# Arabidopsis AAR2, a conserved splicing factor in eukaryotes, acts in microRNA biogenesis

Lusheng Fan<sup>a,b,1</sup>, Bin Gao<sup>b,1</sup>, Ye Xu<sup>b</sup>, Nora Flynn<sup>b</sup>, Brandon Le<sup>b</sup>, Chenjiang You<sup>c</sup>, Shaofang Li<sup>d</sup>, Natalia Achkar<sup>e</sup>, Pablo A. Manavella<sup>e</sup> Zhenbiao Yang<sup>b</sup>, and Xuemei Chen<sup>b,2</sup>

Contributed by Xuemei Chen; received May 16, 2022; accepted August 30, 2022; reviewed by Brian Gregory and Bin Yu

MicroRNAs (miRNAs) play an essential role in plant growth and development, and as such, their biogenesis is fine-tuned via regulation of the core microprocessor components. Here, we report that Arabidopsis AAR2, a homolog of a U5 snRNP assembly factor in yeast and humans, not only acts in splicing but also promotes miRNA biogenesis. AAR2 interacts with the microprocessor component hyponastic leaves 1 (HYL1) in the cytoplasm, nucleus, and dicing bodies. In aar2 mutants, abundance of nonphosphorylated HYL1, the active form of HYL1, and the number of HYL1-labeled dicing bodies are reduced. Primary miRNA (pri-miRNA) accumulation is compromised despite normal promoter activities of MIR genes in aar2 mutants. RNA decay assays show that the aar2-1 mutation leads to faster degradation of pri-miRNAs in a HYL1-dependent manner, which reveals a previously unknown and negative role of HYL1 in miRNA biogenesis. Taken together, our findings reveal a dual role of AAR2 in miRNA biogenesis and premessenger RNA splicing.

microRNA | HYL1 | AAR2 | pri-miRNA | SE

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Small RNAs of 21 to 24 nucleotides (nt), including microRNAs (miRNAs) and small interfering RNAs (siRNAs), are involved in a wide range of biological processes (1). miRNAs are derived from primary transcripts with partially complementary fold-backs while siRNAs are generated from long, perfect, or nearly perfect double-stranded RNAs. Both siRNAs and miRNAs are bound by Argonaute proteins to form RNAinduced silencing complexes (RISCs), which regulate gene expression through either transcriptional gene silencing or posttranscriptional gene silencing in a sequence-specific

In plants, most miRNA genes (MIR) are transcribed as independent transcription units to produce primary miRNAs (pri-miRNAs) that contain a stem loop structure (4). In Arabidopsis, a pri-miRNA is processed in the nucleus by Dicer-like 1 (DCL1) into an miRNA/miRNA\* duplex, which is 2'-O-methlyated at the 3' terminal nucleotides by Hua enhancer 1 (HEN1); the duplex is bound by AGO1 followed by RISC formation upon the ejection of the miRNA\* strand; miRNA-RISC complexes are then exported to the cytoplasm and repress the expression of target genes via transcript cleavage or translational repression (2, 5, 6). In Arabidopsis, several miRNAs target noncoding transcripts from trans-acting sirna (TAS) loci for cleavage and trigger the production of siRNAs from the cleavage fragments (1). These siRNAs are termed ta-siRNAs as they regulate target genes in trans (7).

The precise and efficient processing of pri-miRNAs by DCL1 is enhanced by the double-stranded RNA-binding protein hyponastic leaves 1 (HYL1) and the zinc finger protein serrate (SE); the three proteins form a protein complex called the microprocessor (8, 9). In Arabidopsis, del1 and se null mutations cause embryonic lethality, and hyll null alleles lead to strong developmental defects (10-12). Consistent with their coordinated function in pri-miRNA processing, these three proteins colocalize in nuclear membrane-less speckles, known as dicing bodies (D-bodies), as well as in the nucleoplasm, both of which are thought to be sites of pri-miRNA processing (13, 14). SE contains three intrinsically disordered regions that mediate phase separation, which plays a role in D-body formation (15). Besides its role in miRNA biogenesis, SE is also involved in pre-messenger RNA (pre-mRNA) splicing. Like SE (16), other factors first found to play a role in splicing, such as cap-binding protein 80 (CBP80); cap-binding protein 20 (CBP20); stabilized 1 (STA1); proteins in the MOS4-associated complex (MAC) including MAC3A, MAC3B, PRL1, and PRL2; and serrate-associated protein 1 (SEAP1), were shown to be required for miRNA biogenesis (17-20). Conversely, several factors first found to play a role in miRNA biogenesis, such as protein phosphatase 4 (PP4) regulatory subunit 3 (21), THP1 in the TREX-2 complex (22), and RBV (23), were also found to act in the pre-mRNA splicing for subsets of genes.

## **Significance**

In yeast and humans, AAR2 is involved in pre-messenger RNA (pre-mRNA) splicing through regulating U5 snRNP assembly. This study shows that Arabidopsis AAR2 promotes microRNA (miRNA) accumulation in addition to its conserved role in pre-mRNA splicing. AAR2 is associated with the microprocessor component HYL1 and promotes its dephosphorylation to produce the active form in miRNA biogenesis. The study also reveals a previously unknown role of HYL1 in causing the degradation of the primary precursors to miRNAs (primiRNAs) and a role of AAR2 in protecting pri-miRNAs from HYL1depedent degradation. Taken together, our findings provide insights into the role of a conserved splicing factor in miRNA biogenesis in plants.

Author contributions: L.F., B.G., Z.Y., and X.C. designed research; L.F., B.G., Y.X., N.F., and N.A. performed research; L.F., B.G., Y.X., N.F., B.L., C.Y., S.L., N.A., P.A.M., and X.C. analyzed data; and L.F., B.G., Y.X., N.F., and X.C. wrote the paper.

Reviewers: B.D.G., University of Pennsylvania, and B.Y., University of Nebraska, Lincoln

The authors declare no competing interest.

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<sup>1</sup>L.F. and B.G. contributed equally to this work.

<sup>2</sup>To whom correspondence may be addressed. Email: xuemei.chen@ucr.edu.

This article contains supporting information online at http://www.pnas.org/lookup/suppl/doi:10.1073/pnas. 2208415119/-/DCSupplemental.

Published October 3, 2022.

Cofactors of the Ribonuclease III (RNase III) enzymes involved in miRNA biogenesis are regulated to fine-tune miRNA production (24). In animals, TRBP and DGCR8, cofactors of Dicer and Drosha, respectively, are phosphorylated by mitogen-activated protein kinase (MAPK), which stabilizes the respective protein complexes to promote miRNA biogenesis (25, 26). However, in Arabidopsis, dephosphorylation of HYL1 by CPL1 and CPL2 facilitates the localization of HYL1 in D-bodies and enhances miRNA biogenesis (27). MPK3 was identified as a kinase phosphorylating HYL1 in both rice and Arabidopsis, and a mutation in Arabidopsis MPK3 led to increased miRNA accumulation (28). SMEK1 (suppressor of MEK1) inhibits MAPK-mediated HYL1 phosphorylation and promotes PP4-mediated HYL1 dephosphorylation, which is required for miRNA biogenesis (29). The core photomorphogenic regulator COP1 modulates the degradation of HYL1 in response to the light-to-dark transition (30). The phosphorylated form of HYL1 is inactive and less susceptible to degradation induced by extended periods of darkness. Upon light restoration, HYL1 is quickly reactivated by dephosphorylation, which in turn enables miRNA production and the proper expression of developmentally important genes (31). In response to abscisic acid (ABA) treatment, SnRK2 kinases phosphorylate HYL1, which is important for the stability of HYL1 and miRNA accumulation (32). These findings suggest that dephosphorylated HYL1 is the active form of the protein in miRNA biogenesis, but phosphorylated HYL1 is more stable and may serve as a reservoir from which dephosphorylated HYL1 can be produced.

SE is another important protein that assists DCL1 in primiRNA processing. Intriguingly, while SE is considered a component of the microprocessor, recent studies also point to a negative effect of SE in miRNA biogenesis. CHR2, an ATPase subunit of the SWI/SNF chromatin-remodeling complex, accesses pri-miRNAs through its interaction with SE and remodels the structures of pri-miRNAs to inhibit pri-miRNA processing (33). A MAC component, MAC5A, binds the stemloop region of pri-miRNAs and protects pri-miRNAs from degradation, thus promoting miRNA biogenesis; the degradation of pri-miRNAs in mac5a mutants is dependent on SE and partially caused by 5'-to-3' exoribonucleases XRN2 and XRN3 (34). How the positive and negative roles of SE in miRNA biogenesis are balanced is as yet unknown. SE is phosphorylated by pre-mRNA processing 4 kinase A (PRP4KA), leading to a lower binding affinity to HYL1 as well as higher susceptibility to degradation by the 20S proteasome. Phosphorylation of SE via PRP4KA may define a regulatory mechanism to clear excess SE to maintain its proper levels and promote miRNA production (35). Whether the other microprocessor component, HYL1, has a negative role in miRNA biogenesis is unknown.

In this study, we performed a genetic screen in Arabidopsis using SUC2:atasi-SUL as a reporter to identify genes involved in ta-siRNA or miRNA biogenesis and activity (36). In the SUC2: atasi-SUL transgenic line, atasi-SUL, an artificial ta-siRNA, is engineered in the backbone of TAS1 such that its biogenesis is triggered by miR173 as for endogenous ta-siRNAs from this locus (36). atasi-SUL represses the expression of the sulfur (SUL) gene in mesophyll cells, leading to leaf bleaching. Through this mutagenesis screen, we isolated a mutant showing a global reduction in both ta-siRNA and miRNA production. This mutation is in an uncharacterized gene encoding a protein homologous to AAR2, a U5 snRNP particle assembly factor in yeast and humans (37, 38). We show that aar2 mutations reduce pri-miRNA levels without affecting the transcription of the corresponding MIR genes. The reduction of pri-miRNA levels in aar2 is caused by pri-miRNA degradation that is dependent on HYL1 but not SE,

indicating that HYL1 also has a negative effect in miRNA biogenesis. AAR2 associates with HYL1 and promotes its localization to D-bodies. The ratio between nonphosphorylated and phosphorylated HYL1 is decreased in the aar2 mutant, suggesting that AAR2 promotes HYL1 dephosphorylation. Finally, AAR2 is also involved in pre-mRNA splicing for a subset of introns. Our findings suggest that in addition to its conserved role in premRNA splicing, AAR2 is involved in miRNA biogenesis by preventing HYL1-dependent pri-miRNA degradation and promoting HYL1 dephosphorylation.

### **Results**

Isolation of a mutant with pleiotropic developmental defects and a global reduction in miRNA accumulation. To identify genes required for the biogenesis and/or activity of ta-siRNAs, we took advantage of a visual reporter of ta-siRNA activity, SUC2::atasi-SUL (atasi-SUL) (36), to perform an ethyl methanesulfonate mutagenesis screen. In the atasi-SUL transgenic line, an artificial ta-siRNA targeting the chlorophyll biosynthetic gene SUL (also known as CHLORINA42) leads to the yellowing of leaves. We isolated a suppressor mutant with green leaves, a phenotype indicative of compromised atasi-SUL biogenesis or activity (Fig. 1 A and B). To identify the mutation, we back-crossed the mutant with the parental line atasi-SUL and conducted whole-genome resequencing using pooled F2 segregants with the mutant phenotype. A  $G^{2400}$ -to-A mutation in AT1G66510 was identified that introduced a premature stop codon in the open reading frame (SI Appendix, Fig. S1A). This gene (referred to as AAR2 hereafter) encodes a homolog to AAR2 that is involved in U5 snRNP assembly and consequently pre-mRNA splicing in yeast and humans (37). To examine the developmental phenotypes caused by the aar2 mutation, we crossed this mutant with wild-type plants and obtained the aar2-1 allele without the atasi-SUL transgene. We also acquired an aar2 T-DNA insertion allele, SALK117746 (designated as aar2-2 hereafter) (SI Appendix, Fig. S1A). RNA sequencing (RNA-seq) with these two mutants revealed reduced levels of AAR2 transcripts as compared with wild-type plants (SI Appendix, Fig. S1B). aar2-2 plants showed developmental phenotypes similar to aar2-1 plants, including reduced plant size, shorter primary roots, lower number of lateral roots, and fewer stamens (Fig. 1C and SI Appendix, Fig. S2 A-J). Both aar2 mutants had shorter hypocotyls as compared to wild-type mutants when grown in darkness, and a similar phenotype was also observed for hyl1-2 (SI Appendix, Fig. S2 K-L). To determine whether the aar2-1 mutation was responsible for the reduced leaf yellowing in aar2-1 atasi-SUL, we generated a construct in which a genomic fragment containing the promoter and coding region of AAR2 was fused with yellow fluorescent protein (YFP) and transformed the construct into aar2-1 atasi-SUL. The transgene fully rescued the leaf color phenotype and the developmental defects of aar2-1 atasi-SUL (Fig. 1 A and B). These results demonstrated that the aar2-1 mutation is responsible for these phenotypes.

Given that the aar2-1 atasi-SUL mutant showed compromised atasi-SUL activity, we performed RNA gel blot analysis to determine the levels of atasi-SUL as well as endogenous ta-siRNAs in the aar2-1 atasi-SUL mutant. Both atasi-SUL and endogenous ta-siRNAs showed reduced accumulation in aar2-1 atasi-SUL as compared to atasi-SUL (Fig. 1D). As ta-siRNA production is dependent on their trigger miRNAs, we further examined the levels of miR173 and miR390, the trigger miR-NAs for ta-siRNA production from TAS1, TAS2, and TAS3, as

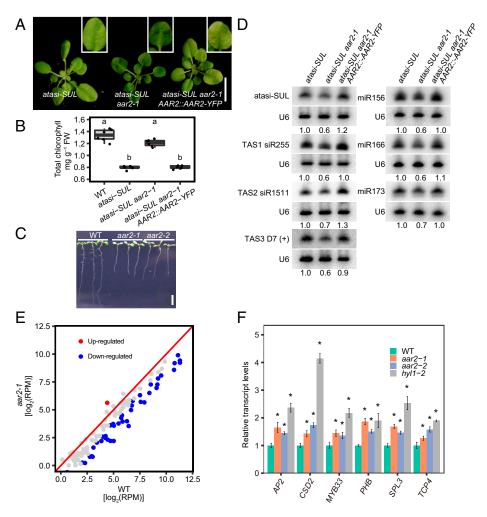


Fig. 1. Isolation of a mutant with pleiotropic developmental defects and a global reduction in miRNA accumulation. (A) Phenotypes of atasi-SUL, atasi-SUL aar2-1, AAR2::AAR2-YFP atasi-SUL aar2-1 plants. Images of rosettes were taken from 3-wk-old plants (Scale bar, 10 mm). (B) Total chlorophyll content was quantified from the rosette leaves of 4-wk-old Arabidopsis seedlings. Six biological replicates were analyzed for each genotype. Letters indicate significant differences (one-way ANOVA followed by Tukey's multiple comparisons; P < 0.001). (C) 9-d-old seedlings of WT, aar2-1 and aar2-2 showing primary root length (Scale bar, 5 mm). (D) RNA gel blot analysis showing reduced accumulation of atasi-SUL, endogenous ta-siRNAs, and miRNAs in the atasi-SUL aar2-1 mutant. Note that expression of the AAR2::AAR2-YFP transgene fully rescued the ta-siRNA and miRNA accumulation defects. U6 was used as the loading control. The numbers below the blots represent the relative amounts of ta-siRNAs and miRNAs. (E) Scatter plot showing differentially accumulated miRNAs in WT and aar2-1. Differentially accumulated miRNAs were identified by DESeq2 with P value < 0.01 and fold change > 2. (F) Determination of miRNA target transcript levels in WT, aar2-1, aar2-2, and hyl1-2 by qRT-PCR. Transcript levels were normalized to UBQ5 and then calculated relative to those in WT. Error bars, SD calculated from three independent replicates. Asterisks indicate significant difference (Student's t test, \*P < 0.05).

well as the levels of other endogenous miRNAs. All examined miRNAs showed a moderate reduction in abundance in aar2-1 atasi-SUL (Fig. 1D and SI Appendix, Fig. S3A). These results indicate that the aar2-1 mutation impaired miRNA production and indirectly affected ta-siRNA biogenesis. Moreover, the AAR2-YFP transgene fully rescued the ta-siRNA and miRNA production defects in aar2-1, further confirming that AAR2 is required for ta-siRNA and miRNA biogenesis (Fig. 1D). To assess the global profiles of small RNAs in the aar2 mutant, we performed small RNA-seq with aar2-1 and wild-type plants. Three biological replicates for each genotype were included, and they were highly correlated (SI Appendix, Fig. S3B). A significant global reduction in 21-nt to 24-nt small RNAs, which consist of most endogenous miRNAs and siRNAs, was found in aar2-1 (Fig. 1E and SI Appendix, Fig. S3C). miRNAs also showed a global reduction in aar2-1, with 39 miRNAs showing a significant decrease in abundance (SI Appendix, Fig. S3D and Dataset S1). In plants, pri-miRNAs are processed in either a base-to-loop or a loop-to-base manner (39, 40). We found that the reduction in miRNA abundance in aar2-1 was regardless of the direction of pri-miRNA processing (SI Appendix, Fig. S4A). When compared to all detected miRNAs, the 39 downregulated miRNAs in aar2-1 did not show a significant difference in the lengths of pri-miRNAs (SI Appendix, Fig. S4B). In comparison to 39 randomly selected miRNAs, the 39 downregulated miRNAs in aar2-1 represented significantly fewer miRNA families (SI Appendix, Fig. S4C), suggesting that some affected miRNAs belong to evolutionarily old miRNA families with more than one member (41). Consistent with such "old" miRNAs having higher abundance (41), the 39 miRNAs showed higher levels as compared to all detected miRNAs in wild-type plants (SI Appendix, Fig. S4D). The reduction in the levels of miRNAs in aar2 mutants correlated with an increase in known miRNA-targeted mRNAs, as expected for an miRNA-deficient mutant (Fig. 1F). To test the possibility that AAR2 affects miRNA production through promoting the expression of key genes in miRNA biogenesis, we examined the transcript or protein levels of known players in miRNA biogenesis or activity, and no significant changes were found in the aar2 mutants (SI Appendix, Fig. S5).

AAR2 associates with microprocessor components and is required for the localization of HYL1 in D-bodies. To further investigate the molecular functions of the AAR2 protein, we first examined its subcellular localization. In the AAR2::AAR2-YFP transgenic line, AAR2-YFP signals were ubiquitous in root and leaf cells and were present in both the nucleus and the cytoplasm (SI Appendix, Fig. S6). In root tip cells of the HYL1-YFP AAR2-mCherry transgenic line, AAR2-mCherry colocalized with HYL1-YFP in the nucleoplasm but was absent from the HYL1-YFP-labeled D-bodies (Fig. 2A). However, we found that HYL1-YFP-labeled D-bodies were significantly reduced in number in aar2-1 as compared to wild-type plants (Fig. 2 B-D), suggesting that AAR2 promotes the localization of HYL1 in D-bodies. Given the fact that DCL1, HYL1, and SE colocalize in D-bodies (13) and aar2-1 is compromised in the D-body localization of HYL1, we speculated that AAR2 may be associated with the microprocessor and act as an accessory component. Thus, we performed bimolecular fluorescence complementation (BiFC) analysis to test the association of AAR2 with DCL1, HYL1, and SE. In this assay, we fused AAR2 with an N-terminal fragment of YFP at its C terminus (AAR2-nYFP) and coexpressed it with DCL1, HYL1, and SE tagged with a C-terminal fragment of YFP at their C termini in Nicotiana benthamiana. We also included HYL1-nYFP and DCL1-cYFP as a positive control, which have been shown to associate with each other in the nucleus (13). As expected, the coexpression of HYL1-nYFP and DCL1-cYFP generated YFP signals mainly in D-bodies in the nucleus (SI Appendix, Fig. S7A). Similarly, the coexpression of AAR2-nYFP with DCL1 and SE also generated YFP signals, but mainly in the nucleoplasm (SI Appendix, Fig. S7A). Interestingly, the coexpression of AAR2-nYFP with HYL1-cYFP gave rise to YFP signals in the cytoplasm, the nucleoplasm, and D-bodies (SI Appendix, Fig. S7A). The nucleoplasmic and cytoplasmic signals were

consistent with the localization of AAR2 and HYL1 in both subcellular compartments (SI Appendix, Fig. S6 [42]). However, the D-body BiFC signals were surprising as AAR2-YFP did not from nuclear foci in the AAR2::AAR2-YFP transgenic line. Overexpression with the 35S promoter probably contributed to the signals, but the absence of D-body BiFC signals between AAR2 and DCL1 or AAR2 and SE suggested that interactions between AAR2 and HYL1 occurs in D-bodies, perhaps transiently. This is consistent with the finding that the aar2-1 mutation affects the formation of HYL1-labeled D-bodies. In a complementary experiment, crude extracts from AAR2:: AAR2-YFP transgenic plants were subjected to size exclusion chromatography. AAR2-YFP was found to partially cofractionate with DCL1, HYL1, and SE in high-molecular-weight fractions (SI Appendix, Fig. S7B). Together, the results are consistent with the association of AAR2 with the microprocessor in the nucleoplasm. The reduced D-body localization of HYL1 and the positive BiFC signals between AAR2 and HYL1 in D-bodies were consistent with the role of AAR2 in enabling the D-body localization of HYL1. However, the interactions between AAR2 and the microprocessor components were likely weak or transient, as we were unable to detect the interactions by coimmunoprecipitation (SI Appendix, Fig. S8).

AAR2 promotes pri-miRNA accumulation without affecting MIR promoter activities. To further investigate how AAR2 promotes miRNA biogenesis, we examined the levels of pri-miRNAs using qRT-PCR and found that levels of all 16 examined primiRNAs except miR167b showed a significant reduction in both aar2 mutants (Fig. 3A). This finding indicates that AAR2 is required for pri-miRNA accumulation. As steady-state primiRNA levels are a result of MIR transcription, pri-miRNA processing, and pri-miRNA degradation, we first examined whether MIR promoter activities were affected in aar2-1. pMIR167a::GUS,

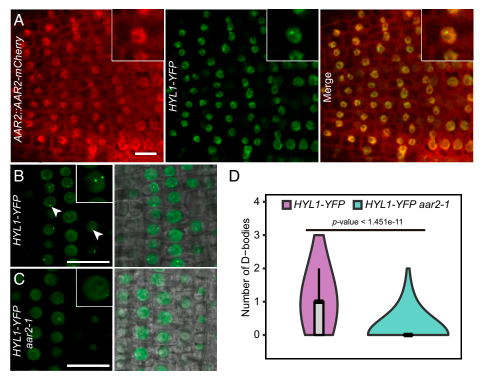


Fig. 2. AAR2 associates with microprocessor components and is required for the localization of HYL1 in D-bodies. (A) Confocal images of root tip cells expressing both HYL1-YFP and AAR2-mCherry. Enlarged nuclei are shown in the insets. (B and C) Representative images of HYL1-YFP in WT (B) and aar2-1 (C). Arrowheads indicate D-bodies (Scale bar, (A-C), 20 µm). (D) Violin plots showing the number of HYL1-YFP-labeled D-bodies per cell in WT and aar2-1. Quantification was performed by counting more than 400 cells from 15 roots for each genotype. P value was calculated by the Wilcoxon test.

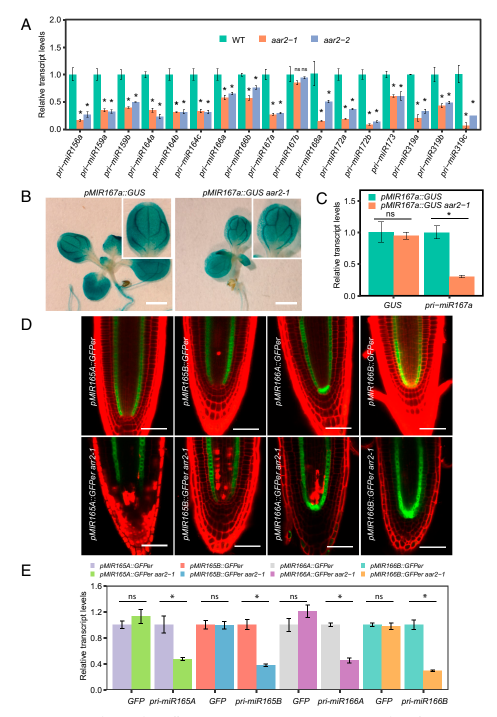


Fig. 3. AAR2 promotes pri-miRNA accumulation without affecting MIR promoter activities. (A) qRT-PCR analysis of pri-miRNAs in WT, aar2-1, and aar2-2. UBQ5 was used as an internal control. (B) Representative images of histochemical staining of GUS in pMIR167a::GUS and pMIR167a::GUS aar2-1 transgenic plants. Fifteen plants for each genotype were analyzed (Scale bar, 1 mm). (C) Levels of the GUS transcript and pri-miR167a in pMIR167a::GUS and pMIR167a: GUS aar2-1 plants as determined by qRT-PCR. (D) Representative images of roots of MIR165A::GFPer, MIR165B::GFPer, MIR166A::GFPer, and MIR166B::GFPer in WT (Top) and aar2-1 (Bottom). We analyzed 15 roots for each genotype. (E) Levels of the GFP transcript and the corresponding endogenous pri-miRNAs in the aforementioned promoter reporter lines as determined by qRT-PCR (Error bars, (A, C, and E), SD calculated from three independent replicates). Asterisks indicate significant difference (Student's t test, \*P < 0.05). ns, no significant difference.

a GUS reporter for MIR167a promoter activity (43), was crossed into aar2-1. GUS histochemical staining showed no obvious difference in terms of GUS activity between aar2-1 and wild-type plants (Fig. 3B). Moreover, qRT-PCR analysis showed that primiR167a was significantly reduced in aar2-1 while GUS transcript levels were comparable between aar2-1 and wild-type plants (Fig. 3C). We further examined four other MIR promoter activity reporters—MIR165a::GFPer, MIR165b::GFPer, MIR166a::GFPer, and MIR166b::GFPer (44)—and found no detectable difference between aar2-1 and wild-type plants in terms of GFP fluorescence intensities or GFP transcript levels, while levels of endogenous primiR165A, pri-miR165B, pri-miR166A, and pri-miR166B were significantly reduced in aar2 as compared to wild-type plants (Fig. 3 D and E). Taken together, these results suggest that AAR2 is required for pri-miRNA accumulation without affecting the transcription of the corresponding MIR genes.

Mutations in AAR2 lead to pri-miRNA degradation in a HYL1-dependent manner. The association of AAR2 with the microprocessor and its requirement for the formation of HYL1labeled D-bodies prompted us to test the genetic interactions between aar2-1 and mutations in HYL1 and SE. The se-1 aar2-1 double mutant showed more-severe developmental phenotypes than either the aar2-1 or se-1 single mutant, while the hyl1-2 aar2-1 double mutant largely resembled hyl1-2 in plant morphology (Fig. 4A). Consistent with these observations, miRNA levels in the se-1 aar2-1 double mutant were further decreased in comparison with those in either the aar2-1 or se-1 single mutant while miRNA levels in the hyl1-2 aar2-1 double mutant were comparable to those in hyl1-2 (Fig. 4B), suggesting that AAR2 and SE contribute additively to miRNA biogenesis while AAR2 contributes to miRNA biogenesis through the HYL1.

The aar2-1 mutation caused a reduction in the levels of primiRNAs without affecting MIR gene transcription. We investigated whether this reduction of pri-miRNA accumulation still occurred in the hyl1 or se background, in which pri-miRNAs accumulate to higher levels due to inefficient processing (45). Levels of examined pri-miRNAs were reduced in the se-1 aar2-1 double mutant relative to se-1 but were similar in the hyl1-2 aar2-1 double mutant as compared to hyl1-2 (Fig. 4C), which indicates that the effects of aar2-1 on pri-miRNA accumulation were independent of SE but dependent on HYL1.

To determine how aar2-1 results in lower pri-miRNA levels, we observed the decay of pri-miRNAs in seedlings treated with the transcription inhibitor cordycepin. When transcription is inhibited, the levels of pri-miRNAs are expected to decrease due to DCL1 processing as well as RNA degradation. Consistent with findings from a previous study (34), in both se-1 and hyl1-2, pri-miRNA levels decreased more slowly than in the wild type, probably due to the lack of pri-miRNA processing (Fig. 4D). The half-lives of examined pri-miRNAs were similar between aar2-1 and the wild type (Fig. 4D). However, the half-lives of pri-miRNAs were shorter in the se-1 aar2-1 double

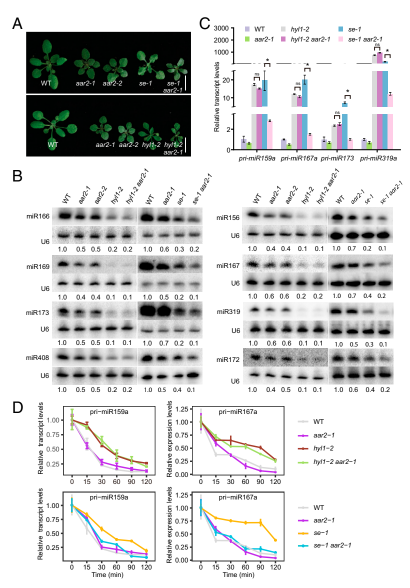


Fig. 4. AAR2 affects miRNA biogenesis in a HYL1- but not SE-dependent manner. (A) Phenotypes of 3-wk-old plants of the indicated genotypes (Scale bar, 10 mm). (B) RNA gel blot analysis of miRNA accumulation in se-1 aar2-1 and hyl1-2 aar2-1 double mutants as compared to the corresponding single mutants. U6 was used as the loading control. The numbers below the blots represent relative miRNA levels. (C) qRT-PCR analysis of pri-miRNAs in hyl1-2 ααr2-1 and se-1 aar2-1 as compared to the corresponding single mutants. UBQ5 was used as an internal control (Error bars, SD calculated from three independent replicates). Asterisks indicate significant difference (Student's t test, \*P < 0.05). (D) Analysis of pri-miRNA decay upon treatment of WT, aar2-1, hyl1-2, se-1, hyl1-2 aar2-1, and se-1 aar2-1 seedlings with the transcription inhibitor cordycepin. UBQ5 was used as an internal control (Error bars, SD from three technical replicates).

mutant as compared to se-1 (Fig. 4D), suggesting that the aar2-1 mutation led to the degradation of pri-miRNAs. Intriguingly, the half-lives of pri-miRNAs were similar in the hyl1-2 aar2-1 double mutant and the hyl1-2 single mutant, suggesting that the accelerated pri-miRNA degradation caused by aar2-1 is dependent on HYL1.

AAR2 is involved in HYL1 degradation. Previous studies showed that HYL1 is an unstable protein with relatively short half-lives, particularly under darkness (30, 46). That AAR2 affects miRNA biogenesis through HYL1 prompted us to explore the possibility that AAR2 plays a role in the regulation of HYL1 protein stability. We transferred 10-d-old seedlings to darkness for 2 d and determined HYL1 protein levels in wild-type and aar2 mutants. Under dark treatment, HYL1 levels decreased in wild-type mutants, which is consistent with previous findings (30). HYL1 protein levels were higher in aar2-1 and aar2-2 relative to wild-type mutants after dark treatment (Fig. 5A). Treatment of seedlings with carbobenzoxy-Leu-Leu-leucinal (MG132) while they were in darkness led to higher levels of HYL1 in both wild-type and aar2 mutants (Fig. 5B). These results indicate that the degradation of HYL1 is impaired in aar2. To test whether AAR2 affects HYL1 degradation under the light condition, we performed a protein decay assay by treating longday grown, 10-d-old seedlings with cycloheximide (CHX) during the light period to block de novo protein synthesis and examined HYL1 protein levels at 2 h after treatment. The levels of HYL1 showed a 50% decrease after 2 h of CHX treatment in wild-type mutants, but the decay rate was much slower in aar2 mutants (Fig. 5C). Moreover, MG132 treatments notably suppressed the degradation of HYL1 in both wild-type and aar2 mutants (Fig. 5D). Therefore, AAR2 is required for the degradation of HYL1.

Because HYL1 is degraded in the cytoplasm under darkness (30), we performed nuclear-cytoplasmic fractionation to examine the distribution of HYL1 in the dark. Results showed that HYL1 levels were increased in aar2-1, specifically in the cytoplasm, as compared to the wild type (Fig. 5E). Moreover, there was no obvious difference in HYL1 levels between aar2-1 and wild type in either total, cytoplasmic, or nuclear fractions under the light conditions (SI Appendix, Fig. S9A), suggesting that aar2-1 does not affect HYL1's nuclear-cytoplasmic partitioning. Similarly, RNA gel blot analysis showed that aar2-1 did not alter the nucleocytoplasmic distribution of miRNAs (SI Appendix, Fig. S9B). Taken together, these results support that AAR2 is involved in HYL1 degradation in the cytoplasm but does not affect miRNA or HYL1's nuclear-cytoplasmic distribution.

AAR2 is required for HYL1 dephosphorylation. Phosphorylated HYL1 is inactive in miRNA biogenesis but forms a reserve pool of HYL1 that is resistant to degradation (31). Because HYL1 is more resistant to degradation in aar2 mutants, we thought that the phosphorylation status of HYL1 might be affected in the mutants. CPL1 is a phosphatase known to dephosphorylate

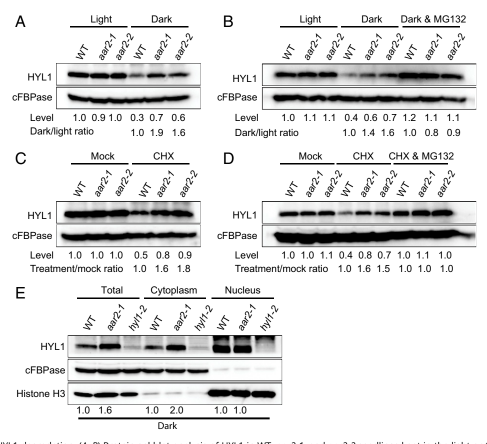


Fig. 5. AAR2 affects HYL1 degradation. (A, B) Protein gel blot analysis of HYL1 in WT, aar2-1, and aar2-2 seedlings kept in the light or transferred to the dark and grown for 2 d (A) or incubated with MG132 during the dark period (B). HYL1 levels were normalized against cFBPase and expressed in values relative to WT in the light. The dark/light ratio represents the ratio of HYL1 levels in the dark and in the light and is relative to WT. (C, D) Protein gel blot analysis of HYL1 in long-day grown WT, aar2-1, and aar2-2 seedlings treated with CHX (C) or CHX together with MG132 (D) for 2 h. HYL1 levels were normalized against cFBPase and expressed in values relative to the WT mock-treated samples. The treatment/mock ratio represents the ratio of HYL1 levels in mock-treated samples and in samples treated with CHX or CHX+MG132 and is relative to WT. (E) HYL1 levels in total, cytoplasmic, and nuclear fractions under dark treatment as described in (A). cFBPase and histone H3 were used as cytoplasmic and nuclear markers, respectively. HYL1 levels were normalized against cFBPase or histone H3 and expressed in values relative to WT.

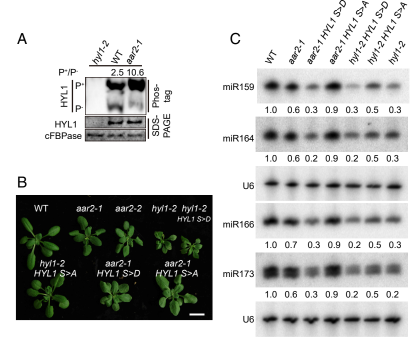


Fig. 6. AAR2 affects HYL1 phosphorylation status. (A) Phosphoprotein mobility shift gel detecting phosphorylated (P+) and nonphosphorylated (P-) HYL1 in the indicated genotypes. SDS-PAGE gels (Middle, Bottom) show the levels of HYL1 and cFBPase (used as a loading control), respectively. (B) Image shows 3-wk-old plants of the indicated genotypes (Scale bar, 10 mm). (C) RNA gel blot analysis of miRNA accumulation in the indicated genotypes. U6 was used as the loading control. The numbers below the blots represent relative abundances.

HYL1 (27). We introduced the cpl1-7 mutation into aar2-1 and found that the aar2-1 cpl1-7 double mutant showed more-severe developmental phenotypes than either aar2-1 or cpl1-7 (SI Appendix, Fig. S10A). The phosphorylation status of HYL1 was analyzed using phos-tag gels, polyacrylamide gels containing phostag, a phosphate-binding molecule that reduces the mobility of phosphorylated proteins, and independent experiments showed that the hypo-phosphorylated form of HYL1 was reduced in aar2-1 as compared to wild-type mutants (Fig. 6A and SI Appendix, Fig. S10B). We examined pri-miRNA accumulation in the aar2-1 cpl1-7 double mutant using qRT-PCR. Results showed that levels of all examined pri-miRNAs were decreased in both aar2-1 and cpl1-7, while a further decrease was observed for some pri-miRNAs in the aar2-1 cpl1-7 double mutant as compared to either aar2-1 or cpl-17 (SI Appendix, Fig. S10C). Moreover, the levels of mature miRNAs were reduced in the aar2-1 cpl1-7 double mutant as compared to either single mutant (SI Appendix, Fig. S10D).

We next tested whether AAR2 affects miRNA biogenesis through HYL1 dephosphorylation. Expression of HYL1 S > A, a nonphosphorylatable HYL1 with all seven predicted serine phosphorylation sites mutated to alanine (27), partially rescued the developmental defects of aar2, while aar2-1 plants expressing HYL1 S > D, a phosphomimetic HYL1 with all seven predicted serine phosphorylation sites mutated to aspartic acid (27), showed more-severe developmental defects (Fig. 6B). We examined miRNA accumulation in HYL1 S > D aar2-1 and HYL1 S > A aar2-1 using RNA gel blotting. As expected, the accumulation of all examined miRNAs was partially rescued in hyl1-2 by HYL1 S > A while HYL1 S > D could not rescue the miRNA biogenesis defects in hyl1-2 (Fig. 6C and SI Appendix, Fig. S10E). Accordingly, pri-miRNA levels were unchanged in hyl1-2 HYL1 S > D and reduced in hyl1-2 HYL1 S > A relative to hyl1-2 (SI Appendix, Fig. S10F). The accumulation of some of the examined miRNAs was also partially rescued in aar2-1 by HYL1 S > A while HYL1 S > D led to a further reduction in the levels of all examined miRNAs in aar2-1 (Fig. 6C and SI Appendix,

Fig. S10E). These results indicate that AAR2 promotes miRNA accumulation by regulating the HYL1 phosphorylation status. The accumulation of phosphorylated HYL1 in aar2 mutants (Fig. 6A and SI Appendix, Fig. S10B) could also explain the increased stability of this protein in the mutants.

AAR2 is required for efficient messenger RNA splicing. Homologs of AAR2 in yeast and mammals are involved in the assembly of U5 snRNP, an essential complex in pre-mRNA splicing (37, 47). In Arabidopsis, a role of AAR2 in splicing has never been characterized. This prompted us to examine whether aar2 mutations show splicing defects. We performed RNA-seq in wild type, aar2-1, and aar2-2 with three biological replicates. In total, we identified 1,162 up-regulated genes (hyper-DEGs) and 1,078 down-regulated genes (hypo-DEGs) in aar2-1, and 647 hyper DEGs and 924 hypo DEGs in aar2-2. 459 up-regulated genes and 604 down-regulated genes were found in both aar2-1 and aar2-2 (SI Appendix, Fig. S11A and Dataset S2). Gene ontology enrichment analysis showed that both common hyper-DEGs and hypo-DEGs were enriched in genes involved in responses to stimuli (SI Appendix, Fig. S11B).

Analyses of the RNA-seq data revealed that both aar2-1 and aar2-2 showed global splicing defects, with 512 and 1,159 intron retention events in aar2-1 and aar2-2, respectively. Two examples (AT4G35760 and AT3G56590) are presented in Fig. 7 A and B and SI Appendix, Dataset S3. HYL1 transcript level or intron splicing was not affected in aar2-1 or aar2-2, which further confirmed that the role of AAR2 in miRNA biogenesis is not through modulating HYL1 expression (SI Appendix, Fig. S12 A and B). We further compared the retained introns between aar2 and mac3a mac3b or prl1 prl2, which are mutations in the MAC complex required for both miRNA biogenesis and pre-mRNA splicing (48). No overlap was found between retained introns in aar2 and either mac3a mac3b or prl1 prl2 (Fig. 7C), suggesting that AAR2 and MAC act on different sets of introns. Next, we examined whether the genes

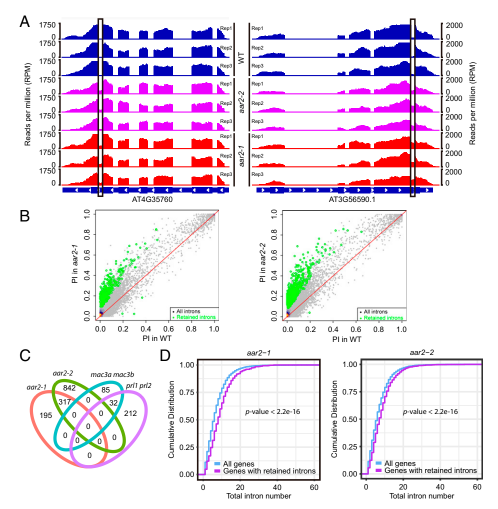


Fig. 7. AAR2 is required for efficient pre-mRNA splicing. (A) Examples of two genes with intron retention defects in the aar2-1 and aar2-2 mutants. RNA-seq reads are shown against the gene models below. In the gene models, exons and introns are marked with rectangles and lines, respectively. The retained introns in the aar2-1 and aar2-2 mutants are marked with black rectangles. (B) Scatterplots showing the percentage of retained introns (PI) in aar2-1 and aar2-2 mutants. The green dots represent introns with statistically significant retention defects in the mutants. (C) Venn diagrams showing the number of retained introns in aar2-1, aar2-2, mac3a mc3b, and prl1 prl2 mutants and the overlap of the introns retained among these mutants. (D) Cumulative density plots of intron numbers in all genes and genes with retained introns in the aar2-1 and aar2-2 mutants. P values were calculated by the Wilcoxon test.

with retained introns in aar2 had any common features. When compared to all genes, the genes with intron retention defects in aar2 tended to have more introns (Fig. 7D).

### **Discussion**

AAR2 is a conserved protein in eukaryotes. Its homologs in yeast and humans globally impact pre-mRNA splicing through a role in U5 snRNP assembly, but AAR2's function in plants has never been characterized. In this study, we show that Arabidopsis AAR2 indeed plays a global role in splicing, with hundreds to thousands of introns being retained in aar2 mutants. We show that Arabidopsis AAR2 is required for miRNA accumulation in addition to its conserved function in splicing. In fact, a global reduction in the levels of 24-nt small RNAs was also found in the aar2-1 mutant, suggesting that AAR2 also promotes siRNA accumulation. Several other splicing factors were also previously shown to impact both miRNA and siRNA accumulation (19, 43, 49), which indicates that mutations in these genes may affect the splicing of a gene(s) involved in siRNA biogenesis and indirectly affect siRNA accumulation. However, it is also possible that these genes are directly involved in Pol IV/RDR2-dependent siRNA biogenesis. We demonstrate that AAR2 promotes miRNA biogenesis through regulating the phosphorylation status of the

microprocessor component HYL1. Because miRNA biogenesis requires double-stranded RNA binding proteins, such as HYL1, TRBP, and DGCR8, in both plants and animals, it is possible that animal AAR2 exerts a similar effect on miRNA biogenesis.

HYL1 is a short-lived protein and is degraded by the cytoplasmic protease HYL1-cleavage subtilase 1 (HCS1) in a lightdependent manner (50). Under light conditions, cytoplasmic constitutive photomorphogenic 1 (COP1) suppresses the activity of HCS1 to prevent HYL1 from degradation, while under dark conditions, the relocation of COP1 to the nucleus releases its repression on HCS1, leading to HYL1 degradation (30, 46, 50, 51). We found that the rate of HYL1 degradation in the dark is slower in aar2 than in the wild type. Furthermore, HYL1 accumulated significantly in the cytoplasmic fraction in aar2 under dark treatment, which is consistent with the current model indicating that HYL1 is degraded in the cytoplasm and suggests that AAR2 is required for HYL1 degradation in the cytoplasm.

The degradation of HYL1 is impacted by its phosphorylation status. Hyper-phosphorylated HYL1 tends to be sequestered in the nucleus and protected from degradation in the cytoplasm in the dark (31). The reduced rate of HYL1 degradation in aar2 mutants in the dark prompted us to examine the phosphorylation status of HYL1 in aar2 mutants. Our results showed that the ratio of nonphosphorylated to phosphorylated HYL1 is significantly

lower in *aar2* even in seedlings grown under long-day conditions. The nonphosphorylated form of HYL1 is the active form in miRNA biogenesis and tends to form D-bodies in the nucleus (27). Thus, the reduction in D-body number and in miRNA accumulation in *aar2* mutants could be explained by defects in HYL1 dephosphorylation. Consistently, the expression of nonphosphorylatable HYL1 partially rescues the miRNA biogenesis defects of *aar2-1*. Taken together, the findings from this study suggest that AAR2 promotes miRNA accumulation through HYL1 dephosphorylation, stability, and localization to D-bodies (*SI Appendix*, Fig. S13).

This study also revealed a previously unknown role of HYL1 in the degradation of pri-miRNAs. In mutants in core microprocessor components, such as dcl1, hyl1, and se, pri-miRNAs accumulate to higher levels as compared to wild-type mutants as a consequence of compromised processing. In aar2 mutants, despite a decrease in the active form of HYL1 and a presumed decrease in pri-miRNA processing, pri-miRNA levels are lower than in wild-type mutants. Moreover, the promoter activities of five examined MIR genes were unaffected in aar2-1 while the corresponding pri-miRNAs were decreased in abundance, which indicates that aar2-1 led to reduced pri-miRNA stability. After MIR gene transcription, both processing by the microprocessor and degradation are expected to contribute to the levels of pri-miRNAs in vivo. In order to examine the effects of aar2-1 on pri-miRNA degradation, we followed the decrease in primiRNA levels in the se-1 background upon the inhibition of transcription to exclude the effects of pri-miRNA processing. Indeed, the aar2-1 mutation led to faster pri-miRNA degradation in the se-1 background. Note that although pri-miRNA levels were reduced in aar2, pri-miRNAs decreased similarly in aar2 and wild-type mutants in the time course of cordycepin treatment (Fig. 4D). One possible explanation is that the processing efficiency of pri-miRNAs is lower in aar2 as compared with wild-type due to the reduced levels of nonphosphorylated HYL1, and the reduced processing would increase the half-life of pri-miRNAs in the assay. Most intriguingly, pri-miRNA levels were similar in the hyl1-2 aar2-1 double mutant and the hyl1-2 single mutant (Fig. 4C), and consistently, the decay of primiRNAs was similar in hyl1-2 aar2-1 and hyl1-2 in the time course of cordycepin treatment (Fig. 4D). We conclude that AAR2 protects pri-miRNAs from HYL1-dependent degradation. This finding, together with those from another study (34), reveal that HYL1 and SE not only promote pri-miRNA processing but also help degrade pri-miRNAs. Perhaps they exert these seemingly opposite functions through distinct interaction partners or at different subcellular locations. Intriguingly, a previous study showed that HYL1 protects pri-miRNAs from degradation by the nuclear exosome (52). Therefore, HYL1 may play opposite roles in primiRNA stability under different conditions or in different forms. Because aar2 mutants accumulate phosphorylated HYL1, we hypothesized that this isoform, so far considered inactive, may have a specific function in promoting pri-miRNA degradation. However, the expression of HYL1 S > D in hyl1-2 did not lead to a reduction in pri-miRNA levels (SI Appendix, Fig. S10F).

Pre-mRNA splicing and miRNA biogenesis can be two independent processes (53). However, several proteins that have been shown to be involved in pre-mRNA splicing also facilitate miRNA biogenesis (17–19, 48). Yeast and mammalian AAR2 are conserved U5 snRNP assembly factors, serving as a component of the cytoplasmic U5 snRNP precursor. Upon import into the nucleus, AAR2 is phosphorylated and replaced by Brr2 to promote the maturation of U5 snRNP (37, 54). We showed that *Arabidopsis* AAR2 is localized in both the cytoplasm and

the nucleus and is required for the splicing of hundreds to thousands of introns. Thus, it is possible that *Arabidopsis* AAR2 functions in U5 snRNP assembly. In *Arabidopsis*, only a subset of pri-miRNAs harbors introns (55, 56). However, miRNA accumulation is indiscriminately affected in *aar2* mutants regardless of whether the pri-miRNAs contain introns or not, which indicates that the miRNA biogenesis function of AAR2 is independent of its function in splicing.

### **Materials and Methods**

Plant materials and growth conditions. All Arabidopsis thaliana strains used in this study are in the Columbia-0 accession. The SUC2::atasi-SUL transgenic line used in the mutagenesis screen is a kind gift from Dr. Detlef Weigel (36). aar2-1 is a new allele isolated from the SUC2::atasi-SUL mutagenesis screening. The T-DNA insertion lines aar2-2 (SALK\_SALK\_117746) and hyl1-2 (SALK\_064863) were obtained from ABRC. se-1 and cpl1-7 were previously described (27, 57). The 35S::HYL1 S > D (seven serine codons [S7, S8, S42, S60, S85, S89, S159] mutated to aspartate) and 35S::HYL1 S > A 7 serine codons ([S7, S8, S42, S60, S85, S89, S159] mutated to alanine) transgenic lines were previously described (31). To generate HYL1-YFP aar2-1, pMIR167a::GUS aar2-1, MIR165A::GFPer aar2-1, aar2-1 was crossed with HYL1-YFP (58), pMIR167a::GUS (43), MIR165A::GFPer, MIR165B::GFPer, MIR166A::GFPer, and MIR166A::GFPer (44), respectively. All primers used for genotyping are listed in SI Appendix, Dataset S4.

Seeds were sown on 1/2 Murashige and Skoog Basal medium containing 1% sucrose, and plants were grown under long-day conditions (16 h light and 8 h dark) at 22  $^{\circ}$ C. The plants were either collected at 14 d for analysis or transferred to soil for phenotype observation. For transient dark treatment, 10-d-old seedlings were grown under long-day conditions and transferred to darkness for 48 h.

Mutagenesis screening and AAR2 identification. The SUC2::atasi-SUL transgenic lines were described previously (36). A transgenic line with a single-locus T-DNA insertion was used for the ethyl methanesulfonate mutagenesis as described previously (36). A mutant with reduced leaf yellowing was isolated and back-crossed with the parental line SUC2::atasi-SUL. In the F2 population, ~100 plants with the SUC2::atasi-SUL aar2-1 phenotype were collected, and genomic DNA was extracted and subjected to library construction. The library was pair-end sequenced on the Illumina platform Hiseq2000 at 50× coverage. A mutation in AT1G66510 (AAR2) was identified and further validated using derived cleaved amplified polymorphic sequences (dCAPS) analysis in the F2 population. Primers for the dCAPS analysis are listed in SI Appendix, Dataset S4.

**DNA constructs and plant transformation.** A 5.2 kb genomic fragment of *AAR2* containing the promoter and coding region without the stop codon was PCR-amplified and cloned into pENTR/D-TOPO (Invitrogen) and then introduced into pGWB40 (59) to generate *AAR2::AAR2-YFP*. For the *AAR2::AAR2-mCherry* construct, the mCherry coding sequence and the *AAR2* genomic fragment were PCR-amplified and recombined into the pCambia1300 vector (Abcam, ab275754). For *35S::AAR2-nYFP*, *AAR2* cDNA was PCR-amplified and inserted downstream of the 35S promoter and fused with nYFP in pXY103 (60). The above constructs were introduced into *aar2-1* by the *Agrobacterium tumefaciens*—mediated floral dipping method (61). The primers used are listed in *SI Appendix*, Dataset S4.

**Small RNA gel blot analysis.** Total RNAs were extracted from 14-d-old seedlings with TRI reagent (Molecular Research Center). Small RNA gel blot analysis was performed as described previously (62). Next, 5 to 10  $\mu$ g total RNA was separated in a 15% urea-PAGE gel and transferred onto a Hybond NX membrane. The membrane was probed by 5'-end  $^{32}$ P-labeled anti-sense DNA oligonucleotides to detect miRNAs or ta-siRNAs. The relative levels of miRNAs or ta-siRNAs were calculated by normalizing against the internal control U6 or total RNA.

**qRT-PCR.** Total RNAs were treated with DNase I (Roche) followed by complementary DNA synthesis using RevertAid reverse transcriptase (Thermo Fisher Scientific) with oligo(dT) primers. qRT-PCR was performed using iQ SYBRGreen Supermix (BioRad) on the BioRad CFX96 system. Primers used are listed in *SI Appendix*, Dataset S4.

RNA stability measurements. RNA stability measurements were carried out as previously described (63). Briefly, 12-d-old seedlings were transferred from 1/2 MS medium agar plates to 12-well plates with 2 mL 1/2 MS liquid medium and incubated overnight to equilibrate the seedlings. Cordycepin was added to a final concentration of 0.6 mM, and seedlings were collected at various time points (0 min, 15 min, 30 min, 60 min, 90 min, 120 min) for RNA extraction. qRT-PCR was performed to determine the levels of pri-miRNAs.

CHX and MG132 treatments. CHX treatment of Arabidopsis seedlings was performed as previously described (30). Briefly, 10-d-old seedlings were treated with 0.5 mM CHX with or without 50  $\mu$ M MG132 for 2 h. Samples were collected and subjected to protein gel blot analysis to determine HYL1 protein levels.

Microscopy and imaging. For GUS staining, whole seedlings from wild-type and aar2-1 seedlings with a homozygous pMIR167a::GUS transgene were incubated with the GUS staining solution (64) at 37 °C in the dark. Tissues were cleared with 70% ethanol before being observed under a stereomicro-

For propidium iodide staining, seedlings were incubated with 10 µM propidium iodide for 10 min and then washed with distilled water. After staining, the roots were excised, mounted in water, and observed under a confocal microscope.

Gel filtration assay. Briefly, 0.5 g 14-d-old AAR2::AAR2-YFP seedlings was ground into fine powder. Next, 1.5 mL 1XPBS supplemented with 1x protease inhibitor mixture (Roche) was added into the powder and the suspension was incubated at 4 °C for 30 min. The suspension was centrifuged at 4 °C for 20 min at 13,200 g and the supernatant was transferred into a new tube. Upon a second centrifugation, the supernatant was collected and filtered through a 0.45 µm filter. Then, 500 µL samples were loaded onto a Superdex 200 Increase 10/300 GL column. Finally, 22 fractions were collected and used for protein detection by protein gel blot analysis.

RNA and small RNA-seq. For small RNA library construction, 50 µg total RNAs were resolved in a 15% urea-PAGE gel and 15- to 40-nt small RNAs were excised from the gel and used as the input for library construction. The libraries from various genotypes and/or replicates were pooled and sequenced on an Illumina HiSEq. 2500 platform. Small RNA-seq data were analyzed with the pRNASeq-Tools pipeline (https://github.com/grubbybio/pRNASeqTools). The raw reads were first trimmed to remove the adapter sequence (AGATCGGAAGAGC) using Cutadapt3.0 (65). The trimmed reads were mapped onto the Arabidopsis thaliana genome Araport11 using Shortstack (66) with the parameters '-bowtie\_m 1000 -ranmax 50 -mmap u -mismatches 0'. Normalization was performed by calculating the reads per million of 45S rRNA reads value. Differentially accumulated small RNAs were analyzed using the R package DESeq2 (67).

For RNA-seq library construction, poly(A) RNAs were isolated from total RNAs using the NEBNext Poly(A) mRNA Magnetic Isolation Module. RNA-seq libraries were prepared using purified poly(A) RNA with the NEBNext Ultra II RNA Library Prep Kit for Illumina and sequenced on an Illumina HiSEq. 2500 platform. Data were analyzed using the pRNASeqTools pipeline (https://github.com/ grubbybio/pRNASeqTools). Briefly, reads were mapped to the Arabidopsis genome Araport11 using STAT v2.6 (68) with default settings. Mapped reads were counted using featureCounts v1.64 (69), and differentially expressed genes were analyzed using the R package DESeq2. The analysis of intron retention in aar2 and WT was performed using SQUID (https://github.com/ sfli001/SQUID). The levels of retained introns were calculated as PI\_Junction: (intron-exon junction reads/[intron-exon junction reads + exon-exon reads]).

Protein gel blot analysis. To determine protein levels, total proteins were extracted from 14-d-old seedlings, resolved in sodium dodecyl-sulfate polyacrylamide gel electrophoresis (SDS) gels, and then transferred to nitrocellulose membranes. The membranes were blocked with 1× PBS supplemented with 5% (wt/vol) nonfat milk and 0.05% Tween 20 and then incubated with primary antibodies (anti-HYL1, 1:2,000, Agrisera, AS06136; anti-cFBPase, 1:5,000, Agrisera, ASO4043; anti-histone H3, 1:3,000, Abcam, ab1791; anti-SE, 1:2,000, Agrisera, AS09532A; anti-GFP, 1:1,000, Sigma-Aldrich, 11814460001; homemade DCL1, 1:2,000) at 4 °C. After 3 washes with 1× PBS supplemented with 0.05% Tween

20, the membranes were incubated with secondary antibodies and then detected with ECL reagent (Amersham, RPN2236). HYL1 phospho-isoforms were detected using phos-tag SDS-PAGE (Fujifilm Wako Pure Chemical Corporation, 190-16721) following the manufacturer's instructions. Briefly, total proteins were separated on a phos-tag SDS-PAGE gel, transferred onto a PVDF membrane using the wet-tank transfer method, and then probed with the HYL1 antibody.

Nuclear-cytoplasmic fractionation. Nuclear-cytoplasmic fractionation was performed as described (70). First, 1 g 14-d-old seedlings was ground into fine powder in liquid nitrogen and the powder was resuspended in 2 mL lysis buffer (20 mM Tris·HCl, pH 7.5; 25% glycerol, 250 mM sucrose, 20 mM KCl, 2.5 mM  $MqCl_2$ , 2 mM EDTA, 5 mM DTT, and 1× protease inhibitor mixture). Then the lysates were filtered through two layers of Miracloth. The flow-through was centrifuged at 1,500 g at 4 °C for 10 min. The supernatant was collected and centrifuged at 10,000 g at 4 °C for 15 min and then was kept as the cytoplasmic fraction. The pellet was washed 6 times with nuclear resuspension buffer (20 mM Tris·HCl, pH 7.5; 2.5 mM MgCl<sub>2</sub>, 25% glycerol, and 0.2% Triton X-100) and then resuspended with 500 µL NRB2 (20 mM Tris·HCl, pH 7.5; 10 mM MgCl<sub>2</sub>, 250 mM sucrose, 0.5% Triton X-100, 5 mM β-mercaptoethanol, and  $1\times$  protease inhibitor mixture) and carefully overlaid on top of 500  $\mu$ L NRB3 (20 mM Tris-HCl, pH 7.5; 10 mM MgCl<sub>2</sub>, 1.7 M sucrose, 0.5% Triton X-100, 5 mM  $\beta$ -mercaptoethanol, and 1 $\times$  protease inhibitor mixture), followed by centrifugation at 16,000 g for 45 min at 4 °C. The nuclear and cytoplasmic fractions were subjected for RNA extraction by TRI reagent (Molecular Research Center). For protein gel blot analysis, the nuclear and cytoplasmic fractions were boiled in 1× SDS sample loading buffer (50 mM Tris-Cl at pH 6.8, 2% SDS, 0.1% bromophenol blue, 10% glycerol, 1% 2-mercaptoethanol) before being resolved in SDS-PAGE gels.

BiFC analysis. BiFC analysis was carried out as previously described (71). HYL1 cDNA fused with an N-terminal fragment of YFP and HYL1, DCL1, and SE cDNA fused with a C-terminal fragment of YFP were previously described (13). The constructs containing the constitutive promoter and gene fusions were transformed into Agrobacterium and coinfiltrated into N. benthamiana leaves. Then, 48 h after infiltration, YFP signals were detected under confocal microscopy (Leica Sp5).

Quantification of chlorophyll content. Chlorophyll content was extracted from 20 mg 4-wk-old Arabidopsis leaves. For each biological replicate, fresh leaves were submerged in 1 mL 95% ethanol and incubated at 80 °C in the dark for 20 min with gentle shaking. Six biological replicates were analyzed for each genotype. The sample was then centrifuged for 30 s at 5,000 g and the supernatant was transferred to a new tube. The absorbance of the supernatant was measured at wavelengths of 664 nm and 648 nm. Total chlorophyll content was calculated using the equations below (72):

```
chlorophyll a (mg/L) = 13.36 * A664 - 5.19 * A648
chlorophyll b (mg/L) = 27.43 * A648 - 8.12 * A664.
Total chlorophyll = chlorophyll a + chlorophyll b
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Data, Materials, and Software Availability. Genomics datasets have been deposited in the National Center for Biotechnology Information Gene Expression Omnibus database (accession numbers GSE201940 and GSE201941) (73,74).

ACKNOWLEDGMENTS. We thank Dr. Detlef Weigel for sharing the SUC2::atasi-SUL transgenic line and Dr. Nicolas Bologna for valuable comments on the manuscript. This work was supported by a grant from the NIH (GM129373) to X.C.

Author affiliations: <sup>a</sup>FAFU-UCR Joint Center for Horticultural Biology & Metabolomics, Haixia Institute of Science and Technology, Fujian Agriculture and Forestry University, Fuzhou, Fujian, 350002, China; <sup>b</sup>Department of Botany and Plant Sciences, Institute for Integrative Genome Biology, University of California, Riverside, CA; 92521; <sup>c</sup>State Key Laboratory of Genetic Engineering and Collaborative Innovation Center of Genetics and Development, Institute of Plant Biology, School of Life Sciences, Fudan University, Shanghai 200438, China; <sup>d</sup>State Key Laboratory for Biology of Plant Diseases and Insect Pests, Institute of Plant Protection, Chinese Academy of Agricultural Sciences, Beijing 100193, China; and <sup>e</sup>Instituto de Agrobiotecnologia del Litoral (CONICET-UNL-FBCB), Santa Fe 3000. Argentina Santa Fe 3000, Argentina

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