Individual single-wall carbon nanotubes (SWNT) have been used to realize molecular-scale electronic devices such as single-electron (1) and field-effect transistors (FET) (2). Several SWNT-based devices have been successfully integrated into logic circuits (3) and transistor arrays (4). However, the difficulty in precise localization and interconnection of nanotubes impedes further progress toward larger-scale integrated circuits.

Self-assembly based on molecular recognition provides a promising approach for constructing complex architectures from molecular building blocks, such as SWNTs, bypassing the need for precise nanofabrication and mechanical manipulations (5). Biology, with its inherent self-assembly capabilities (6–13), is particularly attractive for this task. Biological recognition has been imparted to carbon nanotubes (14–18), but their self-assembly into functional devices and circuits has not yet been demonstrated. We present a framework for self-assembly of carbon nanotube–based electronics using DNA and homologous genetic recombination. A semiconducting SWNT was localized at a desired address on a DNA scaffold molecule using homologous recombination by the RecA protein from *Escherichia coli* bacteria (19). DNA metalization, with the RecA doubling as a scaffold dsDNA molecule serving as a scaffold and a short, auxiliary single-stranded DNA (ssDNA) (20). The assembly process was guided by the information encoded in these DNA molecules. The short ssDNA molecule was synthesized so that its sequence is identical to the dsDNA at the designated location of the FET. RecA proteins were first polymerized on the auxiliary ssDNA molecules to form nucleoprotein filaments (Fig. 1, step i), which were then mixed with the scaffold dsDNA molecules. A nucleoprotein filament bound a dsDNA molecule according to the sequence homology between the ssDNA and the designated address on the dsDNA (Fig. 1, step ii). The RecA later helped localize a SWNT at that address and protect the covered DNA segment against metallization (13). Figure 2A displays an atomic force microscopy (AFM) image of a 500-base-long (~0.25 μm) ssDNA/RecA nucleoprotein filament bound to the homologous section in the middle of a 48,502 base pair long (~16 μm) λ-DNA molecule.

A streptavidin-functionalized SWNT was guided to the right location on the scaffold dsDNA molecule using antibodies to the bound RecA and biotin-streptavidin–specific binding (20) (Fig. 1, step iii), and the SWNTs were solubilized in water by micellization in SDS (21). The solubilized SWNTs were functionalized with streptavidin by nonspecific adsorption (15, 16). Fluorescence microscopy of SWNTs with fluorescence-labeled streptavidin indicates homogeneous coverage of the nanotubes with streptavidin. Primary antibodies to RecA were reacted with the product of the homologous recombination reaction, resulting in specific binding of the antibodies to the RecA nucleopro-
tein filament. Next, biotin-conjugated secondary antibodies, which have high affinity to the primary ones, were localized on the primary antibodies to RecA. Finally, the streptavidin-coated SWNTs were added, leading to their localization on the RecA via biotin-streptavidin–specific binding (Fig. 1, step iii). The DNA/SWNT assembly was then stretched on a passivated oxidized silicon wafer (20). An AFM image of a SWNT bound to a RecA-coated 500-base-long ssDNA localized at the homologous site in the middle of a scaffold k-DNA molecule is shown (Fig. 2B). The conducting carbon nanotube is distinguished from the insulating DNA by scanning conductance microscopy (22, 23). Figure 2, B and C, depicts topographic and conductance images, respectively, of the same area. The evident difference between the two images identifies the SWNT on the DNA molecule; the carbon nanotube is aligned with the DNA, which is almost always the case due to the stiffness of the SWNT and the stretching process. In rare cases (<5%), we find misaligned SWNTs bound to the RecA (fig. S1). In certain cases, we find ropes rather than single nanotubes, due to the difficulty in achieving stable suspensions of individual SWNTs in an aqueous solution (20).

After stretching on the substrate, the scaffold DNA molecule is metallized (20). The RecA, doubling as a sequence-specific resist, protects the active area of the transistor against metallization. We used a previously developed metallization scheme (13) in which aldehyde residues, acting as reducing agents, are bound to the scaffold DNA molecules by reacting the latter with glutaraldehyde. The aldehyde derivatization, done before the homologous recombination reaction, was shown to be compatible with the whole process (13). Highly conductive metallic wires were formed by silver reduction for negative gate voltages indicates resistance in series with the SWNT. The resistance is attributed to the contacts between the gold wires and the SWNT because the resistance of the DNA-templated gold wires is typically smaller than 100 Ω (13). Large-scale circuits will require better contacts (20). In some cases, the FET comprises nanotube ropes. These can be identified by their larger contrast in SEM imaging (Fig. 3B). The rope devices cannot be turned off by gate voltage, probably due to the fact that they contain metallic nanotubes in parallel with the semiconducting ones; both types are present in our samples. The metallic nanotubes cannot be depleted by the available electric field. As the gate bias is made more positive, the rope conduction decreases but saturates to a finite value (fig. S2). Different devices have somewhat different turn-off voltages. They all exhibit hysteresis in the drain-source current as a function of gate bias (fig. S3). Similar behavior is observed in carbon nanotube FETs fabricated by top-down approaches (25). Overall, we measured 45 devices from 3 different batches. Fourteen devices functioned as FETs with partial or full gating, 10 devices conducted but could not be gated (probably metallic SWNT ones, fig. S4), and the rest were mostly disconnected and sometimes shorted to the gate. Our samples contain a mixture of semiconducting and metallic SWNTs, a fact that limits the yield of functional FETs. This drawback can be mitigated by recently developed techniques for separating semiconducting SWNTs from metallic ones (26–28).

The realization of a SWNT FET in a test tube promotes self-assembly as a realistic strategy for the construction of carbon nanotube-based electronics. The approach developed here can be generalized, in principle, to form a functional circuit on a scaffold DNA network. Numerous molecular devices could be localized simultaneously at different addresses on the network and interconnected by DNA-templated wires (12). The RecA-based scheme is robust and general enough to allow flexibility in the integration of other active electronic components into circuits, as long as they are compatible with the biological reactions and the metallization process. Realization of a functional circuit will require improving the electronic properties of the transistor and individual gating to each device. The latter could be achieved by using a three-armed DNA junction as a template (13) with the SWNT localized at the junction and by developing a method for turning one of the arms into a gate.

Fig. 4. Electrical characteristics of the DNA-templated carbon nanotube FET. (A) Schematic representation of the electrical measurement circuit. (B) Drain-source current (I_D) versus drain-source bias (V_D) for different values of gate bias (V_G). V_C = −20 V blue), 1 V (red), 1.5 V (green), 2 V (blue).

References and Notes

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Fig. 3. A DNA-templated carbon nanotube FET and metallic wires contacting it. SEM images of SWNTs contacted by self-assembled DNA-templated gold wires. (A) An individual SWNT. (B) A rope of SWNTs. Bars, 100 nm.
Solid-liquid phase transitions are an everyday occurrence. As one of the state variables (such as temperature or pressure) approaches a phase transition point for the melting of a solid, there are equilibrium fluctuations that lead to density changes commensurate with the new phase. Fluctuations important to a collective phase transition under these conditions occur over distributed time and length scales in which the atomic details are washed out. By using short-pulsed lasers to deposit heat at a rate faster than the thermal expansion rate, it is possible to prepare extreme states of solid matter at temperatures well above the normal melting point (this will be referred to as the strongly driven limit). Under such conditions, the atomic configuration of the entire excited material volume can be modified on the ultrafast time scale, and the melting transition of the prepared state can be seen as a simple model for transition state processes in general.

An atomic-level view of such a process requires that the atomic configuration of the material be observed as it passes from the solid to the liquid state. Ideally, one would like to be able to fully resolve the relative atomic motions during the melting process. The information accessible through time-resolved diffraction experiments, where the observable is intimately connected with the atomic structure of the material (1), can approach such a description. The first experiment along these lines used electron diffraction combined with rapid laser heating, but lacked sufficient temporal resolution (20 to 100 ps) and structural sensitivity for an atomic-level perspective on the process (2). Important applications of time-resolved electron diffraction have recently provided atomic-level structural details of reactive intermediates in the gas phase occurring on a similar time scale (3–5). Approaches based on laser-driven x-ray plasma sources have provided improved temporal resolution (200 to 500 fs) but to date have an insufficient signal-to-noise ratio (structural sensitivity) to adequately resolve the atomic details (6–8). Here we describe an important advance in short-pulsed electron sources that has made it possible to attain femtosecond electron pulses with sufficient electron number density to study the structural evolution of aluminum (Al) as it undergoes an ultrafast laser-induced solid-liquid phase transition. This work gives an atomic-level perspective on one of the simplest transition state processes: the order-to-disorder phase transition involved in melting.

Previous studies of femtosecond laser irradiation of polycrystalline Al at high fluence have observed the onset of liquid-like dielectric properties within 500 fs after laser excitation (9). It was inferred that lattice melting had occurred on this time scale, which is shorter than the time required for the opticaly excited electrons in the solid to relax via lattice phonon scattering. A nonthermal mechanism for melting was thus proposed, wherein the lattice bonding softens because of the photoinduced changes in electron distribution. Similar observations have been made in other materials (10, 11); however, it is not possible to determine detailed quantitative information about the nuclear coordinates from the electronic response (probed in optical measurements), especially for these far-from-equilibrium conditions. Time-resolved diffraction offers a direct probe of the atomic structure of polycrystalline and amorphous samples by providing, through a simple transformation, the time-dependent (atomic) pair correlation function. Such a measurement is able to determine how the lattice structure maps onto the disordered state defining a liquid. In the case of Al, we were able to follow the loss of the long-range order that was present in the crystalline phase and the emergence of the liquid structure where only short-range atomic correlations were present; this transition occurred in 3.5 picoseconds for thin-film aluminum with an excitation fluence of 70 mJ/cm². As opposed to previous studies of strongly driven phase transitions, we were able to determine the onset of the liquid state and find that even in the strong perturbation limit, the phase transition in Al is propagated by thermally sampled