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Construction of gold-siRNA_{NPR1} nanoparticles for effective and quick silencing of NPR1 in Arabidopsis thaliana†

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In recent years, gold nanoparticles (AuNPs) have been widely used as gene silencing agents and therapeutics for treatment of cancers due to their high transfection efficiency and lack of cytotoxicity, but their roles in gene silencing in plants have not yet been reported. Here, we report synthesis of AuNPs-branched polyethylenimine and its integration with the small interfering RNAs (siRNA) of *NPR1* to form a AuNPs-siRNA_{NPR1} compound. Our results showed that AuNPs-siRNA_{NPR1} was capable of infiltrating into *Arabidopsis* cells. AuNPs-siRNA_{NPR1} silenced 80% of the *NPR1* gene in *Arabidopsis*. Bacteriostatic and ion leakage experiments suggest that the *NPR1* gene in *Arabidopsis* leaves was silenced by AuNPs-siRNA_{NPR1}. In Columbia-0 plants, compared with the control group treated with buffer solution, the AuNPs-siRNA_{NPR1} treatment significantly increased the number of colonies and cell death, and the leaves turned yellow, similar to the phenotype of the *npr1* leaves. These results indicated this AuNPs-siRNA_{NPR1} silencing the *NPR1* gene method is simple, effective and quick (3 days), and a powerful tool to study gene functions in plants.

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Introduction

Research on plant gene function heavily relies on agrobacterium-mediated gene gun bombardment and PEGtransformed protoplast regeneration. However, these experiments are time-consuming and sometime not effective. Thus, development of a rapid and effective method for silencing genes will be essential to accelerate research on plant gene regulation. Inorganic nanoparticles (INPs) as gene vectors, have recently attracted great attention for their non-immunogenicity, immunity from pathogens, and low toxicity, although their transfection efficiency is lower than that of viral vectors. INPs have advantages of large loading capacity, high stability, and easy preparation and storage.2-6 Gold nanoparticles (AuNPs) are especially useful for delivery of small interfering RNA (siRNA) and protect RNAs from nuclease degradation.7 The multilayer AuNPs integrated with siRNA effectively silenced the genes, as shown by the assay of luciferase activity.8 In addition, the PEGylated AuNPs containing SH-siRNA showed significant RNA RNAi induced by siRNA does not completely silence the targeted gene⁹ and is thus specially suitable for functional studies of lethal phenotype genes such as *ATG6* in *Arabidopsis*. In this study, we reported the construction of AuNPs-siRNA_{NPR1}, and demonstrated for the first time its effective silencing the *NPR1* gene (nonexpressor of pathogenesis-related gene 1) in *Arabidopsis*.

Materials and methods

Preparation of AuNPs

AuNPs were prepared by chemical reduction. ^{10–12} The solution of 25 mL 0.056% chloroauric acid (Aladdin, Shanghai, China) in a 50 mL conical bottle was added by 0.8 mL 1% branched polyethylenimine (Shyuanye, Shanghai, China) 25 kD (AuNPs-PEI) and stirred at 500 rpm continuously for 24 hours at room temperature (25 °C). When the color of the solution no longer change, the reaction is completed and the solution is filtered using a 0.2 μm microporous filter (Biofil, Guangzhou, China). The AuNPs modified by PEI were stored in the original volume of distilled water and in a refrigerator at 4 °C. We modified the preparation procedure of AuNPs-PEI, used a 30 kDa dialysis bag (MYM, Beijing, China) for dialysis to remove PEI which was not combined with AuNPs to allow minimization of PEI interference. ¹³ The size and zeta potential of the nanomedicines were measured using ZEN3690 zetasizer (Malvern, USA).

inhibition (RNAi) in HuH-7 cells.² However, there is no report on usage of AuNPs in plant gene silencing.

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The actual amount of chloroauric acid in preparation: 0.056% g mL⁻¹ \times 25 mL = 0.014 g; the actual mass of AuNPs: $m_{\rm AuNPs}/m_{\rm chlorogold~acid} = 197/394$, $m_{\rm AuNPs} = 0.007$ g; so we estimated the concentration of nano gold as follows: 0.007 g/25.8 mL = 2.71×10^{-4} g mL⁻¹ ≈ 1 mM.

Preparation of AuNPs-siRNA_{NPR1}

In the pre experiment, we found that the *NPR1* gene could be silenced by mixing siRNA (20 μ M) and AuNPs (1 mM) in a ratio of 1:9 and diluting 15 times with a infiltration buffer (10 mM MgCl₂ and 10 mM MES (2-(*N*-morpholino) ethanesulfonic acid) pH 5.7) for infiltration into the leaves of *Arabidopsis thaliana*.

Infiltrating leaves

In this study, the leaves collected for the experiment were all infiltrated by different solutions (the leaves without infiltration were not used in the experiment). Leaves of 3 weeks-old Arabidopsis were selected for infiltration of AuNPs. AuNPs infiltration was performed by pressure infiltration with a 1 mL syringe through the abaxial leaf surface. In this process, a small amount (about 10 μL) of liquid will gently and slowly be penetrated into the leaves. Infiltration is best performed when the plant stomata are opened in the morning. We chose to infiltrate at 8-10 in the morning. The cultivation conditions of plant materials were 16 h light (6 a.m. to 10 p.m.), 8 h dark (10 p.m. to 6 a.m.), the light intensity is 120 μ mol L⁻¹ m⁻² s, the ambient temperature was controlled at 23 $^{\circ}\text{C}\text{,}$ and the relative humidity of the environment is 82%. Infiltration is a time-consuming process. Through many experiments, we found that it is easy to infiltrate at 8 to 10 a.m. (plants get light from 6 a.m.). The infiltration of tobacco (Nicotiana benthamiana) leaves is the same as that of Arabidopsis, but because the Nicotiana benthamiana leaves are more tender, the infiltration needs to be as gentle as possible to avoid mechanical damage. The standard for successful infiltration is to see the wetting of entire leaves.

Extraction of Arabidopsis protoplasts

The healthy leaves of Arabidopsis thaliana (Col-0) before flowering were selected. SiRNA (20 µM) and AuNPs (1 mM) were mixed in a ratio of 1:9 and diluted 15 times with a infiltration buffer. The protoplasts were extracted after 1-2 hours of infiltration. The enzymatic hydrolysate was prepared with cellulose R-10 (1.5%) (Yakult Honsha, Tokyo, Japan), macrozyme R-10 (0.75%) (Yakult Honsha, Tokyo, Japan), CaCl₂ (10 mM), BSA (0.1%), mannitol (0.5 M) and MES (10 mM, pH = 5.7). The epidermis was peeled off with transparent tape, and several parts of the leaves were cut off and placed in 5 mL enzymatic hydrolysate. After 3 hours of the enzymatic reaction at 26 °C in darkness, under 40-50 rpm oscillation, the solution turned from yellow to green (strictly controlling the temperature of enzymatic hydrolysis). The enzymatic hydrolysate passes through 200 mesh sieve holes, the filtrate is placed in centrifugal tube, and the residue in beaker is washed with W5 that is prepared with NaCl (154 mM), CaCl₂ (125 mM), KCl (50 mM), MES (2 mM) and glucose (5 mM). The mixed was centrifuged at 4 °C, at 60 \times g for 5 minutes, and remove supernatant. The

protoplast was washed with 5 mL W5 and centrifuged at 100 \times g, for 3 min. After three times of W5 wash, add 5 mL W5 and gently pipet it for use. 14,15

Confocal microscopy imaging

The images were obtained in Zeiss LSM 880 confocal laser-scanning microscope (Carl Zeiss Microscopy GmbH, Jena, Germany) and analyzed with ZEN software (Carl Zeiss)¹⁶ to observe infiltration of AuNPs-siRNA $_{NPRI}$ into plant cells. FAM-siRNA $_{NPRI}$ (20 μ M) and AuNPs (1 mM) were mixed in a ratio of 1:9 and diluted 15 times with a infiltration buffer. After infiltrating this compound into *Nicotiana benthamiana* leaves or *Arabidopsis* leaves for 1 hour, images or movies were collected with excitation at 492 and 518 nm wavelengths (green is the fluorescence of FAM; red is the spontaneous fluorescence of chlorophyll).

Western blot (WB)

Leaves were collected at different time points, after the AuNPssiRNA_{NPR1} are infiltrated into the Col-0 and NPR1-GFP. We improved WB method on the basis of the protocols of Yan17 and Chen:18 leaves from 4 weeks-old plants were treated with AuNPssiRNA_{NPR1}. 0.4 g samples were collected, frozen in liquid nitrogen, and homogenized in protein extraction buffer (5 mM Tris-HCl, 15 mM NaCl, 0.5 mM EDTA, 1% (v/v) β-ME, 0.2% (v/v) Triton X-100, 0.5% (v/v) Nonidet P-40, 1 mM phenylmethylsulfonyl fluoride, and 40 μM MG-115). Homogenates were centrifuged at 13 000 g for 17 min at 4 °C. Proteins were then denatured at 75 °C with 5× SDS-PAGE loading buffer supplemented with DL-dithiothreitol (100 mM DTT) for 10 min. Total proteins were separated by SDSpolyacrylamide gel electrophoresis (PAGE) with a constant current of 100 mA and then transferred onto polyvinylidene difluoride membranes at a constant voltage 100 V (Bio-Rad, CA, USA). Skimmed milk (5% (w/v)) was prepared by dissolving milk powder in Tris-buffered Saline Tween-20 buffer and used to block the resulting membrane (Merck, Darmstadt, Germany) and to dilute antibodies for immunoblotting. Antibodies of green fluorescent protein (GFP) (JL-8 Monoclonal Antibody, Fisher Scientific, 632381) and β-actin were purchased from the Beyotime Institute of Biotechnology (Shanghai, China); antibodies of NPR1 were purchased from the Agrisera (Swedish). The β-actin antibody was used as control for analysis.

qPCR analysis

After AuNPs-siRNA_{NPR1} or AuNPs-siRNA_{n.c.} (negative control) were infiltrated into Col-0 and *NPR1-GFP*, leave samples were collected at different time points for evaluation of the silencing effect on *NPR1* gene. Leaves (0.05 g) were collected and frozen in liquid nitrogen and stored at −80 °C. RNA was extracted by plant total RNA isolation kit (Sangon biotech, Shanghai, China) and then treated with primeScript RT Master Mix (Takara, Tokyo, Japan) according to the manufacturer's instructions, to synthesize the cDNA.^{19,20} Refer to Yan's procedure for real time PCR (qPCR).¹⁷ The primers used for *NPR1* are listed as follows: F: GATCGCAAAA-CAAGCCACTATGG and R: ATCGAGCAGCGTCATCTTCAATT.

Growth and inoculation of strains

This study used the *P. syringae* strain (Pst DC3000 (*AvrRps4*)) provided by Dr Yang of South China Normal University. Pst DC3000 (*AvrRps4*) was cultured in King's B medium containing 50 μ g mL⁻¹ rifampicin and 100 μ g mL⁻¹ kanamycin for 18 hours at 28 °C, centrifuged (4000 rpm min⁻¹, twice), washed with 10 mM MgCl₂ twice, and then resuspended in 10 mM MgCl₂ and diluted to the required concentration (OD₆₀₀ = 1.0).¹⁵

Infection experiment of pathogenic bacteria

After 3 days of AuNPs-siRNA $_{NPRI}$ infiltration, bacterial solution of Pst DC3000 (AvrRps4) (OD $_{600}=0.001$ diluted from the stock solution of OD = 1.0) was infiltrated and sampled after 3 days. Two small round leaves were beaten with sterilized puncher (different plants need to be wiped with sterilized water soaked paper before punching), placed in 1.5 mL EP tube, smashed with a grinding rod, and then added with 500 μ L MgCl $_2$ (10 mM) and mashed as fully as possible. In 96-well plate, dilute the sample to 6 concentrations, and then coat on King's B media containing 50 μ g mL $^{-1}$ rifampicin and 100 μ g mL $^{-1}$ kanamycin at 28 °C. The colony count was observed and recorded 2 days later. ^{21,22}

Conductivity experiment

After 3 days of AuNPs-siRNA_{NPR1} infiltration, bacterial solution of Pst DC3000 (*AvrRps4*) with $OD_{600} = 0.02$ (diluted from stock solution of OD = 1.0) was infiltrated and sampled after 3 days.

Six small round leaves with a diameter of 8 mm were obtained with a perforator (75% alcohol disinfection). Then the leaves were placed in a 50 mL centrifugal tube, washed twice with 50 mL ddH₂O, and added 15 mL ddH₂O 10 minutes later. Measure and record data with conductivity meter (Mettler Toledo FE30-FiveEasy, Shanghai, China) every 2 hours.^{21,22}

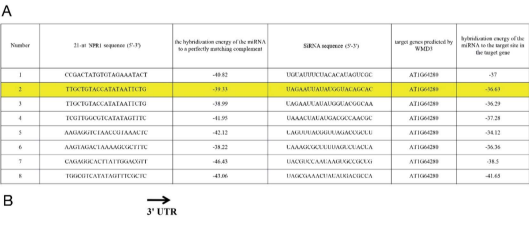
Design of different siRNAs

On the basis of *Arabidopsis* Information Resource (TAIR) genome annotation, we designed siRNA for silencing *NPR1* gene by using the Web MicroRNA Designer platform (WMD3, http://wmd3.weigelworld.org), sequence is: UAGAAUUAUAUGGUA-CAGCAC (Fig. 1A). SiRNA was inserted into 3' untranslated region (UTR) of *NPR1* gene (Fig. 1B).^{23,24} We added the FAM fluorescent sequence to the 5' end of the siRNA to construct FAM-siRNA_{NPR1}. The sequence of "UUCUCCGAACGUGU-CACGUTT" was used as a negative control. SiRNA_{n.c.} is a universal negative control without species specificity. It has been used in mice, humans and other species as a negative control.²⁵⁻²⁷

Statistics of plant resistance to Pst DC3000 (AvrRps4)

Count the number of colonies with corresponding dilution on the solid medium, and then calculate the bacterial growth by the following formula:

Colony-forming units (cfu)/leaf disc =
$$((\text{colonies} \times 10^{\text{dilution}}/10) \times 500)/2$$



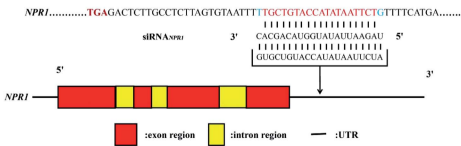


Fig. 1 Design of siRNA_{NPRI}. (A) Selection of siRNA_{NPRI} sequences (21mer) (highlighted by yellow) from the Web MicroRNA Designer platform. (B) The siRNA (red letters) is targeted to the 3'UTR of NPRI gene.

Colonies: the number of colonies counted from the row with the clearest number of colonies. Dilution: the row with the clearest colony number.

Plant material

Paper

Arabidopsis wild type (ecotype Columbia, Col) comes from Professor Kohki Yoshimoto of Japan Plant Science Center. *NPR1-GFP* (with the *npr1-2* mutant as the background) comes from the laboratory of Xinnian Dong of Duke University was obtained by the floral dip method. ** *Nicotiana benthamiana* wild type comes from the laboratory of Xinnian Dong of Duke University also.

Statistical analysis

All tests were repeated at least three times and the results were analyzed by GraphPad Prism 8.¹6 Statistical analyses were performed using Student's t-test (*, p < 0.05, **, p < 0.01, and ***, p < 0.001). Each value was the mean \pm S.D. of three independent replicates.

Result & discussion

Synthesis of AuNPs and siRNA-AuNPs

AuNPs have been used for delivery of siRNA, DNAs and proteins. ²⁹⁻³¹ The following mechanism has been proposed for the function of AuNPs: (1) oligonucleotides are covalently linked to the mercaptan sulfide to AuNPs to make functional surface; (2) the siRNA-AuNPs are recognized by the surface receptors of cells through endocytosis and penetrate cell membrane; and (3) siRNA are released from the particles and entered the nucleus (Fig. 2).⁸

The concentration of AuNPs is about 1 mM and the size of the synthesized AuNPs was in range of 3–45 nm (Fig. 3A) and similar to that reported in ref. 32, and might thus be easily absorbed by cells. In order to explore the optimal binding ratio of AuNPs and siRNA, we tested different combination of AuNPs with siRNA.

We mixed siRNA (20 μ M) and AuNPs (1 mM) in a ratio of 1 : 9 and diluting 15 times with a infiltration buffer (10 mM MgCl₂ and 10 mM MES, pH 5.7) for measure particle size and potential. The same infiltration buffer and AuNPs were mixed at a ratio of 1:9 and diluted 15 times with this infiltration buffer to measure the particle size and potential of AuNPs before binding to siRNA. The average size of AuNPs before binding siRNA is about 34 nm and the zeta potential is 25 mV. The particle size and zeta potential of AuNPs changed after binding with siRNA (Fig. 3B, C, Tables S2 and S3†). The particle size of AuNPs increased by about 4 nm (Fig. 3C and Table S3†). Because the surface of AuNPs is positively charged and the molecular surface of siRNA is negatively charged, the zeta potential reduces from about 25 mV to about 5 mV (Fig. 3B and Table S2†). These changes in particle size and zeta potential can explain the successful formation of the AuNPs-siRNA compounds.33 In addition, only slight change in particle size was found before and after bonding, which indicated that the

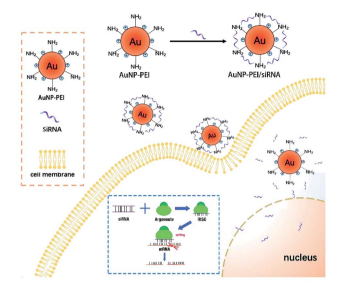


Fig. 2 Mechanisms of siRNA-AuNPs entering cell and nucleus. AuNPs and siRNA are covalently linked and enter cytoplasm through endocytosis. Then siRNA is released and enters the nucleolus through the nuclear pore. siRNA is melted under the action of intracellular RNA helicase, and then antisense siRNA is combined with some enzymes to form RNA-induced silencing complex (RISC) (Argonaute protein is a large class of proteins family, is the main member of the RISCs complex). RISC specifically binds to the homologous region of the mRNA expressed by an exogenous gene. RISC has the function of a nuclease and cleaves the mRNA at the binding site, leading to inhibition of the translation.

combination of AuNPs and siRNA will not cause the AuNPs to become too large and thus affect the infiltration effect.

For their inert and non-toxic properties and good biocompatibility, AuNPs were considered as a good carrier of nucleotides.34 In addition, AuNPs are easily prepared for desired size and dispersibility of biomolecules in complex with oligopeptides or nucleic acids, and may thus be integrated into biological systems.35 The large ratio of surface area/volume of the AuNPs will provide space for loading RNA or DNA.36 As a nucleic acid carrier, AuNPs can improve the survival time of DNA and siRNA in cellular environment, the efficiency of cell uptake from endoplasmic vesicles, and the subsequent release. 30,37 Positively charged AuNPs bind efficiently to negatively charged nucleic acid and also prevent them from being digested by enzymes.38 Through the "proton sponge" effect, the combination of branched PEI and AuNPs can transfer plasmid DNA to cells approximately 15 times more efficiently than unmodified PEI.³⁹ We modified the preparation procedure of AuNPs-PEI, and dialyze the suspension to remove PEI that is not combined with AuNPs to minimize PEI interference.13

AuNPs-FAM-siRNA_{NPR1} was observed to enter plant cells

The FAM-siRNA $_{NPRI}$ was mixed with AuNPs (AuNPs-siRNA $_{NPRI}$) in a ratio of 1:9, diluted three times with the buffer of a buffer of 10 mM MgCl $_2$ and 10 mM MES, incubated at 25 °C under 60 rpm shaking for half an hour, and then diluted five times. In order to investigate whether the silencing complex can enter

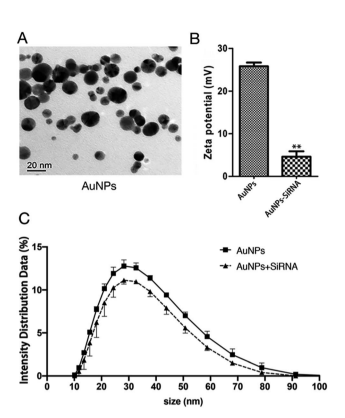


Fig. 3 Synthesis of AuNPs-siRNA $_{NPRI}$. (A) Transmission Electron Microscope (TEM) images of bare AuNPs. Bars = 20 nm. (B) Zeta potential of the AuNPs (left) and siRNA-AuNPs (right). Student's t-test (*, p < 0.05, **, p < 0.01, and***, p < 0.001). (C) The particle size of AuNPs and the change of particle size after AuNPs and siRNA are combined

plant cells, 5 weeks-old leaves of Nicotiana benthamiana were selected for infiltration of AuNPs-FAM-siRNA_{NPR1}. AuNPs-FAMsiRNA_{NPR1} infiltration was performed by pressure infiltration with a 1 mL syringe through the abaxial leaf surface. The criterion for successful infiltration was that the whole leaf was wet without obvious mechanical damage. Considering the quenching of fluorescence signal, we only observed the fluorescence of 1 h injection. In a relatively short period of time, only part of the AuNPs-FAM-siRNA_{NPR1} can enter the cell through endocytosis. So fluorescent signals can be detected in the cytoplasm and extracellular matrix. In order to determine the position of AuNPs-FAM-siRNA_{NPR1} in the cell and eliminate the possibility that the FAM fluorescence of the upper layer overlaps with that of the lower layer, the mesophyll cells were scanned 30 layers from top to bottom, and the images of each layer were decomposed. AuNPs-FAM-siRNA_{NPR1} are small spherical bright spots. FAM fluorescence exists in different positions of different layers, which can effectively explain that AuNPs-FAM-siRNA_{NPR1} are in the inner part of Nicotiana benthamiana mesophyll cells. At the same time, we provide the decomposition diagram of each layer (Fig. S1†). Then we selected the representative third layer, the ninth layer and the seventeenth layer for detailed discussion (Fig. 4A). Because the mesophyll cells of Nicotiana benthamiana are densely packed, we use white or orange to outline the borders of two adjacent cells for easy distinction, and the numbers represent different scanning layers. In order to further determine the location of AuNPs-siRNA_{NPR1}, AuNPs-FAM-siRNA_{NPR1} was infiltrated into Arabidopsis thaliana leaves (4 weeks-old Col-0 plants). After 1-2 hours of infiltration, plant protoplast was extracted from the leaves and the fluorescence in cells was observed by confocal microscope (Fig. 4B). The movies clearly showed that AuNPs-FAM-siRNA_{NPR1} was located in the plants (ESI Movies 1 and 2^{\dagger}).

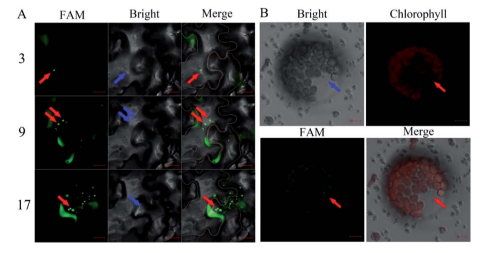


Fig. 4 Infiltration of the AuNPs-FAM-siRNA $_{NPRI}$ into plant cells. (A) Confocal image of AuNPs-FAM-siRNA $_{NPRI}$ fluorescence in *Nicotiana benthamiana* leaves. The cell edge is outlined with white and orange lines, and the number represents different scanning layers. The red arrow indicates that the green light spot is AuNPs-FAM-siRNA $_{NPRI}$, and the blue arrow indicates the same position of these fluorescence in the bright field. Images were taken under the excitation of 492 nm wavelengths (green is the fluorescence of FAM). Bars = 20 μ m. (B) Confocal images of *Arabidopsis* protoplast expressing FAM. The protoplasts were collected after 1–2 hours of infiltration of AuNPs-FAM-siRNA $_{NPRI}$ into *Arabidopsis* leaves. Images were taken with excitation at 492 nm and 518 nm wavelengths. The green light spots represented by the red arrow is AuNPs-FAM-siRNA $_{NPRI}$. The blue arrow indicates the same position of these fluorescence in the bright field and chlorophyll field. Bars = 10 μ m.

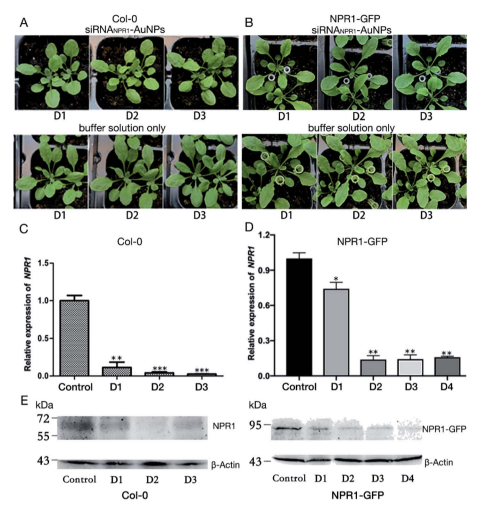


Fig. 5 Characterization and gene silencing effect of Col-0 and NPR1-GFP plants infiltrated with AuNPs-siRNA_{NPR1}. (A) The phenotype of the Col-0 plants infiltrated with AuNPs-siRNA_{NPR1} (top) or with buffer solution as control (bottom). (B) The phenotype of the NPR1-GFP plants infiltrated with AuNPs-siRNA_{NPR1} (top) or with buffer only (bottom). (C) expression of targeted genes in Col-0 plants. After infiltrated AuNPs-siRNA_{NPR1} into Col-0 plants, the leaves were collected every day and RNA was extracted from the leaves. Relative expression levels are shown as means \pm sd from three biological repeats. Student's t-test (*, p < 0.05, **, p < 0.01, and***, p < 0.001). (D) qPCR analysis on expression of targeted genes after infiltrated AuNPs-siRNA_{NPR1} in NPR1-GFP plants. Relative expression levels are shown as means \pm sd from three repeats. Student's t-test (*, p < 0.05, **, p < 0.01, and***, p < 0.001). (E) Western blotting analysis on protein expression in Col-0 plants after AuNPs-siRNA_{NPR1} infiltration. Our target band NPR1 is 66 kD. Col-0 plants infiltrated with buffer were used as positive control, and npr1 mutant was used as negative control.

We also provide an image of protoplasts of *Arabidopsis* leaf epidermal cells 1 hour after injection by AuNPs-FAM-siRNA $_{NPRI}$ (Fig. S2†). Since there is no chloroplast in epidermal cells, we can exclude the possibility that green fluorescence is the light of chloroplast. This shows that these small spherical bright spots must be AuNPs-FAM-siRNA $_{NPRI}$. Previous research shows AuNPs-siRNA can efficiently transfect animal cells, $_{37,40,41}^{37,40,41}$ our experiments show for the first time that the AuNPs-siRNA genesilencing compounds can be infiltrated into plant cells.

AuNPs-siRNA_{NPR1} silence NPR1 gene in Arabidopsis

Optimization on different combination of chemical compounds and their concentration may yield effective binding to the surface of the nanoparticles and thus save test time and cost. Multi-functional AuNPs have been shown to be very stable without acute toxicity and not to impair cell viability, ⁴⁰ and thus

are used in the plant gene silence. We chose Col-0 and *NPR1-GFP* as the experimental plant for test of duration of gene silencing, and used AuNPs-siRNA_{NPR1} to silence the *NPR1* gene for bacteriostatic and ion leakage experiments.

We conducted a series of experiments to explore whether AuNPs-siRNA_{NPR1} can successfully silence the *NPR1* gene of plants. The AuNPs-siRNA_{NPR1} were infiltrated into leaves of the Col-0 and *NPR1-GFP* (overexpressing *NPR1* gene in *npr1-2* background) *Arabidopsis*. The infiltration buffer was used to dilute the silencing complexes of AuNPs-siRNA_{NPR1} and as the blank control. There was no significant difference in phenotypes between the leaves infiltrated with silencing complex or plain buffer, indicating that the AuNPs-siRNA_{NPR1} do not affect the normal growth of both Col-0 and *NPR1-GFP* plants (Fig. 5A and B). To find out whether the compound of AuNPs and siRNA_{NPR1} impacts the health of plants, we observed and

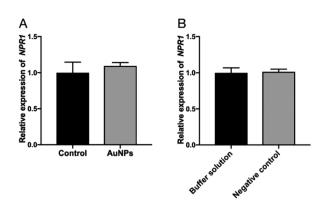


Fig. 6 AuNPs and unrelated siRNA do not cause *NPR1* gene silencing. (A) Effect of AuNPs on the *NPR1* gene expression after three days' infiltration of AuNPs or buffer solution. qPCR analysis showed no significant difference between the infiltration of AuNPs and the blank control. Relative expression levels are shown as means \pm sd from three repeats. (B) SiRNA_{n.c.} did not cause *NPR1* gene silencing. Relative expression levels are shown as means \pm sd from three repeats.

recorded the phenotype of *Arabidopsis* after infiltration of the compound. The results showed that the 1:9 ratio of 20 μ M siRNA_{NPR1} and 1 mM AuNPs is appropriate and has no impact on the normal growth of plants.

The qPCR analysis shows that the *NPR1* gene is effectively silenced after 1 or 2 days of infiltration of AuNPs-siRNA_{NPR1} into *Arabidopsis* Col-0 and *NPR1-GFP* plants. The effect of gene silencing is about 80% (Fig. 5C and D). The NPR1 protein in plant is significantly decreased after 1 and 2 days of infiltration of AuNPs-siRNA_{NPR1} to the Col-0 and *NPR1-GFP* plants (Fig. 5E). These results suggested that we have successfully silenced *NPR1* gene.

The infiltration of AuNPs and unrelated siRNA did not affect the expression of *NPR1* gene

In order to detect whether the AuNPs affect NPR1 gene, the AuNPs were infiltrated into leaves of the NPR1-GFP Arabidopsis. Infiltration buffer was used to dilute the AuNPs and also as the blank control. Then, we combined $siRNA_{n.c.}$ with AuNPs and infiltrated it into NPR1-GFP (the other operations were the same as before). Three days after infiltration, the leaves were collected

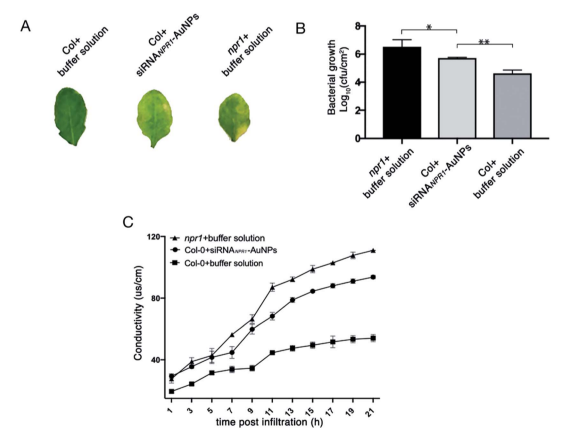


Fig. 7 NPR1 involvement in the regulation of cell death and disease resistance to Pst DC3000 (AvrRps4). (A) Phenotypes of leaves after 3 days infiltration of Pst DC3000 (AvrRps4) with different reagents. (B) Silencing of NPR1 in Col-0 changed plant resistance to Pst DC3000 (AvrRps4). After 3 days of infiltration with AuNPs-siRNA_{NPR1}/buffer solution, the leaves are infiltrated with Pst DC3000 (AvrRps4). The resistance of Col-0 plants infiltrated with AuNPs-siRNA_{NPR1} to Pst DC3000 (AvrRps4) was between positive control (Col-0 plants infiltrated with buffer solution) and negative control (npr1 plants). Student's t-test (*, p < 0.05, **, p < 0.01, and ***, p < 0.001). (C) Statistics of ion leakage of plants infiltrated with different reagents for 3 days. The ion leakage rate of Col-0 plants infiltrated with AuNPs-siRNA_{NPR1} is between the positive control (Col-0 plants infiltrated with buffer solution) and the negative control (npr1 plants).

for qPCR experiment. These experiments suggesting that AuNPs does not affect the expression of *NPR1* gene (Fig. 6A), simultaneously, unrelated siRNA does not silence *NPR1* gene (Fig. 6B), and that siRNA $_{NPR1}$ induced silence of part of the *NPR1* gene (Fig. 5C and D).

Bacterial stress and ion leakage experiments confirmed AuNPs-siRNA_{NPR1} silencing NPR1

NPR1 is involved in the regulation of cell death and disease resistance in response to non-pathogenic pathogens. After 3 days of infiltrating AuNPs-siRNA_{NPR1} into Arabidopsis Col-0 plants (the infiltration buffer was infiltrated in npr1 and Col-0), Pst DC3000 (AvrRps4) with $OD_{600} = 0.001$ was infiltrated. Three days later, the infiltrated leaves were collected and subjected to a bacteriostatic test. The avirulent Pst DC3000 (AvrRps4) infects the leaves of Arabidopsis thaliana, leading to spread of chlorosis symptoms.15 The results showed that the leaves of NPR1 silenced plants with infiltration of Pst DC3000 (AvrRps4) turned to yellow and so do the leaves of npr1 mutant, but the leaves of Col-0 plant remain green (Fig. 7A). The collected leaves were ground, dilute to six concentrations then they were cultured on King's B solid medium resistant to kanamycin and rifampicin. Under Pst DC3000 (AvrRps4) stress, the NPR1 partly silenced plants showed bacterial growth (the number of colony of the plants) more than that of the Col-0 plants, but less than that of the npr1 mutant (Fig. 7B, S3 and Table S1†). Three days after AuNPs-siRNA_{NPR1} were infiltrated into Arabidopsis thaliana Col-0 plants (buffer was infiltrated in npr1 and Col-0), Pst DC3000 (AvrRps4) $OD_{600} = 0.02$ was infiltrated and the conductivity was measured one hour later. The NPR1 partly silenced plants showed cell death more than the Col-0 plants, but less than the *npr1* mutant (Fig. 7C). These results suggested that AuNPs-siRNA_{NPR1} partially silenced NPR1 gene, thus indicating our gene silencing technique is effective.

In short, we show, for the first time, AuNPs-siRNA $_{NPRI}$ can be infiltrated into the leaf cells and partially silence the NPR1 gene in Arabidopsis thaliana. In comparison with the current methods for plant gene silencing, which require pollutant Agrobacterium tumefaciens and are time consuming, our AuNPs-siRNA $_{NPRI}$ for silencing plant genes have advantages of short time, low toxicity, and low dosage. This convenient and rapid method for gene silencing in plants provides a new way to study the gene function of plants.

Conflicts of interest

There are no conflicts to declare.

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