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Tobacco mosaic virus based thin film sensor for detection of volatile organic compounds

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A thin film sensor for the detection of volatile organic compounds (VOC) was fabricated by deposition of oligo-aniline grafted tobacco mosaic virus (TMV) onto a glass substrate. The oligo-aniline motifs were conjugated onto the TMV surface by a traditional diazonium coupling reaction to tyrosine residues followed by Cu(I) catalyzed alkyne-azide cycloaddition (CuAAC) reaction. The modified TMV was easily fabricated into a thin film by directly drop coating onto a glass substrate. Upon integration of the glass substrate into a prototypical device, the virus-based thin film exhibited good sensitivity and selectivity toward ethanol and methanol vapour.

Introduction

Biological nanoparticles (BNPs), such as viruses, ferritins, and other protein complexes, represent a class of structurally well-defined nanoscale assemblies. The sizes of BNPs range from 10 nm to a few hundreds of nanometers with different morphologies. Their three dimensional structures can be characterized at near atomic resolution, which allow genetic manipulation and chemospecific bioconjugation with near atomic precision, providing highly attractive opportunities for the development of nanoscale materials with well-controlled structural features and surface properties.1–13. In these respects, BNPs have a great advantage over synthetic nanoparticles such as carbon nanotubes, quantum dots, dendrimers and polymer vesicles.14–22. Tobacco mosaic virus (TMV) is among the most well-known plant viruses.20,21 Thousands of copies of identical coat proteins of TMV are assembled into a 300 nm helical rod encapsulating its genomic RNA, leaving an interior core with a diameter of 4 nm and an external diameter of 18 nm.22 TMV has been employed as a building block or template to produce materials with applications in light harvesting, electronics, energy storage, and cell culturing.23–27

On the other hand, conducting polymers have found broad applications in electronics, imaging and sensing.28–30 In particular, polyaniline (PANI) has received considerable attention due to its environmental stability, ease of synthesis and its well-known physical properties.31–33 It has been shown that the morphology of PANI is very important to its sensing ability.33–36 Previously, we reported the synthesis of PANI-TMV composite nanowires upon coating TMV surface with PANI via non-covalent interactions.24–27 Although the synthesis was straightforward, the processing of such nanowires was impeded by the formation of bundle-like structures, presumably due to the strong interactions between the surface PANI coatings, which limited its applications. Oligoaniline (OANI) is a small molecule precursor of its parent polymer, PANI, which is electroactive, soluble in aqueous solutions, and can be tailored to include any desired functionality.37 In particular, OANI has shown similar properties to PANI in directing neural cell growth38 and in solution as a TNT sensor.39

In our study, we would like to show the potential application of OANI-TMV conjugates as sensor for volatile organic compounds (VOC), in particular, methanol and ethanol. The extended exposure to methanol and ethanol vapours leads to diseases such as eyesight disturbance, nasal mucous membrane, nerve disease and even death.40,41 Thus the on-line monitoring of these alcohol vapours is very important in many workplaces and laboratories. Recent reports have suggested that BNPs have found applications in chemical and biological sensing.42 In this paper, we show that OANI can be directly conjugated onto the surface of TMV through multiple chemical manipulations. Due to the high-density distribution of OANI on the TMV surface and the good processability of OANI modified TMVs, they can be easily fabricated into a thin film and exhibit good sensitivity and high selectivity toward the ethanol and methanol vapours.

Experimental

Materials

All reagents were used as received. The virus was prepared as previously described.26 Unless otherwise noted, “buffer” refers to 10 mM potassium phosphate buffer, pH 7.8. Sucrose gradient ultracentrifugation separation of virus samples was performed on a 20 mL gradient (made of 40% (w/w) sucrose solution in 10 mM potassium phosphate buffer, frozen at –20 °C and thawed before use) with centrifugation at 90,000 × g for 2 h with a Beckman SW41 rotor using a Beckman Optima™ L90 K ultracentrifuge. The concentration of unmodified virus was measured by absorbance at 260 nm; 0.1 mg/mL of TMV gives a standard absorbance of 0.3.43 Modified virus concentrations were measured using a Modified...
Lowry Protein Assay Kit (Pierce). The molecular weight of a single subunit of wild type TMV coat protein is 17,534 Daltons.

Synthesis of alkyne modified TMV (Alkyne-TMV). The diazonium salt was prepared by mixing the following solutions at 4 °C for 1 h: 400 μL of 0.3 M aqueous p-toluenesulfonic acid monohydrate; 25 μL of 3.0 M aqueous sodium nitrite; and 75 μL of 0.68 M distilled 3-ethylaniline dissolved in acetonitrile. Subsequently, a stock solution of TMV (20 mg/mL, 1.25 mL) was diluted with borate buffer (3.3 mL, pH 8.8, containing 100 mM NaCl). Diazonium salt solution (450 μL) was then added to the mixture. The mixture was place in an ice bath for 3 h while the solution turned a light brown color. Purification of the final product was completed by passing the reaction through a 40% (w/w) sucrose cushion at 160,000 g for 2.5 h. The pellet was redissolved in buffer for future use.

Synthesis of azide modified TMV (Azide-TMV). The bis-1,4-azidobenzene (100 mM in DMSO, 160 μL) and a solution of Alkyne-TMV (15 mg/mL, 200 μL) were mixed with Tris buffer (10 mM, 580 μL, pH 8.0) and DMSO (40 μL). Then solutions of CuSO4 (100 mM, 10 μL) and NaAsc (200 mM, 10 μL) were added and the mixture was incubated at room temperature for 1 h. The reaction mixture was purified via 10–50% sucrose gradient from which the light scattering region was collected. The modified virus was then pelleted using ultracentrifugation at 160,000 × g for 2.5 h. The pellet was dissolved in buffer.

Synthesis of oligoaniline modified TMV (OANI-TMV). The OANIP (100 mM in DMSO, 5.5 μL) and a solution of Azide-TMV (15 mg/mL, 133 μL) were mixed in Tris buffer (10 mM, 547 μL, pH 8.0) and DMSO (294.5 μL). Then solutions of CuSO4 (100 mM, 10 μL) and NaAsc (200 mM, 10 μL) were added and the mixture was incubated at room temperature for 1 h. The reaction mixture was purified via dialysis (100,000 MWCO dialysis tubing) against nanopure H2O. The remaining solution concentration was determined using a Modified Lowry Assay (Pierce).

General protocol for preparation of virus solution and sensing plate. Prior to use, the modified virus was dialyzed against nanopure water using a 100,000 MWCO dialysis tubing (Pierce). A virus solution with concentration of 250 μg/mL was prepared. To that solution, 15 μL p-toluene sulfonic acid (0.1 M, dopant) was added to reduce the pH to a value between 3 and 4. To prepare a sensing plate, 150 μL of the viral solution was spread on a glass slide with gold coating at the edge of the surface as two electrodes (25 mm × 1.5 mm), then allowed to dry in a hood overnight.

General protocol for VOC sensing. After placing the leads on gold electrode parts, the apparatus was assembled as shown in Fig. 3b. There is an inlet and outlet for nitrogen or other gases to flow. The two electrodes are designed to hold the OANI-TMV sensor in place while measuring the current flow through the film. They protrude through the bottom of the container and are hooked up to a Keithley 6487 picoammeter which is used for the current measurements (Labeled A in Fig. 3a). The OANI-TMV sample was evaporated at room temperature onto a custom cut glass slide with dimensions of 1.5 mm × 25 mm. The glass slide was sputter coated with two 40 nm thick strips of gold (Fig. 3c) prior to sample evaporation. These strips of gold were the intermediate contact between the electrodes and the film. Without the gold strips, the sample was easily scraped off of the glass slide by the electrodes. The edges were sealed with additional parafilm to deter moisture from entering the container. N2 was used to flush the container of unwanted chemicals and is used as a blanking standard. After 30 to 60 s of N2 gas, the gas inlet and outlet are sealed with caps and the VOC of choice was injected (50–75 μL) with a syringe. From that point, the sensor was untouched for 15 min while collecting data. After this, N2 was once again used to flush out the container of chemicals.

Characterization
For MALDI-MS analysis, the virus was denatured by adding guanidine hydrochloride (6 M, 6 μL) to the sample (24 μL) and mixing for 5 min at room temperature. Denatured proteins were spotted on MTP 384 massive target plate using Millipore ZipTips®/C18® tips to remove excess salts and assist the binding of protein to the sinapic acid matrix. MALDI-MS analysis was performed using a Bruker Ultra-Flex I TOF/TOF mass spectrometer. For transmission electron microscopy (TEM) analysis, a 20 μL sample solution (0.2 mg/mL) was deposited onto a 300-mesh carbon-coated copper grid for 2 min. The grid was then stained with 20 μL of 2% uranyl acetate for 2 min and was characterized with a Hitachi H-8000 TEM. Tapping mode atomic force microscopy (AFM) images were obtained at ambient conditions using a NanoScope IIIA MultiMode AFM (Veeco). Si tips with a resonance frequency of approximately 300 kHz and a spring constant of about 40 N·m⁻¹, were used for imaging with a scan rate of 0.5 Hz were used. UV-vis absorption studies were performed using an Agilent 8453 UV-vis spectrometer.

Results and discussion
As shown in Scheme 1, TMV was subjected to a three-step bioconjugation protocol to introduce the desired OANI functionality. The initial reaction targeted the phenol side chains of

Scheme 1 Synthesis of OANI modified TMV for VOC sensing.
tyrosine residues of TMV coat proteins to insert a terminal alkyne.\textsuperscript{1,27,44} This was achieved using a diazonium salt generated \textit{in situ} from 3-ethynylaniline then mixed with TMV to form the corresponding alkyne labeled TMV (Alkyne-TMV). MALDI-TOF MS analysis indicated that >95% of the capsid monomers were converted to the alkyne derivatives as seen in Fig. 1 by the disappearance of a peak at 17534 \textit{m/z} and the introduction of a peak at 17664 \textit{m/z}.

The Cu(I) catalyzed azide-alkyne 1,3-dipolar cycloaddition (CuAAC) between Alkyne-TMV and bis-1,4-azidobenzene in the presence of \textit{Cu}\textsuperscript{I}, which was formed \textit{in situ} by the reduction of CuSO\textsubscript{4} with sodium ascorbate, proceeded with >95% efficiency to form the azido functionalized TMV (Azide-TMV). These two steps are necessary to introduce the functional azido group onto the surface of TMV. The final step of conjugation was completed following the same CuAAC reaction protocol with OANI and Azide-TMV as starting materials, which afforded OANI-TMV efficiently as shown in Fig. 1.

It is important to know whether TMV remained intact after a three-step bioconjugation. TEM was used to characterize the morphology and aggregation of TMV before and after modifications. As shown in Fig. 2b, after the second step modification (Azide-TMV), TMVs still keep a similar morphology as Wt-TMV (Fig. 2a). Upon conjugation with OANI units, OANI-TMV still maintains its rod-like structure (Fig. 2c). However, a large amount of aggregation is observed under TEM analysis (Fig. 2c). The aggregation of OANI-TMV is possibly caused by pi-pi interactions and/or the hydrophobic interactions of the OANI motif on the TMV surface between viral particles. To break apart the aggregation of OANI-TMV, DMSO was added into the aqueous solution as the cosolvent to decrease the interactions between OANI units. As visualized by TEM (Fig. 2d), the particles were well-dispersed when the level of DMSO reached 70%, while most of the OANI-TMVs remained intact.

Detection of VOCs with OANI-TMV was pursued based on results from previous studies conducted utilizing polyaniline as the sensing material.\textsuperscript{33,36,45,46} A simple sensing device with sample container was designed (Fig. 3a) and constructed as shown in Fig. 3b. The sample container has a volume of roughly 12 mL. There is an inlet and outlet for nitrogen or other gases to flow. The two electrodes are designed to hold the OANI-TMV sensor in place while measuring the current flow through the film.

Previous investigations utilizing conducting polymers for sensors indicated that the selection of an appropriate dopant is critical for sensor selectivity and sensitivity.\textsuperscript{28,33,45–47} Hence in our study the OANI-TMV was doped with 0.1 M \textit{p}-toluene sulfonic acid prior to evaporation. A general procedure was followed for

![Fig. 1](image1.png)

**Fig. 1** MALDI-TOF MS of the coat protein of Wt-TMV and modified TMV. The values in parenthesis are the expected \textit{m/z}.

![Fig. 2](image2.png)

**Fig. 2** TEM images of (a) Wt-TMV, (b) Azide-TMV, (c) OANI-TMV, and (d) OANI-TMV in 70% DMSO. Scale bar is 200 nm for all images.

![Fig. 3](image3.png)

**Fig. 3** (A) Schematic illustration and (B) image of the sample container used for gas sensing. (C) AFM image confirming that the height of the gold sputtered film is about 40 nm. (D) Height profile across the edge of the film.
sensing experiments; 50 µL of organic solution was injected into the sample holder located in the middle of the sample container. The container was then sealed with parafilm and flushed with nitrogen for one minute to establish a baseline. This was also performed to remove any other contaminants from the atmosphere, such as humidity. The sealed container was then allowed to incubate for 15 min while the atmosphere was filled with the analyte of interest. The concentrations of the analytes were determined by their respective partial pressure at room temperature. Following the incubation time, nitrogen gas was used to flush the container of analyte and the gas desorption was seen as a sharp decrease in the current.

The plots shown in Fig. 4 display the y-values as a Response Current (RC). The RC is the ratio between the measured current (Iₓ) and the maximum current obtained in the presence of only nitrogen (Iₒ). This type of correction is required because there is variability in the film that leads to variability in sensitivity, but not selectivity. Films that cover the majority of the 25 mm × 1.5 mm active area (gap between the gold sputtered films) yield a higher current reading for the same film that may only cover part of the active area.

The following analytes were tested and their RC was plotted as a function of time in Fig. 4a: ethanol, methanol, isobutyl alcohol, acetonitrile, acetone, tetrahydrofuran and toluene. During this period the current of the viral film was affected differently for each analyte. The OANI-TMV shows great selectivity towards ethanol and methanol over the other analytes. Control experiments were performed by using wt-TMV and OANI alkyne thin films. WT-TMV and OANI thin films were prepared by direct deposition of TMV solution and OANI solution onto glass substrates. As shown in Fig. 4b, the RC for WT-TMV and OANI alkyne were significantly lower than OANI-TMV. This indicates that the spatial attribution of OANI ligand on the TMV surface is very critical to the response towards methanol. Two other distinct properties of this particular sensor are the quick response ability for the ethanol/methanol absorption and desorption and multi-time reproducibility (Fig. 4c).

A possible mechanism for the electrical response of OANI-TMV over unmodified TMV is the flexoelectric properties of TMV. It has been shown that when strain is placed on the rod-like particle, a small electrical response can be detected. This implies that if there is a long connected series of viral particles, combined with an outside pressure, then electrons should be able to flow through the viral film. The OANI units cause the viral particle to aggregate and form an interconnected film. The physical absorption of methanol/ethanol causes the film to swell and exerts the necessary external force to cause the desired response. Finally, the electroactive OANI assists in electron transfer through the film.

Conclusions

Using a three step bioconjugation of TMV with an electron rich OANI linker, we have shown that this scaffold provides an excellent VOC sensor. The sensor that has been developed demonstrate high selectivity towards methanol and ethanol over other VOCs. This particular property will be very useful for developing a sensor array for the analysis of real air samples with a multitude of VOCs present. Because of the versatility of the viral scaffold, it is likely that various other ligands with differing chemical properties can be attached to develop sensors with different selectivities.

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Notes and references


