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Evaluation of dsRNA delivery methods for targeting macrophage migration inhibitory factor MIF in RNAi-based aphid control

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Abstract

Macrophage migration inhibitory factors (MIFs) are multifunctional proteins regulating major processes in mammals, including activation of innate immune responses. In invertebrates, MIF proteins participate in the modulation of host immune responses when secreted by parasitic organisms, such as aphids. In this study, we assessed the possibility to use *MIF* genes as targets for RNA interference (RNAi)-based control of the grain aphid *Sitobion avenae* (*Sa*) on barley (*Hordeum vulgare*). When nymphs were fed on artificial diet containing double-stranded (ds)RNAs (*SaMIF*-dsRNAs) that target sequences of the three *MIF* genes *SaMIF1*, *SaMIF2* and *SaMIF3*, they showed higher mortality rates and these rates correlated with reduced *MIF* transcript levels as compared to the aphids feeding on artificial diet containing a control dsRNA (*GFP*-dsRNA). Comparison of different feeding strategies showed that nymphs' survival was not altered when they fed from barley seedlings sprayed with naked *SaMIF*-dsRNAs, suggesting they did not effectively take up dsRNA from the sieve tubes of these plants. Furthermore, aphids' survival was also not affected when the nymphs fed on leaves supplied with dsRNA via basal cut ends of barley leaves. Consistent with this finding, the use of sieve tube-specific YFP-labeled *Arabidopsis* reporter lines confirmed that fluorescent 21 nt dsRNA_{Cy3}, when supplied via petioles or spraying, co-localized with xylem structures, but not with phloem tissue. Our results suggest that *MIF* genes are a potential target for insect control and also imply that application of naked dsRNA to plants for aphid control is inefficient. More efforts should be put into the development of effective dsRNA formulations.

Keywords Macrophage migration inhibitory factor (MIF) · dsRNA · Phloem · Sitobion avenae · Xylem

Introduction

Macrophage migration inhibitory factors (MIFs) are multifunctional proteins regulating major processes in mammals, including activation of innate immune responses (Mitchell and Bucala 2000). MIF proteins also play a role in innate immunity of invertebrates and participate in the modulation of host immune responses when secreted by parasitic organisms such as aphids (Rosani et al. 2019; Ghosh et al. 2020). A broad survey of the presence of *MIF* genes across

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803 species of plants, fungi, protists, and animals identified them in all eukaryotes. MIFs seem to be essential and highly conserved in some kingdoms (e.g., plants), while they appear more dispensable in other kingdoms (e.g., in fungi) or present in several diverged variants (e.g., insects), suggesting potential neofunctionalizations within the protein superfamily (Michelet et al. 2019). MIFs were discovered in 1966 as a product of activated T cells that limited the random migration of macrophages in vitro (David 1966). Subsequently, it was shown that MIFs not only are involved in cell proliferation and apoptosis but play a vital role in the host response against parasitic infection (Calandra and Roger 2003) and vice versa in parasite virulence (Ghosh et al. 2020).

MIFs of aphids also are involved in the response to pathogens and mutualistic symbionts (Dubreuil et al. 2014). Multiple copies of *MIF* genes were found in aphid genomes, including pea aphid (*Acyrthosiphon pisum*, *Ap*) and green peach aphid (*Myzus persicae*, *Mp*) (Dubreuil et al. 2014). MIFs are secreted in aphid saliva during feeding, thereby inhibiting major plant immune responses and therefore are crucial to plant infestation (Naessens et al. 2015). Ectopic expression of *MIFs* in leaf tissues inhibited major plant immune responses, such as the expression of defense-related genes, callose deposition, and hypersensitive cell death. Functional complementation analyses showed that MIF1 is the key member of the MIF protein family that allows aphids to exploit their host plants.

Aphids are one of the largest groups of phloem-feeding pests, which can cause huge losses in agriculture and horticulture worldwide (Jaouannet et al. 2014; Pons et al. 2020). They colonize the leaves and stalks, and migrate later towards the ears and settle among the bracts and kernels in the milky-ripe stage of corn plants. A massive withdrawal of sieve tube components weakens the plant and eventually leads to a reduced overall yield. In most cases, aphids act as important vectors of viruses to spread plant disease (Ng and Perry 2004; Will et al. 2007). More than 5000 aphid species have been described (The International Aphid Genomics 2010).

We investigated the possibility of using *MIF* genes as targets for RNAi-based insect control in plants. Several studies have shown that aphids are sensitive to double-stranded (ds) RNA (Jaubert-Possamai et al. 2007; Pitino et al. 2011) and therefore are amenable to RNAi strategies in crop protection (Christiaens et al. 2020; Liu et al. 2020). In 2015, we showed that dsRNA derived from the gene encoding salivary sheath protein (SHP), when expressed in barley, strongly reduced the survival of the grain aphid *Sitobion avenae* (*Sa*) (Abdellatef et al. 2015). Similar results were obtained when the green peach aphid was grown on transgenic *Arabidopsis thaliana* expressing dsRNA with homology to the *MpC002* gene (Coleman et al. 2015). The *C002* gene was

first described by Mutti et al. (2008) and is predominantly expressed in the salivary glands of aphids.

The degree and the persistence of RNAi in aphids are strong as evidenced by the finding that target genes were also down-regulated in nymphs born from mothers exposed to dsRNA-producing transgenic plants. Notably, *S. avenae* and *M. persicae* aphids reared on transgenic barley (Abdellatef et al. 2015) or *Arabidopsis* (Coleman et al. 2015), expressing dsRNA against salivary protein components, even showed a decline in survival over several generations. These reports strongly support earlier proposals to use RNAi-based strategies for insect control (Price and Gatehouse 2008; Burand and Hunter 2013).

While transgenic strategies using dsRNA-expressing plants have proven successful in insect control, other strategies might also be applicable. Injection and ingestion of dsRNAs also can induce significant levels of gene silencing in insects (Tomoyasu and Denell 2004; Zhu et al. 2011). Thus, it also might be feasible to deliver dsRNA through foliar application (San Miguel and Scott 2016; Gogoi et al. 2017). The purpose of our study was to assess the potential of *MIF* genes as a target for pest control by oral delivery of dsRNAs derived from gene sequences of three *Sitobion avenae MIF* genes. We also compared the efficiency of different dsRNA delivery strategies, including exposure of aphids to artificial diet versus leaf spray application and a sucrose-aid delivery in order to provide theoretical support for future application.

Results

Prediction of MIF genes in Sitobion avenae (Sa)

Genomic MIF sequences of evolutionarily distant species from hemipterans revealed a highly conserved structure (Dubreuil et al. 2014; Michelet et al. 2019). With the aim to deduce MIF gene sequences in Sa from currently available expressed sequence tags (ESTs) in public databases (https://www.ncbi.nlm.nih.gov/), we searched for MIF genes in insect genomes. Based on known peach aphid Myzus persicae and pea aphid Acyrthosiphon pisum sequence data, partial sequences of SaMIF1, SaMIF3, and SaMIF4 were predicted, amplified by PCR using degenerate primers (Table S1) and sequenced. Sequence alignment, which also included the already published SaMIF2 sequence (Dubreuil et al. 2014), confirmed that SaMIFs are highly conserved in aphids' evolutionary history (Fig. S1). The identified SaMIF sequences (Table S2) were cloned and used as a template for dsRNA production.



Detection of fluorescence-labeled dsRNA in aphids' midguts after feeding

We conducted dsRNA feeding experiments to assess the effect of MIF gene silencing on aphid survival. Since

the sucrose concentration in artificial diet is critical, we first tested the optimal concentration of sugar supply. We found that a concentration of 7.5% (w/v) sucrose is optimal for the survival of *Sa* (Fig. S2). Next, we investigated the uptake of fluorescent-labeled dsRNA by *Sa* nymphs from artificial diet. To this end, *SaMIF1*-dsRNA (223 nt; Table S2) labeled with UTP-PEG₅-AF488 during the dsRNA synthesis was added to the artificial diet at

dsRNAAF488 feeding

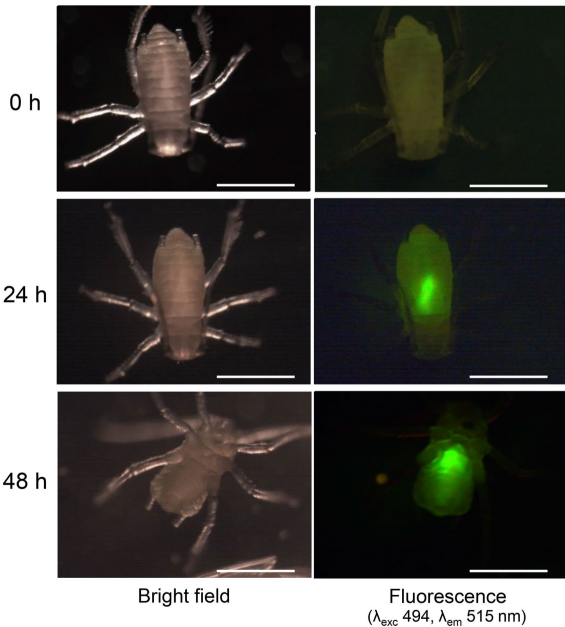


Fig. 1 Uptake of fluorescence-labeled dsRNA from an artificial diet and its spreading inside *Sitobion avenae*. Pictures were taken at 0 h, 24 h and 48 h after onset of feeding. The artificial diet contained 250 ng/µL *SaMIF1*-dsRNA_{A488}. Fluorescence was detected in the

insect gut. Left panels: stereo-microscopic analysis under bright field; right panels: fluorescence stereo microscopic analysis: excitation/emission wavelength (494 nm/515 nm), scale bars = 500 µm



a concentration of 250 ng/ μ L. A fluorescent signal was observed in the midgut of Sa nymphs within 24 h and spread further into the body within 48 h (Fig. 1).

The impact of different SaMIF-dsRNAs on aphids' survival

Aphid MIFs are involved in the regulation of plant immune responses, but it remains largely unknown how the respective members of the MIF family contribute to this activity. In Mp, mainly MIF1 functions as secreted salivary protein to suppress host immunity (Naessens et al. 2015). We investigated the effect of silencing different SaMIF genes on Sa's survival. Since expression of MIF1, MIF2 and MIF3 are strongly induced after immune challenge in Mp (Dubreuil et al. 2014), we placed our focus on these genes. One-day-old Sa nymphs were fed with artificial diet containing dsRNAs directed against SaMIF1 (SaMIF1-dsRNA, 223 nt), SaMIF2 (SaMIF2-dsRNA, 323 nt), SaMIF3 (SaMIF3-dsRNA, 212 nt), and Green fluorescent protein (GFP-dsRNA, 476 nt) (see Table S1) at two different doses, 250 ng/µL and 125 ng/µL. We found that survival rates of aphids treated with either SaMIFdsRNA versus GFP-dsRNA were significant reduced (Kaplan–Meier analysis and log-rank test, $p \le 0.0001$) at day 4 of feeding with 250 ng/µL (Fig. 2a). Feeding with the lower concentration of SaMIF-dsRNAs (125 ng/µL) only resulted in a statistically significant lower survival rate after treatment with SaMIF1-dsRNA (Fig. 2b). This finding also confirms that beyond the anticipated function of MIFs as effector interacting with the plant's defense, MIFs have essential endogenous function in the aphid (Dubreuil et al. 2014).

The impact of *SaMIF*-dsRNA on *MIF* target downregulation

Next, we determined target gene silencing upon feeding aphids with the respective SaMIF-dsRNA (250 ng/ μ L) in artificial diet by 72 h of feeding using RT-qPCR. Consistent with the effects of dsRNA on aphids' survival, transcript levels of all three SaMIF genes were reduced significantly (Student's t-test, p < 0.05) (Fig. 3). These data further substantiate that the effect of SaMIF-dsRNAs on Sa is based on RNAi-mediated gene silencing.

The impact of SaMIF-dsRNA mixtures on aphids' survival

The above data indicate that *SaMIF1* plays a prominent role in the survival of aphids. To further assess *SaMIF1* as a target, we comparatively analyzed the effects of *SaMIF1* silencing versus a triple gene silencing of all three *MIF* genes on the survival of *Sa*. Nymphs were treated with (1) *SaMIF1*-dsRNA (187.5 ng/μL), (2) a mixture of *SaMIF1*-dsRNA, *SaMIF2*-dsRNA, and *SaMIF3*-dsRNA (each at a concentration of 62.5 ng/μL in the artificial diet) and (3) *GFP*-dsRNA (187.5 ng/μL) as control. The relatively low concentration of individual *SaMIF*-dsRNAs in the mixture was chosen because we did not expect a measurable effect on aphid

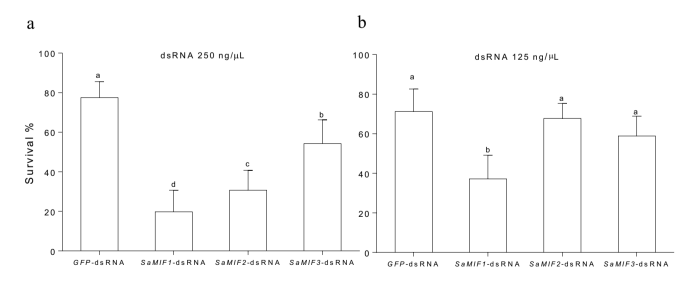


Fig. 2 Sitobion avenae survival rates after four days of feeding on artificial diet supplied with dsRNA present as percent of control (no dsRNA in the diet). SaMIF1-dsRNA, SaMIF2-dsRNA, and SaMIF3-dsRNA were used with concentration of 250 ng/μL (**a**) and 125 ng/μL (**b**) GFP-dsRNA was used as an additional control, since a tar-

get for this dsRNA is lacking in aphids. Survival data were evaluated by Kaplan–Meier analysis and log-rank test based on three biological replicates. Bars represent means (\pm SD) from three independent replicates. Different letters indicate significant differences at p < 0.001



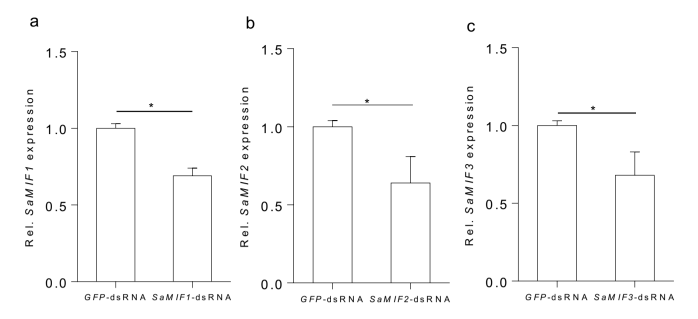


Fig. 3 Relative expression of target genes *SaMIF1*, *SaMIF2* and *SaMIF3* in *Sitobion avenae* fed on an artificial diet containing 250 ng/μL of the respective *SaMIF*-dsRNA. RT-qPCR analysis data for **a** *SaMIF1*, **b** *SaMIF2* and **c** *SaMIF3* were normalized to the aphid's

Ribosomal protein L27 (Rpl27) gene. GFP-dsRNA was used as a control. Bars represent means (\pm SD) from two independent replicates. The asterisks indicate significant differences (Student's *t*-test; p < 0.05)

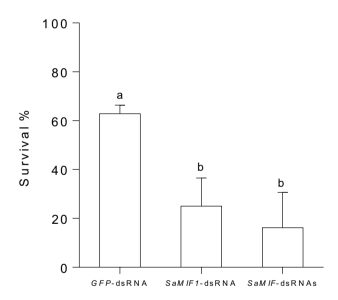


Fig. 4 Aphid survival after five days of feeding on artificial diet supplied with SaMIF1-dsRNA (187.5 ng/ μ L), a mixture of SaMIF1-dsRNA, SaMIF2-dsRNA and SaMIF3-dsRNA (each 62.5 ng/ μ L) or GFP-dsRNA (187.5 ng/ μ L) as a control. Bars represent mean values (\pm SD) of three biological replicates. Survival data were evaluated by Kaplan–Meier analysis and log-rank test. Different letters indicate significant differences at p < 0.0001

survival when administered as single dsRNA doses (see Fig. 2). We found that *Sa*'s survival rates treated with either *SaMIF1*-dsRNA or the mixture of *SaMIF*-dsRNAs were significantly reduced (Kaplan–Meier analysis and log-rank

test, $p \le 0.0001$) after 5 days as compared with *GFP*-dsRNA treatments (Fig. 4). This suggests that the activity of single dsRNAs is not additive but might have a synergistic effect on the aphid mortality instead.

SaMIF1-dsRNA spray application to barley seedlings had no effect on aphids' survival

It has been controversially discussed as to whether application of exogenous dsRNA to plants results in its accumulation in the phloem tissue, which is a prerequisite for the RNAi-based control of phloem-feeding insects (Gogoi et al. 2017; Dalakouras et al. 2018). We investigated the possibility that direct application of SaMIF1-dsRNA to plants have an effect on Sa's survival, when feeding from these plants. Therefore, three barley seedlings per pot were sprayed with 10 μg SaMIF1-dsRNA (500 μL of a 20 ng/μL solution), and seedlings, which were infested 24 h later with 50 one-dayold Sa nymphs, kept in confined jars (Fig. S3a). Compared to GFP-dsRNA-treated control plants, we found no significant differences in the survival rates of aphids feeding on SaMIF1-dsRNA-treated plants (Fig. S3b) and controls. This finding implies that spray application to leaves does not result in the accumulation of sufficient amounts of dsRNA or small RNA duplexes derived from it in the sieve tubes and suggests that spray-treatment of naked, unformulated dsRNA probably does not meet the requirements of efficient crop protection.



Sucrose-aided dsRNA delivery to barley leaves had no effect on aphids' survival

Next, alternative experimental designs were evaluated for simple and rapid screening of potential dsRNA targets for aphid control. Oligodeoxynucleotide (ODN)-directed gene silencing in barley is mediated by passive vascular feeding of ODN through cut leaves in sucrose solution via co-import of sucrose and negatively charged ODN molecules (Sun et al. 2005), suggesting that ODNs reached the leaf symplast and entered living cells. This report, together with accumulating evidence for xylem-to-phloem solute transport (van Bel 1990) and the presence of exo/endocytosis mechanisms in xylem vessels (Botha et al. 2008; Słupianek et al. 2019), prompted us to investigate whether the cut leaf delivery method could also be used to deliver dsRNA molecules to plant cells, including the phloem tissue. Detached leaves from two-week-old barley seedlings were dipped with the

basal end into 1 mL of a solution of 200 mM sucrose and 20 µg SaMIF1-dsRNA (Fig. S4a), and kept in the dark for 24 h. As shown in Fig. 5a–c, dsRNA was taken up through the cut ends as revealed by the detection of fluorescence in upper segments of the detached leaves. In barley leaf cross sections, fluorescence was associated with the vascular bundle, especially the xylem parenchyma cells (Fig. 5d–g). Note that bigger xylem vessels lose their content during preparation of cross sections due to flushing with the fluid set free by the cut cells and thus do not show fluorescence.

Next, the survival of *Sa* on *SaMIF1*-dsRNA *versus GFP*-dsRNA-treated detached barley leaves was recorded after seven days of infestation. Overall, there was no significant difference in the *Sa*'s survival rates on treated and control leaves (Fig. S4b). Consistent with this finding, no difference was found in the expression of the *SaMIF1* target gene in *Sa* fed on *SaMIF1*-dsRNA *versus GFP*-dsRNA treated leaves (Fig. S4c).

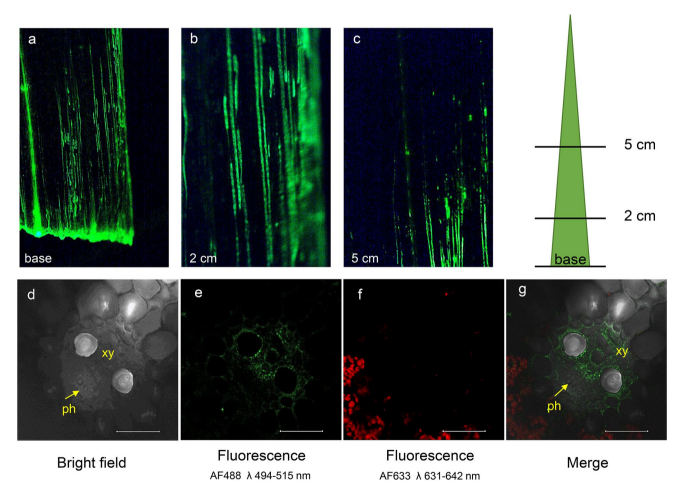


Fig. 5 Confocal images of detached barley leaves having absorbed fluorescence-labeled *SaMIF1*-dsRNA_{A488} through cut basal ends. The leaf base was submerged in 1 mL of 200 mM sucrose solution containing 20 μg dsRNA. Surface views of **a** leaf base; **b** leaf segment 2 cm away from the base at 24 h after onset of soaking. **c** leaf segment 5 cm from the base 48 h after onset of soaking. **d**-**g** leaf cross

section. (3 cm from the cutting), photographs taken at three days after onset of the <code>SaMIF1-dsRNA_A488</code> treatment. The green color represents the fluorescence ($\lambda_{\rm exc}$ 494, $\lambda_{\rm em}$ 515 nm) of the Alexa Flour 488 (AF488) dye. The laser filter AF633 ($\lambda_{\rm exc}$ 631 nm, $\lambda_{\rm em}$ 642 nm) was used for the detection of red fluorescence of chloroplasts. xy, xylem; ph, phloem; bs, bundle sheath



To further substantiate this finding, we conducted the sucrose-aided RNA uptake experiment with *SaSHP*-dsRNA (470 bp; see Table S2), which is known to target the *SaSHP* gene thereby strongly reducing the survival of the aphids on barley (Abdellatef et al. 2015). As shown in Fig. S4b, feeding on *SaSHP*-dsRNA treated leaves also had no effect on aphids' survival and expression of the *SHP* gene in *Sa* was not affected (Fig. S4d).

Petiole-mediated uptake of 21 nt dsRNA_{Cy3} in *Arabidopsis* follows the xylem-route

To further confirm the absence of microscopically detectable exchange of dsRNA between xylem and phloem vessels, when dsRNA is supplied via petioles, we used the *Arabidopsis* reporter line *Arabidopsis* thaliana *SUC2::4xYFP*, in which the promoter of the phloem-specific *SUC2* is fused with *Yellow fluorescent protein* (YFP), allowing visualization of the sieve tubes (Marquès-Bueno et al. 2016). Leaves from thirty-two-day-old plants were inserted with the petioles in nuclease-free water containing fluorescent

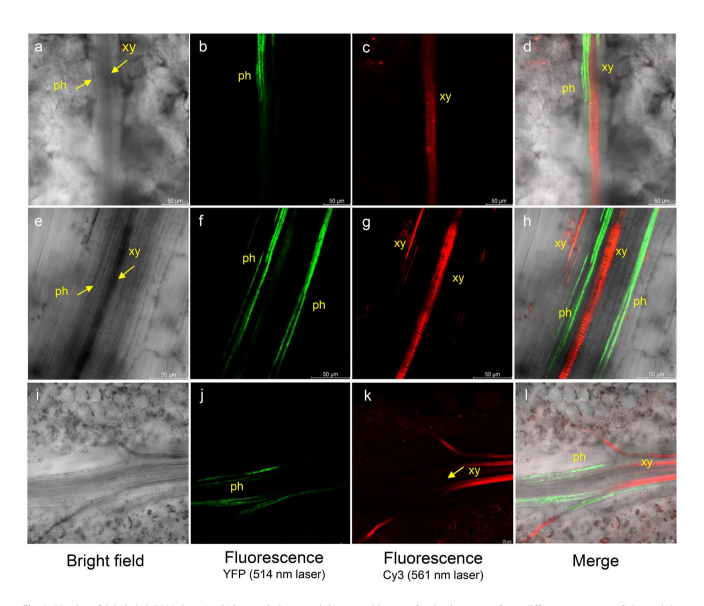


Fig. 6 Uptake of labeled dsRNA into *Arabidopsis thaliana* petioles and leaves. Confocal images of the reporter line *SUC2::YFP.* **a–h** cut petiole ends were submerged for 24 h in nuclease-free water containing 20 μ M 21-nt siRNA_{Cy3} and cross sections were examined at the base (**a–d**) and in the middle of the petiole (**e–h**). **i–l** leaves were dropped with 21 nt dsRNA_{Cy3} (20 μ M) for 24 h. Images were taken

with a confocal microscope from different segments of the petiole. The red color, which is restricted to the xylem vessels, represents Cy3 fluorescence ($\lambda_{\rm exc}$ 561nm, $\lambda_{\rm em}$ 566–635 nm) and the green color represents the phloem-based YFP fluorescence ($\lambda_{\rm exc}$ 514 nm, $\lambda_{\rm em}$ 519–551 nm). xy, xylem; ph, phloem



21 nt dsRNA $_{\text{Cy3}}$ (20 μ M). After 24 h, confocal images were taken from different segments of the petioles. We found that dsRNA $_{\text{Cy3}}$ was localized in the xylem, and its signal did not overlap with the YFP fluorescence of the phloem (Fig. 6a–h). Moreover, sucrose-aid uptake by petioles resulted in the same localization of Cy3 fluorescence in the xylem vessels (Fig. S5). This result is consistent with our finding that the survival of aphids is not negatively affected when they feed on leaves treated with dsRNA supplied via cut leaf ends. Thus, in contrast to reports showing that ODN can be introduced into plant cells via cut leaf ingestion, our data show that this method of introduction does not result in sufficient uptake of dsRNA or small RNA derivatives to affect aphids or be detected by fluorescence techniques.

dsRNA delivery to leaves also follows the xylem route

Finally, we used the *Arabidopsis thaliana SUC2::4xYFP* reporter line to visualize the uptake of fluorescence dsRNA from the leaf surface (Fig. 6i–l). *Arabidopsis* leaves were treated with four 1 μL droplets containing 20 μM dsRNA $_{Cy3}$. After five days, confocal images were taken from different segments of the leaves. We found that dsRNA $_{Cy3}$ was localized in the xylem, and its signal did not overlap with the YFP fluorescence of the phloem. This finding supports our notion that leaf-applied naked dsRNA does not reach the plant symplast and is therefore an inappropriate method for aphid control.

Materials and methods

Plant material and aphids rearing

Spring barley (*Hordeum vulgare* L.) cv. Golden Promise (GP) was used in all experiments. *Arabidopsis thaliana* (Col-0) *SUC2::4xYFP* lines were purchased from NASC (N2106107). Plants were grown under controlled conditions in a climate chamber at 22 °C/18 °C day/night with 65% relative humidity, a 16 h photoperiod and a photon flux density of 240 μmol m⁻² s⁻¹. *Arabidopsis* seedlings were grown in vertical plates containing half-strength MS medium (Murashige and Skoog 1962), 0.5% of sucrose and 0.7% of agar. The grain aphid (*Sitobion avenae*, *Sa*) monoclonal population used in this study was reared on three-week-old GP plants in a climate chamber at 22 °C with a 16 h photoperiod and a photon flux density of 240 μmol m⁻² s⁻¹. One-day-old fresh synchronized nymphs were used for the experiments (Abdellatef et al. 2015).



RT-qPCR was performed with the Applied Biosystems QuantStudio 5 Real-Time PCR system. Amplifications were performed with SYBR® green JumpStart Taq ReadyMix (Sigma-Aldrich). To quantify the target genes expression, the transcript was normalized with *Ribosomal gene L27 (RPL27*, NM_001126221.2) (Table S1) (Zhang et al. 2013). The program was performed with 95 °C for 5 min, 40 cycles (95 °C for 30 s, 57 °C for 30 s, 72 °C for 30 s. Transcript levels of genes were determined via the $2^{-\Delta\Delta Ct}$ method (Livak and Schmittgen 2001) by normalizing to the amount of reference gene transcript.

dsRNA synthesis

The Si-Fi software was used to select the donor sequences for the RNAi design (Luck et al. 2019). SaMIF genes were cloned into pGEM-T-easy vector, using the degenerate primers listed in Table S1, and the resulting plasmids were used as templates for the synthesis of dsRNA. Plasmids pGEM-Teasy-SHP and pGEM-T-easy-GFP contain respective SaSHP and GFP gene sequences (Table S2). The target sequences were amplified from the plasmid DNAs using primers containing T7 polymerase promotor or phi6 polymerase promoter sequences at their 5'-end (Table S1). SaMIF2-, SaSHP- and GFP-dsRNAs were produced using a singletube transcription and replication reaction catalyzed by the T7 DNA-dependent RNA polymerase and the phi6 RNAdependent RNA polymerases (Aalto et al. 2007; Levanova and Poranen 2018). The produced dsRNAs were enriched using stepwise fractionation with LiCl, followed by precipitation with sodium acetate and thorough washing of the resulting pellet with 70% ethanol. Alternatively, SaMIF- and GFP-dsRNAs were generated using MEGAscript T7 Transcription Kit (Thermo Fisher Scientific) following the manufacturer's protocol. The produced dsRNAs were resuspended in RNase-free milliQ-water and stored at -20 °C prior use.

Fluorescence labeling of dsRNA

Fluorescence labeling of *SaMIF1*-dsRNA was performed using the HighYield T7 AF488 RNA Labeling Kit (Jena Bioscience, Germany) following the manufacturer's instruction. Labeled *SaMIF1*-dsRNA_{A488} was used for the uptake experiments. For uptake analysis of small RNA, 21 nt *GAPDH*-dsRNA (provided in the kit) was labeled with CyTM3 utilizing the SilencerTM siRNA Labeling kit (ThermoFisher) according to the manufacturer's instructions.



Feeding of aphids on dsRNA supplemented artificial diet

The rearing method as described by Will et al. (2012) was used with minor modifications. The artificial diet (50 mM L-serine, 50 mM L-methionine, and 50 mM L-aspartic acid; pH 7.2) containing different sucrose concentrations was sealed between two layers of parafilm in a 2 cm diameter feeding tube, and one-day-old *Sa* nymphs were placed on the plates. The plates were covered with a feeding tube. The diet was prepared with RNase-free water. For dsRNA feeding experiments, the dsRNA was mixed with the artificial diet. Ten synchronized nymphs with five replicates for each sample were used. Nymphs were placed at 22 °C under 65% relative humidity, with a photoperiod of 16 h and a photon flux density of 125 µmol m⁻² s⁻¹.

Application of dsRNA

Three-week-old barley seedlings (each pot with three plants) were first sprayed with 0.02% Silwet-77, and 10 min later with 10 μ g dsRNA solved in 500 μ l deionized water. Controls were sprayed with 500 μ L of deionized water. After spraying, the plants were infested with 50 Sa nymphs and stored in closed jars. Seven days later, the number of aphids was counted.

For treatment of *Arabidopsis* leaves, 19-day-old *Arabidopsis SUC2::4xYFP* seedlings grown in vertical plates were treated with 1 μ L drop of nuclease free-water containing 20 μ M dsRNA_{Cy3} at on the top of the leaf. Four leaves were treated. Confocal images were taken 5 days later.

dsRNA delivery via the sucrose-aid method

Ten-day-old barley seedlings were transferred to the dark for 12 h. Leaves were detached and submerged with the basal end into 200 mM sucrose solution containing 20 µg/mL dsRNA for 24 h in the dark. Subsequently, the submerged parts of the leaves were cut and the top segment transferred to agar plates and used for aphid infestation. Thirty-two-day-old Arabidopsis leaves were cut and inserted with the petiole in nuclease-free water containing 20 µM dsRNA $_{\rm Cy3}$. For the sucrose-aid experiment, the solution was supplemented with 200 mM sucrose.

Microscopy

Cross hand-cut sections of barley leaves were analyzed using a confocal laser-scanning microscopy (CLSM, Leica, TCS SP8, Germany). Green fluorescence of dsRNA_{A488} was detect by filter AF488 ($\lambda_{\rm exc}$ 494, $\lambda_{\rm em}$ 515 nm). The laser filter AF633 ($\lambda_{\rm exc}$ 631 nm, $\lambda_{\rm em}$ 642 nm) was used for the detection of red fluorescence of chloroplasts. *Arabidopsis* leaves

were visualized with the CLSM microscope (previously described) for fluorescence YFP ($\lambda_{\rm exc}$ 514 nm, $\lambda_{\rm em}$ 519–551 nm) and Cy3 ($\lambda_{\rm exc}$ 561 nm, $\lambda_{\rm em}$ 566–635 nm). YFP was excited with the 514 nm laser (detection 519–551 nm) and Cy3 with the 561 nm laser (detection 566–635 nm).

Discussion

We show here that members of the Macrophage migration inhibitor factor (MIF) protein family are necessary for the survival of the aphid *Sitobion avenae*. We found that *Sa* contains four *MIF* genes and that silencing of three of them, namely *SaMIF1*, *SaMIF2* and *SaMIF3*, leads to reduced aphid survival on artificial diet. This corroborates findings that MIFs, apart from their roles in suppressing host immunity, also have an endogenous function in the aphid (Naessens et al. 2015). dsRNAs targeting individual *SaMIF* genes were effective at a concentration of 250 ng/μL. At lower concentration (125 ng/μL), only dsRNA directed against the *SaMIF1* transcript reduced target gene expression substantially, suggesting the possibility that *SaMIF1* could be a potential target candidate for aphid control by RNAi.

Functionally redundant MIF gene family members are widespread in eukaryotic genomes, which often hampers the analysis of gene families, due to functional redundancy (Jover-Gil et al. 2014; Martienssen and Irish 1999). For functional analysis, silencing of the entire set of paralogous genes at the same time is a straightforward approach. Simultaneous targeting of three out of the four known SaMIF genes using three SaMIF gene-specific dsRNAs caused a significant reduction in survival, when compared with the activity of a GFP-dsRNA that had no known target in Sa (Fig. 4). Interestingly, when applied in mixtures, SaMIFdsRNAs had a synergistic effect as they affected survival of Sa at a concentration that showed no effects upon single delivery. Overall in spite of these findings, our data suggest that SaMIF1 is a candidate for aphid control and it is probably not required to consider the other SaMIF genes.

In our experiments, different dsRNA delivery strategies were investigated to test the efficiency of RNAi-mediated control of insects. Oral feeding on artificial diet containing SaMIF1-dsRNA showed the highest mortality rate (Fig. 2) and concomitant downregulation of SaMIF1 target transcripts (Fig. 3). In contrast, spraying SaMIF1-dsRNA onto leaves had no effect on the survival rate of nymphs fed on these leaves (Fig. S3). This result can be explained by the fact that the SaMIF1-dsRNA applied to the leaves did not reach the sieve tubes in amounts sufficient to silence the SaMIF1 target gene, though it cannot be excluded that spraying leaves with higher concentration of dsRNA would have an effect on aphid survival. Uptake of dsRNA via the leaf surface has been controversially discussed. Gogoi et al.



(2017) found that aphids take up, among others, a 588 bp long dsRNA from tomato leaves. It should be noted, however, that the dsRNA was applied by gently rubbing the solution onto the upper side of tomato leaflets that were previously carborundum-dusted. Subsequently, the treated leaves were thoroughly washed with 0.05% Triton X-100 for five times in 3 min intervals. Moreover, dsRNA-mediated protection was obtained in tobacco against viral diseases, when leaves were spread with virus-specific dsRNA loaded on non-toxic, degradable, layered double hydroxide (LDH) clay nanosheets (Mitter et al. 2017). Once loaded on LDH, the dsRNA did not wash off, showed sustained release and could be detected on sprayed leaves even 30 days after application. Finally, it was recently reported that strong GFP transgene silencing was accomplished in tobacco and tomato by loading dsRNA into carbon dots (Schwartz et al. 2020). Chemical formulations not only enhance the uptake of RNA from leaves, but could also improve dsRNA penetration into an insect, as shown for a polymer/detergent formulation that improves RNAi-induced mortality in the soybean aphid Aphis glycines (Zheng et al. 2019). In the light of these reports, research should focus on dsRNA delivery strategies that might support more efficient use of RNAi-based plant protection.

Feeding of Sa on barley leaves immersed at the base in SaMIF1-dsRNA containing buffer also did not affect aphids' survival nor could we detect an effect on SaMIF1 target gene expression (Fig. S4). This setup was tested to evaluate an alternative experimental design for simple and rapid screening of potential dsRNA targets for aphid control. In agreement with a lack of effect on aphids, we could not detect fluorescence in phloem tissue when barley leaves had been submerged into fluorescence SaMIF1-dsRNA_{A488} solution. Instead, we detected fluorescence predominantly in the xylem parenchyma cells, mainly the contact cells (Fig. 5a–g). This is in agreement with earlier reports, where apical transport of exogenous dsRNA structurally is located within xylem structures (Dalakouras et al. 2018, 2020). While the latter reports and our investigation support the view that dsRNA application onto leaves and via petioles results in the accumulation of RNA in the xylem, some reports challenge this generalized view: (1) ODN-directed gene silencing in barley is mediated by passive vascular feeding of ODN through cut barley leaves using co-import of sucrose and negatively charged ODN molecules (Sun et al. 2005), resulting in ODN uptake into the leaf symplast and living cells. (2) The importance of the xylem-to-phloem pathway was underscored in a review that summarized work of the precedent two decades (van Bel 1990). Moreover, it is well accepted that exo/endocytosis processes are involved in the uptake of macromolecules from xylem tissue (Botha et al. 2008; Słupianek et al. 2019). (3) Turnip mosaic virus (TuMV) is a single-stranded RNA virus that can cause

diseases in cruciferous plants. Viral RNA can move systemically through both phloem and xylem as membrane-associated complexes in plants (Wan et al. 2015).

Trafficking of vesicles carrying sRNAs has been observed between Arabidopsis and Botrytis cinerea (Cai et al. 2018; Šečić and Kogel 2021). Exosomes derived from Tetraspanin-GFP Arabidopsis line could be visualized as fluorescent dots, demonstrating that sRNA transfer occurs via exosomes. Trafficking of sRNA in vesicular bodies might explain why fluorescence appears in a punctate manner in traversal leaf section (Fig. 5a-c). As RNA transferred from one cell to another via exo/endocytosis would result in vesicles, a dotted fluorescence pattern would indeed occur. The fact that we could not detect dsRNA_{A488} fluorescence in the barley phloem tissue led us to further experiments to substantiate a xylem-associated uptake of dsRNA. We repeated the RNA uptake experiments with the Arabidopsis SUC2::4xYFP reporter line, which is a more sensitive tool to distinguish between transport of solutes in xylem and phloem. When taken up by petioles, we detected 21 nt dsRNA_{Cv3} exclusively in the Arabidopsis xylem, and its signal did not overlap with the YFP fluorescence of the phloem (Fig. 6a-h). This result further substantiates the previous report showing that dsRNA uptake and its acropetal transport follows mainly the apoplastic route via the xylem (Dalakouras et al. 2018). It also shows that a possible exchange of dsRNA from xylem-to-phloem is not efficient enough to be detected in our fluorescence microscopy experiment nor to silence genes from aphids feeding on the phloem at least at the concentrations used here. Nevertheless, soaking roots in dsRNA solution conferred protection in rice and maize against stem-borer (Li et al. 2015), further showing the potential of the approach. We also used the Arabidopsis SUC2::4xYFP reporter line to follow the uptake of 21 nt dsRNA_{Cv3}, upon dropping onto leaves (Fig. 6i-l). In agreement with the results from the barley spray experiments, we could detect fluorescence exclusively in the leaf xylem. While fluorescence imaging is sensitive and a well-accepted method, final proof of the absence of exogenously applied dsRNA in the symplast, e.g., in mesophyll cells and sieve tubes in missing. In particular, the observation that virus-specific dsRNA, when scattered on leaves, is effective in reducing viral infections suggests that dsRNA—possibly assisted by physical means such as formulations and gentle leaf wounding—can lead to symplastic uptake of dsRNA.

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Data availability All data generated or analyzed during this study are included in this published article [and its supplementary information files].

Declarations

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Consent for publication All authors declare consent of publication.

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- (1) Made substantial contributions to the conception or design of the work; or the acquisition, analysis, or interpretation of data; or the creation of new software used in the work;
- (2) Drafted the work or revised it critically for important intellectual content;
- (3) approved the version to be published; and
- (4) AGREE to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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