Homodimerization of HYL1 ensures the correct selection of cleavage sites in primary miRNA

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ABSTRACT

MicroRNA (miRNA) plays an important role in the control of gene expression. HYPONASTIC LEAVES1 (HYL1) is a double-stranded RNA-binding protein that forms a complex with DICER-LIKE1 (DCL1) and SERRATE (SE) to process primary miRNA (pri-miRNA) into mature miRNA. Although HYL1 has been shown to partner with DCL1 to enhance miRNA accuracy, the mechanism by which HYL1 selects the DCL1-targeted cleavage sites in pri-miRNA has remained unknown. By mutagenesis of HYL1 and analysis of in vivo pri-miRNA processing, we investigated the role of HYL1 in pri-miRNA cleavage. HYL1 forms homodimers in which the residues Gly147 and Leu165 in the dsRBD2 domain are shown to be critical. Disruption of HYL1 homodimerization causes incorrect cleavage at sites in pri-miRNA without interrupting the interaction of HYL1 with DCL1 and accumulation of pri-miRNAs in HYL1/pri-miRNA complexes, leading to a reduction in the efficiency and accuracy of miRNAs that results in strong mutant phenotypes of the plants. HYL1 homodimers may function as a molecular anchor for DCL1 to cleave at a distance from the ssRNA–dsRNA junction in pri-miRNA. These results suggest that HYL1 ensures the correct selection of pri-miRNA cleavage sites through homodimerization and thus contributes to gene silencing and plant development.

INTRODUCTION

MicroRNAs (miRNAs) are ∼21 nucleotide (nt) small RNAs produced from partially paired stem-loop regions of primary miRNA (pri-miRNA) transcripts (1). miRNAs regulate gene expression at the post-transcriptional level by inhibiting the expression of mRNAs bearing fully or partly homologous target sequences (1–3). The roles of many miRNAs in developmental regulation, disease resistance, and biotic and abiotic stress response have been demonstrated (4–6). To further understand the molecular mechanism of miRNA regulatory function, it is important to know how miRNA is processed from pri-miRNA.

In animals, pri-miRNAs are processed into miRNA precursors (pre-miRNAs) in nucleus by an RNaseIII-like enzyme named Drosha (7) that selectively cleaves RNA hairpins bearing a large terminal loop (8). After export to the cytoplasm by Exportin-5 (9, 10), the Drosha homolog Dicer cleaves the pre-miRNAs into the miRNA:miRNA* duplex (11). In plants, DICER-LIKE1 (DCL1; the Arabidopsis homolog of Dicer) cleaves both pri-miRNA and pre-miRNA and generates miRNA in the nucleus (12, 13), and then the miRNAs are exported to the cytoplasm and incorporated in the RNA-induced silencing complex (RISC) for negatively regulation to gene expression through miRNA degradation or translation inhibition (13–18).

DCL1, HYPONASTIC LEAVES1 (HYL1) and SERRATE (SE) are all required for the biogenesis of miRNAs (12, 19–22) and function together in dicing bodies (D-bodies) (23). DCL1 catalyzes the processing of pri-miRNAs and pre-miRNAs, consequently producing mature miRNAs (12), whereas HYL1 and SE act as functional partners of DCL1 in the miRNA biogenesis machinery. In vitro miRNA processing assay and in vivo small RNA deep sequencing show that in the absence of HYL1, DCL1 alone generates more inaccurate miRNAs (19, 24). There are some other new proteins that participate in the biogenesis of miRNAs. DAWDLE, a forkhead-associated domain-containing protein, facilitates DCL1 access to or recognition of pri-miRNAs (25). The cap-binding protein complex (CBC) can interact with the 5′ cap of pri-miRNA and affect the loading of pri-miRNAs to dicing complexes with SE (26). HUA ENHANCER 1 (HEN1) methylates miRNA/miRNA* duplexes, protecting them from degradation (27). MOS2 promotes pri-miRNA processing via facilitating the recruitment of pri-miRNAs by the dicing complexes (28). C-TERMINAL DOMAIN PHOSPHATASE-LIKE 1 (CPL1) is required for HYL1 dephosphorylation, which in turn is essential for accurate miRNA processing and strand selection (29).

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HYL1 contains two double-stranded RNA-binding domains (dsRBDS) at its N-terminal end, a putative protein–protein interaction domain at its C-terminal end, and a nuclear localization signal (NLS) in the middle (30,31). The loss-of-function mutant hyl1 exhibits pleiotropic phenotypes. The two N-terminal dsRBDS of HYL1 are sufficient to rescue the phenotypes by restoring the accumulation of miRNAs (31). HYL1 has been reported to regulate the phase transition, establishment of stamen, and the adaxial-abaxial identity of leaf in Arabidopsis by controlling the biogenesis of different miRNA families (32–34). Recently, the crystal structures of dsRBBD1 and dsRBBD2 of HYL1 have been reported (35). Based on their structures, dsRBBD1 likely binds to RNAs and dsRBBD2 may be responsible for protein interactions. Structural analyses also imply that HYL1 forms a dimer and that this dimerization is probably mediated by the dsRBBD2 domain. However, the function of the dimer remains unknown.

In animals, several double-stranded RNA-binding proteins are involved in the two steps of miRNA processing. In the nucleus, DIGEORGE SYNDROME CRITICAL REGION GENE 8 (DGCR8) protein (also known as Pasha in Drosophila melanogaster) functions as the essential cofactor of Drosha (36–39). In the cytoplasm, TAR-RNA BINDING PROTEIN (TRBP) and LOQUACIOUS (LOQS) act as the partners of Dicer (31,40). TRBP contributes to RNA binding, length determination and the assembly of RISCs (41,42). Moreover, TRBP is required for optimal RNA silencing mediated by siRNAs and endogenous miRNAs, and it facilitates cleavage of pre-miRNA in vitro (43). LOQS is known to change the cleavage site targeted by Dicer to produce miRNAs with target specificities different from those generated by Dicer alone or Dicer bound to alternative protein partners (40).

Although several studies have demonstrated the contributions of HYL1 in the accuracy and efficiency of miRNA processing (19,44), the mechanism by which HYL1 recognizes primary RNA substrates and selects cleavage sites has yet to be elucidated. Here, we report that HYL1 regulates miRNA processing via homodimerization mediated by the dsRBBD2 domain. Two residues, Gly147 and Leu165, play key roles in HYL1 dimerization. Disruption of Gly147 and Leu165 in HYL1 dimerization. (A) BiFC analysis showing homodimerization of HYL1 and HYL1 mutants. (B) Pull-down assay results showing protein–protein interaction of HYL1 mutants with themselves. T146E, HYL1 T146E; G147E, HYL1 G147E; I158E, HYL1I158E; L165E, HYL1L165E; L166E, HYL1L166E. (C) Pull-down assay results showing protein–protein interaction of HYL1 mutants with wild-type HYL1.
HYL1 dimerization is not necessary for direct interaction of HYL1 with DCL1 and SE. (A) BiFC analysis showing protein–protein interaction of HYL1 mutants with DCL1 and SE in protoplasts. (B) Pull-down assay results showing protein–protein interaction of HYL1 mutants with DCL1 and SE tagged with 3×Flag.

**MATERIALS AND METHODS**

**Plant materials and growth conditions**

*Arabidopsis thaliana* wild-type and *hyl1-2* (SALK_064863, Columbia ecotype) and *hyl1-3* (Columbia ecotype) mutants were used in this study. The *hyl1-3* mutants were obtained from Dr Detlef Weigel (29).

Seeds were surface-sterilized in 70% ethanol for 30 s and then in 0.1% HgCl2 for 10 min and washed four times in sterile distilled water. For *in vitro* tissue culture, the seeds were mixed in molten 0.1% water agar and plated on top of solid 1% sugar Murashige and Skoog medium. Plates were sealed with parafilm, incubated at 4°C in darkness for 2 days, and then moved to a growth chamber at 22.8°C for 16 h in light. For phenotypic observation, seeds were sown in pots with peat soil and grown in growth chambers in the same conditions as described above.

**Gene cloning and transgenes**

A *HYL1* promoter region (1240 bp upstream of the translation start site) and a full-length coding sequence (1257 bp) were amplified from Columbia seedlings. Then both sequences were cloned into pCAMBIA1301 binary vectors to obtain the *pHYL1::HYL1* constructs. Site-directed mutagenesis was performed, and the primers used for polymerase chain reactions (PCRs) are listed in Supplementary Table S3.

The *Arabidopsis* plants were transformed using a flower-dip method. For selection of transgenic plants, the seeds were sterilized and germinated on agar medium containing 50 mg/ml hygromycin. Seedlings conferring resistance to the hygromycin were transplanted in a greenhouse and grown at 22°C under an 8-h light regimen. The transgenic plants were shelfed for at least three generations, and the seeds from each plant were harvested separately for subsequent observations.

**RNA analysis**

For northern blotting, RNA samples were extracted using TRIzol reagent (Invitrogen, Carlsbad, CA, USA). For northern blotting of small RNA, 50 µg of total RNA was resolved by 15–19% polyacrylamide gel electrophoresis (PAGE) in 1× TBE at 80 V for 4–6 h and transferred to a Hybond membrane (Amersham Biosciences, GE Healthcare) in 0.5× TBE overnight at 28 mA. The UV cross-linked membrane was hybridized in ULTRAhyb® Ultrasensitive Hybridization buffer (Ambion, Austin, TX, USA) using probes of 3′ biotin-labeled DNA oligos (TaKaRa, Otsu, Japan) antisense to the mature miRNA or U6 transcripts.
RNA gel blot analysis was performed according to the methods of Liu et al. (34).

For real-time PCR, total RNA was treated with DNase I (TaKaRa), followed by a phenol/chloroform extraction to remove contaminating DNA. Approximately 4 μg of purified RNA was used for first-strand complementary DNA (cDNA) synthesis using PrimeScript® Reverse Transcriptase (TaKaRa) with oligo(dT) primers. Real-time PCR was performed using the specific primer pairs (Supplementary Table S3) in the MyiQ2 Two-color Real-time PCR Detection System (Bio-Rad, Hercules, CA, USA). The comparative threshold cycle method was used to determine relative transcript levels. Quantitative PCR for each gene was done with at least three biological replicates.

### Analysis of transcriptional activity

Primary mRNAs of some MIRNA genes that contain introns were analyzed to evaluate transcriptional activity according to the methods of Liu et al. (45). The intron-specific primers and oligo(dT) were used for reverse transcription, and then the intron-specific primers for detection of primary mRNA levels of MIRNA genes (Supplementary Table S3).

### Protein analysis

Anti-Flag (Sigma-Aldrich, St Louis, MO, USA; F3165, 1:5000 dilution), anti-GST (Sigma-Aldrich; 1:5000 dilution), anti-MBP (NEB, 1:3000 dilution) and anti-HYL1 (Agrisera, 1:1000 dilution) antibodies were used for western blotting. Secondary antibodies were goat-developed anti-rabbit IgG (GE Healthcare; NA931V, 1:20 000 dilution) and goat-developed anti-mouse IgG (GE Healthcare; NA934V, 1:50 000 dilution).

### Coimmunoprecipitation

For coimmunoprecipitation assays of HYL1, plants of 10-day-old seedlings were ground into a fine powder using liquid nitrogen and then resuspended in a buffer containing 20 mM Tris-HCl, pH 7.4, 200 mM KCl, 40 mM MgCl2, 1 mM EDTA, 20% glycerol and 0.05% Nonidet P-40. After pre-clearing with protein G-agarose beads for 4 h at 4°C, the extracts were incubated with HYL1 antibodies to protein G-agarose beads for 4 h at 4°C. After five washes (20 mM Tris-HCl, pH 7.9, 0.5 M KCl, 10% glycerol, 1 mM EDTA, 5 mM DTT, 0.5% Nonidet P-40 and 0.2 mM phenylmethylsulfonyl fluoride), the proteins in the immunoprecipitates were subjected to northern and western blot analyses.

### Expression and purification of HYL1 proteins from Escherichia coli

For protein expression and purification, full-length and truncated HYL1 was inserted into the expression vectors pGEX4T-1 and pMAL-C2x, respectively, and the recombinant plasmids were transformed into E. coli DH5α. Bacteria were cultured at 37°C overnight in a 20-ml volume, then diluted to 200 ml with 0.8 mM isopropylthiogalactoside and grown at 37°C for 4 h. Nearly 2 L of cultured bacteria was collected and lysed to generate ultrasonic precipitates, and the precipitates were then resolved in a buffer containing 1% Triton X-100, 20 mM Tris, 20 mM EDTA and 1 mM DTT. The GST-fused proteins were purified with glutathione Sepharose resin (GE Healthcare), whereas the MBP-tagged proteins were purified with Amylose Resin (NEB).

### Expression and purification of recombinant DCL1 and SE proteins from human embryonic kidney 293 cells (HEK293)

The cDNAs of DCL1 and SE with 3×Flag were cloned into pN3 vector (from pEGFP-N3 Clontech, but without an enhanced green fluorescent protein tag). The human embryonic kidney 293 (HEK293) cells were maintained in Dulbecco’s modified Eagle’s medium (Invitrogen) supplemented with 10% fetal bovine serum (FBS) (Gibco) and transfected with the DCL1 and SE vectors by Lipofectamine 2000 (Invitrogen) according to the manufacturer’s protocol. The cells were harvested 48 h after transfection and lysed in lysis buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM EDTA, 1 mM DTT, 0.5% Nonidet P-40, 0.2 μg/ml RNaseA, protease inhibitor cocktail) for 15 min. The lysate was then incubated with Flag resin in lysis buffer overnight at 4°C. After five washes with washing buffer (50 mM Tris-HCl, 150 mM NaCl, pH 7.4, 0.005% NP-40), the Flag-fusion proteins were then eluted with 3×Flag peptide (150 ng/μl) in Tris-buffered saline (TBS) (50 mM Tris-HCl, pH 7.4, 150 mM NaCl).

### In vitro pull-down

For MBP pull-down, MBP-tagged proteins were bound to amylose resin (NEB) in binding buffer containing 25 mM Tris, pH 7.4, 1 mM EDTA, 0.01% NP-40 and 2 mM NaCl, and incubated with GST-tagged proteins overnight at 4°C. Then the resin was washed 10 times in the binding buffer and eluted by boiling in sodium dodecyl sulfate (SDS)-PAGE loading dye. Aliquots of eluents (20 μl) were resolved on SDS-PAGE gels for immunoblotting with GST antibody.

### Yeast two-hybrid assays

Yeast transformation, mating and cDNA library preparation were performed according to the Matchmaker™ Gold Yeast Two-Hybrid System User Manual (Clontech). For the initial screen, a pGBKT7+ bait was transformed into Y2H Gold (Clontech) and then mated with the pistil library. Matin mixtures were screened on yeast media lacking -Leu-Trp-Ade+3AT. Inserts were amplified from yeast cultures using colony PCR followed by direct sequencing. The fragments detected were fused in frame with the GAL4 DNA binding domain (pGBK7T) or activation domain (pGAD7). All the recombinant plasmid pairs were cotransformed into the yeast strain AH109. Yeast transformation was performed following the manufacturer’s instructions (Clontech Yeast Protocols Handbook). The cotransformed yeast clones were first grown on SD/-Leu-/Trp medium and subsequently plated on SD/-Ade/-His/-Leu/-Trp medium.
**RESULTS**

**HYL1 forms a homodimer by dsRBD2 domain**

HYL1 acts as an important partner of DCL1 within the D-body. To gain insight into the components of the D-body necessary for accurate miRNA processing, we carried out a yeast two-hybrid screening of HYL1 to investigate the unidentified factors involved in the accuracy of miRNA processing. Several proteins including SE and some functionally unknown proteins were found to interact with HYL1 (Supplementary Table S1). Interestingly, we identified protein–protein interaction of HYL1 with itself, indicating that HYL1 forms a homodimer.

The two dsRBDs, putative protein–protein interaction domain and NLS of HYL1 are indicated in Figure 1A. To determine whether HYL1 forms a homodimer in vitro, we used full-length and truncated HYL1 for a yeast two-hybrid assay. We observed a direct interaction between two HYL1 molecules (Figure 1B). The dsRBD2 fragment interacted with full-length HYL1, whereas the dsRBD1 fragment did not. Moreover, one dsRBD2 fragment interacted with another dsRBD2 but not with dsRBD1. These findings indicate that HYL1 forms a homodimer through the dsRBD2 domains.

To further examine whether HYL1 homodimerizes through dsRBD2, we performed pull-down assays. As expected, MBP-HYL1 pulled down GST-HYL1, and MBP-dsRBD2 pulled down GST-dsRBD2, but MBP-dsRBD1 failed to pull down GST-dsRBD1 (Figure 1C; Supplementary Figure S1). In addition, weak interaction signals were seen between dsRBD1 and dsRBD2. These results confirm that HYL1 dimerizes through the dsRBD2 domains.

To determine which domain is necessary for the interactions of HYL1 with SE and DCL1, we performed yeast two-hybrid assay of truncated HYL1 with SE and DCL1. The dsRBD2 domain rather than dsRBD1 domain interacted with SE and DCL1 (Supplementary Figure S2). Thus, we conclude that the dsRBD2 domain is the core of the HYL1 homodimerization and protein interactions.

**Gly147 and Leu165 are required for HYL1 dimerization**

To identify the residues required for homodimerization of HYL1, we accomplished an extensive mutagenesis analysis. Based on the structure of the HYL1 dsRBD2, we selected seven amino acids within the dsRBD2 domain to generate different HYL1 mutations (Supplementary Table S2). Because they are located in the exposed regions that could be involved in HYL1 dimerization. Eight HYL1 mutations were generated and each HYL1 mutation was expected to potentially impair HYL1 dimerization. In addition to these mutations, three other mutations that may impair the interaction with SE and DCL1 were generated.

Bimolecular fluorescence complementation (BiFC) was used to evaluate the effects of the point mutations on HYL1 dimerization. The same HYL1 mutants were fused with YFP at either the N- or C-terminus and introduced into the Arabidopsis protoplast for transient expression. The HYL1 mutants were visualized via confocal microscopy. Fluorescently labeled HYL1 mutants during the formation of BiFC complexes showed that the fluorescence intensity...
for HYL1G147E (G147E) or HYL1L165E (L165E) was significantly diminished, revealing that the homodimerization of these mutants is weakened (Figure 2A). By contrast, HYL1C122R (C122R), HYL1R130E (R130E), HYL1T146R (T146R), HYL1I158E (I158E), HYL1R162E (R162E) and HYL1L166E (L166E) showed almost the same fluorescence intensity as the wild-type HYL1 did, meaning that they also formed homodimers (Supplementary Figure S3A). Weaker fluorescence intensity was also found when G147E and L165E were cotransformed with wild-type HYL1. To exclude the possibility of transient transfer efficiency, western blotting was carried out to detect the mutated HYL1 fused with YFP. All five HYL1 mutants examined showed almost the same intensity of YFP (Supplementary Figure S4), demonstrating that the difference in transfer efficiency in the BiFC analysis was not significant. Therefore, we conclude that Gly147 and Leu165 are required for HYL1 dimerization. Using this method, we also examined the effects of the other three mutations (R151E, K154E and R162E) on their interaction with SE and DCL1, but found no difference from wild-type HYL1 (data not shown).

To confirm these findings, we used in vitro pull-down experiments to examine the associations of the HYL1 mutants. MBP-HYL1 pulled down GST-HYL1 as expected (Figure 2B). However, MBP-G147E and MBP-L165E pulled down much less GST-G147E and GST-L165E, respectively, and MBP-G147E and MBP-L165E pulled down less GST-HYL1 (Figure 2C). These results provide evidence that the Gly147 and Leu165 mutations impaired HYL1 dimerization.

**HYL1 dimerization is not necessary for direct interaction with DCL1 and SE**

HYL1 is known to interact with DCL1 and SE, but whether non-dimerized HYL1 interacts with DCL1 and SE remains unknown. Therefore, we used BiFC analysis to determine the effects of the point mutations on their interactions with DCL1 and SE. In Arabidopsis protoplasts, all transiently expressed HYL1 mutants, including G147E and L165E, were found to interact with DCL1 and SE, implying that a deficiency in HYL1 dimerization did not affect protein–protein interactions of HYL1 with DCL1 and SE (Figure 3A; Supplementary Figure S3B).

To further test whether disruption of HYL1 homodimerization prevents the direct interaction of HYL1 with SE and DCL1 in vitro, we used pull-down assays to observe the interaction of G147E and L165E with DCL1 and SE. HYL1 was purified from recombinant proteins expressed in Escherichia coli as described previously (31). Recombinant DCL1 and SE were poorly expressed in E. coli but could be expressed in mammalian cells. Therefore, we obtained the recombinant DCL1 and SE proteins using HEK293 cells. DCL1 and SE tagged with 3×Flag were purified by Flag resin and then confirmed by Coomassie Blue staining (Supplementary Figure S5A). Western analysis showed that the strength of interactions between G147E and DCL1 and between L165E and DCL1 did not differ from that between HYL1 and DCL1 (Figure 3B). Similarly, the strength of interactions between G147E and SE and between L165E and SE was equal to that between HYL1 and SE. These results suggest that HYL1 homodimerization does not interfere with the direct interaction of HYL1 with DCL1 and SE.

**HYL1 homodimer is essential for normal plant development**

To examine the biological functions of HYL1 homodimers, we constructed eight mutations of HYL1 under the control of the native promoter and introduced them into the null hyl1-2 mutants. G147E and L165E did not rescue the phenotypes of the hyl1-2 mutants (Figure 4A), but T146E, I158E and the other mutations completely rescued the hyl1-2 mutant phenotype in terms of leaf incurvature. These results indicated that T146E and I158E retained normal function, whereas G147E and L165E were deficient in some aspects of plant development. RT-PCR and western blotting showed that all HYL1 mutants in hyl1-2 plants were expressed equally at both the transcriptional and translational levels (Figure 4B and C).
Figure 5. Accumulation of some miRNAs and expression of their targets in the plants deficient in HYL1 homodimerization. (A) Northern blot showing miRNA accumulation in the hyl1-2 mutants and the transgenic plants. (B) The expression levels of some miRNA-targeted genes in the hyl1-2 mutants and the transgenic plants. (C) Detection of HYL1 homodimers in the hyl1-3 mutants. HYL1 antibody was used to detect HYL1 from the total protein extract of the hyl1-3 plants using the buffer without addition of SDS and urea.

Impairment of HYL1 dimerization disrupts HYL1 function in miRNA biogenesis and miRNA-guided gene silencing

To examine whether the deficiency in HYL1 dimerization affects miRNA biogenesis, we performed northern blotting to detect the accumulation of miRNAs in the hyl1-2 mutants and transgenic plants with HYL1 mutations. In the hyl1-2;G147E, hyl1-2;L165E plants, the accumulation of miR156, miR166, miR172 and miR319 was much less than that in wild-type plants (Figure 5A). By contrast, miR156-targeted SPL9 and miR165/6-targeted REV genes were up-regulated in these plants (Figure 5B). Moreover, the accumulation of these miRNAs in hyl1-2;T146E, hyl1-2;I158E and hyl1-2;L166E was much greater than that in hyl1-2, whereas the target genes SPL9 and REV were down-regulated to the levels almost equal to those of the wild-type.

To examine whether the endogenous G147E in the hyl1-3 plants influences homodimerization as exogenous G147E, we used HYL1 antibody to detect the possible dimerization of G147E in the extracts of Arabidopsis under native condition. As expected, we were able to detect HYL1 homodimer in the wild-type and hyl1-2;I158E plants. (Figure 5C). In the hyl1-3 plants, HYL1 monomer rather than homodimer was detected. This result indicated that endogenous G147E in the hyl1-3 plants impaired homodimerization in vivo. In the hyl1-3 mutants, the expression levels of HYL1 gene and HYL1 protein were not different from the wild type (Figure 6A and B); the accumulation of the miRNAs examined was much less than that in wild type, though it was slightly higher than that in hyl1-2 (Figure 6C), and the genes targeted by these miRNAs were up-regulated (Figure 6D). These results suggest that impairment of HYL1 dimerization affects the biological function of HYL1 in miRNA biogenesis and miRNA-guided gene silencing.
Figure 7. In vitro processing of pri-miRNA to mature miRNA. (A) Pri-miR166g processing using the immunoprecipitates of the transgenic plants expressing HYL1 mutants. (B) In vitro pri-miRNA processing using pri-miR166g and pri-miR167b as the substrates. 146, HYL1T146E; 147, HYL1G147E; 158, HYL1I158E; 166, HYL1L166E. (C) Pri-miR166g and pri-miR167b processing rates of HYL1 mutants. (D) Effects of HYL1 mutants on pri-miR166g processing. −HYL1, absence of HYL1 in the processing complex; +HYL1, HYL1 presence; +G147E, HYL1G147E presence; +I158E, HYL1I158E presence; +L165E, HYL1L165E presence. (E) Distribution of the sequenced small RNAs generated from in vitro processing reactions of different HYL1 mutants.

Dimerization of HYL1 does not affect the accumulation of pri-miRNAs in HYL1/pri-miRNA complexes

To find out how HYL1 regulates the processing of miRNAs through dimerization, we used real-time PCR analysis to examine the accumulation of pri-miRNAs. In the hyl1-3 seedlings, the expression levels of pri-miR160a, pri-miR166a, pri-miR166b, pri-miR168a and pri-miR319a were much greater than those in the wild-type seedlings (Figure 6E). To exclude the possibility that the endogenous GI47E affects the transcriptional activities of MIRNA genes, we targeted the splicing sites (exon–intron junctions) of MIR160a, MIR166a and MIR166b and detected the levels of their unspliced primary transcripts by qPCR using
Figure 8. Correct selection of pri-miRNA cleavage sites in vivo. (A) Cleavage sites on pri-miR168a using primer 321 (an internal primer of the two sequence-specific primers upstream of miR168a*) and primer 74 (an internal primer of the two sequence-specific primers downstream of miR168a*). Arrowheads indicate cleavage sites, and numbers above or below arrowheads list the number of nucleotides between the sites and the first nucleotides of mature miRNA. The numbers in open boxes are the percentages of correct and incorrect cleavage sites. Pearson’s chi-squared test ($\chi^2$) is applied to percentages of correct cleavage sites to evaluate significance of difference between the wild-type, hyfl-2 and hyfl-3 plants.
HYL1 dimerization is necessary for miRNA accuracy

HYL1 has been reported to enhance the accuracy of mature miRNAs (19), and thus, we examined whether the miRNA accuracy is regulated by HYL1 dimerization within in vitro miRNA processing system. Equal amounts of the individual HYL1 mutants were added to each miRNA processing system, which was then incubated with DCL1, SE and pri-miR166g. The small RNAs generated from the pri-miRNAs were cloned, and the resultant small RNA libraries were constructed for sequencing. According to the extent of miRNA accuracy, small RNAs were assigned to three classes: accurate (mature miR166 or miR166*), partially accurate (position shifting by 2 nt or less) and inaccurate (position shifting by more than 2 nt).

Effects of HYL1 absence on miRNA accuracy were obvious. The inaccuracy of miR166 for DCL1+SE complexes was 56% (Figure 7D), which is four times higher than that for DCL1+HYL1+SE complexes. A large proportion of inaccurate miR166 for DCL1+SE complexes were derived from the regions beyond the stem of pre-miRNA (Figure 7E), in contrast with those for DCL1+HYL1+SE complexes in that they were derived from pre-miRNA regions. Like the wild-type HYL1, I158E showed low inaccuracy in miR166 and miR166*. By contrast, G147E and L165E displayed greater inaccuracy than the wild-type HYL1, and their inaccuracy was elevated according to the degree at which HYL1 was absent. The inaccurate small RNAs aris-
HYL1 homodimers contribute to correct selection of pri-miRNA cleavage sites by DCL1

To understand how HYL1 homodimerization affects pri-miRNA cleavage, we used 5′ RACE PCR analysis to detect the cleavage sites in vivo. In wild-type plants, primers 321 and 43 generated the correct cleavage sites at the ends of miRNA* regions on pri-miR168a and pri-miR166g, respectively (Figure 8A and B). In the hyl1-2 and hyl1-3 plants, percentages of the correct cleavage sites generated with these primers were much lower than in wild-type plants. These results show that disruption of HYL1 homodimerization impairs the correct selection of pri-miRNA cleavage sites.

Aberrance of cleavage sites in pri-miRNAs causes the change in sequence of pre-miRNAs, probably leading to the production of inaccurate miRNAs. To detect the cleavage sites at the 3′ ends of miRNA regions, we designed other primers specific for pri-miR168a. Primer 74 generated the correct cleavage sites at the 3′ ends of miR168 regions in the wild-type plants. In hyl1-2 and hyl1-3 plants, percentages of the correct cleavage sites with this primer were much lower than that of wild-type plants, corresponding to the significant increase in percentages of the incorrect cleavage sites. This result reveals that disruption of HYL1 dimerization impairs the correct cleavage site selection at the 3′ ends of miRNA regions.

The secondary structure of a pri-miRNA consists of four regions: 5′ and 3′ single-stranded RNAs (ssRNAs), double-stranded RNA (dsRNA; a stem bearing mature miRNA and miRNA*) and loop. There is a junction between ssRNA and dsRNA. In wild-type plants, primers 321 and 43 generated the correct cleavage sites 15 and 17 nt away from the ssRNA–dsRNA junction, respectively, while primer 74 did 15+21 nt away from the junction (Figure 8A and B). By contrast, these primers produced many incorrect cleavage sites in the hyl1-2 and hyl1-3 mutants. The incorrect cleavage sites of hyl1-2 plants were mainly on the stem and in regions outside the stem, whereas those of the hyl1-3 plants were mainly near the 5′ and 3′ ends of miRNA regions with different distances (1–5 nt shift). These findings reveal that disruption of HYL1 homodimerization increases the incorrect cleavage sites mainly in the dsRNA regions, resulting in the nucleotide shift near the 5′ and 3′ ends of miRNA regions.

DISCUSSION

HYL1 functions through homodimerization

miRNA accuracy is an important factor in miRNA-directed gene silencing. HYL1 has been considered to play an essential role in miRNA accuracy, and we wondered whether some novel proteins help HYL1 accomplish this role. Yeast two-hybrid screening revealed that HYL1 functions in the form of a dimer in miRNA processing. In vitro pull-down assays confirmed the homodimerization of HYL1. From the crystal structure of HYL1, Yang et al. (35) proposed that dsRBD2 harbors a putative dimerization interface. This dimerization interface is primarily located between the β1 strand of one HYL1 dsRBD2 molecule and the β3 strand of its symmetric related molecule in a parallel interaction mode. In another work, however, dsRBD2 behaves exclusively as a monomer in solution (46). We found that HYL1 proteins form homodimers through the dsRBD2 domains. This finding is consistent with the results of Yang et al. (35). In their work, eight triple mutants with point mutations in the dsRBD2 region yielded misfolded proteins or no protein expression at all. We identified two point mutations that disrupted dimerization. Gly147 and Leu165 are critical for HYL1 homodimerization. In our BiFC experiments, the homodimerization of G147E/HYL1 or L165E/HYL1 was hardly detectable compared with that of HYL1/HYL1. In pull-down assays, however, the homodimerization of G147E/HYL1 or L165E/HYL1 was relatively weaker than that of HYL1/HYL1. This indicates that the strength of HYL1 homodimerization with its mutants is different under the in vivo and in vitro conditions.

In animals, Drosha and DGCR8 form a complex termed the Microprocessor to cleave the pri-miRNAs into pre-miRNAs (37–39). The RNA-binding protein DGCR8 is a heme-binding protein and the association with heme promotes dimerization of DGCR8 (47,48). The heme-bound DGCR8 dimer seems to be more active in triggering pri-miRNA cleavage in vitro. Unfortunately, homodimerization of DGCR8 had not been identified in vivo, and its putative role in pri-miRNA processing not defined. Interestingly, HYL1 proteins are able to form homodimers in plants. This reminds us that HYL1 may have the same function as DGCR8 in pri-miRNA processing.

HYL1 homodimers are crucial for correct selection of cleavage sites in pri-miRNA

HYL1-directed miRNA biogenesis is generally divided into five biological steps: (i) interaction of HYL1 with DCL1 and SE, (ii) binding of HYL1 to pri-miRNA, (iii) correct selection of pri-miRNA cleavage sites, (iv) efficient processing of pri-miRNA and (v) maturation of accurate miRNA. The absence of HYL1 in the hyl1-2 mutants actually affects all of these steps because the D-body is aberrant. Unlike hyl1-2 mutants, however, disruption of HYL1 homodimerization in the hyl1-3 allele (or G147E and L165E mutants) causes the incorrect selection of pri-miRNA cleavage sites. This disruption neither interrupts the interaction of HYL1 with DCL1 and SE nor decreases HYL1 binding affinity to pri-miRNA. In this sense, the correct selection of cleavage sites is critical for miRNA accuracy.

Exogenous G147E and L165E generated lower levels of miRNAs than the wild-type HYL1, and endogenous G147E in the hyl1-3 plants led to deficient miRNA biogenesis as well. Either endogenous or exogenous G147E causes the mutant phenotypes because of miRNA inaccuracy. The importance of Gly147 and Leu165 for plant phenotype reveals that HYL1 homodimerization is essential for miRNA accuracy and plant development.
Pri-miRNA processing is an early and critical stage of miRNA biogenesis as it defines the sequence of premiRNAs and mature miRNAs by generating one end of the molecule. In this study, the strand-specific primers uncovered the first cleavage sites of pri-miR168a and pri-miR166g at the 3' ends of miR168* and miR166, respectively. In animal cells, a critical step in human miRNA maturation is the processing of pri-miRNA transcripts by the nuclear RNaseIII enzyme Drosha to generate pre-miRNA (7,49). Drosha selectively cleaves RNA hairpins bearing a large terminal loop, with cleavage sites largely determined by distance along the RNA (8). In addition, LOQs can change the cleavage sites targeted by Dicer, leading to the production of miRNAs with target specificities different from those generated by Dicer alone or Dicer bound to other protein partners (40).

HYL1 homodimers ensure the correct distance from ssRNA–dsRNA junction in pri-miRNA

Aberrance of cleavage site selection in pri-miRNAs can cause changes in the length and/or sequences of pre-miRNAs, leading to the inaccuracy in miRNAs. In animal cells, Drosha – DGCR8 complex cleaves the pri-miRNAs at the sites ~11 bp away from the ssRNA–dsRNA junction (50). In plants, an imperfectly paired stem of ~15 bp below the miRNA:miRNA* duplex is considered to be a key element for the initial pri-miRNA cleavage (51,52). The secondary structure of pri-miRNAs influences the recognition and cleavage pattern during their processing (53). Recently, one report mentioned that DCL1 complexes cleaved premiRNAs at 16–17 bp away from reference ssRNA–dsRNA junctions for either canonical or noncanonical processing (54). Thus, a ‘distance mechanism’ seems to ensure the correct selection of cleavage sites on pri-miRNAs. Our results provide evidence that the correct cleavage at the 5' end of miRNA region in wild-type plants occurs just 15 or 17 nt away from the ssRNA–dsRNA junction. In the hyl1-3 mutants, the correct cleavage sites at the ends of miRNA regions decrease whereas the incorrect ones in dsRNA regions increase. By contrast, a large proportion of the incorrect cleavage sites in the hyl1-2 plants exist in the ssRNA regions, meaning that the absence of HYL1 increase the incorrect cleavage sites in ssRNA regions. This fact suggests that the distance of cleavage sites from ssRNA–dsRNA junction depends on dimerization of HYL1. Possibly, HYL1 homodimers act as a ruler to recognize the ssRNA–dsRNA junction and to direct DCL1 to cleave 15–17 nt away from the junction. The model supposed in Figure 9 suggests that disruption of HYL1 homodimerization impairs the correct selection of cleavage sites in pri-miRNA because the distance from ssRNA–dsRNA junction is altered. The mechanisms by which the correct selection of cleavage sites occurs in pri- and/or pre-miRNA have remained a mystery. Nevertheless, our results indicate that this process may be dependent on HYL1 binding to the correct sites on pri-miRNA or pre-miRNAs. Further study on the binding sites of HYL1 on pri-miRNAs and pre-miRNAs will facilitate an improved understanding of the molecular mechanisms underlying miRNA biogenesis.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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REFERENCES


