MASTER THESIS

High temperature thermal energy storage systems based on latent and thermo-chemical heat storage

Under the direction of

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and

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Submitted in the Technischen Universität Wien

Faculty of Mechanical and Industrial Engineering

from

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ABSTRACT

The increasing of the population and development of the different countries converts the energy topic in one of the most important aspects of our times. The main target due to the limited life of conventional energy sources is the achievement of a sustainable energy mix where thermo solar energy plays an important role. One of the disadvantages of this renewable energy is the fact that energy is not available all the time: the need of heat storage systems appear. In this Master Thesis, a review on the work done until the moment in the frame of latent heat and thermochemical storage systems is presented in the temperature range from 200 to 700 ºC. Finally a design and the first calculations for the modeling of a latent heat storage system for a laboratory device are shown.
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<th>Acronym</th>
</tr>
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<tbody>
<tr>
<td>Australian National University</td>
<td>ANU</td>
</tr>
<tr>
<td>Compressed Expanded Graphite</td>
<td>CEG</td>
</tr>
<tr>
<td>Compact Linear Fresnel Reflector</td>
<td>CLFR</td>
</tr>
<tr>
<td>Cascaded Latent Heat Storage</td>
<td>CLHS</td>
</tr>
<tr>
<td>Concentrated Solar Power</td>
<td>CSP</td>
</tr>
<tr>
<td>German Aerospace Center</td>
<td>DLR</td>
</tr>
<tr>
<td>Direct Steam Generation</td>
<td>DSG</td>
</tr>
<tr>
<td>Expanded Natural Graphite</td>
<td>ENG</td>
</tr>
<tr>
<td>Expanded graphite powder</td>
<td>GFG</td>
</tr>
<tr>
<td>Heat Collector Reactor Element</td>
<td>HCRE</td>
</tr>
<tr>
<td>Heat Transfer Fluid</td>
<td>HTF</td>
</tr>
<tr>
<td>Internal Combustion Engine</td>
<td>ICE</td>
</tr>
<tr>
<td>Latent Heat Storage</td>
<td>LHS</td>
</tr>
<tr>
<td>Latent Heat Thermal Energy Storage</td>
<td>LHTES</td>
</tr>
<tr>
<td>Multi Tower Solar Array</td>
<td>MTSA</td>
</tr>
<tr>
<td>Natural Graphite</td>
<td>NG</td>
</tr>
<tr>
<td>Phase Change Material</td>
<td>PCM</td>
</tr>
<tr>
<td>Plataforma Solar de Almeria</td>
<td>PSA</td>
</tr>
<tr>
<td>Solar thermal electricity generating systems</td>
<td>SEGS</td>
</tr>
<tr>
<td>Sensible Heat Storage</td>
<td>SHS</td>
</tr>
<tr>
<td>Thermal Energy Storage</td>
<td>TES</td>
</tr>
<tr>
<td>Thermal Hybrid System</td>
<td>THS</td>
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NOMENCLATURE

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Unit [IS]</th>
<th>Significance</th>
</tr>
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<tbody>
<tr>
<td>Q</td>
<td>J</td>
<td>Energy accumulated</td>
</tr>
<tr>
<td>m</td>
<td>kg</td>
<td>PCM mass</td>
</tr>
<tr>
<td>$c_p$</td>
<td>J/kg</td>
<td>Specific heat</td>
</tr>
<tr>
<td>$a_m$</td>
<td></td>
<td>Fraction of melted PCM</td>
</tr>
<tr>
<td>$\Delta h_m, h_m$</td>
<td>J/kg</td>
<td>Latent heat of fusion</td>
</tr>
<tr>
<td>$T_i$</td>
<td>K</td>
<td>Initial temperature</td>
</tr>
<tr>
<td>$T_f$</td>
<td>K</td>
<td>Final temperature</td>
</tr>
<tr>
<td>$T_{m_0}, T_f$</td>
<td>K</td>
<td>Melting temperature</td>
</tr>
<tr>
<td>$\rho$</td>
<td>kg/m$^3$</td>
<td>Density</td>
</tr>
<tr>
<td>$k, \lambda$</td>
<td>W/(m·K)</td>
<td>Thermal conductivity</td>
</tr>
<tr>
<td>$X(t)$</td>
<td>m</td>
<td>Position of the solid-liquid interface</td>
</tr>
<tr>
<td>$T_0(t)$</td>
<td>K</td>
<td>Temperature of the infinite slab</td>
</tr>
<tr>
<td>$T(x,t)$</td>
<td>K</td>
<td>Temperature of the liquid region</td>
</tr>
<tr>
<td>$K_{ij}$</td>
<td>W/K</td>
<td>Conductivity between nodes i and j</td>
</tr>
<tr>
<td>$t$</td>
<td>s</td>
<td>Time, time needed for the melting process</td>
</tr>
</tbody>
</table>

NOMENCLATURE FROM ANNEXES

ANNEX A

<table>
<thead>
<tr>
<th>Symbol</th>
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<th>Significance</th>
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<tbody>
<tr>
<td>$eK$</td>
<td>K</td>
<td>Scale change from centigrads to Kelvin</td>
</tr>
<tr>
<td>$T_{dI}$</td>
<td>K</td>
<td>Temperature of the Dowtherm A at the beginning</td>
</tr>
<tr>
<td>$T_{dF}$</td>
<td>K</td>
<td>Temperature of the Dowtherm A at the end</td>
</tr>
<tr>
<td>$T_d$</td>
<td>K</td>
<td>Mean temperature of the Dowtherm A</td>
</tr>
<tr>
<td>$T_m$</td>
<td>K</td>
<td>Melting temperature</td>
</tr>
<tr>
<td>$T_{wall}$</td>
<td>K</td>
<td>Temperature of the inner wall of the tube</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Pa·s</td>
<td>Viscosity of the Dowtherm A</td>
</tr>
<tr>
<td>$c_p$</td>
<td>J/(kg·K)</td>
<td>Specific heat of the Dowtherm A</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>W/(m·K)</td>
<td>Thermal conductivity of Dowtherm A</td>
</tr>
<tr>
<td>$\rho$</td>
<td>kg/m$^3$</td>
<td>Density of the Dowtherm A</td>
</tr>
<tr>
<td>$Pr$</td>
<td></td>
<td>Prandtl number</td>
</tr>
<tr>
<td>$\lambda_s$</td>
<td>W/(m·K)</td>
<td>Thermal conductivity of the steel</td>
</tr>
<tr>
<td>$Q$</td>
<td>m$^3$/s</td>
<td>Volumetric flow</td>
</tr>
</tbody>
</table>
### Symbol Table

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<tr>
<th>Symbol</th>
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<th>Significance</th>
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<tbody>
<tr>
<td>$u$</td>
<td>m/s</td>
<td>Velocity</td>
</tr>
<tr>
<td>$D_i$</td>
<td>m</td>
<td>Inner diameter</td>
</tr>
<tr>
<td>thick</td>
<td>m</td>
<td>Thickness of the tube</td>
</tr>
<tr>
<td>$D_0$</td>
<td>m</td>
<td>Outer diameter</td>
</tr>
<tr>
<td>$P_t$</td>
<td>m</td>
<td>Thermal perimeter</td>
</tr>
<tr>
<td>$P_h$</td>
<td>m</td>
<td>Hydraulic perimeter</td>
</tr>
<tr>
<td>$D_h$</td>
<td>m</td>
<td>Hydraulic diameter</td>
</tr>
<tr>
<td>$S$</td>
<td>m$^2$</td>
<td>Inner cross-section of one tube</td>
</tr>
<tr>
<td>$A_{\text{firstapprox}}$</td>
<td>m$^2$</td>
<td>Total cross-section of the tubes</td>
</tr>
<tr>
<td>MLDT</td>
<td>K</td>
<td>Logarithmic mean temperature difference</td>
</tr>
<tr>
<td>$G$</td>
<td>kg/(m$^2$·s)</td>
<td>Superficial mass flow</td>
</tr>
<tr>
<td>$Re$</td>
<td>-</td>
<td>Reynolds number</td>
</tr>
<tr>
<td>$q$</td>
<td>W/(m$^2$)</td>
<td>Heat flux</td>
</tr>
<tr>
<td>$C_{fm}$</td>
<td>-</td>
<td>Friction coefficient</td>
</tr>
<tr>
<td>$Nu$</td>
<td>-</td>
<td>Nusselt number</td>
</tr>
<tr>
<td>$hd$</td>
<td>W/(m$^2$·K)</td>
<td>Convection coefficient of the Dowtherm A</td>
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<tr>
<td>$A_s$</td>
<td>m$^2$</td>
<td>Inner lateral area</td>
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<tr>
<td>$R_s$</td>
<td>K/W</td>
<td>Thermal resistance from the Dowtherm A to the tube</td>
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<tr>
<td>$R_{ss}$</td>
<td>K/W</td>
<td>Thermal resistance between the two walls of the tube</td>
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<tr>
<td>$U_s$</td>
<td>W/(m$^2$·K)</td>
<td>Global heat transfer coefficient</td>
</tr>
<tr>
<td>$L_i$</td>
<td>m</td>
<td>Length</td>
</tr>
<tr>
<td>$A_i$</td>
<td>m$^2$</td>
<td>Area</td>
</tr>
<tr>
<td>$V_i$</td>
<td>m$^3$</td>
<td>Volume</td>
</tr>
<tr>
<td>$D_{\text{out}}$</td>
<td>m</td>
<td>Outer diameter with the PCM</td>
</tr>
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### ANNEX B

<table>
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<tr>
<th>Symbol</th>
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<tr>
<td>$eK$</td>
<td>K</td>
<td>Scale change from centigrads to Kelvin</td>
</tr>
<tr>
<td>$h$</td>
<td>W/(m$^2$·K)</td>
<td>Convection coefficient of the Dowtherm A</td>
</tr>
<tr>
<td>$T_s$</td>
<td>K</td>
<td>Temperature of the steel</td>
</tr>
<tr>
<td>$\lambda_s$</td>
<td>W/(m·K)</td>
<td>Thermal conductivity of the steel</td>
</tr>
<tr>
<td>$\rho_s$</td>
<td>kg/m$^3$</td>
<td>Density of the steel</td>
</tr>
<tr>
<td>$c_s$</td>
<td>J/(kg·K)</td>
<td>Specific heat of the steel</td>
</tr>
<tr>
<td>$\lambda_a$</td>
<td>W/(m·K)</td>
<td>Thermal conductivity of the aluminum</td>
</tr>
<tr>
<td>$c_a$</td>
<td>J/(kg·K)</td>
<td>Specific heat of the aluminum</td>
</tr>
<tr>
<td>$\rho_a$</td>
<td>kg/m$^3$</td>
<td>Density of the aluminum</td>
</tr>
<tr>
<td>Symbol</td>
<td>Unit</td>
<td>Significance</td>
</tr>
<tr>
<td>----------</td>
<td>------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>$T_{13}$</td>
<td>K</td>
<td>Temperature of the contour in the tube (extremes)</td>
</tr>
<tr>
<td>$D_i$</td>
<td>m</td>
<td>Inner diameter</td>
</tr>
<tr>
<td>thick</td>
<td>m</td>
<td>Thickness of the tube</td>
</tr>
<tr>
<td>$D_0$</td>
<td>m</td>
<td>Outer diameter</td>
</tr>
<tr>
<td>$D_{out}$</td>
<td>m</td>
<td>Outer diameter with the PCM</td>
</tr>
<tr>
<td>$\Delta \varphi$</td>
<td>rad</td>
<td>Parameter of the volume element (see figure 49)</td>
</tr>
<tr>
<td>finthick</td>
<td>m</td>
<td>Total height of the volume element</td>
</tr>
<tr>
<td>$n$</td>
<td>m</td>
<td>Parameter to modify the thickness of the fin</td>
</tr>
<tr>
<td>$\Delta z_1$</td>
<td>m</td>
<td>Height of the aluminum fin</td>
</tr>
<tr>
<td>$\Delta r_1$</td>
<td>m</td>
<td>Distance from the center of node 1 to the outer diameter ($D_0$)</td>
</tr>
<tr>
<td>$V_i$</td>
<td>m$^3$</td>
<td>Volume of node i</td>
</tr>
<tr>
<td>$C_i$</td>
<td>J</td>
<td>Total heat capacity of node i</td>
</tr>
<tr>
<td>$K_{ij}$</td>
<td>W/K</td>
<td>Conductivity between nodes i and j</td>
</tr>
<tr>
<td>$\Delta r_2$</td>
<td>m</td>
<td>Division of the aluminum fin in the radial direction (distance from the center of the node to its extreme)</td>
</tr>
<tr>
<td>$R_{m7}$</td>
<td>K/W</td>
<td>Thermal resistance between the steel and the aluminum in the steel part</td>
</tr>
<tr>
<td>$R_{m8}$</td>
<td>K/W</td>
<td>Thermal resistance between the steel and the aluminum in the aluminum part</td>
</tr>
<tr>
<td>$\Delta t$</td>
<td>s</td>
<td>Time interval for each step of the implicit method</td>
</tr>
</tbody>
</table>
1. Introduction

The increasing on the prices of the conventional energy sources and the environmental awareness have led to increase the use of renewable energies and the energy efficiency. In this scene, storing thermal energy plays a really important role as it allows to improve the dispatchability and the efficiency of different applications such as thermosolar power plants, greenhouses and buildings heating systems.

There are three methods used and still being investigated in order to store thermal energy. One is the sensible heat storage (SHS), the other one is the latent heat storage (LHS) and the last one is the thermo-chemical storage. In the present work the last two ones will be presented in the following sections focusing on the high temperature storage between 200 and 700 °C.

The first method (SHS) is based on raising the temperature of a solid or liquid to store heat and releasing it with the decrease of temperature when it is necessary. The volumes needed to store energy in the scale that world needs are extremely large. That is why the other two methods are being developed. The LHS method is a medium term method and the thermo-chemical storage is a long term method as there have not been so many research work and experiments as with the other two methods.

1.1. LATENT HEAT STORAGE

As it has been said in the introduction, an option to store thermal energy is the latent heat storage. This method is based on the utilization of phase change materials (PCM). These materials store heat when they go from solid to liquid, from liquid to gas or from solid to solid (change of one crystalline form into another without a physical phase change). Then they release energy when they have the reverse phase change. It must be mentioned that until now, the PCM studies and applications have been mainly focused on the solid-liquid phase change.

The storage capacity of a LHS system in the concrete case of solid-liquid transformation is given by equation 1,

\[ Q = \int_{T_i}^{T_m} m \cdot C_p \cdot dT + m \cdot a_m \cdot \Delta h_m + \int_{T_m}^{T_f} m \cdot C_p \cdot dT \]

Equation 1. Storage capacity of a LHS system

where \( T_i \) is the initial temperature, \( T_m \) is the melting temperature, \( m \) is the mass of heat storage medium, \( C_p \) is the specific heat, \( a_m \) is the fraction melted and \( \Delta h_m \) is the heat of fusion per unit mass (\( J/kg \)).

The heat of fusion or the heat of evaporation is much greater than the specific heat. Thanks to that, latent storage materials has got a larger volumetric energy storage capacity than sensible storage materials. Another advantage is the fact that the absorption and release of the energy stored takes place at a constant temperature which makes easier the choice of the material to use in the different applications.
1.1.1. Materials to be used as PCM
1.1.1.1. Properties required

It is possible to find materials with a heat of fusion and melting temperature inside the desired range but a material has to exhibit certain properties to become a feasible PCM. These properties can be classified in 5 groups: thermal properties, physical properties, kinetic properties, chemical properties and economical properties [1] [5].

The **thermal properties** the material must have:

- Suitable phase-transition temperature for the specific application.
- High latent heat of transition in order to occupy the minimum possible volume.
- High thermal conductivity in order to provide the minimum temperature gradients and facilitate the charges and discharges of heat.

The **physical properties** are:

- Favorable phase equilibrium to facilitate the heat storage.
- High density to also occupy the minimum possible volume.
- Small volume change to facilitate the construction of the different needed containers and heat exchangers.
- Low vapor pressure in order to avoid stresses and problems with the containers and heat exchangers needed.

The **kinetic properties** are:

- Sufficient crystallization rate in order to avoid supercooling.
- No supercooling as it makes difficult to control the heat transfer and the truth melting temperature that is in principle given.

The **chemical properties** are

- Long-term chemical stability and complete reversible melt/freeze cycle as it is needed to work during the most cycles possible.
- Compatibility with materials of construction as it is also needed to work the most possible time.
- No toxicity because of safety reasons.
- No fire hazard also because of safety reasons.
- Non-explosive also because of safety reasons.

Finally, from the point of view of **economics**, the material must be abundant, available and cost effective to help into the feasibility of the use of the storage system.

1.1.1.2. Classification of the PCM

The different materials which have the properties mentioned before are classified in different groups. One possible classification is the one shown in figure 1. There are three main groups: organic materials, inorganic materials and eutectics. It must be mentioned that generally the materials do not respect all the properties listed in the previous section and it
has to be compensated with the system design and different enhancement methods such the use of fins or composite materials in the form of matrixes.

![Phase Change Material Diagram](image)

**Figure 1. Classification of PCM [1]**

Inside the organic materials group there are paraffin compounds and non-paraffin compounds. The paraffin compounds are formed by saturated hydrocarbons which correspond to the general formula \(C_nH_{2n+2}\). Paraffin wax is the most PCM material used in this group and it consists in a combination of different hydrocarbons obtained from the petroleum distillation. The properties of the different paraffin compounds are quite similar and that makes easier its study.

The non-paraffin group contains different esters, fatty acids, alcohols and glycols. Each of the materials have different properties and they can be found in the literature in case it is necessary to use one of them.

Inside the inorganic group there are also two main groups: salt hydrate and metallics. Salt hydrates are alloys of inorganic salts and water forming a typical crystalline solid of general formula \(AB\cdot nH_2O\). The metallics group include low melting metals and metal eutectics but it must be mentioned that they have not been really considered as PCM because of their large weight. In spite of this, there are some interesting points to remark: they have got high thermal conductivity, relatively low vapor pressure and high heat of fusion per unit volume.

There are some thermophysical properties from different metal alloys that have been studied as possible PCM in tables 17 to 22 from reference [3].

Finally there is the eutectics group. An eutectic is a minimum-melting composition of two or more components, each of which melts and freezes congruently forming a mixture of the component crystals during crystallization [4]. This group contains also three different
groups: organic-organic, inorganic-inorganic and inorganic-organic, depending on the nature of the components of the composition.

More detailed information about the different groups, its advantages and disadvantages and some tables with materials of each group, melting temperature and heat of fusion can be found in [1].

As the present work is focused on high temperature storage systems between 200 and 700 ºC, as it is shown in figure 2 it is not possible to use all the materials groups as PCM. Only inorganic salts, their eutectic mixtures and if it is possible metallics are going to be considered in the next sections.

Figure 2. PCM groups with their heat of fusion and melting temperature range [2]

Another possible classification similar to the one mentioned before and with some examples for each group is shown in figure 3. There can be observed the groups of most interest for the high temperatures applications as well.
1.1.1.2.1. Inorganic salts and their eutectic mixtures

As given in reference [3] the inorganic salts have a melting temperature that goes from 250 to 1680 °C and a heat of fusion from 68 to 1041 J/g. The most perspective compositions are double and ternary eutectics, especially the ones on the basis of fluorides and chlorides. Some of them are presented in table 2 and table 3 from reference [3]. Besides the fluorides and chlorides, compositions on the basis hydroxides, nitrates, carbonates and other salts are of considerable interest and some of them can also be observed in tables 4 and 5 from the reference [3].

Although it has been mentioned that the most perspective compositions are eutectics on the basis of fluorides and chlorides from reference [27] and from the applications which will be presented in the next chapters the alkali metal nitrates and nitrites, potassium and sodium nitrate and binary mixtures of the two are very suitable, covering the temperature range of 140 °C - 330 °C. In the next figure some of these mixtures with their respective melting temperature and heat of fusion are shown.
It must be mentioned that there are also studies in quaternary and quinary compositions on a basis molybdates, vanadates and sulphates which are also promising. In reference [3] some other tables with salts compositions and different thermophysical properties are given: melting point, heat of fusion and in some cases the thermal capacity, the thermal conductivity and the thermal diffusivity.

Besides the thermophysical properties which must be measured on the compositions of interest, as mentioned in section 1.1.1.1. there are other material properties required. Some aspects of these properties in the specific case of inorganic salts and eutectic mixtures must be considered.

In practice, when a substance solidifies it is possible that it gets to a lower temperature than the melting temperature caused by not having enough crystallization rate and nucleating properties that causes at the same time a lack of crystal nucleus to solidify. That is known as supercooling and there are some possible solutions as mentioned in reference [1]. It is possible to add a nucleating agent facilitating the formation of the crystals nucleus needed to solidify. Another possibility is to retain some crystals, in a small cold region, to serve as nucleus.

As mentioned in reference [3] one of the most challenging features from inorganic salts from the point of view of the design is the change of volume at phase transition. A difficulty added to this area is the lack of information on the temperature dependences of density. Some studies mentioned in reference [3] show some results and there is a change of volume that can exceed the 10% in some cases. There is the need to study this property accurately when it is wanted to use any salt or eutectic composition.

Another important feature is the long-term characteristics of the material as longest possible life-time is desired. Also the life-time of the material depends on the application as told in reference [3]. In space applications the life-time can be about months and years but in power plants 20 to 30 years of life-time must be expected. The studies made in some particular materials show that the melting temperature and the heat of fusion do not change appreciably, but the chemical stability and the amount of thermal cycling which is possible to perform must always be tested.
The last property that must be tested is the compatibility between the salt and the constructional materials. No chemical interaction must happen as the life-time of the design must be guaranteed also in this aspect and corrosions tests must be run. As mentioned in reference [3] the constructional material should have comprehensible mechanical and metallurgical properties. The alloy for the container should be easily processed, be accessible and cheap.

Finally the last feature of salts that must be taken into consideration is their low heat conductivity (nearby 1 W/m) as told in [3] and [7]. This is the biggest technological problem in the use of PCM faced right now.

The two concepts to solve this situation are mainly on one hand the extension of the heat transfer area and on the other hand the use of a composite material compounded by the salt and another material resulting in a higher heat conductivity. Normally the material used to form the composite is graphite. The extension of the heat transfer area between the heat carrier and the PCM can be achieved by the use of fins or by the encapsulation of the PCM. Some studies about the conductivity enhancement will be discussed and presented in section 1.1.4.

1.1.1.2.2. Metallics

Although there have not been so many research as in the case of the other groups, metals and its alloys must been taken into account as they can be used as high temperature storage materials and they, as a rule, are deprived those lacks which are characteristic for salts as mentioned in [3]. These lacks are: low heat conductivity, corrosion activity and high cost of some salts.

Some metal eutectic alloys have been studied as possible phase change materials determining their melting temperature and heat of fusion. These results are shown in tables 17 to 19 from reference [3].

The study of two compositions on the basis of aluminum and silicon performed by Wang et al. [6] must be remarked. One was AlSi$_{12}$ with a melting temperature of 576 ºC and a heat of fusion of 560 J/g and the other one was AlSi$_{20}$ with a $T_m$ of 585 ºC and 460 J/g. As the heat of fusions is greater in the case of AlSi$_{12}$ this material was chosen for further studies. The thermophysical properties of this alloy are shown in table 1. This alloy has been used for the development of a high-temperature isothermal electric heater intended for thermal energy storage at night, when the tariff for the electric power is lower.

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat capacity of solid</td>
<td>1,038</td>
<td>J/g·K</td>
</tr>
<tr>
<td>Heat capacity of liquid</td>
<td>1,741</td>
<td>J/g·K</td>
</tr>
<tr>
<td>Temperature of phase transition</td>
<td>576</td>
<td>ºC</td>
</tr>
<tr>
<td>Heat of fusion</td>
<td>560</td>
<td>J/g</td>
</tr>
<tr>
<td>Density</td>
<td>2,70</td>
<td>g/cm$^3$</td>
</tr>
<tr>
<td>Thermal conductivity</td>
<td>160</td>
<td>W/m·K</td>
</tr>
</tbody>
</table>

Table 1. Thermophysical properties of AlSi$_{12}$ alloy. [6]
1.1.2. Latent heat storage systems

In the temperature range of interest (200-700ºC) the concepts used for the power solar plants can be generalized to use them for the rest of applications. The concepts here presented can also be applied to sensible heat storage systems. As told in [8] there are two main concepts: the active and the passive systems.

In the active systems the storage medium itself circulates through a heat exchanger [8]. They are subdivided into direct and indirect systems. In the first case the heat transfer fluid (HTF) is also the storing medium and in the second case there is a second medium used for storing the heat.

In the case of passive storage systems the HTF is used for charging and discharging the storage material which is a different material to HTF.

1.1.2.1. Active direct systems

As it has been said in these systems the heat transfer fluid and the storage medium are the same. That means a reduction on the costs of the system as there is no need of expensive heat exchangers, the working temperatures can be higher improving the performance of the plant and reducing the levelised electrical cost [8].

The usually system in this group is the two tanks direct system, where there are two tanks. One tank is used to store the hot HTF and the other one to store the HTF when it has been discharged.

The advantages of the system is that cold and heat storage materials are separated, reducing the risk of approach and the high working temperatures that can be performed.

The disadvantages are the high cost of the HTF; high cost of the heat exchanger to produce steam for the Rankine cycle; the need of two tanks; relatively small temperature difference between the hot and the cold fluid in the storage system; high risk of solidification of the storage fluid due to its high freezing temperature; increase of losses in the solar field due to the high working temperature and finally the lowest cost of the thermal energy storage (TES) system design and operation does not correspond to the lowest cost of electricity [8].

1.1.2.2. Active indirect systems

There are two main concepts in this group based on working with different HTF and storage medium. One is the two tanks indirect systems which works as the direct one but as the HTF fluid and the storage material are different there is the need of a heat exchanger between them.

The advantages of this first concept are similar to the two tanks direct system: separation between cold and hot HTF and the storage material flows only between hot and cold tanks, not through the parabolic troughs. The disadvantages are the same as in the two tanks direct systems [8].
The other concept is the single tank system reducing the cost as there is only one tank. The cold and hot fluids are in the same tank and separated because of stratification. This zone is known as thermocline. The hot fluid is at the top and the cold fluid at the bottom and there is also the need of a heat exchanger to transfer the energy between the HTF and the storage material.

The advantages of this concept are the decreasing of the cost in front of the two tank storage system. As for the disadvantages: it is more difficult to separate the hot and cold HTF; in the case of using molten salts it is necessary to keep a minimum system temperature to avoid freezing and salt dissociation as they have high melting temperatures; the outlet temperature is high resulting in an increase of losses in the solar field; the thermocline needs to be controlled with some method (complicated); the design of storage system is complex and from the thermodynamically point of view it is an inefficient power plant [9].

1.1.2.3. Passive systems

In a passive storage system the heat transfer medium passes through the storage only for charging or discharging the system. The storage medium itself does not circulate. Passive systems are generally dual medium storage systems (these systems are also called regenerators). Passive storage systems are mainly solid storage systems (concrete, castable materials and PCM) [8]

In the case of using concrete as a storage material the advantages are the low cost of it; its high heat transfer rate due to a good contact between concrete and piping; the facility of handling the material and the low degradation of heat transfer between the heat exchanger and the storage material. The disadvantages are increase of cost of heat exchanger and of engineering and a long-term instability [8].

In the case of using PCM the advantage is the improvement of storage ratio thanks to the higher heat of fusion than specific heat of materials as it has already been said.

1.1.3. Examples
1.1.3.1. Active direct systems
1.1.3.1.1. DISTOR Project

Solar thermal power plants using parabolic trough collectors are the most economic systems to generate electricity from solar insulation in the MW range. DISS project demonstrated the feasibility of high temperature direct steam generation (DSG) in absorber pipes of parabolic trough which resulted in a cost reduction of the thermal fluid as it is water and also a higher working temperature which improves the efficiency of the plant. INDITEP project, based on the know-how achieved in the DISS project, performed the design of the first pre-commercial DSG solar power plant (5 MWe).

The storage systems play an important role in the improvement and acceptance of these power generation systems as there is a reduction in the dependence on solar insulation, there is a larger efficiency on the electricity production and it facilitates the integration of a solar power plant into an electrical grid, since fluctuations in electricity generation resulting from the actual insulation are reduced. These advantages result in higher revenues [20].
Finally, DSG solar power plants require isothermal energy storage systems and that is the cause of the DISTOR project birth.

The DISTOR project (began in 2004) had the budget of developing thermal storage systems using phase change materials (PCM) in the temperature range from 230°C to 330°C for systems using steam between 30-100bar [19]. The economical goal was to achieve a 20 €/kWh storage system capacity. The project developed different LHS concepts in parallel in order to reach to a solution as fast as possible. These concepts can be classified in three main groups as said in [22] (some of them have already been mentioned in section 1.1.1.2.1.):

- **Extended heat transfer area**: The contact area between the PCM and the containment for the steam is enlarged to reduce the distance for heat transport within the PCM. Possible implementations of this approach use either finned tubes or capsules. The walls of the capsules can be either flexible or stiff to withstand a pressure difference.

- **Composite material with increased thermal conductivity**: A material showing a high thermal conductivity is added to the PCM. The PCM can be infiltrated in a porous matrix made up of the additional material or the two components can be mixed as powders.

- **Intermediate heat transfer medium liquid/gaseous**: Here, the PCM and a heat exchanger are arranged in a container filled with a medium that transfers the energy between these two components. The heat transport involves a phase change of the heat transfer medium. The pressure of the heat transfer medium is significantly below the pressure of the steam in the heat exchanger; the costs for the outer containment can be reduced significantly compared to a system with a direct contact heat exchanger between steam and PCM.

The melting temperature of the PCM must be between the saturation temperature of the steam from the solar field and the saturation temperature in the steam cycle. The integration of the thermal storage in the DSG power plant is shown in figure 5. The material chosen as PCM is an eutectic mixture of KNO$_3$-NaNO$_3$ with a melting temperature of 230°C as due mainly to cost aspects the nitrate salts in eutectic mixtures are the most commonly materials used in LHS systems.

![Figure 5. Basic concept for integration of thermal energy storage into solar thermal parabolic trough power plants using DSG technology [22].](image)
The extended heat transfer area concept includes two different approaches. On the one hand the sandwich concept based on the integration of materials with high thermal conductivity. Thin plates orthogonal to the axis of the steam pipes are used. The first materials contemplated as high thermal conductivity ones were graphite, stainless steel and carbon steel. Due to the costs and good corrosion resistivity graphite foils are chosen instead of steel. The final structure can be observed in figure 6. More details about distances, positions and assembly can be found in [22].

On the other hand the DISTOR project has deepened in the macro-encapsulation concept. The PCM is packed into capsules to reduce the maximum distance for heat transport. Then the capsules are introduced in a vessel where the steam/water circulates. As the PCM undergoes a change of volume of about 10% in the phase change there are two possible options: using a flexible capsule and filling it completely with PCM or using a stiff capsule filling it with PCM and gas.

The flexible capsules for high temperature LHS systems is not a feasible option as the cost of the material needed for the capsules is too high.

For the stiff capsules a cylindrical shape was chosen and the pressure inside the capsule depending on the amount of PCM was studied. There is the need of less than 80% PCM volume in order to avoid a serious cyclic load. The capsules can be observed in figure 7.

In the case of the second group, composite material with increased thermal conductivity, the approach performed is the assembly by compression of a powder mixture of eutectic NaNO$_3$ /KNO$_3$ and expanded graphite. The result can be observed in figure 8. The
content of graphite must be limited as far as it is possible in order to achieve a cost-effectiveness composite material.

Finally, there is not information available of the third group as it needed more research and trials than the other two groups. More detailed information about the different approaches can be found in [22].

The sandwich concept was chosen as the best option and in 2007 a test module of 200 kWh thermal capacity storage system was installed in the *Plataforma Solar de Almería (PSA)* where the DISS Project took place. The storage model and the acquisition devices in order to study its behavior can be observed in figure 9.

![General view of the storage module and the system installed in the PSA DISS plant for its evaluation](image)

*Figure 9. General view of the storage module and the system installed in the PSA DISS plant for its evaluation [23].*

More information about the installation can be found in [23] but it is interesting to remark the design parameters of the test module:

- pressure /nominal temperature: 25 bar/220ºC
- nominal water/steam flow rate: 0.083 kg/s

and the final conclusions of the annual report 2007 [23] concerning the DISTOR project: the technical feasibility of the molten salt thermal storage concept and later heat recovery by crystallization was demonstrated during testing. It was also demonstrated how important tube bundle design is to ensure heat transfer during charging and discharging. The lower velocity of the working fluid (water) during discharging requires the tube bundles to be sized for the right convective heat transfer coefficient without excessive pressure drops in the tube bundle during charging, when the working fluid (steam) density is lower.
Although the project was programmed to end in 2007, due to a delay in the construction the project ended in 2008. It must be mentioned that tests were stopped because there was a leak in the tube bundle and the cause of the leak was detected. It gave valuable information for future prototypes. DLR and CIEMAT which were some of the organisms working in the DISTOR project decided to perform a new LHS test module in PSA out of the DISTOR Project. This new test module is designed to work for a maximum of 80 bar and 300 °C and the salt used is still a nitrate salt. The tests should have been done during 2010 but there is not available information about it [24].

Within the framework of DISTOR project there are some articles which must be mentioned. The first article of interest is [25]. A model of a PCM composite block (figure 10) for heat storage is developed and some simulations and analysis are performed. The use of blocks allows to adjust the heat storage capacity to the working conditions of the DSG solar power plant.

![Figure 10. Composite block of composite EG-PCM studied in [25].](image)

The PCM composite consists of compressed naturally expanded porous graphite matrix (CEG). The manufacture was performed by pouring the expanded graphite powder into a cubic mold of aluminium and then pressed to obtain the porous graphite matrix with the desired bulk apparent density. Then the CEG graphite pores were filled with the binary eutectic KNO₃/NaNO₃ mentioned before. Three different composites were obtained varying the percentage by mass of CEG (3’96, 5’27 and 7’35%) and the thermophysical properties of the resulting composite materials were studied and are shown in table 2. As it can be observed the thermal conductivity in the axial direction is lower than the one in the radial direction due to the axial compression which is used in the manufacturating method.
<table>
<thead>
<tr>
<th>Material</th>
<th>Pure salt</th>
<th>CEG–salt composite (3.96% mass graphite)</th>
<th>CEG–salt composite (5.27% mass graphite)</th>
<th>CEG–salt composite (7.35% mass graphite)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axial thermal conductivity (W/m K)</td>
<td>0.8</td>
<td>1.6</td>
<td>2.6</td>
<td>2.8</td>
</tr>
<tr>
<td>Radial thermal conductivity (W/m K)</td>
<td>0.8</td>
<td>4.1</td>
<td>5.65</td>
<td>8.2</td>
</tr>
<tr>
<td>Apparent latent heat of fusion (J/kg)</td>
<td>94250</td>
<td>90500</td>
<td>89300</td>
<td>87300</td>
</tr>
<tr>
<td>Apparent density of CEG (kg/m3)</td>
<td>-</td>
<td>75</td>
<td>100</td>
<td>140</td>
</tr>
</tbody>
</table>

*Table 2. Measured thermophysical properties of the different composites [25]*

The detailed heat transfer and fluid flow model is detailed in section 3 from [25] but due to its complexity as there is the need to know or fix a great amount of parameters a simplified model is developed in section 4 in order to have an easy and feasible preliminary design. The simplified model is validated through the comparison with the detailed model.

A design procedure for sizing the storage system is presented. The maximum energy storage capacity and average power of system are the most important design criteria and these two parameters are function of the length of the tubes, the number of tubes and the time of charging the system as can be observed in equation 11 from [25]. There is a design procedure explained which is based on fixing the internal tube diameter and solving some of the equations given in [25].

Finally this design procedure is used to design a thermal storage system of 20 kWh with an average power of 10 kW in order to test the different composites. The detailed information of the design is shown in table 2 from [25]. It is interesting to remark the conclusions which are reached. With less than 4% graphite by mass there is a reduction of the overall length of tubes of 74% with an increase in the PCM composite material weight less than 4.5% and a small decrease in the overall heat storage capacity. This is an important point to consider as the steel tubes represent a large part in the total cost of the storage system.

The second article within the framework of the DISTOR project which is interesting to summarize deals with an analysis of a 100 kWth latent heat storage system for DSG using the sandwich concept presented above. The prototype, which is shown in figure 11, was installed in PSA and it proved the feasibility of this latent heat storage system.
The PCM used was an eutectic mixture of 54% wt KNO$_3$/46% NaNO$_3$ and the conductive fins were 490x490 mm$^2$ and 1 mm thick of expanded graphite. More details about the prototype dimensions and configuration are shown in table 2 from [26]. It is interesting to mention the working conditions: a mass flow of 0.08 kg/s, saturated steam at about 235 °C in charge mode and saturated water at about 200 °C in discharge mode.

Different thermocouples were placed in 6 different transversal sections in order to study the behavior of the PCM. The working temperature ranges of the PCM were divided in 4 regions which are specified in [26] both for charge and discharge processes. There were presented also some calculations which are compared to the experimental results in section 4.1 from [26].

There was a mismatch between the experimental and calculation results in some of the regions both for charge and discharge processes. One cause is the different steam production in the central pipes than in the outer pipes. The other cause of the mismatch is an excess of PCM which had no EG foils and then could not transfer heat in an efficient way. This must be corrected in other designs taking care of the TES module volume and tube bundle/PCM arrangement.

There is another disagreement between the theoretical calculations and the experimental results between the energy which can be charged if there is enough time (58 kW$_{th}$h) and the energy which can be recovered (40 kW$_{th}$h). One of the causes is the excess of mass already commented which does not give any energy in discharge process.

The last simulation performed is the application of the quasi static approximation which is presented in detail in [26]. This approximation is based on neglecting the sensible heat exchanged and then the energy equation becomes independent of time. The results of the
calculations of the steam quality shown in [26] lead to the following conclusion: at the beginning of the heat transfer region considered the real HTF condensation is lower than expected from calculations. Some reasons as told in [26] could be the following:

- Not all tubes condense the same amount of vapor.
- Condensation does not occur along the whole tube length.
- The quasi static approximation cannot be applied at the beginning of charging. In other words, it is not possible to neglect the sensible heat exchange at the beginning of charging.

During discharge mode the calculation curve of the steam quality fits the experimental results if it is considered a minor tube length than the real one taking into account that the vaporization does not take place in all the tube length due to a deficient thermal insulation.

Finally it is interesting to include the final suggestions of [26] in order to consider them in future designs:

- PCM mass should not exceed the amount that can be homogeneously melted/frozen by the tube bundle.
- Thermal insulation to environment is crucial and has to be ensured around the whole prototype.
- For a detailed description of TES module with sandwich configuration, it is necessary to improve the quasi static model by introducing a sensible heat term at both beginning and end of latent heat exchange regions.

1.1.3.2. Active indirect systems
1.1.3.2.1. Cascaded latent heat storage systems (CLHS) or multiple PCMs method

In this case, different PCMs are used in order to store the heat as shown in figure 12. This way allows to have a more uniform outlet temperature of the HTF and a higher used percentage of the possible phase change [10][13]. Also almost a constant heat flux is achieved along the device thanks to maintain a nearly constant temperature difference [28]. As it can be observed in figure 12, the melting temperatures of the different PCMs used must be cascaded from the minimum to the maximum operating temperature.

![Figure 12. Proposal of cascaded latent heat storage with 5 PCM according to [11], [10]](image-url)
The study performed in [10] is summarized in this section. It is a cascaded latent heat storage system oriented to be used in a parabolic trough solar power plant with a HTF which works between 290 and 350 °C. The materials selected were: sodium nitrate (NaNO₃), an eutectic mixture of potassium nitrate and potassium chloride (KNO₃/KCl) and potassium nitrate (KNO₃). The selection of these materials was based on experimental and literature data. The properties of these three materials, as given in [10], are shown in Table 3.

<table>
<thead>
<tr>
<th>Property</th>
<th>Unit</th>
<th>NaNO₃</th>
<th>KNO₃/KCl 95.5/4.5</th>
<th>KNO₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>T&lt;sub&gt;m&lt;/sub&gt;</td>
<td>°C</td>
<td>306</td>
<td>320</td>
<td>335</td>
</tr>
<tr>
<td>Δh&lt;sub&gt;f&lt;/sub&gt;</td>
<td>kJ/kg</td>
<td>172</td>
<td>74</td>
<td>95</td>
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<tr>
<td>ρ&lt;sub&gt;s&lt;/sub&gt;</td>
<td>kg/m&lt;sup&gt;3&lt;/sup&gt;</td>
<td>2261</td>
<td>2100</td>
<td>2109</td>
</tr>
<tr>
<td>C&lt;sub&gt;p,s&lt;/sub&gt;</td>
<td>kJ/kg·K</td>
<td>1,10</td>
<td>1,21</td>
<td>0,953</td>
</tr>
<tr>
<td>k&lt;sub&gt;s&lt;/sub&gt;</td>
<td>W/m·K</td>
<td>0,5</td>
<td>0,5</td>
<td>0,5</td>
</tr>
</tbody>
</table>

Table 3. Properties of the materials selected in [10].

The experimental device is shown in Figure 13. It uses a synthetic heat transfer oil (similar to the solar thermal electricity generating systems (SEGS) plants), an electrical heater for the charging period and a cooling tower for the discharging period. The test modules are vertical shell and tube heat exchangers with the heat transfer oil flowing through the inner heat transfer pipes (simple high temperature steel –AISI K01200, outer diameter 0.012 m, wall thickness 0.002 m) and the PCM being enclosed in the annular space between the heat transfer pipes and the shell (simple steel –AISI 1015, inner diameter 0.13 m).

Figure 13. Simplified flow diagram of the thermal oil test facility. Photo on the right side shows test module “B”, before the test modules got insulated. For stressless thermal expansion each module is counter balanced [10].
Due to the PCMs volume change during melting and solidifying the hot oil enters the test module at the top during charging and the somewhat colder oil enters the test module at the bottom during discharging. To determine the heat transfer coefficient during melting a simple double tube test module with a height of 2.5 m was used (Figure 5A). To investigate the behavior of a CLHS three identical test modules with a height of 1.1 m each where placed directly above each other to avoid heat losses of the oil when flowing from one test module to the other (Figure 5B).

Temperature sensors were placed inside the heat transfer pipes to measure the oil’s temperature at the inlet, during the way through the test module and at the outlet. Further temperature sensors were placed in the annular gap around the heat transfer pipes to measure the PCMs temperature.

Consecutive charging-discharging cycles were performed and the temperature distribution of the different PCMs is shown in figure 14. It must be observed that the charging period lasted for 4 hours but the discharging period for 2 hours due to heat losses to the ambient.

![Figure 14. Temperature distribution of a charging period with the scale time on the right side (h:mm).](image)

In this study also LHS were studied in order to compare the CLHS and to study if it is really advantageous. The materials used were NaNO\textsubscript{3} and KNO\textsubscript{3}. In the case of KNO\textsubscript{3}, two studies were performed. In one the temperatures of the oil went from 270 to 330 ºC in order to have a sensible storage system as a reference as this range is below its melting temperature. For the other two cases the temperature range was maintained and in all the cases the temperature difference was the same. The results obtained are shown in table 4.

<table>
<thead>
<tr>
<th>Property</th>
<th>Unit</th>
<th>CLHS</th>
<th>All NaNO\textsubscript{3}</th>
<th>All KNO\textsubscript{3}</th>
<th>All KNO\textsubscript{3} (Sensible)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Q_{\text{theor}}$</td>
<td>MJ</td>
<td>16,5</td>
<td>22,7</td>
<td>14,2</td>
<td>7,0</td>
</tr>
<tr>
<td>$Q_{\text{exp}}$</td>
<td>MJ</td>
<td>9,4</td>
<td>10,2</td>
<td>7,6</td>
<td>5,0</td>
</tr>
<tr>
<td>$Q_{\text{exp}}/Q_{\text{theor}}$</td>
<td>%</td>
<td>57,2</td>
<td>44,9</td>
<td>53,4</td>
<td>70,9</td>
</tr>
</tbody>
</table>

Table 4. Heat storage capacities of four different storage configurations [10].
\( Q_{\text{theor}} \) indicates the theoretical storage capacity of each configuration which is calculated using equation 2. \( Q_{\text{exp}} \) shows the experimental results and the third line is the ratio between these last two values. The interesting points to comment are on the one hand the higher storage capacity of the LHS systems in front the sensible storage system, the higher storage capacity of the All NaNO\(_3\) sample due to its high heat of fusion and the higher ratio \( Q_{\text{exp}}/Q_{\text{theor}} \) of the CLHS in front the other LHS systems which was one of the advantages of this type of system commented at the beginning of the section.

\[
Q_{\text{theor}} = \sum_{i=1}^{n} m_{PCM_i} \cdot \left( \int_{T_{\text{min}}}^{T_{\text{m}}} c_{p,s} \cdot dT + \int_{T_{\text{m}}}^{T_{\text{max}}} c_{p,i} \cdot dT \right) + \Delta h_m
\]

*Equation 2. Theoretical storage capacity of a latent heat storage system.*

A simulation with Dymola-Modelica was also accomplished in order to compare the experimental data. More details about the model can be found in [10] but it must be remarked that all simulation results matched the experimental data well.

The model verified was used to design a CLHS for a 50 MW\(_e\) parabolic trough power plant with a storage capacity of 875 MWh\(_{\text{th}}\) to realize a discharging duration of 6 h (split into two storages with 3 h each for more flexibility) while charged for 2*4 h. The storage operation boundary conditions were:

- **Charging:** \( T_{\text{oil, top}} = 391 \, ^\circ\text{C} \) and \( T_{\text{oil, bottom}} = 290 - 330 \, ^\circ\text{C} \)
- **Discharging:** \( T_{\text{oil, bottom}} = 285 \, ^\circ\text{C} \) and \( T_{\text{oil, out}} = 390 - 350 \, ^\circ\text{C} \)

and the PCMs configuration was like the one shown in figure 4.

More details about the design and simulation are included in [10]. In this case also the study of LHS systems using one PCM was carried out due to comparison reasons and it is interesting to plot the results obtained (table 5 and figure 15).

<table>
<thead>
<tr>
<th>Property</th>
<th>Unit</th>
<th>CLHS</th>
<th>All NaNO(_3)</th>
<th>All KNO(_3)</th>
<th>All MgCl(_2)/KCl/NaCl</th>
</tr>
</thead>
<tbody>
<tr>
<td>( Q_{\text{sim}} )</td>
<td>MJ</td>
<td>64,5</td>
<td>74,0</td>
<td>67,7</td>
<td>56,0</td>
</tr>
<tr>
<td>( m_{\text{phasechange}}/m_{\text{total}} )</td>
<td>%</td>
<td>47</td>
<td>8</td>
<td>30</td>
<td>4</td>
</tr>
</tbody>
</table>

*Table 5. Simulated storage capacities and mass portions to undergo a phase change between charging and discharging.[10]*
The conclusions are that the CLHS has the highest phase change portion and the most uniform temperature compared to a non-cascade LHS system. The effect of the variation on the heat conductivity was also investigated showing a significant improvement when rising it from about 0.5 to 2 W/(m·K). A variation of the separate annular gap width, which contain the PCM, and also the partitioning of the axial segments was done in order to have the same portion of phase change in each PCM. This variation led to a configuration with the same amounts of axial segments for each PCM and an individual annular gap width for each PCM.

More details about the parameters and measurements accomplished and a comparison between the CLHS system and the indirect molten salt system with two tanks designed for the parabolic trough solar power plant AndaSol (Spain) can be found in [10].

The costs and the volume specific heat for the materials used in this experimental device which are given in literature [12] are shown in table 6.

<table>
<thead>
<tr>
<th>PCM</th>
<th>Volume specific heat [kWh/m³]</th>
<th>Media costs per kg [US$/kg]</th>
<th>Media costs per kWh₁ [US$/kg]</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaNO₃</td>
<td>125</td>
<td>0.2</td>
<td>3.6</td>
</tr>
<tr>
<td>KNO₃</td>
<td>156</td>
<td>0.3</td>
<td>4.1</td>
</tr>
</tbody>
</table>

Table 6. Cost and volume specific heat for some of the materials presented in this section [12].

It is important to mention that in the case where there is no axial temperature variation of the HTF the different PCMs should be arranged rather in radial direction than in axial direction in order to extract maximum benefit [28]. From [28] is interesting to show the
multiple PCMs concept developed by Wang et al. which tries the idea just mentioned. The phase change takes place everywhere in the PCM. This means the phase change rate of each component of PCM is same and remains constant during the process. The result is a reduction of the phase change time. This new arrangement is shown in figure 16.

![Diagram of multiple PCMs in LHTS unit of coaxial cylindrical tubes]

*Figure 16. Multiple PCMs in LHTS unit of coaxial cylindrical tubes [28]*

In [28] a study about the right combination of PCMs is summarized. The right combination corresponds to appropriate difference between melting points of PCMs and relative proportions of the PCMs. With larger melting points differences higher melt fractions are achieved. Sets of three PCMs were studied and the conclusion was that when the PCM situated in the middle was decreased the melt fraction increased due to a major proportion of the PCM with lowest melting point. However in all the cases there is an optimal proportion of the PCM in the middle which must be found.

It must be mentioned that although the technical feasibility of the CLHS system has been proven, further development of the concept was hindered because of: the thermodynamic penalty of going from sensible heat to latent heat and back to sensible heat; the complexity of the system; and the uncertainty over the lifetime of PCM [13].
1.1.3.2.2. PCM-based TES for Concentrated Solar Power Tower

As there is the need to store energy in different solar applications: power tower, trough, linear Fresnel collectors... there is the need to develop new technologies in order to reduce the volume and the costs. Terrafore which is a consulting and technology development firm is working on the topic.

The project is called *Heat Transfer and Latent Heat Storage in Inorganic Molten Salts for Concentrating Solar Power Plants*. It started on March 2009 and it is programmed to end on December 2011. The concept which is being developed is shown in figure 17. It works with the thermocline concept presented in the previous section.

![Figure 17. PCM-based TES design being developed by Terrafore [16]](image)

The salt is heated up and inserted on the top of the tank. Then the hot fluid is on the top and the cold fluid at the bottom and when the tank is completely charged all the salt is at the same temperature. For discharging, the salt on the top circulates to the HX-2 in figure 17 (heat exchanger) to generate superheated steam and then reaches a boiler (also a heat exchanger) where the feed water is vaporized resulting in saturated steam. The salt coming out from the boiler is around 40% solid forming a slurry with the liquid phase which is introduced at the bottom of the tank having the solid at the bottom and the liquid on the top.

As the thermocline concept is used, one of the tasks of the project has been the development of an active thermal stratification management controller required for PCM storage and it has been demonstrated that this controller can be applied to any thermocline storage.

As it is said in [17] the objective of the project is to develop and demonstrate a shell and coated tube heat exchanger that significantly improves heat transfer from a freezing salt
mixture as the heat transfer rate is one of the challenges of the LHS systems. The salt melting temperatures studied go from 275 to 350 °C and the operating temperatures to 570 °C.

There were over 700 salt mixtures examined but very few met selection criteria and the salt chosen for the rest of the project is a nitrate based mixture.

The major challenge in the project as told in [16] is that when the salt solidifies on heat exchanger surfaces, the heat transfer rates decreases significantly requiring large heat exchangers. In order to solve this problem, the approach proposed by Terrafore is based on three elements: choosing a dilute composition of salt mixtures that form a eutectic with a specific phase diagram called simple phase diagram, using an additive(s) that will cause the salt to solidify as a slush and that can be easily washed off the heat transfer surface and using a coating on the heat exchanger tubes that discourages strong adhesion of freezing salt [16].

Two coatings have been selected in the first phase of the project in order to perform more tests in the next phases.

There is not more information about the following phases of the project but as the schedule in [17] shows, the next tasks should have been already completed:

- Build a scaled model experiment by July 2010
- Conduct tests with coated tube heat exchangers between July and December 2010
- Quantify adhesion properties of coatings before August 2010
- Characterize salt morphology near and during freezing by October 2010
- Complete modeling and system analysis with PCM TES and Power Tower CSP by November 2010

1.1.3.3. Passive systems
1.1.3.3.1. Combination of sensible and latent storage

In [14] a three-part storage system for a direct steam generation device is proposed using concrete and PCM as it is shown in figure 18. In this concrete studied case the total storage capacity is about 1 MWh. The concrete storage modules are used for preheating and superheating the steam and the PCM storage module is used to evaporate/condensate the heat transfer fluid (water). This last heat transfer represents the 70% of the required energy.
During discharging, feed water enters the preheater (A) and is heated up close to the boiling curve (B), then the water is circulated through the PCM storage, where part of the water evaporates (C). The steam is separated from the water in the steam drum and is superheated in the concrete unit (D), while the remaining water is recirculated through the PCM storage [14].

In the latent storage module the material chosen as PCM was sodium nitrate (NaNO₃) by the sandwich concept using aluminum fins which is detailed in [15]. Its melting temperature is 306 ºC.

Prior to the large scale demonstration PCM storage module, a lab-scale test module with 140 kg NaNO₃ and the sandwich concept for enhancement of heat transfer was successfully tested for more than 4000 h and the tests comprised 172 melting / solidification cycles. A very high level of specific capacity and specific power was achieved, so it was demonstrated that these PCM units are a viable option for steam storage systems [14].

Once the lab-scale was proved, the design was scaled up to about 14 tons of salt with a latent heat capacity of approx. 680 kWh and adapted to the design parameters of the water/steam test loop (128 bar, 400 ºC). The tube register is arranged vertically because the sodium nitrate experiences a volume increase of approx. 10 % during melting. Then an excess volume is provided for the liquid phase at the top of the storage to guarantee the good performance of the module. The module already constructed is shown in figure 19 and detailed information about the construction of the module is presented also in [14].

In charging mode, steam with a temperature slightly above saturation properties (typically about 107 bar and 320 ºC) is routed into the PCM module where it condenses. The flow direction during charging is from top to bottom so that the condensate is removed by gravity. A condensate drain assures that the medium leaves the module only in liquid form. The module is expected to be able to condense the full mass flow of 0.8 kg/s that the test loop can provide. [14]
For discharging, the PCM module is flooded with liquid water at a temperature just below saturation properties (typically about 81 bar and 295 °C). The saturated steam produced by the module during discharge is dried in a steam separator (spherical drum on the tower left of the insulated PCM-module in figure 19). The liquid water from the steam separator is recirculated either by natural recirculation or by a pump. During discharge, the flow direction is from bottom to top. [14]

Figure 19. PCM test module at the test site; left: delivery of the module; right: insulated module [14]

For the concrete modules more detailed information including the description of the material, the design parameters and the construction of the concrete modules can be found in [14]. It is important to remark that only the superheating module was constructed and installed as it is more challenging than the preheating module and it can be used for both purposes.

Different sensors measuring temperature, pressure and mass flow were installed as the design should have undergone different tests during 2009 at the power plant Litoral of Endesa in Carbonera, Spain.

1.1.3.4. Other studies
1.1.3.4.1. PCM in solar thermal concentrating technology based on compact linear Fresnel reflector (CLFR)

Although there is not any specific design presented in [18] it is interesting to summarize the information which is given there. After a literature research on the last years materials suitable for CLFR and Multi tower solar array (MTSA) are presented. These two technologies are included in the recent advances of low cost solar thermal electricity.
The materials considered to be possibly applied in these two applications had a melting temperature in the range of 300 K to 1200 K. Over 700 K, in spite of not being in the temperature range of interest, metals such as Sn and Zn can be used. Also above 500 K molten single salts can be used such as Na₂CO₃ and KCO₃.

For CLFR systems using low pressure turbines at or below 600 K, the salts NaNO₂, NaNO₃ and KNO₃ are interesting candidates.

The heat capacity versus the melting point of the PCM candidates of this study is shown in the following figure.

![Figure 20. Heat capacity of high melting point phase change materials [18].](image1)

And in the following figure, the media cost per kWh calculated from the price of the reagent versus the melting point are presented.

![Figure 21. Media costs of high melting point PCM [18].](image2)
A numerical analysis of charging and discharging the PCM is presented where a metal (Pb) and a salt (KNO$_3$), which have some common characteristics, are compared. The model used is a double tube model as shown in figure 22. The equations followed and boundary conditions are presented in detail in [18] and as for the results of the comparison in both charging and discharging processes the time is much lower in Pb than in KNO$_3$ due to the low thermal conductivity of salts, especially in comparison with metals as it has already been mention in section 1.1.1.2.

![Double tube LHTES](image)

Figure 22. Model for the charging and discharging study [18].

The introduction of LHTES in the commercial solar thermal concentrating technology based on the CLFR at Liddell Power Station is shown in figure 23. In this plant, solar energy is used to preheat the feed water. The same concept will be applied for MTSA technology and although the LHTES design is not presented, it must be taken into account that a CLHS is not necessary in these kind of technologies as there is only one constant temperature to produce the steam that is required.

![CLFR plant design incorporating LHTES](image)

Figure 23. CLFR plant design incorporating LHTES [18].
1.1.3.4.2 Reflux heat transfer storage (RHTS)

In [54] a novel reflux heat transfer storage (RHTS) concept for producing high-temperature superheated steam in the temperature range of 350–400 °C within the frame of parabolic trough solar plants was developed and experimentally tested.

In figure 24, the scheme of the concept is shown. HTF is used to transfer heat between the thermal storage medium – PCM and two heat exchangers (HE) placed externally of the PCM at the bottom and the top and of the storage vessel. The top HE, i.e. steam generator, is fed with high pressure water (return condensate) to produce superheated steam during the storage discharge cycle. The bottom HE is used to charge the thermal storage. It is immersed in the liquid HTF and connected to the solar working fluid, e.g. a flow of solar superheated steam or solar-heated synthetic oil. During heat charge cycles, due to pool-boiling of the HTF, there is an intensive flow of vapors up through the transport channels distributed in the PCM. By condensation on the surfaces of the channels and directly on the surface of the PCM, the latent energy of vapors is transferred across the walls to the PCM. On discharge, the heat flow direction is reversed. The hot PCM causes the liquid HTF to evaporate and the vapors transmit heat to the top located steam generator via the mechanism of condensation [54].

The phase change material is a metallic Zinc-Tin alloy containing 70 wt. % of Zn. The HTF is an eutectic mixture of 26.5% biphenyl and 73.5% diphenyl oxide (Dowtherm-A). Several tests which are presented in detail in [54] were performed to test the compatibility and feasibility of the system with positive results.

It must be remarked that the use of pure zinc is more favorable because it has got a higher heat of fusion than the metallic alloy chosen. Despite of this, a metallic alloy must be chosen, since the melting temperature of zinc is 420 °C which is above the temperature of interest. The resulting melting temperature must be lower than 400 °C to have got a sufficient gradient temperature between the solar steam and the PCM.
In order to extend the operating time of solar thermal electric systems, a thermal buffer (also with the RHTS concept) is added to the system arranged in series to the main one. This buffer serves to settle the time-variable temperature of steam after the main storage caused by the gradually solidification of the alloy (non-isothermal).

Additionally, two different options for installing the thermal storage system are studied: serial connection (using the prime solar system) and parallel connection (using a supplementary solar system). The parallel connection offers a larger thermal capacity but there is an extra investment in the supplementary solar system.

Finally, although the PCM chosen is more expensive than the salts used in other latent heat storage systems, the conclusion is that there are several advantages which can lead to the success of this concept in front of other ones:

- Outstanding chemical stability of the tested Zn70Sn30 – HTF system.
- High substance density and superior thermal conductivity of Zn70Sn30, roughly 50 W/m-K in liquid state that is about two orders of magnitude over molten salts.
- High heat transfer quality demonstrated in the experiments that can be compared to the utility of thermosyphons [54].

1.1.4. Conductivity enhancement

As the conductivity enhancement of PCM is one of the most difficult challenges as it has already been mentioned, it is interesting to dedicate one section to this topic. As it has been said in the DISTOR project section, there are two main approaches to achieve the budget of improving the thermal conduction (although there were considered three approaches, the studies performed are mainly focused in two). On the one hand, it is possible to increase the heat transfer area and on the other hand, it is possible to use PCM composites. In the first case and for high temperature application, there are two options: the macro-encapsulation and the use of a tubular heat exchanger with finned tubes (what is known as sandwich concept). In the following sections, some studies about the conductivity enhancement are presented.

1.1.4.1. Sandwich concept

In reference [27] and as it has already been mentioned in the DISTOR Project, graphite foil, stainless steel, carbon steel, and aluminum are considered as possible materials in order to construct the fins. Although in a preliminary thought, stainless steel would be the chosen one due to its compatibility with the steel tubes of the heat exchangers, graphite foil or aluminum are the chosen ones. The reason is their more favorable heat conductivity which allows a less volume of the material to be used.

As told in [27], the application of fins made of expanded graphite offers several advantages besides good thermal conductivity. Expanded graphite shows good chemical stability in nitrates and nitrites at temperatures up to 250 °C and galvanic corrosion does not occur in contact with steel tubes. Since graphite foils, often used as sealing material, show a high flexibility, a close contact between tubes and fins can easily be realized.
Aluminum fins are applicable for temperatures up to 330 °C. The mounting of the fins on the tubes is an essential issue for the sandwich concept. Galvanic corrosion of aluminum with carbon steel and sodium nitrate has been analyzed, showing no critical corrosion. A good mechanical contact between fins and tubes is important [27].

In the following figure it is possible to observe the improvement which is achieved by the use of fins and also the better behavior of the graphite foils fins in front of the steel ones.

![Figure 25. Comparison of charge state for PCM without fins, with steel fins of different thickness and with graphite fins [27].](image1)

In [27] another study of the sandwich concept is given. Based on previous experiments which are summarized in table 3 from [27], an experimental demonstration is performed with the aim of optimizing the thermo-economic graphite-based sandwich concept. A storage module using sodium nitrate (NaNO$_3$) as PCM and aluminum for fins is chosen due to incompatibility between graphite and NaNO$_3$ in the working temperature range. It is shown in figure 26 and the fins can be observed in figure 27.

![Figure 26. NaNO$_3$ PCM Storage Test Module [27].](image2)

![Figure 27. Fins for heat transfer enhancement [27]](image3)
Thanks to some measurements the theoretical capacity of the storage module was calculated and was of 8.51 kWh. Some cycles (charge and discharge processes) were performed in order to calculate the real storage capacity. The result was a storage capacity of 7.87 ± 0.03 kWh in the charge mode and of 7.20 ± 0.05 kWh in the discharge mode. More details about the tests performed are presented in [27]. The difference between the theoretical and the measured capacity are awarded to uncertainties in the measurement of the oil temperature but in any case the test results and the expectations are in good agreement.

The conclusions of this report are the following: High temperature latent heat storage with high capacity factors was demonstrated at different temperature levels. The sandwich concept using fins made either from graphite or aluminum was proven as the best option to realize cost-effective latent heat energy storage. The application of graphite is preferred for applications up to 250 °C; at higher temperatures aluminum fins are used [27].

1.1.4.2. PCM composite

1.1.4.2.1. Graphite

The thermal conductivity of graphite varies from 24 to 470 W/m·K. The thermal conductivity of PCM (0.2–1 W/m K) can be intensified by a factor of 5–100 depending upon the density of graphite added into the PCM. Cabeza et al. have found that using PCM-graphite compound is the best option in applications like thermally stratified storage tank. The cheapest form of graphite is natural graphite flakes, which are composed of dense staked graphite layers. However, it is found that expanded graphite (EG) can be a better choice as it is of better absorbability due to high porosity. When the EG is mixed with PCM, the melted PCM will be absorbed in the pores of the EG results in composite PCM. The thermal conductivity and thermal capacity of the composite PCM depend upon the relative proportion of graphite and PCM [28].

An article about the conductivity enhancement by the use of PCM composites using graphite can be found in [29]. Three different types of industrial graphite were tested: graphite flakes (NG), expanded natural graphite (ENG) and expanded graphite powder (GFG). Due to their different properties, the same graphite content in the composite results in different properties (thermal conductivity, anisotropy...).

The elaboration route which was normally used is the dispersion and it is detailed in [29]. Four industrial grade composites were studied. They were all composed by the NaNO3/KNO3 eutectic at industrial grade but different graphite types and contents: the NG/400/22, the NG/400/30, the GFG/50/20 and the GFG/500/21. To understand the nomenclature: NG/400/22 is made of 22% wt of NG flakes of 400 nm. Other salts were considered as PCM in a previous study and are presented in table 1 from [29] but due to corrosion aspects, hygroscopic behavior, industrial availability and cost and hazard the nitrate eutectic was selected for further studies.

The properties of the different composites were determined: the melting and solidification temperatures, the latent heat and the thermal conductivity. The results show
that the latent heat of the PCM is not affected by the graphite content but it affects the phase change transition temperature variation which decreases at increasing graphite amount [29].

The different industrial made composites were thermally cycled 200 times and characterized. The latent heat is lowered after the 200 cycles and it depends mainly upon the size of the graphite flakes. A further study of the influence of particle size was performed and the conclusion was that it only affects for the largest amount of graphite flakes.

With respect to thermal conductivity, the results obtained leaded to the following conclusions: different graphite sizes and types lead to a significant variation in conductivity, a larger particle size or a lower density particle (which leads to a larger number of dispersed particles) are favorable to the creation of a better conductive network within the PCM. The influence of temperature on thermal conductivity was also studied and it was proved that the thermal conductivity is strongly temperature dependant [29].

As the thermal conductivity is still low for the solar application (within the framework of DISTOR project the target is a thermal conductivity of 8 w/m·K) it is proposed to use additional conductive graphite fins radially distributed in the volume with respect to the heat transfer tubes. To study the efficiency of this approach a simulation was performed and it is detailed in [29]. The conclusion of the simulation is that each particular case of application (in term of geometry and working parameters) will lead to a specific technico-economic optimum defined by the graphite amount within the composite and the interfoil distance between the graphite fins.

1.1.4.2.2. Graphite and metal foams

In [30] there is another study of the conductivity enhancement in high temperature storage systems. The PCM used is NaNO$_3$ (properties have already been shown in table 3) and there are two improving ways presented: using metal foam or using expanded graphite. An experimental device which is shown in figure 28 was constructed in order to study the conductivity enhancement and its advantages in the charge/discharge processes. More details about the sensors, devices, insulation and container are presented in [30].

![Figure 28. Experimental device for the study of the conductivity enhancement [30].](image)
Three different charging processes and one discharging process are studied. The results show that the conductivity enhancement with the two methods is achieved. Porosity plays an important role and the conclusion is that reducing porosity and cell size significant conductivity enhancement is accomplished. It must be taken into account that the lower porosity implies a lower total heat storage capacity.

One of the charging processes studied correspond to the phase change of the PCM (250 to 350 °C) and the stages and dominant heat transfers mechanisms during the process are explained with detail in [30]. It must be mentioned that NaNO₃ has a solid-solid phase change at 276 °C and this leads to a little step in the temperature curves.

Another point to be remarked is that although metal foam structures enhance conduction, natural convection can be weakened by metal foams. In the case of expanded graphite natural convection can be severely suppressed due to its small pore size and higher viscosity inside the composite [30]. Due to this fact, after the complete melting of the PCM, temperature curves are much more complex with metal foams or expanded graphite than in the case of the pure salt. In some of the tested containers this fact leads to a lower heat transfer in the liquid phase region, namely porous materials do not always enhance heat transfer in every regime.

For the discharging process it is shown that both methods allow a faster discharging process thanks to achieving also an enhancement in the heat transfer process.

Besides the charging and discharging studies, a report about the compatibility in terms of corrosion between NaNO₃ and metal foams is also presented. Heavy corrosion effects in the metal foams can be observed after some tests reducing the positive effect of using metal foams in the enhancement of thermal conductivity of PCM.

Although the overall performance of metal foams is superior to that of expanded graphite, graphite has a good chemical stability to resist corrosion as it has been mentioned before in section 1.1.4.1. The disadvantage of the expanded graphite pointed in [30] is the difficulty of the composite manufacturing (mixing process) with possible voids inside the composites with larger fraction of expanded graphite.

Finally a discussion about the economic impact of the use of composites in TES is presented in [30]. The loss of thermal storage capacity is worth by the enhancement in heat transfer. It is said in [30] that the overall cost of the TES system using expanded graphite can be estimated as a little change or even up to 10% reduction thanks to the reduction of the area of heat exchangers and the relative low cost of the expanded graphite. In respect to the use of metal foams, as they have a higher price than the expanded graphite, it is estimated a 10-20% rise on the cost of the TES system [30].
1.1.4.3.1. Other conductivity enhancement approaches: Heat pipe heat exchanger with latent heat storage

A more recently approach to counteract the low thermal conductivity from the phase change materials is the use of heat pipes. The concept is shown in figure 29. The heat exchanger consists mainly of four parts. The hot fluid flow passage (6), the PCM chamber (7) and the cold fluid flow passage (5) are connected by a number of heat pipes (3). The phase change material (8) is stored in the PCM chamber. In order to enhance the heat transfer process, annular fins made of pure copper (in the case of this reference) are attached to the heat pipes. As one can see from the figure, the sizes of the hot fluid flow passage, the PCM chamber and the cold fluid passage can be designed independently, which presents one of the major advantages over other latent heat thermal storage systems [52].

![Figure 29](image)

In this concept it is possible to work in three different modes in comparison to the two usually modes (charging and discharging) as it is possible to charge and discharge at the same time. This third mode gives a more flexibility to the system in comparison to the other latent heat storage systems presented.

A thermal network model is developed and used to analyze heat transfer in a latent heat storage unit for solar thermal electricity generation in [53]. Two storage configurations are considered; one with PCM surrounding a tube that conveys the heat transfer fluid, and the second with the PCM contained within a tube over which the heat transfer fluid flows. Both melting and solidification are simulated. It is demonstrated that adding heat pipes enhances thermal performance, which is quantified in terms of dimensionless heat pipe effectiveness [53].
Although it has the budget of developing a latent heat storage system for high temperature applications, the experiments performed used water as a HTF and a paraffin with a Tm of 52.1 °C as the PCM. As it is outside the temperature of range of interest, no more details about the model will be presented. It must be remarked that is a concept which must be taken into account as possibly further studies will be available during the next years.

1.2. THERMO-CHEMICAL STORAGE

As it has been said in the introduction the chemical storage is the third approach studied to store heat. This one is the approach which has been developed less. It is based on using reversible thermo-chemical reactions: chemical compounds store energy by an endothermic reaction and the energy is then released in the recombination of the compounds by an exothermic reaction [7].

The most interesting advantage of this method is that the enthalpy of reaction is considerably larger than the specific heat or the heat of fusion. Then the storage density is much better. Some of the compounds and reactions with their respective temperatures and heat of reaction and the storage density are shown in table 4 from [7] and in table 8 from [8].

Another advantage from this approach is that thermal insulation is not required (lower cost) but on the other hand the chemical compounds must be heated and cooled before reactions (higher cost).

1.2.1 Properties required

As told in [7] when selecting the chemical compounds, some properties are required in order to achieve the desired performance:

- **Reversibility**: The reaction must go to completion; no side reactions or changes in reaction rate with time should occur.
- **Reaction rates**: Both forward and reverse reactions must be rapid enough to absorb all the available energy or release it promptly. The reaction rates must not decrease with cycling. Such a decrease can be observed, if structural changes of the storage medium occur (e.g., crushing of particles into powder).
- **Controllability**: The reactions must be controllable because they have to be turned on and off when required. Controllability is achieved by product separation, by controlling the temperature and pressure, or by catalysts.
- **Ease of storage**: The reaction products have to be easily separable prior to storage. Reactions must not occur at storage temperature.
- **Safety**: Toxicity, inflammability, and corrosiveness of the reaction products may pose unacceptable safety hazards.
- **Cost**: Cost should be low; this requires materials which are readily available.
1.2.2. Classification of reactions

There are two different types of reaction which can be used to store thermal energy. On one hand there are the dissociation reactions. In these reactions a solid or a liquid release gas in the endothermic reaction (charging process) and for discharge the gas is recombined with the solid or liquid. The dissociation products have to be separated and stored individually in order to avoid their spontaneous recombination due to a change in the equilibrium by a temperature decrease or a pressure increase [7]. One example of this kind of reaction is the dissociation of Ca(OH)$_2$, which will be presented in detail in section 1.2.3.3.2.

On the other hand there are the catalytic reactions. The name come from the fact that a catalyst is needed in order to increase the reaction rates. These are usually gas-gas reactions. The reaction products are transported in a pipeline from the source (charging process) to the consumer (discharging process). This pipeline is known as chemical heat pipeline or chemical heat pipe. An example of this kind of reaction is the carbon dioxide reforming of methane which is presented in section 1.2.3.2.

Catalytic reactions product do not need to be separated which implies a reduction in their cost. In the dissociation reactions, although separation is easy, when one of the products is gas there are generally some difficulties with the storage (cost, complexity, volume) as it is presented in [7] with some possible options.

The most studied reactions for thermo-chemical energy storage until now are the dissociation of ammonia and the steam and the carbon dioxide reforming of methane. The applications and studies which have been performed will be presented in the next section.

1.2.3. Applications

1.2.3.1. Dissociation of ammonia

The dissociation of ammonia (1) is one of the most investigated reactions for the thermo-chemical energy storage as mentioned before. It has been studied by the Australian National University (ANU) for more than two decades and a summary of the experiments and investigations performed with some results will be presented in this section.

\[
\text{NH}_3 + \Delta H \leftrightarrow \frac{1}{2} \text{N}_2 + \frac{3}{2} \text{H}_2 \quad (1)
\]

There are few reasons for the high interest in this reaction. First of all the use of ammonia in industrial and chemical applications is wide and then the knowledge of the compatible materials, uses and performing of the reaction is also extensive. On the other hand no side reactions take place in and the separation of reactants is spontaneous due to condensation of ammonia fraction at ambient temperature. These are two important advantages in comparison to other reactions considered for thermo-chemical energy storage such as the carbon dioxide reforming of methane. Working at ambient temperature is feasible thanks to the low vapor pressure of the substance. Finally another advantage which must be remarked is the low toxicity of ammonia. [31] [32]

The most important disadvantage of using the ammonia dissociation is its low enthalpy of reaction (66,5 kJ/mol) in comparison to other reactions such as the carbon dioxide reforming of methane (247 kJ/mol).
The recent literature about the ammonia thermo-chemical energy storage is mainly based on the work performed by ANU and is based on a pseudo-homogenous model adapted from the work of Richardson et al. (1988) and the model is described in detail by Lovegrove (1995).

In [33] a study about an endothermic reactor for an ammonia thermo-chemical energy storage system is presented. The most important points about it is the feasibility of reaching pressures up to 30 MPa which provides benefits in the heat transfer coefficients and the volumetric energy storage density. In this technical report also different receiver/reactor concepts are investigated. The details about the tests and results can be found in [33] but it is important to remark the option which is most cost effective is the use of a cavity of directly irradiated catalyst filled tubing as the one shown in figure 30. There are forty-three 500 mm long reactor tubes and the catalyst tube cross-section used was the same used before in the electrically heated dissociation reactor (15.6 mm i.d. 24 mm o.d. Inconel) which was used before the installation of the 20 m$^2$ dish in the ANU campus.

Figure 30. Receiver/reactor concept chosen for the ammonia thermo-chemical storage system developed by ANU [33].

Once the receiver/reactor concept was chosen, the following study which can be found in literature is the one presented in [34]. It presents the results regarding the design, performance modeling and preliminary costing of key system component of a complete 10 MW$_e$ solar thermal power plant system designed for 24-h base-load operation. The conceptual design is shown in figure 31.
Figure 31. Conceptional system design of a solar thermal power plant using an ammonia-based thermochemical closed-loop for energy conversion, storage and transport [34].

The system as told in [34] is based around one or more endothermic reactors (R-101), which are driven by solar heat, and a single exothermic reactor for heat recovery via generation of superheated steam (R-102). Associated with these reactors are counter-flow heat exchangers (E-101 and E-102). They exchange heat between the inlet and outlet reactant flows and so ensure the remainder of the system (e.g. energy storage, transport piping and compressors) operate at ambient temperature. In table 1 from [34] the key operating parameters of the plant are presented.

Details about the different components of the plant:

- Solar collectors: two different technologies have been considered. One is a 400 m² paraboloidal dish solar collector and the other one is a 150 m² stretched-metal-membrane heliostat.
- Ammonia receivers/reactors: there are also two different concepts presented using tubular fixed-bed catalytic reactors. The first one is based on the receiver/reactor presented in figure 30 which was chosen as the most cost effective in the previous study presented in [33]. The second one is a conventional, megawatt-scale tube-bundle reactor which uses hot air to heat up the reactor.
- Ammonia synthesis components which are based on a previous study [32]. More information about the ammonia synthesis reactor based on a 1 kW_chem. laboratory-scale synthesis reactor is presented later.
- The steam Rankine cycle which is a conventional single-stage one due to space and control limitations.
- The heat exchangers which are counter-flow heat exchangers specially developed for this application by FC Consulting.
- Compressors and pumps.
- Storage system: two storage tanks concepts. In the first one the ammonia is stored at 25 MPa which is the working pressure of the reactor and in the second one it is stored in a low-pressure ammonia tank of typically 3 MPa.
- Piping.
with its working parameters, information about the models used and the overall performance are presented in [34].

Finally there a preliminary cost estimation is performed based on 1997 prices, past project experiences and quotes from European and Australian manufacturers. The result of this estimation is shown in table 5 from [34] and the results shown are encouraging as they indicate the potential for economic viability of this solar power plant concept. Besides the positive economical results, it is important to remark that the first receiver/reactor concept is more cost-effective than the second one due to a more direct energy transfer from the sun to the ammonia system [34]. In the laboratory-scale and other experiments for the study of the ammonia thermo-chemical energy storage systems this reactor concept is the one used.

As mentioned before, there is more information available about the ammonia synthesis reactor in the literature [32]. Although different ammonia synthesis reactors have been already used for other applications it is needed to perform a study about it. The reason is that the optimized working conditions for each application vary. In the case of thermo-chemical energy storage it is important to maximize the output of high-temperature heat for electrical power generation in comparison to the maximization of the production rate of ammonia which is desired in other applications.

In order to find the optimal working conditions experimental data have been recorded. A model based on the two-dimensional pseudo-homogeneous packed-bed catalytic reactor model mentioned before has also been validated after performing a calibration with the experimental results. Details about the experimental results and this calibration process can be found in [32]. The 1 kW<sub>chem.</sub> laboratory-scale scheme used in the study can be observed in figure 32.

![Figure 32. Experimental arrangement of ANU's solar laboratory-scale closed-loop energy transfer and storage system [32].](image-url)
Both liquid ammonia and the 3:1 hydrogen/nitrogen gas mixture are stored at ambient temperature and high pressure in a 20 l storage vessel. The ammonia laboratory allows handling and analysis of reactants up to 30 MPa [32].

For dissociation, liquid ammonia is drawn from the storage vessel and passed via a circulation pump to the nitrogen gas backed accumulator. It is subsequently passed to one of the two receivers/reactors mentioned before [32]. The whole system operates at constant volume and so the pressure increases as the ammonia in the vessel is dissociated [35].

For synthesis, gas is drawn from the storage vessel by a compressor pump and stored in a 10 l buffer vessel. Subsequently, it passes through a chilled separator where some fraction of the ammonia vapor condenses and separates from the gas and travels back to the 10 l vessel. The purified gas continues on the heat recovery reactor which is illustrated in figure 33. More detailed information about the laboratory installation and the measuring devices used for testing is presented in [32] and [35].

It is important to mention the conclusions reached about the maximization of the output heat. The external reactor wall temperature plays an important role in the output heat: higher temperature contribute to higher reaction rates while an approach to chemical
equilibrium conditions work to reduce them. Regarding the pressure, thermal output increases almost linearly with operating pressure.

In [35] more studies about the laboratory-scale closed-loop installation presented before are presented. It is important to mention that the dissociation reactor used is an improvement of the one shown in figure 30. The new version is shown in figure 34.

It differs from the previous model in that the endcap is sealed against leaks by welding rather than using a copper gasket. It has a reduced thickness of catalyst bed, as modeling indicated that the significantly improved heat transfer associated with higher gas velocities strongly reduced the reactor wall operating temperature and outweighed the loss of catalyst surface area. The reactant flow direction has also been reversed, ammonia now enters from the base and flows through the catalyst bed toward the tip. This reversal of flow direction allows the hot product gases to give up some of their heat to the bed as they pass back down the centre tube. Then this new reactor uses less catalyst material [35].

An experiment with the details of the working conditions is presented in [35]. It proves the ability of solar ammonia dissociation reactors to operate smoothly and effectively through transients. Also the synthesis reactor was proved to work correctly as the heat was being...
exchanged and the ammonia level at the end of the performance the ammonia level was close to the initial conditions [35].

Finally, some considerations about the scale-up of the system to accept the 15 kW\textsubscript{sol} input from the 20 m\textsuperscript{2} installed at ANU will be mentioned. Twenty 0.5 m long Inconel catalyst filled tubes of similar design to the 1 kW unit are positioned in a conical arrangement around a water-cooled cavity receiver aperture. It is shown in figure 35. The water-cooling can be substituted by ammonia in future designs [35]. The reactor in operation can be observed in figure 36.

![Diagram of cavity receiver with 15 kW\textsubscript{sol} solar dissociation reactor and its assembly on ANU's 20-m\textsuperscript{2} dish without insulation fitted [35].]
The scaled-up heat recovery reactor is constructed from a bundle of 19 tubes virtually identical to the 1 kW prototype. The design and the partially completed assembly are shown in figure 37. Main of the rest of the components of the system function directly with the larger reactors. Some massflow control elements required upgrading and a larger-volume reactant storage vessel has been incorporated to complete the scale-up [35].
It is interesting to remark the conclusions which are presented in [35]. The small laboratory-scale system has shown that ammonia dissociation receiver/reactors are ideally suited for operation through solar transients; ammonia synthesis heat recovery reactors are capable of stable, predictable operation with heat recovery at temperatures suitable for high-quality superheated steam production and reactant storage and handling, for the ammonia system at pressures up to 30 MPa, can be achieved using standard components and manufacturing techniques. These encouraging conclusions have led to the scale-up just presented.

Among the studies performed it is important to mention that there have been run some optimization exergy rate output analysis on the exothermic side of the 1 kW closed-loop. Details about this analysis including the system considered, the exergy calculations and the results can be found in [36].

It is interesting to mention the different optimal working conditions between the maximum thermal power and the maximum net exergy rate output. Whilst equal distance between average composition curve and chemical equilibrium along the reactor gives maximum recovery of heat, maximum net exergy output is achieved with the internal average composition being close to equilibrium at the end of the bed. Also the average outer reactor wall temperature plays an important role in both cases having the maximum net exergy rate output at temperatures slightly higher than for the maximum thermal power. Then it is important to choose and maintain the best average wall temperature taking in consideration both cases [36].

In [38] some tests (experimental and simulation) on an ammonia receiver design for a 500 m² dish which has been constructed in the ANU campus are carried out. Five different receiver geometries were considered and three of them were thoroughly studied and are shown in figure 38. These designs were based on geometric variations to the prototype ammonia receiver used in the 15 kW$_{\text{chem}}$ installation presented before.

![Figure 38. Three receiver configurations viewed looking in to the aperture. Left: Receiver with a 7.5º half cone angle (front shield removed). The indents in the wall insulation indicate the position of the tubes in the frustum with a 17.9º half cone angle. Centre: Receiver in a frustum with base radius 77 mm (front shield removed). Right: Receiver as a frustum with base radius 77 mm, but with minimal insulation on feeder tubes (front shield in place).](image)

The solar to chemical efficiencies were studied for each case varying the mass flow rates of the ammonia through the receiver and then a comparison between the three cases was performed. The efficiencies values are generally quite low due to the masking of the dish
that has effectively considerably reduced its concentration ratio. The maximum difference between the performances of the three cases is only 7% but the receiver with 7,5º performs better than the cone with 77 mm radius base, which performs better than the receiver with 17,5º half cone angle.

It is important to mention that the simulations and experimental results can not be directly compared as the simulation model does not take into account the losses from heat exchanger or due to flux spillage on the shield. Then the heat transfer code of the model which is more detailed in [38] and its references must be revised. Then further work must be done to calibrate the simulations with the experimental data in order to use the model to evaluate different receiver designs for the prototype Generation II 500 m² dish on the ANU campus.

Besides, it is necessary to investigate other methods of increasing the receiver efficiency. The options to be considered include the optimal number of tubes, the optimal tube length and optimal tube diameter [38].

Finally and although ANU have been working on the development of dish concentrators, trough concentrators are the ones which have been more developed and which represent the highest share of installed capacity. Because of that a preliminary study about the application of the ammonia cycle in trough solar power plants is summarized in [37] with encouraging results and remarking the need of further investigation. From [31] there is the scheme of the trough solar power plant using ammonia cycle concept proposed in [37]. It is shown in figure 39.

![Figure 39. Sketch of the parabolic trough driven ammonia system [37].](image)

In this new direction of this new research it is interesting to remark the diploma thesis which is presented in [31]: investigation of the feasibility of a parabolic trough driven solar energy storage system using ammonia. As it is told there, except for the dissociation reactor, all the other components of the system are available on the market for industrial components. Then the diploma thesis focuses on the modeling of a dissociation reactor which fits a parabolic trough solar power plant.
Three different concepts are investigated and presented in detail in [31]: a single tube heat collecting reactor element (HCRE), a tube bundle HCRE and a cavity type HCRE. The advantages and disadvantages of the different approaches and the necessary considerations in order to adapt the concepts to a parabolic trough solar power plant are discussed also in [31]. Some simulations are run in order to test its expectable performance and to demonstrate opportunities for further research within this field.

The design chosen was the cavity type reactor due to reduced heat loss and the ability to expose the reactor elements to a staggered flux distribution. Some simulations varying the diameter and ammonia feeds were performed in order to find the best nominal diameter of the reactor which resulted to be of two inch.

To obtain constructive limits, the limits regarding pressure drop and maximal reactor temperature were chosen, and the maximal achievable reactor length was determined out of the result files.

Depending on the different reactor elements, length, diameter and the reaction extent which could be expected at the corresponding massflow one design and its operation was chosen to simulate a bench mark design for the further modeling. Depending on the results of this bench mark a next step was performed assuming a staggered flux distribution in comparison to the assumption of isothermally operation made in the first steps.

The results obtained from these simulations show that remarkable reaction extents are achievable even for a non optimized reactor design regarding geometry and catalyst used. Then, it can be concluded that the dissociation of ammonia should be achievable using a cavity type reactor operated with the LS-3 type trough considering the assumptions taken within the work presented in [31]. It is also concluded that the new trough design which allow a higher temperature tolerance of the absorber could lead to higher reaction extents and thus to higher overall efficiencies of the system [31].

1.2.3.2. The steam and the carbon dioxide reforming of methane

The reforming processes of methane are catalytic processes. In the case of the carbon dioxide reforming of methane, the overall \( \text{CO}_2/\text{CH}_4 \) reaction is shown in (2) and its enthalpy of reaction is of 247 kJ/mol at 25 °C.

\[
\text{CO}_2 + \text{CH}_4 \leftrightarrow 2 \text{CO} + 2 \text{H}_2 \quad (2)
\]

And the thermo-chemical heat-pipes concepts using this reaction are shown in figure 40. For the storage of heat only the closes loop concept is of interest.
In [45] CSIRO/Pacific Power research on using the carbon dioxide reforming of methane reaction for solar energy storage is described. Fluidised-bed reactors were chosen for both endothermic and exothermic reactions.

Several potential catalysts were prepared and characterized for both reactions. The CSIRO catalysts chosen were used to conduct both reactions in small-scale fluidised-bed reactors under practical reaction conditions without carbon formation. Stable, equilibrium CO₂/CH₄ reforming was obtained over a wide range of operating conditions (650 – 950 °C, 0.1 – 0.7 MPa, 0.8 – 1.2 CO₂/CH₄ feed molar ratio) with thermo-chemical energy storage rates equivalent of up to 30 MWth/tonne of catalyst being achieved. The exothermic reaction was conducted in a fluidised-bed reactor in such a manner than more than 90 % of the reaction heat can be recovered at temperatures around 550 °C, thus allowing the closed loop version of this technology to either be integrated with a conventional Rankine cycle power plant or be used for the production of high grade process heat [46].

For the steam reforming process of methane, the reaction which takes place is shown in (3) and the enthalpy of reaction is 206.2 kJ/mol at 25 °C.

\[
\text{CH}_4 + \text{H}_2\text{O} \leftrightarrow 3\text{H}_2 + \text{CO} \quad (3)
\]

The same concepts as for the carbon dioxide reforming of methane can be applied but as the temperature range needed for the reforming processes is between 750 and 950 °C, no further information about these two thermo-chemical reactions will be presented as it is outside the range of interest. They have been mentioned since they are one of the most studied and a promising technology is being developed around these two thermo-chemical reactions [47].
1.2.3.3. Dehydration/Hydration cycles

1.2.3.3.1. Medium temperature (200-300ºC) chemical heat storage using mixed hydroxides

In [39] an hybrid system with a chemical heat pump and a high-temperature process is proposed. In this case the system is focused on the efficiency enhancement of vehicles as carbon dioxide emission due to vehicle usage has a significant impact on global warming [39].

An electric hybrid system (EHS), which combines an internal combustion engine (ICE) with an electrical battery and a power controller as shown in figure 41, has good energy saving performance. Upon braking, the EHS kinetically converts surplus work into electricity, which is stored in the electrical battery. The stored electricity is used to start the vehicle or to assist during a period of low-efficient operation of the ICE.

![Figure 41. Thermal hybrid system (THS) combined with a medium-temperature chemical heat pump and a high temperature exhaust gas from ICE, in comparison with the electric hybrid system (EHS) currently employed in some hybrid vehicles [39].](image)

The authors are proposing a thermal hybrid system (THS) that combines an ICE with a thermal battery and a thermal power controller for efficient use of exhaust heat from ICE as shown in figure 41 in comparison with the EHS. The thermal battery in THS is used for heat storage of surplus exhaust heat from ICE. The stored heat in the thermal battery is supplied to heat output side for tasks such as cold start, catalyst heating for exhaust gas reduction and vehicle cabin heating. The THS is expected to contribute to load leveling of ICE operation, improvement of fuel consumption efficiency, and reduction of CO₂ emission. The THS concept can also be applied to the cogeneration engine and solar thermal power systems [39].

Different chemical reactions were surveyed and the most promising one was the metal oxide/water reaction (4).

\[ \text{MO} + \text{H}_2\text{O} \leftrightarrow \text{M(OH)}_2 \quad (4) \]

Several reactions of this type are shown in figure 2 from [39] and approach finally chosen was a mixed between magnesium hydroxide and nickel hydroxide as it combines the high reactivity with water vapor from magnesium oxide with the potential as new reaction system of nickel hydroxides.
Several tests were performed and it was found that the mixed hydroxide can expand operation heat storage temperature by changing the composition of the added cation in its. The heat capacity of $\text{Mg}_{0.5}\text{Ni}_{0.5}(\text{OH})_2$ was estimated to be 165 kJ/kg and although the amount of reacted fraction at hydration decreased in the first cycles, it was constant for the following cycles and the repetitive durability of the mixed hydroxide was proved assuming the same behavior as the pure $\text{Mg(OH)}_2$.

The integral reactivity of the developed mixed hydroxide was investigated using a packed bed reactor. It was proven that the mixed hydroxide can be decomposed by storing heat at approximately less than 300 °C and outputting heat at approximately 230 °C by physical sorption and at approximately 185-200 °C by exothermic hydration under the working conditions presented in [39]. The heat output temperature is expected to be increased by increasing the vapor pressure in the hydration process.

Finally, it can be concluded that the mixed hydroxide showed new possibility of chemical heat pump and it was demonstrated the feasibility of a hybrid system combining a medium-temperature chemical heat pump and a high-temperature process for efficient primary energy use. More details of the results and devices used for the tests can be found in [39].

1.2.3.3.2. Dehydration/Hydration cycle of $\text{Ca(OH)}_2/\text{CaO}$

In [41] the study of the dehydration/hydration cycle of $\text{Ca(OH)}_2/\text{CaO}$ in the application of thermal storage is performed by the use of a reactor designed for the purpose. It must be mentioned that normally in the hydration step water is usually added in vapour phase but in the experiments performed liquid water was added at 0 °C. The cycle is shown in (5).

An input energy of 148,6 kJ ($54,0 + 94,6$) can decompose 1 mol of $\text{Ca(OH)}_2$ at 25 °C to $\text{CaO} + \text{H}_2\text{O}$ at 510 °C. The produced $\text{CaO}$ can be easily stored as a solid at 25 °C loosing 23,5 kJ. If water is allowed to escape to the atmosphere, instead of storing it at 510 °C in vapour phase, 61,5 kJ are additionally lost. Therefore, 57,3 % (85,0 kJ) is the portion of input energy that results in sensible heat of the products, and the remaining 42,8 % (63,6 kJ) is actually stored in the form of chemical energy.

When the stored high temperature steam at 510 °C is used in the hydration step 125,1 kJ could be recovered (84,2 % of the initial input of energy), but the recovery of stored heat using liquid water at 25 °C can only provide the already mentioned 63,6 kJ [41].
There are several applications for this cycle but this experiment focuses on the use of the cycle for preheating the combustion engine in motor vehicles. Then the exothermic reactions heat is partially used to heat CaO and water from 0 ºC to 25 ºC (63.6 kJ) and the rest (60.6 kJ) are left for this particular application [41].

Only 20 cycles have been studied because after them the carbonation of CaO and Ca(OH)$_2$ becomes a trouble in system performance. The scheme of the designed thermal prototype is shown in figure 42. Specific heats, reaction rate and enthalpy, mass losses and heat release were monitored and the details of the experimental setup and results can be found in [41].

![Figure 42. Scheme of the designed thermal storage prototype [41].](image)

It is interesting to show the conclusion reached by the experiments carried out which should be considered in further studies.

- Due to carbonation, the process does not reach a full reversibility in the dehydration step, and therefore could be used with a limited number of cycles, that however can go up to 20. A periodic dehydration at 1000 ºC could be necessary to recover full system reversibility.
- Efficiency is higher when the heat recovery solution is changed and a system with renewal of this solution in a prototype of Cu is a better design to heat water from 0ºC up to medium final temperature of 33.3 ºC.
- Hydration water addition should be carried out pouring it slowly with a uniform distribution along the whole mass of CaO as in a spray [41].

In [42] a previous study of the application of the same cycle for the same purpose was performed with the objective of determining how much heat energy can be produced and how long does it take to produce it; the maximum temperature obtainable; and whether it could be chemically recycled in that rig configuration without losing its potential for producing heat; and finally to evaluate the impact of utilizing such a thermal store for reducing energy consumption and pollutants from motor cars.
The scheme of the experimental device used for the experiments is presented in figure 43. The details of the experimental method and the results obtained can be found in [42]. Also a theoretical model was developed and confirmed by comparison to the experimental data. The model can be used for future calculations. The important points to remark from the study are the conclusions itself.

Figure 43. Scheme of the experimental device used in [42].

In all the series of tests conducted, rapid interaction of fresh/dehydrated CaO and water vapor was observed. The maximum temperatures obtained which vary between 207 and 219 °C indicate that CaO is capable of being recycled provided it is well dehydrated and properly stored, thus confirming other established studies on material recyclability. Even though there were slight differences in the maximum temperatures attained, there was no observance of any hindrance to the reaction rate by diffusion of water vapor through the bed. However, insufficient heat transfer due to low effective thermal conductance of the bed and the high thermal contact resistance of the wall was observed. Analysis of the experimental data showed that the average deviation from the theoretical values was about 17% and uncertainties in the primary data accounted for about 8%. 
In conclusion, the experiment has demonstrated that it is feasible to integrate a fixed-bed thermo-chemical energy store in a motor vehicle. However, for effective heat and mass transfer, it would be necessary to redesign the reaction bed as a bonded disc type with extended fins (which were used in the design presented before) and have them arranged in a cascading form. Alternatively, porous vapor tubes can be inserted into the reaction beds to ensure a low flow resistance along the water vapor line for the vapor to diffuse in a more compact structure over short distances only.

More recently, the DLR is studying this reaction to be applied in the temperature range between 400 and 650 °C as the reversibility of the reaction has already been proved. In [45] some simulations and experimental results are presented. Two different reactor concepts are simulated by a 2D-model based on a model previous developed for metal hydride reactions.

An indirectly heated reactor concept as shown in figure 44 a) has been simulated. It uses an external heat exchanger and its performance is strongly limited due to the low thermal conductivity of the fixed bed. The second reactor concept, directly heated reactor with direct heat exchange by flow of a heat transfer fluid through the reaction zone, overcomes this problem and is not limited by heat transfer.

Then an experimental setup as shown in figure 44 b) has been built in order to validate the simulation results of the directly heated reactor concept. First experiments have shown that the reaction can be conducted reversibly and the heat of reaction is partially transferred to the gaseous stream flowing through the solid reactants.

![Figure 44. a) Indirectly heated reactor concept simulated by the DLR b) Experimental setup built for the directly heated reactor concept [45].](image)

It is concluded that reactor kinetics could have an impact on reactor performance and therefore have to be determined experimentally. Due to the thermal capacity of the gaseous stream needed to take-up all of the heat being generated by the reaction the pressure drop through the reaction zone can be a limiting factor in the concept with direct transfer.
Experiments are currently under way to investigate the influence of various operating conditions and to get close to the commercial demonstration of thermo-chemical energy storage.

1.2.3.3. Ni-doped and undoped Mg-MgH₂ materials

In [44] the use of Ni-doped and undoped Mg-MgH₂ materials for thermo-chemical heat storage in the temperature range of 450-500 ºC is also studied. It is proved that both types of materials have excellent cyclic stability and high hydrogenation rates. Several cycling tests are carried out focusing mainly on the causes of hydrogen capacity losses and finding the operational conditions which minimize these losses.

Details about the experimental devices, the samples of the materials and the cycling tests are detailed in [44] with their respective results. It is important to mention that the properties of the different materials are sensitively dependent upon all the details of the experimental cycling conditions: hydrogenation/dehydrogenation pressure and temperature, distance to the equilibrium conditions, time allotted for hydrogenation/dehydrogenation processes, the temperature regime of heating and cooling and the previous cyclization history of the material sample.

The conclusion is that these materials can be applied for applications such as solar generation of heat and cold, heat pumps, hydrogen storage, solar power generation via Stirling engine and storage of industrial heat in the temperature range of 450 to 500 ºC [44].

1.2.3.3.4. Mg₂FeH₆

The reversible Mg₂FeH₆ hydride system has been studied as an option for thermo-chemical storage at approximately 500 ºC. In [43] there is a report about the thermodynamic properties and cyclic stability of this system by an elemental analysis, thermovolumetry, X-ray powder diffraction, PCI measurements and transmission electron microscopy. The preparation of the different samples and experimental devices used and the experiences carried out are presented in [43].

The reaction with its enthalpy of reaction (77.4 kJ/mol H₂) resulted to be the one shown in (6).

\[ 2\text{Mg} + \text{Fe} + 3\text{H} \leftrightarrow \text{Mg}_2\text{FeH}_6 + 77.4 \text{ kJ/mol H}_2 \] (6)

A comparison between this system and the MgH₂ system is also performed reaching to the conclusion that Mg₂FeH₆ shows a lower hydrogen dissociation pressure which is favorable from technical and economical points of view, since the required heat storage containers would be less costly and the H₂ leakage rate would be reduced.

Also the mixed Mg₂FeH₆-MgH₂ is studied and characterized resulting to be also a potential thermo-chemical hydrogenation/dehydrogenation system.

The conclusions are that these two systems have got a higher gravimetric and volumetric thermal energy density than the sensible or latent heat storage materials and the stability in cycle tests at around 500 ºC has been proved. Further advantages include the low
price of the materials, the free choice and constancy of the heat delivery temperature by controlling the applied hydrogen pressure and absence of heat losses with time.

On the other hand further research must be carried out in order to find the optimal working conditions regarding the storage density, the intermediate storage of the desorbed hydrogen and engineering. This studies are especially needed in the mixed Mg$_2$FeH$_6$-MgH$_2$ hydride system with a low proportion of Mg$_2$FeH$_6$.

1.2.3.3.5. Dehydration/hydration cycles of salt hydrates

Salt hydrates are chemicals that release water when heated thereby dissociating into an anhydrous salt. When heated to a charging temperature thermochemical dehydration of the salt hydrate occurs releasing water from the compound. The anhydrous salt has a larger energy content and can be stably stored at ambient temperature for long durations. When thermal energy is required a water vapor flow across the salt once again hydrates it and the stored heat is released to the ambient.

These salts have both a high volumetric heat storage capacity, large thermal conductivity and are also inexpensive. However, liquid supercooling is of major concern and their poor nucleation properties make them inapplicable for reuse.

Although reuse is not possible, they can be considered as a long-term thermal application as for example utilizing solar energy to dissociate a salt hydrate in the summer, storing the anhydrous salt and water vapor separately until the winter and then recombining them to release heat to warm buildings [48].

In [48] a model to study the salt hydrates thermochemical storages is presented using it in the concrete case of the magnesium sulfate heptahydrate (MgSO$_4$-7H$_2$O). The reaction will be the following (7),

$$\text{MgSO}_4 \cdot 7\text{H}_2\text{O} \xrightarrow{\text{Addition of heat (solar)}} \text{MgSO}_4 + 7\text{H}_2\text{O}$$  (7)

and the model can be applied to other salt hydrates in order to study the different parameters which affect the process performance and select the optimal materials for the salt hydrates. Any conclusion about the feasibility of using MgSO$_4$-7H$_2$O is mentioned.

1.2.3.4. Reduction and oxidation reactions (REDOX)

General Atomics in collaboration with the German Aerospace Center (DLR) is studying the feasibility of using REDOX reactions to store and release heat as it is shown in (8):

$$\text{MnO}_{2x+1} \rightarrow \text{MO} + x\text{O}_2 \quad \text{Endothermic reaction (charge)}$$  

$$\text{MO} + x\text{O}_2 \rightarrow \text{MnO}_{2x+1} \quad \text{Exothermic reaction (discharge)}$$  (8)

The project started in April 2009 and it has the budget of demonstrating the feasibility and economics of a solid oxide based TES system by:
• Identifying and select the multivalent oxide based TES systems that have the potential to meet the economical targets.
• Evaluating and demonstrating their feasibility experimentally.
• Determining the process economics based on the data obtained.

Achieving these targets, CSP will be competitive in the intermediate power market by 2015 and baseload market by 2020 [40].

Sixteen pure oxides were found to be able to undergo REDOX in air based on thermodynamics calculation with their respective equilibrium reaction temperatures and energy density storage are shown in figure 45.

![Figure 45. Equilibrium reaction temperature and potential energy storage density for various oxides in air [40].](image)

From these sixteen oxides Cr$_5$O$_{12}$, Li$_2$O$_2$, Mg$_2$O, PtO$_2$, UO$_3$ and Rh$_2$O$_3$ were eliminated based on transition temperatures and raw material costs. Then an experimental protocol which is detailed in [40] was established in order to test and characterize the candidates left.

It is important to mention that in the temperature range of interest (200-700 ºC) no large scale re-oxidation was observed in thermochemical candidates. Nevertheless in the range of 700 to 1100 ºC, Co$_2$O$_4$ was chosen as a feasible candidate as it demonstrated full re-oxidation. Heat charge and discharge in a TES packed bed was shown using the cobalt oxide and further studies will be performed in the next phases of the project.

Finally, re-oxidation kinetics improvement and long term stability of the low cost oxides via secondary oxide incorporation will be key to large scale deployment of this TES concept with CSP and will allow to make feasible other oxides which have been previously mentioned. More details about the project, the time horizon of the different phases and budgets can be found in [40].
2. DESIGN OF A LATENT HEAT STORAGE SYSTEM FOR A LABORATORY DEVICE

Based on the literature research presented in section 1.1. and in order to summarize the most important conclusions, a discussion of the different possible approaches for a high temperature latent heat storage will be discussed.

The latent heat storage in the charging process must store the energy coming from a thermal oil, the Dowtherm A, which is working between 322 and 312 ºC.

During the discharging process, the same thermal oil is used but going from 290 to 300 ºC. Then the suitable phase change material in this range of temperatures and based on the different examples already presented is the sodium nitrate (NaNO₃) as it has got a melting temperature of 306 ºC. It fits the temperature needed and the feasibility of using it as a phase change material for high temperature applications has already been proved.

As for the different possible designs, based mainly in the results from the DISTOR Project, three different approaches must be considered:

- Macro-encapsulation of the PCM
- Using a composite material compounded by the PCM and expanded graphite
- Sandwich concept

Regarding the macro-encapsulation of the PCM approach, cylindrical stiff capsules where chosen in the DISTOR Project due to manufacturating aspects as it has already been said in section 1.1.3.1.1. Nevertheless, the spherical macro-encapsulation must be keeped in mind as several models for low temperature applications of spherical solidification, melting and heat transfer have been performed. The concept itself is based on encapsulating the PCM inside the capsules which are sealed by welding and finally the capsules are integrated into a vessel as it is shown in figure 46.

Figure 46. Macro-encapsulation designing approach [49].
The advantage of this approach is the easy commercialization which can be achieved as some companies can be focused on the manufacturing of PCM capsules providing their features and giving a certain warranty and other engineering companies can just focus on the device or system design. Although for the low temperatures applications there is plenty of information about it, for the high-temperature applications there are some reasons which lead to the rejection of this option:

- The amount of hull material needed for pressure-tight capsules is significant; if steel is used, the mass of steel is almost equivalent to the mass of PCM.
- The volume fraction of PCM in the pressure vessel is less than 40%.
- The filling and sealing procedure with molten PCM is complex.
- A contamination of the steam by PCM due to leakage of capsules must be avoided; this demands high quality standards resulting in a further cost increase [51].

As for the use of composite material compounded by PCM and expanded graphite for LHS design, it must be mentioned that in the case of NaNO₃ the properties from the resulting composite are still under investigation. An article from the Doctor Chang-Ying Zhao (University of Warwick) was summarized in section 1.1.4.2.2. but the properties of the resulting composites were not shown. The investigation is still being performed as can be found in [50]: the research project which is called Thermal Conductivity Enhancement of High-Temperature Thermal Energy Stores for Use With Solar Power Plants was started on 01/10/2008 and is programmed to end on 30/09/2012. It is interesting to trace the project as the results may be useful for next designs.

Regarding the design concept, it has already been shown in section 1.1.3.1.1 but is interesting to plot it again. It is based on the manufacturing of the composite material by compression of a powder mixture of the salt and the expanded graphite as is the manufacturing method which has been chosen in previous studies as shown in figure 47: having the appropriate ducts inside in order to install the pipes where the HTF will flow in the charging and the discharging processes.

*Figure 47. Single segment made of PCM/composite material used for the laboratory-scale storage test unit. Holes are intended for steam pipes [22].*
The shape of the composite block, position of the ducts and dimensions can be varied depending on the working conditions in order to achieve the optimal design. It must be remembered that there is an analytical model with some assumptions presented in [25] which can be useful once the results from Dr. C.Y. Zhao are published. The conclusion is that this option must be rejected as the feasibility of the composite material approach could not be proven for nitrate salts so far [51].

Finally the **sandwich concept** approach which is the one chosen as the best solution in the DISTOR Project was already presented in sections 1.1.3.1.1 and 1.1.4.1. As the melting temperature of the sodium nitrate is 306 ºC, aluminum fins are the ones that must be used in this case.

The feasibility of this approach with high capacity factors, as mentioned in [27], has already been proved in different storage units ranging from 2 to 100 kW. Because of that, this option is the one chosen for the design of the latent heat storage system for the laboratory device which is shown in figure 48. It can be observed several valves, pumps and the heating source and heat since which can be chosen depending on the process required: charging and discharging processes respectively. The temperatures shown in figure 48 are the ones corresponding to the charging process.

![Figure 48. Laboratory device where the LHS will be installed.](image)

\[\text{Figure 48. Laboratory device where the LHS will be installed.}\]
A part from choosing the latent heat storage material and the configuration of the system, the designing parameters are:

- The latent heat storage power: 30 kW.
- The heat transfer fluid: Dowtherm A. The properties of the fluid which have been used for the calculations presented in the following section can be found in [56].
- The volumetric flow: 6 m³/h and a velocity of around 1 m/s.
- The temperatures in the charging process from the Dowtherm A: 322 ºC in the beginning and 312 ºC in the end.
- The temperatures in the discharging process from the Dowtherm A: 290 ºC in the beginning and 300 ºC in the end.
- The maximum dimensions: 2 m³
- Charging and discharging time of 2 hours: energy capacity of 60 kWh.

Once the design has been chosen (sandwich concept) and the main designing parameters are clearly listed, it is possible to model it in order to choose the geometrical parameters and observe the behavior of the system.

Due to the complexity of the problem, the model presented in the following section is a preliminary and approximate model which can be only seen as a way of having an initial idea of how the temperature profiles evolve and a trial to prove the feasibility between the design chosen and the designing parameters. Before building the system for the experimental device, further studies and modeling performances must be carried out, including software and simulation tools such as FLUENT, Mathcad and Matlab.

2.1. Modeling of the latent heat storage system

2.1.1. The Classical Stefan Problem

The calculation of a latent heat storage system is based on the Classical Stefan Problem. The complexity of the problem due to the unknown location of the interface between the solid and liquid results in the need of making several assumptions.

These assumptions are presented in the following table comparing them to the real situation:
### Table 7. Assumptions made in the Classical Stefan Problem [55].

Several models have been built around the Stefan problem which in some cases add some accuracy to the assumptions shown in table 7. For further details about these models, the reviews shown in [57] and [58] and the book mentioned in [59] are interesting.

In this Master Thesis due to time and resources limiting factors, the model performed is a simplified one and has got the target of using an analytical solution of the Stefan Problem: the quasi-static approximation. The effects of sensible heat are neglected and thanks to that the energy equation becomes independent of time and solutions are much easier to obtain.
For the case of interest, the equations which will be used are the ones corresponding to a Cartesian Coordinate geometry as the Stefan Problem is a non-linear problem and it is not possible to obtain the results from the steel cylinder and the aluminum fins and add them at the end. Because of that, the heat transfer from the steel tube is neglected and only the heat transfer from the aluminum fins considering Cartesian coordinates is taken into account.

The general equations governing the process for the quasi-static approximation are the following:

\[ X(t) = \left[2 \cdot \frac{\lambda_t}{\rho \cdot h_{st}} \cdot \int_0^t \left[T_0(t) - T_m\right] \cdot dt\right]^{1/2} \quad \text{for } t \geq 0 \]

*Equation 3. Position of the liquid-solid interface in the melting process of a semi-infinite solid [60]*

\[ T_l(x, t) = T_0(t) - \left[T_0(t) - T_f\right] \cdot \frac{x}{X(t)} \quad \text{in } 0 \leq x \leq X(t) \quad \text{for } t \geq 0 \]

*Equation 4. Temperature profile of the liquid region in the melting process of a semi-infinite solid [60]*

As it can be observed in equations 3 and 4, there is the need to obtain the temperature profile of the aluminum fins in the non-steady state. An analytical solution for a hollow cylinder with radial fins along the cylinder in the external surface does not exist and there is the need to use numerical methods to solve the problem. The finite volume model built for the purpose is presented in the following section.

**2.1.2. The finite volume model**

The finite volume model developed can be observed in figure 49 with the corresponding nodal numeration. It takes into account only one fin and has got 13 nodes, the last three of them are contour nodes and their temperature is already known:

- Node 11 is the phase-change material and it is at its melting temperature as the sensible heat is neglected in the quasi-static approximation and no supercooling is considered in the Classical Stefan Problem as mentioned before.
- Node 12 corresponds to the Dowtherm A and the heat transfer with the steel tube is by convection.
- Node 13 is the contour of the steel tube in its extremities from z direction and it is considered the same temperature in both of the sides (symmetry from the horizontal axis of the fin).
Before performing the numerical method calculation it is necessary to determine the convective coefficient and it is also a good way of checking the feasibility of the design. In order to facilitate the calculations, the convective coefficient is calculated neglecting the fins and for the steady state in an iterative process which is detailed in the annex A.

It is also a way of fixing the steel tubes dimensions which result taking into consideration the volumetric flow, velocity and the European Norm for Steel tubes. The properties of the steel are taken from [61] and regressions using Excel have been done in order to obtain their dependency with temperature.

It is important to mention the volume of PCM needed for the 30 kW device, which is easily calculated taking into account its latent heat of fusion (173 kJ/kg) and density (2261 kg/m$^3$), neglecting the sensible heat exchanged and considering a charging/discharging time of
2 hours. Then the resulting volume of PCM needed is 0.551 m$^3$ which has been placed uniformly as a hollow cylinder in this first calculation (operations shown in annex A if any doubts).

The main results are the need of having four different tubes with 6 steps with an internal diameter of 22.9 mm and a thickness of 2 mm and the mean convective coefficient of the Dowtherm A of 1715 W/(m$^2$·K). The results are considered for the charging process as it was found to be the most limiting process as the longitudes needed were larger than for the discharging process.

Finally it is important to mention that the iterative process shown in the annex A has been done manually until the difference between the new longitude and the old one is smaller than 0.001.

Once the convective coefficient has been found it is possible to proceed with the calculations of the finite volume model. For a general node the equations of the conductivities which correspond to figure 50 are the following:

\[ \Delta V_i = r_i \Delta \phi \Delta r \Delta z \]

\[ K_{i1} = \frac{\lambda \cdot (r_i + \frac{\Delta r}{2}) \cdot \Delta \phi \cdot \Delta z}{\Delta r} \]

\[ E\text{quation 5. Conductivity between the } i\text{-node and the next one in the positive radial direction [62].} \]

\[ K_{i3} = \frac{\lambda \cdot (r_i - \frac{\Delta r}{2}) \cdot \Delta \phi \cdot \Delta z}{\Delta r} \]

\[ E\text{quation 6. Conductivity between the } i\text{-node and the next one in the negative radial direction [62].} \]
\[ K_{i2} = \frac{\lambda \cdot \Delta r \cdot \Delta z}{r_i \cdot \Delta \varphi} = K_{i4} \]

*Equation 7. Conductivity between the i-node and the neighbor in the \( \varphi \) direction (both sides) [62].*

\[ K_{z+} = \frac{\lambda \cdot r_i \cdot \Delta \varphi \cdot \Delta r}{\Delta z} = K_{z-} \]

*Equation 8. Conductivity between the i-node and the neighbor in the z direction (either up and down) [62].*

It is important to mention that due to symmetry the conductivities in the \( \varphi \) direction are not taken into account, there is only heat transfer in two dimensions: \( z \) and \( r \). That leads to use for all the nodes an \( \Delta \varphi \) of \( 2\pi \) and also is important to remark that \( \Phi = \varphi \) in figure 50.

Also there is the need to choose the numerical method resolution: in this case the implicit method as the properties of the different materials are considered constant with temperature and in this method there is not any problems with stability and convergence in the solutions.

Once the conductivities for each node are calculated, it is possible to define the conductivity matrix (\( KT \)), the conductivity contour matrix (\( KC \)), the temperature contour nodes vector (\( TC \)), the CDT matrix which is needed as it is the unsteady state and it is possible to calculate the final temperatures (\( TF \)) after each time interval (2 s) supposing for the first time interval a first temperature vector of the nodes: in this case 579 K as it is the temperature of the steel tube from the calculations presented before (annex A).

The matrix solution is then:

\[ TF = (KT - CDT)^{-1} x(-KCxTC - CDTxTN) \]

*Equation 9. Matrix solution of the unsteady state for the implicit method [62].*

and the details about all the procedure, with the conductivities defined for each node and the matrixes mentioned before can be found in the annex B for the first time interval.

The steady state is reached after 6 seconds and the temperature profile in front of time of the nodes of the aluminum fins (8,9 and 10) are plotted in the following two figures as the results for node 9 and 10 are the same.
As for the contour conditions, the PCM has been considered at a constant temperature: the melting temperature. The results show that the temperature of the nodes of the fin is not enough high as the heat flux between the Dowtherm A and the aluminum fins is not enough.

Then in order to calculate the time needed for the melting of the PCM, after all the assumptions made, it is reasonable enough to consider a constant temperature five degrees above the melting temperature for the aluminum fin which should be the target of future designs.

Then the time it takes for the solid-liquid interface to get to the most extreme area: 2 mm far away from the fin is from equation 3 and considering \( T_0(t) = 311 \, ^\circ C = \text{constant} \):

\[
t = \frac{X(t)^2 \cdot \rho \cdot h_{st}}{2 \cdot k_i \cdot (T_0 - T_m)} = \frac{0.006 \cdot 2 \cdot 2261 \text{kg/m}^3 \cdot 173300 \text{J/kg}}{2 \cdot 0.5 \text{W/(m} \cdot \text{K)} \cdot 5 \text{K}} = 2821 \, s = 0.78 \, h
\]

Then, using equation 3, it is possible to have a bigger distance between fins than the one initially selected as it is possible to have a charging time of 2 hours:

\[
X(7200) = \left[ 2 \cdot \frac{0.5 \text{ W/(m} \cdot \text{K)}}{2261 \text{kg/m}^3 \cdot 173300 \text{J/kg}} \cdot 5 \text{K} \cdot 7200 \text{s} \right]^{0.5} = 0.009 \, m
\]

It must be mentioned that all the calculations presented in both annex A and annex B are based on several chapters from [62].
3. CONCLUSION

A review of the latent heat storage systems which can be found for high temperature applications has been performed. A great progress has already been achieved with several models and laboratory devices which have been built. Nevertheless there is still work to do in order to reach to the commercialization of LHS devices for high temperature.

There are still some challenges which must be overcome and specially the low thermal conductivity of the phase change materials which are normally used. In this direction, different approaches have been studied and the sandwich concept is the most successful at the moment as it has already been mentioned. Nevertheless, the macro-encapsulation of PCM, the use of composites and the use of metallic materials are interesting approaches which must be followed and studied further.

In addition, a review on thermo-chemical storage systems has been shown. Although it is a newer topic than the LHS systems, there is also a great work already developed specially with the dissociation of ammonia, the dehydration/hydration cycles and the reduction and oxidation reactions. As in the case of the LHS systems, there is a long way to reach the commercialization level but to have an idea of when it can be enough developed it is interesting to remark that the reduction and oxidation reactions project already mentioned has got the target of reaching the commercialized level between 2015 and 2020.

Finally, an initial design of a LHS system for a laboratory device has been presented. The model is limited as the phase-change is a non-linear problem which has a complex mathematical modeling. The sandwich concept was the design chosen and some first calculations were performed including a simple finite volume model in order to have a first idea of the behavior and the temperature profiles of the system. Software with adequate simulations tools for the problem must be used in the future to continue the modeling of the system and finish the detailing of the design specially because at the moment the heat flux between the heat transfer fluid and the aluminum fins is not enough. An option to overcome the situation could be the reconsideration of the designing parameters such as the input and output temperature of the heat transfer fluid (Dowtherm A).
ANNEXES

ANNEX A. Calculation of the convective coefficient and first dimensions

All units are in the international system (SI)

eK := 273.15
Td_I := 322 + eK
Td_F := 312 + eK
Tm := 306 + eK

\[
\begin{align*}
\Delta T &= T_F - T_I - T_m \\
VT &= \frac{60 \cdot 3600}{173.32261} \\
V_{PCM} &= 0.551
\end{align*}
\]

Twall := 579.169

Ts := \frac{T_m + Twall}{2} \\
T_s = 579.159

Properties of dowtherm A:

\[
\begin{align*}
\mu(T_{wall}) &= 5 \times 10^{-9} \cdot (T_{wall} - 273.15)^2 - 4 \times 10^{-6} \cdot (T_{wall} - 273.15)^2 + 0.001 \\
c_p(T_d) &= 3 \cdot 10^{-5} \cdot (T_d - 273.15)^3 - 0.0206 \cdot (T_d - 273.15)^2 + 8.0348 \cdot (T_d - 273.15) + 1082.7 \\
\lambda(T_d) &= 10^{-20} \cdot (T_d - 273.15)^2 - 0.0002 \cdot (T_d - 273.15) + 0.1419 \\
\rho(T_d) &= -0.0014 \cdot (T_d - 273.15)^2 - 0.2911 \cdot (T_d - 273.15) + 1020.3 \\
Pr(T_d) &= \frac{\mu(T_d) \cdot c_p(T_d)}{2 \lambda(T_d)}
\end{align*}
\]

Properties of steel

\[
\lambda_s(T_s) := 5.7168 \cdot (T_s - 273.15) - 71.685 \\
N_t := 4
\]

Q := \frac{6}{3600 \cdot N_t} \\
D_1 := 0.0229 \\
thick := 0.002

u := \frac{4 \cdot Q}{D_1^2 \cdot \pi} \\
u = 1.012

D_0 := D_1 + 2 \cdot \text{thick} \\
D_0 = 0.027

\[
\begin{align*}
S &= \frac{\pi \cdot D_i^2}{4} \\
A_{\text{firstapprox}} &= N_t \cdot \frac{\pi \cdot D_0^2}{4} \\
A_{\text{firstapprox}} &= 2.273 \times 10^{-3}
\end{align*}
\]

\[
\begin{align*}
P_t &= \pi \cdot D_i \\
P_h &= \pi \cdot D_i \\
D_h &= 4 \cdot \frac{S}{P_h} \\
\Delta T_I &= T_d_I - T_m \\
\Delta T_F &= T_d_F - T_m
\end{align*}
\]
\[ \text{MLDT} := \frac{\Delta T_I - \Delta T_F}{\ln \left( \frac{\Delta T_I}{\Delta T_F} \right)} \]

\[ G := \rho(T_d)u \]

\[ \text{Re} := \frac{GD_h}{\mu(T_d)} \]

\[ \text{Re} = 7.721 \times 10^4 \quad \text{Turbulent flow} \]

\[ q := Q \rho(T_d) \cdot c_p(T_d) \cdot (T_d_I - T_d_F) \]

\[ \phi := \frac{\mu(T_d)}{\mu(T_{wall})} \]

\[ n := 0.25 \]

\[ C_{fm} := \frac{1}{(1.58 \ln(\text{Re}) - 3.28)^2} \]

\[ L := 6.574 \quad C_{fm} = 4.755 \times 10^{-3} \]

\[ \text{Nu} := \frac{C_{fm}(\text{Re} - 1000) \cdot \text{Pr}(T_d) \cdot \left[ 1 + \left( \frac{D_h}{L} \right)^{\frac{2}{3}} \right]^2}{2 + 17.96 C_{fm}^{0.5} \left[ \left( \frac{2}{\text{Pr}(T_d)} \right)^{\frac{2}{3}} - 1 \right]} \]

\[ h_d := \frac{\text{Nu} \cdot \lambda(T_d)}{D_h} \quad h_d = 1.715 \times 10^3 \]

\[ A_s := \pi \cdot D_1 \cdot L \]

\[ R_s := \frac{1}{h_d \cdot A_s} \quad R_{ss} := \frac{\ln \left( \frac{D_o}{D_1} \right)}{2 \pi \cdot \lambda_s(T_s) \cdot L} \]

\[ U_s := \frac{1}{A_s \left( R_s + R_{ss} \right)} \]

\[ L_{new} := \frac{q}{U_s \cdot \pi \cdot D_1 \cdot \text{MLDT}} \quad L_{new} = 6.574 \]

\[ T_{wall_{new}} := q \cdot R_{ss} + T_m \quad T_{wall_{new}} = 579.169 \]

\[ \text{Error} := L_{new} - L \quad \text{Error} = -1.542 \times 10^{-4} \]

\[ V_{tubes} := \pi \cdot D_0 \cdot L_{new} \quad V_{tubes} = 0.556 \]

\[ V_{total} := V_{tubes} + V_{PCM} \quad n_p := 6 \quad \text{L}_{\text{step}} := \frac{L}{n_p} \]

\[ V_{total} = 1.107 \]
\[ V_{\text{PCM1step}} := \frac{V_{\text{PCM}}}{N_t \cdot n_p} \]

\[ D_{\text{out}} := \sqrt{\frac{4 \cdot L_{\text{step}}}{\pi} \cdot V_{\text{PCM1step}} + D_o^2} \]

\[ A_{\text{transversal}} := N_t \cdot n_p \cdot D_{\text{out}}^2 \]

\[ V_{\text{total2}} := A_{\text{transversal}} \cdot L_{\text{step}} = 0.862 \]
ANNEX B. Numerical model calculation

eK := 273.15  \quad h := 1.715 \times 10^3

Properties of steel

\[ T_s := 579 \]
\[ \lambda_s(T_s) := 5.7168 \left(T_s - 273.15\right) - 71.685 \]
\[ \rho_s := 8000 \]
\[ c_s := 500 \quad \text{Found in the working temperature range of 273 to 373 K} \]

Properties of aluminum

At 600 K as it does not change too much in the temperature working range

\[ \rho_a := 2700 \]
\[ T_{13} := 312 + eK \]

Geometrical parameters:

\[ D_i := 0.0229 \quad \text{thick} := 0.002 \quad D_o := D_i + 2\text{thick} \quad D_{out} := 0.181 \]
\[ \Delta \varphi := 2 \pi \quad \text{fin}\text{thick} := 0.010 \quad n := 0.002 \]

Node 1

\[ \Delta z_1 := \text{finthick} - 2 \cdot n \quad \Delta r_1 := \frac{\text{thick}}{2} \quad V_1 := \pi \cdot \left[ \left(D_i + \Delta r_1\right)^2 - D_i^2 \right] \cdot \Delta z_1 \]

\[ C_1 := \rho_s \cdot V_1 \cdot c_s \]

\[ K_{1.12} := h \cdot \Delta \varphi \cdot \frac{D_i}{2} \cdot \Delta z_1 \]
\[ K_{1.2} := \frac{\lambda_s(T_s) \left(D_i \right) \Delta \varphi}{2} \cdot \frac{\Delta r_1}{\Delta z_1} \]

\[ K_{1.4} := \frac{\lambda_s(T_s) \left(D_i \right) \Delta \varphi}{2} \cdot \frac{\Delta r_1 \cdot \Delta z_1}{\Delta z_1} \]

Node 2

\[ V_2 := V_1 \quad C_2 := C_1 \quad K_{2.12} := K_{1.12} \quad K_{2.1} := K_{1.2} \quad K_{2.3} := K_{1.2} \quad K_{2.5} := K_{1.4} \]

Node 3

\[ V_3 := V_1 \quad C_3 := C_1 \quad K_{3.12} := K_{1.12} \quad K_{3.2} := K_{2.3} \quad K_{3.6} := K_{1.4} \quad K_{3.13} := K_{1.13} \]
Node 4

\[ V_4 := \pi \cdot \left( \frac{(D_1 + 3\Delta r_1)^2 - (D_1 + \Delta r_1)^2}{4} \right) \Delta z_1 \quad C_4 := \rho_s \cdot V_4 \cdot c_s \]

\[ K_{4,1} := K_{1,4} \quad K_{4,5} := \frac{\lambda_s(T_s) \left( \frac{D_1}{2} + \Delta r_1 \right) \Delta \varphi \cdot \Delta r_1}{\Delta z_1} \quad K_{4,13} := \frac{\lambda_s(T_s) \left( \frac{D_1}{2} + \Delta r_1 \right) \Delta \varphi \cdot \Delta z_1}{\Delta z_1} \]

\[ K_{4,11} := \frac{\lambda_s(T_s) \left( \frac{D_1}{2} + \Delta r_1 + \frac{\Delta r_1}{2} \right) \Delta \varphi \cdot \Delta z_1}{\Delta r_1} \]

Node 5

\[ V_5 := V_4 \quad C_5 := C_4 \]

\[ K_{5,2} := K_{2,5} \quad K_{5,6} := K_{4,5} \quad K_{5,7} := \frac{\lambda_s(T_s) \left( \frac{D_1}{2} + \Delta r_1 + \frac{3\Delta r_1}{2} \right) \Delta \varphi \cdot \Delta z_1}{\Delta z_1} \]

\[ K_{5,4} := K_{4,5} \]

Node 6

\[ V_6 := V_4 \quad C_6 := C_4 \]

\[ K_{6,3} := K_{3,6} \quad K_{6,5} := K_{5,6} \quad K_{6,11} := K_{4,11} \quad K_{6,13} := K_{4,13} \]

Node 7

\[ V_7 := \pi \cdot \left( \frac{(D_1 + 4\Delta r_1)^2 - (D_1 + 3\Delta r_1)^2}{4} \right) \Delta z_1 \quad C_7 := \rho_s \cdot V_7 \cdot c_s \]

\[ K_{7,5} := K_{5,7} \quad K_{7,11} := 2 \frac{\lambda_s(T_s) \left( \frac{D_1}{2} + \Delta r_1 + \frac{\Delta r_1}{2} + \frac{\Delta r_1}{4} \right) \Delta \varphi \cdot \Delta z_1}{\Delta z_1} \]

\[ R_{m7} := \frac{\Delta r_1}{\lambda_s(T_s) \left( \frac{D_1}{2} + \Delta r_1 + \frac{\Delta r_1}{2} + \frac{\Delta r_1}{4} \right) \Delta \varphi \cdot \Delta z_1} \]

\[ R_{m8} := \frac{\Delta r_2}{\lambda_a \left( \frac{D_i}{2} + \Delta r_1 + \frac{\Delta r_1}{2} + \frac{\Delta r_1}{4} \right) \Delta \varphi \cdot \Delta z_1} \]

\[ \Delta r_2 := \frac{D_{out} - D_o}{2} \]

\[ K_{7,8} := \frac{1}{R_{m7} + R_{m8}} \]
Nodes from the aluminum fin

Node 8

\[ V_8 := \pi \cdot \frac{\left( D_0 + 4\Delta r_2 \right)^2 - \left( D_0 \right)^2}{4} \cdot \Delta z_1 \quad C_8 := \rho_a \cdot V_8 \cdot c_a \]

\[ K_{8.7} := K_{7.8} \quad K_{8.11} := 2 \cdot \frac{\lambda_a \left( \frac{D_{\text{out}}}{2} - 6\Delta r_2 \right) \cdot \Delta \varphi \cdot 2\Delta r_2}{\Delta z_1} \]

\[ K_{8.9} := \frac{\lambda_a \left( \frac{D_0}{2} + 2\Delta r_2 + \Delta r_2 \right) \cdot \Delta \varphi \cdot \Delta z_1}{2 \cdot \Delta r_2} \]

Node 9

\[ V_9 := \pi \cdot \frac{\left( D_0 + 8\Delta r_2 \right)^2 - \left( D_0 + 4\Delta r_2 \right)^2}{4} \cdot \Delta z_1 \quad C_9 := \rho_a \cdot V_9 \cdot c_a \]

\[ K_{9.8} := K_{8.9} \quad K_{9.11} := 2 \cdot \frac{\lambda_a \left( \frac{D_0}{2} + 3\Delta r_2 \right) \cdot \Delta \varphi \cdot 2\Delta r_2}{\Delta z_1} \]

\[ K_{9.10} := \frac{\lambda_a \left( \frac{D_0}{2} + 3\Delta r_2 + \Delta r_2 \right) \cdot \Delta \varphi \cdot \Delta z_1}{2 \cdot \Delta r_2} \]

Node 10

\[ V_{10} := \pi \cdot \frac{\left( D_0 + 12\Delta r_2 \right)^2 - \left( D_0 + 8\Delta r_2 \right)^2}{4} \cdot \Delta z_1 \quad C_{10} := \rho_a \cdot V_{10} \cdot c_a \]

\[ K_{10.9} := K_{9.10} \]

\[ K_{10.11} := 2 \cdot \frac{\lambda_a \left( \frac{D_0}{2} + 5\Delta r_2 \right) \cdot \Delta \varphi \cdot 2\Delta r_2}{\Delta z_1} + \frac{\lambda_a \left( \frac{D_0}{2} + 5\Delta r_2 + \Delta r_2 \right) \cdot \Delta \varphi \cdot \Delta z_1}{2 \cdot \Delta r_2} \]
\[
\begin{pmatrix}
-K_{1.2} - K_{1.4} - K_{1.12} - K_{1.13} & K_{1.2} & 0 & K_{1.4} & 0 & 0 & 0 & 0 & 0 & 0 \\
K_{2.1} & -K_{2.1} - K_{2.3} - K_{2.5} - K_{2.12} & K_{2.3} & 0 & K_{2.5} & 0 & 0 & 0 & 0 & 0 \\
0 & K_{3.2} & -K_{3.2} - K_{3.16} - K_{3.12} - K_{3.13} & 0 & 0 & K_{3.6} & 0 & 0 & 0 & 0 \\
K_{4.1} & 0 & 0 & -K_{4.1} - K_{4.5} - K_{4.11} - K_{4.13} & K_{4.5} & 0 & 0 & 0 & 0 & 0 \\
0 & K_{5.2} & 0 & K_{5.4} & -K_{5.2} - K_{5.4} - K_{5.6} - K_{5.7} & K_{5.6} & K_{5.7} & 0 & 0 & 0 \\
0 & 0 & K_{6.3} & 0 & K_{6.5} & -K_{6.3} - K_{6.5} - K_{6.11} - K_{6.13} & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & K_{7.5} & 0 & -K_{7.5} - K_{7.8} - K_{7.11} & K_{7.8} & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & -K_{8.7} - K_{8.9} - K_{8.11} & K_{8.9} & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & -K_{9.8} - K_{9.10} - K_{9.11} & K_{9.10} & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & -K_{10.9} - K_{10.11} & K_{10.9} & 0 & 0 \\
\end{pmatrix} = KT
\]
\[ KC := \begin{bmatrix}
0 & K_{1.12} & K_{1.13} \\
0 & K_{2.12} & 0 \\
0 & K_{3.12} & K_{3.13} \\
K_{4.11} & 0 & K_{4.13} \\
0 & 0 & 0 \\
K_{6.11} & 0 & K_{6.13} \\
K_{7.11} & 0 & 0 \\
K_{8.11} & 0 & 0 \\
K_{9.11} & 0 & 0 \\
K_{10.11} & 0 & 0 
\end{bmatrix} \]

\[ \Delta t := 2 \]

\[ CDT := \begin{bmatrix}
\frac{C_1}{\Delta t} & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & \frac{C_2}{\Delta t} & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & \frac{C_3}{\Delta t} & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & \frac{C_4}{\Delta t} & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & \frac{C_5}{\Delta t} & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & \frac{C_6}{\Delta t} & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & \frac{C_7}{\Delta t} & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & \frac{C_8}{\Delta t} & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & \frac{C_9}{\Delta t} & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & \frac{C_{10}}{\Delta t} 
\end{bmatrix} \]
\[ \begin{pmatrix} T_1 \\ T_2 \\ T_3 \end{pmatrix} \]

\[ T_{11} := 306 + eK \]
\[ T_{12} := 322 + eK \]
\[ \mathbf{T} := \begin{pmatrix} T_{11} \\ T_{12} \\ T_{13} \end{pmatrix} \]

\[ \mathbf{T} := (K - CD) \cdot (K - TC - CD \cdot TN) \]

\[
\begin{array}{c|c}
\hline
0 & 579.479 \\
1 & 579.459 \\
2 & 579.479 \\
3 & 579.389 \\
4 & 579.443 \\
5 & 579.389 \\
6 & 579.434 \\
7 & 579.157 \\
8 & 579.147 \\
9 & 579.147 \\
\hline
\end{array}
\]

\[
\begin{array}{c|c}
\hline
0 & 306.329 \\
eK1 & 306.309 \\
eK2 & 306.329 \\
eK3 & 306.239 \\
eK4 & 306.293 \\
eK5 & 306.239 \\
eK6 & 306.284 \\
eK7 & 306.007 \\
eK8 & 305.997 \\
eK9 & 305.997 \\
\hline
\end{array}
\]
Steady state:

\[
T_{\text{steadystate}} := KT^{-1}(-KC+TC)
\]

\[
\begin{array}{c|c}
0 & 579.48 \\
1 & 579.461 \\
2 & 579.48 \\
3 & 579.389 \\
4 & 579.445 \\
5 & 579.389 \\
6 & 579.437 \\
7 & 579.16 \\
8 & 579.15 \\
9 & 579.15 \\
\end{array}
\]

\[
\begin{array}{c|c}
0 & 306.33 \\
eK & 306.311 \\
eK & 306.33 \\
eK & 306.239 \\
eK & 306.295 \\
eK & 306.239 \\
eK & 306.287 \\
eK & 306.01 \\
eK & 306 \\
eK & 306 \\
\end{array}
\]
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