Stable, efficient p-type doping of graphene by nitric acid

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ABSTRACT

We systematically dope monolayer graphene with different concentrations of nitric acid over a range of temperatures, and analyze the variation of sheet resistance under vacuum annealing up to 300 °C. The optimized HNO₃ doping conditions yield sheet resistances as low as 180 Ω/sq, which, under vacuum annealing, is significantly more stable than previously reported values. Raman and photoemission spectroscopy show that this stable graphene doping occurs by a bi-modal mechanism. At mild conditions the dopants are weakly bonded to graphene, but at high acid temperatures and concentrations, the doping is higher and more stable upon post-doping annealing, without causing significant lattice damage. This work shows that large, stable hole concentrations can be induced by transfer doping in graphene.
**Introduction**

Graphene exhibits remarkable electrical, optical and mechanical properties,\(^1,\)\(^2\) which may be advantageous for various applications,\(^3\) including electronics,\(^4\) photovoltaics,\(^5\) energy storage,\(^6\) lighting\(^7\) and displays.\(^8\) In particular, graphene is envisaged as an environmental-friendly and flexible substitute for indium tin oxide (ITO) as transparent conductor,\(^9,\)\(^10\) but obtaining low and stable sheet resistances remains difficult. Pristine graphene exhibits high carrier mobility, but its intrinsic carrier density is close to zero, making its conductivity uncompetitive with ITO. Doping is therefore necessary and can be achieved by various means, such as substitution of atoms in the lattice,\(^11,\)\(^12\) functionalization,\(^13\) and adsorption of atoms and molecules.\(^14,\)\(^15\) Of these, substitutional doping in 2D crystals significantly alters the graphene lattice perturbing the unique properties of graphene;\(^11,\)\(^12\) hence less invasive techniques, like adsorptive doping, are more desirable. The ideal doping induces a high charge carrier density \((> 10^{13} \text{ cm}^{-2})\), remains stable in time across a broad temperature range, and is transparent and homogeneous. Thermal stability is particularly important in optoelectronic devices where significant power may be dissipated. However, to date efficient charge carrier transfer and stability have not been simultaneously achieved. Stable doping can be obtained, for instance, by evaporating high work function oxides like MoO\(_3\)\(^16,\)\(^14,\)\(^17,\)\(^7\) which, whilst effective for carbon nanotube forests where the resistance is lowered by two orders of magnitude,\(^18\) only show a two-fold reduction in sheet resistance (from \(~1210\) to \(~600 \Omega/\text{sq}\)) for monolayer graphene.\(^14\) The charge transfer is thus simply not sufficient for many of the envisioned applications of graphene as a transparent conductor.

In 2010, Bae *et al.*\(^8\) reported that graphene doped with HNO\(_3\) exhibits a remarkably low sheet resistance of \(~125 \Omega/\text{sq}\) for monolayer graphene. Subsequent studies however showed unstable behavior in air at room temperature (RT) over days\(^15\) or on vacuum annealing.\(^7\) The poor stability
of the HNO₃ doping was attributed to the volatility of adsorbed molecules, but a full understanding of the surface chemistry after doping is still lacking. Until now, the stability of HNO₃-doped graphene was measured on samples with various layer numbers, and employing diverse doping procedures and techniques for stability evaluation,⁷,¹⁵ making comparisons of these results difficult. In this report, we systematically analyze the stability of HNO₃-doped graphene, and demonstrate a doping procedure that lowers the sheet resistance of monolayer graphene to 180 Ω/sq and shows stable behavior following annealing in vacuum up to 130 °C. These results represent a step towards graphene with a sheet resistance, which is both stable and low enough to be competitive with ITO. Furthermore, our results provide more general insights into the chemical effects of HNO₃ treatments on graphitic materials, of relevance to processes such as the removal of impurities from carbon nanotubes,¹⁹ and the preparation of graphene oxide.²⁰ We also contrast the behavior of HNO₃ to react with defects in carbon systems, as for example in the purification of carbon nanotubes,²¹ with the pure transfer doping which does not form new defects or react with existing ones.

**Experimental section**

As a substrate for device fabrication, we use silicon wafers with 300 nm of SiO₂, on which we pattern electrodes (Cr/Au 5/45 nm) by electron beam lithography and UVIII resist. Monolayer graphene is grown by chemical vapor deposition on an Alfa Aesar Cu foil (25 μm, 99.8% purity) pre-treated by electropolishing in a home-built cell. The foil is heated up at temperature in Argon atmosphere in a hot-wall CVD furnace to 1000 °C, and exposed to a mixture of 0.08 sccm CH₄, 10 sccm H₂, and 490 sccm Ar for 140 min to grow monolayer graphene on the surface. The foil is then cooled down in a mixture of 10 sccm H₂ and 490 sccm Ar. More detail can be found in a
The graphene is then transferred to patterned electrodes by a wet transfer process, using a poly-methyl methacrylate (PMMA) support layer and an aqueous solution of ammonium persulfate (0.1 M) as etchant. The graphene is then patterned using electron beam lithography with UVIII resist followed by etching with oxygen plasma. To perform resistivity measurements, we fabricate 4 point probe devices with 4 µm wide graphene channels and electrode distances of 5/10/5 µm. For mobility measurements, we fabricate Hall devices with Van der Pauw geometry on a 50×50 µm² piece of graphene with 2.5×2.5 µm² contacts at the corners. To remove residues attached to graphene during transfer and device fabrication, we perform vacuum annealing at 180 °C and 10⁻⁷ mbar for 1 hour.²³

Acid treatment is performed by dipping the chip into 65% HNO₃ diluted in DI water to achieve concentrations from 15-65%. To control the acid temperature, ~5 ml of acid in a 10 ml beaker is placed on a hotplate, while the temperature of the liquid is checked by a type K thermocouple. The sample is then blown with dry nitrogen. Annealing of the sample after doping is performed at 10⁻⁷ mbar with the temperature measured with a type K thermocouple placed on the heater for the samples used for electrical and Raman measurements, while at 10⁻⁸ mbar with the type K thermocouple placed on the sample for the X-ray photoemission spectroscopy (XPS) measurements.

XPS measurements are performed on a Mg X-ray source (hv=1253.6 eV). The photoelectrons are detected with an energy resolution of ~0.8 eV in normal emission geometry. The spectra are fitted with Doniach-Šunjić profiles convoluted with Gaussians after subtracting a Shirley background. The N:C ratio is calculated by evaluating the areas of the N 1s and C 1s photoemission spectra, normalized by the photoionization cross section area of the different elements. The N atom density is then extracted by the known atomic density of C atoms in
graphene (calculated using 0.142 nm as C-C bond). If not stated otherwise, Raman spectra are acquired by a 532 nm laser at ~1.5 mW with a 50× objective (spot size of ~1 μm). The Raman peaks are fitted with Lorentzian profiles.

**Results**

Figure 1 shows that higher HNO₃ concentrations and temperatures are more effective in lowering the sheet resistance of graphene and achieving a more stable resistance. The plots show the change in resistance obtained by dipping the sample in HNO₃ for different (a) acid concentrations, (b) exposure times, and (c) acid temperatures. The change in resistance with acid concentration is measured for samples exposed to acid for 5 min at RT. The measurements show a similar decrease in resistance for both 15% and 30% HNO₃ concentrations to ~0.2× that of the pristine graphene, while at 65% concentration the resistance halves again to give a sheet resistance 0.1× that of the pristine graphene. The doping shows no significant dependence on exposure time for our timescales, as 15 s of dipping time in 65% HNO₃ already reaches a resistance value of 0.12× the original value, while at 1, 5, and 20 min the ratio saturates at 0.10. A further decrease in resistance occurs by increasing the temperature of the acid solution. Keeping both concentration and dipping time constant at 65% and 5 min, the resistance decreases to 0.07× the original resistance when the acid solution is at 52 °C. These ratios were consistently observed for several devices with different initial sheet resistances in the range [2600-3500] Ω/sq. After pre-annealing in vacuum at 180 °C for 1h, the average sheet resistance is 3000 Ω/sq, while after doping with 65% acid for 5 min the sheet resistance decreases to ~310 Ω/sq when the acid is at RT and to ~210 Ω/sq when at 52 °C. The lowest individual-device sheet resistance measured is 180 Ω/sq, after a hot acid treatment. There is a contrast between the one
order of magnitude reduction in graphene resistance by HNO$_3$ treatment reported here and the more modest reductions reported elsewhere, e.g. by three times by Kasry et al.$^{24}$ This can be explained by the higher resistive baseline of pre-annealed graphene, the number of graphene layers analyzed and the optimization of the doping procedure herein.

Figures 1(d,e,f) show the fraction of sheet resistance regained upon heating in vacuum to 70 °C, 130 °C, 200 °C and 300 °C, i.e. the increase in resistance for post-annealing over the reduction in resistance upon doping. Diluted acid (15% and 30%) produces samples that are less stable than that doped with 65% acid solution. The difference is particularly evident after the 200 °C annealing which regains nearly ~40% of the sheet resistance reduction achieved by low concentration acid dipping, while a recovery of just 13% occurs for more concentrated acid. The stability varies little with dipping time, but it depends strongly on the acid temperature. When graphene is doped with 65% HNO$_3$ for 5 min at 52 °C, the doping is maintained even on vacuum annealing up to 130 °C. The percentage of resistance recovery at 130 °C is less than 2% and corresponds to an absolute change in sheet resistance from ~210 to ~250 Ω/sq.

Figure 2 shows the trend of graphene mobility and charge carrier density after transfer, doping, and post-annealing at 200 °C. The data is measured on 6 devices fabricated on three different chips, doped with low and high HNO$_3$ concentration (22% and 65%) at RT, and high concentration (65%) at 52 °C. The as-transferred graphene (before pre-annealing) presents an average Hall mobility of ~5500 cm$^2$V$^{-1}$s$^{-1}$ and charge carrier density of ~1.5×10$^{12}$ cm$^{-2}$. The doping increases the charge carrier density considerably up to ~4×10$^{13}$ cm$^{-2}$ in the case of heated acid, but the mobility decreases to below 900 cm$^2$V$^{-1}$s$^{-1}$. Finally, when post-annealed in vacuum, both mobility and charge carrier density partially recover, increasing and decreasing their values respectively. It is interesting to note that the mobility of the three samples does not diverge
significantly for the different concentrations or temperatures of acid, while the charge carrier density behaves as the conductivity, i.e. it is higher and more stable for higher concentrations and temperatures of acid. We have not included the results on pre-annealed samples, since the measurements on different devices show large variations. The possible reason for this may relate to the Fermi energy being very close to the Dirac point. The residual surface contamination may therefore create regions of $p$-type and $n$-type doping (charge puddles),25 and thus the assumption of uniform doping in Van der Pauw Hall measurements no longer holds.

Figure 3a confirms the non-destructive nature of HNO$_3$ doping. The Raman spectra of pre-annealed graphene show $I_{2D}/I_G$ ratios of ~2.09, and 2D peak FWHM of ~28 cm$^{-1}$, which are typical values for monolayer graphene transferred on silicon dioxide$^{26,27}$. After doping with low concentration HNO$_3$ (23%), high concentration acid (65%) and high concentration heated acid (52 °C), the $I_{2D}/I_G$ ratio decreases to 1.77, 1.69, and 1.59 respectively (still large), the G peak position shifts from 1590 cm$^{-1}$ to 1592 cm$^{-1}$, 1593 cm$^{-1}$, 1594 cm$^{-1}$ and the 2D peak wavenumber shifts from 2681 cm$^{-1}$ to 2682 cm$^{-1}$, 2683 cm$^{-1}$, 2685 cm$^{-1}$ respectively. These variations in the Raman spectra are consistent with increasing doping.$^{28}$ All spectra have a small D peak ($I_D/I_G$ ratio of ~0.03), which indicates the high crystalline quality of the graphene$^{29}$ and the non-destructive nature of HNO$_3$ doping.

Figure 4a shows the C 1$s$ spectrum of graphene before and after doping, fitted to resolve the contributing components. Before doping, the spectrum shows a main graphitic component and three smaller peaks at higher binding energies. The main peak is due to C-C $sp^2$ bonds in the graphene layer, while the other peaks can be attributed to $sp^3$ carbon, C-OH and C=O, corresponding to both defects in the graphene layer and to polymer residue.$^{30}$ After dipping the sample in 65% HNO$_3$ at 52 °C the higher binding energy components become 10% weaker, and
all the peaks shift by ~0.55 eV to lower binding energies. The reduction of intensity of the non-graphitic peaks can be attributed to cleaning of polymer contamination from the graphene surface, as in Kasry et al.,\textsuperscript{24} who observed higher transparency after HNO\textsubscript{3} treatment. The shift indicates a band bending towards \textit{p}-doping. The doping is also confirmed by the increase in asymmetry of the graphitic peak, attributable to an increase in charge carrier density. Figure 4b shows the N 1s spectra of graphene doped with 20\% HNO\textsubscript{3} at RT and 65\% HNO\textsubscript{3} at RT and 52 °C, all of which have been post-annealed at 200 °C. The spectra of doped graphene (dark green and dark blue) show two main components at ~400 eV and ~406 eV, which can be attributed to nitrogen close to carbon atoms and NO\textsubscript{x} radicals, respectively. The peak at ~400 eV is already detected in the pre-annealed (undoped) sample, as in previous reports on other graphitic materials like graphite\textsuperscript{31} or single-walled carbon nanotubes (SWCNTs).\textsuperscript{32} The peak at ~406 eV, seen in HNO\textsubscript{3} treated graphite and SWCNTs, has to our knowledge not been previously reported for monolayer or few-layer graphene.\textsuperscript{33} Upon annealing the peak intensity at ~406 eV lowers, while the C-N peak remains constant. The ratio between nitrogen and carbon atoms on the surface does not exceed 2\%, even for the case of 65\% concentrated acid at RT, which is consistent with a hole density of 10\textsuperscript{13} cm\textsuperscript{-2}.

**Discussion**

The stability of the doping by HNO\textsubscript{3} is expressed as a fraction of the recovered sheet resistance at different temperatures. In a simple model, the observed increase in sheet resistance with annealing temperature can be explained by the desorption of molecules from the graphene surface, for which a similar behavior in the normalized data of figures 1d, 1e, and 1f would be expected. However, our data show significant deviation from this model, suggesting a combination of doping mechanisms is instead involved. From the XPS and Raman spectra, we
infer that the doping is not exclusively related to absorbed NO$_3^-$, as suggested in literature,$^{24}$ but also to a contribution from other adsorbed molecules or atoms, without further significant alteration of the graphene lattice.

We propose the following model based on the evolution of both XPS and Raman spectra, and on the work by Martyna et al.$^{34}$ The HNO$_3$ molecules physisorb onto the graphene sheet, without breaking any C-C bonds. The HNO$_3$ molecule then dissociates into three groups, two radicals NO$_2^0$ and NO$_3^0$ and a water molecule,

$$2\text{HNO}_3 = \text{NO}_2^0 + \text{NO}_3^0 + \text{H}_2\text{O}$$

All three species are physisorbed. The two radicals are calculated to have a singly occupied state below the Fermi energy of the graphene layer.$^{34}$ This allows two electrons to transfer from graphene into these states, creating two holes in the graphene and causing $p$-type doping. The radicals become anions, NO$_2^-$ and NO$_3^-$, and they become bound to the graphene holes by Coulomb attraction. However, they do not form chemisorptive bonds to adjacent carbon atoms, the bond length is too large for this, and the adjacent carbon atoms do not pucker upwards as required to allow the formation of these extra bonds. Thus, the physisorptive sites do not cause an increase in the Raman D peak (as observed).

In addition to NO$_3^-$ adsorption/desorption, the stability assessment reveals an additional factor when doping graphene with concentrated HNO$_3$ at high temperatures. We ascribe this to oxygen-rich groups decorating the graphene layer which are present mostly at higher concentration and temperature of the HNO$_3$ doping process. Concentrated HNO$_3$ is known to be a strong oxidant
which induces the functionalization of graphitic carbon, especially at high temperatures. This was observed on graphite\textsuperscript{35} and on multilayer graphene,\textsuperscript{33} where oxygen is covalently bonded to the graphene lattice as hydroxyl and carboxyl groups. Given our conditions (monolayer graphene transferred onto a silica substrate), we cannot directly determine by XPS the binding of oxygen species to graphene: The O 1s is dominated by the substrate oxygen, while the peaks referring to carbon bound to oxygen in the C 1s spectrum are covered by the overlapping signal coming from residual PMMA after transfer.

Interestingly, we do not observe an increase in the D-peak in the Raman spectrum. We cannot completely exclude that the doping is not introducing any defects in the lattice, as doped graphene presents a lower I_D/I_G in respect to undoped graphene with the same density of defects.\textsuperscript{36} As a control, we immersed a sample of graphene in 65\% at 52 °C for 2 hours, and the D peak is still not found to increase in intensity (Fig. 3b), and neither after annealing at 130 °C. Similar to our case, it has been shown that exposure of graphene to oxygen at high temperatures (below 300 °C) leads to stable adsorption without a significant increase in the D-peak.\textsuperscript{37} Liu et al.\textsuperscript{38} have shown that annealing monolayer graphene in oxygen to 200-300°C for 2 hours, partially etched the layer, but also induced a high hole carrier density (\textasciitilde2\cdot10^{13} \text{ cm}^{-2}). During temperature ramping, the doping manifests first, while the etching (increase of D-peak) occurs at higher temperatures. The doping is therefore not due to damaging of the honeycomb lattice. The stable doping component can be attributed either to 1) oxygen (or an oxygen group) covalently bonded to graphene, consistent with the doping stability but less so with the low D-peak; or to 2) oxygen (or an oxygen group) behaving as a charge-transfer complex, consistent with the low D-peak, but with the doping stability being less expected. Reports in literature describe cases of oxygen functionalization with minor graphene lattice damage.\textsuperscript{39,40}
The doping of graphene by HNO₃ is therefore unstable when only caused by molecular adsorbates of NO₃⁻ but high doping stability is achieved under conditions where the binding or adsorption of oxygen is induced. At acid concentrations below ~50%, the obtained doping is mainly unstable. In contrast, using higher concentrations and temperatures, the doping becomes stable. This explains figure 1d, which show a significant variation of normalized doping stability when graphene is doped with acid concentrations lower or higher than 50%. By increasing the acid temperature, the doping stability increases, indicating that the doping component related to oxidation is increased. This hypothesis is also confirmed by the change in slope of the 52°C acid temperature line in figure 1f after the annealing at 200 °C, in agreement with previous results.³⁹

Figure 5a compares the stability data of HNO₃-doped graphene with similar studies done on I₂- and MoO₃-doped graphene.¹⁴ The plot relates the absolute values of sheet resistance of doped graphene before and after an annealing at 130 °C. MoO₃ and I₂ are two extreme cases of highly stable and highly unstable dopants respectively. Although MoO₃ doping gives high stability,¹⁴ the absolute values of sheet resistance are still relatively high. HNO₃, in the case of annealed and concentrated acid, presents an intermediate stability behavior, along with a significant decrease in absolute sheet resistance.

Figure 5b compares the transparency between the same doping methods. HNO₃-doped graphene enhances the transparency from 97.1% of pristine graphene to 97.4% of transmission at 550 nm; I₂-doped graphene from 97.1% to 97.6% of transmission at 550 nm, while an adlayer of 5 nm of MoO₃ reduces the transparency (from 97.1% to 95.4% of transmission at 550 nm), especially towards the UV region. The improved transparency of the acid can partly be attributed to the removal of polymer contaminations.²⁴
It is worth noting that, even though the monolayer-graphene sheet resistance in this work is comparable with the lowest reported to date, it may be possible to further reduce this value by changing or pretreating the substrate or by modifying the CVD graphene quality.

Conclusion

We have demonstrated that low sheet resistances and stability can be obtained simultaneously in adsorptive-doped graphene. We have analyzed the doping of graphene with HNO$_3$ at various concentrations and temperatures, and its stability upon annealing in vacuum. We have found that graphene doped with concentrated HNO$_3$ at high temperatures exhibits both a sheet resistance as low as 180 $\Omega$/sq and an improved stability, i.e. less than 2% recovery of the doping resistance drop when annealing at 130 °C for 30 min. We find the doping and stability to arise from a combination of doping mechanism. Upon exposure to HNO$_3$ at mild conditions, graphene adsorbs unstable NO$_3^-$ molecules which easily desorb at low temperatures. Conversely, doping at high concentrations and temperatures creates stable $p$-type doping by the adsorption or functionalization of oxygen. This report of both stable and effective doping reveals a promising route toward ITO replacement by graphene.

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Notes

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Figure 1: (a, b, c) sheet resistance of graphene doped with different (a) HNO₃ concentrations, (b) dipping times, and (c) acid temperatures. The left axis refers to values normalized to the undoped sheet resistance, which are consistent for a range of different undoped sheet resistances (2600-3500 Ω/sq.). The right axis shows the indicative absolute sheet resistance of an average device. (d, e, f) variation of sheet resistance upon annealing normalized to the variation of sheet resistance upon doping, i.e. the ratio of sheet resistance recovered upon annealing. The data refers to graphene doped with different (d) HNO₃ concentrations, (e) dipping times, and (f) acid temperatures and the annealing is performed sequentially at 70 °C, 130 °C, 200 °C and 300 °C for 30 minutes. The errors are the standard deviation on the average of measurements taken on different devices.

Figure 2: Mobility (μ) and charge carrier density (n) for graphene as-transferred, doped and post-annealed (200 °C, 30 min), measured with Van der Pauw geometry Hall devices. The graph regards three samples with different doping: 5 min of dipping into 22% (blue) and 65% (green) of HNO₃ solution at RT, and 65% of HNO₃ solution at 52 °C (orange and red).

Figure 3: (a) Raman spectra of graphene before (pre-annealed) and after doping with 22% HNO₃ solution at RT for 5 min, 65% HNO₃ solution at RT for 5 min and 65% HNO₃ solution heated at 52°C for 5 min. (b) Raman spectra of graphene after doping with 65% HNO₃ solution heated at 52°C for 120 min, and after post-annealing at 130°C. The region near the D-peak (~1345 cm⁻¹) has been enlarged by a factor 10. All spectra are normalized by the G-peak intensity and are the average of 3 spectra taken at different positions on the sample. The spectrum in red (post-annealed) is taken with a 5 times higher laser intensity (~7.5 mW).

Figure 4: (a) XPS C 1s spectra of graphene pre-annealed at 180 °C for 1 hour, and after dipping the sample in 65% HNO₃ solution for 5 min at 52 °C. (b) XPS N 1s spectrum of pre-annealed undoped graphene (top spectrum), and XPS N 1s spectrum of graphene doped with 65% HNO₃ solution for 5 min at RT and 52 °C, and with 22% HNO₃ for 5 min at RT, and the same spectra after post-annealing at 200 °C.

Figure 5: (a) Comparison of sheet resistances after doping and after post-doping annealing for graphene doped with HNO₃ (5 min 52°C 65%), and MoO₃ and I₂ from a previous work. The annealing was performed at 130 °C in vacuum (10⁻⁶ mbar) for 30 min for all dopants. (b) UV-VIS transmittance measurements of as-transferred graphene and HNO₃-doped graphene, compared to graphene doped with MoO₃ and I₂ from a previous work. MoO₃ reduces the transparency of graphene; while HNO₃ and I₂ slightly increase the light transmittance, probably due to the removal of PMMA residue.
Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.


