Soft nanocomposite electroadhesives for digital micro- and nanotransfer printing

Sanha Kim1,2, Yijie Jiang3, Kiera L. Thompson Towell2, Michael S. H. Boutiller1, Nigamaa Nayakanti1, Changhong Cao1, Chunxu Chen3, Christine Jacob1, Hangbo Zhao1, Kevin T. Turner3, A. John Hart1*

Automated handling of microscale objects is essential for manufacturing of next-generation electronic systems. Yet, mechanical pick-and-place technologies cannot manipulate smaller objects whose surface forces dominate over gravity, and emerging microtransfer printing methods require multidirectional motion, heating, and/or chemical bonding to switch adhesion. We introduce soft nanocomposite electroadhesives (SNEs), comprising sparse forests of dielectric-coated carbon nanotubes (CNTs), which have electrostatically switchable dry adhesion. SNEs exhibit 40-fold lower nominal dry adhesion than typical solids, yet their adhesion is increased >100-fold by applying 30 V to the CNTs. We characterize the scaling of adhesion with surface morphology, dielectric thickness, and applied voltage and demonstrate digital transfer printing of films of Ag nanowires, polymer and metal microparticles, and unpackaged light-emitting diodes.

INTRODUCTION

Continued advances in electronic devices demand the heterogeneous integration and assembly of components with ever-smaller dimensions. For instance, mechanical sensors including multiaxis inertial sensors and accelerometers are smaller than a millimeter (1), integrated circuit (IC) chips for radio-frequency communication are 0.1 mm or smaller (2), and micro-light-emitting diodes (LEDs) for next-generation displays are 1 to 10 μm in lateral size (3). Furthermore, functional systems for applications in energy conversion, optoelectronics, and biotechnology (4) can be realized by manipulation of complex three-dimensional (3D) micro- and nanomaterials.

At present, automated pick-and-place assembly is the benchmark technique for manufacturing of circuit boards and integration of miniature ICs into systems-on-chip (5–7). However, despite advances in miniature robotic grippers and vacuum nozzles, manipulation is increasingly challenging as object size decreases, because gravity forces diminish more rapidly than van der Waals forces or electrostatic attraction (8). Intricate micromechanical grippers suffer from sticktion and therefore cannot reliably place microscale objects without the aid of secondary adhesive surfaces (9, 10). Therefore, manipulation devices with surfaces that can repeatedly switch between strong adhesion for object pickup and weak adhesion for object placement are necessary for printing at smaller scales. Heterogeneous integration without direct physical manipulation may be possible by self-assembly methods (11, 12), yet precise dimensional control and long-range accuracy of self-assembly are challenging.

Engineering a tunable adhesive surface requires control of both the nanoscale contact mechanics and a reversible mechanism by which attractive forces can be varied between the surface and the target object, ideally without requiring application of shear (13–15), heating (16), or chemical reactions (17, 18). To realize such functionality, we introduce a soft nanocomposite (Fig. 1A) comprising self-assembled nanofibers that, by electrically switchable adhesion, can achieve digitally controllable transfer printing of micro- and nanoscale objects. The soft nanocomposite electroadhesive (SNE) has a very low intrinsic (“off”) dry adhesion through roughness and reduced contact area, while voltage-tunable electrostatic attraction gives strong “on” adhesion force that can overcome the intrinsic adhesion of micro- and nanostructures against donor substrates. The SNE is distinct from well-known electrostatic grippers and electroadhesive films (19–21), as the surface is composed of sparse nanofibers that are nonadhesive yet can be highly compressed. The SNE can therefore conformally contact miniature objects in the absence of an electrostatic force, without adhering. The nanocomposite fibers are conductive but covered with ultrathin dielectric, thus allowing substantial electrostatic pressures to be achieved at low operating voltages (0 to 100 V).

We designed and fabricated SNEs (Fig. 1B) comprising “forests” of vertically aligned multiwalled carbon nanotubes (CNTs; ~10 nm in diameter) (22, 23), grown on a conductive electrode [titanium nitride (TiN), 50 nm in thickness] (24), and then conformally coated with a thin layer of dielectric (Al2O3, ~1 to 10 nm in thickness) by atomic layer deposition (ALD) (25). Further details are given in Materials and Methods and fig. S1. An exemplary pick-and-place sequence using a single SNE pixel is shown in Fig. 1C (movie S1), whereby a 170 μm by 170 μm by 50 μm unpackaged LED chiplet (CREE; detailed geometry in fig. S2) is lifted by the SNE with 30 V applied and placed with 0 V applied.

RESULTS AND DISCUSSION

To study the electrical tunability of adhesion, we measured the adhesive forces of the Al2O3-CNT SNE at different applied voltages using atomic force microscopy (AFM) with a Pt-coated spherical tip (2 μm in radius), which emulates a representative microscale object (Fig. 1D; see fig. S3 for further details). The adhesion, measured by the pull-off force of the cantilever against the SNE, scales quadratically with applied voltage. The off-state adhesion of the SNE is measured to be ~40-fold lower than a reference flat surface (ALD Al2O3 on coated flat TiN on a Si wafer), yet the SNW adhesion increases by >100-fold when 30 V is applied. In micromanipulation, this low off-state adhesion is important to allow objects to be deterministically released from the gripping surface, while the high on-state adhesion enables objects to
be picked from donor substrates that have high intrinsic adhesion to the object (Fig. 1E). Here, we found greater measurement errors using the Pt-coated tips compared to those using uncoated tips. Using a spherical AFM tip comprising a metal sphere directly adhered to a conductive cantilever may improve the measurement accuracy.

The low off-state adhesion of the SNE is due to its low-density fibrous texture and nanoscale roughness, both of which contribute to a low effective contact area against an opposite surface. The self-organization of the CNTs during the CVD process imparts the forest with a top surface consisting of randomly oriented CNTs, supported by well-aligned CNTs underneath (Fig. 2A) (26). As the fibrous SNE makes contact with an object, the CNT forest behaves as a soft foam, where the surface deformation is accompanied by the spring-like behavior of the supporting aligned CNTs (Fig. 2B). The roughness of the Al2O3-CNT SNE surface measured by AFM is approximately 35 nm (Fig. 2C), and the reduced modulus measured by nanoindentation is 86 MPa, which is ~1000-fold less than the flat surface reference (Fig. 2D). Hence, while the surface can deform easily and conformally contact micro- and nanoscale objects with 3D shapes, only a few protruding nanocomposite fibers of the SNE surface will effectively be in

Fig. 1. Picking and placing micro objects using a ceramic-carbon SNE. (A) Schematic of a pick-and-place procedure wherein a micrometer-sized object in contact with an SNE surface can be picked up, while an external voltage is applied and placed back when the voltage is removed. (B) Schematics and scanning electron microscopy (SEM) images of a 200-μm SNE composed of Al2O3-coated carbon nanotubes (CNTs). (C) Snapshots of picking and placing an unpackaged LED chiplet (170 μm by 170 μm by 50 μm) using the Al2O3-CNT nanocomposite electroadhesive (200 μm by 200 μm by 50 μm). (D) Schematic of pull-off force measurement system using a colloidal AFM tip and load-displacement curves for a Al2O3-CNT SNE measured using a ~4-μm-diameter Pt-coated spherical tip with and without a bias voltage of 10 V. The inset SEM image shows the Pt-coated spherical tip. (E) Measured pull-off forces using the Pt-coated spherical tip against a reference surface (flat Al2O3-coated TiN; ~0.8 μN; black dashed line) and against the Al2O3-CNT SNE at different applied voltages (e.g., ~0.02 μN at 0 V and ~2.3 μN at 30 V; blue dashed line), measured by a conductive AFM. Error bars represent the SD in the set of force measurements with each applied voltage.
contact at the interface. Only these contacting fibers contribute to the surface adhesion during retraction, as the surrounding noncontacting fibers do not contribute to the van der Waals adhesion.

The on-state adhesion is determined by electrostatic charge accumulated on the nanofibers both in contact with and in proximity of the object surface, because the electrostatic forces are effective at distances greater than van der Waals forces. As a result, the electrostatic attraction of the SNE to an object can be significantly greater than the van der Waals attraction, especially for rough surfaces. This contrasts the behavior of smooth and flat surfaces, as the van der Waals forces are comparable with the electrostatic forces unless very high voltages (>1 kV) are applied. Accordingly, for the rigid flat surface, the ratio of the on-state adhesive force (with voltage applied) to the off-state (0 V) force is only ~7:1 at 30 V, yet the on/off ratio of the Al₂O₃-CNT SNE is ~100:1 at the same voltage (Fig. 2E and fig. S4).

Surfaces with thinner dielectric coatings exhibit high on/off ratios by reducing the distance between the oppositely charged CNTs and the object surface. A subnanometer Al₂O₃ coating (~0.2 nm) can achieve higher electrostatic attraction forces at a given voltage, although the nonuniformity of such a thin coating (fig. S6), even with ozone-assisted ALD (27), often resulted in breakdown at low voltages. Thus, we found that the minimum thickness conformal dielectric coating was at least ~1 nm. In addition, the density and morphology of the CNT forest surface can be tailored by the CVD growth recipe. For example, a CNT forest with high fiber density exhibits a greater adhesion on/off ratio than a CNT forest with a low fiber density (fig. S7). The electroadhesive properties are independent of forest height beyond ~1 μm (fig. S8), and therefore, the SNE can be designed with fine features by lithographic patterning of the CNTs, or even with spatially varying heights and complex free-form microstructured shapes (28, 29).
Another important factor is the conductivity of the object being manipulated; in the above experiments, both the SNE and the object or probe tip were conductive, and one was grounded. As a charged Al2O3-CNT SNE approaches an object, the local electric field induces charge on the surface of the object. In conductive objects, mobile charges move to the surface to nullify the internal electric field. As a result, the electric field between the CNTs and the object is strong, as is the electrostatic force. In contrast, for nonconductive objects, charge is induced through polarization, i.e., by the electric displacement of bound electron clouds, stretching of polar bonds, and permanent dipole alignment (30). Compared to a conductive object, this results in a lower charge density at a given voltage and a weaker electrostatic force between the CNTs and the nonconductive object (Fig. 3B). However, the stronger electric field created between the SNE and a conductive object will mean that the dielectric coating on the conductive nanofibers will undergo dielectric breakdown at a lower voltage than with a nonconductive object. Thus, while, at a given voltage, the force on a nonconductive object is significantly lower than on a conductive object, significantly higher voltages can be applied to pick up nonconductive objects. This brings the achievable adhesion force for nonconductive objects to within the same range as conductive objects, at the expense of requiring higher voltages.

The scaling of electrostatic attraction compared to van der Waals forces on the SNE can be further understood from a contact mechanics model modified from classical theories (31). We derive the pull-off force \( P_{\text{SNE}} \) against a sphere having radius \( R \) as (details in the Supplementary Materials)

\[
P_{\text{SNE}} = F_{\text{vdW}} + F_{\text{es}} = \frac{3\pi}{2} \eta \varphi \Delta \gamma R + \frac{\pi e^*}{4r^2} \left( \frac{9\eta \varphi \Delta \gamma}{E^*} \right) V^2
\]

where \( \eta \) is the fiber contact ratio; \( \varphi \) is the fiber density; \( \Delta \gamma \) is the work per unit area required to separate two solid smooth surfaces without externally induced electrical charges; \( e^* \) and \( t^* \) are the effective permittivity and effective thickness of the dielectric coating and the air gap, respectively; and \( E^* \) is the reduced modulus of the two contacting surfaces. As given in the first term in Eq. 1, the low fiber contact ratio (\( 0 < \eta < 1 \)) and the low fiber density (\( \sim 0.1 \) or less) are the two major factors that enable the low intrinsic adhesion of the SNE (the first term of the equation). The fiber contact ratio within the nominal contact area depends on the applied load, fiber height variation, and the target object roughness, as owing to height variation, only the most distal nanofiber tips make initial contact (32). Our experiments show that SNE adhesion

![Fig. 3. Switchable adhesion influenced by structural design and object conductivity.](http://advances.sciencemag.org/content/5/8/eaaq4790)

(A) Adhesion on/off ratio (ratio of pull-off force with an external voltage to that without voltage) of Al2O3-CNT SNE with different Al2O3 coating thicknesses (approximately 0.2, 1, and 10 nm) and different fiber densities (sparse and dense). (B) AFM images (top) images of CNTs coated with Al2O3 via ALD with 10 (left) and 50 (right) deposition cycles, respectively, and SEM images (bottom) of SNE top surfaces with different densities. (C) Pull-off forces against a conductive (Pt-coated; \( \sim 4 \mu m \) diameter) and a nonconductive (uncoated SiO2; \( \sim 10 \mu m \) diameter) spherical tip. (D) Schematics of polarization of conductive and dielectric objects when the SNE is in contact with the external voltage on.
increases with preload (fig. S9), whereas the adhesion of solid surfaces is generally independent of preload (fig. S10). The model here only considered normal contact without any shear motion; we experimentally find a decrease in the pull-off force after a shear motion with applied voltages for a spherical object (fig. S11). Therefore, to achieve high maximum SNE adhesion, one should apply normal preload but prevent lateral slip at the interface.

Therefore, proper design of the SNE and understanding of its mechanism of operation enables its use for ambient, electrically switchable manipulation of ultrasmall spherical objects including electronic chiplets, microparticles, and engineered nanostructures. Using Eq. 1, we construct a map (Fig. 4A) that shows the size range of objects that can be manipulated using the demonstrated SNE. For an object of given size, the on-state adhesion of the SNE should exceed the intrinsic adhesion between the object and the donor surface; thus, its size should fall in the “pick” regime. The off-state adhesion should be less than the intrinsic adhesion between the object and the target surface; thus, its size should fall in the “place” regime. For example, we predict that 30 V applied to the Al2O3-CNT SNE can pick an object with a size of ~10 to 100 μm; this is verified by the experiment with a 170 μm by 170 μm by 50 μm micro-LED chiplet (Fig. 1C).

The adhesion measurements and scaling suggest that higher applied voltages (100 V) allow the SNE to pick objects in the far submicrometer size regime. Hence, we demonstrate the use of lithographically patterned SNEs to print complex patterns of micro- and nanomaterials, shown in Fig. 4B, including polymer beads (Fig. 4C) and films of silver nanowires (Fig. 4D). Here, both the donor and target substrates were rigid SiO2/Si wafers, therefore requiring the SNE object adhesion to be switchable to above and below this starting and ending condition. Uniform printing matching the complex shape of the SNE is obtained by simply turning on the DC voltage for pickup and turning off the voltage after the objects are placed in the desired transfer location. However, transfer is not deterministic as some objects transfer without voltage applied to the SNE during pickup (fig. S15). This is attributed to the nonuniformity of the donor layer of objects, which is created by spin coating. Printing is also demonstrated onto flexible polymer films, as in Fig. 4E, which is easier because polymers have stronger surface adhesion than rigid ceramics or metals. The silver nanowires, particularly having a diameter of ~20 nm, could successfully be picked although they are not in the pickable spherical object range. This is due to their wavy structure that induces point contacts (nanometer-scale spherical contacts), while the long-range electrostatic attraction is effective on the whole length of the wire.

In conclusion, the capability of soft, nanostructured surfaces to electrically switch their dry adhesion against both metallic and dielectric objects may provide a new alternative to emerging transfer printing technologies that require thermally or chemically switchable adhesion. Moreover, SNEs may also advance biomimicry of natural surfaces that use polymer-like materials and complex mechanical motions to establish strong adhesion for locomotion. Lithographic patterning of SNE “pixels” in arbitrary patterns, along with their electrical addressability, may enable digital stamps for massive-scale micro- and nanoscale next-generation optical and electronic devices. In addition, SNEs based on 3D CNT microarchitectures (28, 29) could be used to build active legs and pads for miniature climbing robots (33) and microgripper arrays. Applications of SNEs in macroscale manipulation may also be possible, as the measured maximum adhesive strength (force per area) of the SNE (up to ~20 kPa at 225 V) is comparable to planar electroadhesives (at ~5 kV) (20). However, robustness over many cyclic contacts and the influence of charge accumulation and dissipation on the maximum speed of operation require further verification for industrialization.

MATERIALS AND METHODS

Fabrication of SNE

A 50-nm TiN layer was first deposited by sputtering physical vapor deposition (PVD; Endura 5500, Applied Materials) on 4″ (100) silicon wafers with 300 nm of thermally grown SiO2. Then, an Al2O3/Fe catalyst layer was patterned on top of the TiN layer, by lift-off processing using photolithography, followed by ultrasonic agitation in acetone.
The supported catalyst layer, 10 nm of Al₂O₃ and 1 nm of Fe, was sequentially deposited by electron beam PVD (VES-2550, Temescal). The wafer with the deposited catalyst was diced into ~2 cm by 2 cm pieces and placed in a quartz tube furnace for CNT growth. The growth recipe starts with flowing 100/400 SCCM (standard cubic centimeters per minute) of He/H₂ while heating the furnace up to 775°C over 10 min (ramping step) and then holds at 775°C for 10 min with the same gas flow rates (annealing step). Then, the gas flow is changed to 100/400/100 SCCM of C₂H₂/He/H₂ at 775°C for CNT growth for the selected duration. The typical growth rate is ~100 μm/min, and the CNTs are multiwalled with ~10-nm mean diameter with 0.45 coefficient of variation at the top of the forest, as we have previously measured by x-ray scattering (24). After the growth, the furnace is cooled down to <100°C at the same gas flow and lastly purged with 1000 SCCM of He for 5 min. Last, Al₂O₃ was deposited onto CNTs by ALD (GEMStar, Arradiance Corporation). Trimethylaluminum (TMA) and ozone (O₃) were used as the metalorganic and oxidizing precursors, respectively. Using nitrogen as the carrier gas at a flow rate of 40 SCCM, TMA and O₃ were sequentially pulsed into the deposition chamber (2 to 3 torr at 175°C) for 22 and 100 ms, respectively. Following each precursor pulse, the chamber was purged with 90 SCCM of nitrogen for 38 s. Polymide tapes were used to mask the electrical contact points.

Adhesion tests (AFM)

The adhesion tests for a conductive material were done using a standard AFM (Bruker ICON AFM) in conductive mode. The samples were mounted by a vacuum holder and electrically connected to the steel stage. A platinum-coated AFM probe was mounted to the tip holder and connected to an electric control module. The module controlled the voltage bias between the steel stage and the AFM probe. A schematic of the setup is shown in fig. S3. The AFM probes are specially developed products (SD-Sphere-CONT from Nanosensors) with a sphere (~4.5 μm in diameter) at the tip and coated with platinum (~140 nm) by PVD. Before the experiments, the stiffness of the AFM cantilever was calibrated using a thermal tuning method. During each experiment, the AFM probe scanned the sample to map the surface topography. The AFM probe was scanned with a source meter (Keithley 2400). The SNE was mounted on a force sensor with a capacity of 100 mN. Voltages up to 30 V were applied with a source meter (Keithley 2400). The experiment was viewed from the side with a long working distance microscope (Navitar) and a complementary metal-oxide semiconductor camera (D722MU, Pixelink). For LED pick and place, a 200 μm by 200 μm SNE was contacted to the LED chiplet that was resting on a Si substrate; the load cell was used to control the preload. After contact, voltage was applied to the bottom TiN layer of the SNE relative to a ground connected to the stage supporting the Si substrate. The SNE was retracted upward, retrieving the LED. The SNE with the LED was then brought back into contact with the wafer surface, and the voltage was turned off. Upon retraction of the SNE, the SNE separated from the LED with no measurable force. The scale bar is 200 μm. For multiple microparticles and nanowires, a custom-made manual stage was used. First, the microparticle or nanowire dispersion was spin-coated on an Al₂O₃-coated Si wafer at >1500 rpm for >5 min. The particle/wire layer was further dried in an atmospheric environment for >1 day. A patterned Al₂O₃-CNT SNE was brought into contact against the particle/wire layer. A customized manual stage was used to gently bring the nanocomposite onto the layer and be pressed by its own weight. As the spin-coated particle layer was not uniform, a slight lateral motion (few micrometers) was induced to allow most of the SNE surface to make intimate contact with the particles. Then, an external voltage of >40 V was applied by a power supply into the bottom TiN layer attached to the SNE against a

Millimeter-scale force measurements

Millimeter-scale force measurements on the nanocomposite electro-adhesives were performed using custom-built equipment. This equipment used two single-component load cells (LRM200-FSH02899, Futek; ±100 g capacity) attached to a flexure mechanism that decoupled the horizontal and vertical components of force. Attached to the flexure was a flat-tipped probe consisting of a 1.2-mm square silicon wafer die (X040093000W, WaferPro; 300-nm thermal oxide layer, diced on DISCO DAD-3240 diesaw) mounted on a slender aluminum column. The entire probe was sputter-coated with gold (100 nm; MS Q150T ES coater, Quorum) for electrical continuity. Voltage was applied between the tip and CNT forest through copper tape connected to the base of the probe and the forest away from the area of force measurements. The CNT forest sample was leveled relative to the tip on a tip-tilt-rotation stage (TTR001, Thorlabs). The load cell, flexure, and probe assembly was mounted on a two-axis stepper motor–controlled traversing mechanism capable of 100-nm vertical resolution and 1-μm lateral resolution.

Attractive force measurements were performed by positioning the tip at least 30 μm above the CNT forest, applying 100 V (PS375/+20 kV, Stanford Research Systems) between them, and then approaching the surface at a speed of 0.5 μm/s. Force measurements were collected every 100 nm until contact was made with the CNTs. Adhesion measurements were performed by positioning the tip above the CNT forest and approaching the surface at 0.2 μm/s without voltage applied. Tip motion was stopped once the desired preload force was reached. Voltage was then applied, and the tip was retracted at a speed of 0.2 μm/s. The maximum adhesive force recorded during retraction was reported as the adhesion force.

Pick-and-place demonstration

Pick-and-place demonstration of micro-LEDs and stainless steel microparticles (fig. S16) was done on a custom micromechanical test system consisting of a stepper motor–driven stage for controlling motion in the z direction and a two-axis manual translation stage for controlling motion in the x and y directions. The SNE was mounted on a force sensor with a capacity of 100 mN. Voltages up to 30 V were applied to the bottom TiN layer of the SNE relative to a ground connected to the stage supporting the Si substrate. The SNE was retracted upward, retrieving the LED. The SNE with the LED was then brought back into contact with the wafer surface, and the voltage was turned off. Upon retraction of the SNE, the SNE separated from the LED with no measurable force. The scale bar is 200 μm. For multiple microparticles and nanowires, a custom-made manual stage was used. First, the microparticle or nanowire dispersion was spin-coated on an Al₂O₃-coated Si wafer at >1500 rpm for >5 min. The particle/wire layer was further dried in an atmospheric environment for >1 day. A patterned Al₂O₃-CNT SNE was brought into contact against the particle/wire layer. A customized manual stage was used to gently bring the nanocomposite onto the layer and be pressed by its own weight. As the spin-coated particle layer was not uniform, a slight lateral motion (few micrometers) was induced to allow most of the SNE surface to make intimate contact with the particles. Then, an external voltage of >40 V was applied by a power supply into the bottom TiN layer attached to the SNE against a
Supplementary Materials

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/suppl/5/10/eaax4790/DC1

Supplementary Text

Fig. S1. Schematics of the Al2O3-CNT SNE fabrication procedure.

Fig. S8. Pull-off forces of Al2O3-CNT SNEs with different heights according to applied voltages.

Fig. S4. Adhesion on/off ratio (ratio of pull-off force with an external voltage to that without voltages.

Fig. S6. Transmission electron microscopy images of Al2O3 coating on individual CNT fibers via ALD with different numbers of cycles.

Fig. S11. Pull-off forces of SNE measured by AFM indentation using a 10-m by 200-m SiO2 spherical tip.

Fig. S13. Contact mechanics modeling of adhesion between an Al2O3-CNT nanocomposite and a microsphere.

Fig. S16. Picking-and-placing of stainless steel (SS) microparticles, sized 15-45 μm in diameter.

Fig. S14. Adhesive and gravitational forces along the object size.

REFERENCES AND NOTES


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