Molecular mechanisms underlying host-induced gene silencing

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H. Z. K. and R. W. I. outlined, wrote, and edited this review. The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (https://academic.oup.com/plcell) is: Roger W. Innes (rinnes@indiana.edu).

Abstract

Host-induced gene silencing (HIGS) refers to the silencing of genes in pathogens and pests by expressing homologous double-stranded RNAs (dsRNA) or artificial microRNAs (amiRNAs) in the host plant. The discovery of such trans-kingdom RNA silencing has enabled the development of RNA interference-based approaches for controlling diverse crop pathogens and pests. Although HIGS is a promising strategy, the mechanisms by which these regulatory RNAs translocate from plants to pathogens, and how they induce gene silencing in pathogens, are poorly understood. This lack of understanding has led to large variability in the efficacy of various HIGS treatments. This variability is likely due to multiple factors, such as the ability of the target pathogen or pest to take up and/or process RNA from the host, the specific genes and target sequences selected in the pathogen or pest for silencing, and where, when, and how the dsRNAs or amiRNAs are produced and translocated. In this review, we summarize what is currently known about the molecular mechanisms underlying HIGS, identify key unanswered questions, and explore strategies for improving the efficacy and reproducibility of HIGS treatments in the control of crop diseases.

Introduction

The phenomenon of RNA-induced gene silencing (also known as RNA interference [RNAi]) was first described in plants over 30 years ago, when plant scientists attempted to overexpress a gene involved in purple pigment production in petunia (*CHALCONE SYNTHASE1*), but instead silenced this gene through a posttranscriptional mechanism (Napoli et al., 1990). A similar phenomenon was also described in fungi during this time period and was named “quelling” (Romano and Macino, 1992; Cogoni and Macino, 1997). Quelling was originally discovered in the fungus *Neurospora crassa* when the expression of either the *albino-1* or *albino-3* gene resulted in silencing of endogenous *albino-1* or *albino-3*. Although the mechanism of quelling was not clear at the time, the silencing of *albino* genes was correlated with reductions in mRNA levels. Later studies on quelling-defective (QDE) mutants showed that the corresponding genes encoded core proteins/enzymes that are now known to mediate RNAi across eukaryotes: RNA-dependent RNA polymerase (QDE-1; Cogoni and Macino, 1999) and Argonaute (AGO) proteins (QDE-2; Fagard et al., 2000).

RNA-induced gene silencing in eukaryotes is mediated by small RNAs (sRNAs; 20–24 nt long) that associate with AGO proteins and then base-pair with complementary mRNAs. The AGO proteins can then slice the mRNA, inhibit translation of the mRNA, or recruit protein complexes that mediate RNA-directed DNA methylation of complementary DNA (reviewed in Carbonell, 2017). The specificity of such gene silencing complexes is dictated by the sRNAs, which are typically produced from longer self-complementary hairpin
RNAs (hpRNAs) or double-stranded RNAs (dsRNAs; produced using flanking inverted promoters) by dsRNA-cleaving enzymes belonging to the Dicer family (DICER-LIKE [DCL] proteins). This RNAi system is highly conserved among eukaryotes, with functional AGO and DCL proteins known to be expressed in nematodes, insects, fungi, oomycetes, plants, and mammals, including humans (Hammond et al., 2000; Bernstein et al., 2001; Vetekuri et al., 2011; Campo et al., 2016).

In plants, transgenics can be silenced by both transcriptional and posttranscriptional mechanisms. Transcriptional gene silencing (TGS) is associated with the DNA methylation of the target gene, which inhibits the binding of RNA polymerase II (Pol II; Rountree and Selker, 1997; Portela and Esteller, 2010). Post-TGS (PTGS) is a consequence of mRNA degradation or the inhibition of translation (Eamens et al., 2008; Guo et al., 2016). The overexpression of sense or antisense gene constructs can induce homologous gene silencing; however, the expression of dsRNAs or hpRNAs is much more effective in silencing target genes (Waterhouse et al., 1998; Wang and Waterhouse, 2000), with optimal silencing achieved by the inclusion of an intron in the hpRNA construct (Smith et al., 2000; Wesley et al., 2001). These studies were key breakthroughs that led to the wide use of RNAi as a tool for the analysis of gene function in plants (Waterhouse and Helliwell, 2003).

The first evidence that silencing-related RNAs produced in plants might be able to silence genes in another organism came from work done with the free-living nematode Caenorhabditis elegans (Boutla et al., 2002). In that work, total RNA was purified from transgenic Nicotiana benthamiana plants expressing a gene encoding green fluorescent protein (GFP) that had been silenced. RNA gel-blot analyses of this total RNA showed that it included siRNAs complementary to both the sense and anti-sense strands of GFP. Injection of this total RNA into transgenic C. elegans resulted in silencing of GFP, showing that the silencing activity produced by the plant was functional in the nematode. Notably, size fractionation of the RNA showed that the most active fraction was 80-90 nt in length, which suggested that the silencing activity was conferred by longer dsRNAs rather than mature siRNAs.

One of the first applications of RNAi to study plant pests or pathogens involved soaking nonfeeding juvenile (J2) stage soybean cyst nematodes (Heterodera glycines) in a buffered solution containing dsRNA, which was found to be taken up by an oral route (Urwin et al., 2002). Bakhetia et al. (2005) used a similar juvenile feeding approach to silence peroxidase and NADPH oxidase genes in the root knot nematode Meloidogyne incognita. They observed phenotypic changes in nematode development up to 35 days after exposure to dsRNA, and fewer nematodes and eggs were recovered from plants infected with dsRNA-treated versus control-treated nematodes (Bakhetia et al., 2005). These results suggested that transgenic expression of dsRNA targeting nematode virulence genes might be an effective approach to suppressing infection by nematodes, and potentially other plant pests and pathogens.

This hypothesis was confirmed in 2006, when Huang and colleagues reported, for the first time, host-induced gene silencing (HIGS) of a nematode gene (Huang et al., 2006). The authors expressed hpRNA in Arabidopsis thaliana that targeted the parasitism gene 16D10, which encodes a peptide secreted into the saliva of root-knot nematodes. Transgenic Arabidopsis expressing 16D10 hpRNA exhibited effective resistance against four major root knot nematode species, highlighting the promise of using HIGS to engineer disease-resistant crops (Huang et al., 2006; Lilley et al., 2007). Numerous studies soon followed that employed HIGS in crop plants to confer resistance to diverse plant pathogens and pests, including fungi (Nowara et al., 2010; Koch et al., 2013), oomycetes (Govindarajulu et al., 2015; Jahan et al., 2015), and insects (Abdellatief et al., 2015; Zhang et al., 2015b). For example, HIGS has been used in many different crops to target fungal pathogen genes, including barley (Hordeum vulgare: targeting genes in Blumeria graminis and Fusarium graminearum; Nowara et al., 2010; Koch et al., 2013), wheat (Puccinia triticina and F. graminearum; Nowara et al., 2010; Panwar et al., 2013), soybean (Fusarium oxysporum; Kong et al., 2022), maize (Aspergillus flavus; Raruan et al., 2020), and banana (F. oxysporum; Gag et al., 2014). HIGS has also been used against insect pests including aphids (Guo et al., 2014), Colorado potato beetle (Zhang et al., 2015b), and cotton bollworm (Mamta et al., 2016).

Although HIGS represents a promising approach for limiting crop losses caused by pathogens and pests, many questions remain, including how silencing RNAs are secreted from plant cells, and how cells in the pathogen or pest take up these RNAs. It is also not clear which proteins from the plant and pathogen are required for successful HIGS. In this review, we focus on the mechanisms underlying HIGS. We also discuss recent findings highlighting how plants use endogenous RNAi to overcome pathogen infection, how plant cells recognize long dsRNA and produce small interfering RNAs (siRNAs), and how we can improve the efficacy of siRNA production using various dsRNA constructs. We describe possible mechanisms of RNA delivery, including whether extracellular vesicles (EVs) play a role in this process. Lastly, we suggest future directions aimed at addressing the major unanswered questions about how HIGS functions and how it can be improved.

**Molecular mechanisms underlying HIGS**

The molecular mechanisms underlying HIGS may differ between insects and nematodes versus fungi and oomycetes (filamentous pathogens). For the herbivorous insects, it appears that long dsRNAs (including hpRNAs) are taken up directly from the host and then processed using the RNAi machinery within the insect or nematode to induce gene silencing. For filamentous pathogens, the available evidence suggests that siRNAs and microRNAs (miRNAs) produced within the host plant are taken up by the pathogen, which
then induce gene silencing, although in most systems, translocation of long dsRNA has not been ruled out. Here we discuss key experiments addressing these issues within each pest and pathogen group.

**Long dsRNAs and sRNAs both induce gene silencing in insects**

In herbivorous insects, the translocation of dsRNA appears to be more effective at inducing gene silencing than the translocation of siRNAs. This was demonstrated by comparing the HIGS efficacy of dsRNA expression inside chloroplasts versus the nucleus. Chloroplast-expressed dsRNAs are protected against host cell DCL enzymes and thus can accumulate to much higher levels than dsRNAs expressed from nucleus-encoded genes, which are rapidly processed into siRNAs (Zhang et al., 2015b; Bally et al., 2016). Importantly, the efficacy of HIGS against two herbivorous insect species, Colorado potato beetle (*Leptinotarsa decemlineata*) and cotton bollworm (*Helicoverpa armigera*), was much higher when dsRNA was expressed inside chloroplasts, indicating that long dsRNA is the key translocated RNA, rather than siRNAs. Such a plastid-based system is unlikely to work against phloem-feeding insects, however, as these insects are unlikely to ingest whole chloroplasts.

The conclusion that insect's RNAi machinery contributes to efficient HIGS in insects is further supported by a recent study from Bally et al. (2020) in which gene silencing was induced in *H. armigera* by expressing an artificial miRNA (amiRNA) construct in *N. benthamiana* that used the backbone of an insect miRNA precursor gene. This modified amiRNA precursor remained largely unprocessed in *N. benthamiana* due to the lack of recognition by plant DCLs. However, feeding on transgenic leaves expressing this construct led to efficient silencing of *H. armigera* genes and high levels of mortality and growth abnormalities. These results indicate that the unmodified precursor of the amiRNA was taken up by insect cells and processed by the insect’s RNAi machinery.

Although long dsRNAs and hpRNAs can clearly be taken up by insects, miRNAs and siRNAs can also induce silencing in insects. For example, amiRNAs based on plant miRNA precursors and siRNAs derived from the expression of long dsRNAs silence genes in aphids (Pitino et al., 2011; Guo et al., 2014; Abdellatef et al., 2015). Remarkably, in one study, silencing appeared to persist through multiple parthenogenetic generations, even after the aphids (*Sitobion avenae*) were switched back to feeding on nontransgenic plants (Abdellatef et al., 2015). These results suggest that HIGS can induce epigenetic changes in aphids that are heritable, or perhaps that aphids produce secondary siRNAs that can be passed onto subsequent asexual generations.

Once dsRNAs are consumed by insects, they must somehow cross the plasma membrane of intestinal epithelial cells and engage with host cell RNAi machinery. One potential route of uptake is endocytosis. Xiao et al. (2015) showed that inhibiting clathrin-dependent endocytosis significantly reduced the uptake of dsRNA and impaired RNAi in red flour beetles (*Tribolium castaneum*). Consistent with this finding, silencing of the gene encoding the clathrin heavy chain in Colorado potato beetles reduced target gene silencing, as did pharmacological inhibition of endocytosis (Cappelle et al., 2016). In this latter study, however, co-silencing of two putative dsRNA transporter genes (*Systemic Interference Defective 1* (*sid-1*)-like A and C ([SilA and SilC]) also reduced target gene silencing, suggesting that dsRNA may also be taken up via a channel-based mechanism. The *sid-1* gene was first identified in the nematode *Caenorhabditis elegans* when screening for mutants lacking the systemic RNAi phenotype (Feinberg and Hunter, 2003), and the Sid-1 protein was then shown to mediate dsRNA import into cells (Winston et al., 2002). *Sid-1*-like genes were subsequently identified in multiple insect species, including cotton/melon aphid (*Aphis gossypii*; Xu and Han, 2008), grasshopper (*Schistocerca americana*; Dong and Friedrich, 2005), honeybee (*Apis mellifera*; Honeybee Genome Sequencing, 2006), and planthopper (*Nilaparvata lugens*; Xu et al., 2013). However, the work of Capelle et al. is the first to show a functional link between Sid-1 and HIGS in insects.

**Long dsRNAs and sRNAs also induce gene silencing in filamentous pathogens**

How silencing RNAs are taken up by filamentous pathogens is also poorly understood, but recent studies have demonstrated that many species can take up naked dsRNA directly from the environment (Wang et al., 2016; Qiao et al., 2021). The RNA uptake efficiency of six fungal pathogens and the oomycete plant pathogen *Phytophthora infestans* was assessed using fluorescein-labeled dsRNA. Notably, dsRNA uptake efficiency varied among species and correlated with the efficiency of gene silencing mediated by exogenous RNA in each species. This finding suggests that HIGS in at least some filamentous pathogens can be mediated by the direct uptake of naked dsRNAs, as opposed to siRNAs produced by the plant or dsRNAs packaged by the plant. This hypothesis is further supported by the finding that mutating the *DCL1* gene in the necrotrophic fungus *F. graminearum* rendered it insensitive to gene silencing when dsRNA was sprayed on detached barley leaves but had no effect on its sensitivity to silencing when siRNAs were used (Koch et al., 2016). These results demonstrate that dsRNAs are taken up by fungal plant pathogens and can then engage with the fungal RNAi machinery to silence fungal genes. That said, it is also clear that fungal pathogens can take up siRNAs directly from the environment and that these siRNAs can also silence target genes (Wang et al., 2016), bypassing the requirement for fungal DCL proteins (Koch et al., 2016). Thus, for transgenic plants expressing nucleus-encoded hpRNA constructs, it is possible that both dsRNAs and siRNAs contribute to the silencing of pathogen genes, with the relative contribution of each depending on the efficiency at which hpRNAs are processed into siRNAs by plant DCLs and the efficiencies at which hpRNAs and siRNAs are secreted.
Optimizing artificial sRNA production in plants

The processing of long hpRNAs into siRNAs in plants is primarily mediated by DCL2, 3, and 4, which collectively produce 21-, 22-, and 24-nt siRNAs (Figure 1; Fusaro et al., 2006). DCL3 is specifically required for the production of 24-nt siRNAs, and DCL4 is required for the production of 21-nt siRNAs, while DCL2 contributes to the production of 22-nt siRNAs, but only in the absence of DCL4 function (Fusaro et al., 2006). In an Arabidopsis dcl2 dcl4 double mutant background, production of 21- and 22-nt siRNAs derived from the hpRNA is largely eliminated, as is the degradation of target mRNAs within the plant.

In contrast, the elimination of DCL3, which blocks the production of 24-nt siRNAs, enhances PTGS. Although paradoxical, this can be explained by an increase in transcription of the hpRNA due to elimination of RNA-directed DNA methylation of the promoter driving hpRNA expression. This increase in hpRNA production should then increase 21- to 22-nt siRNA production, and thus lead to more efficient degradation of the target mRNA (Fusaro et al., 2006). Thus, 21- and 22-nt siRNAs appear to be the primary mediators of PTGS within a plant. Whether this is also true for gene silencing in plant pathogens is not yet clear. However, 21 and 22-nt sRNAs mediate RNAi in insects such as the fruit fly Drosophila melanogaster (Elbashir et al., 2001).

As mentioned above, amiRNAs have also been successfully used in transgenic plants to silence genes in insects (aphids) and filamentous pathogens. Optimizing the expression of such amiRNAs is thus one avenue toward improving the efficacy of HIGS. Standard amiRNA constructs are based on endogenous primary miRNA (pri-miRNA) genes, which encode RNAs that form complex hairpin structures that are processed by DCL1 in plants to produce 21- or 22-nt mature miRNAs. amiRNAs are generated by replacing just the helical region of the pri-miRNA corresponding to the mature miRNA. Optimizing amiRNA production thus involves optimizing the expression and processing of the pri-amiRNA. For example, the loop region in pri-miRNAs plays an important role in controlling the maturation of pri-miRNAs and can directly contribute to the repression of target mRNAs, at least in human cells (Yue et al., 2011). In addition, for some human pri-miRNAs, the loop sequences can also be processed into functional miRNAs. Indeed, in some miRNA libraries, the read count for loop-miRNAs is higher than that of the corresponding mature miRNAs, suggesting they can be functional, rather than just being the byproducts of pri-mRNA processing (Winter et al., 2013). Consistent with this notion, loop-miRNAs bind to AGO2 protein in human cells and induce the degradation of matching mRNAs (Winter et al., 2013). These observations suggest that in some cases,
the loop regions of pri-miRNAs might trigger gene silencing. Thus, one can envision designing amiRNAs that produce two distinct miRNAs, potentially targeting two different transcripts or two different positions in a single transcript. Whether this is also true for plant pri-miRNAs is not known, but the modification of pri-amiRNA loop sequences might offer one avenue for optimizing amiRNA efficacy.

Because pri-amiRNAs are processed by DCL1, they should not lead to the production of 24-nt siRNAs. Therefore, amiRNA-encoding genes should be less prone to TGS via RNA-directed DNA methylation, which is a known problem for hpRNA constructs (Zilberman et al., 2004). However, a study in wheat revealed a higher frequency of silencing of an amiRNA construct in the T2 generation relative to a hpRNA construct targeting the same gene (Gasparis et al., 2014). This is a single study with germinated seeds and a statistically significant reduction in mRNA levels of the target gene (Gasparis et al., 2014). Both constructs targeted the aphid gene Acetylcholinesterase2, but only the amiRNA construct triggered a statistically significant reduction in mRNA levels of the target gene (Guo et al., 2014). This is a single study with single species, however. Similar comparisons are lacking for oomycete and fungal pathogens.

Loop structures in hpRNAs are also known to affect silencing efficacy. Both loop size and loop sequence can affect hpRNA stability (Grobe and Uhlenbeck, 1988; Antao and Tinoco, 1992; Serra et al., 1993; Vecenie and Serra, 2004; Kuznetsov et al., 2008), which consequently can affect siRNA production. hpRNAs with shorter loops are generally more stable than hpRNAs with longer loops (Grobe and Uhlenbeck, 1988). The stem sequence also affects the stability of hpRNAs, with higher GC content producing more stable hairpins (Grobe and Uhlenbeck, 1988). Thus, the optimization of hpRNA constructs to maximize HIGS may involve optimizing the length of the loop and the sequences within the stem.

The entire stem region of hpRNAs likely contributes to siRNA production, but the length of the hpRNA stem is not critical to efficacy, with stem lengths ranging from 98 to 730 nt all shown to be efficient in triggering target gene silencing (Wesley et al., 2001). More critical to hairpin design is the inclusion of an intron in the construct, which is typically placed at the position of the loop, although an intron located 5′ of the stem structure also appeared to promote silencing (Wesley et al., 2001). Why introns promote target gene silencing is not yet understood but presumably results from more efficient siRNA production. This could be related to the recruitment of spliceosome RNA–protein complexes that somehow promote further processing by DCL proteins.

Optimizing target gene selection for HIGS

The efficacy of HIGS also depends on the genes selected for silencing in the pathogen. Some genes are not suitable as targets for HIGS, because their knockdown does not affect pathogenesis (Govindarajulu et al., 2015). Singh et al. (2018) demonstrated that housekeeping genes can make effective targets for HIGS, at least in insects. The authors targeted housekeeping genes in cotton leafhopper via feeding dsRNAs and observed up to 48% mortality (Singh et al., 2018). A caveat of this approach, however, is that housekeeping genes are usually highly conserved between organisms; thus, targeting these genes could be detrimental to nontarget organisms, and potentially even to the host plant. An ideal target gene would be one that is essential for viability of the pathogen/pest on the host plant but is not found in related beneficial organisms or the host plant.

A second concern when designing HIGS constructs is the potential for the resulting siRNAs to silence off-target genes (Xu et al., 2006; Senthil-Kumar and Mysore, 2011). It is estimated that 50%–70% of gene transcripts in plants have the potential to generate siRNAs that either target more than one gene or target endogenous plant genes (Xu et al., 2006). To minimize this risk, computational approaches have been developed to select a collection of 21-nt sequences from a given target sequence with a low probability of such off-target effects, while still being efficiently loaded into AGO proteins (Ahmed et al., 2020). This approach was validated in N. benthamiana using virus-induced gene silencing (VIGS) targeting two different genes, the phytoene desaturase gene and the ribosomal protein L10 (RPL10) gene. RNA-seq analysis of total transcripts in N. benthamiana following infection with VIGS constructs designed to either maximize or minimize off-target silencing confirmed that the tool was effective at minimizing off-target effects while maximizing target gene silencing (Ahmed et al., 2020).

In the context of HIGS, the efficacy and specificity of candidate siRNA sequences can often be tested on pathogens grown in Petri dishes or other artificial environments prior to the generation of transgenic plants. For example, Guo et al. (2019) tested the effect of artificial siRNAs (asiRNAs) that targeted several different genes in the rice fungal pathogen M. oryzae, including MoAP1, which encodes a transcription factor essential for conidia production. Application of asiRNAs targeting MoAP1 to M. oryzae in a Petri dish inhibited the growth of the pathogen and suppressed pathogenicity on rice (Guo et al., 2019). In contrast, feeding M. oryzae asiRNAs that targeted three genes downstream of MoAP1 failed to affect fungal growth or pathogenicity, revealing that these latter three genes were likely not good targets for HIGS. Based on these analyses, the authors expressed an hpRNA construct in transgenic rice that targeted MoAP1 and observed enhanced resistance to multiple strains of M. oryzae. Thus, direct RNA applications/feeding can be useful for evaluating candidate target genes for HIGS.

Genes that function at an early stage of the infection process appear to be particularly good targets for HIGS. For...
example, MoAPI is required for the formation of appressoria, which are required by *M. oryzae* for the penetration of host cell walls (Guo et al., 2019). Similar observations were made for the fungal pathogens *P. triticina* (Panwar et al., 2013) and *Puccinia striiformis* f. sp. *tritici* (Zhu et al., 2017; Qi et al., 2018), in which genes highly expressed in early stages of infection were targeted by HIGS, resulting in robust resistance. These observations suggest that the siRNAs are transferred into fungi prior to the formation of haustoria, possibly from the leaf surface during the growth of germ tubes. The recent discovery that plants secrete RNA into the leaf apoplast (Zand Karimi et al., 2022) raises the possibility that RNA can be deposited onto the leaf surface.

Although the above approaches should enable the selection of target genes and optimal target sequences, it should be noted that the process of siRNA biogenesis in plants often leads to the spread of siRNA production along a target mRNA. For example, gene silencing can initiate in the 3′-untranslated region (UTR) of a target gene and spread to the coding regions (English et al., 1996; Sijen et al., 2001). As a classic example of this, constructs consisting of a single copy of a target gene fused to an inverted repeat of the 3′-UTR region of the nopaline synthase gene from *Agrobacterium tumefaciens* are highly efficient at silencing target genes in transgenic plants, even though the target genes have a completely different 3′-UTR (English et al., 1996; Sijen et al., 2001). As another example, when a transgenic Arabidopsis plant expressed a full-length green fluorescent protein (GFP) construct, hpRNAs targeted the 5′ half of the GFP gene led to the production of secondary siRNAs from the 3′ half (Tauchy et al., 2017). Such spreading is dependent on the host's RNA-DEPENDENT RNA POLYMERASE 6 (RDR6; Tauchy et al., 2017), indicating that RDR6 plays a central role in the production of secondary siRNAs. These observations also point to the potential for off-target effects resulting from secondary siRNA biogenesis. This may be less of a concern for HIGS constructs if the target gene in the pathogen or pest has no similarity to endogenous plant genes.

**Promoter selection for HIGS constructs can affect both the efficacy and specificity of HIGS**

In addition to selecting the appropriate pathogen gene and target site sequence within that gene, HIGS efficacy is likely dependent on obtaining high levels of hpRNA and/or siRNAs in the tissue/cells colonized by the pathogen or pest. Most HIGS experiments have employed strong constitutive promoters to accomplish this goal, such as the 3SS promoter of cauliflower mosaic virus, which is transcribed by RNA Pol II. An RNA Pol III promoter could also be used to drive high expression of hpRNAs, but such promoters have proven to be less effective in HIGS (Wang et al., 2008). This is likely because Pol II-derived hairpins are mostly processed by DCL4, resulting in the production of 21-nt siRNAs, whereas Pol III-derived siRNAs are processed by DCL3 to produce mostly 24-nt siRNAs (Figure 1; Wang et al., 2008). Since 24-nt siRNAs are known to mediate TGS through DNA methylation, hpRNA constructs transcribed by Pol III are more likely to be silenced and are thus not a good choice for sustained HIGS (Gasciolli et al., 2005; Fusaro et al., 2006; Wang et al., 2008).

Another consideration when designing HIGS constructs is whether to use constructs that generate dsRNAs using flanking inverted promoters to drive the expression of sense and anti-sense strands from the same insert or constructs that generate hpRNAs expressed from a single promoter. A potential problem with the former approach is a higher propensity for transcriptional silencing of the HIGS construct due to the production of RNA from the promoter regions. Nevertheless, such inverted promoter constructs have been used to silence target genes in *F. graminearum* in both barley and Arabidopsis (Koch et al., 2013), although only T2 generations were tested.

An alternative to strong constitutive promoters is to select promoters that drive the expression of hpRNAs specifically in the tissue and cell types being attacked. Alakonya et al. (2012) provided a nice example of this approach, in which they used HIGS to engineer tobacco with resistance to colonization by the parasitic plant *Cuscuta pentagona* (dodder). Most parasitic plants establish vascular connections with their host plants via structures called haustoria. Alakonya et al. (2012) used a phloem-specific promoter from the sucrose transporter gene *SUC2* in Arabidopsis to drive expression of a hpRNA targeting the STM gene of dodder, which is required for haustoria formation. Phloem-specific expression of this hpRNA was highly effective at reducing dodder growth on tobacco. The use of a tissue-specific promoter in this example localizes expression of the hpRNA to areas of likely infection while limiting expression in other parts of the plant, which should reduce potential off-target effects on host genes. Given that HIGS appears to act very early during the fungal infection process, prior to the formation of haustoria (Guo et al., 2019), the use of pathogen-inducible promoters may not be optimal, as siRNA production might occur too late to be effective.

HIGS likely requires AGO proteins, but whether they are supplied by the host or pathogen is not yet known

To silence target genes, siRNAs and miRNAs must associate with AGO proteins. AGO proteins bind the dsRNAs products of DCLs and slice and eject one of the strands, leaving a single-stranded guide RNA. Plant genomes typically contain multiple AGO genes. For example, the Arabidopsis genome (a dicot) contains 10 AGO genes (Zhang et al., 2015a), and the *Brachypodium* genome (a monocot) contains 16 (Secic et al., 2019). Plant AGO genes cluster into three major clades. In Arabidopsis, these include AGO1/5/10 (clade I), AGO2/3/7 (clade II), and AGO4/6/8/9 (clade III; Zhang et al., 2015a).

Individual AGO proteins differ in their binding preferences for sRNAs, especially with regards to preferred 5′-nucleotides and preferred lengths (Zhang et al., 2015a). They also differ in their expression patterns and responses to biotic and
abiotic stress, as well as their mechanism of action (Zhang et al., 2015a; Figure 1). Clade 1 AGOs have primarily been implicated in binding 21-nt miRNAs and function in regulating developmental pathways via posttranscriptional mechanisms (Baumberger and Baulcombe, 2005; Wu et al., 2009). AGO2 (clade II) binds to both 21-nt miRNAs and 21-nt siRNAs and plays a central role in anti-viral and antibacterial immune responses (Zhang et al., 2011; Brosseau and Moffett, 2015). In contrast, AGO4 (clade III) binds to 24-nt siRNAs and functions in RNA-directed DNA methylation to epigenetically silence the transcription of transposable elements (Havecker et al., 2010). Notably, AGO3 (a clade II member derived from a tandem duplication of AGO2) also preferentially binds 24-nt RNAs but appears to function by inhibiting the translation of targeted mRNAs, as it associates with polysomes in the cytoplasm (Jullien et al., 2020). However, AGO3 also partially complemented the ago4 mutant when driven by the AGO4 promoter, suggesting that it also functions in RNA-directed DNA methylation (Zhang et al., 2016b).

It is not yet clear whether plant siRNAs bind to pathogen AGO proteins, or if siRNAs are translocated as plant AGO-siRNA complexes. It is also not yet clear whether silencing in pathogens is achieved primarily through a post-transcriptional mechanism (e.g. cleavage of pathogen mRNAs) or through TGS (e.g. DNA methylation).

To understand the function of plant-derived sRNAs in pathogens, it is necessary to investigate the RNAi pathway in pathogens, and in particular, to assess whether DCL and/or AGO proteins in the pathogen are required for HIGS. In some fungal pathogens such as *Mucor circinelloides*, two classes of 21 and ~25-nt siRNAs are produced from the expression of extra-chromosomal DNA. These two classes of siRNAs cause PTGS of target genes in *M. circinelloides*, indicating that this fungus possesses all the machinery required to carry out RNAi (Nicolas et al., 2003). Similarly, in the fungus *M. oryzae*, expression of hpRNAs results in the production of three different sizes of siRNAs ranging from ~19 to ~23-nt containing both sense and antisense strands (Kadotani et al., 2003). Surprisingly, in *M. oryzae*, all siRNA size classes are generated by just one of the two existing DCL proteins (Kadotani et al., 2004). Although siRNA production in fungi and plants looks similar, it is not known what size siRNAs are responsible for transcriptional or PTGS in fungi. As mentioned above, fungi can take up free dsRNAs from their environment, leading to gene silencing, and thus they must be employing their own DCLs and AGOs, or functional equivalents. Nguyen et al. (2018) investigated the function of three AGO proteins in *M. oryzae*, MoAGO1, MoAGO2, and MoAGO3, and showed that, although knockout mutations in AGO1 and AGO3 reduced the levels of gene silencing, the deletion of MoAGO2 resulted in higher efficiency of gene silencing when triggered by hpRNAs expressed in the fungus. These observations confirm the notion that fungal AGOs function in gene silencing, and they indicate that different fungal AGOs perform different functions, just as observed in plants and mammals.

Fungal DCL proteins have also been shown to function in gene silencing. Knockout of DCL1 and DCL2 in the fungal plant pathogen *Colletotrichum higginsianum* derepressed a dsRNA mycovirus, and immunoprecipitation of sRNAs associated with fungal AGO1 showed abundant loading of viral siRNA. These findings indicate that the RNAi machinery in fungi plays an important role in viral defense, similar to its role in plants (Campo et al., 2016). It is thus tempting to speculate that HIGS of fungal pathogens is mediated by the fungal RNAi machinery, rather than by the transfer of plant AGO–siRNA complexes, but this remains to be tested.

**Pathogen-derived sRNAs are transferred to host cells and contribute to virulence**

The presence of AGO and DICER-LIKE proteins in fungal and oomycete pathogens raises the possibility that sRNAs from these pathogens contribute to virulence. In other words, trans-kingdom RNA-mediated silencing may be bidirectional. The first report documenting the transfer of fungal sRNAs into host cells focused on the necrotrophic pathogen *Botrytis cinerea* during Arabidopsis infection (Weiberg et al., 2013). Evidence for transfer included immunoprecipitation of Arabidopsis AGO1 followed by reverse transcription–PCR. Three different fungal sRNAs associated with Arabidopsis AGO1, but not with Arabidopsis AGO2 or AGO4. In addition, *N. benthamiana* transiently expressing a putative target gene fused to GFP showed reduced GFP accumulation when infected by *B. cinerea*, but not when the target site of the sRNA was mutated. Lastly, overexpression of the *B. cinerea* sRNAs in Arabidopsis using an amiRNA construct rendered Arabidopsis more susceptible to *B. cinerea*. Together, these findings support a role for trans-kingdom RNA silencing in the virulence of *B. cinerea*.

Further support for pathogen-induced gene silencing in hosts comes from recent work with the oomycete *Hyaloperonospora arabidopsidis*, which infects Arabidopsis (Dunker et al., 2020). Immunoprecipitation of host AGO1 protein revealed 133 unique pathogen sRNAs, 34 of which were predicted to target at least one Arabidopsis mRNA based on stringent target prediction criteria. To establish sequence-specific targeting, the authors devised a clever assay that indirectly led to the expression of a beta-glucuronidase (GUS) reporter gene upon cleavage of a target sequence. This assay was used to test two *H. arabidopsidis* sRNAs. When the target sequence was scrambled, no GUS expression was observed, whereas a complementary target sequence led to GUS expression that extended multiple cell layers away from the *H. arabidopsidis* hyphae, suggesting that these sRNAs are secreted and mobile (Dunker et al., 2020). Furthermore, overexpression of a short tandem target site mimic, which was expected to sequester complementary sRNAs, increased the transcript levels of the target genes and rendered Arabidopsis plants more resistant to infection,
showing that these sRNAs contribute to virulence inside the host cell.

A second example of an oomycete sRNA being translocated to host cells comes from work on P. infestans (Hu et al., 2022). The authors immunoprecipitated the P. infestans AGO1 protein from infected potato (Solanum tuberosum) leaves rather than the host AGO1 protein. Among the co-precipitated sRNAs was a miRNA-like RNA named miR8788-3p. This sRNA was predicted to target the potato gene StABH1, which encodes an alpha/beta hydrolase of unknown function. Evidence that miR8788-3p is translocated is indirect: StABH1 transcript levels decreased during infection by P. infestans, but not when miR8788 levels were reduced in P. infestans using a target site mimic. Furthermore, these knockdown strains displayed reduced virulence on potato. It remains unclear, however, whether miR8788-3p is translocated as a complex with P. infestans AGO1 or if it also associates with the host’s AGO1.

Spray-induced gene silencing provides mechanistic insights into HIGS

Although HIGS has shown great potential for controlling pests and diseases in crop plants, public concern about genetically modified organisms (GMOs) and restrictions on releasing GMOs into the environment have encouraged plant scientists to develop new approaches for delivering RNAs into pathogens. In a pioneering study, Koch et al. (2016) demonstrated an effective RNA spraying method, called spray-induced gene silencing (SIGS), for controlling F. graminearum infections on barley (Figure 2).

In this study, the authors sprayed a long dsRNA (CYP3-demethylase genes) that targeted three fungal cytochrome P450 lano-sterol C-14α-demethylase genes required for the biosynthesis of fungal ergosterol. Unexpectedly, they observed efficient spray-induced control of fungal infection in distal (nonsprayed) parts of detached leaves.

The observation of fungal gene silencing in distal tissues suggested that dsRNAs were being absorbed by the plant and that dsRNAs, or siRNAs derived from these dsRNAs, were translocating within the plant. Analysis of fluorescently labeled dsRNAs and RNA gel-blot analyses revealed that dsRNA was indeed moving intact, initially being taken up by xylem vessels in the cut surface of detached leaves but then translocating into the symplasm, including the cytoplasm of mesophyll cells (Koch et al., 2016). Importantly, silencing of target genes in the fungus inoculated on distal (nonsprayed) tissues required the fungal DICER-LIKE 1 gene, which strongly indicates that the fungus was taking up long dsRNA from the plant and that this dsRNA was then triggering the RNAi machinery within the fungus (Koch et al., 2016). Notably, the requirement for the fungal DCL-1 gene could be bypassed by spraying sRNAs instead of long dsRNAs (Koch et al., 2016), which indicates that F. graminearum can also take up sRNAs that likely then engage fungal AGO proteins.

These promising results led to numerous studies on spray application of noncoding RNAs to induce gene silencing in various organisms, including the fungal pathogens F. graminearum (Koch et al., 2016; Gaffar et al., 2019; Werner et al., 2020), Phakopsora pachyrhizi (Hu et al., 2020), Botrytis and Verticillium spp. (Wang et al., 2016), and insect pests such as Colorado potato beetle (San Miguel and Scott, 2016) and aphids (Biedenkopf et al., 2020; Yan et al., 2020). These studies have provided additional mechanistic insights into the movement of siRNAs and dsRNAs within leaves. For example, Biedenkopf et al. (2020) demonstrated that fluorescently labeled dsRNAs could be taken up through the stomata of barley leaves into the apoplast (extracellular space) following spray application and translocated into the phloem where they were able to move into distal, nonsprayed, portions of detached leaves, as well as into the roots of whole plants (Biedenkopf et al., 2020). Importantly, both siRNAs derived from the sprayed dsRNAs and intact dsRNAs were detected in distal phloem fluids collected from aphid styliets. This finding indicates that dsRNAs sprayed on the leaf surface can be taken up by plant cells, processed into siRNAs, and the siRNAs translocated through the phloem. Consistent with this notion, target genes in aphids were effectively silenced when aphids fed on distal portions of barley leaves, demonstrating that silencing RNAs can move systemically in barley leaves. In an independent study, Song et al. (2018) showed that dsRNA applied to cut wheat coleoptiles was efficiently taken up and processed into siRNAs. Notably, the application of dsRNAs in this manner was more efficient when silencing target genes in the fungus F. asiaticum than mixing dsRNAs directly into the fungal growth medium immediately prior to the inoculation of wheat coleoptiles. This observation suggests that SIGS functions via the uptake of RNAs from plant tissues rather than the direct uptake of dsRNAs from the leaf surface by the fungus. If this is indeed the case, SIGS efficacy should be enhanced by treatments that promote the uptake of dsRNAs from the leaf surface into plant cells.

One approach to promoting the uptake of exogenous RNA is to attach the RNA to nanoparticles, which are thought to protect RNAs from degradation and possibly promote endocytosis. In a recent study, however, Zhang et al. (2022) demonstrated that the delivery of siRNA bound to gold nanoparticles (AuNs) does not require the internalization of gold nanoparticles. The infiltration of 10-nm spherical AuNs loaded with siRNAs into the leaves of N. benthamiana plants induced target gene silencing. Notably, the injected AuNs were associated with cell walls or localized within the extracellular space, but not inside cells, indicating that the siRNAs were released from the nanoparticles prior to uptake. Regardless, it is clear that plant cells can take up exogenous RNA from the extracellular space, which suggests that plants employ RNA in intercellular communication.

Although SIGS appears to be a promising strategy, questions remain about its underlying mode of action, overall efficacy, and reproducibility (Dalakouras et al., 2020). For instance, Biedenkopf et al. (2020) reported efficient SIGS of
the aphid (S. avenae) Shp gene (encoding a structural sheath protein) when applying dsRNA to detached barley leaves. However, Liu et al. (2021) reported that the application of naked dsRNA on barley plants was inefficient in silencing different target genes (microphage migration inhibitory factor genes MIF1, MIF2, and MIF3) in the same aphid species, even though nymphs fed an artificial diet containing this dsRNA showed efficient silencing. Furthermore, the authors were unable to detect transport of fluorescently labeled dsRNA into phloem cells, contradicting the findings of Biedenkopf et al. (2020). Why these laboratories obtained conflicting results is unclear.

HIGS appears to be part of the endogenous plant immune system

The examples of HIGS and SIGS described above involved the expression or application of artificial RNA constructs. However, there is compelling evidence that plants also translocate sRNAs from their cells to those of pathogens and pests as part of their normal immune responses. For example, Zhang et al. (2016a) showed that cotton plants upregulate two specific miRNAs when infected by the fungus Verticillium dahliae, miR166 and miR159. These miRNAs are taken up by the fungus, resulting in the silencing of two endogenous genes required for virulence. Verticillium dahliae strains expressing versions of these genes in which the miRNA target sites were modified to no longer base-pair with the miRNAs exhibited enhanced virulence, demonstrating that these miRNAs contribute directly to plant immunity by targeting fungal genes. Although an exciting finding, this work raises questions as to why mutations in these target sites in the fungus have not been selected over the course of evolution given that miR166 and miR159 both function in plant development and are highly conserved across angiosperms (Achard et al., 2004; Williams et al., 2005).

A second example of "natural HIGS" comes from work in the Jin laboratory demonstrating the silencing of genes in the fungus B. cinerea by specific Arabidopsis siRNAs (Cai et al., 2018; He et al., 2021). The authors isolated B. cinerea protoplasts from infected Arabidopsis leaves and subjected the protoplasts to siRNA-seq, which revealed numerous plant siRNAs. These plant siRNAs did not appear to be contaminants, as many of the most abundant siRNAs found
Inside *B. cinerea* protoplasts were not among the most abundant leaf siRNAs, indicating that there was specificity in the siRNAs selected for translocation. Consistent with this hypothesis, these enriched siRNAs were found to be associated with EVs isolated from apoplastic wash fluid (AWF; Cai et al., 2018). Furthermore, disrupting siRNA biogenesis by mutating RNA-DEPENDENT RNA POLYMERASE (RDR6) increased susceptibility to *B. cinerea*, suggesting that siRNAs play a direct role in immunity. In support of this conclusion, predicted target genes of these siRNAs in *B. cinerea* were downregulated during infection, but not in *B. cinerea* collected from infected *rdr6* mutant plants (Cai et al., 2018).

As a third example of HIGS involving endogenous sRNAs, Hou et al. (2019) showed that infecting *Arabidopsis* with the oomycete pathogen *Phytophthora capsici* induced the production of a diverse pool of siRNAs, at least some of which were secreted to the extracellular space. Importantly, disrupting siRNA biogenesis rendered *Arabidopsis* hypersusceptible to infection by *P. capsici*, and transgenic expression of specific siRNAs in this pathogen inhibited its virulence, suggesting that secreted siRNAs contribute directly to plant immunity. Consistent with this notion, a potential target gene in *P. capsici* of one specific siRNA was downregulated during infection in wild-type plants, but not in plants compromised in siRNA production (Hou et al., 2019).

Likely related to HIGS is the recent observation of sRNA transfer from parasitic plants into their hosts (Shahid et al., 2018), although strictly speaking this would be considered pathogen-induced gene silencing. In this study, several *Arabidopsis* mRNAs were shown to be targeted by microRNAs produced by the parasitic plant *Cucusta campestris*. These microRNAs moved into host cells during infection, with the target mRNAs exhibiting reduced accumulation. Furthermore, knockout of these genes rendered *Arabidopsis* significantly more susceptible to *C. campestris* than the control, indicating that *C. campestris* miRNAs play a central role in parasitism by silencing target genes in the host.

**Extracellular RNAs**

The above studies indicated that plants secrete sRNAs into the apoplast but did not provide comprehensive analysis of apoplastic RNA. To address this gap in knowledge, several groups have pursued in-depth analyses of apoplastic RNAs (Baldrich et al., 2019; Tosar and Cayota, 2020; Zand Karimi et al., 2022). Analyses of apoplastic RNA isolated from Arabidopsis leaves have revealed abundant RNA species ranging in size from ~10 nt to >500 nt (Figure 3; Zand Karimi et al., 2022). The great majority of apoplastic RNAs longer than 40 nt could be pelleted by centrifuging the AWF at 100,000g, indicating that long RNAs are associated with particles of some kind. Conversely, the majority of smaller RNAs could not be pelleted, suggesting that they are not associated with particles, including plant EVs, which should all pellet at 100,000g (Rutter and Innes, 2017). Based on protease and RNase protection assays on P100 pellets (materials pelleted by centrifugation at 100,000g for 1 h), the RNAs in these pellets were found to be located outside EVs (Zand Karimi et al., 2022), suggesting that plant EVs do not carry significant amounts of RNA. This conclusion is consistent with quantitative analyses of miRNAs in human EVs, which found that even the most abundant miRNAs are present at concentrations of far less than one miRNA molecule per vesicle, with the majority of extracellular miRNAs located outside of EVs (Arroyo et al., 2011; Chevillet et al., 2014). Based on these findings and those of Zand Karimi et al. (2022), we believe that it is unlikely that EVs play a direct role in the translocation of endogenous RNAs from plant cells to pathogens.

This conclusion runs counter to conclusions reached by Cai et al. (2018) and He et al. (2021), who reported that siRNAs taken up by *B. cinerea* were carried inside EVs. The authors showed that siRNAs purified from the apoplast were protected against degradation by micrococcal nuclease but not when the pellet was pretreated with detergent that should disrupt EV membranes. However, work in our laboratory (Baldrich et al., 2019; Zand Karimi et al., 2022) has shown that most apoplastic siRNAs do not co-pellet with...
EVs (Figure 3), and those that do can be degraded by RNase A if the pellet is pretreated with protease, even though protease treatment does not disrupt EVs. Based on these observations, we concluded that some siRNAs are packaged inside protein complexes that co-purify with EVs, but these are not located inside EVs. It is possible that the detergent treatments described in Cai et al. (2018) disrupted protein–RNA complexes as well as EVs, thus rendering the siRNAs sensitive to nuclease digestion. It should be noted, however, that these methods represent indirect assays for the localization of siRNAs. Final resolution of this debate may require in situ analyses using super resolution or electron microscopy.

Analysis of apoplastic RNAs by RNA-seq have shown that the majority of these RNAs are derived from ribosomal RNA and intergenic regions (Zand Karimi et al., 2022). Notably, apoplastic RNAs also contain a large number of circular RNAs (Zand Karimi et al., 2022), which were recently shown to contribute to immune responses in rice against the fungal pathogen M. oryzae (Fan et al., 2020). We thus speculate that extracellular long RNAs, especially circular RNAs, play important roles in plant–microbe interactions, representing an important issue for further study.

Zand Karimi et al. (2022) revealed that the majority of apoplastic RNA is protected against degradation via association with extracellular proteins. Consistent with this idea, multiple RNA-binding proteins have been identified in AWFs (He et al., 2021), including AGO2 and GLYCINE-RICH RNA-BINDING PROTEIN 7 (GRP7; Zand Karimi et al., 2022). Notably, Arabidopsis grp7 and ago2 single mutants displayed significant changes in apoplastic RNA contents relative to wild-type plants, suggesting that these two proteins contribute directly or indirectly to the secretion and/or stabilization of extracellular RNAs (Zand Karimi et al., 2022). We thus hypothesize that extracellular RNA–protein complexes are the primary mediators of HIGS.

If our hypothesis is correct, several interesting questions arise: How are specific RNA–protein complexes selected for secretion into the apoplast, and how are these complexes taken up by pathogens? Recently, He et al. (2021) identified several RNA-binding proteins in the apoplastic fluids of Arabidopsis, including AGOT1, RNA helicases (RH11 and RH37), and annexins (ANN1 and ANN2). They also reported that pathogen infection increases the secretion of these RNA binding proteins (RBPs) into the apoplast. Using protease protection assays, the authors showed that several of these proteins are protected against protease digestion in the absence of detergent, suggesting that they are located inside EVs. How these proteins are encapsulated inside EVs, and whether EVs release these proteins inside pathogens, are not yet known. These data suggest that these specific RNA-binding proteins contribute to HIGS, but this hypothesis remains to be tested.

RNA uptake by pathogens and pests

How RNAs move from plant cells into the cells of a pathogen or pest is unclear, but this process likely depends on the pathogen’s lifestyle (Koch and Wassenegger, 2021). As described above, insects and nematodes take up plant RNAs from plant cells using their feeding structures, delivering RNA into their digestive systems. Somehow this RNA can survive in the digestive system and be absorbed by intestinal epithelial cells, and from there the RNAi signal spreads into distal tissues via poorly understood mechanisms (Tian et al., 2016; Biedenkopf et al., 2020). Although we have little understanding of RNA uptake by fungi, the observation that some fungal pathogens can take up naked sRNA and long dsRNAs from their environment suggest that RNAs might be taken up via an endocytic process (Wang et al., 2016; Qiao et al., 2021). How the endocytosed RNA is then released intact from endosomes to engage the fungal RNAi machinery represents a major question about the HIGS process that remains to be answered.

Phloem RNA

In addition to the apoplast, another likely location for RNA transfer from plants to pathogens and pests is the phloem. Zhang et al. (2009b) assessed the RNA content of phloem in pumpkin (Cucurbita maxima) and identified several species of RNA, including siRNAs, microRNAs, spliceosomal RNAs, and tRNA fragments. The latter were shown to interfere with translation. Similar results were reported by Lalande et al. (2020), who found that 30–35-nt and 19–25-nt tRNA fragments from Arabidopsis inhibited the in vitro translation of reporter genes. These observations raise the interesting possibility that tRNA fragments also inhibit translation in pathogens and pests when taken up from the phloem or apoplast. It has been suggested that phloem-mobile mRNAs might contain sequence elements in their UTRs that adopt tRNA-like structures that direct RNA into the phloem (Kehr and Buhtz, 2008). Indeed, tRNA-like structures have been detected in viral RNAs that translocate through phloem (Lezhov et al., 2019).

Long-distance RNA translocation through the phloem was confirmed by assessing sRNA/mRNA movement across graft junctions (Ruiz-Medrano et al., 1999; Zhang et al., 2009a; Azumi Kanehira et al., 2010; Notaguchi et al., 2012; Thieme et al., 2015; Zhao et al., 2020). Similarly, mRNA movement through the phloem has been reported in the host–parasite junctions of parasitic plants (David-Schwartz et al., 2008). Interestingly, it appears that RNAs in the phloem are safe from degradation because there is no detectable RNase activity in phloem sap (Sasaki et al., 1998; Doering-Saad et al., 2002; Kehr and Buhtz, 2008).

Although diverse RNA species have been detected in both the phloem (David-Schwartz et al., 2008) and apoplast (Baldrich et al., 2019; Zand Karimi et al., 2022), the mechanism of RNA translocation between the symplast, apoplast, and phloem is not yet clear. Studies on the movement of sRNAs between the maternal tissues, endosperm, and embryos of Arabidopsis showed that sRNAs can move from endosperm cells to embryo cells, even though there are no direct connections between these cells (Martienssen, 2010). This observation indicates that sRNAs must transit through
the apoplast in developing seeds (Martienssen, 2010; Melnyk et al., 2011).

Additional evidence for the apoplastic transport of RNA comes from a study in which hpRNAs and siRNAs were directly injected into the trunks of grapevine (*Vitis vinifera*) and introduced through the base of cut petioles into *N. benthamiana* (Dalakouras et al., 2018). These experiments showed that both classes of RNA can spread systemically over the course of 10 days. Notably, petiole absorption of siRNA labeled with the fluorescent dye CY3 produced fluorescent signal in the guard cells of systemic leaves, but not in mesophyll or other epidermal cells. Since guard cells lack plasmodesmata (Wille and Lucas, 1984), they likely took up RNA directly from the apoplast. Why only guard cells were able to take up RNA from the apoplast is unclear. Consistent with the lack of uptake by mesophyll cells, no silencing of a target GFP transgene was observed.

**Viruses as tools for studying RNA transport**

Plant RNA viruses can spread systemically in the host, which represents a form of intercellular RNA transport. RNA movement appears to occur through both the symplast and apoplast (Omid et al., 2008; Wan and Laliberté, 2015; Movahed et al., 2019). For symplastic movement, a key step is passage through plasmodesmata, which are often modified by viral proteins (Sambade et al., 2008; Epel, 2009). Apoplastic movement is less well-characterized, but Movahed et al. (2019) reported the release of turnip mosaic virus (TuMV) RNA into the apoplast via fusion of multivesicular bodies with the plasma membrane. Whether viral RNAs can use EVs to move between cells is not yet clear, but viral RNA fragments can be detected in apoplastic fluid (Movahed et al., 2019; Hu et al., 2021). Although Movahed et al. reported the presence of TuMV viral particles associated with EVs, Hu et al. (2021) reported that PVX particles found in apoplastic fluid were not associated with EVs. The functional significance of apoplastic viral RNA thus remains enigmatic.

**Apoplastic RNases**

The apoplast contains several species of extracellular ribonucleases (ex-ribonucleases), which have been shown to target viral RNAs (Sangaev et al., 2007; Sugawara et al., 2016; Potrokhov et al., 2021). Some of these ex-ribonucleases are induced in response to wounding and viral infection (Kurata et al., 2002; LeBrasseur et al., 2002). The expression of a wound-induced ex-ribonuclease from *Zinnia elegans* enhanced resistance to potato virus Y in tobacco plants, confirming a role for this ex-ribonuclease in virus resistance. Whether these ex-ribonucleases also affect apoplastic RNAs of plant origin is not clear.

Some fungal pathogens also secrete ribonuclease-like effectors, presumably to target plant-derived RNAs (Pliego et al., 2013; Pennington et al., 2019). However, these effectors appear to lack catalytic activity and thus may function as RBPs rather than RNA degrading enzymes. How dsRNAs/sRNAs are protected against plant and pathogen ribonucleases in the apoplast is unclear, but the presence of RNA-binding proteins in the apoplast suggest that these RNAs may be protected by these proteins (He et al., 2021; Zand Karimi et al., 2022).

**Potential roles of RNA-binding proteins in RNA movement**

Proteomic analyses of phloem sap, xylem sap, and AWFs have shown that these three fluids share overlapping sets of proteins, including multiple RNA-binding proteins (Pallas and Gómez, 2013; Rodríguez-Celma et al., 2016; Rutter and Innes, 2017; Godson and van der Hoorn, 2021). For example, Rutter and Innes (2017) reported that plant EVs contain heat shock proteins, annexins, and a member of the small RBP family (GRP7). Remarkably, some of these proteins appear to be involved in the transport of sRNAs or viral RNAs between cells. Yan et al. (2020) showed that Arabidopsis GRP7 mediates the movement of sRNAs between cells through the plasmodesmata. They also noted that GRP7 binds to single-stranded viral RNAs and transfers them between cells by interacting with plasmodesmatal receptors (Yan et al., 2020). GRP7 belongs to the same family of RNA-binding proteins as HNRNPA2B1 in mammalian cells, which plays a role in loading sRNAs into EVs and in viral RNA movement in mammalian cells (Villarroya-Beltri et al., 2013; Zhou et al., 2020).

Similarly, several RNA-binding proteins in phloem sap have been reported to play critical roles in long-distance RNA translocation (Xoconostle-Cázares et al., 1999; Yoo et al., 2004; Yan et al., 2020). Yoo et al. (2004) identified CmPSRP1 as an RNA-binding protein that binds to sRNAs in the phloem of pumpkin and facilitates intercellular movement. Interestingly, a recent study on sRNA movement in Arabidopsis revealed that sRNAs move long distance and cell-to-cell as double-stranded rather than single-stranded siRNAs and are not bound to AGO proteins (Devers et al., 2020), suggesting that there are specific dsRNA-binding proteins that mediate both local and long-distance sRNA movement.

The above examples mostly focus on the intra-organismal movement of RNAs. However, RNA-binding proteins likely play important roles in inter-organismal movement as well. An interesting example of this has been described in honeybee colonies, where diverse protein-coding and noncoding RNAs are sequestered into royal jelly by worker bees and fed to developing larvae in the hive (Maori et al., 2019). These RNAs are stabilized by an RNA-binding protein named Major Royal Jelly Protein 3 (MRJP-3), which makes up 10%–15% of total proteins in royal jelly. MRJP-3 binds to both single-stranded RNA and dsRNA with a minimum length of 18 nt. This binding occurs in a multivalent fashion, which leads to the formation of large protein-RNA aggregates that are resistant to degradation by RNase A. Notably, MRJP-3 was shown to enhance the uptake of dsRNAs by the nematode *C. elegans*, suggesting that one of its functions is to
enhance the transmission of RNAs from worker bees to larvae (Maori et al., 2019). We speculate that plants may secrete RNA-binding proteins such as GRP7 into the apoplast to enhance the uptake of plant RNAs by microbes.

Future directions for foundational HIGS and SIGS research
Additional investigations are required to understand the mechanisms of RNA uptake by both plants and pathogens, as these insights should lead to improvements in the efficacy of both HIGS and SIGS. Basic questions remain to be answered, such as how specific RNAs and RNA-binding proteins are selected for export, and how they are exported. The roles of pathogen RNAi machineries in HIGS and SIGS also need to be determined. Does SIGS require specific AGO proteins from plants or pathogens? What is the mechanism of RNA uptake and translocation by pathogens? Once RNA is taken up by a pathogen, how is it released into the cytoplasm to engage with pathogen RNAs? Addressing this long list of questions will require a robust HIGS system in which candidate genes can be disrupted in both the host and pathogen, allowing individual genes and gene families to be assessed for their contributions to HIGS and SIGS.

Acknowledgments
We thank Meenu Singla-Rastogi and Brian Rutter for their comments on the manuscript. Figures 1 and 2 were created using BioRender.com.

Funding
This work was supported by the United States National Science Foundation (Grants IOS-1645745 and IOS-1842685 to R.W.I.), Department of Energy (Grant 0000248555 to R.W.I.), and the NovoNordisk Foundation.

Conflict of interest statement. None declared.

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