Indium tin oxide (ITO) has been widely used as electrodes in optoelectronic devices such as organic light-emitting diodes (OLEDs) and photovoltaic cells, due to its high transparency and good conductivity. The price of ITO has gone up significantly in recent years due to the limited indium resources. In addition, their chemical and electronic stability and mechanical flexibility limit their applications in flexible electronics.1 With the emergence of novel flexible electronic devices, there is a growing demand for flexible electrodes, where conventional materials such as metal conductors and ITO may not reach the required bendability.

Two-dimensional networks of carbon nanotubes (CNTs) have been considered as a candidate to replace ITO because they possess excellent mechanical2,3 and electrical properties.4 Recently, there has been tremendous research in the two-dimensional crystal graphene and its derived nanomaterials.5 Graphene sheets can be considered as an unzipped CNT, which has unique physical properties such as high carrier mobility,5 long phase coherence length,6 fractional quantum Hall effect,10 bipolar supercurrent,11 suppression of the weak localization,12 deviation from the adiabatic Born–Oppenheimer approximation,13 etc. On the basis of these unique electronic properties and the two-dimensional planar structure, graphene also appears as a promising material for transparent electrodes.5,14,15 Furthermore, besides good conductivity and transparency of the electrode, the performance and current density for semiconducting electronic devices, such as light-emitting diodes and field-effect transistors, strongly rely on the carrier injection efficiency through the contact between electrodes and semiconducting material layers.16,17 Therefore, the band alignment between electrodes and semiconductors and the surface property of the electrode have been an extensively investigated area.18–20 Finding a certain conductive material with the desired work function for efficient carrier injection may sometimes be difficult. Both theoretical and experimental results have demonstrated that the Fermi energy shift of single-layer graphene (SLG) can be achieved through deliberate doping by aromatic molecules,21,22 gas molecules,23 or electrostatic gating.5,24 These results suggest that controlling the work function of graphene electrodes is possible.

Tremendous progress has been achieved for the synthesis and characterization of graphene in the past few years.25 Transparent graphene electrodes have been formed by either reducing graphene oxide26 or a chemical bottom-up approach, where molecules of giant polycyclic aromatic hydrocarbons cross-link to form graphene nanosheets.27 However, the conductivity of these films is still limited28 and
high enough electrical conductivity is required for good electrodes.

Recently, it has been demonstrated that continuous large-area graphene thin films can be synthesized by vacuum graphitization and ambient pressure chemical vapor deposition (APCVD) methods. In this contribution, we use graphene films formed by chemical vapor deposition (CVD-G) as an ultrathin transparent electrode. We demonstrated that the work function of graphene can be tuned by AuCl₃ doping. AuCl₃ is a commonly used compound in doping organic conductors. The Au⁺ can be reduced to Au particles by the charge transfer from graphene which results in p-type doping behavior of graphene. AuCl₃ has been chosen for the stability of its doping effect. The surface potential change of graphene as a function of doping time was studied by measuring the contact potential difference (CPD) using Kelvin probe microscopy. Silicon photovoltaic devices with graphene electrodes were fabricated as a demonstration of the advantages of using CVD-G electrodes with tunable work function.

RESULTS AND DISCUSSION

The images from the atomic force microscopy (AFM) in Figure 1A–D and from scanning Kelvin probe microscopy (SKPM) in Figure 1A’–D’ show the effect of AuCl₃ doping time on the surface morphology and surface potential of CVD-G, respectively. With longer doping time, it was observed that the size of Au particles became larger and the morphology changed. The formation of Au particles on CVD-G can be understood from an electrochemical perspective since the reduction potential of AuCl₄⁻ ion (1.0 V) is higher than the reduction potential of graphene (0.22 V). In aqueous solutions, AuCl₃ forms a square planar complex, AuCl₄⁻, and the following reaction takes place on graphene:

\[
\text{AuCl}_3^- + 3e^- \rightarrow \text{Au}^{0} + 4\text{Cl}^-
\]  

(1)

We noticed that for samples with 20 s doping time, the Au particles uniformly cover the surface of the graphene film. For less than 20 s doping, Au particles tend to form on the wrinkles and thicker layers of CVD-G. As the doping time increases over 20 s, larger Au particles form and the size distribution of Au particles becomes broader. The SKPM images in Figure 1 indicate that the measured surface potential \(V_{\text{surf}}\) of Au particles is higher than adjacent graphene region (where color contrast is lower), and the \(V_{\text{surf}}\) difference between Au particles and graphene is from ~0.3 to ~0.8 V, depending on the Au particle size. The \(V_{\text{surf}}\) of Au particles is always more positive than that of adjacent graphene regions, suggesting that the doping caused by Au particles is p-type on graphene (i.e., Au particles receive electrons from graphene), which is consistent with the chemical reaction 1. This hole doping process results in the work function shift of the CVD-G. Figure 1E plots the \(\Delta V_{\text{surf}}\) versus various AuCl₃ doping times. Since the CPD method relies on the stability of the work function of AFM tips and is also sensitive to the measurement environment, such as humidity and electric grounding, we took the \(V_{\text{surf}}\) as the averaged value over an area of 1 µm × 1 µm and the as-grown CVD-G film was used as a standard reference. Thus, \(\Delta V_{\text{surf}}\) value are relative to the pristine as-grown sample:

\[
\Delta V_{\text{surf}} = V_{\text{surf}}(\text{sample after doping}) - V_{\text{surf}}(\text{as-grown graphene})
\]  

(2)

For doping time less than 20 s, \(V_{\text{surf}}\) monotonically increases with doping time to about 0.5 V, and as the doping time further increases, the \(V_{\text{surf}}\) reduces slightly. Thus SKPM data indicate that the work function of CVD-G can be tuned by chemical doping and exposing to AuCl₃ results in electrons being withdrawn from CVD-G by the formation of Au particles, which give rise to an up-shift of \(V_{\text{surf}}\). The surface potential measurements were consistent with electric transport measurements of CVD-G field-effect transistors in our previous experiments. The slight down-shift for longer doping time is consistent with the previous results.
time is unclear at this stage and will be subject to further investigation. One mechanism could be the hygroscopic nature of AuCl₃ at higher concentrations. The water molecule could affect the charge transfer behavior between graphene and AuCl₃.

The benefit for having a transparent electrode with tunable work function will be demonstrated subsequently by enhancing the performance of a photovoltaic device. After CVD synthesis, the graphene films were transferred to n-Si substrates by coating a thin layer of PMMA on top of graphene film. The underlying Ni film was etched by Ni etchant (TFB commercial etchant from Transene). Then the graphene/PMMA film was transferred to AuCl₃ solution for chemical doping. After AuCl₃ doping, the sheet resistance of graphene film decreases from 400 to ~100 Ω/sq (see ref 37 for details). The device fabrication procedure is schematically shown in Figure 2. Using CVD synthesis and the PMMA transfer method, the size of graphene conducting film can be easily scaled up without a fundamental limit.

Using the PMMA transferred CVD-G film, we fabricated photovoltaic diodes with a structure as shown in Figure 2. In order to demonstrate the effect of the work function shift on the device performance, the current density of several diodes under varying biases is compared in Figure 3A. These devices were measured with and without AM 1.5 illuminations for pristine CVD-G electrode and the CVD-G film with 20 s AuCl₃ doping. The devices show clear photovoltaic behaviors, which are similar to the previous reported work of carbon nanotube film/n-Si p-n junction solar cells. Without illumination, both devices show a typical diode behavior, but the open circuit voltage ($V_{oc}$) and short circuit current ($I_{sc}$) are dramatically improved for the device with 20 s AuCl₃ doped CVD-G. More detailed AM 1.5 measurements for the devices with various doping time are shown in Figure 3B. The device containing the 20 s AuCl₃ doped CVD-G film displays the highest $V_{oc}$ and $I_{sc}$. For longer doping times, both of the $V_{oc}$ and $I_{sc}$ drop and it can be noticed that for the samples with extremely long doping times, for example, 20 min, the electrical behavior shows no large difference to 30 s doping, which is consistent with the SKPM results. The device performance as a function of AuCl₃ doping time is plotted in Figure 3C. The device with 20 s AuCl₃ doping shows the best photovoltaic performance. This observation can be explained by the SKPM results in Figure 1E. Figure 3D schematically demonstrated the effect of work function change of graphene electrodes on the device performance. The work function of graphene is as large as graphite, ~4.66 eV, and our SKPM measurement shows that with AuCl₃ modification the work function of graphene film can be up-shifted within the range of 0.5 eV. It has been reported that a heterojunction can be formed between carbonaceous (or graphitic) thin films and n-type silicon. Under photoillumination, electron and hole pairs are generated in n-Si, similar to the case of carbon nanotube/n-Si solar cell; the photogenerated electrons are directed toward the n-Si, while holes are injected into the graphene electrodes, due to the built-in electric field at the Si/graphene junction. However, unlike p-type semiconductor carbon nanotube, here graphene is a zero band gap material and it is simply treated as a metal. Therefore, the open circuit voltage corresponds to the amount of forward bias on the solar cell due to the bias of the solar cell junction with the light-generated current. Similar to the case for metal-on Si Schottky barrier solar cell, $V_{oc}$ can be expressed as:

$$V_{oc} = n\phi_B + (kT/q)\ln(I_s/A_eA^*T^2)$$

where $n$ is the diode ideality factor, $\phi_B$ is the barrier height, $T$ is the working temperature, $I_s$ is the diode saturation current, $A_e$ and $A^*$ are the contact area of the diode and the Richardson constant, respectively. Thus, the surface potential modification (or work function engineering) of graphene electrodes results in different barrier heights between Si and graphene, which provides tunability for the $V_{oc}$ in the proposed n-Si/graphene photovoltaic devices. Besides the work function engineering effect, doping with Au particles affects the output characteristics of the photovoltaic device in several other ways. First, the doping also affects the carrier concentration of the graphene film, which further changes the sheet resistance of the graphene film. A reduction in sheet resistance could also contribute to the change of short circuit current. Figure 3C clearly

![Figure 2](image-url)
shows the evolution of $V_{oc}$, $I_{sc}$, and power conversion efficiency (PCE) consistent with the $V_{sur}$ shift in Figure 1E. Also, with a short AuCl$_3$ doping time, the Au particle layer is very thin, and the interface dipole between graphene and Au particles is expected to contribute strongly to the charge-injection process. With the increase of doping time, other effects such as transparency of the graphene film and the morphology-induced sheet resistance change need to be taken into consideration for device integration. A highest PCE achieved is \(~0.08\%\), which is 40 times larger than the device without chemical doping. This value is comparable with the reported multiwalled CNT/n-Si solar cell devices. It is suggested that the PCE of devices could be further improved by inserting a suitable p-type material between n-type Si and CVD-G electrodes or by opening the band gap of graphene. In summary, the chemical doping with Au particles can control the surface properties of graphene films, and the surface potential shift is able to explain the electrical characteristics for the diodes.

**CONCLUSION**

In conclusion, our EFM results show that the chemical doping of CVD-G with AuCl$_3$ causes a surface potential up-shift up to 0.5 eV, which is attributed to the hole doping process from Au particles. When n-type Si is covered by the CVD-G films, photoinduced charge separation can be observed. The CVD-G electrodes offer more flexibility in improving electric properties of the photovoltaic devices due to work function engineering. Our results suggest a new strategy for charge harvesting by graphene films. The detailed photoelectrical behaviors of CVD-G/semiconducting composite devices, particularly the graphene electrode contacts are worthy of further investigation. The capability of tailoring electrical properties of graphene is potentially useful for optoelectronics.

**EXPERIMENTAL SECTION**

Few-layer graphene (FLG) film was synthesized by CVD method using 300 nm Ni-coated Si/SiO$_2$ substrates. The Ni substrates were annealed in the CVD chamber with H$_2$ and Ar (H$_2$/Ar = 400:600 sccm) for 20 min at 1000 °C. CH$_4$ (4 sccm) along with H$_2$ (1400 sccm) was then flowed at 1000 °C for 5 min to carburize the Ni film. Finally, the Ni film was cooled to 500 °C with a cooling rate of 5 °C/min under CH$_4$/H$_2$/Ar = 4:700:700 sccm, in order to control the thickness of graphene films. For the as-grown FLG films, one or two graphene layers were formed in most areas (see Supporting Information Figure S1). Details on the graphene film growth were reported elsewhere. Atomic force microscopy (AFM), Raman spectroscopy, and transmission electron microscopy (TEM) techniques were used to characterize the CVD-synthesized FLG as described in refs 32 and 33. After CVD-G growth, the sample was immersed into AuCl$_3$ aqueous solution with a concentration of 5 mM for various durations. Then the sample was rinsed with DI water and blow-dried by nitrogen gas. In order to have a better grounding, the bottom Ni layer was not removed for the sample used for SKPM measurement. The surface potential ($V_{sur}$) of CVD-G before and after doping with AuCl$_3$ was measured by a dual-pass technique in tapping mode using the scanning Kelvin probe method (SKPM) based on an AFM system Dimension 3000 from Veeco Metrology Group with NanoScope Signal Access Module. The measurement was conducted in air with Olympus (OMCL-AC240TM) Pt-
coated cantilevers. The tip curvature radius is ~15 nm, quality factor ~190, spring constant 2 N/m, resonance frequency ~70 kHz, and cantilever length 240 μm. The CPD value between tip and sample surface was taken as the averaged value over an area of 1 μm × 1 μm. The standard deviation of the experiment was 10 mV.

The surface potential study was performed by a dual-pass technique in tapping mode. Topography information (AFM image) was acquired in the first scan; the second scan was then performed while the tip was maintained at a constant distance (10 nm above the sample surface). Both a DC signal and an AC signal at the resonant frequency of the cantilever are applied to the metal-coated AFM probe while the tip is lifted up. If a potential difference (∆V_p) exists between the tip and the sample surface, the signal creates a varying electrostatic force, causing an oscillating motion in the cantilever. The ∆V_p is measured by adjusting the DC voltage until there is no DC potential difference. The DC voltage is recorded as the CPD value.

Photovoltaic devices were fabricated based on n-Si substrate with a resistivity around 5 Ω·cm. The as-grown CVD-G film was coated with a thin layer of PMMA (~100 nm) by spin coating to avoid the breaking of graphene film.63 After etching the underlying polycrystalline Ni with Ni etchant, the PMMA/CVD-G film was transferred to DI water or additionally 5 mM AuCl₃ aqueous solution for chemical doping. After that, the film was transferred back to water and suspended on the surface of DI water. The film was later picked up by an n-type silicon substrate with an attached insulating glass cover slide. Ti (10 nm) and Au (200 nm) were evaporated on the top side of the glass slide in advance, which forms the upper electrode connects to the PMMA/CVD-G film. The rest of the PMMA/CVD-G film was brought in contact with the Si surface after careful blow drying with N₂ gas. The bottom electrodes were formed by connecting electrical wires to the back side of Si using silver paint. The distance of the area of the PMMA/CVD-G film on top of Si is 3 mm × 3 mm. The photovoltaic devices were irradiated under a solar simulator at AM 1.5 with a light intensity of 100 mW. Electrical testing was performed by an Agilent 4156C precision semiconductor parameter analyzer.

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Supporting Information Available: Figure S1. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES AND NOTES