Solidification of Additive-Enhanced Phase Change Materials in Spherical Enclosures with Convective Cooling

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Abstract

Solidification of eicosane with and without nanoadditives is experimentally investigated in spherical enclosures subject to convective cooling in water and air. The effects of additive volume fraction and external convective cooling conditions (i.e., the heat transfer medium, subcooling, and flow velocity) on the solidification process are examined. The results are compared with a conduction-controlled thermal network model accounting for the enclosure and PCM resistances, as well as the convective subcooling. The experimentally determined solidification time is found to be consistently lower than the model prediction, likely due to asymmetric and dendritic solidification, as well as natural convection inside the enclosure and possible thermocouple position errors. A simple correlation is proposed to predict the solidification time of a phase change material (PCM) in a spherical enclosure subject to convective cooling based on the same enclosure subject to a constant temperature boundary. Results show that the solidification time decreases with the volume fraction of nanoadditives due to the improved PCM conductivity. In addition, the nanoadditives are found to be more effective for solidification in water than in air, due to the large air-side convective resistance that does not benefit from improving PCM conductivity.

Keywords: Phase change materials, nanoadditives, solidification, convective cooling

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1. Introduction

Phase change materials (PCMs), due to their ability to transfer large amounts of heat in a narrow temperature range, are well suited for thermal management systems with mismatching supply and demand thermal loads as well as thermal energy storage systems. Among various configurations, melting and solidification in spherical enclosures is encountered in many applications such as chemical reactors [1], packed bed latent heat energy storage [2], spray freezing [3,4] and advanced PCM slurry heat transfer fluids [5,6].

For several decades, many investigators have modeled the solidification in spherical enclosures. Tao [7] developed a numerical model to investigate the effect of external convection and sensible heat removal from the solidified layer during spherical freezing. Pedroso and Domoto [8] solved the problem of inward spherical solidification subject to a constant wall temperature using a perturbation method. Ismail and Henriquez [9] developed an analytical model for inward solidification in a spherical enclosure subject to either the constant temperature or the convection heat transfer condition on the external surface of the spherical shell. Hill and Kucera [10] utilized a semi-analytical procedure to investigate the inward solidification in a spherical container with the effect of radiation. Wu et al. [11] used the small-time series expansion technique to solve the unsteady heat equation where the effect of curvature on the interface temperature is included. Lin and Jiang [12] used a quasi-steady model for solidification in spherical enclosures with an additional term to account for the transient effect on the temperature distribution. Riley et al. [13] presented an analytical study for inward freezing of a sphere with a two-region analysis to accommodate the singularity at the center of the sphere. McCue at al. [14] developed a model for spherical solidification of superheated PCMs under a constant temperature boundary condition. It was found that, at the first stage of solidification, the temperature in both solid and liquid phases depends on the inverse Stefan number, but at later times the inner liquid core contributes insignificantly to the location of the solidification front [14].

In addition to the mathematical models, experimental and numerical studies have also been performed to investigate the solidification of spherical PCM. Chan and Tan [15] experimentally studied the effects of cooling water temperature and initial PCM superheating on the solidification rate of \( n \)-hexadecane in spherical shells. It was shown that the solidification rate increases by decreasing the cooling water temperature, while the effect of initial superheating of
the liquid PCM is negligible. In a related work, Assis et al. [16] experimentally studied the solidification of a commercial PCM in spherical capsules that were cut at various times to measure the thickness of the solid layer. The morphology of the solidification front was compared with the predictions of their numerical model and a good agreement was observed. A similar experimental study was conducted by Eames and Adref [17] to characterize the freezing and melting processes of deionized water. The solidification rate was determined using the changes in air pressure in the PCM enclosure upon phase change, where it was found that 90% of the total phase change occurs in the first 70% of the total phase change time. Ismail et al. [9] developed a conduction-controlled model to numerically investigate the effect of spherical shell material and size, wall thickness, initial PCM temperature, external fluid temperature, and Biot number on solidification of PCMs in spherical enclosures. The thermal conductivity of the shell material was found to significantly affect the solidification time.

In order to take full advantage of the high energy density of PCMs, the low thermal conductivity of most PCMs must be overcome. Efforts have been made to improve the phase change rates in spherical PCM enclosures by using extended surfaces [18] and dispersion of metallic beads within the PCM [19]. Experimental results by Koizumi [18] showed that insertion of copper fins inside a spherical container reduces the \( n \)-octadecane melting time by 33%. Ettouney et al. [19] conducted experiments to assess the effect of stainless steel beads dispersed within paraffin wax on melting and solidification rates in a 30 mm spherical shell. It was found that total phase change time is reduced by 15%, equivalent to the PCM mass replaced by metallic balls, suggesting negligible improvement for the enclosure size employed. Recently, carbon-based nanoadditives, such as graphite nanofibers [20], exfoliated graphite nanoplatelets (xGnPs) [21,22], and ultrathin graphite foams [23] have been utilized to enhance the thermal conductivity of PCMs. Kalaiselvam et al. [24] conducted experiments on solidification of six types of nanoadditive-enhanced PCMs in spherical capsules under constant wall temperature conditions and reported up to 22% decrease in solidification time compared to that of the pure PCM. They also found that increasing the mass fraction of nanoadditives beyond 7% does not result in further reduction in solidification time. Chandrasekaran et al. [25] experimentally studied solidification of water-based copper oxide nanofluid in a spherical capsule subject to constant wall temperature conditions and reported 35% reduction in solidification time compared to pure water. It was also shown that 50% of the PCM mass is solidified in only 25% of the total
freezing time. In a similar study, Kumaresan et al. [26] conducted experiments on solidification of water-based carbon nanotube nanofluid encapsulated in a spherical shell and observed 20% reduction in solidification time compared to pure water. Altohamy et al. [27] reported experimental results on solidification time of water-based alumina nanofluid in a spherical capsule subject to a constant wall temperature where up to 30% reduction in solidification time was achieved for 2% additive volume fraction.

Review of the literature reveals that most of the analytical and in particular experimental studies have been focused on solidification under constant temperature conditions. However, the majority of the aforementioned applications pertaining to spherical solidification involve convective heat transfer between an external heat transfer fluid and the PCM. One fundamental difference between the constant temperature and convective boundary conditions is that for the former the PCM itself presents the major thermal resistance prevailing the heat transfer. However, for convective cooling, the thermal resistance due to external convection may be comparable or even greater than the PCM thermal resistance. The relative magnitude of the internal (PCM) and external resistances determines the proper choice of heat transfer enhancement to be employed. In this work, solidification of eicosane with xGnP of varying volume fractions in spherical enclosures of 38 and 50 mm is experimentally investigated under constant temperature (submerged in water) and convective cooling (exposed to air flow) boundary conditions. The effects of additive volume fraction, convective subcooling, and external heat transfer coefficient on solidification time are studied. A conduction-controlled thermal network model is developed for the solidification process, and the model predictions are compared with the experimental results. A correlation is proposed to predict the PCM solidification time in a spherical enclosure subject to convective cooling as a function of the Biot number and the solidification time of the same enclosure with a constant wall temperature.

2. Methodology

2.1 Solidification of PCM

As one of the most popular PCMs, eicosane has been used in many experimental studies on PCM thermal properties and phase change behavior [21,28–31]. In the present study, n-eicosane (99%, Sigma Aldrich) is used as the PCM, with a melting point of 36.4 °C, heat of fusion of
247.3 kJ/kg, solid density of 815 kg/m³, liquid density of 780 kg/m³, solid specific heat of 1.92 kJ/(kg·K), and liquid specific heat of 2.46 kJ/(kg·K) [32]. Two sets of solidification experiments were conducted for nanoadditive-enhanced eicosane in spherical enclosures under constant wall temperature and convective cooling conditions, respectively. The constant temperature tests were performed by submerging the PCM enclosures in a temperature-controlled water bath with forced convection (Fig. 1a). For convective cooling tests, the PCM enclosures were placed in a wind tunnel with precise control of the air velocity (Fig. 1b). Stainless steel spherical enclosures with outer diameters of 50 mm and 38 mm and respective wall thicknesses of 2 mm and 3 mm were employed. A thermocouple rake was inserted into the spherical enclosure to measure the temperature distribution of the PCM during solidification. The insets of Fig. 2a and 2b show the thermocouple configurations in the 38 mm diameter sphere and the 50 mm diameter sphere, respectively. Two T-type thermocouples (labeled as TC1 and TC2 in the inset of Fig. 2a) were used in the thermocouple rake in the 38 mm diameter sphere: one in the geometric center of the sphere, and one on the inner surface of the shell. Five T-type thermocouples (labeled as TC1 to TC5 in the inset of Fig. 2b) were used in the thermocouple rake in the 50 mm diameter sphere: TC1 and TC5 were located at the center and on the inner surface of the sphere, respectively; TC2, TC3, and TC4 were located between TC1 and TC5 with spacing of $s_{12} = s_{23} = s_{34} = 6 \text{ mm}$, and $s_{45} = 5 \text{ mm}$. The thermocouples were connected to an Omega Engineering 8-channel OM-CP data logger to record the temperature history. The spherical enclosures filled with PCM are initially submerged in a hot water bath at a temperature of 60°C (well above the melting point of tested PCM) to keep the PCM in liquid state. After thermal equilibrium is reached the PCM spheres are moved to the chiller/wind tunnel to conduct the water/air cooling tests. Once the thermal boundary condition is established the data logger starts to record the temperature history at various locations within the PCM.
Figures 2a and 2b show the example temperature histories during solidification of PCM subject to air cooling at $T_{air} = 25 \, ^\circ C$ and $U_{air} = 25 \, m/s$ in 38 mm and 50 mm diameter spheres, respectively. As shown in Fig. 2a, the PCM is initially superheated, and its temperature decreases quickly after the cooling process starts at $t = 0$. The complete solidification time is determined using the temperature measurement at the inner surface (TC2) and the center (TC1). After cooling starts ($t = 0$), the temperatures at both the inner surface and the center of the sphere drop rapidly. The inflection of the temperature profile at the inner surface (measured by TC2) represents the onset of solidification. The temperature at the center (TC1) also shows an inflection shortly after cooling starts, then remains fairly flat for an extended period of time, and starts to drop rapidly again (showing a second inflection in the curve), indicating the end of phase change. The solidification time is calculated as the time between the inflection of the inner surface temperature (measured by TC2) and the second inflection of the center temperature (measured by TC1), as labeled in Fig. 2a. The greater number of thermocouples in the larger sphere allows for the determination of partial solidification time, where solid volume fractions of 100%, 98%, 86%, and 52% can be measured based on the readings of TC1 to TC4, respectively.
Figure 2. (a) Time history of the temperature of the inner surface and PCM temperature at the center temperatures during solidification of eicosane in spherical enclosure with an outer diameter of 38 mm, (b) Temperature history during solidification of eicosane at various radial locations for solidification of eicosane in spherical enclosures with an outer diameter of 50 mm. Both experiments are performed in air at $T_{\text{air}} = 25 \, ^{\circ}\text{C}$ and $U_{\text{air}} = 25 \, \text{m/s}$. Insets show the configuration of thermocouples.

2.2 Thermal conductivity measurement of PCM

While the heat of fusion of paraffin-based PCMs is reasonably high (~150-250 kJ/kg), the thermal conductivity of these PCMs is usually low (~0.1-0.4 W/(m·K)). The low conductivity of the PCM may lead to large conductive resistance for heat transfer. Experimental studies have shown enhancement in effective thermal conductivity of PCMs with a variety of nano-additives. One of the objectives of the present work is to investigate how nanoadditive concentration affects the effective thermophysical properties of the additive-enhanced PCM. Exfoliated graphite nanoplatelets (xGnPs) from XG Science was used in this study as a thermal enhancement agent dispersed in eicosane, due to its high thermal conductivity and relative low cost and density. The xGnPs used in this work were of 25 μm average diameter and 15 nm average thickness with an effective thermal conductivity of about 98 W/(m·K) [33].

The thermal conductivity of the additive-enhanced PCM was measured using the hot wire method that is suitable for conductivity measurement of low thermal conductivity materials [16-18]. The hot wire method is based on transient one-dimensional cylindrical heat conduction in
the tested material. The thermal conductivity of the tested material is determined based on one-dimensional transient cylindrical heat condition as

\[ k = \frac{q}{4\pi L} \frac{d \ln(t)}{d\Delta T} \]  (1)

where \( q \) is heat flow rate applied to the wire, \( L \) is the length of the wire, \( t \) is the time, and \( \Delta T \) is the transient temperature increase of the hot wire after being subjected to the heat flow \( q \), defined following \( \Delta T(t) = T_{\text{wire}}(t) - T_{\text{wire}}(0) \), where \( t = 0 \) represents the start of the heating.

Figure 3a shows the schematic of the electric circuit to measure thermal conductivity and Fig. 3b shows the zoomed-in schematic of the test section. In the setup, an ultrathin Pt wire with a length of 160 mm and a diameter of 20 μm (aspect ratio = 8000) is used to eliminate the effect of conduction along the wire and ensure one-dimensional heat transfer. The test chamber is made of two concentric cylinders of diameter of 45 mm and 89 mm, respectively. The tested PCM is filled in the inner cylinder where the hot wire is placed in its centerline. The outer cylinder is filled with water and is connected to a chiller for temperature control. The transient temperature of the hot wire (\( \Delta T \)) is determined by measuring the electrical resistance of the hot wire using a Wheatstone bridge circuit, with high precision resistors (i.e. \( R_1 \), \( R_2 \) and \( R_3 \)) at three legs of the bridge, and the hot wire with a potentiometer at the other leg. The potentiometer is used to zero the bridge circuit at the beginning of the tests. In addition to the bridge, the circuit consists of a power supply, a relay connected to a wave generator and an oscilloscope. The input voltage, \( V_s \), and the bridge voltage, \( V_b \), are measured with an oscilloscope. The wave generator is used to synchronize the measurement of the bridge circuit and connecting the circuit. A square wave is pre-set in the wave generator. When the square wave reaches its up edge, the relay will close the loop and meanwhile the signal will be sent to the oscilloscope to start recording the bridge voltage as a function of time. The dynamic electrical resistance of the Pt wire during the tests is calculated using the dynamic voltage recorded by the oscilloscope as following

\[ \Delta R_{\text{w}}(t) = \frac{R_3}{R_1 + R_2} - \frac{V_{b}(t)}{V_s} - \frac{R_1}{R_1 + R_2} \]  (2)

In the present work, the resistance of the three resistors \( R_1 \), \( R_2 \) and \( R_3 \) are set to the same value of 100 Ω. The dynamic temperature of the Pt wire is then calculated as
\[ \frac{\Delta R_w(t)}{R_{w0}} = \alpha \Delta T(t) \]  

(3)

where \( \alpha \) is the temperature coefficient of resistance (\( \alpha = 0.003927 \text{ K}^{-1} \) for platinum [34]). The thermal conductivity of the sample is then determined using Eq. (1) with the transient temperature increase of the hot wire \( \Delta T \). The hot wire setup is validated with two common materials, i.e., air and soybean oil. The measured conductivity of air is 0.0271 W/(m·K) at 25°C that is within 5% relative error with the tabulated value of 0.0257 W/(m·K), and the measured conductivity of soybean oil is 0.147 W/(m·K) at 25°C that is within 7% relative error with the tabulated value of 0.158 W/(m·K) [35].

Figure 3. (a) Schematic of the electric circuit of hot wire method to measure thermal conductivity. (b) Detailed view of the test section of the hot wire setup.

2.3 Thermal network model

In this section, a simple thermal network model is developed for solidification in a spherical enclosure. Figure 4 shows the schematic of PCM solidifying in a spherical enclosure with an outer radius of \( r_o \) and inner radius of \( r_i \). The basic assumptions of the model are:
i) PCM is a homogeneous material with a constant solidification temperature $T_m$;

ii) Liquid PCM is initially at melting temperature $T_m$;

iii) The sensible energy content of PCM is negligible ($\text{Ste} \ll 1$); and

iv) Solid and liquid PCM have the same density.

For solidification in a spherical enclosure subject to convective cooling, there are three major thermal resistances that are in series: resistance due to the external convection, $R_{\text{conv}}$, resistance due to the radial conduction through the wall, $R_{\text{wall}}$, and resistance due to the conduction through the solidified spherical shell, $R_{\text{PCM}}$, given by

\[
R_{\text{conv}} = \frac{1}{hA} \quad (4)
\]

\[
R_{\text{wall}} = \frac{r_o - r_i}{4\pi k_{\text{wall}} r_o^2 r_i} \quad (5)
\]

\[
R_{\text{PCM}} = \frac{r_i - r_s}{4\pi k_{\text{PCM}} r_i^2 r_s} \quad (6)
\]

where $A$ is the area of the external surface of the spherical enclosure and $r_s$ denotes the radial location of the solidification front. The temperature difference between the PCM phase change temperature, $T_m$, and the surrounding temperature, $T_\infty$, induces solidification of the PCM and release of the latent heat of the PCM. The solution procedure is iterative in time. At any time step, the heat transfer rate $q$ is calculated by dividing the driving temperature difference by sum of the thermal resistances $q = (T_m - T_s) / \sum R$. This heat transfer rate is then used to calculate the amount of solidified PCM at the end of the time step, following

\[
\dot{m}_{\text{solid}} = \frac{q}{h_{sf}} \quad (7)
\]

The location of solidification front at the end of time step is then updated using the mass of solidified PCM following
The solidification fraction is then \( V^* = 1 - \left( \frac{r_s}{r_i} \right)^3 \), where \( V_0 \) is the total volume of PCM. The iterations stop when \( V^* > 99.9\% \) or the solidified fraction reaches the desired value.

![Diagram of PCM freezing](image)

Figure 4: (a) Inward freezing of PCM within a spherical enclosure; (b) Thermal resistance network for inward freezing due to convective cooling.

Equations (4)-(6) can be non-dimensionalized by introducing the following dimensionless variables: \( r^* = r/r_o \), \( R^* = R/\left[ r_o/(Ak_{PCM}) \right] \) and \( Bi = 2hr_o/k_{PCM} \), where \( h \) is the external heat transfer coefficient. In addition, in order to focus on the most important parameters affecting the solidification subject to convective cooling, the following non-dimensionalized formulation is further simplified by considering a thin wall \(( r_o = r_i) \) with negligible thermal resistance. Using the above dimensionless variables, the non-dimensional form of the convective and PCM thermal resistances are obtained as

\[
R_{conv}^* = \frac{2}{Bi} 
\]
The evolution of solid volume fraction, $V^*$, versus non-dimensional time, $Fo Ste$, is obtained by non-dimensionalizing Eq. (7) to yield

$$R_{PCM}^* = \frac{1}{r_s} - 1$$

The evolution of solid volume fraction, $V^*$, versus non-dimensional time, $Fo Ste$, is obtained by non-dimensionalizing Eq. (7) to yield

$$\frac{d V^*}{d Ste Fo} = \frac{3}{\left(1 - V^*\right)^{1/3}} - 1 + \frac{2}{Bi}$$

where $Ste$ and $Fo$ are defined as $Ste = c_p(T_m - T_\infty)/h_{sl}$ and $Fo = k_{PCM}/\left(\rho_{PCM}c_p r_s^2\right)$ respectively.

4. Results and Discussion

Figure 5a shows the experimentally-determined effective thermal conductivity of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04%, 1.57% and 2.09% as a function of temperature obtained from the hot wire tests. Eicosane is in solid state at 23 °C, and is in liquid state at 40 °C and 55 °C. The thermal conductivity of pure solid eicosane at 23 °C is 0.407 W/(m·K), consistent with literature [21]. As shown in Fig. 5a, the effective thermal conductivity of eicosane keeps increasing with xGnP volume fraction, and reaches 1.24 W/(m·K) with only 2.09 vol% of xGnP. Furthermore, at all xGnP volume fractions investigated, the thermal conductivity of solid eicosane is higher than that of liquid eicosane. The crystalline structure of eicosane contributes to the higher thermal conductivity in the solid phase. It is observed that the effective thermal conductivity of eicosane in the liquid phase does not change noticeably from 40 °C to 45 °C, indicating that thermal conductivity is not a strong function of temperature in this range.

Figure 5b shows the enhancement of thermal conductivity of eicosane, $k_{eff}/k_{base}$ ($k_{base}$ the thermal conductivity of pure eicosane), as a function of volume fraction of xGnP, $\phi$. The blue squares and red triangles represent solid and liquid eicosane, respectively. The green circles represent the enhancement in effective thermal conductivity of a commercial paraffin PCM (IGI 1230A) from Warzoha and Fleischer [22]. The solid line represents the prediction of effective medium model of Nan et al. [36] following
\[
\frac{k_{\text{eff}}}{k_{\text{base}}} = 1 + \frac{\varphi p}{3} \frac{k_{x\text{GnP}}/k_{\text{base}}}{p + \frac{2a_k}{\delta} k_{x\text{GnP}}/k_{\text{base}}}
\]

where \(p\) is the aspect ratio of xGnP, \(a_k\) is the Kapitza length, and \(\delta\) is the thickness of xGnP. The Kapitza resistance at the interface between graphite nanoplatelet and paraffin is \(R_k \approx 0.9 \times 10^{-7} \text{m}^2\text{K/W}\) [37]. Thus, the Kapitza length is \(a_k = R_k k_{\text{base}} = 36.6 \text{nm}\). The thermal conductivity of xGnP adopted here is based on xGnP paper with \(k_{x\text{GnP}} = 98 \text{W/(m-K)}\) [33], which is much smaller than the intrinsic thermal conductivity of xGnP of 2,000 W/(m-K) [38] due to more phonon-boundary scattering. As shown in Fig. 5b, both present work and Warzoha and Fleischer’s data agree well with the model of Nan et al. With xGnP volume fraction of 2.09%, the thermal conductivity increases by 205% for solid eicosane and 211% for liquid eicosane.

Figure 5. (a) Effective thermal conductivity of eicosane with xGnP as a function of temperature for various xGnP volume fractions, and (b) Enhancement in thermal conductivity relative to pure eicosane compared against theoretical predictions of Nan et al. [36] and experimental measurements of Warzoha and Fleischer [22].

Figure 6a shows the experimental results for solidification time, \(t_{\text{sol}}\), as a function of convective subcooling, \(T_{\text{PCM}} - T_{\text{water}}\), for solidification of eicosane with xGnP volume fractions.
of 0%, 0.52%, 1.04%, 1.57% and 2.09% in spherical enclosures with outer diameter of 38 mm and 50 mm subject to water cooling. For both enclosure sizes, the solidification time decreases with increasing subcooling and increasing xGnP volume fraction. The enhancement in thermal conductivity with increasing nanoadditive volume fraction leads to reduction in solidification time. For example, the thermal conductivity of additive-enhanced eicosane with 1.57 vol% xGnP is 1.05 W/(m·K), which is ~2.5 times of the conductivity of pure eicosane, 0.407 W/(m·K). As a result, a 50% reduction in solidification time is observed for additive-enhanced eicosane with 1.57 vol% xGnP compared to pure eicosane. As shown in Figure 6a, for the same volume fraction of xGnP, the solidification time increases with the diameter of the spherical enclosure. For example, in the subcooling range of 11.4-26.4 °C, the solidification time of pure eicosane in the 50 mm sphere is about 105% greater than that in the 38 mm sphere. The inner diameter of the spherical enclosure (i.e. the diameter of PCM) is 32 mm for the 38 mm sphere, and 46 mm for the 50 mm sphere. Thus, the conduction length of PCM in the 50 mm sphere is ~ 1.43 times of that in the 38 mm sphere. The solidification time roughly following $t_{sol} \propto D_{PCM}^2$ indicates that the solidification of PCM under fixed temperature boundary is a conduction-controlled process, and natural convection does not play a significant role in the process.

It is helpful to generalize the results of the solidification tests by presenting them in terms of dimensionless numbers and compare them with the theoretical modeling. Figure 6b shows the comparison between the experiment-determined and model-predicted non-dimensional solidification time, $F_{sol}$, as a function of $St_e$. It is noted that $F_{sol}$ from the experiments is determined using the inner diameter of the spherical shell, whereas the model does not need any specification of the physical dimensions as long as only the dimensionless quantities are of interest. The solid line represents the prediction of the conduction-controlled model (i.e., Eq. (11)) in the limit of $Bi \to \infty$. The solid and open symbols represent the experimental results for solidification in spherical enclosures with outer diameter of 38 mm and 50 mm, respectively. As shown in Fig. 6b, the solidification time from experiments is smaller than the prediction of the conduction-controlled model, which is consistent with the observation of solidification of PCM in cylindrical enclosures subject to constant temperature boundary [39,28]. The experimental solidification times in this work are determined using the thermocouple at the center and at the edge of the spherical PCM. Dendrite solidification resulting in traces of solid PCM to reach to
the center thermocouple while some amount of liquid PCM still remains in the system may lead to the underestimation of solidification time from experiments [31]. It can also be observed from Fig. 6b that the non-dimensional solidification time, $F_{sol}$, for the 50 mm spherical enclosure is greater than the 38 mm enclosure. The size effect can also be attributed to the dendrite growth as the effect of dendrites can be more pronounced for smaller enclosure size. This trend of increasing $F_{sol}$ with increasing enclosure size is consistent with the observation of solidification of PCM in cylindrical enclosure [28].

The conduction-controlled model is developed based on the four basic assumptions listed in Section 2.3. Here we examine the possible role of these assumptions in the deviation between the model prediction and experimental results. Firstly, the model assumes a constant solidification temperature, which neglects the possible sensible heat accumulation during the solidification process. However, the PCM in the present study has a narrow solidification range of 35-37 °C, leading to a sensible heat of $\sim 4$ kJ/kg (based on solid PCM specific heat of 1.92 kJ/(kg·K)), which is negligible compared to the latent heat of PCM ($\sim 247$ kJ/kg). Secondly, the model assumes that the liquid PCM is initially at the melting temperature $T_m$. In the experiments, the liquid PCM starts at a temperature $\sim 2$-5 °C higher than $T_m$, but the amount of sensible heat due to superheating is $\sim 4$-10 kJ/kg, much smaller than the latent heat of PCM. Thirdly, the model assumes $Ste << 1$. The Stefan number in the experiments is within the range of 0.1-0.205. It has been shown in the literature that for $Ste$ up to 0.235 the effect of sensible cooling of the frozen PCM is negligible [40]. Lastly, the model neglects the density difference between solid and liquid PCM. In our experiments, the density of the solid PCM is slightly ($\sim 4\%$) higher than that of the liquid, which should not lead to a major difference between the experiments and model predictions. The above analysis shows that these four model assumptions may contribute but not be the only root of the disagreement between the experimental results and model predictions.
Figure 6. (a) Solidification time of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04% and 1.57% subject to water cooling as a function of subcooling. (b) Comparison between experiments and model predictions for non-dimensional solidification time, $F_{o_{sol}}$ as a function of Ste for solidification of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04%, 1.57% and 2.09% in water cooling.

Figure 7a shows the experimentally-determined solidification time, $t_{sol}$, as a function of air velocity, $U_{air}$, for solidification of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04%, 1.57% and 2.09% in spherical enclosures with outer diameter of 38 mm and 50 mm subject to air cooling. For both enclosure sizes, the solidification time decreases with increasing air velocity due to enhanced external convection. Moreover, the solidification time decreases with increasing volume fraction of xGnP due to enhanced PCM thermal conductivity. As PCM thermal conductivity increases from 0.407 W/(m·K) for pure eicosane to 1.05 W/(m·K) for eicosane with xGnP volume fraction of 1.57%, an average 33% reduction is observed in solidification time. It is noted that for the same enhancement in PCM thermal conductivity, 50% reduction in solidification time is obtained in water cooling. This is because while conduction resistance of solid PCM is the dominant resistance in water cooling, both the external convection resistance and the conduction resistance of solid PCM are important in air cooling. The addition of nanoadditives reduces the PCM conduction thermal resistance, but does not affect the convection resistance.
Figure 7b shows the comparison between the experimentally-determined and model-predicted non-dimensional solidification time, $F_{o_{solv}}$, as a function of $Bi$. The solid line represents the prediction of the conduction-controlled model developed in Section 2.3. The solid and open symbols represent the experimental results for solidification in spherical enclosures with outer diameter of 38 mm and 50 mm, respectively. The $Bi$ number ($Bi = 2h_{air} r_o / k_{PCM}$) for the experimental results is calculated using the air-side heat transfer coefficient and the measured thermal conductivity of PCM. The air-side heat transfer coefficient is calculated following $h_{air} = Nu_{air} k_{air} / (2r_o)$, and the Nusselt number correlation for flow over a sphere is based on $Nu = 2 + \left(0.4Re^{1/2} + 0.06Re^{2/3}\right)Pr^{0.4} (\mu / \mu_s)^{1/4}$, where the Reynolds number is defined as $Re = 2 \rho_{air} r_o U_a / \mu$, the Prandtl number is given by $Pr = c_{p,air} / k_{air}$, $\mu$ is the dynamic viscosity of the ambient air, and $\mu_s$ is the dynamic air viscosity on the sphere surface.

Consistent with our results in water cooling, the experimentally-determined $F_{o_{solv}}$ in air cooling is smaller than the model prediction. Since the distribution of convection heat transfer coefficient is not completely uniform on the outer surface of the spherical enclosure, solidification of PCM can be asymmetric in the spherical enclosure. It is thus possible that the thermocouple in the center of the spherical enclosure gets in contact with solid PCM while liquid PCM still exists elsewhere in the enclosure. This asymmetric solidification may lead to underestimation of the solidification time from experiments. It can be argued that together with dendrite solidification, the asymmetric solidification is responsible for smaller $F_{o_{solv}}$ than model prediction in air cooling.
Figure 7. (a) Solidification time of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04% and 1.57% in air cooling as a function of air velocity with Ste = 0.0887. (b) Comparison between experiments and model predictions for non-dimensional solidification time, Fo, as a function of Bi for solidification of eicosane with xGnP of varying volume fractions in air cooling.

Figure 8a shows the dynamic solid fraction, $V/V_0$, during the solidification process of eicosane with xGnP of varying volume fractions solidifying in the 50 mm diameter spherical enclosure subject to water cooling at $T_{water} = 25^\circ C$ and air cooling at $T_{air} = 25^\circ C$ and $U_{air} = 25$ m/s. The product of Fourier and Stefan numbers, FoSte is used to represent the non-dimensional time that incorporates the effect of thermal conductivity and subcooling. The black solid line represents the model prediction for water cooling based on Eq. (11) in the limit of Bi $\rightarrow \infty$. The black and green dashed lines represent the model predictions for air cooling at $U_{air} = 25$ m/s for xGnP volume fraction of 0% and 1.57%, respectively. It is observed that the experimental results for water cooling collapse to the model prediction with constant temperature boundary, and the experimental results for air cooling lie within the range of the model predictions for the lowest and highest xGnP volume fractions (i.e. 0% and 1.57%, respectively). The non-dimensional solidification time for air cooling is larger than water cooling due to the effect of convective resistance.
Figure 8. Comparison of experiments and model predictions for solid fraction, $V^*$ as a function of FoSte for solidification of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04% and 1.57% in water at $T_{\text{water}} = 25$ °C and in air at $T_{\text{air}} = 25$ °C and $U_{\text{air}} = 25$ m/s. The model prediction is based on Eq. (11) in the limit of $\text{Bi} \to \infty$ for water cooling, and Eq. (11) at $U_{\text{air}} = 25$ m/s for air cooling.

As shown in Figure 7b and Figure 8, the non-dimensional solidification time under convective cooling, $F_{\text{sol}}$, is a function of Bi, which characterizes the competition between the external convective resistance and the PCM conduction resistance. In the limit of $\text{Bi} \to \infty$ (corresponding to constant temperature boundary), $F_{\text{sol}} \to F_{\text{sol}}|_{\text{Bi} \to \infty}$. As for finite values of Bi (corresponding to convective boundary with finite heat transfer coefficient), $F_{\text{sol}} > F_{\text{sol}}|_{\text{Bi} \to \infty}$ due to the additional convective resistance. By fitting the results of the numerical model, the following simple correlation is obtained

$$\frac{F_{\text{sol}}}{F_{\text{sol}}|_{\text{Bi} \to \infty}} = 1 + \frac{4.5}{\text{Bi}}$$

Equation (13) can be used as a tool to calculate the solidification time of PCM freezing in spherical enclosures subject to convective cooling with known $F_{\text{sol}}|_{\text{Bi} \to \infty}$. The Fourier number
for constant temperature boundary, \( F_{\text{sol}}_{\text{Bi} \to \infty} \), can be solved from Eq. (11) in the limiting case of \( \text{Bi} \to \infty \). And the factor \( \left( 1 + 4.5/\text{Bi} \right) \) can be used to unify the experimental results of solidification time for different cooling conditions.

Figure 9 shows the comparison between the experiments and model predictions for the scaled non-dimensional solidification time, \( F_{\text{sol}}/(1 + 4.5/\text{Bi}) \), as a function of \( \text{Ste} \) for solidification of eicosane with xGnP volume fractions of 0%, 0.52%, 1.04%, 1.57% and 2.09% in water and air. For solidification experiments in air, the non-dimensional solidification time \( F_{\text{sol}} \) is scaled with \( 1 + 4.5/\text{Bi} \), based on Eq. (13), which relates \( F_{\text{sol}} \) for convective cooling to the non-dimensional solidification time for constant temperature cooling, \( F_{\text{sol}} \bigg|_{\text{Bi} \to \infty} \). It is observed that the experimental results for water cooling overlap with the results for air cooling, proving that the scaling factor from Eq. (13) successfully unifies the results for water and air cooling. And despite of the cooling conditions, the scaled dimensionless solidification time \( F_{\text{sol}}/(1 + 4.5/\text{Bi}) \) monotonically decreases with increasing \( \text{Ste} \).
Figure 9. Comparison between the experiments and model predictions for the scaled non-dimensional solidification time, $\frac{F_{0_{\text{sol}}}}{(1 + 4.5/Bi)}$ as a function of Ste.

5. Conclusion

In the present work, the solidification of PCM with xGnP volume fractions of 0%, 0.52%, 1.04%, 1.57% and 2.09% in spherical enclosures subject to air and water cooling has been investigated through experiments and modeling. The experimentally-determined solidification time is \(~30\%\) smaller than the prediction of the simple thermal network model due to possible asymmetric and dendritic solidification, as well as natural convection inside the enclosure. The experiments show that the solidification time decreases with increasing subcooling and increasing external heat transfer coefficient. Furthermore, the solidification time decreases with increasing volume fraction of xGnP due to the xGnP-induced enhancement in thermal conductivity of PCM. The reduction in solidification time is more pronounced in water cooling than air cooling because the PCM conduction resistance limits the heat transfer in water cooling, while external convective resistance is comparable with PCM conduction resistance in air cooling. Based on the conduction-controlled model, a simple correlation for the scaled non-dimensional solidification time, $\frac{F_{0_{\text{sol}}}}{F_{0_{\text{sol}}}} \bigg|_{Bi \rightarrow \infty} = 1 + 4.5/Bi$, has been developed to incorporate
the effect of external heat transfer coefficient, so as to unify the experimental results for water and air cooling.

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