Flexible Dielectric Nanocomposites with Ultra-Wide Zero Temperature Coefficient Window for Electrical Energy Storage and Conversion under Extreme Conditions
Khurram Shehzad†, Yang Xu†‡*, Chao Gao§, Hanying Li§, Zhi-Min Dang∥, Tawfique Hasan†, Jack Luo†∇, and Xiangfeng Duan‡*

†College of Information Science and Electronic Engineering, Zhejiang University, Hangzhou 310027, China
‡Department of Chemistry and Biochemistry and California Nanosystems Institute, University of California, Los Angeles, California 90095, United States
§MOE Key Laboratory of Macromolecular Synthesis and Functionalization, Zhejiang University, Hangzhou 310027, China
∥State Key Laboratory of Power System, Department of Electrical Engineering, Tsinghua University, Beijing 100084, China
† Cambridge Graphene Centre, University of Cambridge, Cambridge, CB3 0FA, UK
∇Institute for Materials Research and Innovation, The University of Bolton Deane Road, Bolton, BL3 5AB, UK
*E.mail: yangxu-isee@zju.edu.cn, xduan@chem.ucla.edu

ABSTRACT
Polymer dielectrics offer key advantages over their ceramic counterparts such as flexibility, scalability, low-cost, and high-breakdown voltages. However, a major drawback that limits more widespread application of polymer dielectrics is their temperature-dependent dielectric properties. To achieve dielectric constants with low/zero temperature coefficient (L/0TC) over a broad temperature range is essential for applications in diverse technologies. Here, we report a hybrid filler strategy to produce polymer composites with an ultra-wide L/0TC window of dielectric constant, as well as significantly enhanced dielectric value, maximum energy storage density, thermal conductivity and stability. By creating a series of percolative polymer composites, we demonstrated hybrid carbon filler based composites can exhibit zero-temperature coefficient window of 200 °C (from −50°C to 150°C), a widest 0TC
window for all polymer composite dielectrics reported to date. We further show the
temperature coefficient of the composites is highly stable against stretching and bending, even
under AC electric field with frequency up to 1 MHz. We envision that our method will push
the functional limits of polymer dielectrics for flexible electronics in extreme conditions such
as in hybrid vehicles, aerospace, power electronics, and underground oil/gas exploration.

Keywords: Dielectric constant; Polymer nanocomposites; Zero temperature coefficient;
Carbon nanotubes; Energy storage and conversion

INTRODUCTION

Dielectrics can deliver a very high power density and represent an attractive rapid
energy storage technology for applications in hybrid vehicles, power electronics, bio-medical
field, and electrical weapon systems.\textsuperscript{1-3} Compared to the ceramic dielectrics, polymer
dielectrics offer a high breakdown voltage, lower processing temperature, easy processibility,
flexibility, and are of increasing interest for diverse applications.\textsuperscript{4} However, polymers
typically have intrinsically low dielectric constants (e.g., $< 10$). It is thus often necessary to
form composites of polymers with dielectric or conductive fillers to raise their dielectric
constant for variety of high energy density applications. Percolative polymer composites
(PPCs), containing a conductive filler, achieve high dielectric constant at relatively low filler
concentrations while retaining the easy processibility and mechanical robustness of polymer
composites, making them a key multi-functional material for applications in energy
conversion and storage,\textsuperscript{5-6} electronics,\textsuperscript{7} and bio-medical fields.\textsuperscript{8-9} Despite their widespread
applications and advantageous properties, PPCs are typically limited by intrinsic drawbacks
of sharp insulator-conductor transition near the percolation threshold, and highly temperature
dependent electric and dielectric properties.\textsuperscript{10-11} Compared to the electrical resistivity ($\rho$), the
dielectric constant ($\kappa$) of PPCs is much more sensitive to the temperature since the filler
concentration of composites with high dielectric constant and low dielectric loss always falls
in the most thermal-sensitive region of percolation curve (near the critical filler concentration).\textsuperscript{10, 12} This intrinsic drawback limits the working temperature of PPCs, and makes them unsuitable for the applications with large temperature swings such as in hybrid vehicles, aerospace, power electronics, and underground oil/gas exploration.\textsuperscript{13-15} In fact, temperature sensitivity of $\kappa$ is a challenge not only for PPCs, but also for traditional ceramic capacitors used in electronic circuits. To achieve the stable function of ceramic capacitors over a wide temperature window, complex temperature compensating electronic circuits are required.

To enhance the applicability of PPCs, low/zero temperature coefficients of dielectric constant (L/0TC$_\kappa$), as well as electrical resistivity (L/0TC$_\rho$), over a broad temperature range is much needed. So far there is no comprehensive strategy available to simultaneously obtain both the wide temperature range L/0TC$_\kappa$ and L/0TC$_\rho$ in PPCs. Strategies such as polymer-filler interface engineering\textsuperscript{16-18} or filler alignment\textsuperscript{19,20} have been employed for this purpose. However, despite considerable efforts with different strategies, L/0TC was achieved only over a relatively narrow temperature window (e.g., 100 °C). A most recent breakthrough reports nanocomposites with relatively high working temperature\textsuperscript{21}, yet with relatively low dielectric constant (~3.5 at maximum), and thus unsuitable for high energy density storage materials.

Herein, we report a novel hybrid filler strategy for tailoring the temperature dependence of dielectric as well as the electrical resistivity of the PPCs. By using hybrid of one-dimensional (1D) carbon nanotubes (CNTs) and common zero-dimensional (0D) carbon black (CB)fillers, we simultaneously achieved L/0TC$_\kappa$ and L/0TC$_\rho$ in the hybrid filler composites over a record ultra-wide temperature window of 200°C (from −50 °C to 150 °C), which is at least 2 times wider than those of single filler based polymer composites reported previously. Moreover, L/0TC properties are highly stable against stretching and bending, even under AC electric field with frequency up to 1 MHz. By controlling the concentration and
relative ratio of hybrid constituent fillers, we demonstrate, for the first time, that PPCs can be engineered to achieve positive, negative and zero temperature coefficients of resistance (PTC, NTC, L/0TC) by design, which can open up many exciting opportunities for practical applications such as temperature sensing and self-regulating heating.

RESULTS AND DISCUSSION

Polymer composites were fabricated by melt mixing the polymers, and the individual (CNT, and CB) as well as their hybrid fillers (CNT-CB). Initially two types of CNTs were studied, but finally one type was chosen due to its better suitability for our studies (details of the process and choice of materials for fabrication of polymer composites are provided in the supporting information). By controlling the relative ratio of hybrid fillers in the composites, it is possible to achieve L/0TC PPC over a wide temperature window. Figure 1a shows the TEM images of CNT, CB, and (CNT+CB) hybrid fillers, while the Fig. 1c describes the schematics of filler networks in the thermoplastic elastomer (TPE) polymer matrix. For CNT+CB hybrid filler, CB particles or clusters are bridged by CNTs as shown by TEM image. Cross sectional SEM images (Fig. 1b) confirm the presence of CNT, CB, and CNT+CB fillers in their respective composites. Figure 2 shows the dependences of of κ and ρ on the volume fraction of different carbon fillers at room temperature. Figure 2 clearly demonstrated a conductor-insulator translation in the volume fraction range of 0.06-0.07, for all carbon fillers. A large enhancement in the electrical conductivity (σ) of several orders of magnitude was observed at critical concentration (percolation threshold), indicating the formation of the continuous conductive network in the nanocomposites. A large enhancement in dielectric permittivity (ε) of nanocomposites was also observed near (below) this critical concentration, which was the result of Maxwell-Wagner-Sillar polarization. Electrical resistivity and dielectric constant of the typical PPCs are generally governed by the following formula: 22-24

\[ \rho \propto (f_{\text{filler}} - f_c)^\gamma, \quad \text{for} \quad f_c \leq f_{\text{filler}} \]  

(1)
\( \kappa \propto (f_c - f_{\text{filler}})^q \), for \( f_c \geq f_{\text{filler}} \) \tag{2} 

where \( f_{\text{filler}} \) is the filler concentration, \( f_c \) is the percolation threshold, and \( q \) and \( t \) are the critical constants. Linear fitting of Eq. 1 and 2 produced the percolation threshold as 0.06, 0.07, and 0.07, values of \( q \) as 0.87, 1.44, and 0.57, and values of \( t \) as 1.06, 4.50, and 4.63, for CNT, CB, and (CNT+CB) based composites, respectively (Fig. S1). For 3D fillers, universal value of \( t \) is in the range of 1.6–2.0 while of \( q \) is 1.0. Non-universal and high value of \( t \) can be attributed to diverging inter-particle distances (caused by variations in aspect ratios, dispersion and the networks of filler in the polymer matrix), yielding a diverging distribution of resistance elements as inter-particle resistance bare the feature of exponential decay of tunneling conductivity with the inter-particle distance. \(^{23,25}\) A lower \( t \) value also means a more sharper percolation curve, and higher \( t \) value corresponds to a more gradual percolation curve. Dielectric loss tangent of the all the polymer composites is provided in Fig. S2a. Dependence of dielectric constant and dielectric loss tangent against the AC frequencies at selected concentration (0.06) of hybrid filler is also provided in Fig. S2b. Percolation threshold of our CNT/polymer composites is relatively high but still consistent with previous studies.\(^{26-27}\) A higher than expected percolation threshold of CNTs is probably result of waviness/flexibility of CNTs which make them bend and consequently lower their effective length and/or aspect ratio. Percolation threshold of hybrid fillers was in close approximation of previously proposed theoretical models which propose the equation \( V_{\text{CNT}}/f_{\text{CNT}} + V_{\text{CB}}/f_{\text{CB}}= 1 \) to calculate the percolation threshold for mixed fillers.\(^{23}\)

Figure 3a and 3b present the effect of temperature on \( \kappa \) and \( \rho \) of all the composites, respectively. For comparison, temperature dependence of pure polymer matrices (TPE and PVDF) is also presented in Supplementary Fig. S3. There is no obvious correlation temperature dependence of dielectric properties between of pure polymers and their composites, as properties of pure polymer at various temperatures are determined by chain
dynamics, while of composites are determined by filler networks. TPE composites with 0D filler (CB) exhibit obvious temperature dependent electrical and dielectric properties, especially near melting temperature of TPE (150 °C) (Fig. S3), where \( \kappa \) decreases and \( \rho \) increases (PTC effect) with increasing temperature (To be consistent with literature and to avoid confusion, the term L/0TC will be used for both \( \rho \) and \( \kappa \), and the terms PTC and NTC in following discussion will be only used with reference to electrical resistivity but not to dielectric constant). Variety of mechanisms (such as volume expansion, tunneling current migration of filler particles, electric field emission, and internal stress) have been reported in the literature to explain the PTC effect. Most probable mechanism in this case may be the thermal expansion of polymer, which took place at the melting point of the polymer matrix, resulted in the break-up of the conducting chains (see schematics in Fig. S4), with a consequent increase in \( \rho \) and decrease in \( \kappa \) value of the composites.\(^{28-30}\) It is important to note that due to the 0D nature of CB particles, contact between CB particles is point-type, which is easily breakable.

In the case of 1D (CNT) filler based polymer composites, compared to CB where inter-filler point-type contact has low contact area, inter-CNT line-type contact, due to the large aspect ratio (AR) of CNTs because of their 1D nature, has larger contact area (the schematics of NTC mechanism for CNT based composites is presented in Fig. S4). It is important to mention that some of the CNTs, when not oriented parallel to each other, may also have point-type contact. However, point-type contact of CNTs is not easy to break because of their higher length. These factors make CNT/polymer composites immune to polymer chain movement induced increase in \( \rho \). Instead, just below the percolation threshold, CNT/polymer composites show a decrease in \( \rho \) and an increase in \( \kappa \) values at higher temperatures (Fig. 3a, 3b). For example, near percolation threshold, the \( \kappa \) value of the composites experiences an increase of at least 75% in the temperature range of −50 °C to 150 °C. The decrease in
resistivity and increase in $\kappa$ with temperature is often ascribed to temperature dependent increase in the conductivity of polymer matrix, thermal fluctuations induced tunneling (FIT),\textsuperscript{31-32} variable range hopping (VRH),\textsuperscript{33} or reparation of the disconnected conductive chains due to the dynamic factors (van der Waals interaction between fillers).\textsuperscript{34-35}

Since CNT and CB based composites display opposite temperature dependence of their electrical and dielectric properties, we hypothesize that a combination of both fillers in appropriate ratio could result in a balance to create a composite with temperature independent electrical/dielectric properties. Indeed, by adding an equal concentration of CB and CNT \([(\text{CNT}_{0.03} + \text{CB}_{0.03})\), where 0.03 in subscript means the volume fraction of each filler], we obtain a balanced composite where CB induced fall in $\kappa$ is compensated by a CNT induced increase in $\kappa$ (Fig. 4a). The figure presents the normalized electrical resistivity $[\rho(T)/\rho_{-50^\circ\text{C}}]$, and the percentage change in $\kappa$ against its value at $-50^\circ\text{C}$. $\rho(T)$ is resistivity at a given temperature and $\rho_{-50^\circ\text{C}}$ is resistivity at $-50^\circ\text{C}$. Normalized resistivity values and change in $\kappa$, with reference to their room temperature value are also presented in Figs. S6-S8.

Figure 4c shows that $\rho$ and $\kappa$ values display no significant change over the entire investigated temperature range of $-50$ to $150^\circ\text{C}$, clearly demonstrating that an unprecedented wide L/0TC$_\rho$ and L/0TC$_\kappa$ window has been achieved in (CNT+CB)/polymer composites with 0.06 filler volume fraction. In contrast to a maximum change of 125% in $\kappa$ for CNT composite and $-42\%$ for CB composite over the temperature range of $-50$ to $150^\circ\text{C}$, the maximum change in $\kappa$ of CNT+CB hybrid composite is only 9.7% (Fig. S5b). By using the room temperature ($25^\circ\text{C}$) as reference, the maximum change of k in the temperature range of $-50$ to $150^\circ\text{C}$ is also only 9.2% (Fig. S6b). Similarly, temperature coefficient of dielectric constant ($\tau_\kappa$) for hybrid filler is 59 parts per million (p.p.m.) per $^\circ\text{C}$ compared to $-640$ p.p.m. per $^\circ\text{C}$ for CB and 2447 p.p.m. for CNT based composites, respectively, within the temperature range of $-50$ to $150^\circ\text{C}$ (Fig. 5). Because the operating temperature range for
most of the electronic devices falls in −55 to 150 °C, the weak-temperature dependence of $\kappa$ in this temperature range is extremely important. Variation of the $\kappa$ against temperature for our L/0TC$_\kappa$ composites fully satisfies the X8P EIA (Electronic Industries Association) specification$^{36}$ which requires the variation of the capacitance in the temperature interval from −55 to 150 °C to be less than ±10% of its room temperature (25 °C) value for a capacitor. Importantly, in this temperature window, percent changes in dielectric loss tangent of our hybrid filler composite were quantitatively on the same level as dielectric constant (<~10 %) and hence also exhibited the L/0TC characteristics (Fig. S9).

Since our polymer matrix is an elastomer, it is interesting to investigate the effects of physical bending and stretching on the L/0TC$_\sigma$ and L/0TC$_\kappa$ characteristics. Multiple bending cycles didn’t show any significant change in the mechanical properties of composites, indicating their good service life (Table S2). Figure 6a shows that composites retain their L/0TC$_\rho$ and L/0TC$_\kappa$ even after 3000 cyclic bending. Compared to bending, where strain is relatively small (usually below 1%)$^{37}$, it is more challenging to retain the L/0TC properties for stretchable devices where the strain is much larger. We stretched the composite samples up to 15% and found that the composites retain their L/0TC characteristics, as shown in Fig. 6b. These results demonstrated that our L/0TC composites have excellent properties and could be very useful for flexible electronics. We further tested the L/0TC$_\sigma$ and L/0TC$_\kappa$ properties at different AC frequencies (100 Hz, 10 kHz and 1 MHz). As evident in Fig. 6c, at lower frequencies and high temperature (>120 °C), L/0TC$_\kappa$ characteristics are relatively poor. With increasing frequency, L/0TC$_\kappa$ improves, probably because of fall in initial $\kappa$ values of the composites. The decrease in the $\kappa$ values of PPCs with frequencies, especially below $10^4$ Hz, is a common phenomenon associated with relaxation in inter–phase polarization$^{22}$ between the conductive filler and the insulating polymer matrix. The changes in the L/0TC$_\rho$ (red lines in Fig. 6c) with frequency may also arise from the frequency dependent variation in $\rho$, as below
the percolation threshold, $\rho$ presents a critical behavior against the AC frequencies. Apart from the L/0TC$\rho$ and L/0TC$\kappa$, our hybrid filler polymer composites, compared to the neat polymer also show significant enhancement of dielectric constant by 50 times (Fig. 1d), reasonable dielectric breakdown strength (Fig. S10), thermal conductivity by over 100% (increased from $\sim 0.12$ W m$^{-1}$ K$^{-1}$ for pure polymer to $\sim 0.30$ W m$^{-1}$ K$^{-1}$ for 0.06 volume fraction hybrid filler composites), and thermal stability by 29% (Fig. S3b).

To test the versatility of our concept for other polymer matrix, we have also fabricated (CNT+CB) hybrid filler PPCs using dielectric polymer polyvinylidene fluoride (PVDF). By adjusting the ratio of CNT and CB in hybrid filler, L/0TC$\rho$ and L/0TC$\kappa$ characteristics of PVDF based composites are obtained at a filler combination of CNT (0.030) + CB (0.035). Compared to TPE based composites, the PVDF based composites display even better L/0TC$\kappa$ properties (Fig. 6d), especially at higher temperatures. The variation in $\kappa$ was less than $\pm 10\%$ at 150 °C, thus fully satisfying the X8P EIA specification. Note that contrary to TPE composites, where L/0TC characteristics are obtained at equal filler concentrations of CNT and CB (e.g. 0.03 volume fraction for each filler), L/0TC$\rho$ and L/0TC$\kappa$ characteristics for (CNT+CB)/PVDF composites were obtained at different filler volume fractions of each filler (0.03, and 0.035 volume fraction of CNT and CB, respectively). Results from PVDF composites indicate that our strategy could be potentially applied to other polymers.

Maximum energy density ($U_{\text{max}}$) of both the TPE and PVDF based composites calculated by expression $U_{\text{max}} = \kappa \varepsilon_0 E_b^2 / 2$, where $\kappa$ is the dielectric constant of the composites, $E_b$ is the electric breakdown field, $\varepsilon_0$ is the vacuum dielectric permittivity ($8.85 \times 10^{-12}$ F/m), was in the range of 0.05–3.2 J/cm$^3$, close to state-of-the art dielectric polymer BOPP (3.5–4 J/cm$^3$). At 0TC concentration, the $U_{\text{max}}$ for both the TPE and PVDF based composites was 0.17, and 2.80 J/cm$^3$, respectively. Main advantage of our composite is reasonably high energy density without any significant change over a wide temperature window. Energy density of
dielectric composites can be further improved by choosing polymer matrix with better breakdown strength and by using the modified core-shell filler structures, however, main focus of this work is to demonstrate the temperature independent properties of dielectric composites over an ultra-wide temperature range. Breakdown strength and maximum energy density of both TPE and PVDF based composites at different hybrid filler concentrations are given in Fig. S10.

For high-κ materials, only L/0TC property is sought and other modes of temperature dependence are not required. While for conductors, all mode of variation in electrical resistivity with temperature (PTCρ, NTCρ, L/0TCρ) are useful for practical applications. We find that by changing the relative concentration of fillers we can engineer the same composites system for any temperature coefficient of electrical resistivity with desired level of ρ and intensity of respective temperature coefficient. Furthermore, it is also possible to tune the initial ρ and κ value of the composites without changing the mode of temperature coefficient (see supporting section S1). We believe that value of temperature coefficient of a single filler based composite at specific filler concentration, rather simply the concentration of filler, may be much more rational parameter while deciding the composition of a hybrid filler composite to achieve the certain mode of temperature coefficient. This is the reason why in the case of PVDF based composites, different concentrations of CB and CNT rather the equal concentration achieved the L/0TC properties. Value of the temperature coefficient of a single filler based composite will depend upon the polymer matrix, and the percolation characteristics such as shape of the percolation curve, percolation threshold and value of the critical constants (supporting section S1).

Figure 7a compares our results with previous literatures on different dielectric materials including ceramics,49-50 ceramic based composites,41-42 and PPCs and shows widest L/0TCκ window for our composites that is nearly two times wider than previously achieved
value. Also compared to earlier reports on the percolative dielectric polymer composites based on particle\textsuperscript{17, 43} or rod-like fillers\textsuperscript{26-27} our hybrid filler approach offers a much wider (almost 2 times) L/0TC window of 200 °C (−50 to 150 °C). Usually, temperature induced variations in electrical or dielectric properties of PPCs are results of thermal transitions in polymers. Our hybrid filler strategy together with polymer matrix which has suppressed thermal transitions, such as recently reported low-$k$ cross-linked polymer nanocomposites with boron nitride nanosheets fillers\textsuperscript{21} can allow even further extension of L/0TC$_{\kappa}$ and L/0TC$_{\rho}$ windows. Such an outcome can possibly lead to applicability of PPCs at even further higher temperatures compared to what we reported here.

We believe that this hybrid filler strategy is not limited only to CNT and CB but could also be extended to other conducting fillers of different shapes (plate, rod, and spheres), dimensions (1D, 2D, or 3D) and chemical nature (organic or inorganic). In particular, considering the recent surge of interest in 2D materials (graphene like)\textsuperscript{44-46} and their composites \textsuperscript{47-50} and the fact that these composites experience the same sort of temperature dependence as other PPCs\textsuperscript{51-52}, our hybrid filler strategy can also be helpful to grant a better control over the temperature dependence of electrical properties of such composites. This strategy can also possibly be applied to layered composites\textsuperscript{53-56}. Generally, composites based on particle fillers observe PTC effect due to the fact that inter-particle contact is point-type which has characteristics of low contact area.\textsuperscript{57} On the other hand, composites based on large contact area type fillers such as with line contacts (for high aspect ratio rod like fillers)\textsuperscript{58} or face contacts (for plate like fillers)\textsuperscript{59} are expected to have the NTC effect (a summary of inter-filler contacts and associated temperature coefficient is provided in Fig. 7b). Therefore, careful choice of relative filler concentration of low and large contact area fillers in the hybrid can open a new rational and universal route to control the temperature dependence of electrical properties of the hybrid filler based composites.
CONCLUSIONS

We have demonstrated a hybrid filler strategy that gives extraordinary control and versatility to the design of dielectric polymer composites. Our approach allows to fabricate the polymer composites with L/0TC characteristic which is independent of physical deformation and AC frequency. Moreover, this strategy is applicable to various fillers and polymers. By changing the total concentration of CNT+CB fillers and by carefully choosing the relative ratio of CNT and CB, we can obtain all three modes of temperature dependence of $\rho$ and $\kappa$ in a single polymer composite system. Such a simple strategy grants extraordinary control and versatility in designing PPCs with controlled temperature dependence of electric and dielectric properties.

EXPERIMENTAL DETAILS

Multi-walled carbon nanotubes (CNTs) were synthesized by chemical vapor deposition method using ferrocene as catalyst precursor and toluene as a carbon source. Amongst two types, CNTs with diameter, length, and specific surface area of 10 – 30 nm, 5 – 15 μm, and 90– 140 m$^2$/g, respectively, were chosen (see Fig. S11). 99.9 % pure Acetylene CB powder (Product No: 39724), and Polyvinylidene fluoride (PVDF) (Product No: 44080) were purchased from Alfa Aesar company. The average size of the CB particles, surface area, and bulk density was 40 nm, 75 m$^2$/g, and 80 – 120 g/l, respectively (supplier data). Dynamically vulcanized polypropylene based thermoplastic elastomer (TPE) matrix [polypropylene and copolymer of ethylene–propylene–diene monomer (PP/EPDM)] in bead form, supplied by Dawn-BH elastomer Co., Ltd., was chosen as polymer matrix.

For composite preparation, three carbon based fillers [CB, CNT and (CNT+CB) hybrid filler] were used. They were mixed with the TPE and PVDF matrix separately in a twin screw HAAKE Polydrive mixer at 180 °C for 20 min. Processing temperature was set at 180 °C, higher than melting temperature of the polymer matrix for better processing. The mixing
speed was 100 rpm. The composites were hot pressed and cut into sheets with a diameter of 12 mm and a thickness of 1 mm, at 20 MPa pressure and 200 °C temperature for 15 min. For electrical measurements, samples were pasted with silver electrodes on both sides to minimize the contact resistance. For planar capacitor structures, interdigitated Cr (15nm)/Au (50nm) electrodes were deposited on polymer sheets by using e-beam evaporation (see Fig. S12). Typical dimensions of the interdigitated electrodes were 300 µm electrode width 300 µm electrode separations, 1 cm electrode length. Dielectric constant of the samples was measured using an HP 4294 A impedance analyzer in the frequency range of 1 kHz to 40 MHz and temperature range of −50 to 150 °C. Temperature scan rate was 5 °C min⁻¹ with soak time of 2 min for each measurement step. The dielectric breakdown strength was measured based on IEC60243-1 standard. Ten samples for each composition were tested and the results were analyzed using Weibull statistical analysis. The DC volume resistance of the samples along the thickness direction was measured by employing a digital multimeter (below 2 × 10⁷ Ω) and a ZC-36 megger (above 2 × 10⁷ Ω) at progressively elevated temperatures. Rate of change in temperature was set as 5 °C min⁻¹. Morphology of the fillers and microstructures of the fractured surface of the composites was characterized using transmission electron microscopy (TEM, Hitachi H-800) and scanning electron microscope (SEM, Hitachi S-4700), respectively. The melting behaviors of the composites were measured by differential scanning calorimetry (DSC, DuPont TA 2910) from 25 °C to 200 °C at heating rate of 5 °C min⁻¹ under N₂. Thermal stability was determined with a Perkin–Elmer TGA-7 thermo gravimetric analyzer over a temperature range of 20–700 °C at a heating rate of 5 °C/min. The thermal conductivity measurements were conducted on a Transient Hot Disk TPS 2500S instrument (Hot Disk AB, Gothenburg, Sweden) according to ISO 22007-2 standard. Mechanical properties were measured with a universal testing machine (Instron 4465) at room temperature and stretching rate of 200mm/min.
Temperature coefficient of dielectric constant

The temperature coefficient of dielectric constant ($\tau_\kappa$), for a given temperature range (from $T_i$ to $T_f$), is defined as:

$$\tau_\kappa = \frac{(k_f - k_i)}{k_{ref}(T_f - T_i)}$$

(3)

where $k_{ref}$ is the dielectric constant at room temperature, $T_i$ and $T_f$ are the low-end and high-end temperatures, respectively, and $k_i$ and $k_f$ are the dielectric constants at $T_i$ and $T_f$, respectively.

SUPPORTING INFORMATION. Text on all mode temperature dependence of electrical resistivity, text on the percolation characteristics of CNT/TPE composites with various aspect ratio CNTs, fitting of percolation curves, thermal transition of pure polymers and composites, schematics of temperature dependence of filler networks in polymer matrix, temperature dependent dielectric and electric properties of composites based on various combinations of hybrid fillers, dielectric loss tangent vs temperature curves, energy storage capacity of polymer composites, schematics and real images of inter-digitated electrodes, data on mechanical properties and temperature coefficients of polymer composites.

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References

36. Electronic Industries Association Eia Rs-198c 1983.
Figure – 1
Figure – 2
Figure – 3
**Figure 4**

**a** \( f_{CB} > f_{CNT} \)

- CB: Blue circles
- CNT: Gray dots
- Polymer: Orange lines

Temperature increase leads to PTC (Positive Temperature Coefficient).

**b** \( f_{CNT} > f_{CB} \)

Temperature increase leads to NTC (Negative Temperature Coefficient).

**c** \( f_{CB} = f_{CNT} \)

Temperature increase leads to OTC (Ohmic Temperature Coefficient).

Graphs show the change in resistance with temperature for CB and CNT.
Figure – 5
Figure – 6
Figure – 7
Figure captions

Fig. 1. (a) TEM images of carbon fillers. (b) SEM micrographs of all three carbon filler/polymer composites. (c) Schematics of carbon filler networks in polymer matrix.

Fig. 2. Percolation curves of dielectric constant @100Hz and electrical resistivity of composites at various filler concentrations.

Fig. 3. (a) Electrical resistivity of composites as a function of temperature at various filler concentrations below, at, and above percolation threshold). (b) Dielectric constant @100Hz of composites as a function of temperature at various filler concentrations near the percolation threshold.

Fig. 4. Schematics and electrical resistivity/dielectric constant plots of CNT+CB hybrid filler composites with (a) PTC, (b) NTC, and (c) L/0TC characteristics.

Fig. 5. Temperature coefficient of dielectric constant of composites in different temperature ranges.

Fig. 6. L/0TC characteristics of electrical and dielectric properties of CNT+CB hybrid filler/TPE polymer composites (a) after various bending cycles, (b) stretching along the length, (c) at different AC frequencies. (d) L/0TC electric characteristics of CNT+CB hybrid filler/PVDF polymer composites.

Fig. 7. (a) Summary of different inter-filler contact types and most probable temperature coefficient of electrical resistivity associated with their corresponding polymer composites. (b) Comparison of our results with previous studies.