Reversible temperature regulation of electrical and thermal conductivity using liquid–solid phase transitions

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Reversible temperature tuning of electrical and thermal conductivities of materials is of interest for many applications, including seasonal regulation of building temperature, thermal storage and sensors. Here we introduce a general strategy to achieve large contrasts in electrical and thermal conductivities using first-order phase transitions in percolated composite materials. Internal stress generated during a phase transition modulates the electrical and thermal contact resistances, leading to large contrasts in the electrical and thermal conductivities at the phase transition temperature. With graphite/hexadecane suspensions, the electrical conductivity changes 2 orders of magnitude and the thermal conductivity varies up to 3.2 times near 18 °C. The generality of the approach is also demonstrated in other materials such as graphite/water and carbon nanotube/hexadecane suspensions.
Reversible temperature tuning of electrical and thermal conductivities of materials is of interest for many applications, such as seasonal regulation of building temperatures, thermal storage, sensing and so on. Although the electrical conductivity can be tuned by temperature using metal-insulator transitions, temperature regulation of thermal conductivity is much more difficult as thermal properties vary little through solid-state phase transitions. In particular, materials capable of near room temperature regulation of both electrical and thermal conductivity could prove useful for new applications.

Liquid–solid phase transitions can be an attractive route for the temperature regulation of electrical and/or thermal properties because of the availability of materials with a wide range of phase transition temperatures. Because a liquid–solid phase transition usually does not cause a metal-insulator transition, nanoparticles can be added to liquids to control the property contrast between the liquid and solid states. In particular, in a crystal-forming liquid loaded with nanoparticles, particles are squeezed towards grain boundaries on freezing\(^7\). The internal stress generated during freezing regulates the contacts among particulates, increasing the electrical and/or thermal conductivities of the composites. The changes of the electrical and/or thermal conductivities at the phase transition temperature can be further optimized by adjusting the concentration of the particulates. By combining these effects, novel composites exhibiting large variations in electrical and thermal conductivities in a narrow room temperature range can be achieved.

In this paper, we demonstrate this first-order phase-transition strategy to regulate electrical and thermal properties. Using a graphite/hexadecane suspension as an example, we achieve a 2-orders-of-magnitude change in the electrical conductivity and a three-fold change in the thermal conductivity around 18°C. The generality of the approach is also demonstrated in other materials such as graphite/water and carbon nanotube/hexadecane suspensions. The critical temperature of such temperature-responsive composites can be readily adjusted for specific applications by selecting host fluids with the right phase transition temperature.

**Results**

**Sample preparation.** We use a hexadecane/graphite suspension as an example. The phase transition temperature of hexadecane is around 18°C, which is convenient for experimentation. Exfoliated graphite flakes have been widely utilized in composite materials because of their high in-plane strength and high electrical and thermal conductivities\(^5\). We followed a process of sulphuric acid intercalation, microwave expansion and ultrasonic dispersion to prepare graphite suspensions\(^6\)–\(^11\). After these processes, natural graphite is exfoliated into graphite flakes, which are then mixed with hexadecane to form a stable suspension.

**Suspension characteristics.** An optical microscope image of 0.2% (volume fraction) graphite/hexadecane suspension is shown in Figure 1a, together with an inset showing a photo of 50 ml of such a suspension after 3 months on the shelf. The graphite suspension is stable and no sediment is found. The graphite flakes have an average diameter of several microns and a thickness from several nanometres to several tens of nanometres (Fig. 1b). Because of the intrinsic stress induced during preparation, most of the graphite flakes are bent, with some of them even rolled up (Fig. 1c). A high-resolution transmission electron microscopy image (inset in Fig. 1c) of a selected flake shows that the flake has \(~30\) atomic layers with an interplanar distance of 0.335 nm, consistent with graphite\(^7\). We measured the mobility of graphite flakes in hexadecane to be \(~0.03\) cm\(^2\) V\(^-1\) s\(^-1\), which means that the graphite flakes are charged in hexadecane.

**Electrical conductivity.** The electrical conductivity of the suspension was measured using a four-point configuration. The electrical conductivities of the graphite suspensions in the liquid state prepared using different volume fractions of the graphite flakes are shown in Figure 2a. The inset of Figure 2a shows a logarithmic plot of electrical conductivity versus \(\phi\), demonstrating a percolation threshold of \(\phi_c = 0.05\%\) (ref. 14). The electrical conductivity of the graphite suspensions varies significantly around 18°C when hexadecane starts to freeze, as shown in Figure 2b for a graphite volume fraction loading between 0.2–1%, above the percolation threshold. In the liquid state, the electrical conductivity of the graphite suspensions varies little with temperature; from 18.5 to 17.5°C, the electrical conductivity increases by 2 orders of magnitude. After the hexadecane is completely frozen, the electrical conductivity stabilizes. In both the solid and the liquid states, the electrical conductivity increases with an increase of the graphite volume fraction. However, the contrast ratio of electrical conductivity, defined as the ratio of electrical conductivity between the solid and the liquid states near the phase transition, peaks at 250 around a volume fraction of 0.8% (Fig. 2b, inset). The peak exists because at a lower graphite volume fraction, the electrical conductivity in the solid state is not sufficiently high. However, at a higher graphite volume fraction, the electrical conductivity in the liquid state is large, also reducing the contrast ratio. We anticipate that the peak value depends on the geometry of the graphite flakes, surface states and liquid properties, and believe that further improvements in the contrast ratio are possible through optimization.

**Thermal conductivity.** The thermal conductivity is measured using a hot-wire method\(^15\). The thermal conductivity of the graphite suspensions has the same trend as the electrical conductivity (Fig. 2c). In the liquid state, thermal conductivity changes little with temperature. As the host solidifies, the thermal conductivity rises sharply. At 0.8% volume fraction, it was observed that the thermal conductivity changes by a factor of 3.2 near the phase transition temperature (Fig. 2c, inset). The smaller contrast in the thermal conductivity change compared with that of electrical conductivity is because of the fact that the thermal and electrical transport mechanisms are different. Electrons are forced to pass only through the graphite flakes, whereas heat can conduct through both the liquid and the solid host phases. Furthermore, the thermal conductivity ratio between graphite and hexadecane is \(~10^4\), whereas for the electrical transport the ratio of conductivities can be on the order of \(10^{12}–10^{16}\).

**Reversibility.** The cycling behaviour of the electrical conductivities as the materials go through repeated melting and solidification processes is shown in Figure 2d. The first freezing cycle shows the largest contrast in electrical conductivities because there are more loose clusters in the liquid states. When the frozen hexadecane remelts, the contact between graphite flakes degrade quickly, leading to sharp reductions in the electrical conductivity. However,
many graphite flakes will be trapped in the percolation network and the electrical conductivity in the liquid state increases from its original as-synthesized state. After the first temperature cycle, the percolation structure becomes stable, and thus the contrast ratio of the electrical conductivity of the suspension approaches a constant. All the experimental data shown in Figure 2b, c are the result of the second freezing cycle of different samples. The cycling behaviour of the thermal conductivity was shown in Figure 2e. The variation of thermal conductivity during the first and subsequent cycles is smaller because unlike electrical conductivity, heat conducts through both the graphite percolation network and the host.

**Discussions**

The experimental results presented in Figure 2b–e were obtained for suspensions that already reached percolation in the liquid state. Thus, the large contrasts observed in the electrical and thermal conductivity changes at the phase transition are not simply due to the percolation structure. The internal stress generated during freezing may be responsible for the large contrasts through regulation of the contact resistances of the percolated nanoparticle network.

We set up an experiment (Fig. 3a) to measure the electrical contact resistance between two highly ordered pyrolytic graphite (SPI-1) flakes in a hexadecane environment. The resistance of the circuit decreases 400 times as the temperature decreases from 18.5 to 17.5 °C (Fig. 3b). The stress distribution (Fig. 3c) measured in the frozen hexadecane (see Method) shows that the stress is unevenly distributed in regions where the average pressure is ~160 p.s.i. We believe the non-uniform pressure distribution is due to the anisotropic growth of hexadecane crystals. The conceptual process is shown in Figure 3d. Similar to exfoliated graphite flakes, naturally peeled highly ordered pyrolytic graphite flakes are uneven and curved. In the liquid state, the contact area is small and hence the electrical resistance between two flakes is high. Hexadecane crystals exhibit strong anisotropic growth kinetics. During the freezing of...
hexadecane needle-like structures are formed, with the aspect ratio depending mainly on the freezing speed. The anisotropic growth of the hexadecane crystals generates stress and increases the contact area of the graphite flakes. Stress can also improve the electrical contacts by reducing the thickness of the insulating liquids in between graphite flakes. After freezing, the contact area and electrical resistance stabilize. When the hexadecane remelts, the pressure on the graphite flakes is released and the contact area and interparticle repulsion. A similar trend exists for the thermal conductivity of materials, using first-order phase transitions between liquid and crystalline solid states through seeding the liquid with particulates to form stable suspensions. During the phase transition, internal stress reduces the electrical and thermal contact resistances in the percolated network formed by the particulates, leading to large changes in the electrical and/or thermal conductivities. The strategy is general and can be tailored for either electrical and/or thermal conductivity or both, and for operation in a desired temperature range. Reversible tuning of electrical and thermal properties near room temperature as demonstrated in this paper has potential applications in temperature regulation and sensing. Extension of the method, such as using high melting point materials, for example, molten salts and ionic liquids, can be applied to solar thermal energy storage and power generation. Temperature regulation of the properties of the materials through percolation networks and stress generation during crystallization as demonstrated in this paper complements other actively controlled suspensions such as magnetorheological fluids and ferrofluids.

Methods

Sample preparation. Exfoliated graphite was prepared from natural graphite (Asbury Carbons) by a chemical intercalation and thermal expansion method. A volume of 85 ml of H₂SO₄ (96%, Alfa Aesar) and 15 ml of hydrogen peroxide (30%, Alfa Aesar) were mixed to form an oxidative agent that was used for the intercalation. For the intercalation process, 2.2 g of natural graphite -1

Figure 2 | Variation of electrical and thermal conductivities around the phase transition point. (a) Electrical conductivity of graphite suspensions as a function of volume fraction in the liquid state. The inset shows a logarithmic plot of σ versus (ϕ−ϕc), which indicates a percolation threshold of ϕc = 0.05%. (b) Electrical conductivity of graphite suspensions as a function of temperature. The inset shows the relationship between the contrast ratio of electrical conductivity and graphite volume fraction. (c) Thermal conductivity at different graphite volume fractions as a function of temperature. The inset plot indicates the relationship between the contrast ratio of thermal conductivity and graphite volume fraction. (d) Cyclic EC (electrical conductivity) versus temperature measurement results of a 0.8% graphite/hexadecane suspension. Blue and red squares indicate the conductivity variance through the processes of freezing and melting, respectively. (e) TC (thermal conductivity) contrast after different cycles. Blue lines indicate the thermal conductivity of 0.8% graphite/hexadecane suspensions at 3 °C during different thermal cycles, whereas red lines show the thermal conductivity at 25 °C during different thermal cycles.
Graphite flakes were reacted with 100 ml of the oxidative agent at room temperature for 60 min. The slurries were rinsed with distilled water (Mallinckrodt Baker) to remove residual salts and acids, followed by filtering and baking on a hotplate at 110 °C for 24 h. The expanded graphite was obtained by thermal expansion of as-prepared graphite intercalation compounds in a 1,100-W commercial microwave oven (SANYO) for 30 s. The expanded graphite was then dispersed in hexadecane and other liquids to create a 1% volume fraction of graphite suspension. Graphite flakes were directly exfoliated from expanded graphite by ultrasonication in hexadecane (Sigma-Aldrich) or distilled water, using a high-intensity ultrasonic probe (Sonics VC-750 (Sonics), 750 W, 80% amplitude) for 1.5 min. The obtained uniform dispersion was then dispersed in hexadecane and other liquids and further ultrasonication was performed for 3 min to obtain 0.2, 0.4, 0.6 and 0.8% suspensions. The dispersions become stable suspensions after they cool to room temperature.

Microstructure characterization. The morphologies of expanded graphite and exfoliated graphite flakes were observed by s.c.m. (JEOL ISM-6320, JEOL). The microstructures of exfoliated graphite were examined using a transmission electron microscopy (JEOL 200cx, JEOL 2011, JEOL) at room temperature. The evolution of the clusters and percolating structures of the graphite flakes in hexadecane was observed by a Nikon multipurpose zoom microscope (AZ100). XPS samples were prepared by pressing graphite flakes into a 5-mm-diameter disc. The sample was analysed by using a Kratos AXIS Ultra Imaging X-ray Photoelectron Spectrometer. The data were processed by using the CasaXPS software.

Electrical conductivity measurement. The electrical resistance of graphite suspensions was measured by pouring graphite gels into a self-made conductivity cell that has two dipping parallel vertical copper electrodes ~4.85 cm apart and with an area of ~1.17 cm². The resistances of the graphite suspensions varied from 1.2–10 MΩ, which is far below the sample resistance. The electrical conductivity of graphite gels is given by the equation \( \sigma = k/R \), where \( R \) is the measured resistance and \( k = 3.83 \text{ cm}^{-1} \) is the cell constant. We estimate that the uncertainty in the electrical conductivity measurement is ~1.6%.

Thermal conductivity measurement. A transient hot-wire method was used to measure the thermal conductivity of the graphite suspensions. A 50-μm diameter Pt wire with a 1.5-μm thick insulating adhesion layer was fully immersed in the samples. The wire was subjected to a current pulse of precisely

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Figure 3 | Mechanisms in reversible temperature control of both electrical and thermal conductivities. (a) Schematic diagram of measurement of electrical resistance between two graphite flakes peeled from highly ordered pyrolytic graphite. The dimensions of the graphite flakes are ~1μm×3 mm×3 mm. Graphite flakes are contacted by 30-μm diameter gold wires. (b) Measured variation of electrical resistance as a function of temperature, from 18.5 to 17.5 °C. The resistance of the circuit decreased about 460 times. Inset 1 is an optical image of graphite flakes submerged in solid hexadecane, and inset 2 shows the graphite flakes submerged in liquid hexadecane. (c) Stress distribution map in frozen hexadecane, showing that the stress is unevenly distributed from 74–400 p.s.i., with an average of ~160 p.s.i. (d) Conceptual illustration of the contact area variation between graphite flakes submerged in hexadecane through the process of hexadecane freezing and remelting. The anisotropic growth of hexadecane crystals generates pressure on the surface of the graphite flakes, which increases the contact area and reduces the flake separation. When the frozen hexadecane remelts, the pressure is released and the contact area decreases.

Figure 4 | The electrical conductivity of three other suspensions. (a) 1% graphite/water suspension. The electrical conductivity decreases about 24 times from ~0.5 to 0.5 °C. (b) 1% carbon nanotube/hexadecane suspension. The electrical conductivity decreases about 40 times from 17.5 to 18.5 °C. (c) 0.4% graphite/polyethylene glycol 400 suspension. Polyethylene glycol 400 (Alfa Aesar) has a melting point between 4 and 8 °C and does not form a crystalline solid. Consequently, no large variation in the electrical resistivity was observed.
known amplitude and the resulting temperature rise was determined as a function of time by monitoring changes in the electrical resistance of the wire. By analysing the temporal temperature profile using solutions to the heat conduction equation, we determined the thermal conductivity of each sample. Our analyses show that the uncertainty of our hot wire thermal conductivity setup is ~1.1%, and can be mostly attributed to systematic uncertainty due to uncertainty in the hot wire length. In addition, we participated in a round-robin to measure the thermal conductivity of nanofluids14, and our experimental results were within 2% of mean values.

Internal stress measurement. The internal stress distribution in hexadecane was measured by a pressure-indicating film (Pressurex from Sensor Products). A scaled 15×10 mm² Pressurex film was submerged in a hexadecane-filled cell. The cell was placed in an ice-water bath for an hour so that hexadecane solidifies. Further, the cell was exposed to the ambient temperature (22 °C) and the Pressurex film was removed after the hexadecane remelted. A greyscale stress distribution profile appeared on the Pressurex film, revealing the relative amount of pressure applied to the film due to solidification. Using the Topaq software (Sensor Products), the greyscale pressure distribution profile is rendered into a full-colour representation of the stress distribution.

References

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Author contributions
R.Z., G.C., and J.G. developed the ideas. R.Z., J.G., and J.W. conducted the experiment. R.Z. and G.C. wrote the manuscript. G.C. supervised the research.

Additional information
Competing financial interests: A U.S. patent application has been filed based on the work ‘Thermal and/or electrical conductivity control in suspensions’ (Application No. 12/720,382).

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