Melamine foam-supported 3D interconnected boron nitride nanosheets network encapsulated in epoxy to achieve significant thermal conductivity enhancement at an ultralow filler loading

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**HIGHLIGHTS**

- Melamine foam was used as a substrate to deposit BNNS via layer-by-layer assembly.
- Melamine skeleton supported 3D BNNS network was fabricated.
- Interconnected BNNS network acts as a heat transfer path in epoxy composite.
- Significant thermal conductivity enhancement was achieved at an ultralow filler loading.

**GRAPHICAL ABSTRACT**

**ABSTRACT**

Realizing high-efficiency thermal conductivity enhancement at low filler loading has a great significance for thermally conductive composite. Herein, three-dimensional (3D) boron nitride nanosheets (BNNSs) wrapped melamine foams (MF@BNNS) were first fabricated by repeated layer-by-layer (L-B-L) assembly using melamine skeleton as substrate and BNNSs as building blocks. The resultant MF@BNNS scaffold with order and interconnected BNNS layer, as a thermally conductive network, was further infiltrated with epoxy resin. As a consequence, a relatively high thermal conductivity of 0.6 W m\(^{-1}\) K\(^{-1}\) was achieved at an ultralow BNNS loading of \(\sim1.1\) vol\%, which is equivalent to a thermal conductivity enhancement of 233% compared to epoxy resin. Besides, the obtained epoxy composite also possesses a good mechanical property and excellent electrical insulativity. This method can be further extended to construct 3D filler network of other 2D layered materials on the melamine foam for high-performance composite.
1. Introduction

With the development of various electronic devices in the trend of higher power, miniaturization and higher integration, using high-efficiency thermal management materials has become more urgent to maintain their reliability [1–3]. Ideal thermal management materials should combine the merits of high thermal conductivity (TC), excellent electrical insulation, low cost and easy preparation [4–7]. Therefore, polymer-based thermally conductive materials show a large potential owing to their easy processing, lightweight and excellent electrical insulation [8,9]. Unfortunately, most of the neat polymers show low TC (around 0.2 W m$^{-1}$ K$^{-1}$) [10,11]. For instance, the TCs of conventional epoxy thermosets are in a quite low range of 0.15–0.21 W m$^{-1}$ K$^{-1}$ because of their isotropic amorphous nature [12]. Although it has been reported that the formation of highly ordered structure can greatly enhance the pristine TC of epoxy thermosets, the used epoxy monomers are high-cost and can’t allow large-scale production [12–14]. Therefore, the most efficient and convenient method to enhance the polymer’s TC is still focused on blending them with many high TC but electrically insulating fillers, such as metal oxides (e.g. Al$_2$O$_3$ [15–17], MgO [18,19]), carbide (e.g. SiC [20,21]) and metal nitride (e.g. AlN [22,23], BN [24–27]). Nevertheless, the impressive TC enhancement always requires a large amount of filler importation because of the low TC enhancement efficiency, which sacrifice the mechanical, processing property and meanwhile cause the high-cost [28,29]. Therefore, obtaining a high TC enhancement at low filler loading is of great significance but also a challenge.

A number of attempts have been made to realize the high-efficiency TC enhancement at a relatively low filler loading, such as filler functionalization [30–32], filler hybrid [33–35], rational orientation of fillers [4–8] and self-assembly into 3D filler network [11,26,35–38]. Recently, preforming 3D interconnected filler network as a thermally conductive path followed by vacuum-assisted impregnation of polymer has been believed a promising strategy to ensure most of the heat transfer along the filler network and finally achieve the high TC enhancement efficiency [24]. In comparison with the composite with randomly distributed fillers, this method requires an extremely low filler loading to form thermally conductive network [36,39]. For instance, Wong et al. prepared a 3D vertically aligned and interconnected graphene network (VAIGN) via an oriented freeze-casting method for epoxy encapsulation and the obtained composite exhibited a high through-plane TC of 2.13 W m$^{-1}$ K$^{-1}$ at an ultralow graphene loading of 0.92 vol% [36]. Taking account of the electrically insulating property of the composite, boron nitride nanosheet (BNNS), an analogue of graphene, is another promising candidate because of its intrinsic high TC and excellent electrical insulativity [40–44]. Similarly, 3D interconnected BNNS networks encapsulated in polymer have also been investigated for the TC enhancement at low BNNS loading. For instance, Xu et al. constructed 3D interconnected NBNS network (VAIGN) via an oriented freeze-casting method for epoxy encapsulation and the obtained composite exhibited a high through-plane TC of 2.85 W m$^{-1}$ K$^{-1}$ at a relatively low BNNSs loading of 9.29 vol% [26]. Huang et al. used cellulose nanofiber-supported 3D interconnected BNNS aerogel as a skeleton to prepare the epoxy composite with large TC enhancement of about 1400% at a low BNNS loading of 9.6 vol% [24].

Although the construction of 3D filler networks can reliably realize the significant TC enhancement at low filler loading, the employed preparation methods (e.g. ice-template [11,36,45], chemical vapor deposition [46–48]) are always time-consumed and processing-complex, hardly meeting the requirement of large-scale production. Recently, commercial polymer foams (e.g. melamine foam, PU foam) with low density and ultrahigh porosity have been extensively investigated as the ideal substrates for the production of flexible electrode materials and water treatment materials via surface functionalization [49–51]. Liu et al. synthesize a macroscopic graphene-wrapped melamine foams (MF-G) by an MF-templated layer-by-layer (L-B-L) assembly to grow polypyrrole (PPy) for flexible supercapacitor [49]. Herein, L-B-L assembly technique enables to form an orderly and compact stacking of graphene nanosheets on the substrate, which is expected to enhance the heat transfer in the graphene layer [52]. It inspires us that by employing BNNSs as building blocks to coat them on the melamine skeleton through L-B-L assembly, it might achieve a compact and well-stacked BNNS layer around the melamine skeleton and then obtain a 3D MF-supported interconnected BNNS network. In comparison with the ice-template and chemical vapor deposition methods, this route is more facile and efficient for the construction of 3D filler network.

Herein, we reported a novel strategy for the fabrication of MF supported 3D BNNS network (MF@BNNS) via multiple L-B-L assembly using BNNS as building blocks. The resultant 3D MF@BNNS scaffold was further encapsulated by epoxy resin to obtain the epoxy composite (EP/MF@BNNS). The through-plane TC of EP/MF@BNNS composites versus L-B-L deposition number of BNNSs was investigated. As a result, a relatively high TC of 0.6 W m$^{-1}$ K$^{-1}$ was achieved at an ultralow BNNS loading of ∼1.1 vol%, corresponding to a TC enhancement of 233% compared to that of neat epoxy. In addition, the composite only shows a slightly decreased mechanical property but maintains an excellent electrical insulativity. These results reveal that this route can reliably realize the significant thermal conductivity enhancement for the polymer composite at an ultralow BNNS loading.

2. Experimental section

2.1. Materials

Polyethyleneimine (PEI, $M_w=10,000$), hydrazine and o-dichloroaniline methane (MOCA) were purchased from Aladdin Chemical Reagent. Isopropanol (IPA) was purchased from Sinopharm Chemical Reagent Co. Ltd. (Shanghai, China). Hexagonal boron nitride powder (h-BN) was purchased from Alfa Aesar. E51 epoxy resin (EP) was provided by Bluestar Wuxi Petrochemical Co. Ltd. (Jiangsu, China). Melamine foam (MF) was purchased from local store.

2.2. Liquid exfoliation of h-BN

2 g of h-BN powders were dispersed into 200 ml IPA (50 vol%)/H$_2$O solution and then sonicated for 12 h to exfoliate h-BN powder by a sonicator (KQ500DB) with an output power of 250 W. Then, the dispersion was centrifuged for 30 min at 1000 rpm to separate the non-exfoliated BN, obtaining the exfoliated BNNSs.

2.3. Preparation of MF@BNNS foam

For the preparation of MF@BNNS, layer-by-layer (L-B-L) assembly method was used to coat BNNS onto the MF skeleton through the electrostatic interactions of positively charged PEI and negatively charged BNNSs. As shown in Fig. 1, in a typical L-B-L assembly cycle, the fresh MF foam was firstly immersed into the PEI aqueous solution (1 mg/ml) for 2 min, rinsed several times with DI Water, and then further immersed into the BNNS dispersion (∼0.5 mg/ml) in IPA (50%)/H$_2$O solution for 2 min, also rinsed several times with DI water. This procedure was repeated multiple times to prepare the MF@BNNS-X foam with different L-B-L deposition cycles (X is the number of deposition cycles).

2.4. Preparation of EP/MF@BNNS composite

The EP/MF@BNNS composites were prepared via the vacuum-assisted infiltration of epoxy. First, 20.1 g of curing agent (MOCA) was homogeneously dissolved in 60 g of E51 epoxy resin monomer under vigorous stirring at 100 °C. Subsequently, the as-prepared MF@BNNS scaffold was completely immersed into the epoxy resin mixture and
then transferred into a vacuum oven at 120 °C for 20 min to remove the air. After that, the resin mixture was cured at 150 °C for 2 h and 180 °C for 2.5 h. The obtained composites were denoted as EP/MF@BNNS-X, where X is the number of deposition cycles. The BNNS loading in final epoxy composite is estimated as (the subtraction of the mass of MF@BNNS and MF before and after L-B-L assembly)/(mass of the final composite after epoxy infiltration). Based on the densities of BNNS and epoxy resin, the estimated mass fraction can be further transformed to volume fraction. Moreover, EP/BNNS composites with randomly distributed BNNS were also prepared by directly adding BNNS into the epoxy resin at an identical curing process.

2.5. Characterization

The morphology of the exfoliated BNNSs was characterized by Transmission electron microscopy (TEM) (JEOL JEM2011 F, Japan) at 200 kV. The thickness of the exfoliated BNNSs was determined by Atomic force microscopy (AFM) (Bruker Multimode 8, Germany) using the tapping mode. The cross-section morphology of the MF@BNNS scaffolds and EP/MF@BNNS composites were observed by Scanning electron microscopy (SEM) (Zeiss Ultra 55, Germany) with EDX. XRD patterns of the h-BN and exfoliated BNNSs were recorded on X’pert PRO Panalytical (Netherland) with Ni-filtered Cu Kα radiation (40 kV, 40 mA). Zeta potentials of the PEI and BNNSs were investigated on Zetasizer Nano-ZS90 (England). The hydrophilicity of the MF and MF@BNNS was characterized by the static water contact angles using OCA15 (Data-physics Co. Ltd., Germany). The tensile curves of the composites were measured on a universal electronic tensile machine UTM4000 (SUNS, China). The volume resistivity of the composites was measured on a Keithley Electrometer. The through-plane thermal diffusivity (α, m² s⁻¹) of the composites were detected by using a laser-flash diffusivity instrument LFA 447 (NETZSCH, Germany). Before the measurement, the square samples (1 cm × 1 cm × 1.4 mm) were firstly spray-coated with a thin layer of fine graphite powder on both sides. The density (ρ, g cm⁻³) of a specimen was calculated by the equation: ρ = m/ν and the eventual thermal conductivity (λ, W m⁻¹ K⁻¹) was calculated by the equation: λ = α × c × ρ.

3. Results and discussion

The formation procedures of MF@BNNS foam and EP/MF@BNNS...
Composite are illustrated in Fig. 1. Briefly, the neat MF skeleton was first wrapped by PEI/BNNS coating via the alternate deposition of anionic BNNSs and cationic PEI by a L-B-L assembly technique and then encapsulated by epoxy resin to prepare the epoxy composite. Considering the greatly larger thermal conductivity of ultrathin BNNS than that of the pristine h-BN, h-BN powder was first exfoliated to few-layered nanosheets through vigorous sonication treatment in a IPA/H_2O mixed solution. After centrifugation and collection, the morphology of the exfoliated BNNS was characterized by transmission electron microscope (TEM).

As shown in Fig. 2a, the obtained BNNSs show a flexible and ultrathin feature. High-resolution TEM image reveals that the BNNS still keep impeccable lattice structure after exfoliation. Furthermore, the size distribution of the exfoliated BNNSs was investigated by counting over 150 pieces of BNNSs in TEM images. One can see from Fig. 2b that the size of the exfoliated BNNSs is mainly distributed in the range of 0.5–1.5 μm. The large size of BNNSs is expected to reduce the interface between BNNSs along the filler network. Atomic force microscopy (AFM) was further employed to determine the thickness of the exfoliated BNNS (Fig. 2c). The corresponding height profile shows that the thickness of the majority of the exfoliated BNNSs is about 3–5 nm, suggesting the multilayered feature of the exfoliated BNNSs. In addition, the crystalline nature of h-BN powder and the exfoliated BNNSs were recorded by X-ray diffraction (XRD) measurement (Fig. S1). The XRD pattern of h-BN and the exfoliated BNNSs both show a series of diffraction peaks, which correspond to (0 0 2), (1 0 0), (1 0 1), (1 0 2) and (0 0 4) planes, respectively [53]. After exfoliation, the peak positions of the BNNSs are almost in accord with those of pristine h-BN powder, suggesting that the hexagonal lattice is not damaged during the exfoliation process. With the (0 0 2) peak as a reference, the intensity of (1 0 0), (1 0 1), (1 0 2) and (0 0 4) diffraction peaks of BNNSs significantly decreases relative to h-BN powder, which might be attributed to the enhanced exposure of (0 0 2) plane because of the successful peel-off of few-layered BNNSs from h-BN along the (0 0 2) plane [29,53].

In this work, PEI and BNNS were used as the cationic and anionic adsorbent, respectively, to alternately coat them on the neat MF skeleton by a conventional L-B-L assembly technique. Therefore, the ζ potentials of cationic PEI in water and BNNSs in IPA/water mixed solution are necessary to measure. As shown in Fig. 3a, the ζ potential of PEI in water is around 56 mV and the ζ potential of BNNSs in IPA/water is about ~18 mV, suggesting the accessible electrostatic assembly of BNNS and PEI. Fig. 3b displays the optical pictures of BNNS suspension containing PEI or not. Without PEI, BNNSs show a homogeneous distribution with a typical Tyndall effect. The adding slight PEI, large aggregates start to form because of strong electrostatic interaction between PEI and BNNS.

Through alternate adsorption of cationic PEI and anionic BNNSs on MF skeleton, MF@BNNS complex foam with controllable PEI/BNNS bilayers can be achieved. As shown in Fig. 3c, the neat MF exhibits an open-cell microstructure with a largely hydrophilic nature for water infiltration. After coated by BNNSs, a visible hydrophilicity-hydrophobicity inversion would be triggered due to the intrinsic hydrophobicity of BNNSs. The obtained MF@BNNS foam shows a large water contact angle of 123° on the surface, indicating the successful coating of BNNS on the MF skeleton. When the MF@BNNS foam was cut into two pieces, the freshly cut face still exhibits a high water repellency with a water contact angle of 118°, suggesting the homogeneous deposition of BNNSs throughout the cell walls. The morphology of the pristine and BNNS-modified MFs is characterized by scanning electron microscopy (SEM). The pristine MF shows a 3D interconnected macro pore structure with an average pore size of ~100 μm and a smooth surface (Fig. 4a and Fig. S1). As expected, SEM images of MF@BNNS foams show that BNNSs are tightly anchored around the cell walls of MF skeleton through the electrostatic interaction-mediated L-B-L assembly, but do not alter the open-cell microstructure of pristine MF (Fig. 4b–d). When the number of L-B-L deposition increases to 20 times, the BNNS plates contact and well stack one by one along the MF skeleton to form a continuous network (Fig. 4e). Further increase of the L-B-L cycles is expected to thicken the BNNS coating and make the thermally conductive pathways more perfect (Fig. 4f). The enlarged image further reveals that BNNSs have a compact stacking of their basal plane parallel to the MF surface. Importantly, the cationic PEI filled the interval between adjacent BNNSs would remove the air gaps and bridge BNNSs via electrostatic interaction, which can reduce the thermal contact resistance of adjacent BNNSs [54]. Moreover, the formation of 3D BNNS
network is further elucidated by the corresponding element mapping image of element boron, which shows a homogeneous, continuous distribution throughout the MF skeleton (Fig. 5). Likewise, MF foam can also be closely packed by graphene nanosheets (GNSs) through the L-B-L assembly of cationic PEI and anionic graphene oxide (GO) (Figs. S3 and S4). As shown in Fig. S5, it can be seen that the MF skeleton is homogeneously wrapped by GNSs and the increasing deposition number causes a rougher surface with a wrinkled feature.

To fabricate MF@BNNS based composites, epoxy resin with low viscosity was used to infiltrate the obtained MF@BNNS foam. Fig. 6a exhibits the optical image of the cured epoxy resin, EP/MF, and EP/MF@BNNS samples. One can see that the epoxy resin has a higher transmittance compared with that of the EP/MF composite. For EP/MF@BNNS composites, the transmittance is further decreased due to the incorporation of BNNS. But they still show a relatively high transmittance. In order to observe the distribution of BNNS in the epoxy matrix, the fractured surface of neat epoxy is very smooth owing to the high brittleness of epoxy resin (Fig. S6). After encapsulation by epoxy, the open pores in MF@BNNS-20 foam have been fully filled with epoxy resin, as shown in Fig. 6b. However, the fractured EP/MF@BNNS remain to keep a relatively smooth surface, indicating a negligible effect of MF@BNNS skeleton on the intrinsic mechanical property of epoxy matrix. Importantly, the enlarged image further shows that the 3D interconnected BNNS network is well maintained throughout the epoxy matrix, which acts as an efficient pathway for heat transfer (Fig. 6c–d). Furthermore, the MF@BNNS foam has a good adhesion with the epoxy resin and no obvious interfacial debonding can be observed. This compact interface is favorable for the decrease of interface thermal resistance between BNNS and epoxy resin. On the other hand, the fractured cross-section of EP/MF@GNS composite was also investigated. Fig. S7 shows that a continuous well-stacked GNS network is distributed at the interface between MF skeleton and epoxy resin as the highway for heat transfer.

The through-plane TC of the prepared EP/MF@BNNS composites was measured by the laser-flash method at room temperature. EP/BNNS composites with randomly distributed BNNSs, as a reference, were also investigated for comparison under same conditions and its fractured surface morphology is shown in Fig. S8. Fig. 7a depicts the
through-plane TC of EP/MF@BNNS composites versus L-B-L deposition numbers. The neat epoxy resin shows a low TC of 0.18 W m\(^{-1}\) K\(^{-1}\) due to its amorphous nature. However, the incorporation of 3D MF@BNNS framework causes a prominent increase of the TC. Below five deposition cycles, the TC of EP/MF@BNNS composites only shows a slight increase because of the insufficient contact of BNNSs. As the increase of deposition cycle, TC of the composite shows a significant enhancement and a high value of 0.53 W m\(^{-1}\) K\(^{-1}\) is achieved after 20 L-B-L deposition cycles, which is nearly two times higher than that of the neat epoxy resin. These results indicate that the formation of 3D BNNS network enables to construct thermally conductive networks inside the epoxy matrix, resulting in the large enhancement of TC of the EP/MF@BNNS composite. Further increase of the deposition cycle can promote the TC (0.6 W m\(^{-1}\) K\(^{-1}\) at 30 L-B-L cycles) but the enhancement efficiency show a remarkable decrease, which is a common phenomenon for the polymer composite above the permeation threshold \([5,55,56]\). Herein, it should be noted that the BNNS content in EP/MF@BNNS-20 and EP/MF@BNNS-30 composites are estimated to be only \(\sim 0.82\) and \(\sim 1.1\) vol\%, respectively. While for the conventional EP/BNNS composite with randomly distributed BNNSs, obtaining a similar level of TC need a higher BNNS loading above 19.7 vol\%, suggesting the large superiority of preforming MF@BNNS foam as a 3D thermally conductive path on the significant enhancement of TC at a ultralow filler loading (Fig. 7b) \([24,55]\). To better compare with the previous investigations focused on the polymer/BN composite, TC enhancement per 1 vol\% filler loading (\(\eta\)) is used to characterize the TC enhancement efficiency of BNNSs to composites, which is defined as

\[
\eta = \frac{TC - TC_0}{100V_{TC}} \times 100\%
\]

where TC and TC\(_0\) represent the thermal conductivity of the composites with given BNNS content and neat polymer matrix, respectively, and V is the volume fraction of BNNS in composites. Table 1 summarizes the \(\eta\) values of the previously reported polymer composites using BNNS or h-BN as the filler. One can see that the \(\eta\) value in this work is higher than those of all previously reported BN-based polymer composites, to the best of our knowledge. Furthermore, compared with the fabrication of 3D porous BNNS network by ice-template or chemical vapor deposition methods, the method reported in this work is relatively much more facile and efficient. Moreover, the TC of the EP/MF@GNS composites versus L-B-L deposition numbers has been also investigated. As shown in Fig. S9, the increased deposition number of GNS is favorable to the enhancement of TC and a high TC of 0.54 W m\(^{-1}\) K\(^{-1}\) is obtained for the EP/MF@GNS composite with 10 deposition cycles. In this case, the GNS loading in EP/MF@GNS-10 composite is estimated to be 0.37 vol\%, suggesting the very high TC enhancement efficiency of GNSs to composites.

To demonstrate the enhanced thermal management ability of EP/MF@BNNS composite, the surface temperature variation of the epoxy composites with time during cooling was captured by an infrared thermal camera. EP/BNNS composite with 19.7 vol\% BNNS, neat epoxy, EP/MF@BNNS-10 and EP/MF@BNNS-20 composites were first
placed on the same hotplate at 80 °C for 30 min to ensure uniform sample temperature, and then the surface temperature variation of these samples as a function of cooling time at room temperature environment were recorded. As shown in Fig. 8a, during the heat dissipation process, one can see that the EP/MF@BNNS-20 composite and EP/BNNS composite with 19.7 vol% BNNS shows much faster temperature decrease with cooling time in comparison with the neat epoxy sample because of their higher through-plane TC.

**Table 1**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fill loading (vol%)</th>
<th>TC (W m⁻¹ K⁻¹)</th>
<th>TC enhancement (%)</th>
<th>η</th>
<th>Test method</th>
<th>Year [Ref.]</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCL/PCL-g-BNNS</td>
<td>−11.2</td>
<td>1.01</td>
<td>400</td>
<td>−35.7</td>
<td>Laser flash</td>
<td>2016 [32]</td>
</tr>
<tr>
<td>PDMS/PVA/BNNS</td>
<td>15.6</td>
<td>1.94</td>
<td>977</td>
<td>62.6</td>
<td>Laser flash</td>
<td>2017 [4]</td>
</tr>
<tr>
<td>EP/BNNs/Fe</td>
<td>38.5</td>
<td>3.59</td>
<td>1690</td>
<td>43.9</td>
<td>Laser flash</td>
<td>2016 [57]</td>
</tr>
<tr>
<td>PMMA/NF-BNNS</td>
<td>−69.1</td>
<td>10.22</td>
<td>4545</td>
<td>65.8</td>
<td>Laser flash</td>
<td>2016 [29]</td>
</tr>
<tr>
<td>PEI/PI-BN</td>
<td>−49.1</td>
<td>1.7</td>
<td>672</td>
<td>13.7</td>
<td>Laser flash</td>
<td>2014 [58]</td>
</tr>
<tr>
<td>TmBNN-silicone</td>
<td>9.2</td>
<td>0.58</td>
<td>−241</td>
<td>26.2</td>
<td>Laser flash</td>
<td>2015 [59]</td>
</tr>
<tr>
<td>EP/VmBNN</td>
<td>−13.5</td>
<td>0.85</td>
<td>420</td>
<td>−31.1</td>
<td>Laser flash</td>
<td>2013 [55]</td>
</tr>
<tr>
<td>EP/P-BN</td>
<td>40</td>
<td>9.0</td>
<td>−4400</td>
<td>−11.0</td>
<td>Laser flash</td>
<td>2017 [25]</td>
</tr>
<tr>
<td>EP/3D-BNNS</td>
<td>9.3</td>
<td>2.4</td>
<td>−1100</td>
<td>118.3</td>
<td>Laser flash</td>
<td>2015 [36]</td>
</tr>
<tr>
<td>GE/BN</td>
<td>19.6</td>
<td>0.64</td>
<td>−156</td>
<td>−8.0</td>
<td>Steady-state</td>
<td>2018 [60]</td>
</tr>
<tr>
<td>PA6/BNNs/graphene</td>
<td>−4.8</td>
<td>0.89</td>
<td>−354</td>
<td>−73.8</td>
<td>Laser flash</td>
<td>2016 [37]</td>
</tr>
<tr>
<td>HDPE/oriented BN</td>
<td>−24</td>
<td>3.57</td>
<td>−865</td>
<td>−36</td>
<td>Laser flash</td>
<td>2017 [27]</td>
</tr>
<tr>
<td>EP/MF@BNNS</td>
<td>−1.1</td>
<td>0.60</td>
<td>239</td>
<td>217.3</td>
<td>Laser flash</td>
<td>This work</td>
</tr>
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Fig. 8. (a) Infrared thermal images with cooling time of EP/BNNS composite with 19.7 vol% randomly distributed BNNS (S1), neat epoxy (S2), EP/MF@BNNS-10 composite (S3) and EP/MF@BNNS-20 composite (S4); (b) Stress-strain curves of neat epoxy, EP/MF@BNNS-20 composite and EP/BNNS composite with 19.7 vol% randomly distributed BNNS. (c) Volume resistivity of epoxy resin, EP/BNNS composite with 19.7 vol% randomly distributed EP/MF and EP/MF@BNNS-20 composite.

It can be seen that incorporation of MF@BNNS-20 skeleton into epoxy has a little negative effect on its mechanical property. However, at the similar level of thermal conductivity, the EP/BNNS composite with 19.7 vol% of BNNS shows a large deterioration of mechanical property owing to the serious aggregation of BNNSs and weak filler/epoxy interfacial adhesion. These results suggest that our method can achieve the significant TC enhancement and simultaneously maintain the good mechanical property of epoxy resin. In addition, a high enough electrical resistivity of the heat conducting materials is important for some applications in which good electrical insulation is another key factor.
need to be considered. Thus the volume electrical resistivity of the neat epoxy and epoxy composites were recorded, as shown in Fig. 8c. In comparison with neat epoxy, incorporation of MF and MF@BNNS-20 skeletons have a negligible influence on the volume electrical resistivity of epoxy matrix. Despite this, all of them exhibit a high volume electrical resistivity over 10¹⁵ Ω cm, indicating an excellent electrical insulation property of them.

4. Conclusions

In summary, a novel epoxy composite with significant TC enhancement efficiency was fabricated by infiltrating 3D BNNS-wrapped melamine scaffold with epoxy resin. The 3D melamine foam supported BNNS network was first fabricated by the repeated L-B-L assembly of anionic BNNSs and cationic PEI on commercial melamine foam driven by electrostatic interaction. As a result, the obtained epoxy composite exhibits a high TC enhancement of 233% in comparison with that of epoxy resin at an ultralow BNNS loading of 1.1 vol%. In addition, this strategy can also be extended to construct 3D interconnected graphene network on the melamine scaffold for the enhancement of the corresponding epoxy composite.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.cjcl.2018.04.196.

References
