Science and applications of wafer-scale crystalline carbon nanotube films prepared through controlled vacuum filtration

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Carbon nanotubes (CNTs) make an ideal one-dimensional (1D) material platform for the exploration of exotic physical phenomena under extremely strong quantum confinement. The 1D character of electrons, phonons and excitons in individual CNTs features extraordinary electronic, thermal and optical properties. Since the first discovery, they have been continuing to attract interest in various disciplines, including chemistry, materials science, physics, and engineering. However, the macroscopic manifestation of such properties is still limited, despite significant efforts for decades. Recently, a controlled vacuum filtration method has been developed for the preparation of wafer-scale films of crystalline chirality-enriched CNTs, and such films immediately enable exciting new fundamental studies and applications. In this review, we will first discuss the controlled vacuum filtration technique, and then summarize recent discoveries in optical spectroscopy studies and optoelectronic device applications using films prepared by this technique.

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I. INTRODUCTION

Carbon is a fundamental element in Group IV of the periodic table of elements, which is not only essential for life on earth but also promising for modern technologies in communication, computation, imaging, sensing, energy, health and space. Each carbon atom has six electrons, occupying the 1s, 2s, and 2p atomic orbitals. Two core electrons (1s2) are strongly bound, while four weakly bound electrons (2s22p2) can form covalent bonds in carbon materials using sp2 hybrid orbitals. Other Group IV elements like Si and Ge can only exhibit sp3 hybridization [1]. Various hybridization states in carbon lead to a versatile chemical nature and determine the properties of carbon allotropes and organic compounds [2]. Two natural carbon allotropes, diamond and graphite with sp3 and sp2 hybridization, respectively, show distinct properties; they were the only known carbon allotropes for a long time until the discovery of zero-dimensional (0D) C60 in 1985 [3], giving birth to nanotechnology. After the advent of C60, a growing family of carbon nanomaterials emerged, including 1D carbon nanotubes (CNTs) [4] and two-dimensional (2D) graphene [5].

Still other members, based on numerous combinations and modifications of carbon atom hybridization, are yet to be discovered. Despite the unique properties of each member of the carbon nanomaterial family, their atomic structures are related to one another. In this sense, graphene can be viewed as a fundamental building block because fullerenes, carbon nanotubes, and graphite can all be derived from graphene [6]. In particular, a single-wall CNT (SWCNT) is a rolled-up version of monolayer graphene, which possesses unique 1D thermal, mechanical, electronic and optical properties [1][2][9].

A. Electronic, optical properties and applications of SWCNTs

The crystal and band structure of SWCNTs are highly related to those of graphene. However, the strong confinement of electronic states along the tube circumference makes SWCNTs a completely different material. Graphene consists of a hexagonal lattice of sp2-bonded carbon atoms, as shown in Figure 1(a). The carbon-carbon bond length aC-C is 1.42 Å. The primitive unit cell contains two carbon atoms, A and B, contributing two π electrons that play fundamental roles in electrical and optical phenomena. Figure 1(b) shows the reciprocal lattice of graphene, which is also a hexagonal lattice, and there are three important high-symmetry points: Γ-point (the center of the Brillouin zone), M-point, and K-point. The a1,2 and b1,2 are the primitive unit vectors in real and reciprocal space, respectively.

The band structure of SWCNTs sensitively depends on the manner in which the graphene sheet is rolled up, which can be uniquely specified by defining the chiral vector C = na1 + ma2, where n and m are positive integers with n ≥ m; see Figure 1(c). A SWCNT is formed by wrapping the graphene sheet in such a way that the two points connected by C meet, with a diameter d. The chiral indices (n,m) fully determine the crystal structure of the SWCNT and the character of the band structure depends on the value ν = (n−m) mod 3:

1. If ν = 0, the SWCNT is metallic;
2. If ν = 1 or 2, the SWCNT is semiconducting with a band gap ~0.7 eV/d (nm).

However, due to the nanotube curvature effect, only armchair nanotubes (n = m) are truly metallic. For other tubes with ν = 0, there is a small, curvature-induced band gap that scales with 1/d2.

As shown in Figures 1(d) and 1(e), 1D SWCNTs have a divergent density of states at the van Hove singularities, which, together with strong 1D excitonic effects, produces strong optical absorption and emission through the excitation and recombination of electron-hole pairs. The strong quantum confinement of electron-hole pairs results in the generation of stable excitons, and these excitons can be excited either optically or electrically and their radiative recombination results in photoluminescence or electroluminescence, respectively. On the other hand, the 1D nature of SWCNTs limits the direction of carrier scattering to occur only along the nanotube axis while in a higher dimensional material system, scattering directions can be arbitrary. For example, metallic SWCNTs have a limited momentum space, resulting in a suppression of back scattering and a very long mean free path [10], which is ideal for quantum wire interconnections [11]. Furthermore, field effect transistors built from semiconducting SWCNTs have demonstrated a strong electric field effect and showed an extremely high room temperature mobility > 105 cm2/Vs [12]. In addition, the optical absorption of semiconducting SWCNTs can be electrically modulated, laying the foundation for optoelectronic device applications.

Although individual CNTs provide an ideal platform for investigating fundamental properties of CNTs and demonstrating proof-of-concept optoelectronic device applications, individual SWCNT devices are still problematic from the viewpoint of practical applications because of a complicated fabrication process, low photoluminescence quantum yields, and an inability to scale up [13]. Individual CNT transistors are considered to be impractical due to difficulty in positioning CNTs precisely, which applies to individual CNT photonic and optoelectronic devices as well. Although light-matter interaction in individual CNTs is unusually strong, further cooperative enhancement can be expected when a large number of CNTs can be assembled in an ordered manner.
FIG. 1. The crystal and band structure of SWCNTs. (a) Real space and (b) reciprocal space representation of the hexagonal lattice of 2D graphene. $a_{C,C}$ is the carbon-carbon bond length. $a_{1,2}$ and $b_{1,2}$ are the primitive unit vectors in real and reciprocal space, respectively. The dashed line parallelogram defines the unit cell of graphene with two different carbon atoms A and B. High-symmetry points in reciprocal space, $\Gamma$, $M$ and $K$, are labeled in the 1st Brillouin zone. (c) A chiral vector $C$ of a SWCNT, defined on the graphene sheet as a linear combination of $a_1$ and $a_2$ with integer coefficients. An index $(n,m)$ represents SWCNT chirality. Different index combinations result in either (d) a metallic SWCNT or (e) a semiconducting SWCNT.

The macroscopic assembly of individual CNTs, in the form of random networks or aligned arrays, is the key to facilitating practical applications. An ensemble assembly can help overcome challenges in individual CNT devices by (1) minimizing device variations, (2) providing a large active area, and (3) not requiring precise individual tube positioning. Currently, there are various ways to prepare a CNT ensemble, including chemical vapor deposition (CVD) [14, 15] and solution-based deposition [16]. The produced ensemble can exhibit either a semiconducting or metallic material behavior and has spurred a wide interest in various applications. For example, CNTs are thermally and chemically stable [17], making them ideal for high-temperature emitters [18] (Figure 2(a)) and compatible with high-$\kappa$ material deposition for high-performance electronics [19] (Figure 2(b)). A large ratio of the surface area to the volume of CNTs enables an efficient charge transfer doping from surrounding chemicals for optimized thermoelectric materials [20] (Figure 2(c)). The excellent mechanical strength of CNTs is perfect for large-area flexible electronic/optoelectronic applications [21] (Figure 2(d)). Large active areas possess the potential for an efficient broadband detector [22, 23] (Figure 2(e)) and enable low-cost, high-efficiency heterogeneous solar cells [24] (Figure 2(f)). Furthermore, a highly conductive metallic SWCNT thin film of a small optical absorption cross section with strong mechanical properties can serve as a stretchable transparent electrode, which is important for certain security applications and backlit displays [25] (Figure 2(g)).

To best preserve the extraordinary properties of individual CNTs, macroscopic ensembles with a single domain of perfectly aligned CNTs, i.e., crystalline CNTs, is highly preferred. Taking SWCNT transistors, for example, the composition and morphology of conducting channels strongly affect the device performance. The degree of
FIG. 2. Representative photonic and optoelectronic devices using CNTs. (a) A current-driven high-speed blackbody light emitter [18]. (b) A high-performance transistor [19]. (c) Thermoelectric materials based on semiconducting SWCNTs with tailored doping profiles [20]. (d) Medium-scale flexible thin-film transistors [21]. (e) A broadband photothermoelectric detector [22]. (f) High-efficiency solar cells based on SWCNT-silicon heterojunctions [23]. (g) Large-scale transparent electrodes [25].

SWCNT alignment and the purification of semiconducting SWCNTs affect two important device parameters: the carrier mobility and the on-off ratio [26]. With a perfectly aligned semiconducting SWCNT ensemble, the mobility and on-off ratio of a large-scale transistor are comparable with transistor devices based on individual CNTs. In contrast, a randomly oriented ensemble leads to a decrease in mobility that is due to stronger intertube scattering [13]. Furthermore, even a small amount of metallic nanotubes (e.g., 1%) dramatically decreases the on-off ratio (e.g., three orders) because of the percolation path formed by metallic tubes [27]. Therefore, it is crucial to separate nanotubes by electronic types and assemble them in an ordered manner to extend the excellent properties of individual CNTs to CNT macroscopic ensembles for real-world applications.
B. Challenges in synthesis of wafer-scale crystalline CNTs

To date, it is still challenging to produce a macroscopic single-domain, highly aligned, densely packed, and chirality-enriched SWCNT film. Current assembly techniques can be separated into two general categories: (1) in-situ direct-growth assembly methods and (2) ex-situ post-growth assembly methods. However, techniques in both categories have inherent limitations.

Direct-growth assembly methods consist of the CVD growth method for either vertical [14, 28-30] or horizontal alignment [31] with respect to growth substrates. If the catalyst density is high enough, a crowding effect leads to vertical alignment of CNTs with heights of ~mm, as shown in Figure 3(a). The as-grown patterns of well-aligned SWCNTs synthesized using the CVD method can be laid over to form horizontally aligned patterns and be transferred to any substrates through a scalable, facile, and dry approach [32]. To directly assemble horizontally aligned arrays of CNTs, multiple external-stimuli-assisted CVD growth methods have been developed, including electric field [33], gas flow [34] and epitaxy [19]. Electric fields can orient CNTs because of the large and highly anisotropic polarizability of CNTs (Figure 3(b)); a feeding gas flow can generate a flow field that can carry CNTs to overcome the gravity to grow along the flow direction (Figure 3(c)); and the oriented van der Waals force along the substrate surface can produce horizontally aligned CNTs (Figure 3(d)).

However, there is a common issue for direct-growth methods: it is still very challenging to control the chirality during the growth procedure. Obtained aligned arrays have mixed electronic types and are not desirable for most applications. Recently, there have been breakthroughs in precisely controlling chirality distribution during synthesis, by designing a catalyst that matches the carbon atom arrangement around the nanotube circumference of a specific chirality [35-37]. However, the packing density of aligned arrays is relatively low. The highest density reported is ~50 CNTs/µm [38], which is quite dilute in a practical sense, especially for photonic and optoelectronic applications.

Post-growth assembly methods mainly consist of solution-based techniques for CNT alignment [39]. It can take advantage of solution-based SWCNT separation techniques, such as the DNA-wrapping method [40, 41], the polymer-wrapping method [42], density gradient ultracentrifugation [43, 44], gel chromatography [45], and aqueous two phase extraction [46, 47]. Existing post-growth assembly techniques consist of methods using external force, self-assembly and liquid crystal phase transition.

External forces used in CNT alignment can be mechanical, magnetic, and electrical. The simplest method is based on mechanical force by dry-spinning CNTs from a CVD-grown CNT forest (Figure 4(a)) [48]; however there is no chirality selectivity in this method. An applied magnetic field can align CNTs based on their magnetic susceptibility anisotropy, but the required magnetic field to achieve meaningful alignment (Figure 4(b)) is typically larger than 10 T [49], impractical for any practical applications. A more affordable and practical method is to align CNTs in liquid by applying an AC electric field, called dielectrophoresis [50], as shown in Figure 4(c). The obtained films have alignment packing densities up to 30~50 SWCNTs/µm, which is still quite low. Furthermore, the alignment quality is relatively poor and full surface coverage is nearly impossible.

Instead, self-assembly methods, including evaporation-driven self-assembly [51, 52], Langmuir-Blodgett [53] and Langmuir-Sheafer [54] assembly, can produce a large-size full-surface-coverage aligned film. As the substrate is immersed inside a CNT suspension during the evaporation-driven self-assembly process, the interplay between the surface friction force, liquid surface tension, and capillary force can align CNTs at the solid-liquid-air interface (Figure 4(d)). Similarly, when the immersed substrate is pulled out of the CNT suspension vertically (Langmuir-Blodgett) or horizontally (Langmuir-Sheafer), the 2D confinement at the air-water interface causes CNTs to orient along certain directions (Figure 4(e)). A semiconducting SWCNT aligned monolayer film of an impressively high packing density of ~500 CNTs/µm with full surface coverage has been demonstrated for significantly improved transistor performance [54]. However, the thickness of fabricated films using these techniques is limited to a few nanometers because of the 2D nature of these techniques, and again the degree of alignment is not very high.

In contrast, a liquid crystal phase transition can lead to alignment in three dimensions. One strategy is to mix CNTs with a liquid crystal polymer matrix [55, 56]. Under certain conditions, the liquid crystal polymer can go through a phase transition spontaneously with macroscopically ordered structures, where the hosted CNTs follow the same transition. The degree of alignment can be very high, but the challenge is how to remove the liquid crystal polymers completely without destroying the alignment of CNTs. As a consequence, it motivated another strategy to use the liquid crystal transition of CNTs based on Onsager’s theory. Different aqueous dispersion systems have been investigated to achieve nematic ordering of CNTs [57, 58] in a short range (Figure 4(f)), but wafer-scale crystalline CNT films through a liquid crystal phase transition has not been achieved.
FIG. 3. In-situ direct-growth assembly methods of CNTs. (a) Water-assisted CVD growth \[29\]. (b) Electric field directed CVD growth \[33\]. (c) Gas flow directed CVD growth \[34\]. (d) Epitaxy CVD growth \[19\].

II. FABRICATION AND CHARACTERIZATION OF WAFER-SCALE CRYSTALLINE FILMS OF CHIRALITY-ENRICHED CNTS

Post-growth assembly methods are suitable for fabricating 3D architectures of macroscopically aligned CNTs, because of the availability of various solution-based chirality separation techniques. However, existing methods suffer from low packing densities, small thicknesses, incompatibility of separation techniques, and local alignment. Currently, there are no available techniques for fabricating large-area multi-domain films of highly aligned, densely packed, and chirality-enriched SWCNTs with controllable thickness. Recently, He and coworkers have developed a new “solution-based” alignment technique based on controlled vacuum filtration \[60\], which can provide a uniform, wafer-scale SWCNT film of an arbitrarily and precisely controllable thickness (from a few nm to \(\sim\)100 nm) with a high degree of alignment (nematic order parameter \(S \sim 1\)) and packing (\(\sim 3.8 \times 10^5\) tubes in a cross-sectional area of 1 \(\mu\)m\(^2\)) in a simple, well-controlled, and reproducible manner, regardless of the synthesis method, type, or chirality of the SWCNTs used. Furthermore, the produced films are compatible with standard micro/nanofabrication processes used to fabricate various electronic and photonic devices.

The controlled vacuum filtration process starts with a well-dispersed CNT suspension, where individual CNTs are separated and suspended. In order to prepare an aqueous suspension, the sidewalls of SWCNTs have to be functionalized either by chemical functional groups or amphiphiles with a hydrophobic tail and a hydrophilic head \[61\].
Ultrasonication is a widely used method for general nanoparticle dispersion, agitating molecules either in a bath or probe (or horn) sonicator. The generated ultrasound wave is delivered to a CNT bundle in a surfactant solution, and individual CNTs are peeled off from the outer part of the CNT bundle and wrapped by surfactant molecules to prevent future rebundling. The obtained suspension is then ultracentrifuged to remove any undissolved bundles, and the supernatant is collected. The prepared suspension is poured into a vacuum filtration system, as shown in Figure 5(a). A filter membrane with a pore size smaller than the CNT length blocks the passage of CNTs and allows water to penetrate through, under a differential pressure applied across the membrane that is maintained by a vacuum pump underneath. During the deposition process of CNTs, the penetrability of the filter membrane reduces gradually. This self-limiting process guarantees a uniform film [25]; see Figure 5(b). The obtained film can be readily transferred onto any substrate, such as a quartz wafer in Figure 5(c), by dissolving the filter membrane in organic solvents. Furthermore, as long as CNTs are well-dispersed in the suspension, the controlled vacuum filtration can apply to any CNT species, enabling us to take advantage of solution-based chirality separation techniques. For example, filtrated films made from mono-dispersed SWCNTs separated using density gradient ultracentrifugation show different colors due to subtractive coloration [62].

It has been believed that the vacuum filtration technique can only produce randomly oriented CNTs for many years. Figure 5(e) demonstrates an atomic force microscopy (AFM) image of randomly oriented films of CNTs prepared in a standard way. However, He et al. showed that this technique can be adapted to produce instead crystalline structures of CNTs [60]. Previous work [59] revealed some local ordering utilizing the liquid crystal phase transition of CNTs, as CNTs accumulate near the surface of the filter membrane during the filtration process. He et al. empirically identified three crucial conditions to achieve macroscopically aligned films: (1) the surfactant concentration must be below the critical micelle concentration; (2) the CNT concentration must be below a threshold value; and (3) the filtration process must be well controlled at a low speed. Furthermore, the surface chemistry of the filter membrane and CNT structural parameters have important influence on optimal conditions for macroscopic alignment. The scanning
FIG. 5. Wafer-scale crystalline CNT films fabricated using controlled vacuum filtration. (a) A CNT suspension goes through a vacuum filtration system. (b) A wafer-scale, uniform CNT film is formed on the filter membrane. (c) Optical image of the produced film after being transferred onto a transparent substrate. (d) Separated mono-dispersed suspension using the density gradient ultracentrifugation method and fabricated semitransparent colorful films [63]. (e) A typical AFM image of random networks of CNTs fabricated using the conventional vacuum filtration technique [25]. (f) A high-resolution SEM image and (g) a cross-sectional TEM image of the film. Adapted from Ref. [60]. (h),(i) Cross-sectional TEM images of a film produced by Falk and coworkers using the controlled vacuum filtration technique, showing crystalline CNTs with a high packing density ~3.8 x 10^5 CNTs/µm². (j) A TEM diffraction image of a selected area in (i), showing a hexagonal lattice. (k) Grazing-incidence X-ray diffraction spectrum of a crystallized CNT film (blue curve), and a control film of randomly oriented CNTs (black curve). Adapted from Ref. [64].

Electron microscopy (SEM) image in Figure 5(f) shows perfectly aligned SWCNTs, and the cross-sectional transmission electron microscopy (TEM) image in Figure 5(g) shows crystalline ordered SWCNT structure. Furthermore, Falk and coworkers [61, 65] reproduced the work of He et al. [60] and performed studies of crystal structures in obtained films using cross-sectional TEM and X-ray diffraction analyses. Cross-sectional TEM images in Figures 5(h) & (i) show that densely packed CNTs assembled in controlled vacuum filtration form a crystalline structure, and selected-area TEM diffraction patterns in Figure 5(j) confirm a hexagonal lattice. Figure 5(k) displays the X-ray diffraction spectra for both aligned and randomly oriented films, and a prominent peak at 2θ = 5.8° is only present in the aligned film. This peak corresponds to a lattice constant of 1.74 nm, an excellent agreement with the summation of the average diameter of CNTs (1.4 nm) and the constant spacing between carbon atoms (0.34 nm). Based on this information, the packing density in these films was estimated to be ~3.8 x 10^5 tubes in a cross-sectional area of 1 µm² [64].
Quantitative characterization of alignment quality can be made by calculating the nematic order parameter, \( S \). It is defined as the ensemble average value of \( \frac{3\cos^2(\theta) - 1}{2} \) in 3D and \( 2\cos^2(\theta) - 1 \) in 2D, where \( \theta \) is the angle deviation from the macroscopic alignment direction. Figure 6(a) shows the angle distribution of SWCNTs in obtained films made of arc-discharge SWCNTs with an angle standard deviation of \( \sim 1.5^\circ \) across an area \( \sim 1\text{mm}^2 \), indicating \( S \sim 1 \). This densely packed SWCNTs feature a strong anisotropic optical attenuation in the entire electromagnetic spectrum, from the terahertz (THz) to the visible, as shown in Figure 6(b). Specifically, there is no attenuation for the perpendicular polarization in the entire THz/infrared range (\(< 1\text{ eV}\)), whereas there is a prominent, broad peak at \( \sim 0.02\text{ eV} \) in the parallel direction because of the plasmon resonance in finite-length CNTs [66]. Figure 6(c) plots the same spectra in the infrared range with the energy axis on a linear scale. The two interband transitions in semiconducting nanotubes (\( E_{11}^S \) and \( E_{22}^S \)) and the first interband transition in metallic nanotubes (\( E_{11}^M \)) are clearly observed. According to the selection rules that will be discussed later in detail, these peaks are completely absent for the perpendicular polarization case. Instead, a broad absorption feature is observed between the \( E_{11}^S \) and \( E_{22}^S \) peaks, which can be attributed to the crosspolarized and depolarization-suppressed \( E_{12}^S / E_{21}^S \) absorption peak [67–69]. However, due to the mixed electronic types inside this film, the assignment is still speculative but the investigation of highly aligned single-chirality SWCNTs will be deterministic.

**FIG. 6.** Characterization of crystalline films of arc-discharge SWCNTs of mixed electronic types. (a) Angular distribution of CNTs within a 1 cm\(^2\) area of the film determined by SEM image analysis. (b) Polarization-dependent attenuation spectra in a wide spectral range, from the THz/far-infrared to the visible. (c) Expanded view of (b), showing interband transitions. (d) Time-domain THz waveforms of transmitted THz radiation for parallel and perpendicular polarizations. (e) Attenuation as a function of the angle between the THz polarization and the nanotube alignment direction. (f) Voltage-current relationship when the current flow is parallel and perpendicular to the CNT alignment direction. Adapted from Ref. [60].
parallel to the nanotube axis direction, which is valid for THz response of individual SWCNTs. Figure 6(d) shows time-domain waveforms of THz radiation transmitted through an aligned arc-discharge SWCNT film on an intrinsic silicon substrate for polarizations parallel and perpendicular to the alignment direction, together with a reference waveform obtained for the substrate alone. The data for the perpendicular case completely coincide with the reference trace; that is, no attenuation occurs within the SWCNT film. On the other hand, there is significant attenuation for the parallel case. Figure 6(e) shows a more detailed polarization-angle dependence of THz attenuation. The calculated $S$ is approximately 1, which is consistent with SEM analysis. Moreover, the obtained film shows strong anisotropic electronic conductivity (Figure 6(f)), with the conductivity along the alignment direction $\sigma_\parallel \sim 2500$ S/cm, the conductivity perpendicular to the alignment direction $\sigma_\perp \sim 42$ S/cm and the ratio between these two $\sigma_\parallel / \sigma_\perp \sim 60$ at room temperature.

FIG. 7. Macroscopically aligned chirality-enriched SWCNT films and devices. (a) Chirality separated semiconducting (10,3) and (6,5) SWCNTs using gel chromatography and aqueous two phase extraction methods, respectively. (b) A typical SEM image of fabricated aligned chirality-enriched films. Polarized photoluminescence of aligned (c) (6,5) and (d) (10,3) films, when optically pumped on resonance with the first interband transition. (e) False-colour SEM images of a thin-film transistor with a channel width of $\sim 5$ $\mu$m and channel length of $\sim 30$ $\mu$m made from an aligned and (6,5)-enriched SWCNT film. (f) Source-drain current versus source-drain voltage at zero gate voltage of the transistor, showing anisotropic conductivities of the aligned (6,5)-enriched thin-film transistor. (g) Source-drain current at a source-drain voltage of 1 V versus gate voltage of the (6,5)-enriched transistor. The on-current density is enhanced by a factor of 50 in a transistor made from larger-diameter semiconductor-enriched arc-discharge SWCNTs. Adapted from Ref. [60]
than that of their counterparts made from random films. As shown in Figures 7(f) & (g), the on-current density of the transistor in the parallel (perpendicular) direction is \( \sim 2 \text{ nA} \mu\text{m}^{-1} \) (\( \sim 80 \text{ pA} \mu\text{m}^{-1} \)), indicating that the on-current density can be improved by aligning the CNTs in one direction. The on-current density can be further enhanced by using larger diameter nanotubes, which is also demonstrated by our transistors based on semiconductor-enriched arc-discharge SWCNTs with an average diameter of 1.4 nm (Figure 7(g)). The device shows an enhancement of on-current density by \( \sim 50 \) times compared to the (6,5) CNT transistor at the same drain-source voltage. All fabricated devices display decent on-off ratios \( \sim 1 \times 10^3 \), which can be improved by using a high-\( \kappa \) dielectric material or using a top-gate structure.

![Diagram](image_url)

**FIG. 8.** 3D architectures made of aligned SWCNT films. (a) Nematic order parameter (\( S \), left axis) and the extinction ratio (ER, right axis), as a function of film thickness. (b) 1-inch and 2-inch aligned films made using controlled vacuum filtration. (c) A multilayer structure by stacking thin films of aligned SWCNTs and sandwiching doping molecules in-between. (d) Height profile of stacked films from one layer to four layers. (e) ER versus frequency in the THz range for 1–4 layers. (f) ER versus frequency in the THz range before and after doping. Adapted from Ref. [60, 70].

Obtained films can scale up both vertically and laterally and intercalate external molecules in between to form 3D architectures as shown by Komatsu et al. [70]. By adjusting the filtrated volume during the controlled vacuum filtration process, the thicknesses of obtained films can be simply controlled, and they all display good alignment (Figure 8(a)). The extinction ratio (ER) of THz polarized attenuation can be as high as \( \sim 120 \text{ dB/\mu m} \), thanks to the high packing density in the films. On the other hand, the lateral size of obtained films is determined by the filtration system size. For instance, a 2-inch system can produce a larger film, compared to 1-inch systems. These 2-inch macroscopic crystalline SWCNT films enable us to build 3D architectures, through both layer stacking and molecule intercalation, as shown in Figure 8(c). This process consists of simple manual layer stacking with careful control of angles between layers and chemical doping by either electron donors or acceptors for \( n \)-type or \( p \)-type doping, respectively [70]. These films are compatible with a wide spectrum of dopants, such as HNO\(_3\) [71], H\(_2\)SO\(_4\) [71, 72], NH\(_4\)S\(_2\)O\(_8\) [72], HCl [71], H\(_2\)SO\(_3\) [73], iodine solution [73], and benzyl viologen [74]. Figure 8(d) shows the height profile using AFM for four stacked layers, demonstrating this method as a consistent and well-controlled technique. The ER in the THz range increases as the film thickness and doping level increase, as shown in Figures 8(e) & (f). As the doping level (and thus Fermi level) is increased, the first interband transition is suppressed and eventually completely quenched; see Figure 8(g).
III. ABSORPTION OF PERPENDICULARLY POLARIZED LIGHT

Extremely confined 1D excitons with huge binding energies in semiconducting SWCNTs lead to a variety of optical properties with light polarized along the SWCNT axis, while there are limited investigations of optical properties for light polarized perpendicular to the SWCNT axis, especially in a macroscopic SWCNT ensemble. The macroscopic crystalline films fabricated through controlled vacuum filtration provide a unique opportunity for such study. In this section, two examples – intersubband plasmons and crosspolarized excitons – are given.

Optical selection rules govern the observable transitions under far-field excitation. For light polarized parallel and perpendicular to the SWCNT axis, selection rules are different. Figure 9(a) schematically shows the band structure of metallic and semiconducting SWCNTs, respectively. Arrows of different colors show representative allowed optical transitions for light polarized along the tube axis (blue) and perpendicular to the tube axis (red and yellow). Conduction and valence subbands in metallic (semiconducting) nanotubes are indicated as $C_M$ and $V_M$ ($C_S$ and $V_S$), respectively, where $i = 1, 2, 3, \cdots$. Each subband has a well-defined angular momentum with quantum number $n$ and an eigenvalue of the Pauli matrix $\sigma_x$ \cite{68, 80}. Light with polarization parallel to the nanotube axis can cause transitions with $\Delta n = 0$. For light polarized perpendicular to the nanotube axis, the following selection rules have to be satisfied:

1. $\Delta n = \pm 1$, meaning that the angular momentum quantum number has to change by $\pm 1$
2. $\Delta \sigma_x = 0$, meaning that the eigenvalue of the Pauli matrix $\sigma_x$ has to be conserved.

FIG. 9. Optical properties for light polarized perpendicular to the CNT axis. (a) Interband transitions for parallel polarization (blue arrows), interband transitions (yellow arrows) and intersubband transitions (red arrows) for perpendicular polarization in metallic (left) and semiconducting (right) SWCNTs. (b) The joint density of states for interband and intersubband transitions. (c) UV-vis-near infrared optical absorption spectra of an arc-discharge SWCNT film doped with increasing AuCl$_3$ concentration \cite{81}. (d) Cross-polarized photoluminescence of a CNT suspension \cite{82}. (e) Circular dichrosim of (6,5) SWCNT enantiomers \cite{83}. 

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In undoped or lightly doped SWCNTs, optical absorption spectra are dominated by \( i \)-conserving transitions (for example, the \( M_{11} \), \( S_{11} \), and \( S_{22} \) transitions) for parallel polarization. Transitions for perpendicular polarization, such as the lowest-energy transitions \( M_{12} \) and \( S_{12}/S_{21} \), are strongly suppressed due to the depolarization effect [67, 69]. These transitions generate crosspolarized excitons. In heavily doped SWCNTs, where the Fermi level \( (E_F) \) is inside the conduction (valence) band, perpendicularly polarized light can excite the \( C_{M1} \rightarrow C_{M2} \) \( (V_{M1} \rightarrow V_{M2}) \) transition in metallic SWCNTs and the \( C_{S1} \rightarrow C_{S3} \) and \( C_{S2} \rightarrow C_{S4} \) \( (V_{S1} \rightarrow V_{S3} \) and \( V_{S2} \rightarrow V_{S4} \) transitions) in semiconducting SWCNTs. As has been shown in III-V semiconductor quantum wells, such as GaAs quantum wells [84], these intersubband transitions are intrinsically collective, deviating from the single-particle picture and involving many-body effects, and are thus termed intersubband plasmons (ISBPs). Unlike ISBPs in quantum wells with oblique incident light excitation, ISBPs in 1D SWCNTs can be excited under normal incidence. Also, different from crosspolarized excitons, ISBPs are expected to be strong due to the concentrated joint density-of-states (see Figure 9(b)) as well as the suppression of the depolarization effect at high \( E_F \) [85, 86].

Previous studies of heavily doped SWCNTs have revealed a new peak [81, 87–89] in absorption spectra. For example, as SWCNT films are heavily doped by AuCl, a new peak emerges at \( \sim 1.12 \) eV, as shown in Figure 9(c). However, in these cases, the nanotubes were randomly oriented, precluding definitive interpretation of the origin of this peak. On the other hand, crosspolarized excitons are significantly reduced, and it is challenging to observe this feature in single-CNT spectroscopy and studies of randomly oriented CNTs. For example, crosspolarized photoluminescence of a CNT suspension [82] (Figure 9(d)) and circular dichroism of (6,5) SWCNT enantiomer suspensions [83] (Figure 9(e)) have revealed these excitons, but any quantitative analysis such as oscillator strength estimation has not been possible. The crystalline SWCNT films produced by the controlled vacuum filtration technique are compatible with various dopants and electrolyte gating techniques and allow for a direct measurement using macroscopic spectroscopy techniques. These samples enable us unambiguous determination of these optical features occurring when light is polarized perpendicular to the nanotube axis.

First, we discuss recently reported direct evidence for ISBPs in gated and aligned SWCNTs through polarization-dependent absorption spectroscopy by Yanagi et al., which elucidates the origin of the observed unknown feature in these previous studies on doped SWCNTs [80]. A film of macroscopically aligned and packed SWCNTs with an average diameter of 1.4 nm, containing both metallic and semiconducting SWCNTs, was prepared using controlled vacuum filtration. The \( E_F \) was tuned using electrolyte gating techniques [88, 89]. A schematic illustration of the experimental setup is shown in Figure 10(a). Figure 10(b) shows parallel-polarization optical absorption spectra for the film at three gate voltages, \( V_G = -2.0, 0.0 \) and \(+4.3 \) V. At \( V_G = 0.0 \) V, the interband transitions \( S_{11} \) and \( S_{22} \) in semiconducting nanotubes and the interband transition \( M_{11} \) in metallic nanotubes are clearly observed. As the gate voltage is increased in the positive (negative) direction, electrons (holes) are injected into the nanotubes through formation of an electric double layer on their surfaces. As a result, the \( S_{11}, S_{22}, \) and \( M_{11} \) peaks disappear at \( V_G = -2.0 \) and \(+4.3 \) V because of Pauli blocking.

Completely different behavior is observed in perpendicular-polarization spectra, as shown in Figure 10(c). As \( |V_G| \) is increased, a new absorption peak due to ISBP appears at \( \sim 1 \) eV, grows in intensity, and dominates the spectrum at the highest \( |V_G| \). The properties of the ISBP peak are entirely orthogonal to those of the \( S_{11}, S_{22}, \) and \( M_{11} \) peaks. Figure 10(d) shows detailed changes of the ISBP peak for this aligned film as the gate voltage is varied on the electron injection side. The peak first appears at 1 eV when the gate voltage reaches 3 V, and then its intensity gradually increases with increasing \( V_G \). In addition, the peak position blue shifts with further increasing \( V_G \), reaching 1.05 eV at 4.3 V.

Second, a systematic polarization-dependent optical absorption spectroscopy study of a macroscopic film of highly aligned single-chirality (6,5) SWCNTs was performed for the observation of crosspolarized excitons [90]. Recent advances in pH-controlled gel chromatography [91, 92] enables the preparation of a large-scale suspension of chirality-aligned single-chirality (6,5) SWCNTs was performed for the observation of crosspolarized excitons [90]. Recent industry-scale separation [91] is highly desirable for the vacuum filtration technique to produce 

\[
\begin{align*}
\text{A} & \quad \text{and subtracted by} \quad \text{A}, \quad \text{is presented. Further detailed analysis based on nematic order parameters revealed that the oscillator strength for} \quad E_{12} \quad (f_{12}) \quad \text{is} \quad 1/10 \quad \text{of the oscillator strength for} \quad E_{11} \quad (f_{11}) \quad [90].
\end{align*}
\]
FIG. 10. ISBPs in gated aligned SWCNT films. (a) Schematic diagram for the setup for the polarization-dependent absorption spectroscopy experiments on an aligned SWCNT film gated through electrolyte gating. $V_G$: gate voltage, $R$: reference voltage. (b) Parallel-polarization and (c) perpendicular-polarization optical absorption spectra at different gate voltages for an aligned SWCNT film containing semiconducting and metallic nanotubes with an average diameter of 1.4 nm. The ISBP peak appears only for perpendicular polarization for both electron and hole doping, corresponding to $V_G = -2.0$ and $4.3$ V, respectively. (d) Optical absorption spectra showing how the ISBP peak evolves as the gate voltage increases. Adapted from Ref. [80]

IV. MICROCAVITY EXCITON-POLARITONS IN ALIGNED SEMICONDUCTING SWCNTS

Strong coupling of photons and excitons inside a microcavity produces hybrid quasiparticles, microcavity exciton-polaritons, which continue to stimulate much interest. In a general model, a two-level system interacts with photons inside a microcavity, with the coupling strength $g$ and photon (matter) decay rate $\kappa$ ($\gamma$). Resonant coupling leads to a splitting into two normal modes with an energy separation, $\hbar \Omega_R$, known as the vacuum Rabi splitting (VRS), with $2g \approx \Omega_R$ when $g \gg \kappa, \gamma$. Depending on the relationship between $g$ and $\kappa, \gamma$, there are different regimes. The strong coupling regime is achieved when $4g^2/(\kappa \gamma) > 1$. Furthermore, the ultrastrong coupling regime arises when $g/\omega_0 > 0.1$, where $\omega_0$ is the resonance energy, whereas the deep strong coupling regime is defined as $g/\omega_0 > 1$.

Microcavity exciton-polaritons based on semiconductor quantum wells have been a model system for highlighting the difference between light-atom-coupling and light-condensed-matter-coupling. However, they remain in the strong coupling regime typically with $g/\omega_0 < 10^{-2}$. Furthermore, the fabrication of such devices usually requires sophisticated molecular beam epitaxy, and measurements have to be done at cryogenic temperatures. Excitons in organic semiconductors, possessing large binding energies and oscillator strengths, have displayed larger VRS at room temperature. Moreover, nanomaterials, such as transition metal dichalcogenides and semiconducting SWCNTs, have recently emerged as a new platform for studying strong-coupling physics under extreme conditions.
quantum confinement. Large oscillator strengths in these materials help relax the stringent requirement for a high quality factor (high-$Q$) photonic cavity. In particular for SWCNTs, Graf et al. first reported near-infrared microcavity exciton-polaritons by incorporating a film of polymer-selected chirality-enriched (6,5) SWCNTs of random orientation inside a Fabry-Pérot microcavity, simply consisting of two parallel metal mirrors [101]. Cavities of similar structures are widely used in organic molecule exciton-polaritons. A very large VRS, exceeding 100 meV, was observed to scale proportionally to $\sqrt{N}$, where $N$ is the number of SWCNTs inside the cavity, displaying a cooperative enhancement [101]. Furthermore, electrical pumping of exciton-polaritons was achieved by the same group [102], paving the way toward carbon-based polariton emitters and lasers.

More recently, Gao and coworkers developed a unique architecture in which excitons in an aligned single-chirality (6,5) CNT film interact with cavity photons in a polarization-dependent manner [103]. Again, they prepared aligned (6,5) films by vacuum filtrating suspensions separated using the aqueous two phase extraction method. The microcavity exciton-polariton devices were created by embedding the obtained aligned film with thickness $d$ inside a Fabry-Pérot cavity. By changing the incidence angle ($\theta$ in Figure 12(a)), the resonance frequency of the microcavity was tuned to resonate with the $E_{11}$ or $E_{22}$ of (6,5) SWCNTs. Furthermore, adjustment of the angle between the incident light polarization direction and the SWCNT alignment direction ($\phi$ in Figure 12(a)) provided a convenient knob to probe the directional variation of $g$. The angle $\theta$ has a one-to-one correspondence with the in-plane wave vector $k_{\parallel}$, so the $\theta$ dependence of transmission peaks can be converted into $k_y$ or $k_z$ dependence for $\phi = 0^\circ$ or $90^\circ$, respectively, to map out polariton dispersions. Figures 12(b) & (c) show experimental and simulated in-plane dispersion relationships for a device with $d = 8$ nm for $\phi = 0^\circ$ and $90^\circ$, respectively. A prominent VRS of $137 \pm 6$ meV ($g/\omega_0 > 5.5\%$) is observed, while no splitting is seen at $\phi = 90^\circ$. Here, $\omega_0$ is the resonance frequency of $E_{11}$. Furthermore, any value of VRS between zero and its maximum value can be selected, on demand, by setting the polarization angle $\phi$. 

![FIG. 11. Cross polarized excitons in aligned single-chirality (6,5) SWCNT films. (a) A photograph of separated high-purity (6,5) suspension using the gel chromatography technique. (b) Absorption spectrum in the near-infrared and visible range for this suspension. The purity is estimated to be $> 97\%$. (c) Polarization-dependent attenuation spectra for the aligned (6,5) SWCNT film at polarization angles of $0^\circ$, $30^\circ$, $45^\circ$, $60^\circ$, and $90^\circ$ with respect to the nanotube alignment direction. (d) Comparison of spectra at $0^\circ$ ($A_\parallel$) and $90^\circ$ ($A_\perp$). The blue line indicates $3.2A_\perp - A_\parallel$. Adapted from Ref. [90]
FIG. 12. Microcavity exciton polaritons based on aligned semiconducting SWCNTs. (a) The two angles that are continuously scanned in the present experiments - the incident angle $\theta$ and the polarization angle $\phi$. The latter is the angle between the SWCNT alignment direction (x axis) and the incident light polarization direction. (b) Anisotropic dispersions of microcavity exciton-polaritons in the $E_{11}$ region for $\phi = 0^\circ$ and $\phi = 90^\circ$. A clear VRS is observed at $\phi = 0^\circ$, while a photon dispersion with no splitting is observed at $\phi = 90^\circ$. (c) Corresponding simulated anisotropic dispersions in the $E_{11}$ region. (d) Experimental transmittance spectra at zero detuning ($\theta \sim 39^\circ$) for various polarization angles $\phi$ from 0° to 90° for a device working in the $E_{11}$ region using an aligned (6,5) SWCNT film with $d = 8$ nm. (e) Dispersion surfaces of the upper polariton (UP) and lower polariton (LP) (colored surfaces) for the device in (d). (f) VRS at $\phi = 0^\circ$ versus the square root of the film thickness, demonstrating the $\sqrt{N}$-fold enhancement of collective light-matter coupling. Adapted from Ref. [103].

Since the in-plane wavevector ($k_x, k_y$) has one-to-one correspondence with ($\phi, \theta$), the continuous adjustment of $\phi$ and $\theta$ maps out the full anisotropic SWCNT exciton polariton dispersion surfaces, as demonstrated in Figure 12(e) for the 8-nm-thick device. The most striking feature of Figure 12(e) is the appearance of two circular arcs on which the energy is constant; namely, the upper polariton (UP) and lower polariton (LP) branches coalesce to form these constant-energy arcs in momentum space. Furthermore, Gao et al. analyzed the obtained spectra using quantum Langevin equations, including losses, and demonstrated that the end points of each arc are exceptional points, that is, spectral singularities that ubiquitously appear in open or dissipative systems. Finally, Figure 12(f) confirms that VRS scales with $\sqrt{d}$ and thus $\sqrt{N}$. The VRS value for the 64-nm-thick device was 329 meV ($g/\omega_0 > 13.3\%$), which is the largest value ever reported for exciton polaritons based on Wannier excitons.

V. PLASMON RESONANCES AND HYBRIDIZATION IN PATTERNED ALIGNED SWCNTS

The longitudinal confinement of free carriers in finite-length SWCNTs leads to a prominent broad absorption peak in the THz and infrared ranges [60, 66], as shown in Figure 6(b). Temperature-dependent broadband spectroscopy unambiguously revealed the plasmonic nature in films of randomly oriented SWCNTs [66]. Resonance energy depends on the length of SWCNT and can be tuned by adjusting the nanotube length through sonication processes of different
FIG. 13. Plasmon resonances and plexcitons in patterned aligned SWCNTs. (a) Schematic diagram of plasmon resonances in patterned aligned SWCNTs. A SEM image shows a fabricated device. (b) Attenuation spectra for strips of aligned SWCNTs with different $L$. The peak energy is $\nu_p$. (c) Attenuation spectra for strips of aligned SWCNTs with different doping levels. (d) The extinction spectra of patterned films of highly doped aligned SWCNTs (top) and annealed aligned SWCNTs (bottom). (e) Resonance energies as a function of wavevector $q$, which is defined as $\pi/L$. The solid line is a fit to $\omega_p \propto \sqrt{q}$ (orange lines). Peak energies of the plexcitons, showing an anticrossing (blue lines). Adapted from Ref. [65] and Ref. [64].

duration [104]. However, due to a broad length distribution in these films, the linewidth of the resonance is usually broad and the resonance peak position is up to $\sim 400 \text{ cm}^{-1}$ in the far-infrared. Crystalline SWCNT films produced using the controlled vacuum filtration method, which are compatible with various micro/nanofabrication techniques such as lithography and etching, provide a unique platform for unifying CNT lengths for high quality plasmon resonators. The longitudinal confinement, determined by the lithography-defined dimensions, can range from tens to hundreds of nanometers. In combination with chemical doping techniques, the resonance frequency can be broadly tuned from the THz to near-infrared.

Figure 13(a) illustrates the bounded electron oscillation in SWCNTs and a representative SEM image of patterned strips of aligned films of SWCNTs produced through controlled vacuum filtration reported by Chiu et al. [65]. For strips of a 165-nm-thick film, there are strong attenuation peaks ($\nu_p$) in the mid-infrared range. As the lateral dimension ($L$) decreases from 800 nm to 100 nm at fixed film thickness ($t$), $\nu_p$ blue-shifts and $Q$ increases up to 9; see Figure 13(b). This increase of the $Q$ factor reflects the increased length uniformity in smaller-$L$ devices. Furthermore, exposure to
strong dopants, such as HNO$_3$, increases the film conductivity and pushes $\nu_p$ further to the near-infrared, as shown in Figure 13(c). The presence of dopants, however, reduces the $Q$ factor from 4.2 to 2.4, because of an increase of scattering.

As dopants are introduced and the $E_F$ is elevated, the lowest-energy excitons in semiconducting SWCNTs, $S_{11}$, in these films are switched off; see Figure 8(g) and Figure 9(a). There is a single resonance peak as $L$ is tuned, as shown in the top panel of Figure 13(e). After films are annealed to remove excess dopants, exciton transitions are activated with the $S_{11}$ transition energy at $\omega_0 = 0.66$ eV. As $L$ is continuously adjusted and thus $\nu_p$ is continuously tuned across $\omega_0$, strong coupling between plasmon and exciton resonances emerges; see the bottom panel of Figure 13(d). The formed new quasiparticles are known as plexcitons. Figure 13(e) demonstrates the peak energies for the LP branch, UP branch, and bare plasmon resonances in highly doped films. An extremely large VRS $\hbar\Omega \sim 485$ meV ($g/\omega_0 \sim 36.7\%$) is observed, indicating ultrastrong coupling. A Hopfield-like Hamiltonian can fully capture the polariton dispersions [64]. Moreover, the coupling strength can be tuned by adjusting the doping level and film thickness. The increase of doping level, thus film conductivity, decreases the oscillator strength of $S_{11}$ and the coupling strength.

![Figure 14](image14.png)

**FIG. 14.** Phonon-plasmon-polaritons in patterned aligned SWCNTs. (a) The attenuation spectra of strips of aligned SWCNTs with different lengths on the 40-nm diamondlike carbon substrate. (b) An expanded view of the dashed rectangle in Figure 14(a), along with fits to Fano functions (black dashed lines). (c) The resonant frequencies, $\nu_1$ and $\nu_3$, as a function of $q$, on substrates with different thicknesses of diamondlike carbon. Adapted from Ref. [105].

In addition to plexcitons, these plasmon resonances in strips of aligned SWCNTs can strongly couple with other quasiparticles, such as phonons. Falk et al. showed strong coupling between longitudinal optical phonons in SiO$_2$ substrates (attenuation peak $\nu_3$ in Figure 14(a)), infrared-active $E_1$ and $E_{1u}$ phonons in CNTs [106, 107] (attenuation peak $\nu_2$ in Figure 14(a)) and plasmon resonances (attenuation peak $\nu_1$ in Figure 14(a)), by putting patterned strips with various $L$ of aligned SWCNT films on SiO$_2$ substrates. The strong coupling enhances usually weak infrared-active phonons and leads to an asymmetric lineshape, which can be fit very well with a Fano resonance formula [105], as shown in Figure 14(b). Peak positions of $\nu_1$ and $\nu_3$ demonstrate a clear anticrossing behavior, as displayed in Figure 14(c).
VI. CONCLUSION

This review summarizes the recent achievement of fabricating wafer-scale crystalline CNTs and the discovery of new science and applications in these new materials, specifically in photonics and optoelectronics. These wafer-scale crystalline CNTs with control of chirality distribution fabricated through controlled vacuum filtration open new pathways toward both fundamental studies and various applications in a diverse set of disciplines. This technique addresses the grand "bottom-to-top" challenge in the CNT community for utilizing extraordinary properties of individual CNTs at the nanoscale to macroscopic applications. We anticipate that the initial studies described in this review will further stimulate interest in CNT research and push toward real-world applications employing CNTs.
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