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Graphene-decorated silica stabilized stearic acid as a thermal energy storage material†

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Novel thermal energy storage materials were synthesized from graphene-decorated silica (SG) and stearic acid (SA) by vacuum impregnation method. Three kinds of SG (SG₁, SG₅, and SG₁₀) were prepared by decorating silica with different contents of graphene and were then used to stabilize SA to prepare SA/ SG₁, SA/SG₅, and SA/SG₁₀ composites. The structures and thermal energy storage performances of the SA/SG composites were investigated. It is of interest that the thermal energy storage behaviors of the SA/ SG composites were dramatically changed with different contents of graphene, presenting more than one endothermal or exothermal peak in the differential scanning calorimetry (DSC) curves while pure SA had only one. The SA in SA/SG_1 and SA/SG_5 showed higher crystallinity (F_c , 84.44% and 84.39%) and greater effective energy storage per unit mass (E_{ef} , \sim 150 J g⁻¹) than that of SA in SA/SG₁₀. These thermal energy storage behaviors and properties were revealed to be related to the pore structures of the SG. The thermal stability of the SA/SG composites was analyzed by a thermogravimetric analyzer (TGA), and the SA/SG composites have good thermal stability. Addition of graphene was beneficial to the enhancement in thermal conductivity of the SA/SG composite, which could reach $0.90~\mathrm{W}~\mathrm{m}^{-1}~\mathrm{K}^{-1}$, 1.05 W m $^{-1}$ K $^{-1}$, and 1.12 W m $^{-1}$ K $^{-1}$ for SA/SG₁, SA/SG₅, and SA/SG₁₀, respectively; and were 246%, 304%, and 331% higher than pure SA, respectively. SA/SG₅ has potential for application in thermal energy storage, especially in thermal gradients due to it having both high $E_{\rm ef}$ and thermal conductivity.

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1. Introduction

Energy storage is nowadays considered as a key element in the modern energy supply chain, and research on energy storage technologies has been gaining more and more attention. 1-5 Thermal energy storage (TES) has been recognized as one of the most efficient ways to enhance the energy efficiency and to relieve the mismatch between energy supply and demand.6-8 TES works by heating thermal energy storage materials (TESMs) during a charging period and then releasing the heat when the energy is needed.8 Thus, TESMs play a crucial part in thermal energy storage systems, including thermal management application.9,10 Finding new TESMs has become more and more important. Phase change materials (PCMs) are a good choice for thermal energy storage systems because the PCMs have high energy storage density and can storage energy in isothermal process.11 Nevertheless, PCMs still have some disadvantages, including the leakage during the phase change process and low thermal conductivity of most PCMs, which limit their wide application for thermal energy storage.11-13 Stabilizing PCMs

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with supporting matrix (e.g., carbon materials, metal or their oxides, porous structure materials, nanomaterials, etc.) into form-stable composite PCMs is the most promising practical method to overcome these problems.^{6,11,14-21}

Silica-based materials has been used as supporting matrix due to it can make liquid PCMs easy to handle and protect PCMs from harmful interactions with the surrounding materials and environment during the phase change process.22 Fang et al. successfully used silicon dioxide or silica shell to stabilize nhexadecane,23 lauric acid,24 and octadecane,11 respectively, and the thermal storage capacities of the prepared composite PCMs were 100-230 J g⁻¹. Chung et al.²⁵ successfully impregnated octadecane and BioPCM into the pore of micronized silica and the prepared composite PCMs also had considerable thermal storage capacities (86.07-108.1 J g⁻¹). The octadecane and BioPCM molecules were held easily into the pore of micronized silica by these physical interactions and thus leakage of the melted PCM from the porous was prevented. Wang et al.26 skillfully impregnated stearic acid into the silica fume matrix and its thermal storage capacity was 84.47 J g-1 while the maximum percentage of stearic acid in the composite was 46 wt%. Jeong et al.27 also utilized the silica fume to support hexadecane, octadecane, and paraffin, respectively. The thermal storage capacities of the prepared composite PCMs were 50-90 J g^{-1} and they were useful for applying to building materials. Min et al.22 for the first time used the radial mesoporous silica to

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support polyethylene glycol, and their thermal storage capacities were 60-120 J g⁻¹ with different contents of PEG. Song et al.²⁸ used the silica shell microencapsulated caprice-stearic acid with a thermal storage capacity of 91.48 J g^{-1} while the encapsulation ratio was 56.7 wt%, and prepared composite PCMs showed good thermal reliability. Also, the silica shell encapsulated phase change materials were successfully fabricated by Liu et al.29 and the thermal storage capacities were range from $30-110 \, \mathrm{J g^{-1}}$ with different encapsulation ratios. These above mentioned silicabased composite PCMs had considerable thermal storage capacities. Therefore, the silica is a promising supporting matrix to stabilize PCMs and prevent the melted PCMs leakage. Also the thermal conductivities of PCMs were improved to 0.6-0.8 W m⁻¹ K⁻¹ by silica.^{29,30} As an advanced heat transfer organic materials with excellent thermal conductivities, the PCMs preferably has a thermal conductivity above 1 W m⁻¹ K⁻¹, which broadens its wide application for thermal energy storage. 31,32

In this paper, in order to enhance the thermal properties of silica-based composite PCMs, we firstly decorated silica by graphene, which has attracted a great deal of attention in recent years because of its high thermal conductivity.33 Then we synthesized novel composite PCMs (SA/SG) from graphenedecorated silica (SG) and stearic acid (SA) by a simple blending and impregnation process. The structures and thermal energy storage performances of the SA/SG composites were investigated. It is of interest that the thermal energy storage behaviors of SA/SG composites were dramatically changed along with different contents of graphene, presenting more than one endothermal or exothermal peaks in the DSC curves while pure SA only one of that. The thermal conductivity of the SA/SG reached 1.12 W m⁻¹ K⁻¹, which was 331% higher than that of pure SA.

2. **Experimental**

2.1. Materials

Silica sol (SiO2, 30 wt%) was supplied by Research Institute Of Tsinghua University In Shenzhen, China. Graphite was purchased from Qingdao Graphite Co., Ltd., China. KMnO₄, H₂O₂ and NaNO₃ were purchased from Xilong Chemical Co., Ltd. H₂SO₄ and HCl was purchased from Research Institute Of Chemical Industry In Zhuzhou. (3-Aminopropyl)-triethoxysilane (APTS) was purchased from Nanjing Compton Shuguang Organosilicon Chemical Co., Ltd. (China). Stearic acid (SA, CH₃(CH₂)₁₆COOH) was supplied by Tianjin Hengxing Chemical Reagent Co., Ltd., China.

2.2. Preparation procedures of the composites

Typical procedures were depicted in Fig. 1, including preparation of the SG and the SA/SG composites.

2.2.1. Preparation of the SG. The SG were prepared from the GO and SN. The GO dispersion was prepared by simple sonication of graphite oxide that was obtained from natural graphite using a modified Hummers' method. 34,35 Graphite (2.5 g) and NaNO₃ (1.25 g) were mixed with a 95% H₂SO₄ (60 mL) in a 250 mL flask. The mixture was stirred for 30 min in an ice

bath. Under vigorous stirring, 7.5 g of KMnO₄ was added to the suspension. The rate of addition was controlled to keep the reaction temperature lower than 20 °C. The ice bath was then removed, and the mixture was stirred at 35 °C for 5 h. Afterward, 150 mL of deionized H₂O was slowly added to the mixture still under vigorous agitation. The reaction temperature was observed to rapidly increase to 98 °C. The diluted suspension was stirred for 12 h. Then, 25 mL of 30% H₂O₂ was added to the mixture. To purify, the mixture was washed with 5% of HCl and then with deionized H₂O for several times to obtain the graphite oxide sample. The final GO dispersion was prepared by diluted and sonication of graphite oxide, and concentration of GO is 1 mg mL^{-1} .

The SN was prepared by silanizing of SiO₂. Silanized of SiO₂ with the APTS coupling agent: the silica sol (300 g) was mixed into the 15 mL APTS dissolved in 584 mL DI H₂O by stirred at 50 °C for 12 h to ensure completion of the silanization reaction. The obtained product was SiO₂-NH₂ and labelled as SN (SiO₂,

And then, the silanized silica suspension (SN, 50 g) was slowly added into the GO aqueous dispersion and stirred for 2 h. The obtained slurry was dried at 60 °C for 24 h. The SNfunctionalized GO (GO ratio of 1%, 5%, and 10%) was designated as SNGO₁, SNGO₅, and SNGO₁₀, respectively. Three kinds of SNGO (SNGO₁, SNGO₅, and SNGO₁₀) were then pyrolyzed at 800 °C under nitrogen atmospheres, generating the SG₁, SG₅, and SG₁₀, respectively.

2.2.2. Preparation of the SA/SG composites. The composites were prepared using vacuum impregnation method: 3.9 g support (SG₁, SG₅, and SG₁₀) and 2.1 g SA were placed inside a conical flask (the content of SA in the composites was designed as 35 wt%), and a device of preventing of backward suction was used to connect the conical flask with vacuum pump. The vacuum of conical flask was evacuated to -0.1 MPa for 5 min. Then, the conical flask was placed in thermostatical water bath at 95 °C for 30 min. The vacuum pump was then turned down and allowed the air to enter the flask again, with ultrasonic heating at 80 °C for 10 min. After cooling, composites were obtained and named as SA/SG₁, SA/SG₅, and SA/SG₁₀, respectively.

Characterization

The differential scanning calorimetry (DSC) analysis of SA and SA/SG composites was performed using TA Instruments Q2000 at a heating rate of 5 °C min⁻¹ from 20 °C to 80 °C under a constant stream of nitrogen at the atmospheric pressure. Liquid nitrogen was used for cooling the sample during the freezing period. Thermo-gravimetric analysis (TGA) and differential scanning calorimetry (DSC) analysis were conducted at a heating rate of 10 °C min⁻¹ up to 550 °C in nitrogen atmosphere, using α -Al₂O₃ crucibles in NETZSCH STA 449F3. X-ray diffraction (XRD) was carried out by using a Rigaku D/max-rA analyzer (Cu-Ka) under the following conditions: voltage of 40 kV; current of 40 mA; scan range from 5° to 80° and step size of 0.02°. The thermal conductivity of the samples at room temperature was measured by means of steady-state heat flow method using a thermal conductivity tester (DRX-II-RW,

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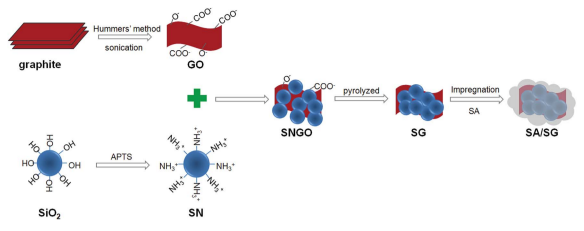


Fig. 1 Schematic illustration of the preparation process of the SG and SA/SG composites

Xiangtan Huafeng Instrument Manufacturing Co., Ltd., Hunan, China). The microstructures of the samples were investigated by a JEOL JSM-6360LV scanning electron microscopy (SEM). Fourier transform infrared spectroscopy (FTIR) spectra were recorded by using a Thermo Electron Corporation Nicolet 6700 FTIR spectrometer in the range of 4000–400 cm⁻¹. Nitrogen gas adsorption–desorption isotherms for the SG were measured by an ASAP 2020 surface area and porosimetry system, and the SG samples were outgassed prior to the measurement at 523.15 K and 10⁻⁴ mbar for 12 h.

3. Results and discussions

3.1. XRD and FTIR characterization for preparing SG

The XRD patterns of the graphite before and after modified Hummers method were illustrated in Fig. S1.† It is seem that GO showed two diffraction peaks at $2\theta = 10.4^{\circ}$ and 11.3° , indicating the complete exfoliation of the graphite.34 The XRD patterns of the SiO2 before and after silanized with APTS were illustrated in Fig. S1,† suggesting that the SiO2 still maintained amorphous silica. Fig. S2† showed the XRD patterns of the SNGO before and after pyrolyzed at 800 °C under nitrogen atmospheres. It indicated that the samples kept the amorphous form after pyrolyzed under nitrogen atmospheres. The silica present amorphous form or crystalline form at different atmospheres.37,38 The silica used here pyrolyzed at 800 °C under nitrogen atmosphere and air atmosphere were amorphous form and crystalline form, respectively (Fig. S3†), and these results were in accord with the References. 37,38 Fig. S4† showed the FTIR spectrum of the graphite, GO, SiO₂, and SN. Compared with the FTIR spectrum of graphite, that of GO exhibited the bands at around 3410 cm⁻¹, 1726 cm⁻¹, 1623 cm⁻¹, 1145 cm⁻¹, and 1114 cm⁻¹, which can be attributed to the oxygen containing functional groups on GO.39 In the FTIR spectrum for the SiO2 and SN, the weakened band at 1384 cm⁻¹ indicated that the APTS bonded to SiO₂. The FTIR spectrum of the SNGO and SG were shown in Fig. S5.† It is seem that the bands at 3450 cm⁻¹ and 1630 cm⁻¹ in the SNGO had disappeared after pyrolyzed (see the FTIR spectrum of SG in Fig. S5†), verifying the reduction of GO to G under nitrogen atmospheres.40

3.2. Crystallization characteristics of the SA/SG composites

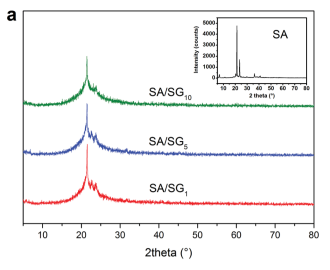
The XRD patterns of the SA/SG composites with different contents of graphene were illustrated in Fig. 2a. The reflections at $2\theta=21.2^\circ$ and 23.5° were characteristic of the SA (inset of Fig. 2a). After the SA was impregnated into the SG, the characteristic reflections of the SA were observed in the XRD patterns of the SA/SG composites. Comparing the XRD patterns of pure SA (inset of Fig. 2a) and SG (Fig. S2†), a new weak peak emerged at $2\theta=22.6$, 22.6, and 23.0 in the SA/SG₁, SA/SG₅, and SA/SG₁₀, respectively, should be noticed. These peaks suggested a novel interaction between the SA and the SG matrix, which could had an effect on its phase change process.

3.3. FTIR analysis of the SA/SG composites

FTIR spectra of the SA, SG, and SA/SG composites were depicted in Fig. 2b. From the spectra of pure SA, the characteristic vibration peaks of the SA were at 2920 cm⁻¹ (-CH₃ group symmetrical stretching), 2850 cm⁻¹ (-CH₂ group symmetrical stretching), 1704 cm⁻¹ (C=O stretching), 1465 cm⁻¹ (-OH bending), 1382 cm^{-1} (-OH bending), 1113 cm^{-1} (C=O stretching), 723 cm⁻¹ (-OH swinging), and 680 cm⁻¹ (-OH bending).41,42 From the spectra of the SG1, SG5, and SG10, the characteristic vibration peaks of the SG were at 1096 cm⁻¹ (Si-O-Si asymmetry stretching and), 804 cm⁻¹ (symmetric stretching mode of the Si-O-Si bond), and 463 cm⁻¹ (Si-O-Si bending).38 According to the FTIR spectra of the SA/SG composites, the peak at 804 cm⁻¹ of the SG weakened in the SA/SG composites and the peak at 1096 cm⁻¹ shifted to a higher wavenumber of 1113 cm⁻¹. This phenomenon also suggested a novel interaction between the SA and the SG matrix.

3.4. Morphological investigation of the SA/SG composites

The SEM images and EDS patterns of the SG_5 and SA/SG_5 were shown in Fig. 3. The irregular particles of SG_5 were observed from Fig. 3a and emerged with the size ranges of 0.5–30 μ m. A lot of small particles dispersed around the large particles. The EDS patterns of the SG_5 (inset of Fig. 3a) suggested that the graphene and the silica were successfully mixed together and the content of the graphene in the SG_5 approached the designed value (5%).



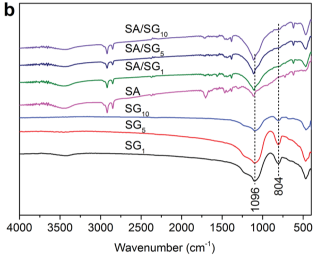


Fig. 2 (a) XRD patterns of the SA/SG composites and (b) FTIR spectra of the samples.

After the impregnation, the sizes of irregular particles were enlarged by introducing the SA and ranged from 5 μm to 50 μm , indicating from SEM images of the SG5 (Fig. 3b). From the EDS patterns of the SG5 (inset of Fig. 3b), the mass ratio of C element increased due to the added SA and also approached the designed value. By comparing Fig. 3a with 3b, it can be concluded that the surfaces of SG5 were occupied by SA in the SA/SG5, and the SG particles were surrounded by the SA and formed aggregates. Interestingly, some flakes emerged in the smaller particle of the SA/SG5 (Fig. 3c) and also in larger particle of the SA/SG5 (Fig. 3d). It illustrated that the SA was fully adsorbed onto the SG5 and the mass ratio of SA was no overdose.

3.5. Thermal stability of the SA/SG composites

For the application of PCMs, thermal stability determines their practical environment and should be thermally stable over its operating temperature range. So, good thermal stability always is a critical parameter for TES applications. TGA is the most commonly used method to characterize the thermal stability of

PCMs.43 Fig. 4 illustrated the TGA-DSC curves of the SA and SA/ SG composites. There was a single degradation process for the SA and SA/SG composites due to the decomposition of SA, and no decomposition was observed for the SA/SG composites within 200 °C. In this study, the thermal stability of the SA/SG composites was evaluated with respect to their degradation temperatures $(T_{5\%})$ at which the mass losses were higher than 5% (Fig. 4a). The SA/SG composites degraded at a high temperature of above 247 °C ($T_{5\%}$) and evaporated completely at around 500 °C, while for the SA, a low $T_{5\%}$ at 242.3 °C was found and evaporated completely at around 350 °C, indicting SA/SG composites had higher thermal stability than pure SA. That is to say, the thermal stability of the SA in the SA/SG composites was enhanced by the SG matrix. With the content of graphene in SG composites increasing, the $T_{5\%}$ of the SA/SG composites also rose and were 247.5 °C, 260.1 °C, and 265.1 °C for SA/SG₁, SA/ SG₅, and SA/SG₁₀, respectively. It indicated that the graphene had an advantage for thermal stability of the SA/SG composite. Compared with the $T_{5\%}$ of pure SA, the $T_{5\%}$ of the SA/SG₁, SA/ SG₅, and SA/SG₁₀ were about 5.2 °C, 17.8 °C, and 22.8 °C higher than that of SA, respectively. Moreover, the degradation temperature of the SA/SG composites were much higher than their phase change temperatures. It can be concluded that the SA/SG composites had good thermal stability in their operating temperature range or above their melting points. From the corresponding DSC curves (Fig. 4b), a endothermic peak at around 321 °C in the DSC curve of pure SA indicated the SA decomposed. The SA/SG₁ and SA/SG₅ had same tendency in the DSC curves. But, the DSC curve of SA/SG₁₀ was found to be clearly different from that of the SA/SG1 and SA/SG5: a broad endothermic peak at around 150 °C appeared in the curve while no evident peak in that of the SA/SG1 and SA/SG5 at this temperature interval. This may be related to the interaction between the SA and the SG matrix. Moreover, the exact content of the SA in the SA/SG composites had been confirmed by the TGA. The content of SA in the SA/SG₁, SA/SG₅, and SA/SG₁₀ were 34.3 wt%, 34.3 wt%, and 34.8 wt%, respectively, which approached the designed value (35 wt%).

3.6. Thermal energy storage behaviors and properties of the SA/SG composites

The thermal energy storage behaviors and properties of the SA and SA/SG composites were determined by DSC thermal analysis (Fig. 5a). The onset temperature of each DSC peak was obtained from the intersection of a line at the point of maximum slope of the leading edge of the DSC peak and the extrapolated baseline on the same side of the peak.⁴⁴ The detailed thermal energy storage properties of the SA and SA/SG composites were given in Table 1. The curve of pure SA (inset of Fig. 5a) showed a melting temperature ($T_{\rm m}$) at 54.11 °C in the endothermic curve and a freezing temperature ($T_{\rm f}$) at 53.22 °C in the exothermic curve. The latent heats of melting ($\Delta H_{\rm m}$) and freezing ($\Delta H_{\rm f}$) of pure SA were 177.3 J g⁻¹ and 173.8 J g⁻¹, respectively.

The thermal energy storage behaviors are a crucial factor for composite PCMs. Compared with pure SA and the SA/SG **RSC Advances**

Fig. 3 SEM images and EDS patterns of the (a) SG_5 and (b) SA/SG_5 ; magnified SEM images of (c) smaller particle of the SA/SG_5 and (d) larger particle of the SA/SG_5 .

composites, it was surprising to find that more than one peak in the endothermic or exothermic curves of the composites (Fig. 5a). The thermal characteristics of the composites were very different from that of pure SA (inset of Fig. 5a), indicating a novel interaction between the SA and the SG matrix, 45 which was also reflected in the XRD, FTIR, and TG-DSC results. SA/SG₁ had three phase change processes: the phase changes were at 47.15 °C, 52.16 °C, and 54.03 °C in endothermal curve, respectively; the phase changes were 48.18 °C, 52.51 °C, and 54.92 °C in the exothermal curve, respectively. There was a similar phenomenon in the DSC curve of the SA/SG₅. But the comparison between the SA/SG₁ and the SA/SG₅ indicated a tiny difference: the phase change temperature at 47.15 °C, 52.16 °C, and 54.03 °C in endothermal curve shifted to lower temperatures of 45.55 °C, 51.53 °C, and 47.91 °C, respectively (Table 1). Nevertheless, SA/SG₁₀ had two phase change processes: the phase changes were at 39.03 °C and 52.58 °C in endothermal curve, respectively; the phase changes were 39.96 °C and 50.28 °C in the exothermal curve, respectively. A large separation of about 10 °C emerged in the DSC curve of SA/SG₁₀ and the phase change peaks became more defined. For pure SG, there was no endothermal or exothermal peak in the DSC curves (Fig. S6†) in the same temperature range, indicating that SG had no contribution to the phase change peak of the SA/SG

composites. From what is discussed above, the thermal energy storage behaviors of the SA were changed by impregnating it into the SG matrix. As further increasing the adding ratio of the graphene in the SG composites (SG_1 , SG_5 , and SG_{10}), the thermal energy storage behaviors of the SA/SG composites (SA/SG_1 , SA/SG_5 , and SA/SG_{10}) were dramatically changed.

The latent heats of melting and freezing were $51.35\,\mathrm{J}\,\mathrm{g}^{-1}$ and $49.83\,\mathrm{J}\,\mathrm{g}^{-1}$ for SA/SG₁, $51.32\,\mathrm{J}\,\mathrm{g}^{-1}$ and $49.18\,\mathrm{J}\,\mathrm{g}^{-1}$ for SA/SG₅, and $37.14\,\mathrm{J}\,\mathrm{g}^{-1}$ and $33.87\,\mathrm{J}\,\mathrm{g}^{-1}$ for SA/SG₁₀, respectively, which were less than their corresponding theoretical values (Table 1). The decrease in the latent heats of the composites was not only attributable to the lower fraction of SA within the composites, but could also be related to the crystallinity of the SA in the composites. Crystallization of the SA in the composites may be inhibited by interactions between the SA and the supports, which would also reduce the latent heats of the composites. The crystallinity of the SA (F_c) in the composite was calculated by: $^{46-49}$

$$F_{\rm c} = \frac{\Delta H_{\rm composite}}{\Delta H_{\rm pure} \beta} \times 100\% \tag{1}$$

where $\Delta H_{\rm composite}$ and $\Delta H_{\rm pure}$ were the latent heats of the composites and pure SA, respectively, and β represented the loading of SA in the composites. The crystallinity of the SA in the

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a 100 80 60 Mass (%) 40 SA/SG SA/SG 20 SA/SG₁₀ 0 100 300 400 500 600 Temperature (°C)

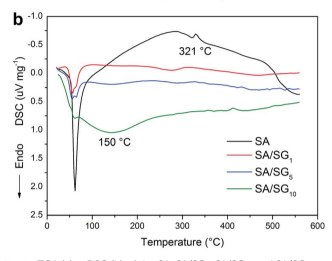
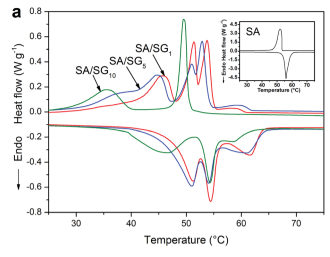


Fig. 4 $\,$ TGA (a) - DSC (b) of the SA, SA/SG1, SA/SG5, and SA/SG10.

SA/SG₁ (84.44%) was close to that in the SA/SG₅ (84.39%), and they were clearly higher than that in the SA/SG₁₀ (60.19%). According to our previous research, ^{13,49} it meant that more confined SA (disordered) in the SA/SG₁₀ composite, comparing with more free SA (ordered) in the SA/SG₁ and SA/SG₅ composites. That is to say, with increasing the adding ratio of the graphene in the SG composites, more SA were confined by the SG. The confined SA cannot crystallize and will not work for thermal energy storage. ⁴⁹ The effective energy stored per unit mass of the SA ($E_{\rm ef}$) was used to evaluate the effectiveness of the SA in different composites (Table 1). The SA in the SA/SG₁ and in the SA/SG₅ had a adjacent $E_{\rm ef}$ (~150 J g⁻¹) which higher than that of the SA/SG₁₀ (106.7 J g⁻¹).

Furthermore, the thermal energy storage properties of the SA/SG composites with different contents of SA were investigated by increasing SA dosage (designed as 40% and 45%) for the SA/SG $_5$ composite. Fig. 5b demonstrated the DSC curves of the SA $^+$ /SG $_5$ and SA $^{++}$ /SG $_5$, and the content of SA in the SA $^+$ /SG $_5$ and SA $^{++}$ /SG $_5$ were determined as 39.5% and 44.8% by TGA (inset of Fig. 5b). The detailed thermal energy storage properties of the SA $^+$ /SG $_5$ and SA $^{++}$ /SG $_5$ were given in Table S1.† Increasing the content of SA to 39.5%, the DSC curve of the SA $^+$ /SG $_5$ reduced to two phase change processes (Fig. 5b). Compared to



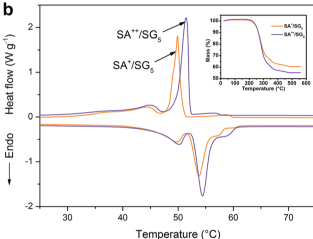


Fig. 5 (a) DSC curves of the (a) SA, (b) SA/SG $_1$, (c) SA/SG $_5$, and (d) SA/SG $_{10}$; (b) DSC curves and TG (inset) of the SA $^+$ /SG $_5$ and SA $^{++}$ /SG $_5$.

the SA/SG₅, the phase change peaks of SA⁺/SG₅ at high temperatures became a single peak and enhancive. As further increasing the content of SA to 44.8%, the phase change peaks at low temperatures shifted to higher temperature zone. Meanwhile, the phase change peaks at high temperatures became more enhancive, which manifested that the incremental SA in the SA⁺⁺/SG₅ had mainly contributed to the phase change enthalpy occurring at the high temperatures. With increasing the content of SA, the crystallinities of SA in composites were enhanced to 86.25% and 88.44% for SA⁺/SG₅ and SA⁺⁺/SG₅, respectively.

The phenomenon: (a) more than one endothermal or exothermal peak in the DSC curves of SA/SG composites; (b) the $E_{\rm ef}$ decreased while increasing the adding ratio of graphene in the SA/SG₁₀ indicated an interesting question between the SA and the SG matrix. This interesting question may be connected to the special pore structure of the SG matrix on the basis of our previous research.^{13,49} To unlock this interesting question and further clarify this hypothesis, the pore structure of the SG matrix were depicted in Fig. 6, and the specific surface areas and porous properties of them were listed in Table 2. The specific

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Table 1 Thermal properties of the SA and SA/SG composites^a

Samples	Loadage (β, wt%)	Melting temperature $(T_{ m m}, {^{\circ}{ m C}})$	Freezing temperature $(T_{\rm f},^{\circ}{ m C})$	Latent heat of melting $(\Delta H_{\rm m}, \ \ \ \ \ \ \ \ \ \ \ \)$	Latent heat of freezing $(\Delta H_{\rm f}, \ {\rm J} \ {\rm g}^{-1})$	Theoretic values of ΔH_{m} $(\Delta H_{\mathrm{th}}, \mathrm{J} \ \mathrm{g}^{-1})$	Crystallinity of SA $(F_c, \%)$	Efficient energy per unit mass of SA $(E_{ef}, J g^{-1})$
SA	100	54.11	53.22	177.3	173.8	_	100	_
SA/SG ₁	34.3	47.15/52.16/54.03	48.18/52.51/54.92	51.35	49.83	60.81	84.44	149.7
SA/SG ₅	34.3	45.55/51.53/47.91	47.24/53.14/54.03	51.32	49.18	60.81	84.39	149.6
SA/SG ₁₀	34.8	39.03/52.58	39.96/50.28	37.14	33.87	61.70	60.19	106.7

^a Note: $\Delta H_{\rm th} = \Delta H_{\rm pure} \times \beta$; $E_{\rm ef} = \Delta H_{\rm pure} \times F_{\rm c}$.

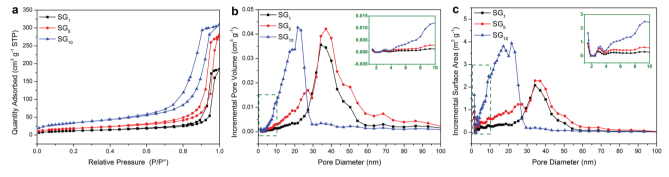


Fig. 6 (a) Nitrogen adsorption—desorption isotherms of the SG, and BJH pore size distribution for (b) incremental pore volume and (c) incre-

surface areas of the SG1, SG5, and SG10 were calculated to be $41.29 \text{ m}^2 \text{ g}^{-1}$, $65.29 \text{ m}^2 \text{ g}^{-1}$, and $120.27 \text{ m}^2 \text{ g}^{-1}$, respectively. The results also exhibited that the adsorption cumulative volume of pores and surface area of pores of SG₁₀ were 0.49 cm³ g⁻¹ and 136.38 m² g⁻¹, respectively, which larger than that of the SG₁ and SG₅. With increasing the adding ratio of graphene in the SG composites, the adsorption average pore widths of the SG decreased and were 26.16 nm, 24.00 nm, and 14.50 nm for SG₁, SG_5 , and SG_{10} , respectively. Compared with the pore size distributions in incremental pore volume and in incremental surface area of the SG₁ and SG₅, the SG₁₀ had significantly different pore size distributions while the SG₁ and SG₅ presented approximately status (Fig. 6b and c). The most pore volume and surface area of the SG₁₀ distributed in the size range of 5 to 30 nm, while that of the SG₁ and SG₅ appeared in larger size range (20–60 nm). Observing the pore diameters from 5 nm to 30 nm, the SG₅ had slightly more pore volume and surface area than that of the SG₁, resulting in a tiny difference in the DSC curves (Fig. 5a). After introduced to the supporting matrix, the SA molecules (long hydrocarbon chains) were adsorbed in

different size pore. Previous researches have been testified that the phase change temperatures of organic phase change materials decrease as the supporting spaces become narrower. 45,50,51 Thus, the SA molecules adsorbed in serial narrower pore exhibited different lower temperature of phase change just like SA/SG₁ and SA/SG₅ (Fig. 5a). With increasing the adding ratio of graphene in the SG composites, the more pore volume distributed in the small size, the more SA was supported in these pore-size ranges. It resulted in the phase change temperatures further shifted to lower temperature due to the rotation of the hydrocarbon chains require more low thermal energy in this condition. 50,51 Compared with the pore size distribution in incremental pore volume of the SG₁ and SG₅ (20-60 nm and 40 nm), the SG₁₀ had a smaller and concentrated size range (5-30 nm and 25 nm), generating a defined phase change peaks occurring 10 °C lower. Also, the interaction (including surface tension forces and capillary forces) occurred between the SA and the SG matrix would be strong enough to disturb its phase change process when the pore can confine the crystallization of the PCM. 49,52 Colligated the above discussions, the

Table 2 Specific surface areas and porous properties of the SG

Samples	BET surface area (cm ² g ⁻¹)	BJH adsorption cumulative volume of pores (cm ³ g ⁻¹)	BJH adsorption cumulative surface area of pores $(cm^2 g^{-1})$	BJH adsorption average pore diameter (nm)
SG ₁	41.29	0.29	44.53	26.16
SG_5 SG_{10}	65.29 120.27	0.43 0.49	72.40 136.38	24.00 14.50

different pore structure of the SG matrix could be a reason to more than one endothermal or exothermal peak in the DSC curves of the SA/SG composites. From the inset of Fig. 6b and c, the SG₁₀ had larger pore volume and surface area than that of the SG₁ and SG₅ in the size range of 1-10 nm, signifying more SA hydrocarbon chains were confined in the SA/SG₁₀. Moreover, the interaction between the SA and the SG matrix of capillary and surface tension forces could affect the crystallization of the SA,49 the larger capillary and surface tension forces between the SA and the SG₁₀ was contributed to the larger BET surface area of the SG₁₀ (Table 2). And then the interaction between the SA and the SG matrix in the SA/SG₁₀ is more than that in the SA/SG₁ and SA/SG₅. This is consistent with the observation from the thermal stability results (Fig. 4b): the SA/SG₁₀ need greater energy to out of the shackles of the surface of SG₁₀ and then a broad endothermic peak at around 150 °C appeared while no

Table 3 Thermal conductivities of the SA and SA/SG composites

Samples	SA	SA/SG_1	SA/SG_5	SA/SG_{10}	SA/SiO ₂
$\lambda (W m^{-1} K^{-1})$	0.26	0.90	1.05	1.12	0.58

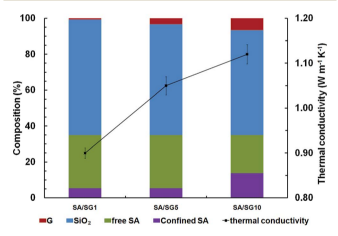


Fig. 7 The compositions and thermal conductivities of the SA/SG composites.

evident peak in that of the SA/SG_1 and SA/SG_5 at this temperature interval. These then dramatically decreased the crystallinity of the SA in the SA/SG_{10} composite, comparing with that in the SA/SG_1 and SA/SG_5 composites. As mentioned above, the confined SA cannot crystallize and will not work for thermal energy storage. So, the $E_{\rm ef}$ decreased while increasing the adding ratio of graphene in the SA/SG_{10} .

3.7. Thermal conductivities of the SA/SG composites

The thermal conductivity values were 0.26, 0.90, 1.05, and 1.12 W m⁻¹ K⁻¹ for the SA, SA/SG₁, SA/SG₅, and SA/SG₁₀, respectively (Table 3). For comparison, the silica without decorated by graphene was prepared and used to support the SA by the same process described in experimental, labeling as SA/SiO₂ with a thermal conductivity of 0.58 W m⁻¹ K⁻¹. The comparison between the thermal conductivities of pure SA and the SA/SiO2 demonstrated that SiO₂ substrate could improve the thermal conductivity of pure SA by 123%. In order to further enhance the thermal conductivities of the composites, the SiO₂ substrate was decorated by graphene. The enhancements in the thermal conductivity compared with that of SA/SiO2 were 55%, 81%, and 93% for the SA/SG₁, SA/SG₅, and SA/SG₁₀, respectively. The enhancements in the thermal conductivity compared with that of SA were 246%, 304%, and 331% for the SA/SG₁, SA/SG₅, and SA/SG₁₀, respectively. From the SA/SG₁ to the SA/SG₅, the thermal conductivity of composites was enhanced by 17%; from the SA/SG₅ to the SA/SG₁₀, the thermal conductivity of composites was only enhanced by 7%. These illustrated that with increasing the graphene ratio, thermal conductivity enhancing ability of graphene decreased, and the reason was explained in the following paragraph.

Thermal conductivities of SA/SG composites were governed by phonon propagation, because the composites were blends of the nonmetal (SG) and the organic materials (SA) (Fig. 1).⁵³ For a nonmetallic composite, the ordered or disordered crystals, grain boundary, and interfaces significantly affected the phonon transport.^{54–57} In particular, the disorder in crystals usually interferes with heat conduction in most materials; thus disordered component tend to have lower thermal conductivity because of strong phonon scattering.⁵⁸ The components of the

Table 4 Comparison of the thermal properties of the as-synthesized composite with the reported materials

Samples	Melting temperature, $T_{\rm m}$ (°C)	Freezing temperature, $T_{\rm f}$ (°C)	Latent heat of melting, $\Delta H_{\rm m}$ (J g ⁻¹)	Latent heat of freezing, $\Delta H_{\rm f}$ (J g ⁻¹)	Thermal conductivity $(W m^{-1} K^{-1})$	Reference
Poly(ethylene glycol) octadecyl ether (60 wt%)/ porous silica	33.1	35.0	49.7	35.9	_	59
<i>N</i> -octadecane (51.2 wt%)/ silica shell	27.14	23.53/16.72	108.8	98.61	0.4403	60
<i>N</i> -tetradecane (37.21 wt%)/ polystyrene-silica	2.13	0.39	83.38	79.37	0.4035	61
N-octadecane (36 wt%)/ silica wall	28.43	23.73/20.55	40.35	37.96	0.874	62
SA (34.3 wt%)/SG ₅	45.55/51.53/47.91	47.24/53.14/54.03	51.32	49.18	1.05	This work

SA/SG composites were exhibited in Fig. 7. All the SA/SG composites had a almost same SA loadage (~35 wt%), but showed a different proportion of the confined SA. The SA/SG. This

composites had a almost same SA loadage (\sim 35 wt%), but showed a different proportion of the confined SA. The SA/SG₅ had a very similar proportion of the confined SA to that of the SA/SG₁, and the SA in the SA/SG₁ and in the SA/SG₅ had a comparative thermal conductivity. The thermal conductivity of the SA/SG₅ was obviously (17%) enhanced with increasing the content of graphene (G, 0.65% to 3.25%). But with increasing the content of G, the confined SA in the SA/SG₁₀ reached 13.93%, there are more disordered SA in the composites. The SA in the SA/SG₁₀ had a lower thermal conductivity due to stronger phonon scattering. It deservedly affected the thermal conductivity of the SA/SG₁₀ were smaller (7%) enhanced with increasing the content of G (3.25% to 6.50%). Thus, thermal conductivity enhancing ability of G in the SA/SG₁₀ decreased.

In addition, Table 4 showed the comparison of the latent heats and the thermal conductivities of the prepared composite PCMs with that of silica-based composite PCMs in the literature, $^{59-62}$ the SA/SG $_5$ composite showed some obvious advantages over the reported materials: added phase change temperatures, considerable latent heat capacities with less PCMs dosage, and higher thermal conductivity (more than 1.0 W m $^{-1}$ K $^{-1}$), which broadens its wide application for thermal energy storage. It is remarkably noted that the prepared SA/SG $_5$ composite has potential for application in the thermal energy storage, especially in the thermal gradient using.

4. Conclusions

RSC Advances

Novel thermal energy storage materials (SA/SG₁, SA/SG₅, and SA/ SG₁₀) were synthesized from graphene-decorated silica (SG) and stearic acid (SA) by vacuum impregnation method. The structure and thermal energy storage performances of the SA/SG composites were in detail investigated by XRD, FTIR, SEM-EDS, TG-DSC, DSC, BET, and thermal conductivity. The SA/ SG₁, SA/SG₅, and SA/SG₁₀ composites have good thermal stability and the thermal degradation temperature increased by 5.2 °C, 17.8 °C, and 22.8 °C against pure SA. It is of interest that the thermal energy storage behaviors of the SA/SG composites were dramatically changed along with different contents of graphene, presenting more than one endothermal or exothermal peaks in the DSC curves while pure SA only one of that. The SA in the SA/SG₁ and in the SA/SG₅ showed greater effective energy storage per unit mass ($E_{\rm ef}$, ~ 150 J g⁻¹) than that of SA in the SA/SG₁₀. These thermal energy storage behaviors and properties were illuminated to be related to the special pore structure of SG matrix by nitrogen gas adsorption-desorption isotherms. Decorating silica by graphene was beneficial to the enhancement in thermal conductivity of the SA/SG composite, and 55%, 81%, and 93% higher than that of SA/SiO₂ (without decorated by graphene), respectively; and 246%, 304%, and 331% higher than that of pure SA, for SA/SG₁, SA/SG₅, and SA/ SG₁₀, respectively. The SA/SG₅ has potential for application in thermal energy storage, especially in the thermal gradient using due to both the considerable $E_{\rm ef}$ and the high thermal conductivity of 1.05 W m⁻¹ K⁻¹.

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