Experimental and numerical study of non-stationary solid-liquid phase transitions of n-tetradecane

Viktoria Zakharova1*, Pavel Kuznetsov1, Daniyar Kalimzhanov1, and Aleksandr Baranenko1

1ITMO University, Faculty of Cryogenic Engineering, 197101, St. Petersburg, Kronversky pr. 49, Russian Federation

Abstract. During the experiments, four spherical flasks with n-tetradecane with a total mass of 0.344 kg were placed in a test container inside which an aqueous solution of propylene glycol with a mass concentration of 50% and an initial temperature of 25 °C was circulating. As a result of studies, the values of heat flows at different time intervals during the solid-liquid phase transition of n-tetradecane were obtained. Based on the modeling of the processes, the distribution of temperature and velocity of the heat carrier in the test container as well as the values of the local heat transfer coefficients on the surface of the flasks were established. Numerical calculations of the process of phase transition being studied according to the authors' model were also performed. Numerical calculations of the process under study have been performed using the authors' model. A satisfactory convergence of the experimental and calculated values of the heat flow has been obtained. The results can be used in the development of thermal energy accumulators with n-tetradecane.

1 Introduction

Non-stationary liquid-solid-liquid phase transitions are rather common processes. They can be implemented in various technical devices. In particular, they occur in thermal energy accumulators with phase change materials (PCM) [1, 2].

Thermal energy accumulators are used in heating, hot water supply, cooling and air conditioning systems with variable load. The energy stored in accumulators is used at the maximum consumption of the corresponding resource. This allows installing equipment of lower power, equalizing daily electricity consumption, reducing electricity costs when double-tariff accounting is used and when charging thermal energy storage devices takes place at night. The accumulators can use renewable energy sources such as solar energy and night-time radiation cooling, which reduces electricity consumption [3, 4].

In comparison with accumulators with liquid, the advantage of thermal energy accumulators with PCM is that they have an order of magnitude higher volume and mass density of stored energy. This is due to the fact that in them, the main contribution to the thermal energy accumulation is made by the heat of the phase transition.

Also, in such accumulators, the process of accumulating and spending energy stored is carried out at a constant temperature corresponding to the temperature of the phase transition of the PCM. In a more common design of these devices, the PCM is placed inside the container in capsules, which can be of arbitrary shape, including balls or cylinders. The heat carrier circulates in the remaining free internal volume of the accumulator [5].

A fairly large number of theoretical and experimental papers are devoted to the study of processes in such thermal energy storage devices [6-12]. In [6], a detailed experimental and theoretical study of two-dimensional temperature fields in spherical capsules was carried out (axisymmetric problem). In [7], a calculation model for a numerical solution based on Navier–Stokes equation using the enthalpy approach is presented. In [8], the calculation model was applied to study the influence of fins located inside cylindrical capsules on heat exchange. In [9], an experimental and numerical study for elliptical capsules was carried out. However, in these papers, the determination of such quantities as the total heat flow, primarily necessary for the design of cooling systems, occurs indirectly from the results of calculating temperature fields. In [11], the authors investigated the processes in dense backfills of steel capsules. Paper [12] is devoted to modeling a system with thermal energy storage devices as a whole.

For cold accumulation in air conditioning systems, PCM with a phase transition temperature of 5-10 °C are of interest. Among such substances are paraffins which advantage of which is stability during repeated multidirectional phase transitions, a relatively high specific heat of the phase transition and a slight difference in density in the liquid and solid states.

2 Research and results

The authors have performed an experimental and numerical study of the solid-liquid phase transition of n-
tetradecane (chemical formula C14H30). According to the passport, the mass fraction of the main substance is 99.89%, the melting point is 6.1 °C. The main properties [13, 14] are the following: specific mass of melting (λ_{m}) is 215 kJ/kg, liquid phase density (ρ_{lc}) at 25 °C is 762.8 kg/m³, dynamic viscosity at 25 °C is 2.13·10⁻³ Pa·s, thermal conductivity coefficient of the liquid phase is 0.143 W/(m·K), specific heat of the liquid phase (C_{p,lc}) is 2198 J/(kg·K).

Experimental studies were carried out in an isolated test container with dimensions of 240x150x115 mm (Fig. 1). The PCM was placed in round-bottom flasks of a spherical shape with an outer diameter of 66 mm, the flasks were filled to the base of the neck. In the experiments, 4 flasks were used; in them, the PCM mass was 0.344 kg. In the test container, the flasks were located in a strictly defined order in all the experiments as it is shown in Fig. 1. The heat carrier movement was carried out in a closed circuit and with a mass flow rate of 2.1 l/m. The heat carrier volume in the circulation circuit was 4.15 l. An aqueous solution of propylene glycol with a mass concentration of 50% was used as a heat carrier. In all the experiments, the initial temperature of the heat carrier was 25 °C.

The movement of the heat carrier in the test container is shown in Fig. 2, it is modeled in the Fluent package. Modeling is carried out for the steady-state operation mode of the stand. When modeling the movement of the heat carrier, the natural convective movement of the liquid is also considered, since preliminary calculations have shown its noticeable influence in this design of the stand. When modeling, the external walls of the test container are assumed to be thermally insulated, since the temperature of the heat carrier corresponds to the ambient temperature, which eliminates the heat flow between them. The temperature of the walls of the flasks is assumed to be constant and equal to the melting temperature of the SPT.

Before starting the experiments, the flasks with the PCM were cooled until the complete transition of the PCM to a solid state. In the course of these experiments, the following parameters were determined and recorded. At the beginning and end of the experiment, the temperature of the PCM as well as of the phase transition; the heat carrier temperature in regular time intervals, the time of the beginning and end of the experiment, the heat gain from the pump. The experiments were over with the complete transition of the PCM to a liquid state. To measure the temperature, electronic thermometers with an accuracy of 0.2 °C were used. In the experiments, the phase transition temperature of the PCM was 6.1 °C, which corresponds to its passport value.

![Fig. 1. Temperature distribution in the test container during the experiment.](image1)

![Fig. 2. Heat carrier movement in the test container.](image2)

Numerical calculations have also shown that in the test container, during the experiment, the temperature gradient of the heat carrier can reach 10°C. The lowest temperatures of the heat carrier are set in the lower part of the test container, despite the fact that the heat carrier moves in it at a certain speed. This indicates the feasibility of supplying the heat carrier from below. Local heat transfer coefficients differ by an order of magnitude (Fig. 3). Their maximum value is 3600 W/(m²·K), the average value is 380 W/(m²·K). The maximum values of the local heat transfer coefficients take place on the side of the flasks along the flow of the heat carrier of their inlet pipe in the central zone of the test container.

| Flask number | 1   | 2   | 3   | 4   |
|--------------|-----|-----|-----|-----|
| Average heat transfer coefficient, W/( m²·K) | 345 | 453 | 341 | 384 |

When processing experimental data, the convergence of thermal balances was checked. The heat absorbed by the PCM (Q_{PCM1}) should be equal to the heat released from the heat carrier (Q_{PCM2}). Q_{PCM2} includes the heat of the heat carrier (Q_{hc}), flask (Q_{f}) and pump (Q_{p}), kJ.
The average heat flow from the heat carrier for a certain time interval, kW

\[ q = q_{hc} + q_f + q_p \]  

(7)

\[ q_{hc} = \frac{V_{hc} \rho_{hc} C_{p, hc} \Delta \tau_{hc}}{\Delta \tau_i}, \quad q_f = \frac{m_f C_{p, f} \Delta \tau_f}{\Delta \tau_i} \]  

(8)

\[ \Delta \tau_i \] is the change in temperature of the heat carrier and flask for a certain time interval \( \Delta \tau_i \).

The values of \( q \) calculated from experimental data are presented in Fig. 4. This figure also shows the calculated dependence \( q_{PCM} = f(\tau) \) computed by the authors’ model [15].

At the initial moment of time, the heat flow increases sharply and after a fairly short time, it begins to decrease due to an increase in the thermal resistance of the liquid phase of the PCM (since its thickness increases) and a decrease in the contact surface of the phases of the PCM. The results obtained correspond to the regularities of the process under study as well as to the data of other authors [16]. At the same time, the calculated and experimental values of the duration of the solid-liquid phase transition process differ by only 2%. The greatest interest present the data on the values of the heat flow in the time interval of the phase transition process of 5-30 min, when the dependence of \( q_p \) on time is monotonous. Based on these data, it is most advisable to model the calculations of thermal energy accumulators.

3 Conclusions

The duration of the experiment was 36-38 minutes. The heat flow reached its maximum value of 26 W in 90 seconds after the start of the experiments. In the interval of 5-30 minutes, the dependence of the heat flow on time is monotonous. The results obtained by the authors have confirmed the adequacy of the previously developed
model of the phase transition process. They can be used in the development of methods for calculating thermal energy accumulators with the PCM.

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