Preparation and characterization of myristic acid/expanded graphite composite phase change materials for thermal energy storage

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Myristic acid/expanded graphite (MA/EG) composite phase-change material (CPCM) was prepared by absorbing liquid MA (as the PCM) into EG (as the supporting material). Its chemical structure, microstructure, and thermal properties were characterized and studied. In the MA/EG CPCM, the largest mass content of MA was 93.5% by using the diffusion–exudation circle method for the first time. Fourier transform infrared spectroscopy (FTIR) analysis indicated that the MA and EG were a pure physical mixture of which the structure does not change, and they undergo no chemical reaction. Differential-scanning-calorimetry (DSC) analysis results showed that the melting and freezing temperatures of the MA/EG CPCM were 53.3 and 52.4 °C, respectively, and the melting and freezing latent heats were 189.5 and 187.8 J/g, respectively. After several heat-cycle accelerations, the material still had good thermal-energy-storage effect. MA/EG CPCM thermoconductivity was greatly enhanced after adding EG, and the results of thermal-storage/-release experiments indicated that the thermal-storage and -release ratios of the MA/EG phase-change unit was greatly improved when compared with that of MA. These results indicated that the MA/EG CPCM was a suitable low-temperature thermal-energy-storage material.

Because of the wide applications of thermal storage technology in many fields, such as solar-energy utilization, the recycling and utilization of industrial exhaust heat, peak-load shifting, and building heating ventilation and air conditioning, it has attracted increasing attention1–4. Thermal storage is generally classified into three methods, latent-heat thermal storage, sensible heat-thermal storage, and chemical-energy storage5. Of the three, latent-heat thermal storage has advantages such as stable temperature output, a distinct energy-saving effect, high-density energy storage, and wide use in solar-energy systems, including air conditioning, building-energy conservation, aerospace, and power systems6–11.

The core technology of latent-heat thermal storage is phase change materials (PCMs). Multiple inorganic, organic, and composite PCMs have been studied for building-energy efficiency, such as paraffins2,13, polyhydric alcohols14,15, inorganic salts16, and fatty acids17. In these materials, fatty acids are organic PCM that have attracted more attention due to their large latent heat, nontoxicitiy, suitable transformation temperature, noncorrosiveness, low degree of supercooling, good thermal stability, and zero or minimal volume change. The transformation temperature of fatty acids that are commonly used as phase-change energy-storage material ranges from 30.1 to 70.7 °C, and its phase-change latent heat ranges from 149.1 to 222.8 J/g, which can be used for an energy-storage system of corresponding temperature20,21. However, the thermoconductivity of fatty acids is relatively low, and leakage of the liquid phase occurs in the solid–liquid phase-change process, which severely restricts its application in thermal-energy storage. The most widely used method for improving the thermal-conductivity coefficient of fatty acid PCMs is the addition of a high-thermal-conductivity medium to obtain composite PCMs with good thermoconductivity. Substances that are generally used as a high thermal conductivity medium mainly include metal particles, metal oxides, and carbon materials (such as expansion ink and carbon nanotubes)22,23. Liquid-phase leakage can be avoided by...

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encapsulating PCMs into stable composite PCMs in consideration of the inorganic porous medium as matrix. The final composite PCMs are prepared by absorbing fatty acid PCMs into their pore structure\(^5\). A porous matrix has a large specific surface area and multiple microporous structures in which the core material is distributed. The liquid-phase-change core material cannot easily overflow from the micropores due to the capillarity of micropores and the chemical bond force between the phase change core material and the matrix. Expanded graphite (EG) is widely used as a form-stable material because of its low density, high thermal conductivity, and multiple pores that can not only prevent liquid leakage, but also sharply enhance the thermal conductivity property of PCMs\(^25,26\).

Wang et al.\(^27\) prepared stearic acid/EG (SA/EG) composite PCMs with an SA mass content of 85%, and the corresponding phase-change temperature and latent heat were 128 \(^\circ\)C and 187 J/g, respectively. Wu et al.\(^28\) prepared EG/SA composite PCMs, and found that the best condition was when SA mass content was 75% with packed density of 900 kg/m\(^3\). Fang et al.\(^29\) prepared SA/EG form-stable PCMs with different mass rates, and found that, when EG content was 17%, the melting and freezing temperatures were 53.12 and 54.28 \(^\circ\)C, respectively. Moreover, the corresponding phase-change latent heats were 155.50 and 155.70 kJ/kg, and thermoconductivity increased almost 10 times compared to that of pure SA. SA in EG’s porous network was well-absorbed, and no SA leakage was found in the composite material, even in a molten state. Hu et al.\(^30\) prepared palmitic acid/EG (PA/EG) composite PCMs with an EG mass fraction of 1%, 3%, 5%, and 8%, and found that PA/EG composite PCMs preserve multiple pores of the EG material, the phase-change temperature of which is close to that of PA. Zhang et al.\(^31\) prepared CA–PA–SA/EG composite PCMs, of which EG quality content was 10%. Yang et al.\(^32\) prepared MA–PA–SA/EG composite PCMs, and found that PCM thermoconductivity was improved for the high thermoconductivity of the EG. Their research results proved that PCM thermoconductivity can be increased by adding EG.

MA, which melts at 53.6 \(^\circ\)C with a latent heat of 199.4 kJ/kg and freezes at 51.8 \(^\circ\)C with a latent heat of 199.0 kJ/kg, is beneficial organic PCM for thermal-energy storage. EG is a porous material with high thermoconductivity. However, studies of MA/EG composite PCMs for thermal-energy storage are rare. Cao et al.\(^33\) prepared MA/EG composite PCMs and confirmed that the best mass ratio of MA and EG was 15:1, and the transformation temperature and latent heat of MA/EG composite PCMs were 53.19 \(^\circ\)C and 191.75 J/g, relatively, but some thermal properties have not been studied, such as thermoconductivity, thermal stability, and thermal reliability. Very little research has focused on the thermal properties of MA/EG composite PCMs. Hence, in this paper, MA/EG composite PCMs were prepared by absorbing liquid MA into EG. Their chemical structure, microstructure, and thermal properties were characterized and studied. The prepared MA/EG composite PCMs can be used for solar-energy-storage systems, that can store solar heat by daylight and release heat at night or on rainy days. This composite material can also be used for heat storage of a building exterior; it absorbs heat from its surroundings or solar radiation and releases it at night to reduce the load in air-conditioning systems.

Experiments

Materials. MA (\(\geq 98\) purity) as thermal-energy-storage material was supplied by Shanghai Zhunyun Chemical Co., Ltd., Shanghai, China. The grain fineness number 350 meshes of the expandable graphite (Carbon content > 99, expansion coefficient 100 mL/g, 350 meshes) was purchased from Qingdao Hengrunda Graphite Products Co., Ltd., Qingdao, China.

MA/EG composite CPCM preparation. Expanded graphite was obtained by heating the dried expandable graphite in a porcelain crucible for 40 s in a 900 \(^\circ\)C muffle furnace. A quantity of prepared expanded graphite and the MA were fully mixed in a beaker and then heated in boiling water that was constantly stirred until the MA melted. The melted MA was absorbed by EG. Then, the beaker was placed in a 100 \(^\circ\)C vacuum-drying oven for dehydration. The mixture was stirred every 2 h until it cooled down to the indoor temperature to ensure the uniform mixing and absorption of MA and EG. Finally, MA/EG composite PCMs were obtained.

MA/EG CPCM characterization. The transformation temperature (melting and freezing temperatures) and the phase-change latent heat (melting and freezing latent heat of the MA and the MA/EG eutectic mixtures before and after thermal cycling) were determined by differential scanning calorimetry (DSC; DSC instrument, STA2500, Germany) adjusted with inheatdium standard in a temperature range between 0 and 100 \(^\circ\)C. The growth ratio of DSC temperature measurement was 5 \(^\circ\)C/min, and the samples were cooled by liquid nitrogen. Same-sample DSC measurements were performed three times. The accuracy of phase-change temperature was \(\pm 0.1\)% and \(\pm 4\)% for latent heats. The extrapolated onset temperatures on the DSC curves were the transformation temperature of the PCMs.

The effect of the thermal-cycle-index on thermal properties was investigated by heating the MA/EG eutectic mixtures from a solid to a liquid state, and then cooling from a liquid to a solid state using a heat controller. When the numbers were 50, 100, and 200, the thermal cycling process was stopped. The property evolution of the MA/EG mixtures was surveyed by DSC, thermogravimetric analysis (TGA), and Fourier transform infrared spectroscopy (FTIR) techniques.

The thermal stability of the MA/EG was analyzed with TGA technique (TGA instrument, STA2500, Germany) with accuracy of \(\pm 0.2\)%.

The morphology and microstructure of the MA/EG composites were observed using a scanning electron microscope (SEM, S-4800, Hitachi Inc., Japan). Structural analysis of the MA/EG composites was conducted via FTIR (FTIR instrument, Thermo Scientific Nicolet iS5, USA). PCMs thermoconductivity was measured using a thermoconductivity analyzer (DRL-III, Xiangtan Xiangxi Instrument Co. LTD., China) with an accuracy of \(\pm 3\)%.
Results and discussion
Maximal absorption ratio of MA in MA/EG CPCMs. Generally, the higher the mass of fatty acids in composite PCMs is, the better the thermal-energy storage of composite materials is. In the MA/EG composite PCM, EG was the carrier material in which MA was absorbed. When phase change occurs, MA changes from a solid state to a liquid state. However, due to the micropore capillarity and chemical-bond force between the phase-change core material and the matrix, MA will not leak out of the composite PCM, and the composite PCM remains in its original form, i.e. a solid state. However, beyond the maximal absorption of the carrier materials, redundant fatty acids may attach to the surface of the composite PCMs, which can melt and lead to leakage during the phase-change process. EG content slightly affects the phase-transition temperature. Hence, the largest mass ratio of aliphatic acids in composite PCMs must be confirmed.

The diffusion–exudation circle method was applied to confirm the large mass rate of aliphatic acids in composite PCMs. The test diagram is shown in Fig. 1, and the test flow was as follows34. Composite PCMs samples were prepared with different MA contents (from 92 to 95%), and then equably dispersed in the test area with a diameter of 30 mm on the filter-paper center (as shown in Fig. 1a). Afterward, the specimens were moved to the vacuum drying-oven, heated at homeothermy of 70 °C for 30 min, and removed. The leakage degree of the phase change of the working substance component outside the self-tempering unit test area was observed (as shown in Fig. 1b). The largest and smallest diameters of the exudation circles that formed outside the exudation test areas were measured, and the average was calculated. Table 1 shows a series of regulated leakage percentages as the seepage-stability-assessment standard. The percentage of the excess of the average exudation-circle diameter to the test-area diameter was calculated, and the values were compared with the standard ones (Table 1) to assess stability. All values not higher than the standard indicated that stability reached the corresponding standards.

Figure 2 shows all photos of the prepared composite PCMs before (a) and after (b) testing. Leakage was evident in the sample with a 94% MA mass content, whereas no exudation was found in that with 93% MA mass content. Hence, the largest MA mass content was approximately 93–94%. Therefore, we prepared multiple samples with such mass content and applied the diffusion–exudation circle method to assess seepage stability; results are shown in Table 2. The largest MA mass content in the composite PCMs was approximately 93.5%. The MA/EG composite materials adopted all contents of MA at 93.5%.

FTIR analysis of MA/EG CPCMs. The consistency and structural stability of the MA mixture with EG were investigated by conducting an FTIR test on pure MA and MA/EG composite materials; results are shown in Fig. 3. The infrared spectral curve showed a stretching vibration crest value at 1748 cm⁻¹, of which the strong crest value was C=O on fatty acid carboxyl. Peaks occurred at 2945 and 2877 cm⁻¹ due to the stretching vibration of the C–H bond in −CH₃ and −CH₂−35. The spectra of MA and MA/EG were the same, and the positions of the

| Item                              | Leakage percentage Φ (%) | Stability         |
|----------------------------------|--------------------------|-------------------|
| Considered no exudation          | Φ ≤ 10                   | Very stable       |
| Microscale exudition             | 10 < Φ ≤ 15              | Stable            |
| Slight exudation                 | 15 < Φ ≤ 30              | Basically stable  |
| Medium amount of exudation       | 30 < Φ ≤ 50              | Basically stable  |
| Large amount of exudation        | Φ > 50                   | Extremely unstable|

Table 1. Seepage-stability-assessment standard.
characteristic peaks corresponded to one, indicating that the combination of MA and EG was a pure physical mixture. The structure did not change, indicating that no chemical reaction occurred.

**MA/EG CPCMs microstructure.** Figure 4 shows the microstructure of the EG and MA/EG composite materials, observed using SEM. The SEM picture of EG (Fig. 4a) shows that the EG pores were mainly at the
microlevel, the wormlike porous structure caused EG to have a relatively large specific surface area and surface activity, and the molten MA was easier to absorb. In Fig. 4b, the SEM picture of MA/EG shows that MA was evenly dispersed in the porous structures of EG, in which the capillary tubes and surface tension could prevent the leakage of molten MA. On a macro level, the appearance of the MA/EG CPCM did not change during the phase change, indicating a certain form-stable effect. These observations revealed that, when the composite material was at the phase change, the molten MA in the pore network structure of EG had relatively good adhesive quality and compatibility, and MA was always equably distributed in EG without any leakage.

**MA/EG CPCM thermal properties.** The DSC curve of the MA/EG composite PCMs is shown in Fig. 5, and the thermal properties are shown in Table 3. As shown in Table 3, the phase-change characteristics of the MA/EG composites were very near those of the MA.

![Microscope images of EG and MA/EG](image)

**Figure 4.** SEM pictures of EG (a) and MA/EG (b) CPCMs.

![DSC curves](image)

**Figure 5.** DSC curves of MA and MA/EG CPCMs.

| PCM    | Melting | | | Freezing | | |
|--------|---------|---------|---------|----------|---------|---------|
|        | Onset temperature (°C) | Peak temperature (°C) | Latent heat (kJ/kg) | Onset temperature (°C) | Peak temperature (°C) | Latent heat (kJ/kg) |
| MA     | 53.6    | 59.5    | 199.4   | 51.8     | 46.3     | 199.0   |
| MA/EG  | 53.3    | 58.0    | 189.5   | 52.4     | 48.1     | 187.8   |

**Table 3.** DSC data of MA and MA/EG CPCMs.
Table 3 shows that the melting-point temperature of MA/EG was 53.3 °C, and the freezing point temperature was 52.4 °C. After a comparison, the melting temperature of the composite material was found to be slightly lower than that of MA, and freezing temperature slightly increased. However, the difference was small because MA was evenly dispersed in the mesh pore structure of EG, and EG has relatively high thermoconductivity. The high thermal-conductivity coefficient of the support material accelerated the heat-transfer rate of the PCMs from the external to the internal36, thereby resulting in lower phase-transition temperature.

Table 3 shows that the melting and freezing latent heat of MA/EG were 189.5 and 187.8 kJ/kg, respectively. Phase-change latent heat could also be calculated using Formula (1)29. Results are shown in Table 4. The difference between the experiment result and the calculation was less than 2%

\[
\Delta H_{PCM} = \eta \Delta H_{MA}
\]

where \(\Delta H_{PCM}\) is the calculated latent heat of the composite PCMs, \(\eta\) is the mass percentage of MA in the composite PCM, and \(\Delta H_{MA}\) is the latent heat of the MA as measured by the DSC.

The phase-change latent heat of the MA/EG was relatively lower than that of MA because in the MA/EG composite PCMs, EG only supported the internal MA, but did not influence the melt crystallization properties of the PCMs. Hence, the phase-change latent heat of the MA/EG CPCMs in the mass unit dwindled. The melting latent thermal energy of the MA/EG composite PCMs was reduced by 4.96%, and its freezing latent heat was reduced by 5.63%, which still had relatively high phase-change latent heat.

### Thermoconductivity measurement and thermal-storage/release evaluation of MA/EG CPCMs.

MA belongs to organic PCMs with low thermoconductivity, and EG has relatively high thermoconductivity. Hence, the thermoconductivity of MA/EG composite material was greatly enhanced after adding EG. In this study, the MA/EG composite PCMs cylindrical blocks compressed with a smooth surface were formed by dry-pressing. Thermoconductivity-test data of the MA/EG composite PCM cylindrical samples with different densities are shown in Fig. 6. Results showed that the thermoconductivity of the samples increased with the increase of density. When sample density was 610.7, 710.5, 796.6, 864.2, and 1007.5 kg/m³, thermoconductivity was 1.495, 1.720, 1.847, 1.957, and 2.095 W/m K, respectively. Formula (2) was fitted from the data in Fig. 6 that showed a nearly linear relationship between the thermoconductivity (y) and packing density (x) of the MA/EG composite PCMs. The relation between thermoconductivity and packing density with different composite PCMs after adding EG can be found in relevant references37,38. This was probably caused by the extension of the contact surface area for the composite particles, and the reduction of the void space within the CPCM39. The higher the packing density of the CPCM was, the smaller the porosity in EG, the closer the contact among particles, the faster the heat-transfer speed, and the higher the thermal conductivity of the CPCMs were

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y = 0.0015x + 0.6281 \quad (R^2 = 0.9683)
\]
EG addition enhanced the thermoconductivity of the composite PCMs. This finding could be proven by comparing the thermal-storage/-release characterization of MA and MA/EG composite PCMs. The test installation is shown in Fig. 7. The thermal-storage/-release rate could be assessed by the time when the temperature of the PCMs center reached the setting temperature during the phase-change processes. The temperature of the storage process was set 80 °C, and that of the release process was set at 10 °C. Results are shown in Fig. 8. Figure 8a shows the melting-temperature curves of the PCMs. The two curves indicate that the time required for heating MA/EG and MA from 10 to 80 °C was 58 and 118 min, respectively. The required time to complete the heat storage of MA/EG was 50.8% shorter than that of MA. Figure 8b shows the freezing-temperature curves of the PCMs, and indicating that the required time for cooling MA/EG and MA from 80 to 10 °C was 62 and 123 min, respectively.

Figure 7. Experiment diagram for thermal-storage/-release measurement.

Figure 8. Storage (a) and release (b) curves of MA and MA/EG CPCMs.
The required time to complete the heat release of MA/EG was 49.6% shorter than that of MA. Therefore, the thermal-storage/-release rate of the MA/EG composite PCMs was greatly improved when compared with that of MA because EG had a high thermal-conductivity coefficient. The thermal-conductivity coefficient of the composite PCMs was enhanced compared with that of MA, thereby accelerating the efficiency of heat transfer.

**MA/EG CPCM thermostability and reliability.** The thermostability of the MA/EG composite materials was assessed via TGA. The TGA curve is shown in Fig. 9. In the test, the MA/EG composite material was heated from indoor temperature to 80 °C and preserved for 60 min under such temperature with a constant weight-loss rate. Figure 9 shows that MA began losing weight from a temperature of approximately 120 °C. Quality sharply decreased with increasing temperature. The weight-loss percentage reached the largest value at 210.83 °C, and was almost completely volatilized at a temperature of approximately 244 °C. This analysis indicated that, in a working environment of less than 120 °C, even if the melting temperature of MA were surpassed, the MA/EG composite material would not lose MA. Hence, the MA/EG composite material has good heat stability in cryogenic applications of less than 100 °C.

The thermal-cycle reliability of PCMs means the attenuation of heat-storage performance after repeating heat-storage/-release processes. This parameter is important in evaluating the useful life of PCMs. Thermal-cycle-acceleration experiment usually test for the thermal-cycle reliability of PCMs, which majorly concerns two important thermodynamic variables, namely, phase-transition temperature before and after a thermal cycle and phase-change latent heat. The DSC curves and heat performance, such as melting and freezing temperatures, and the latent heat of melting freezing of MA/EG after 50, 100, and 200 thermal cycles, are presented in Fig. 10 and Table 5. Figure 10 shows that the DSC curves of the MA/EG composite material before and after a thermal cycle were very close to each other. Table 5 shows that after 50, 100, and 200 times thermal cycles, the phase-change temperatures changed to 0.3, 0.1, and 0.5 °C, and the phase-change latent heat changed to 0.1%, 0.9% and 4.5%. This outcome indicated that the MA/EG PCMs have good thermal-cycle reliability.

**Conclusion**

In consideration of MA as PCM and EG as the carrier, and on the basis of the melt-adsorption method, an MA/EG composite phase-change energy-storage material was prepared with the largest MA quality content of 93.5%. FTIR test results indicated that no chemical reaction occurred before and after mixing MA and EG; EG only played the role of absorption, and a physical mixture was obtained. SEM characterization showed that MA could equally disperse in the EG network structure, and no seepage of liquid MA was found. DSC tests showed that the melting phase-change temperature and phase-change latent thermal energy of MA/EG PCMs were 53.3 °C and 189.5 J/g, and those of freezing were 52.4 °C and 187.8 J/g. The change of thermal properties was relatively smaller than that of pure MA. The TGA test proved that this phase-change energy-storage material has excellent stability within its operating temperature range. The thermal-cycle acceleration test showed that after 50, 100, and 200 cycles, the changes in the phase-change temperature and latent thermal energy of the ME/EG composite material could be neglected, and it has good reliability for long-term operation. The thermoconductivity of MA/EG CPCMs was greatly enhanced after adding EG, as verified by the thermal-storage/-release experiment. On the basis of this excellent performance, MA/EG composite PCMs can be widely used in low-temperature thermal-energy-storage systems, such as air-conditioning system, solar-energy storage, construction energy saving, and surplus-heat utilization.
Figure 10. DSC curves of MA/EG CPCMs before and after thermal cycles.
Table 5. Thermal performance of MA/EG PCMs before and after thermal cycles.

| Number of thermal cycling | Melting Temperature (°C) | Melting Latent heat (kJ/kg) | Freezing Temperature (°C) | Freezing Latent heat (kJ/kg) | Extent of supercooling (°C) |
|---------------------------|---------------------------|-----------------------------|---------------------------|-----------------------------|-----------------------------|
| 0                         | 53.3                      | 189.5                       | 52.4                      | 187.8                       | 0.9                         |
| 50                        | 53.0                      | 189.3                       | 52.6                      | 188.0                       | 0.4                         |
| 100                       | 53.2                      | 187.8                       | 52.4                      | 185.0                       | 0.8                         |
| 200                       | 52.8                      | 180.9                       | 52.2                      | 179.9                       | 0.6                         |

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References
1. Souayfane, F., Fardoun, F. & Biwole, P.-H. Phase change materials (PCM) for cooling applications in buildings: A review. *Energy Build.* 129, 396–431. https://doi.org/10.1016/j.enbuild.2016.04.006 (2016).
2. Jamekhorshid, A., Sadrameli, S. & Farid, M. A review of microencapsulation methods of phase change materials (PCMs) as a thermal energy storage (TES) medium. *Renew. Sustain. Energy Rev.* 31, 531–542. https://doi.org/10.1016/j.rser.2013.12.033 (2014).
3. Li, G., Hwang, Y. & Radermacher, R. Review of cold storage materials for air conditioning application. *Int. J. Refrigerat.* 35, 2053–2077. https://doi.org/10.1016/j.ijrefrig.2012.06.003 (2012).
4. Jose, P. D. C. & Eames, P. Thermal energy storage for low and medium temperature applications using phase change materials—A review. *Appl. Energy* 177, 227–238. https://doi.org/10.1016/j.apenergy.2016.05.097 (2016).
5. Li, T. X., Lee, I., Wang, R. Z. & Kang, Y. T. Enhancement of heat transfer for thermal energy storage application using stearic acid nanocomposite with multiwall carbon nanotubes. *Energy* 55, 752–761. https://doi.org/10.1016/j.energy.2013.04.010 (2013).
6. Alva, G., Liu, L.-K., Huang, X. & Fang, G.Y. Thermal energy storage materials and systems for solar energy applications. *Renew. Sustain. Energy Rev.* 68, 693–706. https://doi.org/10.1016/j.rser.2016.10.021 (2017).
7. Pandey, A. K. et al. Novel approaches and recent developments on potential applications of phase change materials in solar energy. *Renew. Sustain. Energy Rev.* 82, 281–323. https://doi.org/10.1016/j.rser.2017.09.043 (2018).
8. Zeinelabdein, R., Omer, S. & Gan, G. Critical review of latent heat storage systems for free cooling in buildings. *Renew. Sustain. Energy Rev.* 82, 2843–2868. https://doi.org/10.1016/j.rser.2017.10.031 (2018).
9. Tang, F., Su, D., Tang, Y. & Fang, G. Synthesis and thermal properties of fatty acid eutectics and diatomite composites as shape-stabilized phase change materials: A review. *Appl. Energy* 205, 1–12. https://doi.org/10.1016/j.apenergy.2018.01.009 (2018).
10. Zhou, Z. et al. Research on the application of phase-change heat storage in centralized solar hot water system. *J. Clean. Prod.* 198, 1262–1275. https://doi.org/10.1016/j.jclepro.2018.06.281 (2018).
11. Lin, W., Ren, H., Ma, Z. & Yang, L. Using fuzzy clustering and weighted cumulative probability distribution techniques for optimal design of phase change material thermal energy storage. *J. Clean. Prod.* 233, 1259–1268. https://doi.org/10.1016/j.jclepro.2019.05.040 (2019).
12. Jiang, N. & Pulskamp, H. Investigating thermal properties of using nano-tubular zn powder in paraffin as phase change material compound for thermal energy storage. *Compos. Part B Eng.* 126, 68–93. https://doi.org/10.1016/j.compositesb.2017.06.006 (2017).
13. Zhang, Z. et al. Preparation and thermal energy storage properties of paraffin/expanded graphite composite phase change material. *Appl. Energy* 91, 426–431. https://doi.org/10.1016/j.apenergy.2011.10.014 (2012).
14. Alva, G., Liu, L.-K., Huang, X. & Fang, G.Y. Thermal energy storage for low and medium temperature applications using phase change materials—A review. *Appl. Energy* 91, 426–431. https://doi.org/10.1016/j.apenergy.2011.10.014 (2012).
15. Song, X., Su, D., Tang, Y. & Fang, G. Preparation and performance of form-stable polyethylene glycol/silicon dioxide composites as solid-liquid phase change materials. *Appl. Energy* 86, 170–174. https://doi.org/10.1016/j.apenergy.2007.12.003 (2009).
16. Xu, X., Cui, H., Memon, S. A., Yang, H. & Tang, W. Development of novel composite pcm for thermal energy storage using CaCl2·6H2O with graphene oxide and SrCl2·6H2O nanocomposite with multiwalled carbon nanotubes. *Energy Build.* 60, 472–510. https://doi.org/10.1016/j.enbuild.2015.02.023 (2015).
17. Yuan, Y., Zhang, N., Tao, W., Cao, X. & He, Y. Fatty acids as phase change materials: A review. *Renew. Sustain. Energy Rev.* 29, 482–498. https://doi.org/10.1016/j.rser.2013.08.107 (2017).
18. Akeiber, H. et al. A review on phase change material (PCM) for sustainable passive cooling in building envelopes. *Renew. Sustain. Energy Rev.* 40, 1470–1497. https://doi.org/10.1016/j.rser.2016.03.036 (2016).
19. Sun, N. & Xiao, Z. Synthesis and performance of phase change materials microcapsules with a polymer/Bn/Tio hybrid shell for thermal energy storage. *Energy Fuels* 31, 10186–10195. https://doi.org/10.1021/acs.energyfuels.5b01271 (2017).
20. Kalnese, S. E. & Jelle, B. P. Phase change materials and products for building applications: A state-of-the-art review and future research opportunities. *Energy Build.* 94, 150–176. https://doi.org/10.1016/j.enbuild.2015.02.023 (2015).
21. Zhang, N. et al. Latent heat thermal energy storage systems with solid–liquid phase change materials: A review. *Adv. Eng. Mater.* 20, 1700753. https://doi.org/10.1002/adem.201700753 (2018).
22. Wu, S., Yan, T., Kuai, Z. & Pan, W. Thermal conductivity enhancement on phase change materials for thermal energy storage: A review. *Energy Storage Mater.* 17, 100215 (2019).
23. Zhang, P., Xiao, X. & Ma, Z. A review of the composite phase change materials: Fabrication, characterization, mathematical modeling and application to performance enhancement. *Appl. Energy* 165, 472–510. https://doi.org/10.1016/j.apenergy.2015.12.043 (2016).
24. Sari, A. & Karaiepekli, A. Preparation, thermal properties and thermal reliability of palmitic acid/expanded graphite composite as form-stable pcm for thermal energy storage. *Solar Energy Mater. Solar Cells* 93, 571–576. https://doi.org/10.1016/j.solmat.2012.08.057 (2009).
25. Tang, F., Su, D., Tang, Y. & Fang, G. Synthesis and thermal properties of fatty acid eutectics and diatomite composites as shape-stabilized phase change materials with enhanced thermal conductivity. *Solar Energy Mater. Solar Cells* 141, 218–224. https://doi.org/10.1016/j.solmat.2015.03.045 (2015).
26. Zhang, Z., Shi, G., Wang, S., Fang, X. & Liu, X. Thermal energy storage cement mortar containing n-octadecane/expanded graphite composite phase change material. *Renew. Energy* 50, 670–675. https://doi.org/10.1016/j.renene.2012.08.024 (2013).
27. Wang, S. et al. A novel sebacic acid/expanded graphite composite phase change material for solar thermal medium-temperature applications. *Solar Energy* 99, 283–290. https://doi.org/10.1016/j.solener.2013.11.018 (2014).
28. Wu, S., Li, T., Yan, T., Dai, Y. & Wang, R. High performance form-stable expanded graphite/stearic acid composite phase change material for modular thermal energy storage. Int. J. Heat Mass Transfer 102, 733–744. https://doi.org/10.1016/j.ijheatmasstransfer.2016.06.066 (2016).

29. Fang, G., Li, H., Chen, Z. & Liu, X. Preparation and characterization of stearic acid/expanded graphite composites as thermal energy storage materials. Energy 35, 4622–4626. https://doi.org/10.1016/j.energy.2010.09.046 (2010).

30. HU, P., YU, J.-J. & WANG, X.-W. Preparation and thermal performance of palmitic acid/expanded graphite composite phase change materials. J. Eng. Thermophys. 38, 464–469 (2017).

31. Zhang, H. et al. A capric-palmitic-stearic acid ternary eutectic mixture/expanded graphite composite phase change material for thermal energy storage. Compos. Part A Appl. Sci. Manuf. 87, 138–145. https://doi.org/10.1016/j.compositesa.2016.04.024 (2016).

32. Yang, X., Yuan, Y., Zhang, N., Cao, X. & Liu, C. Preparation and properties of myristic-palmitic-stearic acid/expanded graphite composites as phase change materials for energy storage. Solar Energy 99, 259–266. https://doi.org/10.1016/j.solener.2013.11.021 (2014).

33. Cao, X., Yuan, Y., Wang, X., Zhang, N. & Yang, X. Preparation and thermal properties of myristic acid/expanded graphite composite as phase change material. Acta Energ. Solar. Sin. 35, 1493–1498. https://doi.org/10.3969/j.issn.0254-0096.2014.08.029 (2014).

34. Sun, J. & Wu, Z. Study on evaluation method of phase change material leaky degree for building. New Build. Mater. 7, 43–46. https://doi.org/10.3969/j.issn.1001-702X.2004.07.015 (2004).

35. Fu, L., Dong, F., He, P. & Yang, Y. Study on preparation and properties of capric-myristic acid/diatomite form-stable phase change energy storage materials. J. Funct. Mater. 44, 1465–1468. https://doi.org/10.3969/j.issn.1001-9731.2013.10.021 (2013).

36. Wang, Y., Zheng, H., Feng, H. X. et al. Effect of preparation methods on the structure and thermal properties of stearic acid/activated montmorillonite phase change materials. Energy Build. 47, 467–473. https://doi.org/10.1016/j.enbuild.2011.12.021 (2012).

37. Zhang, Q., Wang, H., Ling, Z., Fang, X. & Zhang, Z. R1100/expand graphite composite phase change material with excellent structure stability, photo-thermal performance and good thermal reliability. Solar Energy Mater. Solar Cells 140, 158–166. https://doi.org/10.1016/j.solmat.2015.04.008 (2015).

38. Ling, Z. et al. Thermal conductivity of an organic phase change material composite phase change graphite composite across the phase change temperature range and a novel thermal conductivity model. Energy Conv. Manag. 102, 202–208. https://doi.org/10.1016/j.enconman.2014.11.040 (2015).

39. Huang, Z., Gao, X., Xu, T., Fang, Y. & Zhang, Z. Thermal property measurement and heat storage analysis of lin03/kcl-expanded graphite composite phase change material. Appl. Energy 115, 265–271. https://doi.org/10.1016/j.apenergy.2013.11.019 (2014).

40. Sarı, A. & Karaipekli, A. Fatty acid esters-based composite phase change materials for thermal energy storage in buildings. Appl. Thermal Eng. 37, 208–216. https://doi.org/10.1016/j.applthermaleng.2011.11.017 (2012).

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Author contributions
Z. D. and L. Y. conceived and designed the experiments. Z. D., Y. J. and Z. Y. performed the experiments and analyzed the results. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

Additional information

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