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# Development of multi-target dsRNAs targeting *PcNLP* gene family to suppress *Phytophthora capsici* infection in *Nicotiana benthamiana*

Minsu Park<sup>1,2†</sup>, Yujin Kweon<sup>1†</sup>, Jihyun Eom<sup>1</sup>, Minsun Oh<sup>1</sup> and Chanseok Shin<sup>1,2,3,4\*</sup>

### **Abstract**

Phytophthora capsici, which causes diseases in solanaceous crops, secretes necrosis and ethylene-inducing peptide 1-like proteins (NLPs) that induce plant defense responses and leaf necrosis. In this study, we used RNA interference (RNAi) technique, a proven strategy for crop protection and gene regulation in plants, to suppress P. capsici infection through the inhibition of PcNLPs. In the RNAi mechanism, Dicer processes double-stranded RNA (dsRNA) into smaller entities known as small interfering RNAs (siRNAs). These siRNAs subsequently integrate into the RNA-induced silencing complex to form sequence-specific base pairing with complementary regions of the target mRNA. This interaction effectively initiates the degradation process of the target mRNA. We designed and synthesized dsRNAs targeting the "AIMY" and "GHRHDWE" conserved motifs of PcNLP gene family, which are predicted to be key elements for the expression of NLPs and pathogen infection. After infiltration of dsRNAs targeting the motifs and inoculation with P. capsici, we confirmed a significant suppression of P. capsici infection and downregulation of the PcNLP gene family. These findings imply that the dsRNA-mediated RNAi technique holds potential for mitigating a wide range of pathogens, while simultaneously suppressing the expression of a particular gene family using dsRNA targeting functional conserved motifs in the gene family.

Keywords RNA interference, Phytophthora capsici, NLP, Multi-target double-stranded RNA

Chanseok Shin

cshin@snu.ac.kr

### Introduction

RNA interference (RNAi) has emerged as a promising tool for crop protection and gene regulation in plants. Following the exogenous introduction of double-stranded RNA (dsRNA) molecules, the RNAi pathway triggers a gene silencing mechanism, offering a potential means to control plant genes and pathogens [1]. In the cellular system of plants, dsRNA is cleaved by Dicer-like endonucleases into small interfering RNAs (siRNAs). These siRNAs are then incorporated into the Argonaute protein, forming the RNA-induced silencing complex, which can specifically target and silence complementary sequences in the target mRNA [1–3].

The oomycete plant pathogen *Phytophthora capsici* (*P. capsici*) targets solanaceous crops, including pepper,



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<sup>&</sup>lt;sup>†</sup>Minsu Park and Yujin Kweon have contributed equally to this work.

<sup>\*</sup>Correspondence:

<sup>&</sup>lt;sup>1</sup> Department of Agricultural Biotechnology, Seoul National University, Seoul 08826, Republic of Korea

<sup>&</sup>lt;sup>2</sup> Research Institute of Agriculture and Life Sciences, Seoul National University, Seoul 08826, Republic of Korea

<sup>&</sup>lt;sup>3</sup> Plant Genomics and Breeding Institute, Seoul National University, Seoul 08826, Republic of Korea

<sup>&</sup>lt;sup>4</sup> Research Center for Plant Plasticity, Seoul National University, Seoul 08826, Republic of Korea

which causes root rot and results in substantial yield losses [4, 5]. The secretion of necrosis and ethyleneinducing peptide 1-like proteins (NLPs) by oomycetes induces both the plant defense response and leaf necrosis [6, 7]. In *P. capsici*, significant upregulation of *PcNLP2* and PcNLP6 was observed during the infection stage. Furthermore, when PcNLP2 and PcNLP6 were introduced in agroinfection assays, Capsicum annuum and Nicotiana benthamiana leaves showed the most extensive necrotic areas, indicating the crucial role of these genes in promoting virulence throughout the infection phases [8]. Most of the NLPs contain the "AIMY" motif and highly sequence-conserved "GHRHDWE" motif. Mutations in the "AIMY" motif in NLP1 reduce the production of reactive oxygen species and suppress Colletotrichum orbiculare infection in cucumber [9]. Additionally, the "GHRHDWE" motif is a requirement for the activity of NLPs. In addition, this motif is critically involved in cavity formation on the protein surface, which is important for the necrosis-inducing activity of Pectobacterium carotovorum [10]. These results suggest that the "AIMY" and "GHRHDWE" motifs in PcNLP2 and PcNLP6 may play a particularly important role in P. capsici infection.

In our previous studies, we suppressed pathogens, including the pepper mottle virus and P. capsici, using dsRNAs varying position of their targets [11–13]. We confirmed that P. capsici infection was suppressed by dsRNAs targeting PcNLP2 and PcNLP6 of P. capsici. Significant differences in the suppression of P. capsici infection were observed depending on the regions of PcNLP2 and PcNLP6 that were targeted by the dsRNA [13]. Interestingly, the dsRNAs that effectively suppressed P. capsici infection targeted regions containing the "AIMY" and "GHRHDWE" motifs. Therefore, to investigate the role of the "AIMY" and "GHRHDWE" motifs in P. capsici infection, we used dsRNAs specifically targeting regions containing these motifs in PcNLP2 and PcNLP6, as well as dsRNAs targeting regions not containing the motifs. Furthermore, we examined whether dsRNAs targeting regions containing the motifs also suppressed the expression of other *PcNLPs*. Through this approach, we aimed to ascertain whether our designed multi-target dsRNA, which targets the functional conserved motifs of a particular gene family, can effectively regulate the expression of that gene family.

### Materials and methods

### Plant growth conditions and P. capsici maintenance

The wild-type *Nicotiana benthamiana* was grown at 25 °C and 50% humidity, exposed to a daily photoperiod of 16 h of light followed by 8 h of darkness, all within a growth chamber. For the experiments, *N. benthamiana* 

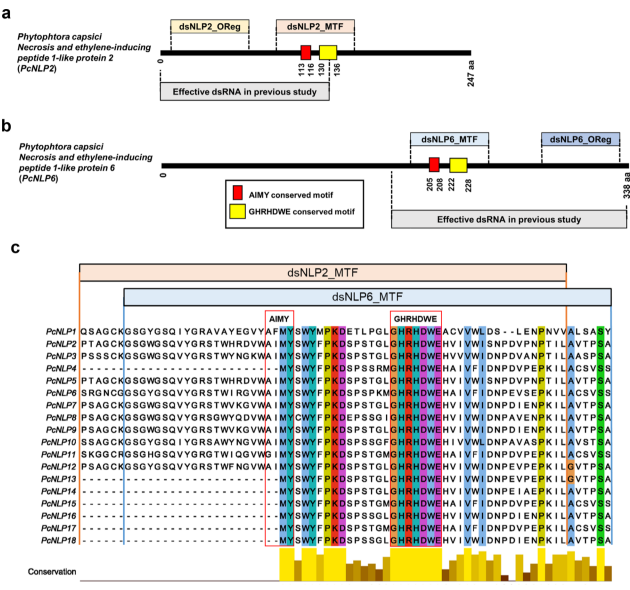
leaves were selected when plants were 3 weeks old. The KACC 40476 strain of *P. capsici*, generously supplied by Dr. Doil Choi's Laboratory (Seoul National University, Seoul, Republic of Korea), was grown on V8 juice agar medium for 8 days, maintained in continuous darkness at 23 °C.

### Design and synthesis of dsRNAs

Two dsRNAs, each 200 bp in length, were designed to target the regions containing the "AIMY" and "GHRH-DWE" motifs of PcNLP2 and PcNLP6, and regions not containing the motifs, respectively (Fig. 1a, b). The conservation of amino acids and RNA sequences was measured using Jalview software [14]. A 200 bp dsRNA, designed to target the Renilla luciferase gene, was employed as a mock in the experiments. As reported in our previous study [13], for the synthesis of dsRNAs, we first prepared the corresponding DNA templates, incorporating T7 promoter sequences (5'-TAATAC GACTCACATATAAGAGAG-3'). This was achieved through a polymerase chain reaction (PCR) utilizing Phusion High-Fidelity DNA polymerase (Thermo Scientific, United States), in accordance with the manufacturer's protocol. The resulting PCR products were subsequently employed in the dsRNA synthesis process utilizing MEGAscript RNAi Kit (Invitrogen, United States), in accordance with the manufacturer's protocol. The purity of dsRNAs was shown in Additional file 1: Fig. S1. The primers used for PCR are listed in Additional file 1: Table S1.

## Administration of dsRNAs and *P. capsici* inoculation in *N. benthamiana*

As reported in our previous study [13], 2 days prior to P. capsici inoculation, we introduced 500 µL of dsRNAs (200 nM) into the abaxial side of N. benthamiana leaves using a needle-free syringe. A day before the P. capsici inoculation, the mycelium of P. capsici, cultured on V8 juice agar medium, was scrapped and left to incubate overnight under continuous exposure to light at 23 °C to facilitate sporangia formation. On the day of inoculation, the culture plate was flooded with 10 mL of distilled water, followed by a 1 h incubation at 4 °C to collect the zoospores. When the concentration of the zoospores reached at the  $5 \times 10^4$  zoospores mL<sup>-1</sup>, 12  $\mu$ L of collected zoospore suspension was inoculated in the abaxial side of the N. benthamiana leaves. Following inoculation, the leaves were incubated in darkness at 23 °C for approximately 24 h. The confirmation of P. capsici infection was conducted through phenotypic observation prior to sampling of leaves.



**Fig. 1** Design of double-stranded RNAs (dsRNAs) targeting the "AIMY" and "GHRHDWE" motifs of *PcNLP2* and *PcNLP6*. **a** dsNLP2\_MTF and dsNLP2\_OReg were designed to target the regions containing the "AIMY" and "GHRHDWE" motifs of *PcNLP2*, and the outer region not containing the motifs, respectively. The length of each dsRNA was 200 bp. **b** dsNLP6\_MTF and dsNLP6\_OReg were designed to target the region containing the "AIMY" and "GHRHDWE" motifs of *PcNLP6*, and the outer region not containing the motifs, respectively. The length of each dsRNA was 200 bp. **c** Conservations of amino acids in *PcNLPs* in the regions targeted by dsNLP2\_MTF and dsNLP6\_MTF was measured using Jalview software. MTF: Motif, OReg: Outer region. Red boxes: "AIMY" and "GHRHDWE" motifs region

# Assessment of chlorophyll fluorescence expression in *N. benthamiana* leaves

The infection of *P. capsici* on *N. benthamiana* leaves was confirmed by chlorophyll fluorescence expression using FOBI fluorescence in vivo imaging system (Neoscience, Republic of Korea), as previously described [13]. ImageJ was used to quantify the size of lesion [15].

# Total RNA extraction and complementary DNA (cDNA) synthesis

Following assessment of chlorophyll fluorescence expression, *N. benthamiana* leaves were collected and ground in liquid nitrogen. Total RNA extraction was performed using RiboEx (GeneAll, republic of Korea), and the extracted total RNA was treated with recombinant

DNase I (Takara, Japan) to remove both single-stranded and double-stranded DNAs, in accordance with the manufacturer's instructions.

To synthesize cDNA, we utilized 1  $\mu$ g of each RNA sample. This process was executed using PrimeScript Reverse Transcriptase (Takara, Japan) with oligo (dT) primers (Thermo Scientific, United States), in accordance with the manufacturer's instructions.

### Quantitative real-time PCR (qRT-PCR)

Light Cycler 480 (Roche, United States) with AccuPower 2X GreenStar qPCR Master Mix (Bioneer, Republic of Korea) was used for performing qRT-PCR, along with cDNA and gene-specific primers. The Ct values for the target genes were normalized to those of the house-keeping gene NbEF1a, as a control. To calculate relative expression levels, the  $\Delta\Delta$ Ct method was employed. Statistical analysis was performed using Student's t-test (\*P<0.05; \*\*P<0.01; \*\*\*P<0.001). Due to high homology among PcNLPs, we designed qRT-PCR primers considering potential off-target effects and the motif regions affected by dsRNAs. The primer sequences are listed in Additional file 1: Table S2. The qRT-PCR target regions within PcNLPs are depicted in Additional file 1: Fig. S2.

### **Results**

## Design of dsRNAs targeting "AIMY" and "GHRHDWE" motifs of PcNLP2 and PcNLP6

In our previous study [13], plants treated with dsRNAs targeting 5' and 3' regions of PcNLP2 and PcNLP6 sequence yielded different effects on the target gene and P. *capsici* infection. The dsRNAs that targeted the 5'-region of PcNLP2 and the 3'-region of PcNLP6 resulted in effective suppression of target gene expression and P. capsici infection. Interestingly, the target regions of the effective dsRNAs contained the "AIMY" and "GHRHDWE" motifs (Fig. 1a, b), which are predicted to be key elements in *P*. capsici infection and the expression of PcNLPs, based on previous studies [9, 10]. Therefore, we designed two dsRNAs for each gene: dsNLP2\_MTF and dsNLP6\_MTF (MTF; Motif) targeting both the "AIMY" and "GHRH-DWE" motifs of PcNLP2 and PcNLP6, respectively, and dsNLP2\_OReg and dsNLP6\_OReg (OReg; Outer region) targeting regions that do not contain motifs, which were the outer regions of effective dsRNA in previous study [13], respectively (Fig. 1a, b). dsNLP2\_MTF and dsNLP6\_MTF are designed to target the "AIMY" motif on nine PcNLPs and the "GHRHDWE" motif on all PcN-*LPs* (Fig. 1c). In addition, the RNA sequences of "AIMY" and "GHRHDWE" motifs, which are targets of dsRNA, are identical in most *PcNLPs* (Additional file 1: Fig. S3). In total four dsRNAs were synthesized to evaluate ability to suppress *P. capsici* infection and alter the expression of *PcNLPs*.

# Suppression of *P. capsici* infection via dsRNAs targeting "AIMY" and "GHRHDWE" motifs of *PcNLP2* and *PcNLP6*

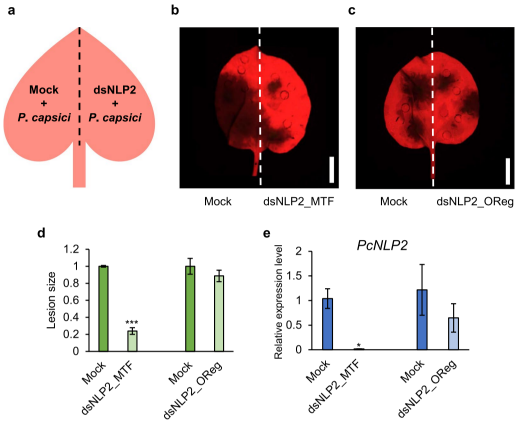
Consistent with the previous study [13], we introduced dsRNAs targeting the motif regions and the outer region of PcNLP2, PcNLP6, or mock into the N. benthamiana leaves to assess the effect of dsRNAs on P. capsici infection (Figs. 2a, 3a). We measured the size of the infected lesions and the expression of PcNLP2 and PcNLP6 in P. capsica-inoculated leaves at 24 h post-inoculation (hpi) using FOBI and qRT-PCR, respectively. At 24 hpi, infected lesions were significantly suppressed in dsNLP2\_MTF-treated leaves compared with mocktreated leaves (Fig. 2b, d), but there was less suppression in dsNLP2\_OReg-treated leaves than the dsNLP2\_MTFtreated leaves (Fig. 2c, d). Similarly, there was significant suppression of lesions in dsNLP6\_MTF-treated leaves (Fig. 3b, d), whereas this did not occur in dsRNA6\_ORegtreated leaves (Fig. 3c, d).

Additionally, we analyzed the levels of *PcNLP2* and *PcNLP6* transcript in leaves treated with dsRNAs, respectively. Compared with mock-treated leaves, *PcNLP2* expression was 57-fold lower in dsNLP2\_MTF-treated leaves, and *PcNLP6* expression was 6,133-fold lower in dsNLP6\_MTF-treated leaves (Figs. 2e, 3e). These results indicate that dsRNAs targeting the motifs suppress *P. capsici* infection and suppress the expression of *PcNLP2* and *PcNLP6*. However, dsRNAs targeting the outer region did not significantly affect the expression of *PcNLP2* and *PcNLP6* (*P*>0.05), causing only a slight reduction (Figs. 2e, 3e). In addition, they had no effect on *P. capsici* infection (Figs. 2d, 3d).

# Expression of *PcNLPs* in *P. capsici* treated with dsRNAs targeting "AIMY" and "GHRHDWE" motifs of *PcNLP2* and *PcNLP6*

We performed qRT-PCR on *PcNLPs* of *P. capsici* to investigate the multi-target effects of dsNLP2\_MTF and dsNLP6\_MTF. Interestingly, compared with mocktreated leaves, the expression of *PcNLPs* decreased from 2-fold to 362-fold in dsNLP2\_MTF-treated leaves (Fig. 4a), and from 5-fold to 6,133-fold in dsNLP6\_MTF-treated leaves (Fig. 4b). However, there was no significant decrease in the expression of *PcNLPs* in dsNLP2\_OReg and dsNLP6\_OReg-treated leaves (Additional file 1: Fig. S4). A Ct value for *PcNLP8* was not detected in mocktreated and dsNLP6\_MTF-treated leaves (Fig. 4b). This was attributed to the significantly lower expression level of *PcNLP8* at the *P. capsici* infection stage compared with other *PcNLPs*, resulting in the non-detection of *PcNLP8* using qRT-PCR. (Additional file 1: Fig. S5). Overall, these

Park et al. Applied Biological Chemistry



**Fig. 2** Regulation of *Phytophthora capsici* infection and *PcNLP2* expression via dsNLP2\_MTF and dsNLP2\_OReg. **a** The experimental scheme for dsRNA treatment and *P. capsici* infection. **b** The phenotype of *P. capsici* infection lesion introduced with mock and dsNLP2\_MTF, respectively, in 3-week-old *Nicotiana benthamiana* leaves, determined using a FOBI in vivo fluorescence imaging system. Scale bar = 1 cm. **c** The phenotype of *P. capsici* infection lesion introduced with mock and dsNLP2\_OReg, respectively. Scale bar = 1 cm. **d** Quantification of *P. capsici* infection lesion size using ImageJ. **e** Relative expression level of *PcNLP2* using quantitative real-time PCR (qRT-PCR). Mock: Treatment with dsRNA targeting *Renilla luciferase*; dsNLP2\_MTF: Treatment with dsNLP2\_OReg: Treatment with dsNLP2\_OReg. Outer region. Data represent the mean ± standard error of mean (SEM; *N* = 3). Statistical significance is determined by Student's *t*-test (\**P* < 0.05 and \*\*\*\**P* < 0.001)

results suggest that multi-target dsRNAs targeting the "AIMY" and "GHRHDWE" motifs of *PcNLPs* not only suppress *P. capsici* infection, but also reduce the expression of *PcNLPs*.

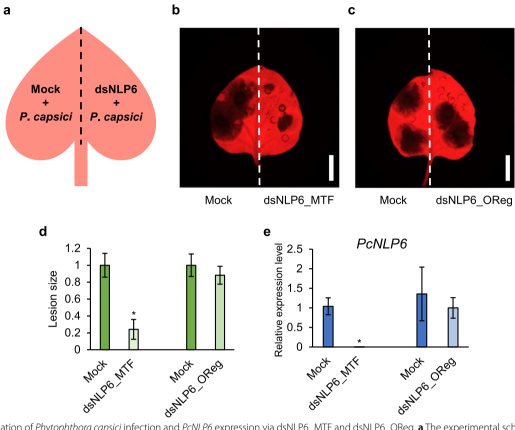
### Discussion

Traditional fungicides have conventionally controlled *P. capsici* [5, 16], but their prolonged use and accumulation may induce fungicide-resistant pathogens and unknown mutations in the pathogen. dsRNA-mediated RNAi technology is now extensively employed to target plant genes, insects, viruses, and fungi [1]. This approach allows the fastest response to the mutation of plant pathogens as dsRNAs can be targeted to the specific mutated gene.

In a previous study [13], we showed the effective suppression of *P. capsici* by dsRNAs targeting specific regions of the *P. capsici* effector genes *PcNLP2* and *PcNLP6*. In this study, we confirmed that the "AIMY"

and "GHRHDWE" motifs were included in the target regions of the effective dsRNAs used in the previous study. The dsRNAs were designed and synthesized by dividing the target regions of the effective dsRNAs used in the previous study into regions containing motifs (dsNLP2\_MTF, dsNLP6\_MTF) and regions not containing motifs (dsNLP2\_OReg, dsNLP6\_OReg) (Fig. 1a, b). dsNLP2 MTF and dsNLP6 MTF, which targeted the motif regions, effectively suppressed P. capsici infection (Figs. 2, 3) and the expression of other *PcNLPs* (Fig. 4), including PcNLP2 and PcNLP6. However, dsNLP2\_ OReg and dsNLP6\_OReg did not significantly affect the expression of PcNLP2, PcNLP6, and P. capsici infection, respectively (Figs. 2, 3). The region targeted by dsNLP2\_ OReg does not contain conserved motifs and the RNA sequences of PcNLPs compared with the region targeted by dsNLP2\_MTF (Additional file 1: Figs. S6, S7). However, the region targeted by dsNLP6\_OReg contains

Park et al. Applied Biological Chemistry

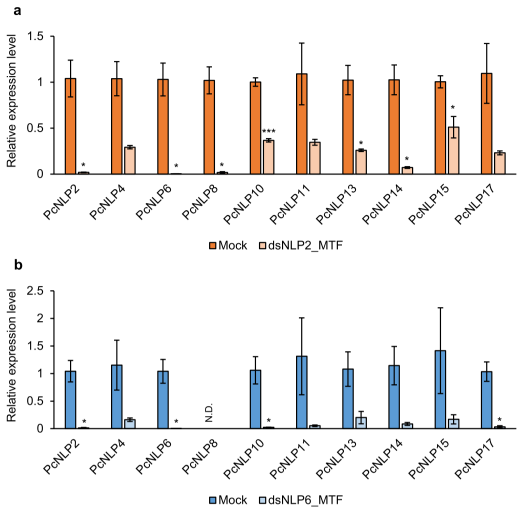


**Fig. 3** Regulation of *Phytophthora capsici* infection and *PcNLP6* expression via dsNLP6\_MTF and dsNLP6\_OReg. **a** The experimental scheme for dsRNA treatment and *P. capsici* infection. **b** The phenotype of *P. capsici* infection lesion introduced with mock and dsNLP6\_MTF, respectively, in 3-week-old *Nicotiana benthamiana* leaves, determined using a FOBI in vivo fluorescence imaging system. Scale bar=1 cm. **c** The phenotype of *P. capsici* infection lesion introduced with mock and dsNLP6\_OReg, respectively. Scale bar=1 cm. **d** Quantification of *P. capsici* infection lesion size using ImageJ. **e** Relative expression level of *PcNLP6* using quantitative real-time PCR (qRT-PCR). Mock: Treatment with dsRNA targeting *Renilla luciferase*; dsNLP6\_MTF: Treatment with dsNLP6\_OReg: Treatment with dsNLP6\_OReg. MTF: Motif, OReg: Outer region. Data represent the mean ± standard error of mean (SEM; *N*=3). Statistical significance is determined by Student's *t*-test (\**P* < 0.05)

conserved amino acid residues and RNA sequences except for the "AIMY" and "GHRHDWE" motifs (Additional file 1: Figs. S8, S9). The regulation of the target region except for the motifs by dsNLP6\_OReg suggests that it cannot control *P. capsici* infection (Fig. 3c, d) or the expression of other *PcNLPs* (Additional file 1: Fig. S4b), even if some conserved amino acid residues or RNA sequences are regulated by dsRNA. These results showed that using only dsNLP2\_MTF and dsNLP6\_MTF to target the "AIMY" and "GHRHDWE" motifs can suppress *P. capsici* infection and the expression of *PcNLPs* (Figs. 2, 3, 4). In addition, when dsNLP2\_OReg and dsNLP6\_OReg were used, the suppression of *PcNLP2* and *PcNLP6* expression was weaker than when dsNLP2\_MTF and

dsNLP6\_MTF were used (Figs. 2e, 3e), suggesting that "AIMY" and "GHRHDWE" motifs are key elements for *P. capsici* infection and *PcNLPs* expression, similar to previous studies [9, 10]. In future research, it will be necessary to separate the dsRNA targeting both the "AIMY" and "GHRHDWE" motifs into dsRNAs targeting only one of these motifs to compare the degree of suppression of *P. capsici* infection and *PcNLPs* expression. This approach will help to uncover the roles of the "AIMY" motif and "GHRHDWE" motif, respectively, in regulating *P. capsici* infection and *PcNLPs* expression.

In conclusion, this study demonstrated that multitarget dsRNAs targeting conserved motifs in the *PcNLP* gene family could suppress *P. capsici* infection and the Park et al. Applied Biological Chemistry



**Fig. 4** Changes in the expression of the *PcNLP* gene family after treatment with dsRNAs targeting dsNLP2\_MTF and dsNLP6\_MTF. Quantification of the expression of the *PcNLP* gene family after treatment with mock, **a** dsNLP2\_MTF, and **b** dsNLP6\_MTF using qRT-PCR. Mock: Treatment with dsRNA targeting *Renilla luciferase*; dsNLP2\_MTF: Treatment with dsNLP2\_MTF; dsNLP6\_MTF: Treatment with dsNLP6\_MTF. MTF: Motif. N.D.: Not detected. Data represent the mean  $\pm$  SEM (N=3). Statistical significance is determined by Student's t-test (\*P<0.05 and \*\*\*P<0.001)

expression of *PcNLPs*. In addition, we provided further evidence that the "AIMY" and "GHRHDWE" motifs could be key elements regulating *P. capsici* infection and the expression of *PcNLPs*. These results can contribute to the development of RNAi technology that can control the entire gene family using one specific multi-target dsRNA.

### Abbreviations

NLP Necrosis and ethylene-inducing peptide 1-like protein

RNAi RNA interference
dsRNA Double-stranded RNA
siRNA Small interfering RNA
hpi Hours post-inoculation
cDNA Complementary DNA

qRT-PCR Quantitative real-time polymerase chain reaction

### Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13765-023-00828-9.

Additional file 1: Table S1. Primer sequences used in dsRNA synthesis. Table S2. Primer sequences used in qRT-PCR. Figure S1. The purity of Mock, dsNLP2\_MTF, dsNLP2\_OReg, dsNLP6\_MTF, and dsNLP6\_OReg. M: 100 bp marker, 1: After transcription, 2: After nuclease treatment, 3: After purification. MTF: Motif, OReg: Outer region. Figure S2. Schematic diagram of target regions in PcNLPs using qRT-PCR. Figure S3. RNA sequences conservation of the region containing the "AlMY" and "GHRHDWE" motifs of PcNLPs targeted by dsNLP2\_MTF and dsNLP6\_MTF. MTF: Motif. Figure S4. Quantification of the expression of PcNLP family genes after treatment with mock, (a) dsNLP2\_OReg, and (b) dsNLP6\_OReg using qRT-PCR. Mock: Treated with dsRNA targeting Renilla luciferase; dsNLP2\_OReg: Treated with dsNLP2\_OReg; dsNLP6\_OReg: Treated with dsNLP6\_OReg. OReg: Outer region. Data represent mean ± SEM (N = 3). Figure S5. Dot plot representation of the delta Ct values of PcNLPs in P. capsica-infected wild-type leaves using qRT-PCR. The dot represents

outliers of replicated samples, and whiskers represent standard deviation. of mean. **Figure S6.** Amino acid sequences conservation of the region targeted by dsNLP2\_OReg. OReg: Outer region. **Figure S7.** RNA sequences conservation of the region targeted by dsNLP2\_OReg. OReg: Outer region. **Figure S8.** Amino acid sequences conservation of the region targeted by dsNLP6\_OReg. OReg: Outer region. **Figure S9.** RNA sequences conservation of the region targeted by dsNLP6\_OReg. OReg: Outer region.

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#### **Author contributions**

CS conceived the project. MP, YK, JE, MO performed experiments. MP, YK, and CS wrote the manuscript. All authors read and approved the final manuscript.

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### Availability of data and materials

Not applicable.

### **Declarations**

### Competing interests

The authors declare that they have no competing interests.

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