Numerical Solution of Transonic Wet Steam Flow in Blade-to-Blade Cascade with Non-equilibrium Condensation and Real Thermodynamics

Vladimír Hric¹,a and Jan Halama²,b

1 Department of Technical Mathematics, FME CTU Prague, Karl. ná́m. 13, CZ-12135 Prague 2, Czech Republic

2 Institute of Thermomechanics, AS CR, Dolejškova 5, CZ-18200, Prague 8, Czech Republic

Abstract. We present an engineering approach to mathematical modeling and numerical solution of 2D inviscid transonic flow of wet steam in a steam turbine cascade channel of penultimate stage at rotor tip section in full Eulerian framework. Our flow model consists of the Euler system for the mixture (dry steam + homogeneously dispersed water droplets) and transport equations for moments of droplet number distribution function known as moments. Thermodynamic properties of vapor steam are provided by set of IAPWS equations. For equation of state for vapor phase valid both in superheated and wet (meta-stable) region we adopted recently developed equation in CFD formulation for low pressures provided by Hrubý et al. [9], [8], [10]. For extraction of vapor parameters from the mixture ones we implemented simple relations in polynomial form describing thermodynamic properties of saturated liquid state. Nucleation model is resorting to modified classical nucleation theory. Linear droplet growth model is implemented for calculation of liquid sources. Numerical method is simple: cell-centered finite volume approach, 1st-order AUSM+ scheme for spatial derivatives, symmetrical fractional step method for separation of convection and condensation part, explicit 2-stage 2nd-order Runge-Kutta method for time integration. Geometry of blade profile and experimental results are provided by Bakhtar’s work [22], [23]. Results were obtained for one subsonic inlet/subsonic outlet regime and gave quite reasonable accordance with experiment.

1 Introduction

By the wet steam flow we mean just one of the various regimes of multiphase flow – the flow in which a large number of tiny spherical liquid droplets of various size are homogeneously dispersed in a continuous gas phase consisting only of pure vapor. Steam turbines are mechanical devices that transfer thermal energy from expanding steam to electrical power. It is expected that they will continue to play their prominent role in the ensuing decades in this respect. Nucleation process is accompanied with the irreversible (latent) heat release which results in additional total pressure loss and influences thermodynamic and flow parameters. Rapid expansion of initially superheated dry steam leads to the crossing of equilibrium saturation curve, but the first droplets occur only after certain supercooling is attained (non-equilibrium process). Physical theory, experimental measurements and numerical simulations are more complicated in the presence of the second phase. In practice many nucleation models rely on almost 80 years old Becker-Döring classical nucleation theory which with some added corrections gives in case of steam quite reasonable results. Accurate simulations require real thermodynamic concept even if low pressure conditions prevail, as condensation models are very sensitive to the thermodynamic parameters. Our approach is based on the homogeneous mixture non-equilibrium model which approximates pressure of the mixture by the vapor pressure and neglects slip between phases.

2 Mathematical models

2.1 Mixture model

Equations describing dynamics of fluid flow are based on conservation laws of mass, momentum and total energy in Eulerian frame of reference. We consider that there is no velocity slip between phases and that the phases share the same pressure field. Therefore, instead of balancing conservative properties for both phases separately – so-called two-fluid model [7], we treat both phases as a one mixture – so-called mixture (or single-fluid) model. Detailed assessment of single- and two-fluid model can be found e.g. in [6]. Provided that viscosity, heat conduction and gravitational effects are negligible, governing equations for mixture dynamics can be expressed in the form of inviscid Euler system of equations

\[
\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{u}) = 0, \]

\[
\frac{\partial (\rho \mathbf{u})}{\partial t} + \nabla \cdot (\rho \mathbf{u} \mathbf{u} + I p) = 0, \]

\[
\frac{\partial (\rho E)}{\partial t} + \nabla \cdot (\rho \mathbf{u} H) = 0, \]

where \( \rho \) denotes mixture mass density, \( \mathbf{u} \) is velocity vector, \( E \) is total energy of mixture, \( p \) is static pressure, \( H = E + p/\rho \) is total mixture enthalpy and \( I \) is identity matrix. The system needs closure in the form of vapor phase equation of state for pressure. Benefit of the mixture approach is reduction of computationally demands, but a main drawback is that vapor properties are not directly accessible from the...
mixture ones. For this purpose we extract vapor density and vapor internal energy in the following way

\[ \rho_v = \frac{1 - y}{1/\rho - y/\rho_v}, \quad e_v = \frac{e - y e_\ell}{1 - y}. \]  

(4)

Further, we can consider thermodynamic states of liquid phase to be close to saturation liquid boundary. Therefore, we implemented simple relations in polynomial form with temperature as independent variable for liquid properties like density \( \rho_\ell \), enthalpy \( h_\ell \) and speed of sound \( w_\ell \), such that they fit liquid saturation data. Specific internal energy of liquid phase \( e_\ell \) is considered to comprise bulk part only (energy of surface is not accepted). More elegant and thermodynamically consistent method is outlined in [9]. It is assumed that at lower pressures \( (p < 1 \text{ bar}) \) internal energy stored in liquid phase differs not too much from enthalpy of liquid phase, hence we adopted simplification \( e_\ell = h_\ell \).

### 2.2 Equation of state for vapor phase

Most of CFD codes still uses only perfect gas relation for pressure \( p = R T / \gamma \) and internal energy \( e = \gamma R T / (\gamma - 1) \), sometimes with modified specific gas constant \( R \) and ratio of specific isobaric to specific isochoric heat capacity \( \kappa = c_p / c_v \). However, modeling of thermodynamic states in meta-stable region requires using more general equation of state. State-of-the-art equation of state in CFD formulation is applicable only for lower pressures. The low pressure boundary is determined by the value of entropy \( s \geq 7.5 \text{ kJ kg}^{-1} \text{ K}^{-1} \). Below the entropy limit, deviations from experimental data grow significantly.

### 2.3 Equations for liquid parameters

For liquid density \( \rho_\ell = \rho_i(T) \), liquid enthalpy \( h_\ell = h_i(T) \) and liquid sound speed \( w_\ell = w_i(T) \) we used following approximations of saturation data. For liquid density

\[ \rho_\ell = -2.031 \theta^6 + 3.056 \theta^5 + 3.297 \theta^4 - 3.15 \theta^3 - 18 \theta^2 - 50.99 \theta + 966.1 \]  

(15)

with scaled temperature \( \theta = (T - 362.4)/79.46 \), for liquid enthalpy \( (\approx \text{ liquid internal energy}) \)

\[ h_\ell = \max \left\{ 0, 8.599 e^{-3} T^3 - 9.069 T^2 + 7.36 e^3 T - 1.511 e^6 \right\} \]  

(16)

and for liquid sound of speed

\[ w_\ell = -4.168 \theta^4 + 12.64 \theta^3 - 58.37 \theta^2 + 84.99 \theta + 1529 \]  

(17)

with scaled temperature \( \theta = (T - 386.3)/66.12 \).

### 2.4 Evolution of liquid phase

Evolution of liquid phase can be described by mass conservation law for droplets in phase space \((x,r)\). Instead of expressing balance equation in the general discrete kinetic form which takes interaction between all monomers (clusters containing one molecule) into account, we use its reduced continuum version for balance of droplet number continuous distribution function \( f = f(x,r,t) \). Then, the total number of droplets in unit volume with radii smaller than \( r \) at the location \( x \) and at the time \( t \) is given by integral

\[ \int_0^r f(x,r,t) \, dr. \]

For much more details about droplet size distributions we refer to [1]. This reduction assumes
that droplets with very small number of molecules are neglected. Then, balance equation takes the form

$$\frac{\partial f}{\partial t} + \nabla \cdot (f u) = -\frac{\partial (rf)}{\partial r} + \delta (r - r_c) J.$$  (18)

The first term on the RHS is related to the droplet growth as a result of capturing new molecules from the surrounding vapor and the second source concerns creating of new droplets at the critical radius, $\delta$ is Dirac delta function. De-nucleation source and droplet growth owing to liquid compressibility is not included. Coagulation and segregation of molecules are not respected as well. Because the distribution function $f$ is not known a priori and we do not intend to model its shape, we reduce the balance equation further by its multiplication by $r^{k}$ and subsequent integration \( \int_{0}^{\infty} (\cdot) dr \). With the definition of the k-th moment of the droplet number distribution function

$$\mu_k = \int_{0}^{\infty} r^k f dr$$  (19)

and applying integration by parts in droplet growth source term with boundary conditions

$$f(r = r_c) = 0, \quad \lim_{r \to \infty} f = 0,$$  (20)

we get the efficient so-called moment version of balance equation for moments

$$\frac{\partial \mu_k}{\partial t} + \nabla \cdot (\mu_k u) = k \int_{0}^{\infty} r^{k-1} f dr + r_c^k J.$$  (21)

In our CFD code we utilized only the first four moments, i.e. $k = 0,1,2,3$. This is sufficient in practical applications for basic modeling of polydispersed droplet spectra. In the classical method of moments droplet growth law (equation for $\dot{r}$) cannot be used in the general form with respect to radial coordinate $r$. We decided to provide its linear approximation in the vicinity of average droplet radius, hence $\dot{r} = a + br$, where constants $a, b$ have to be determined properly and are functions of vapor properties only. For average droplet radius we used Hill’s definition $\bar{r} = \sqrt{\mu_2/\mu_0}$. With the mentioned closure method we can rewrite moment equations as follows

$$\frac{\partial \mu_k}{\partial t} + \nabla \cdot (\mu_k u) = k (a \mu_{k-1} + b \mu_k) + r_c^k J.$$  (22)

where $k = 0 : 3$. The last equation ($k = 3$) can be transformed to the equation for wetness factor by its multiplying by factor $4\pi r^2 / 3$ and using definition for wetness fraction $\rho_y = 4\pi r^2 \mu_3 / 3$ and utilizing $\rho_I = \text{const.}$

$$\frac{\partial (\rho y)}{\partial t} + \nabla \cdot (\rho y u) = 4\pi \rho_I \left( 3\mu_2 + r_c^2 J \right) + 3b \rho y.$$  (23)

Moments of droplet number distribution function $\mu_k$ are occasionally expressed by means of so-called liquid moments $Q_k$ and holds

$$\mu_k = \rho Q_k.$$  (24)

### 2.5 Creating of new water droplets

Rapid expansion of initially superheated steam into the metastable region causes its supercooling, but steam remains dry until from the energetic point of view it is more advantageous to start nucleation process. Driving force for nucleation is so-called supersaturation ratio $S = p/p_{\text{sat}}$ or equivalently supercooling $\Delta T = T_{\text{sat}} - T_c$. Condensation (phase change) process-initiating work can be expressed by means of thermodynamic potential Gibbs free enthalpy $G$. The change in free enthalpy during condensation process that poses the minimal work required to form a spherical water droplet is derived for general equation of state in [3] as follows

$$\Delta G = 4\pi r^2 \sigma + \frac{4}{3} \pi r^3 (p - p_{\text{sat}})$$

$$- \frac{4}{3} \pi r^3 \rho_I (RT_c \ln S + \Delta g'),$$  (25)

where $\sigma$ is surface tension of water and is modeled by IAPWS release [20], which allows its applicability for temperatures below 273.15K (possibility of its variation with droplet radius is not considered), $r$ is droplet radius, $R$ is specific gas constant of water, $R = 461.51805 \ J \ K^{-1}$ and $\Delta g' = g'(p,T_c) - g'(p_{\text{sat}},T_c)$ is difference of specific free enthalpy between actual and saturated state at constant vapor temperature for which we utilized IAPWS-97 formulation [19]. Then, we can write

$$\Delta g' = RT_c \sum_{i=1}^{13} n_i (\tau - 0.5)^i \left( p^i - p_{\text{sat}}^i \right),$$  (26)

where $\tau = T_c/T, \pi = p/(\text{MPa})$ and $n_i, I_i, J_i$ are constants. The radius at which the change in free enthalpy attains maximum is called critical radius $r_c$ (system is in unstable equilibrium) and it results from applying condition for local extremum of function $AG$ with respect to radius

$$r_c = \frac{2\sigma}{\rho_I (RT_c \ln S + \Delta g') - (p - p_{\text{sat}})}.$$  (27)

Clusters with radii larger than critical tend to grow contrary to clusters with radii smaller than critical which tend to decay (system tends to minimize its free enthalpy).

Nucleation rate $J$ is the rate at which number of new cores of droplets (nuclei) in unit volume is spontaneously generated from supersaturated vapor steam at the onset of homogeneous condensation. Because we consider steam to be perfectly pure, no foreign particles exist in the flow field an therefore we neglect heterogeneous condensation altogether. Model for nucleation rate is for engineering applications frequently provided by classical kinetic nucleation theory which assumes capillarity approximation and works with macroscopic quantities. Classical steady-state nucleation rate of Becker and Döring is expressed as

$$J_{\text{st}} = K \exp \left( \frac{-AG_c}{k_B T_c} \right).$$  (28)

where so-called nucleation barrier $AG_c = 4\pi r_c^2 \sigma / 3$ is given by substituting critical radius (27) into Eq. (25). Group $k_B T_c$ represents thermal energy with $k_B$ Boltzmann constant, $k_B = 1.3806488 \times 10^{-23} \ J \ K^{-1}$ and $K$ is kinetic prefactor
expressed as
\[ K = \sqrt{\frac{2\sigma}{\pi m_1^4}} \left( \frac{p}{RT_c} \right)^{2} \frac{1}{\rho_v} \]  
(29)
in which \(m_1 = M/N_A\) is mass of single steam molecule, \(M = 18.015268 e^{-3} kg/mol^{-1}\) is molecular mass of water and \(N_A = 6.02214129 e^{23} mol^{-1}\) is Avogadro constant. Classical nucleation rate \(J_c\) is frequently corrected by Courtney’s and Kantrowicz’s correction factors which take into account effects of partial pressure of the clusters and temperature difference between clusters and vapor phase. According [21] joint Courtney’s and Kantrowicz’s correction is
\[ C_{corr} = \frac{1}{S} \left( 1 + \frac{\rho R}{\alpha} \sqrt{\frac{RT}{2\pi}} \left( \frac{h_v}{h_v} - \frac{1}{2} \right) \right), \]  
(30)
where \(\alpha\) is convective heat transfer coefficient for critical cluster and is defined later, Eq. (38). Then, corrected nucleation rate is given by
\[ J = C_{corr} J_c. \]  
(31)

### 2.6 Growth of existing water droplets

Temperature of water droplets differs from surrounding vapor phase temperature the more the faster expansion takes place. Faster expansion implies faster droplet growth, hence there is less time to diminish temperature difference. The rate of change in radial direction of water droplet can be determined by balance of energy over a surface of spherical droplet (absence of mass transfer). Only a part of released latent heat by condensing vapor steam is used for droplet heating. Simplified energy balance equation can be written as follows
\[ (h_v - h_c) \frac{dm}{dt} = \Delta s_r (T_r - T_v), \]  
(32)
where the term on LHS is the rate of local specific latent heat released to the droplet and the RHS term is a portion of heat flowing toward vapor by convection mechanism between droplet and surrounding vapor steam. Heat transfer by conduction throughout droplet is neglected. Substituting \(m = 4 \pi r^3 \rho/3\) for mass and \(A = 4 \pi r^2\) for surface of single droplet we arrive at the default equation for droplet growth
\[ \frac{dr}{dt} = \frac{\alpha(T_r - T_v)}{(h_v - h_c)\rho_v}. \]  
(33)

Important quantity in determining droplet growth is Knudsen number \(Kn\) defined as a ratio of mean free path of vapor molecules \(\ell_c\) to the droplet diameter
\[ Kn = \frac{\ell_c}{2r} = \sqrt{\frac{\pi \rho_v}{2RT_v \rho_v}}, \]  
(34)
where \(\ell_c\) is given by kinetic theory. Regime with \(Kn \gg 1\) (\(Kn > 4.5\)) is considered as free molecular and with \(Kn \ll 1\) (\(Kn < 0.01\)) as continuum, while in between there is the transition regime. Since size of droplets in steam turbines covers quite wide range, providing droplet growth model covering entire range of Knudsen number is useful from practical point of view.

Gyarmathy postulated that temperature of sub-micron water droplets can be estimated utilizing capillarity approximation in the following way
\[ T_f = T_s - (T_s - T_v) \frac{r_e}{r}. \]  
(35)

Hence, critical droplets have temperature of surrounding vapor and the more the droplets are larger then their critical size the closer is their temperature to saturation temperature. For the convective heat transfer coefficient he presented empirical relation [5]. Here we adopted its slightly modified form by utilizing Langmuir droplet model which defines interface radius \(r_i = r + \beta \ell_c\). We distinguish following regimes
\[ r \leq r_i \quad \text{... continuum regime}, \]
\[ r > r_i \quad \text{... free molecular regime}. \]

Choice \(\beta = 0.75\) is frequently suggested. Gyarmathy-Langmuir heat transfer coefficient takes the form
\[ \alpha = \frac{1}{1 + \beta e^{Kn}} + \frac{2 \sqrt{\pi}}{1.5 Kn} \frac{1}{1 + \beta e^{Kn}} \]  
(36)
We rewrite it as follows
\[ \alpha = \frac{1}{1 + e^{Kn}} + \frac{2 \sqrt{\pi}}{1.5 Kn} \frac{1}{1 + e^{Kn}} \]  
(37)
by utilizing relation for Prandtl number \(Pr = \mu c_p/\lambda\) and isobaric specific heat capacity for perfect gas \(c_p = kR/(K - 1)\). Substituting \(K = 1 + \rho v^2/\rho_t \approx 1.332633\) from [18] we arrive at
\[ \alpha = \frac{1}{1 + e^{Kn}} + \frac{2 \sqrt{\pi}}{1.5 Kn} \frac{1}{1 + e^{Kn}} \]  
(38)
where \(c = 1.032663666979 e^{-3}\). In the continuum regime \(\ell \ll r\), relation (38) is in accordance with conduction theory for which \(\alpha = \lambda/\ell\). For our implementation we used only temperature dependent ideal part of vapor thermal conductivity \(\lambda_v = \lambda(T_v)/\sqrt{T_v}\) and the same holds for vapor phase dynamic viscosity \(\mu_v = \mu(T_v)\), both stated in [4].

With Gyarmathy and Langmuir assumptions final form of droplet growth takes the form
\[ \frac{dr}{dt} = A \frac{1 - r_i/r}{r + \beta \ell_c} + B \]  
(39)
where \(A, B\) depend only on vapor parameters
\[ A = \frac{\lambda_v(T_{sat} - T_v)}{(h_v - h_c)\rho_v}, \]  
(40)
\[ B = 6.024548294942 e^{-5} \frac{\lambda_v}{\rho_v \sqrt{T_v}}. \]  
(41)

Another (probably better) growth model presents e.g. White and Young in [11] and Dykas and Wróblewski in [7].

Coefficients of linear approximation of droplet growth \(g(r) \approx a + br\) are given as follows
\[ a = \lambda(r) - \lambda'(r)r, \quad b = \lambda'(r), \]
where \(\lambda'(r)\) is radial derivative of droplet growth evaluated at average radius \(\bar{r}\) and in our CFD code is implemented exactly.
2.7 Final form of governing equations

Following set of partial differential equations in conservative form is solved
\[
\frac{\partial W_i}{\partial t} + \frac{\partial F_i}{\partial x} + \frac{\partial G_i}{\partial y} = S_i, \quad i = 1:8,
\]
where indexes \(i = 1:4\) form Euler equations for mixture and indexes \(i = 5:8\) form equations for liquid phase, \(W_i\) are conservative variables, \(F_i\) are components of flux vector in \(x\)-direction, \(G_i\) are components of flux vector in \(y\)-direction and \(S_i\) are liquid phase sources. In more detail
\[
\begin{bmatrix}
\rho \\
\rho u \\
\rho v \\
\rho E \\
\rho Q_0 \\
\rho Q_1 \\
\rho Q_2 \\
\rho y
\end{bmatrix}
+ \frac{\partial}{\partial x}
\begin{bmatrix}
\rho u \\
\rho u^2 + p \\
\rho u v \\
(pE + p)u \\
\rho Q_{0u} \\
\rho Q_{1u} \\
\rho Q_{2u} \\
\rho yu
\end{bmatrix}
+ \frac{\partial}{\partial y}
\begin{bmatrix}
\rho v \\
\rho u v \\
\rho v^2 + p \\
(pE + p)v \\
\rho Q_{0v} \\
\rho Q_{1v} \\
\rho Q_{2v} \\
\rho vy
\end{bmatrix}
= \begin{bmatrix}
0 \\
0 \\
0 \\
0 \\
r_sJ + apQ_0 + bpQ_1 \\
r_s^2J + 2apQ_1 + 2bpQ_2 \\
\frac{4}{3}\pi
\end{bmatrix}.
\]

This system is closed by additional equations
- equation of state for vapor phase
- nucleation model
- droplet growth model
- liquid saturation properties
- other properties for vapor phase

3 Numerical method

3.1 Problem formulation

We are interested in stationary solution \(W : \mathbb{D} \rightarrow \mathbb{R}^8\) obtained by the time marching method, where \(\mathbb{D} \subset \mathbb{R}^2\) is domain of solution, see figure 1. Solution vector \(W\) has to fulfill the integral form of the system Eq. (42)
\[
\frac{\partial}{\partial t} \int_{\mathbb{D}} W \, dx + \int_{\mathbb{D}} [\nabla \cdot (\mathbf{F}, \mathbf{G}) - S] \, dx = 0
\]
for any subset \(\tilde{\mathbb{D}} \subset \mathbb{D}\) and boundary conditions prescribed at boundaries of solution domain.

3.2 Calculation of vapor parameters from conservative parameters

In order to calculate conservative vector \(W^{n+1}\) in new time level \((n + 1)\) we have to determine static pressure, vapor temperature and sound speed of mixture from the conservative vector in previous time level \(W^n\). Pressure and vapor temperature are calculated from vapor density and vapor specific internal energy utilizing equation of state for vapor phase. Mixture sound speed is calculated according [15]
\[
w^2 = \frac{u}{\rho} + \frac{1}{\rho(\rho - 1)}(\frac{d\rho}{dp})
\]
but in our case it is calculated with frozen composition, which results in vanishing the last term in denominator. Note that we calculated liquid density, liquid specific internal energy, cf. (4), and liquid sound speed from vapor temperature in previous time level \(T_{n}^{v}\) otherwise we would had to solve non-linear algebraic equations simultaneously.

3.3 Boundary conditions for subsonic inlet and subsonic outlet

Due to equation of state in CFD formulation, no iteration is needed in order to calculate static pressure and/or vapor temperature in inner cells, but there is different situation in case of fluxes calculations through inlet and outlet boundary. Inlet ghost cells work with
- first order extrapolation of static pressure \(p\) from the first neighboring cell
- entropy \(s\) and enthalpy \(h\) equal to their stagnant values and calculated from prescribed total pressure \(p_{st}\) and total temperature \(T_{st}\) during preprocessing stage
- prescribed flow angle \(\alpha\)
- assumption of dry steam: \(W_4 = W_5 = W_6 = W_7 = 0\)

Therefore, numerical solution of system of two nonlinear algebraic equations (from given pressure and entropy calculate density and temperature) is necessary provided we intend to keep the same state equation. Newton-Raphson method is implemented with appropriate stop criterion. Properties of vapor phase are equal to their mixture values, as we consider dry steam at inlet.

Similar situation is in outlet ghost cells, where we solve one nonlinear equation – vapor temperature is calculated from prescribed average static back pressure and vapor density which is extracted from first order extrapolation of mixture density and wetness factor from the first neighboring cell.

In the inviscid case we implemented slip condition on the wall boundaries. Normal component of velocity vector is zero, hence fluxes are composed from pressure part only. Static pressure on the wall is approximated by the value of static pressure in the first neighboring cell.

On periodic boundaries we implemented point-to-point periodicity.
3.4 Solution procedure

In order to tackle stiff character of the system due to different time scales of convection and condensation, we split the system into two subsystems by fractional step method. Because we implemented this method in symmetrical form [12], conservative variables in new time level \((n+1)\) are calculated from the ones in old time level \((n)\) by following successive steps

\[
W^n_0 = W^n_i, \\
\frac{\partial W^n_i}{\partial t} = S^n_i, \quad \Delta t = 0.5 \Delta t, \\
\frac{\partial W^n_{ii}}{\partial t} = -\frac{\partial F^n_{i}}{\partial x} - \frac{\partial G^n_{ii}}{\partial y}, \quad W^n_{ii} = W^n_i, \quad \Delta t = \Delta t, \\
\frac{\partial W^n_{iii}}{\partial t} = S^n_{iii}, \quad W^n_{iii} = W^n_{ii}, \quad \Delta t = 0.5 \Delta t, \\
W^{n+1}_i = W^{n+1}_{ii}.
\]

Time step \(\Delta t\) in inviscid case is estimated as in [2] and we used \(CFL = 0.75\). Time integration in case of condensation part is repeated \(k\)-times with reduced time step, where \(k\) is given as

\[
k = 1 + (int) \left( \frac{\Delta t}{\tau^c} \right) \quad (46)
\]

and condensation time scale \(\tau^c\) is estimated by Wegener formula [14]

\[
\tau^c = \frac{\sigma}{\rho \ln(S)} \sqrt{\frac{8\pi}{RT_g}} \quad (47)
\]

Convective fluxes of conservative variables through cell interfaces are calculated applying mid-point rule and AUSM+ scheme [16]. Only constant distribution of variables in cell volume is considered, that is why only 1st order spatial accuracy is attained. The scheme uses arithmetic averaged speed of sound in mixture \(w\) from left and right cells. For time integration we implemented explicit 2nd-order 2-stage Runge-Kutta method in case of both parts.

3.5 Calculation of liquid sources

We distinguished four cases in calculation of liquid sources \(S\). Our algorithm is as follows: Calculate supersaturation ratio \(S = p/p_{sat}(T_s)\) and average droplet radius \(\bar{r}\)

\[
\bar{r} = \left\{ \begin{array}{ll} \sqrt{W_6/W_i} \ldots W_7 > W_{sat} \ldots 0 \ldots \text{otherwise} \end{array} \right. \quad (48)
\]

where we decided for \(W_{sat} = 1 \times 10^{-12}\). Then proceed with the following branching

1. \((S > S_b)\) and \((\bar{r} > r_c)\)
   - nucleation and droplet growth
   - calculate \(J\) and \(\dot{r}\)
2. \((S > S_b)\) and \((\bar{r} \leq r_c)\)
   - only nucleation
   - set \(\dot{r} = 0\) \((a = b = 0)\) and calculate \(J\)
3. \((S < 2.0 - S_b)\) and \((\bar{r} > r_{min})\)
   - droplets evaporation
   - set \(J = 0\) and calculate \(\dot{r}\) with \(r_c = 0\)
4. (otherwise)

4 Results of numerical simulation

4.1 Test case and remarks

We followed the validation test case no. 27 in [22], [23]. So, we prescribed at inlet boundary

\[
p_{sat} = 0.0999 \, MPa, \quad T_{sat} = 360.8 \, K, \quad \alpha = 38^\circ,
\]

\[
W_4 = W_5 = W_6 = W_7 = 0
\]

and at outlet boundary average pressure ratio

\[
P_{out} = P_{out} = 2.34.
\]

Only one period of blade cascade was meshed. We used unstructured mesh configuration with quadrangle elements. Free tool Gmsh [13] was used for this purpose. Mesh consists of 30,687 cells, 61,917 cell faces and 31,230 nodes, see figures 2, 3. Trailing edge was modified due to inviscid model. Original circular end of the trailing edge was
replaced by the sharp wedge end – it reduces partly unphysical strong expansion at the trailing edge.

Implementation of linear model of droplet growth gave good results in case of flow in nozzles (with slightly worse convergence comparing to constant droplet growth). However its using in case of blade-to-blade cascades resulted in very different results comparing to experiment data and results obtained by several authors. We blame unrealistic expansion in the vicinity of sharp trailing edge for that. Therefore, we simplified our model by consideration of constant droplet growth $(b = 0)$ in evaluating liquid sources.

### 4.2 Results with commentary

Figure 4 presents distribution of supercooling $\Delta T = T_{\text{sat}} - T_v$ and figure 5 supersaturation ratio $S = p/p_{\text{sat}}$ of expanding steam in the blade cascade with some representative contours of constant value. Saturation contour $S = 1$ is showed as well. Region with $S < 1$ represents superheated region where evaporation of droplets occurs. This region is initiated by shock wave behind which vapor temperature crosses saturation temperature. Figure 6 depicts some contours of nucleation rate $J$. Unphysical strong expansion at the trailing edge is the reason of maximum nucleation rate $(J > 1\times10^4 \frac{m^3}{s^3},$ region MAX in the figure). Static pressure is presented in figure 7. Figure 8 shows Mach number in the domain. Maximum local mach number is 1.28. Contour with $M = 1$ is presented as well (thick blue line in the figure). Calculated surface-averaged droplet radius in the domain is in figure 9. The biggest droplets with radius about $0.03 \mu m$ are located in the middle of the channel. In the region behind the shock wave droplets are partly shrunk. Wetness factor is in figure 10 and has maximum value about 0.058.

Along the blade surface we present following: Figures 11, 12 shows supercooling and supersaturation, respectively. Nucleation rate is presented in figure 13. Ratio of static pressure to the total inlet pressure is plotted in figure 14. Figure 15 presents Mach number. Figure 16 and figure 17 plots droplet radius and wetness factor, respectively. The distribution of static pressure on the pressure side is in very good accordance with experiment [22], [23]. The most interesting is the suction part of the blade. First nucleation occurs in the initial part of the suction surface as a result of
Figure 7. Static pressure in domain: 25 contours

Figure 8. Mach number in domain: 50 contours, contour $M = 1$ (thick blue line)

Figure 9. Surface-averaged droplet radius in domain: 25 contours

Figure 10. Wetness factor in domain: 25 contours
curved blade profile. Second nucleation part occurs in the middle of channel, where pressure knee is visible. However, the magnitude of pressure rise is smaller and location is shifted upstream comparing to experiment data. Small location of trailing edge on the pressure side exhibits extrema of nearly all parameters resulting from unphysical over-expansion stemming from using inviscid model.

5 Conclusion

The main contribution of the paper is implementation of iteration-free vapor equation of state in CFD formulation. We based our work on the full Eulerian approach which combines the Euler inviscid system for the whole mixture and the method of moments for liquid phase. Presented flow model was validated on one selected regime.
of transonic wet steam flow in the steam turbine cascade channel of penultimate stage at the rotor tip section provided by Bakhtar. Taking limitations which arise from inviscid approach, our numerical results are in satisfied accordance with the experiment data and numerical simulations provided by the others authors. Our implementation of real thermodynamics of steam was successful. Attention requires implementation of one additional term in liquid sources resulting from linear approximation of droplet growth, which is our goal for future work.

Acknowledgments

The support of the grant GA ČR 101/11/1593 and the grant No. SGS13/174/OHK2/3T/12 of Grant Agency of the Czech Technical University in Prague is gratefully acknowledged.

References

1. R.S.R. Sidin, *Droplet size distribution in condensing flow* (University of Twente, Enschede, 2009)
2. J. Blazek, *Computational fluid dynamics - principles and applications* (Elsevier, 2006)
3. K. Fakhari, *Numerical modeling and investigation of unsteady phenomena in condensing flows of industrial steam turbines* (University of Bochum, 2010)
4. W. Wagner, H.-J. Kretzschmar, *International steam tables* (Springer, 2008)
5. M.J. Moore, C.H. Sieverding, *Two-phase steam flow in turbines and separators* (McGraw-Hill, 1976)
6. S. Dykas, W. Wróblewski, International Journal of Multiphase flow 37, 1245 (2011)
7. S. Dykas, W. Wróblewski, International Journal of Heat and Mass Transfer 60, 88 (2013)
8. J. Hrubý, EPJ Web of Conferences 25, (2012)
9. J. Hrubý, M. Duška, EPJ Web of Conferences 67, (2014)
10. J. Hrubý, J. Pátek, M. Duška, Journal of Power and Energy 228, 120 (2014)
11. A.J. White, J.B. Young, J. Propul. Power 9, 579 (1993)
12. G. Strang, SIAM Journal of Numerical Analysis 5, 506 (1968)
13. C. Geuzaine, J.-F. Remacle, International Journal for Numerical Methods in Engineering 79, 1309 (2009)
14. P.P. Wegener, L.M. Mack, ZAMM – Journal of Applied Mathematics and Mechanics 38, 489 (1958)
15. J.R. Travis, W. Breitung, D. Piccioni Koch, T. Jordan, International Journal of Hydrogen Energy 37, 17373 (2012)
16. M.-S. Liou, Journal of Computational Physics 129, 364 (1996)
17. W. Wagner, A. Pruss, J. Phys. Chem. Ref. Data 31, 387 (2002)
18. IAPWS, Revised release on the IAPWS formulation 1995 for the thermodynamic properties of ordinary water substance for general and scientific use, (2009)
19. IAPWS, Revised Release on the IAPWS Industrial Formulation 1997 for the Thermodynamic Properties of Water and Steam, (2007)
20. IAPWS, Revised Release on Surface Tension of Ordinary Water Substance, (2014)
21. F. Bakhtar, A.J. White, H. Mashmoushy, Proceedings of the Institution of Mechanical Engineers 219, 1335 (2005)
22. Bakhtar F., Ebrahimi M., Webb R.A., Proceedings of the Institution of Mechanical Engineers 209, 115 (1995)
23. F. Bakhtar, M. Ebrahimi, B.O. Bamkole, Proceedings of the Institution of Mechanical Engineers 209, 169 (1995)