A fungal effector suppresses the nuclear export of AGO1–miRNA complex to promote infection in plants

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Communication between interacting organisms via bioactive molecules is widespread in nature and plays key roles in diverse biological processes. Small RNAs (sRNAs) can travel between host plants and filamentous pathogens to trigger transkingdom RNA interference (RNAi) in recipient cells and modulate plant defense and pathogen virulence. However, how fungal pathogens counteract transkingdom antifungal RNAi has rarely been reported. Here we show that a secretory protein VdSSR1 (secretory silencing repressor 1) from V. dahliae, a soil-borne phytopathogenic fungus that causes wilt diseases in a wide range of plant hosts, is required for fungal virulence in plants. VdSSR1 can translocate to plant nucleus and serve as a general suppressor of antifungal RNAi. Our findings reveal a mechanism by which phytopathogenic fungi antagonize antifungal RNAi-dependent plant immunity and expand the understanding on the complex interaction between host and filamentous pathogens.

Significance
Increasing evidence demonstrates that small RNAs can serve as trafficking effectors to mediate bidirectional transkingdom RNA interference (RNAi) in interacting organisms, including plant–pathogenic fungi systems. Previous findings demonstrated that plants can send microRNAs (miRNAs) to fungal pathogen V. dahliae to trigger antifungal RNAi. Here we report that V. dahliae is able to secret an effector to the plant nucleus to interfere with the nuclear export of AGO1–miRNA complexes, leading to an inhibition in antifungal RNAi and increased virulence in plants. Thus, we reveal an antagonistic mechanism that can be exploited by fungal pathogens to counteract antifungal RNAi immunity via manipulation of plant small RNA function.
EVs and are believed to silence *Phytophthora* genes during natural infection (18). However, whether such a suppressive phenomenon is widespread in the transkingdom RNAi between plant and filamentous pathogens and whether there are alternative mechanisms are still unclear.

Here we demonstrated that the secretory protein VdSSR1 from *V. dahiae*, a causal agent of plant wilt diseases, is a general suppressor of RNA silencing in plant cells. VdSSR1 sequencers ALY family proteins, the adaptors of the TREX (Transcription-Export) miRNA transport complex, to interfere with TREX/TREX-2 complexes–mediated nuclear export of AGO1–miRNA complex and mRNAs, leading to a reduced accumulation of transkingdom miR159 and miR166 in fungal cells and increased expression of fungal virulence genes. Our findings identify a mechanism by which phytopathogenic fungi antagonize RNA silencing-dependent plant immunity and advance our understanding of the arms race between plants and phytopathogenic fungi.

**Results**

The *V. dahiae* Secretory Protein VdSSR1 Can Translocate to the Plant Nucleus. Recent studies reported that plants can send small RNAs to interacting organisms, including phytopathogenic fungi, to trigger transkingdom RNAi (8, 9). In the interacting system between plant and *V. dahiae*, a causal agent of plant wilt diseases that cause very serious harm to numerous crops, miR159 and miR166 can be sent to fungal cells to trigger transkingdom RNAi and antagonize fungal virulence. We are interested in the question of whether fungi can send effector to plant cells to antagonize this process. It is well known that miRNAs are processed mainly in nucleus. To answer this question, we first performed mass spectrometry (MS) assay in the culture filtrates of a virulent defoliating *V. dahiae* isolate V592 from cotton and identified potential effectors from the MS data that can be located in plant nuclei and possess silencing suppression activity in plants. In this way, we identified that a 247-amino acid protein VdSSR1, which contains a conserved RRM_Aly_REF_like domain (cd12418) (19) and a C/NLS motif (Fig. 1A and Dataset S1), is a potential candidate. VdSSR1 is expressed in both mycelia and spores and displays a higher expression in spores (*SI Appendix, Fig. S1*). VdSSR1 is present in both the nucleus and cytoplasm of fungal cells but localized mainly in the nucleus of plant cells (Fig. 1B). VdSSR1 mainly localizes to the cytoplasm in fungal cells (Fig. 1B). VdSSR1 is encoded by gene VDAG_04215. According to the annotation of the Fungal Secretome Database, VDAG_04215 in VdLs. 17 strain is a class NS secreted protein that lacks a typical signal peptide ([http://fsl.sn.u.ac.kr/cgi-bin/pv?protein_dv&bid=533&ov=VDAG_04215&ovf=SEQUENCE_NAME](http://fsl.sn.u.ac.kr/cgi-bin/pv?protein_dv&bid=533&ov=VDAG_04215&ovf=SEQUENCE_NAME)).

To test whether VdSSR1 can be secreted into plant cells during infection, we first examined the accumulation of VdSSR1 protein in the culture filtrates of VdSSR1-GFP–expressing V592 strain via Western blotting as previously described (20). Culture filtrates from the strains expressing free GFP and VdPDA1-GFP, a characterized secretory protein of *V. dahiae* (21), were also used for nonsecretory and secretory controls. As shown in Fig. 1C, similar to VdPDA1-GFP, VdSSR1-GFP protein was detected in the culture filtrates of VdSSR1-GFP strain but not in that of V592, suggesting that VdSSR1-GFP protein can be secreted into the extracellular space from fungal cells. To confirm the secretion, a cellophane penetration assay was performed as previously reported (22). As shown in Fig. 1D, both VdSSR1-GFP and VdPDA1-GFP can penetrate into the cellophane and were observed in the infectious structure, which also functions as a secretory structure (22). Considering that VdSSR1 lacks a typical signal peptide, we asked whether it is secreted via EVs. To answer this question, EVs were isolated from the VdSSR1-GFP–expressing strain. Unfortunately, we cannot detect the accumulation of VdSSR1 (*SI Appendix, Fig. S2*). We speculate that VdSSR1 may be secreted through an unconventional method as reported previously (23). To further verify that VdSSR1 could be translocated from *V. dahiae* to plant cells, conidial spores of VdSSR1-GFP and free GFP strain were inoculated on onion epidermal cells. At 7 d postinoculation (dpi), a nuclear fluorescence signal was observed in the nucleus of onion epidermal cells inoculated with VdSSR1-GFP conidial spores (Fig. 1E). In contrast, in the free GFP strain–inoculated cells, fluorescence was only observed in conidial spores (Fig. 1F). Combined with these data, we concluded that VdSSR1 could translocate from *V. dahiae* to the plant nucleus.

**VdSSR1 Is Required for *V. dahiae* Virulence in Host Plants.** To investigate whether VdSSR1 has RNA silencing suppressor activity, VdSSR1 was transiently coexpressed with the reporter gene GFP in *Nicotiana benthamiana* leaves. The reported silencing suppressors 2b (26) from cucumber mosaic virus (CMV-2b)
and P19 from tomato bushy stunt virus (TBSV) served as positive controls, and β-glucuronidase (GUS) protein served as a parallel control. As shown in Fig. 3A, compared with the GUS control, dramatically enhanced GFP fluorescence signals were observed in VdSSR1-, CMV-2b-, and P19-infiltrated leaf regions. Consistently, greatly increased levels of GFP mRNA and protein were detected (Fig. 3B). These results suggested that VdSSR1 could suppress sense exogenous gene-triggered RNA silencing. To further characterize the suppressor activity of VdSSR1, the Myc-VdSSR1 transgene was crossed into a SUC:SUL silencing reporter system (Suli) (27). In this system, an inverted repeat of the endogenous SULFUR (SUL) gene is expressed in phloem companion cells and leads to the production of siRNAs (siSULs) and the ensuing RNA silencing, resulting in a leaf bleaching phenotype in the silenced cells (28). The results demonstrated that the expression of VdSSR1 greatly inhibited the spread of SUL silencing (Fig. 3C) and fully rescued the reduced levels of SUL mRNA in Suli plants (Fig. 3D). Moreover, the siSUL levels were obviously reduced in VdSSR1-expressing plants compared with Suli plants (Fig. 3E). Taken together, these data demonstrated that VdSSR1 not only suppressed sense exogenous gene-triggered silencing but also interfered with inverted repeat dsRNA-mediated RNA silencing. Consistent with the RNA silencing suppressor activity, VdSSR1 plants exhibited higher sensitivity to CMV infection (SI Appendix, Fig. S5).

We next asked whether nucleus targeting is indispensable for VdSSR1-dependent silencing suppression and fungal virulence in plants. To this end, NLS deletion protein VdSSR1Δnls and VdSSR1Δnls-expressing strain in Δvdssr1 background were generated, and silencing suppression and fungal inoculation assays were performed. As shown in SI Appendix, Fig. S6A, NLS deletion greatly reduced the nuclear accumulation of VdSSR1 but dramatically increased its cytoplasmic accumulation. The RNA silencing suppression activity of VdSSR1Δnls was almost lost (SI Appendix, Fig. S6B and C), and expression of VdSSR1Δnls cannot rescue the reduced virulence of Δvdssr1 strain (SI Appendix, Fig. S6D). These results indicated that nucleus targeting is indispensable for VdSSR1-dependent silencing suppressor activity and fungal virulence in plants.
VdSSR1 Suppresses Transkingdom RNAi in the Plant–V. dahliae System. In some plant–pathogenic fungus systems, including the plant–V. dahliae system, small RNAs can be trafficked bidirectionally to guide transkingdom RNA silencing, resulting in the modulation of plant immunity and fungal virulence (2, 6, 8, 9). Cotton and Arabidopsis plants have been shown to export miR159 and miR166 to V. dahliae cells to trigger silencing of the fungal genes Vdhic-15 and VdClp-1, respectively, thereby antagonizing the virulence of V. dahliae (8). To test whether VdSSR1 plays a role in the transkingdom silencing mediated by plant trafficking miRNAs, we examined the accumulation of plant miR159 and miR166 in V. dahliae cultured from hyphae recovered from infected cotton plants (V. dahliae-Cotton) as previously reported (8). Consistent with previous findings, RNA blotting results demonstrated that trafficking miR159 and miR166 were detected in the cultured V. dahliae recovered from the infected cotton plant (V592-Cotton) but not in the uninfected V592 (Fig. 3F). Intriguingly, much higher levels of trafficking miR159 and miR166 were detected in Δvdssr1-Cotton compared with V592-Cotton (Fig. 3F). In line with the increased accumulation of trafficking miRNAs, the levels of Vdhic-15 and VdClp-1 mRNAs, the targets of plant miR159 and miR166, respectively, were significantly reduced in Δvdssr1-Cotton (Fig. 3G), demonstrating that transkingdom RNA silencing was enhanced by the knockout of VdSSR1. Importantly, no significant difference in Vdhic-15 and VdClp-1 mRNA levels was detected between the uninfected V592 and Δvdssr1 strains (Fig. 3G), implying that the down-regulation of Vdhic-15 and VdClp-1 mRNA levels observed in Δvdssr1-Cotton was attributed to the increased accumulation of trafficking miRNAs but not a direct effect of VdSSR1 knockout on these two genes. Altogether, these data demonstrated that VdSSR1 could suppress the transkingdom RNAi of fungal virulence genes by inhibiting the accumulation of trafficking plant miRNAs in recipient cells.

VdSSR1 Affects the Nucleocytoplasmic Partitioning of sRNAs in Plants. To decipher the mechanism underlying VdSSR1-mediated silencing suppression, we first examined small RNA accumulation in transgenic Myc-VdSSR1 plants by RNA blotting. To our surprise, in contrast to the strong suppression activity of VdSSR1, the transgenic expression of VdSSR1 only had a weak repression effect on the accumulation of four representative miRNAs, miR159, miR166, miR171, and miR164, and two trans-acting siRNAs (tasiRNAs), tasi5D8 and tasi1151 (29) (SI Appendix, Fig. S7). We next measured primary miRNA (pri-miRNA) levels and found that the selected pri-miRNAs exhibited no obvious change or slightly reduced levels in the Myc-VdSSR1 plants compared with Col-0 (SI Appendix, Fig. S8). To gain further insight into the effect of VdSSR1 on plant sRNA biogenesis, small RNA sequencing was performed in Col-0 and Myc-VdSSR1#6 plants. The results indicated that VdSSR1 exerted a weak effect on global small RNA patterns,
including the composition of sRNAs in different genomic elements and sRNA length distribution (SI Appendix, Fig. S9). Consistent with the RNA blotting results, most of the annotated miRNAs had a similar accumulation in Col-0 and VdSSR1 plants (SI Appendix, Fig. S10).

Nuclear export is a necessary step in miRNA maturation. We next examined whether the nucleocytoplasmic partitioning of the tested sRNAs was affected by VdSSR1. RNA was extracted from the nucleus and cytoplasm of Col-0 and Myc-VdSSR1 plants to measure small RNA levels. Small RNA blotting results demonstrated that the cytoplasmic accumulation of all the tested miRNAs as well as tasiRNAs was significantly reduced in Myc-VdSSR1 plants compared with Col-0 plants. In contrast, the nuclear levels were increased (Fig. 4A and SI Appendix, Fig. S11), leading to a great reduction in the ratio of cytoplasmic/nuclear (C/N) sRNA levels (Fig. 4B).

In line with the reduction in cytoplasmic miRNAs, an obvious increase in the mRNA levels of representative target genes was observed in Myc-VdSSR1 plants (Fig. 4C). In addition to miRNAs and tasiRNAs, we also measured the nucleocytoplasmic distribution of siSuls from Suli and Myc-VdSSR1/Suli plants and found that the C/N siSul ratio was also reduced in Myc-VdSSR1/Suli compared with Suli plants (Fig. 4D). Intriguingly, unlike tested miRNAs and tasiRNAs, nuclear siSuls were reduced in Myc-VdSSR1/Suli plants. This result is consistent with the obvious reduction of total siSuls in Myc-VdSSR1/Suli plants (Fig. 3E). We speculate that this discrepancy may be due to a different biogenesis of miRNA and artificial siRNAs, as reported previously (28). In brief, these data support the notion that VdSSR1 suppresses RNA silencing by modulating the nucleocytoplasmic partitioning of sRNAs.

VdSSR1 Associates with TREX/TREX-2 Complexes In Vivo by Interacting with ALY Family Adaptors. To decipher how VdSSR1 affects the nucleocytoplasmic partitioning of sRNAs, a VdSSR1 immunoprecipitation assay followed by MS (IP–MS) was performed in Myc-VdSSR1 transgenic Arabidopsis. Intriguingly, three RNA export complex–related proteins, ALY4, DSS1-V, and SAC3B, were copurified with Myc-VdSSR1 protein (Fig. 5A and Dataset S2). Arabidopsis encodes four ALY family proteins (ALY1–4), which have been shown to function as adaptor proteins in the TREX complex–mediated mRNA export pathway (30). DSS1-V and SAC3B are conserved components of the TREX-2 mRNA export receptor complex (30–32). In yeasts and metazoans, the TREX complex associates with nascent transcripts, leading to the recruitment of export adaptors that bind to the mRNA and deliver it to the TREX-2 export receptor complex to facilitate translocation of the mRNP through the nuclear pore complex (NPC) (30). Intriguingly, phylogenetic analysis indicated that VdSSR1 is evolutionarily related to Arabidopsis ALYs but displays closer evolutionary relationship to its fungal homologous proteins (SI Appendix, Fig. S12). Supporting this notion, the RRM domains of VdSSR1 and ALY family proteins displayed sequence similarity (SI Appendix, Fig. S13). The copurification of VdSSR1 with RNA export-related proteins suggested that VdSSR1 might interfere with the TREX/TREX-2–related pathway to regulate miRNA nucleocytoplasmic shuttling. To test our hypothesis, protein interactions were examined between VdSSR1 and copurified RNA export proteins. The yeast two-hybrid (Y2H) data indicated that VdSSR1 directly interacted with all four ALY proteins but not DSS1-V and SAC3B (Fig. 5B). The interaction between VdSSR1 and ALYs was further supported by evidence from the split luciferase assay in N. PNAS 2022 Vol. 119 No. 12 e2114583119 https://doi.org/10.1073/pnas.2114583119 5 of 12
**VdSSR1 affects the nucleocytoplasmic partitioning of siRNAs in plant cells.** (A) RNA blotting results showing the nuclear and cytoplasmic accumulation of representative siRNAs in Col-0 and Myc-VdSSR1 plants. U6 and tRNA<sup>Met</sup> served as nuclear and cytoplasmic RNA loading controls, respectively. One representative result of three biological replicates (SI Appendix, Fig. S11) was shown. (B) The relative C/N ratios of siRNAs as determined in three RNA blotting results (Fig. 4A and SI Appendix, Fig. S11). Data are the means ± SD of three biological repeats. An unpaired one-tailed t test was performed. *P < 0.1, **P < 0.01. (C) The relative transcript levels of representative miRNA target genes in Col-0 and two Myc-VdSSR1 lines. The mRNA levels were first normalized to ACT2 and then to Col-0. Data are the means ± SD of three biological repeats. An unpaired one-tailed t test was performed. *P < 0.01, **P < 0.01. (D) Small RNA blotting results showing the nuclear and cytoplasmic accumulation of siSuls from Suli and VdSSR1/Suli plants. (Lower) The relative C/N ratios of siRNAs as determined in Upper.

**VdSSR1 Sequesters ALY Adaptors to Inhibit Its Association with UAP56 and Disrupt mRNA Export.** In *Arabidopsis*, metazoa, and yeast, ALY export adaptors are recruited by the DEAD-box RNA helicase UAP56 (Sub2 in yeast) (30, 35, 36). We found that *V. dahliae* encoded one UAP56-like protein (VdUAP56L) and three ALY-like proteins (SI Appendix, Fig. S15), including VdSSR1. Y2H data demonstrated that all four Arabidopsis ALY proteins could interact with UAP56 (Fig. 6A). However, VdSSR1 could interact neither with *V. dahliae* VdUAP56L nor with *Arabidopsis* UAP56 (Fig. 6A and 6B).
In addition, RNA binding activity was indispensable for the function of ALY adaptors in RNA export (37–39). We examined the RNA binding activity of VdSSR1 in Arabidopsis ALY4 as a positive control. Compared with ALY4, VdSSR1 lost its RNA binding activity (Fig. 6B). The above evidence supports the conclusion that VdSSR1 may be an unfunctional analog of ALY adaptors, or act as a different mechanism from Arabidopsis adaptors. Based on these data, another hypothesis was proposed: VdSSR1 may mimic and sequester ALY adaptors to interfere with the functioning of TREX/TREX-2 complexes. To test this hypothesis, an immunoprecipitation assay was performed to determine the effect of VdSSR1 on the ALY-UAP56 interaction in vivo. In this assay, myc-tagged ALY4 was coexpressed with GFP-tagged UAP56 in tobacco leaves. A recombinant VdSSR1 protein purified from Escherichia coli was added to the lysate of tobacco leaves. α-GFP immunoprecipitation results demonstrated that ALY4 could be pulled down by UAP56 (Fig. 6C). Importantly, the pulled-down ALY4 level was greatly reduced by the addition of recombinant VdSSR1 protein to the lysate. In contrast, addition of the MBP control protein had no obvious effect on the pulled-down ALY4 level. These results support our hypothesis that VdSSR1 sequesters ALY adaptors to inhibit its interaction with UAP56.

It has been reported that knockout of ALY adaptors leads to a nuclear retention of mRNA (37). We next asked whether VdSSR1 influences the nucleocytoplasmic mRNA transport. To this end, a fluorescence in situ hybridization assay was performed in Col-0, Myc-VdSSR1 plants, and a mutant of Nuclear Pore Anchor (NUA), in which mRNA retention has been reported (40). Consistent with previous finding (40), a great increase of nuclear mRNA signal was observed in the nua mutant (Fig. 6D). As expected, increased fluorescence signal in the nucleus was also observed in VdSSR1 plants, suggesting that VdSSR1-ALY interaction has a negative impact on the mRNA export process.
**VdSSR1 Interferes with the Nucleocytoplasmic Shuttling of the AGO1–miRNA Complex.** In plants, miRNA biogenesis undergoes stepwise nuclear maturation before engaging cytosolic target transcripts. During nuclear maturation, the pri-miRNAs are first processed by a dicing complex containing DCL1, HYL1, and SE, and the resulting miRNA–miRNA* duplexes are 2′-O–methylated by the small RNA methyltransferase HUA ENHANCER1 (HEN1) (41, 42). Mature miRNA duplexes were originally believed to translocate to the cytosol in an EXPORTIN5 homolog HASTY (HST)-dependent manner and to be incorporated into AGO1 (43). Recent studies have revealed that HST participates in the miRNA pathway independent of its cargo-exporting activity in Arabidopsis because hst mutants with impaired shuttling exhibit a normal subcellular distribution of miRNAs (47). The current literature supports a model of AGO1–miRNA complex-dependent nucleocytoplasmic shuttling (47, 48). In this model, miRNA is incorporated into AGO1 in the nucleus, and the resulting AGO1–miRNA complexes are then exported to the cytosol through an interaction of the AGO1 nuclear export signal (NES) with EXPORTIN1 (EXP1/XPO1) in the nuclear pore (48). Consistently, TREX is associated with the NPC, and a recent study has reported that the Arabidopsis TREX-2 RNA export complex is essential for multiple steps in miRNA biogenesis, including transcription and nuclear export, through its interactions with RNA polymerase II and the nucleoporin protein NUP1, respectively (49). Dysfunction of the TREX-2 complex leads to increased retention of AGO1 and compromised nuclear export of miRNAs (49).

Here the compromised nucleocytoplasmic shuttling of miRNAs, together with the interaction of VdSSR1 with TREX complex adaptor, inspired us to propose that VdSSR1 may influence the nuclear export of the AGO1–miRNA complex through the connection between TREX and TREX-2 complexes. To test this hypothesis, the nucleocytoplasmic distributions of the AGO1 proteins were measured by Western blotting in Col-0 and *Myco-VdSSR1* plants as well as the *aly2/3*, *aly4*, and *thp1* mutants, which is a component of TREX-2 complex (SI Appendix, Fig. S16). The data revealed a similar accumulation of total AGO1 proteins in all tested genotypes (Fig. 6B). Importantly, AGO1 protein showed increased accumulation in the nuclear fraction in the tested mutants and *Myco-VdSSR1* plants compared with Col-0. In contrast, cytoplasmic AGO1 protein levels were greatly reduced. In line with the increased retention of nuclear AGO1, compromised miRNA export (Fig. 6F and SI Appendix, Fig. S17) and increased accumulation of representative miRNA target genes (SI Appendix, Fig. S18) were also detected in the *aly2/3* and *aly4* mutants. We noted that the reduction of C/N sRNA levels in *aly4* mutant is weaker than in *aly2/3* mutant (Fig. 6G), which may be attributed to the functional redundancy between ALY family adaptors (37). In addition, like *Myco-VdSSR1* plants, the levels of tested pri-miRNAs were slightly reduced in *aly2/3* and *aly4* mutants (SI Appendix, Fig. S8).

The similar inhibition of AGO1–miRNA nuclear export inspires us to investigate the global effects of VdSSR1 and ALY dysfunction on miRNA target genes. To this end, RNA sequencing was performed in Col-0, VdSSR1#6, and *aly2/3* mutant plants. The results indicate that most of the miRNA target genes were commonly regulated in VdSSR1#6 and *aly2/3* mutant plants compared with Col-0 (SI Appendix, Fig. S19), among which 370 genes were commonly up-regulated and 270 genes were commonly down-regulated. The expression pattern of miRNA target genes may attribute to a comprehensive effect of VdSSR1–ALY interaction on the inhibition of nuclear export of both AGO1–miRNA and miRNA.

**VdSSR1-Mediated Interference of AGO1–miRNA Nuclear Export Contributes to *V. dahliae* Virulence in Plants.** Next, we asked whether VdSSR1–ALY interaction-mediated interference of plant AGO1–miRNA nuclear export contributes to *V. dahliae* virulence in plants. To this end, *Myc-VdSSR1* plants were crossed into *aly2/3* mutant to generate *Myc-VdSSR1/aly2/3* plants. The sensitivities of *Myc-VdSSR1*, *aly2/3*, and *Myc-VdSSR1/aly2/3* plants to *V. dahliae* were measured. As shown in Fig. 7A and B, all tested genotypes exhibited similar sensitivity to *V. dahliae* compared with Col-0. Compared with *Myc-VdSSR1* and *aly2/3* plants, no obvious additive effect was observed in *Myc-VdSSR1/aly2/3* plants, suggesting that VdSSR1 promote *V. dahliae* infection in host plants mainly through hijacking ALY functions.

To further verify the contribution of inhibiting AGO1–miRNA nuclear export in VdSSR1-dependent virulence regulation, mutants of XPO1A, a nuclear export receptor in Arabidopsis (50), and NUP96, a component of NPC, as well as *ago1-27*, were subjected to fungal inoculation assay. Consistent with previous reports that XPO1A and NUP1, another component of NPC, were indispensable for the nuclear export of AGO1–miRNA complex (48, 49), *xpo1a* and *nup96* were more sensitive to both V592 and Δvdsr1 strains compared with Col-0 (SI Appendix, Fig. S20). Intriguingly, we found that *ago1-27* mutant was more resistant to both V592 and Δsrr1 strains compared with Col-0 (SI Appendix, Fig. S20). This result is in line with the previous findings of *ago1* in *V. dahliae* (51) and *B. cinerea* (6) inoculation assays. The enhanced resistance of *ago1* mutant to Δsrr1 strain conflicts with the notion that VdSSR1 promotes *V. dahliae* virulence by interfering with AGO1–miRNA nuclear export. We speculate that this may be due to the multifaceted function of AGO1 in plant–fungus interaction. AGO1 is indispensable not only for sRNA-mediated posttranscriptional silencing of target genes, including defense genes, but also for transkingdom RNAi triggered by fungal sRNAs (6). In the latter case, AGO1 dysfunction will lead to a disability of fungal sRNA-mediated transkingdom silencing of plant defense genes, thereby enhancing plant resistance against fungal pathogens (6). Therefore, we think that the increased resistance of *ago1* to Δsrr1 strain is a comprehensive result of the interaction of multiple AGO1-dependent processes.

Based on the above evidence, we proposed an AGO1–miRNA nuclear export-dependent mechanism of VdSSR1-mediated virulence regulation in the plant–*V. dahliae* system (Fig. 7C). In this mechanism, VdSSR1 is translocated from fungal cells to the plant nucleus, possibly through a noncanonical secretion pathway. Nucleus-localized VdSSR1 sequesters ALY adaptors to inhibit their association with the UAP56–TREX complex, which directly/indirectly interferes with TREX-2 complex-dependent nuclear export of the AGO1–miRNA complex and mRNAs. Therefore, VdSSR1 is a general suppressor of plant miRNAs, including the transkingdom miR159 and miR166 in the plant–*V. dahliae* system. The compromised accumulation of cytoplasmic miR166 and miR159 results in attenuated transport to *V. dahliae* cells, finally suppressing the transkingdom silencing of fungal virulence genes and promoting infection. In addition, VdSSR1-mediated inhibition of the mRNA export of certain defense genes may also contribute to the enhanced virulence in plants.

**Discussion**

In this study, we show that the fungal effector VdSSR1 from the phytopathogenic fungus *V. dahliae* sequesters host ALY adaptors to interfere with the nuclear export of the AGO1–miRNA complex and mRNAs, leading to a reduction...
in the levels of cytoplasmic miRNAs, including mobile miRNAs, and succedent attenuation of transkingdom RNAi. With this mechanism, V. dahliae antagonizes transkingdom sRNA-dependent plant immunity and establishes an efficient infection in hosts.

Some biomolecules, such as proteins, antimicrobial peptides, and metabolites, can move from pathogens and pests to hosts or vice versa to modulate cellular processes in recipient organisms (52–55). Recently, increasing evidence has shown that regulatory sRNAs can also move between hosts and pathogens to modulate plant immunity and pathogenic virulence in recipient organisms (3, 15). In this mechanism, plant regulatory sRNAs are assembled into EVs and are transported into the recipient cells of filamentous pathogens to confer transkingdom RNAi of virulent genes (3). Thus, efficient transkingdom RNAi mediated by host sRNAs is beneficial for plants in establishing a successful defense system. It is well known that encoding silencing suppressors is a general strategy for viruses to counteract plant antiviral RNA silencing, and diverse mechanisms are adopted by viruses to hijack host RNA silencing (14). Recently, a few studies have shown that such a strategy can also be adopted by filamentous pathogens to counteract plant antiviral RNA silencing, and diverse mechanisms are adopted by viruses to hijack host RNA silencing (14). Recently, a few studies have shown that such a strategy can also be adopted by filamentous pathogens to counteract plant antiviral RNA silencing, and diverse mechanisms are adopted by viruses to hijack host RNA silencing (14).

In this study, we report a mechanism of fungal effector-mediated suppression of transkingdom RNAi-dependent plant immunity: interfering with the nuclear export of the AGO1–miRNA complex. In this case, our evidence supports the notion that VdSSR1 serves as a general regulator of plant sRNA biogenesis. VdSSR1 expression in plants inhibits the accumulation of mature sRNAs in cytoplasm, leading to an attenuation in the levels of mobile miRNAs and succedent transkingdom silencing of virulent genes in fungal cells. As a result, the sensitivity of VdSSR1-expressing plants to V. dahliae is enhanced. Nuclear export is essential for the maturation of regulatory sRNAs and ensuing targeting. A recent report has shown that miRNAs are mature, methylated, loaded into AGO1 in the nucleus, and exported to the cytoplasm as AGO1–miRNA complexes in a CRM1(EXPO1)/NES-dependent manner (48). Moreover, a recent study from another group further revealed that the TREX-2 complex coordinates with the NPC to link the key steps of miRNA biogenesis, including transcription, processing, and nuclear export, in an AGO1-dependent manner (49). In this study, VdSSR1 interacted with ALY family proteins, adaptors of the TREX complex, to interfere with nucleocytoplasmic shuttling of both the AGO1–miRNA complex and mRNAs. In line with our findings, previous reports have shown that a number of TREX complex mutants display defects in miRNA and siRNA biosynthesis, including tasiRNAs (57, 58). In the tex1 mutant, inverted repeat siRNAs and tasiRNAs are down-regulated, but miR173/miR173* levels are not obviously affected (58). In the case of THO2, miRNAs, tasiRNAs, and 24-nt siR1003 are down-regulated in its mutant (59). This evidence suggests that different mutations of the TREX complex have specificity in the regulation of sRNA biogenesis. In this study, we showed that the total levels of miRNAs and tasiRNAs were slightly
down-regulated in VdSSR1 plants (*SI Appendix*, Fig. S7A), and compromised nucleocytoplasmic shuffling of miRNAs, taSiRNAs, and AGO1 proteins was observed in both VdSSR1 and *aly* mutants (Fig. 6F). These findings suggest that the TREX complex is also involved in the nuclear transport of the AGO1–miRNA complex, although the detailed mechanism remains elusive. In the TREX-2 complex, THP1, one of the core subunits, has been shown to interact and colocalize with the nucleoporin protein NUP1 at the nuclear envelope (49). Thus, we speculate that the TREX complex may coordinate with the TREX-2 complex to facilitate the nuclear export of the AGO1–miRNA complex. This phenomenon may explain the increase in nuclear sRNAs in VdSSR1 plants. Notably, the transcript levels of tested pri-miRNAs are only slightly reduced in *Myc-VdSSR1* plants and *aly* mutants (*SI Appendix*, Fig. S8) compared with an obvious reduction in TREX-2 mutant (49). We propose that the TREX complex or ALY adaptors have a different effect on transcription with TREX-2 complex.

Here we also observed increased nuclear retention of mRNAs in *Myc-VdSSR1* plants, which is consistent with a previous report showing that ALY family proteins are required for nucleocytoplasmic mRNA transport (37). Thus, it is reasonable to propose that a reduced cytoplasmic accumulation of certain defense-related mRNAs may also contribute to the enhanced sensitivity of VdSSR1 plants to *V. dahliae* infection. This issue is worth addressing in future research.

Although our evidence supports that VdSSR1 is a secretory protein, the pathway through which VdSSR1 is secreted remains not completely understood. We speculate that VdSSR1 is secreted through an uncanonical method, as reported previously in a study of two effectors from *P. sojae* and *V. dahliae* (23). In addition, considering the reduced melanin production in Δvdssr1 strain (Fig. 2E) and the increased accumulation of V592 in VdSSR1-expressing plants compared with Δvdssr1 (Fig. 2D), the possibility that VdSSR1 may possess dual functions in both plants and fungi cannot be ruled out, although a previous study has shown that *V. dahliae* virulence in plants is less affected by the production of melanin (25). Indeed, VdSSR1 and *Arabidopsis* ALYs belong to similar protein families but display closer evolutionary relationship with fungal homologs (*SI Appendix*, Fig. S12). Importantly, VdSSR1 lose the ability of RNA-binding activity and interaction with UAP56 (Fig. 6 A and B). There are two possibilities: one is that VdSSR1 may function in a different way from *Arabidopsis* ALYs, and the other is that VdSSR1 has evolved into an effector without ALY-like function. All of these questions deserve further clarity in future studies. Altogether, we revealed an antagonizable mechanism adopted by filamentous pathogens in counteracting transkingdom sRNA-dependent plant immunity. Considering that nuclear export is an essential process for sRNA biogenesis, this mechanism may also be used by other pathogens.

**Materials and Methods**

**Materials.** *Arabidopsis* (Col-0 ecotype) and *N. benthamiana* plants were grown in a growth chamber at 22 ± 2 °C with a 16-h light/8-h dark photoperiod. *G. symptomaticum* hyphae that grew in a growth chamber at 25 ± 2 °C with a 16-h light/8-h dark photoperiod. The virulent *V. dahliae* strain V592 (from diseased cotton in Xinjiang Province, China) was grown on PDA plates at 16-h light/8-h dark photoperiod. The virulent defoliating *N. benthamiana* strain, the coding sequence was ligated into the TefGFP vector with a CloneExpress II One Step Cloning Kit (Vazyme), producing a construct for homologous recombination. For the VdSSR1com strain, the coding sequence was ligated into the TefGFP vector with a CloneExpress II One Step Cloning Kit (Vazyme). All constructs were transformed into *Agrobacterium tumefaciens* strain EHA105 individually. The *Agrobacterium*-mediated transformation method (24) was used for knockout and complementation transformation in the V592 and Δvdssr1 strains, respectively. All the primers used in this study were listed in Dataset S3.

**Fungal Inoculation Assay.** Plants were inoculated by the root-dipping inoculation method as described previously (51, 61). At 18 to 21 dpi, *Arabidopsis* rosette leaves and cotton stem tissues were harvested per plant and flash-frozen in liquid nitrogen. The DNA from each sample (100 mg) was isolated for fungal biomass analysis through RT-qPCR calculation of *V. dahliae*-specific gene expression (51). *AtRubIsCo* gene was used for normalization.

**Suppressor Activity Assay.** Suppressor activity assays were carried out according to a previous report (26). In brief, 4-wk-old *N. benthamiana* leaves were coinfiltrated with cultures of *A. tumefaciens* GV3101 carrying different constructs. The final concentration of each culture in the mixture (per mL) was adjusted to an OD600 of 0.5.

**Purification of Recombinant Proteins and EMSA Assay.** The recombinant plasmids were transformed into BL21 cells and induced with 0.3 mM isopropyl b-D-1-thiogalactopyranoside (Sigma-Aldrich) in Luria-Berti (LB) medium at 16 °C for 12 h. Fusion proteins were purified using Glutathione Sepharose 4B (GE Healthcare) and amylose-conjugated agarose beads (NEB), according to the manufacturer’s instructions. For the EMSA, synthesized RNA oligos were radiolabeled in 50-pmol quantities with 0.3 mM [γ-32P] ATP and 20 units of T4 PNK (NEB) and annealed. Binding reactions were performed according to a previous method (26). After 1 h of electrophoresis at 4 °C, the gel was dried and exposed to a storage phosphor screen (GE Healthcare). The image was scanned using an Amersham Typhoon (GE Healthcare).

**RNA Isolation, Gene Quantification, and RNA Blotting.** Total RNA was isolated using the RNAprep Pure Plant Kit (TIANGEN). The cDNA was synthesized using HiScript II Q Select RT SuperMix (Vazyme). Quantitative PCR was performed with AceQ qPCR SYBR Green Master Mix (Vazyme). The relative gene expression level was normalized to *AtAct2* or *VdEF1*. RNA blotting assay was performed as reported previously (62). For mRNA blotting, 10 μg of total RNA was separated by electrophoresis on 1.2% agarose gels with formaldehyde and transferred to nylon membranes through capillary transfer. For sRNA blotting, 60 μg of total RNA was separated by electrophoresis on 17% PAGE gels and electrophoretically transferred to nylon+ membranes. [γ-32P] ATP was used to label gene-specific oligos sequences by polynucleotide kinase (NEB) for hybridization (8). After washing and drying, the membrane was exposed to a storage phosphor screen (GE Healthcare). Images were scanned using an Amersham Typhoon (GE Healthcare).

**Secretion Assay and Confocal Observation.** For the effector secretion assay (20), the VdSSR1-GFP strain was inoculated on the inner layer of onion epidermal cells and incubated on 1% agar plate at room temperature for 5 to 7 d, followed by confocal imaging (TCS SP8, Leica). To collect fungal culture filtrates and ultrafiltrates for protein isolation and Western blot, 1 mL fungal spore (106 spore mL−1) was cultured in 50 mL liquid Czapek-Dox medium for 7 d. After centrifugation at 1,614 × g for 10 min. The pellets of fungal tissues were frozen and stored at −80 °C for protein isolation. The supernatants were filtered through 0.22-μm-diameter filters to obtain culture filtrates and ultrafiltrated by Amicon Ultra-15 (3 kDa NMWCO) (Millipore) at 4 °C. Approximately 50 mL supernatant of each sample was ultrafiltrated to 100 μL and stored at −80 °C for immunoblotting.

**Fungal Recovery Assay.** The fungus was recovered from infected cotton stems or *Arabidopsis* leaves as previously described (8). *V. dahliae* hyphae that grew from cotton stems or *Arabidopsis* leaves were transferred to PDA medium for...
further growth for another 2 to 3 d. Then, the hyphae were cut from the agar and placed in CM liquid medium for another 3 d of culture. The final large number of hyphae were collected for RNA blotting and gene expression examination.

**IP–MS Assay.** Twelve-day-old Arabidopsis seedlings were ground to a fine powder in liquid nitrogen, and the powder was resuspended in IP buffer (40 mM Tris-HCl, pH 8.0, 300 mM NaCl, 4 mM MgCl2, 2% glycerol, 0.1% Triton-X100, 1 mM PMSF, and 1X mixture) and incubated for 30 min with gentle rotation at 4 °C. The protein suspensions were centrifuged at 14,000 rpm for 20 min to remove debris. IP–MS was performed as previously reported (63). In brief, the supernatant was incubated with Myc-conjugated beads (Invitrogen) for 2 h at 4 °C and then was washed with IP buffer two times following a fivewash with PBS buffer. The pellet was subjected to MS analysis.

**Y2H and Split Luciferase Assay.** The Y2H assays were performed according to the Matchmaker Gold Y2H system user manual (Clontech). The pGBK7 and pGADT7 plasmids containing tested genes were transformed into the AH109 yeast strain. Each combination of single colonies from the SD-LW medium culture was further transferred to synthetically defined (SD) medium lacking leu- cine, tryptophan, histidine (SD-LWH) or lacking leucine, tryptophan, histidine, alanine (SD-LWHA) culture media. For the split luciferase assay (64), the proteins fused to split luciferase were coinfiltrated into 4-wk-old N. benthamiana leaves. The luciferase activity was determined using a charge-coupled device (CCD) cam- era equipped with Winview software (Princeton Instruments).

**Fluorescence in Situ Hybridization Assay.** To determine the relative distri- bution of mRNA in nuclei and cytosol, 6-d-old seedlings grown on solid MS medium were used as previously described (65). Hybridization was performed in PerfectHyb plus solution (Sigma) with an Alexa Fluor 488-labeled 48-nucleotide oligo (dT) probe. Fluorescent signals of seedling root tip cells were imaged using a Leica SP8 microscope.

**Co-IP Assay.** Four-week-old N. benthamiana leaves were infiltrated with a cul- ture of A. tumefaciens GV3101 carrying 35S-ALY4-Myc and 35S-UPA56-GFP or 35S-UPA56-GFP and 35S-GUS. After 2 d, the protein extraction and Co-IP proce- dures were performed as previously described (66), excluding the addition of MBP or MBP-SSR1 before the GFP-conjugated bead Co-IP.

**Nuclear–Cytoplasmic Fractionation.** The nucleocytoplasmic fractionation assay was conducted as previously described (49). Five grams of 12-d-old seedlings were used for each sample. The 5-mL cytoplasmic fraction and final nuclear pellet were used for RNA isolation. For the protein immunoblot in nucleocyto- plasmic fractionation, 12-d-old seedlings were cross-linked in formaldehyde as previously described (49). The frozen tissue was then subjected to the same procedure as described above.

**Data Availability.** All study data are included in the article and/or supporting information. The small RNA and mRNA sequencing data have been deposited in the GEO with the accession code GSE190467.

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