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Modeling dynamic systems: Contribution to the Unsteady Behavior of a Condenser Based on the Pseudo-Bond Graph Approach

M. Kebdani^{*a}, G.Dauphin-Tanguy^b, A. Dazin^a

^aArts et Métiers Paris Tech/ LML UMR CNRS 8107, Boulevard Louis XIV, 59000 Lille. France.

^bEcole Centrale de Lille/ CRISAL UMR CNRS 9189, CS 20048, 59651 Villeneuve d'Ascq. France.

**For correspondence: Email: kebdani.mp@hotmail.fr*

Abstract

This article is devoted to the dynamic study of the brazed plate condenser (BPC). The proposed model is based on the bond graph theory. The proposed model is based on the bond graph theory because of its energetic approach and multi-physics character of the studied system. The model is discretized into five control volumes. The resolution of mass and energy equations is done by Runge-Kutta method embedded in 20sim software. Analyses of simulation results show that the model has a good ability to transcribe the time evolution of the temperature and pressure in both regimes, transitional and permanent. Also, the model is experimentally validated without any fitting of the set of thermal exchange coefficients.

Keywords: Bond graph theory; Dynamic systems modeling; Multiphysic coupling; Two-phase flow; Heat transfers; Transient; Condenser.

Nomenclature

c_p	Specific heat	$J / kg.K$	λ	Thermal conductivity	$W / m \cdot K$
D_h	Hydraulic diameter	m	μ	Dynamic viscosity	$Pa \cdot s$
e	Wall thickness	m	ρ	Density	kg / m^3
G	Flow density	W	β	Chevron angle	degre
H	Latent heat of condensation	J / kg	f	Friction coefficient	---
\dot{H}	Flow enthalpy	J / s			
g	Acceleration	m	acc	Accumulation	
l	Length	m	amb	Ambiant	
L	Channels length	m	col	Colector	
m	Mass	kg	conv	Convection	
\dot{m}	Mass flow rate	kg / s	dis	Distributor	
N_{can}	Channels number	---	fric	Friction	
P	Pressure	Pa	in	Inlet	
Re	Reynolds number	---	grav	Gravity	
x	Vapor quality		liq	Liquid	
p_{co}	Pitch corrugation	m	out	Outler	
Pr	Prandtl number	---	red	Reduced	
S	Surface	m^2	sat	Saturation	
t	Time	s	vap	Vapor	
T	Temperature	K	Δ	Difference	

1 Introduction

It turns out that the Brazed Plates Condenser (BPC) technology, based on thermal plates with corrugated geometry, allows high performances. This is due to the compact size, and effective heat transfers. These advantages extend the use of BPC. For instance, they equip power electronics cooling systems [1]. A dynamic study of this condenser is necessary to better understand the transitional behavior of this exchanger. Hence the purpose of this article is the development of an original dynamic model based on bond graph approach. The model is experimentally validated using a test rig specially developed for this need.

The article starts with a succinct presentation of the condenser. Afterward, is

presented a state of the art of BPC modeling, convective coefficients correlations and pressure losses correlations. In the second section is given a detailed mathematical description of the proposed thermo hydraulic model. The article ends up with a discussion of experimental results leading to conclude on the validity of the model.

2 General description of a BPC

2.1 Technology and advantages of BPC

The BPC is composed of corrugated plates. The set of thermal plates is brazed with copper, some connections called “ports” are positioned at the corners of the cover plates, see Figure 1. During the brazing operation, the copper is melted and perfectly

conducted to connection points between plates, forming thereby a unique functional block. This process ensures a high level of impermeability and guarantees a structural integrity which allows BPC to withstand important operating pressures up to 45 bar.

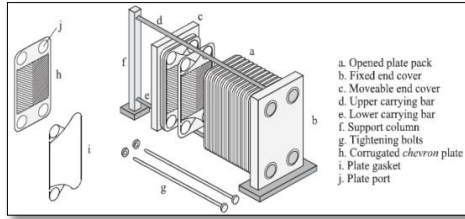


Figure 1: Exploded view of a BPC.

The type of the flow pattern adopted in this work is a counter-current. The two coolants (water) flow in opposite directions, as shown schematically in Figure 2 (a). Indeed, this configuration promotes better heat exchanges, compared to performance provided by a co-current condenser. It is proven that the outlet temperature of the fluid T_{out_PC} may be lower than the outlet temperature of the secondary T_{out_SC} , as shown in Figure 2 (b).

The BPC possesses a number of benefits. They may exist with a volume up to 30% less than that of a traditional tubular condenser [2] and [3]. Also, channel design provides efficient heat transfers. In fact, the multiple fluid streams intersect at the central channel, provoking a turbulent flow [4] and [5], which boosts great heat transfers [6] and [2]. Furthermore, turbulence and secondary flows developed in the exchanger, inhibit the formation of the fouling layers. For more details see [2].

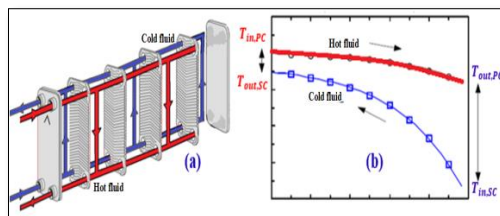


Figure 2: (a) Flow direction of cold and hot fluid in a counter-current BPC
(b) Spatial evolution of the temperature of cold and hot fluids in a counter-current BPC

The BPC is used in a wide range of applications: air conditioning, refrigeration, food and chemical industry. This component is very convenient in monophasic and two phase flow applications. References [2], [5], [6] and [7] contain more details about the different uses of the BPC.

2.2 State of the art on BPC modeling

Unlike tubular condensers, nowadays the scientific literature identifies very few modeling works specific to BPC. On the other side, the majority of disclosed researches works give the main equations used for calculation, important details are not always available. *Moreover BPC dynamic models based on bond graph methodology do not exist in the open literature.* Furthermore, the literature revue shows that the dynamic modeling contributions are mainly proprietary of manufacturers and are confidential.

Two examples of works related to BPC

Method of Arman and Rabas [8]: authors developed a stationary model, able to predict the thermo hydraulic state of steam, condensate film and coolant liquid. The BPC iterative model is based on an incremental procedure.

Model of Gut [9]: they present an algorithm allowing the simulation of flow in the steady state. The algorithm is based on the resolution of energy conservation equation. The assumptions are:

- the model treats only steady states.
- no heat losses to be considered.
- the author does not deal with the phase change problems.
- the overall heat transfer coefficient is meant to be constant.
- the flow is assumed to be “plugged” along the exchanger.

Comparison and scientific contributions

Based on the comparison with the existing modeling works, the contributions are:

1. *The use of bond graph tool to model the BPC is new. The bond graph is particularly useful in writing the system equations and in assessing the effects of changes in the model even before the equations are written [10] and [11].*
2. *The proposed model takes into account the dynamic behavior, unlike the majority of classical models limited to the permanent regime.*
3. *The transitional model pays particular attention to the heat transfer coefficients. The novelty, compared to previous work, is twofold: first, coefficients vary with the evolution of local thermodynamic conditions. Second, the thermal behavior of the condenser is experimentally validated without any use of recalibration of the set of used coefficients.*
4. *Unlike most existing models, the proposed model considers the heat exchange with the outside and the longitudinal conduction along the walls.*
5. *Gaussian and polynomial correlations of thermo physical properties are specially developed on the basis of data provided by NIST (site 1).*

State of the art on the thermal convection coefficients specific to BPCs

The major problems encountered when modeling the BPC are:

- Find out the correct heat transfer coefficients to be used in both primary and secondary circuits.
- Choose the right method to calculate the pressure losses coefficients.

There are no universal and reliable correlations. The reason is due to the strong dependence of coefficients to many parameters such as: plate geometry, mass flow rate, thermo-physical properties of the fluid, vapor quality, pressure and temperature, Reynolds number and dependence of the coefficient h_{conv} on the pattern flow.

Chosen correlations: The scientific literature provides access to a wide range of

heat transfer and friction coefficients. References [12] and [13] present an excellent synthesis work. Nevertheless, we should keep in mind that each correlation remains closely related to the operating conditions under which it has been developed. That is why correlations tested and adapted to our work are given in Appendices B and C.

State of the art on the friction coefficients related to BPCs

The pressure drop in two-phase flows inside a BPC is mainly the sum of three contributions:

- Pressure variation due to gravitation.
- Pressure drops at ports.
- Pressure drops due to friction of the two-phase flow.

This section aims at presenting the pressure losses correlations applied for two-phase flows.

Pressure variation due to gravitation: the change in pressure due to gravity is determined by the hydrostatic equation:

$$\Delta P_{grav} = \pm \rho_{aver} \cdot g \cdot L \quad (1)$$

where the average density ρ_{aver} of the two phase flow is given by:

$$\rho_{aver} = \left[\frac{x}{\rho_{vap}} + \frac{1-x}{\rho_{liq}} \right]^{-1} \quad (2)$$

Pressure drop in distributors and collectors: the losses generated inside distributors (inflow) and collectors (outflow) are empirically estimated by Shah and Focke [3]:

$$\Delta P_{col/dis} = \frac{1.5 \cdot G^2}{2 \cdot \rho_{aver}} \quad (3)$$

Pressure losses due to friction of the two-phase flow: the friction coefficient in two-phase flow is complicated to determine. The Table 2 shows a number of correlations available in the literature.

Chosen correlations: The correlations feeding the proposed model are chosen as follows:

Monophasic	Diphasic		
	Re < 100	Transition	Re > 1500
Primary circuit	Laminar		Turbulent
• Heavner	• Nusselt		• Isashenko
• Muley and Manglik	• Hagen-Poiseuille		• Shaw
Secondary circuit			
• Muley and anglik	$f = \frac{64}{Re}$		• Yan
• Kim			• Blasius: $f = \frac{0.3164}{Re^{0.25}}$

Figure 3: Classification of correlations in function of flow regime.

3 Bond graph model of the BPC.

3.1 About the theory of bond graphs

Industrial thermo fluids systems are multidisciplinary. Their functioning involves systematically several domains of the physic (mechanic, hydraulic, thermal, chemical, and electrical fields). This phenomenological complexity makes it difficult to develop a standard model based on partial derivative equations (CFD¹). The bond graph methodology is based on differential equations. It allows to represent graphically the power exchanges between components of a complex system. A bond graph model is a unified graphical representation for all areas of engineering science. It is based on the principle of analogy between different fields of physics [14] and [15]. Also, a bond graph model is both structural and behavioral which allows simulation and analysis of the model properties [16]. The bond graph allows modeling adapted to the growing technological needs. In fact, the model can be easily modified by adding or removing elements, permitting a modeling approach called ‘bottom-up’ or ‘top down’.

3.2 Assumptions

The model is based on the following hypotheses:

- The fluid is unidirectional.
- The fluid in the secondary circuit (SC) flows with a constant mass flow rate.

- The two-phase volume connoted v_6 in Figure 5 occupies half of the primary circuit volume (PC).
- The refrigerant used for the simulation is water, while the validation tests are conducted with water containing 4% of PAG²
- The temperature gradient in the condensate film is assumed to be linear: this hypothesis is plausible given the small hydraulic diameter of channels.

3.3 Control volumes

The BPC is composed of channels in both circuits: hot, called primary circuit (PC), and the cold one also called secondary circuit (SC), as shown in Figure 4.

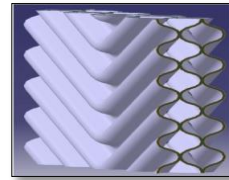


Figure 4: internal geometry of flow channels in a BPC.

In our modeling work the BPC is the union of six control volumes:

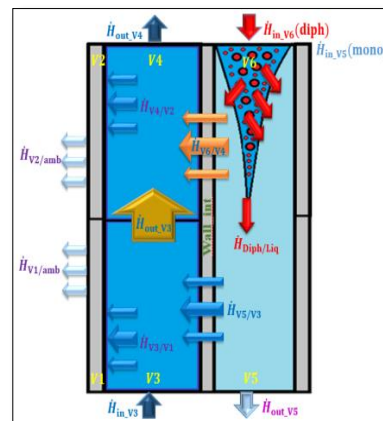


Figure 5: Volume discretization of the BPC, and distribution of heat flows occurring during the condensation phenomenon.

The primary circuit (PC) is assimilated to a simple pipe. In case of two-phase flow, the PC is divided into two volumes separated by a condensation interface as shown Figure 5:

¹ CFD: Computational Fluid Dynamics

² Polyalkylene Glycol

V6 a diphasic flow: homogeneous mixture of liquid and vapor.

V5 a monophasic flow: constituted of condensate film and the liquid “drained” down the condenser.

The **secondary circuit (SC)** has a cylindrical form and contains water in single phase state. This circuit is represented by two volumes V3 and V4 as shown in Figure 5.

The material body of the condenser is represented by two volumes V1 and V2.

3.4 General structure of the bond graph model.

The structure of the bond graph model is shown in Figure 6. The six sub models shown in Figure 6 refer to the six volumes mentioned above and are represented by the following bond graph elements:

Volume	designation	Bond graph element
V1 and V2	outer wall of the BPC	C elements with 1 thermal port
V3 and V4	Secondary circuit	RC elements with 2 ports (thermal and hydraulic).
V5	liquid portion occupying the primary circuit	RC element with 2 ports (thermal and hydraulic).
V6	two-phase portion present in the primary circuit	C element with 2 ports (thermal and hydraulic).

Blue lines correspond to the hydraulic power exchanges, while the orange lines represent the thermal power transfers. All studied heat transfer phenomena (convection and conduction) are modeled by dissipative elements

3.5 Mathematical description of the dynamic bond graph model

A detailed description of conservation equations (mass and energy) governing the thermo hydraulic behavior of the condenser is proposed here. The resolution of the equations is done by Runge-Kutta method embedded in 20sim software (site 2). The chosen "effort / flow" variables are pressure

and mass flow rate for hydraulic domain and temperature and enthalpy flow (for convection) or heat flow (for conduction) for thermal domain. This is why the proposed model is a pseudo bond graph. Moreover it is possible to use volume flow rate [17] and entropy flow rate instead of mass flow rate and enthalpy flow rate to build a real Bond Graph, however the mathematical formalism will be more complicated. Note, that interpolations giving the evolution of thermo physical properties are specifically developed for the current project with high accuracy, and are given in [18].

Volume V1 of the outer wall

The temperature $T_{V1}(t)$ is calculated using the energy balance. Furthermore, this volume is submitted to two thermal flows, as shown in Figure 5, $\dot{H}_{V3/V1}$ modeled in bond graph by $R_{Wall_V1_INT}$ and $\dot{H}_{V1/amb}$ modeled by the element $R_{Wall_V1_EXT}$. The energy balance is written at the O_5 junction:

$$\dot{H}_{Wall_V1_acc} = \dot{H}_{V3/V1} - \dot{H}_{V1/amb} \quad (4)$$

The enthalpy stored inside v_1 is then calculated in the element C_{Wall_V1} as:

$$H_{Wall_V1_acc}(t) = \int_t \dot{H}_{Wall_V1_acc}(t) dt + H_{Wall_V1,0} \quad (5)$$

where, $H_{Wall_V1,0}$ is the initial enthalpy in the wall v_1 .

The constitutive law of the element C_{Wall_V1} giving the temperature of the wall is then:

$$T_{V1}(t) = \frac{H_{Wall_V1_acc}}{m_{V1} \cdot c_{p,V1}} \quad (6)$$

$c_{p,V1}$: Specific heat of the wall (inox316).

Volume V2 of the outer wall

The temperature $T_{V2}(t)$ inside the volume v_2 is obtained in a similar way from the constitutive law of the element C_{Wall_V2}

$$T_{V2}(t) = \frac{H_{Wall_V2_acc}}{m_{V2} \cdot c_{p,V2}} \quad (7)$$

with: $c_{p,V2} = c_{p,V1} = 416 \text{ (J / kg.K)}$ et $m_{V2} = m_{V1}$.

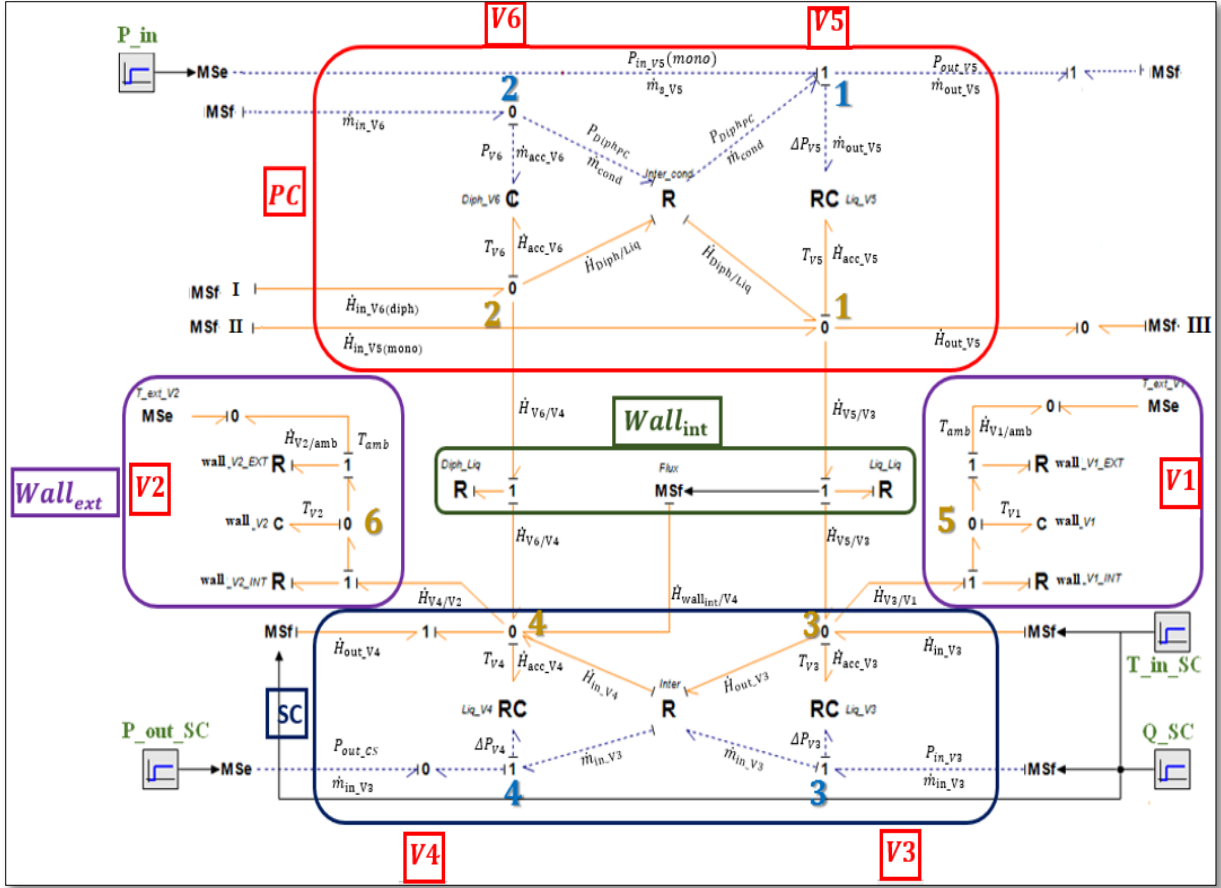


Figure 6: Dynamic bond graph model of the BPC.

Volume V3 of the liquid in the CS

Mass conservation equation

The coolant mass flow rate \dot{m}_{in_V3} is considered constant. Which means that there is no mass storage in the volume V3.

Energy balance equation

The energy balance is written in the 0₃ junction:

$$\dot{H}_{acc_V3} = \dot{H}_{in_V3} + \dot{H}_{V5/V3} + \dot{H}_{V3/V1} - \dot{H}_{s_V3} + \Delta P_{V3} \cdot \frac{\dot{m}_{in_V3}}{\rho_{liq_V3}} \quad (8)$$

The enthalpy stored in v_3 is then calculated in the element RC_{Liq_V3} as:

$$H_{acc_V3}(t) = \int_t \dot{H}_{acc_V3}(t) dt + H_{liq_V3,0} \quad (9)$$

where, $H_{liq_V3,0}$ is the initial enthalpy in the wall v_3 . The constitutive law of the element RC_{Liq_V3} giving the temperature of the wall is then:

$$T_{V3}(t) = \frac{H_{acc_V3}(t)}{\frac{\dot{m}_{in_V3}}{2} \cdot c_{p,liq,V3}} \quad (10)$$

The pressure drop ΔP_{V3} modeled by the 2-ports element RC_{Liq_V3} is obtained using equations: (11), (12), (13) and (14). Frictional pressure losses induced in single phase flow ΔP_{fric} are calculated using Darcy-Weisbach formula.

$$\Delta P_{V3} = \Delta P_{grav} + \Delta P_{fric} + \Delta P_{col/dis} \quad (11)$$

where :

$$\Delta P_{grav} = \rho_{liq_V3} \cdot g \cdot L_{V3} \quad (12)$$

$$\Delta P_{col/dis} = \frac{1.5 \cdot G^2}{2 \cdot \rho_m} \quad (13)$$

$$\Delta P_{fric} = \frac{1}{2} \cdot \rho_{liq_V3} \cdot v^2 \cdot \frac{L_{V3}}{D_h} \cdot f = 2 \cdot \frac{\dot{m}_{in_V3}^2}{\rho_{liq_V3} \cdot \pi^2 \cdot D_h^5} \cdot f \quad (14)$$

If : $Re < 100$ (laminar regime according to Wanniarachchi [5]), then Hagen- Poiseuille formula is used:

$$f = \frac{64}{Re} \quad (15)$$

If not (transient or turbulent regime), the formula of Blasius for turbulent flow regime gives:

$$f = \frac{0.3164}{Re^{0.25}} \quad (16)$$

Now it remains to determine the hydraulic diameter: D_h . For a better understanding of the geometric problem, parameters used in the formula (17) are denoted in Figure 7.

$$D_h = \frac{4 \cdot S_{V3}}{Per_{V3}} = \frac{4 \cdot b \cdot l}{2 \cdot l \cdot \Phi} = \frac{2 \cdot b}{\Phi} \quad (17)$$

with :

- $\Phi = 1.17$, a value given by the manufacturer SWEP.
- $b = p - t$. Where “p” is the corrugation depth and “t” the corrugation thickness.

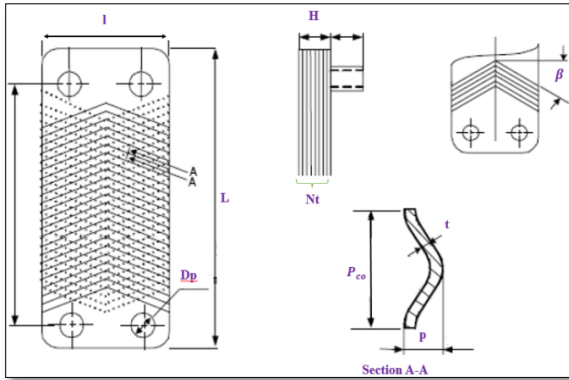


Figure 7: Specification of geometric parameters on the thermal plate of a BPC.

Volume V4 of the liquid in the CS

The liquid temperature is calculated as:

$$T_{V4}(t) = \frac{H_{acc_V4}(t)}{\frac{m_{in_V4}}{2} \cdot c_{p,liq,V4}} \quad (18)$$

Volume V5 of the liquid in the PC

As long as the liquid temperature does not exceed the saturation temperature corresponding to the pressure in the volume V5, the condenser contains only the liquid phase. Henceforth, a phase indicator is defined as follows:

- if $T_{liq_V5} < T_{sat}(P_{evap})$: then the fluid is liquid in a single-phase state (no vapor trace exists in the liquid) and $ind_{PC} = 0$. In this case, the whole PC is modeled only by

the element RC_{Liq_V5} while the element C_{Diph_V6} has no physical impact.

- if $T_{liq_V5} > T_{sat}(P_{evap})$, then: $ind_{PC} = 1$. It is assumed that the total volume of the PC is divided into two equal volumes V5 and V6, successively represented by the elements: RC_{Liq_V5} and C_{Diph_V6} .

Case I: $ind_{PC} = 0$: PC, containing only liquid, is modeled by the element: RC_{Liq_V5} .

Mass conservation equation

During the single-phase flow it is assumed that the mass flowrate \dot{m}_{out_V5} of the refrigerant through the PC is constant.

Energy balance equation

The energy balance is written in the 0₁ junction:

$$\dot{H}_{acc_V5} = \dot{H}_{in_CP(mono)} + \dot{H}_{Diph/Liq} - \dot{H}_{V5/V3} - \dot{H}_{out_V5} + \Delta P_{V5} \cdot \frac{\dot{m}_{out_V5}}{\rho_{liq_V5}} \quad (19)$$

The enthalpy stored within V5 is then calculated in the element RC_{Liq_V5} as:

$$H_{acc_V5}(t) = \int_t \dot{H}_{acc_V5}(t) dt + H_{liq_V5,0} \quad (20)$$

where, $H_{liq_V5,0}$ is the initial enthalpy in the volume V5.

The constitutive law of the element RC_{Liq_V5} giving the temperature of the wall is:

$$T_{V5}(t) = \frac{H_{acc_V5}(t)}{m_{out_V5} \cdot c_{p,liq_V5}} \quad (21)$$

The pressure drop ΔP_{V5} in the volume V5 represented by the multiport element RC_{Liq_V5} is calculated like ΔP_{V3} .

Case II: $ind_{PC} = 1$: the liquid volume V5 is modeled by the element: RC_{Liq_V5} . Equations specific to RC_{Liq_V5} are identical to those presented in the case above, but with a volume ($V_5/2$) and a pipe length ($L_{V5}/2$).

Volume V6 of the two-phase mixture

The two-phase volume v_6 constituted of half the total volume of the PC, is modeled by the element C_{Diph_V6} whose mathematical description is:

Mass conservation equation:

This equation is written in the thermal 0_{h2} junction

$$\dot{m}_{acc_V6} = \dot{m}_{in_V6} - \dot{m}_{cond} \quad (22)$$

- \dot{m}_{in_V6} is the mass flow rate of the fluid entering into the condenser. It is calculated by solving the momentum equation at the condenser inlet.
- The condensate mass flow rate \dot{m}_{cond} in the condenser is established in [18]. Its expression is:

$$\dot{m}_{cond} = \rho_{liq} \cdot \frac{(\rho_{liq} - \rho_{vap}) \cdot g \cdot \delta_{film}^3}{3 \cdot \mu_{liq}} \cdot \left(2 \cdot \pi \cdot \frac{D_h}{2} \right) \quad (23)$$

$$\delta_{film} = \left[\frac{4 \cdot \mu_{liq} \cdot \lambda_{liq} \cdot L / 2 \cdot (T_{wall} - T_{vap})}{g \cdot \Delta H \cdot \rho_{liq} \cdot (\rho_{liq} - \rho_{vap})} \right]^{1/4} \quad (24)$$

The mass stored in the volume v_6 is calculated in the multiport element C_{Diph_V6} as:

$$m_{acc_V6}(t) = \int_t \dot{m}_{acc_V6}(t) dt \quad (25)$$

Energy balance equation

The energy balance is written in the 0_{l2} junction:

$$\dot{H}_{acc_V6} = \dot{H}_{in_V6(diph)} - \dot{H}_{Diph/Liq} - \dot{H}_{V6/V4} \quad (26)$$

The enthalpy stored inside v_6 is then calculated in the element C_{Diph_V6} as:

$$H_{acc_V6}(t) = \int_t \dot{H}_{acc_V6}(t) dt \quad (27)$$

The constitutive laws of the element C_{Diph_V6} giving the pressure P_{V6} and the temperature T_{V6} of the two-phase flow are the result of a complex mathematical calculation that requires, at each time step, a connection between Matlab and 20sim. The connection is meant to solve a polynomial equation of 12th order whose solution is the pressure of the two-phase mixture. On the other hand, this method detailed in appendix A is able

to predict the temporal evolution of the vapor quality $x(t)$ of the two-phase mixture.

Temperature of the two-phase mixture:

this temperature is obtained directly from the pressure $P_{V6}(t)$, since it is assumed that the vapor-liquid mixture is saturated. Pressure and temperature are related by the following interpolation, especially developed for the project:

$$T_{V6}(t) = 367.5 \cdot \exp\left(-\left(\frac{P_{V6}(t) - 1.078}{1.081}\right)^2\right) + 125 \cdot \exp\left(-\left(\frac{P_{V6}(t) - 0.173}{0.4797}\right)^2\right) + 38.87 \cdot \exp\left(-\left(\frac{P_{V6}(t) - 0.03264}{0.1967}\right)^2\right) \quad (28)$$

3.6. Expression of conducto-convective flows

Heat flows are modeled using dissipative R-elements, used formula are listed in Appendix D: Conduction flows. According to Newton formula we have:

$$\Phi = h_{conv} \cdot S_{ech} \cdot \Delta T \quad (29)$$

Convected enthalpy is : $\Phi = \dot{m} \cdot c_p \cdot T$, with:

BG element	heat flow Φ	Mass flow rate \dot{m}	Specific heat c_p	T
$MSf(\dot{H}_{in_V3})$	\dot{H}_{in_V3}	\dot{m}_{in_V3}	$c_{p,in_V3}(T_{in_CS})$	T_{in_CS}
R_{inter}	\dot{H}_{out_V3}	\dot{m}_{in_V3}	$c_{p,V3}(T_{V3})$	T_{V3}
$MSf(\dot{H}_{out_V4})$	\dot{H}_{out_V4}	\dot{m}_{in_V3}	$c_{p,V4}(T_{V4})$	T_{V4}
$MSf(I)$	$\dot{H}_{in_V5(mono)}$	\dot{m}_{out_V5}	$c_{p,in_V5}(T_{V5})$	T_{V5}
$MSf(II)$	$\dot{H}_{in_V6(diph)}$	\dot{m}_{in_V6}	$c_{p,in_V6}(T_{V6})$	T_{V6}
R_{inter_cond}	$\dot{H}_{Diph/Liq}$	\dot{m}_{cond}	$c_{p,diph,V6}(T_{V6})^*$	T_{V6}
$MSf(III)$	\dot{H}_{out_V5}	\dot{m}_{out_V5}	$c_{p,V5}(T_{V5})$	T_{V5}

Table 1: convection flows.

where : $c_{p_diph,V6}(T_{V6})^* = x \cdot c_{p,vap}(T_{V6}) + (1-x) \cdot c_{p,liq}(T_{V6})$

4 Experimental validation

4.1 Sensors

The Figure 8 gives a schematic representation of the BPC, as well as the

distribution of different sensors used to validate the model. A first set of four sensors provides experimental data (pressure and temperature) considered as inputs to the developed model: P_{in} , P_{out_SC} , Q_{SC} , T_{in_SC} . A second set of five sensors is used to validate the model simulations: P_{in_PC} , T_{in_PC} , P_{out_PC} , T_{out_PC} , T_{out_SC} .

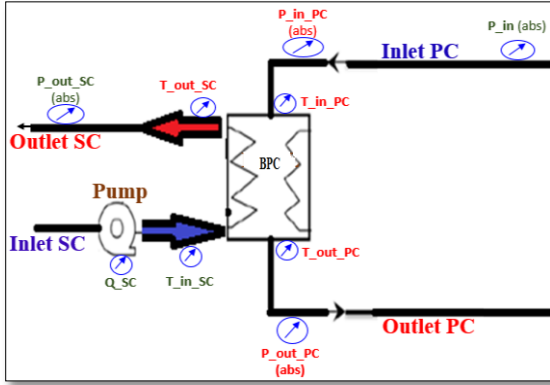


Figure 8: Position of pressure, temperature and mass flow sensors mounted on the BPC.

All geometric data required to launch a simulation are reported in Appendix E.

Description of the validation test

The three experimental data (P_{in} , $\dot{m}_{SC} = \frac{Q_{SC}}{\rho_{liq}}$, T_{in_SC}) represent the boundary conditions

applied to the condenser and are plotted in Figure 9. The pressure P_{out_SC} is atmospheric. The test is composed of four steps reported in Figure 9:

- The interval referenced **II** in the Figure 9 corresponds to the heating of the refrigerant moving through the condenser. During this phase, the fluid heats up but remains liquid.
- The beginning of interval **III** is marked by a sudden peak of pressure P_{in} (highlighted by a circle). Indeed, this peak reflects the birth of vapor bubbles.
- Furthermore, the pressure P_{in} is fluctuating throughout intervals **III** and **IV**. These oscillations are due to the presence of vapor bubbles.
- During interval **IV** the circulation of the coolant fluid contained in the secondary circuit is triggered. The corresponding mass flow rate \dot{m}_{dot_SC} is given by graph 2.

The model validation is based on comparison of the time evolution of simulated and measured values of: temperatures and pressures in volumes: V5 (liquid) and V6 (two-phase mixture), shown schematically in Figure 5.

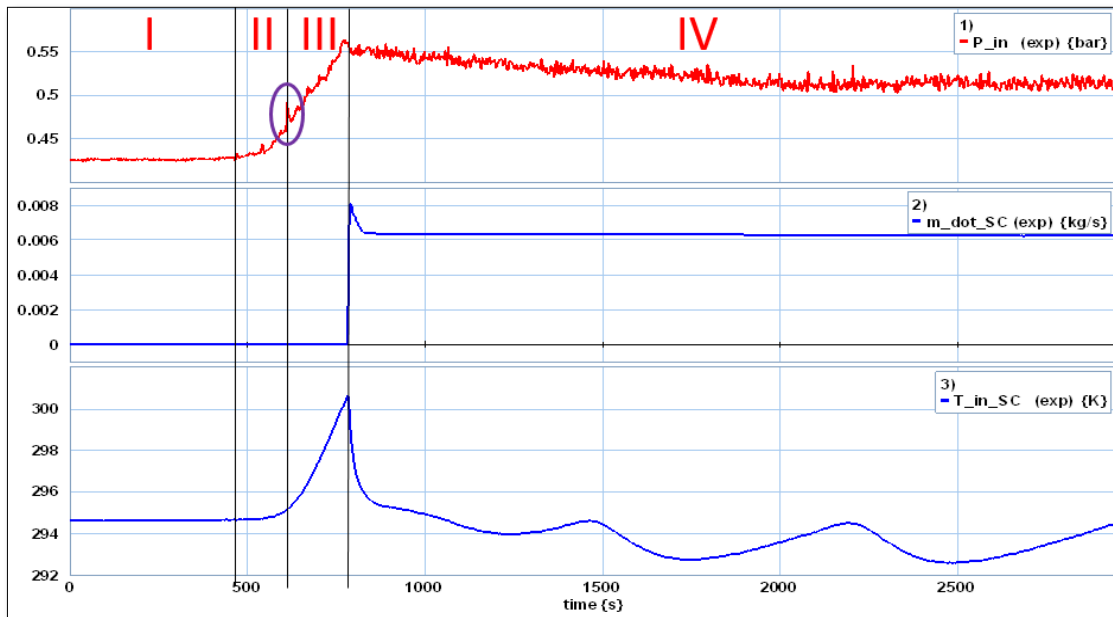


Figure 9: Solicitations applied to the BPC.

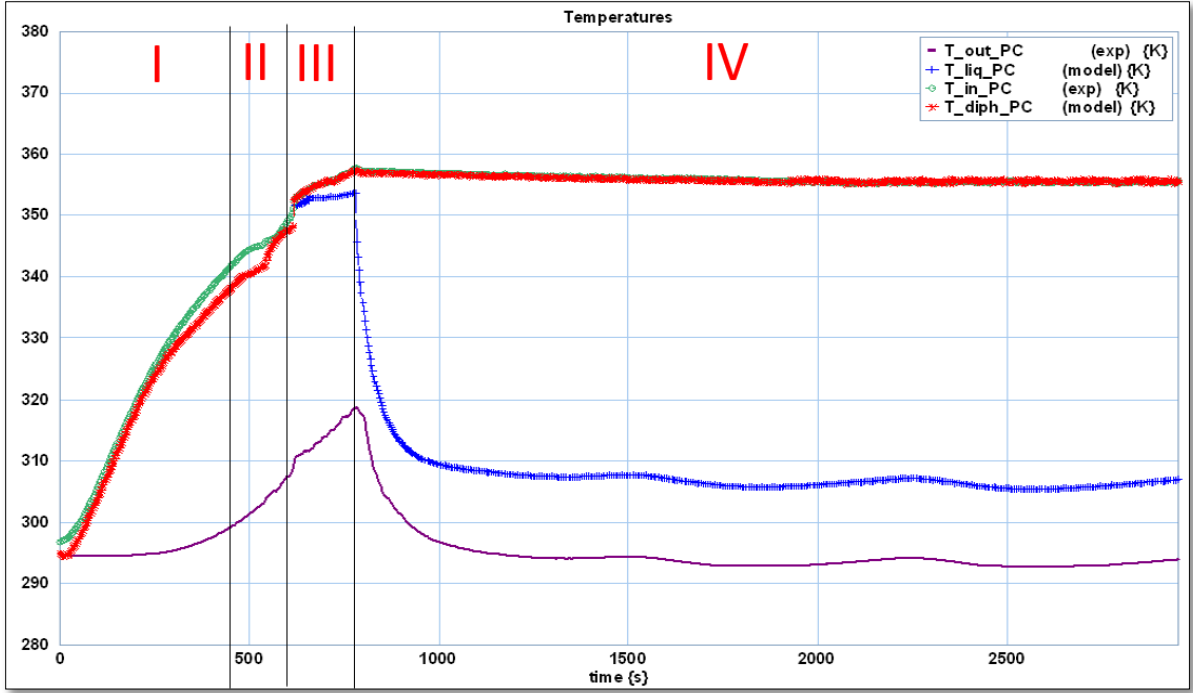


Figure 10: Time evolution of the temperatures calculated in the condenser and measured at the terminals

4.2 Temperatures

Figure 10 gives the time evolution of:

- Liquid temperature, measured at the condenser outlet: T_{out_PC} .
- Liquid temperature, calculated inside the volume V5 of the condenser: T_{liq_PC} .
- Temperature of the mixture, measured at the condenser inlet: T_{in_PC} .
- Temperature of the mixture, calculated inside the volume V6 of the condenser: T_{diph_PC} .

Monophasic intervals (I and II)

First, it should be noted that the temperature of the mixture T_{diph_PC} is considered equal to the liquid temperature T_{liq_PC} as long as the two-phase flow does not appear ($ind_{PC} = 0$). Generally, during this heating phase, it is found that the global dynamic translated by the calculated temperature T_{liq_PC} is very similar to that one measured in the condenser inlet T_{in_PC} .

However, discrepancies remain between the two quantities. Actually the temperature sensor T_{in_PC} , see Figure 8, is mounted at the input port of the condenser, while the liquid

temperature T_{liq_PC} is calculated in the primary circuit represented by the volume V5, see Figure 5. This difference is then justified by the temperature gradient existing between the measurement point and the calculation volume.

The temperature difference between the condenser outlet T_{out_PC} and T_{liq_PC} can be explained:

- The temperature sensor is positioned on the outer wall of the pipe connected to the condenser outlet. Then, the temperature gradient in the wall is not negligible.
- The sensor is not thermally insulated from the outside. Therefore, it is obvious that the measurement of this temperature is disturbed by convective heat exchange with ambient air.
- Finally, the sensor used has an accuracy of $1.5^\circ C$.

Diphasic intervals (III and IV)

Once the two-phase behavior occurs, the condenser model is treated as two volumes, separated by a condensation interface, see

Figure 5. This justifies the presence of two calculated temperatures: $T_{\text{diph_PC}}$ and $T_{\text{liq_PC}}$.

In two-phase state, water is saturated, so its temperature is a function of the pressure. However, the pressure at the inlet of the condenser, just in front of the temperature sensor $T_{\text{in_PC}}$, is close to that prevailing inside the condenser (volume V6 of Figure 5), the small difference is due to low pressure drop. Therefore, the measured temperature of the diphasic mixture $T_{\text{in_PC}}$ and the calculated temperature $T_{\text{diph_PC}}$ are meant to be very close, which is found by the transient model

(red curve compared to the green one in Figure 10).

As to the liquid temperature $T_{\text{liq_PC}}$, the thermal dynamics translated by the temperature sensor is retrieved. However, the discrepancies are due to the sensor position. Also, once the fluid in the secondary circuit flows, the model predicts a decrease of the liquid temperature $T_{\text{liq_PC}}$ with a correctly estimated slope. Finally, required time to reach the second steady regime seems to be the same for the model as for the experiment.

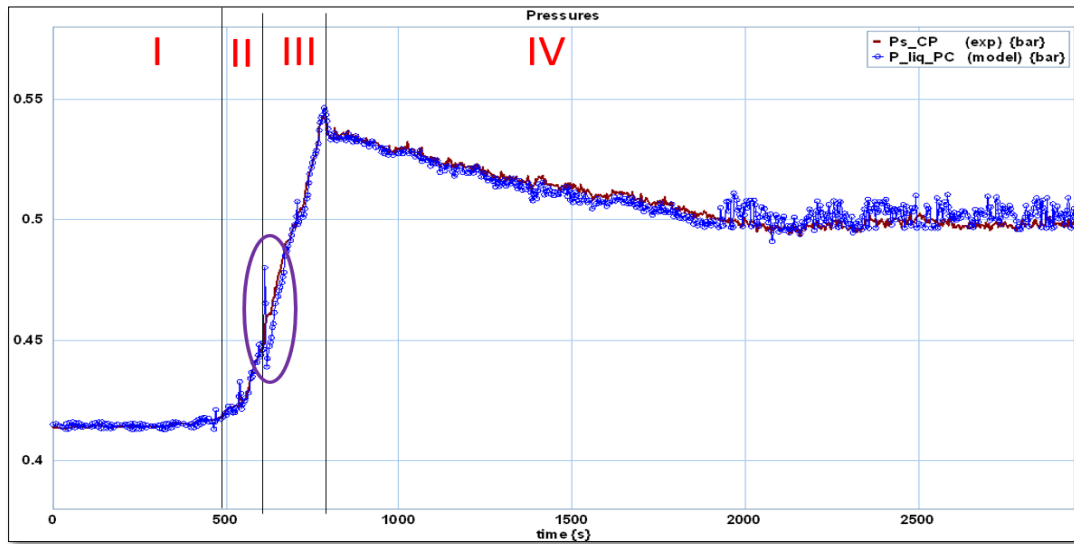


Figure 11: Time evolution of the liquid pressures, calculated and measured in the BPC.

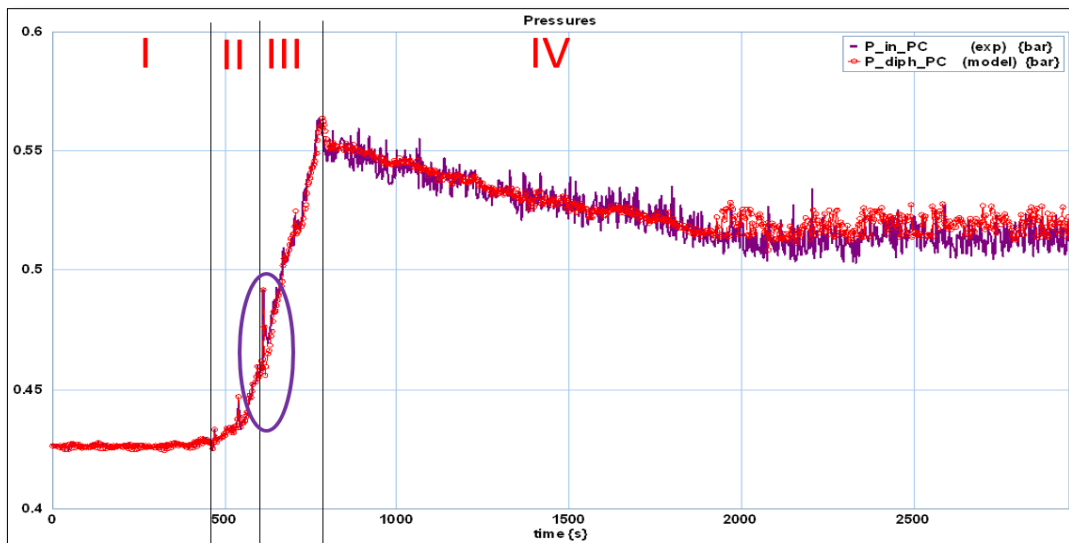


Figure 12: Time evolution of the two-phase mixture pressures, calculated and measured in the BPC.

4.3 Fluid pressure in the primary circuit

The time evolutions of the liquid pressure, calculated P_{liq_PC} and measured P_{out_PC} are shown in Figure 11. While the pressure of the diphasic mixture, calculated P_{diph_PC} and measured P_{in_PC} are plotted in Figure 12.

Pressures P_{liq_PC} and P_{diph_PC} are well transcribed by the transient model in both states: steady and transient which highlights the ability of the model to predict, with good accuracy, the transition from single-phase to two-phase regime. This transition is controlled using the phase indicator introduced in paragraph (3.5).

Result of the validation test

Comparison between simulation results and experimental data reveals that the dynamic model is able to predict, with good agreement, the thermo hydraulic dynamic of the brazed plate heat exchanger. Despite the modeling simplification hypotheses, the model does not react as a low pass filter and allows the identification of unsteady phenomena.

5 Conclusion

The article starts with a succinct description of the brazed plates condenser. It turns out that the technology based on corrugated plates, gives to the exchanger competitive performances. This is due to its compactness, and efficient heat transfers which explain the wide use of this exchanger, particularly in cooling loop used in space applications and constituting the heart of our works.

In the first section two literature reviews have been conducted. The first one, devoted to study existing models, is used to highlight the originality of the proposed model. The second one allows establishing a synthesis of correlations related to thermal convection coefficient and pressure losses, classically used for the two regimes: monophasic and biphasic.

The second section describes the mathematical model of BPC. The model is based on the bond graph methodology. The modularity of this approach gives the model an evolutionary feature, where it is possible to rewrite equations (for a finer modeling) without changing the original structure. The model is discretized into five volumes and is based on the resolution of mass and energy conservation equations. Finally, modeling of heat exchanges is engaged with a noticeable attention. Indeed, the model is validated using several tests without any experimental recalibration of exchange coefficients.

A final section is devoted to the comparison of the simulation results obtained by the unsteady model with experimental recordings from the test bench. An analysis of modeling results leads to conclude that the model has a good ability to transcribe the time evolution of temperatures and pressures, in both regimes, transitional and permanent.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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Appendix A: Mathematical formalism

Graphical scheme of the pressure calculation:

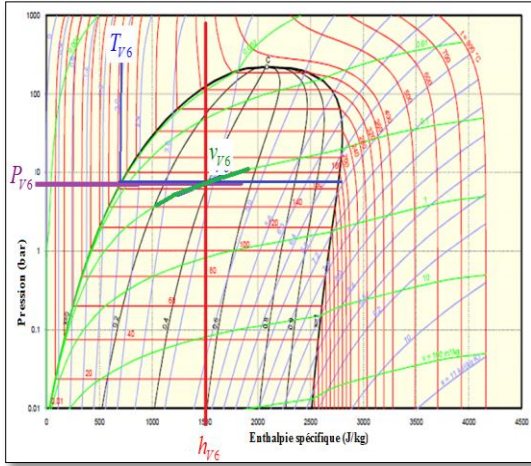


Figure 13: P-h diagram of Mollier.

Theoretical developments

Specific enthalpy of the two-phase mixture:

$$h_{v6}(t) = \frac{H_{acc_v6}(t)}{m_{acc_v6}(t)} \quad (30)$$

Specific volume of the two-phase mixture

$$v_{v6}(t) = \frac{V6}{m_{acc_v6}(t)} \quad (31)$$

Construction of the 12 order polynomial

The specific volume $v_{v6}(t)$ of the mixture can also be expressed as:

$$v_{v6}(t) = x \cdot v''(P_{v6}(t)) + (1-x) \cdot v'(P_{v6}(t)) \quad (32)$$

The specific enthalpy $h_{v6}(t)$ of the mixture can be expressed as:

$$h_{v6}(t) = x \cdot h''(P_{v6}(t)) + (1-x) \cdot h'(P_{v6}(t)) \quad (33)$$

Note about the thermodynamic functions:

v'' , h'' , v' and h' are respectively the specific volume and enthalpy of the vapor portion in the mixture and specific volume and enthalpy of the liquid portion in the mixture. These four thermodynamic functions are all expressed in terms of the pressure $P_{v6}(t)$ using polynomial interpolation. From the two equations (32) and (33), and some other arrangements, we are able to get the 12-order polynomial equation (34). The real, positive and minimum root of this equation is the researched pressure $P_{v6}(t)$

$$\begin{aligned} & v_{v6}(t) \cdot \left[\left(\sum_{i=0}^6 h_i'' \cdot P_{v6}^i(t) \right) - \left(\sum_{i=0}^6 h_i' \cdot P_{v6}^i(t) \right) \right] - \\ & \left\{ \left(\sum_{i=0}^6 v_i'' \cdot P_{v6}^i(t) \right) \cdot \left[h_{v6}(t) - \left(\sum_{i=0}^6 h_i' \cdot P_{v6}^i(t) \right) \right] \right\} \\ & \dots - \left\{ \left(\sum_{i=0}^6 v_i' \cdot P_{v6}^i(t) \right) \cdot \left[\left(\sum_{i=0}^6 h_i'' \cdot P_{v6}^i(t) \right) - h_{v6}(t) \right] \right\} = 0 \end{aligned}$$

Vapor quality of the two-phase flow

The resolution of the equation (34) provides, at each time step, a value of the pressure $P_{v6}(t)$. Based on this value, determining the Vapor quality becomes easy:

$$x(t) = \frac{v_{v6}(t) - \left(\sum_{i=0}^6 v_{i,j}'' \cdot P_{v6}^i(t) \right)}{\left(\sum_{i=0}^6 v_{i,j}' \cdot P_{v6}^i(t) \right) - \left(\sum_{i=0}^6 v_{i,j}'' \cdot P_{v6}^i(t) \right)} \quad (35)$$

Appendix B: friction coefficients

Author	Comments	formula
Muley and Manglik [19]	Valid for water. For Prandtl : $Pr \in [2.4; 4.5]$	$Re \in [2; 200]$ $f = \left[\left(\frac{40.32}{Re} \right)^5 + \left(\frac{8.12}{Re^{0.5}} \right)^5 \right]^{-0.2}$ $Re \geq 1000$ $f = 1.274 \cdot Re^{-0.15}$
Darcy Weisbach	Valid for water	Hagen- Poiseuille for laminar flow $f = \frac{64}{Re}$ Blasius for turbulent flow $f = \frac{0.3164}{Re^{0.25}}$

Table 2: Friction coefficients correlations used for single-phase flow.

Author	Comments	formula
Yan [20]	BPC in a vertical position. BPC composed of two channels	$f \cdot Re^{0.4} \cdot Bo^{-0.5} \cdot P_{red}^{0.8} = 94.75 \cdot Re_{eq}^{-0.0467}$ $Bo = \frac{\dot{m}}{G \cdot \Delta H}$ $Re_{eq} = \frac{G_{eq} \cdot D_h}{\mu_{liq}}$ $G_{eq} = G \cdot \left[1 - x + x \cdot \left(\frac{\rho_{liq}}{\rho_{vap}} \right)^{1/2} \right]$ $G = \frac{\dot{m}}{N_{can} \cdot b \cdot l^*}$
Kuo [21]	CPB composed of three brazed plates	$f = 21500 \cdot Re_{eq}^{-1.14} \cdot Bo^{-0.085}$
Yan [5]	Ge_3 and Ge_4 take into account the geometry of the BPC. Formula valid for: $Re_{eq} \in [300; 4000]$	$f = Ge_3 \cdot Re_{eq}^{Ge_4}$ $Ge_3 = 352.1 \cdot \left(\frac{p_{co}}{D_h} \right)^{4.17} \cdot \left(\frac{\pi}{2} - \beta \right)^{-7.75}$ $Ge_4 = -1.024 \cdot \left(\frac{p_{co}}{D_h} \right)^{0.0925} \cdot \left(\frac{\pi}{2} - \beta \right)^{-1.3}$ $Re_{eq} = \frac{G_{eq} \cdot D_h}{\mu_f}$ $G_{eq} = G \cdot \left[1 - x + x \cdot \left(\frac{\rho_{liq}}{\rho_{vap}} \right)^{1/2} \right]$ $G = \frac{\dot{m}}{N_{can} \cdot b \cdot l^*}$

Table 3: Friction coefficient correlations used for a two-phase flow.

Appendix C: heat transfer coefficients

Author	Comments	formula																																				
<div> <div>Heavner</div> <div>[19]</div> </div>	Valid for water.	<div> $J = \frac{Nu}{Pr^{1/3} \cdot \left(\frac{\mu}{\mu_{wall}} \right)^{0.17}} = b \cdot Re^m$ <table> <tr> <th>Chevron Angle Combination</th> <th>Mean Chevron Angle</th> <th>a</th> <th>n</th> <th>b</th> <th>m</th> </tr> <tr> <td>45°/0°</td> <td>22.5°</td> <td>1.715</td> <td>0.0838</td> <td>0.278</td> <td>0.683</td> </tr> <tr> <td>67°/0°</td> <td>33.5°</td> <td>1.645</td> <td>0.1353</td> <td>0.308</td> <td>0.667</td> </tr> <tr> <td>45°/45°</td> <td>45°</td> <td>0.810</td> <td>0.1405</td> <td>0.195</td> <td>0.692</td> </tr> <tr> <td>67°/45°</td> <td>56°</td> <td>0.649</td> <td>0.1555</td> <td>0.118</td> <td>0.720</td> </tr> <tr> <td>67°/67°</td> <td>67°</td> <td>0.571</td> <td>0.1814</td> <td>0.089</td> <td>0.718</td> </tr> </table> </div>	Chevron Angle Combination	Mean Chevron Angle	a	n	b	m	45°/0°	22.5°	1.715	0.0838	0.278	0.683	67°/0°	33.5°	1.645	0.1353	0.308	0.667	45°/45°	45°	0.810	0.1405	0.195	0.692	67°/45°	56°	0.649	0.1555	0.118	0.720	67°/67°	67°	0.571	0.1814	0.089	0.718
Chevron Angle Combination	Mean Chevron Angle	a	n	b	m																																	
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67°/45°	56°	0.649	0.1555	0.118	0.720																																	
67°/67°	67°	0.571	0.1814	0.089	0.718																																	
<div> <div>Boyko and Kruzhilin</div> <div>(Sánta 2012)</div> </div>	Valid for water.	<div> $h_{conv} = 0.0021 \cdot Re_{liq}^{0.8} \cdot Pr_{liq}^{0.43} \cdot \left(\frac{\lambda_{liq}}{D_h} \right)$ </div>																																				
<div> <div>Muley and Manglik</div> <div>[22]</div> </div>	Valid for water. $Pr_{eau} \in [5;10]$ $Re_{eau} \in [200;1200]$	<div> $h_{conv} = 0.277 \cdot \left(\frac{\lambda_{liq}}{D_h} \right) \cdot Re_{liq}^{0.766} \cdot Pr_{liq}^{0.333}$ </div>																																				
<div> <div>Kim</div> <div>[23]</div> </div>	Valid for water.	<div> $h_{conv} = 0.295 \cdot \left(\frac{\lambda_{liq}}{D_h} \right) \cdot Re^{0.64} \cdot Pr^{0.32} \cdot \left(\frac{\pi}{2} - \beta \right)^{0.09}$ </div>																																				

Table 4: Correlations of heat transfer coefficients used for single-phase flow.

Author	Comments	formula
Nusselt [24]	Laminar film. Condensation controlled by gravity. Vertical surface. Linear temperature gradient in the film. Constant Steam temperature.	$h_{conv} = \left[\frac{\rho_{liq}^2 \cdot g \cdot \Delta H \cdot \lambda_{liq}^3}{4 \cdot \mu_{liq} \cdot l^* \cdot (T_{sat} - T_{wall})} \right]^{1/4}$
Boyko And Kruzhilin [25]	Takes into account water vapor. Valid for Reynolds number: $Re \in [1500;15000]$ Valid for annular flow.	$\alpha_{liq} = 0.021 \cdot Re_{liq}^{0.8} \cdot Pr_{liq}^{0.43} \cdot \frac{\lambda_f}{D_h}$ $h_{conv} = \alpha_{liq} \cdot \left(1 + x \cdot \left(\frac{\rho_{liq}}{\rho_{vap}} - 1 \right) \right)^{0.5}$
Shaw [12]	Condensation controlled by shearing. Valid for water vapor.	$Nu = 0.023 \cdot Re_{liq}^{0.8} \cdot Pr_{liq}^{0.4} \cdot \left[(1-x)^{0.8} + \frac{3.8 \cdot x^{0.76} \cdot (1-x)^{0.04}}{P_{red}^{0.38}} \right]$ $P_{red} = \frac{P}{P_{crit}}$ $h_{conv} = \frac{Nu \cdot \lambda_{liq}}{D_h}$

Table 5: Correlations of heat transfer coefficients used for two-phase flow.

Appendix D: Conduction flows

BG element	heat flow: Φ	Convection coefficient h_{conv}	Exchange surface S_{ech}	ΔT
$R_{Paroi_V1_INT}$	$\dot{H}_{V3/V1}$	Alfa Laval : $Nu = 234 \cdot \lambda_{liq} \cdot Pr^{1/3} \cdot \left[\frac{\rho_{liq} \cdot \Delta P_{V3}}{\mu_{liq}} \right]^{0.3274}$ ΔP_{V3} : Pressure drop in (kPa) μ_{liq} : Viscosity in (cP).	$2 \cdot (l \cdot L)$ The liquid in the SC exchanges with the ambient through the two cover plates.	$T_{V3} - T_{V1}$
$R_{Paroi_V1_EXT}$	$\dot{H}_{V1/amb}$	$h_{ext} = 1.42 \cdot \left(\frac{T_{V1} - T_{amb}}{L/2} \right)^{0.25}$	$2 \cdot (l \cdot L)$	$T_{V1} - T_{amb}$
$R_{Paroi_V2_INT}$	$\dot{H}_{V4/V2}$	$Nu = 234 \cdot \lambda_{liq} \cdot Pr^{1/3} \cdot \left[\frac{\rho_{liq} \cdot \Delta P_{V3}}{\mu_{liq}} \right]^{0.3274}$ ΔP_{V4} : Pressure drop in (kPa) μ_{liq} : viscosity en (cP).	$2 \cdot (l \cdot L)$	$T_{V4} - T_{V2}$
$R_{Paroi_V2_EXT}$	$\dot{H}_{V2/amb}$	$h_{ext} = 1.42 \cdot \left(\frac{T_{V1} - T_{amb}}{L/2} \right)^{0.25}$	$2 \cdot (l \cdot L)$	$T_{V2} - T_{amb}$
R_{Liq_Liq}	$\dot{H}_{V5/V3}$	$h_{conv} = \frac{1}{\frac{e_{wall}}{\lambda_{wall}} + \frac{1}{h_{V5}} + \frac{1}{h_{V3}}}$ Boyko and Kruzhilin : $h_{V5} = 0.021 \cdot Re_{V5}^{0.8} \cdot Pr_{V5}^{0.43} \cdot \left(\frac{\lambda_{liq}}{D_h} \right)$ Alfa Laval : $h_{V3} = 234 \cdot \lambda_{liq} \cdot Pr_{liq}^{1/3} \cdot \left[\frac{\rho_{liq} \cdot \Delta P_{V3}}{\mu_{liq}} \right]^{0.3274}$	$0.0240m^2$ Value given by the manufacturer of BPC used in this work.	$T_{V5} - T_{V3}$
R_{Diph_Liq}	$\dot{H}_{V6/V4}$	$h_{conv} = \frac{1}{\frac{e_{wall}}{\lambda_{wall}} + \frac{1}{h_{V5}} + \frac{1}{h_{V6}}}$ Boyko and Kruzhilin : $\alpha_f = 0.021 \cdot Re_{V6}^{0.8} \cdot Pr_{V6}^{0.43} \cdot \left(\frac{\lambda_{liq}}{D_h} \right)$ $h_{V6} = \alpha_f \cdot \left(1 + x \cdot \left(\frac{\rho_{liq}}{\rho_{vap}} - 1 \right) \right)^{0.5}$	$0.0240m^2$ Value given by the manufacturer of BPC used in this work.	$T_{V6} - T_{V4}$

Table 6: Modeled conduction flows.

Appendix E: Geometric parameters of the BPC

Parameter	notation	value	Parameter	notation	value
Number of plates	N_t	6	Volume of SC	V_{CS}	0.0741 dm3
Plate Thickness	t	0.0003 m	Volume of PC	V_{CP}	0.0494 dm3
Plate Length	L	0.154 m	Exchange surface	S_{ech}	0.048 m².
Plate Height	H	0.0174 m	material		Inox 316
Plate Width	l	0.076 m	Pitch of corrugation	p_{co}	0.001 m
Angle of chevron	β	30°	Total number of channels	$N_{can} = N_t - 1$	5

Table 7: Summary of geometric parameters required for a simulation.