Ultrafast Optical Spectroscopy of Micelle-Suspended Single-Walled Carbon Nanotubes

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Abstract We present results of wavelength-dependent ultrafast pump-probe experiments on micelle-suspended single-walled carbon nanotubes. The linear absorption and photoluminescence spectra of the samples show a number of chirality-dependent peaks, and consequently, the pump-probe results sensitively depend on the wavelength. In the wavelength range corresponding to the second van Hove singularities (VHSs), we observe subpicosecond decays, as has been seen in previous pump-probe studies. We ascribe these ultrafast decays to intraband carrier relaxation. On the other hand, in the wavelength range corresponding to the first VHSs, we observe two distinct regimes in ultrafast carrier relaxation: fast (0.3-1.2 ps) and slow (5-20 ps). The slow component, which has not been observed previously, is resonantly enhanced whenever the pump photon energy resonates with an interband absorption peak, and we attribute it to radiative carrier recombination. Finally, the slow component is dependent on the pH of the solution, which suggests an important role played by H+ ions surrounding the nanotubes.

1 Introduction

Single-walled carbon nanotubes (SWNTs) provide a variety of new opportunities for the exploration of one-dimensional (1-D) quantum physics as well as novel device applications. Although their mechanical and electrical properties have been extensively studied during the past decade, there has been only limited success in exploring their optical, magnetic, and magneto-optical properties. In addition to what is commonly expected for excitons in 1-D systems [12,3], SWNTs are expected to show a new class of optical phenomena that arise from their unique tubular structure with varying diameters and chiral angles. Linear and nonlinear optical coefficients are expected to be diameter- and chirality-dependent, and an external magnetic field is expected to induce drastic modifications on their band structure via the Aharonov-Bohm phase [4,5,6,7]. Furthermore, nonlinear harmonic generation is expected to be highly selective for creating certain orders of high harmonics [8].

However, these predicted optical phenomena and properties have not been probed experimentally. This is because in standard production methods SWNTs appear in the form of bundles (or ‘ropes’) due to their strong van der Waals forces. This ‘roping’ results in significant broadening of electronic states, smearing out any chirality-dependent features in optical spectra [9,10,11,12,13,14,15]. Very recently, a new technique for producing individually suspended SWNTs has been reported [16]. These samples revealed, for the first time, a number of clearly observable peaks in linear absorption and photoluminescence (PL) spectra, corresponding to interband transitions in different types of tubes. A subsequent PL excitation (PLE) spectroscopy study successfully provided detailed peak assignments [17,18]. These seminal studies have provided opportunities to systematically study other predicted, unique optical properties of SWNTs.

Ultrafast laser spectroscopy is one of the best methods for measuring carrier distribution functions and relaxation mechanisms. Using a wide variety of techniques employing femtosecond pulses, one can directly examine dynamical processes after creation of electron-hole pairs across the band gap, i.e., carrier dephasing processes, carrier-carrier scattering, carrier-phonon scattering, exciton formation, and radiative and non-radiative recombination dynamics. There have been several ultra-
fast optical studies of SWNTs [19, 20, 21, 22]. Hertel and Moos [19] used ultrafast photoemission spectroscopy to determine thermalization times in metallic SWNTs to be \( \sim 200 \text{ fs} \). Three groups [20, 21, 22] independently observed subpicosecond to a ps lifetimes for semiconducting SWNTs in pump-probe spectroscopy. Although such ultrashort lifetimes can be expected for intraband relaxation towards the band edge, the microscopic origin of the fast interband decay (which is likely to be non-radiative) is still unclear. One important issue is that these studies were performed on bundled SWNTs, which did not show any PL. It is thus desired to carry out ultrafast spectroscopy on SWNT samples that show chirality-assigned absorption and PL peaks.

In this article, we report results of a femtosecond pump-probe study of chirality-assigned SWNTs. The linear absorption and PL spectra of such micelle-suspended SWNT samples show a number of chirality-dependent peaks, and as a result, the pump-probe data sensitively depends on the wavelength used. In the wavelength range corresponding to the second van Hove singularities (VHSs), we observe ultrafast (sub-picosecond) decays, as has been reported previously [20, 21, 22]. We ascribe these ultrafast decays to intraband carrier relaxation. On the other hand, in the wavelength range corresponding to the first VHSs, we observe two distinct regimes in ultrafast carrier relaxation: fast (0.3-1.2 ps) and slow (5-20 ps). The slow component has not been observed previously, is resonantly enhanced when the pump photon energy matches an absorption peak, and is attributed to radiative carrier recombination. The slow component is also strongly dependent on the pH of the solution, especially in large diameter tubes.

2 Samples Studied

The SWNTs studied in the present work were dispersed in aqueous sodium dodecyl sulfate (SDS) surfactant, sonicated, and centrifuged, which left micelle-suspended nanotube solutions. Details of the sample preparation method were described previously [16].

Typical linear absorption and photoluminescence (PL) spectra for such micelle-suspended SWNTs are shown in Fig. 1(b), together with a schematic density of states corresponding to the second van Hove singularities (VHSs), consisting of three overlapping bands: \( E_1 \), \( H_1 \) and valence (\( H_2 \)) subbands. The absorption spectrum shows peaks from the near-infrared (\( \sim 0.9-1.4 \text{ eV} \)) and are due to transitions involving the first conduction (\( E_1 \)) and valence (\( H_1 \)) subbands. The absorption spectrum shows peaks from the near-infrared to the visible range, consisting of three overlapping bands: \( E_1 H_1 \) transitions in semiconducting tubes (0.78-1.55 eV), \( E_2 H_2 \) transitions in semiconducting tubes (1.38-2.26 eV), and transitions in metallic tubes (2.07-3.11 eV). Detailed analyses and interpretations of these linear absorption/emission features are described in [17, 18].

3 Experimental Setup

We performed wavelength-dependent, degenerate pump-probe measurements using \( \sim 150 \text{ fs} \) pulses from an optical parametric amplifier (OPA) pumped by a chirped pulse amplifier (Clark-MXR CPA2010). We used a low pulse-repetition rate (1 kHz) for minimizing the average power and reducing any thermal effects while keeping the fluence high. To detect small photoinduced changes in probe transmission, we synchronously chopped the pump beam at 500 Hz and measured the transmission with (\( T \)) and without (\( T_0 \)) the pump using two different gates of a box-car integrator. The smallest detectable differential transmission \( \Delta T/T_0 \) was \( \sim 10^{-4} \). We used a noncollinear geometry with a pump beam diameter of \( \sim 220 \mu \text{m} \) in the overlap area. We tuned the OPA throughout the range of first subband transitions (i.e., \( E_1 H_1 \) transitions; see Fig. 1 with photon energies \( h\nu = 0.8 \text{ eV} \) to 1.13 eV (wavelengths \( \lambda = 1.1 \mu \text{m} \) to 1.55 \( \mu \text{m} \)). In addition, by directly using the CPA beam (\( h\nu = 1.60 \text{ eV} \), \( \lambda = 775 \text{ nm} \)), we probed a region of second subband transitions (\( E_2 H_2 \)).
4 Experimental Results

4.1 First vs. Second Subband Excitations

Figure 2 shows typical differential transmission ($\Delta T/T_0$) data as a function of time delay. Two traces are shown, taken at 0.89 eV (1393 nm) and 1.60 eV (775 nm), corresponding to $E_{1H_1}$ and $E_{2H_2}$ ranges, respectively (refer to Fig. 1). Both show a positive change (or an increase) in transmission, i.e., photoinduced bleaching, which is consistent with band filling. An exponential fit reveals a fast, single decay time of 770 fs for the $E_{2H_2}$ transition, which is consistent with earlier reports and can be explained by the very fast intraband carrier relaxation towards the band edge [22]. On the contrary, data in the range of $E_{1H_1}$ transitions exhibit multiple exponential decays. The major decay of the photoinduced signal happens in the first picosecond (with decay time $\tau_1$), which is followed by a much slower relaxation process. For the particular data shown in Fig. 2, we obtained an exponential decay time of $\tau_2 \approx 10$ ps. This long decay time has not been reported previously for either metallic [19] or semiconducting SWNTs [20,21,22].

4.2 Pump Fluence Dependence

For both first and second subband transitions, the pump fluence dependence of the maximum value of $\Delta T/T_0$ reveals clear saturation at high fluences, as shown in Fig. 3. This implies that, in the saturation regime, most of the carrier states are filled up and thus the sample absorption is nearly completely quenched. A careful analysis of the differential transmission decays for the $h\nu = 0.89$ eV case showed that relaxation dynamics are not dependent on the pump fluence, including the saturation regime. As an example, in Fig. 4 we show pump-probe data at a wavelength of 1402 nm for four different pump fluences. This precludes the possibility of any nonlinear recombination process such as the Auger recombination.
Fig. 5  (a) Linear absorption in the $E_{1}H_{1}$ range. The numbers (1-16) correspond to the 16 photon energies at which pump-probe measurements were made. (b) The peak value of $\Delta T/T_0$ (left axis) and the ratio of slow to fast components (right axis) vs. photon energy.

4.3 Resonant vs. Non-resonant Excitations

To study any differences between resonant and non-resonant excitations in carrier relaxation as well as any diameter- and/or chirality-dependent phenomena, we scanned the photon energy from 0.8 eV to 1.1 eV, corresponding to the $E_{1}H_{1}$ transitions of 0.82-1.29 nm diameter tubes [17]. For all the photon energies, the pump fluence was kept constant at 1 mJ/cm$^2$, which is below the saturation regime [see Fig. 3]. Figure 5(a) shows the linear absorption spectrum in the $E_{1}H_{1}$ transition range. The photon energies at which we performed pump-probe measurements are labeled 1 – 16, covering both peaks and valleys in absorption. Figure 5(b) shows the maximum value of $\Delta T/T_0$ as a function of photon energy; it loosely follows the absorption curve in (a). Also shown in Fig. 5(b) (right vertical axis) is the ratio of the slow component (defined as $\Delta T/T_0$ at 5 ps) to the fast component (defined as $\Delta T/T_0$ at 0 ps) as a function of photon energy; it also follows the absorption curve in (a), indicating that the slow component is resonantly enhanced at absorption peaks.

Fig. 6 Wavelength-dependent (one-color) pump-probe spectroscopy data, corresponding to the photon energies labeled #2, #3, #4, #6, #7, #8, #11, and #13 in Fig. 5(a), covering both absorption peaks and valleys. For the energies corresponding to absorption peaks [(a), (c), (e), and (g)], the chirality indices $(n,m)$ of the SWNTs probed at those energies are (a): (10,2), (c): (12,1) and (8,6), (e): (10,3) and (10,5), and (g): (11,4).
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5 Discussion

As we presented in the last section, our wavelength-dependent single-color pump-probe spectroscopy data on micelle-suspended SWNTs revealed two different decay components, suggesting two different carrier relaxation processes. One component is fast (0.3-1.2 ps) and the other is slow (5-20 ps). In the following we discuss possible origins of these two dynamical processes as well as the intriguing pH dependence of the pump-probe data we obtained.

Both the positive sign of the differential transmission and the initial ultrafast (~1 ps) relaxation agree with the recent reports for semiconducting SWNTs [20,21,22]. The positive sign can be interpreted as state filling as the cause of the photo-bleaching signal and is consistent with the saturation at high fluences (Fig. 3). Namely, the pump-induced carriers fill states and reduce the probe absorption. We believe that the ultrafast decay mechanism is non-radiative (i.e., phonon and/or impurity-mediated) and intraband. It is likely to be non-radiative because this signal exists even for samples that do not luminesce [20,21,22]. It is likely to be intraband relaxation (as opposed to interband relaxation) since it exists both in resonant and non-resonant cases. Note that even when the photon energy is resonant with the \(E_{2H_1}\) absorption peak of a particular tube type, it creates non-resonant carriers in other types of tubes whose band gaps are smaller than that of the resonant tube. Note also that we are not seeing any pump-power-dependent decay times, as shown in Fig. 4 which precludes carrier-density-dependent non-radiative processes such as Auger...
recombination as the origin of the fast decay component. In addition, it is likely that there is some coherent contribution to the pump-probe signal in this ultrashort time scale. However, since we have not observed any four-wave mixing signal, we do not have an estimate on the dephasing time and thus will not discuss how large this contribution is.

On the other hand, the slow decay signal, which was resonantly enhanced when the photon energy coincided with an interband absorption peak, has not been reported previously. We interpret this slow component as interband carrier recombination. Note that in bundled samples band edges are not well defined for individual tubes due to electronic coupling, and thus photo-created carriers cannot show clear interband dynamics in pump-probe measurements. In other words, there is no clear distinction between inter-tube dynamics and intraband dynamics in bundled tubes because van Hove singularities of different tube types form a van Hove absorption band. We further argue that the slow component we observed is probably related to radiative recombination. An important fact for supporting this claim is that in the previous work no PL was observed whereas our sample exhibits PL peaks at the same energies as EtH1 absorption peaks (see Fig. 3(a)). In addition, reducing the pH of the solution destroys PL and absorption peaks while at the same time the slow decay also vanishes (see Fig. 4(a)), indicating an intimate relationship between PL and the slow decay signal. The actual value of the radiative recombination lifetime $\tau_r$ depends on the value of the radiative efficiency (or quantum yield) $\eta = \tau_r^{-1}/(\tau_r^{-1} + \tau_{nr}^{-1})$, where $\tau_{nr}$ is the nonradiative lifetime, since what we measure experimentally is the total decay rate, $\tau^{-1} = \tau_r^{-1} + \tau_{nr}^{-1}$. Direct measurements of $\tau_r$ through time-resolved PL are in progress.

Finally, we propose possible scenarios for the drastic pH dependence we observed. Decreasing the pH leads to an increase in the density of H$^+$. First, adsorbed H$^+$ ions on the nanotube surface could add a relaxation channel by the creation of ultrafast carrier trapping centers (defects). This should make the fast nonradiative recombination dominant over the slower radiative recombination if the density of such trapping centers is high. In addition, as previously noted, H$^+$ ions effectively act as acceptors in SWNTs, and thus the Fermi energy ($E_F$) depends on the H$^+$ density. When the density is so high that $E_F$ lies inside the valence band, interband absorption peaks disappear, irrespective of whether the peaks are due to excitons or van Hove singularities. This can be viewed as a 1-D manifestation of the well-known Burstein-Moss effect. Larger-diameter tubes are expected to have smaller acceptor binding energies due to their smaller effective masses, and thus should be more susceptible to pH changes. In a heavy doping regime, there is a degenerate hole gas in the valence band, which can completely quench excitonic processes.

### 6 Summary

In summary, we have carried out an ultrafast optical study of micelle-suspended single-walled carbon nanotubes. We have observed two relaxation regimes in the dynamics of micelle-suspended single-walled carbon nanotubes under resonant excitations. We interpret the previously observed shorter decay time $\tau_1$ as nonradiative intraband relaxation and the previously unobserved larger decay time $\tau_2$ as related to radiative interband recombination. The relaxation of photoexcited carriers can be made faster by increasing the density of H$^+$ ions in the solution while simply increasing the number of photoexcited carriers does not change the dynamics. The sensitive pH dependence provides a novel means to chemically control carrier states and dynamics in nanotubes.
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