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Predictive approach of heat transfer for the modeling of large-scale latent heat storages

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Abstract – Thermal energy storage systems based on phase change materials are interesting candidates to handle the difficulties raised by intermittent renewable sources or by batch processes. Among these systems, many rely on the use of steam, as for instance in concentrating solar power plants or district heating, or in the pharmaceutical or food industries. Today, there is no systematic method to design such systems quickly and easily since complex heat transfer is observed due to the influence of the geometry and to the dual characteristics associated with solid/liquid and liquid/gas transitions. The aim of the present work is thus to propose a multi-scale modelling methodology of a latent heat storage system for the storage of steam. It mainly involves two different simulation models with different scales for the heat transfer fluid and the phase change material. Furthermore, it relies on the use of a heat transfer correlation based on specific non-dimensional numbers, which is deduced from previous simulations of the phase change material's behavior, obtained with fine 3D computational fluid dynamics calculations. Consequently, a reduced model is built to simulate the whole system. This model does not need to be tuned against experiments. This model is then directly used to compare the numerical results with measurements coming from a prototype scale latent heat storage available at CEA Grenoble. The results are very promising and show that an *a priori* approach that is more physically consistent and not based on any model tuning can lead to acceptable results. Moreover, the computational time can be divided by 10 to 40, thus allowing future design and real performance evaluations of latent heat storage modules.

Keywords – Thermal energy storage; liquid-vapor; solid-liquid; model reduction; non-dimensional correlation; heat transfer

Highlights

- Development of a reduced multi-scale modelling approach for the charge of a steam storage module involving a PCM.
- Deduction of a heat transfer correlation from fine 3D CFD computations.
- No experimental tuning.
- Dual two-phase storage with liquid/gas HTF inside the tubes and solid/liquid PCM outside.
- A local heat transfer correlation is built.

1

2 Nomenclature

3 Latin letters

| | |
|--------------|---|
| A | factor in the damping term of the momentum equation ($\text{kg}\cdot\text{m}^{-3}\cdot\text{s}^{-1}$) |
| A_{mush} | mushy zone constant ($\text{kg}\cdot\text{m}^{-3}\cdot\text{s}^{-1}$) |
| b | constant term in the damping term |
| c | tube perimeter (m) |
| c_p | heat capacity at constant pressure ($\text{J}\cdot\text{kg}^{-1}\cdot\text{K}^{-1}$) |
| D | thermal diffusivity ($\text{m}^2\cdot\text{s}^{-1}$) |
| e_l | liquid layer thickness (m) |
| g, \vec{g} | acceleration of gravity ($\text{m}\cdot\text{s}^{-2}$) |
| h | mass enthalpy ($\text{J}\cdot\text{kg}^{-1}$) |
| l_c | characteristic length (m) |
| L | latent heat ($\text{J}\cdot\text{kg}^{-1}$) |
| \dot{m} | mass flow-rate ($\text{kg}\cdot\text{s}^{-1}$) |
| Nu | Nusselt number (–) |
| P | pressure (Pa) |
| r | radial coordinate (m) |
| R | radius (m) |
| Ra | Rayleigh number (–) |
| S | tube section (m^2) |
| S_e | heat exchange surface (m^2) |
| t | time (s) |
| T | temperature ($^{\circ}\text{C}$ or K) |
| \vec{U} | velocity ($\text{m}\cdot\text{s}^{-1}$) |
| V | volume (m^3) |
| x | volume fraction (–) |
| Y | mass fraction (–) |
| z | axial coordinate (m) (–) |

4

5 Greek symbols

| | |
|----------------|--|
| α | heat transfer coefficient ($\text{W}\cdot\text{m}^{-2}\cdot\text{K}^{-1}$) |
| β | thermal expansion coefficient (K^{-1}) |
| ρ | density ($\text{kg}\cdot\text{m}^{-3}$) |
| μ | dynamic viscosity ($\text{kg}\cdot\text{m}^{-1}\cdot\text{s}^{-1}$) |
| ν | kinematic viscosity ($\text{m}^2\cdot\text{s}^{-1}$) |
| ε | void fraction (–) |
| λ | thermal conductivity ($\text{W}\cdot\text{m}^{-1}\cdot\text{K}^{-1}$) |
| λ_{eq} | equivalent thermal conductivity ($\text{W}\cdot\text{m}^{-1}\cdot\text{K}^{-1}$) |
| φ | heat flux ($\text{W}\cdot\text{m}^{-2}$) |
| Φ | heat flow-rate (W) |

6

7 Superscript and subscripts

| | |
|----------------|-----------------|
| \overline{A} | mean value of A |
| eff | effective |

| | |
|-------------|---|
| <i>ext</i> | external boundary |
| <i>fin</i> | fins and inserts |
| <i>h</i> | homogeneous material |
| <i>i, j</i> | cell indexes (axial and radial direction) |
| <i>l</i> | liquid phase |
| <i>liq</i> | liquidus |
| <i>pc</i> | phase change |
| <i>ref</i> | reference |
| <i>s</i> | solid |
| <i>sat</i> | saturation |
| <i>sens</i> | sensible |
| <i>sol</i> | solidus |
| <i>t</i> | tube |
| <i>text</i> | outer tube wall |
| <i>tint</i> | inner tube wall |
| <i>v</i> | vapor phase |
| <i>w</i> | wall |

1

2 Abbreviations

| | |
|-----|--------------------------|
| CSP | concentrated solar power |
| DSG | direct steam generation |
| HTF | heat transfer fluid |
| LHS | latent heat storage |
| PCM | phase change material |

3

1

2 1 Introduction

3 1.1 Forewords

4 Thermal energy storage systems are known to be key components of solar thermal power plants, principally because
5 they reduce the probabilities of breakdown [1], [2] and thus lead to more stable regimes. At the same time, many
6 industrial applications make a huge use of steam, as for example in Direct Steam Generation (DSG) concentrating solar
7 power (CSP) plants, where the two-phase heat transfer fluid (HTF) avoids the installation of heat exchangers and
8 extends the 400°C limit usually associated with common thermal oils. Again, this simplifies the design and operations,
9 and increases overall efficiency [3], [4]. Nonetheless, the thermal storage of DSG-CSP must be adapted to the two-
10 phase feature of the HTF. Generally, a steam accumulator is used and the water vapor is stored in pressurized vessels,
11 in a saturated state in thermal equilibrium with a saturated liquid water volume [5]. Unfortunately, such technologies
12 are rather expensive when large capacities are needed [6], [7]. Therefore, the underlying research question of the
13 present contribution concerns the development of latent heat storages (LHSs), using phase change materials (PCMs) to
14 store the thermal energy of the steam in the latent heat of a solid-liquid phase transition. In such storage systems
15 where the phase change occurs in a narrow temperature range, the steam pressure drop between charge and
16 discharge cycles is minimized compared to sensible storages [8]. The most mature steam storage technology with PCM
17 is the shell-and-tube concept [9], [10], where the HTF flows in tubes that go through a PCM-filled tank. During the
18 charge, the steam condenses inside the tubes and the initially solid PCM progressively melts, while during the
19 discharge, liquid water fed to the tubes evaporates and regenerates steam due to the heat released during the
20 solidification of the PCM. Practically, metallic or graphite fins are often disposed around these tubes to enhance the
21 heat transfer, in order to cope with the low thermal conductivities of usual PCM [11]. Moreover, the PCM shell does
22 not need to be pressurized, which simplifies the storage design and reduces its costs. Generally, there are two
23 materials that are regularly used as PCMs with storages designed for CSP applications, namely sodium nitrate NaNO_3
24 [6]-[8], [12], [20] and a eutectic mixture of potassium nitrate and sodium nitrate $\text{KNO}_3/\text{NaNO}_3$. [11], [14]. For
25 prototypes or laboratory scale bench-top modules, all kinds of PCM can be found, such as for instance RT35-HC [9]-
26 [10] or n-octadecane [16]-[17], as well as salt hydrates [19]. Several prototype-scale shell-and-tube LHS modules
27 involving a two-phase liquid water – steam HTF have been built and successfully operated by several teams around the
28 world [6], [12], [92], demonstrating the feasibility and the relevance of this concept for the storage of steam. Among
29 numerical studies, a few 1D or 2D simulation models intended for module design or quick performance estimation can
30 be found in the literature [7], [8], [20]. Due to the small number of large-scale modules being built, and to the
31 complexity of the heat transfer phenomena inside a module, no design model has yet been validated on experimental
32 data from large-scale modules. In summary, since the selection of a PCM is known to be strongly oriented by the
33 storage application, DSG-CSP plants require a PCM whose phase transition is around 300°C (saturation temperature of
34 steam at a pressure level suitable for turbine operations). In this temperature range, inorganic salts (such as sodium
35 nitrate) are an interesting solution thanks to their favorable physical and chemical properties (stable, non-corrosive)
36 and low cost. Their major drawback is their low thermal conductivity, which implies either very low charging and
37 discharging thermal powers, or a small distance between tubes. To achieve acceptable charge and discharge times
38 (typically about six hours for CSP applications), without increasing the number of tubes and thus the cost of the heat
39 exchanger, an improvement in the heat transfers is needed.

40 In general, the heat transfer coefficient between the condensing/evaporating HTF and the inner tube wall is high.
41 Consequently, the heat transfer dynamics are mainly driven by the phenomena occurring on the PCM side. Here, heat
42 transfer is influenced by both the geometric shape and dimensions of the fins, and by the possible natural convection
43 movements inside the PCM liquid phase. Indeed, several studies have shown that natural convection could have a
44 non-negligible influence on the thermal performances of a storage module [12]–[15]. Since computational fluid
45 dynamics (CFD) calculations, which are capable of representing this kind of problem, are often too long and/or
46 expensive, one needs simpler engineering tools to perform fast and multiple computations. Generally, one tries to rely
47 on heat transfer correlations [16]-[17]; however, these correlations often lack generality and show caveats when they
48 are simply fitted on experimental data [16]–[19]. In contrast, other studies propose to use CFD simulations to estimate

1 a non-dimensional correlation, which can then be used later on in a reduced system model. Thus, Garcia *et al.* [12] and
2 Johnson *et al.* [20] implemented such a technique in a 3D purely conductive regime of fusion/solidification to account
3 for the role of fins and inserts. Another similar approach can be found in Mostafavi Tehrani *et al.* [21] who modeled
4 the convective-driven melting of a PCM around a single tube without fins and inserts. They deduced from these
5 simulations a non-dimensional correlation for the heat transfer, which was then used in a reduced system model. In
6 fact, the underlying goals of these analyses are to allow for better predictions of the main features of the storages,
7 namely their energy capacity and associated efficiency and their charging/discharging times. These characteristics are
8 of paramount importance, yet they are singularly different from one case to another. For instance, in [6, 99] a NaNO_3 -
9 based storage system with an average storage capacity of 700 kWh and a discharge thermal power of 200 to 500 kW
10 was investigated both experimentally and numerically. Its charging time was around 1-3h. The same PCM, with the
11 same methodology, were used in the LHASSA module [12] where full cycles for the charge and discharge lasted
12 approximately 8 h and 6 h respectively, with a storage efficiency of 90%. The equivalent powers were thus around 35
13 kW and 47 kW. Along the same lines, [97] performed a numerical study of the effectiveness of a multi-tube storage,
14 with a NaCl-MgCl_2 eutectic mixture as a PCM, where the length and diameter were varied (between 10 to 150 m, and
15 between 1 and 2 inches and 1 and 2 and 3 inches for the inner diameter of the tubes and the space between them).
16 Results showed a variation in the efficiency of the order of 70%. With an insulated steel container housing a fin-tube
17 heat exchanger filled with HDPE, [96] demonstrated experimentally and numerically the achievement of a 20-25 kWh
18 storage in 1-2 h. In the case of smaller volumes or with preliminary prototype modules, the same research questions
19 regularly arise. In [98], three 0.5 m concentric tubes with internal fins containing a tube with internal fins, and PCM
20 between them, was experimentally investigated, to show that charging could occur in 60 to 160 min. In [15],
21 concentric tubes, with varying lengths and radii for the PCM zone, were tested together with three different PCM
22 (H325, H425 and H525). Whatever the PCM, the charging (respectively discharging) time was shown to vary up to
23 three (resp. 2) times depending on the conditions, while the maximum variation between the various PCM could go up
24 to six in both modes. In [19], a prototype of a 140 L shell-and-tube storage permitted to measure a 5 kW power charge
25 and 2-3.5 kW power discharge with RT70, and a 3.55 kW power charge and 2-3 kW power discharge with $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$.
26 Lastly, an interesting concept of a new fin design, for a shell containing extended finned tubes surrounded by NaNO_3
27 as PCM, was simulated in [20]. The discharge thermal power per tube length varied from 3 to 10 kWm^{-1} , to fulfill the 15
28 min constraint initially set in their project.

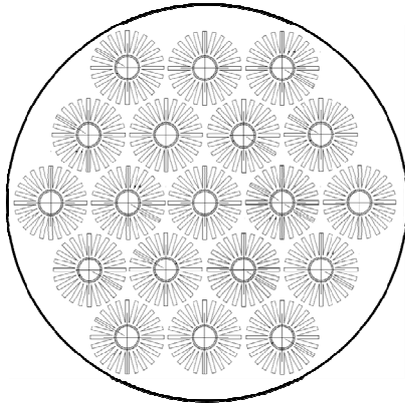
30 1.2 Novelty of the study

31 This work presents the development of a reduced multi-scale modelling approach for the charge of a steam storage
32 module involving a PCM. Similarly to what was proposed by Mostafavi Tehrani *et al.* [21], a heat transfer correlation
33 during the melting is deduced on the PCM side from fine 3D CFD computations. Indeed, as it is judiciously highlighted
34 in [sec. 1.2 p. 4, 21], prior correlations too often rely on experimental fitting. The contributions of this paper are
35 threefold. First, it proposes to continue apace the development of such a methodology since it has the potential to
36 avoid any experimental tuning and to overcome the limits of corresponding methods. In fact, the implicit goal is to
37 further improve the knowledge concerning heat transfer correlations for solid/liquid phase change, since it still lags far
38 behind what can be found for liquid/gas phase change, e.g. [33]. Secondly, to the authors' knowledge, this is the first
39 time ever that such a method is used for multi-tubes with fins and inserts, and for a dual two-phase storage with
40 liquid/gas HTF inside the tubes and solid/liquid PCM outside. Thirdly, a local correlation is built, and not an average
41 one. Since phase changes, from both the HTF and PCM, can occupy only part of the tube's length, the heat transfer is
42 indeed far from being homogeneous. Here again, this is the first time that such a procedure is considered.

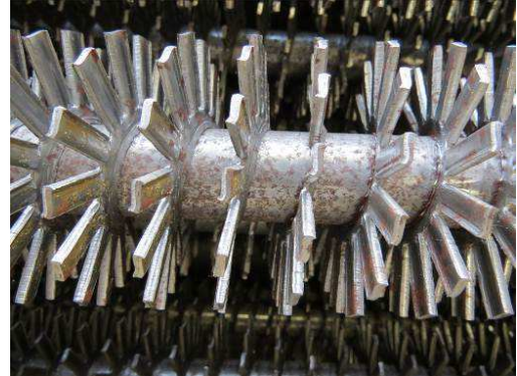
43 2 Methodology

44 As stated above, the current situation deals with a prototype shell-and-tube latent heat steam storage module
45 available at CEA Grenoble [93, 95]. This module is made of a cylindrical shell, containing a bundle of 19 finned tubes
46 (see Figure 1-(a)), with melded rectangular fins following a helical pattern (see Figure 1-(b)). This storage design does
47 not include any inserts. Both tubes and fins are made of carbon steel SA334 Grade 6. To be cost competitive, storage

1 modules are designed using off-the-shelf standard fins. The closer the tubes or the higher the heat exchange surface of
 2 the fins, the higher the thermal performances of the storage system and the higher its cost: consequently, the spacing
 3 between the tubes is logically a trade-off between system performances and costs. The HTF flows inside the tubes,
 4 whereas the shell is filled with approximately 1700 kg of sodium nitrate (NaNO_3). The immersed tube length is 4.08 m
 5 when the PCM is fully liquid.



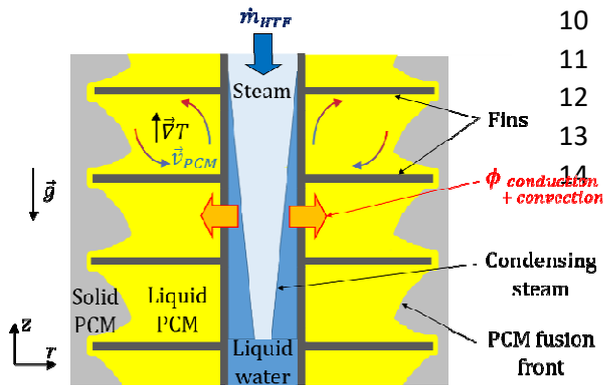
(a) tubes bundle distribution (black circle represents the outer shell of the module)



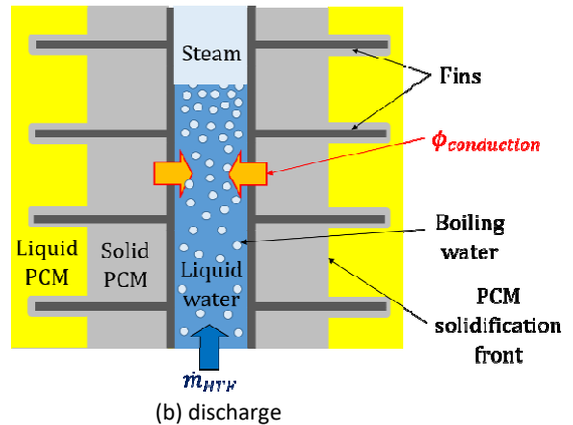
(b) side photograph of the finned tubes

Figure 1: Description of the prototype storage module

7 Schematically, the present problem is summarized in Figure 2. The two-phase HTF (liquid water / steam) and the PCM
 8 (solid / liquid) exchange heat and three thermal resistances can be identified, which need to be assessed. First, a
 9 convective heat exchange takes place between the flowing HTF and the inner tube wall.



(a) charge



(b) discharge

Figure 2 : Schematic view of the physical phenomena occurring in a shell-and-tube steam storage system based on LHS

19 Firstly, a 3D CFD model is used to simulate the phase transition of the PCM, in presence of conductive and convective
 20 heat transfer. Then, several simulations are performed to build a non-dimensional correlation for the heat transfer,
 21 considering both the presence of fins and the phase change. Following the classic approach for non-reactive
 22 convective-only flows, this correlation is searched under the following form [22], [23]:

$$\overline{Nu} = C Ra^n \quad (1)$$

23 Here, the Rayleigh and Nusselt numbers need to be properly defined, since their expression is not unequivocal [24],
 24 [16], [17], [25], [21], [26]:

$$Ra = g \left(\frac{\beta}{\nu D} \right)_{PCM} (T_w - \bar{T}_{PCM}) l_c^3 \quad (2)$$

$$Nu = \frac{\phi}{S_e (T_w - \bar{T}_{PCM})} \frac{l_c}{\lambda_{PCM,l}} \quad (3)$$

1 There are indeed various possible definitions for the characteristic length l_c and the mean PCM temperature \bar{T}_{PCM} .
 2 Besides, these definitions must be local since a local correlation is searched for; so, they should involve the altitude
 3 inside the tube. Several possibilities will be discussed and compared, as shown in section 3.2.

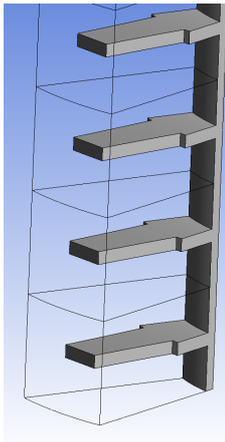
4 Secondly, a reduced system model is built to represent the whole dual two-phase storage module. The goal being to
 5 develop a fast and stand-alone tool, the outside of the tube (PCM, fins, inserts if any) is approximated as a
 6 homogeneous material, and natural convection inside the PCM is handled through an effective thermal conductivity.
 7 Practically, this latter is based on the previous Nusselt correlation, built from prior numerical experiments with the
 8 previous CFD model.

9 Eventually, the proposed methodology was then used for the prototype shell-and-tube latent heat steam storage
 10 module presented in Figure 1, to carry out three simulations of charging mode.

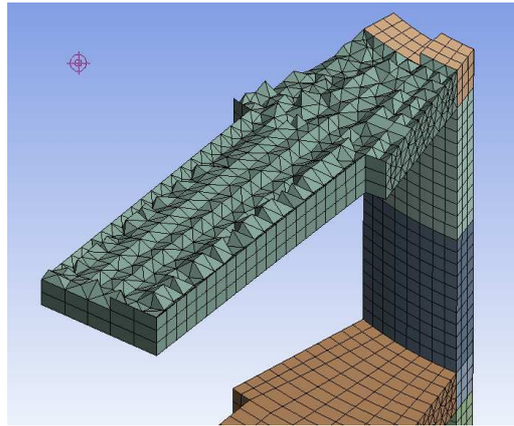
11 3 Determination of the heat transfer correlation

12 3.1 CFD modelling

13 In the present calculations, for the sake of limiting the computational time, only an angular sector of 22.5° and a tube
 14 length of 0.54 m (27 fins) are represented (see Figure 3-(a)). For the associated boundary conditions, a fixed
 15 temperature is set at the inner tube wall and a symmetry condition for the lateral and outer boundaries, while a wall
 16 condition is imposed for the others. The entire mesh is composed of 539 046 cells, whose mean size is 0.83 mm, and
 17 its maximum skewness is 0.81 (see Figure 3-(b)). Grid independence was verified by comparing punctual PCM
 18 temperature evolutions at several distances from the external tube wall to those obtained with a finer mesh
 19 containing 774 098 cells, for a mean size of 0.63 mm. The coarser mesh was found to give acceptable results, and
 20 furthermore, its features are similar to those used in [14-15]. The time step adopted is 0.025 s or 0.05 s, depending on
 21 the simulations, and, here again, the independence of the results on the time step was checked by dividing it regularly
 22 by 2.



23 a) fraction of the prototype simulated



(b) details of the mesh in the wall and fins

Figure 3 : Description of the geometry and mesh retained

24 Next, the PCM is modelled using the enthalpy-porosity approach, described by Voller and Prakash [30], which assumes
 25 an incompressible regime and uses the Boussinesq approximation to account for the buoyancy forces. The balance
 26 equations for mass, momentum and energy conservations are written:

$$\nabla \cdot \vec{U} = 0 \quad (4)$$

$$\rho \left(\frac{d\vec{U}}{dt} + \vec{U} \cdot \nabla \vec{U} \right) = -\vec{\nabla} P + \vec{\nabla}(\mu \nabla \vec{U}) - \rho \beta (T - T_{ref}) \vec{g} + A \vec{U} \quad (5)$$

$$\rho c_p \left(\frac{dT}{dt} + \vec{U} \cdot \nabla T \right) = \vec{\nabla}(\lambda \nabla T) - \rho L \left(\frac{dY_l}{dt} + \vec{U} \cdot \nabla Y_l \right) \quad (6)$$

1 The porosity approach consists in adding a damping term at the end of the right-hand side of the momentum
2 equation, whose role is to cancel the velocity field in solid phase. It expresses:

$$A = -A_{mush} \frac{(1 - Y_l)^2}{Y_l^3 + b} \quad (7)$$

3 with $b = 10^{-3}$.

4 In the tube wall and fins, only the energy equation applies:

$$\rho c_p \frac{\partial T}{\partial t} = \lambda \Delta T \quad (8)$$

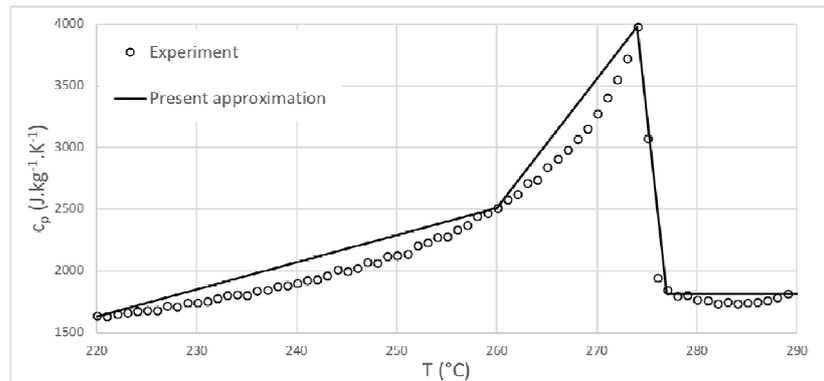
5 The physical properties of the PCM and the steel are given in Table 1, and in Figure 4 and Figure 5. The value of the
6 mushy zone constant was set to $A_{mush} = 10^8 \text{ kg.m}^{-3}.\text{s}^{-1}$, using a prior experiment where the temperature field was
7 compared with corresponding measurements [93, 94]. For the specific heat capacities and the thermal conductivities,
8 linear regressions were performed.

9

Table 1: Material properties for the CFD model

| Material | NaNO ₃ [9], [10],[31],[32] | Steel |
|---|---|-------|
| $\rho \text{ (kg.m}^{-3}\text{)}$ | 1927 | 7764 |
| $\beta \text{ (K}^{-1}\text{)}$ | $3.8 \cdot 10^{-4}$ | -- |
| $c_p \text{ (J.kg}^{-1}.\text{K}^{-1}\text{)}$ | Variable solid liquid 1704 | 542.8 |
| $\lambda \text{ (W.m}^{-1}.\text{K}^{-1}\text{)}$ | Variable solid liquid Variable | 50.33 |
| $\mu \text{ (Pa.s)}$ | $2.8 \cdot 10^{-3}$ | -- |
| $L \text{ (kJ.kg}^{-1}\text{)}$ | 173.3 | -- |
| $T_{sol} \text{ (}^\circ\text{C)}$ | 303.3 | -- |
| $T_{liq} \text{ (}^\circ\text{C)}$ | 306.6 | -- |

10



11

12

Figure 4 : Specific heat capacity of NaNO₃

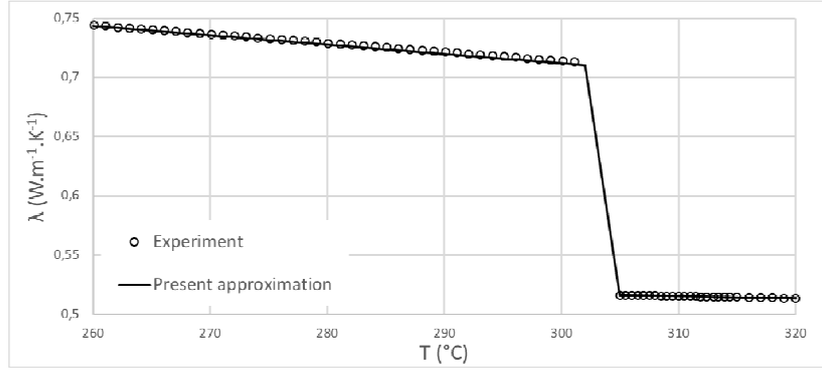


Figure 5 : Thermal conductivity of NaNO_3

The numerical experiments were performed with ANSYS Fluent® v. 17.2 software. Due to the incompressible feature of the partial differential equations system (4-6), a pressure-based formulation was chosen. Then, the PISO scheme was adopted for the pressure-velocity coupling together with, respectively, a second order upwind scheme for the convective terms in the momentum and energy equations and a least-square cell-based scheme for the diffusive terms. Finally, the pressure equation was discretized with the PRESTO! approach; for the residuals' convergence, relative decrease values of 10^{-3} were taken for the continuity and momentum equations but the energy residual was set to 10^{-11} [91].

3.2 Non-dimensional relation

Three CFD computations were performed with an imposed temperature $T_w = 310, 318$ and 328°C respectively at the inner tube wall. The PCM is initially solid at 290°C . The meshed domain was then divided into 27 *representative volumes*, centered on the individual fins (the boundaries between these volumes can be seen on Figure 3-(a)). Regularly, the heat flow-rate entering the PCM was recorded as a function of time, along with associated physical variables, e.g. temperature field, molten fraction...

From these values, expressions were investigated for the characteristic length $\overline{l_c}$ and the mean PCM temperature $\overline{T_{PCM}}$ involved in Eqs. (2-3). Thus, four and three different relations were tested for $\overline{l_c}$ and $\overline{T_{PCM}}$ respectively:

- Characteristic length $\overline{l_c}$
 1. Altitude, taken at the center of the representative volume considered:

$$\overline{l_c} = z_i \quad (9)$$

2. Thickness of the annular space outside the tube:

$$\overline{l_c} = R_{ext} - R_{text} \quad (10)$$

3. Local thickness of the liquid PCM layer:

$$\overline{l_c} = -R_{ext} + \sqrt{R_{ext}^2 + (R_{text}^2 - R_{ext}^2) \overline{Y_{l,i}}} \quad (11)$$

where $\overline{Y_{l,i}}$ is the mean mass liquid fraction in the representative volume considered:

$$\overline{Y_l} = \frac{\sum_{i=1}^{nb \text{ PCM cells}} \rho_i Y_{l,i} V_i}{\sum_{i=1}^{nb \text{ PCM cells}} \rho_i V_i} \quad (12)$$

4. Present complete thickness of the liquid PCM layer on the whole tube height:

$$\overline{l_c} = -R_{ext} + \sqrt{R_{ext}^2 + (R_{text}^2 - R_{ext}^2) \overline{Y_l}} \quad (13)$$

where \overline{Y}_l is computed in the same way as $\overline{Y}_{l,i}$, but by averaging the liquid fraction on the whole computational domain, and not only on the representative volume considered.

- Mean PCM temperature \overline{T}_{PCM}
 1. Phase change temperature:

$$\overline{T}_{PCM} = T_{pc} = \frac{1}{2}(T_{sol} + T_{liq}) \quad (14)$$

2. Mean temperature of the liquid PCM layer in the representative volume considered:

$$\overline{T}_{PCM} = \frac{\sum_{i=1}^{nb \text{ PCM cells}} c_{p,i} T_i Y_{l,i} V_i}{\sum_{i=1}^{nb \text{ PCM cells}} c_{p,i} Y_{l,i} V_i} \quad (15)$$

3. Temperature of the first mesh cells near the outer tube wall. It is computed as the mean temperature of six PCM cells and one fine mesh cell near the wall, as shown on Figure 6.

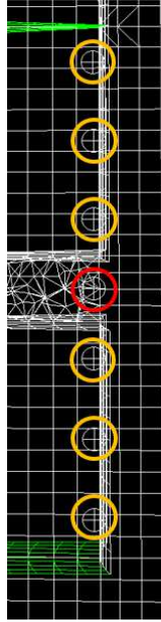
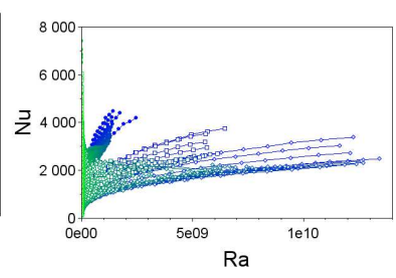
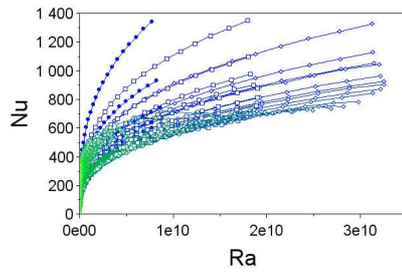
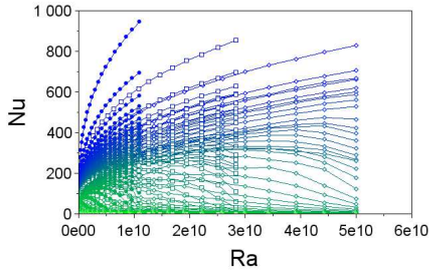
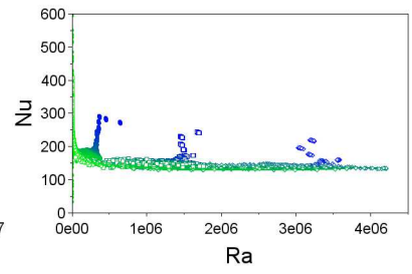
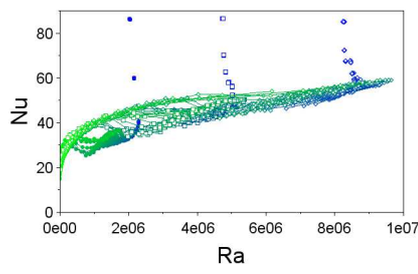
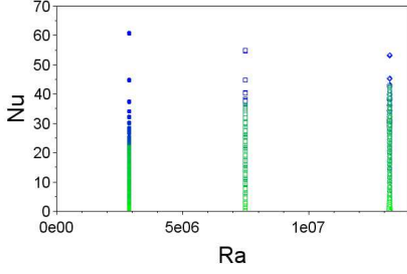


Figure 6: Mesh cells near the outer wall of the inner tube (yellow: PCM cells; red: fin cells)

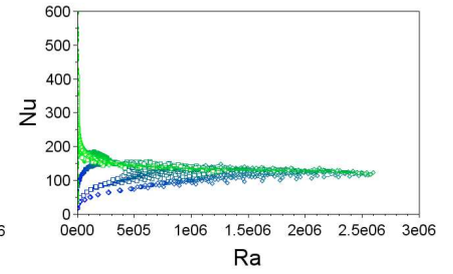
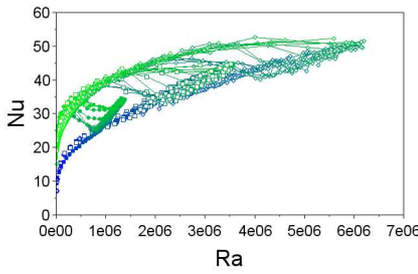
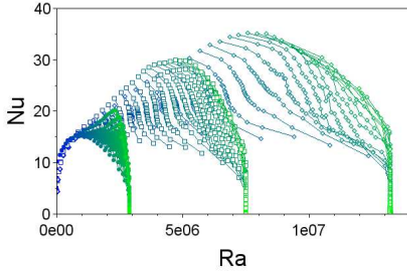
This enabled 12 scatter plots to be drawn, to assess whether or not the above expressions allow the Rayleigh and Nusselt numbers to correlate. The corresponding graphs are shown in Figure 7. From the results available on Figure 7, one can easily note that some definitions for the characteristic length \overline{l}_c and the mean PCM temperature \overline{T}_{PCM} do not lead to a good correlation between the Rayleigh and Nusselt numbers. This is especially the case for Eq. (14), whatever the definition of \overline{l}_c , and similarly for Eq. (9), whatever the definition of \overline{T}_{PCM} . For the six remaining candidates, two behaviors can be observed. Either the points gather relatively well around a single curve, except at the beginning of melting, or two curves are to be found (one for the lowest Nusselt numbers at the beginning of melting – in blue –, and then another one for the highest Nusselt numbers at the end of melting – in green –). Given that the heat transfer should go from a mainly conductive regime to a convection-dominated one, it is preferable to select the latter case. Therefore, one can use either Eqs. (11) or (13) for the characteristic length \overline{l}_c and the mean PCM temperature \overline{T}_{PCM} should be taken as that of the liquid layer Eq. (15) at the given height.



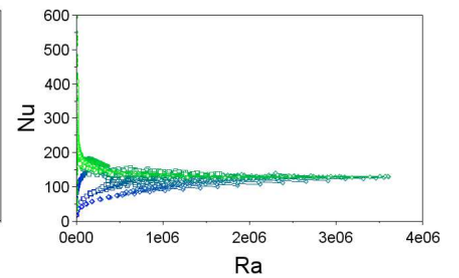
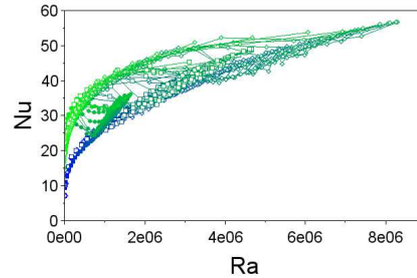
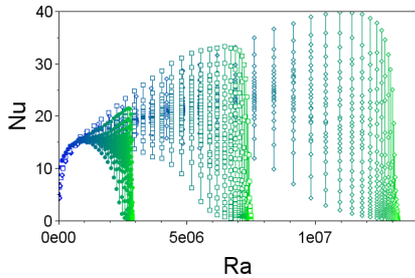
(a) Characteristic length given by Eq. (9)



(b) Characteristic length given by Eq. (10)



(c) Characteristic length given by Eq. (11)



(d) Characteristic length given by Eq. (13)

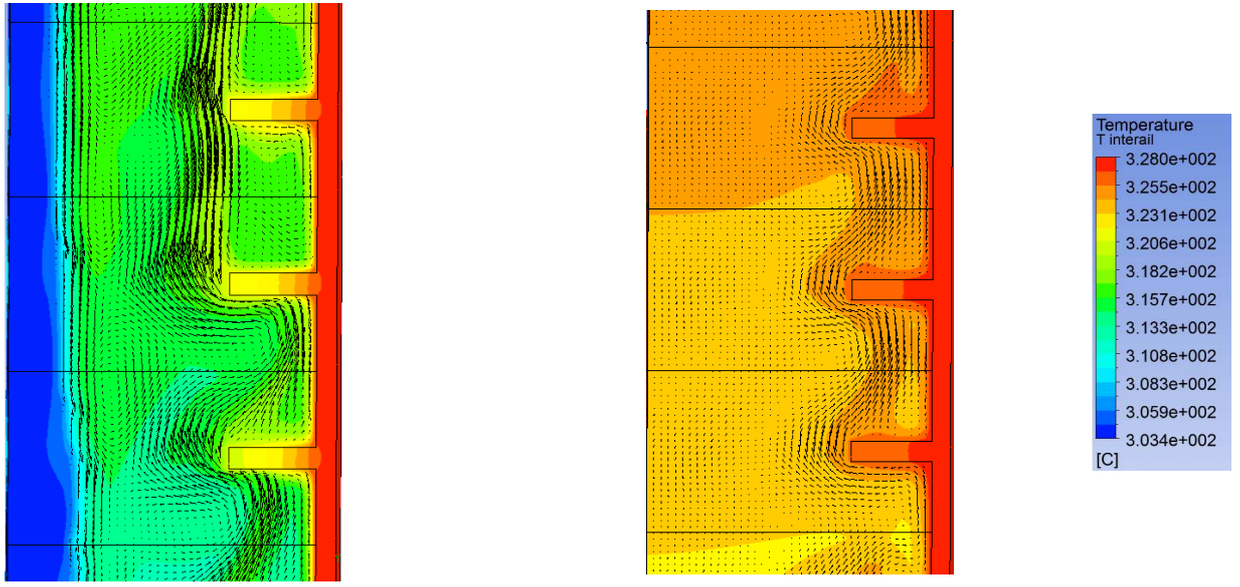
Figure 7: Accuracy of the correlation between the Rayleigh and Nusselt numbers depending on the definitions taken for $\sqrt{V_c}$ and $\sqrt{V_{PCM}}$: phase change temperature (left), mean liquid temperature at the considered height (middle), mean temperature near the wall (right)

To further highlight this difference between low and high Nusselt numbers, for a given similar Rayleigh number, it is proposed to look at the temperature and velocity fields at a time when the 27 representative volumes are split between both curves. Figure 8 presents the corresponding results in these two zones, in a vertical plane situated in the middle of the domain. Depending on the vertical position in the tube, the Nusselt number can either undergo small values, corresponding to a velocity maximum at $17 \text{ mm}\cdot\text{s}^{-1}$ (see Figure 8.a), or high values, for an equivalent maximum velocity at $9 \text{ mm}\cdot\text{s}^{-1}$ (see Figure 8.b).

Then, it can be noted that melting is still ongoing in volumes whose (Ra, Nu) values gather around the first curve, whereas melting of the PCM is complete in volumes whose (Ra, Nu) couples follow the second curve. Besides, one can remark the highest velocity magnitudes in the first case. This is due to the higher temperature gradient inside the PCM, between the phase front remaining at the melting temperature and the inner wall (also at a fixed temperature).

1 When the PCM is molten, heat transfer is singularly reduced since this temperature gradient is reduced. Finally, these
 2 two situations just entail two different regimes of convective heat transfer, logically associated with two different
 3 scaling laws. Last but not least, it was verified that points lying between these two asymptotic behaviors show liquid
 4 fraction values between 0 and 1 at the external boundary of the domain.

5



(a) lower portion of the tube with ongoing melting

(b) higher portion of the tube where fusion is over (only liquid phase here)

Figure 8: Temperature and velocity fields at $t = 2390$ s for the case $T_w = 328^\circ\text{C}$

6

7 In summary, using Eqs. (11) or (13) and Eq. (15) for, respectively, the characteristic length $\overline{L_c}$ and the mean PCM
 8 temperature $\overline{T_{PCM}}$ gives to obtain a scaling law between the Nusselt and Rayleigh numbers. Depending on the time
 9 and the altitude, that is to say depending on the completeness of the phase change, this scaling law will be bounded
 10 by two asymptotic behaviors. In this respect, a local heat transfer correlation will be preferred for Eq. (1) and,
 11 consequently, final choices will rely on Eqs. (11) or (15). It leads to the following scaling laws for the two limiting cases:

$$\overline{Nu} = 0.402 Ra^{0.306} \tag{16}$$

$$\overline{Nu} = 2.614 Ra^{0.196} \tag{17}$$

12 As can be seen on Figure 9, a very good agreement with the numerical results is observed.

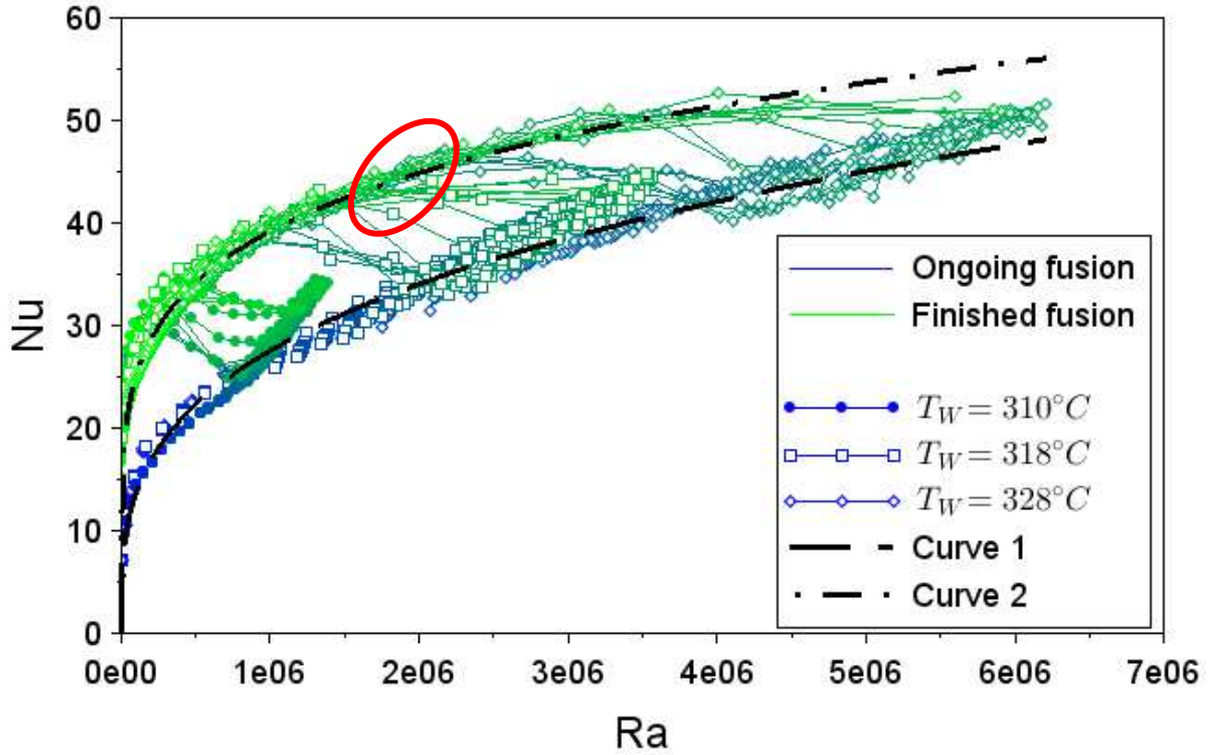


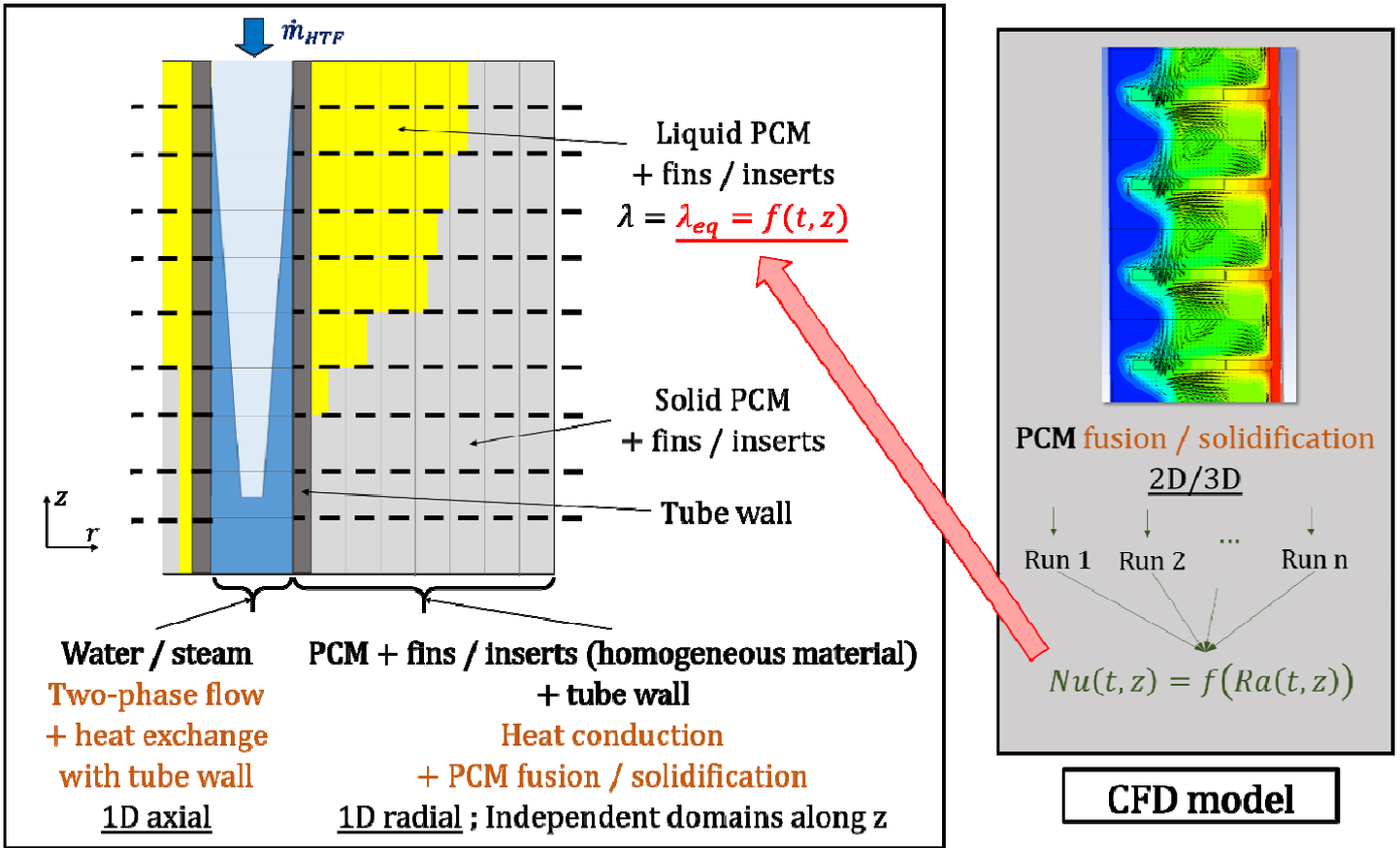
Figure 9: Comparisons of the (Ra, Nu) scatter plot obtained from the CFD results, using Eqs. (11) and (15) for the definitions of l_c and T_{PCM} , with correlations (16) and (17) for the heat transfer

From a practical point of view, one will need a criterion to know when to go from one expression to the other. Indeed, a continuous formulation is required since the real physical heat transfer is not discontinuous and, second, because it is more suitable in a reduced model to avoid oscillations and unstable behaviors. As the proper definition of this threshold is beyond the scope of the present article, the following definition was chosen:

$$\begin{cases} Nu_i = 0.402 Ra_i^{0.306} & 0 < \overline{Y}_{l,i} < 0.98 \\ Nu_i = \frac{\overline{Y}_{l,i} - 0.98}{0.02} (2.614 Ra_i^{0.196}) + \frac{1 - \overline{Y}_{l,i}}{0.02} (0.402 Ra_i^{0.306}) & 0.98 < \overline{Y}_{l,i} < 1 \\ Nu_i = 2.614 Ra_i^{0.196} & \overline{Y}_{l,i} = 1 \end{cases} \quad (18)$$

4 Application

As mentioned in Section 2, the idea is now to use the Nusselt correlation thus obtained to build an effective thermal conductivity that can be implemented in the reduced model, as summarized in Figure 10. The proposed methodology will be tested on the three previous fusion experiments.



System model

Figure 10 : Structure and characteristics of the proposed multi-scale model

4.1 System modelling

The present model will stand for the two-phase HTF inside the tube, for the tube, and for the fins, inserts and PCM approximated together as a homogeneous material. For each of these materials, 1D and 2D discretization are adopted respectively.

On the HTF side, the flow has been shown to be in forced regime [10]. Moreover, the liquid and vapor phases are assumed to be in thermal and dynamical equilibrium, *i.e.* with the same local velocity and temperature, and axial conduction is neglected. Therefore, the PDE system for the HTF, incorporating directly the wall boundary condition in the energy equation [28], is written:

$$S_{tint} \frac{\partial \rho_{HTF}}{\partial t} + \frac{\partial \dot{m}}{\partial z} = 0 \tag{19}$$

$$S_{tint} \rho_{HTF} \frac{\partial h_{HTF}}{\partial t} + \frac{\partial (\dot{m} h_{HTF})}{\partial z} - c_{tint} \varphi_{tint} = 0 \tag{20}$$

To compute the HTF density, it is assumed that liquid and vapor phases are at saturation in the two-phase region:

$$\rho_{HTF} = \varepsilon \rho_{HTF,v}^{sat} + (1 - \varepsilon) \rho_{HTF,l}^{sat} \tag{21}$$

with

$$\varepsilon = \frac{1}{1 + \frac{1 - Y_{HTF,v}}{Y_{HTF,v}} \frac{\rho_{HTF,v}^{sat}}{\rho_{HTF,l}^{sat}}} \tag{22}$$

$$Y_v = \frac{h_{HTF} - h_{HTF,l}^{sat}}{h_{HTF,v}^{sat} - h_{HTF,l}^{sat}} \quad (23)$$

1 Finally, the heat flux at the wall is calculated by means of a classic thermal resistance:

$$\varphi_{tint} = \frac{T_t - T_{HTF}}{\frac{1}{\alpha_{HTF}} + \frac{R_{tint} \ln\left(\frac{1}{2} + \frac{R_{ttext}}{2 R_{tint}}\right)}{\lambda_t}} \quad (24)$$

2 Next, the tube simply obeys the heat equation. Following the thermal resistance analogy of Eq. (24), the energy
3 balance is written (using enthalpy for the sake of clarity with other PDE systems):

$$S_t \rho_t \frac{\partial h_t}{\partial t} = S_t \lambda_t \frac{\partial^2 T_p}{\partial z^2} + c_{ttext} \varphi_{ttext} - c_{tint} \varphi_{tint} \quad (25)$$

4 and logically, one gets for the outer heat flux:

$$\varphi_{ttext} = \frac{T_h - T_t}{R_{ttext} \left(\frac{\ln\left(\frac{2 R_{ttext}}{R_{tint} + R_{ttext}}\right)}{\lambda_t} + \frac{\ln\left(1 + \frac{\Delta r}{2 R_{ttext}}\right)}{\lambda_{h,eq}} \right)} \quad (26)$$

5 Last but not least, one needs to write the modelling of the homogeneous material. Since natural convection is
6 accounted for by means of an effective thermal conductivity, it only relies on the energy balance equation. To handle
7 the latent term, the previous source term approach [29] is still considered. For the sake of simplicity, the heat transfer
8 in the axial direction is neglected and thermal losses are also assumed to be negligible. Thus, the energy equation can
9 be written:

$$\rho_h \frac{\partial h_{h,sens}}{\partial t} = \lambda_h \left(\frac{\partial^2 T_h}{\partial r^2} + \frac{1}{r} \frac{\partial T_h}{\partial r} \right) - \rho_{PCM} x_{PCM} L \frac{\partial Y_{PCM,l}}{\partial t} \quad (27)$$

10 where the average properties are given by:

$$\rho_h = x_{PCM} \rho_{PCM} + (1 - x_{PCM}) \rho_{fin} \quad (28)$$

$$c_{p,h} = Y_{PCM} c_{p,PCM} + (1 - Y_{PCM}) c_{p,fin} \quad (29)$$

11 Then, the paramount parameter that needs to be further detailed is the effective thermal conductivity. It is generally
12 deduced from the Nusselt number, or equivalently, from the Rayleigh number. During melting, the situation
13 encountered can be represented as depicted in Figure 11: thermal conductivity of the homogeneous material
14 corresponds to the effective thermal conductivity λ_{eff} in the liquid molten layer and to the average thermal
15 conductivity in the external solid part.

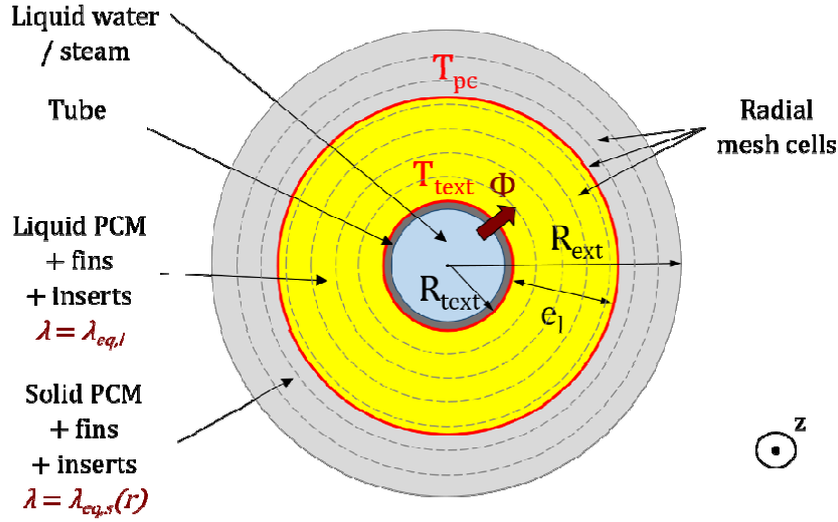


Figure 11: General case for the definition of the effective thermal conductivity

By definition, the effective thermal conductivity is such that the total amount of heat is equal to the heat exchanged in the convective case:

$$\phi = 2\pi\Delta z\lambda_{eq,l} \frac{(T_{text} - T_{pc})}{\ln\left(1 + \frac{e_l}{R_{text}}\right)} \quad (30)$$

To express the thickness of the liquid layer e_l , a mass balance is used, involving the mean mass liquid fraction at the considered altitude \bar{Y}_l :

$$e_l = -R_{text} + \sqrt{R_{text}^2 + (R_{ext}^2 - R_{text}^2)\bar{Y}_l} \quad (31)$$

$$\bar{Y}_l = \frac{\sum_{j=1}^{nb \text{ mesh cells, sub-domain}} Y_{l,j} \rho_{PCM} x_v V_j}{\sum_{j=1}^{nb \text{ mesh cells, sub-domain}} \rho_{PCM} x_v V_j} \quad (32)$$

In (32), V_j represents the volume of the j -th cell in the radial direction.

Then, at a given altitude z along the tube, the local Nusselt and Rayleigh numbers are expressed as follows:

$$Nu(z) = \frac{\phi}{2\pi R_{text} \Delta z (T_{text}(z) - T_{PCM}(z)) \lambda_{PCM,l}} \frac{l_c(z)}{l_c(z)} \quad (33)$$

$$Ra(z) = g \left(\frac{\beta}{\nu D} \right)_{PCM} (T_{text}(z) - T_{PCM}(z)) l_c(z)^3 \quad (34)$$

By combining equations (30), (31), (33) and (34), one gets:

$$\lambda_{eq,l}(z) = \lambda_{PCM,l} \frac{T_{text}(z) - T_{PCM}(z)}{T_{text}(z) - T_{pc}} \frac{R_{text}}{l_c(z)} \times \ln \left(1 + \frac{-R_{text} + \sqrt{R_{ext}^2 + (R_{ext}^2 - R_{text}^2)\bar{Y}_l(z)}}{R_{text}} \right) \times f \left(g \left(\frac{\beta}{\nu D} \right)_{PCM} (T_{text}(z) - T_{PCM}(z)) l_c(z)^3 \right) \quad (35)$$

In the solid part, a classic mean calculation holds:

$$\lambda_{eq,s,j} = Y_{PCM,s,j} \lambda_{PCM,s} + (1 - Y_{PCM,s,j}) \lambda_{fin} \quad (36)$$

1 Finally, in each cell, the equivalent thermal conductivity $\lambda_{eq,j}$ is computed with a weighted sum between the effective
 2 thermal conductivity in the liquid layer and the homogeneous equivalent thermal conductivity in the solid layer:

$$\lambda_{eq,j} = Y_{l,j}\lambda_{eq,l} + (1 - Y_{l,j})\lambda_{eq,s,j} \quad (37)$$

3 The values of all the other variables, and of the operational parameters, are provided in Table 2.

4 *Table 2: Material and operational properties for the system model*

| Geometrical configuration | |
|----------------------------------|---|
| Total PCM volume fraction | 0.936 |
| Total PCM mass fraction | 0.785 |
| Material properties | |
| ρ_h | 2299 kg.m ⁻³ |
| $C_{p,h}$ | solid liquid |
| | 1540 J.kg ⁻¹ .K ⁻¹ 1454 J.kg ⁻¹ .K ⁻¹ |
| λ_{PCM} | solid liquid |
| | 0.720 W.m ⁻¹ .K ⁻¹ 0.515 W.m ⁻¹ .K ⁻¹ |
| $\lambda_{eq,s}$ | |
| | $R_{text} < r < R_{text} + 10$ mm $R_{text} + 10$ mm $< r < R_{text} + 30$ mm $r > R_{text} + 30$ mm |
| | 6.92 W.m ⁻¹ .K ⁻¹ Variable (bounded values: 5.61 – 3.48 W.m ⁻¹ .K ⁻¹) 0.720 W.m ⁻¹ .K ⁻¹ |
| Mesh properties | |
| Number of cells | axial direction radial direction total |
| | 10 9 110 |

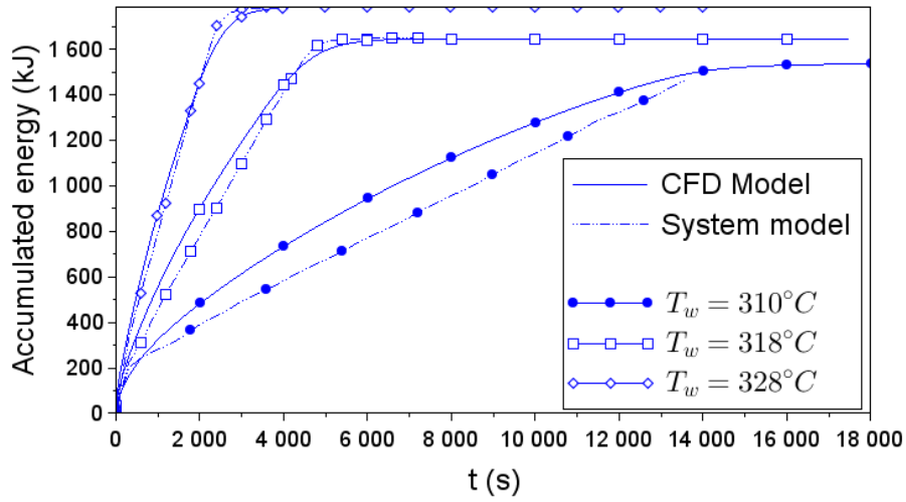
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6 4.2 Results

7 The present computations deal with the same test cases as the ones presented in Section 3. In the current situation,
 8 the space steps are $\Delta z = 54.0$ mm and $\Delta r = 1.85$ mm. The HTF pressure is chosen so that the saturation temperature is
 9 equal to the desired wall temperature, and accordingly, the internal heat transfer coefficient is set to $\alpha_{int} = 10^6$ W.m⁻².K⁻¹.
 10 Water vapor, with a slight superheating of 0.1°C, is injected at the top with a 0.05 kg.s⁻¹ mass flow-rate.

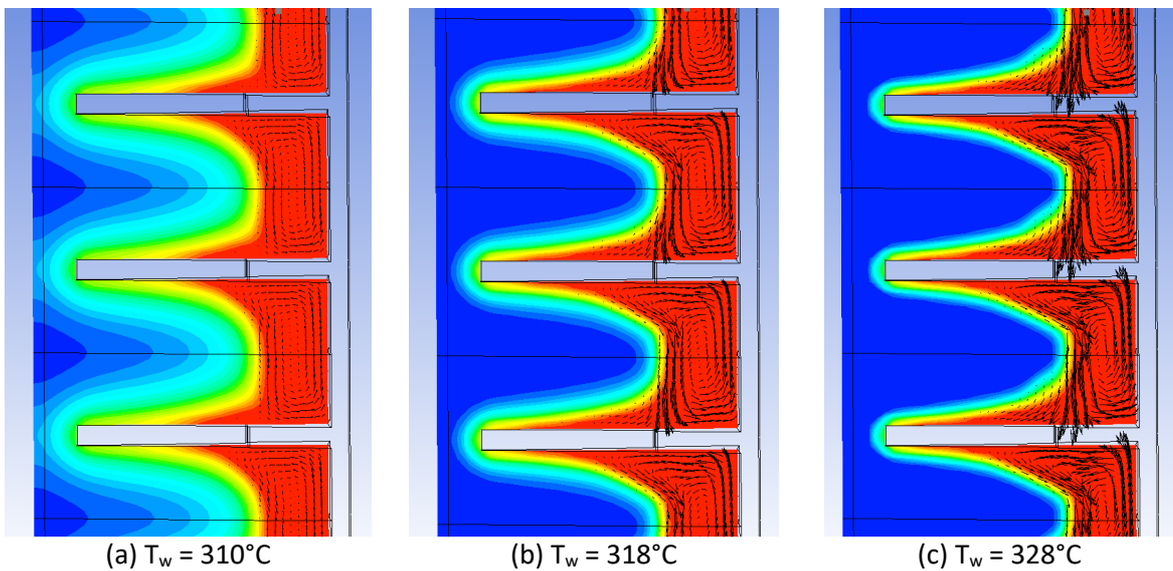
11 The whole system is solved with an in-house code, in Scilab language. For Eqs. (19), (20), (25) and (27), a classic finite
 12 volume method is used. The temporal integration is fully implicit. The numerical resolution is done separately for the
 13 HTF on the one hand, and for the tube thickness and the homogeneous material on the other hand. For these two
 14 blocks, the convergence criterion is based on the variation of the solved quantities from one iteration to another. On
 15 the HTF side, convergence is reached when the relative variation of mass enthalpy and mass flow in each mesh cell is
 16 below 10⁻¹². For the second block, the total mass enthalpy is normalized by its initial value, and convergence is
 17 assumed to be reached when the variation of this normalized mass enthalpy is below 10⁻⁸.

18 The main focus is very much on the ability to reproduce the storage behavior, that is to say to represent the variation
 19 of energy inside the module. Therefore, the temporal evolution of the energy accumulated by the PCM is represented
 20 on Figure 12 for the three cases, for both the CFD model and the reduced model. Overall, the agreement between the
 21 system model and the CFD results is good: the mean discrepancy between the two models is 4.8% for $T_w = 328^\circ\text{C}$, 6.8%
 22 for $T_w = 318^\circ\text{C}$, and 14% for $T_w = 310^\circ\text{C}$. In the latter case, it is important to mention that this discrepancy probably
 23 comes from the intrinsic limit of the Nusselt correlation that fundamentally holds only for the liquid region. However,
 24 as can be seen in Figure 13, the mushy region is far more pronounced for the lower HTF temperature. As a
 25 consequence, the method is here less reliable.



1
2
3

Figure 12: Comparisons of the energy accumulated in the PCM, the tube and the fins during melting at a constant and uniform wall temperature, for the CFD and system models



4
5

Figure 13: Liquid fraction and velocity fields in a vertical plane for the three CFD computations

6
7
8

Shapes of the phase front are presented in Figure 14 at two different instants for the case $T_w = 328^\circ\text{C}$. Though the initial CFD results logically match the fins' shape, the mean front position predicted by the system model is in good agreement with these results. This evidences the capability of the reduced model to reproduce the melting rate.

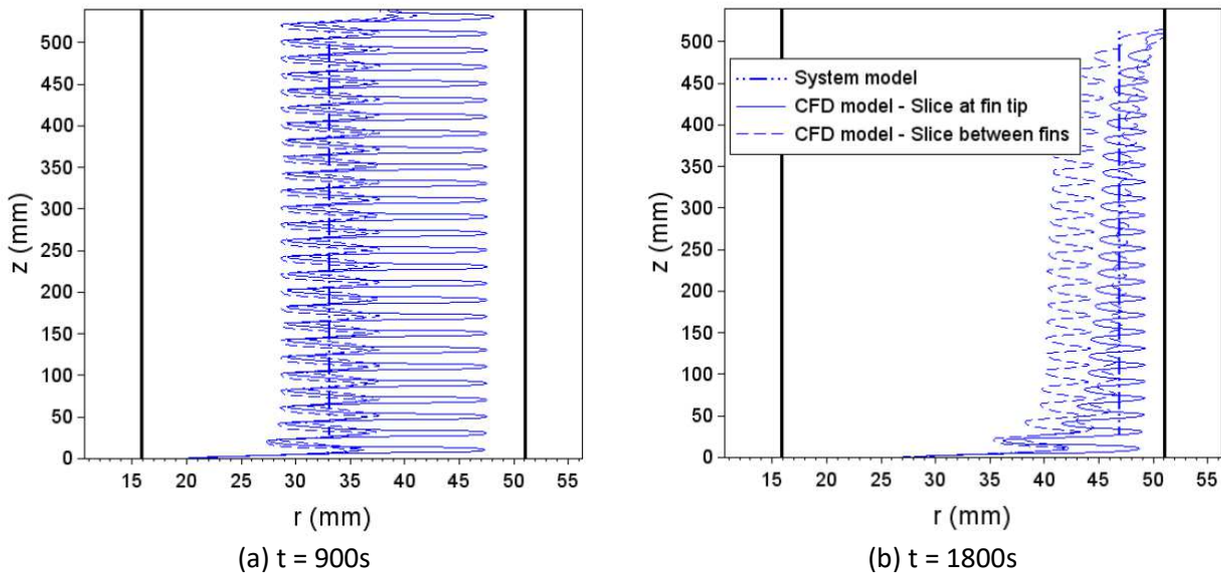


Figure 14: Comparisons of the phase fronts predicted by the CFD and system models for $T_w = 328^\circ\text{C}$ (black lines represent the boundaries of the PCM domain. Left: outer tube wall. Right: external domain boundary.)

All in all, the overall agreement between the reduced model and the complete CFD model is relatively good, and these results are very promising. Finally, to further highlight the assets of the present method, comparisons of the computational times needed to obtain the results are presented in Table 3. It is straightforward to notice that a considerable reduction can be achieved, resulting in 10 to 40 times lower CPU costs.

Table 3: Computational times for the CFD model and the reduced system model

| | $T_w = 310^\circ\text{C}$ | $T_w = 318^\circ\text{C}$ | $T_w = 328^\circ\text{C}$ |
|--------------|----------------------------|---------------------------|---------------------------|
| CFD model | 289 h (≈ 12 days) | 132 h (5.5 days) | 158 h (≈ 7 days) |
| System model | 29 h | 8 h | 4 h |

5 Conclusion

A reduced model has been built to carry out fast and accurate simulations of a dual two-phase storage of steam, composed of a shell containing a PCM crossed by several finned tubes containing a liquid/vapor heat transfer fluid. The corresponding system is based on an effective thermal conductivity which is determined using a heat transfer correlation with specific non-dimensional numbers. This correlation mainly involves the Nusselt and Rayleigh numbers which are linked from previous simulations of the phase change material's behavior, obtained with fine 3D computational fluid dynamics calculations. A specific study was made to determine the best suited definitions for the characteristic length and the mean PCM temperature involved in the Nusselt and Rayleigh numbers. In summary, the present method proposes to avoid the usual tuning of endogenous parameters; instead, a Nusselt correlation is determined and then used directly in the system model without any other *a posteriori* calibration.

When comparing the results of the two approaches, the agreement obtained is rather good for the three test cases considered (three different temperatures of the inner tube wall $T_w = 310, 318$ and 328°C with the PCM initially solid at 290°C). The evolution of the accumulated energy over time is predicted by the system model with an average deviation of 4 to 14% from the CFD results. The time required for the simulation of a melting is 10 to 40 times less for the reduced model than for the CFD model.

Thus, the current method appears to be very promising, making it possible to consider future easy and fast designs of latent heat storage systems, and/or multiple sensitivity analysis that would be too prohibitive otherwise. In the near future, it is planned to consolidate first the determination of the Nusselt correlation. Then, the reduced model will be further developed to remove its basic assumptions, that can easily be avoided; at the same time, it will be validated on numerous test cases in order to bound its precision, together with the savings in computational cost.

1 Last but not least, an important prospect will be the development of a similar multi-scale methodology for steam
2 discharge, involving PCM solidification, and boiling of the heat transfer fluid. During solidification in a PCM heat
3 storage module, the heat transfers are generally mainly conductive. The reduced model will have to be adapted.
4

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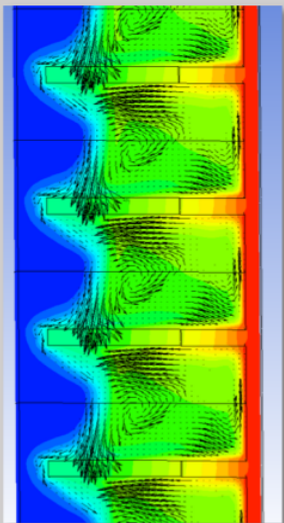
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PCM

Fusion / solidification

$$Nu(t, z) = f(Ra(t, z))$$

CFD Model

Liquid PCM
+ fins / inserts

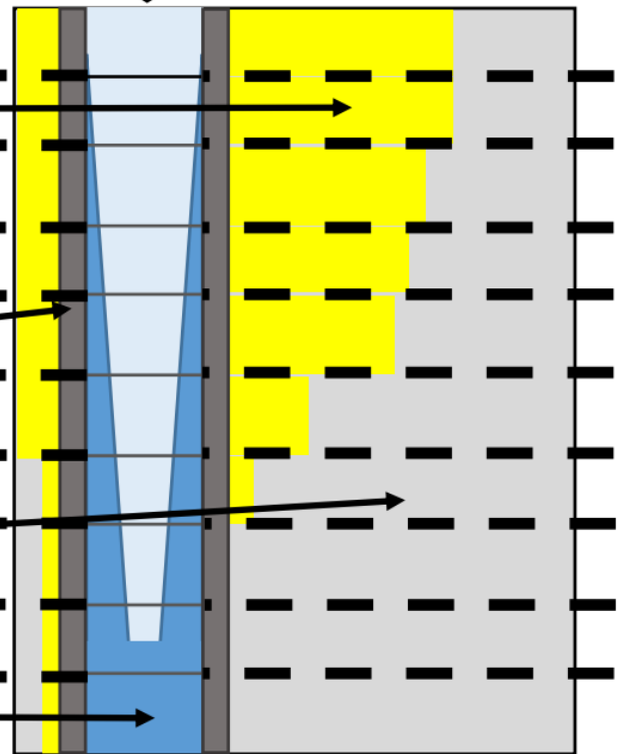
$$\lambda_{eq} = f(t, z)$$

Tube wall

Solid PCM +
fins / inserts

Water / steam

\dot{m}_{HTF}



System Model