

Study and development of a solidification model using CFD

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MASTER THESIS

Study and development of a solidification model using CFD

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Declaration of Authorship

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Abstract

Faculty Name Escola Tècnica Superior d'Enginyeria de Camins, Canals i Ports de Barcelona

Master Thesis

Study and development of a solidification model using CFD

by Aitor Bazán Escoda

Phase change materials (PCMs) are of great interest within the automotive industry field. Not only when used in thermal management applications but also in different areas where these materials are of vital importance for both a safe and comfortable driving. For such objective, the present project arises from the idea of understanding solidification processes in windshield washer tanks. In this context, this master's thesis produces a comprehensive state of the art of some of the current numerical methods to effectively represent water solidification.

An OpenFOAM 21.12. solver based on a multi-phase solver, multi-component incompressible solver based on a volume of fluid method is adapted to deal with diffusive-convective phase change. So as to reach this goal, an implementation of the enthalpy-porosity techinque is carried out. The work of Voller is closely followed, and a detailed explanation of the used equations and the assumptions taken is given. Validation of the model is accomplished by comparing the results with the authors in Bourdillon and Kowaleski and Rebow.

On a second stage, a 2D semi-empirical model based on the work of Lee is adapted to account for the nucleation characteristics during the process of the water phase change. Validation of the model is done by comparing the obtained results to Neumann solutions for classical Stefan problem.

Finally, the current work is extended to couple a fluid region in which the liquid undergoes a phase-change and a solid region. This is done in the context of a conjugate heat transfer environment.

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Contents

De	eclara	tion of	Authorship	iii
Al	ostrac	et .		v
A	cknov	vledger	nents	vii
1	Intr	oductio	n	1
	1.1	Thesis	Statement. Background and motivation	1
	1.2	Phase-	Change Process	1
		1.2.1	Phase diagram of ice	1
		1.2.2	Properties of ice	
		1.2.3	Freezing phenomena	2
	1.3	Mecha	nisms of Heat Transfer. Heat convection	2
		1.3.1	Stefan Problem	3
	1.4	Conjus	gate Heat Transfer. Heat conduction	3
		1.4.1	Governing Equations	3
			1.4.1.1 Governing Equations for the Fluid	3
			1.4.1.2 Governing Equations for the Solid	
2	Nur	nerical i	Methods for Phase-Change Phenomena	5
_	2.1		of Art. Numerical Methods	5
	2.2		ication methods	5
	2.2		Volume-of-Fluid Method: General Aspects	6
			Enthalpy-Porosity Model. Governing Equations	6
			Lee model	7
		2.2.0	2.2.3.1 Momentum Equation	
			*	
			07 1	8
		224	2.2.3.3 Classical nucleation theory. The coefficient C_f Let a property and also	
		2.2.4	Interphase porosity models	11
			2.2.4.1 Surface tension model	12
3			Simulation of Solidification Process	13
	3.1	-	FOAM. General Aspects	
		3.1.1	Boundary Conditions Directory	
		3.1.2	Constant Properties Directory	14
		3.1.3	System Directory	14
	3.2		ication process. Methodology	15
	3.3	OpenF	FOAM: BuoyantBoussinesqPimpleFOAM. Natural Convection	
		solver		16
		3.3.1	Control Loop	16
		3.3.2	Governing Equations	17
			3.3.2.1 Momentum Equation	17
			3.3.2.2 Temperature Equation	17

	hlioa	raphy		31
A	Freq A.1	uently How c	Asked Questions do I change the colors of links?	29 29
6		re Wor		27
5		clusion		25
_	Com	alucio		25
		4.2.5	Validation of Results and Conclusions	23
		4.2.4	Case Setup	23
		1.4.0	4.2.3.1 Energy Equation	23
		4.2.3	Governing Equations of the Solid Region	23
			4.2.2.3 Energy Equation	23
			4.2.2.1 Momentum Equation	23 23
		4.2.2	Governing Equations of the Fluid Region	23
		4.2.1		23
	4.2		FOAM: chtMultiphaseInterFOAM. Conjugate Heat Transfer	23
	4.1		dology	23
4	Nun		Simulation of Heat Transfer	23
		3.0.1		
		3.5.4	Validation of Results and Conclusions	22
		3.5.2	Case Setup	20
		3.5.2	Hypotheses And Assumptions	20
	3.5	3.5.1	Description	20
	2 5	Casa	3.4.4.3 Energy Equation	20 20
			3.4.4.2 Pressure Equation	20
			3.4.4.1 Momentum Equation	20
		3.4.4	Governing Equations	20
		3.4.3	Mass transfer models	20
		3.4.2	Phase models	20
		3.4.1	Control Loop	20
		cess .		20
	3.4	OpenF	FOAM: IcoReactingMultiphaseInterFOAM. Phase-Change Pro-	
		3.3.7	Validation of Results and Conclusions	19
		3.3.6	Case Setup	18
		3.3.5	Code implementations	18
		3.3.4	Hypotheses And Assumptions	18
		3.3.3	Case Description	17

List of Figures

1.1	Process of crystallization of water.	3
2.1	Crystallization rate versus temperature	11
3.1	General structure of an OpenFOAM case	14
3.2	Flowchart of integration procedure. buoyantBoussinesqPimpleFoam	16

List of Tables

3.1	Boundary conditions for natural convection case	18
3.2	Water properties for natural convection	18
3.3	Discretization schemes	18
3.4	Solvers for the discretised equations	19
3.5	Parameters for the discretised equations	19
3.6	Boundary conditions for natural convection case	21
3.7	Boundary conditions for natural convection case	21
3.8	Water properties for natural convection	21
3.9	Discretization schemes	22
3.10	Solvers for the discretised equations	22
3.11	Parameters for the discretised equations.	22

List of Abbreviations

LAH List Abbreviations HereWSF What (it) Stands For

Physical Constants

Speed of Light $c_0 = 2.99792458 \times 10^8 \,\mathrm{m \, s^{-1}}$ (exact)

xix

List of Symbols

a distance r

P power $W(J s^{-1})$

 ω angular frequency rad

Chapter 1

Introduction

1.1 Thesis Statement. Background and motivation

During the last decade, the use of phase change materials has been growing in the automotive industry.

These substances release or absorb large amounts of latent heat when they go through a change in their physical state, as the material reaches its specific phase change temperature. Thus, in the process of latent heat release or absorption, the temperature of the PCM remains constant. Therefore, PCMs are considered to be efficient in terms of thermal storage.

On the other hand, the use of these materials is not exclusive for thermal management. It is of relevant importance when used in conjunction with soap in the windshield wiper system of the car. In such zone, problems involving solidification are of considerable relevance. And this is mainly due to a volumetric expansion originated by the thermal effects within the PCM which at its turn, generates stresses in the polymeric tank in which is bottled up.

Therefore, this master's thesis main objective aims to study different numerical techniques to represent solidification process and, specially, pure water phase change. This is accomplished by first imlementing an enthalpy-porosity technique within the frame of a multi-phase incompressible solver based on volume-of-fluid (VOF) method for interface tracking. The objective of this first stage is to apply sensible and latent heat as source terms in the energy equation. On a second term, a 2D semi-empirical model based on the work of Lee is adapted to account for the nucleation characteristics occured during the water phase transition. The final stage of this thesis is devoted to an implementation of a multiregion solver to calculate conjugate heat transfer problems between solid and fluid zones with the singularity of being, the fluid zone, capable of handling phase change materials.

1.2 Phase-Change Process

A complex interaction of the molecular forces generate water to behave in a curious way when it gets frozen into ice. The vast majority of substances, when they are cooled, become more dense in the frozen state than when liquid. However, when cooled under a specific temperature, water begins to expand and, once it starts freezing, it becomes less dense than water.

1.2.1 Phase diagram of ice

Ice crystals undergo different kinds of structures. Called ice Ih, in the form of hexagonal ice and, manifested in six cornered snow flakes, is the natural ice found

in earth. However, at lower pressures below 2 kbar, many other ice structures may exist.

The ice phase diagram shown below, points out the conditions of stability for all ice phases. As it is cleared out, the line between the water and ice Ih is an equilibrium line with a negative slope, consequence of having, the solid, lower density than the liquid. These equilibrium lines extend in the form of metastable phase boundaries into the area of stability of other ice phases. Although there are at least 11 crystalline ice shapes, the only which is found in naturally on earth is the hexagonal form. As a remark, the implication that there is a rise on the pressure would not propitiate ice formation at 0°C, instead water would need to be cooled down.

1.2.2 Properties of ice

Ice, when subjected to visible light conditions, is transparent and has the lowest index of refraction for the sodium spectrum of any known crystalline material[]. Mechanically, ice behaves like a viscoelastic material with a non linear law. Pollycrystalline ice subjected to stress, deforms elastically, followed by a transient creep and finally, a secondary creep in the form of steady viscous flow is obtained. As pointed out in [], the surface of ice Ih near the melting point has many dangling broken bounds that boost the presence of a liquid-like layer and as a consequence, low friction on such surface. Variation of density of ice with phase at 110 K and some physical properties of ice Ih at 0°C are described in the tables below.

1.2.3 Freezing phenomena

Time-temperature diagram for freezing of pure water (ABCDE) and aqueous solutions (AB'C'D'E') show the physical process that occurs during the solidification. The first stage, from A to B, belongs to undercooling, also called supercooling, and it is arisen below the freezing point T_f , which is equal to the melting point, T_m . This point is referred to a non equilibrium point and it is analogous to an activation energy necessary for the nucleation process. Before nucleation process, pure water may need to be cooled down several degrees. At point B, the system nucleates and releases its latent heat faster than the heat which is being removed from itself. From C to D, the horizontal axis shows the evolution of the crystal growth in time. At C, there exists the nucleation point and, from there through D, latent heat gets removed out of the system at constant temperature. In this way, the mixture, which is in a partially frozen state, does not cool until all the potentially freezable water has crystallized.

1.3 Mechanisms of Heat Transfer. Heat convection

The process of water freezing in enclosures is common in engineering. When there exist temperature gradients within the liquid phase in the process of solidification, a natural buoyancy driven flow is initiated and such behavior is determined to affect the shape of the liquid/solid interface as well as the progress of solidification. Indeed, these temperature differences in the liquid cause density variations so that the natural motion occurs. Boussinesq approximation can be validly used for fluids whose density varies linearly with temperature. However, pure water exhibits a maximum in its density when it ranges between 0°C and 4°C. Beyond this last

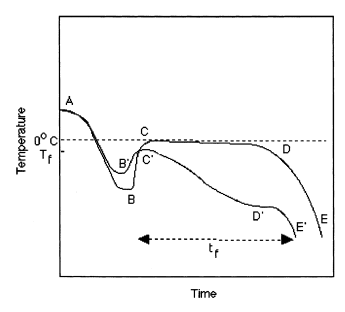


FIGURE 1.1: Process of crystallization of water.

temperature, and known as density inversion point, density decreases in a nonlinear manner as the temperature passes through the freezing point. In convective heat transfer, surroundings of the temperature where the aforementioned maximum happens to be, behave in a complex manner leading to fully control the process of growth of the solid phase.

1.3.1 Stefan Problem

1.4 Conjugate Heat Transfer. Heat conduction

1.4.1 Governing Equations

1.4.1.1 Governing Equations for the Fluid

1.4.1.2 Governing Equations for the Solid

$$\frac{\partial}{\partial x_i} \left(\rho u_i T \right) = \frac{\partial}{\partial x_i} \left(\frac{k}{C_p} \frac{\partial u_j}{\partial x_j} \right) \tag{1.1}$$

Chapter 2

Numerical Methods for Phase-Change Phenomena

2.1 State of Art. Numerical Methods

Considering the PCM density as constant in the model might be thought as a reasonable assumption in some cases, in others where thermo-mechanical coupling between the fluid and its container is intended, it makes impossible to account for some physical behaviors which may result from expansion or contraction during the phase change of the material. However, the main goal of this thesis is not to present a method that represents thermo-mechanical coupling but a technique that ensures volume expansion due to density changes through the fluid domain. To reach this point, it is important to point some of the numerous researches that have been conducted in order to investigate the problem of solidification and melting. At the present, the main numerical methods representing the treatment of liquid-solid phase change are divided into these categories:

- Surface tracking methods,
- Volume tracking methods,
- Moving mesh methods,

Surface tracking methods Volume tracking methods Moving mesh methods

2.2 Solidification methods

The challenge of a numerical investigation of a solidification process is to capture the free surface for the flow of the phase change material and, at the same time, account for the moving boundary induced by the phase change within the PCM. The free surface may be handled by the volume-of-fluid (VOF), originally introduced by [ref Hirt]. VOF relies on the definition of a transport indicator function within the finite volume method's framework. Simultaneously, and in order to account for the phase changes, some of the used models are based on meso-scale. This is the phenomena occurring between microscopic and continuum length scales and, in the current context, the complex micro structure generated during the solidification is approximated as liquid, mushy (intermediate state), and solid regions. Mushy region is thereby described as an averaged value of the liquid and solid properties.

One of the most used methods is the enthalpy-porosity technique, originally developed by Voller and Prakash [ref. Voller], which uses the typical conservation equations on a fixed Eulerian grid. The main concepts underlaying such method are: on the one side, an additional source term to the energy conservation equation is applied to describe the release of latent heat. On the other side, the solidification effects on the mass transport are modelled as a porosity variable and this is introduced as a Darcy-type source term to the momentum equation.

Some of the studies found on this topic, the coupling of both VOF and enthalpy-porosity methods are mainly related to casting processes. Rösler and Brüggermann [ref] introduced a numerical model for a solid-liquid phase change inside a latent heat thermal energy storage. Richter et al. [ref] worked out a method for the simultaneous moult filling and solidification process which settles the developing of free surface flow and the liquid-solid phase transition under the volume-of-fluid and enthalpy-porosity methods. However, no adaptation of these methods to purely solidification processes has been found. Therefore, and, the objectives of this research are:

- To introduce a new solver based on the coupling of VOF and enthalpy-porosity techniques which covers the relevant physical effects during the process of solidification.
- To validate simulation results by using benchmark cases found in the literature.

Numerical methods commented are briefly described next.

On the other hand, semi-empirical methods ...

2.2.1 Volume-of-Fluid Method: General Aspects

The volume of fluid method takes relevance when fluids coexist with other phases. An example could be the ice (solid phase) advancing front within the liquid phase. The surface in between both phases needs to be solved by means of the volume of fluid technique.

This is sometimes seen as the conservation of the mixture components along the path of a fluid region. The equation which allows that is described as:

$$\frac{\partial \gamma_{\text{phase}}}{\partial t} + \frac{\partial \left(\gamma_{\text{phase}} u_j\right)}{\partial x_j} = 0 \tag{2.1}$$

In which α_{phase} corresponds to the phase fraction and it applies

$$\gamma_{phase} = \begin{cases} 0 & = \text{ solid PCM} \\ 0 < \gamma_{phase} < 1 & = \text{ cell contains the interface} \\ 1 & = \text{ liquid PCM} \end{cases}$$
 (2.2)

2.2.2 Enthalpy-Porosity Model. Governing Equations

The energy equation based on the enthalpy formulation for convective-diffusive heat transfer states that,

$$\frac{\partial \rho h}{\partial t} + \frac{\partial}{\partial x_j} \left(\rho u_j h \right) = \frac{\partial}{\partial x_j} \left(\lambda \frac{\partial}{\partial x_j} T \right) \tag{2.3}$$

where u is the velocity component and λ is the thermal conductivity of the fluid. h can be expressed as a function of its latent heat and the specific sensible parts, However, the enthalpy-porosity method describes the enthalpy h of the mixture by its sensible part and the latent heat of solidification. The release of the latent heat is dependent on the stage of the phase change, and must be restricted to the phase change material.

$$h = \int_{T_r}^T c_p dT + \alpha_\ell L \tag{2.4}$$

where the latent heat is is driven by the evolution of the liquid α_l . The phase transition is modelled by expressing the liquid volume fraction as a function of the temperature,

$$\alpha_{l} = \begin{cases} 1 & T > T_{liq} \\ \frac{T - T_{sol}}{T_{liq} - T_{sol} + \varepsilon} & T_{sol} < T < T_{liq} \\ 0 & T < T_{sol} \end{cases}$$
(2.5)

For seek of brevity on the following expressions, it is adapted the term γ_{phase} to γ_l If expression 2.4 is replaced in 2.3,

$$\frac{\partial \left(\rho c_p T + \gamma_l \alpha_l L\right)}{\partial t} + \frac{\partial \left(\rho u_j c_p T + u_j \gamma_l \alpha_l L\right)}{\partial x_j}
= \frac{\partial}{\partial x_j} \left(\lambda \frac{\partial T}{\partial x_j}\right)$$
(2.6)

Rearranging terms, it yields,

$$\frac{\partial(\rho C_p T)}{\partial t} + \frac{\partial(u_j \rho C_p T)}{\partial x_j} + L \left[\frac{\partial(\rho \alpha_l \gamma_l)}{\partial t} + \frac{\partial(u_j \rho \alpha_l \gamma_l)}{\partial x_j} \right] = \frac{\partial}{\partial x_j} \left(\lambda \frac{\partial T}{\partial x_j} \right)
S = -L \left[\frac{\partial(\rho \alpha_l \gamma_l)}{\partial t} + \frac{\partial(u_j \rho \alpha_l \gamma_l)}{\partial x_j} \right]$$
(2.7)

The momentum equation is discussed in detail in the sub-chapter *Interphase porosity models*.

2.2.3 Lee model

The Lee model is based in the liquid-vapour mass transfer. Governed by the vapour transport equation 2.8, this model is applicable during melting or solidification of a fluid.

$$\frac{\partial}{\partial t} \left(\alpha_i \rho_i \right) + \nabla \left(\alpha_i \rho_i u_i \right) = S_{m_i} \tag{2.8}$$

 ρ_i and u_i are the fluid density and fluid velocity of the *ith* phase. Moreover, S_{m_i} is the mass source which takes on a zero value at the interface. During melting, $T_l > T_{sat}$,

$$\frac{dm_{sl}}{dt} = C_f \rho_s \alpha_s \left(\frac{T_s - T_{sat}}{T_{sat}} \right) \tag{2.9}$$

During solidification, $T_l < T_s at$

$$\frac{dm_{ls}}{dt} = C_f \rho_l \alpha_l \left(\frac{T_{sat} - T_l}{T_{sat}} \right) \tag{2.10}$$

The coefficient C_f might be interpreted as a time rate and must be empirically tunned. Its magnitude is expressed in $\frac{1}{s}$. α represents the phase volume fraction. $\frac{dm_i}{dt}$ are the mass transfer rates from one phase to another. The subscripts "s", "l", refer to solid and liquid phases respectively. T_{sat} , is the phase transition temperature which, in case of pure water would be 273.15 K. The source term of 2.8 is then calculated as,

$$S_{m_i} = \begin{cases} \frac{dm_{sl}}{dt} - \frac{dm_{ls}}{dt}, & \text{for water phase} \\ \frac{dm_{ls}}{dt} - \frac{dm_{sl}}{dt}, & \text{for ice phase} \end{cases}$$
(2.11)

2.2.3.1 Momentum Equation

In the momentum equation, the flow is modelled as,

$$\frac{\partial (\rho u_i)}{\partial t} + \frac{\partial (\rho u_i u_j)}{\partial x_j}
= -\alpha_i \nabla p + \frac{\partial}{\partial x_i} \left(\mu \frac{\partial u_i}{\partial x_j} \right) + F_{\sigma i} + S_{u_i}$$
(2.12)

The source term for the momentum equation can be written as,

$$S_{u_i} = \begin{cases} \frac{dm_{sl}}{dt} u_l - \frac{dm_{ls}}{dt} u_s, & \text{for water phase} \\ \frac{dm_{ls}}{dt} u_s - \frac{dm_{sl}}{dt} u_l, & \text{for ice phase} \end{cases}$$
 (2.13)

where u_l and u_s are the liquid and solid velocity components accordingly. The source terms related to interphase porosity (??) may be added to the momentum equation presented here for the Lee model 2.13.

2.2.3.2 Energy Equation

The energy equation for the Lee model can be described as,

$$\frac{\partial(\rho C_p T)}{\partial t} + \nabla \cdot \left(u_j \rho C_p T\right) = \nabla \cdot (k_i \nabla T_i) + S_{H_i} \tag{2.14}$$

where the heat source term due to mass transfer in the energy equation is calculated as,

$$S_{h_i} = \begin{cases} \frac{dm_{sl}}{dt} H_L, & \text{for water phase} \\ \frac{dm_{ls}}{dt} H_L & \text{for ice phase} \end{cases}$$
 (2.15)

where H_l is the latent heat induced by the phase transition and k_i , the thermal conductivity.

2.2.3.3 Classical nucleation theory. The coefficient C_f .

The coefficient C_f that appears on Equations 2.9 and 2.10 is computed accordingly to the work of Huang, Wang, and Li, 2020. In these work, the Lee model is used and the nucleation rate is introduced for the calculation of mass transfer rate between phases.

The concept behind the *Classical Nucleation Theory*, CNT, as described in Ickes et al., 2015 resides in the idea of droplet freezing. This is initiated in the fluctuation of molecules of a supercooled liquid due to thermal vibration which lead, at its turn, to spontaneous formation of ordered solid molecule clusters (ice embryos). The size of

these embryos oscillates as individual water molecules are crystallized or lost from the liquid phase. When the size of the embryo reaches a critical value, it leads a faster and auspicious thermodynamic joining of further water molecules to the crystal lattice. This means the critical embryo enhancing the "parent phase", supercooled liquid, to undergo a macroscopic phase transition: droplet freezing.

And this is what CNT aims to describe; the freezing process in terms of temperature-dependent nucleation rate by joining two components: thermodynamic and kinetic. These components, briefly described in the following chapters, are based on the theory found at Wu, Lai, and Zhang, 2015 and Huang, Wang, and Li, 2020, and adapted in Huang, Wang, and Li, 2020.

As a remark, in this thesis a brief introduction of this theory is given. However, for further details on the assumptions used refer to the literature.

Thermodynamic component

This thermodynamic component seeks for the number of critical embryos formed per unit of volume at a specific temperature. A decrease in the enthalpy, and consequently a change in Gibbs free energy required to form an ice embryo containing water molecules generates an energy barrier to nucleation. However, for ice embryo formation, this barrier needs to be overcome.

$$\Delta G_C = \underbrace{\Delta G_V}_{\text{volume term}} + \underbrace{\Delta G_S}_{\text{surface term}}$$
 (2.16)

$$\Delta G_c = -\frac{4}{3} \cdot \frac{\pi r^3}{\Omega} \cdot \Delta g_v + 4\pi r^2 \gamma_{sf}$$
 (2.17)

where r is the radius of a simplified spherical embryo, γ_{sf} the interfacial tension between phases Ω the volume of a single molecule ($\Omega = V_{m,w}/N_A$), $V_{m,w}$ is the molar volume and Δg_v represents the decrease in volume of the Gibbs free energy of a molecule and is defined as:

$$\Delta g_v = \frac{\Delta_m H_1}{N_A} \frac{\Delta T}{T^*} \tag{2.18}$$

where $\Delta_m H_1$ is the molar latent heat of crystallization, N_A is the Avogadro's number, T^* is the freezing temperature and $\delta T = T^* - T$, the degree of supercooling. The radius has an influence on the change in Gibbs free energy. This is when:

- $r < r_{crit} \Rightarrow \Delta G_c > 0 \mid \mid \Delta G_c \uparrow \Rightarrow r \uparrow \Leftarrow \text{ endothermic process}$
- $r > r_{crit} \Rightarrow \Delta G_c < 0 \mid \mid \Delta G_c \downarrow \Rightarrow r \uparrow \Leftarrow$ exothermic process

The critical radius exists when the global enthalpy variation gets negative. By differentiating Eq. 2.17 and setting $\frac{d(\Delta G_c)}{dr} = 0$, the critical radius is defined as:

$$r_{crit} = \frac{2\gamma_{sf}T^*V_{m,w}}{\Delta_m H_1 \Delta T} \tag{2.19}$$

Then, if subsituting Eq. 2.18 and 2.19 in Eq. 2.17, it is obtained the energy barrier:

$$\Delta G_{crit} = \frac{16\pi}{3} \cdot \frac{\gamma_{sf}^3 V_{m,w}^2 T^2}{\Delta_m H_1^2 \Delta T^2} = \frac{1}{3} \left(4\pi r_{crit}^2 \gamma_{sf} \right)$$
 (2.20)

In Huang, Wang, and Li, 2020, the expression concerning the variation of Gibbs function for the phase change does not include the molar volume of water but a

shape coefficient of nucleation. It involves the influence of the contact angle when going from a uniform state to an inhomogeneous. This shape factor is defined as:

$$\alpha_{ey} = \frac{2 - 3\cos\theta + \cos\theta^3}{4} \tag{2.21}$$

Temperature and saturation dependent number of ice embryos per unit volume of water may be expressed in a Boltzmann distribution form using ΔG :

$$N_{\text{embryo}} \left[\mathbf{m}^{-3} \right] = N_{\text{l}} \cdot \exp \left(-\frac{\Delta G}{k_{\text{B}} T} \right)$$
 (2.22)

where N_1 is a volume-based number density of water molecules in the liquid phase.

Kinetic component

The kinetic part of the nucleation rate is introduced in the form of water molecules flux. This is expressed as a Boltzmann distribution such that:

$$\Phi = \frac{k_B T}{h} \cdot \exp\left(-\frac{\Delta g}{k_B T}\right) \tag{2.23}$$

where h is the Planck's physical constant, and Δg the activation energy for the transfer of a water molecule across the phase boundary.

The rate at which the water molecules are transferred into an ince embryo is defined as:

$$K = n_{\rm s} \cdot 4\pi r_{\rm embryo}^2 \cdot Z \cdot \Phi \tag{2.24}$$

where n_s is the number of molecules and $4\pi r_{\rm embryo}^2$ is the surface area of the critical embryo and Z a kinetic prefactor. For seek of simplification, the authors of the theory suggest that the product of these terms are close to unity. Thus, considering this change, the equation yields as:

$$K = \Phi \tag{2.25}$$

Nucleation rate

Combining the thermodynamic component Eq. 2.22 and the kinetic one 2.25, the formulation of the nucleation rate can be expressed as:

$$J_{\text{hom}} \left[\mathbf{m}^{-3} \cdot \mathbf{s}^{-1} \right] = \underbrace{K}_{\text{Kinetics}} \cdot \underbrace{N_{\text{l}} \cdot \exp\left(-\frac{\Delta G}{k_{\text{B}}T} \right)}_{\text{Number of embryos}}$$
(2.26)

As a final step, inserting Eq. 2.25 into Eq. 2.26, the nucleation rate is expressed in the form of:

$$J_{\text{hom}} \left[\mathbf{m}^{-3} \cdot \mathbf{s}^{-1} \right] = \frac{k_B T}{h} \cdot \underbrace{\exp\left(-\frac{\Delta g^{\#}}{k_B T} \right)}_{\text{diffusion of molecules effect}}$$

$$\cdot N_1 \cdot \underbrace{\exp\left(-\frac{\Delta G}{k_B T} \right)}_{\text{nucleation effect}}$$
(2.27)

Fig. 2.1 characterizes the variation of the crystallization rate in function of the temperature. In the image, the dotted line shows how the nucleation effect is 0 close to the cooling, and as the temperature values decrease, this lines tend to 1. Moreover, the dashed line shows how the effect of the diffusion of the molecules increases as the temperature does. This prompts out the ease of the embryo formation when at low temperatures since the molecules cannot overcome the energy barrier to enter the embryo. However, the crystal growth becomes harder. This may be seen as constant search of equilibrium among the nucleation and the crystal growth. Finally,

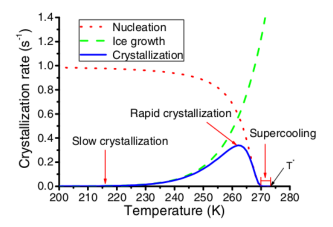


FIGURE 2.1: Crystallization rate versus temperature.

the coefficient C_f that appears on Equations 2.9 and 2.10 as commented above, is defined for the Lee model as:

$$C_f = J_{\text{hom}} \cdot V_l \tag{2.28}$$

where V_l is the volume of water in each cell.

2.2.4 Interphase porosity models

Interphase porosity models add an aritificial momentum source over the interface between phases to compute the sink of velocity in the solidified region. Therefore, influencing the behavior of the physics during the process of solidification or melting.

The model implemented in OpenFOAM is *Voller Prakash method* and it defines the source terms, S_y and S_z such that when along the fluid domain these terms take on a value of zero, the momentum equations are driven by the actual values of the velocities. On the other side, when it comes to treat the mushy region (i.e. porous region), the value of these source terms dominate convective, diffusive and transient terms and the momentum equation tends to approximate de Darcy law.

The two source terms as specified above,

$$\begin{cases}
S_y = -Av \\
S_z = -Aw
\end{cases}$$
(2.29)

Then, to specify a term for the function A, it is used the *Carman-Koseny equation*, which is derived from the Darcy law. The former expresses the gradient for the pressure as a combination of the velocity, \mathbf{u} , and the porosity, λ . The coefficient C

depends on the morphology of the medium.

$$gradP = -\left(\frac{C(1-\lambda)^2}{\lambda^3}\right)\mathbf{u} \tag{2.30}$$

To avoid division by zero, q is added to the equations shown

$$A = -\left(\frac{C(1-\lambda)^2}{\lambda^3 + q}\right) \tag{2.31}$$

The source terms S_y and S_z are added in the Eq. 2.32 and 2.33. The source term S_b corresponds to the body forces of the fluid and will be discused later on this thesis.

$$\frac{\partial(\rho v)}{\partial t} + \operatorname{div}(\rho \mathbf{u}v) = \operatorname{div}(\mu \operatorname{grad} v) - \frac{\partial P}{\partial y} + S_y$$
 (2.32)

$$\frac{\partial(\rho w)}{\partial t} + \operatorname{div}(\rho \mathbf{u}w) = \operatorname{div}(\mu \operatorname{grad} w) - \frac{\partial P}{\partial z} + S_z + S_b$$
 (2.33)

2.2.4.1 Surface tension model

The surface tension is only specified on a phase pair basis. In this version of OpenFOAM, it is present a constant model for a given σ .

Chapter 3

Numerical Simulation of Solidification Process

3.1 OpenFOAM. General Aspects

OpenFOAM is a free open-source software written in C++ and mainly conceived to perform computational fluid dynamics (CFD) simulations based on a finite volume discretization. In this first section, a brief introduction on the structure and functioning of the OpenFOAM software is given. In the folder structure tree shown in Fig. ??, it is shown a typical case setup for a phase change problem using <code>icoReactingMultiphaseFoam</code> solver.

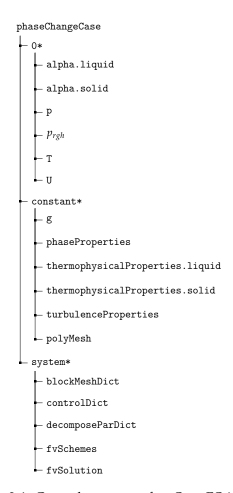


FIGURE 3.1: General structure of an OpenFOAM case.

3.1.1 Boundary Conditions Directory

The "0" directory gathers all the boundary conditions at time zero and the initial conditions to set up the case. As the simulation starts running, the information of these fields is saved in folders at every timestep.

3.1.2 Constant Properties Directory

The "constant" directory contains all the information typically regarding the physical properties which are kept constant through the simulation. Moreover, once the dictionary *blockMeshDict* is run, OpenFOAM creates a folder called *polyMesh* containing all the information relevant to the mesh (points, faces,...)

3.1.3 System Directory

This folder contains the files required by the control of the solver and the solution itself. The most common files are:

• **blockMeshDict**: in this file the parameters required to build up the computational domain, the mesh and the boundaries are found. The command **blockMesh** executes this dictionary creating the *polyMesh* folder commented above.

- **controlDict:** Time parameters associated to the computation are set in this file.
- decomposeParDict: In the realization of this thesis, the help of parallel computing is required. Thus, in this file, parameters regarding the decomposition of the mesh are configured. It is executed by means of the decomposePar application implicit in OF. The mesh is afterwards reconstructed by using reconstructPar
- **fvSchemes:** Schemes selected for the discretization of the derivative terms are defined. Among others, time schemes, gradient schemes, laplacian schemes, divergent schemes, interpolation schemes can be declared here.
- **fvSolution:** contains sub-dictionaries used to control the solvers and the solution algorithms. It also allows the definition of the fields resolution.

3.2 Solidification process. Methodology

A convection solver is used to represent the flow behavior generated by the density difference due to existing temperature gradients whithin the volume of control. A polynomic water density is implemented in the native OpenFOAM solver and compared with the standard Boussinesq approximation. The current model is validated against numerical results form the literature. The solution of this convection solver is later used as a boundary condition, before solidification phenomena plays a role.

3.3 OpenFOAM: BuoyantBoussinesqPimpleFOAM. Natural Convection solver

In a natural convection environment, the motion of the fluid is mainly driven by the density difference within the fluid volume of control. At its turn, the differences in the density, responsible for buoyancy forces, are generated by the existing temperature gradients. Within a physical context, the fluid near a hot heat source gets warmed up and, as a result, it becomes less dense moving up inside a domain. Consequently, the fluid in contact of the cold heat source is pushed from its zone to replace the hot fluid location. At this point, the cycle starts again repeating the physical phenomena.

3.3.1 Control Loop

The buoyantBoussinesqPimpleFoam is a solver used to solve non-steady buoyancy-driven fluids by using the Boussinesq approximation as a coupling between density and temperature fields. It considers the fluid as incompressible and uses the PIM-PLE algorithm for the pressure-velocity coupling. The flowchart of the integration procedure for the solver is presented below:

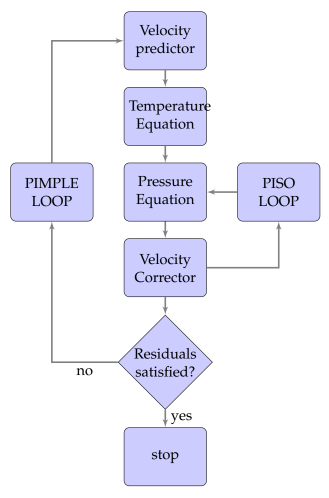


FIGURE 3.2: Flowchart of integration procedure. buoyantBoussinesqPimpleFoam

3.3.2 Governing Equations

In this section, the governing equations for the used solver are described first. The conservation of mass states that the mass flowing into the volume of control (CV) must be equal to the mass flowing out of such volume.

$$\frac{\partial v}{\partial y} + \frac{\partial w}{\partial z} = 0 \tag{3.1}$$

3.3.2.1 Momentum Equation

Throughout the CV the momentum of the fluid flow is preserved and here below it is expressed for the y-direction and z-direction.

$$\frac{\partial(\rho v)}{\partial t} + \operatorname{div}(\rho \mathbf{u}v) = \operatorname{div}(\mu \operatorname{grad} v) - \frac{\partial P}{\partial y}$$
(3.2)

$$\frac{\partial(\rho w)}{\partial t} + \operatorname{div}(\rho \mathbf{u}w) = \operatorname{div}(\mu \operatorname{grad} w) - \frac{\partial P}{\partial z} + S_b$$
(3.3)

where in the case of the *Boussinesq approximation* where the density variation is linear:

$$S_b = g \cdot \rho_r [1 - \beta (T - T_r)] \tag{3.4}$$

in the case of the implemented polynomial density which accounts for the inversion point as in Bourdillon, 2016:

$$S_b = g \cdot [\rho_r - \rho(T)] \tag{3.5}$$

where the polynomial expression from ρ is:

$$\begin{split} \rho(T) &= 999.840281167108 + 0.0673268037314653 \times T \\ &- 0.00894484552601798 \times T^2 \\ &+ 8.78462866500416.10^{-5} \times T^3 - 6.62139792627547.10^{-7} \times T^4 \end{split} \tag{3.6}$$

3.3.2.2 Temperature Equation

The temperature equation representing the convection phenomena yields as:

$$\frac{\partial T}{\partial t} + \frac{\partial (u_j T)}{\partial x_j} = \frac{\partial}{\partial x_j} \left(\gamma \frac{\partial T}{\partial x_j} \right) \tag{3.7}$$

where the thermal diffusivity, γ , is defined as:

$$\gamma = \frac{\lambda}{\rho_r c_p} \tag{3.8}$$

3.3.3 Case Description.

Here a squared geometry is created.

3.3.4 Hypotheses And Assumptions

3.3.5 Code implementations

3.3.6 Case Setup

Boundary conditions

Boundary	Conditions
Left	$T_l = 283, v_l = 0$
Right	$T_r=273, v_r=0$
Upper	$\frac{\partial T_u}{\partial n} = 0, v_u = 0$
Bottom	$\frac{\partial T_b}{\partial n} = 0, v_b = 0$

Table 3.1: Boundary conditions for natural convection case.

Thermophysical properties

Water properties	Symbol	Values	Units
Density	ρ_r	999.8	$kg.m^{-3}$
Dynamic viscosity	μ	0.001003	$kg.m^{-1}.s^{-1}$
Thermal conductivity	λ	0.6	$W.m^{-1}.K^{-1}$
Heat capacity	C_p	4182	$J.kg.K^{-1}$
Gravitational acceleration	8	9.81	$m.s^{-2}$
Thermal diffusivity	γ	1.435e-7	$m^2.s^{-1}$
Thermal expansion coefficient	β	6.734e-5	K^{-1}
Laminar Prandtl number	P_r	6.99	-
Reference temperature	T_r	6.734e-5	K

TABLE 3.2: Water properties for natural convection.

Solver parameters

Modeling Term	Keyword	Scheme	Remarks
Time derivatives Divergence term	ddtSchemes	Euler	
Gradient term Laplacian term Others	gradSchemes laplacianSchemes snGradSchemes interpolationSchemes	Gauss linear uncorrected uncorrected linear	

TABLE 3.3: Discretization schemes.

Equation	Linear Solver	Smoother/Preconditioner	Tolerance
Pressure correction equation	PCG	DIC	1e-8
Momentum equation	PBiCGStab	DILU	1e-6
Temperature equation	PBiCGStab	DILU	1e-6

TABLE 3.4: Solvers for the discretised equations.

Parameter	Value	Remarks
momentumPredictor	no	
nOuterCorrectors	1	
nNonOrthogonalCorrectors	0	
nCorrectors	2	

TABLE 3.5: Parameters for the discretised equations.

3.3.7 Validation of Results and Conclusions

In order to compare consistently the obtained results with those of the literature, the following dimensionless values are pointed out:

$$\tilde{T} = \frac{T - T_{\text{cold}}}{T_{\text{hot}} - T_{\text{cold}}} = \frac{T - 273}{10}$$
 (3.9)

$$\tilde{x} = \frac{x}{\ell} = \frac{x}{38 \times 10^{-3}} \tag{3.10}$$

$$\tilde{v} = \frac{v\ell}{\gamma} = \frac{v38 \times 10^{-3}}{1.435 \times 10^{-7}} \tag{3.11}$$

$$\tilde{u} = \frac{u\ell}{\gamma} = \frac{u38 \times 10^{-3}}{1.435 \times 10^{-7}} \tag{3.12}$$

$$\tilde{t} = \frac{t\gamma}{\ell^2} = \frac{t \times 1.435 \times 10^{-7}}{1.444 \times 10^{-6}} \tag{3.13}$$

$$\tilde{y} = \frac{y}{\ell} = \frac{y}{38 \times 10^{-3}} \tag{3.14}$$

3.4 OpenFOAM: IcoReactingMultiphaseInterFOAM. Phase-Change Process

IcoReactingMultiphaseInterFoam solver is a multiphase, multicomponent incompressible solver based on volume of fluid method. The solver captures the interfaces and includes contact angle and surface tension effects for each phase. Moreover, this solver supports mass and heat transfer across phases.

3.4.1 Control Loop

3.4.2 Phase models

The solver presents three phase model types:

- pureStaticSolidPhaseModel: For pure static phase, like a solid.
- pureMovingPhaseModel: For pure moving phase, like a fluid.
- multiComponentMovingPhaseModel: For multi-component moving phase, like a multi-component fluid.

3.4.3 Mass transfer models

For each pair of phases, two mass transfer models might be used:

- Lee model: Used for solid melting and liquid solidification.
- **KineticGasEvaporation:** Used for condensation and evaporation.

In this thesis, only the Lee model will be considered for further explanation.

3.4.4 Governing Equations

- 3.4.4.1 Momentum Equation
- 3.4.4.2 Pressure Equation
- 3.4.4.3 Energy Equation

3.5 Case Description.

For the solidification process, two geomtries are created: A squared and cylindrical plane geometries.

3.5.1 Hypotheses And Assumptions

3.5.2 Code implementations

3.5.3 Case Setup

Boundary conditions

For the squared cavity: The initial conditions for the velocity and temperature fields are inherited from the last timestep of the natural convection case. *inletOutlet is normally the same as zero gradient but it switches to fixed value if the velocity vector next to the boundary aims inside the domain. For the cylinder:

Boundary	Conditions
Left	$\frac{\partial \alpha_l}{\partial n} = 0, \frac{\partial \alpha_s}{\partial n} = 0$
Right	$\alpha_l = inletOutlet * (1), \alpha_l = inletOutlet(0)$
Upper	$rac{\partial lpha_l}{\partial n} = 0, rac{\partial lpha_s}{\partial n} = 0 \ rac{\partial lpha_l}{\partial n} = 0, rac{\partial lpha_s}{\partial n} = 0$
Bottom	$\frac{\partial \alpha_l}{\partial n} = 0, \frac{\partial \alpha_s}{\partial n} = 0$

TABLE 3.6: Boundary conditions for natural convection case.

Boundary	Conditions
Left	$T_l = 283, v_l = 0$
Right	$T_r=273, v_r=0$
Upper	$\frac{\partial T_u}{\partial n} = 0, v_u = 0$
Bottom	$\frac{\partial T_b}{\partial n} = 0, v_b = 0$

TABLE 3.7: Boundary conditions for natural convection case.

Thermophysical properties

Water properties	Symbol	Values	Units
Water density	ρ_l	999.8	$kg.m^{-3}$
Ice density	$ ho_s$	916.8	kg.m ⁻³ kg.m ⁻³
Water kinematic viscosity	ν_l	1.79e-6	$m^2.s^{-1}$
Ice kinematic viscosity	$ u_s$	2.0e-6	$m^2.s^{-1}$
Water thermal conductivity	λ_l	0.56	$W.m^{-1}.K^{-1}$
Ice thermal conductivity	λ_s	2.26	$W.m^{-1}.K^{-1}$
Heat capacity	$C_{p_l} = C_{p_s}$	4202	$J.kg.K^{-1}$
Gravitational acceleration	8	9.81	$m.s^{-2}$
Thermal diffusivity	$\dot{\gamma}$	1.435e-7	$m^2.s^{-1}$
Thermal expansion coefficient	β	6.734e-5	K^{-1}
Latent heat	$\stackrel{\cdot}{L}$	335000	$J.K^{-1}$
Laminar Prandtl number	P_r	6.99	-
Reference temperature	T_r	6.734e-5	K
Darcy's constant	D_c	10e8	-

TABLE 3.8: Water properties for natural convection.

Lee's model thermophysical parameters:

Two phase properties

Within a multiphase framework, a model reflects a jump in properties through the interphase. Thus, a smooth transition between phase properties must be achieved.

$$\lambda = \lambda_{\ell} \alpha_{\ell} + \lambda_{s} f_{s} \tag{3.15}$$

$$C_p = C_{p_\ell} \alpha_\ell + C_{p_s} f_s \tag{3.16}$$

$$\mu = \mu_{\ell} \alpha_{\ell} + \mu_{s} f_{s} \tag{3.17}$$

In the case of polynomial density variation it is settled in a similar manner. The polynomial is not thought to suit negative temperatures, and when the problem is within this range, the density should take ice's density.

$$\rho(T)' = \rho(T)\alpha_{\ell} + \rho_s f_s \tag{3.18}$$

Solver parameters

Modeling Term	Keyword	Scheme	Remarks
Time derivatives Divergence term	ddtSchemes		
Gradient term Laplacian term Others	gradSchemes laplacianSchemes snGradSchemes interpolationSchemes		

TABLE 3.9: Discretization schemes.

Equation	Linear Solver	Smoother/Preconditioner	Tolerance
Pressure correction equation Momentum equation Volume fraction equation	smooth Solver	DIC symGaussSeidel symGaussSeidel	

TABLE 3.10: Solvers for the discretised equations.

Parameter	Value	Remarks
nAlphaCorr nAlphaSubCycles cAlpha momentumPredictor nOuterCorrectors nNonOrthogonalCorrectors nCorrectors	PCG smoothSolver smoothSolver	DIC symGaussSeidel symGaussSeidel

TABLE 3.11: Parameters for the discretised equations.

3.5.4 Validation of Results and Conclusions

Chapter 4

Numerical Simulation of Heat Transfer

- 4.1 Methodology
- 4.2 OpenFOAM: chtMultiphaseInterFOAM. Conjugate Heat Transfer
- 4.2.1 Control Loop
- 4.2.2 Governing Equations of the Fluid Region
- 4.2.2.1 Momentum Equation
- 4.2.2.2 Pressure Equation
- 4.2.2.3 Energy Equation
- 4.2.3 Governing Equations of the Solid Region
- 4.2.3.1 Energy Equation
- 4.2.4 Case Setup
- 4.2.5 Validation of Results and Conclusions

Chapter 5

Conclusions

Chapter 6

Future Works

Appendix A

Frequently Asked Questions

A.1 How do I change the colors of links?

The color of links can be changed to your liking using:

\hypersetup{urlcolor=red}, or

\hypersetup{citecolor=green}, or

\hypersetup{allcolor=blue}.

If you want to completely hide the links, you can use:

\hypersetup{allcolors=.}, or even better:

\hypersetup{hidelinks}.

If you want to have obvious links in the PDF but not the printed text, use:

\hypersetup{colorlinks=false}.

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