Isolation and Identification of Post-Transcriptional Gene Silencing-Related Micro-RNAs by Functionalized Silicon Nanowire Field-effect Transistor

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Many transcribed RNAs are non-coding RNAs, including microRNAs (miRNAs), which bind to complementary sequences on messenger RNAs to regulate the translation efficacy. Therefore, identifying the miRNAs expressed in cells/organisms aids in understanding genetic control in cells/organisms. In this report, we determined the binding of oligonucleotides to a receptor-modified silicon nanowire field-effect transistor (SiNW-FET) by monitoring the changes in conductance of the SiNW-FET. We first modified a SiNW-FET with a DNA probe to directly and selectively detect the complementary miRNA in cell lysates. This SiNW-FET device has 7-fold higher sensitivity than reverse transcription-quantitative polymerase chain reaction in detecting the corresponding miRNA. Next, we anchored viral p19 proteins, which bind the double-strand small RNAs (ds-sRNAs), on the SiNW-FET. By perfusing the device with synthesized ds-sRNAs of different pairing statuses, the dissociation constants revealed that the nucleotides at the 3′-overhangs and pairings at the terminus are important for the interactions. After perfusing the total RNA mixture extracted from Nicotiana benthamiana across the device, this device could enrich the ds-sRNAs for sequence analysis. Finally, this bionanoelectronic SiNW-FET, which is able to isolate and identify the interacting protein-RNA, adds an additional tool in genomic technology for the future study of direct biomolecular interactions.

Micro-ribonucleic acids (miRNAs) with lengths of 21 to 22 nucleotides have sequence specificities that guide RNA-induced silencing complexes to cleave the complementary messenger RNAs (mRNAs), resulting in post-transcriptional gene silencing (PTGS)1,2. This is an important mechanism in controlling the expression of specific genes during the development of an organism3. Most miRNA detection methods, such as microarray, Northern blot, and reverse transcription-quantitative polymerase chain reaction (RT-qPCR), are designed for probing single-stranded RNAs4,5 but are time-consuming and laborious.

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The viral p19 proteins form dimers and sequester double-strand small RNA duplexes (ds-sRNAs, including miRNA/miRNA*, where miRNA* represents the complementary anti-sense miRNA) in host cells, leading to blocking the host cell RNA interference (RNAi) defense mechanism and prevent the viral RNAs from being digested by gene silencing mechanism. Therefore, p19 is a good candidate for concentrating the ds-sRNAs in an RNA mixture to characterize the miRNAs expressed in a cell or organism. Some reports have adopted p19 as a screening tool to study miRNA expression in cancer cells or tissues; however, these approaches are not reusable and require isotopes to enhance sensitivity. Additionally, they lack real-time detection.

To characterize the miRNA profile expressed in cells, we applied silicon nanowire field-effect transistor (SiNW-FET) biosensors that provide ultra-sensitive, real-time, and reversible detection. In previous studies, SiNW-FETs were used to detect the interaction between biomolecules and to sense the virus DNA or the presence of a specific miRNA from cell extracts with the femtomolar sensitivity by modifying the SiNWs with complementary oligonucleotides. Therefore, SiNW-FET has the potential in detecting the miRNA expressed in minute levels from an extracted mixture.

To rapidly screen the target-receptor interactions, we applied reusable SiNW-FET devices with reversible surface functionalization based on our previously developed strategy. The conductance of the SiNW-FET is determined by the electric field generated from the molecules surrounding the nanowire; henceforth, the SiNW-FET is very sensitive in monitoring the interaction among biomolecules in a real-time mode. In addition, the anchorage of the receptor molecules on the SiNW-FET surface is reversible; therefore, we could not only analyze the target-receptor interaction but also elute the bound target-receptor complex for analysis.

Results and Discussion

The expressions of RNAs are under strict regulation and usually in a small quantity except those house-keeping genes. RT-qPCR is the most sensitive technique to identify the expression of a specific RNA from a mixture owing to its precise amplification procedure. To verify the detection sensitivity and target selectivity of a SiNW-FET in probing the miRNAs of interest from total extracted RNA, we anchored the single-strand DNAs (ss-DNAs) onto a 3-mercaptopropyl-trimethoxysilane (MPTMS)-modified SiNW-FET (referred to as SH/SiNW-FET) via disulfide bonding (referred to as DNAprobe/SiNW-FET) and perfused the DNAprobe/SiNW-FET with extracted RNA for the selective binding of the complementary miRNA to the DNAprobe. We then eluted the bound DNAprobe miRNA complexes using dithiothreitol (DTT) to reduce the disulfide bonds and returned the SiNW-FET surface to its original state for device reusability.

To detect the miR159 (5′-UUUGGAUUGAAGGGAGCUCUA-3′), a miRNA which regulates the plant development and fertility, from the total RNA of Arabidopsis, we linked a synthesized miR159probe onto the SiNW-FET. In the repeated cycles of electrical measurements (Fig. 1B), the reproducible changes in conductance (ΔGs) of an SH/SiNW-FET in the serial processes of immobilizing miR159probe, introducing the total RNA, and eluting the miRNA159/miR159probe complex demonstrated the reusability of the SiNW-FET biosensor. To confirm and quantify the miRNA of interest present in the eluted mixtures and the total extracted RNA, we performed RT-qPCR with specific primers against miR159 (Fig. 1C). Due to the small sensing surface area of the SiNW-FET, the bound miR159 was 0.18% of the miR159 in the total extracted RNA. As a control test, miR168 presents in the total extracted RNA, but not in the eluted complexes. These results demonstrate the specificity of the DNAprobe/SiNW-FET in recognizing the target miRNA and we could elute the bound oligonucleotides for qPCR amplification or sequencing analysis.

MiRNAs guide the RNA-induced silencing complex to cleave the complementary messenger RNAs, resulting in the PTGS to control the expression level of a specific gene. Henceforth, the information about the relative amount of a specific miRNA in different cells is valuable in understanding the genetic control. To verify that the SiNW-FET is capable of monitoring differential expression of the same miRNA in different cells, we quantified miR21, which is an miRNA that is highly expressed in breast cancer, from the total RNA isolated from an MCF-7 cell line (derived from breast cancer) and an M10 cell line (derived from normal mammary epithelial cells). Figure 1D shows that, using RT-qPCR, MCF-7 had an ~5-fold higher miR21 expression level than M10, while the detection limits in both cell lines are ~0.07 μg/μl of the total RNA. In contrast, the miR21probe/SiNW-FET was able to detect miR21 from both samples with a total RNA concentration at 0.01 μg/μl (Fig. 1E) and MCF-7 had an ~3-fold higher miR21 level than that of M10. Therefore, both approaches (RT-qPCR and SiNW-FET) demonstrate a similar difference in the expression level of miR21 in these two cell lines. Using RT-qPCR to quantitate the amount of an RNA sample, we need to convert the extracted RNA to cDNA first and then amplify the oligonucleotide segments of interest with specific primers. The detection limit and target sensitivity might be lost during the conversion process but are greatly improved by the amplification procedure. In contrast, a SiNW-FET monitors the direct interaction between the designate miRNA in the total RNA extract and the complementary DNAprobe modified on the SiNW-FET; henceforth, a SiNW-FET exhibits high probing sensitivity and detection efficiencies in sample amount and time spent. A recent report even shows that SiNW-FET could have a limit of detection of 1 zeptomole in detecting the miRNA from cancer cell extracrs. These measurements demonstrate that the detection sensitivity of a SiNW-FET biosensor is comparable to RT-qPCR in monitoring differentially expressed genes and support that this SiNW-FET device can be applied to detect the minute expression of a specific miRNA from a mixture.
Figure 1. Detection of the endogenous miRNA by SiNW-FET. (A) A flow diagram of a reusable DNA probe/SiNW-FET device. The MPTMS-modified SiNW-FET (SH/SiNW-FET) provides reversible disulfide bonding sites for the DNA probe tagged with a thiol group at the 3' end (DNA probe/SiNW-FET). After the targeted miRNAs bind to the DNA probe/SiNW-FET, the bound DNA probe-miRNA complex can be eluted by flushing dithiothreitol (DTT) to break the disulfide bond, returning the device surface to SH/SiNW-FET. (B) The electrical conductance changes (ΔGs) of SH/SiNW-FET during repeated cycles of DTT-buffer washing, miR159 probe modification, and RNA binding (0.3 μg/μL total RNA extracted from Arabidopsis). (C) miRNA159 in the Input and Eluted mixtures. After using miR159 probe/SiNW-FET, the relative amounts of miR159 (upper panel) and miR168 (lower panel) in the bound fraction (Eluted) to the total RNA (Input) extracted from Arabidopsis were analyzed by RT-qPCR. (D,E) Comparison of the detection limits between RT-qPCR and SiNW-FET. We determined the amounts of miR21 expressed in different concentrations of total RNA extracted from cancer cell lines, MCF-7 and M10, by (D) RT-qPCR with specific primers or (E) miR21 probe/SiNW-FET.
To demonstrate the sequestration of the ds-sRNAs by p19, we modified the SiNW-FET with glutathione (referred to as GSH/SiNW-FET) to support the binding of glutathione S-transferase-conjugated p19 (referred to as p19/SiNW-FET) (Fig. 2A)\(^{17,18,24}\). We performed electric measurements in phosphate solution (containing 240 \(\mu\)M NaH\(_2\)PO\(_4\) and 760 \(\mu\)M Na\(_2\)HPO\(_4\) at pH 7.4 with NaOH) with a

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**Figure 2.** Discriminating the secondary structures of ds-sRNA by p19/SiNW-FET. (A) A flow diagram of a reusable p19/SiNW-FET using the GSH/GST association-dissociation. The process includes the immobilization of GST-p19 on a GSH/SiNW-FET to form p19/SiNW-FET, the application of ds-sRNA (miRNA/miRNA*) to bind p19, and the elution of the GST-p19-ds-sRNA complexes with \(\geq 1\) mM GSH. (B) The responses of p19/SiNW-FET to various forms of nucleic acids. The normalized \(\Delta G\)s were measured by introducing 1 \(\mu\)M of a 21-nucleotide solution in (i) perfectly matched ds-sRNA form (ds-sRNA-0), (ii) ss-sRNA form (ss-sRNA-0), (iii) ds-sDNA form (ds-sDNA-0) to a p19/SiNW-FET; (iv) p19 replaced by a binding-deficient mutant (p19 mut); and (v) p19 absent in the tests. (C) The dissociation constant \((K_d)\) of p19 to various 21-nucleotide nucleic acids with different mismatch pairings. The \(\Delta G\)s of p19/SiNW-FET to different concentrations of nucleic acids were used to determine the \(K_d\) values (Supplementary Fig. S5), which were then normalized to that of the ds-sRNA-0 \((K_{rel})\). Nucleotides marked in red indicate the mispaired bases.
signal-to-noise (S/N) were increased largely to 117 for ds-sRNA-1 with three mismatched bases at nucleotide positions of 9–11, were mutated to S, G, S, and G, respectively (Supplementary Fig. S3)26,29. The lack of observable p19 binding (Fig. 2B(iv)) indicated negligible binding between ds-sRNA and p19 mut; in addition, the p19mut could not bind the fluorescence-tagged ds-sRNA as p19 did (Supplementary Fig. S4). These results reveal that p19 selectively binds ds-sRNA and the 2′ hydroxyl group of RNA plays a significant role in the association with p19.

To characterize whether p19/SiNW-FET could differentiate the mismatched pairing in ds-sRNA, we synthesized five ds-sRNAs (ds-sRNA-1~5 with their structures being listed in Fig. 2C) with mismatched nucleotides or blunt ends and calculated the dissociation constants (Kd) between the p19/SiNW-FET and ds-sRNA-x (x = 0–5). By plotting the measured ∆Gs as functions of the sample concentration and fitting these data to the Langmuir adsorption isotherm model (Fig. S5)30,31, the Kd of the p19-ds-sRNA-0 complex was 16±5 nM, similar to the previously reported values32,33,34, and the detection limit was 1 nM with signal-to-noise (S/N) ≥ 3. For ds-sRNA-1 with three mismatched bases at nucleotide positions of 9–11, the Kd was slightly increased to 55±11 nM; for ds-sRNA-2 and ds-sRNA-3, of which both have mismatched bases at the termini, the Kd values were greatly increased to 227±46 nM and 1138±164 nM, respectively. For ds-sRNA-4 with a single blunt end and ds-sRNA-5 with two blunt ends, the Kd values were increased largely to 117±25 and 722±71 nM, respectively. These results support previous reports that the two tryptophans of a p19 monomer interact with the overhangs at the 3′-ends of miRNA/miRNA* duplexes29. In addition, the termini of the RNA duplex are more critical in binding to p19 than the central region, and our p19/SiNW-FET biosensor exhibits binding affinities that can distinguish various ds-sRNA secondary structures.

To verify that p19/SiNW-FET could selectively sequester ds-sRNAs from an RNA mixture, we perfused the p19/NW-FET with synthesized ds-sRNA-0 (Fig. 3A) or extracted RNA from Arabidopsis (Fig. 3B) and analyzed the relative amount of RNA (the sense vs. anti-sense strands of ds-sRNA-0) by RT-qPCR. Perfusion of the synthesized ds-sRNA-0 or total extracted RNA across the p19/NW-FET induced a decrease in the conductivity (Supplementary Fig. S6). Figure 3A shows that the normalized amounts of the sense and anti-sense strands of ds-sRNA-0 in the input mixture were 100 and 67.8%, respectively; whereas the relative amounts in the eluted sample were 21.7 and 20.6%, respectively. These results revealed that the sense and anti-sense strands of ds-sRNA-0 are not all paired in the input mixture; however, after p19 sequestration, the amounts of the sense and anti-sense strands are almost of equal amounts suggesting that the RNA are mostly paired. Unlike the synthesized ds-sRNA-0, endogenous ds-sRNAs degraded quickly; so the total RNA (input) contains very few miR168* (4.9%) relative to the miR168 (regarded as 100%) as shown in Fig. 3B; however, the eluted sRNAs contains 1.8 and 1.1% of miR168 and miR168*, respectively. The nearly equal amounts of sense and anti-sense miRNAs in the eluted samples demonstrate the binding specificity of p19 to ds-sRNA.

To verify p19/SiNW-FET is useful in enriching the miRNA/miRNA* duplexes specifically in the total RNA extracted from N. benthamiana, we dissociated the GSH-GST association to release the p19-ds-sRNA complexes from the SiNW-FET surface and used a next generation sequencing (NGS) technique to analyze the sequence of each ss-sRNA in the total (Input) and captured (Eluted) samples (Fig. 3C). The abundance of a specific ss-sRNA is determined by the times of the identified sequences (read counts). In total extracted RNA, 80% of the 4,653,979 ss-sRNAs identified was sequenced one time (1 read count), and only 1.1% had counts of ≥ 16. For the eluted ss-sRNAs (total count is 199,001), 39% was sequenced one time and > 50% had counts of ≥ 16. The decrease and increase in the fractions of low and high sequence counts, respectively, support the ability of p19 in selectively enriching certain sRNAs. To demonstrate that the eluted sRNA contains ds-sRNA, we paired the possible ds-sRNA from the sequences obtained according to the following algorithms: (1) read counts of > 30, (2) two nucleotides at the 3′ overhang on both termini, (3) less than four mismatched pairing, and (4) permissible G:U pairing in the RNA structure. The above requirements (1)–(3) were decided according to the results of
Fig. 2C which shows the different association constants between p19 and ds-sRNAs with different pairing statuses; the requirement (4) is a general rule for RNA pairing. Using these criteria, we identified 459 and 3,752 possible ds-sRNA pairs present in the input and eluted sRNA samples, respectively. Considering the total number of ss-sRNA sequenced in the input (4,653,979) and eluted (199,001) fractions, the ds-sRNA fraction dramatically increased after p19 enrichment (i.e., 459/4,653,979 in the input and 3,752/199,001 in the eluted). These results confirm that most of the enriched sRNAs after p19/SiNW-FET sequestration are ds-sRNAs that were derived from the sRNAs with low abundances in the cells.

Single RNA strands, like transfer RNAs or ribosomal RNAs, form special hairpin structures for various functions. To generate the miRNAs for gene silencing, each transcribed pre-miRNA forms a hairpin structure with the sense and anti-sense strands of ds-sRNA-0 in the total and eluted fractions. (B) The relative amounts of miR168 (sense) and miR168* (anti-sense) in the extracted and bound fractions. (C) Proportions of the analyzed sRNAs with different read counts. The number of each sRNA sequenced was counted (read counts) and binned with a power of 2; the number of sRNAs in each binning group was normalized to the total number of sRNA counted. (D) Predicted structures of three pre-miRNAs yielding the corresponding paired ds-sRNAs captured on p19/SiNW-FET. According to the genome of *N. benthamiana*, nine genes (Table S1) might transcribe pre-miRNAs which have 2nd structures yielding paired ds-sRNAs (bases in bold) identified from the Elute fraction. The red lines indicate the matured miRNA forms. The digits beside each sRNA segment (bases in bold) are the counts of the corresponding sRNA segment appearing in the Input (I) or Eluted (E) sRNA sample. The data (mean ± standard deviations) were the averages of three independent experiments and ** indicates the *p*-value < 0.01. A magnified image of Fig. 3D is shown in Supplementary Fig. S7 for easier reading.
The counts of these three miRNA/miRNA* pairs in the total extracted RNA were 299/23, 207/4, and 194/24, respectively, indicating that after digestion of the pre-miRNAs in cells, most miRNA* were degraded. However, after p19 sequestration, the miRNA/miRNA* ratio tends to approaching unity, i.e. 79/36, 49/39, and 172/117, respectively, for the three predicted structures. These results further support that the p19/SiNW-FET binds ds-sRNAs with high affinity and specificity.

In conclusion, the reversible design of a SiNW-FET device facilitates the isolation and identification of ds-sRNAs from samples with low biologically relevant concentrations, demonstrating that a SiNW-FET can be used as a biosensor for the fast analysis of sRNAs in genomic studies. In addition, the direct interaction between a nucleic acid probe, immobilized on the SiNW-FET, and the target nucleic acid provides a biosensing tool with a sensitivity at the femtomolar level, which is suitable for most biomedical diagnoses. Although we have improved the reusability of a SiNW-FET device in detecting biomolecular interactions via a reversible surface modification method for calibratable and quantitative analyses, we could not ignore the technical challenges and environmental confrontations for SiNW-FETs, e.g., neutral molecule detection, sensing in a high salt buffer, and high-throughput multiplexed sensing.

In this report, we have shown that the p19/SiNW-FET can not only offer highly sensitive detection for miRNA but also probe the low abundance of intermediate forms of miRNA/miRNA* duplexes with high affinity and specificity. Therefore, with appropriate modifications to SiNW-FET, our system has great potential for biomedical applications in detecting protein-RNA interactions with high sensitivity and reusability. Finally, this bionanoelectronic device, which is able to isolate and identify the interacting protein-RNA, adds an additional tool in genomic technology for the future study of direct biomolecular interactions.

**Methods**

**Fabrication of SiNW-FET devices.** The SiNW-FETs were fabricated from silicon-on-insulator (SOI) wafers, composed of a layer of high-quality single-crystal silicon (device layer), a layer of buried oxide (BOX layer), and the bulk substrate. These SOI wafers are currently state-of-the-art in the production of metal-oxide-semiconductor field-effect-transistors (MOSFETs). In this study, 4-inch n-type SOI wafers containing a 50-nm-thick device layer and a 400-nm-thick BOX layer with a device layer resistivity of approximately 100 Ω-cm and a phosphor doping concentration of 4 × 10^{19} cm^{-3} were used. After standard electron-beam lithographic and photolithographic fabrication procedures, the SOI chip was thermally oxidized to form a SiO₂ insulating layer (~10 nm in thickness) on top of the SiNW surface to prevent the charge transfer between analyte molecules and the SiNW-FET. The advantages of this SiO₂ dielectric layer are not only providing an insulator layer to prevent electrical leakage to the aqueous buffer solution during biosensing measurements, but also facilitating the coupling of a PDMS channel onto the SiNW-FET chip for subsequent experiments without contaminating the SiNW-FET devices. Each chip (14 × 14 mm² in dimension) contains eight SiNW-FET devices (L 6 μm × W 200 nm × T 50 nm for each SiNW, Supplementary Fig. S1). The contact pads (Ni/Au = 10/50 nm) were fabricated by photolithography and were attached directly to the Si device layer.

**Reversible surface modification of the SOI chip.** The reversible surface modification of SiNW-FETs can be realized via either the formation of disulfide bonds, or the association-dissociation of GSH/Si (PDMS) microfluidic channel (L 6.25 × W 0.5 × H 0.05 mm³), which was designed to couple the SiNW-FET device arrays, with a syringe pump at a flow rate of 300 μL/hr for 30 min. After MPTS was modified on the SiNW-FET (denoted by SH/SiNW-FET), ethanol was guided into the PDMS microfluidic channel to wash the FET devices for 10 min. It is noted that ethanol solution might swell the PDMS channel and MPTS could also be modified on the PDMS channel walls; to minimize this disadvantage, the swollen PDMS channel could be replaced with a new one for the following experiments. Subsequently, 500 mM dithiothreitol (DTT) in 1× PBS (containing 138 mM NaCl, 2.7 mM KCl, 8 mM Na₂HPO₄, 1.5 mM KH₂PO₄, pH 7.4 with NaOH) was pumped into the PDMS channel for 30 min to reduce any possible disulfide bonding between the thiol groups of MPTS. The SH/SiNW-FET was then flushed with 1× PBS for 20 min. Next, a 1× PBS solution containing 1 μM DNA (disulfate DNA-dimer) and 100 mM DTT was pumped into the PDMS microfluidic channel. Finally, the sulphydryl group of the DNA probe reacted with the thiol group of MPTMS to anchor the DNA probe on the MPTMS/SiNW-FET (denoted by SH/SiNW-FET). After each electrical measurement, where miRNAs bound specifically to the DNA probe/SiNW-FET, the DNA-miRNA complexes were removed by cleaving the disulfide bond with DTT (500 mM in 1× PBS) for 2 hr to return the device surface to SH/SiNW-FET, thus making the SiNW-FETs reusable.

In the modification of GSH on the SiNW-FET surface (Fig. 2A), the SOI chip containing eight SiNW-FET devices was immersed in 1% 3-aminopropyltrimethoxysilane (APTS) in ethanol for 30 min and then heated at 110°C for 10 min to form a self-assembled layer of APTMS on the SOI surface. A linker of 3-maleimidobenzoic acid N-hydroxysuccinimide ester (MBS), prepared in a 1:9 mixture of dimethyl sulfoxide and 1× PBS, was applied to react APTMS through the formation of an amide bond. Subsequently, the SOI chip was immersed in 1× PBS containing 1 mM GSH for 30 min to induce bonding...
between the sulphydryl group of GSH and the maleimide group of MBS (to form a GSH/SiNW-FET), washed with 1× PBS, and blow-dried with N₂.

**Electrical measurement.** In the electrical measurements with a SiNW-FET, the sample molecules were diluted in 0.1× PS (phosphate solution, containing 240 µM Na₂HPO₄, 760 µM Na₂HPO₄, pH 7.4 with NaOH) with the Debye–Hückel screening length of λₑ = 6.1 nm. The sample solution, pumped into a PDMS microfluidic channel with a syringe pump at a flow rate of 0.3 mL/h, was delivered to the SiNW-FET surface. All sensing measurements were conducted using a detection system that combined a current preamplifier (1211, DL Instruments) and a lock-in amplifier (SR830, Stanford Research Systems) operating at a source-drain voltage (V₀) of 30 mV, a modulation frequency of 79 Hz, and a time constant of 100 ms. An Ag/AgCl electrode (MF2052, BASi) connected to the PDMS microfluidic channel could be used as a solution gate. To minimize noise, the Ag/AgCl electrode was maintained at ground potential throughout the electrical measurements.

The charge sensitivity of a SiNW-FET is in principle the same as that of thin-film FETs, provided that the thickness of the thin film does not exceed the Debye length of the semiconducting channel. In that case, the operation of both SiNW-FETs and thin-film FETs are in the fully depleted regime.

**Amplification and quantification of the sRNA sequence using RT-qPCR.** The RT-qPCR method used to amplify and quantify the gene expression levels has been described previously. For the reverse transcription (RT) reaction, a combination of 4.5 µL of the p19/SiNW-FET eluents, 50 nM of a stem-loop RT primer, 0.25 mM of dNTP, 20 units reverse transcriptase, 1× reverse transcriptase buffer, 10 mM of DTT, and 4 units of an RNAse inhibitor was prepared. The reactions were incubated at 16 °C for 30 min, followed by pulsed RT for 60 cycles at 30 °C for 30 s, 42 °C for 30 s, and 50 °C for 1 s to produce the first-strand cDNA. The RT reaction was terminated by heating at 85 °C for 5 min.

RT-qPCR was performed using the Universal Probe Library assay (Roche Diagnostics, Indianapolis, Indiana, USA) to amplify and quantify the cDNA synthesized from the miRNA captured by the p19/SiNW-FET. For the PCRs, 2.7 µL of the cDNA mixture was combined with 0.5 µM of each forward and reverse primer, 0.1 µM of Universal Probe Library Probe no. 21, and 1× Master Mix solution. The PCR amplification curves were generated using an initial denaturation at 95 °C for 10 min, followed by 45 cycles of 95 °C for 5 s, 60 °C for 30 s, and 72 °C for 1 s. The data were collected and analyzed using a LightCycler 480 real-time PCR instrument (Roche Molecular Systems, Pleasanton, CA, USA).

For the PCR reaction, the universal reverse primer was 5′-GGTGCAGGGTCCGAGGTATTCGCACCAGAGCCAACACCGTA-3′, and 5′-GTGATGTAGCCTCCCCTCACAGGCTACTCGCCTTTCTTGTC-3′ for the PCRs, 2.7 µL of the cDNA mixture was combined with 0.5 µM of each forward and reverse primer. The RT-qPCR method used to amplify and quantify the gene expression levels has been described previously. For the reverse transcription (RT) reaction, a combination of 4.5 µL of the p19/SiNW-FET eluents, 50 nM of a stem-loop RT primer, 0.25 mM of dNTP, 20 units reverse transcriptase, 1× reverse transcriptase buffer, 10 mM of DTT, and 4 units of an RNAse inhibitor was prepared. The reactions were incubated at 16 °C for 30 min, followed by pulsed RT for 60 cycles at 30 °C for 30 s, 42 °C for 30 s, and 50 °C for 1 s to produce the first-strand cDNA. The RT reaction was terminated by heating at 85 °C for 5 min.

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**Dissociation constant.** The dissociation constant (Kd) of the p19-ds-sRNA complex was determined by a least-squares fit of the Cds-sRNA/ΔG vs. Cds-sRNA data (Supplementary Fig. S5C) to the Langmuir adsorption isotherm model of Eq. (1,31)

\[
\frac{C_{ds-sRNA}}{\Delta G} = \frac{1}{\Delta G_{\text{max}}} \cdot C_{ds-sRNA} + \frac{1}{\Delta G_{\text{max}}} \cdot K_d
\]

where \(\Delta G_{\text{max}}\) is the saturated \(\Delta G\) at \(C_{ds-sRNA} \geq 200 \text{nM}\) (Supplementary Fig. S5B). As plotted in Supplementary Fig. S5C, the fit to the experimental data yielded \(K_d = 15.9 \pm 4.8 \text{nM}\) of the p19-ds-sRNA complex.

**Molecular biology.** We amplified the full-length p19 from infectious clones of the CymRSV12,38 by PCR using the p19-F1 (5′-CCCCGAGTCTCATGAAAGCAGCTACAAAGG-3′) and p19-R1 (5′-GCGGCCCTCGAGCTTCTTCTTTTCTTGAAGGC-3′) primers and inserted into the pGEX-4T-1 bacterial plasmid (GE Healthcare, Waukesha, WI, USA) for protein expression. To construct the mutated p19 (p19mut), we used p19-Mut-FW (5′-GACGAAAGTCCGAGTGGCTTGAGGGAGGTCTACATCAG-3′) and p19-Mut-Rev (5′-GTGATGTAGCCTCCCCTCGAGCTACAACTTCTTGTCG-3′) primers for PCR-mediated mutagenesis and inserted the product into pGEX-4T-1. To express the proteins, we transformed the BL21 (DE3) strain of Escherichia coli with these constructs and induced the protein expression by adding isopropyl-ß-D-1-thiogalactopyranoside (IPTG, 1 mM) into the lysogeny broth (LB) medium.

We extracted the total RNA from the leaves of 2-week-old Arabidopsis thaliana (col-0) seedlings using the Trizol reagent (Invitrogen, Carlsbad, CA, USA). The 21-nucleotides positive-stranded ds RNA was synthesized by RT-qPCR method used to amplify and quantify the gene expression levels has been described previously. For the reverse transcription (RT) reaction, a combination of 4.5 µL of the p19/SiNW-FET eluents, 50 nM of a stem-loop RT primer, 0.25 mM of dNTP, 20 units reverse transcriptase, 1× reverse transcriptase buffer, 10 mM of DTT, and 4 units of an RNAse inhibitor was prepared. The reactions were incubated at 16 °C for 30 min, followed by pulsed RT for 60 cycles at 30 °C for 30 s, 42 °C for 30 s, and 50 °C for 1 s to produce the first-strand cDNA. The RT reaction was terminated by heating at 85 °C for 5 min.

For the PCR reaction, the universal reverse primer was 5′-GGTGCAGGGTCCGAGGTATTCGCACCAGAGCCAACACCGTA-3′, and 5′-GTGATGTAGCCTCCCCTCACAGGCTACTCGCCTTTCTTGTC-3′ for the PCRs, 2.7 µL of the cDNA mixture was combined with 0.5 µM of each forward and reverse primer, 0.1 µM of Universal Probe Library Probe no. 21, and 1× Master Mix solution. The PCR amplification curves were generated using an initial denaturation at 95 °C for 10 min, followed by 45 cycles of 95 °C for 5 s, 60 °C for 30 s, and 72 °C for 1 s. The data were collected and analyzed using a LightCycler 480 real-time PCR instrument (Roche Molecular Systems, Pleasanton, CA, USA).

The amplification reaction was performed using the Universal Probe Library assay (Roche Diagnostics, Indianapolis, Indiana, USA) to amplify and quantify the cDNA synthesized from the miRNA captured by the p19/SiNW-FET. For the PCRs, 2.7 µL of the cDNA mixture was combined with 0.5 µM of each forward and reverse primer, 0.1 µM of Universal Probe Library Probe no. 21, and 1× Master Mix solution. The PCR amplification curves were generated using an initial denaturation at 95 °C for 10 min, followed by 45 cycles of 95 °C for 5 s, 60 °C for 30 s, and 72 °C for 1 s. The data were collected and analyzed using a LightCycler 480 real-time PCR instrument (Roche Molecular Systems, Pleasanton, CA, USA).
sRNA (5′-CGUACGCGGAUACUUCGAU-3′) and negative-stranded sRNA (5′-UUGCAUGCGCUCUAGAAGCU-3′) were synthesized by MDBio (Taipei, Taiwan). The 5′-ends of the sRNAs were phosphorylated and a Cy3 fluorescent label was added to the 5′-end of the positive-stranded sRNA. The related 21-nucleotide DNA primers, including the positive-stranded DNA oligo (5′-CGTACGCGGAACTTTCGATT-3′) and negative-stranded DNA oligo (5′-TTGCACTGGCCTTATGAGCT-3′), were synthesized and used as controls for the p19 binding assay.

**sRNA analysis and miRNA prediction.** We sequenced the sRNAs either isolated from *N. benthamiana* or eluted from the p19/SiNW-FET by a next generation sequencer (NGS), illumina HiSeq 2000 system (Illumina, San Diego, CA, USA). We first removed the sequence errors in the reads by filtering through the *N. benthamiana* genome version 0.44 contigs (Niben_genome.v0.4.4.contig.fasta)69; the clean reads were used to determine the read counts (weight) and the identity of the sRNA species.

**References**

1. Hannon, G. J. RNA interference. *Nature* **418**, 244–251 (2002).
2. Tomari, Y. & Zamore, P. D. Perspective: machines for RNAi. *Gene Dev.* **19**, 517–529 (2005).
3. Bartel, D. P. MicroRNAs: Genomics, biogenesis, mechanism, and function. *Cell* **116**, 281–297 (2004).
4. Wu, H.-W., Lin, S.-S., Chen, K.-C., Yeh, S.-D. & Chua, N.-H. Discriminating mutations of HC-Pro of zucchini yellow mosaic virus with differential effects on small RNA pathways involved in viral pathogenicity and symptom development. *Mol. Plant-Microbe Interact.* **23**, 17–28 (2010).
5. Wu, R.-M., Wood, M., Thrush, A., Walton, E. F. & Varkonyi-Gasic, E. Real-Time PCR quantification of plant miRNAs using Universal ProbeLibrary Technology. *Biochimica et Biophysica Acta* **12**, 12–15 (2007).
6. Danielson, D. C. & Pezacki, J. P. Studying the RNA silencing suppressor with the p19 protein. *FEBS Lett.* **587**, 1198–1205 (2013).
7. Campuzano, S. et al. Magnetobiosensors based on viral protein p19 for miRNA determination in cancer cells and tissues. *Angew. Chem. Int. Ed. Engl.* **53**, 6168–6171 (2014).
8. Jin, J. & McReynolds, L. A., Gullerova, M. p19-mediated enrichment and detection of siRNAs. *Methods Mol. Biol.* **1173**, 99–111 (2014).
9. Chen, K.-I., Li, B.-R. & Chen, Y.-T. Silicon nanowire field-effect transistor-based biosensors for biomedical diagnosis and cellular recording investigation. *Nano Today* **6**, 131–154 (2011).
10. Cui, Y., Wei, Q., Park, H. & Lieber, C. M. Nanowire biosensors for highly sensitive and selective detection of biological and chemical species. *Science* **293**, 1289–1292 (2001).
11. Li, B.-R., Chen, C.-C., Kumar, U. R. & Chen, Y.-T. Advances in nanowire transistors for biological analysis and cellular investigation. *Analyst* **139**, 1589–1608 (2014).
12. Noor, M. O. & Krull, U. J. Silicon nanowires as field-effect transducers for biosensor development: a review. *Anal. Chim. Acta* **825**, 1–25 (2014).
13. Stern, E. et al. Label-free immunodetection with CMOS-compatible semiconducting nanowires. *Nature* **445**, 519–522 (2007).
14. Gao, A. et al. Silicon-nanowire-based CMOS-compatible field-effect transistor biosensors for ultrasensitive electrical detection of nucleic acids. *Nano Lett.* **11**, 3974–3978 (2011).
15. Zhang, G.-J., Chua, J. H., Chee, R.-E., Agarwal, A. & Wong, S. M. Label-free direct detection of MiRNAs with silicon nanowire biosensors. *Biosens. Bioelectron.* **24**, 2504–2508 (2009).
16. Chiang, P.-L. et al. Nanowire-based ultrasensitive virus detection with reversible surface functionalization. *Chem. Asian J.* **7**, 2073–2079 (2012).
17. Lin, S.-F. et al. A reversible surface functionalized nanowire transistor to study protein-protein interactions. *Nano Today* **4**, 235–243 (2009).
18. Lin, T.-W. et al. Label-free detection of protein-protein interactions using a calmodulin-modified nanowire transistor. *Proc. Natl. Acad. Sci. USA* **107**, 1047–1052 (2010).
19. Li, Y., Li, C., Ding, G. & Jin, Y. Evolution of MIR193/319 microRNA genes and their post-transcriptional regulatory link to siRNA pathways. *BMC Evol. Biol.* **11**, 122 (2011).
20. Tan, L. X. et al. Knockdown of miR-21 in human breast cancer cell lines inhibits proliferation, in vitro migration and in vivo tumor growth. *Breast Cancer Res.* **13**, R12 (2011).
21. Ribas, J. et al. A novel source for miR-21 expression through the alternative polyadenylation of VMP1 gene transcripts. *Nucleic Acids Res.* **40**, 6821–6833 (2012).
22. Fix, L. N., Shah, M., Efferth, T., Farwell, M. A. & Zhang, B. MicroRNA expression profile of MCF-7 human breast cancer cells and the effect of green tea polyphenon-60. *Cancer Genomics Proteomics* **9**, 261–277 (2012).
23. Lu, N. et al. CMOS-compatible silicon nanowire field-effect transistors for ultrasensitive and label-free microRNAs sensing. *Small* **10**, 2022–2028 (2014).
24. Lin, T.-Y. et al. Improved silicon nanowire field-effect transistors for fast protein-protein interaction screening. *Lab Chip* **13**, 676–684 (2013).
25. Nishida, M., Harada, S., Noguchi, S., Satow, Y., Inoue, H. & Takahashi, K. Three-dimensional structure of Escherichia coli glutathione S-transferase complexed with glutathione sulfonate: Catalytic roles of Cys10 and His106. *J. Mol. Biol.* **281**, 135–147 (1998).
26. Vargason, J. M., Szititya, G., Burgyan, J. & Hall, T. M. Size selective recognition of siRNA by an RNA silencing suppressor. *Cell* **115**, 799–811 (2003).
27. Lin, M.-C. et al. Control and detection of organosilane polarization on nanowire field-effect transistors. *Nano Lett.* **7**, 3656–3661 (2007).
28. Chu, C.-J. et al. Improving nanowire sensing capability by electrical field alignment of surface probing molecules. *Nano Lett.* **13**, 2564–2569 (2013).
29. Ye K., Malmina, L. & Patel, D. J. Recognition of small interfering RNA by a viral suppressor of RNA silencing. *Nature* **426**, 874–878 (2003).
30. Li, B.-R., Chen, C.-W., Yang, W.-L., Lin, T.-Y., Pan, C.-Y. & Chen, Y.-T. Biomolecular recognition with a sensitivity-enhanced nanowire transistor biosensor. *Biosens. Bioelectron.* **45**, 252–259 (2013).
31. Li, B.-R., Hsieh, Y.-J., Chen, Y.-X., Chang, Y.-T., Pan, C.-Y. & Chen, Y.-T. An ultrasensitive nanowire-transistor biosensor for detecting dopamine release from living PC12 cells under hypoxic stimulation. *J. Am. Chem. Soc.* **135**, 16034–16037 (2013).
32. Lakatos, L., Szititya, G., Sillhavy, D. & Burgyan, J. Molecular mechanism of RNA silencing suppression mediated by p19 protein of tombusviruses. *EMBO J.* **23**, 876–884 (2004).
33. Cheng, J., Sagan, S. M., Jakubek, Z. J. & Pezacki, J. P. Studies of the interaction of the viral suppressor of RNA silencing protein p19 with small RNAs using fluorescence polarization. *Biochemistry* 47, 8130–8138 (2008).
34. Sagan, S. M., Koukiekolo, R., Rodgers, E., Goto, N. K. & Pezacki, J. P. Inhibition of siRNA binding to a p19 viral suppressor of RNA silencing by cysteine alkylation. *Angew Chem. Int. Ed. Engl.* 46, 2005–2009 (2007).
35. Krol, J. & Krzyzosiak, W. J. Structure analysis of microRNA precursors. *Methods Mol. Biol.* 342, 19–32 (2006).
36. Zhang, G. J. & Ning, Y. Silicon nanowire biosensor and its applications in disease diagnostics: a review. *Anal. Chim. Acta* 749, 1–15 (2012).
37. Chang K. S. *et al.* Monitoring extracellular K+ flux with a valinomycin-coated silicon nanowire field-effect transistor. *Biosens. Bioelectron.* 31, 137–143 (2012).
38. Omarov, R. T., Ciomperlik, J. J. & Scholthof, H. B. RNAi-associated ssRNA-specific ribonucleases in Tombusvirus P19 mutant-infected plants and evidence for a discrete siRNA-containing effector complex. *Proc. Natl. Acad. Sci. USA* 104, 1714–1719 (2007).
39. Bombarely, A., Rosli, H. G., Vrebalov, J., Moffett, P., Mueller, L. A. & Martin, G. B. A draft genome sequence of Nicotiana benthamiana to enhance molecular plant-microbe biology research. *Mol. Plant-Microbe Interact.* 25, 1523–1530 (2012).

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**Author Contributions**

K.I.C. carried out the p19 experiment, experimental data analysis, and manuscript writing. C.Y.P. finalized the manuscript writing. K.H.L. and Y.C.H. performed the p19 experiment and data analysis. C.W.L. and C.Y.T. did the bioinformatic analysis and miRNA prediction. Y.W.S. fabricated SiNW-FET devices. L.W.T. did the miRNA experiments included in Fig. 1. K.C.T. and C.Y.L. were involved in the p19 experiment. C.D.C. was in charge of the SiNW-FET devices fabrication and experimental design. S.S.L. and Y.T.C. are the corresponding authors in charge of the project design, experimental data analysis, and manuscript writing.

**Additional Information**

**Accession codes:** The raw sRNA reads reported in this paper are available in the NCBI Short Read Archive under accession numbers SRR1555764 (input sample), and SRR1555765 (eluted sample).

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