microRNA390 modulates *Nicotiana attenuata*’s tolerance response to *Manduca sexta* herbivory

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**Abstract**

miR390 is a highly conserved miRNA in plant lineages known to function in growth and development processes, such as lateral root development, and in responses to salt and metal stress. In the ecological model species, *Nicotiana attenuata*, miR390’s biological function remains unknown, which we explore here with a gain-of-function analysis with plants over-expressing (OE-) *N. attenuata* miR390 (Na-miR390) in glasshouse and natural environments. OEmiR390 plants showed normal developmental processes, including lateral root formation or reproductive output, in plants grown under standard conditions in the glasshouse. OEmiR390 plants did not have dramatically altered interactions with arbuscular mycorrhizal fungi (AMF), *Fusarium* pathogens, or herbivores. However, Na-miR390 regulated the plant’s tolerance of herbivory. Caterpillar feeding elicits the accumulation of a suite of phytohormones, including auxin and jasmonates, which further regulate host-tolerance. The increase in Na-miR390 abundance reduces the accumulation of auxin but does not influence levels of other phytohormones including jasmonates (JA, JA-Ile), salicylic acid (SA), and abscisic acid (ABA). Na-miR390 overexpression reduces reproductive output, quantified as capsule production, when plants are attacked by herbivores. Exogenous auxin treatments of herbivore-attacked plants restored capsule production to wild-type levels. During herbivory, Na-miR390 transcript abundances are increased; its overexpression reduces the abundances of auxin biosynthesizing *YUCCA* and *ARF* (mainly *ARF4*) transcripts during herbivory. Furthermore, the accumulation of auxin-regulated phenolamide secondary metabolites (caffeoylputrescine, dicaffeoylspermidine) is also reduced. In *N. attenuata*, miR390 functions in modulating tolerance responses of herbivore-attacked plants.

**KEYWORDS**

ARF4, auxin, herbivore resistance, *Manduca sexta*, miR390, *Nicotiana attenuata*, tolerance, *YUCCA*
microRNAs (miRNAs) form an essential layer of regulators of gene expression in nearly all eukaryotes. They are noncoding RNA molecules, produced from the stem regions of hairpin precursor molecules. miRNAs are produced from “miRNA genes” in an RNA-polymerase II dependent manner (Song et al., 2019). Most miRNA genes are found in intergenic regions and are transcribed as independent units. Several miRNAs are also intronic and cotranscribed with their host protein coding regions (Song et al., 2019). The size of mature miRNAs ranges from 18–25 nucleotides (nt), with 21–22 nt long miRNAs being the most abundant in plants. miRNAs recognize their targets through sequence complementarity, where they facilitate mRNA cleavage, translational repression, and DNA methylation (Pandey et al., 2019; Song et al., 2019). miRNAs are thought to play indispensable roles in a plant’s phenotypic plasticity, being involved in the regulation of differentiation, development and reproduction, and mediating responses to environmental fluctuations (Brant & Budak, 2018; Li et al., 2017; Song et al., 2019). In studies that have manipulated the expression of different components of the small-RNA (smRNA) pathways, evidence for roles in the diverse plastic responses that allow plants to adapt to environmental challenges in natural habitats has been reported (Pandey et al., 2008; Pandey & Baldwin, 2007; Pradhan et al., 2017, 2020).

The effectors of miRNA action, the Argonaute (AGO) proteins, are conserved across all lineages of photosynthetic organisms (Pradhan et al., 2020), indicating that miRNA-driven regulation of gene expression could be ancient. However, individual miRNAs/miRNA-families appear to evolve rapidly, as suggested by their frequent birth and loss in lineage- and species-specific manners (Cuperus et al., 2011; Nozawa et al., 2012). Most miRNA genes appear to be relatively young (Cuperus et al., 2011), and only a few miRNA families are conserved across plant lineages (Cuperus et al., 2011; Nozawa et al., 2012). miRNA930 (miR390) is one of these, being highly conserved across lineages of vascular and nonvascular plants (Morozov et al., 2018; Xia et al., 2017). Studies focusing on evolution of miR390 have highlighted significant variations in its regulatory circuit (Xia et al., 2017). Yet, the exploration of diversification of miR390 biological functions has remained limited.

At a molecular level of analysis, miR390 acts as an activator of the trans-acting small interfering RNA (tasiRNA) pathway, with two known target sites in the transcript of the tasiRNA precursor gene, TAS3 (Axtell et al., 2006). Following miR390-directed cleavage, tasiRNAs are produced from TAS3 transcripts, which regulate auxin response factors (ARFs). ARFs are a class of transcription factors that modulate (repressing or activating) the expression of genes regulated by the plant hormone, auxin. In this way, miR390 functions as a central component of the highly conserved “miR390-TAS3-ARF regulatory pathway” (Xia et al., 2017), which is expected to modulate auxin signaling. In turn, in Arabidopsis, miR390 expression itself is regulated by auxin levels (Yoon et al., 2010).

miR390 has been regarded as a key regulator of developmental events in several plant species. miR390 has been described to mediate lateral root growth, developmental timings, and patterning in Arabidopsis and Physcomitrella patens (Cho et al., 2012; Marin et al., 2010). During potato development, miR390 regulates a calcium-dependent protein kinase (CDPK) that phosphorylates auxin efflux carriers (Santín et al., 2017). Interestingly, miR390 has also been implicated in abiotic stress responses, in particular, salt and metal stress, by regulating lateral root growth (Ding et al., 2016; Dmitriev et al., 2017; He et al., 2014, 2018; Lu et al., 2018). The miR390/TAS3 regulatory module also plays an important role in gall formation when Arabidopsis roots are attacked by the nematode Meloidogyne javanica (Cabrera et al., 2016). Infection by root-knot nematodes induces the expression of miR390/TAS3 smRNA, which is necessary for gall formation, possibly through ARFs (Cabrera et al., 2016). Legume root colonization by symbiotic rhizobia also requires the regulatory miR390/TAS3 module to control rhizobial root colonization and nodule formation. Overexpression of miR390 promotes lateral root growth while negatively regulating nodule organogenesis, rhizobial infection, and induction of nodulation signaling pathway (NSP) 1 and 2 genes (Hobecker et al., 2017). These studies indicate a role of miR390 not only in developmental processes but also during different stress responses and symbiotic interactions. However, the role of miR390 in modulating host responses to other stress factors, such as pathogen and herbivore attack, as well as interactions with arbuscular mycorrhizal fungi (AMF) have not been thoroughly studied. Moreover, the functional role of miR390 has not been examined in plants growing in their natural habitats.

Nicotiana attenuata is an annual plant native to the southwestern United States and is an ecological model as its natural history and its interactions with herbivores, pollinators, and microbes have been extensively studied not only under glasshouse conditions but also in its natural habitats (Baldwin, 2001; Navarro-Quezada et al., 2020). This wild tobacco species grows in the agricultural primordial niche: the post-fire habitat. After wild fires, N. attenuata germinates from long-lived seed banks and establishes mono-culture populations, while encountering several ecological challenges. The plant is colonized by AMF (Pandey et al., 2018) and is also attacked by pathogens such as Fusarium and Alternaria species (Schuck et al., 2014). Furthermore, N. attenuata faces an unpredictable attack by a diverse suite of herbivore species, which is thought to represent an important ecological challenge for this species; consequently, N. attenuata recruits several direct and indirect defenses (Baldwin, 2001; Navarro-Quezada et al., 2020). Additionally, N. attenuata has a remarkable capacity to tolerate herbivore attack to minimize fitness losses (Schwachtje et al., 2006). N. attenuata’s resistance and tolerance strategies, the signaling cascades, and defense-related secondary metabolites have been well studied. Moreover, the composition and role of the RNA-directed RNA polymerase (RdR), DCL, and AGO components of the smRNA machinery during adaptation to these natural challenges have also been studied (Navarro-Quezada et al., 2020). The composition of miRNAs in N. attenuata’s genome (Xu et al., 2017) and its reprogramming during herbivore attack, pathogen infection, and AMF colonization are also known (Pandey et al., 2018; Pradhan et al., 2017, 2020).
miR390 has been identified in *N. attenuata*: its expression is up-regulated during simulated herbivory (Bozorov et al., 2012), whereas it is down-regulated during interactions with AMF (Pandey et al., 2018). Yet, the biological function of this highly conserved miRNA remains poorly studied in *N. attenuata* and is the subject of this investigation. We performed gain-of-function studies by stable over-expression (OE-) of *N. attenuata* miR390 (Na-miR390) to evaluate changes in performance when plants engage with other organisms, specifically, with AMF, a native pathogen (*Fusarium brachygibbosum*) and a native herbivore (*Manduca sexta*).

### RESULTS

#### 2.1 miR390 does not regulate development events in *N. attenuata*

In *N. attenuata*, miR390 sequences have been reported in two previous independent investigations (Bozorov et al., 2012; Pandey et al., 2018). Na-miR390a-i (Bozorov et al., 2012) and Na-miR390-5p-ii (Pandey et al., 2018) are nearly identical as they vary in only a single nucleotide (A/G) at the 19th position of the mature sequence (Figure S1). We compared the *N. attenuata* miR390 sequences to those of other species (Figure S1). Both sequence variants have been reported in other plant species: For instance, Gm-miR390 contains an “A” at 19th position, whereas Mt- and At-miR390s contain the “G” (Figure S1). As there are no variations in 20 of 21 nucleotides, including the important residues of target recognition and cleavage site within the first 12 nucleotides, we do not expect significant effects of this single nucleotide variation in the regulation of targets of this miRNA.

To evaluate the biological function of this miRNA, we overexpressed the originally identified Na-miR390a-i in transgenic lines (OEmiR390) with the transformation vector shown in Figure S1 that carries the miRNA sequence as well as backbone sequence. After transformation, we obtained 20 independently transformed lines and screened them for homozygosity, occurrence of single insertions, completeness of the insertions (no truncations or border over-reads), and successful overexpression. For further experiments, we selected two homozygous lines, A-12-392-63-71 and A-12-385-42-51, which showed strong overexpression and harbored a single complete insertion (Figures 1a and S1).

Based on previous results for lines altered in miR390 expression (Marin et al., 2010), we measured seedling root lengths and the number of lateral roots (Figure 1b). No significant reductions in root length (one-way ANOVA, *F* = .621, *P* > .05) were observed for the two *N. attenuata* lines overexpressing miR390. Furthermore, in contrast to the results of studies with other plant species (e.g., rice (Lu et al., 2018) and Arabidopsis (Marin et al., 2010)), the number of lateral roots in OEmiR390 *N. attenuata* plants was also not significantly altered (one-way ANOVA, *F* = .65, *P* > .05) as compared to those of WT seedlings (Figure 1b).

As miR390 is thought to play a key role in the juvenile-to-adult phase transition of many species, we grew the two miR390-overexpressing lines in individual pots with full nutrient supply and documented growth parameters indicative of changes in phase transitions. The two OEmiR390 lines did not show any significant differences in rosette size (one-way ANOVA, *F* = 1.17, *P* > .05), stalk length (one-way ANOVA, *F* = 2.91, *P* = .183), flower (one-way ANOVA, *F* = 1.228, *P* > .05) and capsule numbers (two-way repeated measures ANOVA with Bonferroni correction, *F* = 1.87, *P* > .05 and Kruskal-Wallis test) when compared with WT plants (Figure 1c,d). These results indicated that overexpression of miR390 does not dramatically change plant development or fitness in *N. attenuata* under standard, unstressed growth conditions of the glasshouse.

#### 2.2 Na-miR390 does not affect resistance, colonization, or reproductive output during plant-pathogen and plant-AMF interactions in *N. attenuata*

Previous studies have reported that the expression of miR390 is altered during a plant’s interaction with microbes (Hobecker et al., 2017; Pandey et al., 2018), and this motivated our search for its role in plant-AMF and plant-pathogen interactions. To examine if miR390 is a crucial factor for AMF root colonization, similar to its role in symbiotic nodulation in legumes (Hobecker et al., 2017), we inoculated plants grown under reduced inorganic phosphate (Pi) fertilization regimes with AMF, observed growth and development, and determined root AMF colonization by measuring the levels of 11-carboxyblumenol-C glucoside in leaves as a proxy for AMF colonization (Wang et al., 2018). Blumenol levels were not significantly different among the WT and miR390-overexpressing lines (Figure 2a). Furthermore, rosette diameters, flower and capsule numbers were not different between WT and OEmiR390 plants (Figure S2). Only stalk lengths were marginally higher for one of the two lines (Figure S2). Similar results were obtained for plants grown individually in 1 L pots as well as for OEmiR390 plants competing with an initially size-matched WT neighbor (Figure S2). The marginal increase in stalk length had no fitness consequences as seed capsule production did not differ between the WT and the miR390 overexpressing plants.

To evaluate if Na-miR390 is involved in the resistance response to a natural fungal pathogen, single leaf inoculations with *Fusarium brachygibbosum* Utah 4 isolates were performed (Pradhan et al., 2020), and the development of visible disease symptoms was recorded. No significant differences in the diameter of chlorotic/necrotic lesions of WT and miR390-overexpressing plants were observed during infection with the *Fusarium* pathogen (Figures 2b and S3).

These results suggest that Na-miR390 does not play a significant role in these plant-fungal (pathogen and AMF) interactions.

#### 2.3 Na-miR390 has a role in host tolerance response during herbivory

miR390 transcripts accumulate 1 h after simulated herbivory in *N. attenuata* (Bozorov et al., 2012). We confirmed this upregulation of
**Figure 1** Phenotypic characterization of two independent, homozygous OEmiR390 lines (392, 385). (a) miR390 was overexpressed in *N. attenuata* (OEmiR390 plants) in two independent lines. Wild type seeds and seeds of the two OEmiR390 lines (392, 385) were germinated on GB5 medium. Transcript accumulation of miR390 was analyzed in 10 day-old seedlings by qPCR with miR390 specific primers (*N* = 3, mean ± SE). Different letters indicate significant differences based on one-way ANOVA with Tukey’s HSD. (b) The two lines do not differ in root length and the number of lateral roots 10 days after germination (*N* = 5–6, mean ± SE). Each replicate represents 5 pooled seedlings from an independent GB5 germination Petri dish. n.s., no significant difference based on ANOVA analyses, *p* > .05. (c) Growth and (d) fitness data were recorded of non-treated OEmiR390 and WT plants grown under glasshouse conditions in potting soil. Stalk lengths, rosette diameters and flower numbers were taken after 9 weeks, and capsule numbers after 11 weeks (*N* = 4, mean ± SE). n.s., no significant difference based on ANOVA and Kruskal–Wallis test.

**Figure 2** miR390 does not play a role in arbuscular mycorrhizal root colonization, or in pathogen defense in *N. attenuata*. (a) The two lines overexpressing miR390 (392, 385) were grown in 2 L pots in competition with wild type (WT) plants with AMF inoculum and the levels of leaf 11-carboxyblumenol-C glucoside as marker of arbuscule colonization were measured 6 weeks after inoculation (*N* = 8–9, mean ± SE). (b) In rosette-stage plants grown individually in 1-L pots, leaves were infiltrated with a spore suspension of *F. brachygibbosum*. Lesion diameters were measured and photographed 7 days after infection (*N* = 10, mean ± SE); n.s., no significant difference based on ANOVA analyses.
wounding and application of *M. sexta* oral secretion (simulated herbivory) in WT plants, while the expression was not further enhanced in the overexpressing lines (Figure 3a). To examine Na-miR390’s functional role in plant–herbivore interactions, we used two established approaches to evaluate both the herbivore and plant sides of the interaction: (i) We measured the growth of *M. sexta* larvae feeding on WT and the two lines overexpressing Na-miR390. This “caterpillar performance bioassay” evaluates if the defenses and/or nutritional capacity of host plants is altered (Pandey & Baldwin, 2007; Pradhan et al., 2017). (ii) We recorded plant growth and reproductive output/plant fitness in WT and OEmiR390 plants during herbivory. The maintenance of reproductive output in herbivore-attacked plants reflects a plant’s tolerance of herbivore attack (Schwachtje et al., 2006).

(i) Caterpillar growth was not affected when consuming Na-miR390 overexpressing plants. Independent of plant genotype, *M. sexta* 1st to 4th instar caterpillars gained equivalent amounts of mass (two-way repeated measures ANOVA, *F* = .224, *P* = .78) when feeding on WT and OEmiR390 plants (Figures 3b and S4). They also appeared to consume similar amounts of plant material from both the genotypes (Figure S4).

(ii) *M. sexta*-damaged OEmiR390 plants had taller stems than similarly damaged WT plants (one-way ANOVA with Bonferroni correction, *F* = 15.60, *P* = .00008; Figure 3c). However, herbivore-damaged OEmiR390 plants produced significantly fewer seed capsules (one-way ANOVA, *F* = 5.68, *P* < .01 and Kruskal-Wallis test followed by Dunn’s test) than did WT plants (Figure 3d), a decline seen in both the number of open and closed capsules (Figure S4). Independent feeding experiments confirmed these results (Figure S4). The significant reductions in capsule production of attacked plants suggested that miR390 expression influences *N. attenuata*’s tolerance of herbivory.

### 2.4 Overexpression of miR390 reduces the accumulation of auxin and auxin-regulated phenolamides during herbivory

To further understand the physiological basis of reduced herbivore tolerance of OEmiR390 plants, we quantified phytohormone levels in...
response to caterpillar feeding. Insect attack changes the accumulation of a suite of phytohormones that includes jasmonates, salicylic acid (SA), abscisic acid (ABA), and auxin. Auxin modulates herbivore-induced tolerance-defense trade-offs (Machado et al., 2013; Machado et al., 2016) along with jasmonates that also regulate induced plant defenses during herbivory (Howe & Jander, 2008). Auxin levels were significantly reduced (by up to nine-fold: to 10.6% and 28.8% of WT levels for lines 385 and 392, respectively) in herbivore attacked OEmiR390 plants as compared to WT plants (Figure 4a). In contrast, levels of jasmonic acid (JA) and its active conjugate, jasmonoyl-isoleucine (JA-Ile), did not significantly differ between the two OEmiR390 lines and the WT during herbivory (Figure 4a). Furthermore, SA and ABA levels were also not significantly different between the OEmiR390 and the WT plants (Figure S5).

Auxin regulates the accumulation of phenolamides, such as caffeoylputrescine and dicafeoyl spermidine, during herbivory (Machado et al., 2016). The reduced auxin accumulations in OEmiR390 plants during herbivory motivated us to investigate whether these metabolites were also affected. Indeed, both metabolites were significantly reduced in herbivore-damaged OEmiR390 plants as compared to the WT (Figure 4c). From these results, we infer that overexpression of miR390 specifically inhibits the accumulation of auxin and auxin-regulated metabolites during herbivore attack.

Additionally, we determined how the expression of GAL83 was affected by miR390. GAL83 is a regulatory subunit of an SnRK1-related kinase that is involved in plant tolerance to herbivory by regulating source-sink relationship of carbon assimilates (Schwachtje et al., 2006). No differences were observed in GAL83 levels in WT and OEmiR390 plants during herbivory (Figure 4c), or during constitutive conditions (no-herbivory controls; Figure S6).

### 2.5 Exogenous application of IAA restores tolerance to herbivory

The reduced accumulations of auxin as well as auxin-regulated phenolamides suggested that auxin was the critical factor in the reduced tolerance of OEmiR390 to herbivory. To test this hypothesis, we determined how the complementation of herbivore-attacked plants with exogenously supplied auxin (indole acetic acid [IAA], 1 mM as a lanolin paste to petioles) effects the plant’s reproductive performance. Indeed, when treated with exogenous IAA, herbivore-attacked OEmiR390 plants were able to restore their capsule production to the WT levels (Figure 5). The OEmiR390 plants that were fed by only M. sexta for 9 continuous days (no IAA complementation; feeding controls) had reduced capsules as compared to similarly attacked WT plants (Figure 5), whereas no differences in capsule productions were seen among the three lines when no herbivore stress (no-herbivory treatment controls) was applied (Figure 5). Both of these control datasets are consistent with the observations described above.

### 2.6 Overexpression of miR390 down-regulates expression of auxin biosynthesis (YUCCA) and auxin response factor (ARF) genes during herbivory

The rate-limiting step in auxin biosynthesis involves oxidative decarboxylation of indole-3-pyruvic acid (Figure 6). This reaction is catalyzed by the flavin monooxygenase-like proteins encoded by YUCCA genes (Machado et al., 2016). Expression of YUCCA genes, like YUCCA1 and YUCCA3, is elicited quickly within 60 min of simulated M. sexta attack in N. attenuata (Machado et al., 2016). We evaluated how the expression of these YUCCA genes were altered after overexpression of Na-miR390. OEmiR390 plants accumulated significantly fewer transcripts of YUCCA1 (reductions by 50% of WT levels) and YUCCA3 (reductions by 52%) during herbivory (Figure 6). No differences in constitutive levels (in no-herbivory controls) were observed between the WT and OEmiR390 plants (Figure S6).

ARFs are a demonstrated target of the miR390/TAS3 pathway, which modulate auxin signaling. Furthermore, ARFs, specifically ARF4 (Marin et al., 2010; Yoon et al., 2010), also often have feedback/homeostatic roles for miR390 expression during auxin signaling. Therefore, we analyzed the accumulation of ARFs in the Na-miR390 overexpression lines, 9 days after continuous herbivore feeding. The up-regulation of miR390 correlated with a down-regulation of ARFs, in particular of ARF4 (Figure 7): the levels of ARF4 mRNA were significantly down-regulated (by up to 3-fold reductions in comparison to the levels of WT plants) in herbivore-damaged OEmiR390 plants, whereas reductions in ARF2 were more modest and those of ARF3 were only marginally lower (Figure 7). The constitutive transcript levels of ARFs in the uninduced WT and the OEmiR390 plants were similar (Figure S6). These results were consistent with the inference that overexpression of miR390 had down-regulated ARF expression and auxin signaling during herbivory.

Overall, we infer that miR390 in N. attenuata plays a central role in optimizing plant fitness/tolerance responses during M. sexta herbivore attack by modulating auxin signaling (Figure 8).

### 2.7 Performance of OEmiR390 plants under natural conditions

The biological function of any miRNA has yet to been examined under field conditions. Our two decade long experience of planting transgenic plants into natural habitats has revealed that some phenotypes only become apparent when plants are grown under field conditions (Kessler et al., 2004), while other phenotypes, clearly seen in the glasshouse, are lost in the field, due to the multiplicity of environmental signals that plants are exposed to under natural conditions (Kaur et al., 2012). We therefore planted one of the miR390 overexpressing lines in the plant’s native habitat (in the Great Basin Desert of Utah, USA) in the 2019 field season, in size-matched pairs with WT plants and observed herbivore damage as well as growth and fitness parameters for 8 weeks of growth in the field. Rosette diameters were similar between the WT and the OEmiR390 plants (repeated measures
ANOVA, \( n = 17, F = 2.36, P = .115 \), while stalk lengths of OEmiR390 plants were marginally taller than WT plants (repeated measures ANOVA, \( n = 17, F = 10.702, P = .0395 \)) at the end of the field season (Figure S7). Overall, the WT and the OEmiR390 plants encountered mild attack rates from the natural herbivore community, resulting in only 10%–12% losses of rosette area after 3 and 4 weeks, which decreased to nearly 7% for the rest of the season. When herbivore damage (at 4 weeks after planting) was correlated with the number of capsules, OEmiR390 plants exhibited a steeper decline in capsule numbers at the higher early herbivore damage rates (Figure S7). It should be noted that for the 2019 field season, Manduca species did not arrive while the OEmiR390-WT plant pairs were in the field.

**FIGURE 4** Overexpression of miR390 reduces the production of auxin and auxin-dependent tolerance responses but not jasmonates during herbivory. (a) Levels of auxin, jasmonic acid (JA) and JA-isoleucine (JA-Ile) hormones were quantified in plants continuously fed on by M. sexta for 9 days. Auxin levels were strongly attenuated in miR390-overexpressing plants; one-way ANOVA, \( F = 43.17, n = 10–15 \), **** significantly different at \( P < .001 \). n.s., no significant differences in JA or JA-Ile amounts were observed between WT and OEmiR390 plants. (b) Evaluation of factors related to tolerance response of *N. attenuata* during herbivory. Peak areas of caffeoylputrescine (upper panel) and dicaffeoylspermidine (middle panel). Levels were measured in the leaves after 9 d of *M. sexta* feeding. One-way ANOVA, \( F = 1.8 \) for caffeoylputrescine and \( F = 4.54 \) for dicaffeoylspermidine. \( n = 10–15 \), * and ** represent significant differences from WT at \( P \leq .05 \) and \( P \leq .01 \). (c) Lower panel shows transcript abundance of GAL83 after 9 days of *M. sexta* feeding on WT and two miR390 overexpressing plants. No significant differences among genotypes were observed

### DISCUSSION

Here, we examined the biological function of miR390 in the ecological model plant species, *N. attenuata*. We show that Na-miR390 plays a role in modulating tolerance responses during *M. sexta* attack, likely by regulating auxin signaling, but does not play a major role when plants are colonized by beneficial or pathogenic fungi. Overexpression of Na-miR390, one of the highly conserved plant miRNAs, resulted in reduced plant reproductive output only when plants were attacked by *M. sexta* caterpillars. The overexpression of miR390 did not significantly influence caterpillar performance (caterpillar mass), and also did not significantly influence plant development or its reproductive output when grown under unstressed glasshouse conditions.
We propose the following model for the ecological role of miR390 in modulating host tolerance responses to herbivory (Figure 8): attack of M. sexta larvae elicits an auxin burst (Machado et al., 2013, 2016) parallel to the elicitation of jasmonates. The attack also elicits miR390 expression, which regulates auxin biosynthesis (such as YUCCA gene expressions), its levels and the expression of ARFs (e.g., ARF4), which, in turn, modulates auxin-responsive changes in secondary metabolism and tolerance responses (Figure 8). This miR390-auxin regulated tolerance response is independent of (and plausibly synergistic to) JA and GAL83 (Schtachtje et al., 2006) regulations (Figure 8). Whether the miR390-ARF4-auxin module mediates the oft-discussed tolerance-defense tradeoffs during herbivory in N. attenuata requires additional research.

Tolerance of plants to insect attack is a distinct type of response to herbivory, yet little is known about its mechanistic basis (Koch et al., 2016). Tolerance is characterized as a host plant’s ability to recover from injuries of herbivore attack through compensatory physiological processes (Koch et al., 2016). Resistance to insect attack is often categorized into antibiosis (traits that impact pest biology), antixenosis (non-preference traits impacting pest behavior), and tolerance. Tolerance is distinctive from the other two types of resistance as these traits do not necessarily interfere with the performance (growth, physiology, behavior, or population) of the attacking insects but only reduce the fitness impact of herbivory on the host after it has occurred (Koch et al., 2016). In agro-ecological terms, tolerant plants are able to maintain their fitness to a greater degree when
attacked by insects compared to susceptible plants, in a nonreciprocal manner that involves only plant response and not pest responses (Peterson et al., 2017). Tolerance is often measured in terms of reproductive output or yield (which is a currency of “fitness”) of plants after herbivore attack as compared to that of unattacked plants.

Although the underlying mechanisms are largely uncharacterized, five physiological processes are thought to contribute to herbivory tolerance: alterations in photosynthesis, relative growth rate, branching or tillering, preexisting carbon reserves in roots, and the ability to relocate carbon from roots to shoots after herbivory (Koch et al., 2016; Peterson et al., 2017). At a molecular level, only a few genes, such as GAL83 SNF1-related kinase for resource allocation during the feeding of the chewing insect M. sexta (Schwachtje et al., 2006), peroxidases and catalases in response to chinch bug attack, and those involved in ROS detoxification mechanisms important for Hemipteran pests tolerance, have been associated with host tolerance to herbivory (Koch et al., 2016; Peterson et al., 2017). Furthermore, ROS signaling is thought to overlap with phytohormone (JA and SA) signaling pathways, which may be key hubs through which signals might be propagated for long-term responses, leading to mitigation and resumption of growth after herbivory (Koch et al., 2016).

*N. attenuata* has coevolved with the attack of its specialist herbivore, *M. sexta* (Lepidoptera). This host plant deploys a suite of direct and indirect defenses which are not always effective and a *M. sexta* larvae is often able to completely defoliate its host plant. Therefore, *N. attenuata* benefits from tolerance responses as they complement the elaborate direct and indirect defense traits elicited during *M. sexta* attack (Schwachtje et al., 2006). By down-regulating GAL83 expression in response to *M. sexta* attack, *N. attenuata* is able to shunt recently fixed photoassimilates to roots which are later remobilized to shoots to sustain reflowering and to tolerate herbivory (Schwachtje et al., 2006). Furthermore, *N. attenuata* deploys auxin signaling, in addition to jasmonate signaling, to regulate regrowth and maximize plant fitness during herbivory (Machado et al., 2013). Overexpression of miR390 reduced tolerance, as revealed by significant reductions in seed capsule production, without affecting the growth of *M. sexta* larvae. This reduced tolerance could be attributed to reductions in auxin accumulation during herbivory (for instance due to limited elicitation of *YUCCA* genes), along with reduced accumulations of *ARF4* transcripts. On the other hand, overexpression of miR390 did not affect jasmonate accumulations (or other phytohormones such as SA or ABA) or the expression of GAL83. A minor compensatory growth response as seen in stalk elongation was noticed in OEmiR390 plants after caterpillar feeding, but these increases failed to compensate for the reduced capsule production. The reduction in capsule production of *M. sexta* attacked OEmiR390 plants could be restored to WT levels when plants were exogenously treated with IAA.

**FIGURE 7** Overexpression of miR390 down-regulates accumulation of transcripts of auxin response factors (ARFs) during herbivory. After 9 days of caterpillar feeding, transcript abundances in WT and the two miR390 overexpressing lines (OEmiR390) were evaluated with quantitative real-time PCR assays. *ECI* was used as internal reference. Transcript levels in WT were set to 1 and relative abundances in the two transgenic lines were determined. Four biological replicates were randomly chosen; values represent mean ± SDs. Transcripts of ARF4 were significantly down-regulated in both the miR390-overexpressing lines (one-way ANOVA, *F* = 7.032, *P* < .01). Down regulation of ARF2 was significant in one of the OEmiR390 lines (one-way ANOVA, *F* = 3.296, *P*< .05), whereas there were no significant differences (n.s.) in ARF3 levels among the three genotypes. (a) and (b) indicate statistically significant differences across genotypes.

**FIGURE 8** A model for miR390 function in regulating plant fitness responses during herbivory by modulating auxin but not jasmonate signaling. Dashed line: the role of ARFs in tolerance of plants to herbivory is postulated but not studied in *N. attenuata*.
AM fungi (Koch et al., 2016; Peterson et al., 2017; Tao et al., 2016). In order to tolerate attack from sucking insects, plants deploy ROS-detoxification and photosynthetic rate changes, whereas attack from defoliating chewing insects engages mechanisms related to over-compensation through the production of new tissues, changes in architecture (e.g., increased branching), or resource relocation (Koch et al., 2016). Under field conditions, N. attenuata plants face an unpredictable level of herbivore attack every year (Baldwin, 2001; Pandey & Baldwin, 2007; Pradhan et al., 2020). The composition of the herbivore community is known to vary across years and may be responsible for the substantial amount of adaptive plasticity observed in this species (Navarro-Quezada et al., 2020). During the 2019 field season, OEmiR390 plants were exposed to the natural herbivore community, which at the time of the experiments did not include a significant Manduca component, we saw only a mild fitness consequence of herbivory, likely due to the overall paucity of folivory. We predict that had Manduca been a significant part of the herbivore community, and the plants had suffered significant folivory, we would have observed considerable fitness differences between WT plants with their functional miR390-auxin regulatory module and the OEmiR390 lines in which this module was disabled. In order to abide by US Department of Agriculture Animal and Plant Health Inspection Service (APHIS) regulations for field release of transgenic plants, we had to terminate our study sooner than the glasshouse studies to prevent any spread of transgenic seeds in nature. Therefore, a fully detailed quantification of reproductive output of OEmiR390 plants in the field could not be made, and only an early account could be recorded. Nevertheless, this is one of the first field tests of miRNA function.

Molecularly, miR390, ARFs, and auxin form the components of an auto-regulatory circuit in a cell (Marin et al., 2010; Yoon et al., 2010). miR390 transcription has been shown to be induced at high IAA concentrations, and miR390 expression is regarded as a regulatory component that senses auxin concentrations for modulating ARF4 expression (Yoon et al., 2010). Further, positive and negative feedback regulation of miR390 by ARFs have also been proposed (Marin et al., 2010). In the context of our study, recruitment of such a regulatory circuit is highly plausible because in response to herbivore attack, IAA levels are rapidly induced (Machado et al., 2013). Herbivore attack also induces the expression of miR390 in the attacked leaves. We hypothesize that miR390 regulates the expression of ARF4, which might exert a feedback to optimize auxin sensing and in turn its biosynthesis by YUCCAs (Figure 8). miR390-modulated expression of ARF transcription factors could regulate the downstream tolerance response related to modulation of primary and secondary metabolite accumulations that are produced in response to herbivory and ultimately provide the metabolic basis for tolerance (Figure 8).

miR390 has been a widely conserved miRNA in eudicots (Xia et al., 2017) that often acts as a critical factor in auxin signaling. The miR390-ARF regulatory pathway has undergone a dynamic evolutionary route (Xia et al., 2017). Therefore, it should not be surprising that the biological functions of such a component would also undergo diversification as a consequence of changing ecological and environmental selection pressures. Nearly half of the miRNA gene families are frequently lost and the “repertoires of miRNA genes” have often “dynamically” changed during the evolution (Nozawa et al., 2012). Indeed, several divergent biological functions of miR390 have been identified in addition to its well-studied roles in developmental processes: for example, miR390 has been implicated in gall formation by Meloidogyne (Cabrera et al., 2016), cadmium and aluminum tolerance in rice and flax (Ding et al., 2016; Dmitriev et al., 2017), symbiotic nodulation in Medicago (Hobeker et al., 2017), and salt stress induced developmental plasticity of popular (He et al., 2018). Here, we extend the functional arena of miR390 to a role in herbivore induced tolerance responses in N. attenuata. The next step would be to gain a deeper insight into molecular mechanism of action that warrants future investigation. Further research is needed to determine how miR390 responds to herbivory signals and regulates auxin signaling to ultimately provide the observed tolerance responses.

4 | MATERIALS AND METHODS

4.1 | Plant material, generation of miR390-overexpressing plants, and growth conditions

N. attenuata wild type (WT) plants (22nd inbred generation) were transformed according to Krügel et al. (2002) using the plasmid depicted in Figure S1, carrying the miR390 precursor sequence (Figure S1). N. attenuata transcriptome sequencing on the GS FLX Titanium platform (Gase & Baldwin, 2012) revealed the sequence of the miR390 precursor transcript (GBGF01029632) (Bozorov et al., 2012). Based on this sequence information, we designed primers SRNA1-39 (5’-GGGCGGCTGAGATCCATTCATTTATAGTACATCTTTTAG-3’) and SRNA2-42 (5’-GGGCGGGTGTCACCGTTATCATGATTGAAGAACGATACACATATAC-3’). For cDNA synthesis, total RNA was isolated from N. attenuata leaves harvested 1 h after treatment with wound and oral secretion from M. sexta larvae using the TRIZOL method as recommended by the manufacturer (http://www.invitrogen.com). Reverse transcription was done with SuperScript Reverse Transcriptase following the protocol of the same manufacturer. PCR was performed with the designed primer pair and Phusion DNA Polymerase (www.thermofisher.com) according to the instructions of the manufacturer to amplify the pre-miR390 gene from the synthesized cDNA. The amplified PCR fragment (Figure S1) was then cloned in the 9.7 kb Xhol-BstEII fragment of the binary plant transformation vector pRESC800 (Gase & Baldwin, 2012); JQ354897), yielding pRESC9MIR390. T1 seeds of 20 independently transformed lines were screened on hygromycin according to (Gase et al., 2011). The number of insertions was checked by evaluating the number of reads of different target sites of the transformation vector as well as three internal genes using the nCounter Nanostring procedure, as previously described (He et al., 2019). Lines “385–42” and “392–63” showed a single complete insertion (Figure S1).

For glasshouse experiments, WT seeds (31st inbred generation) were germinated on GB5 medium and OEmiR390 lines (A-12-392-63-71, A-12-385-42-51) were germinated on GB5
supplemented with hygromycin (Krügel et al., 2002; Onokoseung et al., 2012); 10–12 days after germination, plants were transferred to Teku pots either filled with soil or sand for the AMF experiment (Krügel et al., 2002; Pandey et al., 2018); 10–12 days later plants were transferred to soil or AMF inoculum (active AMF inoculum, BiomyC Vital, www.biomycc.de, diluted 1:10 with expanded clay, particle size: 2–4 mm). Plants were grown either as single plants in 1-L pots or under competition with a size- and age-matched WT neighbor (Pandey et al., 2008), as indicated in the text. Plants grown in soil received the optimal amount of nutrients, while AMF inoculated plants were fertilized with hydroponic solution with 1/10 of the regular Pi content during rosette stage, and starting with the elongation stage with 1/4 Pi concentration (Pandey et al., 2018). Plant performance measurements of rosette diameter, chlorophyll content, stalk length, flower number, total capsule number, and shoot and root fresh mass were recorded at respective time points.

4.2 | miR390 accumulation and transcripts measurements by quantitative real-time PCR

Overexpression of miR390 was evaluated by quantitative real-time PCR (qPCR) assays, as was the abundance of YUCCA1, YUCCA3, ARF2–4, and GAL83 transcripts. Total RNA was extracted from frozen seedlings using the lithium chloride method (Kistner & Matamoros, 2005) and adapted to N. attenuata (Pradhan et al., 2017), and DNA traces were removed with DNase I (DNA-free kit; Ambion). For analysis of miRNA abundances, a total of 1 μg of total RNA was converted to complementary miRNA using miScript™ II RT Kit (Qigen, Cat. No. 218,161) according to the manufacturer’s instruction (www.qiagen.com). qPCR was performed with the help of miScript SYBR Green PCR kit (Qigen, Cat. No. 218073), by following the manufacturer’s protocol, adapted to N. attenuata and previously described (Pradhan et al., 2020), where 10 ng of complimentary miRNA was used. For analysis of targets, 1 μg of total RNA was converted to cDNA according to the manufacturer’s instruction using the SuperScript Reverse Transcriptase (Invitrogen). Using 10–50 ng of cDNA, qPCR was performed with the qPCR core kit for Taykon™ SYBR master mix (Eurogentec) on a Mx3005P qPCR system (Stratagene, Santa Clara, CA, USA; http://www.stratagene.com). The 2−ΔΔCT method was used for data analysis, and the 5S rRNA gene was used for normalization of miRNA (Pradhan et al., 2020), whereas the ECI gene was used for normalization of YUCCA, ARF, and GAL83 transcripts (Bubner et al., 2004). Sequences of all primers (genes and miRNAs) used in the study are provided in Table S1.

4.3 | Blumenol analysis to estimate root colonization by AMF

11-Carboxyblumenol-C-glucoside is a reliable quantitative leaf marker for root arbuscule colonization (Wang et al., 2018). Leaf samples were powdered with two steel balls with an automated tissue homogenizer (SPEX SamplePrep) for 60 s at 1150 strokes min−1. About 100 mg plant tissues were extracted with 800 μl of 80% MeOH spiked with internal standard (10 ng ABA per sample). After shaking in the tissue homogenizer, samples were centrifuged for 20 min with 1913 × g at 4°C; the supernatant was collected and analyzed by ultrahigh performance liquid chromatography (UHPLC) coupled to a triple quadrupole mass spectrometer equipped with a heated electrospray ionization (ESI) source (EVOQ, Bruker, Bremen, Germany) as described previously (Mindt et al., 2019; Wang et al., 2018).

4.4 | Determination of susceptibility against Fusarium brachygibbosum

The F. brachygibbosum pathogenesis bioassay was conducted as previously described (Pradhan et al., 2020). The Padwick Utah 4 strain, isolated from diseased N. attenuata plants growing in the native habitats (Schuck et al., 2014), was used for determining whether the susceptibility of OEmiR390 plants to a natural pathogen was altered. A pure fungal culture was regularly maintained on potato dextrose agar (PDA, Fluka Analytical, Steinheim, Germany) at 25°C in the dark (Pradhan et al., 2020). A droplet (10 μl) of spore suspension (106 spore ml−1) of F. brachygibbosum, harvested from 12- to 15-day-old pure cultures grown on PDA, was inoculated on the “+2-leaves” of rosette-stage (Pradhan et al., 2020) N. attenuata plants under high humidity in a climate controlled growth chamber (day/night cycle of 16 h (26°C)/8 h (22°C) with 80 ± 5% humidity. First disease symptoms were visible within 3–4 dpi and lesion size was measured at 5 dpi (Pradhan et al., 2020); 10–15 independent biological replicates per genotype were used.

4.5 | Herbivore performance assay

M. sexta performance assays were conducted on stably transformed lines as described earlier (Pandey & Baldwin, 2007; Pradhan et al., 2017); 20 replicate plants of WT and the two OEmiR390 lines were germinated, transferred to Teku pots and grown on soil in 1-L pots in the glasshouse as described above. Two neonate larvae were placed on the lower surface of the +2 leaf of rosette stage plants; after 2 days, one neonate larva per plant was allowed to feed for 11 to 14 days. Caterpillar biomass was recorded every 3 to 4 d. Rosette size and stalk length were recorded weekly. All capsules per plant were counted 9, 10, and 11 weeks after potting. Closed (unripe) and open (ripe) capsules were counted separately. The caterpillar performance assay and growth and fitness measurements were repeated in an independent biological experiment with a similar setup using 10 independent biological replicates.

4.6 | Auxin complementation assays

For auxin complementation assays, 1 mM IAA (Pandey et al., 2008) was exogenously supplied to herbivore-attacked plants. Fully
expanded WT and OEmiR390 plants were subject to M. sexta neonates as described above. After 3 days of caterpillar feeding, IAA, dissolved in lanolin paste (.1752 mg/ml) and 20 μl/petiole of this IAA-lanolin paste was applied to the petioles of +2 leaves as previously described (Machado et al., 2013). Plants were grown until seed set. Recovery of plant fitness was evaluated by counting seed capsules as described above.

4.7 | Quantification of phytohormones and secondary metabolites

To determine differences in herbivore-induced phytohormone and secondary metabolite accumulation between the WT and OEmiR390 plants, leaves were harvested after continuous caterpillar feeding for 9 days, immediately frozen in liquid nitrogen and stored at –80°C until further processing. Leaf material was ground into a fine powder and 100 mg of the frozen tissue were aliquoted into 96-well biotubes and samples were extracted following a published protocol (Schäfer et al., 2016). After solid phase extraction samples were reconstituted in acidified MeOH (.2 N formic acid [FA] in 80% MeOH) and analyzed on an UHPLC-ESI-Triple-Quadrupole-MS (EVOQ Elite, Bruker) using a Zorbax Eclipse XDB-C18 column (100 x 2 mm, particle size 1.8 μm, Agilent) as previously described (Schäfer et al., 2016). Post-run data analysis was performed in the MS Data Review software (Bruker). Compound concentrations were calculated using the corresponding labeled internal standards as previously described (Schäfer et al., 2016).

A second aliquot (100 mg) of the frozen leaf material was extracted in 800 μl of 80% MeOH containing the internal standard testosterone and used for a qTOF analysis. After incubation for 30 min at –20°C, samples were homogenized in a Genogrinder (60 s at 1150 strokes min⁻¹), centrifuged at 1913×g for 20 min at 4°C and 150 μl of supernatant were transferred to a skirted 96-well plate and subsequently sealed with sticky film. Separation was performed using a UHPLC system (UltiMate3000 RSLC, Thermo Scientific) equipped with a 150 mm × 2.1 mm, 2.2 μm, Acclaim RSLC C18 column (ThermoFisher). The following binary solvent gradient (A: water, .1% [v/v] acetonitrile, .05%FA; and B: acetonitrile, .05% FA) was applied: 0 to .5 min, isocratic 90% A, 10% B; .5 to 23.5 min, linear ramp to 10% A, 90% B; 23.5 to 25 min, isocratic 10% A, 90% B. The flow rate was .4 ml min⁻¹ and 2 μl of leaf extract were injected. Mass detection was carried out on a Bruker ESI-qTOF-MS as previously described (Gaquerel et al., 2010). Post-run analysis was performed in the Compass DataAnalysis software (Bruker).

4.8 | Field experiments

For field-grown plants, seeds of the transformed N. attenuata lines (OEmiR390, EV) were imported and released under US Department of Agriculture Animal and Plant Health Inspection Service (APHIS) notification numbers 18–282-103r with genotypic descriptors NA418 and NA056, respectively. The field experiment was conducted at the Lytle Ranch Preserve, located in the Great Basin Desert of Southwestern Utah, USA (latitude 37.146, longitude 114.020). WT and OEmiR390 (A-12-392-63-71) seeds were germinated directly on borax-soaked Jiffy 703 pots (AlwaysGrows), and transplanted into the field plot 3 to 4 weeks later. The two genotypes were grown in pairs; in total 18 replicate plants per genotype were analyzed. Rosette diameter, stalk length, flower number and the number of immature seed capsules were determined at the indicated time-points. Seed capsules were removed after counting to fulfill safety regulations. Herbivore damage per plant was estimated 3, 4, and 7 weeks after planting.

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CONFLICT OF INTEREST

The authors declare no conflict of interests.

AUTHOR CONTRIBUTIONS

S.P.P. and I.T.B. conceived the research plan and designed experiments, M.P., C.R., R.H. and S.P.P. conducted research and analyzed data, M.P., C.R., R.H. and S.P.P. conducted field work. S.P.P., M.P. and I.T.B. provided most of the resources.

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