Book Chapter

A Thermal-Hydraulic Coolant Channel Module (CCM) for Single- and Two-Phase Flow

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Abstract

A theoretical ‘drift-flux based thermal-hydraulic mixture-fluid coolant channel model’ is presented. It yields the basis for a corresponding digital ‘Coolant Channel Module (CCM)’. The for this purpose derived ’Separate-Region Mixture Fluid Approach’ should yield an alternative platform to the currently dominant ‘Separate-Phase Models’ where each phase is treated separately. Here, contrary to it, a direct procedure could be established. Its aim is to make it possible to simulate in an as general as possible way the steady state and transient behaviour of characteristic single- and/or (now not necessarily separated) two-phase parameters of fluids flowing within any type of heated or non-heated coolant channels. And has besides the advantage of being able to utilize conservation laws now also the needed constitutive equations can be based directly on corresponding well-proofed correlations for drift flux, friction and heat transfer. And is thus not any longer bounded on sometimes very speculative exchange terms between both two phases but also each of the phases and the wall.

For the case of water/steam flowing along a heated or cooled channel (named ‘basic channel’) a possible sudden transition from single- to two-phase flow and vice versa has the consequence that mathematical discontinuities within the solution procedure have to be expected. To circumvent these difficulties these basic coolant channel (BC) will, in the here presented approach, be assumed to be subdivided into a number of sub-channels (SC-s), each of them being occupied exclusively by only a single or a two-phase flow regime. Hence, after an appropriate spatial discretization and nodalization of the BC (and thus its SC-s) a 'modified finite volume method' (together with a special polygon approximation procedure PAX) could be established. Applied to the conservation laws of thermal-hydraulics (in form of a set of non-linear partial differential equations) and together with corresponding constitutive relations the wanted set of non-linear ordinary differential equations of 1-st order (for each SC and, if extended to the entire channel, BC type) could be derived. Obviously, special attention had to be given now to the varying SC entrance or outlet positions.
Parameters which describe the movement of boiling boundaries or mixture levels along a channel, including even the possibility of SC-s to disappear or be created anew.

The universally applicable code package CCM can thus act as a fundamental element for the simulation of the thermal-hydraulic situations over each type of coolant channels (distinguished by their key numbers) within a complex physical system (Such as different constructions of heat exchangers and steam generators as being in use in both conventional but also nuclear power stations, 1D and 3D nuclear reactor cores etc). By combining all of them with other differential and algebraic equations resulting from the remaining parts of this system this (thermal-hydraulic) part can finally be extended to the demanded overall set of equations representing the theoretical model.

The validity of this alternative approach could be confirmed by a wide range of verification and validation runs, showing very satisfactory results.

**Keywords**

Applied Mathematics; Non-Linear Partial Differential Equations of First Order; Thermal -Hydraulics of Single- and Two-Phase Flow; Separate-Region Mixture-Fluid Model Concept

**1. Introduction**

A large number of thermal-hydraulic (TH) models and, based on them, effective computer codes has been developed and is still in progress, needed for the theoretical and computational description of heat and fluid transport along different types of sometimes very complex technical systems. Their objective is to describe in an adequate way both the steady state and transient behaviour of characteristic key parameters of a single- or two-phase fluid flowing along different types of loops of such a system. Various forms of heated or non-heated coolant channels play an important part within these loops. As, for example, being demanded in nuclear but also conventional power plants.
Obviously many discussions have and will continue to take place among experts about the question which sort of theoretical approach should be chosen for the correct description of such thermal-hydraulic two-phase problems when looking at the wide range of applications. What is thus the most appropriate way to deal with such a special thermal-hydraulic problem?

1.1 Separate-Phase Model Concept

With the derivation of a ‘Separate-Phase Model Concept’ already very early an efficient way was found how to circumvent the above mentioned upcoming difficulties. This by proposing a solution method with the intention to separate the two-phases within the basic equations of such a mixture-flow either partially or even completely from each other. In this case approaches where the phases are treated separately yield a system of 4, 5 or sometimes even 6 equations by splitting each conservation equation into two so-called ‘field equations’. Compared to the four independent parameters characterising the mixture fluid the separate-phase systems demand a much higher number of additional variables and often very speculative assumptions. This has the additional consequence that a number of very complex relations had to be incorporated into the theoretical description of such a module. An enormous amount of CPU-time has to be expended for the solution of the resulting sets of differential and analytical equations within a computer code.

Additionally, according to these assumptions, the interfacial relations both between the (heated or cooled) wall but also between each of the two phases had to be completely rearranged. This raises the difficult question of how to describe in a realistic way the direct heat input into and between the phases and the movement resp. the friction of the phases between them. In such separate-phase approaches this problem is solved by introducing corresponding exchange (=closure) terms between the equations based on special transfer (=closure) laws. Since they can, however, not be based on fundamental laws but also not on experimental measurements this approach requires a significant effort to find an adequate formulation of these exchange terms.
between the phases. It must therefore be recognised that the quality of the basic equations (and especially their boundary conditions) will be intimately related to the (rather artificial and somehow speculative) assumptions with respect to the original conservation laws of the basic 3-equation system and their constitutive equations as well. The problem of a correct description of the interfacial reactions between the phases and the wall remains.

Another problem arises from the fact that special methods have to be foreseen to describe the moving of a boiling boundary or a mixture level (or at least to estimate their ‘condensed’ levels) in such a mixture fluid (See, for example, the ‘Level Tracking’ method in TRAC [1]). Additionally, these methods show often deficiencies in describing extreme situations such as the treatment of single- and two-phase flow at the ceasing of natural circulation, the power situations if decreasing to zero etc. The codes are sometimes very inflexible, especially if they have to provide to a very complex physical system also elements which belong not to the usual class of ‘thermal-hydraulic coolant channels’. These can, for example, be nuclear kinetic considerations, heat transfer out of a fuel rod or through a tube wall, pressure build-up within a compartment, time delay during the movement of an enthalpy front along a downcomer, natural circulation along a closed loop, parallel channels, inner loops etc.

However, the ‘Separate-Phase Models’ have become (despite of these difficulties) increasingly fashionable and dominant in the last decades of thermal-hydraulics as demonstrated by the widely-used codes TRAC [1,2], CATHENA [3], RELAP [4,5], CATHARE [6], ATHLET [7,8].

1.2 Separate-Region Mixture Fluid Model Concept

Obviously among other international institutes also at the Gesellschaft für Anlagen- und Reaktorsicherheit (GRS) at Garching/Munich very early activities within the scope of reactor safety research have been started with the aim to develop thermal-hydraulic models and digital codes which could have the potential to describe in a detailed way the overall transient and
accidental behaviour of fluids flowing along a reactor core but also within the main components of different Nuclear Power Plant (NPP) types. For one of these components, namely the natural circulation U-tube steam generator together with its feedwater and main steam system an own theoretical model and a first version of the digital code UTSG [9] has been derived. Based on the experience of many years of application both at the GRS and a number of other institutes in different countries but also due to the rising demands coming from the safety-related research studies this UTSG theory and code has been continuously extended, ending finally in the establishment of a very satisfactory and mature code version UTSG-2 [10]. This could be used both in a stand-alone way but also as part of more comprehensive transient codes, such as the thermal-hydraulic GRS system code ATHLET [11]. Together with a high level simulation language GCSM (General Control Simulation Module) it could be taken care of a manifold of balance-of-plant (BOP) actions too.

During the research work for the development of an enhanced version of the code UTSG-2 it arose finally the idea to establish an own basic module which makes it possible to simulate the thermal-hydraulic mixture-fluid situation within any geometrical set-up of cooled or heated channels in a very general way. This with the aim to be applicable for any modular construction of complex thermal-hydraulic assemblies of pipes and junctions.

Finally a very special and unique alternative theoretical approach has been found, differing both in its form of application but also in its theoretical background. Thereby each coolant channel should be seen as a (basic) channel (BC) which can, according to their different flow regimes, be subdivided into a number of sub-channels (SC-s). It is clear that each of these SC-s can consist of only two types of flow regimes. A SC with just a single-phase fluid, containing exclusively either sub-cooled water and superheated or even supercritical steam, or a SC with a two-phase mixture. The theoretical considerations of this ‘Separate-Region Approach’ can then (within the class of mixture-fluid models) be appointed to only one of these two regimes. Hence, for each SC type, the ‘classical’ three conservation equations for
mass, energy and momentum can be treated in a direct way (and now without mathematical restrictions). In the case of a SC with mixture flow these basic equations had to be supported by a drift flux correlation (which should, however, also take care of stagnant or counter-current flow situations), yielding an additional relation for the appearing fourth variable, namely the steam mass flow.

The problem of the application of such an approach has thus shifted to the fact that now also varying SC entrance and outlet boundaries (marking the time-varying phase boundary positions) have to be considered. And causing in special situations the additional difficulty that along a channel such a SC can even disappear or be created anew. The solution procedure is based on the fact that after an appropriate nodalization of such a BC (and thus also it’s SC-s) a 'modified finite volume method' (among others based on the Leibniz Integration Rule) had to be derived for the spatial discretization of the fundamental (non-linear) partial differential equations (PDE-s) of first order which represent the basic conservation equations of thermal-hydraulics for each SC.

Additionally, to link within the integration procedure the resulting mean nodal with their nodal boundary function values an adequate quadratic polygon approximation method (PAX) had to be found too. A method representing a central part of the here described ‘Separate-Region Approach’. Together with the suggestion to subdivide a (basic) channel into regions of different flow type and the very thoroughly tested packages for drift flux and single- and two-phase friction factors. Thus yielding an elegant method how to solve this essential thermal-hydraulic problem in a direct way.

The procedure should derive, at the start of a calculation, for each SC (and finally also the entire BC) a set of steady state non-linear algebraic equations and then, for the following transient case, at each time step the wanted set of non-linear ordinary thermal-hydraulic differential equations (ODE-s) of 1-st order. All of them supported with their algebraic constitutive relations.
For more details see section 8, comparing there these two approaches with each of them.

### 1.3 Remarks

The here presented article is part and the continuation of a publication series of four papers being already published in two INTECH and one Applied Mathematics Open Access Books ([12-14] and ([43]). It is, according to a continuous research work, updated to the newest status in this field and concentrated in an advanced and very condensed form mainly to the theoretical background of this basic coolant channel model and to the special (and generally applicable) mathematical methods to solve this sort of resulting system of equations in a corresponding digital module.

### 2. Fundamental Equations (Thermal-Hydraulic Drift-Flux Based Mixture Fluid Approach)

Thermal-hydraulic single-phase or mixture-fluid models for coolant channels or their sub-channels are based on a number of fundamental physical laws, i.e., they obey genuine conservation equations for mass, energy and momentum. They are controlled at single-phase flow conditions by the three independent parameters ‘total mass flow G, temperature T and pressure P’. At two-phase flow conditions one more variable is asked, namely the ‘total and steam mass flow G and G, void fraction α and pressure P’. And are supported by adequate constitutive equations, such as packages for thermo-dynamic and transport properties of water and steam, for single- and two-phase friction coefficients and (in case of two-phase flow) for drift flux.

#### 2.1 Thermal-Hydraulic Conservation Equations

##### 2.1.1 Mass Balance (Single- and two-phase flow)

\[
\frac{\partial}{\partial t} \left[ A(1-\alpha) \rho_w + \alpha \rho_s \right] + \frac{\partial}{\partial z} G = 0
\]  

(1)

containing density terms \( \rho_w \) for sub-cooled or saturated water and \( \rho_s \) for saturated or supercritical steam, the void fraction \( \alpha \)
and the cross flow area $A$ which can (as discussed in section 7) eventually be locally varying along the coolant channel. It determines, after a nodalization, the total mass flow $G=G_W+G_S$ at node outlet in dependence of its node entrance values.

2.1.2 Energy Balance (Single- and two-phase flow)

$$\frac{\partial}{\partial t} \{ A[(1-\alpha)\rho_W h_W+\alpha \rho_S h_S - P] \} + \frac{\partial}{\partial z} [G_W h_W+G_S h_S] = q_{TWL} \begin{equation} = U_{TW} q_{TWF} = A q_D \end{equation}$$

with the enthalpy terms $h_W$ for sub-cooled or saturated water and $h_S$ for saturated, superheated or even supercritical steam. As boundary values either the ‘linear power $q_{TWL}$’, the ‘heat flux $q_{TWF}$’ along the heated (or cooled) tube wall (with its heated perimeter $U_{TW}$) or the local ‘power density term $q_D$’ (transferred into the coolant channel with its cross section $A$) are demanded to be known. The power terms having a positive sign if they are directed into the coolant (See also sections 6.1, 6.4.2 and 9.1).

2.1.3 Momentum balance (Single- and two-phase flow)

$$\frac{\partial}{\partial t} \left( G_F \right) + \left( \frac{\partial P}{\partial z} \right)_A = \left( \frac{\partial P}{\partial z} \right)_S + \left( \frac{\partial P}{\partial z} \right)_C + \left( \frac{\partial P}{\partial z} \right)_X \begin{equation} \end{equation}$$

describing either the pressure differences (at steady state) or (in the transient case) the change in total mass flux ($G_F=\frac{G}{A}$) along a channel (See section 6.5).

The general pressure gradient ($\frac{\partial P}{\partial z}$) can be determined in dependence of terms for

- mass acceleration

$$\left( \frac{\partial P}{\partial z} \right)_A = - \frac{\partial}{\partial z} (G_F v_W + G_S v_S) \begin{equation} \end{equation}$$

with $v_S$ and $v_W$ denoting steam and water velocities (their definitions see nomenclature)

- static head

$$\left( \frac{\partial P}{\partial z} \right)_S = - \cos(\Phi_{ZG}) g \left[ \alpha \rho_S + (1-\alpha) \rho_W \right] \begin{equation} \end{equation}$$
with $\Phi_{ZG}$ representing the angle between z-axis and flow direction. Hence $\cos(\Phi_{ZG}) = \pm \Delta z_{EL} / \Delta z_L$, with $\Delta z_L$ denoting the nodal length and $\Delta z_{EL}$ the nodal elevation height (having a positive sign at upwards flow).

- the single- and/or two-phase friction relation

$$\left( \frac{dP}{dz} \right)_F = - f_R \frac{G_L |G_L|}{2m_f \rho_f}$$  \hspace{1cm} (6)

with a friction factor derived from corresponding constitutive equations (see section 2.2.2)

and finally

- the direct outside perturbation terms $\left( \frac{dP}{dz} \right)_x$, arising either by starting an external pump or considering a pressure adjustment due to mass exchange between parallel channels.

### 2.2 Constitutive Equations

For the exact description of the steady state and transient behaviour of single- or two-phase fluids a number of mostly empirical constitutive correlations are, besides the above mentioned conservation equations, demanded. To bring a structure into the manifold of existing correlations established by various authors, to find the best fitting ones for the different fields of application and to get a smooth transfer from one to another of them special and effective correlation packages had to be developed. Their validities can be and has been tested out-of-pile by means of adequate driver codes. Obviously, by means of this method improved correlations can easily be incorporated into the existing theory.

#### 2.2.1 Thermodynamic and transport water/steam properties

The different thermodynamic properties for water and steam (and their derivatives with respect to P and T, but also P and h) as demanded by the conservation and constitutive equations have to be determined by applying adequate water/steam tables ([15], [16], [17]). Additionally, since from the constitutive equations
also the time-derivatives of these thermodynamic properties are asked which can be represented as

$$\frac{d}{dt} h(z,t) = \frac{d}{dt} h[T(z,t), P(z,t)] = h^T \frac{d}{dt} T_m(z,t) + h^p \frac{d}{dt} P_m(z,t)$$  (7)

Additionally, specific thermodynamic transport properties such as the ‘dynamic viscosity’ and ‘thermal heat conductivity’ (and thus the ‘Prandl number’) are needed too. This is, for light-water systems, realized in the code package MPP [15] (see also [12]) containing the code packages MPPWS and MPPETA. All of them can be derived on the basis of tables given by [16] and [17].

Obviously, the CCM code is also applicable for other coolant systems (superficial flow, heavy water, gas, oil) if adequate thermodynamic tables for this type of fluids are available.

2.2.2 Single- and two-phase friction factors

The friction factor $f_R$ needed in (6) can, as proposed by [18], in case of single-phase flow be set equal to the Darcy-Weisbach single-phase friction factor. For two-phase the corresponding (Martinelli-Nelson) coefficient flow has to be extended by means of a two-phase multiplier $\Phi_{2PF}$ as recommended by [19].

2.2.3 Drift flux correlation

The additional independent variable $\alpha$ appearing in the conservation equations (1), (2) and (3) for a SC with only two-phase flow demands that the set has to be completed by a further relation. This can be achieved by any two-phase correlation, acting thereby as a ‘bridge’ between $G_S$ and $\alpha$. For example, by applying a slip correlation. However, to take care of stagnant or counter-current flow situations too an effective drift-flux correlation seemed here to be more appropriate. For this purpose an own package has been established, named MDS (see [20, 21, 22 and 23]).

Regarding the different requirements in the application of CCM it turned out that it has many advantages to choose the
Sonnenburg ‘flooding-based full-range’ correlation [20] as applied for MDS. This correlation combines the common drift-flux procedure being formulated by Zuber-Findlay [24] and expanded by Ishii-Mishima [25] and (Ishii [26]) etc. with the modern envelope theory. The correlation in the final package MDS had, however, to be rearranged in such a way that also the special cases of $\alpha \to 0$ or $\alpha \to 1$ are included and that, besides their absolute values and corresponding slopes, also the gradients of the approximation function can be made available for CCM. Additionally, an inverse form had to be installed (needed, for example, for steady state conditions) and, eventually, also considerations with respect to possible entrainment effects must be taken care.

The resulting package MDS presents in case of a vertical non-heated or, for the case of low void fractions, heated wall ($L_{HEATB} = 0$ or $= 1$) a relation for the drift velocity $v_D$ with respect to the void fraction $\alpha$

$$v_D = 1.5 \left[ \frac{v_{WLIM}}{C_0} \left( 1 + C_0 \right)^{\frac{3}{2}} - 1.5 + C_0 \right]$$

with $v_D \to v_{D0}$ if $\alpha \to 0$ and $v_{D0} = -\frac{G_F}{\rho} \frac{D_0 V_0}{W_0}$

(if $L_{HEATB} = 0$) or $v_{D0} = 0$ (if $L_{HEATB} = 1$)

and, independently if the wall is heated or not,

with $v_D \to v_{D1} = 1$ if $\alpha \to 1$

introducing as (dimensionless) coefficient

$$C_0 = \frac{2}{3} \frac{v_{SLIM}}{v_{WLIM}} \frac{1-C_0}{C_0}$$

which means that there are needed

- additional relations for the limit velocities $v_{SLIM}$ and $v_{WLIM}$ (both independent of the total mass flow $G$, a fact which is important for the theory below).

and

- adequate correlations for the phase distribution parameter $C_0$ with their limit values $C_0 \to 0$ or $\to 1$ (if $L_{HEATB} = 1$ or $= 0$) and $\alpha \to 0$ resp. (independent of $L_{HEATB}$) $C_0 \to 1$ if $\alpha \to 1$. 
All of them depend on the input parameters 'system pressure $P$', 'hydraulic diameter $d_{HY}$' (with respect to the wetted surface $A_{TW}$ and its inclination angle $\Phi_{ZG}$), specifications about the geometry type ($L_{GTYP}$) and, for low void fractions, the information whether the channel is heated or not ($L_{HEATB} = 1$ or $= 0$).

Finally, from the drift flux theory above also interrelations between different two-phase parameters can be established (now already in dependence of $G$) by starting from their definition equations. The steam mass flow (or flux) can, for example, be represented by

$$G_s = \frac{\rho''}{\rho'} \alpha \left( C_0 G + A \rho' v_a \right) = A G_{vs} = X G$$

the steam velocity by

$$v_s = \frac{1}{C_{GC}} \left( C_0 v_{w_0} + v_D \right) \quad \text{with} \quad v_{w_0} = \frac{G_F}{\rho'}$$

if introducing the coefficient

$$C_{GC} = 1 - (1 - \frac{\rho''}{\rho'}) \alpha C_0 \rightarrow 1 \quad \text{if} \quad \alpha \rightarrow 0 \quad \text{and}$$

$$\rightarrow \frac{\rho''}{\rho'} \quad \text{if} \quad \alpha \rightarrow 1$$

The relations of all the other characteristic two-phase parameters are given in more detail in [22, 23]. These could be the 'phase distribution parameter $C_0$', the 'water and steam mass flows $G_W$ and $G_S$', the 'drift, water, steam and relative velocities $v_D$, $v_W$, $v_S$ and $v_R$' with special values for $v_S \rightarrow v_{S0}$ (if $\alpha \rightarrow 0$) and $v_W \rightarrow v_{W1}$ (if $\alpha \rightarrow 1$) and eventually the 'steam quality $X$'. Their interrelations are presented, for example, in the corresponding tables. Especially the determination of the steam mass flow gradient

$$G_s^\alpha \rightarrow G_{s0}^{\alpha} = \frac{\rho''}{\rho'} \left( C_{00} G + A \rho' v_{D0} \right) = A \rho'' v_{s0} \quad \text{or}$$

$$\rightarrow 0 \quad \text{if} \quad \alpha \rightarrow 0 \quad \text{for both} \quad L_{HEATD} = 0 \quad \text{or} \quad 1$$

$$\rightarrow G_{s1}^{\alpha} = \frac{\rho''}{\rho'} A \left( 1 + C_{01} \right) \left( G - \rho'' v_{SLIM} \right) = A \rho' v_{w1} \quad \text{if} \quad \alpha \rightarrow 1$$

(13)
plays (as shown, for example, in (57)) an important part, if looking to the special situation that the entrance or outlet positions of a SC can cross a BC node boundary (with $\alpha \rightarrow 0$ or $\rightarrow 1$). This possibility makes the drift-flux package MDS to an indispensable part in the nodalization procedure of the mixture-fluid mass and energy balance.

For some application cases it seems to be preferable to have the possibility to avoid counter-current flow (CCF) situations. This can be achieved by limiting (within the general drift-flux theory) the steam quality value along the $X - \alpha$ curve by a value $X_{\text{LIM}}$ which remains within the range $0 \leq X_{\text{LIM}} \leq 1$. This means that according to (10) also either $v_D$ or $C_0$ are restricted. The phase distribution parameter $C_0$ can, however, be also restricted directly, for example by choosing as limit function to $C_0$

$$C_{\text{0LIM}} = C_{00}^\alpha \alpha + (1 - C_{00}^\alpha) \alpha^2 \quad (\text{if setting } L_{\text{NOCCF}} = 1 \text{ at the input}) \quad (14)$$

(with $C_{00}^\alpha$ representing the slope of $C_0$ at $\alpha = 0$)

In the case of a SC with two-phase flow the total and the steam mass flow terms $G$ and $G_S$ are available at steady state conditions as input terms and can now be taken as the basis for further two-phase considerations. The (nodal) void fraction $\alpha$ itself and thus all the other two-phase parameters ($v_D$, $C_0$) can then be calculated from an inverse (INV) form of the drift-flux correlation (10). For the transient case after applying an integration and then the PAX procedure (as shown later-on in the chapters 6.3 and 4.2) the mean and the boundary node values of void fraction ($\alpha_{Mn}$ and $\alpha_{Nn}$) together with the superheating boundary are produced, allowing now from (10) to determine $G_S$ and thus all the other two-phase parameters too.

3. Coolant Channel Geometry Data (BC and SC-s)

The ‘basic’ coolant channel (BC) will, as sketched in fig.1, be assumed to consists of a tube with an as general as possible geometrical form. For discretization purposes the entire BC (with its total length $z_{\text{BT}} = z_{\text{BA}} - z_{\text{BE}}$) must be subdivided into a number of (not necessarily equidistant) $N_{\text{BT}}$ nodes. Their nodal
positions are \( z_{BE}, z_{Bk} \) (with \( k=1,N_{BT} \)), the elevation heights \( z_{ELBE}, z_{ELk} \), the nodal length \( \Delta z_{Bk}=z_{Bk}-z_{Bk-1} \), the nodal elevations \( \Delta z_{ELBk}=z_{ELBk}-z_{ELBk-1} \), with eventually also locally varying cross flow and average areas \( A_{Bk} \) and \( A_{BMk}=0.5 \left( A_{Bk}+A_{Bk-1} \right) \) and their slopes \( A_{Bk}^z=(A_{Bk}-A_{Bk-1})/\Delta z_{Bk} \), a hydraulic diameter \( d_{HYBk} \) and corresponding nodal volumes \( V_{BMk}=\Delta z_{Bk} A_{BMk} \).

Figure 1: Subdivision of a ‘basic channel (BC)’ into ‘sub-channels (SC-s)’ according to their flow regimes and their discretization

The theoretical considerations will, as channel (BC) can according to their flow regimes and (characterized already pointed-out, take advantage of the fact that, as sketched in fig.1, a ‘basic’ coolant by the logical \( L_{TYPE} \) be subdivided into a number \( N_{SCT} \) of sub-channels (SC-s). Each of these SC-s will be distinguished by their characteristic key numbers \( N_{SC} \). Thereby it has, obviously, to be taken into account that their entrance and outlet SC-s can now have variable entrance and/or outlet positions.

As a consequence of the discretization of the BC each of the sub-channels (SC-s) is then subdivided too, now into a number of \( N_{CT} \) SC nodes. Their geometry data are identical to the corresponding BC values, except, of course, at their entrance and outlet positions. The SC entrance position \( z_{CE} \) and their function
f_{CE} are either identical with the BC entrance values \( z_{BE} \) and \( f_{BE} \) or equal to the outlet values of the SC before. The SC outlet position \( (z_{CA}) \) is either limited by the BC outlet \( (z_{BA}) \) or characterized by the fact that the corresponding outlet function has reached an upper or lower limit \( (f_{LIMCA}) \). This term represents then either a function at the boiling boundary, a mixture level or the start position of a supercritical flow. Such a function follows from the given BC limit values and will, in the case of single-phase flow, be equal to the saturation temperature \( T_{SATCA} \) or saturation enthalpies \( (h' \text{ or } h'') \) if \( L_{TYPE} = 1 \) or 2. In the case of two-phase flow \( (L_{TYPE} = 0) \) it has to be set equal to a void fraction of \( \alpha = 1 \) or \( \alpha = 0 \). The moving SC inlet and outlet positions \( z_{CE} \) and \( z_{CA} \) can (together with their corresponding BC nodes \( N_{BCE} \) and \( N_{BCA} = N_{BCE} + N_{CT} \)) be determined according to the conditions \((z_{BNk-1} < z_{CE} < z_{BNk} \text{ at } k = N_{BCE})\) and \((z_{BNk-1} < z_{CA} < z_{BNk} \text{ at } k = N_{BCA})\). Then also the total number of SC nodes \( (N_{CT} = N_{BCA} - N_{BCE}) \) is given, the connection between \( n \) and \( k \) \((n = k - N_{BCE} \text{ with } n = 1, N_{CT})\), the corresponding positions \( (z_{Nn}, z_{ELCE}, z_{ELNn}) \), their lengths \( (\Delta z_{Nn} = z_{Nn} - z_{Nn-1}) \), elevations \( (\Delta z_{ELNn} = z_{ELNn} - z_{ELNn-1}) \), volumes \( (V_{Mn} = z_{Nn} A_{Mn}) \), nodal boundaries and mean nodal flow areas \( (A_{Nn}, A_{Mn}) \).

4. Mathematical tools needed for the solution of (non-linear) PDE-s of 1-st order along a SC with varying entrance and outlet positions

For the realization of the theoretical thermal-hydraulic model in a corresponding digital module (CCM) a special generally applicable integration procedures had to be derived.

4.1 Spatial discretization of PDE-s of 1-st order (Modified finite element method)

Based on the above presented nodalization the spatial discretization of the fundamental equations (1) to (3) can be performed by applying of a ’modified finite element method’.
This means that if a partial differential equation (PDE) of 1-st order having the general form with respect to a solution function \( f(z,t) \)

\[
\frac{\partial}{\partial t} f(z,t) + \frac{\partial}{\partial z} H[f(z,t)] = R[f(z,t)]
\]  

(15)

is integrated over the length of a SC node three types of discretization elements can be expected:

- Integrating a function \( f = f(z,t) \) over a SC node \( n \) yields the nodal mean function values \( f_{Mn} \),
- integrating over the gradient of a function \( f(z,t) \) yields a difference of functions values \( f_{Nn} - f_{Nn-1} \) at their node boundaries
- and, finally,
  - integrating over a time-derivative of this function \( f(z,t) \) (by applying the 'Leibniz' integration rule)

\[
\frac{z_{Nn}(t)}{z_{Nn-1}(t)} \frac{\partial}{\partial t} f(z,t)dz = \Delta z_{Nn}(t) \frac{d}{dt} f_{Mn}(t) \frac{d}{dt} f_{Mn}(t) - [f_{Nn}(t) - f_{Mn}(t)] \frac{d}{dt} z_{Nn}(t) - [f_{Mn}(t) - f_{Nn-1}(t)] \frac{d}{dt} z_{Nn-1}(t) \quad (n=1, N_{CT})
\]  

(16)

### 4.2 Quadratic polygon approximation procedure PAX

Considering the above described three different types of possible discretization elements for the steady state case the solution of the set of algebraic equations will produce directly function values \( f_{Nn} \) at node boundaries \( z_{Nn} \). On the other hand, in the transient case, the solution of the set of ordinary differential equations will yield the time-derivatives of the mean nodal functions \( f_{Mn} \) and, for the last SC node (if \( z_{CA} < z_{BA} \)), of the SC outlet position \( z_{CA} \) and, after an integration procedure, also their absolute values.

It is obvious that an appropriate method had to be found which can help to establish then, on the basis of these different sets of available input data, also the corresponding stationary mean nodal \( f_{Mn} \) resp. the transient nodal boundary \( f_{Nn} \) function
values. And giving thus to the entire discretization a better ‘fine structure. Besides them also their nodal slopes \( f^s_{Nn} \) and \( f^s_{Mn} \) together with their gradients \( f^z_{Nn} \) are asked. These terms are needed, as to be explained later-on in the sections 6.2 to 6.4, by relations describing situations where the length of a SC node could eventually tend to zero so that slopes have to be replaced by their gradients

\[
f^s_{Nn} = \frac{(f_{Nn} - f_{Nn-1})}{\Delta z_{Nn}} \rightarrow f^z_{CEI} \quad (=\text{input}) \quad \text{or} \quad \rightarrow f^z_{Nn-1} \quad \text{if} \quad \Delta z_{Nn} \rightarrow 0 \quad (17)
\]

(at \( n = N_{CT} \) with \( N_{CT} = \) or > 1)

\[
f^s_{Mn} = 2 \frac{(f_{Mn} - f_{Nn-1})}{\Delta z_{Nn}} \rightarrow f^z_{CEI} \quad (=\text{input}) \quad \text{or} \quad \rightarrow f^z_{Nn-1} \quad \text{if} \quad \Delta z_{Nn} \rightarrow 0 \quad (18)
\]

(at \( n = N_{CT} \) with \( N_{CT} = \) or > 1)

For this purpose a special and, as it turned out, very effective ‘quadratic polygon approximation’ procedure, named 'PAX', had to be developed. A procedure differing considerably from some of the 'separate-phase' methods where, as already pointed out, very often only the collapsed levels of a mixture fluid can be calculated. This is different to the ‘separate-phase’ model approach where mostly a method is applied (called ‘upwind or donor cell differencing scheme’) with the mean parameter values to be shifted (in flow direction) to the node boundaries.

It can be stated that the PAX procedure plays, together with the in (16) presented Leibniz rule, an outstanding part in the development of the resulting ‘mixture-fluid model’, helping, in particular, to solve the difficult task of how to take care of a varying SC boundary (which can eventually cross a BC node boundary) in an appropriate and exact way.

4.2.1 Construction of an adequate approximation function

The PAX procedure is based on the assumption that the solution function \( f(z) \) of a PDE (for example temperature or void fraction) along the SC part of a basic (= BC) coolant channel is split into a number of \( N_{CT} \) nodal SC functions \( f_n(z,t) \). Each of them has then to be approximated by a specially constructed quadratic polygon
which can be represented as \( f(z,t) = f_{Nn-1} + a_{Nn} \Delta z + b_{Nn} (\Delta z)^2 \) with \( f_{Nn} = f(\Delta z = \Delta z_{Nn}, t) \) (Corresponding terms for the mean values, slopes and gradients can then be derived too). Thereby, such an effective and adequate quadratic approximation function along a SC (and thus the corresponding nodes) has to fulfil a number of requirements (and helping thus, as demonstrated already in [12], to determine the needed approximation coefficients \( a_{Nn} \) and \( b_{Nn} \)):

- The node entrance functions \( (f_{Nn-1}) \) must be either equal to the SC entrance function \( (f_{Nn-1} = f_{CE} \text{ if } n = 1) \) or to the outlet function of the node before \( (\text{if } n > 1) \).
- The mean function values \( f_{Mn} \) over each of the SC nodes have to be preserved \( (\text{otherwise the balance equations could be hurt}) \).
- It is obvious that the gradients of the nodal entrance functions must not necessarily be equal to the gradients of the SC node outlets before \( (\text{except for the entrance to the last node, i.e. at } n=N_{CT}) \). However, with the objective to guarantee stable behaviour of the approximated functions \( (\text{for example in order to exclude 'saw tooth-like behaviour' of them}) \) it will, as a speciality of this approach, in an additional assumption be demanded that the outlet gradients of the first \( N_{CT} - 1 \) nodes should be set equal to the slopes between their neighbour mean function values. The entrance gradient of the last node \( (n = N_{CT}) \) should be set either equal to the outlet gradient of the node before \( (\text{if } n = N_{CT} > 1) \) or equal to a given SC input gradient \( (\text{for the special case } n = N_{CT} = 1) \).

This very important and characteristic step of this approach can thus be expressed by

\[
f_{Nn}^Z = \left( \frac{\partial f}{\partial z} \right) = 2 \frac{(f_{Mn+1} - f_{Mn})}{\Delta z_{Nn+1} + \Delta z_{Nn}} \quad (n=1, N_{CT}-1, \text{ if } N_{CT} > 1)
\]

\[
= \frac{2}{\Delta z_{Nn}} \left( 2f_{Mn} - 3f_{Mn} + f_{Nn-1} \right) \quad \text{or} \quad = f_{Nn-1} \quad (19)
\]

(\text{if } n = N_{CT} > 1 \text{ and } \Delta z_{CA} > 0 \text{ or } \approx 0)

with its value at SC entrance

\[
f_{CE}^Z = \frac{2}{\Delta z_{Nn}} \left( 3f_{Mn} - 2f_{CE} - f_{Nn} \right) \quad (\text{if } n = 1 \text{ and } N_{CT} > 1 \text{ or } = 1 \text{ and } \Delta z_{CA} > 0)
\]
\[ f^{z}_{\text{CEI}} \quad \text{(if } n = N_{\text{CT}} = 1 \text{ and } \Delta z_{\text{CA}} \approx 0) \tag{20} \]

Note: The second part of the set of equations above results directly from a rearrangement of the definition equations for the quadratic approximation functions of \( f_{Nn} \) and \( f_{Mn} \) (See, for example, also [12]).

This means, the corresponding approximation function reaches not only over the node \( n \). Its next higher one \((n+1)\) has to be included into the considerations too. Except, of course, for the last node. There (in case of \( N_{\text{CT}} > 1 \)) the quadratic approximation function (and thus also its corresponding approximation coefficients) are valid for both (the last and last but one) node.

The procedure has to take care also of the possibility that a SC can consist of only a single node \((N_{\text{CT}} = 1)\), with the (quadratic) approximation function having then to turn into a straight line. Its slope should (as shown in (20) above) reflect two characteristic aspects. Namely to be, for the case that \( \Delta z_{\text{CA}} \rightarrow \Delta z_{\text{BA}} \), equal to the slope (= gradient) \( f^{z}_{\text{BCA}} \) at the corresponding BC outlet node. And, secondly, for the special case that \( \Delta z_{\text{CA}} \rightarrow 0 \) and thus for situations where during a transient either the first or last SC of a BC starts to disappear or to be created anew (i.e. \( z_{\text{CA}} \rightarrow z_{\text{BE}} \) or \( z_{\text{CE}} \rightarrow z_{\text{BA}} \)), to demand as an additional input to PAX (instead of the now not anymore or not yet available term \( f^{z}_{Mn} \)) the gradient \( f^{z}_{\text{CEI}} \) at SC entrance. Representing thereby either (at single-phase flow conditions) the coolant temperature BC entrance gradient \( T^{z}_{\text{CEI}} \) or (for two-phase flow) the void fraction entrance gradient \( \alpha^{z}_{\text{CEI}} \). Since these parameters are usually not directly available they can, for example, be estimated by combining the mass and energy balance equations at SC entrance in an adequate way (See [10, 33]).

These generally valid assumptions make the PAX procedure very effective (and stable). It is a conclusive onset in this method which helps to smooth the curve, guarantees that the gradients at the upper or lower SC boundary do not show abrupt changes for the case that a SC node boundary cross a BC one, avoids ‘saw-tooth’ like behaviour of the parameters along the coolant channel and has the effect that perturbations at channel entrance do not directly affect corresponding parameters of the upper BC nodes.
4.2.2 Nodal parameters needed as input to PAX

After having solved the basic set of non-linear algebraic equations (as presented later-on in the sections 6.2 to 6.4) it is expected that as input to PAX the following nodal boundary parameters should be provided:

- Geometry data such as the SC entrance \( z_{CE} \) and node boundary positions \( z_{Nn} \) and thus also the SC outlet boundary \( z_{CA} \) (as explained in section 3) determining then in PAX the number of SC nodes \( N_{CT} \),
- the SC entrance function \( f_{N0} = f_{CE} \) and (at least for the special case \( n=N_{CT}=1 \)) its gradient \( f_{CE}^{(2)} \) and finally either
  - the nodal boundary functions \( f_{Nn} \) \( (n=1,N_{CT}) \) with \( f_{CA} = f_{Nn} \) at \( n = N_{CT} \) (Note: \( f_{CA} = f_{LIMCA} \) if \( z_{CA} < z_{BA} \)) for the steady state situation
  or
  - the mean nodal functions \( f_{Mn} \) \( (n=1,N_{CT}) \) with \( f_{CA} = f_{Nn} \) at \( n = N_{CT} \) (Note: \( f_{CA} = f_{LIMCA} \) if \( z_{CA} < z_{BA} \)) for the transient situation.

**Steady state case:**

Here, based on inputs from steady state considerations the procedure PAX yields the wanted mean nodal function values

\[
f_{Mn} = \frac{1}{2} (f_{CA} + f_{CE}) = f_{CE} + \frac{1}{2} \Delta z_{CA} f_{CA}^{(2)}
\]

with \( f_{CA} = f_{LIMCA} \) if \( z_{CA} < z_{BA} \) \( (n = N_{CT}, N_{CT} = 1) \)

\[
= \frac{1}{3} (f_{CA} + 2f_{Nn-1}) + \frac{1}{6} \frac{\Delta z_{CA}}{\Delta z_{CA} + \Delta z_{Nn-1}} (f_{CA} - f_{Nn-2})
\]

with \( f_{CA} = f_{LIMCA} \) if \( z_{CA} < z_{BA} \) \( (n = N_{CT}, N_{CT} > 1) \)

\[
= \frac{1}{3 \Delta z_{Nn+1} + 2 \Delta z_{Nn}} [(\Delta z_{Nn} + \Delta z_{Nn+1})(2f_{Nn} + f_{Nn-1}) - \Delta z_{Nn} f_{Mn+1}]
\]

\( (n = 1, N_{CT} - 1, N_{CT} > 1) \)

These functions can, in turn, then be applied as initial values for the transient case
Transient case:
After the integration of the set of non-linear ordinary differential equations (ODE-s) (as presented in the sections 6.2 to 6.4) it can be expected that as input to PAX the SC outlet position ($z_{CA} = z_{Nn}$) and thus also the total number $N_{CT}$ of SC nodes are directly available. Thereby it has to be distinguished (with regard to the transient situation) between two cases of possible input data sets available for PAX:

If the SC outlet position is identical with the BC outlet ($z_{CA} = z_{BA}$) as input to PAX the mean nodal function values $f_{Mn}$ for all $N_{CT}$ nodes ($n=1,N_{CT}$) are available

- If this known SC outlet position is identical with the BC outlet ($z_{CA} = z_{BA}$) as input to PAX the mean nodal function values $f_{Mn}$ for all $N_{CT}$ nodes ($n=1,N_{CT}$) are available

- If this known SC outlet position is identical with the BC outlet ($z_{CA} = z_{BA}$) as input to PAX the mean nodal function values $f_{Mn}$ for all $N_{CT}$ nodes ($n=1,N_{CT}$) are available

or, on the other hand,

- if (due to the transient conditions) the SC outlet is moving within the BC ($z_{CA} < z_{BA}$) the corresponding SC outlet function is known, since restricted by a limit function, i.e. $f_{nn} = f_{CA} = f_{LIMCA}$ at $n = N_{CT}$. This limit is usually the saturation temperature at single-phase flow resp. void fraction equal to 1 or 0 at mixture flow. The last mean nodal function value $f_{Mn}$ ($n = N_{CT}$) is thus interconnected to the movement of the SC outlet position $z_{CA}$. Hence only one of them is needed to be determined by the integration procedure. Hence it can be expected that besides the first $N_{CT} -1$ mean nodal function values $f_{Mn}$ (at $n=1$, $N_{CT} -1$) instead of the last one now the (transient) outlet position $z_{CA}$ will be directly available from the integration procedure. The missing last SC mean nodal function $f_{Mn}$ (at $n = N_{CT}$ and $z_{CA} < z_{BA}$) can then be determined from a relation as presented already in (21) for the steady state input situation.
Hence, it follows for the transient case if the SC outlet is either moving within the BC

\[ f_{Nn} = \frac{1}{2} \left( 3f_{Mn} - f_{Nn-1} \right) + \frac{1}{2} \left( \frac{\Delta z_{Nn}}{\Delta z_{Nn-1} + \Delta z_{Nn}} \right) \left( f_{Mn+1} - f_{Mn} \right) \]

\[ (n = 1, N_{CT} - 2, N_{CT} > 2 \text{ if } z_{CA} < z_{BA}) \]

\[ = \frac{1}{2} \left( 3f_{Mn} - f_{Nn-1} \right) + \frac{1}{4} \left( \frac{\Delta z_{Nn}}{\Delta z_{CA} + \Delta z_{Nn-1}} \right) \left( f_{LIMCA} - f_{Nn-1} \right) \]

\[ (n = N_{CT} - 1, N_{CT} > 1 \text{ if } z_{CA} < z_{BA}) \]

\[ = f_{LIMCA} \quad (n = N_{CT}, N_{CT} = 1 \text{ or } 1 \text{ and } z_{CA} < z_{BA}) \] (22)

or, if not, for the functions of the last SC within the BC

\[ f_{Nn} = \frac{1}{2} \left( 3f_{Mn} - f_{Nn-1} \right) + \frac{1}{2} \left( \frac{\Delta z_{Nn}}{\Delta z_{Nn-1} + \Delta z_{Nn}} \right) \left( f_{Mn+1} - f_{Mn} \right) \]

\[ (n = 1, N_{CT} - 1, N_{CT} > 1 \text{ if } z_{CA} = z_{BA}) \]

\[ = f_{CA} - \frac{1}{3\Delta z_{CA} + 2\Delta z_{Nn-1}} \left[ 2(\Delta z_{CA} + \Delta z_{Nn-1})(3f_{Mn} - 2f_{Nn-1}) + \Delta z_{CA} f_{Nn-2} \right] \]

\[ (n = N_{CT} > 1 \text{ if } z_{CA} = z_{BA}) \]

\[ = f_{CA} = 2f_{Mn} - f_{CE} = f_{CE} + \Delta z_{CA} f_{(s)}^{(CA)} \quad (n = N_{CT} = 1 \text{ if } z_{CA} = z_{BA}) \] (23)

**Final results for both cases:**

All these nodal input function values can together with its entrance parameters \( f_{CE} \) and \( z_{CE} \) and the nodal SC positions \( z_{Cn} \) (at \( n = 1, N_{CT} \)) then (in both cases) be applied as basic points for the PAX procedure. Yielding now, after rearranging the definition equations of the approximation function in an adequate way, all the other not directly known nodal SC function parameters.

The slopes and gradients can finally be determined from (17), (18) and (20). They are indispensable for the special finite element method where due to the moving SC entrance or outlet boundaries they can cross the BC boundaries, i.e., their intervals can go to zero ( \( \Delta z_{CE} \) or \( \Delta z_{CA} \rightarrow 0 \)).

The corresponding time-derivative of the last mean node function which is needed for the determination of the SC
boundary time-derivative (see section 6.4) follows (as long as $z_{CA} < z_{BA}$) by differentiating the relation (21b) above

$$\frac{d}{dt} f_{Mn} = f_{PAXCA}^t + f_{PAXCA}^z \frac{d}{dt} z_{CA}$$

(n = N_{CT} if $z_{CA} < z_{BA}$) (24)

with the coefficients

$$f_{PAXCA}^t = \frac{1}{3} (\frac{d}{dt} f_{LIMCA} + 2 \frac{d}{dt} f_{CE}) - \frac{1}{6} (f_{CEI}^z \frac{d}{dt} z_{CE} - \Delta z_{CA} \frac{d}{dt} f_{CEI}^z)$$

(n = N_{CT} = 1 if $z_{CA} < z_{BA}$) (25)

$$f_{PAXCA}^z = \frac{1}{6} f_{CEI}^z$$

or

$$= 0 \text{ (n = N_{CT} = 1 or } > 1 \text{ and } z_{CA} < z_{BA})$$

(26)

The differentials $\frac{d}{dt} f_{Mn-1}$, $\frac{d}{dt} z_{CA}$, $\frac{d}{dt} f_{LIMCA}$ are directly available from CCM and, if $N_{CT} = 2$, the term $\frac{d}{dt} f_{Nn-2} = \frac{d}{dt} f_{CE}$ from input too. For the case that a SC contains more than two nodes only their corresponding mean values are known, the needed term $\frac{d}{dt} f_{Nn-2}$ has thus to be estimated by establishing the time-derivatives of all boundary functions at all nodes below $N_{CT} < 2$.

### 4.2.3 Code package PAX

Based on the above established set of equations a routine PAX had to be developed with the objective to calculate automatically either the nodal mean or nodal boundary values (in case of an either steady state or transient situation). The procedure should allow also determining the gradients and slopes at SC entrance and outlet (and thus also outlet values characterizing the entrance parameters of an eventually subsequent SC). Additionally, contributions needed for the calculation of the time-derivatives of the boiling boundary or mixture level can be gained (See later-on (70) and (71)).

Before incorporating the subroutine into the overall coolant channel module the validity of the presented PAX procedure has been thoroughly tested. By means of a special driver code
(PAXDRI) different characteristic and extreme cases have been calculated. The resulting curves of such a characteristic example are plotted in fig.2. It presents the two approximation curves of an artificially constructed void fraction distribution $f(z) = \alpha(z)$ along a SC with two-phase flow both for the steady state but also transient situation. Both curves (on the basis of $f_{Mn}$ and $f_{Nn}$) should be (and are) identical.

4.2 Decoupling of mass and energy balance from momentum balance equations

Treating the three conservation equations in a direct way would produce a set of ‘stiff’ ODE-s. This is due to elements with fast pressure wave propagation (and thus being responsible for very small time constants). It has the consequence that their solution turns out to be enormously CPU-time consuming.

Figure 2: Approximation function $f(z)$ along a SC for both steady state and transient conditions after applying PAX (Example).
Hence, to avoid this costly procedure CCM has been developed with the aim to decouple the mass and energy from their momentum balance equations. This can be achieved by determining the thermodynamic properties of water and steam in the energy and mass balance equations on the basis of an estimated pressure profile $P(z,t)$. Thereby the pressure difference terms from a recursive (or a prior computational time step) will be added to an eventually time-varying system pressure $P_{SYS}(t)$, known from boundary conditions (See section 6.5). After having solved the two conservation equations for mass and energy (now separately from and not simultaneously with the momentum balance) the different nodal pressure gradient terms can (by the then following momentum balance considerations) be determined according to (4), (5) and (6).

It can additionally be assumed that according to the very fast (acoustical) pressure wave propagation along a coolant channel all the local pressure time-derivatives can be replaced by a given external system pressure time-derivative, i.e.,

$$\frac{d}{dt} P(z,t) \approx \frac{d}{dt} P_{SYS}$$  \hspace{1cm} (27)

By applying the above explained ‘intelligent’ (since physically justified) simplification in CCM the small, practically negligible, error in establishing the thermodynamic properties on the basis of such an estimated pressure profile can be outweighed by the enormous benefit substantiated by two facts:

- Avoidance of the very time-consuming solution of stiff equations,
- the calculation of the mass flow distribution into different channels resulting from pressure balance considerations can, in a recursive way, be adapted already within each integration time step, i.e. there is no need to solve the entire set of differential equations for this purpose (See ‘closed channel’ concept in section 9.3).

5. **BC input data demanded by CCM**

Besides the BC geometry data (as described in section 3) obviously additional mostly BC function values are needed as inputs to CCM. Selected with the intention to apply only easily
available BC values. They will then, within CCM, be automatically translated into corresponding SC values.

5.1 Initial conditions
For the start of the transient calculations adequate steady state parameters have to be available as initial conditions.

5.2 Boundary conditions and (perturbation) parameters at BC entrance and/or outlet:
In the steady state case and especially to get boundary conditions for transient calculations the following input parameters are expected to be known:

- BC power profile. This means that either the nodal heat flux terms \( q_{FBE} \) and \( q_{FBk} \) (at BC entrance and each node \( k (=1, N_{BT}) \))
or
- \( q_{FBE} \) and the nodal power terms \( Q_{BMk} \) are expected to be known, either directly from input or (as explained in section 9.1) by solving the appropriate ‘Fourier heat conduction equation’. From the relation

\[
Q_{BMk} = \frac{1}{2} \Delta z_{Bk} \left( q_{Lbk} + q_{Lbk-1} \right) = \frac{1}{2} \Delta z_{Bk} \left( U_{TWE} q_{FBE} + U_{TWE} q_{FBE-1} \right) = V_{BME} q_{BMk}
\] (28)

with \( q_{LBE} = U_{TWE} q_{FBE} \) and \( q_{DBE} = \frac{U_{TWE}}{A_{BE}} q_{FBE} \) (\( k=1, N_{BT} \))

Hence, also the other BC nodal terms (\( q_{FBk} \) or \( Q_{BMk} \), \( q_{LBk} \) and \( q_{DBk} \)) can be determined [27, 28 and 29].

- For normalization purposes at the starting calculation (i.e., at the steady state situation) as an additional parameter the total nominal (steady state) heat power \( Q_{NOM,0} \) is asked.
- Channel entrance temperature \( T_{BEIN} \) (or enthalpy \( h_{BEIN} \))
- System pressure \( P_{SYS} \) and its time-derivative (\( dP_{SYS}/dt \)), situated at a fixed position either along the BC (entrance, outlet) or even outside of the ensemble. Due to the fast pressure wave propagation each local
pressure time-derivative can then be set equal to the change in system pressure (as described in section 9.3).

- Total mass flow $G_{\text{BEIN}}$ at BC entrance together with pressure terms at BC entrance $P_{\text{BEIN}}$ and outlet $P_{\text{BAIN}}$. These three parameters are needed for steady state considerations (and partially used for normalization purposes). In the transient case only two of them are demanded as input. The third one will be determined automatically by the model. These allows then to distinguish between the situation of an ‘open’ or ‘closed channel’ concept as this will be explained in more detail in section 9.3.

- Steam mass flow $G_{\text{SBEIN}}$ at BC entrance ($=0$ or $= G_{\text{BEIN}}$ at single- or $0 < G_{\text{SBEIN}} < G_{\text{BEIN}}$ at two-phase flow conditions). The corresponding entrance void fraction $\alpha_{\text{BE}}$ will then be determined automatically within the code by applying the inverse drift-flux correlation.

- Eventually needed time-derivatives of such (perturbation) entrance functions can either be expected to be known directly from input or be estimated from their absolute values.

By choosing adequate boundary conditions then also thermal-hydraulic conditions of special situations can be simulated. For example, that of several channel assembles (of nuclear power plants, test loops etc.) which can consist of a complex web of pipes and branches (represented by different BC-s, all of them distinguished by their key numbers KEYBC). Obviously, also the case of an ensemble consisting of inner loops (as this is needed if describing parallel channels in a reactor core) can be treated in an adequate way by applying the concept of a ‘closed channel’ procedure (see section 9.3).

6. Theoretical thermal-hydraulic separate-region approach

6.1 Power profile along a SC (based on corresponding BC input data)
In order to be able to determine automatically the power profile along each of the SC-s too only the corresponding BC power profile, i.e. the nodal BC power $Q_{BMk}$ together with the linear power term $q_{LBE}$ at BC entrance (known from input, as explained in section 5.2) is asked. In connection with the SC outlet position $z_{CA}$ (and thus the length $\Delta z_{CA}$ of its last node). In the transient case (and if $z_{CA} < z_{BA}$) directly known from the integration procedure.

Hence, if assuming linear behaviour of the ‘linear nodal BC power terms’ within the corresponding BC nodes it follows for

‘the nodal linear SC power’ term

$$q_{LNn} = q_{LCE} = q_{LBE} \quad (=\text{BC entrance}) \quad \text{or}
$$

$$= (q_{LCA}) \text{ of the last node of the SC before} \quad (n=0 \text{ if } z_{CE} = \text{ or } > z_{BE})$$

$$= q_{LBk} \quad (n=1, N_{CT} \text{ and } k=n+N_{BCE} \text{ if } L_{FTYPE}=2) \quad \text{or}$$

$$= q_{LCA} = q_{LBk+1} + (q_{LBk} - q_{LBk-1}) \frac{\Delta z_{CA}}{\Delta z_{Bk}} \quad (n=N_{CT} \text{ and } k=N_{BCA} \text{ if } z_{CA} < z_{BA})$$

$$= Q_{Mn} = \frac{1}{2} \Delta z_{Nn}(q_{LNn} + q_{LNn-1}) \quad (n=1, N_{CT})$$

$$= Q_{BMk} - (Q_{MCA}) \text{ of the last node of the SC before} \quad (n=1 \text{ with } k=1+N_{BCE})$$

$$= Q_{BMk} \quad 2, N_{CT}-1 \text{ and } k=n+N_{BCE}(n=)$$

$$= Q_{MCA} = \Delta z_{CA} [q_{LBk-1} + \frac{1}{2} (q_{LBk} - q_{LBk-1}) \frac{\Delta z_{CA}}{\Delta z_{Bk}}] \quad (n=N_{CT} \text{ and } k=N_{BCA} \text{ if } z_{CA} < z_{BA} \text{ and transient conditions})$$

$$= Q_{M0,0} = Q_{MCA,0} \quad (n= N_{CT} \text{ and steady state conditions}) \quad (30)$$

with $Q_{MCA,0}$ occupied in (45) or (63) if $L_{FTYPE} > \text{ or } = 0$

At steady state conditions, the term $Q_{MCA,0}$ can be determined from energy balance considerations ((44) and (63)) so that
Finally $\Delta z_{CA,0}$ could be calculated, in a reverse manner, according to (69).

Finally it follows for the ‘mean nodal’ and ‘nodal boundary SC power density’ terms ($q_{Mn}$ and $q_{Nn}$)

$$q_{Mn} = \frac{Q_{Mn}}{V_{Mn}} \quad \text{and} \quad (\text{independent of the node length})$$

$$= \frac{1}{2A_{Mn}} (q_{LN_{n-1}}+q_{LN_{n}}) \quad (n=1,N_{CT}) \quad (31)$$

$$q_{Nn} = q_{CE} = q_{BE} \quad \text{or} \quad = (q_{CA}) \quad (\text{of the last node of the SC before})$$

$$(n=0 \text{ if } z_{CE} = \text{ or } > z_{BE})$$

$$= 2q_{Mn} - q_{Nn-1} \quad (n=1,N_{CT}) \quad (32)$$

These terms are, since independent of $\Delta z_{Nn}$, very useful for the solution of special equations where $\Delta z_{Nn} \rightarrow 0$ (as demanded later-on by (32) and (49)).

Hence, the ‘total SC power’ can be presented as

$$Q_{CT} = \sum_{n=1}^{N_{CT}} Q_{Mn} \quad (33)$$

6.2 SC with single-phase fluid ($L_{FTYPE} > 0$)

**Transient situation**

The spatial integration of the two PDE-s representing the conservation equations (1) and (2) over a (single-phase) SC node $n$ (and if taking into account the rules from section 4.1, the relations from (7) and (27), the possibility of a locally changing nodal cross flow area along the BC and the fact that eventually $V_{Mn} \rightarrow 0$) yields for the transient case relations for

- the total nodal mass flow

$$G_{Nn} = G_{Nn-1} - V_{Mn}(\rho_{Mn}^T \frac{d}{dt} T_{Mn} + \rho_{Mn}^p \frac{d}{dt} P_{SYS}) + (\rho_{Nn} - \rho_{Mn})A_{Nn} \frac{d}{dt} z_{Nn}$$

$$+ (\rho_{Mn}^T - \rho_{Nn-1}) A_{Nn-1} \frac{d}{dt} z_{Nn-1} \quad (n=1,N_{CT}), L_{FTYPE} > 0 \quad (34)$$

and, if eliminating the term $G_{Nn}$ by inserting from the equation above into the discretized energy balance equation
the time-derivative of the mean nodal coolant temperature:

\[
\frac{d}{dt} T_{Mn} = T^i_{Tn} + T^z_{TCA} \frac{d}{dt} z_{Nn} \quad \text{(n=1, N\_CT and } z_{CA}=z_{BA}) \quad \text{or} \quad (n=1, N\_CT-1 \text{ and } z_{CA}<z_{BA}), L_{\text{FTYPE}}>0
\]

with \( \frac{d}{dt} z_{Nn} = \frac{d}{dt} z_{CA} \) or \( = 0 \) if \( n = N\_CT \) or \( < N\_CT \)

containing the also for the special situation of \( \Delta z_{CA} \rightarrow 0 \) resp. \( V_{Mn} \rightarrow 0 \) valid part

\[
T^i_{Tn} = \frac{1}{P_{Tm}} (Q_{Mn} - Q_{Gn} + q_{Pn} ) + T^z_{TCE} \frac{d}{dt} z_{Nn-1}
\]

\[
= \frac{1}{V_{\text{min}} P_{Tm}} (Q_{Mn} - Q_{GMn} + Q_{PMn}) + T^z_{TCE} \frac{d}{dt} z_{Nn-1}
\]

with \( \frac{d}{dt} z_{Nn-1} = \frac{d}{dt} z_{CE} \) or \( = 0 \) if \( n = 1 \) or \( > 1 \)

and the coefficients

\[
P_{Tm} = \rho_{Mn} h^T_{Mn} C_{TMn}
\]

with \( C_{TMn} = 1 - \frac{\rho_{Mn}}{\rho_{Mn}} \frac{h_{Nn}}{h^T_{Mn}} = 1 - \frac{\rho_{Mn}}{\rho_{Mn}} (T_{Nn} - T_{Mn}) \)

\[
Q_{GMn} = G_{Nn-1} (h_{Nn} - h_{Nn-1}) = V_{Mn} q_{Gn}
\]

with \( q_{Gn} = \frac{G_{Nn-1}}{A_{Mn}} h^{(s)}_{Nn} \)

\[
Q_{PNn} = V_{Mn} q_{Pn}
\]

with \( q_{Pn} = C_{\text{RHPn}} \frac{d}{dt} P_{\text{SYS}} \) and

\[
C_{\text{RHPn}} = 1 - \rho_{Mn} h^p_{Mn} + \rho^p_{Mn} (h_{Nn} - h_{Mn})
\]

\[
q_{z_n} = \frac{1}{2A_{Mn}} \left\{ A_{Mn} \rho_{Mn} h^{(s)}_{Nn} \frac{d}{dt} z_{Nn} \right. + A_{Nn}(\rho_{Mn} h^{(s)}_{Mn} - 2(\rho_{Mn} - \rho_{Nn-1}) h^{(s)}_{Nn}) \left. \frac{d}{dt} z_{Nn-1} \right\}
\]

\[
T^z_{TCE} = \frac{1}{2} [T^{(s)}_{Mn} - 2(1 - \frac{P_{CE}}{P_{Mn}}) T^{(s)}_{Nn}] \frac{A_{CE}}{A_{Mn} C_{TMn}} \quad \text{or}
\]
= T^z_{TCEI} = \frac{q_{CE}}{G_{FCE}h_{CE}^1} \\
(\text{if either } z_{CE} > z_{BE} \text{ or } z_{CE} \approx z_{BE} \text{ at } n = 1 \text{ and } L_{FTYPE} > 0)

T^z_{TCA} = \frac{1}{2} \frac{A_{n}}{A_{MN}} \frac{T^{(s)}_{n}}{T_{n}} \text{ with } T^{(s)}_{n} \rightarrow T^z_{CA} \text{ if } z_{CE} \rightarrow z_{CA} \quad (42)

(n < N_{CT} \text{ and } z_{CA} < z_{BA} \text{ or } n = N_{CT} \text{ and } z_{CA} \approx z_{BA}, L_{FTYPE} > 0)

In the transient case the mean nodal coolant temperature value \( T_{Mn} \) is at the begin of each (intermediate) time step known. This either from steady state considerations (in combination with PAX) or as a result of the integration procedure. Hence, other characteristic parameters needed in the relations above can be determined too. From the PAX procedure it follow also the SC nodal terms \( T^{(s)}_{n} \), \( T^{(s)}_{n} \) and the slopes \( T^{(s)}_{n} \) resp., for the case that \( \Delta Z_{n} \rightarrow 0 \), their gradients. Finally, using the water/steam tables [15] also their nodal enthalpies are fixed.

Among other results also the SC outlet position \( z_{CA} \) is provided by the integration procedure allowing then to determine the total number of SC nodes (\( N_{CT} \)) too. The situation that \( z_{CA} = z_{BT} \) means the SC nodal boundary temperature values have, within the entire BC, not yet reached their limit values (\( T_{\text{LIMNn}}=T_{\text{SATNn}} \)) and \( N_{CT}=N_{BCA} \) with \( N_{BCA}=N_{BT}-N_{BCE} \). Otherwise, if \( z_{CA} < z_{BT} \), this limit is reached (at node n), then \( N_{CT} = n \). Obviously, the procedure above yields also the time-derivative of the SC outlet position moving within this channel (As described in section 6.4).

\textbf{Steady state}

The steady state part of the total nodal mass flow (charaterized by the index 0) follows from the basic non-linear algebraic equation (34) if setting there the time-derivative equal to 0:

\[ G_{Nn,0} = G_{CA,0} = G_{CE,0} = G_{BA,0} = G_{BE,0} \quad (n=1,N_{CT}, L_{FTYPE} >0) \quad (43) \]

Treating (35) in a similar way and multiplying the resulting relation by \( V_{Mn,0} \) yields the steady state nodal temperature resp. corresponding enthalpy terms

\[ h_{Nn,0} = h_{Nn-1,0} + \frac{Q_{Mn,0}}{G_{BE,0}} \leq h'_{Nn,0} \quad \text{or} \quad \geq h''_{Nn,0} \quad (44) \]

(if \( L_{FTYPE} = 1 \text{ or } = 2 \text{ at } n = 1, N_{BT}-N_{BCE} \text{ and } h_{Nn-1,0}= h_{CE,0} \text{ at } n=1)\]
limited by their saturation values. Then, demanded by (30), which results from energy balance considerations, the nodal power term for the last SC node has to obey to the relation

\[ Q_{MCA,0} = Q_{Mn,0} = (h_{CA,0}^{'} - h_{Nn-1,0}) G_{BE,0} \]

(45)

\[ = (h_{Nn-1,0} - h_{CA,0}^{''}) G_{BE,0} \]

(if \( n = N_{CT} < N_{BCA} \) at \( L_{TYPE} = 1 \))

(45)

\[ (if \ n = N_{CT} < N_{BCA} \) at \( L_{TYPE} = 2 \))

Thus, for the steady state case, the number \( N_{CT} \) is fixed too:

\[ N_{CT} = n < N_{BCA} = N_{BT} - N_{BCE} + 1 \]

and

\[ z_{CA,0} (= z_{Nn,0}) < z_{BA} \]

(46)

\[ (if \ h_{Nn,0} = h_{Nn,0}^{'} \at \( L_{TYPE} = 1 \)) \]

\[ = n = N_{BCA} \] and \( z_{CA,0} = z_{BA} \) (if \( h_{Nn,0} < h_{Nn,0}^{'} \at \( L_{TYPE} = 1 \))

with similar relations for the case \( L_{TYPE} = 2 \).

From the resulting steady state enthalpy value \( h_{Nn,0} \) then (by using the thermodynamic water/steam tables) the corresponding coolant temperature value \( T_{Nn,0} \) follows (with \( T_{Nn,0} = T_{SATNn,0} \) if \( n = N_{CT} \) and \( z_{CA} < z_{BA} \)) and, by applying the PAX procedure, their mean nodal temperature and enthalpy values \( T_{Mn,0} \) and \( h_{Mn,0} \), parameters which are needed as start values for the transient calculations. Obviously, due to the non-linearity of the basic steady state equations, this procedure has to be performed in a recursive way.

It can additionally be stated that both the steady state and transient two-phase mass flow parameters get the trivial form

\[ G_{SNn} = G_{SNn,0} = 0 \text{ resp. } G_{WNn} = G_{CE} \text{ and } G_{WNn,0} = G_{BE,0} \]

(47)

\[ G_{SNn} = G_{SNn,0} = G_{BE,0} \text{ resp. } G_{WNn} = G_{WNn,0} = 0 \]

(47)

\[ (n = 1, N_{CT}, \text{ if } L_{TYPE} = 1) \]

\[ (n = 1, N_{CT}, \text{ if } L_{TYPE} = 2) \]

and

\[ \frac{d}{dt} \alpha_{Mn} = 0 , \alpha_{Nn} = \alpha_{Nn,0} = \alpha_{Mn,0} = 0 \text{ or } = 1 \] and

(48)
\[ X_{Nn,0} = \frac{h_{Nn,0} - h'_{Nn,0}}{h_{SWNn,0}} \quad (n=1, N_{CT} \text{ if } L_{FTYPE} = 1 \text{ or } 2) \]

**6.3 SC with Mixture Fluid (L_{FTYPE} = 0)**

The spatial integration of the two PDE-s of the conservation equations (1) and (2) (now over the mixture-phase SC nodes \( n \)) can be performed by applying again the rules from section 4.1, the relations from (7) and (27), considering the possibility of a locally changing nodal cross flow area along the BC) and the fact that eventually \( V_{Mn} \to 0 \). This yields then relations for

- the total nodal mass flow term

\[
G_{Nn} = G_{Nn-1} + V_{Mn} (\rho' - \rho'') h_{Mn} \left( \frac{d}{dt} \alpha_{Mn} - \alpha_{Gp}^{t} - \alpha_{Gz}^{t} \right) \quad (n=1, N_{CT}, L_{FTYPE} = 0)
\]

with the coefficients (neglecting thereby the small differences between mean and nodal saturation thermodynamic values)

\[
\alpha_{Gp}^{t} = \frac{1}{\left(\rho' - \rho''\right)_{Mn}} \left[ (1 - \alpha) \rho / \rho \right]_{Mn} \frac{d}{dt} P_{SYS}
\]

\[
\alpha_{Gz}^{t} = \alpha_{CE} \frac{d}{dt} z_{Nn-1} + \alpha_{CA} \frac{d}{dt} z_{Nn} \quad (n = 1, N_{CT})
\]

with \( \alpha_{Gz}^{t} = \alpha_{Az}^{t} \) of (50)

\[
= \alpha_{CE} \frac{d}{dt} z_{CE} = \frac{1}{2} \frac{A_{CE}}{A_{Mn}} \alpha_{Mn}^{s} \frac{d}{dt} z_{CE}
\]

\[
= 0 \quad (1 < n < N_{CT})
\]

\[
= \alpha_{CA} \frac{d}{dt} z_{CA} = \frac{A_{CA}}{A_{Mn}} \left( \alpha_{CA}^{s} - \frac{1}{2} \alpha_{Mn}^{s} \right) \frac{d}{dt} z_{CA}
\]

\[
(n = N_{CT}, N_{CT} > 1 \text{ and } z_{CA} < z_{BA})
\]

- the mean nodal void fraction time-derivatives

\[
\frac{d}{dt} \alpha_{Mn} = \alpha_{An}^{t} + \alpha_{Az}^{t} \quad \text{with } \alpha_{An}^{t} = \alpha_{As}^{t} - \alpha_{Ap}^{t}
\]

\[
(n=1, N_{CT} \text{ and } z_{CA} = z_{BA}) \quad \text{or} \quad (n=1, N_{CT}-1, N_{CT} > 1 \text{ and } z_{CA} < z_{BA}), \quad L_{FTYPE}=0
\]

and the coefficients

\[
\alpha_{As}^{t} = \alpha_{Aq}^{t} - \alpha_{Aq}^{t} = \frac{1}{\rho_{Mn}^{s}} [q_{Mn} - h_{SWMn} \frac{G_{Mn}^{s}}{A_{Mn}}]
\]
\[
\alpha_{AP_n}^t = C_{RHP_n} \frac{d}{dt} P_{SYS} \quad \text{with} \\
C_{RHP_n} = \frac{1}{(\rho/h_{sw})_{Mn}} \left[ (1-\alpha) \rho^{h'} + \alpha (\rho^{h'} h_{sw} + \rho^{h'} h_{sw}) - 1 \right]_{Mn}
\]
\[
\alpha_{AZ_n}^t = \alpha_{GP_n}^t \quad \text{(see (51))}
\]
\[
G_{SN}^s = \frac{\Delta G_{SN}}{\Delta z_{Mn}} \rightarrow G_{SN}^z = G_{SN}^s \alpha_{N_n}^s
\]

If \( \alpha_{N_n} \rightarrow \alpha_{CE} = 0 \) (at \( n = 1 \)) or \( \alpha_{N_n} \rightarrow \alpha_{CA} = 1 \) (at \( n = N_{CT} \))

At the begin of each (intermediate) time step the mean nodal void fraction values \( \alpha_{Mn} \) are known. This either from steady state considerations (at the start of the transient calculations) or as a result of the integration procedure. Hence, the additional parameters needed in the relations above can be determined too. From the PAX procedure it follow their nodal boundary void fraction terms \( \alpha_{NV}^s \) together with their slopes \( \alpha_{N_n}^s \) and \( \alpha_{Mn}^s \) resp. gradient \( \alpha_{N_n}^z \) and thus, as shown both in section 2.2.3 but also in the tables given in [22, 23], all the other characteristic two-phase parameters (steam, water or relative velocities, steam qualities etc). Obviously, due to the non-linearity of the basic equations the steady state solution procedure has to be performed in a recursive way.

If, in the transient case, the void fraction \( \alpha_{N_n} \) at SC nodal boundary does (within the entire BC) not reach its limit value (\( \alpha_{LIMN_n}=1 \) or 0) the total number of SC nodes is given as \( N_{CT} = N_{BT} - N_{BCE} \) and \( z_{N_n} \) (at \( n=N_{CT} \)) = \( z_{CA} = z_{BT} \). Otherwise, if this limit is reached (at node \( n \)) then \( N_{CT} = n \) and \( \alpha_{N_n}=1 \) (or =0) with \( z_{CA} (< z_{BT}) \) resulting from the integration. Then, from the procedure above also the time-derivative of the boiling boundary, moving within the BC, can be established (as this will be discussed in section 6.4 too).

With regard to the drift-flux package [21] a relation for the steam mass flow gradient (as already derived in (13)) is required. It is based on the form

\[
G_{SN}^{(a)} = A_{CE} v_{S0} \rho^{h'}_{N_n} \quad \text{with} \quad v_{S0} = v_S \quad \text{(at} \quad \alpha_{N_n} = 0) \\
\text{\( (n=1 \) and \( \alpha_{N_n} \rightarrow \alpha_{CE} = 0) \)}
\]
= A_{CA} v_{W1} \rho_{\Delta n} \text{ with } v_{W1} = v_W (\text{at } \alpha_{Nn} = 1) \quad (n=N_{CT} \text{ and } \alpha_{Nn} \rightarrow \alpha_{CA} = 1) \quad (57)

Eliminating in (49) the term $\frac{d}{dt} \alpha_{Mn}$ by inserting from (52) yields a relation between $G_{SNn}$ and $G_{Nn}$

$$G_{Nn} + (\frac{\rho'}{\rho_{\alpha}} - 1)_{Mn} G_{SNn} = G_{Xn} \quad (n=1,N_{CT}, L_{FTYPE} = 0) \quad (58)$$

The resulting ‘auxiliary’ mass flow term $G_{Xn}$ refers only to already known values (for example the power profile or parameters from the node before) and is thus directly available

$$G_{Xn} = G_{Nn-1} + (\frac{\rho'}{\rho_{\alpha}} - 1)_{Mn} (G_{SNn-1} + \frac{Q_{mn}}{\rho_{SWMn}}) - V_{Mn} (\rho' - \rho''')_{Mn} (\alpha_{APn} + \alpha_{GPn}) \quad (n=1,N_{CT}, L_{FTYPE} = 0) \quad (59)$$

A similar relation can be established from the drift flux correlation (10) by taking advantage of the fact that the needed drift velocity $v_{DNn}$ and the phase distribution parameter $C_{\theta Nn}$ are independent from the total mass flow $G_{Nn}$ (and can thus be determined before knowing $G_{Nn}$). This term ($G_{Nn}$) results then by combining (58) and (10)

$$G_{Nn} = \frac{G_{Sn} - (A_{av} \rho' C_{DC})_{\lambda_{Sn}}}{1 + (\alpha C_{DC})_{\lambda_{Sn}}} = \frac{G_{Xn} - G_{SNn}}{C_{SSn}} \quad (n=1,N_{CT}) \quad (60)$$

using the coefficient

$$C_{DCNn} = (\frac{\rho'}{\rho_{\alpha}} - 1)_{Mn} (\frac{\rho'''}{\rho C_{GC}})_{Nn} \quad (61)$$

From the drift flux correlation package [22] and [23] follow then all the other characteristic two-phase parameters. These are especially the nodal steam mass flow $G_{SNn}$ and, eventually, the slope $\alpha_{SNn}$ resp., according to (56), $G_{SNn}$. Then, finally, from (52) (or (49)) the mean nodal void fraction time-derivative $\frac{d}{dt} \alpha_{Mn}$ will result, needed for the next integration step.

Obviously, the mean nodal temperature and enthalpy terms of a mixture flow are equal to their saturation values.
\[ T_{Mn} = T_{\text{SAT}}(P_{Mn}) \text{ resp. } h'_Mn = h'(P_{Mn}) \text{ or } h''_Mn = h''(P_{Mn}) \quad (62) \]

\[ (n = 1, N_{CT} \text{ and } L_{\text{FTYPE}} = 0) \]

and are thus only dependent on their local pressure value.

Relations for the steady state case follow by setting in (49) and (52) (resp. (53)) the time-derivatives equal to 0. For the total mass flow parameters a similar relation as already given for the single-phase flow (see (52)) is valid, yielding \( G_{Nn,0} = G_{BE,0} \).

Hence, one obtains a relation for the (steady state) nodal steam mass flow

\[ G_{SNn,0} = G_{SNn-1,0} + \frac{Q_{Sn,0}}{h_{\text{SWM}s,0}} \leq G_{Nn,0} = G_{BE,0} \quad (63) \]

\[ (n = 1, N_{BT} - N_{BCE} \text{ and } L_{\text{FTYPE}} = 0) \]

if knowing the term \( Q_{MCA,0} = Q_{Mn,0} \) at \( n = N_{CT} \) (see (30)).

Then the total number \( (N_{CT}) \) of SC nodes follows too:

\[ N_{CT} = n < N_{BT} - N_{BCE} \text{ and } z_{CA,0} = z_{Nn,0} < z_{BA} \]

\[ (\text{if } G_{SNn,0} = G_{BE,0} \text{ and } L_{\text{FTYPE}} = 0) \]

\[ = N_{BT} - N_{BCE} \text{ and } z_{CA,0} = z_{BA} \]

\[ (\text{if } G_{SNn,0} < G_{BE,0} \text{ and } L_{\text{FTYPE}} = 0) \quad (64) \]

yielding now also the corresponding steam quality parameters

\[ X_{Nn,0} = \frac{G_{SNn,0}}{G_{Nn,0}} \quad (n = 1, N_{CT} \text{ if } L_{\text{FTYPE}} = 0) \quad (65) \]

The nodal boundary void fraction values \( \alpha_{Nn,0} \) can be determined by applying the inverse drift-flux correlation and the mean nodal void fraction value \( \alpha_{Mn,0} \) from the PAX procedure. All of them are needed as starting values for the transient calculation.
6.4 SC Entrance and Outlet Boundaries

6.4.1 SC entrance position:

The SC entrance position $z_{CE} (= z_{Nn} \text{ at } n=0)$ is (for both the steady state but also transient case) either equal to BC entrance $z_{BE}$ (for the first SC within the BC) or equal to the outlet boundary of the SC before (See also section 3).

6.4.2 SC outlet position:

Steady state:

The SC (steady state) outlet position $z_{CA,0} (= z_{Nn} \text{ at } n= N_{CT})$, i.e. the boiling boundary $z_{BB,0}$ or mixture level $z_{ML,0}$, can for the trivial case of $z_{CA,0} \geq z_{BA}$ be represented as

$$z_{CA,0} = z_{BA} \quad (n = N_{CT} \text{ and } z_{CA,0} = z_{BA}) \quad (66)$$

Otherwise, the length of the last (SC) node ($\Delta z_{CA,0}$) and thus also $z_{CA,0} = z_{Na-1} + \Delta z_{CA,0}$ can be determined if taking into account that the corresponding steady state total number $N_{CT}$ of nodes of a (single- or two-phase) SC is already determined by (46) or (64). The nodal power term $Q_{MCA,0}$ of the last SC node are then given by (45) or (63). Hence, if dividing (30) by $\Delta z_{Bk}q_{LBk-1,0}$ yields an algebraic quadratic equation of the form

$$\frac{1}{2} \frac{q_{LBk,0} - q_{LBk-1,0}}{q_{LBk-1,0}} (\frac{\Delta z_{CA,0}}{\Delta z_{Bk}})^2 + \frac{\Delta z_{CA,0}}{\Delta z_{Bk}} - \frac{Q_{MCA,0}}{q_{LBk,0} q_{LBk-1,0}} = 0 \quad (68)$$

where $(n = N_{CT} \text{ and } k= N_{BCA} \text{ if } N_{CT} = N_{BT})$

yielding finally as solution

$$\Delta z_{CA,0} = \Delta z_{Bk} \quad (n = N_{CT} \text{ and } k= N_{BCA} \text{ if } N_{CT} = N_{BT})$$

$$= \Delta z_{Bk} \frac{q_{LBk-1,0}}{q_{LBk,0}} \left[1 - \sqrt{1 - 2\left(1 - \frac{q_{LBk,0}}{q_{LBk-1,0}}\right) \frac{Q_{CMA,0}}{q_{LBk,0} q_{LBk-1,0}}}ight]$$

$\quad (k= N_{BCA} \text{ if } N_{CT} < N_{BT})$
\[ \frac{Q_{MCA,0}}{q_{LBk,-1,0}} \left[ 1 + \frac{1}{2} \left( 1 - \frac{q_{LBk,0}}{q_{LBk,-1,0}} \right) \frac{Q_{MCA,0}}{\Delta z_{Bk} q_{LBk,-1,0}} \right] \]

if \( q_{LBk,0} \rightarrow q_{LBk,-1,0} > 0 \)

Then, from the relations in section 6.1, also the other characteristic steady state power terms can be calculated.

**Transient situation:**

The transient SC outlet boundary \( z_{CA} \) boundary \( z_{CA} \) (representing the boiling boundary, the mixture or the supercritical level) follows, as already pointed out, directly from the integration procedure. Thus also \( \Delta z_{CA} \) and \( N_{CT} \) are determined. A boundary which can move along the entire BC (and thereby also cross BC node boundaries). A SC can even shrink to a single node (\( N_{CT} = 1 \), start to disappear or to be created anew. Then, if \( \Delta z_{CA} \rightarrow 0 \), in the relations above the slope in the vicinity of such a boundary has to be replaced by a gradient (determined in PAX).

The mean nodal coolant temperature or, if \( L_{FTYPE}=0 \), void fraction of the last SC node is interrelated by the PAX procedure with the locally varying SC outlet boundary \( z_{CA} \). Hence, in a transient situation the time-derivative of only one of these parameters is demanded. The second one follows then after the integration from the PAX procedure.

If combining (in case of single-phase flow) (24) and (35) the wanted relation for the SC boundary time derivative can be expressed by

\[ \frac{d}{dt} z_{CA} = \frac{d}{dt} z_{BB} = \frac{T_{PAXCA}^i - T_{CA}^i}{T_{CA}^i - T_{PAXCA}^i} \quad \text{or} \quad = 0 \]

\[ (n = N_{CT}, z_{CA} < z_{BA} \quad \text{or} \quad z_{CA} = z_{BA} \quad \text{if} \quad L_{FTYPE} > 0) \]

and if taking for the case of a mixture flow (24) and (52) into account

\[ \frac{d}{dt} z_{CA} = \frac{d}{dt} z_{ML} = \frac{\alpha_{PAXCA}^i - \alpha_{ML}^i}{\alpha_{ML}^i - \alpha_{PAXCA}^i} \quad \text{or} \quad = 0 \]

\[ (n = N_{CT}, z_{CA} < z_{BA} \quad \text{or} \quad z_{CA} = z_{BA} \quad \text{if} \quad L_{FTYPE} = 0) \]
If $z_{CA} < z_{BA}$, the corresponding time-derivatives $\frac{d}{dt} T_{Mn}$ or $\frac{d}{dt} \alpha_{Mn}$ of the last SC node (at $n=N_{CT}$) follow by inserting the terms above into (35) or (52). After the integration procedure then the SC outlet boundary $z_{CA}$ (= boiling boundary $z_{BB}$ or mixture level $z_{ML}$) and thus also the total number $N_{CT}$ of SC nodes are given.

### 6.5 Pressure Profile along a SC (and thus also BC)

After having solved the mass and energy balance equations, separately and not simultaneously with the momentum balance, the now exact nodal SC and BC pressure difference terms ($\Delta P_{Nn} = P_{Nn} - P_{Nn-1}$ and $\Delta P_{BNn}$) can be determined for both single- or two-phase flow situations by discretizing the momentum balance equation (13) and, if applying a modified ‘finite element method’, integrating (4) to (6) over the corresponding SC nodes. The total BC pressure difference $\Delta P_{BT} = P_{BA} - P_{BE}$ between BC outlet and entrance follows then from the relation

$$\Delta P_{BT} = \Delta P_{PBT} - \Delta P_{GBT}$$

(with $\Delta P_{GBT,0} = 0$ at steady state conditions)

and the part

$$\Delta P_{PBT} = \Delta P_{SBT} + \Delta P_{ABT} + \Delta P_{XBT} + \Delta P_{FBT} + \Delta P_{DBT}$$

(with $\Delta P_{PBT,0} = \Delta P_{BTIN,0}$ at steady state conditions)

comprising, as described in section 2.1.3, terms from static head ($\Delta P_{SBT}$), mass acceleration ($\Delta P_{ABT}$), wall friction ($\Delta P_{FBT}$) and external pressure accelerations ($\Delta P_{XBT}$ due to pumps or other perturbations from outside) and an (only in the transient situation needed) term, $\Delta P_{GBT}$, being represented as

$$\Delta P_{GBT} = \int_{0}^{z_{BT}} \frac{d}{dt} G_{FB}(z,t) \, dz = z_{BT} \frac{d}{dt} G_{FBMT} \text{ at transient conditions}$$

$$= 0 \text{ at steady state conditions}$$

This last term describes the influence of time-dependent changes in total mass flux along a BC (caused by the direct influence of changing nodal mass fluxes) and can be estimated by introducing
a ‘fictive’ mean mass flux term $G_{FBMT}$ (averaged over the entire BC)

$$G_{FBMT} = \frac{1}{z_{BF}} \int_{0}^{z_{BF}} \frac{d}{dt} G_{FB}(z,t) \, dz$$

$$\approx \frac{1}{z_{BF}} \sum_{n=1}^{N_{CT}} \sum_{N_{CT}} \Delta z_{Nn} G_{FBMn} = \frac{1}{2} \frac{1}{z_{BF}} \sum_{1}^{N_{BT}} \frac{\Delta z_{Bk}}{A_{Bk}} (G_{Bk} + G_{BK-1})$$ (75)

Its time-derivative can then be represented by

$$\frac{d}{dt} G_{FBMT} \approx \frac{G_{FBMT} - G_{FBMTB}}{\Delta t} = (\frac{d}{dt} G_{FBMT})_{at \, t_B}$$

if $\Delta t = t - t_B = 0$ (Index B = begin of time-step)

Looking at the available friction correlations, there arises the problem how to consider correctly contributions from spacers, tube bends, abrupt changes in cross sections etc. as well. The entire friction pressure decrease ($\Delta P_{FBT}$) along a BC can thus never be described in a satisfactory manner solely by analytical expressions. To minimize these uncertainties a further friction term, $\Delta P_{DBT}$, had to be included into these considerations.

Since the total steady state BC pressure difference $\Delta P_{BT,0} = \Delta P_{PBT,0}$ is known from input the corresponding steady state additive pressure difference term $\Delta P_{DBT,0}$ is (according to (73) and (74)) fixed. It seems, however, to be reasonable to treat this term as a ‘friction’ (or at least the sum with $\Delta P_{FBT}$) and not as a ‘driving’ force. Thus it must be demanded that these terms should remain negative. Otherwise, input terms such as the entire pressure difference along the BC or corresponding friction factors have to be adjusted in an adequate way.

Describing the general additive pressure difference term $\Delta P_{DBT}$ as to have the form

$$\Delta P_{DBT} = (f_{FMP,0} - 1) \Delta P_{FBT} + \Delta P_{FADD}$$ (77)

means that $\Delta P_{DBT}$ is either supplemented with a direct additive term (index FADD) or the friction part is provided with a
multiplicative factor \((f_{FMP,0} - 1)\). For the additive part it will be assumed to be the \((1 - \varepsilon_{DPZ})\)-th part of the total additional pressure difference term and to be proportional to the square of the total coolant mass flow, e.g., at BC entrance

\[
\Delta P_{FADD} = - f_{ADD} z_{BT} \left( \frac{G_f |G_f|}{\rho d_{BH}} \right)_{BE} = (1 - \varepsilon_{DPZ}) \Delta P_{DBT} \quad \text{and} \quad (78)
\]

\[
= 0 \quad (\text{i.e., } \varepsilon_{DPZ} = 1) \quad \text{if } \Delta P_{DBT,0} > 0
\]

with the input coefficient \(\varepsilon_{DPZ} = \varepsilon_{DPZI}\) governing from outside which of them should prevail.

From the known steady state total ‘additional’ term \(\Delta P_{DBT,0}\) the corresponding additive friction factor \(f_{ADD,0}\) follows then directly from the equation above. From (77) then the multiplicative one

\[
f_{FMP,0} = 1 + \varepsilon_{DPZ} \frac{\Delta P_{DBT,0}}{\Delta P_{FST,0}} \quad \text{(setting } \varepsilon_{DPZ} = 1 \text{ if } \Delta P_{DBT,0} > 0) \quad (79)
\]

There arises now the question how the validity of both correction factors could be expanded to the transient case too. In the here presented approach this is done by assuming that these factors should remain time-independent, i.e., that \(f_{ADD} = f_{ADD,0}\) and \(f_{FMP} = f_{FMP,0}\). This allows finally also to determine the wanted nodal pressure decrease term \(\Delta P_{DBT}\) of (73) for the transient case.

By adding the resulting nodal BC pressure difference terms to the (time-varying) system pressure \(P_{SYS}(t)\), given from outside as boundary condition with respect to a certain position (in- or outside of the BC), then finally also the absolute nodal pressure profile \(P_{BE}\) along the BC can be established (This term is needed at the begin of the next time step within the constitutive equations).

7. Digital Coolant Channel Module CCM

Starting from the above presented ‘drift-flux based mixture-fluid theory’ an universally applicable (1D) thermal-hydraulic coolant channel module, named CCM, could be established. It was derived with the intention to provide the authors of different and sometimes very complex multi-channel thermal-hydraulic codes with a general and easily applicable tool needed for the simulation of the steady state and transient behaviours of the
most important single- and two-phase parameters along any type of heated or cooled coolant channel.

For the realization of the theoretical thermal-hydraulic model in the corresponding (digital) module CCM very specific methods had to be achieved by taking into account the following points:

- The code has to be easily applicable, demanding only a limited amount of directly available input data. It should yield as output all the necessary time-derivatives and constitutive parameters of the basic coolant channels required for the establishment of an overall thermal-hydraulic code.

- It was the intention of CCM to act as a complete system in its own right, requiring only BC (and not SC) related, and thus easily available input parameters (geometry data, initial and boundary conditions, parameters resulting from the integration etc.). The partitioning of BC-s into SC-s is done at the beginning of each recursion or time-step automatically within CCM, so no special actions are required of the user.

- The quality of such a model is very much dependent on the method by which the problem of the varying SC entrance and outlet boundaries can be solved. Especially if they cross BC node boundaries during their movement along a channel. For this purpose a special ‘modified finite element-method’ has been developed which takes advantage of the ‘Leibniz’ rule (16) for integration.

- For the support of the nodalized differential equations along different SC-s a ‘quadratic polygon approximation’ procedure (PAX) was constructed in order to interrelate the mean nodal with the nodal boundary functions. Providing additionally also nodal entrance gradients which are required for the calculation of varying SC entrance and outlet boundaries (See section 4.2).

- Several correlation packages such as, for example, packages for the thermodynamic properties of water and steam, single- and two-phase friction coefficients and drift flux correlations had to be developed and
implemented (See sections 2.2.1 to 2.2.3). And, for the case that the power distribution term along the channel is not directly available also a heat transfer correlation package (See section 9.1).

- Knowing the characteristic parameters at all SC nodes (within a BC) then the single- and two-phase parameters at all node boundaries of the entire BC can be determined. And, in the transient case, also the corresponding time-derivatives of the characteristic averaged parameters of coolant temperatures resp. void fraction over these nodes. This yields a final set of (steady-state) algebraic resp. ODE-s and constitutive equations.

- In order to be able to describe also thermodynamic non-equilibrium situations it can be assumed that each phase is represented by an own with each other interacting BC. For these purpose in the model the possibility of a variable cross flow area along the entire channel had to be considered as well.

Within the CCM procedure two further aspects play an important role. They are, however, not essential for the development of mixture-fluid models but can help enormously to enhance the computational speed and applicability of the resulting code when simulating natural circulation within a complex net of coolant pipes:

- The solution of the energy and mass balance equations at each intermediate time step will be performed independently from momentum balance considerations (Section 4.3). Hence the heavy CPU-time consuming solution of stiff equations can be avoided.

- The decoupling allows then also the introduction of an ‘open’ and ‘closed channel’ concept (see section 9.3). Such a special method can be very helpful in describing complex physical systems with eventually inner loops. As an example the simulation of a 3D compartment by parallel channels see [27, 28].

- It represents thus a valuable and very generally applicable tool for the establishment of complex thermal-hydraulic computer codes. And this by being
able to describe any physical system containing a variety of complicated single-and mixture fluid regions. Which can be represented by a number of different types of (basic) coolant channels. It provides automatically an overall set of nodal non-linear algebraic resp. ordinary differential equations and corresponding constitutive relations needed for each of these sub-channels and thus basic channels. Hence, this direct method can be seen as a real counterpart to the currently preferred and dominant ‘separate-phase models’.

The application of a direct mixture-fluid technique follows a long tradition of research efforts. Ishii [26], a pioneer of two-fluid modelling, states with respect to the application of effective drift-flux correlation packages in thermal-hydraulic models: ‘In view of the limited data base presently available and difficulties associated with detailed measurements in two-phase flow, an advanced mixture-fluid model is probably the most reliable and accurate tool for standard two-phase flow problems’. There is no new knowledge available to indicate that this view is invalid.

For more details about the construction of the module CCM see [13], [29-34] and [43].

8. Comparison between the ‘Separate-Region’ and ‘Separate-Phase’ concepts

The here presented theoretical ‘separate-region’ approach can thus disclose a new way in describing thermal-hydraulic problems. The resulting ‘mixture-fluid’ technique can be regarded as a very appropriate way to circumvent the uncertainties apparent from the separation of the phases in a mixture flow. The starting equations are the direct consequence of the original fundamental physical laws for the conservation of mass, energy and momentum, supported by well-tested heat transfer and single- and two-phase friction correlation packages (and thus avoiding also the sometimes very speculative
derivation of the ‘closure’ terms). In a very comprehensive study [27]) a variety of arguments for the here presented type of approach is given.

Generally, the mixture-fluid approach is in line with Fabic [35] who names three strong points arguing in favour of this type of drift-flux based mixture-fluid models:

- The application of (genuine) constitutive equations have the advantage that these relations are supported by a wealth of test data,
- they do not require unknown or untested closure relations concerning mass, energy and momentum exchange between phases (they are mostly very speculative, influencing thus the reliability of the codes),
- they are much simpler to apply,

and, it can be added

- discontinuities during phase changes can be avoided by having to concentrate to SC-s with one fluid type only and by deriving special solution procedures for the simulation of the movement of the SC entrance and outlet boundaries,
- the possibility to circumvent a set of ‘stiff’ ODE-s saves an enormous amount of CPU time which means that the other parts of the code can be treated in much more detail.

In contrast to the currently very dominant separate-phase models, the existing theoretical inconsistencies in describing a two-phase fluid flowing along a coolant channel if changing between single-phase and two-phase conditions and vice versa can be circumvented in the ‘separate-region’ mixture-fluid approach presented here in a very elegant way. A very unique technique has been established built on the concept of subdividing a basic channel (BC) into different sub-channels (SC-s), thus yielding exact solutions of the basic drift-flux supported conservation equations. This type of approach shows, as discussed in [27], distinct advantages vs. ‘separate phase’ codes, especially if taking into account
the quality of the fundamental equations with basic conservation equations on the one side following directly from physical laws supported by experimentally based constitutive equations vs. split ‘field’ equations on the other side with artificial closure terms,

the special solution methods due to the detailed interpolation procedure from PAX allowing to calculate the exact movement of boiling boundaries and mixture (or dry-out) levels, different to the ‘donor-cell averaging’ methods yielding mostly only ‘condensed’ levels,

the easy replacement of new and improved correlations within the different packages without having to change the basic equations of the theory (for example the complicated exchange terms of a ‘separate-phase’ approach),

the possibility to take advantage of the ‘closed-channel concept’ for an overall system (as explained in the next section), This concept can, for example, be needed for thermal-hydraulic 3D considerations, allowing thus to decouple a characteristic (‘closed’) channel from other parts of a complex system of loops,

the speed of the computation,

the derivation of the theory in close and parallel connection with the establishment of the planned code by incorporating demands coming from both sides,

its easy applicability,

thermal non-equilibrium situations can be taken care too because of the introduction of varying cross sections along the BC into the concept. Two separate phases of such a fluid can then be simulated by two with each other interacting basic channels (for example if injecting sub-cooled water rays into a steam dome).

the maturity of the module, which is continuously enhanced by new application cases.

taking advantage of the considerable effort that has been done in verifying and checking CCM (besides an extensive V & V procedure), with respect to the
applicability and adjustment and also for very extreme situations

- and the benefits coming from the fact that most of the development work for the coolant channel thermal-hydraulics has already been shifted to establishment the here presented module, including the special provisions for extreme situations such as stagnant flow, zero power or zero sub-cooling, test calculations for the verification and validation of the code etc.).

The existence of the resulting widely verified and validated module CCM represents an important basic element for the construction of a variety of other comprehensive thermal-hydraulic models and codes as well. Such models and modules can be needed for the simulation of the steady state and transient behaviour of different types of steam generators, of 3D thermal-hydraulic compartments consisting of a number of parallel channels (reactor cores, VVER steam generators etc.). It shows special advantages in view of the determination of the mass flow distribution into different coolant channels after non-symmetric perturbations (see [27, 28]) , a problem which is far from being solved in many of the newest 3D studies.

It has, however, to be noted that there exist essential differences in applying these two types of approaches. The modular codes (such as TRAC, RELAP, ATHLET etc.) combine the different parts of a complex physical system by means of special input prescriptions. Being thus more susceptible to input errors. In the here presented separate-region approach the very well tested-out module CCM yields only for each compartment the needed characteristic algebraic and differential equations for the overall code (e.g. UTSG-3). Which allows more flexibility in describing a complicated physical system but demands also from the user that he is competent in establishing digital codes.

9. Overall Thermal-Hydraulic Multi-Channel Model and Digital Code

In order to be able to incorporate CCM in an overall thermal-hydraulic multi-channel code the interaction between the
different compartments of the (sometimes very) complex physical system has to be defined. For example the heat transfer by means of boundary conditions.

And it has to be considered how the inner circles within this system work. For this purpose usually three more conservation laws have to be taken into account. They yield the necessary initial and boundary conditions to CCM with CCM providing, vice versa, the feedbacks in return.

The first two laws, namely the volume and mass balance relations, allow then to calculate the transient behaviour of the water (and steam volume) and of the pressure (and thus also system pressure $P_{SYS}$) within a fixed (but also even a changing) overall volume $V_T$.

The third physical law is based on the (trivial) fact that the sum of all pressure decrease terms along a closed loop must be zero. And thus in the transient case the entire pressure difference of at least one of the BC-s within this chain of loops is fixed. This allows then to determine the absolute values of the mass flow profile along the entire loop (for example at least at the entrance to this special BC to determine.

9.1 Power Profile Along a Coolant Channel (Heat Transport into and out of a BC)

The nodal BC heat power terms $Q_{BMk}$ (and the corresponding density $q_{BCE}$ at BC entrance) into or out of the coolant are needed (as explained in section 6.1) as boundary condition for the energy balance equation (2) for each of the channels.

These terms can either be directly available (as this is the case for electrically heated loops) or be provided from other parts of the overall system.

In this later case the nodal power distribution (and the wall temperature $T_{TW}$) have to be determined by solving the adequate Fourier heat conduction equations (a PDE of 2-nd order) at each
BC node. Demanding thereby adequate boundary condition, such as, for example,
\[ q_F = \alpha_{TW} (T_{TW} - T) = \frac{q_{L,TW}}{U_{TW}} = \frac{A}{U_{TW}} q_D \]

for the case of heat transfer out of a fuel rod resp. out or into a U-tube (with respect to a coolant having the temperature T). Such a procedure is, for example, presented in [30, 13] for the case of heat conduction through a U-tube wall).

For this purpose adequate heat transfer coefficients (\( \alpha_{TW} \)) are demanded. This means a method had to be found for getting these coefficients along a coolant channel at different flow regimes. In connection with the development of the UTSG code (and thus also of CCM) an own very comprehensive heat transfer coefficient package, called HETRAC [37], has been established.

Obviously, since the resulting set of heat transfer coefficients can never represent the reality in a correct way the wetted surface of the involved heat transfer elements has, in view of the given nominal total steady state power value \( Q_{NOM,0} \), to be provided with a corresponding correction factor. A factor which will be assumed to remain unchanged also in the transient case.

This classic method is different to a ‘separate-phase’ concept where it has to be assumed that the heat is transferred both directly from the wall to each of the two possible phases but also exchanged between them. There arises then the question how the corresponding heat transfer coefficients for each phase should look like.

### 9.2 Mass and Volume Balance along the Entire Overall System

Calculating, as explained above, the transient behaviour of the water (and steam volume) and of the system pressure \( P_{SYS} \). Together with the local pressure difference terms then also the pressure profile along each corresponding BC can be determined.
9.3 BC Entrance Mass Flow (‘Open and closed channel Concept’)

It can be expected that at steady state conditions for each BC in a consecutive arrangement of BC-s the two pressure entrance and outlet values \( P_{BE} = P_{BEIN}, \ P_{BA} = P_{BAIN} \) and the entrance mass flow \( G_{BE} = G_{BEIN} \) are known from input. The entire friction pressure decrease along a BC can, however, never be described in a satisfactory manner solely by analytical expressions as based only on the available friction correlations. Hence (as explained already in section 6.3) in order to consider correctly contributions from spacers, tube bends, abrupt changes in cross sections etc. as well correction terms \( f_{FMP,0} \) and \( \Delta P_{FADD,0} \) had to be introduced. In the case of parallel channels one of them will be taken as the representative one. Since it can usually be assumed that all these channels will have a common friction behaviour (otherwise a special procedure has to be foreseen), the correction factors for this channel can be assumed to be valid for all the other channels too. This allows then to calculate also the different mass flow distribution into these channels (See for example [29])

In the transient case only two of the three characteristic variables (BC entrance mass flow, BC entrance and outlet pressure) can be expected to be known from input. Hence, in these considerations it will have to be distinguished between two special concepts. Is the mass flow at a BC entrance known as input, either as outside perturbation (resulting from a pump) or as an outlet value from a BC below) or not? Assuming that in both cases the resulting (steady state) correction factors will remain unchanged then in the 1-st (‘normal’) case (called the ‘Open Channel Concept’) the mass flow and pressure development along the channel (and thus also the missing pressure term at ether BC outlet or entrance) can be determined in the usual way (See section 3). In the other case (named ‘Closed Channel Concept’) there has, however, to be found a method how to estimate an appropriate entrance mass flow value (or its time-derivative) by adjusting it to the given now fixed BC pressure difference.
This last case plays an important role in the simulation of the natural circulation behaviour of masses within the closed loops of complex physical systems. Due to the fact that the sum of the entire pressure decrease terms along such a closed circuit must be zero. Then the pressure difference (and thus also the absolute entrance and outlet values) are at least of one of the BC channels are defined. And with special methods the needed coolant mass flow terms can be determined.

Usually, in the common thermal-hydraulic codes (see for example the ‘separate-phase’ approaches) this problem is handled by solving the three (or more) fundamental equations for the entire complex system simultaneously. A procedure which affords mostly immense computational times and costs. In the here presented module (based on a separate treatment of momentum from mass and energy balance) a more elegant method could be established. Thus introducing an additional very efficient aspect into the theory of CCM.

This last and still very provisional method has been applied within the UTSG-3 code [13] for the simulation of the natural-circulation behaviour of the secondary steam generator loop. Similar considerations have been undertaken for a 3D case where the automatic mass flow distribution into different entrances of a set of parallel channels is asked (See e.g. [28, 29]). The experience of such calculations should help to decide which of the different possible procedures should finally be given preference.

The ‘open/closed channel concept’ makes sure that measures with regard to the entire closed loop do not need to be taken into account simultaneously but (for each channel) separately. Its application can be restricted to only one ‘characteristic’ channel of a sequence of channels of a complex loop. This additional tool of CCM can in such cases help to handle the variety of closed loops within a complex physical system in a very comfortable way.
10. Solution procedure

10.1 Steady State

The characteristic steady state parameters are determined in a direct way, i.e. calculated by setting all the time-derivatives equal to zero and solving the resulting set of non-linear algebraic equations for SC-s (as being presented in the sections 6.2, 6.3 and 6.4). Thereby, due to the nonlinearities in the set of the (steady state) constitutive equations a recursive procedure in combination with and controlled by the main program has to be applied until a certain convergence in the solution vectors can be stated. The results are then combined to BC parameters and transferred again back to the main (= calling) program.

The (seemingly trivial) demand that the solution vectors should remain unchanged at the begin of calculation and still zero-power conditions is a very valuable and important check of the quality of the set of transient basic equations.

10.2 Transient Situation

For the transient case, as a result of the integration (performed within the calling program and thus outside of CCM) the solution parameters of the set of ODE-s are transferred after each intermediate time step to CCM. These are (as described in detail also in section 7) mainly the mean nodal SC and thus BC coolant temperatures, mean nodal void fractions and the resulting boiling or superheating boundaries. These last two parameters allow then to subdivide the BC into SC-s number (N_{CT}) of SC nodes. Finally, the needed SC (and thus BC) time-derivatives can then be determined within CCM yielding the corresponding constitutive parameters and the total and nodal length (z_{Ns} and \Delta z_{Nn}) of these SC-s and thus also their total (as described in the sections 6.2 to 6.4) and then transmitted again to the calling program where the integration for the next time step can take place.
11. Natural-Circulation Thermal-Hydraulic U-Tube Steam Generator Code UTSG-3

The construction of such an overall code has, for example, be exercised by establishing a new and advanced version of UTSG, now called UTSG-3, being based, similarly as in the previous code UTSG-2, on the same U-tube system layout (heat transfer through U-tubes, top plenum with the main steam system, downcomer (with feedwater injection) and natural circulation along the secondary loop included). But now replacing, among other essential improvements, the three characteristic channel elements of the code UTSG-2 (i.e. the primary and secondary side of the heat exchange zone and the riser) by adequate CCM modules.

The very successful application of the ‘separate-region coolant channel concept’ within the code combination UTSG-3/CCM demonstrates the ability to find an exact and direct solution for the basic equations of a ‘non-homogeneous drift-flux based thermal-hydraulic mixture-fluid coolant channel model’. And it allows to check the performance and validity of the code package CCM and to verify it. For more details see [31, 33 and 13].

12. Verification and Validation (V & V) Procedures

During the course of development of the different versions of the code combination UTSG-3/CCM the module has gone through appropriate verification and validation (V&V) procedures (with continuous feedbacks being considered in the more and more advanced formulation of the theoretical model).

CCM is (similar as done in the separate-phase models) constructed with the objective to be used only as an element within an overall code. Hence, further V&V steps could be performed only in an indirect way, i.e. in combination with such overall codes. This has been achieved, as already explained in the section above, in a very successful way by means of the
natural-circulation U-tube steam generator code UTSG-3. Thereby the module CCM could take advantage from the experiences been gained in decades of years work with the construction of an effective non-linear one-dimensional theoretical model and, based on it, corresponding digital code UTSG-2 for vertical, natural-circulation U-tube steam generators ([9, 10, 11, 33 and 34], [39, 40]) and now also the new advanced code version UTSG-3 (27, 28).

The good agreement of the test calculations with similar calculations of earlier versions applied to the same transient cases demonstrates that despite of the continuous improvements of the code UTSG and the incorporation of CCM into UTSG-3 the newest and advanced version has still preserved its validity.

A more detailed description over these general V&V measures demonstrated on one characteristic test case can be found in [13].

### 13. Conclusions

The universally applicable coolant channel module CCM has been established with the aim to describe the thermal-hydraulic situation of a coolant flowing up-, horizontal or downwards along any channel with fluids changing between sub-cooled, saturated and supercritical conditions. It must be recognized that it represents a complete system in its own right, which requires only BC-related, and thus easily available input values (geometry data, initial and boundary conditions, resulting parameters from integration). The partitioning of a basic channel into SC-s is done automatically within the module, requiring no special actions on the part of the user. At the end of a time-step the characteristic parameters of all SC-s are transferred to the corresponding BC positions, thus yielding the final set of ODE-s together with the parameters following from the constitutive equations of CCM.

The resulting equations for different channels appearing in a complex physical system can be combined with other sets of algebraic equations and ODE-s coming from additional parts of such a complex model (heat transfer or nuclear kinetics considerations, top plenum, main steam system and downcomer
of a steam generator etc.). The final overall set of ODE-s can then be solved by applying an appropriate time-integration routine [27, 33, 34, 35 and 13].

The enormous efforts already made in the verification and validation of the code UTSG-3, its application in a number of transient calculations at very extreme situations (fast opening of safety valves, dry out of the total channel with SC-s disappearing or created anew) brings the code and thus also CCM to a very mature and (what is important) easily applicable state.

There is, however, not yet enough experience to judge how the potential of the mixture -fluid models and especially of CCM can be expanded to other extreme cases (e.g., water and steam hammer). Is it justified to prefer separate-phase models versus the drift-flux based (and thus non-homogeneous) mixture fluid models? This depends, among other criteria, also on the quality of the special models and their exact derivation.

Considering the arguments presented above it can, however, be stated that in general the here presented ‘Separate-Region Mixture Fluid’ concept offers a very satisfactory alternative to the currently dominant ‘Separate-Phase Model’ concepts.

A very detailed derivation of the in this paper established set of characteristic model equations for both the polygon approximation procedure PAX (as a part of CCM) but also of the Coolant Channel Module CCM is given in [41,42].

### 10. Nomenclature

| Symbol | Definition |
|--------|------------|
| $A_{Bk}$ | BC cross sectional area (at BC node boundary k) |
| $A_{Nn}$, $A_{Mn}$ | SC cross sectional area (at SC node boundary n and its mean value) |
| $A_{TWMn}$ | Surface area of a (single) tube wall along a node n |
| $C$ | Dimensionless constant |
| $C_0$, $C_\alpha^{\alpha_0}$ | Phase distribution parameter and its slope at $\alpha = 0$ |
| $d_{HY}$ | Hydraulic diameter |
| $f(z,t)$, $f_{Nn}$, $f_{Mn}$ | General and nodal (boundary and mean) solution functions |
| $f_{LIMCA}$ | Upper or lower limit of the approx. function $f(z,t)$ |
\[ G = G_W + G_S \]

\[
G_F = \frac{G_A}{A} = v_p \alpha = G_{FW} + G_{FS} \quad \text{kg} \quad \text{sm}^2 \]

\[
h, \ h^p, \ c_p = h^T \quad \frac{j}{kg} \quad \frac{m^3}{kg} \quad \text{Specific enthalpy and its partial derivatives with respect to pressure and temperature} \quad (= \text{specific heat})
\]

\[
h_{SW} = h'' - h', \ h''', \ h' \quad \frac{j}{kg} \quad \text{Latent heat, saturation steam and water enthalpy}
\]

\[
K_{EYBC} \quad - \quad \text{Characteristic key number for each channel (BC) within an overall design}
\]

\[
L_{FTYPE} = 0, 1, 2 \text{ or } 3 \quad - \quad \text{SC with saturated water/steam mixture, sub-cooled water, saturated or supercritical steam}
\]

\[
L_{FTBE} \quad \text{of} \quad 1\text{-st} \quad - \quad L_{FTYPE} \text{ of } 1\text{-st SC within BC}
\]

\[
L_{HEATB} = 0 \text{ or } 1 \quad - \quad \text{Non-heated or heated wall}
\]

\[
L_{NOCCF} = 0 \text{ or } 1 \quad - \quad \text{Eventual CCF situation allowed or not allowed}
\]

\[
N_{BT} = N_{BCA} - N_{BCE} + 1 \quad - \quad \text{Total number of BC nodes}
\]

\[
N_{BCA} = N_{CT} + N_{BCE} - 1, \quad - \quad \text{BC node numbers containing BC outlet or entrance}
\]

\[
N_{BCE} \quad - \quad \text{Total number of SC nodes}
\]

\[
N_{CA}, N_{CE} \quad - \quad \text{BC node number containing SC outlet and entrance and total number of SC nodes}
\]

\[
N_{SC} = N_{SCE}, N_{SCA} \quad - \quad \text{Characteristic number of each SC, setting } N_{SCA} = 1, 2, 3, 0, 1, 2 \text{ or } 3. \text{ Then } N_{SCA} = N_{SCE} + N_{SCT} - 1
\]

\[
P, \ \Delta P_T = P_A - P_E \quad Pa = \frac{j}{m^2} \quad \text{Pressure and pressure difference (in flow direction)}
\]

\[
Q_{BT}, Q_{BMK} \quad W \quad \text{Total and nodal power into BC node k}
\]

\[
q_{BMK} = \frac{U_{BK}}{A_{BMK}} q_{LBMK} \quad \text{Mean nodal BC power density into the fluid}
\]

\[
q_{FBK} = A_{BK} \frac{q_{MK}}{U_{BK}} = q_{LBK} \frac{U_{TWBk}}{U_{TW}} \quad \text{Heat flux from (heated) wall to fluid at BC node k}
\]

\[
q_{LBK} = \frac{Q_{BK}}{A_{BK}} = A_{BL} q_{BK} \quad W \quad \text{Linear power along BC node k}
\]

\[
q_{Mn} = \frac{Q_{Mn}}{V_{Mn}} \quad W \quad \text{Mean nodal SC power density into fluid}
\]

\[
= \frac{1}{2} \pi A_{MN} (q_{LNn} + q_{LNn-1}) \quad (= \text{volumetric \ heat transfer rate})
\]

\[
T, \ t \quad C, \ s \quad \text{Temperature, time}
\]

\[
U_{TW} \quad m \quad \text{Perimeter of a heated (single) tube wall}
\]

\[
V_{Mn} = \frac{1}{2} (A_{Nn} + A_{Nn-1}) \Delta z_{Nn} \quad m^3 \quad \text{Mean nodal SC volume}
\]

\[
v_W = \frac{G_{FW}}{(1-\alpha)P_W} \to v_{W0} \quad m \quad \text{Water velocity}
\]
if $\alpha \to 1$

$$v_S = \frac{G_{FS}}{\alpha_{FS}} \to v_{S1} \quad \text{if } \alpha \to 0 \quad \text{m/s}$$

$$X = \frac{G_s}{c} \quad \text{or} \quad \frac{n-k}{b_{sw}}$$

- Steam velocity

- Steam quality (2- resp. extended to 1-phase flow)

- Local position, SC node length ($z_{Nn-1}=z_{CE}$ at $n=0$)

- BC and SC outlet and entrance positions, total length

- Boiling boundary, mixture level resp. supercritical boundary within a BC

- Void fraction

- Heat transfer coefficient along a BC wall surface

- Nodal differences

- Coefficient controlling the additional friction part

- Abs. roughness of tube wall ($\varepsilon_{TW}/d_{HW}=$rel. value)

- Two-phase multiplier

- Angle between upwards and flow direction

- Density and their partial derivatives with respect to (system) pressure and temperature

- Time constant

- Partial derivative

**Subscripts**

- 0, 0 (=E or BE) Steady state or entrance to SC or BC ($n$ or $k=0$)
- A, E, T (=AE) Outlet, entrance, total (i.e. from outlet to entrance)
- B, S Basic channel or sub-channel (=channel region)
- A, S, F, D, X Acceleration, static head, direct and additional (P=A+S+F+D+X) friction, external pressure differences (in connection with $\Delta P$) and pressure differences due to changes in mass flux
- Mn, BMk Mean values over SC or BC nodes
- Nn, Bk SC or BC node boundaries ($n=0$ or $k=0$: Entrance)
- D Drift
- S, W Steam, water
- P, T Derivative at constant pressure or temperature
- TW Tube wall surface

**Superscripts**

- /, // Saturated water or steam
- P, T Partial derivatives with respect to P or T
- ($G_s$), $\alpha$ Partial derivatives with respect to $G_s$, $\alpha$
- or $z$ (=gradient), slope
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