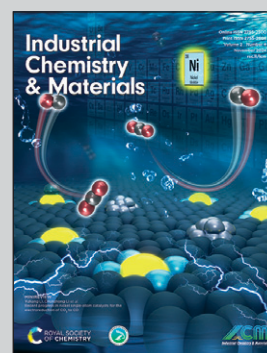


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Preparation of inorganic molten salt composite phase change materials and study on their electrothermal conversion properties

The EG conductive pathway enables CPCM to achieve direct electrical heating for energy storage and to regulate the temperature module uniformity through electric field control.

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Preparation of inorganic molten salt composite phase change materials and study on their electrothermal conversion properties†

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Due to their limitations in conductivity and shape stability, molten salt phase change materials have encountered obstacles to effectively integrating into electric heating conversion technologies, which are crucial in energy storage and conversion fields. In this study, we synthesized an inorganic molten salt composite phase change material (CPCM) with enhanced conductivity and shape stability using a gas-phase silica adsorption method. Our findings revealed the regularities in thermal properties modulation by expanded graphite (EG) within CPCM and delved into its characteristics of electric heating conversion. The study elucidated that a conductive network is essentially formed when the EG content exceeds 3 wt%. Following the fabrication of CPCM into electric heating conversion modules, we observed a correlation between the uniformity of module temperature and the quantity of EG, as well as the distribution of electrode resistance and external voltage magnitude. Building upon this observation, we proposed a strategy to adjust the module temperature field with an electric field. Comparing the proposed direct electrical heating energy storage method with traditional indirect electrical heating methods, the energy storage rate increases by 93.8%, with an improved temperature uniformity. This research offers valuable insights for the application of molten salt electric heating conversion CPCM.

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1 Introduction

In recent years, solar energy, wind energy, and other green renewable energy sources have emerged as the preferred options for clean power generation.¹ However, the intermittency of wind and solar energy has resulted in a serious issue of surplus electricity that cannot be effectively utilized.^{2,3} To address this challenge, one solution is to convert the electricity generated by intermittent renewable energy sources into thermal energy for storage, which can then be supplied to users when needed.^{4,5} This effectively resolves the uneven distribution of electrical energy over time and space, thereby enhancing

energy utilization. Phase change energy storage technology utilizes latent heat during phase transitions of phase change materials (PCMs) for thermal storage, offering advantages such as high energy storage density and stable phase change temperatures, making it an effective method for heat storage.⁶

Molten salt PCM have significant application prospects in the field of latent heat energy storage, due to their wide range of phase change temperatures, high density, and low cost. The authors of the paper synthesized $\text{LiNO}_3\text{-KCl/SiO}_2$ composite PCM, achieving an energy storage density during the phase change stage of $3.4 \times 10^8 \text{ J m}^{-3}$, which is 2.5 to 4 times higher than that reported in previous studies.⁷⁻⁹ Qi *et al.* proposed a hydroxylation method, resulting in a phase change enthalpy of the molten salt composite PCM reaching 355.52 J g^{-1} .¹⁰ Shuai *et al.* utilized porous silicon carbide ceramics and solar salt to form a composite PCM, resulting in a more uniform temperature distribution during phase change, with the maximum temperature difference reduced from $148 \text{ }^\circ\text{C}$ to $130 \text{ }^\circ\text{C}$, and an overall phase change rate increased by 42.9%.¹¹ However, due to the limitations in electrical conductivity and shape stability of molten salt PCM, heat storage is typically achieved through indirect

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electrical heating (IEH). This involves converting electrical energy into thermal energy using external devices such as electric heaters, which is then transferred to the PCM for storage. Chuan *et al.* investigated the charging performance of shell-and-tube devices containing molten salt-based PCMs. Compared to devices containing pure salt, the melting process in devices containing salt composite materials can be shortened by approximately 79%.¹² Li *et al.* further established a high-temperature packed bed heat storage system encapsulating molten salt PCM, which exhibited an overall efficiency 1.9–2.4 times higher than shell-and-tube thermal storage systems.¹³ However, the longer energy transfer path in IEH leads to increased energy losses due to intermediate heat exchange processes. In contrast, direct electrical heating (DEH) enables energy storage by directly heating the phase change material through electric current. This transition eliminates intermediate heat transfer processes, thereby improving system efficiency.

Currently, some researchers have explored the electrothermal conversion performance of PCM under DEH. Maleki *et al.* prepared multilevel porous carbon foams (CFs) as conductive frameworks, achieving an energy conversion-storage efficiency of up to 74% during the phase change.¹⁴ Xue *et al.* fabricated hybrid aerogels composed of melamine foam (MF) and graphene oxide (GO)/graphene nanoplatelets (GNPs), with a phase change enthalpy of 161.7 J g⁻¹ and a thermal conductivity of 1.46 W m⁻¹ K⁻¹, achieving an energy conversion-storage efficiency of 62.5%.¹⁵ Zhang *et al.* found that by adding 20 wt% carbon black to polyethylene glycol–calcium chloride PCM, an energy conversion-storage efficiency of 64.7% could be achieved.¹⁶ Due to the poor thermal conductivity of most PCMs, PCM and framework materials alone cannot meet the high thermal conductivity requirements of materials, so the addition of conductive fillers can improve the thermal conductivity of composite PCMs.¹⁷ Sun *et al.* employed carbon nanotubes as thermal conductive fillers, resulting in a reduction of electrical resistivity from 9500 Ω m to 90 Ω m and an increase in thermal conductivity to 0.91 W m⁻¹ K⁻¹ in the composite PCM. As a result, the energy conversion-storage efficiency reached 70.2%.¹⁸ Aftab *et al.* infiltrated polyethylene glycol PCM into carbon nanotube arrays, achieving an energy conversion-storage efficiency as high as 94%, with the capability of rapid response to electrical heating under low voltage conditions.¹⁹ It is evident that these studies all use organic substances as PCM. However, organic PCMs usually have phase-change temperatures below 100 °C, lower density, higher costs, and a certain level of flammability. These factors result in issues such as low heat storage density, a narrow application temperature range, and poor safety, thereby restricting their practical use. Furthermore, the dimensions of the composite PCM samples prepared in the mentioned studies are relatively small, with a volume of approximately 5 cm³. Due to the small sample size, the material often shows good temperature uniformity in the electric heating process, but in the large-scale heat storage system, the amount of electric heating conversion material may reach tens of kilograms or even higher, and the size of the system will also be

an order of magnitude difference. This may lead to local overheating of the system. If the temperature monitoring device fails to accurately monitor the overall temperature of the system, there is a risk of local melting of the heat exchange tube and chemical decomposition failure of the material. However, the uniformity of temperature in larger-sized PCMs during the electrothermal conversion process has not been validated.

This paper utilizes a mixed molten salt of LiNO₃ and KCl as the PCM and employs hydrophilic gas-phase silica (SiO₂) as a porous adsorption carrier. Through the molten adsorption method, a shaped molten salt/SiO₂ composite phase-change material (CPCM) is prepared. We added expanded graphite (EG) to the CPCM to provide a conductive and thermally conductive network, thereby imparting electrical conductivity and high thermal performance to the CPCM. Subsequently, we investigated the relationship between the added amount of EG and the thermal conductivity, enthalpy, and apparent specific heat capacity of the CPCM. This exploration aimed to uncover how EG regulates the thermal properties of the CPCM. Finally, we established an electrothermal conversion testing system for a CPCM device with dimensions of 170 mm in length, 35 mm in width, and 80 mm in height. The system allowed us to conduct comprehensive tests on the electrothermal conversion performance of the CPCM, covering both sensible heat storage and latent heat storage stages. Throughout our study, we identified key factors influencing the temperature uniformity of the electrothermal conversion device incorporating the CPCM. Moreover, by combining experimental research with numerical simulation, we effectively evaluated the electrothermal conversion capability of the CPCM within the device. This integrated approach contributes valuable insights and guidance for the practical application of molten salt electrothermal conversion PCMs.

2 Results and discussion

2.1 Experimental results analysis

2.1.1 Structural analysis of CPCM. To investigate the changes in the composition of LiNO₃–KCl/SiO₂/EG composite phase change material during the preparation process, structural analysis was conducted using SEM, FT-IR, and XRD for LiNO₃–KCl/SiO₂, EG, and LiNO₃–KCl/SiO₂/EG (6 wt%).

2.1.1.1 SEM. From Fig. S1,† it can be observed that the microstructure of EG exhibits a lamellar morphology, with relatively large pores between the layers, making it suitable for adsorbing phase change materials. After compounding EG with LiNO₃–KCl/SiO₂, LiNO₃–KCl/SiO₂ adheres to the layers of EG, as illustrated in Fig. S1d.†

2.1.1.2 FT-IR and XRD. From Fig. 1a, it can be inferred that, in EG, the absorption peak within the 1558 cm⁻¹ region is typically associated with the C=C stretching vibration peak. The absorption peaks at 1650 cm⁻¹ and 3450 cm⁻¹ correspond to the –OH characteristic peaks. Possible reasons for this may include the introduction of trace amounts of moisture during the sample pressing process, and the



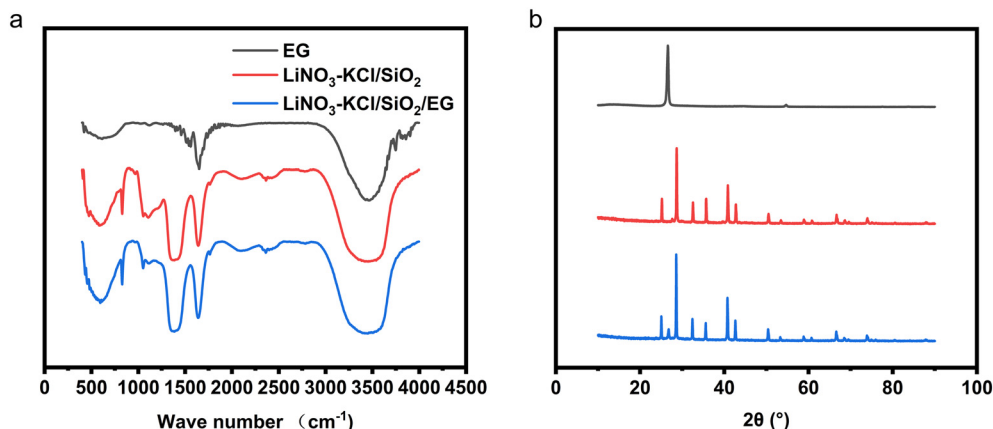


Fig. 1 (a) Infrared patterns of the materials involved; (b) XRD.

presence of a small amount of hydroxyl groups on the surface of EG, which contribute to the existence of this peak.

In $\text{LiNO}_3\text{-KCl/SiO}_2$, the absorption peak in the 595 cm^{-1} region is typically associated with Li-O vibrations. The absorption peak in the 809 cm^{-1} region is usually attributed to the symmetric stretching vibration of Si-O-Si bonds, a significant characteristic of the silica sample. The absorption peak in the 1091 cm^{-1} region corresponds to the antisymmetric stretching vibration of Si-O-Si bonds. The absorption peak in the 1450 cm^{-1} region is induced by the stretching vibration of N-O bonds. The absorption peak at 1650 cm^{-1} is attributed to water adsorbed by LiNO_3 , and the strong absorption peak at 3400 cm^{-1} corresponds to the stretching vibration of O-H bonds in physically adsorbed water in the sample.

In the XRD spectrum, EG exhibits a diffraction peak at 26.59° , corresponding to the (002) crystal plane of standard graphite (JCPDS no. 41-1487), which is consistent with the results reported in literature.²⁰⁻²² The relatively strong intensity of this diffraction peak indicates that the atomic arrangement on this crystal plane is ordered and dense. However, in the XRD spectrum of $\text{LiNO}_3\text{-KCl/SiO}_2/\text{EG}$

composite phase change material (CPCM), the intensity of the corresponding EG diffraction peak is weaker. This is attributed to the disruption of EG morphology during the preparation process of the CPCM, where EG and $\text{LiNO}_3\text{-KCl/SiO}_2$ undergo stirring and crushing. As a result, the atomic spacing on the crystal plane of EG increases, and the structure becomes looser.

Compared to $\text{LiNO}_3\text{-KCl/SiO}_2$ and EG, the FT-IR and XRD spectra of $\text{LiNO}_3\text{-KCl/SiO}_2/\text{EG}$ CPCM do not exhibit new signal peaks. This observation suggests that the composite process involves physical blending, and no new substances are formed.

2.1.2 Thermal and electrical properties of CPCM

2.1.2.1 DSC and apparent specific heat capacity testing. The DSC curves of CPCM_x under different amounts of EG are depicted in Fig. 2a. As per my previous work, the melting enthalpy of $\text{LiNO}_3\text{-KCl/SiO}_2$ is established as 188.9 J g^{-1} , with a phase transition temperature of 159.8°C . Upon the addition of EG, the melting enthalpy of CPCM_x gradually decreases. This reduction is attributed to the fact that EG does not contribute latent heat to the composite phase change material. As shown in Table 1, the relative error between the actual enthalpy values

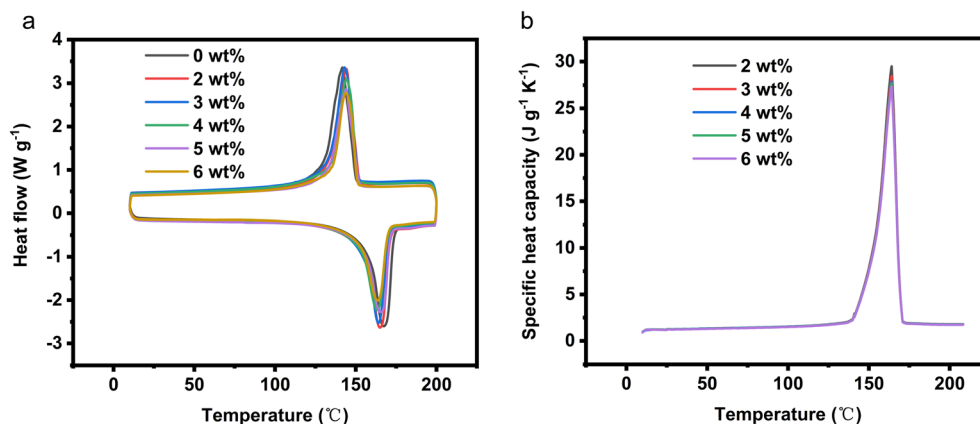


Fig. 2 (a) DSC curves of CPCM_x; (b) apparent specific heat capacity curves of CPCM_x.



Table 1 Phase change physical property data of CPCMX

Concentration of EG (wt%)	Enthalpy of fusion (J g^{-1})	Discrepancy from the theoretical value (%)	Enthalpy of solidification (J g^{-1})	Melting temperature ($^{\circ}\text{C}$)	Freezing temperature ($^{\circ}\text{C}$)
0	188.9	—	188.3	156.8	170.5
2	186.5	0.28	182.6	155.7	174.6
3	184.6	0.34	185.8	154.1	175.8
4	182.8	0.46	188.5	155.0	173.5
5	181.9	1.1	179.9	153.6	175.9
6	176.7	0.77	181	152.4	170.0

and the theoretical enthalpy values of CPCMX is less than 2%, falling within a reasonable range. The specific heat capacity of a material is an important parameter for measuring its thermal storage capacity. The apparent specific heat capacity curve of $\text{LiNO}_3\text{-KCl/SiO}_2/\text{EG}$ CPCMX is shown in Fig. 2b. The peak shapes of the apparent specific heat capacity curves for CPCMXs with different EG content are similar. The apparent specific heat capacity of the material, both before and after phase transition, represents the specific heat capacity of the material during sensible heat. During the phase transition of the material, the value of the apparent specific heat capacity can be calculated using an equivalent method. It can be observed from the figure that the addition of EG reduces the specific heat capacity of the material. This is because the specific heat capacity of the eutectic molten salt is greater than $1 \text{ J g}^{-1} \text{ K}^{-1}$, while the specific heat capacity of EG at room temperature is only $0.71 \text{ J g}^{-1} \text{ K}^{-1}$.

2.1.2.2 Volume resistivity and thermal conductivity. As shown in Fig. 3a, with the increase in the amount of EG, the thermal conductivity of the composite block shows an increasing trend, while the volume resistivity exhibits a decreasing trend. The opposite trends in volume resistivity and thermal conductivity are attributed to the high electrical

and thermal conductivity of EG. EG is composed of layered carbon atoms, with each layer of carbon atoms bonded together by covalent bonds into hexagonal rings. This layered structure enables rapid conduction of electrons and heat along the plane direction in EG. Additionally, the porous microstructure of EG provides additional conduction pathways, facilitating the transfer of electrons and heat within the material.

The change in volume resistivity of the CPCMX block with EG content exhibits a nonlinear relationship. This is attributed to the presence of the percolation effect of EG. As shown in Fig. 3b, when the amount of EG exceeds a critical value, a complete conductive network forms within the material,^{23,24} leading to a sharp decrease in the material's resistivity. With the increase in EG content, the CPCMX block undergoes a rapid transition from electrical insulator to conductor, and the electrical percolation threshold of EG is around 2 wt%. This is in close agreement with the results reported in literature.²⁵

2.1.3 Analysis of electrothermal conversion performance during the sensible heat storage phase. As shown in Fig. 4a, under a constant electric heating power of 80 W, the temperature rise rate of ETHCM2 is significantly higher than that of other EG content ETHCMs, while the temperature rise rates of other EG content ETHCMs are similar. This difference may be attributed to the uneven distribution of resistance in ETHCM2, leading to poor temperature uniformity in the module.

The calculation results of energy conversion-storage efficiency are shown in Table 2. It can be observed that the energy conversion-storage efficiency of ETHCM2 is significantly higher than that of other ETHCMs. This discrepancy is likely due to the fact that the temperatures measured by the thermocouples cannot accurately reflect the average temperature of ETHCM2, leading to calculated energy conversion-storage efficiency values deviating from normal values.

2.1.4 Analysis of temperature uniformity during the sensible heat storage phase

2.1.4.1 Temperature uniformity analysis of EthCMx. From Fig. 4b, it can be observed that, for ETHCM2, there is a significant temperature difference of up to $17.6 \text{ }^{\circ}\text{C}$ between the temperatures measured at points T_1 and T_3 , while for ETHCM6, this temperature difference is only $0.2 \text{ }^{\circ}\text{C}$. This discrepancy is attributed to the highly thermally conductive and porous structure of EG. When the amount of EG is low, the main part of the CPCMX is still formed by the insulating

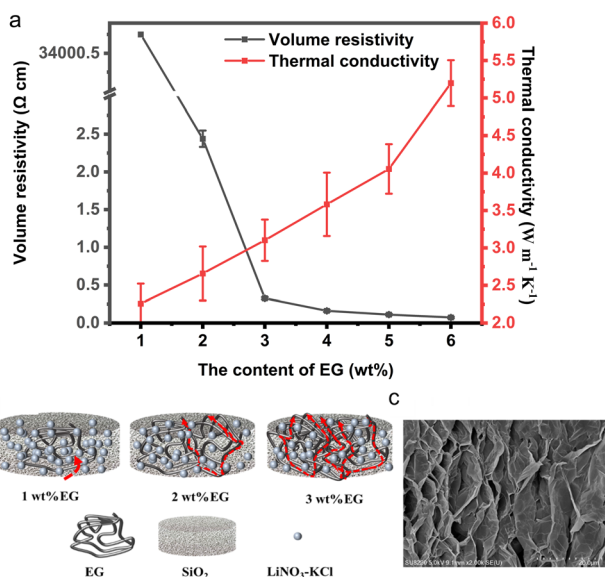


Fig. 3 (a) Analysis of volume resistivity and thermal conductivity of CPCMX; (b) schematic diagram of conductive network formation; (c) SEM diagram of porous microstructure of EG.



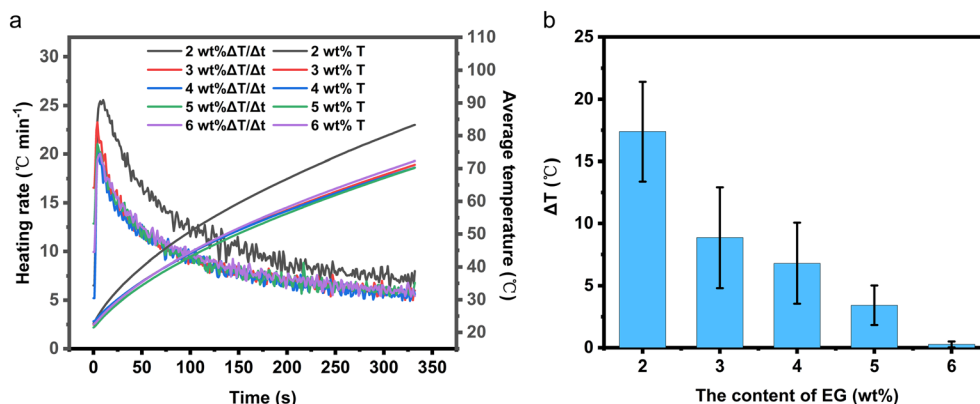


Fig. 4 (a) Temperature rise and temperature rise rate curve of EThCMx; (b) temperature difference statistics at T_1 and T_3 of EThCMx.

Table 2 Data statistics of EThCMx energy conversion-storage efficiency

Concentration of EG (wt%)	Initial average temperature (°C)	Final average temperature (°C)	Energy conversion and storage efficiency (%)	Heating rate (°C min ⁻¹)
2	22.63	86.01	95.2	11.52
3	22.37	71.12	76.1	8.86
4	22.44	70.33	74.7	8.71
5	21.5	70.27	75.1	8.87
6	22.48	72.31	77.1	9.06

and low thermally conductive SiO₂ skeleton network, and EG serves as a conductive and thermal filling material inside the CPCM. During sample preparation, uneven mixing of materials may lead to temperature anomalies during the electric heating process. When the amount of EG is high, the main part of the CPCM becomes the conductive and thermally conductive EG, while the insulating and low thermally conductive LiNO₃-KCl/SiO₂ is filled in the interlayer gaps of EG. At this point, the conductive and thermally conductive network of the CPCM has been established.

2.1.4.2 Analysis of the impact of electrode resistance distribution on temperature uniformity of the EThCM. During the experimental process, it was observed that the temperature in the middle part (T_2) of EThCM is slightly lower than the temperatures on both sides (T_1 and T_3) in most cases. To investigate the cause of this phenomenon, this section includes a comparative experiment. As shown in Table 3, all graphite electrodes have the same specifications.

In the experimental group, the resistivity of one pair of graphite electrodes differs by a factor of two, while in the control group, it is equal. To control variables, CPCM6 was selected, and the electric heating power was set to 80 W.

From Fig. 5, it can be observed that the temperature distribution of the electrothermal conversion module varies gradually, first decreasing and then increasing, as the positions of points A, B, C, D, E, F, and G change. This phenomenon arises due to the inconsistent temperature rise rates between the graphite electrode and the CPCM block. The temperature rise rates of the graphite electrode and CPCM block can be calculated using the following formula:

$$I^2Rt = C_p m \Delta T \quad (1)$$

Upon conversion, the following results can be obtained:

$$v = \frac{\Delta T}{t} = \frac{I^2 R}{C_p m} \quad (2)$$

Table 3 Statistical table of graphite electrode parameters

	Experimental group		Control group	
	The electrode near the T_1 region	The electrode near the T_3 region	The electrode near the T_1 region	The electrode near the T_3 region
Mass (g)	9.12	9.08	8.90	9.15
Total resistance (mΩ)	31.63	16.00	16.80	16.40
Resistivity ($\times 10^{-3}$ Ω cm)	1.52	0.77	0.81	0.79
Specifications (mm ³)	100 × 16 × 3			



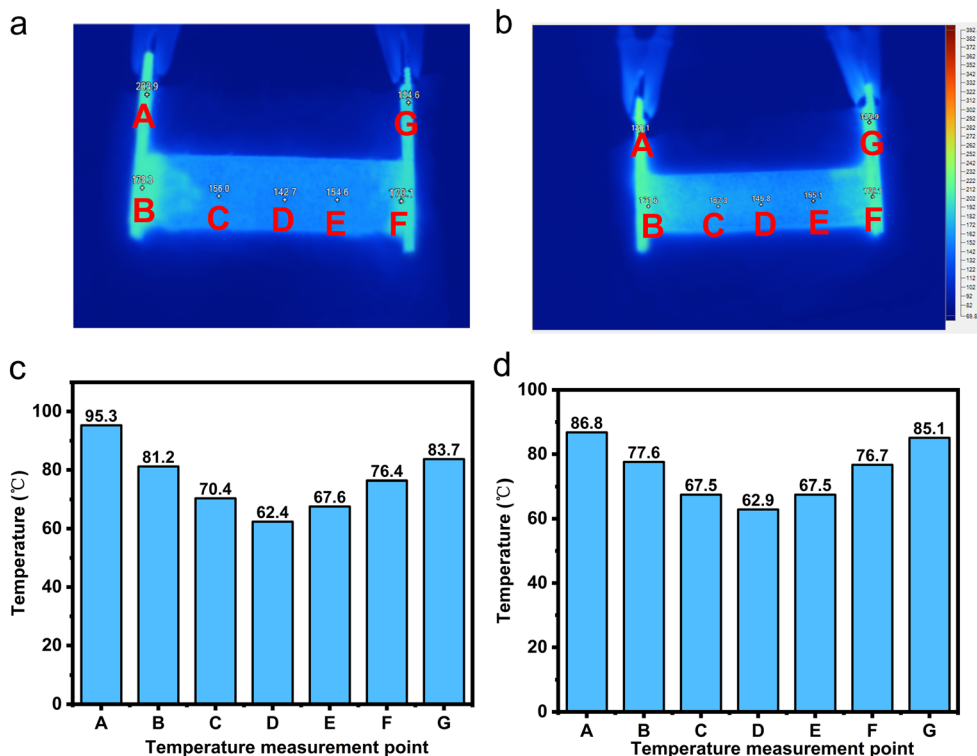


Fig. 5 (a) Infrared thermal imaging of the experimental group; (b) infrared thermal imaging of control group; (c) the temperature-location analysis diagram of the experimental group; (d) temperature-location analysis diagram of control group.

Where ν is the heating rate, ΔT is the temperature rise, t is the duration of electric heating, I is the current passing through the material, R is the resistance of the material, c is the specific heat capacity of the material, and m is the mass of the material.

As the graphite electrode and CPCM block are connected in series, the ratio of the temperature rise rates between the graphite electrode and the CPCM block can be utilized to eliminate the common term I^2 in the formula.

$$\frac{\nu_1}{\nu_2} = \frac{R_1}{R_2} \times \frac{C_{P,2}}{C_{P,1}} \times \frac{m_2}{m_1} \quad (3)$$

Where the subscript 1 represents the parameters of the graphite electrode, and subscript 2 represents the parameters of the CPCM block. The specific heat capacity of graphite $C_{P,1}$ is approximately $0.71 \text{ J g}^{-1} \text{ K}^{-1}$, and within the temperature rise range, the specific heat capacity of CPCM $C_{P,2}$ is $1.45 \text{ J g}^{-1} \text{ K}^{-1}$. The mass of a single graphite electrode m_1 is approximately 9.5 g, and the mass of the CPCM block m_2 is 138 g. The experimentally measured resistance of the CPCM block R_2 is 216 m Ω . From these parameters, the following conclusions can be drawn:

(1) When the resistance of the graphite electrode is 7.28 m Ω , ν_1 equals ν_2 . The temperature rise rate of the graphite electrode is equal to that of the CPCM block, resulting in uniform temperatures throughout the electrothermal conversion module.

(2) When the resistance of the graphite electrode exceeds 7.28 m Ω , ν_1 is greater than ν_2 . At this point, the temperature rise rate of the graphite electrode is higher than that of the CPCM block.

(3) When the resistance of the graphite electrode is less than 7.28 m Ω , ν_1 is less than ν_2 . In this case, the temperature rise rate of the graphite electrode is lower than that of the CPCM block.

2.1.4.3 Analysis of the influence of applied voltage magnitude on temperature uniformity of the electrothermal conversion module. As shown in Fig. 6, as the externally applied voltage increases, the temperature difference inside EThCM becomes larger. This is because a higher voltage results in greater heating power in the graphite electrode, leading to a larger difference in heating power between the graphite electrode and the CPCM block, causing an increase in the temperature difference between them. Additionally, due to the presence of contact resistance between the graphite electrode and the CPCM block, as the input voltage increases, the heating power at the contact interface also sharply increases, resulting in a larger temperature difference between the average temperatures at T_1 and T_3 and the temperature at T_2 within the EThCM. After heating for a certain period, the temperature difference in EThCM tends to stabilize. This is because the graphite electrode is exposed to the environment, and as the temperature increases, its heat dissipation power also increases. When the heat dissipation power balances with the electric heating power, the temperature difference also tends to balance.



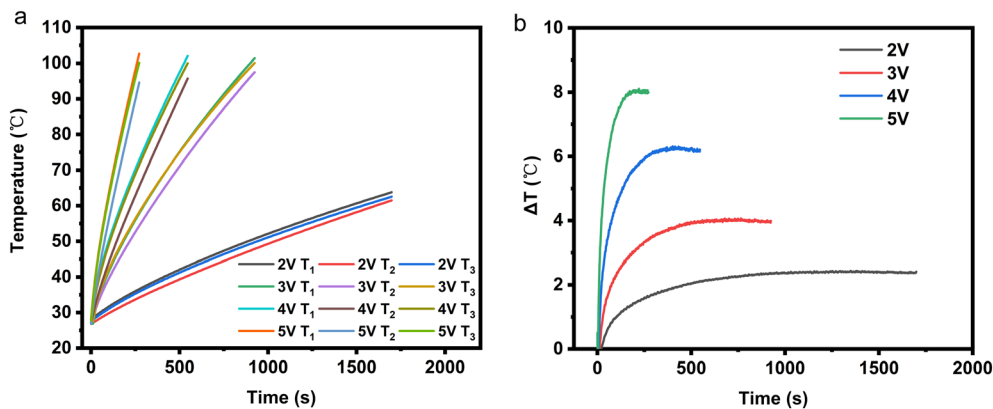


Fig. 6 (a) The temperature–time relationships at various positions under different input voltages; (b) the temperature difference–time relationships at various positions under different input voltages.

Through an exploration of the above influencing factors, the optimal experimental conditions for EThCM can be summarized as follows:

(1) The amount of EG in LiNO₃–KCl/SiO₂/EG composite phase change material is 6 wt% (CPCM6). At this point, the conductive and thermally conductive network of the composite phase change material has been established, resulting in relatively uniform heating throughout EThCM.

(2) The resistivity of the graphite electrodes on both sides of EThCM is equal. In this scenario, the heating power and heat transfer power of the graphite electrodes at both ends of EThCM are equal, and the temperature gradient exhibits a symmetric distribution with the module's central axis as the reference.

(3) When the graphite electrodes and CPCM block form EThCM, there exists a critical resistance value for the graphite electrodes. When the resistance of the graphite electrode is at the critical value, the temperature distribution throughout EThCM is uniform during the electric heating process.

2.1.5 Analysis of electrothermal conversion performance during the latent heat storage phase. In accordance with the optimal experimental conditions summarized earlier, electrothermal conversion experiments were conducted. However, due to experimental constraints, the minimum resistivity of the graphite electrode remained at 16 mΩ, which exceeds the critical resistance value of 7.28 mΩ. To balance the temperature uniformity of the material and the electrothermal storage efficiency, the input voltage was set to 4.5 V. At this voltage, it is anticipated that the temperature difference during the sensible and latent heat storage phases would be around 6 °C, with a heating rate exceeding 9 °C min⁻¹, indicating rapid heat storage capabilities. As depicted in Fig. 7a, under a constant voltage of 4.5 V, the resistivity of EThCM fluctuates during the electric heating process. At 500 s, the resistivity of the material begins to slowly rise, with an increase of approximately 10%. This is attributed to the onset of phase transition in the material. Post-transition, the resistivity of the material is higher than that before the phase transition, consistent with conclusions found in the literature.²⁶

As shown in Fig. 7b, with the progression of the heating process, the electrothermal conversion module undergoes both sensible and latent heat storage phases. The module's heating rate exhibits a trend of initially decreasing and then increasing, while the temperature difference shows a pattern of initially increasing, followed by a decrease with a peak value.

When the electric heating progresses to 630 s (point A), the heating rate exhibits a small plateau, followed by a sharp decrease. This is because EThCM enters the phase change

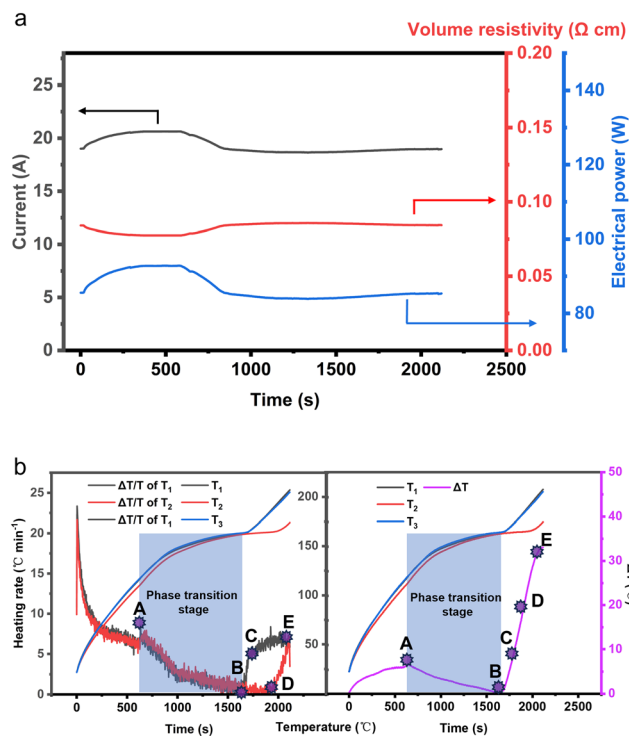


Fig. 7 (a) Analysis of electric performance parameters in electric heating process; (b) the relationship between temperature, temperature difference and heating rate and time of each part in the process of electric heating.



stage at this point. The apparent specific heat capacity of the material sharply increases, and the heat obtained from electrothermal conversion is stored in the material in the form of latent heat storage. Similarly, in the temperature difference–time relationship graph, at 630 s, the internal temperature difference within EThCM reaches its maximum value. Subsequently, as the material enters the latent heat storage phase, the internal temperature difference within EThCM decreases.

When electric heating progresses to 1676 s (point B), due to the position at T_1 being closer to the graphite electrode than T_2 , the material at T_1 completes the phase transition process first. At this point, the apparent specific heat capacity of the material at this position sharply decreases, and the heating rate rapidly increases, reaching $5.78\text{ }^\circ\text{C min}^{-1}$. The apparent temperature of the material also rises, while the material at T_2 is still in the phase change stage. The internal temperature difference within EThCM gradually increases.

When electric heating progresses to 1712 s (point C), the rate of change of the heating rate at T_1 decreases. This is because there is a temperature difference between T_1 and T_2 , and the heat absorbed at T_1 is not only used to raise its own temperature but is also conducted in the form of heat transfer to the lower temperature at T_2 .

When electric heating progresses to 1922 s (point D), the material at T_2 completes the phase transition, and the heating rate of the material in the central part of EThCM also increases. At this point, the entire EThCM completes latent heat storage and enters the sensible heat storage phase. The heat absorbed by the material at T_2 is entirely used to raise its own temperature, without the need to conduct to a lower temperature, so the rate of change of the heating rate at T_2 remains constant and is greater than the rate of change of the heating rate at T_1 .

When electric heating progresses to 2110 s (point E), the heating rate at T_2 exceeds that at T_1 for the first time, and a plateau appears in the temperature difference between T_1 and T_2 .

2.2 Simulation results analysis

2.2.1 Simulation analysis of the influence of electrode resistance distribution on temperature uniformity

2.2.1.1 Setting of monitoring positions and simulation parameters. The monitoring point positions of EThCM are

shown in Fig. S2.† Through the analysis of grid independence, the final calculation model adopts a total number of grids of 1447, and the time step is set to 0.1 s.

In the simulation process, it is necessary to determine the heating density of the material. The physical properties of the materials are listed in Table S1.† Since in the EThCM, the “electrode AB-block CDE-electrode GF” forms a series system, the current is equal everywhere. According to the power formula $I^2 = P/R$, it is known that the heating power of the material is proportional to the resistance of the material. Therefore, by setting the total input power to 80 W, the heating power of each part can be calculated, and thus the heating density of each part can be calculated. To verify the calculation results of the critical resistance of the graphite electrode and the correctness of the conclusions in the previous text, four operating conditions are set as shown in Table 4. Among them, in conditions 1 and 2, the resistance of the graphite electrode is higher than the critical resistance of the graphite electrode, 7.28 mΩ. In condition 3, the resistance of the graphite electrode is lower than this value, and in condition 4, the resistance of the graphite electrode is equal to this value.

2.2.1.2 Analysis of simulation results. Numerical simulations were conducted for 120 seconds on the unsealed EThCM under different operating conditions. The temperature contour map and the temperatures at various monitoring points obtained from the simulation are shown in Fig. 8a–d. As shown in Fig. 8d, under the conditions of condition 4, when the resistance of the graphite electrode is at the critical value, the heating of the graphite electrode and the CPCM block is uniform, and the temperature differences at various monitoring points are small. This confirms the existence of the critical resistance of the graphite electrode, and the calculation results are consistent with theoretical values.

From Fig. 8e and f, it can be observed that the temperature contour maps for conditions 1 and 2 generally match the infrared thermal images in Fig. 8a and b, showing a trend of lower temperature in the middle and higher temperatures on both sides. However, through the comparison between simulation and experiment, there are still some differences. Specifically, the simulated temperatures are higher than the actual temperatures. This is attributed to convective heat transfer between the graphite electrode and the air in actual conditions, causing the graphite electrode's heating rate to be

Table 4 The set value of each parameter under four working conditions

Operating condition	Parameter	Left-side graphite electrode	Right-side graphite electrode	CPCM block
1	Resistance (Ω)	0.032	0.016	0.216
	Heat density (W m^{-3})	2.12×10^6	1.06×10^6	8.58×10^5
2	Resistance (Ω)	0.016	0.016	0.216
	Heat density (W m^{-3})	1.13×10^6	1.13×10^6	9.13×10^5
3	Resistance (Ω)	0	0	0.216
	Heat density (W m^{-3})	0	0	1.05×10^6
4	Resistance (Ω)	0.00728	0.00728	0.216
	Heat density (W m^{-3})	5.62×10^5	5.62×10^5	9.79×10^5



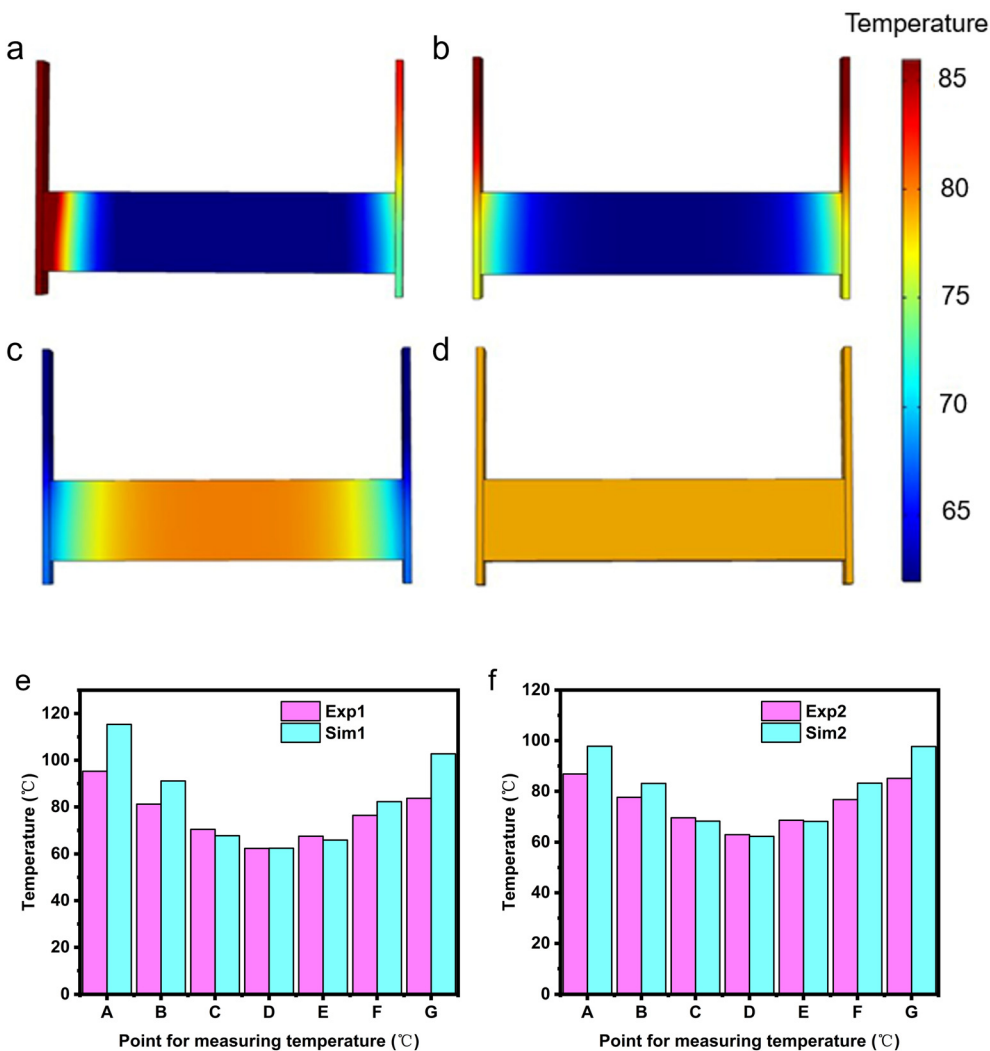


Fig. 8 (a) The temperature distribution map of the module under operating condition 1; (b) the temperature distribution map of the module under operating condition 2; (c) the temperature distribution map of the module under operating condition 3; (d) the temperature distribution map of the module under operating condition 4; (e and f) the comparison between simulation and experimental results under operating condition 1; (f) the comparison between simulation and experimental results under operating condition 2.

lower than the theoretical value. Additionally, due to the longer heating time in actual conditions, the temperatures at points C and E are slightly higher than the simulated values.

2.2.2 Simulation analysis of the electric heating process.

Due to the external encapsulation of CPCMs with a layer of sealing glue, it is not possible to accurately determine the temperature distribution of the sealing glue through experimental methods, making it challenging to precisely calculate the energy storage density of EThCM. Therefore, numerical simulation analysis is employed, based on existing data, to model the system and analyze the temperature distribution of EThCM during electric heating. In the simulation conducted in this section, the total number of grid points used in the model is 1722, and the time step is set to 1 s. The parameter settings for each component under the simulated operating conditions are listed in Table S2.† The variation of heat density for each component is depicted in Fig. S3.† This approach allows for the calculation of energy

storage density at different locations within EThCM and the examination of the impact of varying input power on the energy storage rate. In this section, COMSOL software is utilized for numerical simulation analysis.

To verify the accuracy of the simulation results, a comparison is made between the simulated and experimental results. As shown in Fig. 9, the temperatures of T_1 and T_2 are calculated using the apparent specific heat model. The simulated temperature trends closely resemble the observed temperature trends. Additionally, the relative errors for T_1 and T_2 are within reasonable limits, with T_1 's relative error not exceeding 1.3% and T_2 's relative error not exceeding 4%. This indicates that the model established in this section is capable of accurately calculating the heating process of EThCM.

At the end of the electrical heating process, the parameters of each component of EThCM are summarized in Table 5. From the table, it can be observed that despite the larger volume of



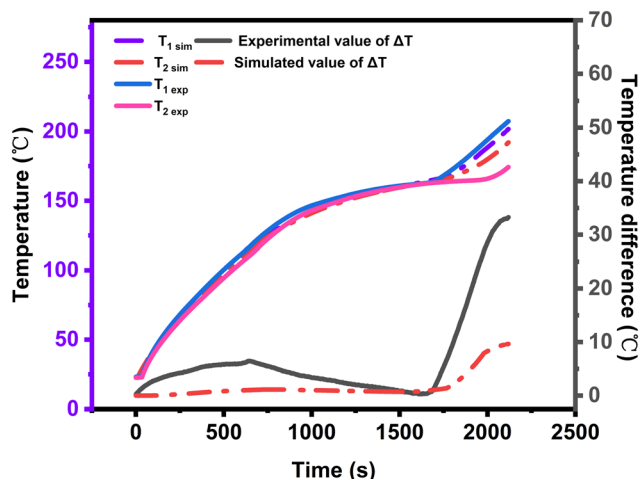


Fig. 9 Experimental verification of simulation model.

the sealing gel and its lower energy storage density, it still stores a considerable amount of energy. The heat generated during electrical heating is stored by both the CPCMC block in the form of sensible and latent heat storage and the sealing gel through sensible heat storage. The table also indicates that, compared to sensible heat storage materials, CPCMC offers the advantage of higher energy storage density with a smaller volume, further highlighting the promising prospects of applying CPCMC in the field of electrothermal conversion.

2.2.3 Comparative simulation analysis of DEH and IEH. In order to highlight the advantages of rapid and efficient thermal energy storage in CPCMC, a comparative analysis of the energy storage rates between DEH and IEH for EThCM was conducted. DEH involves the application of electrical power directly to CPCMC to achieve thermal energy storage, while IEH employs an electrically heated plate to transfer heat to CPCMC, enabling thermal energy storage in the composite phase-change material. For ease of calculating the module's energy storage rate, numerical simulation analysis was employed, utilizing established data for modeling purposes. The temperature distribution during the electrical heating process was analyzed for both models, and a comparison of the thermal storage rates between direct and indirect heating was conducted.

2.2.3.1 Model and mesh partitioning. Numerical simulation and analysis were conducted using COMSOL software. The position of the electrical heating plate and the mesh model for the indirect heating scenario are illustrated in Fig. 10a and b. The parameter settings for each component

under the two simulated operating conditions are presented in Table 6.

During the simulation, it is crucial to monitor the temperature of various parts of EThCM. The locations of monitoring points are depicted in Fig. 10c and d. For comparison purposes, the input power of the direct heating model was set to 130 W. The curves depicting the variation of the heat generation density of the CPCMC with temperature in both the direct heating and indirect heating models are illustrated in Fig. 10e. This curve was computed based on the apparent specific heat capacity function of the composite phase-change material block (CPCMC6). As indicated in the figure, the heating process remained stable, with the heat generation density around $1.465 \times 10^6 \text{ W m}^{-3}$, and the overall deviation was less than 1%. Furthermore, the dimensions and volumes of the CPCMC and the electric heating plate were consistent between the two conditions, indicating that their heat generation powers were identical. In the indirect heating model, the electric heating plate was in full contact with the CPCMC, with the thermal resistance within it being neglected.

2.2.3.2 Comparative simulation results and discussion. From Fig. 10f, it is evident that under the condition of consistent heating density–temperature curves, the material's heating rate is higher in the DEH model compared to the IEH model. In the IEH model, the time required for the material to reach 200 °C is 2038 s, with a heating rate of 5.3 °C min^{-1} . In the DEH model, the time required for the material to reach 200 °C is 1051 s, with a heating rate of $10.27 \text{ °C min}^{-1}$. Adopting direct electrical heating shortens the material's energy storage time by 48.4%, and the energy storage rate nearly doubles. This improvement is attributed to the real-time nature of direct electrical heating, where electrical energy is instantly converted into material thermal energy, unlike the IEH model, where heat transfer rate is limited by temperature differentials.

From Fig. 11d, it is evident that when the IEH model operates for 2050 s, certain local positions within the CPCMC block have not reached 200 °C. Conversely, as depicted in Fig. 11h, when the DEH model operates for 1100 s, the entire CPCMC block has already reached 200 °C, providing further evidence that the heat storage rate of DEH is significantly higher than that of IEH. Additionally, as illustrated in Fig. 11b, the temperature scale range for the IEH model at 1000 s ranges from 120 °C to 160 °C, with a temperature difference of 40 °C. In comparison, Fig. 11h shows that when the DEH model operates for 1100 s, the temperature range is 200 °C to 220 °C, with a reduced temperature difference of 20 °C. This indicates

Table 5 Performance parameters of the simulation model when running 2119 s

Object	Average temperature (°C)	Heating rate (°C min ⁻¹)	Volume (cm ³)	Stored energy (J)	Energy density (×10 ³ J m ⁻³)
Graphite	225.9	5.8	4.95	1302.9	0.263
CPCMC block	201.8	5.14	76.5	109 063	1.425
Sealing adhesive	170.3	4.25	393.7	105 384	0.267



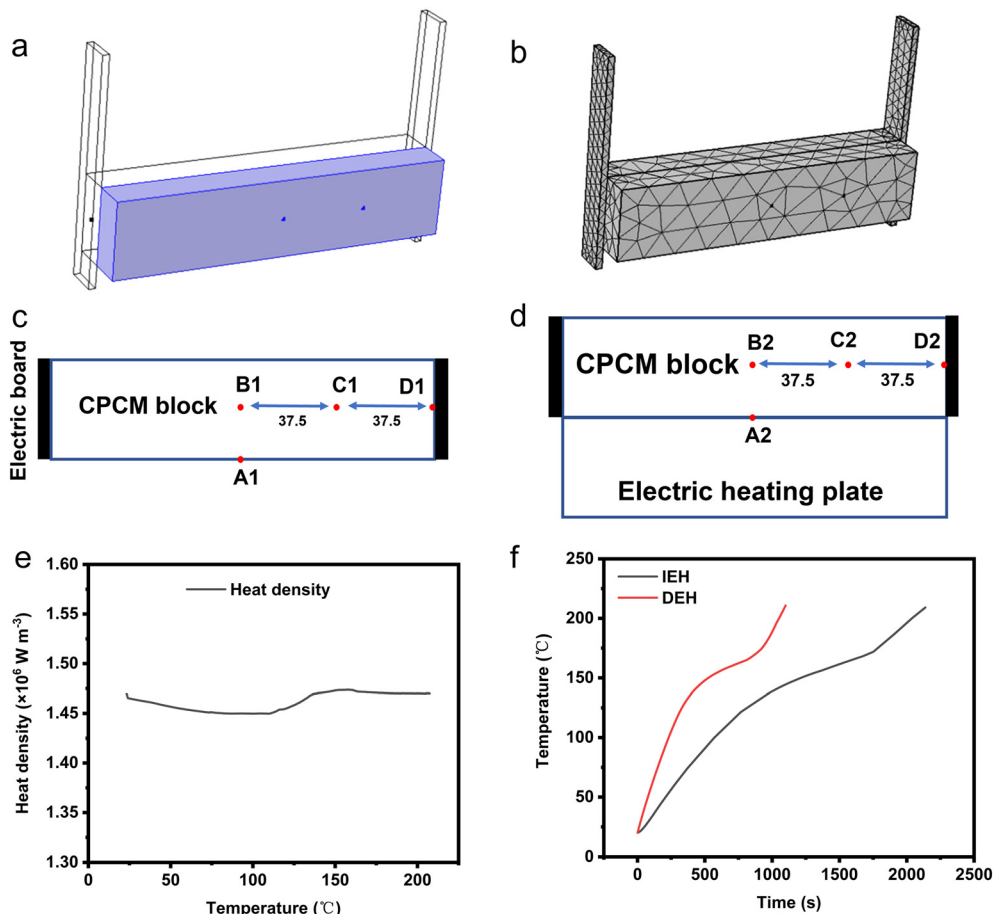


Fig. 10 (a) Position of electric heating plate; (b) grid model; (c) distribution of temperature monitoring points in IEH models (top view); (d) distribution of temperature monitoring points in DEH models (top view); (e) curve of heating density of electric heating plate and composite phase change material with temperature; (f) mean temperature–time curves for IEH and DEH.

Table 6 Parameter setting values of each component under two simulated working conditions

	IEH	DEH
	Indirectly heated electric heating plate	Directly heating composite phase transition material
Specific heat capacity ($\text{J g}^{-1} \text{K}^{-1}$)	0.5	Apparent specific heat capacity function
Density (g cm^{-3})	7.98	1.8
Thermal conductivity ($\text{W m}^{-1} \text{K}^{-1}$)	16.2	5.2
Heat generation density (W m^{-3})	$1.465 \times 10^6 \pm 1\%$	

that the temperature uniformity of the DEH model is superior to that of the IEH model. Moreover, as discussed in the previous section, by controlling the electric field, it is possible to effectively regulate the temperature field of the material. Therefore, through optimization, the temperature difference of the DEH model can be further reduced, improving its temperature uniformity.

3 Conclusion

This study employed physical blending to create $\text{LiNO}_3\text{-KCl/SiO}_2\text{/EG}$ composite phase-change materials (CPCM x) with varied EG content. EG modulation influenced CPCM's electrical

resistivity and thermal conductivity, providing electric-thermal conversion properties. CPCM2 formed a preliminary conductive network, while CPCM3 established a substantial network. CPCM6 exhibited a high melting enthalpy of 176 J g^{-1} and a phase transition temperature of $152 \text{ }^\circ\text{C}$. Following this, the establishment of a CPCM electrothermal conversion system (EThCM) revealed through electric heating experiments that temperature uniformity in EThCM is influenced by EG content, electrode resistance distribution, and external applied voltage. Higher EG content, smaller differences in resistance between electrodes, closer electrode resistance to the critical value, and lower external voltage result in improved temperature uniformity in EThCM.



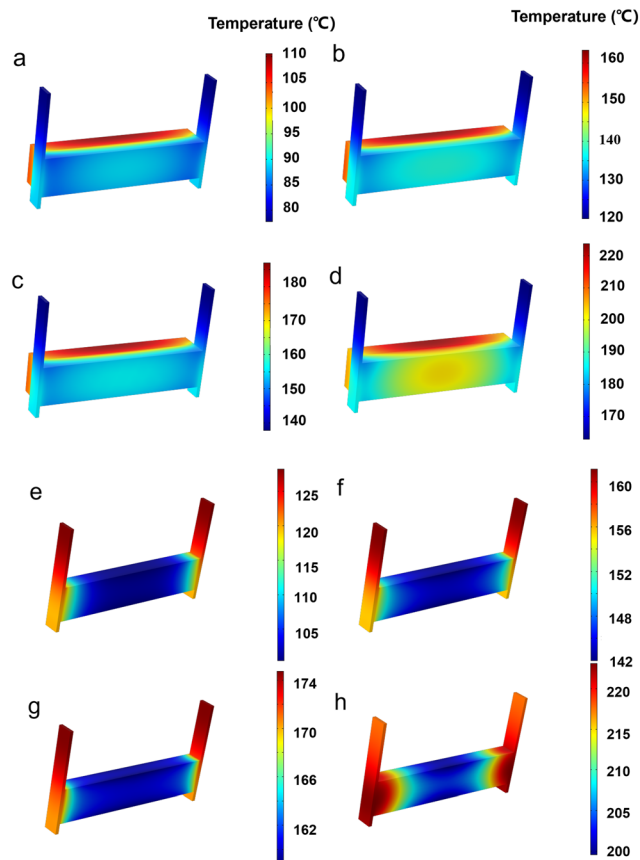


Fig. 11 Temperature distribution maps of different heating models at various operational durations. (a) IEH, 500 s; (b) IEH, 1000 s; (c) IEH, 1500 s; (d) IEH, 2050 s; (e) DEH, 250 s; (f) DEH, 500 s; (g) DEH, 750 s; (h) DEH, 1100 s.

Through the establishment of relevant models and simulation analysis, we analyzed the electric heating process in EThCM, validating the existence of the critical resistance value in the graphite electrode and revealing the circuit's regulation on the temperature field. Experimental verification led to strategies for regulating temperature uniformity. Additionally, simulations showed that direct heating significantly improves energy storage, reducing storage time by 48.4%, doubling the storage rate compared to indirect heating, and achieving better temperature uniformity. The error between simulation and experimentation is within 4%, confirming the model's accuracy. Simulation suggests further improvements in temperature uniformity through electric field regulation.

4 Experiment

4.1 Materials

Lithium nitrate (LiNO_3 , analytical grade) was purchased from Shanghai Macklin Biochemical Technology Co., Ltd.; potassium chloride (KCl, analytical grade) was obtained from Guangzhou Chemical Reagent Factory; hydrophilic gas-phase silica (SiO_2 , 99.8%) was purchased from Hubei Huifu

Table 7 Composition design ratio of composite phase transition materials

	Concentration of $\text{LiNO}_3\text{-KCl}$ (wt%)	Concentration of SiO_2 (wt%)	Concentration of EG (wt%)
CPCM2	88.2	9.8	2
CPCM3	87.3	9.7	3
CPCM4	86.4	9.6	4
CPCM5	85.5	9.5	5
CPCM6	84.6	9.4	6

Nanomaterials Co., Ltd.; expanded graphite (EG, industrial grade, 50 mesh) was obtained from Qingdao Furuite Graphite Co., Ltd.

4.2 Preparation of CPCM

The material preparation process is illustrated in Fig. S4.† Materials were weighed according to a mass ratio of 6:4 for LiNO_3 and KCl, and heated at a constant temperature of 220 °C for 1 hour to achieve co-melting. Subsequently, SiO_2 was added in a mass ratio of 9:1 to the eutectic salt to prepare the $\text{LiNO}_3\text{-KCl/SiO}_2$ composite phase-change material. To regulate its electrothermal performance, $\text{LiNO}_3\text{-KCl/SiO}_2\text{/EG}$ composite phase-change material (CPCM) was prepared with mass fractions of expanded graphite (EG) ranging from 2% to 6%. The specific steps are as follows: the $\text{LiNO}_3\text{-KCl/SiO}_2$ composite phase-change material was heated for 1 hour above the phase-change temperature. Different mass fractions of EG were separately weighed and added to a beaker, and the two components were stirred with a glass rod until they were uniformly mixed, resulting in $\text{LiNO}_3\text{-KCl/SiO}_2\text{/EG}$ composite phase-change materials with varying EG ratios. As shown in Table 7, when distinguishing between different EG addition amounts, they were denoted as CPCM $_x$, where x represents the mass fraction of EG. While still hot, a certain amount of powdered material was weighed into a mold, and a flat vulcanizing machine was used to press the composite phase-change material into blocks (150 mm in length, 15 mm in width, and 34 mm in height) at a temperature of 150 °C.

4.3 Characterization

4.3.1 Structural analysis. The surface morphology of SiO_2 and EG was characterized using the SU8220 scanning electron microscope (SEM) to observe the microstructure of the materials.

4.3.2 Chemical characteristics. The chemical structure of the experimental raw materials and CPCM was analyzed using the Nicolet IS50-Nicolet Continuum Fourier-transform infrared (FTIR) spectrometer. The measurements were conducted in the wavenumber range of 400–4000 cm^{-1} using the potassium bromide (KBR) pellet method.

The experimental raw materials and the CPCM were characterized using the X'pert Powder X-ray diffractometer to analyze their crystalline phases.



4.3.3 Thermal and electrical properties. The thermal and electrical properties of CPCMX with varying amounts of added EG were characterized using the Q20 differential scanning calorimeter (DSC) from TA Instruments, USA. The measurements included determining the phase transition temperatures and latent heat of the samples. The specimens were prepared as cylindrical blocks ($\Phi 30$ mm \times 10 mm). The thermal conductivity was measured using the TPS2500 Hot-Disk thermal constant analyzer.

Using the three-step method with a Differential Scanning Calorimeter (DSC), the apparent specific heat capacity of the $\text{LiNO}_3\text{-KCl/SiO}_2\text{/EG}$ composite phase change material was measured within the temperature range of 10 °C to 200 °C. The calculation formula is expressed as eqn (4).

$$C_{P,\text{PCM}} = C_{P,\text{SP}} \times \frac{(P_{\text{PCM}} - P_{\text{bas}}) \times m_{\text{SP}}}{(P_{\text{SP}} - P_{\text{bas}}) \times m_{\text{PCM}}} \quad (4)$$

Where $C_{P,\text{PCM}}$ is the measured apparent specific heat capacity of the sample, $C_{P,\text{SP}}$ is the specific heat capacity of sapphire, measured in $\text{J g}^{-1} \text{K}^{-1}$; P represents the heat flux density, and m is the mass. Subscripts PCM denote the $\text{LiNO}_3\text{-KCl/SiO}_2\text{/EG}$ composite phase change material, SP represents the sapphire reference sample, and bas indicates the baseline.

The samples were prepared as rectangular prisms with dimensions of 150 mm \times 15 mm \times 34 mm. The electrical performance was tested using the TH2511A DC low-resistance tester. The formula for calculating resistivity is given by $\rho = \left(\frac{R}{S}\right)/L$, where R is the measured resistance of

the material, and S and L represent the cross-sectional area and length of the electrothermal conversion module (EThCM), respectively.

4.3.4 Fabrication of electrothermal conversion modules and system setup. As depicted in Fig. S5,[†] graphite plates were affixed as electrodes at both ends of the CPCMX block, and conductive silver paste was utilized to reduce contact resistance. For accurate measurement of the internal temperature of the CPCMX block and assessment of temperature uniformity during the electric heating process, a handheld electric drill was used to create three equidistant holes on the block's surface. The hole depth was half of the block's height, and thermocouples were inserted. To prevent moisture from entering the material during electric heating, the block was sealed using an organosilicone potting compound. After sealing, demolding resulted in the composite phase change material electrothermal conversion module (EThCM). To differentiate between different amounts of added EG, they were designated as EThCM x , where x represents the mass fraction of EG. The external part of the electrothermal conversion module (EThCM) was enveloped in 30 mm thick insulation to minimize heat loss during the electric heating experiments. As depicted in Fig. 12a, the thermocouples inside the EThCM were connected to an Agilent data logger for internal temperature monitoring. The positive and negative terminals of the power source were clamped to the electrodes at both ends of the EThCM, completing the system setup.

4.3.5 Monitoring module temperature uniformity. The electrical properties of the module were monitored using a LAND Test System. EThCM was subjected to electric heating,

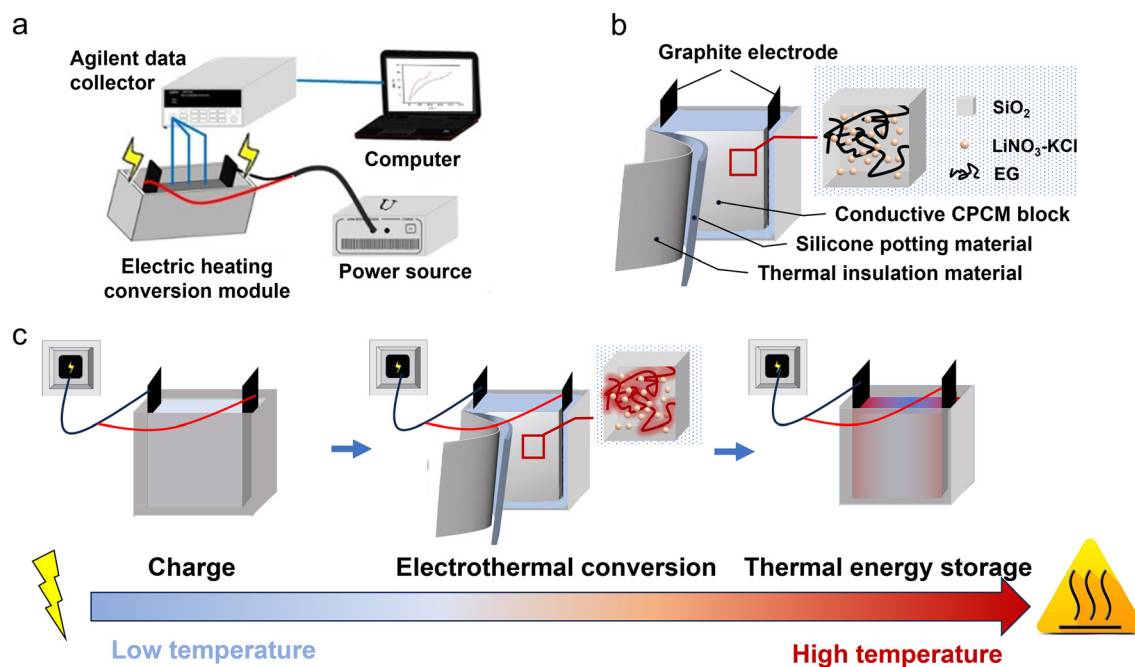


Fig. 12 (a) Schematic diagram of electrothermal conversion; (b) schematic diagram of electrothermal conversion module; (c) schematic of electrothermal conversion.



and an infrared thermal imaging camera, along with an Agilent temperature data logger, was employed to monitor the temperature of the CPCM block during the electric heating process. This approach allowed for the assessment of the temperature uniformity of the material during electric heating.

4.3.6 Electrothermal conversion testing. By measuring the material's temperature and incorporating the numerical values of the apparent specific heat capacity, one can calculate the energy conversion-storage efficiency of the material during electrothermal conversion. The formula for calculating this efficiency is presented as eqn (5):

$$\eta = \frac{m \int_{T=T_0}^{T_1} C_p(T) dT}{\int_{t=t_0}^{t_1} P(t) dt} \times 100\% \quad (5)$$

In the expressions, C_p represents the apparent specific heat capacity, and $\int_{T=T_0}^{T_1} C_p(T) dT$ denotes the integral of the apparent specific heat capacity with respect to temperature. The numerical value of this integral reflects the energy stored as the material's temperature increases from T_0 to T_1 , measured in joules (J). The variable m denotes the mass of the material, measured in grams (g). The symbol P represents electrical power, and $\int_{t=t_0}^{t_1} P(t) dt$ represents the integral of electrical power with respect to time. The numerical value of this integral indicates the energy output from the external power source during the time interval from t_0 to t_1 , equivalent to the energy absorbed by the test material.

4.4 Numerical simulation

By conducting simulation and analysis of the EThCM, we aim to validate experimental results and simulate conditions that are currently impractical to achieve in reality. Additionally, utilizing simulation software for EThCM allows for high accuracy, ease of operation, quick data processing, and resource-saving under the premise of a well-designed simulation. In this study, the EThCM model was designed using Solidworks software, and numerical simulation and analysis calculations were performed using COMSOL software. This approach provides a reliable means to supplement experimental findings and explore scenarios that may be challenging or impossible to implement physically.

4.4.1 Model structure. The structure of the electrothermal conversion module (EThCM), designed using Solidworks software, is illustrated in Fig. S6.†

This paper involves simulation content in three parts: electric heating simulation before and after sealing of the EThCM, and thermal simulation of indirect heating. Since the focus of simulation varies for these three parts, there are differences in the description of the physical models. Additionally, throughout the entire experimental process,

heat conduction, convective heat transfer, and thermal radiation coexist. To simplify the calculation process, a series of assumptions need to be made for the computational model, and there are variations in the assumptions made for the three parts.

4.4.2 Description of the physical model. For the three models, their common assumptions are as follows:

(1) Radiation heat transfer is neglected, and only conduction heat transfer is considered. (2) Contact thermal resistance and contact electrical resistance between the graphite electrodes and the composite phase-change material are ignored. (3) The physical properties of the graphite electrode are constant and do not change with an increase in temperature. (4) The graphite electrode and the composite phase-change material are assumed to be uniform and isotropic.

For the electrothermal conversion module model before sealing, the additional assumptions are as follows:

(1) Convective heat transfer is neglected, and the boundary condition for the outer wall of the module is adiabatic. (2) The physical properties of the composite phase-change material block are constant and do not change with an increase in temperature.

For the electrothermal conversion module model after sealing, the additional assumptions are as follows:

(1) Convective heat transfer is considered, and the outer wall of the module is subject to a heat flux boundary condition with a specified heat transfer coefficient of $4 \text{ W m}^{-2} \text{ K}^{-1}$. (2) The physical properties of the sealing compound are constant and do not change with an increase in temperature. The apparent specific heat capacity of the composite phase-change material block is considered a function of temperature. (3) Contact thermal resistance between the sealing compound and the composite phase-change material, as well as between the sealing compound and the electrodes, is neglected.

For the model of indirect heating, the additional assumptions are as follows:

(1) The electric heating plate is made of 316 stainless steel and is uniform and isotropic. (2) Contact thermal resistance between the electric heating plate and the composite phase-change material is neglected. (3) Convective heat transfer is considered, and a convective heat flux boundary condition is applied with a specified heat transfer coefficient of $10 \text{ W m}^{-2} \text{ K}^{-1}$.

4.4.3 Description of the control equations. For the three models, the common energy conservation equation is:

(1) Material energy conservation equation

$$H = \int_{T_0}^T C_p dT \quad (6)$$

In the equation, H represents the energy stored in the material, T_0 denotes the initial temperature, and C_p is the material's specific heat capacity at constant pressure.



(2) System energy conservation equation

$$\frac{\partial}{\partial t}(\rho H) = k\nabla^2 T + \dot{q} \quad (7)$$

In the equation, ρ represents the material density, k denotes the material's thermal conductivity, T and t represent temperature and time, and \dot{q} is the heat generation density of the material.

(3) Initial conditions

Under the initial conditions, it is assumed that the temperature of the electrothermal conversion module is uniform, with an initial temperature $T_0 = 20$ °C

$$T(x, y, z) = T_0 \quad (8)$$

(4) Boundary conditions at the interface between graphite electrode and composite phase change materials

$$-k_g \frac{\partial T}{\partial n} = -k_{\text{PCM}} \frac{\partial T}{\partial n} \quad (9)$$

In the expressions, k_g represents the thermal conductivity of the graphite electrode, and k_{PCM} represents the thermal conductivity of the CPCM block.

For the pre-encapsulation electrothermal conversion module, its supplementary governing equation is as follows:

Boundary conditions: during the simulation process, it is assumed that the outer wall of the electrothermal conversion module is adiabatic.

$$\frac{\partial T}{\partial x} = 0, \quad \frac{\partial T}{\partial y} = 0, \quad \frac{\partial T}{\partial z} = 0 \quad (10)$$

For the post-encapsulation electrothermal conversion module, its supplementary governing equation is as follows:

Boundary conditions: the wall adopts a heat flux boundary condition with a specified heat transfer coefficient of $4 \text{ W m}^{-2} \text{ K}^{-1}$.

Boundary conditions at the interface between the graphite electrode and the encapsulation material are as follows:

$$-k_g \frac{\partial T}{\partial n} = -k_p \frac{\partial T}{\partial n} \quad (11)$$

Boundary conditions at the interface between the CPCM and the encapsulation material are as follows:

$$-k_{\text{PCM}} \frac{\partial T}{\partial n} = -k_p \frac{\partial T}{\partial n} \quad (12)$$

Where k_p represents the thermal conductivity of the encapsulation material.

For the module incorporating the bonded heating element, its supplementary governing equation is as follows:

Boundary conditions: the wall adopts a heat flux boundary condition with a specified heat transfer coefficient of $10 \text{ W m}^{-2} \text{ K}^{-1}$.

Boundary conditions at the interface between the CPCM and the electric heating element are as follows:

$$-k_{\text{PCM}} \frac{\partial T}{\partial n} = -k_E \frac{\partial T}{\partial n} \quad (13)$$

Where k_E represents the thermal conductivity of the electric heating element.

Data availability

Data will be made available on request.

Author contributions

Jiandong Zuo: data curation, investigation, writing – original draft. Hongjie Luo: investigation, validation, visualization. Ziyue Ling: writing – review & editing, supervision. Zhengguo Zhang: conceptualization, project administration. Xiaoming Fang: methodology, supervision. Weiwei Zhang: funding acquisition, resources.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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