Theoretical formulation and numerical simulation of thermal performance enhancements for cascade thermal energy storage systems

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Abstract: Cascaded Thermal Energy Storage (CTES), a term that refers to a thermal energy storage system with multiple phase change materials (PCMs), has been suggested as a solution for heat transfer reduction through the process of heat exchange by reducing temperature differences. The PCMs used are thus paraffin waxes with different melting temperatures. A numerical simulation was made to determine the optimum length of a CTES system compared with use of a single PCM. The enthalpy-porosity theory was utilised to simulate the phase transition of the PCM, and the simulations then used to mimic the charging and discharging of thermal energy storage of optimum length at different heat transfer fluid flow rates. The results indicated that heat transfer can be greatly enhanced, and melting and solidification time significantly reduced, by using multiple PCMs as compared with using a single PCM.

1. Introduction
In recent years, energy consumption has expanded dramatically because of global economic development. As a result, there has also been an increase in greenhouse gases emission and environmental pollution. These factors have led to increased efforts improve energy efficiency so as to reduce overall energy consumption. Thermal energy storage (TES) is an effective method of balancing energy demand and energy supply, allowing the energy system to be more stable and efficient by storing excess thermal energy during high production hours and using it during low production hours [1].
TES can be categorized into three main divisions: storage of sensible heat, storage of latent heat, and storage of thermochemical energy [2]. Storage of latent heat is an attractive technique due to its ability to supply high density energy storage over a relatively constant temperature range analogous to the phase change temperature of the storage material [3]. The materials used in the storage of latent heat are referred to as phase change materials (PCMs), and such PCMs have been studied in several different applications, including solar power plant energy saving buildings [4], systems of waste heat recovery [5], the cooling of electronic devices [6], greenhouses [7], and solar cookers [8].
During charging, the PCMs absorb heat and their temperatures rise; thus, PCMs perform as sensible storage materials. Solid PCMs thus convert to liquid PCMs and the thermal energy is conveyed in this change. The PCMs absorb thermal energy at even small differences in temperatures and store 5 to 14 times more energy.
than sensible storage materials [9]. The process reverses on release of heat during the discharge process and the PCM liquids completely revert into solids.

The high capacity of storage for small temperature differences is the major advantage of latent heat storage over storage of sensible heat; however, this does not mean that storage of latent heat cannot be beneficial at wider temperature differences. Nevertheless, it is reasonable to split a wide temperature difference into smaller divisions, each wrapped in a different PCM with an appropriate phase transition temperature [10]. A combination that uses multiple PCMs with various melting temperatures is called a Cascaded Thermal Energy Storage (CTES) system. Another objective for using CTES is that the charge and discharge time is limited in the most common functional case, and the heat must be released or stored rapidly. During the charging of storage systems with single PCMs, the heat transmits to the PCM from the heat transfer fluid (HTF) very rapidly. Thus, the temperature of the HTF decreases, and the temperature difference between the HTF and the PCM decreases, leading to reduced heat transfer at the storage end. Consequently, the PCM melts quickly at the part where the HTF enters the storage, but very slowly at the storage end where the HTF exits the storage. The problem is the same for discharging: the PCM at the storage ending might not be utilised in raising the HTF's temperature. This problem can be solved by creating a CTES [11]. Farid et al. [12, 13] experimentally and numerically constructed a model that included several cylindrical capsules filled with three PCMs with different melting points; the results showed enhancements in both charging and discharging. Gong and Mujumdar [14, 15] evolved a finite element model for a high-temperature thermal storage system of a slab [15] and a tube [14] that used five PCMs, finding a considerable improvement in heat transfer compared with the use of one PCM. Wang et al. [16] experimentally analysed the charging of a cylindrical thermal storage capsule containing multiple PCMs, discovering that the melting process was 15 to 25% faster than in systems containing one PCM. Shaikh and Lafdi [17] conducted a simulation to study the effects of utilising various arrangements of multiple PCMs slab configurations with various melting points, and they demonstrated significant enhancements in heat transfer. Fang and Chen [18] investigated a shell and tube storage system with multiple PCMs, noting that presence of an optimal assembly of multiple PCMs offered extreme thermal energy charging rates. Wang et al. [19, 20] numerically studied a heat exchange device with a zigzag arrangement including multiple PCMs throughout the discharge [19] and charge [20] process; the results indicated that intensified charging and discharging in comparison with the use of one PCM.

The present paper thus aims to numerically investigate the optimum length of CTES, as compared with a TES with one PCM, and then to perform simulations for charging and discharging CTES with this best length.

2. Physical problem

To determine the optimum length of a CTES, a comparison between Single-stage Thermal Energy Storage (STES) and CTES for different lengths was completed. A STES was formed with only PCM2 (Figure 1), while the CTES was constituted of three PCMs along the flow direction of HTF: PCM 1, 2, and 3 (Figure 2). The PCMs used were paraffin waxes and their thermo-physical properties are recorded in Table 1 [21]. The melting temperature of PCM2 (in STES) has an intermediate value between PCMs 1 and 3, to give comparability between STES and CTES. The HTF is air, and its properties are listed in Table 2 along with other relevant system parameters. The CTES with the best length was then studied with regard to the charging and discharge process. For the charging process, a PCM with a lower melting temperature was positioned at the end of the storage to ensure that the temperature difference between the HTF and PCMs was sufficiently large to ensure melting of all PCMs; this arrangement was inverted during the discharging process, as shown in Figure 3.
Figure 1. STES

Figure 2. CTES

Figure 3. PCM arrangement

Table 1. Thermo-physical properties of PCMs [21]

| PCMs       | PCM1     | PCM2     | PCM3     |
|------------|----------|----------|----------|
| Melting temperature [K] | 331      | 336      | 340      |
| Solidus Density [kg/m³]  | 840      | 850      | 860      |
| Liquidus Density [kg/m³] | 765      | 766      | 767      |
| Specific Heat [J/kg K]   | 2763     | 2817     | 2871     |
| Thermal Conductivity [W / m K] | 0.21  | 0.212    | 0.214    |
| Dynamic Viscosity [kg/m s] | 0.0116  | 0.01215  | 0.0127   |
| Thermal expansion coefficient [1/K] | 0.00031 | 0.000305 | 0.0003   |
| Latent Heat [J / kg]     | 267670   | 270715   | 273760   |
| Solidus Temperature [K]  | 321      | 326      | 330      |
| Liquidus Temperature [K] | 335      | 340      | 344      |

Table 2. System parameters

| HTF properties                      | 1.225               | System dimensions |
|-------------------------------------|---------------------|-------------------|
| Density [kg/m³]                     | 1006.43             | h₁ = 20mm, h₂ = 25mm |
| Specific Heat [J/kg K]              | 0.0242              | L₁ = L₂ = L₃      |
| Thermal Conductivity [W / m K]      | 1.7894e-05          |                   |
| Dynamic Viscosity [kg/m s]          | 373                 |                   |
| Inlet temperature during charge [K] | 294                 |                   |
| Inlet temperature during discharge [K] |                   |                   |
3. Numerical model

A simulation of the melting and solidification processes of PCMs was applied using the enthalpy-porosity method [22]. In this method, there is no explicit tracking for the liquid-solid interface; instead, there is a mixed liquid-solid region, referred to as a mushy zone. This zone is modelled as a porous medium, with a porosity specified as equivalent to liquid fraction (β). The value of β is given by [23]

\[
\beta = 0 \quad \text{for } T < T_{\text{solidus}}
\]

\[
\beta = \frac{(T - T_{\text{solidus}})}{(T_{\text{liquidus}} - T_{\text{solidus}})} \quad \text{for } T_{\text{liquidus}} < T < T_{\text{solidus}}
\]

\[
\beta = 1 \quad \text{for } T > T_{\text{liquidus}}
\]

where T is the local temperature. Thus, the velocity will be affected as follows:

\[ V = V_1 \quad \text{in the liquid phase} \]

\[ V = \beta V_1 \quad \text{in the mushy zone} \]

\[ V = 0 \quad \text{in the solid phase} \]

where \( V \) is the superficial velocity and \( V_1 \) is the actual velocity [22]. A Boussinesq approximation is utilised to simulate the natural convection in the PCM, and thus the density of the PCM varies with temperature [24]:

\[
\rho = \rho_0 \left[ 1 - \gamma(T - T_0) \right]
\]

where \( \rho \) is the local density of the PCM, \( T_0 \) and \( \rho_0 \) are the operating temperature and density, and \( \gamma \) is the thermal expansion coefficient. Assuming the flow of the liquid PCM is Newtonian, incompressible, and laminar, the governing equations are continuity, momentum, and energy. The continuity equation can therefore be illustrated as [25]

\[
\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \vec{V}) = 0
\]

where \( t \) is the time. The pressure losses of flow produced by the existence of the solid PCM can be estimated using the source terms in the momentum equation [26]:

\[
\rho \left( \frac{\partial \vec{V}}{\partial t} + (\nabla \cdot \vec{V}) \right) \vec{V} = \mu \nabla^2 \vec{V} - \nabla P + \vec{S}
\]

where \( \mu \) is the viscosity of the PCM, \( P \) is the pressure, and the vector \( \vec{S} \) is a global source term given by the following form:

\[
\vec{S} = \frac{(1-\beta)^2}{(\beta + \varepsilon)^3} A_{mush} \vec{V} + \rho \vec{g} \gamma(T - T_0)
\]
The first term on the right-hand side appears due to the existence of solid PCM in the mixed region, where $\epsilon$ is set as a small number (less than 0.001) to prevent division by zero [27]. $A_{mush}$ is the mushy zone constant, which acts as a damping factor of velocity through the solidification of the PCM. Its value affects the PCM melting rate, and here it is set to $10^3$ kg/m$^3$s [28]. The second term is the Boussinesq approximation, which is essential for modelling the natural convection in the liquid phase of the PCM. The vector $\vec{g}$ is the gravitational acceleration, which is equal to 0 m/s$^2$ in the x-direction and -9.81 m/s$^2$ in the y-direction. The energy equation can be described as [20]

$$\frac{\partial}{\partial t}(\rho_h) + \nabla \cdot (\rho \vec{V} h) = \nabla \cdot (k \nabla T) + S_h$$

(7)

where $h$ is the sensible enthalpy, which can be expressed as [29]

$$h = h_{ref} + \int_{T_{ref}}^{T} C_p dT + \beta L$$

(8)

where $h_{ref}$ is the enthalpy at the reference temperature $T_{ref}$, and $C_p$ and $L$ are the specific heat at a constant pressure and the latent heat of the PCM. $S_h$ is the energy source term, given by [20]

$$S_h = \frac{\partial (\rho \Delta h)}{\partial t} + \nabla \cdot (\rho \vec{V} \Delta h)$$

(9)

The flow of HTF is assumed to be turbulent flow; to consider the turbulence effect, the $(K - \epsilon)$ model [28] was thus implemented in the numerical model. Two-dimensional numerical simulations were made with ANSYS Fluent. For the grid independence solution, three different grid densities were tested, with 21,600, 84,000, and 336,000 elements, respectively. The results showed that the 84,000 elements density was most suitable, because it represented the best compromise between solution accuracy and computational cost (see Figure 4). A comparison with the work in [30] was undertaken for model validation: the same characteristics were used, and the two models gave similar results (see Figure 5), demonstrating good agreement between them.

*Figure 4. The grid independence solution.*
4. Results and discussion

4.1. Best length of CTES

In order to determine the best length of CTES containing the selected PCMs, a comparison between the CTES and STES was carried out for different lengths (750; 1,000; 1,200; 1,500; and 2,000 mm) during the discharge process. Figure 6 illustrates the percentage of improvement in solidification time of each CTES over STES, categorised by length. As shown, increases in length lead to increases in the percentage of improvement in solidification time to a length of 1,200 mm, where the highest increase in the percentage of improvement is seen; after that, the percentage of improvement decreases. This means that a length of 1,200 mm is optimal for using these PCMs.

Figure 6. The best length of CTES.
In Figure 7, the liquid fraction contours for both STES and CTES at the best length can be seen at the same instant of time (4,000 s), for both charging and discharging processes. The red region is the liquid phase, while the blue region is the solid phase. Alternating colours represent the mushy zone. Clearly, the charging and discharging processes are faster and more uniform in CTES compared with STES, as CTES ensures
almost uniform temperature difference between the HTF and PCMs along the storage. Figure 8 also shows a comparison of the average liquid fractions; it can be seen that the liquid fraction tends to rise faster with CTES during the charging process (Figure 8 (a)), where the melting time for STES is 6,860 s, while for CTES it is 4,595 s. Similarly, for the discharge process (Figure 8 (b)), the solidification time for STES was 4,685 s, while for CTES it was 4,257 s. This means that CTES significantly improves heat transfer for both charging and discharging processes, although more improvement is seen for charging process. This finding is critical, and common in all literature dealing with this subject. The main cause of this phenomenon is that during the discharging process, the upper layers of the PCM solidify first, and thus act as additional resistance to heat transfer, reducing any enhancement. However, in the charging process, the upper layers of the PCM melt first, enhancing the heat transfer and allowing natural currents to be developed. As shown in Fig. 8, the effect of CTES is clearly shown in the final stages of the charging and discharging processes; this is because the temperature difference between the PCM and HTF reduces at the final stages in STES, while CTES works to increase this difference and accelerate heat exchange.

4.2. Influence of HTF velocity variation on CTES
The influence of HTF velocity variation on outlet air temperature for the charging and discharging processes is illustrated in Figure 9. During the charging process (Figure 9 (a)), the minimal HTF velocity (v=1 m/s) was used to give the minimal outlet temperature and maximum difference between inlet and outlet temperatures; this is because at lower velocity, the air has more time to flow over the PCMs and enable more heat exchange. The maximum HTF velocity (v=15 m/s) similarly gives the maximum outlet temperature, ensuring minimal difference between inlet and outlet temperatures due to the lower amount of time that the passing air is exposed to PCMs. For the discharging process (Figure 9 (b)), the maximum velocity again gave a minimal difference between outlet and inlet temperatures, and the minimal velocity give the minimal difference between outlet and inlet temperatures, for the reasons mentioned above. Each curve ends when melting (Figure 9 (a)) or solidification (Figure 9 (b)) is complete; this indicates that the HTF velocity affects the melting and solidification time, as shown in Figure 10, where it is noted that increased velocity leads to a significant reduction in both melting and solidification time. It should be noted that the melting process at (v=1 m/s) is never completed; this velocity is not sufficient to melt the PCMs, and thus higher velocities were used throughout.
5. Conclusions
The numerical investigation into thermal performance of CTES was accomplished; the enthalpy-porosity theory was applied to simulate the phase change of the PCMs, and the comparison between STES and CTES for different lengths was established, with the results indicating that the length of CTES was 1,200 mm for the selected PCMs. The CTES with the best length was seen to be greatly enhanced in terms of heat transfer for both charging and discharging processes, especially in the final stage, and the melting and solidification times were significantly reduced. Simulations of the charging and discharging processes for a CTES with the best length under the influence of HTF velocity variations of outlet air temperature were accomplished, and the effects on the melting and solidification times were studied. The results showed that increased velocity reduced the difference between air inlet and outlet temperatures, thus reducing the process time.

Figure 9. The influence of HTF velocity variation on outlet air temperature for (a) charging and (b) discharging process.

Figure 10. The influence of HTF velocity variation on melting and solidification time.
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