | ALKBH1C | At3g14160 | | | | |
| Erasers | ALKBH1D | At5g01780 | | | | |
| Erasers | ALKBH2 | At2g22260 | | | LOC_Os06g17830 | ALKBH2 |
| Erasers | ALKBH6 | At4g20350 | | | LOC_Os10g28410 | ALKBH6 |
| Erasers | ALKBH8 | At1g36310 | tRNA mcm^5U | | LOC_Os04g51360 | ALKBH8 |
| Erasers | ALKBH8A | At1g31600 | tRNA mcm^5U | | LOC_Os11g43610 | |
| Erasers | ALKBH8B | At4g20485 | | | | |
| Erasers | ALKBH9A | At1g48980 | | | LOC_Os06g04660 | ALKBH5 |
| Erasers | ALKBH9B | At2g17970 | m^6A | Viral infection | LOC_Os05g33310 | |
| Erasers | ALKBH9C | At4g36090 | | | | |
| Erasers | ALKBH10A | At2g48080 | | | LOC_Os10g02760 | |
Wilm’s tumor 1-associating protein functions as a stabilizer for the heterodimer localized in nuclear speckle (Ping et al., 2014; Lence et al., 2016). VIRMA plays a role in guiding the methyltransferase complex to the selective target region of mRNAs (Niessen et al., 2001; Yue et al., 2018). Arabidopsis VIR and FIP37 were identified as an ortholog of VIRMA and WTAP, respectively (Zhong et al., 2008; Bodi et al., 2012; Shen et al., 2016; Růžička et al., 2017).

Recently, zinc finger CCCH domain-containing protein 13 (ZC3H13), the latest component of methyltransferase complex, was found to be essential for localization of methyltransferase complex in mammals and Drosophila (Guo et al., 2018; Knuckle et al., 2018). However, the existence and molecular function of ZC3H13 in plants remain unknown. Interestingly, Arabidopsis contains E3 ubiquitin ligase HAKAI as an additional m^6^A writer component (Růžička et al., 2017; Table 1). Although knockdown of its expression can decrease m^6^A level (Růžička et al., 2017), the primary role of HAKAI in methyltransferase complexes has yet to be investigated.

**Erasers**

Removal of methylation marks on RNAs is carried out by α-ketoglutarate-dependent dioxygenase (AlkB) homolog (ALKBH) proteins that can erase alkyl and methyl groups from DNAs, RNAs, and proteins (Fedele et al., 2015; Alemu et al., 2016). Mammals have nine ALKBH family members: ALKBH1 to ALKBH8 and fat mass- and obesity-associated protein (FTO) (Ougland et al., 2015; Table 1). Although ALKBH2 and ALKBH3 have been identified as main DNA repair enzymes, ALKBH3 also shows activity on m^1^A and m^2^C of RNAs (Ueda et al., 2017). Interestingly, ALKBH1 acts on a wide range of substrates in DNAs, RNAs, and histones (Westbye et al., 2008; Ougland et al., 2012; Wu et al., 2016). In addition to its role in cytoplasm, human ALKBH1 targets several m^1^A methylated tRNAs in mitochondria, influencing the organellar translation and function (Kawarada et al., 2017; Müller et al., 2018). ALKBH8, another tRNA DMT, interestingly contains both methyltransferase and demethylase domains, unlike other family members (Pastore et al., 2012).

Only two m^6^A erasers, ALKBH5 and FTO, have been identified in animals so far. Both enzymes were originally shown to be involved in demethylation of m^6^A (Jia et al., 2011; Zheng et al., 2013). However, recent studies have suggested that FTO has a much higher activity toward N^6^, 2′-O-dimethyladenosine (m^6^A_m) compared to that for m^6^A (Meyer and Jaffrey, 2017; Mao et al., 2017; Mao and Jaffrey, 2018). ALKBH5 and FTO have been found to be involved in alternative splicing, 3′-UTR processing, mRNA stability, translation, and amino-acids deprivation response pathway (Zheng et al., 2013; Zhao et al., 2014; Bartosovic et al., 2017; Tang et al., 2018). Arabidopsis contains several putative m^6^A eraser ALKBH family proteins (Table 1), among which only two eraser proteins ALKBH9B and ALKBH10B have been functionally characterized in viral infection and floral transition (Duan et al., 2017; Martínez-Pérez et al., 2017). In summary, although increasing number of erasers targeting specific methylation marks have been identified, the activity and substrate RNAs of most ALKBH family members in plants and animals are yet to be determined.

**Readers**

Interpretation of methylation marks is tightly related to posttranscriptional regulation of mRNA metabolism which requires reader proteins to recognize methylated transcripts and ultimately determine their fates. In recent years, several RNA-binding proteins (RBPs) that can recognize m^6^A marks on mRNAs have been identified in animals by RNA-protein immunoprecipitation using synthetic m^6^A-containing RNAs (Dominissini et al., 2012; Xu et al., 2014; Arguello et al., 2017; Edupuganti et al., 2017; Wu et al., 2017). YTH521-B homology (YTH) domain family (YTHDF) protein was first identified as an m^6^A-binding protein (Xu et al., 2014). Recently, human and mouse YTHDF proteins including YTHDF1, YTHDF2, YTHDF3, YTHDC1, and YTHDC2 were found to possess a specific binding pocket for m^6^A nucleotides and exhibit significantly high affinity to methylated RNAs, suggesting their role as m^6^A readers (Dominissini et al., 2012; Hsu et al., 2017; Xiang et al., 2017; Liao et al., 2018). YHHDH2 can bind to m^6^A-modified RNAs and play a distinct role in mRNA degradation by recruiting the CCR4-NOT deadenylase complex (Wang et al., 2014; Zhou et al., 2015; Du et al., 2016). YTHDF1 was found to recognize the 5′UTR of m^6^A-modified mRNAs in the cytosol, which promotes translation of target transcripts in a cap-independent manner (Wang et al., 2015b; Shi et al., 2017). YTHDC1 is involved in exon-selective gene splicing in the nucleus (Xiang et al., 2017). Interestingly, YTHDC2 also contains RNA helicase domain (Jain et al., 2018). Arabidopsis and rice genomes encode 13 and 12 YTHD proteins, respectively (Li et al., 2014a; Table 1). Contrary to extensive study on YTHD proteins in animals, only three evolutionarily conserved c-terminal region (ECT) family proteins have recently been functionally characterized in Arabidopsis as YTHD homologs (Arribas-Hernández et al., 2018; Scutenaire et al., 2018; Wei et al., 2018; Table 1).

Besides YTHD proteins, two other proteins containing different RNA-binding domains that can recognize m^6^A marks in animal cells have been reported. One is a heterogeneous nuclear ribonucleoprotein A2/B1 (HNRNP2B1) which regulate RNA splicing in the nucleus through a well-characterized RNA-recognition motif (Alarcon et al., 2015). Notably, instead of directly binding to m^6^A site as YTHD proteins, HNRNP2B1 might bind to altered structures right after the m^6^A site (Alarcon et al., 2015). Insulin-like growth factor 2 mRNA-binding protein (IGF2BP) contains tandem K-homology (KH) domains to recognize m^6^A sites and enhance target mRNA stability, storage, and translation in an m^6^A-dependent manner (Nicastro et al., 2015; Huang et al., 2018). Eukaryotic initiation factor 3 (eIF3) can also promote translation of m^6^A-modified mRNAs depending on m^6^A modification (Meyer et al., 2015). Clearly, more reader proteins recognizing other RNA modifications as well as m^6^A marks should be uncovered to fully understand
cellular roles of epitranscriptomic RNA modifications in both plants and animals.

**RNA METHYLATION IN ANIMAL DEVELOPMENT AND DISEASES**

m\(^6\)A methylation has been demonstrated to affect all fates of mRNA metabolism, including pre-mRNA processing and intron splicing in the nucleus, nucleus-to-cytoplasm RNA export, translation, and RNA decay in the cytoplasm (Figure 2). Analysis of different mettl mutants demonstrated the essential role of m\(^6\)A methylation in cell development, proliferation, differentiation, and motility by regulating mRNA stability and splicing pattern of diverse transcripts (Wang et al., 2014; Chen et al., 2015; Geula et al., 2015; Park and Hong, 2017; Widagdo and Anggono, 2018).

Loss of FTO can inhibit differentiation of primary myoblasts and skeletal muscle in mice, suggesting that m\(^6\)A demethylase FTO plays a crucial role in somatic and neural stem cell differentiation (Wang et al., 2017). A larger number of gene encoding clock genes and clock output genes are enriched in m\(^6\)A methylation (Fustin et al., 2013; Hastings, 2013) and changes in m\(^6\)A levels can affect circadian rhythms, cellular growth, and survival (Fustin et al., 2018).

Notably, recent studies have demonstrated that alteration in m\(^6\)A levels is closely associated with various diseases, especially cancer (reviewed in Dai et al., 2018; Pan et al., 2018). For example, FTO affects m\(^6\)A level and translation of Angptl4 mRNA, which regulates fatty acid mobilization in adipocytes and body weight (Wang et al., 2015a). Low m\(^6\)A level in total RNA is related to type 2 diabetes mellitus (Shen et al., 2014). Considering that aberrant cell growth and differentiation cause cancer, it is worth noting that cancer cells may improve their survival rate and progression by modulating aberrant methylation of target RNAs. Several studies have shown that expression of FTO or ALKBH5 can decrease m\(^6\)A level, resulting in enhanced cancer cell growth (Zhang et al., 2016; Li et al., 2017; Zhang et al., 2017). METTL3 acts as an oncogene in cancer cells, enhancing the translation of cancer-inducing genes by interacting with translation initiation factor (Lin et al., 2016).

In addition to m\(^6\)A methylation, m\(^5\)C is also involved in cell development and diseases. This modification is deeply associated

![FIGURE 2](image-url) Diverse cellular processes affected by m\(^6\)A RNA methylation. Splicing of mRNAs in the nucleus and diverse RNA metabolism in cytoplasm, including cap-dependent and cap-independent translation, RNA decay in cytosol and P-body, and RNA storage, is affected by m\(^6\)A RNA methylation. Specific “reader” proteins recognizing m\(^6\)A marks on mRNAs play essential roles in these cellular processes. Writers (MTA, MTB, FIP37, VIR, and HAKAI), erasers (ALKBH9B/10B), and reader (YTH09) identified in Arabidopsis are shown.
with testis differentiation and tumor cell proliferation. A previous study has shown that NOP2/sun RNA methyltransferase family member 2 (NSUN2), an m^6^A writer, is highly expressed in tumor cells and its depletion decreases levels of Ddx4, Miwi, and Tudor domain-containing proteins, suggesting an essential role of m^5^C RNA methylation in male germ cell differentiation (Frye and Watt, 2006). Moreover, loss of NSUN2 causes an accumulation of progenitors, decreases in upper-layer neurons, and increases in tRNA fragment accumulation in the brain, resulting in damage to neural stem cell differentiation and motility (Flores et al., 2017). Although these studies clearly demonstrate the importance of m^6^A and m^5^C in cell proliferation and diseases, biological functions of other RNA methylations in animal development and pathogenesis are yet to be elucidated.

**RNA METHYLATION IN PLANT DEVELOPMENT AND ABIOTIC STRESS RESPONSES**

Although our understanding of writers, readers, and erasers in plants is far behind their animal counterparts, accumulating reports clearly demonstrate that these factors are essential for plant growth and abiotic stress responses. Herein, we will summarize and discuss characterized and potential writers, readers, and erasers (Table 1) in plants.

### m^6^A Writers

Genome-wide m^6^A methylation patterns have been mapped in barley, *Arabidopsis*, and rice (Li et al., 2014b; Luo et al., 2014). However, key enzymes responsible for this methylation have only been studied in *Arabidopsis*. Analysis of mta (*Arabidopsis* ortholog of human METTL3) knockdown mutants has revealed that MTA is required for m^6^A mRNA methylation and essential for normal growth and development, such as shoot and root growth as well as leaf and floral development (Zhong et al., 2008; Bodi et al., 2012). Moreover, MTA was found to interact with MTB, an *Arabidopsis* ortholog of human METTL14. Knockdown of MTB showed a similar but less severe phenotype compared to *mta* mutants, indicating that both writers are essential for plant development (Růžička et al., 2017). The *Arabidopsis* m^6^A writer complex also includes an ortholog of human WTAP named FIP37. Depletion of FIP37 results in embryo lethality while its partial loss causes a graft overproliferation of shoot meristems by increasing the stability of shoot meristemless (*STM*) and WUSCHEL (*WUS*) (Shen et al., 2016). Vir and Hakai are other m^6^A writer components in *Arabidopsis*. They affect root and shoot growth as well as cotyledon development, similar to other m^6^A writer mutant phenotypes (Růžička et al., 2017). However, the molecular mechanism underlying Vir and Hakai functions is yet to be elucidated.

Despite increasing understanding of the roles of m^6^A writers in plant growth and development, reports demonstrating their involvement and functions in plant response to abiotic stresses are lacking. Our analysis of publically available microarray data using GENEVESTIGATOR revealed that expressions levels of writers in *Arabidopsis* and rice are differently modulated by diverse abiotic stresses (Figure 3). In *Arabidopsis*, levels of most m^6^A writer components were not significantly modulated by abiotic stresses. Levels of MTA and FIP37 were only marginally increased by cold and heat stress, respectively. In rice, the level of OsFIP was increased by cold stress whereas levels of OsMTA, OsMTB, and OsVIR were decrease by cold, drought, or salt stress. The constant expression of m^6^A writer components under normal and stress conditions suggests the fundamental role of m^6^A methylation in plant development and stress responses.

### m^5^C Writers

Although m^5^C methylation in DNA has been studied for many years, its cellular and molecular functions in RNAs is just beginning to be uncovered. Due to advancement in RNA sequencing, m^5^C RNA methylation could be mapped to mRNAs in both animals and plants (Schafer et al., 2009; Hussain et al., 2013; Song et al., 2018). Overall, m^5^C RNA methylation is a less abundant modification of RNA than m^6^A methylation. In *Arabidopsis*, two enzymes, TRM4A and TRM4B, are responsible for m^5^C RNA methylation. Both enzymes are orthologs of human m^5^C methyltransferase NSuns2. However, TRM4A contributes to tRNA m^5^C methylation while TRM4B targets mRNA for m^5^C methylation. Loss of TRM4A does not exhibit any visible phenotype while loss of Trm4B reduces root length, suggesting the role of mRNA m^5^C methylation in root development regulation (David et al., 2017). In accordance to this, loss of m^5^C RNA methylation affects the stability of short hypocotyl 2 (SHY2) and indoleacetic acid-induced protein 16 (IAA16), two critical genes related to root development (Cui et al., 2017). Our analysis showed that expression levels of *Arabidopsis* TRM4B are marginally increased by cold stress, although they decrease under heat stress. In contrast, expression levels of rice TRM4A and TRM4B are not altered in response to abiotic stresses (Figure 3). Although these expression patterns suggest potential roles of m^5^C writers in abiotic stress response, the relevance of m^5^C methylation to abiotic stress responses awaits further investigation.

### m^6^A Erasers

Among protein factors involved in RNA methylation in plants, erasers are so far the least studied, although new knowledge is gained rapidly. Thirteen *Arabidopsis* ALKBH family members have been identified by bioinformatic analysis (Mielecki et al., 2012). However, only a few of them have been studied so far (Table 1). Among them, ALKBH9A, 9B, 9C, 10A, and 10B show the highest amino acid sequence similarity with human ALKBH5. Other family members are numbered based on their sequence similarity to human orthologs (Table 1). Like animal counterparts, most erasers are localized in the nucleus and cytoplasm whereas ALKBH1D is also present in chloroplasts. Interestingly, some of them show relocation to the nucleus in response to methylating agents (Mielecki et al., 2012). ALKBH10B was identified as the principal mRNA m^6^A eraser influencing floral transition by controlling transcript levels of SPL3, SPL9, and *FLOWERING LOCUS T* (Duan et al., 2017). Another demethylase, ALKBH9B, was shown to revert m^6^A from single-stranded RNA in vitro (Martinez-Perez et al., 2017).
Although alk9b knockout mutants do not show differences in plant RNA m6A methylation level (Duan et al., 2017), its depletion results in hypermethylation of alfalfa mosaic virus (AMV) RNA, mediating systemic infection by interacting with viral cap proteins (Martínez-Pérez et al., 2017).

Expression of ALKBH9A is highly induced in roots under salt stress but not in response to ABA (Ma et al., 2006). Its level is much lower than ALKBH9 and ALKBH10 under normal conditions (Duan et al., 2017). ALKBH10A is down-regulated by heat stress (Merret et al., 2015) whereas ALKBH10B is up-regulated in response to karrikins (Nelson et al., 2010). Although these previous studies suggest a specific role of ALKBHs in stress responses as well as plant development, nothing is known about their actual roles. Our analysis showed that expression levels of ALKBH members were marginally up- or down-regulated in Arabidopsis by different abiotic stresses (Figure 3). Notably, levels of ALKBH1 in rice were highly increased upon drought, cold, or ABA treatment whereas expression levels of ALKBH6, ALKBH8B, and ALKBH10A were decreased by drought, ABA, or cold (Figure 3). These data suggest that ALKBHs could play important roles in abiotic stress responses, although this awaits further investigation.

m6A Readers

Although several RBPs interpreting m6A marks have been identified in animals, roles of only three YTHD m6A reader proteins have very recently been determined in Arabidopsis (Arribas-Hernández et al., 2018; Scutenaire et al., 2018; Wei et al., 2018). YTHD09 (ECT2) is involved in trichome development. Moreover, cytoplasmic-localized YTHD09 relocates to stress granules upon heat exposure, suggesting its role in mRNA fate control under stress conditions (Scutenaire et al., 2018). By using single and double mutants, it has been demonstrated that YTHD09 (ECT2), YTHD13 (ECT3), and ECT4 regulate the timing and execution of plant organogenesis (Arribas-Hernández et al., 2018). Moreover, a molecular study revealed that ECT2 targets a large number of m6A-containing transcripts, including TTG1, ITB1, and DIS2, which are involved in trichome development (Wei et al., 2018). Further sequencing analysis suggested that ECT2 increases the stability of these
transcripts and influences trichome development (Wei et al., 2018). Although these studies clearly point to important roles of YTHD readers in plant development, more in-depth and focused efforts are needed to identify and characterize potential reader proteins (Table 1) that can recognize not only m^6^A modification, but also other methylation marks in plants.

No reports demonstrating the involvement or functions of any RNA methylation reader proteins in plant response to abiotic stresses have been published so far. However, a previous study and our current analysis showed that the expression of YTHDs in Arabidopsis and rice is highly regulated by different abiotic stresses (Li et al., 2014a; Figure 3). In Arabidopsis, levels of YTHD05, YTHD06, and YTHD07 are increased by heat, cold, hypoxia, or submergence stress. In contrast, the expression level of YTHD10 decreases under cold, drought, salt, or osmotic stress whereas YTHD08 level is reduced by heat stress. In rice, YTHDs responded differently to various abiotic stresses (Li et al., 2014a; Figure 3). Expression levels of YTHD05, YTHD06, YTHD07, and YTHD09 are downregulated by cold stress whereas levels of YTHD03 and YTHD08 increase under submergence and heat stress, respectively. Notably, none of these rice YTHDs showed altered expression under salt stress whereas YTHD01, YTHD02, YTHD03, YTHD04, or YTHD08 does not respond to cold stress. The fact that m^6^A reader proteins respond more to abiotic stresses than writers and erasers suggests that decoding of methylation marks is much more important than introducing or removing these marks during stress adaptation process in plants. It would be interesting to characterize roles of reader proteins in RNA metabolism and its consequence in stress responses.

**CONCLUDING REMARKS AND PERSPECTIVES**

Chemical modifications of RNAs are invaluable ways to expand decoding capacity of RNA transcripts beyond genetic information inherent to genome sequences. They are crucial for posttranscriptional gene regulatory events such as mRNA splicing, stability, and translation. The ability to regulate the fate of RNA molecules through nucleotide modifications is vital to plant survival and fitness under adverse as well as favorable environmental conditions. Despite the increasing discovery of cellular components essential for chemical modification and decoding of modified RNA molecules, our knowledge regarding physiological roles of proteins involved in these processes is far from sufficient. Several key questions remain to be further investigated. Are there any other internal RNA modifications not identified so far? How these components are regulated depending on developmental stages and/or in response to changing environmental cues? What guides the specificity of interactions between these components with target transcripts? Are these components conserved between dicots and monocots, especially in crop species? Addressing these questions will greatly expand our knowledge on the process of chemical modifications of RNAs and its effects on plant survival and fitness under stressful conditions. Such studies could provide potential new targets for engineering crop plants with higher adaptability to adverse environmental conditions.

**AUTHOR CONTRIBUTIONS**

HK designed the concept. JH and SM compiled and analyzed data. JH, SM, and HK contributed to the writing of this review.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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