Improved latent heat storage properties through mesopore enrichment of a zeolitic shape stabilizer

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ABSTRACT

Latent heat storage systems are applied to keep temperature of a local environment within a constant range. The process takes place via release/storage of latent heat during freezing/melting of a corresponding phase change material embedded in a shape stabilizer, which is the scaffold keeping the phase change material stationary in its molten form. In this work, a highly siliceous ZSM-5 and modified versions thereof were chosen as shape stabilizers for molecular and polymeric phase change materials (namely lauric acid and polyethylene glycol), to be impregnated using solvent assisted vacuum impregnation. The dominantly microporous analogues, parent ZSM-5 and its acid-treated derivative, were limited to 40% uptake for each phase change material. Contrastingly, a mesopore rich analogue (as formed under basic conditions) reached 65% impregnation for lauric acid and 70% for polyethylene glycol, without any leakage at 70 °C, resulting in latent heats of 106.9 J/g and 118.6 J/g for each composite, respectively. A simple prototypical real-world application demonstrated that the prepared lauric acid and polyethylene glycol composites of mesopore enriched ZSM-5 could maintain their temperatures up to 27% and 22% lower than the ambient environment under solar heating, as well as up to 20% and 26% higher when solar heating stops. The presented findings indicate mesopore enrichment improves phase change material uptake in these low cost, non-toxic zeolitic shape stabilizers, hence making them good candidates as isolation materials to address energy loss during heating/cooling of household environments.

1. Introduction

Producing energy in a sustainable manner, without causing pollution is necessary to keep the Earth as a livable planet since conventional production processes result in significant environmental issues. Unfortunately, environmentally unfriendly energy sources are still generally preferred essentially as a consequence of costs, with significant improvements in efficiency and reductions in costs at scale required for green energy production alternatives [1–3]. Positively, a considerable volume of research has been reported with regard to addressing challenges and demonstrating improved performance of alternative energy systems, a viable solution is unlikely to be presented in the near future [4]. Therefore, reducing energy consumption is considered as a highly beneficial complementary approach, leading to a reduction in pollution resulting from excess energy use, but also a decrease the required energy-per-capita to be reached by the prospective alternative energy solution [5].

Heating is the most energy-consuming process according to International Energy Agency [6], with the use of facile techniques such as proper isolation leading to reductions in energy demand [7]. In this context, Phase Change Materials (PCMs) are interesting since their use in latent heat storage systems aims to keep the local environment within a constant temperature window [8–10]. Essentially, heat buffering process occurs via consumption of excess heat at temperatures above the melting point or the release of the stored heat at temperatures below the freezing point of the corresponding PCM. Since the latent heat of the PCM compensates for the heat change, the PCM should be selected in accordance with the aimed application. Thus, organic compounds with

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long alkyl chains such as fatty acids or polymeric materials like polyethylene glycols with different molecular weights can be utilized as useful PCMs since several of their derivatives offer different melting/freezing points [11]. Additionally, their non-toxicity, easy processability, relatively low-costs, and inert nature also make them desirable for such use [12].

The biggest challenge in the application of PCMs is keeping the molten form stationary at elevated temperatures [13]. To solve this issue, PCMs are embedded in shape stabilizers, which are mostly porous scaffolds such as zeolites, porous carbons, metal-organic frameworks, or porous polymers [14–19]. Aside from having excellent thermal stability and conductivity, a good shape stabilizer should be able to host a large amount of PCM to have better thermal properties since the latent heat is stored by those compounds. In the literature, mesopore-rich, hierarchically porous materials are promoted as useful shape stabilizers since the mobility of long alkyl chain PCMs can be limited for microporous materials due to steric hindrance, while keeping the adsorbed PCM in the shape stabilizer (i.e. avoiding leakage) can be hard for macroporous scaffolds due to poor capillary effect [20–23].

Since their discovery in the 19th century, zeolites have been used in various fields from water softening to high-temperature catalytic cracking [24–27]. Indeed, there are examples of zeolites used in latent heat storage systems as shape stabilizers. For instance, Goitandia et al. [23] compared Zeolite Y with various inorganic shape stabilizers possessing different pore sizes. They reported the domination of the mesoporous silica over ultra-microporous zeolite Y with an impregnation of 45% for hexadecane as PCM without any leakage [23]. In another study, a frequently used zeolite, namely ZSM-5, was used as the shape stabilizer for polyethylene glycol and showed a remarkable performance with a 50% impregnation and excellent thermal conductivity [28]. In this context, introduction of mesopores among the skeleton of ZSM-5 seems interesting. Thus, in this work, we present the effect of mesopore enrichment on PCM uptake for ZSM-5 and in turn the final latent heat storage properties. Comparing the shape stabilizing performance of a pristine microporous ZSM-5 with its post-synthetically modified versions possessing altered porosity allow us to investigate specifically the effect of pore size on latent heat storage properties via eliminating the effects of other variables (e.g. thermal conductivity difference) since the scaffolds have essentially the same backbone. Moreover, the choice of a molecular fatty acid (lauric acid (LA)) and a polymeric material (polyethylene glycol 4000 (PEG)) as PCMs facilitate the monitoring of structural differences such as size, shape and the number of heteroatoms on PCM loading and latent heat storage properties (see Fig. 1).

2. Experimental section

2.1. Materials

ZSM-5 type Zeolite (SiO\textsubscript{2}/Al\textsubscript{2}O\textsubscript{3} ratio of 469) was received from Acros Organics. Poly (ethylene glycol) with an average molecular weight 3500–4500 g/mol (PEG4000), lauric acid and potassium hydroxide pellets were purchased from Merck. Extra pure methanol (99.8%) was purchased from Tekkim Ltd. (Bursa/Turkey). Pure water was obtained using a Milli-Q Plus system. All chemicals were used without further purification.

2.1.1. General procedure for synthesis of ZSM-A

The preparation and characterization of ZSM-A and ZSM-B were explained in detail elsewhere [29]. The acidic treatment of ZSM-5 for the dealumination was performed in 2.5 M HCl (aq). Briefly, 50.0 g ZSM-5 was placed in a flask and 250 mL 2.5 M HCl (aq) was added. Following stirring for 6 h at 150 °C, the mixture was diluted two-fold with water. The solid was filtered and washed two times with 250 mL deionized water. Finally, the solid was dried at 120 °C overnight (under vacuum) and ZSM-A was obtained.

2.1.2. General procedure for synthesis of ZSM-B

Desilicated ZSM-5 was prepared by using nearly the same method above. 50.0 g ZSM-5 and 250 mL 2 M KOH (aq) were put in a flask, and the reaction performed 110 °C for 6 h. Subsequently, the solution was diluted with 250 mL deionized water, and the precipitate then filtered. The collected precipitate was twice washed with 250 mL water and dried at 120 °C overnight (under vacuum) to yield ZSM-B.

2.1.3. General procedure for preparation of PCM composites

Solvent assisted impregnation under vacuum was used to obtain ZSM-5, ZSM-A and ZSM-B-based PCM (LA and PEG) composites. By adjusting the total amount to 0.3 g for desired composite percentage (i.e. 0.12 g of PCM to 0.18 g shape stabilizer for 40% loading), the mixture of corresponding PCM and dried ZSM-X (in vacuum oven at 150 °C for 24 h) were added into 7.5 mL MeOH in 50 mL centrifuge tube. The charged tube was then placed in an ultrasonic water bath for 15 min and transferred to the vacuum impregnation system. Solvent removal was conducted gradually by reducing the pressure to 7–9 mbar first at room temperature by stopping the vacuum after 5 min and stirring for 5 min and repeated three times. Heating to 70 °C was then used to remove the solvent completely (in order to improve the impregnation). After solvent removal, the obtained powders were further dried in a vacuum oven for 24 h at 40 °C.

2.2. Methods

2.2.1. Instruments

Surface area measurements were performed by using the Quantachrome Instrument Nova4000e (Florida, USA). Fourier Transform Infrared (FTIR) spectroscopy with an ATR system was utilized for the analysis of PCM composites. Amount of embedded PCM in composite was measured with Shimadzu Corp. DTG-60H (TGA/DTA) instrument by heating samples up to 1000 °C with a rate of 20 °C/min. Thermal properties of the PCM composites were investigated using differential scanning calorimetry (DSC; Thermal Analysis TAQ2000). Heat storage performance of PCM composites were carried out at the rate of 10 °C/min with a heating and cooling cycle between 0 and 100 °C. Stability tests were also performed using DSC from 10 to 80 °C at a rate of 20 °C/min for each cycle. All thermal measurement was carried out under nitrogen atmosphere. Thermal behavior of PCM composites under solar power light were evaluated using an Oriel LCS-100 solar simulator at 0.404 W/cm\textsuperscript{2}. Temperature changes were measured by a Hanna thermometer attached to a K-type thermocouple.
2.2.2. Solar simulator experiments

0.1 g of ZSMB/LA65 and ZSMB/PEG70 were placed into a 2 mL polypropylene micropipette tip and temperature change was monitored using a K-type thermocouple (as inserted into the powder).

The prepared LA and PEG containing PCM composites were irradiated in the solar simulator at 0.404 W/cm$^2$ power. Time-temperature curves were obtained by recording the temperatures of environment (blank), neat ZSM-B, and the corresponding composite every 2 s during heating and cooling. The measurements were repeated three times, and the calculated average values were used for time-temperature graphs.

3. Results and discussion

3.1. Comparison of shape stability of ZSM-5, ZSM-A and ZSM-B PCM composites

The properties of zeolitic structures can change significantly after simple dealumination or desilication of the framework [30]. Such modifications might lead to hierarchical porosity, which results in altered diffusion dynamics as a consequence of removal of the corresponding network unit. In this context, introducing mesopore-rich hierarchical porosity to the backbone of ZSM-5 is expected to provide favorable properties in latent heat storage systems since mesoporosity is an asset for such scaffolds in latent heat storage systems [23], and also because pristine ZSM-5 has been previously highlighted as a promising shape stabilizer for PCMs due to its good thermal conductivity, even in its microporous form [28].

Alteration to the parent ZSM-5 after acidic and basic treatments were investigated thoroughly, and explained in our previous publication [29]. Briefly, dealumination of the highly siliceous ZSM-5 (Si/Al ratio = 938) was not successful, yet the acid exposure resulted in highly microporous ZSM-A. On the other hand, base treatment of the parent material to obtain ZSM-B yielded a hierarchically porous material possessing a dominantly mesoporous character and containing less silicon. Moreover, a lower specific surface area (BET) for ZSM-B (180 m$^2$g$^{-1}$) was recorded in comparison to ZSM-5 (356 m$^2$g$^{-1}$) and ZSM-A (359 m$^2$g$^{-1}$) probably due to increased mesoporosity (Figs. S2 and S3).

Shape stabilizers were loaded with LA and PEG through vacuum impregnation method by mixing ZSM-X (X = 5 or A or B) and corresponding quantities of PCMs dissolved in methanol at 40–80%, (w/w) ratios, followed by ultrasonication and solvent removal under reduced pressure. Composite ZSMX/PCM powders obtained are shown in Fig. 2A.

Since impregnating shape stabilizers with the highest amount PCM...
without any leakage is a highly desired feature to yield optimum latent heat storage properties [31], the non-exudative maximum PCM uptake capacities of the shape stabilizers were determined via performing a leakage test at elevated temperatures. Briefly, ZSMX/PCM composites were placed on a regular paper and heated in an oven at 70 °C for 30 min, and the area below the powder was checked whether it was wet or not. As can be seen in Fig. 2B, impregnation of ZSM-A and ZSM-5 with both PCMs at ratios of 40% and above resulted in leakage, whereas the ZSM-B composites did not leak any LA and PEG even at 65% and 70% impregnation, respectively. Moreover, the actual amounts of organic PCMs loaded into ZSM-B composites were verified by thermal gravimetric analysis (TGA) under ambient atmosphere. Impregnation ratios determined by the weight losses in ZSMB/LA and ZSMB/PEG due to decomposition of organic components (for LA around 220 °C and for PEG near 410 °C) were in good agreement with theoretical values (Figure S4 and Table S1). Based on these results, the composites formed from ZSM-B were conducted for further investigations.

The exudation stability of ZSMB/LA65 and ZSMB/PEG70 were evaluated by performing DSC experiments to confirm the results after the abovementioned macro-level leakage tests. Briefly, the samples were placed in a DSC sample holder and scanned for four times, during which the powders were removed from the DSC pan and put in the oven at 70 °C for 30 min after each cycle. The melting and solidifying patterns of ZSMB/LA65 and ZSMB/PEG70 were nearly the same for each cycle (Fig. 3A and C). Both melting and cooling curves of ZSMB/LA65 became slightly narrower with respect to the first cycle. This was probably due to the small amount of LA that evaporated from the composite powder at elevated temperatures during the DSC scan due to weak capillary forces [23,32]. Nevertheless, the change on the melting/solidifying enthalpies can be regarded as insignificant (Fig. 3B). On the other hand, such evaporation did not occur in case of ZSMB/PEG70 which is composed of a polymeric PCM (Fig. 3C and D).

Extended thermal stability tests of ZSMB/LA65 and ZSMB/PEG70 were employed by subjecting samples to 45 consecutive thermal cycles, during which the samples were transferred to a new pan after every 15th cycle (Table 1, Figure S5). The melting temperature of ZSMB/LA65 was detected to be only 1.5 °C lower than its initial state while its solidifying temperature remained nearly constant. On the other hand, solidifying temperature of ZSMB/PEG70 changed by only 0.9 °C while its melting temperature remained almost the same. Moreover, the melting and solidifying enthalpies also changed insignificantly, where they decreased by 4.3 J/g and 4.1 J/g, for ZSMB/LA65 and by 9.1 J/g and 5.7 J/g for ZSMB/PEG70. The negligible changes (% ΔH values; Table 1) showed that both of the prepared composites were stable after more than several melting/solidifying processes, indicating their long-term stability. Additionally, the infrared spectra of the composites before and after 45 thermal cycles also demonstrated the structural stability of the PCMs by showing nearly the same responses, particularly at ca. 2900 cm⁻¹ and 900 cm⁻¹ which are the vibrations of the C-H groups of the organic PCMs (Figure S6).
3.2. Thermal energy storage performance of ZSM-B based PCM composites

Latent heat storage properties of the prepared ZSM-B composites containing LA and PEG at impregnation ratios of 40% and above were investigated by DSC in order to monitor the effect of higher PCM uptake on latent heat storage properties (Fig. 4). The melting and solidifying temperatures of all composites was found to slightly shift to lower values with respect to pure LA and PEG. This was potentially caused by weak molecular interactions of PCMs with the porous shape stabilizer (e.g. capillary forces, hydrogen bonding interactions, etc.), which affect the required energies for phase transitions [33, 34]. Moreover, melting and

![Fig. 4. DSC curves of (a) ZSMB/LA and (b) ZSMB/PEG during melting and solidifying process; enthalpies of melting and solidifying of (c) ZSMB/LA and (d) ZSMB/PEG with different PCM contents.](image)

![Fig. 5. Heat loss percentages of (A) ZSMB/LA and (B) ZSMB/PEG composites.](image)
solidifying enthalpies of the prepared composites changed proportionally with respect to their PCM impregnation ratios, confirming the successful impregnation to ZSM-B (Fig. 4C and D). The melting enthalpies of composites prepared at highest impregnation ratios, ZSMB/LA65 and ZSMB/PEG70, reached 106.9 J/g and 118.6 J/g, respectively, which are comparable enthalpies reported for similar shape stabilized PCM composites (Table S3) [28,35–41]. Especially, the composite prepared in this work via impregnation of ZSM-B with PEG presented 35.5% higher latent heat than the composite prepared with ZSM-5 [28] (also with PEG 4000) confirming the potential of ZSM-B as a high capacity PCM shape stabilizer. Notably, the actual enthalpies of all ZSM-B composites were found to be lower than the theoretical values to different extents. A significant difference was observed for ZSMB/LA40, for which the LA molecules should be hindered in the core of the scaffold as a result of the low mass fraction. Therefore, the big deviation can either be due to the possible gap between the PCM and the exterior area of the scaffold [42,43] or altered phase change behavior of the LA in such highly confined spaces [44].

Theoretical enthalpies of ZSMB/PCM composites were calculated by using Eq. (1). The actual enthalpies of pure PCMs (100% PCM content) were accepted as their theoretical enthalpies to make comparison with the ZSMB/PCM composite system.

\[ \Delta H_{\text{theo.}} = (1-w) \Delta H_{\text{m/s}} \]  

(1)

where “\( \Delta H_{\text{theo.}} \)” is theoretical latent enthalpy, “\( \Delta H_{\text{m/s}} \)” is the melting or solidifying enthalpy of pure PCMs, and “w” denotes the content of ZSM-B.

On the other hand, the variations between actual and theoretical values were less significant for the composites of PEG. Presumably, the higher heteroatom content (oxygen) of PEG or its bulkier polymeric structure (Mw: \( \approx 4000 \text{ g/mol} \)) result in strong interactions with the skeleton of zeolitic framework might reduce the strength of the capillary forces to confine the material through the core of the scaffold as deep as LA. Nevertheless, such interactions should also be limiting the (re)crystallization of the molten phase in the low loadings, hence yielding lower actual enthalpies than the theoretical [34].

The differences between melting and solidifying enthalpies (heat loss) of the prepared composites were calculated to monitor their relative latent heating and cooling performances (Fig. 5). The heat loss calculated for pure LA was considerably higher than the heat loss calculated for its composites indicating that the capillary forces of the shape stabilizer limits its evaporation [45] at elevated temperatures. Indeed, the composite with the lowest loading of LA, ZSMB/LA-40, showed nearly minimal heat loss as a consequence of the hindrance of such low amount of LA through the network. In case of the composites of the polymeric PEG, the heat loss occurs due to energy differences between melting and crystallization, which even occurs in case of the neat PEG [22,41].

Fig. 6. Thermal behavior of (A–B) ZSMB/LA65 and (C–D) ZSMB/PEG70 composites under solar simulator irradiation.
3.3. Latent heat storage performances of ZSM-B based PCM composites under solar irradiation

The composites containing the highest amount of the PCMs were subjected to heating-cooling cycles by using a solar simulator as a prototype of real-world applications. Temperatures of ZSMB/LA65 and ZSMB/PEG70 composite powders were recorded under solar irradiation and also after the simulator was switched off. Even though the composites demonstrated nearly homogeneous heat buffering until their saturation points, maximum performances were highlighted in Fig. 6 (see Table S4 for more information).

The temperatures of both composites were compared to the temperature of the environment in the absence of the composites and the temperature of PCM-free ZSM-B powder as references under the same solar irradiation conditions. Accordingly, when the environment control temperature reached 55.6 °C, the temperature of ZSMB/LA65 powder was only 49.8 °C, providing 27% cooling. At this point, the temperature of neat ZSM-B powder was 66.9 °C showing that the medium of ZSMB/LA65 composite powder was 39% cooler. Similarly, the temperature of ZSMB/PEG70 composite powder was 38.5 °C, at the time when the temperature of the environment reached 47.5 °C meaning that ZSMB/PEG70 composite powder keeps system 22% cooler than the ambient temperature.

Thermal behavior of both composites after the solar simulator was switched off was also evaluated. For ZSMB/LA65, at the time the environment temperature was 32.9 °C, the temperature of ZSMB/LA65 was 39.6 °C which corresponds to a 20% warmer medium than the environment. Similarly, ZSMB/PEG70 composite powder allowed a 26% warmer medium since its temperature was 40.9 °C while it was 32.4 °C for the blank. The results indicated that ZSMB/LA65 and ZSMB/PEG70 successfully perform latent heat storage by providing significantly cooler environments during solar heating and significantly warmer environments when the solar heating is disrupted through corresponding phase transitions.

4. Conclusions

Desilication was applied to a microporous zeolitic shape stabilizer, ZSM-5, in order to yield a mesopore rich hierarchically porous backbone in the framework to increase the amount of PCM impregnation. Even though the specific surface area dropped after the treatment, mesopore rich ZSM-B was able to hold nearly twice the amount of PCMs inside the framework with respect to other microporous analogues. The optimum non-exudative composites of ZSM-B containing 65% LA and 70% PEG were able to store latent heat up to 106.9 J/g and 118.6 J/g, respectively. Moreover, the solar simulator tests of ZSMB/LA65 and ZSMB/PEG70 indicated the latent heat storage potential of the composites via buffering the temperature change up to 27% and 22% when the solar simulator was on as well 20% and 26% during the cooling, respectively. Our results demonstrate that the introduction of mesopore rich hierarchical porosity to microporous ZSM-5 improved the PCM impregnation capacity, hence the latent heat storage properties of the zeolitic shape stabilizer. Additionally, the solar simulator tests indicate how promising the latent heat storage systems are to keep an environment in a constant temperature to save energy spent for heating and its significant lab-to-fab potential via hereby presented combination of non-toxic PCMs with low-cost shape stabilizers.

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Declaration of competing 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.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.solmat.2020.110677.

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