Probing thermal expansion of graphene and modal dispersion at low-temperature using graphene nanoelectromechanical systems resonators

Vibhor Singh¹, Shamashis Sengupta¹, Hari S Solanki¹, Rohan Dhall¹, Adrien Allain¹, Sajal Dhara¹, Prita Pant² and Mandar M Deshmukh¹

¹ Department of Condensed Matter Physics, TIFR, Homi Bhabha Road, Mumbai 400005, India
² Department of Metallurgical Engineering and Materials Science, IIT Bombay, Powai, Mumbai 400076, India

E-mail: deshmukh@tifr.res.in

Received 25 November 2009, in final form 23 February 2010
Published 30 March 2010
Online at stacks.iop.org/Nano/21/165204

Abstract

We use suspended graphene electromechanical resonators to study the variation of resonant frequency as a function of temperature. Measuring the change in frequency resulting from a change in tension, from 300 to 30 K, allows us to extract information about the thermal expansion of monolayer graphene as a function of temperature, which is critical for strain engineering applications. We find that thermal expansion of graphene is negative for all temperatures between 300 and 30 K. We also study the dispersion, the variation of resonant frequency with DC gate voltage, of the electromechanical modes and find considerable tunability of resonant frequency, desirable for applications like mass sensing and RF signal processing at room temperature. With a lowering of temperature, we find that the positively dispersing electromechanical modes evolve into negatively dispersing ones. We quantitatively explain this crossover and discuss optimal electromechanical properties that are desirable for temperature-compensated sensors.

Online supplementary data available from stacks.iop.org/Nano/21/165204/mmedia

1. Introduction

Electronic properties of graphene have been studied extensively [1, 2] since the first experiments probing the quantum Hall effect [3, 4]. In addition to the electronic properties, the remarkable mechanical properties of graphene include a high in-plane Young's modulus of \( \sim 1 \) TPa probed using nanoindentation of suspended graphene [5], force extension measurements [6] and electromechanical resonators [7–9]. NEMS (nanoelectromechanical systems) devices using nanostructures like carbon nanotubes [10–16], nanowires [17, 18] and bulk micromachined structures [19–21] offer the promise of new applications and allow us to probe fundamental properties at the nanoscale. NEMS [22]-based devices are ideal platforms to harness the unique mechanical properties of graphene. Electromechanical measurements with graphene resonators [7, 8] suggest that, with improvement of the quality factor \( (Q) \), graphene-based NEMS devices have the potential to be very sensitive detectors of mass and charge. Additionally, the sensitivity of graphene to chemical-specific processes [23] offers the possibility of integrated mass and chemical detection. The large surface-to-mass ratio of graphene offers a distinct advantage over other nanostructures for such applications. In order to better understand the potential of graphene-based electrically actuated and detected resonators [8] and the challenges in realizing strain-engineered graphene devices [24, 25], we experimentally measure the coefficient of thermal expansion of graphene \( (\alpha_{\text{graphene}}(T)) \) as a function of temperature. Our measurements indicate...
\( \alpha_{\text{graphene}}(T) < 0 \) for 30 K \(< T < 300 \text{ K} \) and larger in magnitude than theoretical prediction [26]. We also probe the dispersion, or the tunability, of mechanical modes using the DC gate voltage at low temperatures and find that the thermal expansion of graphene, built-in tension and added mass play an important role in changing the extent of tunability of the resonators [27] and the resonant frequency—parameters that are critical for various applications. Additionally, measuring temperature-dependent mechanical properties [28] of suspended structures down to low temperatures will give an insight into strain engineering of graphene-based devices [24, 25] and also help in understanding the role of rippling in degrading carrier mobility at low temperatures. Our experiments probe these issues in detail and suggest that monitoring the resonant frequency as a function of temperature could provide important information about nanoscale stress that is useful for probing phase transitions.

2. Experimental details

2.1. Device fabrication

To fabricate monolayer graphene electromechanical resonators, we suspended graphene devices using the previously reported [8, 29–31] process which starts with micromechanical exfoliation of graphene from graphite crystal using an adhesive tape on a 300 \( \mu \text{m} \) thick degenerately doped silicon wafer coated with 300 nm thick thermally grown \( \text{SiO}_2 \). Following the location of monolayer graphene flakes using optical microscopy, electron beam lithography is used to pattern resist for fabricating electrodes for electrical contact. The electrodes are fabricated by evaporation of 10 nm of chromium and 60 nm of gold following patterning of the resist. To release the graphene from the \( \text{SiO}_2 \) substrate, a dilute buffered-HF solution is used to selectively etch an area around the graphene device by masking the rest of the substrate using a polymer resist. Following an etch for 5 min 30 s which results in a 170 nm deep trench in \( \text{SiO}_2 \), the device is rinsed in DI water and isopropanol. Critical point drying, to prevent collapse of the device due to surface tension, is the final step in the fabrication of suspended graphene devices. A scanning electron microscope (SEM) image of a suspended graphene device is shown in figure 1(a).

2.2. Measurement scheme

The electrical actuation and detection is done by using the suspended graphene device as a heterodyne mixer [8, 10, 32]. The scheme for electrical actuation and detection is also shown in figure 1(a) superimposed on the SEM image of the device. We use electrostatic interaction between the graphene membrane and the back-gate electrode to actuate the motion in a plane perpendicular to the substrate. A radio-frequency (RF) signal \( V_g(\omega) \) and a DC voltage \( V_g^{\text{DC}} \) are applied at the gate terminal using a bias-tee. Another RF signal \( V_{SD}(\omega + \Delta \omega) \) is applied to the source electrode, figure 1(a). The RF signal applied at the gate \( V_g(\omega) \) modulates the gap between graphene and substrate at frequency \( \omega \) and \( V_g^{\text{DC}} \) alters the overall tension and carrier density in the membrane. The

![Figure 1](image)

(a) Tilted angle scanning electron microscope (SEM) image of a suspended monolayer graphene device and the electrical circuit for actuation and detection of the mechanical motion of the graphene membrane. The scale bar indicates a length of 2 \( \mu \text{m} \). (b) A plot of the mixing current \( I_{\text{mix}}(\Delta \omega) \) as a function of frequency \( f(=\frac{\omega}{2\pi}) \) at 7 K with the DC gate voltage \( V_g^{\text{DC}} = -5 \text{ V} \) for device (D1). The sharp feature in the mixing current corresponds to the mechanical resonance. (c) Colorscale plot of the mixing current as a function of frequency \( f \) and DC gate voltage \( V_g^{\text{DC}} \) at 7 K for device (D1). The dashed line shows the position of the line scan shown in (b). The dotted–dashed line indicates the position of the Dirac peak for the graphene device. The measured dispersion of this mode is negative. The maxima (blue) and minima (red) of the colorbar correspond to 2.6 and 0 nA, respectively. (d) Plot of the measured \( Q \) for the data shown in (c). The \( Q \) drops as \( |V_g^{\text{DC}}| \) increases and around the Dirac point the \( Q \) shows a dip. (e) Colorscale plot of \( I_{\text{mix}}(\Delta \omega) \) as a function of frequency \( f \) and the DC gate voltage for device (D2) at 300 K. Two positively dispersing modes of the device are seen.
amplitude of the current through the graphene membrane at the difference frequency ($\Delta \omega$), also called the mixing current $I_{\text{mix}}(\Delta \omega)$, can be written as $[8, 10, 16, 33, 34]$

$$I_{\text{mix}}(\Delta \omega) = \frac{1}{2} \frac{dG}{dV} \left( \frac{dC}{dz} \right) \left[ C_g V_g(\omega) + C_V V_g(\omega) \right] V_{SD}(\omega + \Delta \omega),$$

where $G$ is the conductance of the graphene device, $q$ is the charge induced by the gate voltage, $C_g$ is the capacitance between the gate electrode and graphene and $z(\omega)$ is the amplitude of oscillation at the driving frequency $\omega$ along the $z$ axis (perpendicular to the substrate). The difference frequency signal at ($\Delta \omega$) arises from the product of the modulation signals of $V_{SD}(\omega + \Delta \omega)$ and $G(\omega)$. At the mechanical resonance of the membrane, the first term in equation (1) contributes significantly and the second term, which does not depend on the mechanical motion of the graphene membrane, provides a smooth background.

### 2.3. Results and discussions

Figure 1(b) shows the result of such a measurement at 7 K for a suspended graphene device (D1) while $V_{gDC}$ is set at $-5$ V. Using a Lorentzian lineshape for the resonance curve $[10, 35]$, we can extract the $Q \sim 1500$ of the resonator (variation of $Q$ with temperature is shown in supplementary information available at stacks.iop.org/Nano/21/165204/mmedia). Figure 1(c) shows the colorscale plot of $I_{\text{mix}}(\Delta \omega)$ as a function of $V_{gDC}$ and $f = \omega/2\pi$ at 7 K. The resonant frequency is lowered as the magnitude of $V_{gDC}$ is increased—mechanical mode disperses negatively with $|V_{gDC}|$. This is well understood in terms of mode softening due to the capacitive contribution to the energy of the resonator $[18, 36]$. The Dirac peak for our device is shifted ($V_{D} = 13$ V) from the expected position (0 V) due to unintentional doping during the fabrication process.$^3$ Also seen in figure 1(c) is the information regarding the amplitude of the mixing current. The overall amplitude of the mixing current at resonance scales with $V_{gDC}$ (the first term in equation (1)) and, as a result, the amplitude of the mixing current is very small near $V_{gDC} = 0$ V. However, at $V_{gDC} = 13$ V, the amplitude of the mixing changes abruptly due to the Dirac peak ($V_{D}$) $[3, 4]$. At this point, the term $\frac{dG}{dV}$ appearing in equation (1) becomes zero. Figure 1(d) shows the quality factor ($Q$) dependence on $V_{gDC}$ calculated from the data shown in figure 1(c). The quality factor of the device is largest around $V_{gDC} = 0$ V and decreases as the gate voltage is swept away from zero. Also, at low temperatures, $f_0$ decreases with the increase in $|V_{gDC}|$ and $Q$ follows a similar trend. As negative dispersion—caused by softening of the spring constant—is accompanied by the movement of the graphene towards the gate electrode, it leads to an increase in the modulated capacitance and results in larger dissipative current $[36]$. In the neighborhood of $V_{gDC} = V_D$, $Q$ is smaller than the value expected from the trend from the higher $V_{gDC}$.

$^3$ A shift in the Dirac peak away from $V_{gDC} = 0$ V is desirable in our experiments because the actuation efficiency is feeble at $V_{gDC} = 0$ V and that would make the observation of physics near the Dirac point inaccessible for electromechanical measurements.

side. One possible reason for this behavior might be that, in the vicinity of the Dirac point, large device resistance leads to increased ohmic losses $[35]$. An additional mechanism that can result in dissipation is the mechanical deformation modifying the distribution of charge puddles of electrons and holes in the graphene sheet. These phenomena warrant further detailed investigation $[37, 38]$. Such a measurement could be used to study charge inhomogeneity. Figure 1(e) shows a similar measurement for a device (D2) at 300 K. Here, we observe two mechanical modes which disperse positively because of the increase in tension in the graphene membrane due to the electrostatic attraction from the back gate. Most of our graphene devices exhibit an in-built tensile stress at 300 K due to the fabrication process. Multiple ripples on the membrane are also seen in figure 1(a) $[28]$. To better understand the modal dispersion quantitatively, we model the graphene membranes within the elastic continuum regime and in a limit where tension in the membrane dominates over the flexural rigidity $[5, 7, 39]$. As a result, the resonant frequency $f_0$ of the graphene membrane can be written as

$$f_0(V_{gDC}) = \frac{1}{2L} \sqrt{\frac{\Gamma}{\rho t w}},$$

where $L$ is the length of the membrane, $w$ is the thickness, $\rho$ is the mass density, $\Gamma_0(T)$ is the in-built tension, $\Gamma$ is the tension at a given temperature $T$ and $V_{gDC}$. The functional form of $\Gamma(\Gamma_0(T), V_{gDC})$ is dependent on the details of the model used to take into account the electrostatic and elastic energies. (Our model is described in detail in the supplementary information available at stacks.iop.org/Nano/21/165204/mmedia.) Resonant frequency at zero gate voltage is given by $f_0(0) = \frac{1}{2L} \sqrt{\frac{\Gamma_0(T)}{\rho t}}$; therefore, from $f_0(0)$ an independent estimation of $\Gamma_0(T)$ and $\rho$ is not possible. However, resonant frequency dispersion with gate voltage $f_0(V_{gDC})$ allows us to estimate $\rho$ and $\Gamma_0(T)$ simultaneously. Using fits based on the continuum model (calculations described in supplementary information available at stacks.iop.org/Nano/21/165204/mmedia) we estimated the mass density $\rho = 7.4 \rho_{\text{graphene}}$ and the in-built tension $\Gamma_0 (300 \text{ K})$ to be 68.6 nN for the fundamental mode, where $\rho_{\text{graphene}}$ is the mass density of pristine graphene for the device data shown in figure 1(e). We attribute this extra mass and in-built tension to the resistive residue that can get deposited on the graphene membrane during the fabrication process and these values of mass density are similar to ones reported by Chen et al $[8]$. We note that the dispersion of the two modes shown in figure 1(e) is different and our modal calculations give $\rho = 7.3 \rho_{\text{graphene}}$ and $\Gamma_0(300 \text{ K}) = 107.6 \text{ nN}$ for the upper mode. However, the simple model of a rectangular membrane predicts higher-order modes at much higher frequencies—this is a limitation of the simple assumptions we have made. The presence of other resonant modes at small intervals of frequency $f$ seen in figure 1(e) (also seen in figure 4(a)) indicates a deviation from this simplified picture of the membrane under tension due to either edge modes $[9]$ or due to the rippling/curling of the membrane $[28, 40]$. In the later part
of this report, we discuss in detail the reasons for positively or negatively dispersing modes and temperature-induced change in dispersion.

3. Probing thermal expansion of graphene

We now consider how the resonant frequency \( f_0 \) evolves as a function of the temperature. Figure 2(a) shows the result of an evolution of a mode as a function of temperature at \( V_{DC} = 15 \) V. The resonant frequency increases as the device is cooled from room temperature. This increase has been seen in all the devices we have studied. The degree of frequency shift varies from one device to another, depending on the device geometry. The origin of this frequency shift with temperature is the increase in tension in graphene due to the expansion/contraction of substrate, gold electrodes and graphene. The frequency shift can be understood by taking into account the contribution of various strains as the device is cooled below 300 K. The three main contributions are—firstly, the thermal strain in unconstrained graphene \( \epsilon_{\text{graphene}}(T) = \int_T^{300} \alpha_{\text{graphene}}(t) dt \) due to the coefficient of thermal expansion of graphene \( \alpha_{\text{graphene}}(T) \); second, the thermal strain due to the gold electrodes \( \epsilon_{\text{gold}}(T) = \int_T^{300} \alpha_{\text{gold}}(t) dt \) and lastly the contribution of the strain induced by the substrate \( \epsilon_{\text{substrate}}(T) \); here \( \epsilon_{\text{gold}}(T) \) is the coefficient of thermal expansion for gold [41].

The strain in gold electrodes plays an important role due to the geometry of the device. The under-etch that releases the graphene membrane also etches under the graphene covered by the electrodes—resulting in the graphene membrane being suspended off the gold electrodes [8] (see supplementary information for SEM images available at stacks.iop.org/Nano/21/165204/mmedia). The geometry of the resulting device is shown in the inset to figure 2(a). The elastic strain in the curved substrate can be calculated using Stoney’s equation [42]. Its contribution to the strain in the graphene membrane is very small (see supplementary information available at stacks.iop.org/Nano/21/165204/mmedia) and therefore, upon cooling, a change of tension in the graphene is due to contraction of the gold electrode and expansion/contraction of graphene. We assume that Young’s modulus of graphene does not vary significantly over the temperature range [43]. At the interface of the gold electrodes supporting the graphene membrane, the net force must balance to zero; however, the stresses are different considering the cross-sectional area of gold electrodes (~500 nm x 60 nm) and graphene (~500 nm x 0.3 nm). The large difference in cross-sectional area implies that the effective stiffness of gold electrodes is large compared to the stiffness of graphene. As a result, to a very good approximation, the total elastic strain at a given temperature in graphene that is confined by ‘rigid’ gold electrodes is

\[
\epsilon_{\text{graphene clamped}} = \epsilon_{\text{graphene}}(T) + \epsilon_{\text{substrate}}(T) - \epsilon_{\text{gold}}(T)\,,
\]

where \( \omega_{\text{electrode}} \) is the average of the width of gold electrodes holding the suspended flake. The change in tension in the membrane can be written as a function of temperature as

\[
\Delta \Gamma_{\text{g}}(T) = \omega_{\text{electrode}} \epsilon_{\text{graphene clamped}}(T) E_{\text{graphene}},
\]

where \( E_{\text{graphene}} \) is Young’s modulus of graphene. Measuring the tension \( \Gamma_{\text{g}}(T) \) as a function of temperature offers a way to track the thermal strain in graphene membrane. Figure 2(a) shows the evolution of resonant frequency as a function of temperature from a device (D2) at \( V_{DC} = 15 \) V, where the increase in frequency is largely due to the contraction of the gold electrodes. However, this rate of increase of resonant frequency is significantly reduced due to the negative \( \epsilon_{\text{graphene}} \) for all \( T < 300 \) K from the case of frequency change including only the gold’s contraction. Using such a measurement of frequency shift while assuming uniform expansion of all the materials and using equation (2), it is possible to calculate the expansion coefficient from the frequency at \( V_{DC} = 0 \) V as

\[
\alpha_{\text{graphene}}(T) = -2 f_0(0) \frac{\mathrm{d} f_0(0)}{\mathrm{d} T} \times \frac{(2L)^2 \rho}{E_{\text{graphene}}} \frac{\epsilon_{\text{substrate}}(T) - \epsilon_{\text{gold}}(T) \omega_{\text{electrode}}}{L}.
\]
in the vicinity of a temperature coefficient of thermal expansion. This can possibly be one reason for its deviation from the theoretically predicted values. In our calculation, we have assumed that the presence of a residue does not affect the expansion of graphene, which may not be the complete description of the experimental system. However, these values of $\alpha_{\text{graphene}}$ still remain valuable for device engineering as the complete elimination of the resist residue from the device is difficult.

4. Modal dispersion of graphene NEM resonator

We next consider the dispersion of the resonant modes with temperature. Figure 3(a) shows the dispersion of resonant mode for a device (D3) at various temperatures down to 6 K. Upon cooling, the resonant frequency increases and its tunability using gate voltage is reduced, an undesirable feature. Further, at low temperatures, resonance frequency first decreases with $|V_{g}^{DC}|$ and then it increases. (Similar evolution of modal dispersion with $V_{g}^{DC}$ has been observed by Chen et al [8].) We now try to understand the evolution of the modal dispersion as a function of temperature. The two limiting cases of modal dispersion—pure positively and negatively dispersing modes—can be respectively understood by considering the limits where the tension (or in the case of flexural modes mechanical rigidity) dominates with increasing $V_{g}^{DC}$ and the case when capacitive pulling leading to the softening of the spring constant dominates [18, 35, 36]. A simple model [18] to understand the intermediate regime where these two interactions compete is to model the resonator with an intrinsic spring constant $K_i = k + \alpha(V_{g}^{DC})^2 + \beta(V_{g}^{DC})^4 + H O(V_{g}^{DC})$ with $\alpha$ and $\beta$ as constants of the system (when $\beta > 0$, this is consistent with the positive dispersion of modes with an increased $V_{g}^{DC}$). A second contribution due to the electrostatic interaction (capacitive coupling) softens the intrinsic spring constant $K_i$ so that the effective spring constant

$$K_{\text{eff}} = K_i - \frac{1}{2} \left( V_{g}^{DC} \right)^2 \frac{\partial^2 C_{\text{eff}}}{\partial z^2}.$$
with $C_g$ being the capacitive coupling of the membrane with gate electrode. As $V_{g}^{DC}$ is varied the $K_{eff}$ varies and the modal dispersion changes from a negatively dispersing mode to a positively dispersing mode at large $V_{g}^{DC}$—the value of $V_{g}^{DC}$ at which the crossover happens is a measure of the relative contribution of capacitive and elastic energies. This phenomenological model describes the generalized modal dispersion and can be connected to the properties of the resonator (described in supplementary information available at stacks.iop.org/Nano/21/165204/mmedia). $\alpha$ and $\beta$ depend on temperature through the coefficient of thermal expansion, and also on the elastic constant of graphene; as a result they change with temperature, leading to the evolution of modal dispersion as a function of temperature. Additionally the capacitive contribution to energy is expected to change as a function of temperature. The observed modal dispersion behavior at all temperature can be qualitatively understood from the contraction of the suspended gold electrodes with temperature, which leads to increased tension in the membrane. This increase in in-built tension is accompanied by the reduction in tunability with $V_{g}^{DC}$ (model described in the supplementary information available at stacks.iop.org/Nano/21/165204/mmedia). Results of our calculation (figure 3(c)) clearly show that we can successfully model the temperature evolution of dispersion, using only a single fit parameter in our model. It is critical to understand the origin of observed dispersion of resonant frequency as a function of temperature due to the desirable property of the resonator—its tunability. Additionally, if the loss mechanisms are frequency-independent such tunability can increase the $Q$ of the system [35].

In order to use graphene electromechanical resonators for applications like mass and charge sensing it is important to understand the microscopic origin of the modes and to test the suitability of continuum models [7, 8, 39] beyond basic properties such as dispersion. Scanned probe measurements to image modes [9] indicate that the origin of these modes is complex—for instance, some of the modes are associated with the edges of the graphene membrane. The reason this is critical is that the notion of effective mass of mechanical modes [22, 45] is associated with the spatial distribution of the mode$^6$. We try to understand the nature of modes in graphene resonators by studying devices with multiple resonances in the measurement frequency window. Figure 4(a) shows two higher-order modes along with a fundamental mode for a graphene resonator at 7 K. The dispersions of these three modes are different. Though it is difficult to determine the exact nature of these modes using our technique, a semi-quantitative explanation is provided here. At lower temperatures, negative dispersion can be understood in terms of spring constant softening as discussed earlier. Following equation (4), the effective spring constant $K_{eff}$ can be used to calculate the modal dispersion with gate voltage ($V_{g}^{DC}$):

$$f^2 = f_0^2 - \frac{1}{8\pi^2 m_{eff}} V_{g}^{DC} \left( \frac{d^2 C_g}{dz^2} \right)^2,$$

where $f_0 = \frac{1}{2\pi} \sqrt{\frac{K_{eff}}{m_{eff}}}$ and $m_{eff}$ is the effective mass for a given mode [22, 45].

Using equation (5), we can describe the higher modes also by using $f_0$ and modal mass ($m_{eff}$) as the fitting parameters. For a sheet under tension with uniform loading, the effective modal mass ($m_{eff}$) for different modes does not change, unlike the case of flexural modes, and is equal to $m_{eff} = 0.785\rho Lw$ [45]. However, our fitting gives different effective masses for different modes, which is expected as these are not the higher-order harmonics of the fundamental mode (in the model described above, the higher-order modes will be integral multiples of the fundamental mode). This suggests that the simple picture of a rectangular membrane under tension within the continuum description does not work well to describe the system. This could be due to four main reasons:

(a) non-uniform loading of the membrane (due to resist residue) [8] can modify the $m_{eff}$ for different modes,
(b) due to the presence of edge-dependent modes [9],
(c) due to the curvature of membrane [40] or rippling of graphene [28] (as seen in figure 1(a)) the effective stiffness of the modes could be a very complex quantity with a tensor nature and could be significantly different from the ideal value of $\sim 1$ TPa as observed for the case of rippled carbon nanotubes [46–48] and
(d) the capacitance to gate ($C_g$) for the graphene membrane is likely to be mode-dependent.

Further experiments with pristine unrippled graphene resonators are needed to clarify our understanding of the microscopic origin of modes. The presence of multiple closely spaced modes can be potentially useful for mass spectrometry as the position of the added mass can be extracted accurately [49].

5. Conclusions

To conclude, our measurements of $Q$, for graphene resonators, as a function of DC gate voltage indicate that larger dissipation in the resonator can occur around the Dirac point. Using these NEMS devices measurement of the coefficient of thermal expansion of graphene at low temperatures is possible. Our measurements indicate that $\alpha_{\text{graphene}}$ is negative for all temperatures between 300 and 30 K and larger in magnitude than the numbers predicted by theoretical calculations [26]; this could be critical in designing strain-engineered devices using graphene [24, 25]. Additionally, the modal dispersion of graphene resonators is affected by the thermal expansion of graphene, reducing the tunability. The continuum description of the mechanics of a rectangular graphene membrane is inadequate for explaining the resonances due to the presence of non-uniform impurities and rippling of the membrane.

Acknowledgments

The authors thank the Government of India for the financial support. The authors would like to thank Smita Gohil and Subramanian S for their help during the experiments.
References

[1] Castro Neto A H, Guinea F, Peres N M R, Novoselov K S and Geim A K 2009 The electronic properties of graphene Rev. Mod. Phys. 81 109–54

[2] Geim A K 2009 Graphene: status and prospects Science 324 1530–4

[3] Zhang Y, Tan Y-W, Stormer H L and Kim P 2005 Experimental observation of the quantum Hall effect and berry’s phase in graphene Nature 438 201–4

[4] Novoselov K S, Geim A K, Morozov S V, Jiang D, Katsnelson M I, Grigorieva I V, Dubonos S V and Firsov A A 2005 Two-dimensional gas of massless dirac fermions in graphene Nature 438 197–200

[5] Lee C, Wei X, Kysar J W and Hone J 2008 Measurement of the elastic properties and intrinsic strength of monolayer graphene Science 321 385–8

[6] Gomez-Navarro C, Burghard M and Kern K 2008 Elastic properties of chemically derived single graphene sheets Nano Lett. 8 2045–9

[7] Bunch J S, van der Zande A M, Verbridge S S, Frank I W, Lau C N and von Klitzing K and Yacoby A 2008 Observation of the quantum hall effect and berry’s phase in graphene by strain engineering Nat. Phys. 5 109–54

[8] Garcia-Sanchez D, van der Zande A M, San Paulo A, Lassagne B, McEuen P L and Bachtold A 2008 Imaging mechanical vibrations in suspended graphene sheets Nano Lett. 8 1399–403

[9] Sazonova V 2006 A tunable carbon nanotube electromechanical resonator Nano Lett. 6 2045–9

[10] Novoselov K S, Geim A K, Morozov S V, Jiang D, Katsnelson M I, Grigorieva I V, Dubonos S V and Firsov A A 2005 Two-dimensional gas of massless dirac fermions in graphene Nature 438 197–200

[11] Lee C, Wei X, Kysar J W and Hone J 2008 Measurement of the elastic properties and intrinsic strength of monolayer graphene Science 321 385–8

[12] Gomez-Navarro C, Burghard M and Kern K 2008 Elastic properties of chemically derived single graphene sheets Nano Lett. 8 2045–9

[13] Bunch J S, van der Zande A M, Verbridge S S, Frank I W, Lau C N and von Klitzing K and Yacoby A 2008 Observation of the quantum hall effect and berry’s phase in graphene by strain engineering Nat. Phys. 5 109–54

[14] Bachtold A 2008 Ultrasensitive mass sensing with a nanotube electromechanical resonator Nano Lett. 8 3735–8

[15] Du X, Skachko I, Barker A and Andrei E Y 2008 Approaching ballistic transport in suspended graphene Nanotech. 3 491–5

[16] Bolotin K I, Sikes K J, Jiang Z, Klima M, Fudenberg G, Hone J, Kim P and Stormer H L 2008 Ultrahigh electron mobility in suspended graphene Solid State Commun. 146 351–5

[17] Sazonova V 2006 A tunable carbon nanotube resonator PhD Thesis Cornell University

[18] Kozinsky I, Ch Postma H W, Bargatin I and Roukes M L 2006 Tuning nonlinearity, dynamic range, and frequency of nanomechanical resonators Appl. Phys. Lett. 88 253101–3

[19] Martin J, Akerman N, Ulbricht G, Lohmann T, Smet J H, Kozinsky I, Ch Postma H W, Bargatin I and Roukes M L 2006 A tunable carbon nanotube resonator Nano Lett. 6 2045–9

[20] Unterreithmeier Q P, Weig E M and Kotthaus J P 2009 Universal transduction scheme for nanomechanical systems based on dielectric forces Nature 458 1001–4

[21] Naik A K, Hanay M S, Hiebert W K, Feng X L and Roukes M L 2009 Towards single-molecule nanomechanical mass spectrometry Nat. Nanotechnol. 4 445–50

[22] Schedin F, Geim A K, Morozov S V, Hill E W, Blake P, Katsnelson M I and Novoselov K S 2007 Detection of individual gas molecules adsorbed on graphene Nat. Mater. 6 652–5

[23] Pereira V M and Castro Neto A H 2009 Strain engineering of graphene’s electronic structure Phys. Rev. Lett. 103 046801

[24] Guinea F, Katsnelson M I and Geim A K 2009 Energy gaps and a zero-field quantum hall effect in graphene by strain engineering Nat. Phys. advance online publication

[25] Bachtold A 2009 Coupling mechanics to charge transport in carbon nanotube electromechanical resonator Appl. Phys. Lett. 81 532–4

[26] Hui H-Y, Hung P, Ch Postma H W and Bockrath M 2008 Atomic-scale mass sensing using carbon nanotube resonators Nano Lett. 8 4342–6

[27] Feng X L, He R, Yang P and Roukes M L 2007 Very high frequency silicon nanowire electromechanical resonators Nano Lett. 7 1953–9

[28] Solanki H S, Sengupta S, Dhara S, Singh V, Dhall R, Patil S, Parpia J, Bhattacharyya A and Deshmukh M M 2010 Tuning mechanical modes and influence of charge screening in nanowire resonators Phys. Rev. B 81 115459

[29] Carr D W, Evoy S, Sekaric L, Craighead H G and Parpia J M 1999 Measurement of mechanical resonance and losses in nanometer scale silicon wires Appl. Phys. Lett. 75 920–2

[30] Katsnelson M I and Novoselov K S 2007 Detection of individual gas molecules adsorbed on graphene Nat. Mater. 6 652–5

[31] Guinea F, Katsnelson M I and Geim A K 2009 Energy gaps and a zero-field quantum hall effect in graphene by strain engineering Nat. Phys. advance online publication

[32] Kozinsky I, Ch Postma H W, Bargatin I and Roukes M L 2006 Tuning nonlinearity, dynamic range, and frequency of nanomechanical resonators Appl. Phys. Lett. 88 253101–3

[33] Atalaya J, Isacsson A and Kinaret J M 2008 Continuum elastic modeling of graphene resonators Nano Lett. 8 4196–200

[34] Shivaraman S et al 2009 Free-standing epitaxial graphene Nano Lett. 9 3100–5

[35] Nix F C and MacNair D 1941 The thermal expansion of pure metals: copper, gold, aluminium, nickel, and iron Phys. Rev. 60 597

[36] Freund L B and Suresh S (ed) 2003 Thin Film Materials (Cambridge: Cambridge University Press)
[43] Jiang J-W, Wang J-S and Li B 2009 Young’s modulus of graphene: a molecular dynamics study Phys. Rev. B 80 113405

[44] Ho G K, Sundaresan K, Pourkamali S and Ayazi F 2005 Low-motional-impedance highly-tunable i2 resonators for temperature-compensated reference oscillators MEMS 2005: 18th IEEE Int. Conf. on Micro Electro Mechanical Systems pp 116–20

[45] Karabacak D M 2007 Resonant operation of nanoelectromechanical systems in fluidic environments PhD Thesis, Boston University

[46] Poncharal P, Wang Z L, Ugarte D and de Heer W A 1999 Electrostatic deflections and electromechanical resonances of carbon nanotubes Science 283 1513–6

[47] Arroyo M and Belytschko T 2003 Nonlinear mechanical response and rippling of thick multiwalled carbon nanotubes Phys. Rev. Lett. 91 215505

[48] Liu J Z, Zheng Q and Jiang Q 2001 Effect of a rippling mode on resonances of carbon nanotubes Phys. Rev. Lett. 86 4843

[49] Dohn S, Svendsen W, Boisen A and Hansen O 2007 Mass and position determination of attached particles on cantilever based mass sensors Rev. Sci. Instrum. 78 103303