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Graphene-based field-effect transistor biosensors functionalized using gas-phase synthesized gold nanoparticles

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ABSTRACT
Research has focused on graphene for developing the next generation of label-free biosensors, capable of highly sensitive and specific detection of DNA or other biomolecules. The binding of charged analytes to the one-atom thick layer of graphene can greatly affect its electronic properties. However, graphene is highly chemically inert, thus surface functionalization through chemical treatment is typically necessary to immobilize receptors of the target biological analyte on the graphene. In this work, we use gas-phase synthesized gold nanoparticles (Au NPs) to functionalize and bind a DNA aptamer to the graphene surface. The graphene is employed in a liquid gated field-effect transistor (FET) configuration to detect the hybridization of the complementary DNA strand, as well as the protein streptavidin, at attomolar level (aM, $10^{-18}$ mol L$^{-1}$). The sensor shows a high dynamic detecting range from aM to picomolar (pM) levels ($10^{-12}$ to $10^{-15}$ mol L$^{-1}$), can discriminate between a complementary strand and a single nucleotide polymorphism (SNP) containing strand, and achieves a detection limit as low as 15 aM. The high detection limit suggests that decorating biosensors with Au NPs synthesized from magnetron sputtering inert gas condensing technique is a promising method for biosensor functionalization, particularly for larger-area sensors that employ two-dimensional materials such as graphene.

1. Introduction

Field-effect transistors (FETs) have attracted widespread attention for their biosensing applications; particularly for DNA detection, as this capability is of fundamental interest in clinical diagnostics, environmental monitoring, forensics, and biomedical research [1–5]. DNA can also be developed into aptamers designed to bind to specific biomolecules for early disease detection [6,7]. Unlike traditional optical DNA sensing methods, FET based sensors do not require sophisticated fluorometric or extensive fluorescent labeling processes. This is because of FETs are solid state devices in which the channel conductance between the source and drain electrodes is controlled via the electrostatic gating effect from a third electrode [8]. In a bio-sensor FET, charged molecules may dope or apply a virtual gate bias upon binding to the semiconducting channel [4,9–11]. By functionalizing the FET with a specific biological recognition element, the binding of a target analyte produces a change in the channel conductance and a sensor response which is highly specific.

Graphene is an attractive nanomaterial in biosensors due to its excellent physical and chemical properties such as high carrier mobility and ease of functionalization [5]. Its one-atom thick nature, large surface area to volume ratio, and high conductivity allow graphene-based FETs (G-FETs) to be highly sensitive, with limits of detection (LoD) for nucleic acids in the high aM to low femtomolar range [7,11,12]. G-FETs have been employed as sensors for a wide variety of biological analytes, including small-molecule biomarkers, amino acids, enzymes, glucose, and nucleic acids [3,11,13–15]. In addition, two dimensional materials such as graphene are more compatible with standard planar technology and microfabrication at the wafer scale to produce arrays of multiple biosensors [16,17].

The selectivity of FET biosensors is typically defined by immobilizing receptor molecules on the device surface to uniquely bind to a targeted analyte, such as an antibody or a complementary DNA strand. As graphene is relatively inert, it must be chemically treated to generate active groups for the binding of biomolecules. Noble metal nanoparticles (NPs) deposited on graphene are widely used as binding sites, and have been shown to greatly enhance DNA detection sensitivity [11,15,18–20]. Gold NPs have been especially popular due to

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their biocompatibility and well-known functionalization chemistry. Au has a high chemical affinity with thiol groups, enabling easy binding of thiol-terminated DNA to act as receptor molecules [10].

In this work, we successfully demonstrate the integration of Au NPs produced via a magnetron sputtering inert gas condensing technique in G-FET arrays for DNA sensing. This is the first demonstration of such NPs decorated G-FETs for bio-sensing applications and the method avoids the use of residual capping ligands or reactants present in solvent based methods that could interfere with the Au-thiol binding [21,22]. Gas-phase synthesis also allows for a high degree of control over the Au NP size and shape while ensuring a uniform coverage over a large area during deposition [23–25]. Combined with a scalable lamination procedure to transfer graphene grown using chemical vapor deposition (CVD) to Si:SiO2 substrates [16], this method can efficiently produce highly specific sensors with sensitivity down to the aM level at a large scale. A DNA aptamer is used to detect both a target DNA strand and streptavidin, a protein commonly used in biomedical research due to its high affinity for biotin. High selectivity is demonstrated by comparing the sensor response between fully complementary strands and those with a single nucleotide polymorphism (SNP).

2. Materials and methods

2.1. Graphene FET fabrication

Graphene was synthesized in a CVD system (Easytube 2000, FirstNano, USA) on a 30 μm thick commercial copper foil (Nilaco, Japan). The copper foil was sonicated in acetone and isopropyl alcohol (IPA) before CVD growth. The copper foil was first annealed for 1 h at 990 °C at 760 torr, with gas mixture of Ar and H2 at flow rates of 4875 and 125 standard cubic centimeter per minute (sccm), respectively. The CVD growth step was then performed for 15 min at 1000 °C at 760 torr, with gas mixture of Ar, H2, and CH4 with flow rates of 4875, 125, and 2.5 sccm, respectively. The sample was subsequently cooled to room temperature under an Ar flow rate of 5000 sccm.

Graphene transfer from the copper foil to Si:SiO2 substrates was performed using a Poly(vinyl alcohol) (PVA) lamination method [16]. PVA films (Cubic Coating) were initially rinsed in IPA and dried with N2. Copper foil pieces with CVD grown graphene were placed in deionized (DI) water for 8 h to facilitate the intercalation of graphene from the Cu or Cu2O surface [26]. After drying, the PVA film is laminated on top of the graphene covered copper foil at 110 °C and a speed of 15 mm/s using a commercial laminator (Meiko Shokai THS 330). The PVA/graphene adhesion. Once the PVA film is peeled off the copper foil, it removes the graphene as well. The graphene on PVA can be transferred to the desired substrate using a second lamination step under the same conditions. Then the substrate is baked at 110 °C for 1 min, followed by peeling off the paper support of the PVA film while the sample is still on the hot plate. The PVA layer is removed by placing the substrate in room temperature DI water overnight, leaving a monolayer graphene layer on top of the target substrate.

Si:SiO2 substrates covered with monolayer graphene were patterned into biosensor devices using photolithography and oxygen plasma etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching. The Ti/Au contacts (5/50 nm) were deposited by e-beam evaporation and de-etching.

A PDMS solution well was attached to the G-FET to protect the source and drain electrodes during liquid-gate measurements. Liquid PDMS was placed on the underside of a hollow PDMS cylinder, then cured at 90 °C for 1 h for good adhesion to the FET surface. The solution well enclosed 3 G-FETs, each with an area of 1 mm2. A schematic of a completed enclosed Au NP decorated G-FET is shown in Fig. 1.

2.2. Gold nanoparticles decoration

Before attaching the PDMS wells, G-FET devices were decorated with Au NPs using a direct-current (DC) magnetron-sputtering inert-gas aggregation system (Fig. 2a [23,24,27–29]). A high purity (99 %) Au target was bombarded with an Ar plasma, freeing Au atoms for nanocluster formation in an initial chamber (aggregation zone). In this chamber, Au atoms nucleate into small clusters via collisions with Ar atoms and other Au atoms. Adjusting deposition parameters such as the plasma power and the pressure difference between the aggregation zone and deposition chamber, allowed us to control the nanoparticle size and crystallinity as they form from coalescing Au clusters. The aggregation zone is held at a higher pressure than the deposition chamber, forcing Au NPs to migrate into the deposition chamber and land on the target substrate. The particles also passed through a quadrupole mass filter (QMF) between the two chambers for size control. We selected Au NPs with an average diameter of 2.5 nm. The base pressure in the deposition chamber and aggregation zone before deposition was maintained below ~1.5 × 10–7 and ~2.5 × 10–6 mbar, whereas the process pressures were ~1.4 × 10–3 and ~4.5 × 10–1 mbar, respectively. Depositions were performed using an Ar flow rate of 100 sccm, a DC power of ~6 W and the length of the aggregation zone was 125 mm. The substrate holder was rotated at 2 rpm to ensure uniform NP decorating density. An AFM image of the Au NPs decorated graphene is shown in Fig. 2b After 1 h of deposition, the NP density is approximately 5 × 1010 cm–2, therefore each graphene device will present 5 × 108 binding sites to the biological solution.

2.3. Biological solution preparation

The dimeric form of the streptavidin-binding aptamer was derived from the monomeric form (StrepApt5) reported by Ruigrok VJ et al. [30]

Aptamer: 5′-thiol-modified/ AAAGGAAAAACCGATCGAGGT TT CCCATAAACAGCCGATCGAGGTATTG TCATCGTGTTCTTT-3′ (60 nt)

Complementary strand (cDNA): 5′- TTTATGGGAAACCTGCGATCG GTGCGTGTTCTTT-3′ (34 nt)

SNP strand M1 (cDNA): 5′- TTTATGGGAAACCTGCGATCGGTGCC GTTCCCGTT3′ (34 nt)

SNP strand M4 (cDNA): 5′- TTTATGGGAAACCTGCGATCGGTGCC GTTCCCGTT3′ (34 nt)

The disulfide in a thiol-modified aptamer solution was reduced to monothiol using tris (2-carboxyethyl) phosphine (TCEP 20 mM, 2 h at RT). A solution containing 1X TBE buffer (89 mM Tris, 89 mM boric acid, 2 mM EDTA, pH 8.0), 1 μM TCEP, and 50 mM NaCl was used for cleaving the ss bond of the 10 nM aptamer. This solution was kept at ~20 °C. The following oligonucleotides were used for preparing 2000 bp DNA with 5′-thiol modification:

PCR primer 1: 5′-thiol-modified/GTCTCGCGCGTTTCGGTGAT-3′ (20 nt)

PCR primer 2: GAACCGAGCTGAATGAAGCC-3′ (21 nt)

The 2000 bp DNA with 5′-thiol modification was prepared by PCR.
amplification of the pUC19 plasmid. The plasmid (10 ng) plus primers 1 and 2 (0.5 μM each) were PCR amplified using 2 × Phusion Master Mix (New England Biolabs) in 25 μL reaction volume. After initial denaturation at 98 °C for 30 s, 25 cycles of 5 s at 98 °C followed by 45 s at 72 °C and 30 s at 70 °C were used. The resulting PCR product was column purified using a DNA Clean and Concentrator Kit (Zymo Research). A solution containing 3 nM of thiol-modified duplex DNA was subjected to TCEP treatment as mentioned previously. Streptavidin (10 nM) solution was prepared in 1x phosphate-buffered saline (PBS) buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, pH 7.4).

2.4. Electrical measurements & characterization

Atomic force microscopy (AFM) was used to measure the Au NPs density on graphene devices. AFM topography measurements were performed on a conventional Multimode 8 scanning probe microscope (Bruker, USA) in PeakForce tapping mode. The high-resolution AFM probe (ScanAsyst-Air) from Bruker with nominal tip radius ~2 nm, resonant frequency 70 kHz, and low spring constant 0.4 N/m were used for all AFM measurements. The high resolution (512 by 512 pixels) AFM images were captured at a scan rate of 0.5 Hz and further processed by using the Nanoscope Analysis software (Ver 9).

The transfer characteristics of the liquid-gated G-FETs were measured using a semiconductor parameter analyzer (Keithley 4200-SCS-A) under a constant drain to source (Vds) bias of 0.1 V. A silver wire (1 mm diameter) was used as the liquid gate electrode [10]. The DNA aptamer was bound to the Au NPs on the G-FET by placing a 40 μL droplet of 10 μM DNA in buffer in the solution well overnight in a wet chamber at 4 °C. The device was rinsed with 1x PBS and DI water to remove unbound DNA strands. Before cDNA detection experiments, the graphene surface was passivated by incubating a 100 μM solution of ethanolamine in 0.01X PBS for 1 h at room temperature [11]. Before streptavidin detection experiments, BSA was used as a passivation layer instead by incubating an extra 2 μM solution in DI water for 1 h. The schematic in Fig. 1 shows the reference electrode configuration and DNA bound to the Au NPs present during measurement.

3. Results

Graphene was grown on Cu foil using a CVD process and transferred to Si-SiO2 substrates via a PVA lamination procedure described in Shivayogimath A et al. [16] This is a convenient, non-toxic, and scalable procedure that avoids etching or electrochemical delamination techniques. A typical microscope image of the resulting transferred graphene is shown in Fig. 3a, with small regions of darker purple color distributed over a lighter colored background. The darker zones, which suggest areas of thicker graphene [31], are arranged in a line style normal to the direction of laminate peeling. The linear arrangement is possible due to the stress accumulation during the peeling process, which indicates that the resulting device graphene thickness does not only depend on the CVD growth parameters, but also on the peeling procedure. Raman spectra acquired from these samples show two trends, seen in Fig. 3b. The top spectrum (black) was measured with the laser focus on the dark purple zone, while the bottom spectrum (red) was from the light background. In both spectra, four peaks were detected: D, G, D + D’, and 2D modes. The G (ca. 1590 cm−1) and 2D (ca. 2680 cm−1) peaks are the typical Raman modes of graphene, from the first order in-plane vibration modes and second order in-plane transverse optical phonons, respectively. The D (1350 cm−1) and D + D’ (2450 cm−1) peaks are mostly due to grain boundaries, surface contaminations, or other defects and strains formed during the transfer process. Both peaks are quite low and comparable to previous reports; indicating a low defect density in the transferred graphene [32–34]. It is well known that the 2D/G height ratio and the 2D peak symmetry can be used for estimating the thickness of graphene: a ratio of 2 for monolayer and 1 for bilayer graphene [35]. 2D/G peak height ratios were calculated to be 1.42 and 1.9 for the dark and light-colored zones, respectively. Deviation of the 2D/G ratio from the ideal value of 2.0 is due to the relatively large Raman laser spots, so that signals from both zones would inevitably be collected. The same theory also explains the value of 1.42 for the darker zones, as the height ratio should be ca. 1.0 for bilayer graphene. A Raman mapping image illustrating the 2D/G height ratio distribution is shown in Fig. 3c, in which regions of low ratio (blue) interspersed with regions of high 2G/D (red). This observation from Raman mapping agrees with the optical microscope measurements that the laminate transferred graphene consists of an overall monolayer with isolated islands of bilayer graphene. Further optimization of the lamination or peeling procedure could suppress the formation of these regions, but large enough areas of monolayer graphene are present for ultrasensitive DNA detection via gas-phase synthesized Au NPs.

After patterning the monolayer graphene into an array of devices, Au NPs decoration was performed using a previously described magnetron sputtering inert gas condensing method [23,25]. The Au NPs were size selected to have an average diameter of 2.5 nm (Supplementary Fig. S1) which illustrate the G-FET output and transfer characteristics, respectively. The Id-Vd curve decreases with a slight reduction in
magnitude of the applied liquid gate voltage, indicating the sensitivity of the device to the gate bias and verifying that a good ohmic contact exists between the graphene and gold electrode \([37,38]\). The G-FETs are operated at low voltages so that any electrochemical processes or gate currents are negligible; Fig. S2b shows that under normal G-FET operation, \(I_g\) is approximately 3 orders of magnitude smaller than \(I_d\). Due to the electronic band structure of graphene, its conductivity cannot be switched off at room temperature. However, at some applied liquid gate voltage, the current will be at a minimum as the ambipolar device switches from the hole conducting regime to the electron conducting regime (moving from more negative to more positive voltages). This voltage is known as the charge neutrality potential, \(V_{\text{CNP}}\), which can be affected by electrostatic gating or doping from molecules bound to the graphene. The shift in the \(V_{\text{CNP}}\) due to binding of a target analyte is the sensor response of the G-FET.

Fig. 4a shows the shift in transfer characteristics of the Au NPs decorated G-FET after DNA functionalization and ethanolamine passivation. Initially, the Au NPs decorated G-FET has a \(V_{\text{CNP}}\) of approximately 0.44 V, heavily p-doped from the ideal case of 0 V due to the p-type doping from the Au NPs deposition and adsorbates from the environment \([11,39]\). After exposure to a 10 \(\mu\)M solution of the thiol-terminated DNA aptamer, numerous single strands are strongly bound to the Au NPs via the Au-S bond. This causes a large leftward, negative shift in the \(V_{\text{CNP}}\) to approximately 0.3 V as the electron-rich nucleobases in the DNA n-dope the graphene. Electrostatic gating from the negatively charged bound DNA aptamer would instead result in a positive shift, therefore our results can more easily be explained by n-doping of the graphene channel from interactions between the DNA backbone and the graphene surface \([4]\). From the method described by Xu et al. 2017, we estimate the density of bound aptamers on the G-FET to be \(3.5 \times 10^{10} \text{ cm}^{-2}\), comparable to the density of deposited Au NPs seen in the AFM imaging experiments \([40]\). This leftward shift in response to DNA binding is commonly seen in G-FETs that utilize Au NPs \([4,10,11,39]\), in contrast to chemically functionalized G-FETs which report a positive \(V_{\text{CNP}}\) shift from DNA binding due to electrostatic gating \([9,12]\). The G-FET is then passivated using an ethanolamine solution to deactivate and block reactive groups on the graphene surface, preventing non-specific binding.

After the passivation step, we performed the DNA sensing experiments. Beginning with the lowest concentration, a droplet of buffered DNA fully complementary to the aptamer (cDNA) was placed on the G-FET channel and interacted with the Au NPs bound aptamer for 30 min to allow hybridization \([11,12,15,41]\). Afterwards, the device was rinsed with 1X PBS and DI water to remove weakly bound or unhybridized strands, and the transfer characteristics were measured again in 0.01x PBS. Fig. 4b shows the set of G-FET transfer curves for different concentrations of cDNA from 1 nM to 100 pM. As the cDNA concentration increases, a steady leftwards shift in \(V_{\text{CNP}}\) is observed, indicating increasing n-doping of the graphene channel by the cDNA. For cDNA concentrations greater than 1 pM, \(V_{\text{CNP}}\) does not significantly change, as the DNA aptamer strands bound to the Au NPs have all been hybridized and no further doping is possible. The slight increase in minimum current with increased cDNA concentration can be explained with charge doping and the interaction of the charged molecules with charged impurities on the graphene surface \([7,42]\). Due to doping from the cDNA, electron density in the graphene channel increases, resulting in an upward shift of the Fermi level. Electron mobility is preserved due to increased scattering of hole carriers in the graphene channel with the negatively charged cDNA \([7]\). Positively charged impurities in the graphene channel or defects are screened by counterions in the solution \([43,44]\).

Fig. 5 shows the shift in the charge neutrality point (\(\Delta V_{\text{CNP}}\)) for two series of measurements on Au NPs decorated and bare G-FET devices exposed to cDNA concentrations of 1 nM to 100 pM. Each data point in Fig. 5 is an average of 5 different G-FETs and the error bars are one standard deviation. The Au NP functionalized sensor exhibits a logarithmic response between 1 nM and 1 pM before saturating at \(\Delta V_{\text{CNP}} \approx 50 \text{ mV}\) in the picomolar cDNA concentration range. This corresponds to a sensitivity to six orders of magnitude in analyte concentration, or a
Au NPs decoration (roughly $10^{10}$ cm$^{-2}$) [12]. Extending the Au NPs deposition time or functionalization methods which normally result in DNA probe denaturation, the sensor was tested by comparing the activity of biosensor G-FETs.

The specificity of the sensor was tested by comparing the $\Delta V_{\text{CNP}}$ upon exposure to 100 pM of the cDNA, and two different sequences which each exhibit a different single nucleotide polymorphism (M1 and M4 SNP). In Fig. 6, it is apparent that the one-base mismatched SNP exhibit a significantly reduced $\Delta V_{\text{CNP}}$ at 100 pM, roughly equivalent to the shift produced by the fully complementary strand at 1 aM ($\approx$ 10 mV). This shift in $\Delta V_{\text{CNP}}$ is also similar in magnitude to the shift seen in G-FET hysteresis and extended bias stress measurements (Supplementary Fig. S2c and d), which is due to carrier density enhancement in the graphene caused by capacitive gating [46]. We can use the maximum signal produced by the M1 SNP to set the limit of detection (LoD) of the sensor to $\approx 15$ aM, a significant improvement over many G-FETs functionalized by chemical methods or solvent-based Au NPs [10–12,45,47]. The achieved LoD indicates that using graphene transferred via the laminate method does not intrinsically limit the sensitivity of biosensor G-FETs.

We can also employ the G-FETs functionalized with the DNA aptamer to sense streptavidin in a 0.01x PBS solution. After passivation with 5 μM of BSA, solutions of streptavidin in 0.01x PBS were placed on the G-FET channel and left to interact with the DNA aptamer bound to the Au NPs. The transfer characteristics of the devices were measured after 30 min. without an intervening rinsing step, to prevent the weakly bound streptavidin from being detached from the DNA aptamer. Fig. 7 shows the $\Delta V_{\text{CNP}}$ for two series of measurements on Au NPs decorated and bare G-FET devices exposed to streptavidin concentrations of 1 aM to 100 pM. As before, each data point in Fig. 7 is an average of 5 different devices and the error bars are one standard deviation. Streptavidin is known to have a slight negative charge in neutral pH solutions [48,49], therefore a leftward shift due to the n-doping of graphene that increases with solution concentration is again observed. As for the cDNA case, the G-FET sensor is capable of analyte detection in the attomolar range, using the maximum response of the bare graphene sensor to the streptavidin solutions as the noise limit for the sensor. We estimate an LoD of $\approx 9$ aM and a sensitivity of approximately 14 mV/decade. However, the G-FET rapidly saturates upon exposure to 1 fM concentrations of streptavidin (dynamic range of $10^5$) and is less stable at higher concentrations. Due to streptavidin’s much lower charge in solution than the complementary strand [49], a higher population of binding sites may be required to improve this dynamic range.

4. Conclusion

In summary, we have developed an Au NPs-decorated graphene FET biosensor capable of label-free and ultrasensitive (in the attomolar range) detection of DNA and streptavidin using a liquid gate measurement. The sensor was fabricated using a simple graphene lamination method and functionalized via solvent-free gas-phase synthesized Au NPs, a novel combination. Underdecorated G-FETs show no sensor response to the complementary DNA strand or streptavidin. The decorated G-FET can detect full hybridization of the complementary strand down to 15 aM; solutions with SNP-containing DNA only produce an equivalent response at concentrations 7 orders of magnitude higher.
CRediT authorship contribution statement


Declaration of Competing Interest

The authors declare no competing financial interest.

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Appendix A. Supplementary data

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