Temperature dependent separation of metallic and semiconducting carbon nanotubes using gel agarose chromatography

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ABSTRACT

Post-synthesis separation of metallic (m-SWNTs) and semiconducting (s-SWNTs) single-wall carbon nanotubes (SWNTs) remains a challenging process. Gel agarose chromatography is emerging as an efficient and large scale separation technique. However, the full (100%) separation has not been achieved yet, mainly due to the lack of understanding of the underlying mechanism. Here, we study the temperature effect on the SWNTs separation via gel agarose chromatography, for four different SWNT sources. Exploiting a gel agarose micro-beads filtration technique we achieve up to 70% m-SWNTs and over 90% s-SWNTs, independent of the source material. The process is temperature dependent, with yields up to 95% for s-SWNT (HiPco) at 6°C. Temperature affects the sodium dodecyl sulfate surfactant-micelle distribution along the SWNT sidewalls, thus determining the effectiveness of the SWNTs sorting by electronic type. The sorted SWNTs are then used to fabricate transistors with very low OFF-currents (~10^{-13} A), high ON/OFF current ratio (>10^6) and charge carriers mobility ~ 40 cm^2 V^{-1} s^{-1}.

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1. Introduction

Carbon nanotubes (CNTs) attract much attention in view of potential applications that can exploit their transport properties [1], ranging from large area networks [2], to sensors [3] and individual SWNT devices [4–7], just to cite a few.

However, these often require SWNTs of specific electronic type: either metallic (m-SWNTs) or semiconducting (s-SWNTs). To date, the vast majority of synthesized SWNTs are composed of a mixture of the two electronic species. Growth of pure (100%) m-SWNTs or s-SWNTs is still being pursued. Refs [8–16] reported SWNT enrichments based on electronic type [8–12] and specific chiralities [15,16]. However, achieving on-demand control of SWNTs type is still far from being solved. The rise of graphene [17–19] and other two-dimensional (2d) crystals [19–22], has added further pressure to realize the promise of SWNTs in electronic devices, since 2d crystals do not require a sorting process to define their electronic nature (although
some 2D crystals, e.g. MoS₂, show indirect band gap in bulk and direct in monolayer configuration [20,21], a step otherwise necessary in the case of SWNTs [23–27]. On the other hand, selectivity of un-sorted SWNTs is not always required and, e.g., the heterogeneity of un-sorted SWNTs can be exploited in ultrafast ac-dielectrophoresis [23], DNA wrapping [24], selective breakdown of m-SWNTs [25], density gradient ultracentrifugation (DGU) [26,31,32], amine extraction [33] and gel agarose electrophoresis [27] have been exploited for such goal. These separation strategies have been reviewed in detail in Refs. [34,35]. Although DGU is by far the most complete approach, and to date has permitted the separation by length [36], diameter [37], metallic vs. semiconducting [26], chirality [31], and handedness [38], it is time and resource consuming, and has low throughput at laboratory scale [37,39]. On one hand it is possible to scale-up the DGU process and improve yield using large capacity centrifuges, but the costs involved are high. Routinely, DGU separations require up to 20 h [26,37,32] centrifugation time, which reflects on energy consumption. Other separation methods based on chemical approaches [40] involve irreversible covalent functionalization [41] with chemicals or biological compounds that not only affect the intrinsic SWNT properties [34,42,43], but may also have health and environmental impacts [44,45].

Agarose gel chromatography has emerged as a promising alternative to DGU. Ref. [46] reported that agarose gel beads can be exploited for ms-separation, due to simplicity, low cost, short process time and scalability [46]. The processing time for gel agarose beads separation is ~20 min [34] compared to ~10–20 h [26,37] needed for the DGU. The agarose beads filtration requires only a single step and, as in the case of DGU [26,31], the SWNT’s dispersion is in aqueous surfactant solutions. Moreover, the separation purity level (i.e. the percentage of m- and/or s-SWNTs) is now comparable (e.g. ~90% for (6,5), (7,6), (8,6) [47] with DGU ~95% both for m-SWNTs [31] and s-SWNTs [26,27]). It is possible to achieve purities up to ~99% by multiple agarose gel filtration iterations [48], but, also in this case, the process is time and resource heavy. Although the mechanism of separation by hydrogels (e.g., agarose and sephacryl) was demonstrated to be an entropy-driven process [49], it is not fully understood yet. Furthermore, there are still unexplored process parameters, such as the temperature, which is crucial in order to optimize purity.

Here, we report ms-separation of SWNTs from four different sources using gel agarose beads filtration. We investigate the effect of temperature, showing that this affects the purity of sorted SWNTs. The separation is shown to be influenced by the surfactant micelles formation around the SWNTs’ surface, which in turns is affected by temperature. We demonstrate that there is a trade-off between the purity the enriched s-SWNTs and m-SWNTs fractions for separation carried out at varying temperature. The percentage of s-SWNTs is the highest at low temperature (~6 °C) and gradually drops as the temperature is increased (~50 °C), whereas for m-SWNTs, the reverse is observed. To demonstrate a direct application, the enriched s-SWNTs are then used to fabricate CNT-based field effect transistors (CNT-FETs) with superior performance in terms of ON/OFF current ratio (>10⁶) with respect to devices based on un-sorted SWNTs (<2).

2. Experimental methods

2.1. Nanotube sources

Four types of SWNTs are used. The first is prepared by enhanced direct injection pyrolytic synthesis [50,51] (DIPS-CNT). These have diameters in the range 0.8–3 nm, and are obtained from Nikkiso Corp. Japan. The second is arc-discharge SWNT (AD-CNT) from Nanocarblab, with diameters in the 1.2–1.4 nm range [52]. The third is CoMoCAT SWNTs, with diameter distribution between 0.7 and 0.9 nm [53,54]. The fourth is high pressure carbon monoxide (HiPco) SWNTs [55], with 0.8–1.2 nm diameter [55]. All SWNTs are from raw material, except for DIPS–CNT, which is purified by the supplier and contains >90% SWNTs.

2.2. Dispersion

SWNT dispersions are prepared by adding 1 mg/10 ml weight to volume ratio of each SWNT source to an aqueous solution of 2% sodium dodecyl sulfate (SDS, Sigma–Aldrich) in deionized water (DIW). De-bundling is obtained via ultrasonication using a Cole Parmer Ultrasonic Processor for 4 h (40% of 750 kW at 20 kHz). The dispersions are then ultracentrifuged using a Beckman L8–70M Ultracentrifuge at 200,000 g for 20 min at 18 °C to remove SWNT bundles and other impurities, such as amorphous carbon, catalyst residual etc [31]. The supernatant is collected using pipettes and used for the gel filtration separation process.

2.3. Separation process

Disposable 2.5 ml syringes, used as continuous separation columns, are plugged with cotton, and filled with 1.5 ml agarose micro-beads suspended in ethanol (Sepharose 2B, 2% Agarose, 60–200 μm, Sigma–Aldrich). The syringe column is then washed and equilibrated with the surfactant aqueous solution by pouring 5 ml of 1% SDS in DIW into it and letting the solution pass through the agarose gel beads to remove the ethanol. ~0.5 ml SWNT dispersion is then poured into the column. Subsequently, 1.5 ml 1% SDS solution is added. This causes a displacement of the SWNT dispersion along the column. A portion of the SWNTs are trapped at the top layer of the agarose beads, creating the top band, and another portion moves down the column, forming the bottom band, as for Fig. 1. The bottom band is eluted from the column when an additional 1 ml of 1% SDS is added, and then collected for spectroscopic characterization. Depending on the SWNT source, the top and bottom bands formed in the column have...
Optical absorption spectra (OAS) reveal various properties of SWNT dispersions. The characterization is carried out by optical absorption, photoluminescence excitation (PLE), and Raman spectroscopy. The assignment of the optical transitions is based on the empirical Kataura plot of Ref. [64]. This gives values of optical transition frequencies as a function of chirality for SWNTs in aqueous surfactant dispersions, and is more appropriate than Kataura plots theoretically derived from tight binding and other models [64,66]. We use the empirical Kataura plot of Ref. [64] also for chirality assignment in PLE and Raman spectroscopy.

2.4. Characterization

The characterization is carried out by optical absorption, photoluminescence excitation (PLE), and Raman spectroscopy.

2.5. Optical absorption spectroscopy

Optical absorption spectra (OAS) reveal various properties of SWNT dispersions, such as transition energies [56–68], bundling [57–59], and concentration [60]. The determination of the relative concentration of chiralities is carried out taking the derivative of absorption, as for Refs. [26,31] to determine the peak absorption. For quantitative analysis, if we assume that the absorption spectrum obeys the Beer–Lambert law [61], $A_k = \varepsilon_k \frac{d \lambda}{C}$, where $A_k$ is the absorbance at wavelength $\lambda$ [nm], $\varepsilon_k$ [L mol$^{-1}$ cm$^{-1}$] the absorption coefficient at the same wavelength, $l$ [cm] the length of optical path and $C$ [mol L$^{-1}$] is the concentration of material, there will be a similar relation between concentration and absorbance for the first-order derivative: $\frac{dA_k}{dC} = \varepsilon_k \frac{dx}{dC}$.

Fig. 1 – Syringe column filled with agarose gel beads. The example shown here is the separation of a HiPco sample. (A color version of this figure can be viewed online.)

The OAS are acquired in a Varian Cary 5000. Measurements are carried out in the range 400–1300 nm, limited by the absorption of water. This range is sufficient to cover the first and second excitonic transitions of s-SWNTs [56,62,63] and the first transition of m-SWNTs, for CoMoCAT and HiPco [36,62,63]. For DIPS–CNT and AD-CNT, this covers the second and third transitions of s-SWNTs and first of m-SWNTs [64,65]. Absorption from solvent and surfactants is subtracted, by measuring solutions with only solvent and surfactant.

The assignment of the optical transitions is based on the empirical Kataura plot of Ref. [64]. This gives values of optical transition frequencies as a function of chirality for SWNTs in aqueous surfactant dispersions, and is more appropriate than Kataura plots theoretically derived from tight binding and other models [64,66]. We use the empirical Kataura plot of Ref. [64] also for chirality assignment in PLE and Raman spectroscopy.

2.6. Photoluminescence excitation spectroscopy

Photoluminescence excitation spectroscopy (PLE) is one of the most used techniques to monitor SWNT dispersions [66–68]. The $(\epsilon h_i, \epsilon h_1)$ resonances ($i = 1, 2, \ldots$ etc) from different SWNTs appear as sharp features $(\lambda_{ex}, \lambda_{em})$, where $\lambda_{ex}$ and $\lambda_{em}$ are the excitation and emission wavelengths, respectively. Other peaks can be observed, either due to excitonic-phonon sidebands [69], exciton energy transfer (EET) [57,70] or bright phonon sidebands (BS) of dark K-momentum excitons [71].

In general, the PL intensity is proportional to the concentration of a species, its absorption cross-section [72] at the excitation wavelength [73], and its fluorescence quantum yield [74]. Thus, it does not directly reveal the relative abundance of SWNTs [73]. However, the relative PL intensity of different chiralities can be used to compare the separation process at different temperatures. The PLE maps are recorded using a Horiba Jobin-Yvon excitation–emission spectrophotometer (Fluorolog3) with 10 nm slits for the double grating excitation monochromator and 14 nm for the single grating emission. The scan step is 5 nm for excitation, with a range from 440 to 850 nm for CoMoCAT and HiPco SWNTs, and from 350 to 700 nm for DIPS–CNT samples, respectively. The emission is collected by a liquid-nitrogen-cooled InGaAs detector using a right angle scattering mode, in the ranges: 900–1200 nm for CoMoCAT, 850–1450 nm for HiPco and 1250–1550 nm for DIPS–CNT. PLE measurements are not carried out for the AD-CNTs due to their large diameter distribution that prevents the emission to be detected by our spectrometer. Note that $\sim 1600$ nm is our upper detection limit. This restricts the detection of PL from SWNTs with diameter larger than $\sim 1.3$ nm [64]. The raw PL data are corrected by the excitation profile.

2.7. Raman spectroscopy

Raman spectroscopy can be used to probe SWNTs within dispersions. In the low frequency region, the Radial Breathing Modes (RBMs) are observed [75]. Their position, $\text{Pos}$(RBM), is inversely related to SWNT diameter, $d$ [76–78], as given by...
Pos(RBM) = $\frac{d}{\lambda} + C$. Combining Pos(RBM), with excitation wavelength and the ‘Kataura plot’ [64,65], it is, in principle, possible to derive the SWNT chirality [79,80].

Matching the diameter given by Pos(RBM) with excitation wavelength in the Kataura plot also gives information on the s- or m- character. A variety of $C_1$ and $C_2$ were proposed for this relation [25–81]. Here we use the $C_1 = 214.4 \text{ cm}^{-1} \text{nm}$ and $C_2 = 18.7 \text{ cm}^{-1}$, from Ref. [76]. These were derived by plotting the resonance energy as a function of inverse RBM frequency without additional assumptions. However, we also validated our results by using the parameters proposed in Refs. [77,81,82]. Raman spectroscopy also probes possible damage via the D peak [83–85]. The typical Raman spectrum in the 1500–1600 cm$^{-1}$ region consists of the G$^+$ and G$^-$ bands.

In s-SWNTs, they originate from the longitudinal (LO) and tangential (TO) modes, respectively, derived from the splitting of the $E_{2g}$ phonon of graphene [85]. The positions of the G$^+$ and G$^-$ peaks, Pos(G$^+$), Pos(G$^-$), are diameter dependent and the separation between them increases with decreasing diameter [86,87]. In m-SWNTs, the assignment of the G$^+$ and G$^-$ bands is the opposite, and the full width at half maximum of the G$^+$ peak, FWHM(G$^+$), is larger and Pos(G$^-$) down-shifted with respect to the semiconducting counterpart [86,88]. Thus, a wide, low frequency G$^-$ is a fingerprint of m-SWNTs. On the other hand, the absence of such a feature does not necessarily imply that only s-SWNTs are present, but could signify that m-SWNTs are off-resonance.

Doping could also modify positions and FWHMs [89,90]. In m-SWNTs, a Pos(G$^+$) blueshift, accompanied by a FWHM(G$^-$) decrease, is observed with electron or hole doping [91,92]. In s-SWNTs doping upshifts Pos(G$^+$), but does not affect FWHM(G$^+$) [89,91].

A large number of excitation wavelengths are in principle necessary for a complete characterization of SWNTs [78,81]. Nevertheless, useful information can be derived even with few excitations, especially for process monitoring, when Raman spectroscopy is used to compare the “raw” material with end product.

Raman spectra are taken both on the un-sorted and sorted dispersions deposited on a Si substrate, with a Renishaw system at 514 nm (2.41 eV), 633 nm (1.96 eV) and 782 nm (1.58 eV), using a 50X objective and less than 1 mW on the sample. The RBM detection is limited by the cut-off of the notch and edge filters. These are at 120, 100 and 110 cm$^{-1}$ for 514, 633 and 782 nm, respectively, limiting the detection of tubes with diameter up to $\sim 1.9$ nm. We use Lorentzians to fit the RBM, and G$^+$ and G$^-$ peaks.

2.8. CNT-based transistors

After separation, the SWNTs are used for CNT-FETs. The devices fabricated from each SWNT samples are based on a spin coating enriched s-SWNTs dispersion to form a network of SWNTs on n-type Si substrate (acting as the device gate) covered with 250 nm thermally grown SiO$_2$ as gate dielectric. After annealing at 200 °C for 20 min, Au electrodes are patterned, via electron beam lithography, to form the source and drain. The devices are then annealed again to evaporate residual solvents at 200 °C for 5 min, and characterized with a Keithley 4200 Analyzer in ambient conditions (both pressure and temperature).

3. Results and discussion

3.1. Room temperature separation

The optical absorption data of the four different sets of SWNT before and after the separation process at RT is shown in Fig. 2. The absorption peaks are denoted as $M_{11}$, $eh_{11}(S_{11})$, $eh_{22}(S_{22})$, and $eh_{33}(S_{33})$, and assigned to the first transition of m-SWNTs and the first, second and third excitonic transition of s-SWNTs, respectively [64,65]. The spectrum for the unsorted DIPS–SWNTs (Fig. 2a) shows broad bands for the $M_{11}$, $S_{22}$, and $S_{33}$ peaks due to the “nominally” large diameter distribution [93]. Moreover, the lack of sharp features for the excitonic transitions of both m- and s-SWNTs is a signature of bundles [59]. Contrary, the top and bottom bands of the sorted samples eluted through the column show more intense and “sharp” absorption features. The top band has an increased absorption in the $eh_{22}$ and $eh_{33}$ region. These bands are well resolved, unlike the spectrum of the starting material. We estimate, by using the derivative of the absorption, $\sim$76% s-SWNTs, the rest being m-SWNTs, with absorption peaks at $\sim$801 and 865 nm, that we assign [65] to (13,10) and/or (17,10) and (17,8) and/or (22,1) tubes.

Better resolved, compared to the DIPS–CNT samples, absorption spectra are obtained for the AD-CNT tubes in Fig. 2b. The top fraction shows an increase of the absorption peaks corresponding to $eh_{11}$ in the 800–1100 nm range, with respect to the starting material. We assign [65] the peaks at $\sim$655, $\sim$700 and $\sim$747 nm detected in the $M_{11}$ region, to (10,10), (13,7) and/or (12,9), (19,1) and/or (18,3), respectively. By analyzing the derivative of the absorption we get $\sim$90% s-SWNTs in this fraction. The spectrum of the bottom fraction has increased absorption in the region 550–800 nm, with a broad and not resolved band also detected in the region 800–1200 nm, due to $eh_{22}$.

The OAS of HiPco (Fig. 2c) and CoMoCAT (Fig. 2d) SWNTs have sharper peaks than the DIPS and AD-CNT samples (both for starting material and sorted samples). The top band of sorted HiPco SWNTs (blue curve in Fig. 2c) has a reduced absorption in the $M_{11}$ region and an enhancement of the peaks in the $eh_{11}$ region with respect to the starting material (black curve in Fig. 2c). The peaks in the region 400–600 nm are assigned [64,65] to the $eh_{22}$ of (7,3), 505 nm; (9,2), 552 nm; (6,5), 566 nm; (8,4), 591 nm; (11,1), 611 nm; and $M_{11}$ of (6,3) and (5,5) at 440 and 411 nm, respectively. There is also an increase in $eh_{11}$ absorption with respect to the starting material.

The calculated percentage of s-SWNTs in this fraction is $\sim$90%, with 10% m-SWNTs. The spectrum of the bottom fraction shows an increase in absorption in the region 400–600 nm. We assign [65] the peaks to the following m-SWNTs: (5,5), 411 nm; (6,4) and (7,6) 455 nm; (7,7) and (9,0) 507 nm; (8,8) and (11,2) 558 nm; (9,9), 598 nm; (10,10), 653 nm; (11,11), 733 nm.

However, some of the peaks are quite broad, being the results of a combination of $M_{11}$ and $eh_{22}$ transitions. Indeed,
absorption features from s-SWNTs are detected in the $e_{11}$ region (800–1300 nm), with broad absorption peaks associated to $(8,3)\ 952\ \text{nm};\ (6,5)\ 983\ \text{nm};\ (7,3)\ 990\ \text{nm};\ (7,5)\ 1020\ \text{nm};\ (8,1)\ (10,2)\ (8,4)\ (8,6)$ around 1020 nm; other larger diameter SWNTs at $/C24 1150\ \text{nm}$[64,65]. The estimated percentage of s- and m-SWNTs in the bottom fraction is $/C24 38\%\ and\ 62\%$, respectively.

Similar results to the HiPco one are obtained with the CoMoCAT SWNTs (Fig. 2d). The top band has a reduced $M_{11}$ absorption and an enhancement in the $e_{11}$ and $e_{22}$ regions with respect to the starting material. In particular, there is a reduction of $(6,6)$ absorption with respect to the un-sorted sample. The bottom band shows an increase of absorption in the region 400–550 nm, with peaks assigned to $(6,6)$ and $(7,4)$ at 458 nm; $(7,7)$ and $(9,9)$ at 507 nm; $(9,9)$ at 598 nm; $(10,10)$ at 658 nm; $(11,11)$ at 733 nm [65]. However, these bands are quite broad due to the combination of $M_{11}$ and $e_{22}$. s-SWNTs are detected via the $e_{11}$ absorption in the region 800–1300 nm. We estimate that in the top fraction there are $\sim 78\%$ s-SWNTs, while in the bottom fraction there is an enrichment of m-SWNTs of $\sim 65\%$.

The Raman spectra of un-sorted and sorted samples are analyzed and compared in Figs. 3–6. Fig. 3a and b plots the spectra at 514.5 nm for DIPS. This wavelength is expected to be in resonance mostly with the third excitonic transition of s-SWNT [64,65]. The spectrum in the RBM region of the starting material shows features related to m- and s-SWNTs [76]. In particular RBMs of s-SWNTs $(16,6),\ (13,8),\ (14,7),\ (13,5),\ (15,2)$ and m-SWNTs $(8,5)$ and $(9,3)$ are detected. After sorting, we still detect the RBMs of s-SWNTs in the top band. On the contrary, in the bottom band we detect only RBM assigned to $(8,5)$ and $(9,3)$. In the $G$ peak region (Fig. 3b), the un-sorted sample shows the typical $G^+$ and $G^-$ peaks expected from a mixture of s-and m-SWNTs [76,87,94]. FWHM($G^+$) of the top band is $\sim 47\%$ narrower with respect to the un-sorted sample. FWHM($G^-$) of the bottom band is $\sim 28\%$ and $\sim 89\%$ larger with respect to the un-sorted sample and the top band, respectively. The value of FWHM($G^-$) is an indication of higher s- and m-SWNTs content in the top and bottom bands, respectively. Doping could also modify the peaks’ shape [89,91]. We do not expect doping to play a major role here for two reasons. First, both samples contain the same surfactants, even if with different concentration, so the doping contribution should be similar for both. Second, Pos($G^-$) should upshift and become narrower as a consequence of both p or n doping [89,91].
Fig. 3 – Raman spectra of DIPS CNTs. (a) RBM and (b) G region measured at 514.5 nm, (c) RBM and (d) G region measured at 633 nm and (e) RBM and (f) G region measured at 782 nm. A peak from the Si substrate is marked with * in (c). (A color version of this figure can be viewed online.)
Fig. 4 – Raman spectra of AD-CNT SWNTs. (a) RBM and (b) G region measured at 514.5 nm, (c) RBM and (d) G region measured at 633 nm and (e) RBM and (f) G region measured at 782 nm. A peak from the Si substrate is marked with *. (A color version of this figure can be viewed online.)
Fig. 5 – Raman spectra of HiPco SWNTs. (a) RBM and (b) G region measured at 514.5 nm, (c) RBM and (d) G region at 633 nm and (e) RBM and (f) G region measured at 782 nm. (A color version of this figure can be viewed online.)
Fig. 3c and d plot spectra at 633 nm. The RBM region (Fig. 3c) shows an enhancement of the peak associated with the (13,4) s-SWNT in the top fraction. This peak is reduced in intensity in the bottom fraction. In the same fraction there is also an increase in intensity of the peaks assigned to (14,2), (15,0) and (13,1). The FWHM($G^*$) of the top fraction
(23.4 cm$^{-1}$, Fig. 3d) is $\sim$70% narrower with respect to the bottom band, which could indicate a higher percentage of s-SWNTs.

Evidence of ms-separation is seen also at 782 nm. While the RBM region (Fig. 3e) of the un-sorted materials shows peaks associated both to s- and m-SWNTs, in the top fraction there is a decrease in intensity of peaks at $\sim160$ cm$^{-1}$, associated to (19,1) and (18,3), and an enhancement of peaks in the region 200–235 cm$^{-1}$, due to (14,1), (9,7), (8,7), (11,3) and (12,1) [65,76]. The latter peaks, observed both in the starting material and the top band, have very low intensity in the bottom fraction.

Fig. 3e also indicates the presence of m-SWNTs such as (19,1) and (18,3). In the G peak region (Fig. 3f), the FWHM of the top band is $\sim$66% narrower with respect to the un-sorted sample. While FWHM(G$^+$) for the top band is $\sim$26% and $\sim$106% larger with respect to the un-sorted sample and the top band, respectively. This indicates higher s- and m-SWNTs content in the top and bottom band, respectively.

Fig. 4a and b plots the spectra at 514.5 nm for AD-CNTs. These are not significantly different. This is due to the resonant excitation of s-SWNTs at this wavelength [65,76]. The Raman results agree with the absorption data shown in Fig. 2b. Indeed, the spectrum of the bottom band has FWHM(G$^+$) only $\sim$3 cm$^{-1}$ larger than the top band, too small to be statistically significant of a higher m-content. Similar spectra are also measured at 633 nm. However, at this wavelength, mostly m-SWNTs are in resonance. The RBM region (Fig. 4c) of the bottom fraction shows an enhancement of the peaks at $\sim200$ cm$^{-1}$ (assigned to m-SWNTs) with respect to the un-sorted sample and the top band, respectively. In the G peak region (Fig. 4d), the FWHM(G$^+$) of the top band sample is $\sim$35% narrower than the un-sorted sample and the bottom band. The Pos(G$^+$) is also $\sim$10 cm$^{-1}$ (1559 cm$^{-1}$) upshifted with respect to the un-sorted sample (1550 cm$^{-1}$) and bottom band (1548 cm$^{-1}$). Evidence of ms-separation is seen at 782 nm. The RBM region (Fig. 4e) in the top fraction shows an intensity reduction of the peaks $\sim165$ cm$^{-1}$, due to (15,6), and (12,8) [65,76], while the peaks at $\sim210$ and $\sim238$ cm$^{-1}$ due to s-SWNTs are enhanced in the top band and decrease ($\sim210$ cm$^{-1}$) or disappear ($\sim238$ cm$^{-1}$) in the bottom band. FWHM(G$^+$) is broader ($\sim$24%) in the bottom band, with respect to both the un-sorted sample and the top band, fingerprint of m-separation, see Fig. 4f.

The Raman spectra for HiPco samples are reported in Fig. 5. The spectra in the RBM region at 514.5 nm after enrichment are rather different with respect to the un-sorted samples (Fig. 5a). The top band shows peaks in the range 170–215 cm$^{-1}$, while these have a reduced intensity in the bottom band. These peaks are mainly assigned to s-SWNTs: (11,9), (16,2), (10,9) and (14,3). The top band shows a decrease in intensity of the peaks in the range 250–320 cm$^{-1}$. In particular, no peak is detected at $\sim316$ cm$^{-1}$, assigned to (8,2) [65,76], in the top fraction, while this is enhanced with respect to the un-sorted sample in the bottom band.

In Fig. 5b, the FWHM(G$^+$) of the top band is $\sim$60% narrower with respect to the un-sorted sample and the bottom band. Moreover, Pos(G$^+$) is $\sim$22 cm$^{-1}$ upshifted with respect to the bottom band. This is an indication that the contribution of Pos(G$^+$) in the top band is the combination of m- and s-SWNTs [91], the latter having a diameter $\sim$1.3 nm [86]. The bottom band has Pos(G$^+$) $\sim$1543 cm$^{-1}$ with main contributions from m-SWNTs with 0.7–0.9 nm diameter [86]. Fig. 5c and d shows Raman spectra measured at 633 nm. At this excitation both m- and s-SWNTs are expected to be in resonance [65]. From Fig. 5c, m-SWNTs are detected in the 150–225 cm$^{-1}$ range, while s-SWNTs in the range 240–350 cm$^{-1}$. The spectra of the bottom band show an increase in the RBMs intensity at $\sim$150–225 cm$^{-1}$, in particular at $\sim$190 cm$^{-1}$, assigned to (12,3) [65,76], and a reduction of the RBM peaks in the 240–350 cm$^{-1}$ region with respect to the un-sorted sample and the top band. The latter has the opposite behavior, with an enhancement of the peaks assigned to s-SWNTs. In the G peak region (Fig. 5d), evidence of splitting of the G$^+$ peak is seen in all spectra. The peak at lower wavenumber ($\sim$1542 cm$^{-1}$), due to the LO mode of metallic tubes with diameter $\sim$1.0–1.1 nm [86], has the same position in the three spectra, but the FWHM is $\sim$63% larger in the bottom band with respect to the starting material. The peak at $\sim$1555 cm$^{-1}$, due to a combination of LO of m-SWNTs with diameter of $\sim$1.2 nm and TO of s-SWNTs with diameter $\sim$1.0 nm [86], has both the same position and FWHM in the three spectra. FWHM(G$^+$) is reduced both in the top (63%) and bottom (32%) fractions, with respect to the starting material.

Fig. 5e and f report the spectra in the RBM and G peak region, respectively, measured at 782 nm for the HiPco samples. This excitation wavelength is in resonance mainly with the s-SWNTs e_{32} [65]. The RBM region shows an enhancement of the peaks in the spectral range 200–250 cm$^{-1}$ for the top fraction, with respect to the un-sorted sample, while the same peaks have a reduced intensity in the bottom fraction. The detection of mainly s-SWNTs in all three samples is also evidenced by the G peak region. Pos(G$^+$) and Pos(G$^-$) have are at $\sim$1590 and $\sim$1559 cm$^{-1}$ for the three samples. Moreover, FWHM(G$^+$) has a similar width ($\sim$15 cm$^{-1}$) in the top and bottom bands.

As for the HiPco samples, also in CoMoCAT SWNTs the 514.5 nm excitation probes both m- and s-SWNTs [65]. The RBM spectra are reported in Fig. 6a. The spectra show two distinct regions: 150–200 cm$^{-1}$, assigned predominantly to m-SWNTs, and the other (with more intense peaks) in the 225–320 cm$^{-1}$ range, associated with resonance of both m- and s-SWNTs [65]. The spectra of the top band have lower intensity than the bottom one. There is also a signal associated to s-SWNTs (i.e. (8,4)). This chirality as well as (7,6) and (10,2), are also detected in the absorption spectrum of the bottom band, Fig. 2d. The higher content of s- and m-SWNTs in the top and bottom fractions, respectively, with respect to the un-sorted sample is evidenced by the analysis of the G peak (Fig. 6b).

Indeed, FWHM(G$^+$) of the top band is $\sim$54% narrower and Pos(G$^+$) is $\sim$21 cm$^{-1}$ upshifted with respect to the un-sorted sample, while we detect a $\sim$23% widening of FWHM(G$^-$) and $\sim$13 cm$^{-1}$ downshift of Pos(G$^-$) with respect to the un-sorted sample. At 633 and 782 nm, mostly s-SWNTs are in resonance and no clear separation is detected from the Raman analysis. Absorption measurements in Fig. 2d indicate s-SWNTs (i.e. (7,5), (7,6), (10,2), (9,4), (8,4) etc.) [64,65], in the bottom band as well. These tubes are also detected in the RBM region.
To understand the temperature influence on the sorting process, we carry out the process between 6 and 50 °C, respectively. For DIPS–CNT, there is no clear color distinction between the top and bottom band of HiPco and CoMoCAT, the separation process at RT causes the different colors to overlap. The colors of SWNTs are black or gray. This is due to the large tube diameter distribution that results, where an enrichment of s-SWNTs is detected in the top fraction (∼78%), while the bottom fraction, although enriched in m-SWNTs, still contains ∼35% s-SWNTs. FWHM(G+) for the top band is narrower compared to the unsorted one (11 instead of 14 cm⁻¹), see Fig. 6f. However, this reduction is not sufficient to indicate a narrower diameter distribution. Due to the diameter dependence of Pos(G+) [86,87], the removal of tubes with large difference in diameter, will reduce FWHM(G+) [86,87]. Pos(G⁻) is 3 cm⁻¹ downshifted and FWHM(G⁻) ∼47% larger in the bottom with respect to the top band.

To summarize, Figs. 3–6 show that after sorting using gel agarose filtration, we have an enrichment of s- and m-SWNTs in the top and bottom bands, respectively. The separation is more effective for s- than m-SWNTs, in agreement with the absorption spectra in Fig. 2.

### 3.2. Temperature dependence

To understand the temperature influence on the sorting process we carry out the process between 6 and 50 °C, Fig. 7. The temperature range is chosen based on the instability of SDS and gel agarose gel below 6 °C and above 50 °C, respectively. For AD-CNT, HiPco and CoMoCAT, the separation process at RT shows distinct coloration between the top and bottom band [33,37,95]. For DIPS–CNT, there is no clear color distinction between the top and bottom bands, they both are black or gray. This is due to the large tube diameter distribution that causes the different colors to overlap. The colors of SWNTs with different chirality depend on the optical absorption peak [39]. The colors of sorted SWNTs in Fig. 7 change according to the separation temperature. This could be ascribed to the presence of diverse SWNT species, due to the sorting process, that absorb at different wavelengths.

For different separation temperature, the variation in fraction concentration between the top and bottom band is observable, indicating a difference between the quantities of tubes in the respective bands after separation. Their relative concentrations can be determined from the optical absorption analysis presented in the supporting information (Fig. S5). Generally, the bottom bands are more concentrated for low temperatures top bands are more concentrated at high temperatures. The relative concentration is dependent on the SWNTs source. From Fig. S5, the maximum bottom band to top band concentration ratio is ∼10 and ∼0.1 at 6 and 50 °C, respectively.

The effectiveness of the ms-separation changes with process temperature both for the top and bottom bands, as can be seen from the OAS in Fig. 8. The bottom band (Fig. 8a) extracted for DIPS–CNT shows the highest enrichment in m-SWNTs at RT, while the top band (Fig. 8b) has the highest (∼80%) percentage of s-SWNTs at 10 °C. A similar trend is also observed for AD-CNTs. The bottom band for AD-CNTs (Fig. 8c) has the highest percentage of m-SWNTs at 30 °C, while the top bands (Fig. 8d) has the highest percentage of s-SWNTs (∼95%) at 6 °C. The same trend, with a better enrichment of m-SWNTs, is detected in the bottom bands at RT or slightly higher, and s-SWNTs in the top bands at low temperatures is also valid for HiPco samples, Fig. 9a and b. As for AD-CNTs, HiPco nanotubes show a higher enrichment (∼70%) of m-SWNTs in the bottom fractions, Fig. 9a, at 40 °C, while the s-SWNTs in the top fractions increase (∼95%) at 6 °C, see Fig. 9b.

The effect of temperature for CoMoCAT tubes is less pronounced with respect to the other samples. However, Fig. 9c and d show higher m-SWNT enrichment in the bottom band at 40 °C (Fig. 9c) while, contrary to the other tube sources, the top band (Fig. 9d) shows higher enrichment of s-SWNTs at RT.

The highest separation purity for s-SWNTs (6 °C) and m-SWNTs (40 °C) are ∼95% and ∼70%, respectively. While the highest s-SWNT purity (∼95%) is comparable to ref. [46] using the same separation method, while the purity for m-SWNT achieved here (∼70%) is lower. Depending on the electronic type required, i.e. s-SWNT or m-SWNT, one has to optimize the temperature parameter accordingly. Furthermore, the correlation between process temperature and separation purity is consistent for all four different SWNT sources.

The Raman spectra of the sorted fractions for sources are reported in supplementary information. The Raman analysis confirms the data from OAS, showing higher enrichment of s-SWNTs at low temperatures and m-SWNTs at temperatures higher than RT.

PLE measurements are used to analyze the content of s-SWNTs in the sorted fractions and how their composition changes with temperature. Due to the diameter range of DIPS–CNT and AD-CNT samples, the PL emission mostly fall in the infra-red region, therefore cannot be measured by our detector. Fig. 10 shows the PLE contour maps of sorted HiPco and CoMoCAT, at 6, 24 and 50 °C. The PLE maps for the un-sorted materials are reported as well. The emission intensity contours of these maps are normalized using the same factor used for the OAS. Figs. 10 and 11 show that the PLE signals are lower in the bottom fractions. Also, the PLE intensity in the bottom fractions decreases as temperature increases.

Figs. 10 and 11 are PLE maps of the starting materials and the sorted sample dispersions for HiPco and CoMoCAT, respectively. Fig. 10a is a PLE map of un-sorted HiPco. PL from exciton–exciton resonances of tubes with different chiralities, phonon sidebands, and EET [57,58] are detected. The strongest EET (εh_bar_22, εh_bar_33) features, i.e. from (6,5) to (7,5), from (7,5) to (10,2) and from (7,6) to (8,6) have PL emission comparable to the (εh_bar_22, εh_bar_33) emission of other chiralities. The EET features are an indication of the presence of bundles [57,58]. Fig. 10b shows a decrease in PL emission from smaller diameter tubes, such as (6,5), (8,3) and (7,5) in the top fraction of the sorted sample at 6 °C with respect to the un-sorted dispersion. An increase in PL emission from larger tubes, such as (8,6), (8,7), (9,5), etc. is detected in the same sample with
respect to the un-sorted dispersion. The PLE intensity of the sorted top fraction of HiPco SWNTs at 24 and 50 °C (Fig. 10c and d), decrease with temperature. EET features are also detected in these PLE maps. Fig. 10e–g plots the PLE maps of the bottom fraction of sorted HiPco SWNTs at 6, 24 and 50 °C, respectively. The absorption and Raman measurements

Fig. 7 – Bottom (B) and top (T) band fractions for three different process temperatures. (a) DIPS–CNTs, (b) AD-CNTs, (c) HiPco, (d) CoMoCAT. The colors of the dispersions vary moving from 6 to 50 °C and for different SWNT sources. (A color version of this figure can be viewed online.)

Fig. 8 – Optical absorption spectra for DIPS–CNTs (a) bottom and (b) top band, AD-CNTs (c) bottom and (d) top band. The peaks between 950 and 1150 nm in (b) and (d) are instrument originated noise associated with changing of source and detector. (A color version of this figure can be viewed online.)
Fig. 9 – Optical absorption spectra for HiPco (a) bottom and (b) top band and CoMoCAT (c) bottom and (d) top band. (A color version of this figure can be viewed online.)

Fig. 10 – PLE maps of (a) un-sorted and sorted HiPco at 6, 24 and 50 °C, for (b–d) top and (e–g) bottom fractions. The emission intensity contour are normalized with the same factor used for the OAS in Fig. 9a. (A color version of this figure can be viewed online.)
indicate that these fractions are enriched in m-SWNTs. However, we still detect PL emission from these dispersions, although with lower intensity with respect to both the unsorted dispersion and the top fractions (enriched in s-SWNTs). E.g., the (7,5) PL intensity is ~20% and ~10% (Fig. 10e and f), of that from tubes of the same chirality in the un-sorted dispersions. Unlike the top fraction (s-SWNTs enriched), the bottom fraction has more m-SWNTs for higher temperature. While at 6 and 24 °C we still detect PL emission from a number of chiralities, the bottom fraction sorted at 50 °C does not have any PL features. These two observations are in agreement with the Raman and OAS measurements.

The PLE maps of CoMoCAT samples are reported in Fig. 11. As for the HiPco samples, the un-sorted CoMoCAT dispersions also show bundles [57,58]. Indeed, strong EET features are revealed together with the exciton–exciton resonances of tubes with different chiralities and phonon sidebands. Moreover the PLE maps of the top fraction (Fig. 11b–d) processed at different temperatures show a decrease in PL emission from smaller diameter tubes, such as (6,5), (8,3) and (7,5) in the top fraction of the sorted sample at 6 °C with respect to the un-sorted dispersion.

An increase in PL emission from larger tubes, such as (7,6), (8,4), (9,4), etc. is detected in the same samples with respect to the un-sorted dispersion. The PL emission is reduced with increasing process temperature, as shown in the PLE maps of the sorted top fraction of CoMoCAT at 24 and 50 °C (Fig. 11c and d). EET features are also detected in these PLE maps. The PLE maps of the bottom fraction, for the separation process carried out at 6, 24 and 50 °C, show a low intensity PL emission, Fig. 10e–g. As for HiPco samples, the PL intensity decreases with increasing process temperature. However, unlike the HiPco sample (Fig. 10g) where no PL emission is detected at 50 °C, for CoMoCAT, a low intensity PL is still detected from (6,5), (7,5), (8,5) and (7,6), Fig. 11g. This confirms the data from OAS and Raman, pointing to a more effective ms-separation with HiPco compared to CoMoCAT.

The optical characterization shows that there is a correlation between the percentage of m- or s-SWNTs and temperature. The top fractions have increasing s-SWNTs content with decreasing temperature (up to 95% at 6 °C for HiPco SWNTs). The m-SWNT purity in the bottom fraction increases with increasing temperature. Ref. [48] reported that the affinity or interaction strength between s-SWNTs and the agarose gel surface increases with decreasing tube diameter. In the case of DIPS–CNT and AD-CNT, having larger average diameter with respect to CoMoCAT and HiPco, the weaker agarose-SWNT affinity results in more s-SWNT getting eluted along with m-SWNT when the 1% SDS solution is poured into the separation column, resulting in the bottom band having lower m-SWNTs compared to HiPco and CoMoCAT (~70%).

3.3. Separation mechanism

The separation mechanism is connected with the wrapping and encapsulation [31] of the SDS surfactant molecules around SWNTs. SDS molecules discriminate between SWNTs of different electronic structure by forming different assembly micelles structures on s- and m-SWNTs [96]. The interaction is strongly dependent on pH condition [97] and concentration of SDS molecules [49,98]. Ion-dipole forces were proposed as the primary interaction mechanism responsible for SWNTs adsorption on agarose gel [99]. Ref. [100] suggested that, due to the electrostatic properties of SWNTs, SDS molecules form flatly assembled and randomly oriented structures around s-SWNTs, and cylindrical micelles around
m-SWNTs. This difference in encapsulation orientation is responsible for the separation process [100].

The agarose gel surface was shown to have very low physical affinity toward SDS molecules [33]. s-SWNTs with random flat SDS shell have less surfactant coverage. This is responsible for the ineffectiveness of the shielding between s-SWNTs and the agarose gel, driven by the stronger affinity of s-SWNTs toward agarose gel [100]. In contrast, m-SWNTs with ordered cylindrical micelles have high density of surfactants around the surface, forming a brush-like structure [100,101]. The latter results in a steric hindrance between SWNTs sidewalls and agarose surface. Fig. 12 schematically represents this process. Upon flushing 1% SDS solution into the agarose-SWNTs mixture the m-SWNTs are eluted, while the s-SWNTs remain physisorbed onto the agarose surface. By adding a different surfactant aqueous solution, such as SDC, SC, SDS, Tween 20 and TX-100, all with bigger molecular structures compared to SDS [102], the s-SWNTs can then be removed from the agarose surface and eluted through the separation column [46]. The added surfactants substitute and displace the SDS in the agarose gel column.

Our results are in agreement with the proposed separation mechanism described in Ref. [100]. In our case, the adsorption of SDC molecules with the subsequent formation of secondary micelles [31] enhances the steric hindrance between agarose and SWNTs. The s-SWNTs are then eluted following the same process as the metallic counterpart. This proposed separation mechanism could explain why the percentage of m-SWNT in the bottom band is lower than the percentage of s-SWNT in the top band (70% vs. 95%), as reported in Section 3.1.

In addition, our data shows evidence that, by varying the separation process temperature, the relative size and distribution of micelle structures around the SWNTs are affected. Consequently, the resulting separation purity is also affected and dependent on the process temperature variation, for a given agarose concentration (2% agarose). The purity of s-SWNTs in the top band increases with decreasing temperature, while m-SWNTs show the opposite behavior.

Ref. [103] reported a correlation between the purity of the separated SWNTs and the concentration of agarose in the gel matrix, which is related to the pore size. It was suggested that the interaction between SWNTs and agarose gel occurs within the pores in the gel matrix, leading to the separation of SWNTs by electronic type [103]. However, from our study, we suggest that the interaction between SWNTs and agarose gel occurs primarily around the surface of the gel beads [46], rather than within the gel matrix as described in Ref. [103]. Here, the agarose gel beads column acts as a sieve through which the SWNT solution is filtered by means of gravitational pull alone, similar to gel column chromatography [46]. The SWNTs are not embedded in the agarose gel matrix, in which case electrophoresis [27] or gel compression [103] must be employed to separate and release them. Consequently, the variance in separation purity with temperature here may be attributed to the surfactant micelles structure and density around the tubes, rather than the change of gel agarose morphology or pore size.

There is still a considerable debate regarding the wrapping mechanism and orientation of surfactant molecules around SWNTs. The selectivity of SDS toward m- or s-SWNTs has not been fully clarified yet. Both theoretical [101,104,105] and experimental [100,102,106] investigations concur that surfactants can have different orientations around SWNTs depending on concentration [101] and type [102], as well as tube diameter [101,105] and electronic type [100]. SDS can either lie randomly on the surface of SWNTs [100,105], or can form cylindrical micelles perpendicular to the SWNT circumference, with the hydrophobic tail attached to the SWNT surface, and the hydrophilic head pointing outwards.

![Fig. 12 – Scheme of ms-separation process. Blue tubes are m-SWNTs encapsulated with SDS forming cylindrical micelles; red tubes are s-SWNTs with low concentration of random flat SDS adsorbed on their sidewalls; textured green masses represent agarose gel microbeads. The m-SWNTs remain mobile and suspended in the solvent while the s-SWNTs are physisorbed on the surface of the agarose gel microbeads. (A color version of this figure can be viewed online.)](image-url)
[100–102]. It was also shown that SDS micelles are affected by temperature. When the temperature is increased, the SDS micelles’ size reduces, the micellar shape becomes distorted and the aggregation number reduces [107].

Therefore, we propose that the separation process temperature affects the distribution of SDS micelles around the SWNT sidewalls, resulting in the variation of m- and s-SWNTs percentage. At low temperatures (6–10 °C), the micelle structures around the SWNTs are bigger and the SDS molecules are closely packed, possibly making them less susceptible to be physisorbed on the gel agarose surface. This mechanism could explain why most SWNTs are eluted from the separation column after flushing with a SDS solution. Only the s-SWNTs with low enough SDS encapsulation density can physisorb on the agarose surface with limited physisorption sites, resulting in a higher percentage of s-SWNTs being eluted from the separation column after flushing with 1% SDC. At RT or higher, the SDS micelles structure encapsulating the SWNTs becomes smaller and loosely packed in a way that increases the affinity toward SWNTs by providing more physisorption sites. Consequently, SWNTs with low micelles density and random micelles structure, which includes some m-SWNTs, are more susceptible to being physisorbed and trapped onto the agarose surface even after successive addition of 1% SDS solution into the separation column. Only m-SWNTs with large surfactant micelles density will be eluted, the rest being trapped in the gel column, yielding higher m-SWNTs purity in the bottom band. Upon adding the 1% SDC solution, the remaining trapped SWNTs in the top band are eluted. Thus, the collected top band will contain higher amounts of m-SWNTs, therefore yielding lower s-SWNTs purity than its low temperature counterpart.

3.4. CNT-based transistors

CNT-FETs are then fabricated to assess the electrical characteristics of the sorted s-SWNTs and to demonstrate their direct application in electronic devices. Each device consists of a SWNTs network contacted by Au metal source and drain electrodes with channel width ~500 nm on heavily doped p-type Si, with 250 nm thick thermally grown SiO2 dielectric. We prepared devices based on the s-SWNT fraction from each of the four SWNT samples after ms-separation at 6 °C, i.e. DIPS–CNT (Dev-1), AD-CNT (Dev-2), HiPco (Dev-3) and CoMoCAT (Dev-4). For comparison, we also fabricate a device based on a DIPS–CNT sample (Dev-5) processed in the same way as the ones used for Dev-1 but without filtration (enrichment). Fig. 13a shows a representative scanning electron micrograph (SEM) image of an array of devices from a single SWNT source (Dev-1). The measurements are carried out on one of the electrode pairs for each SWNT sources. The densities of the SWNT networks of all devices are controlled to be ~5 tubes per μm². The zero gate bias I-V characteristic of the five devices are shown in Fig. 13b. Both Dev-1 and Dev-2 show non-linear I-V curves with maximum conductance ~2.7 × 10⁻⁸ S. For Dev-2, which consist of an AD-CNT channel, the I-V curve is asymmetrical, possibly due to the rectifying effect of an unbalanced Schottky barrier on one of the electrode-tube interfaces. The I-V curves for Dev-3 and Dev-4 are also non-linear, but they exhibit lower conductance ~6.2 × 10⁻¹⁰ and ~2.1 × 10⁻⁹ S, respectively, with respect to Dev-1 and Dev-2. A non-linear I-V curve is expected for a typical s-SWNTs network [2]. However, Dev-5 has an Ohmic linear I-V curve with higher maximum conductance (~2.2 × 10⁻⁷ S) with respect to the other devices, which reflects the characteristics of an un-sorted SWNTs channel where carrier transport is markedly affected by the presence of m-SWNTs.

The 3-terminal field effect output and transfer characteristics are shown in Figs. 14 and 15, respectively. The output characteristic of Dev-1 in Fig. 14b indicates how the gate-source voltage bias, V₉GS, modulates the drain-source current, I_DS. An increase of V₉GS in the negative range corresponds to an increase of I_DS (ON state). A gradual decrease of I_DS, toward values approaching zero (OFF state), is seen when V₉GS increases in the positive range, which indicates p-type FET behavior [108]. The output characteristics of Dev-2, Dev-3 and Dev-4 are very similar to Dev-1. The un-sorted SWNTs channel of Dev-5, by comparison, produces very slight gate bias modulation and higher I_DS with respect to Dev-1 (Fig. 14b). This is another indication that the carrier transport in the channel is dominated by m-SWNTs, which cannot be fully depleted, thus the devices cannot be turned off. The

Fig. 13 – (a) SEM of a representative CNT-FETs array. The scale bar is 2 μm. (b) I-V curves at zero gate bias for devices based on s-SWNTs enriched channels: DIPS–CNTs (Dev-1, red), AD-CNTs (Dev-2, blue), HiPco (Dev-3, magenta), CoMoCAT (Dev-4, green), and un-sorted DIPS–CNTs channel (Dev-5, black). (A color version of this figure can be viewed online.)
analysis of the output characteristics, and the comparison with Dev-5, further confirms the spectroscopic analysis, indicating that the sorted SWNT network in Dev-1, Dev-2, Dev-3, Dev-4 containing mainly s-SWNTs.

The transfer characteristics of the devices in Fig. 15 demonstrate current modulation and switching for all four SWNT sources. On the other hand, the transfer curve for Dev-5 with unsorted SWNTs shows almost no switching. The transfer curve for Dev-1 has ambipolar behavior, attributed to the intrinsic properties of DIPS–CNTs. The low turn-on voltage observed for Dev-1 (~10 V) is due to the large average tube diameter (d) of DIPS–CNTs, hence smaller bandgaps, compared to the other SWNT sources.

Dev-1 and Dev-2 have \( I_{ON}/I_{OFF} \sim 10^5 \) and ~10^6, respectively. Dev-3 and Dev-4 have \( I_{ON}/I_{OFF} \sim 10^4 \). \( I_{ON} \) up to ~10^3 have been reported for CNT-FETs [109], using thin molecular high-k dielectric as gate oxide [109]. CNT-FETs fabricated from sorted SWNTs via DGU [37] and gel filtration [103,110,111] have shown lower \( I_{ON}/I_{OFF} \) of ~10^2–10^4. The very high \( I_{ON}/I_{OFF} \) of Dev-2 is predominantly due to the low off-current ~10^-13 A, compared to that of Dev-1, ~10^-11 A. Dev-3 and Dev-4 also have low off-current of ~10^-12 A, but the on-currents are lower than in Dev-1 and Dev-2. This results in a lower \( I_{ON}/I_{OFF} \). For Dev-2, the average off-current is ~10^-12 A. This can be ascribed to the fact that the carrier transport in the device channel is dominated by s-SWNTs and can therefore be modulated by the field generated by gate bias. The \( I_{ON}/I_{OFF} \) of Dev-5 is less than 2, with high off-current ~4 \( \mu \)A, which is ~7 orders of magnitude higher than Dev-2. This is related to the presence of m-SWNTs, which dominate the carrier transport. The transfer characteristic of Dev-1 shows asymmetric ambipolar behavior, p-type dominant, attributed to the intrinsic properties of the DIPS–SWNTs source. SWNTs synthesized via the DIPS method were shown to have ambipolar behavior when exploited as the active channel of a transistor [112], while SWNTs in general exhibit p-type behavior in air [5].

There are intrinsic physical limitations to the maximum \( I_{ON}/I_{OFF} \) obtainable for a given supply voltage [22]. At best, \( I_{DS} \) increases exponentially from \( I_{OFF} \) to \( I_{ON} \) with a rate described by the subthreshold swing (SS), expressed in mV dec^{-1} (i.e., mV of incremental gate voltage required to change the drain current by one decade) [22,108].

Therefore, a low SS is desirable for efficient device switching [108]. From Fig. 15, SS for Dev-1 to Dev-4 is estimated to be 700, 1100, 1570 and 7700 mV/dec, respectively. These values are higher than the best SS reported for CNT-FET devices with similar structures (~200 mV/dec) [113]. Our higher SS can be attributed to the thick 250 nm SiO_2, and lower concentration of SWNTs in the channel network [114], as well as charge in the SiO_2 and surface traps. Another indicator of CNT-FET switching performance is the transconductance \( g_{m} \), defined as \( g_{m} = \frac{\Delta I_{DS}}{\Delta V_{GS}} \), where \( I_{OUT} \) is the current variation at the output and \( V_{IN} \) the voltage variation at the input [108]. Dev-1 and Dev-2 have \( g_{m} \sim 2.45 \times 10^{-8} \) S for both devices. Dev-3 and Dev-4 exhibit much lower \( g_{m} \sim 0.3 \times 10^{-8} \) S and ~0.1 \times 10^{-8} S, respectively. As for SS, this low \( g_{m} \) can be attributed to the thick dielectric layer, which introduces high parasitic capacitances and charge trapping between channel and gate. However, the variation of SS and \( g_{m} \) between the four devices investigated here can be attributed to the respective SWNT sources, with varying average tube diameter, and hence band gap [115–117].

The mobility \( \mu \) of our CNT-FETs can also be determined from the 3-terminal electrical characteristic measurements. We estimate the “device mobility” or “field effect mobility”, \( \mu_{FE} \), which takes into account the scattering effects of the

![Fig. 14 – Output characteristics for (a) devices with un-sorted DIPS–CNTs, and (b) device with sorted DIPS–CNTs. (A color version of this figure can be viewed online.)](image-url)

![Fig. 15 – Transfer characteristics for device with sorted DIPS–CNTs channel (Dev-1), sorted AD-CNTs channel (Dev-2), sorted HiPco channel (Dev-3), sorted CoMoCAT (Dev-4) and un-sorted DIPS–CNTs channel (Dev-5). The source-drain voltage, \( V_{DS} \), is 1 V for all curves. (A color version of this figure can be viewed online.)](image-url)
SWNT channel and the metal-SWNT contacts, rather than the SWNT intrinsic mobility [118,119]. A single device is measured between a pair of source-drain electrodes with channel length \( \sim 500 \text{ nm} \). The active channel of the devices consists of either a single or a few SWNTs. Therefore we assume a one-dimensional system to simplify the calculations. \( \mu_{FE} \) can be calculated directly by using the formula:

\[
\mu_{FE} = \frac{L}{C_g} \frac{\partial G}{\partial V_{GS}}
\]

where \( L \) is the SWNT channel length, \( C_g \) is the gate capacitance, and \( G \) is the device conductance. \( C_g \) can be calculated using:

\[
C_g = L \left[ C_G^{-1} + \frac{\ln(2 \frac{L_{ox}}{\theta_0})}{2 \varepsilon_0 \varepsilon_r d_i} \right]^{-1}
\]

where \( C_G^{-1} \) is the quantum conductance (0.4 nF.m\(^{-1}\)) [120], \( \varepsilon_{ox} \) is the oxide thickness (\( \sim 250 \text{ nm} \)) and \( \varepsilon_r \) and \( \varepsilon_0 \) are vacuum permittivity and SiO\(_2\) relative permittivity, respectively. \( d_i \) is the SWNT diameter, which is approximated by the average diameter of the respective SWNT sources. This gives \( \mu_{FE} \) of \( \sim 49 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) for Dev-1 (DIPS–CNT). The maximum \( \mu_{FE} \) for device Dev-2 (AD-CNT), Dev-3 (HiPco) and Dev-4 (CoMoCAT) are 5.2, 4.2 and 2.3 cm\(^2\) V\(^{-1}\) s\(^{-1}\), respectively. These values are typical for devices from solution-suspended SWNTs, which are generally inferior to as grown CVD SWNT devices [119]. This difference can be attributed to the high SWNT-electrode contact resistance and strong scattering in the channel region from structural defects [121] likely induced by the sonication process.

4. Conclusions

We demonstrated the effect of process temperature in the sorting of m- and s-SWNTs via column chromatography, exploiting agarose gel beads. The sorting procedure is of general validity, being almost independent of nanotubes source. The enrichment of s-SWNTs is more effective at low temperatures (6–10 °C) while m-SWNTs are sorted more effectively in the range 30–40 °C. The best performances is achieved for HiPco: \( \sim 95\% \) s-SWNTs at 6 °C. The temperature dependence helps understanding the sorting mechanism. We suggested that the separation process temperature influences the SDS micelles’ structure size and distribution density, consequently affecting the separation purity. The high purity achieved by our separation protocol allowed us to fabricate CNT-FETs, using sorted s-SWNTs, reaching \( I_{ON}/I_{OFF} \sim 10^6 \), a further indication that the channels consist predominantly of s-SWNTs.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.carbon.2015.05.036.

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