

Four-Point Resistance of Individual Single-Wall Carbon Nanotubes

B. Gao,¹ Y. F. Chen,² M. S. Fuhrer,² D. C. Glattli,^{1,3} and A. Bachtold^{1,4,*}

¹*Laboratoire Pierre Aigrain, Ecole Normale Supérieure, 75231 Paris 05, France*

²*Department of Physics and Center for Superconductivity Research, University of Maryland, College Park, Maryland 20742, USA*

³*SPEC, CEA Saclay, F-91191 Gif-sur-Yvette, France*

⁴*CNM-CSIC, Campus Universitat Autònoma de Barcelona, E-08193 Bellaterra, Spain*

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We have studied the resistance of single-wall carbon nanotubes measured in a four-point configuration with noninvasive voltage electrodes. The voltage drop is detected using multiwalled carbon nanotubes while the current is injected through nanofabricated Au electrodes. The resistance at room temperature is shown to be linear with the length as expected for a classical resistor. This changes at cryogenic temperature; the four-point resistance then depends on the resistance at the Au-tube interfaces and can even become negative due to quantum-interference effects.

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Transport measurements are a powerful technique to investigate electronic properties of molecular systems [1]. Most often, individual molecular systems are electrically attached to two nanofabricated electrodes. However, such two-point experiments do not allow the determination of the intrinsic resistance that results from scattering processes involving, e.g., phonons or disorder. Indeed, the resistance is mainly dominated by poorly defined contacts that lie in series. A solution to eliminate the contribution of contacts has been found with scanning probe microscopy techniques [2–6], which enable the measurement of resistance variations along long systems such as nanotubes, but these techniques have only been applied at room temperature. The standard method to determine the intrinsic resistivity of macroscopic systems is the four-point measurement. The application of this technique to molecular systems is challenging, however, since the electrodes used so far have been invasive. For example, nanofabricated electrodes were shown to divide nanotubes into multiple quantum dots [7].

We report four-point resistance measurements on single-wall carbon nanotubes (SWNTs) using multiwalled carbon nanotubes (MWNTs) as noninvasive voltage electrodes [Fig. 1(a)]. We find that SWNTs are remarkably good one-dimensional conductors with resistances as low as 1.5 k Ω for a 95 nm long section. The nanotube resistance is shown to linearly increase with length at room temperature, in agreement with Ohm's law. At low temperature, however, the resistance can become negative and the amplitude then depends on the resistance at the Au-SWNT interfaces. In this regime, four-point measurements can be described by the Landauer-Büttiker formalism taking into account quantum-interference effects.

Before discussing these measurements, we briefly review the basic physics of four-point measurements [Fig. 1(b)]. In the diffusive incoherent limit, the four-point resistance R_{4pt} of SWNTs, characterized by 4 conducting channels, is given by

$$R_{4pt} = \frac{h}{4e^2} \frac{L}{l_e} \quad (1)$$

with l_e the elastic mean-free path and L the separation between the voltage electrodes. Equation (1) describes the intrinsic resistance generated by the electronic backscattering along the nanotube.

Equation (1) requires the transmission to the voltage electrodes to be weak to suppress an additional resistance contribution that results from electrons entering the voltage electrodes. Such electrons are replaced by electrons that scatter into either direction of the tube, enhancing the resistance [8].

By introducing quantum-interference effects, four-point measurements can give striking results such as negative R_{4pt} . This is best seen using the Büttiker formula [9–11], which is convenient for the description of multiterminal conductors. The current I_α in each electrode is related to

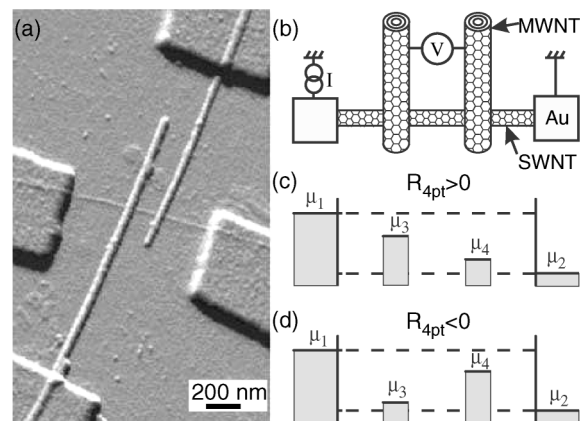


FIG. 1. (a) Atomic force microscopy image of a SWNT contacted by 2 MWNTs and 2 Au electrodes. (b) Schematic of the R_{4pt} measurement. (c), (d) Levels of the electrochemical potential for the 4 electrodes that give a positive R_{4pt} in (c) and a negative R_{4pt} in (d).

the electrochemical potential μ_β of other electrodes by $I_\alpha = 4e^2/h \sum_\beta T_{\beta\alpha} \mu_\alpha - T_{\alpha\beta} \mu_\beta$ with $T_{\alpha\beta}$ the total transmission between the α and the β electrodes [Figs. 1(c) and 1(d)]. The condition $I_3 = 0$ for a voltage probe gives $\mu_3 = (T_{31}\mu_1 + T_{32}\mu_2)/(T_{31} + T_{32})$. The transmission between electrodes 3 and 4 has been neglected since it corresponds to a second-order process. The potential of the voltage electrode μ_3 can thus take any value between μ_1 and μ_2 . Since the same holds for μ_4 , R_{4pt} can be negative [see Figs. 1(c) and 1(d)]. Using $R_{2pt} = (\mu_1 - \mu_2)/I$ and $R_{4pt} = (\mu_3 - \mu_4)/I$, we find that R_{4pt} takes any value between [10]

$$-R_{2pt} \leq R_{4pt} \leq R_{2pt}. \quad (2)$$

The remarkable prediction of $R_{4pt} < 0$ is difficult to observe experimentally. Some works on ballistic one-dimensional conductors fabricated in semiconductors showed that R_{4pt} can become slightly negative [12,13]. However, we believe that our data show for the first time significant negative R_{4pt} that approaches R_{2pt} .

We have fabricated nanotube circuits with a new layout for four-point measurements. First, ~ 1 nm diameter SWNTs grown by laser-ablation [14] or chemical-vapor deposition [15] are selected with atomic force microscopy (AFM). Voltage electrodes are then defined by positioning two MWNTs above the SWNT using AFM manipulation. We choose such voltage electrodes since the electric transmission between two crossed nanotubes is low [16,17]. Then, Cr/Au electrodes are patterned for electric connection with standard electron-beam lithography techniques [see Fig. 1(a)]. Characteristics of the different devices are summarized in Table I. Measurements are always in the linear-transport regime.

We first use Coulomb blockade (CB) measurements to investigate the degree to which the MWNT probes are invasive. Nanofabricated electrodes have been reported to create barriers along SWNTs and to divide tubes in multiple quantum dots [7]. In those previous measurements most of the CB peaks are completely suppressed. Figure 2 shows a series of CB peaks that appear regularly when sweeping the gate voltage V_g on the backgate. Peaks are at the same V_g for measurements between different pairs of

electrodes. This indicates that Au and MWNT electrodes probe the same quantum dot. Coulomb diamonds measurements (not shown) give that the charging energy $E_c \approx 5$ meV. It has been shown that $E_c \approx \frac{5 \text{ meV}}{L[\mu\text{m}]}$ for similarly prepared samples [18]. The dot length is thus $\sim 1 \mu\text{m}$, which is consistent with the 600 nm separation between Au electrodes. These measurements suggest that MWNTs are sufficiently noninvasive to not divide the SWNT in multiple quantum dots.

The question of invasiveness can further be tested by measuring the length dependence of R_{4pt} . Two types of measurements have been performed. In Fig. 3(a), an AFM tip has been used to change the separation between two MWNTs. In Fig. 3(b), 6 MWNTs have been placed on a long SWNT, enabling the resistance measurement of multiple portions. R_{4pt} linearly increases with the length, and R_{4pt} tends to zero as the length is reduced to zero. A significant resistance contribution from the MWNTs would give a finite R_{4pt} at zero length, in opposition to the measurements. This suggests that MWNTs are mainly noninvasive. Note, however, that measurements in Fig. 3(a) are rather scattered. This may come from the AFM manipulation that stretches the tube, deposits or removes some molecules adsorbed on the SWNT, or modifies the pressure applied by the MWNT on the SWNT. Thus, we cannot exclude a small resistance perturbation from the MWNTs and/or the AFM manipulation.

Our four-point measurement with no (or little) invasive voltage electrodes allows for the first direct measurement of the intrinsic resistance of a nanotube. The lowest resistance that we obtained is 1.5 k Ω for a 95 nm long section [Fig. 3(a)]. Such a low resistance is remarkable, since the resistance of quasi one-dimensional conductors is expected to be dramatically enhanced with the presence of disorder or phonons. This agrees with previous two-point measurements [19–22] since R_{2pt} was shown to approach the quantum resistance $h/4e^2 = 6.5$ k Ω , which suggests low intrinsic resistance.

Further insight into transport properties is obtained by decreasing the temperature T . Figure 4 shows that R_{4pt} does not change for T above ~ 80 K, suggesting that the intrinsic resistance is related to some static disorder and not to phonons. At lower T , the resistance becomes larger and

TABLE I. Device characteristics at 300 K. L is the separation between the MWNTs and $L_{\text{Au-Au}}$ between the Au electrodes.

	R_{4pt} (k Ω)	L (nm)	l_e (nm)	$L_{\text{Au-Au}}$ (μm)		
Device 1	1.5	95	408	2.7	SGS ^a	CVD
Device 2	37.0	100	17	5.6	SC ^b	LA
Device 3	2.7	150	358	1.0	metal	CVD
Device 4	6.3	140	143	0.6	metal	LA
Device 5	12.7	590	300	1.4	metal	LA

^aSmall-gap semiconductor with the current reduction occurring at $V_g > 2$ V.

^bLarge-gap semiconductor with the threshold voltage at ~ 40 V. LA = laser ablation. CVD = chemical vapor deposition.

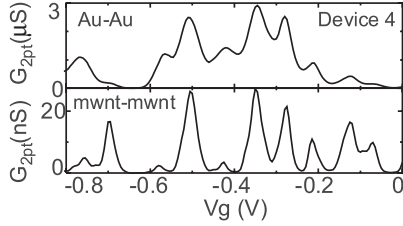


FIG. 2. Two-point conductance as a function of V_g at 1.4 K. CB peaks measured between different electrodes appear also at the same V_g 's for Devices 1 and 3.

starts to depend on the gate voltage. This will be discussed in more detail elsewhere.

Measurements above have shown that the four-point resistance is linear with length, in agreement with Ohm's law and Eq. (1) for a classical resistor. At 1.4 K this is, however, different, and R_{4pt} can even become significantly negative. The inset of Fig. 5(a) shows that R_{4pt} is $-29 \text{ M}\Omega$ near zero V_g . However, R_{4pt} significantly drops to 0 for V_g between -0.05 and -0.1 V when the two-point conductance between Au electrodes increases, as shown in Fig. 2. This is in agreement with Eq. (2). Figs. 5(a) and 5(b) show that the modulations of R_{4pt}/R_{2pt} are significant, with absolute values that reach as high as 0.6.

We now discuss the origin of the negative R_{4pt} . It might come from narrow diameter MWNTs or SWNT bundles [23,24]. The current pathway through different SWNTs may be complicated giving rise to negative R_{4pt} . However, such a classical effect should persist at higher T , which is not the case since $R_{4pt} > 0$ at $T \geq 10 \text{ K}$. Moreover, complicated current pathways give a finite non-local R at high T as recently shown [23]. We have not observed this for devices in this Letter.

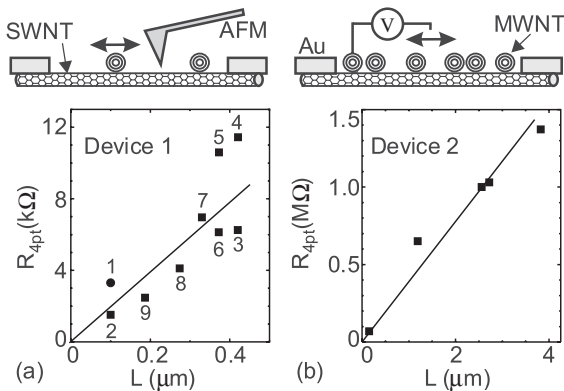


FIG. 3. Length dependence of R_{4pt} at room temperature and $V_g = 0$. (a) SWNT contacted by 2 MWNTs. One MWNT is displaced back and forth with an AFM tip. Points are numbered to describe the measurement sequence. Point 1 has been acquired one week before in the cryostat. Most points have been recorded while decreasing L , so that the R enhancement with L is not due to a structural degradation during the manipulation. (b) SWNT contacted by 6 MWNTs.

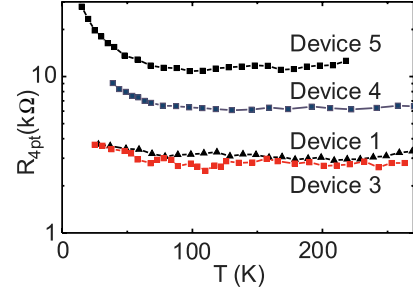


FIG. 4 (color online). Temperature dependence of R_{4pt} measured at $V_g = 0$.

The resistance ratio R_{4pt}/R_{2pt} is given by [10]

$$\frac{R_{4pt}}{R_{2pt}} = \frac{T_{31}T_{42} - T_{32}T_{41}}{(T_{31} + T_{32})(T_{41} + T_{42})}. \quad (3)$$

Part of the fluctuations are expected to come from the Coulomb blockade observed at 1.4 K that leads to oscillations in T_{ij} transmissions. However, regular $V_g = 75 \text{ mV}$ CB oscillations cannot alone account for the rapid $V_g \sim 10 \text{ mV}$ fluctuations of R_{4pt}/R_{2pt} in Fig. 5(a). We rather attribute those fluctuations to quantum-interference terms [25,26] that are contained in T_{ij} transmissions and that may arise from the superposition of different electronic paths between i and j electrodes [27]. Indeed, the disorder along the SWNT leads to different possibilities in the pathway between 2 electrodes [11]; see the inset of Fig. 5(b). It is also likely that the sign change originates from those interferences. Variations of T_{ij} transmissions with V_g are

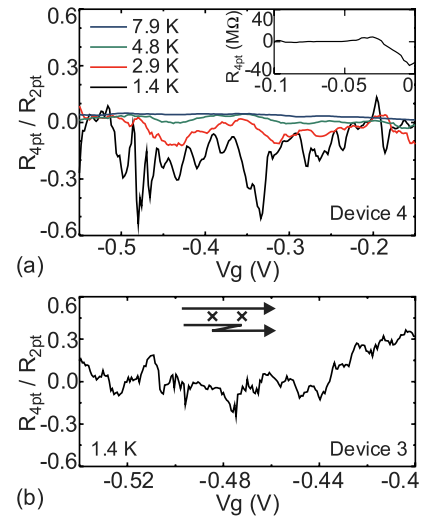


FIG. 5 (color online). Negative four-point resistance at low temperature. (a), (b) R_{4pt}/R_{2pt} as a function of V_g . Similar results are also obtained for Device 1. R_{2pt} measured between Au electrodes is checked to be lower than R_{2pt} between MWNT electrodes. The inset of (a) shows $R_{4pt}(V_g)$ at 1.4 K. R_{4pt} is close to 0 for V_g between -0.1 and -0.6 V . The inset of (b) shows two scattering centers that generate interference.

then uncorrelated, enabling the sign reversal of the numerator in Eq. (3).

Note that the SWNT may have entered the regime of strong localization (SL) [11]. This is expected for SWNTs when the phase-coherence length is longer than a few times l_e . This may be the case since l_e of Device 4 is 143 nm and negative R_{4pt} suggests coherence over at least $L = 140$ nm. However, similar R_{4pt} measurements are obtained for devices that are farther from the SL regime [Fig. 5(b), $l_e = 358$ nm]. Moreover, the two-point conductance modulation of Devices 1 and 3–5 is quite regular when V_g is swept (Fig. 2), in opposition to SL predictions [28]. We note that these devices are rather short (not much longer than l_e). This may explain that deviations from the diffusion regime due to SL are not significant.

So far, we have not taken into account the electron-electron interaction, which is expected to form a Luttinger liquid (LL) [17]. R_{4pt} of a LL has not been calculated yet. However, interference due to impurities in a LL should also lead to negative R_{4pt} .

We have shown the transition of R_{4pt} between the Ohm's law at 300 K and the quantum-interference regime at low temperature. The deviation from Ohm's law can become so dramatic that R_{4pt} is negative. Hence it is likely that inclusion of these quantum-mechanical interference effects will ultimately be required in the design of practical multi-terminal intramolecular devices.

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*Corresponding author.

Email address: bachtold@lpa.ens.fr

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