



Carbon Nanotube Actuators

Ray H. Baughman, *et al. Science* **284**, 1340 (1999); DOI: 10.1126/science.284.5418.1340

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recently reported by Gumbsch and Gao (26). However, such processes, although similar to shear crack growth in the sense that they are driven by shear stresses, are also clearly different in the sense that they do not involve the breaking of material bonds and the subsequent creation of new surfaces. Moreover, no experimental confirmation of such intersonic processes is currently available. To date, our experimental observations of intersonic processes have been limited to cracks along bimaterial interfaces (27) and along the fibers in unidirectional composites (28). However, in both situations the crack speed was subsonic with respect to one of the two constituents, the material with higher wave speed in the case of bimaterials and the fibers in the case of unidirectional composites. The experiments presented here provide unambiguous evidence that shearing mode conditions near a propagating crack tip can drive the crack to intersonic speeds, which are found to be possible even in purely homogeneous systems with only one distinct set of wave speeds.

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strength of the bond was systematically varied from 30 to 50% of the strength of the bulk material. This shear bond strength was measured with the use of a conventional losipescu shear test fixture.

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Carbon Nanotube Actuators

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Electromechanical actuators based on sheets of single-walled carbon nanotubes were shown to generate higher stresses than natural muscle and higher strains than high-modulus ferroelectrics. Like natural muscles, the macroscopic actuators are assemblies of billions of individual nanoscale actuators. The actuation mechanism (quantum chemical-based expansion due to electrochemical double-layer charging) does not require ion intercalation, which limits the life and rate of faradaic conducting polymer actuators. Unlike conventional ferroelectric actuators, low operating voltages of a few volts generate large actuator strains. Predictions based on measurements suggest that actuators using optimized nanotube sheets may eventually provide substantially higher work densities per cycle than any previously known technology.

The direct conversion of electrical energy to mechanical energy through a material response is critically important for such diverse needs as robotics, optical fiber switches, optical displays, prosthetic devices, sonar projectors, and microscopic pumps. Ferroelectric and electrostrictive materials are especially useful for direct energy conversion. However, applications are restricted by the maximum allowable operation temperature, the need for high voltages, and limitations on the work density per cycle.

Conducting polymer actuators based on electrochemical dopant intercalation were proposed a decade ago (1) and have been pioneered in many different laboratories (2, 3). Faradaic processes for these batterylike devices involve solid-state dopant diffusion and structural changes that limit rate, cycle life, and energy conversion efficiencies.

The new actuators we describe here use

*To whom correspondence should be addressed. Email: ray.baughman@alliedsignal.com carbon single-walled nanotube (SWNT) sheets (4) as electrolyte-filled electrodes of a supercapacitor. No dopant intercalation is required for device operation. Changing the applied voltage injects electronic charge into a SWNT electrode, which is compensated at the nanotube-electrolyte interface by electrolyte ions (forming the so-called double layer). The new actuators use dimensional changes in covalently bonded directions caused by this charge injection, which originate from quantum chemical and double-layer electrostatic effects. For low charge densities, calculations (5) and experimental results (6) for charge transfer complexes of graphite and conducting polymers (Fig. 1) show that the strain due to quantum mechanical effects (changes in orbital occupation and band structure) changes sign from an expansion for electron injection to a contraction for hole injection. However, expansion results from both quantum chemical effects and electrostatic double-layer charging for high-density charge injection of either sign (5, 6).

Like natural muscle (7), the nanotube sheet actuators are arrays of nanofiber actuators. A novel type of actuation results because the SWNTs add high surface area to the mechanical properties, electrical conductivity, and charge transfer properties of graphite. The results presented here are for nanotube sheets ("nanotube paper") composed of mats of nanotube bundles joined by mechanical entangle-

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ment and van der Waals forces along incidental points (and lines) of contact. These observations enable prediction of the actuator properties that are ultimately obtainable for nonbundled nanotubes, which have extraordinarily high surface area and mechanical properties (8). The concept of using either nanotube sheets or single nanotubes as actuators, and the effect of nanotube bundling, are schematically illustrated in Fig. 2.

The commercially obtained nanotubes were made by the dual pulsed-laser vaporization method and purified by a published method that involves nitric acid reflux, cycles of washing and centrifugation, and cross-flow filtration (4, 9). Materials made by this method consist of hexagonally packed bundles of carbon nanotubes that have diameters of 12 to 14 Å, an intertube separation within a bundle of ~ 17 Å, an average bundle diameter of ~100 Å, and lengths of many micrometers (4, 10). The nanotube sheets were formed by vacuum filtration of a nanotube suspension on a poly(tetrafluorethylene) filter, and the dried nanotube sheets were peeled from the filter (4, 9). These freestanding sheets (composed of highly entangled nanotube bundles) were used as they were obtained, without any effort to optimize the mechanical properties, surface area, or electrical conductivity that are important for the actuator application.

The electrochemical charge injection process was characterized by measurement of the gravimetric capacitance ($C_{\rm G}$) of the nanotube paper in various electrolytes [all potentials are versus a saturated calomel electrode (SCE)]. $C_{\rm G}$ was derived from the slope of specific current versus voltage scan rate in a cell containing nanotube paper for both a working electrode and a counterelectrode. After a few cycles to ensure the maximum degree of electrode wetting, the measured $C_{\rm G}$ increased up to about 15 F/g for aqueous solutions of both 1 M NaCl and 38% by weight H₂SO₄. Similar $C_{\rm G}$ s were observed for 1 M LiClO₄ in acetonitrile (12 to 17

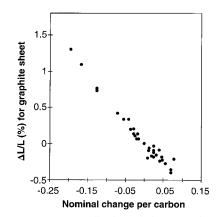


Fig. 1. Experimentally derived changes in basal plane strain as a function of nominal charge derived from data (6) for graphite intercalation compounds.

F/g) or in propylene carbonate (13 to 14 F/g), and a higher $C_{\rm G}$ was observed for a 5 M aqueous KOH solution (30 F/g), which readily wet the nanotube paper.

These measurements and estimates of the available surface area suggest that double-layer charge injection for the nanotube bundles is similar to that for the basal plane of graphite. The gravimetric surface area of the nanotube paper ($A_{\rm G}$) measured by the Brunauer-Emmett-Teller (BET) method is about 285 m²/g (11); this value is near the 300 m²/g surface area of 100 Å diameter cylinders having the crystallo-graphically derived bundle density of 1.33 g/cm³ (10), which suggests that the N₂ in the

BET measurement does not access significant surface area in the space between nanotubes in a bundle. An areal capacitance (C_A) of 4 to 10 μ F/cm² results for the above range of observed gravimetric capacitances and an A_G of 300 m²/g. This value is in the approximate range of measured C_A s for the basal plane of graphite (12), which has a minimum at ~3 μ F/cm² (for either 0.9 M aqueous NaF or 0.2 M tetrapropylammonium tetrafluoride solution in acetonitrile).

Our first demonstration of a carbon nanotube actuator was surprisingly simple to produce. Similar to a design for a polyaniline conducting polymer actuator (3) (Fig. 3), this ac-

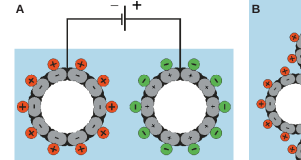
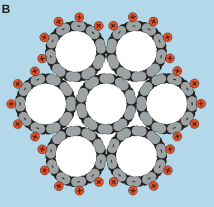
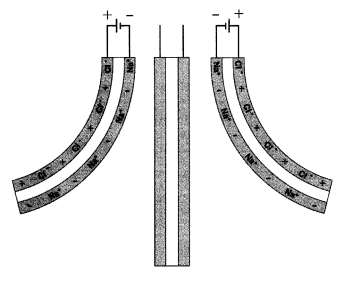


Fig. 2. Schematic illustration of charge injection in a nanotube-based electromechanical actuator and the effect of nanotube bundling. (A) An applied potential injects charge of opposite



sign in the two pictured nanotube electrodes, which are in a liquid or solid electrolyte (blue background). These indicated charges in each electrode are completely balanced by ions from the electrolyte (denoted by the charged spheres on each nanotube cylinder). The pictured single nanotube electrodes represent an arbitrary number of nanotubes in each electrode that mechanically and electrically act in parallel. Depending on the potential and the relative number of nanotubes in each electrode, the opposite electrodes can provide either in-phase or out-of-phase mechanical deformations. (B) Charge injection at the surface of a nanotube bundle is illustrated, which is balanced by the pictured surface layer of electrolyte cations. Although penetration of electrolyte and gases into interstitial sites and nanotube cores might occur for the investigated bundled nanotubes, this has no noticeable effect on the observed BET surface area (11) or the measured electrode capacitance.

Fig. 3. Schematic edgeview of a cantilever-based actuator operated in aqueous NaCl, which consists of two strips of SWNTs (shaded) that are laminated together with an intermediate layer of double-sided Scotch tape (white). The Na $^+$ and Cl $^-$ ions in the nanotube sheets represent ions in the double laver at the nanotube bundle surfaces, which are compensated by the indicated charges that are injected into the nanotube bundles. The equality between the lengths of the two nanotube sheets (center) is disrupted when a voltage is applied, causing the indicated actuator displace-



ments to the left or right. The possible existence of a small amount of double-layer charge before the application of a voltage is ignored.

tuator response was approximately linear, with

applied voltage between -0.4 and 0.1 V (with

negligible hysteresis) (Fig. 5), but both a non-

linear voltage dependence of displacement on

potential and hysteretic behavior occurred at

higher potentials. This actuator has a resonance

peak at ~ 25 Hz when driven in the liquid

electrolyte with an ac voltage (~ 2 V from peak

to peak between the working electrode and the

cantilever actuators depends on the difference

in the electrically induced expansion of oppo-

site actuator electrodes, characterization of

these devices does not directly provide the elec-

tromechanical response of the nanotube paper.

Thus, we directly measured the change in

length of the nanotube paper versus the poten-

tial in 1 M NaCl. One end of a vertical nanotube

paper strip was anchored to the bottom of the

electrolyte bath, and the other end was attached

to a horizontal cantilever (Fig. 4B). Changes in the length of the nanotube paper strip deflected

the cantilever, whose movement was measured

with an attached mirror and the above optical

sensor. Like the field dependence for ferroelec-

trics, plots of deformation versus potential (Fig.

6A) show increasing hysteresis with increasing

potential range during cyclic voltametry. The

direction of strain change reverses at 0.56 V

when potential is increased and at 0.27 V when

potential is decreased for the highest voltage

scan. Figure 6, B and C, show that the actuator

devices can be effectively switched in 1 s.

Figure 6C shows that an actuator response,

albeit small, can be observed to >1 kHz. Cy-

cling a strip of nanotube paper at a constant

scan rate for 140,000 cycles (between ± 0.5 V

at 1 Hz) resulted in a \sim 33% decrease in actua-

tor stroke. A stress (0.75 MPa) significantly

higher than the peak capacity of human skeletal

muscle (0.3 MPa) was generated during the

isometric contraction of a nanotube sheet [using

Optical

sensor

Mirror

SCE

Electrolyte

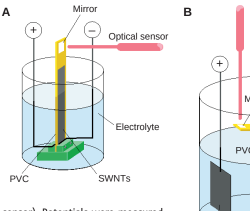
Because the response of the above bimorph

counterelectrode).

tuator consisted of cut strips of nanotube paper (3 mm by 20 mm and 25 to 50 µm thick) that were adhesively applied to opposite sides of a 10% larger piece of Scotch Double Stick Tape. An insulating plastic clamp at the upper end of the three-layer strip supported the device vertically in 1 M NaCl electrolyte and held two platinum foil electrical leads in contact with the nanotube paper sheets. When a dc potential of a few volts or less was applied to these leads, a deflection of up to about a centimeter was observed for the cantilever tip, which reversed on reversal of the potential (Fig. 3). When a square wave potential was applied, oscillation was visually observed up to at least 15 Hz as the actuator pushed the electrolyte back and forth. Similar operation was observed for this type of actuator made from nanotube paper that had been annealed at 1100°C for 1 hour in flowing argon.

The above actuator is called a bimorph cantilever actuator because the device response depends on the difference in the electrically induced expansion of opposite actuator electrodes. In order to obtain more quantitative results, we made a bimorph cantilever actuator in which two nanotube sheets having the above strip dimensions worked to bend a strip of poly(vinylchloride) (PVC) film that was an order of magnitude thicker (215 µm thick) (Fig. 4A). This thick film (10% wider and 20% longer than the nanotube paper strips) was laminated on both sides with nanotube paper strips by means of an intermediary layer of Scotch Double Stick Tape. The lower end of the actuator (containing the nanotube paper strips) was mounted in the bottom of the electrolyte bath (1 M NaCl) with a clamp that provided electrical contacts. The PVC strip from the actuator core extended above the electrolyte and supported a sputtered gold mirror on one strip face. The movement of this mirror as a result of actuator tip displacement was measured as a function of electrode potential with a

Fig. 4. (A) The apparatus used for characterizing an actuator, consisting of strips of SWNT paper that are symmetrically laminated (with an intermediate layer of doublesided Scotch tape) on opposite sides of a strip of PVC. Differential changes in the lengths of the nanotube strips bend the actuator, thereby causing a displacement of the mirror (which is



measured with the optical sensor). Potentials were measured versus a reference SCE, which is not pictured. (B) The apparatus used for the measurement of the dependence of SWNT sheet length on applied potential (versus a SCE). Changes in the length of the nanotube strip bend the PVC cantilever that

carries an attached mirror, whose displacement is measured with the optical sensor.

a 0.1-Hz square well potential between -0.5 and 1.5 V and the measurement apparatus described by Della Santa *et al.* (2)].

The dependence of strain on potential for the nanotube sheets (Fig. 6A) is quasi-parabolic, like that observed in electrocapillary measurements on electrodes of activated carbon and exfoliated graphite (12). This response is expected if surface energy changes due to doublelayer charging provide the mechanical stresses that elongate the material. If electrostatic effects dominate, the electrode dimension would be a minimum at the potential of zero charge (pzc), and either positive or negative charge injection would result in approximately equivalent expansion. In contrast, the minimum length occurs for the nanotube paper at a much higher potential (0.27 V for reduction and 0.56 V for oxidation) than the expected pzc for the 1 M NaCl solution [about -0.2 V versus a SCE, corresponding to the pzc for the basal plane of graphite (12)]. The implication is that quantum chemical effects are greater than classical coulombic effects for the carbon nanotube sheets. This result is consistent with the observation that the high-frequency tangential vibration mode of SWNTs increases frequency upon acceptor doping and decreases frequency upon donor doping (13), corresponding to bond shrinkage and expansion, respectively. Theory also predicts that quantum chemical effects dominate over electrostatic effects for charge transfer complexes of conducting polymers and graphite (5). This quantum chemical effect is large for metals having conjugated π bonded backbones and is negligible for elemental metals.

Although nanotube bundling reduces strain, the maximum observed actuator strains are still large: >0.2%. The dimensional changes of the covalent bond network along the nanotube direction are the likely origin of these strains. Evidence that these observed changes in the sheet dimension are proportional to

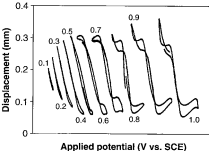
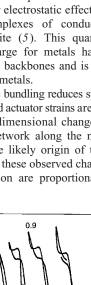


Fig. 5. Measured cantilever displacement (arbitrary origin) versus electrode potential (in 1 M NaCl) for a cantilever bimorph (Fig. 4A) in which two 15-µm-thick nanotube paper films bend a 215-µm-thick film of PVC. The potential range was symmetric about 0 V (versus a SCE), and the scan rate was 50 mV/s. The labels on the curves indicate the maximum applied potential.



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SWNTs

changes in the nanofiber length is provided by the observation of an in-plane thermal expansion coefficient that is an order of magnitude smaller than expected for a van der Waals bonded direction (14).

The actuator response can be described by a strain-voltage coefficient $S_{\rm v}$, which is the ratio of change in actuator strain to the change in applied electrode potential. The data in Fig. 6A yield an S_V of 0.032%/V in the potential range from -0.2 to 0.2 V, which increases to 0.11%/Vin the nearly linear region between -0.8 and -0.2 V. Ignoring the dependence of observed specific capacitance on potential (which appears to be small), these $S_{\rm V}$ values can be used to derive the strain-charge coefficient (ΔL / $L\Delta y$), where $\Delta L/L$ is the fractional change in length caused by a Δy change in the concentration of injected charge per monomer unit (1). The relation used is $S_{\rm V} = C_{\rm A}A_{\rm G}M(\Delta L/L\Delta y)/N$, where M is the monomer molecular weight and N is the number of coulombs of charge in a mole of electrons. Using the observed gravimetric capacitance for a 1 M NaCl solution ($C_{\rm G}$ $= C_A A_G = 15$ F/g) and the above strain coefficients, the calculated $\Delta L/L\Delta y$ increases from 0.17 (for -0.2 to 0.2 V) to 0.59 (for -0.8 to -0.2 V). Even the former coefficient is greater than that derived for the basal plane of graphite from diffraction measurements on donor complexes (0.066) (5, 6). The existence of a larger strain-charge coefficient for SWNTs than for graphite is supported by the observation of greater stiffening of the tangential mode for SWNTs (relative to that for the graphite intralayer mode) upon acceptor doping (13).

We believe that only the nanotubes close to the bundle surface undergo substantial charging, so it is conceivable that these nanotubes elongate whereas the nanotubes in the bundle interior undergo little dimensional change. If different expansion of surface and interior nanotubes occurs, the above derived straincharge coefficients would be upper bounds on the correct values. However, considering the very high aspect ratio of the nanotubes, it seems more likely that interfiber contact over long fiber distances will ensure that surface nanotubes and interior nanotubes undergo comparable elongations. In such a case, nonuniform charge distribution will affect the strain-voltage coefficient only to the extent that the straincharge coefficient varies between differently charged regions of a nanotube bundle.

These results enable prediction of the actuator performance that may ultimately be obtained for separate (nonbundled) SWNTs. Isolated nanotubes having the observed van der Waals diameter of 17 Å would have a surface area of ~1600 m²/g, which is about five times greater than for the investigated sheets consisting of nanotube bundles. If we use this surface area, the largest observed specific interfacial capacity (10 μ F/cm² for 5 M KOH), and the strain-charge coefficient ($\Delta L/L\Delta y = 0.59$ for -0.8 to -0.2 V) derived with the 1 M NaCl electrolyte, we obtain $C_{\rm G} = 160$ F/gm and S = 1.2%/V for nanotube sheets composed of nonbundled nanotubes. The predicted maximum strain ($\varepsilon_{\rm m}$) of ~1% for operation in aqueous electrolytes (with a 1-V operation range) and the substantially higher strain predicted for organic electrolytes are consistent with the 2% maximum variation in basal plane dimension that is observed for graphite intercalation compounds (Fig. 1). For comparison, presently used high-modulus ferroelectric ceramics provide a maximum strain of ~0.1%.

In addition to having a large displacement, a high-performance actuator material should have both high strength and high modulus. Based on theoretical and experimental results (15), a close-packed SWNT bundle composed of the investigated nanotubes should have a Young's modulus (Y) of ~ 0.64 TPa (6.1×10^6 kgf/cm²), which is more than two orders of magnitude greater than measured for our primitively constructed nanotube sheets (~ 1.2 GPa). The volumetric and gravimetric work densities per cycle for a matched mechanical load are proportional to $\frac{1}{2}Y\varepsilon_m^2$ and $\frac{1}{2}Y\varepsilon_m^2/\rho$, respectively, where ρ is the density (1.33 g/cm³) of the nanotube bundles (16). The above extraordinarily high modulus inherent for the single nanotubes, combined with the above predicted actuator strain of ~ 1 %, yield a predicted volumetric work capacity that is ~ 29 times higher than has been tabulated (16) for the best known ferroelectric, electrostrictive, and magnetostrictive materials. This predicted advantage of an ideal nanotube actuator increases to a factor of \sim 150 for gravimetric work capacity, which might be approached for nanoscale actuators based on separated nanotubes.

The origin of such spectacular predicted work capacities requires further explanation. As for ferroelectrics (17), the maximum work capacity per cycle is the product of the input electrical energy and a factor that increases from $\mathbf{k}^2/4$ to $\mathbf{k}^2/2$ with an increasing electromechanical coupling factor (\mathbf{k}^2) . Because \mathbf{k}^2 is the gravimetric ratio of stored mechanical energy to input electrical energy, the upper limit is approximately $\mathbf{k}^2 = (\frac{1}{2} S^2 V^2 Y/\rho)/$ $(\frac{1}{2}C_{\rm G}V^2) = S^2 Y / \rho C_{\rm G}$, which is linearly proportional to $A_{\rm G}$. If we use the above \hat{S} = 1.2%/V and $C_{\rm G}$ = 160 F/gm for separated nanotubes and the same Y/ρ as for bundled nanotubes, $\mathbf{k}^2 = 0.43$, which is comparable to that of high-performance ferroelectrics (17). The giant predicted work capacities per cycle are almost entirely a consequence of the greater electrical energy storage densities as compared with those of ferroelectrics.

Even if the mechanical property optimization for sheets of nonbundled nanotubes results in a Y/ρ value that is two orders of magnitude lower than experimentally derived (15) for individual nanotube bundles, the predicted gravimetric work capacity still exceeds that of any alternative actuator material. However, it must be emphasized that the present nanotube sheets are primitively constructed initial materials that are far from providing the surface areas and mechanical properties required for such predicted performance. Also, high power densities are required for many applications, and the presently achieved response rates of nanotube sheets are problematic for high-frequency applications. Additionally, although the observed 0.01-

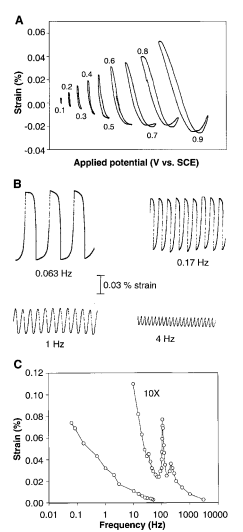


Fig. 6. Actuator characteristics measured by using the length changes of a nanotube sheet electrode to bend a cantilevered beam (Fig. 4B). Electrode potentials are versus a SCE for a 1 M aqueous NaCl electrolyte. (A) The strain (arbitrary origin) of a nanotube sheet versus applied potential. The potential range was symmetric about 0 V (versus a SCE), and the scan rate was 100 mV/s. The labels on the curves indicate the maximum applied potential. (B) The time dependence of the strain in a nanotube sheet for an applied square wave potential of ± 0.5 V having the indicated frequencies. (C) The frequency dependence of strain in a nanotube sheet electrode for an applied square wave potential of ± 0.5 V. The frequency dependence of strain at higher frequencies (where the strain scale is amplified 10-fold) show resonances arising from the cantilever beam.

ms time constant of a micrometer-sized electrode (18) (having a large counterelectrode) suggests that very high-frequency operation may be achievable for single nanotube actuators, this remains to be experimentally demonstrated. Our nanotube actuators are first prototypes, and much more work is required before the potential advantages of this technology can be experimentally assessed in practical devices.

Nevertheless, relatively modest improvements in the presently demonstrated actuator properties may enable such applications as electrically activated microcantilevers for medical catheter applications, where a low operation voltage is a major advantage. Using high-temperature electrolytes (molten or solid state), it may be possible to make nanotube actuators that function at temperatures far above those usable for ferroelectrics, which might be important for such applications as airflow control in jet engines. Also, demonstrations of nanotube actuator operation in seawater and in saline solutions, which are electrochemically like blood, suggest various possible marine and biomedical applications. Finally, nanotube actuators could be run in reverse to convert mechanical energy to electrical energy for mechanical sensor and energy-conversion devices. Ferroelectrics generate low currents at high voltages, but the nanotubes would provide high currents at low voltages. This effect is desirable for minimizing the effect of lead capacitances for remotely located sensors; for example, so that sensor-response amplifiers need not be located down-hole when doing seismology for oil exploration. If nanotubelike mechanical properties can be exploited to provide large inputs of mechanical energy, the output volumetric (or gravimetric) electrical energy per cycle would be much higher than that produced by alternative technologies. This feature, as well as the feasibility of operation at low frequencies, may eventually enable such applications as the conversion of the mechanical energy of ocean waves to electrical energy.

References and Notes

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- 9. The SWNTs were commercially obtained as an aqueous suspension from Tubes@Rice (Rice Univ., Houston, TX). The nanotube sheets were typically made by vacuum filtration of ~20 ml of a ~0.6 mg/ml nanotube suspension through a poly(tetrafluoroethylene) filter (Millipore LS, 47 mm in diameter, 5-μm pores). The nanotube sheet (formed over the clear funnel area, which was 37 mm in diameter) was washed with ~200 ml of deionized water and then 100 ml of methanol to remove residual NaOH and surfactant, respectively. These sheets were allowed to dry under continued vacuum purge for ~1 hour before being peeled from the filter. The typical nanotube sheet was between 15 and 35 μm thick and

weighed ${\sim}12$ mg, providing a bulk density of 0.3 to 0.4 g/cm³ and a four-point probe electrical conductivity of ${\sim}5000$ S/cm.

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The Magnetic Excitation Spectrum and Thermodynamics of High-T_c Superconductors

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Inelastic neutron scattering was used to study the wave vector– and frequencydependent magnetic fluctuations in single crystals of superconducting YBa₂Cu₃O_{6+x}. The spectra contain several important features, including a gap in the superconducting state, a pseudogap in the normal state, and the muchdiscussed resonance peak. The appearance of the pseudogap determined from transport and nuclear resonance coincides with formation of the resonance in the magnetic excitations. The exchange energy associated with the resonance has the temperature and doping dependences as well as the magnitude to describe approximately the electronic specific heat near the superconducting transition temperature (T_c).

The parent compounds of the high transition temperature (high- T_c) cuprate superconductors are antiferromagnetic (AF) insulators characterized by a simple doubling of the crystallographic unit cells in the CuO₂ planes. When holes are doped into these planes, the long-range AF-ordered phase disappears and the lamellar cop-

To whom correspondence should be addressed. Email: piq@ornl.gov per oxide materials become metallic and superconducting with persistent short-range AF spin correlations (fluctuations). Although spin fluctuations in cuprate superconductors are observed for materials such as YBa₂Cu₃O_{6+x} [denoted as $(123)O_{6+x}$] at all hole-doping levels, x (1-6), the role of such fluctuations in the pairing mechanism and superconductivity is still a subject of controversy (7). The most prominent feature in the fluctuations for $(123)O_{6+r}$ is a sharp resonance, which for highly doped compositions appears below the superconducting transition temperature T_c at an energy of 41 meV (1-3). We show that the temperaturedependent resonance intensity is correlated with the electronic part of the specific heat of $(123)O_{6+x}, C^{el}(x,T)$ (8, 9), including the extended fluctuation regime ("pseudogap phase"), and that the pseudogap temperature T^ deter-

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