1 **REVIEW ARTICLE**

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Beyond transcription: compelling open questions in plant RNA 3 biology 4

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2 Short title: Open questions in plant RNA biology

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5 Abstract

The study of RNAs has become one of the most influential research fields in 6 7 contemporary biology and biomedicine. In the last few years, new sequencing 8 technologies have produced an explosion of new and exciting discoveries in the field 9 but have also given rise to many open questions. Defining these questions, together with old, long-standing gaps in our knowledge, is the spirit of this article. The breadth of 10 topics within RNA biology research is vast, and every aspect of the biology of these 11 molecules contains countless exciting open questions. Here, we asked twelve groups to 12 discuss their most compelling question among some plant RNA biology topics. The 13 14 following vignettes cover RNA alternative splicing: RNA dynamics: RNA translation: RNA structures; R-loops; epitranscriptomics; long noncoding RNAs; small RNA 15 production and their functions in crops; small RNAs during gametogenesis and in cross-16 kingdom RNA interference; and RNA-directed DNA methylation. In each section, we will 17 18 present the current state-of-the-art in plant RNA biology research before asking the guestions that will surely motivate future discoveries in the field. We hope this article will 19 spark a debate about the future perspective on RNA biology and provoke novel 20 reflections in the reader. 21

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23 Introduction

24 (Written by Pablo A. Manavella)

25 In all living organisms, DNA is the molecule storing all genetic information, while RNA carries this data to the ribosomes to be translated into proteins. While DNA is 26 27 omnipresent in our imagination, making star appearances in movies, TV shows, and books, the contribution of RNA to life is less recognized by society. However, as a 28 consequence of the recent COVID pandemic, the word "RNA" has reached most people 29 on the planet as they learned about the RNA-based genome of the virus and the 30 therapeutic use of RNA vaccines. Thus, the concept of 'information flow', that is the 31 decoding of DNA to protein using an RNA intermediate, has suddenly become the 32 center of attention and conversations. What remains largely unknown to the general 33 34 audience is that the advent of sequencing technologies has made it clear that RNA is 35 not only a coding molecule but also has various other functions, mostly in the form of cellular non-coding RNA transcripts. 36

The study of RNAs has emerged as a particularly important research field in contemporary biology, especially in plant biology, where these molecules execute many actions during development and response to the environment. Advances in sequencing technologies have allowed the global analysis of RNA modifications, the resolution of RNA secondary structures, the mapping of epigenetic modifications, the identification of 1 RNA-edited sequences, and the discovery of novel classes of RNAs resulting in a 2 revolution in molecular biology that is just starting.

In this article, we gathered twelve experts in different aspects of plant RNA biology to discuss some of the most compelling open questions in the field. Each section discusses long-standing open questions of the field as well as questions that have only begun to emerge from breakthrough discoveries. We hope this article helps stimulate the community and sparks new ideas and research projects that will expand the frontiers of RNA biology knowledge in plants.

10 How does light control RNA alternative splicing in plants?

11 (Written by Micaela Godoy Herz and Alberto Kornblihtt)

Plants rely on light as their main source of energy, but light also regulates many developmental and physiological responses during the plant life cycle (Arsovski et al., 2012). Furthermore, light signals induce a massive reprogramming of gene expression in plants (Tognacca et al., 2020). Alternative splicing produces multiple mRNA variants from a single locus. Splicing and alternative splicing are coupled with transcription, and factors that regulate transcription also affect alternative splicing (Kornblihtt et al., 2013).

18 Our laboratory showed how light regulates plant alternative splicing through the chloroplast (Petrillo et al., 2014). Light and dark conditions affect alternative splicing of a 19 subset of Arabidopsis (Arabidopsis thaliana) genes preferentially encoding proteins 20 involved in RNA processing. This effect requires functional chloroplasts: treatment of 21 Arabidopsis seedlings with drugs that impair the chloroplast photosynthetic transport 22 23 chain inhibit the effect of light on alternative splicing. Moreover, the effect of light is also observed in roots when communication with leaves -the photosynthetic tissue- is not 24 interrupted (Petrillo et al., 2014). Light, sensed by the chloroplast, indeed triggers a 25 retrograde signal that regulates alternative splicing not only in leaves, but also in roots. 26

How does light cause splicing responses in roots? In a recent work, Riegler and collaborators investigated this shoot-to-root signaling: they showed that alternative splicing responses in roots are not directly caused by light, but are instead most likely triggered by sugars. The kinase TARGET OF RAPAMYCIN (TOR) plays a key role in this signaling pathway. Sugars activate the TOR pathway and act as mobile signals to coordinate alternative splicing responses throughout the plant (Riegler et al., 2021).

These results afforded us a better understanding of how mobile signals regulate alternative splicing throughout the entire plant in response to light. One remaining outstanding open question is what happens in the nucleus: that is, what are the mechanisms involved in this regulation of alternative splicing in plants?

We performed different experiments to address the role of transcription elongation and determined that the light control of alternative splicing responds to a kinetic coupling mechanism (Godoy Herz et al., 2019). Briefly, the kinetic coupling model explains how changes in RNA Polymerase II (Pol II) elongation rate influence alternative splicing.

Each splice site consists of a consensus sequence that is recognized by spliceosomal 1 components, although 'strong' splice sites (those that are close to the consensus 2 sequence) are more efficiently recognized than 'weak' splice sites, which are 3 suboptimal. In the example illustrated in Figure 1, there is an alternative splicing event 4 5 with two 3' splice sites: a weak upstream 3' splice site, and a strong downstream splice site. If Pol II elongation rate is fast, both sites are presented to the splicing machinery at 6 the same time, and the strong 3' splice site is recognized by the splicing machinery 7 more efficiently, resulting in exon skipping. However, if Pol II transcription rate is slow, 8 the splicing machinery will recognize the upstream, weaker, 3' splice site first, and 9 afterwards the strong 3' splice site, which leads to exon inclusion (Godoy Herz et al., 10 2019). We showed by different experimental approaches that light promoted 11 transcription elongation in Arabidopsis, while Pol II elongation was slower in darkness. 12 Furthermore, the light control of alternative splicing and elongation was abolished in 13 14 plants lacking function for TRANSCRIPTION FACTOR II S (TEIIS) in a previous report (Dolata et al., 2015): These TFIIS mutant plants did not respond to light signaling on a 15 group of alternative splicing events. This result demonstrated that coupling between 16 transcription and splicing is important for a whole organism to respond to environmental 17 18 cues (Figure 1).

Plant lines with higher Pol II transcription activity were recently generated by introducing point mutations in NRPB2, the second largest subunit of Pol II. As a result, an accelerated Pol II elongation rate increased the polymerase signal in gene bodies, which appeared to modulate alternative splicing choices (Leng et al., 2020)..

Even though our knowledge of alternative splicing in plants has grown significantly in the last decade, many important open questions remain. It has been shown that, in response to light, sugars activate the TOR pathway, which in turn regulates alternative splicing. But how does TOR regulate alternative splicing in the nucleus? In the chloroplast, the exact nature of the chloroplast retrograde signal that regulates alternative splicing remain unknown, although it may be triggered by the oxidation state of the plastoquinone pool connecting both photosystems (Petrillo et al., 2014).

Moving forward, inside the nucleus, how light promotes Pol II elongation is unknown: 30 31 what makes Pol II elongation faster in the light, and slower in darkness? There are 32 many possible mechanisms that might explain how the chloroplast regulates transcription elongation. Furthermore, the role of chromatin modifications on the 33 regulation of alternative splicing in plants remains an interesting field to investigate. 34 35 Previous studies in mammalian systems have shown that histone post-translational modifications play a key role in the regulation of alternative splicing decisions. Treating 36 cell cultures with drugs that open chromatin structure promoted changes in alternative 37 splicing by facilitating Pol II elongation and exon skipping (Schor et al., 2009). By 38 contrast, cell differentiation results in an increase in intragenic silencing chromatin 39 40 marks that raised the rate of higher exon inclusion (Schor et al., 2013). In our work,

histone acetylation mimics the effect of light on alternative splicing, but light does not
affect the levels of this histone modification (Godoy Herz et al., 2019). Future
experiments will be needed to address the role of chromatin structure in splicing
regulation in plants.

5 Moreover, coupling between transcription elongation and alternative splicing may also 6 act in response to other environmental stimuli, like temperature. A recent work shows 7 that the TFIIS elongation factor is required for thermal adaptation in Arabidopsis 8 (Szadeczky-Kardoss et al., 2022). Furthermore, analyses of plant native elongation 9 transcript sequencing (plaNET-seq) experiments in response to cold showed changes in 10 Pol II promoter-proximal stalling and at the 3' end of genes (Kindgren et al., 2020).

Finally, it would be interesting to study if these mechanisms of gene expression regulation are also conserved in other plants and other photosynthetic organisms like

algae. Future work from different groups will be needed to address these guestions.

14

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23 The invisible world of RNA dynamics

24 (Written by Reed Sorenson and Leslie E. Sieburth)

Transcriptomics has transformed our understanding of molecular responses to signals. 25 The abundance of many mRNAs can be robustly upregulated or downregulated, and 26 27 many regulated genes bring about changes in development or physiology. Indeed, measurements of RNA abundance are so ingrained in our thinking that changes in RNA 28 29 levels are frequently referred to as gene expression or transcriptional responses. However, alongside regulatory events that lead to changes in mRNA levels, there lurks 30 31 the largely unseen layer of mRNA decay rate regulation. In addition to RNAs with 32 modified rates of decay and changes in their abundance, rates of decay can also be modified for mRNAs whose abundances are held steady. This largely invisible dynamic 33 regulation is just beginning to be investigated, and so there are numerous unanswered 34 35 questions, including why decay rates are modified independently of changes in abundance, how this modulation occurs, and whether this regulation has implications for 36 mRNAs that do show changes in their abundance. 37

RNA abundances are influenced by both synthesis (transcription) and decay, and the rate of RNA turnover is called flux (Figure 2A). Wide variations in flux have been observed in all deep RNA decay analyses, but whether flux rates affect RNA abundances and/or regulation is still an open question. A special case of mRNA flux
regulation occurs when both the transcription and decay rates of an mRNA are
modulated, and yet the mRNA abundance does not change. This phenomenon is called
'RNA buffering' because transcription and decay rates are balanced to maintain steady
abundances (Figure 2B). RNA buffering has been documented in Arabidopsis, but we
are at the very beginning of understanding all aspects of this process, including both
how and why some mRNAs become buffered.

8 The system where RNA buffering is best understood is budding yeast (Saccharomyces) cerevisiae). A mysterious observation led to its discovery: mutants with defects in either 9 RNA decay or transcription were found to maintain normal mRNA abundances. It turned 10 out that the initial defect, in e.g. RNA decay, was accompanied by a compensatory 11 change (e.g. in transcription). That is, normal abundances of mRNAs in many 12 transcription and RNA decay mutants were maintained by precisely balanced changes 13 through RNA buffering (Haimovich et al., 2013; Sun et al., 2013a; Timmers and Tora, 14 2018; Hartenian and Glaunsinger, 2019). Because most mRNA decay occurs in the 15 cytoplasm, while transcription takes place in the nucleus, RNA buffering requires not 16 just precise regulation, but also communication between the nucleus and the cytoplasm. 17 18 Mechanisms underlying this regulation are still emerging and somewhat controversial, but studies in yeast have revealed RNA decay proteins relocating to the nucleus and 19 displaying novel functions. For example, Sun and colleagues showed that the yeast 20 $5' \rightarrow 3'$ EXORIBONUCLEASE 1 (XRN1) moves from the cytoplasm to the nucleus, where 21 it binds DNA and influences transcription of buffered RNAs (Sun et al., 2013a). 22 23 Similarly, upon nuclear RNA exosome dysfunction, RNA buffering was activated by global attenuation of transcription via stabilization of the mRNA encoding HISTONE 24 SIRTUIN DEACETYLASE (HST3) (Bryll and Peterson, 2022). RNA buffering has also 25 been observed in Drosophila (Drosophila melanogaster), where it was used for gene 26 27 dosage compensation (Faucillion et al., 2022).

It was a similarly mysterious observation that led us to discover RNA buffering in 28 Arabidopsis (Sorenson et al., 2018). Cytoplasmic mRNA decay initiates through 29 deadenylation, and decay in the $3' \rightarrow 5'$ direction can be catalyzed by either the RNA 30 31 exosome or SUPPRESSOR OF VARICOSE (SOV)/ DIS3-LIKE EXONUCLEASE 2 (DIS3L2), while decay in the $5' \rightarrow 3'$ direction is initiated by decapping followed by 32 exoribonucleolytic digestion by XRN4 (Labno et al., 2016). Because the popular wild-33 type accession Columbia-0 (Col-0) harbors a sov loss-of-function mutation (Zhang et 34 35 al., 2010) possible functions of this decay pathway were mysterious. To understand why sov mutants did not show an abnormal phenotype, and also identify mRNA substrates 36 of decapping and SOV, we compared genome-wide RNA decay rates for wild type and 37 RNA decay mutants. In varicose (vcs) mutants (which lack mRNA decapping) the 38 expectation that mRNA decapping substrates would decay more slowly was largely 39 40 observed. The most common decay pattern (seen in >7,000 RNAs) was half-lives that

were longer in vcs, and longer still in vcs sov double mutants, indicating that these 1 RNAs were typically degraded by VCS, but upon loss of VCS, SOV provided back-up. 2 However, many mRNAs in sov mutants showed a surprising shift to shorter half-lives. 3 Moreover, mRNA decapping (via VCS) was required to sustain these shorter half-lives. 4 5 This unusual decay rate shift had no significant effect on RNA abundances, indicative of RNA buffering and explaining the lack of phenotypic consequences in sov mutants in 6 Col-0. Data suggestive of RNA buffering was also identified in an Arabidopsis study of 7 cold response (Arae et al., 2017). We do not know whether plants use a mechanism 8 similar to that of yeast for RNA buffering; however there are no reports of XRN4 being 9 found in the nucleus (suggesting that RNA buffering in Arabidopsis might differ 10 mechanistically from yeast), and the shifting of SOV substrates to decapping via VCS 11 12 has not been described previously.

Conventional views of gene expression place all the action on those mRNAs whose 13 14 abundances are altered. However, RNA buffering turns this conventional view on its head by demonstrating that many mRNAs with stable unchanging abundances also 15 undergo complex regulation. And this observation leads to an even bigger open 16 question: how are some mRNAs buffered so that their abundances do not vary, while 17 18 others appear to be able to freely increase or decrease in abundance? If only specific mRNAs are buffered, perhaps they share a common sequence motif, e.g. for a 19 regulatory RNA-binding protein. Alternatively, buffering of RNA abundances may be a 20 default state, and some sort of licensing might be required to allow mRNAs to undergo 21 changes in abundance (Figure 2B). What distinguishes RNAs to be buffered from those 22 23 licensed to undergo alterations in their abundance? Attractive candidates can be found in the expanding world of RNA modifications, from differing caps to covalent 24 modifications such as N⁶-methyladenosine (m⁶A) or structure (Kwok et al., 2013; Mauer 25 et al., 2017; Anderson et al., 2018; Reichel et al., 2019; Wang et al., 2019; Zhang et al., 26 27 2019a). Addressing these many open questions will require much deeper understanding of RNA kinetics. 28

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Stabilization of mRNA and translational regulation by stress granules in response to environmental conditions

36 (Written by Kentaro Nakaminami and Motoaki Seki)

Current technologies used to analyze gene expression have enabled a high-level of resolution on the expression of thousands of genes. Advancements in proteomic

- technologies have also greatly improved the comprehensive analysis of proteins. Thus,
- it has become possible to analyze plant physiology and metabolism in great detail using

various analytical methods. Collectively, these studies have empirically indicated that gene and protein abundance patterns are not always identical based on the results of multiomics analyses. Major factors contributing to the observed differences between mRNA and protein patterns are post-transcriptional regulation of mRNA and translational regulation of proteins. Both mechanisms fine-tune which mRNAs are translated into proteins to regulate the physiology and metabolism of living organisms.

Various events occur between mRNA transcription and translation via the activity of 7 RNA-binding proteins (RBPs) that determine whether proteins are synthesized (Burjoski 8 and Reddy, 2021). These events begin with quality control of transcribed mRNAs, 9 followed by degradation of unnecessary or aberrant mRNAs, or protein translation. 10 Additionally, mRNAs can be temporarily stored via a stabilization system for subsequent 11 activation in response to environmental changes and other stimuli, Although mRNA 12 abundance is affected by the balance between transcription and degradation, the 13 14 amount of protein is not always proportional to mRNA abundance, and is affected by post-transcriptional regulatory mechanisms such as the speed of translation and 15 translational inhibition. mRNA degradation is regulated by mRNA-protein (mRNP) 16 complexes called processing bodies (PBs); translation is carried out by ribosome 17 complexes (poly-ribosomes or polysomes), while mRNA stabilization or storage occurs 18 in stress granules (SGs) (Chantarachot and Bailey-Serres, 2018; Maruri-Lopez et al., 19 2021). These granules are not organelles but rather membraneless RNA granules 20 formed via liquid-liquid phase separation (LLPS) (Emenecker et al., 2020). They have 21 been reported to be present in both animals and plants and are becoming a growing 22 23 research focus. This section of the present review discusses the nature of SGs, which are responsible for the mechanisms of translational regulation, and how mRNAs are 24 stabilized and stored in these bodies. 25

Plants suppress mRNA translation when they are subjected to severe stress 26 (Merchante et al., 2017). This strategy reduces energy expenditure under stress 27 conditions, as only essential proteins are synthesized. Since translation requires 28 29 considerable energy, reducing energy requirements during stress contributes to increased survival rates. Importantly, active but selective translation must operate 30 31 during stress response in plants since essential proteins are still translated. The temporary storage of mRNA in SGs during stress conditions can be rapidly reversed, 32 with mRNAs being released in a translationally active form (polysomes) as plants 33 recover from stress conditions (Kosmacz et al., 2019). The mechanisms responsible for 34 35 determining target selectivity and translation timing by mRNP complexes, however, have not been clearly elucidated. 36

SG complexes that form in the cytoplasm during stress are conserved in eukaryotes
(Maruri-Lopez et al., 2021). SG formation in plants is triggered by a variety of stresses,
including high temperature, hypoxia, high salinity, and darkness (Chantarachot and
Bailey-Serres, 2018; Hamada et al., 2018). An SG is composed of translationally

arrested mRNAs and proteins related to the initiation of translation, such as translation 1 initiation factors, small subunits of ribosomal RNA (rRNA), poly(A)-binding proteins, as 2 well as regulatory RBPs that inhibit translation. Recently, hundreds of proteins have 3 been characterized as SG components by combining immunoprecipitation (IP) and 4 5 genome-wide mRNA-binding interactome capture methods with proteomic analyses (Chantarachot and Bailey-Serres, 2018; Kosmacz et al., 2019; Marondedze et al., 2019; 6 Gutierrez-Beltran et al., 2021; Maruri-Lopez et al., 2021). These results have suggested 7 that SGs are formed not only upon heat and hypoxia stresses, but also by drought 8 stress, resulting in translational repression. The components discovered in these 9 studies were not revealed based on their homology to SG components in animals and 10 veasts as in previous studies, but rather were directly identified by the indicated 11 methodologies as components of SGs. Although many SG components have been 12 isolated with this approach, proteins within SGs also include translation-promoting 13 14 proteins such as translation initiation factors, and not all are related to translation inhibition. It is necessary to consider the components and functions of SGs, including 15 spatiotemporal factors such as the dynamics of SG formation/dissociation, timing and 16 localization. SGs suppress translation and protect transcribed mRNAs from degradation 17 by temporarily storing selected mRNAs. SGs can be disassembled during stress 18 recovery and the stored mRNAs then become accessible for immediate translation. This 19 rapid reactivation is believed to be a response to environmental changes. Previous 20 studies have identified SG-regulated target mRNAs by analyzing RBPs present in SGs. 21 The identification of untranslated target mRNAs stored in SG has provided information 22 23 on the translational control or selective translation mechanism that occurs in response to stress. In previous studies using hypoxic and heat stress samples, various direct 24 target mRNA identifications have been performed with multiomics analyses such as 25 RNA immunoprecipitation followed by sequencing (RIP-seq) analysis, transcriptome 26 27 analysis, translatome and mRNA degradation rate analysis (Sorenson and Bailey-Serres, 2014; Nguyen et al., 2016; Tian et al., 2022; Zhu et al., 2022). Although many 28 mRNA targets that are thought to be regulated by SGs have been revealed, their 29 subsequent fates, such as when translation-inhibited mRNAs are finally translated into 30 31 proteins, still remains unclear at this time. 32 While multiomics analyses, such as a RIP analysis combined with translatome and

polysomal analyses, have enabled the identification of SG components and direct target 33 mRNAs that are SG-regulated, there are still many open questions that are await 34 35 clarification. For example, OLIGOURIDYLATE BINDING PROTEIN 1b (UBP1b), an SG component, localizes to the nucleus under non-stress conditions. UBP1b-stress 36 granules (UBP1b-SG) are induced to form in the cytoplasm in response to heat stress, 37 and candidate target mRNAs for UBP1b have been identified. UBP1b is present in both 38 the nucleus and cytoplasm, but its precise location where it exerts its mRNA 39 40 stabilization role remains unclear (Figure 3). In addition, PBs and SGs co-localize,

suggesting that their constituent components might interact (Hamada et al., 2018).
Although many mRNAs have been described as SG targets, not all will be
translationally inhibited. It is plausible that some targets might be degraded by PBs
rather than stabilized by SGs; the underlying mechanism of recruitment of target
mRNAs remains to be elucidated. Future studies should elucidate how targeted mRNAs
are exactly regulated by SGs. Clarifying the mechanism(s) of selective translation will
be a major step forward in understanding stress responses in plants.

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17 18

19 The pervasive function of RNA structure in plant growth and development

20 (Written by Yiliang Ding)

Plant growth and development is a continuous process starting with embryogenesis, 21 and the formation of the embryonic root and shoot, followed by organogenesis of 22 23 diverse organs such as roots, leaves, branches, and flowers. Plants rely on gene expression regulation to achieve specific cell differentiation and elongation to form 24 different organs. This extremely high coordination of gene expression at both temporal 25 and spatial levels requires diverse regulatory mechanisms to achieve evolutionary 26 27 fitness. Furthermore, plants have evolved to adapt to wide-ranging environmental conditions, acquiring highly dynamic regulation of gene expression in response to 28 29 different environmental factors. In addition to gene sequence content, RNA structure is 30 another important property of genes that can dynamically regulate gene expression at 31 the post-transcriptional level (Zhang and Ding, 2021).

32 Recent advances in RNA structure studies have enabled unprecedented opportunities to determine the functional importance of RNA structure across varied aspects of plant 33 growth and development. For instance, the antisense long noncoding RNA (IncRNA), 34 35 COOLAIR, folds into a complex RNA structure (Hawkes et al., 2016) that was suggested to suppress transcription of the key flowering gene, FLOWERING LOCUS C 36 (FLC) and promote flowering following vernalization. Another key regulator of plant 37 vascular development, JULGI (JUL), was shown to limit phloem differentiation through 38 its direct interaction with an RNA tertiary structure motif, RNA G-quadruplex, on the 5' 39 40 untranslated regions (5' UTRs) of SUPPRESSOR OF MAX2 1-LIKE4/5 (SMXL4/5)

mRNA to suppress their translation (Cho et al., 2018). Other studies have shown that 1 RNA G-quadruplex affects plant root growth and development (Foley et al., 2017; 2 Zhang et al., 2019b; Yang et al., 2020a). Extensive studies have indicated that RNA 3 structural conformations change in response to temperature (Su et al., 2018; Chung et 4 5 al., 2020), light (Gawronski et al., 2021), salinity stress (Kramer et al., 2020; Tack et al., 2020), and phosphate starvation (Reis et al., 2021). These changes subsequently alter 6 gene expression at the post-transcriptional level such as translation and RNA 7 degradation (Su et al., 2018; Chung et al., 2020; Kramer et al., 2020; Tack et al., 2020; 8 Gawronski et al., 2021; Reis et al., 2021). These recent studies have focused on either 9 identifying a specific RNA structural element on a specific transcript, or determining 10 global associations between RNA structure features and corresponding molecular 11 functions, and further support the growing evidence that highlights the importance of 12 RNA structure across diverse aspects of plant growth and development. 13

14 Since every mRNA is capable of folding into a particular RNA structure, this question has stimulated interest to explore the pervasive role of RNA structure in individual 15 genes to gain a more comprehensive understanding of RNA structure-mediated 16 regulation in plant growth and development. To achieve this in-depth understanding, the 17 strategies employed for studying RNA structure functionality need to reach a new level. 18 A promising approach may be the capability of achieving specific cell type resolution. In 19 plants, although stem cells are pluripotent, their cellular trajectories are limited in scope 20 because the identity of any given cell depends on its position relative to its neighbors. 21 For instance, root growth starts from sets of stem cell initials in the quiescent center 22 23 (QC), which generate continuous parallel files of epidermal cells that divide in a transverse, anticlinal orientation. Cells then divide in the meristematic zone before 24 starting to elongate into the differentiation zone of the mature root. After division, cells 25 remain in the same position and belong to the same lineage (Costa, 2016). 26 27 Interestingly, all these cell types share the same genetic information encoded in their DNA, but with diverse cellular conditions. The folding status of RNA structure is highly 28 dependent on cellular conditions such as ion concentrations and interacting proteins 29 (Zhang and Ding, 2021). Thus, it is likely that RNAs may fold differently to specify gene 30 31 function in different cell types, resulting in unique cell identities (Figure 4). Future research could focus on dissecting the extent of RNA structure diversities across 32 individual cell types. Indeed, the development of single-cell RNA structure profiling will 33 advance our understanding of RNA structural dynamics in plant cells. 34

Another future perspective could be to elucidate how RNA structures serve as environmental sensors. During growth, plants are constantly challenged by fluctuating environmental conditions such as biotic and abiotic stresses. Other abiotic stresses such as flooding and drought are likely to affect the folding status of RNA structures due to changing molecular concentrations in the cells (Zhang and Ding, 2021). During pathogen infection, many metabolites are significantly altered that may also influence

RNA folding (Zhang and Ding, 2021). Additional research could focus on dissecting the 1 detailed mechanisms of RNA structure-mediated stress responses including 2 comprehensive assessments of different stresses, or different degrees and duration of 3 stress. Finally, it may be possible to assess the evolution of RNA structures across the 4 5 plant kingdom. Previous studies have illustrated the evidence of evolutionary selection of certain RNA structure motifs (Yang et al., 2020a) and distinguished RNA structure 6 features in specific species (Deng et al., 2018; Yang et al., 2021). Studies of 7 evolutionary RNA structures may shed novel insight into understanding nucleotide 8 9 diversities in noncoding regions and at synonymous codon positions. Extension of RNA structure studies with a phylogenomic perspective may provide an evolutionary 10 perspective on RNA structure functionality. 11

With the rise of transcriptome-wide RNA structuromes, large volumes of RNA structure 12 data now provide the necessary scope for deep learning applications with the potential 13 14 for translating fundamental knowledge into RNA structure-based molecular design. For instance, from transcriptome-wide RNA structure and RNA stability data, we can now 15 learn and predict what kind of RNA structure features are responsible for RNA stability. 16 It may be possible to customize these RNA structure features into more or less stable 17 18 RNAs of interest. Where RNA structure acts as a post-transcriptional regulator, directly affecting protein production, RNA structure-guided molecular design may offer the 19 potential for new avenues in synthetic biology. 20 Recent technological advances have significantly pushed the discovery of RNA 21 structure functionalities forward. Further innovations in PacBio and Nanopore 22

structure functionalities forward. Further innovations in PacBio and Nanopore technologies to study RNA structures may offer more accurate RNA structure information at single-base resolution. These upcoming developments will invigorate RNA structure views to individual RNA structure conformations rather than the familiar bulk conformation. RNA structures may be a type of hidden "codon" embedded in every gene that facilitates the complexity of gene expression regulation. The rapid growth of RNA structure research may ultimately reveal the regulatory power of RNA structures in every aspect of plant growth and development.

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1 2

3 Sensing, regulation, and functions of R-loops in plants

4 (Written by Qianwen Sun)

5 The R-loop, a three-stranded chromatin structure comprising one single-stranded DNA molecule and one RNA:DNA hybrid duplex, is widely distributed in the genome, with 6 essential roles in multiple cellular and disease processes (Garcia-Muse and Aquilera, 7 2019; Brickner et al., 2022; Petermann et al., 2022). Recent advances in genome-wide 8 9 detection methods have broadened our understanding of the distribution and dynamic patterns of R-loops (Xu et al., 2022b). R-loops are involved in many biological 10 processes related to genome regulation, including transcription, replication, DNA 11 damage and repair, and chromatin organization (Zhou et al., 2022a). The biological 12 study of R-loops in plants began in 2013 when we discovered that an R-loop formed on 13 14 the promoter of the antisense IncRNA COOLAIR and affected the expression of FLC (Sun et al., 2013b). In 2017, following the development of ssDRIP-seq (single-strand 15 DNA ligation-based library construction after DNA:RNA hybrid immunoprecipitation, 16 followed by sequencing), the localization of R-loops in the Arabidopsis genome was 17 18 revealed (Xu et al., 2017). The R-loop profiles of other plants have since been disclosed (Figure 5). Through analysis of genome-wide data (Xu et al., 2017; Xu et al., 2020b), 19 some unique features of R-loop distribution in the nuclear genomes of plants have 20 emerged, prompting intriguing research directions in plant R-loop biology. 21

While analyzing the genome-wide distribution of R-loops, we identified a unique group 22 23 of R-loops formed by antisense IncRNAs near transcription start sites (TSS) named asTSS_R-loops, (Xu et al., 2017). Similar patterns of R-loop distribution have also been 24 observed in other plant species, such as rice (Oryza sativa) and maize (Zea mays) 25 (Figure 5, and summarized in (Zhou et al., 2022a)). These conserved patterns raise 26 27 several questions: what are the functions of these asTSS R-loops; how are they transcribed, and is the transcriptional initiation of the antisense IncRNAs specific to 28 particular physiological or pathological responses? Another notable finding was the 29 prevalence of transfer RNA (tRNA)-promoted sense R-loops throughout the genome 30 31 (Xu et al., 2017). We discovered that these intragenic R-loops orchestrated 32 transcriptional interference between Pol II and Pol III, thus regulating the expression of oxidative-responsive genes (Liu and Sun, 2021). Surprisingly, a large proportion of R-33 loops is located in constitutive pericentromeric heterochromatin and overlaps with 34 35 H3K9me2 and H3K27me1 heterochromatic marks in Arabidopsis (Xu et al., 2017). This observation raises intriguing guestions about the functions of R-loops 36 in heterochromatin formation and organization in plants. 37

R-loops play important roles in cellular reprogramming in mammals (Li et al., 2020b;
Yan et al., 2020). During the lifecycle of Arabidopsis, R-loops showed a range of
dynamic changes during generational switches (such as flowering and germination) and

during recovery from long-term heat-stress treatment (Xu et al., 2020b). During the 1 transition from vegetative growth to flowering, R-loop formation decreased dramatically, 2 whereas from flower development to germination, R-loop formation gradually increased, 3 suggesting that the global reprogramming of R-loops also occurs in Arabidopsis. These 4 5 dramatic changes in R-loop formation likely co-occur with other events of genome regulation, such as DNA replication and transcriptional reprogramming. It will be 6 important to explore the biological functions and regulatory mechanisms of R-loop 7 reprogramming during key developmental transitions in plants. Conversely, R-loops 8 likely function in transcriptional reprogramming during physiological and pathological 9 processes. Interestingly, Moore et al. proposed a model of R-loop-mediated 10 transcriptional reprogramming during plant defense responses (Moore et al., 2011), 11 although experimental evidence is still lacking. 12

Most R-loops form and function in *cis*. However, *trans*-formed R-loops may also play 13 14 important roles in plants. For example, the IncRNA APOLO promoted trans-R-loop formation and altered chromatin loop conformation (Ariel et al., 2020). Current detection 15 methods cannot provide information about whether R-loops form in cis or in trans, 16 underscoring the need to develop a high-throughput technique for distinguishing *cis*- or 17 trans-R-loops globally. Moreover, it would be useful to alter the levels of R-loops at 18 specific genomic loci (Liu and Sun, 2021), but there is currently no efficient way to 19 modulate an entire group of R-loops (such as asTSS R-loops) jointly. asTSS R-loops 20 were recently proposed to promote co-transcriptional micro RNA (miRNA) processing 21 (Gonzalo et al., 2022). However, the lack of tools for modulating R-loops makes it 22 23 challenging to study the functions of particular groups of R-loops with similar distribution patterns in the genome. Alternatively, identifying the specific regulators of a particular 24 group of R-loops could help solve this problem. 25

To date, several R-loop modulators have been identified in plants (Zhou et al., 2022a). 26 27 Among these, the evolutionarily conserved RNase H1 proteins specifically remove the RNA moiety in RNA:DNA hybrids, thus resolving R-loops efficiently. The Arabidopsis 28 genome encodes three RNase H1 proteins: AtRNH1A, AtRNH1B, and AtRNH1C (Yang 29 et al., 2017). While AtRNH1B and AtRNH1C are involved in stabilizing the genome 30 31 integrity of semi-autonomous organelles (mitochondria and chloroplasts) (Yang et al., 2017; Cheng et al., 2021; Wang et al., 2021b), the biological function of nucleus-32 localized AtRNH1A is still unclear. The biological functions of RNase H1 proteins and 33 other R-loop regulators in different plant species also need to be further explored. 34

Organisms must integrate and coordinate the activities of different tissues and cell types. Precisely analyzing the genome-wide patterns of R-loops from ultra-low input samples is difficult using current methods (Zhou et al., 2022a). It is imperative to establish ultralow-input (or even single-cell) R-loop profiling techniques to systematically explore the functions of R-loops in critical genomic events in complex tissues. Such tools could be powerful for dissecting R-loop distribution and dynamics in specific cell

types and during specific differentiation programs, such as double fertilization in plants.
As the topological state of the genome could affect R-loop formation, it would be useful
to develop tools to quantitatively measure topological conformation and explore how the
3D genome organization influences R-loop formation. Advanced computational
predictions of R-loops genome-wide could complement experimental approaches for
species with available genome sequence information.

Chloroplasts and mitochondria are semi-autonomous organelles of endosymbiotic origin 7 with their own genetic materials. In the face of complex external environmental 8 conditions and internal growth and developmental factors, how these organelles 9 maintain their genome stability has long been unclear. We recently discovered that R-10 loops act as regulatory centers in determining the stability of organelle genomes (Figure 11 5). R-loops play both positive and negative roles in maintaining the stability of the 12 organellar genome, which not only causes genome instability by modulating head-on 13 transcription-replication conflicts (Yang et al., 2017; Yang et al., 2020b) but also 14 promotes DNA damage repair (Cheng et al., 2021; Wang et al., 2021b). However, our 15 knowledge about how R-loop levels are sensed and adjusted to maintain normal 16 organellar function is still in its infancy. For example, in cells lacking mitochondrion-17 localized AtRNH1B, chloroplast-localized AtRNH1C sensed high R-loop levels and 18 relocalized to mitochondria via an unknown mechanism (Figure 5). This observation 19 raises the following intriguing question: Do plants sense R-loops during chloroplast-20 mitochondria communication? Furthermore, how do plants coordinate and adjust R-loop 21 levels inside and between cells, and how is this process managed in response to 22 23 physiological and pathological processes?

24

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30

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37 Epitranscriptomic mRNA modification: a potent regulatory mechanism in plant

38 development and stress responses

39 (Written by Hunseung Kang)

Epitranscriptomic RNA modifications, which are analogous to epigenetic regulation that 1 involves DNA methylation and histone modifications, are emerging as a new layer of 2 gene regulation. These modifications play a pivotal role in fine-tuning plant development 3 and fitness to changing environmental cues. At least 160 mRNA modifications have 4 been identified to date, among which N⁶-methyladenosine (m⁶A), N¹-methyladenosine 5 (m¹A), and 5-methylcytidine (m⁵C) are common and abundant internal modifications 6 observed in coding RNAs; m⁶A is the most prevalent internal modification in eukaryotic 7 mRNAs (Boccaletto et al., 2018). Methyltransferases (referred to as "writers"), 8 demethylases (referred to as "erasers"), and RNA-binding proteins (referred to as 9 10 "readers") are cellular components responsible for the installation, removal, and interpretation of m⁶A marks, respectively (Figure 6). Recent transcriptome-wide m⁶A 11 mapping, as well as the identification and characterization of m⁶A writers, readers, and 12 erasers in Arabidopsis and model crops, have enhanced our understanding of the 13 dynamics, distribution, regulatory mechanisms, and biological functions of m⁶A 14 methylation in plant development and stress responses. 15 Transcriptome-wide analyses of m⁶A methylation patterns in plants have led to the 16

identification of an RR($m^{6}A$)CH (R = A/G; H = A/C/U) motif found in all eukaryotes (Luo 17 et al., 2014; Duan et al., 2017; Hu et al., 2021) and a URU(m⁶A)Y (Y = C/U) motif 18 unique to plants (Arribas-Hernandez et al., 2018; Arribas-Hernandez et al., 2021a; 19 Arribas-Hernandez et al., 2021b; Hu et al., 2021; Hu et al., 2022). The presence of the 20 plant-specific m⁶A motif, as well as the common m⁶A motif conserved across all 21 eukaryotes, suggests that m⁶A modifications exert multifaceted functions in plants. The 22 m⁶A writers responsible for these modifications include METHYLTRANSFERASE A 23 (MTA), MTB, FKBP12-interacting protein 37 (FIP37), VIRILIZER (VIR), the E3 ubiquitin 24 ligase HAKAI, and FIONA1 (FIO1) (Ruzicka et al., 2017); reviewed in (Hu et al., 2019; 25 Xu et al., 2022a). The three m⁶A erasers, AlkB homolog 2 (ALKBH2), ALKBH9B, and 26 ALKBH10B, have been confirmed as m⁶A demethylases (Duan et al., 2017; Martinez-27 Perez et al., 2017, Zhou et al., 2019) (Figure 6). YT521-B homology (YTH)-domain 28 proteins have been characterized as m⁶A readers that recognize m⁶A marks and affect 29 stability, translation, nucleus-to-cytoplasm movement, and 30 the alternative polyadenylation of m⁶A-modified transcripts (Arribas-Hernandez et al., 2018; Scutenaire 31 et al., 2018; Wei et al., 2018; Hu et al., 2019; Arribas-Hernandez et al., 2021a; Arribas-32 Hernandez et al., 2021b; Song et al., 2021; Hou et al., 2022) (Figure 6). Dynamic and 33 reversible m⁶A methylation play vital roles in embryogenesis, morphogenesis, trichome 34 morphology, root development, and fruit ripening (Ruzicka et al., 2017; Arribas-35 Hernandez et al., 2018; Hu et al., 2019; Zhou et al., 2019; Hu et al., 2022) (Figure 6). 36 Accumulating evidence has highlighted the pivotal roles of m⁶A modifications in plant 37 growth and development. However, several guestions, including the mechanism by 38 which m⁶A is added to, or removed from, mRNA transcripts in a growth stage-39

dependent manner and differentially regulates the abundance of transcripts crucial for
 plant development, remain unanswered.

Mapping and characterization of mRNA modifications in plant stress responses are 3 currently at the nascent stage. Bioinformatics analyses revealed that the expression 4 levels of m⁶A writers, erasers, and readers change differentially in response to diverse 5 stresses (Hu et al., 2019), suggesting a vital role for m⁶A methylation in plant stress 6 responses. Recent molecular evidence has established a link between mRNA 7 modifications and transcript levels involved in plant stress responses (Hou et al., 2021; 8 Hu et al., 2021; Hou et al., 2022). Notably, m⁶A modifications play crucial roles in plant 9 responses to diverse stresses, including salt, drought, and nutrient (nitrate) starvation, 10 by affecting mRNA stability, alternative polyadenylation, and translation efficiency of 11 stress-responsive genes (Hou et al., 2021; Hu et al., 2021; Hou et al., 2022). However, 12 the precise mechanism underlying RNA modification-mediated gene regulation during 13 14 stress adaptation requires further investigation. Therefore, the crucial aspects that remain unexplored are the mechanisms by which RNA modification patterns vary under 15 specific stress conditions and the association of these modifications with stress-induced 16 alterations in transcript and protein levels. 17

Most studies conducted thus far have focused on the cellular components responsible 18 for RNA methylation and their roles in the nucleus and cytoplasm. Chloroplast and 19 mitochondrial RNAs are highly m⁶A-methylated, accounting for 98–100% and 86–90% 20 of the transcripts in chloroplasts and mitochondria, respectively (Luo et al., 2014; Wang 21 et al., 2017b). Therefore, RNA methylation might likely exert crucial roles in plant 22 23 organelles. However, the nature and identity of writers, erasers, and readers in chloroplasts and mitochondria, except m⁴C and m₂⁶A rRNA writers in chloroplasts, are 24 largely unknown (reviewed in (Manduzio and Kang, 2021). Analysis of chloroplast 25 proteomes by liquid chromatography-tandem mass spectrometry and prediction of 26 organelle-localized proteins have revealed that the m⁶A writer components MTA, MTB, 27 and FIP37 found in plant nuclei were also possibly localized in chloroplasts and 28 mitochondria and several putative S-adenosyl methionine (SAM)-dependent 29 methyltransferase proteins are present in the chloroplasts of Arabidopsis (reviewed in 30 31 (Manduzio and Kang, 2021). Further verification of the methyltransferase activity of these putative writer proteins, as well as the previously unknown erasers or readers in 32 chloroplasts and mitochondria, will help elucidate the significance of RNA modifications 33 in plant organelles. 34

Rapid progress in transcriptome-wide mapping and the identification of writers, readers,
 and erasers have unraveled the regulatory roles of m⁶A modification in plant
 development and stress responses. Nonetheless, many challenges remain in mapping
 m⁶A modifications at single-base resolution using recently advanced sequencing
 methods, including Nanopore direct transcriptome deep sequencing (RNA-seq),
 MAZTER-seq, m⁶A-REF-seq (m⁶A-sensitive RNA-Endoribonuclease-Facilitated

sequencing), and miCLIP-seq (m⁶A individual-nucleotide-resolution cross-linking and 1 immunoprecipitation combined with high-throughput sequencing). Furthermore, 2 characterizing novel cellular components of writers, readers, and erasers in crops will 3 help firmly establish the molecular link between m⁶A, crop productivity, and stress 4 adaptation. Recent findings have suggested that m⁶A is associated with LLPS, which 5 expands the repertoire of regulatory mechanisms crucial for cellular responses to 6 developmental and environmental cues (Scutenaire et al., 2018; Ries et al., 2019; Song 7 et al., 2021). Integrating these molecular insights to the regulatory roles of m⁶A 8 modification with novel genome-editing technologies, including A-to-G base editing to 9 modify potential m⁶A sites and clustered regularly interspaced short palindromic repeat 10 (CRISPR)/CRISPR-associated nuclease 13 (Cas13)-based targeted RNA methylation 11 (Liu et al., 2019; Li et al., 2020a), will greatly facilitate epitranscriptomics research and 12 lead to the development of a potential strategy for breeding stress-tolerant crops via 13 14 precisely engineered RNA modifications. Further exploration of this field is warranted, and we anticipate exciting discoveries in the near future. 15

16

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20

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29 Decoding the grammar of plant long noncoding RNAs

30 (Written by Federico D. Ariel and Martin Crespi)

31 The inspection of the presence and combination of domains within a protein is generally a good starting point to infer its potential molecular action. This information is then 32 complemented with subcellular localization studies, biochemical characterization, 33 analysis of expression patterns of the encoding gene across multicellular organisms and 34 35 genetic approaches to propose a biological role of the given gene in plants. By contrast, the comprehensive functional characterization of IncRNAs (Wierzbicki et al., 2021) is a 36 challenging task that should take into account (i) their promiscuous or specific 37 interaction with other molecules based on their sequence and/or structure; (ii) their 38 redundancy with other unrelated transcripts; (iii) their subcellular localization; (iv) their 39 40 role within molecular regulatory networks; and (v) an eventual RNA biological activity

(Figure 7). In the last 15 years, thousands of IncRNAs have been annotated from a 1 growing number of plant species, although their functional characterization lags behind, 2 thus severely hindering the differentiation between transcriptional noise and biologically 3 relevant noncoding transcripts. Identifying general molecular features linking specific 4 5 IncRNAs with their targets have uncovered certain mechanisms. For instance, target mimicry of miRNAs (RNA molecules acting as decoy of miRNAs blocking their activity) 6 was demonstrated for INDUCED BY PHOSPHATE STARVATION 1 (IPS1) and could 7 be later predicted in silico for other IncRNAs across species (Franco-Zorrilla et al., 8 2007). However, for the large majority of IncRNAs acting through other molecular 9 mechanisms, there is no evident features to define their targets in order to dissect the 10 molecular basis governing their action in plants. 11

Upon extensive annotation of IncRNAs across species, future screenings for biological functions, likely based on systematic CRISPR-derived approaches, may empower the selection of novel relevant IncRNAs for in-depth molecular characterization. In addition, integration of IncRNA expression patterns from transcriptomic data of multiple wild-type plants, mutants, and natural accessions in response to environmental and developmental cues will position the IncRNA of interest within particular regulatory networks driving plant development and/or adaptation to the environment.

Specific IncRNAs have been shown to interact with protein partners in ribonucleoprotein 19 (RNP) complexes (modulating their stability, subcellular localization, or their activity), 20 DNA (forming RNA-DNA duplexes known as R-loops), or other transcripts (such as 21 antisense RNAs, forming paired RNA regions triggering mRNA degradation or 22 23 promoting translation) (Lucero et al., 2021). Future research to generalize these interactions may include global identification of RNAs forming R-loops or interacting with 24 specific RNP complexes involved in splicing modulation (Rigo et al., 2020) or the 25 translational machinery (Bazin et al., 2017). Another emerging mechanism is the 26 27 interaction of IncRNAs with chromatin-related proteins linked to epigenetic regulations such as LIKE HETEROCHROMATIN PROTEIN 1 (LHP1), CURLY LEAF (CLF), 28 MEIOTIC F-BOX (MOF), ARABIDOPSIS TRITHORAX-LIKE PROTEIN1 (ATX1) or 29 WD40 REPEAT 5A (WDR5a) (Fonouni-Farde et al., 2021) although their binding 30 31 specificity remains uncertain, or even with transcription factors (e.g. WRKY42, (Moison et al., 2021)). In addition, the identification of nascent RNAs (Kindgren et al., 2020) as 32 well as chromatin-associated IncRNAs based on chromatin isolation and high 33 throughput sequencing techniques will further contribute to creating a matrix of IncRNA 34 35 features underlying their function. Altogether, mapping IncRNA interactions with DNA, chromatin, and proteins involved in a wide range of mechanisms in model and crop 36 plants should set the stage for a comprehensive classification of IncRNAs enabling the 37 search of singularities and commonalities behind the functions of noncoding transcripts. 38

The identification of protein partners and IncRNA-interacting nucleic acids using biotinylated probes for the purification of IncRNA-containing complexes followed by mass spectrometry or DNA sequencing is an initial key goal to define the IncRNA interactome, despite the potential artifacts linked to these approaches (Machyna and Simon, 2018). Alongside the genome-wide identification of IncRNAs participating in alternative RNP complexes, the detailed characterization of selected IncRNA actions on these complexes remains essential to better understand the diversity of regulatory mechanisms involving noncoding transcription.

Another major question in IncRNA biology and biochemistry concerns transcript 7 structure (Zhu et al., 2021). Secondary and tertiary structures of RNAs are very likely 8 determinant features for their dynamic interaction with proteins and other partners. 9 Considering that plants cannot modulate their body temperature, the structure of 10 IncRNAs may serve as potential versatile molecules acting as thermosensors in order to 11 rapidly adjust epigenomic features and alternative splicing, two major processes 12 affected by ambient temperature (John et al., 2021; Perrella et al., 2022). A growing 13 14 number of prediction tools based on classical and machine learning approaches have shed light on this field (Bugnon et al., 2022), although the biochemical characterization 15 of individual or groups of plant IncRNAs is just starting. In general, genome-wide 16 approaches for the mapping of double-stranded RNAs (dsRNAs) or chemical 17 18 degradation profiles to reconstruct transcript structures fail to deliver enough data about low-abundance IncRNAs. However, in vitro transcription of selected IncRNAs followed 19 by biochemical approaches ignores the enormous collection of epitranscriptomic 20 modifications as well as their *in vivo* interaction with partner molecules, which are likely 21 to affect RNA structure (Miller et al., 2022). 22

23 Similar to the study of metazoan IncRNAs, cell biology techniques, notably single molecule RNA (smRNA) fluorescence in situ hybridization (FISH) (Duncan et al., 2017), 24 can contribute to our understanding of the mechanisms involving specific IncRNAs. As a 25 complement to subcellular fractionation studies followed by high-throughput 26 27 sequencing, smRNA FISH may not only indicate whether a given IncRNA accumulates in the nucleus or the cytoplasm, but also reveal its distribution in "speckles", or 28 localization in specific loci, in subcellular compartments, in non-membranous organelles 29 or particles. However, the technical difficulties related to the presence of cell wall 30 31 barriers in plant tissues prevents the accessibility of fluorescent oligonucleotide probes, 32 thus delaying the massive use of this approach by most plant RNA biology groups, in comparison to labs working on mammalian cell culture models. 33

The fields of plant RNA biology and biochemistry will need to integrate cell biology, RNP proteomics, genomic and genetic approaches to unveil the function and evolution of the noncoding transcriptome, in particular during differentiation and environmental stress responses. Evolutionary analysis at a global level (e.g. involving synteny) of lncRNAs exhibiting common features (e.g. integration into specific RNPs), together with the indepth characterization of specific leading cases, will achieve a better understanding of the structures and sequences (likely very short) setting the specificity rules of their 1 interaction with partner molecules. As the RNA interactome ultimately determines their

2 function, these integrated approaches will hopefully help us uncover the grammar of

- 3 plant IncRNAs.
- 4

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7 8

9 Emerging and long-standing questions about miRNA biogenesis in plants

10 (Written by Axel Giudicatti and Pablo A. Manavella)

From the point of view of RNA biology, miRNAs are exciting molecules. Not only do 11 mature miRNAs target other RNA molecules to block their translation or trigger their 12 degradation, but their precursors undergo nearly all the regulatory features described in 13 14 this article. For instance, many MIRNA genes contain introns that affect the processing of the primary transcripts (pri-miRNAs) (Stepien et al., 2017); the ribonucleotides of 15 miRNA precursors can be modified or edited to change their regulatory outcome 16 (Mingardi et al., 2018; Bhat et al., 2020); pri-miRNA secondary structure fluctuations 17 define miRNA biogenesis (Wang et al., 2018b; Re et al., 2019); even asTSS_R-loops 18 were recently shown to promote co-transcriptional processing of miRNAs (Gonzalo et 19 al., 2022). These features make miRNAs a unique entity where many aspects of RNA 20 biology converge. Even after more than 20 years of research, all these aspects of 21 miRNA biology present unresolved questions and intriguing gaps in our knowledge. For 22 23 instance, although we know that there is a crosstalk between splicing and pri-miRNA processing (Stepien et al., 2017), it is unclear how these two processes interact. The 24 transcription and processing of pri-miRNAs is coupled (Fang et al., 2015; Gonzalo et al., 25 2022). This observation opens the possibility that the crosstalk between pri-miRNA 26 27 processing and splicing only exists for miRNAs processed co-transcriptionally where both machineries, the spliceosome and microprocessor, meet. In this scenario, it is 28 unclear whether the miRNA processing factors and splicing factors act cooperatively or 29 simply interfere entropically with each other over the nascent pri-miRNAs during 30 31 maturation. Advances in RNA sequencing technologies, especially of nascent RNAs, will help our understanding of how these two processes are connected (Figure 8C). 32

On its own, the discovery of coupling between transcription and miRNA processing, 33 initially suggested by the ground-breaking work of Fang et al (Fang et al., 2015) and 34 35 further confirmed in 2022 (Gonzalo et al., 2022), opened many exciting new avenues of inquiry. The recruitment of the microprocessor to MIRNA loci is a well-reported 36 phenomenon (Fang et al., 2015; Cambiagno et al., 2021). However, how the 37 microprocessor specifically recognizes these loci over any other Pol II-transcribed 38 region remains an enigma. Still, the association of the microprocessor to MIRNAs 39 40 requires the presence of the pri-miRNA transcript (Fang et al., 2015). Thus, it is possible

that the Dicing complex recognizes the stem-loop structure within pri-miRNA transcripts, 1 thereby giving specificity to the system. Co-transcriptional miRNA processing appeared 2 favored in those loci containing asTSS R-loops (Gonzalo et al., 2022). These three-3 stranded chromatin structures may also provide an initial signal promoting the 4 5 recruitment of the microprocessor to these loci, although their functions in this process are still merely hypothetical. Nevertheless, this result raises the possibility that the 6 three-stranded hybrid is the platform upon which the microprocessor is built. It will be 7 interesting to study whether any of the proteins proposed to link the microprocessor to 8 chromatin have affinity for R-loops, either for the single-stranded DNA or the RNA/DNA 9 hybrid (Figure 8B). The assembly of the processing complex also presents a 10 challenging, but very relevant, problem to solve; which is the hierarchical order of 11 recruitment of the microprocessor components to MIRNA loci? Another compelling 12 question raised from the discovery of the processing of nascent pri-miRNAs is whether 13 14 co-transcriptionally processed miRNAs have distinct functions. In this sense, it was recently shown that the protein HASTY (HST) is required for both the assembly of the 15 microprocessor at MIRNA loci and to promote the non-cell-autonomous function of 16 miRNAs (Brioudes et al., 2021; Cambiagno et al., 2021). It is therefore possible that 17 miRNAs processed during transcription take a particular road that makes them mobile 18 molecules (Figure 8E). Perhaps this pool of miRNAs somehow avoids loading into 19 ARGONAUTE 1 (AGO1), an event proposed to lock miRNAs inside the cell, preventing 20 their movement (Devers et al., 2020; Fan et al., 2022; Voinnet, 2022). It is curious that 21 the precise mechanisms of miRNA movement between cells and whether such 22 movement is chaperoned, still remains unknown. In fact, this question is probably one of 23 the longest-standing questions in the field. 24

Among the four DICER-Like (DCL) enzymes in Arabidopsis, DCL1 is the main actor in 25 miRNA processing, due to its nuclear localization and preference to process imperfect 26 27 stem-loop folded RNAs. Within the pri-miRNA stem-loop, DCL1 recognizes structural features that guide processing to release a unique miRNA duplex (Bologna et al., 2013; 28 Manavella et al., 2019). It was recently shown that the folding of pri-miRNAs can be 29 altered, consequently modifying processing efficiency (Wang et al., 2018b). In addition, 30 31 nucleotides at pri-miRNAs can be modified and even edited, although the influence that these events have over the miRNA processing were not demonstrated in plants 32 (Mingardi et al., 2018; Bhat et al., 2020). The role of RNA editing, modification, and re-33 folding in miRNA processing are just emerging as important regulatory mechanisms and 34 deserve our attention. It is expected, as plants are nonthermogenic organisms, that the 35 secondary structure of plant pri-miRNAs will fluctuate with ambient temperature, likely 36 affecting their processing. Thus, it can be envisioned that some miRNAs may even act 37 as thermosensors (Figure 8A). Although we do have some evidence that temperature 38 changes how miRNAs are processed (Re et al., 2019), much more needs to be done on 39 40 this subject.

D-bodies are one of the most intriguing elements in the miRNA pathway. These discrete 1 membraneless nuclear speckles are the typical localization of many fluorescently-2 tagged miRNA biogenesis proteins (Fang and Spector, 2007). The localization of these 3 proteins led to the proposal that D-bodies are the center of miRNA processing in plants. 4 5 A recent study showed that D-bodies arise through SERRATE (SE)-mediated phase separation (Xie et al., 2021). Disruption of SE phase separation, and thus D-body 6 formation, by deleting the N-terminal intrinsically disordered region (IDR) of SE reduces 7 miRNA accumulation, supporting the idea that D-bodies are sites of pri-miRNA 8 processing. The role of D-bodies in miRNA processing is also supported by several 9 studies showing a correlation between D-body formation and miRNA production. 10 Intriguingly, other studies have shown that the disappearance of D-bodies does not 11 affect the ability of the cell to produce miRNAs (discussed by (Mencia et al., 2022)). 12 This observation suggests that D-bodies are not the sole place of miRNA processing 13 14 and raises the possibility that compensatory mechanisms act to offset the reduction of miRNAs caused by the loss of D-bodies. While we now know that miRNA can be 15 processed co-transcriptionally (Fang et al., 2015; Cambiagno et al., 2021; Gonzalo et 16 al., 2022), many pri-miRNAs are partially or entirely processed in the nucleoplasm, likely 17 in D-bodies (Gonzalo et al., 2022). Thus, a balance between these two processing sites 18 may buffer any fluctuation in processing and maintain stable levels of miRNAs. Given 19 the current data, it is hard to simply categorize D-bodies as the only place where miRNA 20 biogenesis occurs. It is also possible that D-bodies are not unique entities but rather a 21 collection of small micro-reactors of different compositions and functions (Figure 8D). 22 23 This idea goes along with the finding that despite localizing to D-bodies, some miRNA factors do not co-localize with each other (Tomassi et al., 2020). We previously 24 discussed several possible scenarios for D-body functions (Mencia et al., 2022). 25 Defining the nature and role of D-bodies and their crosstalk with co-transcriptional 26 27 processing is at the frontier of miRNA research, although it is a technically challenging goal. Future studies applying state-of-the-art in vivo immunostaining and biochemical 28 approaches will certainly surprise us with new discoveries about these nuclear 29 speckles. 30 31 These are only a few of the many open questions regarding how miRNAs are produced

and do not even consider the equally large number of questions we have regarding how miRNAs act once loaded into the RNA-induced silencing complex (RISC) and how these molecules are stabilized or degraded when necessary.

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7 **Cross-kingdom RNAi**

8 (Written by Qiang Cai and Hailing Jin)

Over the years, studies on extracellular RNAs, including small RNAs (sRNAs), have 9 focused mostly on their movement between cells and tissues within an organism (Liu 10 and Chen, 2018; Huang et al., 2019). Naturally occurring sRNA trafficking across 11 organismal boundaries between interacting organisms that induces gene silencing in 12 the counter party, a biological phenomenon named cross-kingdom/species RNA 13 14 interference (RNAi), was first described in plant-fungal interactions (Weiberg et al., 2013). During infection, the gray mold fungal pathogen Botrytis cinerea delivers its 15 sRNAs, called sRNA effectors, into plant cells and hijacks the plant RNAi machinery to 16 silence those host genes that are involved in plant immunity (Weiberg et al., 2013). 17 Similar phenomena were later observed in many plant pathogens and parasites. For 18 example, sRNAs from the fungal pathogens Verticillium dahlia (causing verticillium wilt 19 of cotton [Gossypium hirsutum]) and Puccinia striiformis (causing stripe rust of wheat 20 [Triticum aestivum]) can move into their plant host and silence plant defense genes 21 (Wang et al., 2016; Wang et al., 2017a). Similarly, oomycete pathogens, such as 22 23 Hyaloperonospora arabidopsidis (causing downy mildew of Arabidopsis) (Dunker et al., 2020), also utilize cross-kingdom RNAi to achieve aggressive infection. Furthermore, 24 miRNAs from parasitic plant dodders (Cuscuta campestris) act as cross-species 25 regulators of gene expression in their plant hosts, suggesting that mobile sRNAs act as 26 27 virulence factors during parasitism (Shahid et al., 2018). Cross-kingdom RNAi is not limited to pathogenic interactions but also exists in symbiotic interacting systems. A 28 recently discovered fungal miRNA from the beneficial ectomycorrhizal fungus *Pisolithus* 29 microcarpus enters Eucalyptus grandis root cells and stabilizes the symbiotic interaction 30 31 by silencing several nucleotide-binding (NB)-ARC domain-containing proteins from the host (Wong-Bajracharya et al., 2022). Even for prokaryotic pathogens that do not have 32 a canonical RNAi pathway, rhizobial tRNA-derived short RNAs act as functional sRNAs 33 moving into plant cells to silence nodulation-related target genes (Ren et al., 2019). 34 35 Most strikingly, the molecular mechanism underlying cross-kingdom RNAi is also conserved. The sRNAs from the fungal pathogens B. cinerea and V. dahlia, the 36 oomycete pathogen H. arabidopsidis, and the rhizobium were all found to be loaded into 37 the plant host AGO1 to silence host target genes (Weiberg et al., 2013; Wang et al., 38 2016; Ren et al., 2019; Dunker et al., 2020). 39

Recent studies have shown that cross-kingdom RNAi is bidirectional, and many plant 1 species can also transport endogenous sRNAs into their interacting pathogens (Cai et 2 al., 2021; Liu et al., 2021). For example, Arabidopsis plants send miRNAs, phased 3 secondary small interfering RNAs (phasiRNAs), and other endogenous short interfering 4 5 RNAs (siRNAs) into interacting B. cinerea cells (Cai et al., 2018). These transported host sRNAs can silence *B. cinerea* virulence-related genes, many of which are involved 6 in fungal vesicle-trafficking pathways (Cai et al., 2018). Cross-kingdom sRNA trafficking 7 from host plants into pathogens was also observed in other plant-fungal systems, such 8 as cotton-V. dahliae and wheat-F. graminearum interaction systems (Cai et al., 2021). 9

It has been demonstrated that plant sRNAs are transported into fungal cells mainly by 10 extracellular vesicles (EVs) (Cai et al., 2018). EVs are heterogeneous membrane-11 encapsulated structures that transport different RNA and protein cargoes between cells 12 (Mathieu et al., 2019). EVs play an important role in sRNA trafficking between cells and 13 tissues in both animal and plant systems (Cai et al., 2019; Mathieu et al., 2019). Like 14 animal cells, a heterogeneous population of EVs exists in plants (Cai et al., 2019; 15 Huang et al., 2021b). In Arabidopsis, a distinct class of EVs, called tetraspanin (TET)-16 positive exosomes, are responsible for secretion and transport of functional sRNAs and 17 play a significant role in cross-kingdom RNAi and plant-microbial interactions (Cai et al., 18 2018; He et al., 2021). 19

How specific plant sRNAs are selectively loaded into EVs has long remained poorly 20 understood. A recent study identified a list of EV-localized RNA-binding proteins, 21 including AGO1, DEAD-box RNA helicases (RH11, RH37, and RH52), and ANNEXIN 1 22 23 and 2 (ANN1 and ANN2) (He et al., 2021). AGO1, RH11, and RH37 were shown to selectively bind to a set of sRNAs that are found in EVs, and contribute to selective 24 sRNA loading into EVs, mostly TET-positive exosomes, whereas ANN1/2 bind to RNAs 25 non-specifically. The level of sRNAs is reduced in EVs isolated from ann1 ann2 26 27 mutants, which indicates that ANN1/2 are involved in sRNA stabilization in EVs, although they do not contribute to selective sRNA loading (He et al., 2021). 28

29 Research on cross-kingdom RNAi and sRNA trafficking is still in its infancy, and increasing studies demonstrate that mobile RNA molecules are important regulatory 30 31 elements of the interaction between hosts and interacting organisms (Huang et al., 2019). Bidirectional cross-kingdom RNAi has been developed during the co-32 evolutionary arms race between hosts and pathogens, which has become a widespread 33 molecular regulatory mechanism in plant-microbial interaction and plays a significant 34 35 role in host immunity and pathogen virulence (Huang et al., 2019). Current studies show that EVs are essential in transporting sRNAs from the plant hosts to pathogens (Cai et 36 al., 2021). EV-mediated sRNA transport has evolved in both plant and animal systems, 37 suggesting that it is likely a conserved mechanism for cell-to-cell communication. The 38 current understanding of cross-kingdom RNA transport is just the tip of the iceberg. 39 40 Many questions remaining to be answered in this field are: i) Can pathogen sRNAs act

as effector molecules, and can plants sense them as pathogen-associated molecular 1 patterns (PAMPs)? ii) Do plant EVs also transport other classes of RNAs, i.e., mRNAs 2 and IncRNAs, into pathogen cells to inhibit virulence? iii) Are there other mechanisms 3 by which plant RNAs are selectively loaded into EVs? iv) Do fungal pathogens also 4 5 utilize EVs to deliver RNAs into host plants? v) Besides EVs, do other pathways contribute to cross-kingdom RNA transport? A better understanding of RNA 6 communications between interacting organisms will contribute to the development of 7 new strategies for disease control and crop protection, such as EV-based sRNA 8 9 fungicides.

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16 Why does germline development require specialized small RNAs?

17 (Written by Xiaoqi Feng)

Small interfering RNAs (siRNAs) move between cells and exert regulatory functions 18 during plant and animal development (Chen and Rechavi, 2022). Specialized, 19 somatically produced siRNAs play essential roles during plant germline development. 20 Similarly, a special army of siRNAs operates in the animal germline, called Piwi-21 interacting sRNAs (piRNAs) (Ozata et al., 2019). A central question arising from these 22 23 reproductive-cell-specific siRNAs is why such specificity? What is intrinsic about sexual reproduction that requires specialized siRNAs? This is arguably one of the most exciting 24 questions in RNA and reproductive biology. As these siRNAs have diverse, pleiotropic 25 roles during reproductive development, investigation of multiple eukaryotic lineages is 26 27 necessary to resolve this question.

Pioneering evidence of soma-germ siRNA movement in plants came from Arabidopsis 28 pollen where 21-nt siRNAs associate with derepressed transposable elements (TEs) in 29 the sperm companion cell, the vegetative cell (Slotkin et al., 2009) (VC; Figure 9). 30 31 These siRNAs, but not the TE transcripts, accumulate in the sperm cell, suggesting that VC siRNAs can move into the sperm to reinforce TE silencing (Figure 9). Such TE 32 reactivation (and hence associated siRNAs) is likely confined to gamete companion 33 cells, as it is largely driven by a DNA demethylase, DEMETER (DME) (He et al., 2019), 34 35 whose encoding gene is specifically expressed in companion cells (Feng et al., 2013). Indeed, TEs that are demethylated by DME in the VC are hypermethylated in sperm 36 (where DME is not expressed) in a DME-dependent manner (Ibarra et al., 2012). DME 37 is also expressed in the egg companion cell, the central cell, and likely activates siRNAs 38 moving into the egg (Ibarra et al., 2012; Feng et al., 2013) (Figure 9). 39

Since the above-mentioned study by Slotkin et al., it has become clear that somatic 1 cells surrounding the germline produce distinct populations of siRNAs. An example is a 2 variant form of the small RNA-directed DNA methylation pathway (RdDM) in meiocyte 3 nurse cells (the tapetum; Figure 9). RdDM methylates TEs using 24-nt siRNAs 4 5 transcribed by RNA Polymerase IV (Pol IV), which is recruited by putative chromatin remodelers, CLASSY1-4 (CLSY1-4). Somatic tissues mainly express CLSY1 and 6 CLSY2, and their proteins recruit RdDM to thousands of repeats. In tapetal cells and 7 ovules, CLSY3 is expressed at much higher levels than CLSY1/2, leading to a distinct 8 24-nt siRNA profile with the vast majority of siRNAs coming from a few hundred loci 9 (Long et al., 2021; Zhou et al., 2022b). 10

Although these germline siRNAs were discovered due to their roles in TE silencing, 11 increasing evidence links them to gene regulation, for example during pollen 12 development in Capsella (Wang et al., 2020). 24-nt siRNAs produced by tapetal cells 13 14 methylate genes with similar but not identical sequences in male meiocytes (Walker et al., 2018; Long et al., 2021) (Figure 9), thereby regulating the splicing of a meiotic gene 15 and facilitating meiosis (Walker et al., 2018; Long et al., 2021). As the TE-silencing and 16 gene regulatory functions of germline siRNAs go hand in hand, it is tantalizing but 17 difficult to tease apart which is the primary function, if such a distinction is possible. 18

Compounding the complexity, germline siRNA biogenesis varies among plant species. 19 Although Arabidopsis meiotic 24-nt siRNAs are produced by Pol IV and RdDM, similarly 20 abundant 24-nt phased secondary siRNAs (phasiRNAs) in maize and rice tapetal cells 21 are produced by cleavage of noncoding Pol II transcripts by a miRNA (Liu et al., 2020). 22 23 Monocot anther wall cells also accumulate an earlier wave of 21-nt phasiRNAs. Both 21-nt and 24-nt phasiRNAs have been proposed to move into meiotic cells and are 24 important for male fertility, especially under certain environmental conditions, although it 25 is still unclear why (Liu et al., 2020; Zhou et al., 2022c). 26

27 Another challenge is to elucidate the link between siRNA-mediated gene regulation and germ cell differentiation. The most well-understood example is the differentiation of 28 female mejocytes, called megaspore mother cells (MMCs). Normally, only one 29 subepidermal (L2) cell adopts MMC fate and undergoes meiosis in each ovule (Figure 30 31 9). Multiple MMCs differentiate in mutants of RdDM or 21-22 nt trans-acting siRNA (tasiRNA) pathways (Olmedo-Monfil et al., 2010; Su et al., 2020). Key components of 32 both pathways are specifically expressed in apical epidermal (L1) cells, suggesting that 33 these L1-produced siRNAs are essential for suppressing MMC fate in L2 cells (Figure 34 35 9). Importantly, causal links were made between L1-produced tasiRNAs, the repression of AUXIN RESPONSE FACTOR 3 (ARF3) in L2 cells, and the suppression of MMC fate 36 (Su et al., 2017; Su et al., 2020). However, this is unlikely the sole regulatory 37 mechanism for MMC differentiation, as mutations of other epigenetic pathways, such as 38 METHYLTRANSFERASE 1 (MET1)-mediated DNA methylation maintenance (Li et al., 39 40 2017), also cause a similar supernumerary MMC phenotype. An indirect mechanism is

also plausible, e.g. failure of epigenetic silencing interferes with MMC meiosis or
function, which activates neighboring cells to adopt MMC fate as a compensating
mechanism.

A converging feature of germline siRNAs is their non-cell-autonomy, which raises the 4 5 question of why germ cells do not produce the siRNAs themselves, but instead rely on neighboring companion/nurse cells. Many ideas have arisen: perhaps siRNA 6 biosynthesis exposes certain risks as it generally involves transcription of TEs, or nurse 7 cells might afford to sensitize their chromatin environment to unfurl their genome and 8 reveal potentially hazardous TEs (Feng et al., 2013), or maybe it is a question of why 9 not, as nurse cells are already geared to provide a wide range of nutrients and other 10 molecules to germ cells. These are exciting concepts ripe for exploration. 11

For Arabidopsis tapetal siRNAs, non-cell-autonomy may allow more precise control of 12 germline transcriptional regulation. Canonical RdDM is self-reinforcing, as DNA 13 14 methylation promotes the generation of methylation-inducing siRNAs by recruiting Pol IV. The methylation arm of RdDM is tuned more aggressively in meiocytes to target 15 broader sequences, which allows the targeting of genes and fast-evolving TEs (Long et 16 al., 2021). However, given the self-reinforcing nature of RdDM, this broad-targeting 17 18 ability needs to be tightly controlled to prevent the long-term establishment of RdDM at aenomic Such control achievable 19 inappropriate regions. is bv cellular compartmentalization: 24-nt siRNA biogenesis is confined to the tapetum, whereas 20 broad-targeting competence is restricted to male meiocytes (Long et al., 2021). 21

Understanding how germline siRNAs move between cells remains technically challenging. Plasmodesmata provide symplastic connections between daughter cells and are known to prevail in several scenarios of germline siRNA movement (Liu et al., 2020; Long et al., 2021). However, in which form(s) and how does the silencing signal move (Chen and Rechavi, 2022)? Furthermore, one cannot exclude the possibility of an apoplastic transport mechanism (reviewed before in the context of cross-kingdom RNAi), warranting further investigation.

Finally, germline siRNAs undoubtedly have functions beyond those in germ cells. 29 siRNAs in sperm can act as quantitative measures of paternal genome dosage, whose 30 31 imbalance with maternal dosage causes seed abortion (Wang et al., 2018a). Similarly, 32 encountering gamete siRNAs in the zygote could, in theory, assess the compatibility of parental genomes, leading to hybridization barriers (Bourc'his and Voinnet, 2010). 33 Although debated, endosperm siRNAs have also been proposed to move into the 34 35 embryo, where they may exert a transgenerational effect (Bourc'his and Voinnet, 2010). siRNA pathways are known to be environmentally sensitive and malleable. Thus, 36 germline siRNAs might be inherited by the next generation to facilitate memory of the 37 environment and regulate the development of the offspring accordingly. The 38 transgenerational effect of siRNAs (if any) remains an exciting area for future 39 40 investigation.

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11 The roles of small RNAs in the regulation of agronomic traits of crops

12 (Written by Yijun Qi)

Our knowledge of the biogenesis, action mode and biological roles of small RNAs has mostly been obtained from studies in Arabidopsis. However, findings in Arabidopsis cannot always be reasonably extrapolated to crops. Studies in crops, despite still being limited, have revealed that small RNAs play unexpected roles, particularly in the regulation of traits of agronomic significance.

Dozens of miRNAs have been shown to regulate crop development, metabolism and 18 stress responses. For instance, miR156, one of the most conserved miRNAs among 19 plant species, regulates juvenile to adult transition in Arabidopsis (Wang et al., 2009; 20 Wu et al., 2009a). However, in rice, miR156 not only helps shape plant architecture but 21 also regulates grain development and filling (Jiao et al., 2010). The conserved miR396, 22 23 which in Arabidopsis regulates plant development, targets and regulates the transcription factor gene HaWRKY6 in sunflower (Helianthus annuus) during heat 24 response (Giacomelli et al., 2012). There are many species-specific miRNAs in crops. 25 For example, miR528, a monocot-specific miRNA, targets a number of genes involved 26 27 in a variety of developmental processes or biotic and abiotic stress responses (Chen et al., 2019). How conserved miRNAs gain more regulatory functions and how species-28 29 specific miRNAs have been acquired by certain crops remain to be fully elucidated. Dissection of diversified roles of miRNAs in crops will greatly improve our understanding 30 31 of the range of miRNA-mediated regulation.

32 In addition to canonical 21-nt miRNAs, there is a distinct class of 24-nt long miRNAs, referred to as ImiRNAs, in rice (Wu et al., 2009b). ImiRNAs regulate transcription via 33 directing DNA methylation at target sites (Wu et al., 2010). It remains unclear how 34 35 prevalent ImiRNAs are among crops. ImiRNAs that have been functionally characterized were all found to regulate rice biotic stress responses (Jiang et al., 2020a; Zhou et al., 36 2020; Campo et al., 2021). This result raises the question as to whether ImiRNAs 37 evolved for plant stress responses and adaptation to environmental changes. 38 Systematic identification of ImiRNAs and their target genes in different crops will be 39 40 necessary for a better understanding of ImiRNA evolution and function.

Twenty-four-nt siRNAs are produced mainly from TEs and direct DNA methylation at 1 target loci through RdDM. While Arabidopsis mutants lacking RdDM do not show 2 obvious phenotypes, rice RdDM mutants have pleiotropic alterations, including 3 dwarfism, an increase in rice tillering and a reduction in rice panicle size (Wei et al., 4 5 2014; Xu et al., 2020a). In maize, loss of 24-nt siRNAs leads to dwarfism, altered leaf polarity and development of feminized tassels (Alleman et al., 2006). These findings 6 indicate that 24-nt siRNAs are important regulators of agronomic traits in crops. The 7 more prevailing regulatory role of 24-nt siRNAs in rice and maize could be explained by 8 the fact that TEs are very abundant and dispersed in euchromatic regions in these 9 plants, which greatly increases the likelihood that RdDM at TEs regulates nearby genes. 10 Indeed, increased tillering in rice RdDM mutants is attributed to loss of RdDM at 11 miniature inverted-repeat transposable elements (MITEs) near MIR156d/j and D14, 12 which control rice tillering (Xu et al., 2020a). Interestingly, it has recently been shown 13 14 that 24-nt siRNA can direct DNA methylation at imperfectly matched targets in Arabidopsis and cabbage (Brassica rapa) (Fei et al., 2021; Long et al., 2021; Burgess et 15 al., 2022), which may greatly increase the range and complexity of RdDM-mediated 16 gene regulation. For most 24-nt siRNAs, their tissue-specific expression, their targets, 17 18 and the effects of their loss remain unknown.

PhasiRNAs, secondary siRNAs that are produced following miRNA-directed target 19 mRNA cleavage, can be 21 or 24 nt in length, depending on the miRNA trigger. 20 PhasiRNAs are the predominant type of small RNAs in anthers in monocots, suggesting 21 that they play a pivotal role in crop reproduction. Supporting this notion, loss of 21-nt 22 23 phasiRNAs, or their activity, in rice leads to pollen sterility (Jiang et al., 2020b), and overproduction of 21-nt phasiRNAs at the *Pms1* locus results in photoperiod-sensitive 24 male sterility, which allows the establishment of a two-line system for hybrid rice 25 breeding (Fan et al., 2016). 21-nt phasiRNAs were found to facilitate the progression of 26 27 meiosis by directing target mRNA cleavage (Jiang et al., 2020b; Zhang et al., 2020). As these targets are regulated for successful meiosis, investigation of their functions could 28 29 be a shortcut to discovering genes and mechanisms important for crop reproduction. Loss of 24-nt phasiRNAs causes reduced pollen fertility and seed-setting rate in rice 30 31 and temperature-sensitive male sterility in maize. There is some evidence supporting 32 the idea that 24-nt phasiRNAs direct DNA methylation in *cis* (Zhang et al., 2021). Whether they can direct DNA methylation in *trans* and whether DNA methylation, if 33 established, can be passed to next generation and regulates grain development remain 34 35 to be explored.

tRNA-derived small RNAs (tsRNAs) and rRNA-derived small RNAs (rsRNAs) are two classes of small RNAs that have recently been identified. Whereas we still have limited information about the expression profile, modes of action, and biological roles of rsRNAs in plants, tsRNAs have been profiled in Arabidopsis (Ma et al., 2021). tsRNA levels appear to undergo dynamic changes in response to abiotic and biotic stresses. A 19-nt 5' tsRNA produced from tRNA-Ala regulates anti-fungal defense in Arabidopsis
 (Gu et al., 2022). tsRNAs have not been well characterized in crops and their functions
 remain to be revealed. It will be also interesting to investigate whether they are widely
 involved in stress responses in crops.

5 Because many agronomic traits are controlled by small RNAs, manipulation of small RNA-mediated gene regulation has emerged as an important strategy to achieving desired agronomic traits. Unlike overexpressing or knocking out a gene, manipulation of small RNA activity allows us to fine-tune or precisely control the expression of a gene. Such changes in gene expression can be more physiologically relevant and may overcome side effects induced by all-or-nothing approaches. Thus, this offers a great new strategy to improve agronomic traits in crops.

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21 Open questions in the study of RNA-directed DNA methylation

22 (Written by Craig S. Pikaard)

23 Eukaryotic cells protect themselves against TEs, viruses and other selfish genetic elements using RNAi pathways dependent on siRNAs. In plants, siRNAs range in size 24 from 21-24 nt and mediate both post-transcriptional gene silencing (PTGS) and 25 transcriptional gene silencing. RdDM is an important aspect of transcriptional gene 26 27 silencing, involving siRNAs to bring about cytosine methylation of complementary DNA sequences (Erdmann and Picard, 2020). Chemical modifications of histone proteins 28 29 also occur, in crosstalk with DNA methylation (Law and Jacobsen, 2010). Collectively, DNA and histone modifications result in chromatin environments that suppress 30 31 promoter-dependent gene activation, but exactly how is not clear.

32 Most of what we know about RNA-dependent silencing in plants comes from studies of Arabidopsis. At least two pathways contribute to RdDM: an initiation pathway that acts 33 on transcriptionally active transposons or invading viruses and a maintenance pathway 34 35 that perpetuates cytosine methylation at thousands of transposon loci throughout the genome (Figure 10). The establishment pathway overlaps with the pathway for PTGS 36 (Nuthikattu et al., 2013) and begins with transposon, virus or transgene transcripts that 37 are somehow recognized as being different from other cellular RNAs (Hung and Slotkin, 38 2021), triggering their conversion into double-stranded RNA (dsRNA) by RNA-39 40 DEPENDENT RNA POLYMERASE 6 (RDR6). The dsRNAs are then cut (diced) into 21-

or 22-nt siRNAs by the Dicer-like endonucleases DCL4 or DCL2 and loaded into an 1 Argonaute family protein, primarily AGO1 or AGO6 (Ariel and Manavella, 2021). siRNA-2 AGO1 complexes can bind complementary target mRNAs to cause their destruction or 3 interfere with their translation, thus achieving PTGS. In parallel, 21-22-nt siRNAs bound 4 5 to AGO6 guide low-level cytosine methylation at complementary DNA sequences in partnership with multisubunit RNA Polymerase V (Pol V) and the DNA 6 methyltransferase DOMAINS REARRANGED METHYLTRANSFERASE 2 (DRM2). 7 Low-level methylation is not sufficient for transcriptional gene silencing but serves as a 8 signal to recruit the machinery of the maintenance pathway, which accounts for the vast 9 majority of RdDM activity (Figure 10). This pathway involves RNA Polymerase IV (Pol 10 IV), RNA-DEPENDENT RNA POLYMERASE 2 (RDR2), DCL3, 24-nt siRNAs, AGO4, 11 Pol V, DRM2 and numerous helper activities implicated in Pol IV or Pol V recruitment or 12 chromatin modification and is dependent on 24-nt siRNAs (Figure 10). 13

The biogenesis of 24-nt siRNAs is understood in some detail, having been recapitulated 14 in vitro (Singh et al., 2019) using purified enzymes whose structures have recently been 15 resolved (Fukudome et al., 2021; Huang et al., 2021a; Wang et al., 2021a), yet 16 questions still remain. Pol IV acts first in the pathway, presumably initiating RNA 17 biosynthesis within the context of a melted DNA transcription bubble, as is the case for 18 other DNA-dependent RNA polymerases. However, Pol IV is unable to sustain 19 transcriptional elongation over more than ~12-16 nt into the double-stranded DNA 20 beyond the initiation bubble (Singh et al., 2019), for reasons that are not yet clear. This 21 behavior causes the polymerase to stall and then retreat, sliding backward along the 22 23 DNA template as the template and non-template strands reanneal (Fukudome et al., 2021; Huang et al., 2021a), a phenomenon known as polymerase backtracking. As Pol 24 IV backtracks, the 3' end of its short (~30 nt) transcript becomes unpaired from the 25 template DNA strand and is extruded and becomes engaged by RDR2 (Huang et al., 26 27 2021a), which uses the RNA as a template and initiates transcription 1-2 nt internal to its 3' end (Fukudome et al., 2021). Whether the physical interactions of RDR2 with 28 specific Pol IV subunits stimulates Pol IV backtracking and disfavors Pol IV elongation 29 remains unclear, but is testable. Upon completing transcription of the Pol IV strand to 30 31 generate a dsRNA, RDR2 has an intrinsic terminal transferase activity that adds an extra untemplated nucleotide to the 3' end of its transcript, and then RDR2 releases the 32 resulting dsRNA (Singh et al., 2019). Due to initiation by RDR2 internal to the 3' end of 33 the Pol IV transcript and its addition of an untemplated nucleotide to the 3' end of its 34 35 transcript, the resulting dsRNA has 3' overhangs of 1-2 nt at each end. These overhangs, together with 5' nucleotide preferences, program alternative DCL3 dicing 36 reactions from either end of the dsRNAs, yielding siRNA duplexes that consist of a 24-nt 37 strand paired with a 23-nt strand or a pair of 24-nt strands (Loffer et al., 2022) (Figure 38 10). In the case of 24/23 duplexes, the 23-nt RNAs serve as so-called passenger 39 40 strands that help specify that the paired 24-nt strands are loaded into AGO4 to serve as

guide strands (Wang et al., 2022). The passenger strand is then sliced by AGO and 1 partially released. It is not clear how, or why, 24-nt siRNAs are specifically loaded as 2 guide strands given that 21-, 22-, 23- or 24-nt RNAs can be loaded into recombinant 3 AGO4 and guide slicing of target RNAs with similar efficiency (Wang et al., 2022). One 4 5 speculation is that a dsRNA-binding chaperone activity that can discriminate between 3' overhangs of 1 or 2 nt orients the siRNA duplex such that the strand with the 2-nt 6 overhang is loaded into AGO4 as the guide strand. In the case of asymmetric 24/23 7 duplexes, the 24-nt strand would be oriented to become the guide whereas for 8 symmetrical 24/24 duplexes, with 2-nt overhangs at each end, guide strand choice 9 would presumably be random. Experiments are needed to test this hypothesis. 10

What happens following AGO4-siRNA loading is not clear. Early studies showed that 11 AGO4 localization at RdDM loci is dependent on Pol V transcription, that AGO4 can be 12 chemically crosslinked to Pol V transcripts (Wierzbicki et al., 2009) and that cvtosine 13 methylation occurs where siRNAs overlap sites of Pol V occupancy (Wierzbicki et al., 14 2012). Other studies have revealed that AGO4 can bind the C-terminal domain (CTD) of 15 the Pol V largest subunit and/or the Pol V-associated protein, SPT5L (Suppressor of Ty 16 insertion 5-like) (EI-Shami et al., 2007; Bies-Etheve et al., 2009). Thus, AGO4-RNA and 17 AGO4-protein interactions are both likely to be important, but whether they occur 18 simultaneously or sequentially is unknown. And how does DNA methylation, and/or the 19 histone modifications that correlate with DNA methylation, ensue from these siRNA-20 AGO4-Pol V interactions? There is co-immunoprecipitation evidence that DRM2 and 21 AGO4 can directly interact (Zhong et al., 2014), but RdDM has not yet been achieved in 22 23 vitro. Biochemical and structural studies that could reveal the spatial positions of the proteins, RNAs and DNA strands when RdDM occurs would be break-through studies 24 for the field. 25

Other major unanswered questions pertain to how Pol IV and Pol V transcription is 26 27 initiated. Bacterial and archaeal multisubunit RNA polymerases, as well as eukaryotic RNA polymerases I, II and III require DNA-binding transcription factors that recruit the 28 polymerase to promoters, melt the DNA in the vicinity of the start site and position the 29 polymerase to initiate transcription of one of the two DNA strands. However, 30 31 conventional transcription factors and promoters have not been implicated in Pol IV or Pol V transcription. Instead, the evidence suggests that pre-existing chromatin 32 modifications serve as recruitment signals, with cytosine methylation in the CG context, 33 requiring MET1 and HISTONE DEACETYLASE 6 (HDA6) (Blevins et al., 2014), methyl 34 cytosine binding by SUPPRESSOR OF VARIEGATION 3-9 HOMOLOG PROTEIN 2/9 35 (SUVH2/9), or binding of methylated histone H3 lysine 9 (H3K9) by SAWADEE 36 HOMEODOMAIN HOMOLOG 1 (SHH1) implicated in Pol IV and/or Pol V recruitment 37 (Figure 10) (Erdmann and Picard, 2020). ATP-dependent DNA translocases are also 38 implicated, including the CLSY protein family in the case of Pol IV and DEFECTIVE IN 39 40 RNA-DIRECTED DNA METHYLATION 1 (DRD1) in the case of Pol V. However, there

is currently no biochemical evidence to suggest how promoter-independent DNA
 melting, polymerase positioning or transcription initiation occurs for Pol IV or Pol V.
 Once again, *in vitro* experiments with purified components will be needed to move from
 knowing the list of proteins involved to knowing what they do and how they work.

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7 **REFERENCES**

Alleman, M., Sidorenko, L., McGinnis, K., Seshadri, V., Dorweiler, J.E., White, J.,
Sikkink, K., and Chandler, V.L. (2006). An RNA-dependent RNA polymerase is
required for paramutation in maize. Nature 442, 295-298.

Anderson, S.J., Kramer, M.C., Gosai, S.J., Yu, X., Vandivier, L.E., Nelson, A.D.L.,

12 Anderson, Z.D., Beilstein, M.A., Fray, R.G., Lyons, E., and Gregory, B.D. (2018).

13 N(6)-Methyladenosine Inhibits Local Ribonucleolytic Cleavage to Stabilize mRNAs in

14 Arabidopsis. Cell Rep **25**, 1146-1157 e1143.

Arae, T., Isai, S., Sakai, A., Mineta, K., Yokota Hirai, M., Suzuki, Y., Kanaya, S.,

16 Yamaguchi, J., Naito, S., and Chiba, Y. (2017). Co-ordinated Regulations of mRNA

17 Synthesis and Decay during Cold Acclimation in Arabidopsis Cells. Plant Cell Physiol

18 **58,** 1090-1102.

Ariel, F., Lucero, L., Christ, A., Mammarella, M.F., Jegu, T., Veluchamy, A.,

20 Mariappan, K., Latrasse, D., Blein, T., Liu, C., Benhamed, M., and Crespi, M.

21 (2020). R-Loop Mediated trans Action of the APOLO Long Noncoding RNA. Mol Cell 77,

22 1055-1065 e1054.

Ariel, F.D., and Manavella, P.A. (2021). When junk DNA turns functional: transposonderived non-coding RNAs in plants. J Exp Bot **72**, 4132-4143.

25 Arribas-Hernandez, L., Bressendorff, S., Hansen, M.H., Poulsen, C., Erdmann, S.,

and Brodersen, P. (2018). An m(6)A-YTH Module Controls Developmental Timing and

27 Morphogenesis in Arabidopsis. Plant Cell **30**, 952-967.

Arribas-Hernandez, L., Rennie, S., Koster, T., Porcelli, C., Lewinski, M., Staiger, D.,

Andersson, R., and Brodersen, P. (2021a). Principles of mRNA targeting via the Arabidopsis m(6)A-binding protein ECT2. Elife **10**.

31 Arribas-Hernandez, L., Rennie, S., Schon, M., Porcelli, C., Enugutti, B.,

Andersson, R., Nodine, M.D., and Brodersen, P. (2021b). The YTHDF proteins ECT2

33 and ECT3 bind largely overlapping target sets and influence target mRNA abundance,

- 34 not alternative polyadenylation. Elife **10**.
- Arsovski, A.A., Galstyan, A., Guseman, J.M., and Nemhauser, J.L. (2012). Photomorphogenesis. Arabidopsis Book **10**, e0147.

37 Bazin, J., Baerenfaller, K., Gosai, S.J., Gregory, B.D., Crespi, M., and Bailey-

38 Serres, J. (2017). Global analysis of ribosome-associated noncoding RNAs unveils new

modes of translational regulation. Proc Natl Acad Sci U S A **114**, E10018-E10027.

Bhat, S.S., Bielewicz, D., Gulanicz, T., Bodi, Z., Yu, X., Anderson, S.J., Szewc, L.,
 Bajczyk, M., Dolata, J., Grzelak, N., Smolinski, D.J., Gregory, B.D., Fray, R.G.,
 Jarmolowski, A., and Szweykowska-Kulinska, Z. (2020). mRNA adenosine
 methylase (MTA) deposits m(6)A on pri-miRNAs to modulate miRNA biogenesis in

- 5 Arabidopsis thaliana. Proc Natl Acad Sci U S A **117**, 21785-21795.
- 6 Bies-Etheve, N., Pontier, D., Lahmy, S., Picart, C., Vega, D., Cooke, R., and
- 7 Lagrange, T. (2009). RNA-directed DNA methylation requires an AGO4-interacting

8 member of the SPT5 elongation factor family. EMBO Rep **10**, 649-654.

- 9 Blevins, T., Pontvianne, F., Cocklin, R., Podicheti, R., Chandrasekhara, C.,
- 10 Yerneni, S., Braun, C., Lee, B., Rusch, D., Mockaitis, K., Tang, H., and Pikaard,
- C.S. (2014). A two-step process for epigenetic inheritance in Arabidopsis. Mol Cell 54, 30-42.
- 13 Boccaletto, P., Machnicka, M.A., Purta, E., Piatkowski, P., Baginski, B., Wirecki,
- 14 T.K., de Crecy-Lagard, V., Ross, R., Limbach, P.A., Kotter, A., Helm, M., and
- 15 Bujnicki, J.M. (2018). MODOMICS: a database of RNA modification pathways. 2017
- update. Nucleic Acids Res **46**, D303-D307.
- Bologna, N.G., Schapire, A.L., and Palatnik, J.F. (2013). Processing of plant
 microRNA precursors. Brief Funct Genomics 12, 37-45.
- 19 Bourc'his, D., and Voinnet, O. (2010). A small-RNA perspective on gametogenesis,
- fertilization, and early zygotic development. Science **330**, 617-622.
- Brickner, J.R., Garzon, J.L., and Cimprich, K.A. (2022). Walking a tightrope: The
 complex balancing act of R-loops in genome stability. Molecular cell.
- Brioudes, F., Jay, F., Sarazin, A., Grentzinger, T., Devers, E.A., and Voinnet, O. (2021). HASTY, the Arabidopsis EXPORTIN5 ortholog, regulates cell-to-cell and vascular microRNA movement. EMBO J **40**, e107455.
- 26 Bryll, A.R., and Peterson, C.L. (2022). Functional interaction between the RNA
- exosome and the sirtuin deacetylase Hst3 maintains transcriptional homeostasis. Genes Dev **36**, 17-22.
- 29 Bugnon, L.A., Edera, A.A., Prochetto, S., Gerard, M., Raad, J., Fenoy, E., Rubiolo,
- 30 M., Chorostecki, U., Gabaldon, T., Ariel, F., Di Persia, L.E., Milone, D.H., and
- 31 Stegmayer, G. (2022). Secondary structure prediction of long noncoding RNA: review
- and experimental comparison of existing approaches. Brief Bioinform **23**.
- 33 Burgess, D., Chow, H.T., Grover, J.W., Freeling, M., and Mosher, R.A. (2022). Ovule
- siRNAs methylate protein-coding genes in trans. Plant Cell **34**, 3647-3664.
- Burjoski, V., and Reddy, A.S.N. (2021). The Landscape of RNA-Protein Interactions in
 Plants: Approaches and Current Status. Int J Mol Sci 22.
- 37 Cai, Q., He, B., and Jin, H. (2019). A safe ride in extracellular vesicles small RNA
- trafficking between plant hosts and pathogens. Curr Opin Plant Biol **52**, 140-148.

- 1 Cai, Q., Qiao, L., Wang, M., He, B., Lin, F.M., Palmquist, J., Huang, S.D., and Jin, H.
- 2 (2018). Plants send small RNAs in extracellular vesicles to fungal pathogen to silence
- 3 virulence genes. Science **360**, 1126-1129.
- 4 Cai, Q., He, B., Wang, S., Fletcher, S., Niu, D., Mitter, N., Birch, P.R.J., and Jin, H.
- 5 (2021). Message in a Bubble: Shuttling Small RNAs and Proteins Between Cells and
- 6 Interacting Organisms Using Extracellular Vesicles. Annu Rev Plant Biol **72**, 497-524.
- 7 Cambiagno, D.A., Giudicatti, A.J., Arce, A.L., Gagliardi, D., Li, L., Yuan, W.,
- 8 Lundberg, D.S., Weigel, D., and Manavella, P.A. (2021). HASTY modulates miRNA
- 9 biogenesis by linking pri-miRNA transcription and processing. Mol Plant **14**, 426-439.
- 10 Campo, S., Sanchez-Sanuy, F., Camargo-Ramirez, R., Gomez-Ariza, J., Baldrich,
- 11 P., Campos-Soriano, L., Soto-Suarez, M., and San Segundo, B. (2021). A novel
- 12 Transposable element-derived microRNA participates in plant immunity to rice blast
- disease. Plant Biotechnol J **19**, 1798-1811.
- 14 Chantarachot, T., and Bailey-Serres, J. (2018). Polysomes, Stress Granules, and
- 15 Processing Bodies: A Dynamic Triumvirate Controlling Cytoplasmic mRNA Fate and
- 16 Function. Plant Physiol **176**, 254-269.
- 17 **Chen, C., Liu, Y., and Xia, R.** (2019). Jack of Many Trades: The Multifaceted Role of 18 miR528 in Monocots. Mol Plant **12**, 1044-1046.
- 19 **Chen, X., and Rechavi, O.** (2022). Plant and animal small RNA communications 20 between cells and organisms. Nat Rev Mol Cell Biol **23**, 185-203.
- Cheng, L., Wang, W., Yao, Y., and Sun, Q. (2021). Mitochondrial RNase H1 activity
 regulates R-loop homeostasis to maintain genome integrity and enable early
 embryogenesis in Arabidopsis. PLoS Biol 19, e3001357.
- 24 Cho, H., Cho, H.S., Nam, H., Jo, H., Yoon, J., Park, C., Dang, T.V.T., Kim, E., Jeong,
- J., Park, S., Wallner, E.S., Youn, H., Park, J., Jeon, J., Ryu, H., Greb, T., Choi, K.,
- Lee, Y., Jang, S.K., Ban, C., and Hwang, I. (2018). Translational control of phloem
- development by RNA G-quadruplex-JULGI determines plant sink strength. Nat Plants 4,
 376-390.
- 29 Chung, B.Y.W., Balcerowicz, M., Di Antonio, M., Jaeger, K.E., Geng, F., Franaszek,
- 30 **K., Marriott, P., Brierley, I., Firth, A.E., and Wigge, P.A.** (2020). An RNA 31 thermoswitch regulates daytime growth in Arabidopsis. Nat Plants **6**, 522-532.
- 32 **Costa, S.** (2016). Cell identity: a matter of lineage and neighbours. New Phytol **210**,
- 33 1155**-**1158.
- 34 Deng, H., Cheema, J., Zhang, H., Woolfenden, H., Norris, M., Liu, Z., Liu, Q., Yang,
- **X., Yang, M., Deng, X., Cao, X., and Ding, Y.** (2018). Rice In Vivo RNA Structurome
- 36 Reveals RNA Secondary Structure Conservation and Divergence in Plants. Mol Plant
- **11,** 607-622.
- 38 Devers, E.A., Brosnan, C.A., Sarazin, A., Albertini, D., Amsler, A.C., Brioudes, F.,
- Jullien, P.E., Lim, P., Schott, G., and Voinnet, O. (2020). Movement and differential

- 1 consumption of short interfering RNA duplexes underlie mobile RNA interference. Nat
- 2 Plants 6, 789-799.
- 3 Dolata, J., Guo, Y., Kolowerzo, A., Smolinski, D., Brzyzek, G., Jarmolowski, A., and
- 4 **Swiezewski, S.** (2015). NTR1 is required for transcription elongation checkpoints at alternative exons in Arabidopsis. EMBO J **34**, 544-558.
- 6 Duan, H.C., Wei, L.H., Zhang, C., Wang, Y., Chen, L., Lu, Z., Chen, P.R., He, C., and
- 7 Jia, G. (2017). ALKBH10B Is an RNA N(6)-Methyladenosine Demethylase Affecting
- 8 Arabidopsis Floral Transition. Plant Cell **29**, 2995-3011.
- Duncan, S., Olsson, T.S.G., Hartley, M., Dean, C., and Rosa, S. (2017). Single
 Molecule RNA FISH in Arabidopsis Root Cells. Bio Protoc 7, e2240.
- 11 Dunker, F., Trutzenberg, A., Rothenpieler, J.S., Kuhn, S., Prols, R., Schreiber, T.,
- 12 Tissier, A., Kemen, A., Kemen, E., Huckelhoven, R., and Weiberg, A. (2020).
- 13 Oomycete small RNAs bind to the plant RNA-induced silencing complex for virulence.
- 14 Elife **9**.
- 15 El-Shami, M., Pontier, D., Lahmy, S., Braun, L., Picart, C., Vega, D., Hakimi, M.A.,
- Jacobsen, S.E., Cooke, R., and Lagrange, T. (2007). Reiterated WG/GW motifs form
- 17 functionally and evolutionarily conserved ARGONAUTE-binding platforms in RNAi-
- related components. Genes Dev **21**, 2539-2544.
- Emenecker, R.J., Holehouse, A.S., and Strader, L.C. (2020). Emerging Roles for
 Phase Separation in Plants. Dev Cell 55, 69-83.
- Erdmann, R.M., and Picard, C.L. (2020). RNA-directed DNA Methylation. PLoS Genet
 16, e1009034.
- Fan, L., Zhang, C., Gao, B., Zhang, Y., Stewart, E., Jez, J., Nakajima, K., and Chen,
- 24 X. (2022). Microtubules promote the non-cell autonomous action of microRNAs by
- inhibiting their cytoplasmic loading onto ARGONAUTE1 in Arabidopsis. Dev Cell 57,
 995-1008 e1005.
- Fan, Y., Yang, J., Mathioni, S.M., Yu, J., Shen, J., Yang, X., Wang, L., Zhang, Q.,
- 28 Cai, Z., Xu, C., Li, X., Xiao, J., Meyers, B.C., and Zhang, Q. (2016). PMS1T,
- 29 producing phased small-interfering RNAs, regulates photoperiod-sensitive male sterility
- 30 in rice. Proc Natl Acad Sci U S A **113**, 15144-15149.
- Fang, X., Cui, Y., Li, Y., and Qi, Y. (2015). Transcription and processing of primary microRNAs are coupled by Elongator complex in Arabidopsis. Nat Plants 1, 15075.
- 33 Fang, Y., and Spector, D.L. (2007). Identification of nuclear dicing bodies containing
- proteins for microRNA biogenesis in living Arabidopsis plants. Curr Biol **17**, 818-823.
- Faucillion, M.L., Johansson, A.M., and Larsson, J. (2022). Modulation of RNA
- 36 stability regulates gene expression in two opposite ways: through buffering of RNA
- 37 levels upon global perturbations and by supporting adapted differential expression.
- 38 Nucleic Acids Res **50**, 4372-4388.
- Fei, Y., Nyiko, T., and Molnar, A. (2021). Non-perfectly matching small RNAs can induce stable and heritable epigenetic modifications and can be used as molecular

markers to trace the origin and fate of silencing RNAs. Nucleic Acids Res 49, 19001913.

Feng, X., Zilberman, D., and Dickinson, H. (2013). A conversation across
 generations: soma-germ cell crosstalk in plants. Dev Cell 24, 215-225.

5 Foley, S.W., Gosai, S.J., Wang, D., Selamoglu, N., Sollitti, A.C., Koster, T., Steffen,

6 A., Lyons, E., Daldal, F., Garcia, B.A., Staiger, D., Deal, R.B., and Gregory, B.D.

7 (2017). A Global View of RNA-Protein Interactions Identifies Post-transcriptional

8 Regulators of Root Hair Cell Fate. Dev Cell **41**, 204-220 e205.

- 9 Fonouni-Farde, C., Ariel, F., and Crespi, M. (2021). Plant Long Noncoding RNAs:
- 10 New Players in the Field of Post-Transcriptional Regulations. Noncoding RNA 7.
- 11 Franco-Zorrilla, J.M., Valli, A., Todesco, M., Mateos, I., Puga, M.I., Rubio-Somoza,
- 12 I., Leyva, A., Weigel, D., Garcia, J.A., and Paz-Ares, J. (2007). Target mimicry
- provides a new mechanism for regulation of microRNA activity. Nat Genet **39**, 1033-1037.
- 15 Fukudome, A., Singh, J., Mishra, V., Reddem, E., Martinez-Marquez, F., Wenzel, S.,

16 Yan, R., Shiozaki, M., Yu, Z., Wang, J.C., Takagi, Y., and Pikaard, C.S. (2021).

- 17 Structure and RNA template requirements of Arabidopsis RNA-DEPENDENT RNA
- 18 POLYMERASE 2. Proc Natl Acad Sci U S A **118**.
- Garcia-Muse, T., and Aguilera, A. (2019). R Loops: From Physiological to Pathological
 Roles. Cell 179, 604-618.
- 21 Gawronski, P., Enroth, C., Kindgren, P., Marquardt, S., Karpinski, S., Leister, D.,

Jensen, P.E., Vinther, J., and Scharff, L.B. (2021). Light-Dependent Translation
 Change of Arabidopsis psbA Correlates with RNA Structure Alterations at the
 Translation Initiation Region. Cells 10.

25 Giacomelli, J.I., Weigel, D., Chan, R.L., and Manavella, P.A. (2012). Role of recently

- 26 evolved miRNA regulation of sunflower HaWRKY6 in response to temperature damage.
- 27 New Phytol **195**, 766-773.
- 28 Godoy Herz, M.A., Kubaczka, M.G., Brzyzek, G., Servi, L., Krzyszton, M., Simpson,
- 29 C., Brown, J., Swiezewski, S., Petrillo, E., and Kornblihtt, A.R. (2019). Light
- 30 Regulates Plant Alternative Splicing through the Control of Transcriptional Elongation.
- 31 Mol Cell **73**, 1066-1074 e1063.
- 32 Gonzalo, L., Tossolini, I., Gulanicz, T., Cambiagno, D.A., Kasprowicz-Maluski, A.,

33 Smolinski, D.J., Mammarella, M.F., Ariel, F.D., Marquardt, S., Szweykowska-

34 Kulinska, Z., Jarmolowski, A., and Manavella, P.A. (2022). R-loops at microRNA

- encoding loci promote co-transcriptional processing of pri-miRNAs in plants. Nat Plants
 8, 402-418.
- 37 Gu, H., Lian, B., Yuan, Y., Kong, C., Li, Y., Liu, C., and Qi, Y. (2022). A 5' tRNA-Ala-
- derived small RNA regulates anti-fungal defense in plants. Sci China Life Sci 65, 1-15.
- 39 Gutierrez-Beltran, E., Elander, P.H., Dalman, K., Dayhoff, G.W., 2nd, Moschou,
- 40 P.N., Uversky, V.N., Crespo, J.L., and Bozhkov, P.V. (2021). Tudor staphylococcal

- nuclease is a docking platform for stress granule components and is essential for 1 SnRK1 activation in Arabidopsis. EMBO J 40, e105043. 2
- Haimovich, G., Medina, D.A., Causse, S.Z., Garber, M., Millan-Zambrano, G., 3
- Barkai, O., Chavez, S., Perez-Ortin, J.E., Darzacq, X., and Choder, M. (2013). Gene 4
- 5 expression is circular: factors for mRNA degradation also foster mRNA synthesis. Cell
- 6 **153,** 1000-1011.
- Hamada, T., Yako, M., Minegishi, M., Sato, M., Kamei, Y., Yanagawa, Y., Toyooka, 7
- K., Watanabe, Y., and Hara-Nishimura, I. (2018). Stress granule formation is induced 8
- by a threshold temperature rather than a temperature difference in Arabidopsis. J Cell 9 Sci 131. 10
- Hartenian, E., and Glaunsinger, B.A. (2019). Feedback to the central dogma: 11
- cytoplasmic mRNA decay and transcription are interdependent processes. Crit Rev 12
- Biochem Mol Biol 54, 385-398. 13
- 14 Hawkes, E.J., Hennelly, S.P., Novikova, I.V., Irwin, J.A., Dean, C., and
- Sanbonmatsu, K.Y. (2016). COOLAIR Antisense RNAs Form Evolutionarily Conserved 15
- 16 Elaborate Secondary Structures. Cell Rep 16, 3087-3096.
- He, B., Cai, Q., Qiao, L., Huang, C.Y., Wang, S., Miao, W., Ha, T., Wang, Y., and Jin, 17
- H. (2021). RNA-binding proteins contribute to small RNA loading in plant extracellular 18 vesicles. Nat Plants 7, 342-352. 19
- He, S.B., Vickers, M., Zhang, J.Y., and Feng, X.Q. (2019). Natural depletion of histone 20
- H1 in sex cells causes DNA demethylation, heterochromatin decondensation and 21 transposon activation. Elife 8, e42530. 22
- Hou, N., Li, C., He, J., Liu, Y., Yu, S., Malnoy, M., Mobeen Tahir, M., Xu, L., Ma, F., 23 and Guan, Q. (2022). MdMTA-mediated m(6) A modification enhances drought 24 tolerance by promoting mRNA stability and translation efficiency of genes involved in 25
- lignin deposition and oxidative stress. New Phytol 234, 1294-1314. 26
- Hou, Y., Sun, J., Wu, B., Gao, Y., Nie, H., Nie, Z., Quan, S., Wang, Y., Cao, X., and 27
- Li, S. (2021). CPSF30-L-mediated recognition of mRNA m(6)A modification controls 28
- 29 alternative polyadenylation of nitrate signaling-related gene transcripts in Arabidopsis.
- Mol Plant 14, 688-699. 30
- 31 Hu, J., Manduzio, S., and Kang, H. (2019). Epitranscriptomic RNA Methylation in Plant
- Development and Abiotic Stress Responses. Front Plant Sci 10, 500. 32
- Hu, J., Cai, J., Umme, A., Chen, Y., Xu, T., and Kang, H. (2022). Unique features of 33
- mRNA m6A methylomes during expansion of tomato (Solanum lycopersicum) fruits. 34 35 Plant Physiol 188, 2215-2227.
- Hu, J., Cai, J., Park, S.J., Lee, K., Li, Y., Chen, Y., Yun, J.Y., Xu, T., and Kang, H. 36
- (2021). N(6) -Methyladenosine mRNA methylation is important for salt stress tolerance 37 in Arabidopsis. Plant J 106, 1759-1775. 38
- Huang, C.Y., Wang, H., Hu, P., Hamby, R., and Jin, H. (2019). Small RNAs Big 39
- 40 Players in Plant-Microbe Interactions. Cell Host Microbe 26, 173-182.

- 1 Huang, K., Wu, X.X., Fang, C.L., Xu, Z.G., Zhang, H.W., Gao, J., Zhou, C.M., You,
- 2 L.L., Gu, Z.X., Mu, W.H., Feng, Y., Wang, J.W., and Zhang, Y. (2021a). Pol IV and
- 3 RDR2: A two-RNA-polymerase machine that produces double-stranded RNA. Science
- 4 **374**, 1579-1586.
- 5 Huang, Y., Wang, S., Cai, Q., and Jin, H. (2021b). Effective methods for isolation and
- 6 purification of extracellular vesicles from plants. J Integr Plant Biol **63**, 2020-2030.
- 7 Hung, Y.H., and Slotkin, R.K. (2021). The initiation of RNA interference (RNAi) in
- 8 plants. Curr Opin Plant Biol **61**, 102014.
- 9 Ibarra, C.A., Feng, X., Schoft, V.K., Hsieh, T.F., Uzawa, R., Rodrigues, J.A.,
- 10 Zemach, A., Chumak, N., Machlicova, A., Nishimura, T., Rojas, D., Fischer, R.L.,
- 11 **Tamaru, H., and Zilberman, D.** (2012). Active DNA demethylation in plant companion
- cells reinforces transposon methylation in gametes. Science **337**, 1360-1364.
- 13 Jiang, G., Liu, D., Yin, D., Zhou, Z., Shi, Y., Li, C., Zhu, L., and Zhai, W. (2020a). A
- 14 Rice NBS-ARC Gene Conferring Quantitative Resistance to Bacterial Blight Is
- 15 Regulated by a Pathogen Effector-Inducible miRNA. Mol Plant **13**, 1752-1767.
- Jiang, P., Lian, B., Liu, C., Fu, Z., Shen, Y., Cheng, Z., and Qi, Y. (2020b). 21-nt phasiRNAs direct target mRNA cleavage in rice male germ cells. Nat Commun **11**, 5191.
- Jiao, Y., Wang, Y., Xue, D., Wang, J., Yan, M., Liu, G., Dong, G., Zeng, D., Lu, Z.,
- **Zhu, X., Qian, Q., and Li, J.** (2010). Regulation of OsSPL14 by OsmiR156 defines ideal plant architecture in rice. Nat Genet **42**, 541-544.
- John, S., Olas, J.J., and Mueller-Roeber, B. (2021). Regulation of alternative splicing in response to temperature variation in plants. J Exp Bot **72**, 6150-6163.
- Kindgren, P., Ivanov, M., and Marquardt, S. (2020). Native elongation transcript
 sequencing reveals temperature dependent dynamics of nascent RNAPII transcription
 in Arabidopsis. Nucleic Acids Res 48, 2332-2347.
- Kornblihtt, A.R., Schor, I.E., Allo, M., Dujardin, G., Petrillo, E., and Munoz, M.J.
- (2013). Alternative splicing: a pivotal step between eukaryotic transcription and
 translation. Nat Rev Mol Cell Biol 14, 153-165.
- 30 Kosmacz, M., Gorka, M., Schmidt, S., Luzarowski, M., Moreno, J.C., Szlachetko, J.,
- Leniak, E., Sokolowska, E.M., Sofroni, K., Schnittger, A., and Skirycz, A. (2019).
- Protein and metabolite composition of Arabidopsis stress granules. New Phytol **222**, 1420-1433.
- Kramer, M.C., Janssen, K.A., Palos, K., Nelson, A.D.L., Vandivier, L.E., Garcia,
 B.A., Lyons, E., Beilstein, M.A., and Gregory, B.D. (2020). N(6)-methyladenosine and
 RNA secondary structure affect transcript stability and protein abundance during
- 37 systemic salt stress in Arabidopsis. Plant Direct **4**, e00239.
- 38 Kwok, C.K., Ding, Y., Tang, Y., Assmann, S.M., and Bevilacqua, P.C. (2013).
- Determination of in vivo RNA structure in low-abundance transcripts. Nat Commun **4**, 2971.

- Labno, A., Tomecki, R., and Dziembowski, A. (2016). Cytoplasmic RNA decay
 pathways Enzymes and mechanisms. Biochim Biophys Acta 1863, 3125-3147.
- 3 Law, J.A., and Jacobsen, S.E. (2010). Establishing, maintaining and modifying DNA
- 4 methylation patterns in plants and animals. Nat Rev Genet **11**, 204-220.
- 5 Leng, X., Ivanov, M., Kindgren, P., Malik, I., Thieffry, A., Brodersen, P., Sandelin,
- 6 A., Kaplan, C.D., and Marquardt, S. (2020). Organismal benefits of transcription speed
- 7 control at gene boundaries. EMBO Rep **21**, e49315.
- 8 Li, J., Chen, Z., Chen, F., Xie, G., Ling, Y., Peng, Y., Lin, Y., Luo, N., Chiang, C.M.,
- 9 and Wang, H. (2020a). Targeted mRNA demethylation using an engineered dCas13b-
- 10 ALKBH5 fusion protein. Nucleic Acids Res **48**, 5684-5694.
- Li, L., Wu, W., Zhao, Y., and Zheng, B. (2017). A reciprocal inhibition between ARID1
- and MET1 in male and female gametes in Arabidopsis. J Integr Plant Biol **59**, 657-668.
- Li, Y., Song, Y., Xu, W., Li, Q., Wang, X., Li, K., Wang, J., Liu, Z., Velychko, S., Ye,
- R., Xia, Q., Wang, L., Guo, R., Dong, X., Zheng, Z., Dai, Y., Li, H., Yao, M., Xue, Y.,
- 15 Scholer, H.R., Sun, Q., and Yao, H. (2020b). R-loops coordinate with SOX2 in
- regulating reprogramming to pluripotency. Sci Adv **6**, eaba0777.
- Liu, G., Kang, G., Wang, S., Huang, Y., and Cai, Q. (2021). Extracellular Vesicles:
 Emerging Players in Plant Defense Against Pathogens. Front Plant Sci 12, 757925.
- 19 Liu, K., and Sun, Q. (2021). Intragenic tRNA-promoted R-loops orchestrate
- transcription interference for plant oxidative stress responses. Plant Cell **33**, 3574-3591.
- Liu, L., and Chen, X. (2018). Intercellular and systemic trafficking of RNAs in plants.
- 22 Nat Plants **4**, 869-878.
- Liu, X.M., Zhou, J., Mao, Y., Ji, Q., and Qian, S.B. (2019). Programmable RNA N(6)methyladenosine editing by CRISPR-Cas9 conjugates. Nat Chem Biol **15**, 865-871.
- Liu, Y., Teng, C., Xia, R., and Meyers, B.C. (2020). PhasiRNAs in Plants: Their Biogenesis, Genic Sources, and Roles in Stress Responses, Development, and Reproduction. Plant Cell **32**, 3059-3080.
- Loffer, A., Singh, J., Fukudome, A., Mishra, V., Wang, F., and Pikaard, C.S. (2022).
- A DCL3 dicing code within Pol IV-RDR2 transcripts diversifies the siRNA pool guiding
 RNA-directed DNA methylation. Elife **11**.
- Long, J., Walker, J., She, W., Aldridge, B., Gao, H., Deans, S., Vickers, M., and Feng, X. (2021). Nurse cell--derived small RNAs define paternal epigenetic inheritance
- in Arabidopsis. Science **373**.
- Lucero, L., Ferrero, L., Fonouni-Farde, C., and Ariel, F. (2021). Functional classification of plant long noncoding RNAs: a transcript is known by the company it keeps. New Phytol **229**, 1251-1260.
- Luo, G.Z., MacQueen, A., Zheng, G., Duan, H., Dore, L.C., Lu, Z., Liu, J., Chen, K.,
- Jia, G., Bergelson, J., and He, C. (2014). Unique features of the m6A methylome in
- Arabidopsis thaliana. Nat Commun **5**, 5630.

- 1 Ma, X., Liu, C., Kong, X., Liu, J., Zhang, S., Liang, S., Luan, W., and Cao, X. (2021).
- 2 Extensive profiling of the expressions of tRNAs and tRNA-derived fragments (tRFs)
- reveals the complexities of tRNA and tRF populations in plants. Sci China Life Sci 64,
 495-511.
- 5 **Machyna, M., and Simon, M.D.** (2018). Catching RNAs on chromatin using 6 hybridization capture methods. Brief Funct Genomics **17**, 96-103.
- 7 Manavella, P.A., Yang, S.W., and Palatnik, J. (2019). Keep calm and carry on: miRNA
- 8 biogenesis under stress. Plant J **99**, 832-843.
- Manduzio, S., and Kang, H. (2021). RNA methylation in chloroplasts or mitochondria in
 plants. RNA Biol 18, 2127-2135.
- 11 Marondedze, C., Thomas, L., Lilley, K.S., and Gehring, C. (2019). Drought Stress
- 12 Causes Specific Changes to the Spliceosome and Stress Granule Components. Front
- 13 Mol Biosci **6**, 163.
- Martinez-Perez, M., Aparicio, F., Lopez-Gresa, M.P., Belles, J.M., Sanchez-Navarro,
- **J.A., and Pallas, V.** (2017). Arabidopsis m(6)A demethylase activity modulates viral
- 16 infection of a plant virus and the m(6)A abundance in its genomic RNAs. Proc Natl Acad
- 17 Sci U S A **114**, 10755-10760.
- 18 Maruri-Lopez, I., Figueroa, N.E., Hernandez-Sanchez, I.E., and Chodasiewicz, M.
- 19 (2021). Plant Stress Granules: Trends and Beyond. Front Plant Sci **12**, 722643.
- Mathieu, M., Martin-Jaular, L., Lavieu, G., and Thery, C. (2019). Specificities of secretion and uptake of exosomes and other extracellular vesicles for cell-to-cell communication. Nat Cell Biol **21**, 9-17.
- Matzke, M.A., and Mosher, R.A. (2014). RNA-directed DNA methylation: an epigenetic
 pathway of increasing complexity. Nat Rev Genet 15, 394-408.
- Mauer, J., Luo, X., Blanjoie, A., Jiao, X., Grozhik, A.V., Patil, D.P., Linder, B.,
- Pickering, B.F., Vasseur, J.J., Chen, Q., Gross, S.S., Elemento, O., Debart, F.,
- Kiledjian, M., and Jaffrey, S.R. (2017). Reversible methylation of m(6)Am in the 5' cap
 controls mRNA stability. Nature 541, 371-375.
- 29 Mencia, R., Gonzalo, L., Tossolini, I., and Manavella, P.A. (2022). Keeping up with
- the miRNAs: current paradigms of the biogenesis pathway. J Exp Bot.
- 31 Merchante, C., Stepanova, A.N., and Alonso, J.M. (2017). Translation regulation in
- plants: an interesting past, an exciting present and a promising future. Plant J 90, 628 653.
- 34 Miller, H.E., Ilieva, M., Bishop, A.J.R., and Uchida, S. (2022). Current Status of
- 35 Epitranscriptomic Marks Affecting IncRNA Structures and Functions. Noncoding RNA 8.
- Mingardi, J., Musazzi, L., De Petro, G., and Barbon, A. (2018). miRNA Editing: New
- Insights into the Fast Control of Gene Expression in Health and Disease. Mol Neurobiol
- 38 **55,** 7717-7727.
- 39 Moison, M., Pacheco, J.M., Lucero, L., Fonouni-Farde, C., Rodriguez-Melo, J.,
- 40 Mansilla, N., Christ, A., Bazin, J., Benhamed, M., Ibanez, F., Crespi, M., Estevez,

J.M., and Ariel, F. (2021). The IncRNA APOLO interacts with the transcription factor WRKY42 to trigger root hair cell expansion in response to cold. Mol Plant **14**, 937-948.

3 Moore, J.W., Loake, G.J., and Spoel, S.H. (2011). Transcription dynamics in plant

4 immunity. Plant Cell **23**, 2809-2820.

5 Nguyen, C.C., Nakaminami, K., Matsui, A., Kobayashi, S., Kurihara, Y., Toyooka,

6 K., Tanaka, M., and Seki, M. (2016). Oligouridylate Binding Protein 1b Plays an

7 Integral Role in Plant Heat Stress Tolerance. Front Plant Sci **7**, 853.

8 Nuthikattu, S., McCue, A.D., Panda, K., Fultz, D., DeFraia, C., Thomas, E.N., and

9 Slotkin, R.K. (2013). The initiation of epigenetic silencing of active transposable

- 10 elements is triggered by RDR6 and 21-22 nucleotide small interfering RNAs. Plant
- 11 Physiol **162**, 116-131.
- 12 Olmedo-Monfil, V., Duran-Figueroa, N., Arteaga-Vazquez, M., Demesa-Arevalo, E.,
- Autran, D., Grimanelli, D., Slotkin, R.K., Martienssen, R.A., and Vielle-Calzada, J.P.
- 14 (2010). Control of female gamete formation by a small RNA pathway in Arabidopsis.
- 15 Nature **464**, 628-632.
- 16 Ozata, D.M., Gainetdinov, I., Zoch, A., O'Carroll, D., and Zamore, P.D. (2019). PIWI-
- interacting RNAs: small RNAs with big functions. Nat Rev Genet **20**, 89-108.

18 **Perrella, G., Baurle, I., and van Zanten, M.** (2022). Epigenetic regulation of 19 thermomorphogenesis and heat stress tolerance. New Phytol **234**, 1144-1160.

- Petermann, E., Lan, L., and Zou, L. (2022). Sources, resolution and physiological relevance of R-loops and RNA-DNA hybrids. Nat Rev Mol Cell Biol.
- 22 Petrillo, E., Godoy Herz, M.A., Fuchs, A., Reifer, D., Fuller, J., Yanovsky, M.J.,

23 Simpson, C., Brown, J.W., Barta, A., Kalyna, M., and Kornblihtt, A.R. (2014). A

- chloroplast retrograde signal regulates nuclear alternative splicing. Science **344**, 427430.
- Pikaard, C.S., Haag, J.R., Pontes, O.M., Blevins, T., and Cocklin, R. (2012). A
- transcription fork model for Pol IV and Pol V-dependent RNA-directed DNA methylation.
 Cold Spring Harb Symp Quant Biol 77, 205-212.
- 29 Re, D.A., Lang, P.L.M., Yones, C., Arce, A.L., Stegmayer, G., Milone, D., and
- 30 **Manavella, P.A.** (2019). Alternative use of miRNA-biogenesis co-factors in plants at low
- 31 temperatures. Development **146**.
- 32 Reichel, M., Koster, T., and Staiger, D. (2019). Marking RNA: m6A writers, readers,
- and functions in Arabidopsis. J Mol Cell Biol **11**, 899-910.
- Reis, R.S., Deforges, J., Schmidt, R.R., Schippers, J.H.M., and Poirier, Y. (2021).
- An antisense noncoding RNA enhances translation via localized structural rearrangements of its cognate mRNA. Plant Cell **33**, 1381-1397.
- 37 Ren, B., Wang, X., Duan, J., and Ma, J. (2019). Rhizobial tRNA-derived small RNAs
- are signal molecules regulating plant nodulation. Science **365**, 919-922.
- 39 Riegler, S., Servi, L., Scarpin, M.R., Godoy Herz, M.A., Kubaczka, M.G., Venhuizen,
- 40 P., Meyer, C., Brunkard, J.O., Kalyna, M., Barta, A., and Petrillo, E. (2021). Light

- regulates alternative splicing outcomes via the TOR kinase pathway. Cell Rep 36,
 109676.
- 3 Ries, R.J., Zaccara, S., Klein, P., Olarerin-George, A., Namkoong, S., Pickering,
- B.F., Patil, D.P., Kwak, H., Lee, J.H., and Jaffrey, S.R. (2019). m(6)A enhances the
 phase separation potential of mRNA. Nature 571, 424-428.
- 6 Rigo, R., Bazin, J., Romero-Barrios, N., Moison, M., Lucero, L., Christ, A.,
- 7 Benhamed, M., Blein, T., Huguet, S., Charon, C., Crespi, M., and Ariel, F. (2020).
- 8 The Arabidopsis IncRNA ASCO modulates the transcriptome through interaction with
- 9 splicing factors. EMBO Rep 21, e48977.
- 10 Ruzicka, K., Zhang, M., Campilho, A., Bodi, Z., Kashif, M., Saleh, M., Eeckhout, D.,
- 11 El-Showk, S., Li, H., Zhong, S., De Jaeger, G., Mongan, N.P., Hejatko, J., Helariutta,
- 12 Y., and Fray, R.G. (2017). Identification of factors required for m(6) A mRNA
- 13 methylation in Arabidopsis reveals a role for the conserved E3 ubiquitin ligase HAKAI.
- 14 New Phytol **215**, 157-172.
- Schor, I.E., Fiszbein, A., Petrillo, E., and Kornblihtt, A.R. (2013). Intragenic
 epigenetic changes modulate NCAM alternative splicing in neuronal differentiation.
 EMBO J 32, 2264-2274.
- 18 Schor, I.E., Rascovan, N., Pelisch, F., Allo, M., and Kornblihtt, A.R. (2009).
- 19 Neuronal cell depolarization induces intragenic chromatin modifications affecting NCAM
- 20 alternative splicing. Proc Natl Acad Sci U S A **106**, 4325-4330.
- Scutenaire, J., Deragon, J.M., Jean, V., Benhamed, M., Raynaud, C., Favory, J.J.,
- Merret, R., and Bousquet-Antonelli, C. (2018). The YTH Domain Protein ECT2 Is an m(6)A Reader Required for Normal Trichome Branching in Arabidopsis. Plant Cell **30**,
- 24 986-1005.
- 25 Shahid, S., Kim, G., Johnson, N.R., Wafula, E., Wang, F., Coruh, C., Bernal-
- Galeano, V., Phifer, T., dePamphilis, C.W., Westwood, J.H., and Axtell, M.J. (2018).
- MicroRNAs from the parasitic plant Cuscuta campestris target host messenger RNAs.
 Nature 553, 82-85.
- 29 Singh, J., Mishra, V., Wang, F., Huang, H.Y., and Pikaard, C.S. (2019). Reaction
- 30 Mechanisms of Pol IV, RDR2, and DCL3 Drive RNA Channeling in the siRNA-Directed
- 31 DNA Methylation Pathway. Mol Cell **75**, 576-589 e575.
- 32 Slotkin, R.K., Vaughn, M., Borges, F., Tanurdzic, M., Becker, J.D., Feijo, J.A., and
- Martienssen, R.A. (2009). Epigenetic reprogramming and small RNA silencing of transposable elements in pollen. Cell **136**, 461-472.
- 35 Song, P., Yang, J., Wang, C., Lu, Q., Shi, L., Tayier, S., and Jia, G. (2021).
- 36 Arabidopsis N(6)-methyladenosine reader CPSF30-L recognizes FUE signals to control
- polyadenylation site choice in liquid-like nuclear bodies. Mol Plant **14**, 571-587.
- 38 Sorenson, R., and Bailey-Serres, J. (2014). Selective mRNA sequestration by
- 39 OLIGOURIDYLATE-BINDING PROTEIN 1 contributes to translational control during
- 40 hypoxia in Arabidopsis. Proc Natl Acad Sci U S A **111**, 2373-2378.

1 Sorenson, R.S., Deshotel, M.J., Johnson, K., Adler, F.R., and Sieburth, L.E. (2018).

- 2 Arabidopsis mRNA decay landscape arises from specialized RNA decay substrates,
- decapping-mediated feedback, and redundancy. Proc Natl Acad Sci U S A 115, E1485 E1494.
- 5 Stepien, A., Knop, K., Dolata, J., Taube, M., Bajczyk, M., Barciszewska-Pacak, M.,
- 6 Pacak, A., Jarmolowski, A., and Szweykowska-Kulinska, Z. (2017).
- Posttranscriptional coordination of splicing and miRNA biogenesis in plants. Wiley
 Interdiscip Rev RNA 8.
- 9 Su, Z., Tang, Y., Ritchey, L.E., Tack, D.C., Zhu, M., Bevilacqua, P.C., and Assmann,
- 10 S.M. (2018). Genome-wide RNA structurome reprogramming by acute heat shock
- 11 globally regulates mRNA abundance. Proc Natl Acad Sci U S A **115**, 12170-12175.
- 12 Su, Z., Wang, N., Hou, Z., Li, B., Li, D., Liu, Y., Cai, H., Qin, Y., and Chen, X. (2020).
- 13 Regulation of Female Germline Specification via Small RNA Mobility in Arabidopsis.
- 14 Plant Cell **32**, 2842-2854.
- 15 Su, Z., Zhao, L., Zhao, Y., Li, S., Won, S., Cai, H., Wang, L., Li, Z., Chen, P., Qin, Y.,
- and Chen, X. (2017). The THO Complex Non-Cell-Autonomously Represses Female
 Germline Specification through the TAS3-ARF3 Module. Curr Biol 27, 1597-1609
- 18 e1592.
- 19 Sun, M., Schwalb, B., Pirkl, N., Maier, K.C., Schenk, A., Failmezger, H., Tresch, A.,
- and Cramer, P. (2013a). Global analysis of eukaryotic mRNA degradation reveals
 Xrn1-dependent buffering of transcript levels. Mol Cell 52, 52-62.
- 22 Sun, Q., Csorba, T., Skourti-Stathaki, K., Proudfoot, N.J., and Dean, C. (2013b). R-
- loop stabilization represses antisense transcription at the Arabidopsis FLC locus.
 Science **340**, 619-621.
- 25 Szadeczky-Kardoss, I., Szaker, H.M., Verma, R., Darko, E., Pettko-Szandtner, A.,
- 26 **Silhavy, D., and Csorba, T.** (2022). Elongation factor TFIIS is essential for heat stress 27 adaptation in plants. Nucleic Acids Res **50**, 1927-1950.
- Tack, D.C., Su, Z., Yu, Y., Bevilacqua, P.C., and Assmann, S.M. (2020). Tissue specific changes in the RNA structurome mediate salinity response in Arabidopsis. RNA
 26, 492-511.
- Tian, X., Qin, Z., Zhao, Y., Wen, J., Lan, T., Zhang, L., Wang, F., Qin, D., Yu, K.,
- 32 Zhao, A., Hu, Z., Yao, Y., Ni, Z., Sun, Q., De Smet, I., Peng, H., and Xin, M. (2022).
- 33 Stress granule-associated TaMBF1c confers thermotolerance through regulating
- 34 specific mRNA translation in wheat (Triticum aestivum). New Phytol **233**, 1719-1731.
- **Timmers, H.T.M., and Tora, L.** (2018). Transcript Buffering: A Balancing Act between mRNA Synthesis and mRNA Degradation. Mol Cell **72**, 10-17.
- ³⁷ Tognacca, R.S., Kubaczka, M.G., Servi, L., Rodriguez, F.S., Godoy Herz, M.A., and
- **Petrillo, E.** (2020). Light in the transcription landscape: chromatin, RNA polymerase II
- and splicing throughout Arabidopsis thaliana's life cycle. Transcription **11**, 117-133.

- 1 Tomassi, A.H., Re, D.A., Romani, F., Cambiagno, D.A., Gonzalo, L., Moreno, J.E.,
- 2 Arce, A.L., and Manavella, P.A. (2020). The Intrinsically Disordered Protein CARP9
- 3 Bridges HYL1 to AGO1 in the Nucleus to Promote MicroRNA Activity. Plant Physiol **184**,
- 4 316-329.
- Voinnet, O. (2022). Revisiting small RNA movement in plants. Nat Rev Mol Cell Biol 23,
 163-164.
- 7 Walker, J., Gao, H., Zhang, J., Aldridge, B., Vickers, M., Higgins, J.D., and Feng, X.
- 8 (2018). Sexual-lineage-specific DNA methylation regulates meiosis in Arabidopsis. Nat
 9 Genet 50, 130-137.
- 10 Wang, B., Sun, Y., Song, N., Zhao, M., Liu, R., Feng, H., Wang, X., and Kang, Z.
- 11 (2017a). Puccinia striiformis f. sp. tritici microRNA-like RNA 1 (Pst-milR1), an important
- 12 pathogenicity factor of Pst, impairs wheat resistance to Pst by suppressing the wheat
- 13 pathogenesis-related 2 gene. New Phytol **215**, 338-350.
- 14 Wang, F., Huang, H.-Y., Huang, J., Singh, J., and Pikaard, C.S. (2022). Mechanisms
- of AGO4 slicing-enhanced RNA-directed DNA methylation. BioRxiv.
- 16 Wang, G., Jiang, H., Del Toro de Leon, G., Martinez, G., and Kohler, C. (2018a).
- 17 Sequestration of a Transposon-Derived siRNA by a Target Mimic Imprinted Gene
- Induces Postzygotic Reproductive Isolation in Arabidopsis. Dev Cell 46, 696-705 e694.
 Wang, J.W., Czech, B., and Weigel, D. (2009). miR156-regulated SPL transcription
- Wang, J.W., Czech, B., and Weigel, D. (2009). miR156-regulated SPL transcription
 factors define an endogenous flowering pathway in Arabidopsis thaliana. Cell 138, 738749.
- Wang, M., Weiberg, A., Lin, F.M., Thomma, B.P., Huang, H.D., and Jin, H. (2016).
 Bidirectional cross-kingdom RNAi and fungal uptake of external RNAs confer plant
- protection. Nat Plants **2**, 16151.
- 25 Wang, Q., Xue, Y., Zhang, L., Zhong, Z., Feng, S., Wang, C., Xiao, L., Yang, Z.,
- Harris, C.J., Wu, Z., Zhai, J., Yang, M., Li, S., Jacobsen, S.E., and Du, J. (2021a).
- Mechanism of siRNA production by a plant Dicer-RNA complex in dicing-competent conformation. Science **374**, 1152-1157.
- 29 Wang, W., Li, K., Yang, Z., Hou, Q., Zhao, W.W., and Sun, Q. (2021b). RNase H1C
- 30 collaborates with ssDNA binding proteins WHY1/3 and recombinase RecA1 to fulfill the
- 31 DNA damage repair in Arabidopsis chloroplasts. Nucleic Acids Res **49**, 6771-6787.
- Wang, Y., Li, S., Zhao, Y., You, C., Le, B., Gong, Z., Mo, B., Xia, Y., and Chen, X.
- (2019). NAD(+)-capped RNAs are widespread in the Arabidopsis transcriptome and can
 probably be translated. Proc Natl Acad Sci U S A **116**, 12094-12102.
- 35 Wang, Z., Tang, K., Zhang, D., Wan, Y., Wen, Y., Lu, Q., and Wang, L. (2017b). High-
- throughput m6A-seg reveals RNA m6A methylation patterns in the chloroplast and
- mitochondria transcriptomes of Arabidopsis thaliana. PLoS One **12**, e0185612.
- 38 Wang, Z., Butel, N., Santos-Gonzalez, J., Borges, F., Yi, J., Martienssen, R.A.,
- 39 Martinez, G., and Kohler, C. (2020). Polymerase IV Plays a Crucial Role in Pollen
- 40 Development in Capsella. Plant Cell **32**, 950-966.

- 1 Wang, Z., Ma, Z., Castillo-Gonzalez, C., Sun, D., Li, Y., Yu, B., Zhao, B., Li, P., and
- **Zhang, X.** (2018b). SWI2/SNF2 ATPase CHR2 remodels pri-miRNAs via Serrate to
 impede miRNA production. Nature **557**, 516-521.
- 4 Wei, L., Gu, L., Song, X., Cui, X., Lu, Z., Zhou, M., Wang, L., Hu, F., Zhai, J.,
- 5 Meyers, B.C., and Cao, X. (2014). Dicer-like 3 produces transposable element-
- 6 associated 24-nt siRNAs that control agricultural traits in rice. Proc Natl Acad Sci U S A
- 7 **111,** 3877-3882.
- 8 Wei, L.H., Song, P., Wang, Y., Lu, Z., Tang, Q., Yu, Q., Xiao, Y., Zhang, X., Duan,
- 9 H.C., and Jia, G. (2018). The m(6)A Reader ECT2 Controls Trichome Morphology by
- Affecting mRNA Stability in Arabidopsis. Plant Cell 30, 968-985.
 Weiberg, A., Wang, M., Lin, F.M., Zhao, H., Zhang, Z., Kaloshian, I., Huang, H.D.,
- weiberg, A., Wang, W., Lin, F.W., Zhao, H., Zhang, Z., Kaloshian, L., Huang, H.D.,
- and Jin, H. (2013). Fungal small RNAs suppress plant immunity by hijacking host RNA
- 13 interference pathways. Science **342**, 118-123.
- 14 Wendte, J.M., and Pikaard, C.S. (2017). The RNAs of RNA-directed DNA methylation.
- 15 Biochim Biophys Acta Gene Regul Mech **1860**, 140-148.
- Wierzbicki, A.T., Blevins, T., and Swiezewski, S. (2021). Long Noncoding RNAs in
 Plants. Annu Rev Plant Biol 72, 245-271.
- 18 Wierzbicki, A.T., Ream, T.S., Haag, J.R., and Pikaard, C.S. (2009). RNA polymerase
- 19 V transcription guides ARGONAUTE4 to chromatin. Nat Genet **41**, 630-634.
- 20 Wierzbicki, A.T., Cocklin, R., Mayampurath, A., Lister, R., Rowley, M.J., Gregory,
- B.D., Ecker, J.R., Tang, H., and Pikaard, C.S. (2012). Spatial and functional
 relationships among Pol V-associated loci, Pol IV-dependent siRNAs, and cytosine
 methylation in the Arabidopsis epigenome. Genes Dev 26, 1825-1836.
- 24 Wong-Bajracharya, J., Singan, V.R., Monti, R., Plett, K.L., Ng, V., Grigoriev, I.V.,
- 25 Martin, F.M., Anderson, I.C., and Plett, J.M. (2022). The ectomycorrhizal fungus
- Pisolithus microcarpus encodes a microRNA involved in cross-kingdom gene silencing
 during symbiosis. Proc Natl Acad Sci U S A 119.
- 28 Wu, G., Park, M.Y., Conway, S.R., Wang, J.W., Weigel, D., and Poethig, R.S.
- (2009a). The sequential action of miR156 and miR172 regulates developmental timing
 in Arabidopsis, Cell 138, 750-759.
- 31 Wu, L., Zhang, Q., Zhou, H., Ni, F., Wu, X., and Qi, Y. (2009b). Rice MicroRNA 32 effector complexes and targets. Plant Cell **21**, 3421-3435.
- Wu, L., Zhou, H., Zhang, Q., Zhang, J., Ni, F., Liu, C., and Qi, Y. (2010). DNA methylation mediated by a microRNA pathway. Mol Cell **38**, 465-475.
- 35 Xie, D., Chen, M., Niu, J., Wang, L., Li, Y., Fang, X., Li, P., and Qi, Y. (2021). Phase
- 36 separation of SERRATE drives dicing body assembly and promotes miRNA processing
- in Arabidopsis. Nat Cell Biol **23**, 32-39.
- 38 Xu, L., Yuan, K., Yuan, M., Meng, X., Chen, M., Wu, J., Li, J., and Qi, Y. (2020a).
- 39 Regulation of Rice Tillering by RNA-Directed DNA Methylation at Miniature Inverted-
- 40 Repeat Transposable Elements. Mol Plant **13**, 851-863.

- 1 Xu, T., Wu, X., Wong, C.E., Fan, S., Zhang, Y., Zhang, S., Liang, Z., Yu, H., and
- 2 Shen, L. (2022a). FIONA1-Mediated m(6) A Modification Regulates the Floral
- 3 Transition in Arabidopsis. Adv Sci (Weinh) **9**, e2103628.
- 4 Xu, W., Li, K., Li, Q., Li, S., Zhou, J., and Sun, Q. (2022b). Quantitative, Convenient,
- 5 and Efficient Genome-Wide R-Loop Profiling by ssDRIP-Seq in Multiple Organisms.
- 6 Methods Mol Biol **2528**, 445-464.
- Xu, W., Xu, H., Li, K., Fan, Y., Liu, Y., Yang, X., and Sun, Q. (2017). The R-loop is a
 common chromatin feature of the Arabidopsis genome. Nat Plants 3, 704-714.
- 9 Xu, W., Li, K., Li, S., Hou, Q., Zhang, Y., Liu, K., and Sun, Q. (2020b). The R-Loop
- 10 Atlas of Arabidopsis Development and Responses to Environmental Stimuli. Plant Cell
- 11 **32,** 888-903.
- 12 Yan, P., Liu, Z., Song, M., Wu, Z., Xu, W., Li, K., Ji, Q., Wang, S., Liu, X., Yan, K.,
- 13 Esteban, C.R., Ci, W., Belmonte, J.C.I., Xie, W., Ren, J., Zhang, W., Sun, Q., Qu, J.,
- and Liu, G.H. (2020). Genome-wide R-loop Landscapes during Cell Differentiation and
- 15 Reprogramming. Cell Rep **32**, 107870.
- 16 Yang, X., Cheema, J., Zhang, Y., Deng, H., Duncan, S., Umar, M.I., Zhao, J., Liu, Q.,
- 17 Cao, X., Kwok, C.K., and Ding, Y. (2020a). RNA G-quadruplex structures exist and
- 18 function in vivo in plants. Genome Biol **21**, 226.
- 19 Yang, X., Yu, H., Sun, W., Ding, L., Li, J., Cheema, J., Ramirez-Gonzalez, R., Zhao,
- 20 X., Martin, A.C., Lu, F., Liu, B., Uauy, C., Ding, Y., and Zhang, H. (2021). Wheat in
- vivo RNA structure landscape reveals a prevalent role of RNA structure in modulating
- translational subgenome expression asymmetry. Genome Biol **22**, 326.
- Yang, Z., Li, M., and Sun, Q. (2020b). RHON1 Co-transcriptionally Resolves R-Loops
 for Arabidopsis Chloroplast Genome Maintenance. Cell Rep 30, 243-256 e245.
- 25 Yang, Z., Hou, Q., Cheng, L., Xu, W., Hong, Y., Li, S., and Sun, Q. (2017). RNase H1
- 26 Cooperates with DNA Gyrases to Restrict R-Loops and Maintain Genome Integrity in
- 27 Arabidopsis Chloroplasts. Plant Cell **29**, 2478-2497.
- **Zhang, H., and Ding, Y.** (2021). Novel insights into the pervasive role of RNA structure
- in post-transcriptional regulation of gene expression in plants. Biochem Soc Trans 49,1829-1839.
- Zhang, H., Zhong, H., Zhang, S., Shao, X., Ni, M., Cai, Z., Chen, X., and Xia, Y.
- (2019a). NAD tagSeq reveals that NAD(+)-capped RNAs are mostly produced from a
 large number of protein-coding genes in Arabidopsis. Proc Natl Acad Sci U S A 116,
- 34 12072-12077.
- Zhang, M., Ma, X., Wang, C., Li, Q., Meyers, B.C., Springer, N.M., and Walbot, V.
 (2021). CHH DNA methylation increases at 24-PHAS loci depend on 24-nt phased
- 37 small interfering RNAs in maize meiotic anthers. New Phytol **229**, 2984-2997.
- **Zhang, W., Murphy, C., and Sieburth, L.E.** (2010). Conserved RNasell domain protein
- 39 functions in cytoplasmic mRNA decay and suppresses Arabidopsis decapping mutant
- 40 phenotypes. Proc Natl Acad Sci U S A **107**, 15981-15985.

- ¹ Zhang, Y., Yang, M., Duncan, S., Yang, X., Abdelhamid, M.A.S., Huang, L., Zhang,
- 2 H., Benfey, P.N., Waller, Z.A.E., and Ding, Y. (2019b). G-quadruplex structures trigger
- 3 RNA phase separation. Nucleic Acids Res **47**, 11746-11754.
- Zhang, Y.C., Lei, M.Q., Zhou, Y.F., Yang, Y.W., Lian, J.P., Yu, Y., Feng, Y.Z., Zhou,
- 5 K.R., He, R.R., He, H., Zhang, Z., Yang, J.H., and Chen, Y.Q. (2020). Reproductive
- 6 phasiRNAs regulate reprogramming of gene expression and meiotic progression in rice.
- 7 Nat Commun **11**, 6031.
- 8 Zhong, X., Du, J., Hale, C.J., Gallego-Bartolome, J., Feng, S., Vashisht, A.A.,
- 9 Chory, J., Wohlschlegel, J.A., Patel, D.J., and Jacobsen, S.E. (2014). Molecular
- mechanism of action of plant DRM de novo DNA methyltransferases. Cell **157**, 1050 1060.
- **Zhou, J., Zhang, W., and Sun, Q.** (2022a). R-loop: The new genome regulatory
 element in plants. J Integr Plant Biol.
- 14 **Zhou, L., Tian, S., and Qin, G.** (2019). RNA methylomes reveal the m(6)A-mediated
- regulation of DNA demethylase gene SIDML2 in tomato fruit ripening. Genome Biol 20,156.
- 17 Zhou, M., Coruh, C., Xu, G., Martins, L.M., Bourbousse, C., Lambolez, A., and Law,
- **J.A.** (2022b). The CLASSY family controls tissue-specific DNA methylation patterns in Arabidopsis. Nat Commun **13**, 244.
- Zhou, S.X., Zhu, Y., Wang, L.F., Zheng, Y.P., Chen, J.F., Li, T.T., Yang, X.M., Wang,
- H., Li, X.P., Ma, X.C., Zhao, J.Q., Pu, M., Feng, H., Li, Y., Fan, J., Zhang, J.W.,
- Huang, Y.Y., and Wang, W.M. (2020). Osa-miR1873 fine-tunes rice immunity against
- Magnaporthe oryzae and yield traits. J Integr Plant Biol **62**, 1213-1226.
- Z4 Zhou, X., Huang, K., Teng, C., Abdelgawad, A., Batish, M., Meyers, B.C., and
- 25 Walbot, V. (2022c). 24-nt phasiRNAs move from tapetal to meiotic cells in maize
- anthers. New Phytol **235**, 488-501.
- Zhu, J., Li, C., Peng, X., and Zhang, X. (2021). RNA architecture influences plant
 biology. J Exp Bot 72, 4144-4160.
- 29 Zhu, S., Gu, J., Yao, J., Li, Y., Zhang, Z., Xia, W., Wang, Z., Gui, X., Li, L., Li, D.,
- 30 Zhang, H., and Liu, C. (2022). Liquid-liquid phase separation of RBGD2/4 is required
- for heat stress resistance in Arabidopsis. Dev Cell **57**, 583-597 e586.
- 32

1 Figure Legends

Figure 1. Light, sensed by the chloroplast, triggers a retrograde signal that
 regulates alternative splicing in the nucleus.

- 4 In the light, RNA polymerase II (Poll II) elongation rate is fast, resulting in exon skipping.
- 5 Leaf cells produce sugars that act as mobile signals to coordinate alternative splicing
- 6 responses throughout the whole plant, thus reaching root cells.
- 7
- Figure 2. RNA buffering is a flux-mediated regulatory mechanism that maintains
 some mRNAs at a stable abundance.
- 10 A. RNA abundance is influenced by the balance between RNA synthesis and 11 degradation. B. RNA flux describes the turnover rate of an mRNA, RNA buffering occurs 12 when the flux of an RNA shifts, but not its abundance.

13

14 Figure 3. Hypothetical model of stress granule function.

- UBP1b localizes to the nucleus under non-stress conditions. UBP1b-stress granules (SGs) are induced to form in the cytoplasm in response to heat stress. UBP1b-SGs protect target mRNA from degradation during stress. Elucidation of the mechanism of target mRNA recruitment and the timing of the translation of the protected mRNA will provide critical information on the selective translation mechanisms utilized in plants in response to stress.
- 21

Figure 4. RNA structure may pervasively function in plant growth and development.

- 24 RNAs may fold into diverse RNA structures in different cell types and under different 25 environmental conditions. These dynamic and diverse RNA structures facilitate the 26 regulatory specificities of gene expression at post-transcriptional levels.
- 27

Figure 5. R-loops in plant cells.

- Left, different distribution patterns of nuclear R-loops along the gene body in the genomes of Arabidopsis, maize, and rice.
- Right, chloroplast-localized AtRNH1C restricts RNA:DNA hybrid formation to release head-on transcription-replication conflicts (TRCs) and to promote homologous recombination (HR) repair in chloroplasts. Mitochondrion-localized AtRNH1B inhibits homologous recombination at repeats in the mitochondrial genome by suppressing RNA:DNA hybrid formation. In the absence of AtRNH1B, high levels of mitochondrial Rloops stimulate the relocation of AtRNH1C to mitochondria.
- 37

Figure 6. Regulatory roles of N⁶-methyladenosine (m⁶A) writers, erasers, and readers in RNA metabolism, plant development, and stress responses.

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The cellular components responsible for installation, removal, and interpretation of m⁶A marks are methyltransferases ("writers"), demethylases ("erasers"), and RNA-binding proteins ("readers"), respectively. The m⁶A reader proteins YTH5, YTH9, and YTH13 are also known as ECT4, ECT2, and ECT3, respectively. RNA methylation affects all aspects of RNA metabolism, including stability, export, intron splicing, and translational control, which are crucial for plant development and stress responses. Several potential m⁶A erasers and readers are yet to be identified.

8

9 Figure 7. Plant IncRNA grammar is determined by the transcript interactome.

Multiple features contribute to the interaction of IncRNAs with DNA, protein partners or 10 other RNA molecules. First, their expression pattern and their subcellular localization 11 will restrict the range of potential partners. Second, the IncRNA interacting capacity 12 depends on its sequence, post-transcriptional modifications, and secondary and tertiary 13 structure adopted, which is, in turn, modulated by the interaction with partner molecules. 14 Third, the resulting IncRNA interactome participates in the regulatory networks behind 15 plant development and adaptation to the environment as all these factors can be 16 responsive to environmental cues. 17

18

19 Figure 8. Unanswered questions of miRNA biogenesis.

A. Can the alterations of processing efficiency caused by pri-miRNA refolding upon

- 21 temperature change act as thermosensors during the plant response to heat?
- B. Can proteins specifically binding to the ssDNA or RNA/DNA strands of R-loops act as
- 23 scaffold to recruit the microprocessor to *MIRNA* loci?
- 24 C. How does the microprocessor and spliceosome interact?
- D. Can we define different D-bodies? And if so, can we establish the precise biochemistry within D-bodies during their maturation?
- E. Are co-transcriptionally processed miRNAs functionally different from their siblings produced post-transcriptionally, perhaps defining mobile miRNAs?
- 29

Figure 9. siRNA movement during male and female germline development in Arabidopsis.

Here we use a more relaxed definition of the germline to indicate the cell lineage that undergoes meiosis and produces the gamete(s). The germline as strictly defined, is marked in red, ie. the generative and sperm cells (male) and the egg cell (female). Arrows mark the direction of the proposed siRNA movement. PMC, pollen mother cells; MMC, megaspore mother cells.

37

Figure 10. Establishment and maintenance of DNA methylation by RdDM

21- and 22-nt siRNAs that are generated by DCL4 and DCL2 can bind to AGO1 to

40 target mRNAs for post-transcriptional silencing (PTGS) or bind to AGO6 to initiate

RdDM in partnership with Pol V and DRM2. The latter enzymes are also key to the 1 major RdDM pathway that maintains silencing of thousands of loci and requires 24 -t 2 siRNAs that are generated by the Pol IV-RDR2 complex and DCL3 and loaded primarily 3 into AGO4. CG maintenance methylation, requiring MET1 and HDA6, is important for 4 5 both Pol IV and Pol V recruitment, and correlates with histone H3 lysine 9 dimethylation (H3K9me2) among associated nucleosomes. Proteins that interact with these marks 6 and are implicated in Pol IV or Pol V transcriptional activity are indicated, as are histone 7 modifying enzymes involved in establishing repressive chromatin environments. The 8 figure is an update of the transcription fork model originally published in 2013 (Pikaard 9 et al., 2012), revised in 2017 (Wendte and Pikaard, 2017) and also adapted by other 10 authors (Matzke and Mosher, 2014). 11

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Figure 5 339x190 mm (x DPI)





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