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An assessment of in-tube flow boiling correlations for ammonia-water mixtures and their influence on heat exchanger size

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Abstract

Heat transfer correlations for pool and flow boiling are indispensable for boiler design. The correlations for predicting in-tube flow boiling heat transfer of ammonia-water mixtures are not well established in the open literature and there is a lack of experimental measurements for the full range of composition, vapor qualities, fluid conditions, etc.

This paper presents a comparison of several flow boiling heat transfer prediction methods (correlations) for ammonia-water mixtures. Firstly, these methods are reviewed and compared at various fluid conditions. The methods include: (1) the ammonia-water specific flow boiling correlations from the open literature, (2) the ammonia-water specific pool boiling correlations from the open literature extended to flow boiling by using the pure fluid correlation by Gungor and Winterton, and (3) the classical wide-boiling correlations. Secondly, their influence on the required heat exchanger size (surface area) is investigated during numerical design. For this purpose, two case studies related to the use of the Kalina cycle are considered: a flue gas based heat recovery boiler for a combined cycle power plant and a hot oil based boiler for a solar thermal power plant.

The results indicate that the nucleate boiling contribution to flow boiling is small compared to the flow boiling contribution for the investigated conditions. Furthermore, the use of the different flow boiling correlation methods resulted in evaporator size differences within 6\% for the heat recovery boiler and 28\% for the oil based boiler.

Keywords: Ammonia-water, Flow boiling, Heat exchanger design, Heat transfer, Kalina cycle, Modeling, Zeotropic mixture
Nomenclature

Roman

Bo boiling number

$D$ mass diffusivity [$m^2 s^{-1}$]

d diameter [m]

$E_0$ enhancement factor

$F$ two-phase multiplier

$F_e$ pool boiling mixture correction factor

$G$ mass flux [$kg m^{-2} s^{-1}$]

g gravitational acceleration [$m s^{-2}$]

$h_{LV}$ latent heat of evaporation [$kJ kg^{-1}$]

$h$ heat transfer coefficient [$W m^{-2} K^{-1}$]

$h_I$ ideal heat transfer coefficient [$W m^{-2} K^{-1}$]

$k$ thermal conductivity [$W m^{-1} K^{-1}$]

$p$ pressure [Pa]

$Pr$ Prandtl number

$q$ heat flux [$W m^{-2}$]

$Re$ Reynolds number

$S$ suppression factor

$T$ temperature [$K$]

$X$ vapor quality [-]

$Z$ bulk mass fraction of ammonia

$X_{HI}$ Lockhart-Martinelli parameter

$x, \bar{x}$ liquid mass and mole fraction of ammonia

$y, \bar{y}$ vapor mass and mole fraction of ammonia

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Greek

1. $\alpha$ thermal diffusivity [m$^2$ s$^{-1}$]
2. $\rho$ density [kg m$^{-3}$]
3. $\Delta T_{bp}$ boiling range [K]
4. $\Delta T_I$ ideal wall superheat [K]
5. $\mu$ dynamic viscosity [Pa s]
6. $\sigma$ surface tension [N m$^{-1}$]

Subscripts

9. bub bubble point
10. $c$ cold
11. dew dew point
12. $h$ hot
13. L liquid
14. cb convective boiling
15. nb nucleate boiling
16. $s$ saturation
17. V vapor

Abbreviations

18. EV evaporator
19. EC economizer
20. HRB heat recovery boiler
21. LL linear law
22. MP mixture properties
23. OBB oil based boiler
24. RSD relative standard deviation
25. SH superheater
1. Introduction

The ammonia-water mixture has been used as working fluid for decades in various applications utilizing low-grade heat such as absorption refrigeration systems, heat pumps and power cycles. Reliable heat transfer correlations are always needed for the design of heat exchangers in such systems as both the plant efficiency and the cost (heat exchangers size) are important when comparing the economic feasibility.

The ammonia-water mixture is a zeotropic mixture having a non-isothermal phase-change. One of the main benefits of the non-isothermal phase-change is the possibility of temperature matching with the heat source/sink temperature profiles, thus decreasing thermal irreversibilities and improving plant efficiency compared to pure fluids. On the other hand, the required heat transfer surface area increases because of the reduced temperature difference between the working fluid and the heat source/sink. In addition, the mixture flow boiling heat transfer coefficients are typically smaller than their pure counterparts [2, 3]. The resulting area increase influence negatively in the eventual economic advantage of a more effective plant [4] and it may result in an unfeasible plant economy. The heat transfer degradation in the nucleate boiling region is mainly due to the additional mass transfer resistance, while in the convective boiling region, it is mainly due to the changes in thermodynamic and transport properties [2].

The thermodynamic properties of ammonia-water mixtures are experimentally established [5] and may be accessed by databases such as REFPROP 9.12 [6]. The transport properties of ammonia-water mixtures are less established and different experimental sources show remarkable inconsistencies [7]. The influence of using different transport property methods for the estimation of viscosity and thermal conductivity of ammonia-water mixtures have recently been evaluated during numerical heat exchanger design [8]. The results suggested that the different transport property methods, traditionally used for the ammonia-water mixtures, show small variation in the estimated heat exchanger sizes. Although the heat exchanger size shows very little sensitivity towards the different transport property estimation methods, it remains to assess the sensitivity of the various flow boiling heat transfer correlations on the predicted heat exchanger size.

Only two in-tube flow boiling heat transfer correlations were found developed specifically for
ammonia-water mixtures [9, 10] in the open literature. These two correlations employ the simple
flow boiling correlation by Mishra et al. [11] but with very different constants. Three other studies
on flow boiling of ammonia-water mixtures have been found [12–14], however these consider
plate heat exchangers, whose heat transfer correlations are not applicable in the general case of
in-tube flow boiling. Additionally, the pressures in all these studies are well below that typically
encountered in Kalina cycle boilers. From the low availability in the open literature, it is clear that
general heat transfer prediction methodologies for wide boiling mixtures (e.g. ammonia-water)
must be used in order to predict the flow boiling heat transfer of ammonia-water mixtures and thus
the evaporator size. To the authors’ knowledge, no previous analysis has been conducted about the
influence of using different flow boiling heat transfer correlations for ammonia-water mixtures.

This paper compares a set of various in-tube flow boiling heat transfer correlations applied to
the ammonia-water mixture. The two aforementioned ammonia-water specific flow boiling corre-
lations [9, 10] are compared with two classical wide-boiling correlations [2, 15]. Additionally, a
few ammonia-water specific pool boiling correlations are extended to flow boiling following the
approach outlined by Palen [16] and Thome [17], using the pure fluid flow boiling correlation by
Gungor and Winterton [1]. Then the required evaporator surface area is investigated using each
correlation during numerical design. The objective is to clarify whether the different heat transfer
correlations have a significant influence on the designed evaporator size. Two Kalina cycle case
studies are considered for this purpose: a flue gas based heat recovery boiler (HRB) for a combined
cycle power plant and a hot oil based boiler (OBB) for a solar thermal power plant. The two cases
represent boilers with either poor or good heat transfer characteristics on the hot fluid side. The
boiler is usually the largest component in the Kalina cycle, taking up about half of the total surface
area and destroying the most exergy [18]. In total, 7 correlations are used during the numerical
design and simulated at different ammonia mass fractions and pressures in order to include several
design conditions.

The contributions to state-of-the-art of the paper are a thorough review of applicable flow boil-
ing correlations for ammonia-water mixtures, and a comprehensive comparison of these correla-
tions and their individual effect on the required heat exchanger size (surface area) during numerical
design of the two Kalina cycle case studies.
The outline of the paper is as follows: Section 2 introduces the different correlations. Section 3 describes the design models of the two case studies. Section 4 presents the results constituting the estimated heat exchanger sizes when using the different flow boiling heat transfer correlations. Finally, the results are discussed in Section 5 and followed up by the conclusions in Section 6.

2. Literature survey

Heat transfer correlations for pool boiling and flow boiling of binary mixtures are widely available for various organic mixtures, aqueous solutions and cryogenic fluids, and have been researched and reviewed by many [3, 15, 17, 19–22]. Such correlations are generally developed by modifying existing pure fluid correlations to account for mixture effects such as the liquid mass diffusion in the nucleate boiling region, a phenomenon absent in pure fluids. The liquid mass diffusivity is typically an order of magnitude smaller than the liquid thermal diffusivity and controls the nucleate boiling process [3]. The more volatile component becomes depleted in the surrounding liquid of a vapor bubble, causing a local increase of the boiling point at the liquid/vapor interface and loss of wall superheat. In contrast, the convective boiling contribution is typically well predicted with that in a pure fluid flow boiling correlation using mixture thermophysical properties. Nevertheless, the various heat transfer correlations for binary mixtures are generally mixture specific, and are only valid within the data bank limits from which they were developed [3].

Several studies have been conducted on pool boiling of ammonia-water mixtures while only a few studies have considered flow boiling. Also in these studies, the main aim was to develop better correlations to predict the boiling heat transfer by modifying already existing pure fluid correlations. However, the pressures in these experiments ($p < 15$ bar) are much lower than those of interest in Kalina cycles [18, 23, 24], which may be supercritical with respect to pure ammonia ($p > 113$ bar, $T > 132.3^\circ$C). In such conditions, the pure ammonia nucleate boiling coefficient cannot exist and many mixture heat transfer correlations fail to compute the ideal heat transfer coefficient, see Equation 7.

The current work is concerned with in-tube flow boiling of ammonia-water mixtures. However, both flow boiling and nucleate pool boiling have been considered. The latter is useful for flow boiling also as pointed out by Thome [17]: any existing pure fluid flow boiling correlation may
be modified in the nucleate boiling contribution to flow boiling by using the pool boiling mixture correction factor. As examples Thome [17] showed how to apply the pool boiling mixture correlation (Thome and Shakir [25]) in the pure fluid flow boiling correlations by Gungor and Winterton [1, 26]. Moreover, indefinite correlation combinations may be chosen by the correlation developer to fit the flow boiling data bank best, and errors become induced by both the failure to predict the nucleate boiling mixture correction as well as the failure of the general pure fluid correlation.

The aim of this literature survey is to establish a number of suitable flow boiling correlations for the ammonia-water mixture.

2.1. Flow boiling of ammonia-water mixtures

Rivera and Best [9] were the first authors to conduct flow boiling experiments of ammonia-water mixtures. They measured the heat transfer coefficient in a vertical smooth tube ($d = 25.8$ mm) at ammonia mass fractions from 0.38 to 0.48, pressures from 9.4 to 12.4 bar, mass fluxes from 4.2 to 9.0 kg m$^{-2}$ s$^{-1}$ and heat fluxes from 11.2 to 18.1 kW m$^{-2}$. No information was given on the vapor quality range. They argued that the flow boiling correlation by Mishra et al. [11] is advantageous, when there is a lack of transport property information available, e.g. ammonia-water mixtures. Hence, the Mishra et al. [11] equation was used to correlate their data:

$$h = C \left( \frac{1}{X_{tt}} \right)^m \text{Bo}^n$$  \hspace{1cm} (1)

where $C$, $m$ and $n$ were modified to be 65, 0.5 and 0.15, respectively. $h_L$ is the liquid convective heat transfer coefficient, estimated by the Dittus and Boelter [27] correlation, $X_{tt}$ is the Lockhart-Martinelli parameter and Bo is the boiling number, all defined by:

$$h_L = \frac{k_L}{d} \left[ 0.023 \text{Re}_L^{0.8} \text{Pr}_L^{0.4} \right]$$  \hspace{1cm} (2)

$$X_{tt} = \left( \frac{1 - X}{X} \right)^{0.9} \left( \frac{\rho_v}{\rho_L} \right)^{0.5} \left( \frac{\mu_L}{\mu_v} \right)^{0.1}$$ \hspace{1cm} (3)

$$\text{Bo} = \frac{q}{G h_L}$$ \hspace{1cm} (4)

where $k$, $\rho$, $\mu$ and $X$ are the thermal conductivity, density, dynamic viscosity and vapor quality, respectively. Re and Pr are the Reynolds and Prandtl numbers, $G$ is the mass flux (or mass velocity),
\( q \) is the heat flux, \( d \) is the diameter, and \( h_{LV} \) is the latent heat of evaporation. Furthermore, subscripts \( L \) and \( V \) denote the saturated liquid and vapor, respectively. The liquid Reynolds number is defined in terms of the liquid mass flux (\( \text{Re}_L = \frac{G(1 - X)}{\mu_L} \)).

The correlation was shown to predict the experimental data with an average deviation of 25%. Additionally, the nucleate boiling was shown to be suppressed at high values of \( X_{nt}^{-1} \) (independence of heat flux). Despite the simplicity, the Mishra et al. [11] correlation shows a major drawback as the constant and exponents in Equation 1 change with composition. In the original work by Mishra et al. [11], the authors recommended two sets of the constant and exponents ([\( C, m, n \) = 5.64,0.23,0.05] and [21.75,0.29,0.23]) at two different compositions for the R12-R22 mixture. Moreover, Celata et al. [28] showed the inadequacy of the Mishra et al. [11] correlation as more than 50% of their data was predicted with an error of more than 35%.

Khir et al. [10] performed flow boiling experiments with ammonia-water in a vertical stainless steel co-axial tube (\( d = 28 \text{ mm} \)). Three ammonia mass fractions (0.42, 0.55 and 0.61), three mass fluxes (707, 1590 and 2688 kg m\(^{-2}\) s\(^{-1}\)) and three heat fluxes (8.2, 14.8 and 18.5 kW m\(^{-2}\)) were varied simultaneously in their experiments, while the pressure ranged from 1.5 to 20 bar and the vapor qualities ranged from 0 to 0.6. They compared their experimental data with the correlations by Mishra et al. [11], Bennett and Chen [29] and the pure fluid correlation by Jung et al. [2], and found that only the Mishra et al. [11] correlation with [\( C, m, n \) = 5.64,0.23,0.05] gave reasonable predictions (errors less than 12%). Comparing these coefficients with those proposed by Rivera and Best [9] show remarkable inconsistencies (coefficients \( C, m \) and \( n \) are 91%, 54% and 67% smaller than those of Rivera and Best [9]), however, the mass flux range in these experiments differs by more than two orders of magnitude.

These flow boiling experimental studies do not provide sufficient information for the development of a general in-tube flow-boiling correlation for the ammonia-water mixtures. Most importantly, the composition was barely varied and the reduced pressure was too low for high pressure Kalina cycles. The fact that the composition was barely varied favors simple boiling expressions such as the correlation by Mishra et al. [11].
2.2. Pool boiling of ammonia-water mixtures

Nucleate pool boiling correlations for mixtures are generally expressed as follows (see Table 1):

\[ \frac{h}{h_1} = \frac{\Delta T_I}{\Delta T} = \frac{1}{1 + K} \]  \hspace{2cm} (5)

where \( h_1 \) and \( T_I \) denote the ideal heat transfer coefficient and ideal wall superheat, respectively, and \( K \) is a factor representing the heat transfer degradation. In general, it depends on the thermophysical and transport properties of the mixture, the liquid-vapor equilibrium relationship, the slope of vapor pressure curve and the non-linear dependence on composition. The latter may be incorporated by using either the vapor-liquid composition difference \( |\tilde{y} - \tilde{x}| \) or the boiling range \( (\Delta T_{bp} = T_{dew} - T_{bub}) \). The ideal wall superheat \( \Delta T_I \) is typically based on the molar average of the pure fluid wall superheats, i.e. the linear molar mixing law:

\[ \Delta T_I = \sum_{j=1}^{N} \tilde{x}_j \Delta T_j = \frac{q}{h_1} \]  \hspace{2cm} (6)

for which a division by the heat flux \( q \) leads to

\[ \frac{1}{h_1} = \sum_{j=1}^{N} \frac{\tilde{x}_j}{h_j} \]  \hspace{2cm} (7)

Here an appropriate pure fluid pool boiling correlation is used to predict the pure fluid nucleate boiling heat transfer coefficients. Alternatively, \( h_1 \) may be evaluated using mixture properties directly in the pure fluid correlation. Equation 7 seems to be applied mostly in the open literature, however if boiling occurs in between each pure fluid critical pressures or in between each triple point pressures, Equation 7 fails, whereas the alternative method is generally applicable. Additionally, for multi-component mixtures, the evaluation of each component boiling coefficients becomes cumbersome.

Inoue et al. [42] were found to be the first researchers to conduct pool boiling experiments with ammonia-water mixtures. They employed a horizontal heated platinum wire \( (d = 0.3 \text{ mm}) \) and measured heat transfer coefficients over all ranges of composition, below heat fluxes of 2000 kW m\(^{-2}\) and at pressures of 4 and 7 bar, respectively. The results showed a remarkable decrease in the heat
Table 1: Existing nucleate pool boiling correlations for mixtures

<table>
<thead>
<tr>
<th>Author</th>
<th>Correlation (generally applied)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stephan and Körner [30]</td>
<td>( \frac{h}{h_0} = \left( 1 + A_0(\bar{y} - \bar{x})(0.88 + 0.12\rho[\text{bar}]) \right)^{-1} ), ( A_0 = 1.53 ) (8)</td>
</tr>
<tr>
<td>Calus and Rice [31]</td>
<td>( \frac{h}{h_0} = \left( 1 + [\bar{y} - x] \left( \frac{\alpha L}{D_L} \right)^{0.5} \right)^{-1} ) (9)</td>
</tr>
<tr>
<td>Jungnickel et al. [32]</td>
<td>( \frac{h}{h_0} = \left( 1 + K_0(\bar{y} - \bar{x})(\rho \nu / \rho L)^{0.48 + 0.11\nu} \right)^{-1} ) (10)</td>
</tr>
<tr>
<td>Schlünder [33]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{h_0}{q} (T_{cl2} - T_{cl}) (\bar{y} - \bar{x}) \left( 1 - \exp \left( \frac{-B_0 q}{\rho L \nu L} \right) \right)^{-1} ), ( B_0 = 1, \beta_L = 0.0002 ) (11)</td>
</tr>
<tr>
<td>Thome and Shakir [25]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{h_0}{q} \frac{\Delta T_{bp}}{\Delta T_1} \left( 1 - 0.8 \exp \left( -\frac{q}{10^2} \right) \right)^{-1} \right) ) (12)</td>
</tr>
<tr>
<td>Fujita and Tsutsui [34]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} \left( 1 - \exp \left( \frac{-60 \rho}{\rho L \nu L} \left( \frac{\rho L}{\nu L} \right)^{1/4} \right) \right)^{-1} \right) ) (13)</td>
</tr>
<tr>
<td>Fujita and Tsutsui [35]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} \left( 1 - \exp \left( \frac{-60 \rho}{\rho L \nu L} \left( \frac{\rho L}{\nu L} \right)^{1/4} \right) \right)^{-1} \right) ) (14)</td>
</tr>
<tr>
<td>Inoue et al. [36]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} \left( 1 - 0.75 \exp \left( -\frac{0.75 q}{10^2} \right) \right)^{-1} \right) ) (15)</td>
</tr>
<tr>
<td>Vinayak Rao and Balakrishnan [37]</td>
<td>( \frac{h}{h_0} = \left( 1 - (\bar{y} - \bar{x}) \sqrt{\frac{\rho L}{\nu L}} \right)^{0.5} ) (16)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Author</th>
<th>Correlation (ammonia-water specific)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arima et al. [38]</td>
<td>( \frac{h}{h_0} = \left( 1 + A_1 \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} \right) + A_2(\bar{y} - \bar{x})(0.88 + 0.12\rho[\text{bar}])^{-1} ), ( A_1 = 0.134, A_2 = 2.07 ) (17)</td>
</tr>
<tr>
<td>Táboas et al. [39]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} \left( \frac{T_{cl2} - T_{cl}}{\Delta T_1} \right)^2 \left( 1 - \exp \left( \frac{-B_0 q}{\rho L \nu L} \right) \right) + \frac{A_1}{2} \frac{\Delta T_{bp}}{\Delta T_1} \left( 1 - \exp \left( \frac{-B_0 q}{\rho L \nu L} \right) \right) \right)^{-1} ) (18)</td>
</tr>
<tr>
<td>Inoue and Monde [40]</td>
<td>( \frac{h}{h_0} = \left( 1 + A_1 \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} \right) + A_2(\bar{y} - \bar{x})(0.88 + 0.12\rho[\text{bar}])^{-1} ), ( A_0 = 3.10, A_1 = 0.15, A_2 = 0.65 ) (19)</td>
</tr>
<tr>
<td>Sathyabhama and Babu [41]</td>
<td>( \frac{h}{h_0} = \left( 1 + \frac{k_1}{\rho L} \frac{\Delta T_{bp}}{\Delta T_1} + A_3 \frac{\Delta T_{bp}}{\Delta T_1} \left( \frac{T_{cl2} - T_{cl}}{\Delta T_1} \right)^2 \left( 1 - \exp \left( \frac{-B_0 q}{\rho L \nu L} \right) \right) \right)^{-1} ), ( A_1 = 0.15, A_3 = 0.25 ) (20)</td>
</tr>
</tbody>
</table>

1 transfer performance of the mixture close to the pure components. Five nucleate pool boiling mixture correlations (Equation 8, 10, 11, 13 and 15) were compared to their experimental data, but all of them failed to predict the measured coefficients satisfactory. Inoue et al. [42] noted that the non-linear variation in the heat transfer coefficient with composition was different than the non-linear variation of both \( \bar{y} - \bar{x} \) and \( \Delta T_{bp} \) with composition, which is typically in contrast to other
fluid mixtures. No alternative correlation was proposed by the authors.

Arima et al. [38] investigated pool boiling of ammonia-water mixtures on a horizontal flat heated silver surface at both low pressure and low ammonia mass fraction (2, 4 and 7 bar; 0 < x < 0.27) and high pressure and high ammonia mass fraction (7, 10 and 15 bar, 0.5 < x < 1). The splitting of concentration range and pressures resulted from their experimental apparatus’ upper temperature limit of 160 °C. They compared their results with two correlations (Equation 8 and 15). Equation 8 uses |\tilde{y} - \bar{x}| to express the non-linear variation with concentration while Equation 15 uses ΔT_{bp}. Again, none of the correlations were found to give satisfactory predictions, however, Arima et al. [38] proposed a revised correlation (Equation 17) that essentially is a combination of Equation 8 and 15. No uncertainty measures of the new correlation were given. Additionally, the ideal heat transfer coefficient were computed by Equation 7 with modified constant and exponents in the Nishikawa and Fujita [43] pure fluid correlation.

Táboas et al. [39] used the experimental data obtained by Inoue et al. [42] and Arima et al. [38], for heat fluxes below 1MW m$^{-2}$ and compared those data with three correlations (Equation 11, 12 and 14). All these correlations under-predicted the experimental heat transfer coefficients. Equation 7 was used for the ideal heat transfer coefficient, where it was found that the Mostinski [44] correlation was the most adequate to predict both the pure water and ammonia pool boiling coefficients compared to others [45–47]. Finally, a new revised correlation (Equation 18) was proposed that is also a combination of two correlations, i.e. Equation 11 and 12. The correlation reproduces the experimental data well within a 40% accuracy range.

Inoue and Monde [40] later compared three correlations (Equation 8, 11 and 15) with their improved data collection and found that the combinations of Inoue et al. [36] with either the correlation by Stephan and Körner [30] or Schlünder [33] were able to predict 97% of their data within 20% accuracy below 1MW m$^{-2}$ at 4 bar (Equation 19 and 20). Again, Equation 7 was used where the pure component pool boiling coefficients were obtained by the most recent Nishikawa and Fujita [48] correlation with original coefficients except for the nucleation factor (0.39) for ammonia.

Sathyabhama and Babu [41] investigated pool boiling of ammonia-water on a cylindrical stainless steel rod (d = 6 mm). The experiments were performed at heat fluxes from 360 to
2000 kW m\(^{-2}\), ammonia mass fractions from 0 to 0.3 and pressures of 4, 6, and 8 bar. They compared seven different existing correlations (Equation 8, 9, 11, 14, 16, 19 and 20) with their measurements and found that the correlations (Equation 8, 9 and 20) predicted their data within 20% accuracy. They further modified the exponent in the Calus and Rice [31] correlation (Equation 21), and compared it to the experimental data from Arima et al. [38] and Inoue and Monde [40]. The modified Calus and Rice [31] correlation was found to predict most of this literature data within 40% limit for all ranges of mass fraction. Interestingly, this correlation has no dependency on the heat flux. Sathyabhama and Babu [41] compared several pure fluid pool boiling correlations for water, and found that the correlations by Stephan and Abdelsalam [45] and Nishikawa et al. [49] gave reasonable results. An earlier study by the same research group [50] acknowledged the Mostinski [44] correlation amongst others for ammonia, however, no exact indication of how to calculate the ideal heat transfer coefficient was given.

2.2.1. Comparison of pool boiling correlations

From the above literature survey, it may be concluded that most authors use a combination of correlations that use either the concentration difference $|\tilde{y} - \tilde{x}|$ or the boiling range $\Delta T_{bp}$, respectively, to correlate their experimental data. Only the proposed correlation by Sathyabhama and Babu [41] (Equation 21) uses $|\tilde{y} - \tilde{x}|$ solely to express the reduction in the nucleate pool boiling, however, this correlation relies on transport properties of ammonia-water mixtures (thermal diffusivity and mass diffusivity), which are not well established in the open literature. Additionally, the survey shows that correlations in terms of concentration difference best represent the experimental data trend at low ammonia concentration, while correlations in terms of boiling range best represent the experimental data trend at high concentration.

All of the aforementioned pool boiling correlations use Equation 7 to compute the ideal heat transfer coefficient. In order to understand the effects of using different pure fluid pool boiling correlations at higher pressures, several well-known correlations for both fluids are evaluated in Figure 1. These correlations are the most recent by Nishikawa and Fujita [48] (with a nucleation factor of 0.39 for ammonia), the modified Nishikawa and Fujita [43] correlation by Arima et al. [12], the Nishikawa et al. [49] version used by Sathyabhama and Babu [41], and the correlations by
Figure 1: Pure ammonia and water pool boiling predictions as function of heat flux at different pressures

Cooper [46], Gorenflo and Kenning [47], Mostinski [44] and Stephan and Abdelsalam [45]. The equation developed for refrigerants was used for ammonia in the Stephan and Abdelsalam [45] correlation. Also the influence of surface roughness was neglected in the correlations by Cooper [46] and Gorenflo and Kenning [47].

The results indicate that the pool boiling correlations tend to differ more dramatically for ammonia than for water at higher pressures, thus different pressure dependencies are estimated by the
Figure 2: Pool boiling mixture correction factors (correlations) as function of ammonia mass fraction at various pressures and heat fluxes; (LL) is using the linear law (Equation 7) with the Mostinski [44] correlation; (MP) is using the Mostinski [44] correlation directly with mixture properties to evaluate $h_I$.

correlations for the pure ammonia. Táboas et al. [39] also compared a set of pool boiling correlations for ammonia to the available experimental data sets in the literature. They found relatively large discrepancies among the various experimental data sets, and noted that some correlations were developed from only some of the data sets, thus giving rise to discrepancies in the correla-
tions also. Furthermore, only 11 data points from the same data set goes beyond 15 bar. For these reasons, one must be careful when selecting a pool boiling correlation for ammonia.

Following the recommendations by Táboas et al. [39], Inoue and Monde [40] and Sathyabhama and Hegde [50], the pool boiling of ammonia may be satisfactorily predicted by the correlations of Mostinski [44], Stephan and Abdelsalam [45] and the most recent correlation by Nishikawa and Fujita [48] with a nucleation factor of 0.39 at low pressures. These correlations tend to coincide at higher pressures also. The correlation by Mostinski [44] is rather simple and it has also been recommended by Táboas et al. [39] for water. For these reasons, the correlation by Mostinski [44] was used for both the pure ammonia and water pool boiling predictions in the current work.

Figure 2 depicts the pool boiling mixture correction factor \( F_c = h/h_I \) using Equation 17 to 21 from Table 1.

In addition to the linear law (LL) (Equation 7), an example is included (Táboas et al. (MP)) for which the mixture properties (MP) have been used directly in the pool boiling correlation by Mostinski [44] to compute the ideal heat transfer coefficient. It is observed that the significance in the computation of the ideal heat transfer coefficient is small even at high pressure (Táboas et al. (MP) vs. Táboas et al. (LL)).

The general trend in Figure 2 is that the pool boiling mixture correction factor decreases (mass transfer resistance increase) as either pressure or heat flux increases. It is well known that the pool boiling heat transfer coefficient increases with both pressure and heat flux, thus causing the mass transfer resistance to increase. On the other hand, if the heat transfer coefficient tends to zero, so does the heat flux and the mixture correction factor must tend to unity. This dependency is in fact a property of most of the correlations presented in Table 1. The correlations by Arima et al. [38] and Inoue and Monde [40] (Equation 17 and 19) are however weak functions of heat flux compared to the correlations by Táboas et al. [39] and Inoue and Monde [40] (Equation 18 and 20), whereas the correlation by Sathyabhama and Babu [41] (Equation 21) is not a function of the heat flux. The differences in the predicted pool boiling mixture correction factor tend to be larger at low heat flux and smaller at high heat flux, excluding the correlation by Sathyabhama and Babu [41] which show a much different result at high pressures. Relatively large reductions in the pool boiling predictions are possible with respect to the ideal value, i.e. up to 70 % at 4 bar and 90 %
at 100 bar.

2.3. Pool boiling extensions to flow boiling

Following the approach by Palen [16] and Thome [17, 51], the general pure fluid flow boiling correlations may be modified in the nucleate boiling contribution to flow boiling by using the pool boiling mixture correction factor ($F_c = h/h_L$) defined in the previous section.

Palen [16] modified the widely quoted Chen [52] correlation to include the mixture boiling correction factor by Palen and Small [53] and obtained

$$h = h_L F + h_{nb} S F_c$$

where $F$ is the two-phase multiplier, $h_{nb}$ is the nucleate boiling coefficient, $S$ is the boiling suppression factor and $F_c$ is given as

$$F_c = \exp\left(-0.027\Delta T_{bp}\right)$$

with a minimum value such that $F_c \geq 0.1$. The other parameters are computed using the same equations as in the original Chen [52] correlation using mixture properties.

Thome [51] provided another modification of the Chen [52] correlation by using Equation 12 for $F_c$ in Equation 22. In a later review on boiling of new refrigerants, Thome [17] showed how to apply the same modification in the Chen-type correlation by Gungor and Winterton [1]

$$h = h_L E_0 + h_{nb} S F_c$$

where $E_0$ is the convective enhancement factor analogous to the two-phase multiplier $F$ in Equation 22. Again all parameters except $F_c$ are evaluated by the original Gungor and Winterton [1] correlation with mixture properties. Similarly, the simplified Gungor and Winterton [26] correlation was modified for mixtures as

$$h = h_L E$$

$$E = 1 + 3000 (\text{Bo} F_c)^{0.86} + 1.12 \left(\frac{X}{1-X}\right)^{0.75} \left(\frac{\rho_L}{\rho_{LV}}\right)^{0.41}$$
The heat flux to use for evaluating the mixture boiling correction factor, $F_c$, is the nucleate boiling heat flux and not the total heat flux as pointed out by Thome [17]. The total heat flux may be used as an approximation for small boiling range mixtures only, which does not apply for ammonia-water mixtures.

2.4. Classical wide-boiling correlations

Grauso et al. [22] gathered many experimental studies on wide-boiling mixtures and defined “wide-boiling mixtures” as mixtures with a boiling range of more than 5 K. Most of the considered mixtures had a maximum boiling range of less than 30 K, while that of the ammonia-water mixture is 94.5 K at 1 bar and 66.7 K at 100 bar. Only a single experimental study [54] was found in their literature survey with a fluid mixture (R22-DMAC) having a boiling range more than ammonia-water (over 200 K). In this study, it was found that the nucleate boiling was suppressed and that the Chen [52] correlation with negligible nucleate boiling contribution was found to predict the experiments best. It may indicate that the nucleate boiling is much suppressed for mixtures with very wide boiling range such as the ammonia-water mixture. On the other hand, Grauso et al. [22] compared their experimental results obtained for CO$_2$-propane blends including the data banks by Wettermann and Steiner [15] and Cho et al. [55], and found that the correlation by Wettermann and Steiner [15] performed best and predicted 61% of the heat transfer coefficients within 30% error.

The correlation by Wettermann and Steiner [15] was specifically developed for mixtures with wide-boiling range (e.g. R116-R134a with maximum $\Delta T_{bp} = 28.6$ K at 11 bar). It is also an extension of a well-known pure fluid flow boiling correlation by Steiner and Taborek [56], who used the asymptotic model to weight the nucleate and convective boiling contributions:

$$ h = \frac{1}{3} (F_{\text{cb}} h_{\text{cb}})^{\frac{1}{3}} + (F_{\text{hb}} h_{\text{hb}})^{\frac{1}{3}} $$  \hspace{1cm} (27)

Wettermann and Steiner [15] employed the nucleate boiling correlation by Thome and Shakir [25] (Equation 12) with $h_1$ defined by Equation 7 to compute the nucleate boiling heat transfer coefficient $h_{\text{nb}}$. Note that $F_{\text{nb}}$ is a nucleate boiling correction factor rather than a suppression factor, which is not needed in asymptotic models. Wettermann and Steiner [15] used $F_{\text{nb}} = 1$, 17
where the other parameters were computed similarly to the original pure fluid correlation with mixture properties. Note that the correlation by Wettermann and Steiner [15] also involves the determination of a high boundary heat flux \( q_B \) and the onset of nucleate boiling \( q_{onb} \). When \( q < q_{onb} \) the nucleate boiling contribution is neglected, and when \( q > q_B \) the nucleate boiling contribution is calculated by means other than Equation 12.

Another classical wide-boiling correlation is given by Jung et al. [2]. The correlation is also a Chen-type correlation and is given by:

\[
h = C_{me} F h_L + \frac{N}{C_{UN}} h_{UN}
\]

Here \( C_{me} \) considers the mass transfer resistance in the convective boiling region, \( 0.9 < C_{me} \leq 1 \), \( F \) is the enhancement factor, \( N \) is the suppression factor while \( C_{UN} \) and \( h_{UN} \) are the inverse mixture correction factor \((1/F_c)\) and nucleate pool boiling heat transfer coefficient, respectively, as proposed by Ünal [57]. The ideal heat transfer coefficient is computed by Equation 7 with the Stephan and Abdelsalam [45] correlation.

### 2.5. Choice of correlations

Table 2 summarizes the ammonia-water flow boiling correlations that are chosen for further investigations regarding the influence on the heat exchanger size. Two ammonia-water specific flow boiling correlations are considered as introduced in Section 2.1, which both rely on the simple correlation by Mishra et al. [11] with different coefficients. Three ammonia-water specific pool boiling correlations extended to flow-boiling are considered using the pure fluid flow boiling correlation by Gungor and Winterton [1] as described in Sections 2.2 and 2.3. The ammonia-water specific pool boiling correlation by Sathyabhama and Babu [41] is excluded from this analysis, because it is independent of the heat flux and gives results very different from the others at high pressures. Furthermore, the first correlation by Inoue and Monde [40] (Equation 19) is also excluded, because it coincides with the correlation by Arima et al. [38]. Finally, two classical wide-boiling correlations are used as introduced in Section 2.4.

Figure 3 shows the predicted heat transfer coefficients using these correlations vs. the vapor quality at 50 % bulk ammonia mass fraction, 4 bar, mass fluxes of 50, 150 and 400 \( \text{kg m}^{-2}\text{s}^{-1} \) and
Table 2: Ammonia-water flow boiling correlations chosen for further investigations in the current work

<table>
<thead>
<tr>
<th>Correlation</th>
<th>Proposed by</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mishra et al. [11], Equation 1</td>
<td>Rivera and Best [9] with ([C, m, n] = [65, 0.5, 0.15])</td>
</tr>
<tr>
<td>Mishra et al. [11], Equation 1</td>
<td>Khir et al. [10] with ([C, m, n] = [5.64, 0.23, 0.05])</td>
</tr>
</tbody>
</table>

Ammonia-water specific nucleate pool boiling correlations extended to flow boiling

<table>
<thead>
<tr>
<th>Pool boiling mixture correction factor</th>
<th>Ideal heat transfer coefficient method</th>
<th>Flow boiling correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arima et al. [38], Equation 17</td>
<td>Equation 7 with Mostinski [44]</td>
<td>Gungor and Winterton [1]</td>
</tr>
<tr>
<td>Taboas et al. [39], Equation 18</td>
<td>Equation 7 with Mostinski [44]</td>
<td>Gungor and Winterton [1]</td>
</tr>
<tr>
<td>Inoue and Monde [40], Equation 20</td>
<td>Equation 7 with Mostinski [44]</td>
<td>Gungor and Winterton [1]</td>
</tr>
</tbody>
</table>

Classical wide-boiling correlations

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Wettermann and Steiner [15]</td>
<td></td>
</tr>
<tr>
<td>Jung et al. [2]</td>
<td></td>
</tr>
</tbody>
</table>

heat fluxes of 10 and 100 kW m\(^{-2}\) for a vertical tube having an inner diameter of 34 mm.

Figure 3 indicate that the ammonia-water specific flow boiling correlations by Rivera and Best [9] and Khir et al. [10] are around an order of magnitude different from each other. Compared to the other correlations, the one by Rivera and Best [9] over-predicts while the one by Khir et al. [10] under-predicts the heat transfer coefficients. The general trend in Figure 3 is that an increase in the mass flux and heat flux increases the heat transfer coefficient. However, for the more elaborate methods (the ammonia-water specific pool boiling extensions to flow boiling and the classical wide-boiling correlations) the dependency on heat flux becomes suppressed at high vapor qualities and mass fluxes. The convective contribution to flow boiling, as predicted by Gungor and Winterton [1], is also presented in Figure 3. Compared to the pool boiling correlation extensions to flow boiling, it shows that the nucleate boiling contribution is rather small and mostly present at small vapor qualities, low mass fluxes and high heat fluxes. In other words, the different pool boiling mixture correction factors (see Figure 2) do not matter much when they are used to predict flow boiling. The classical wide-boiling correlations by Wettermann and Steiner [15] and Jung et al. [2] predict reasonably similar results and are close to the pool boiling extensions to flow boiling in most conditions.

Figure 4 shows the predicted flow boiling heat transfer coefficients at 100 bar. Here the cor-
Figure 3: Flow boiling heat transfer predictions vs. vapor quality at 50% bulk ammonia mass fraction, 4 bar and various mass and heat fluxes; the tube is vertical and has an inner diameter of 34 mm.
boiling correlations indicate that the nucleate boiling contribution is small and mostly present at low vapor qualities and mass fluxes. However, at both high pressure and high heat flux the nucleate boiling is almost fully suppressed. The general trend as pressure increases is that the boiling heat transfer coefficient decreases as compared with Figure 3. Again the classical wide-boiling correlations give similar predictions and are close to the pool boiling extensions to flow boiling at
most conditions.

3. Model development

This section presents the models of the flue gas based heat recovery boiler (HRB) for a combined cycle power plant and the oil based boiler (OBB) for a solar thermal power plant. The models represent Kalina cycle boilers with either poor or good heat transfer characteristics on the hot fluid side. The models are identical to the models used in Kærn et al. [8], except that the boiling heat transfer coefficients are computed differently. Note that the design of the boilers includes not only the evaporator, but also the economizer and the superheater. This is chosen so as to give a full overview of the boilers, i.e. how much the different boiling heat transfer correlations affect the evaporator area and the total area including the economizer and the superheater.

The thermodynamic properties of the ammonia-water mixture are evaluated using the NIST standardized reference database (REFPROP 9.12 [6]) for which the latest so-called "Ammonia (Lemmon)" fluid file is used for the ammonia-water mixture [58]. The fluid file contains a refit of the ammonia equation of state that is compatible with the latest excess Helmholtz mixture model. The new ammonia-water mixture formulation is more stable and faster while giving outputs that are very close to that of the earlier Tillner-Roth and Friend [5] formulation [59]. Additionally, the transport properties are evaluated with the formulations proposed by Conde-Petit [7]. The models are implemented in Dymola 2014 [60] using the Modelica language, for which the freely available Modelica-REFPROP interface [61] is used.

3.1. HRB model

A sketch of the HRB is shown in Figure 5 including the economizer (EC), the evaporator (EV) and the superheater (SH). The HRB is recognized as a once-through boiler having the cold fluid flowing inside a vertical tube bundle with external circular fins and the hot fluid flowing in a counter-cross flow arrangement. The hot fluid is simplified to be the product of a stoichiometric burning of methane and air (21 % O₂, 79 % N₂). The properties of the flue gas are evaluated using the Modelica.Media library, i.e. an ideal gas mixture. The inlet gas temperature and mass flow rate are 524 °C and 100 kg s⁻¹ replicating the exhaust gas conditions of a gas turbine. The
cold medium is the ammonia-water mixture with an ammonia mass fraction, inlet temperature and pressure of 0.7, 25 °C and 40 bar, respectively. The pinch point temperature difference is 15 K and the approach temperature difference (hot fluid inlet, cold fluid outlet) is 20 K. The ammonia mass fraction and pressure are varied from 0.5 to 0.9 and 40-100 bar, respectively, which are typical inlet conditions in Kalina cycle boilers [18, 23, 24].

![Heat recovery boiler](image)

Figure 5: Heat recovery boiler

With these assumptions, for a given ammonia mass fraction, the temperature profiles of both fluids as function of the heat flow rate may be constructed, see Figure 7a. In other words, the heat exchange in the economizer, the evaporator and the superheater is known by these design conditions, while the heat exchanger surface area remains unknown.

The HRB model computes the local heat transfer by discretization with respect to the heat flow rate. The specific geometry of the HRB is listed in Table 3 and the applied heat transfer and pressure drop correlations other than the boiling heat transfer correlation are shown in Table 4. Additionally, the tube layout is staggered and the tube material is steel (constant thermal conductivity of 42 W m⁻¹K⁻¹ is assumed).

The solution progress as well as a convergence analysis are described in Kærn et al. [8]. Based on the convergence analysis, the number of cells for the discretization is chosen to be 2000.

3.2. OBB model

A sketch of the OBB is shown in Figure 6. The OBB is recognized as several shell and tube heat exchangers having 1 shell pass for the hot fluid and 1 tube pass for the cold fluid. The tubes have circular fins and no tubes are placed in the window section. These features serve to enhance the heat transfer on the oil side while maintaining a low pressure drop. The hot fluid is Therminol
Table 3: Main geometry of the HRB

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tube length</td>
<td>5 m</td>
</tr>
<tr>
<td>Inner tube diameter</td>
<td>34 mm</td>
</tr>
<tr>
<td>Outer tube diameter</td>
<td>40.64 mm</td>
</tr>
<tr>
<td>Fin density</td>
<td>236 m$^{-1}$</td>
</tr>
<tr>
<td>Fin height</td>
<td>20 mm</td>
</tr>
<tr>
<td>Fin thickness</td>
<td>1.5 mm</td>
</tr>
<tr>
<td>Transverse tube pitch</td>
<td>100 mm</td>
</tr>
<tr>
<td>Longitudinal tube pitch</td>
<td>90 mm</td>
</tr>
<tr>
<td>Tubes per row / cold fluid circuits</td>
<td>60</td>
</tr>
<tr>
<td>Frontal area</td>
<td>30 m$^2$</td>
</tr>
<tr>
<td>Minimum free flow area</td>
<td>13.56 m$^2$</td>
</tr>
</tbody>
</table>

Table 4: Heat transfer and pressure drop correlations

<table>
<thead>
<tr>
<th>Fluid Type</th>
<th>Heat transfer</th>
<th>Friction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hot fluid</td>
<td>Briggs and Young [62]</td>
<td>Robinson and Briggs [64]</td>
</tr>
<tr>
<td>Fin efficiency</td>
<td>Shah and Sekulic [63]$^a$</td>
<td></td>
</tr>
<tr>
<td>Cold fluid single phase</td>
<td>Gnielinski [65]</td>
<td>Blasius [66]</td>
</tr>
<tr>
<td>Cold fluid two-phase</td>
<td>see Table 2</td>
<td>Müller-Steinhagen and Heck [67]</td>
</tr>
</tbody>
</table>

$^a$ approximative function ± 1 % accuracy.

66 and the properties are obtained from Solutia Inc. [68]. The conditions are almost the same as those of the HRB, except that the cold fluid and the hot fluid inlet temperatures are 50°C and 320°C, respectively. Again, the temperature profiles of both fluids as function of the heat flow rate may be constructed (see Figure 7b), and the OBB remains to be designed to find the required heat exchanger surface area.

The model of the OBB is identical to the HRB from previous section, except that the heat trans-
The heat transfer coefficient and pressure drop on the shell side are computed differently. The Bell-Delaware method as extended by Taborek [69] has been used to establish the heat transfer coefficient and pressure drop on the shell side. In fact the ideal heat transfer coefficient and ideal pressure drop (ideal tube bundle cross flow through the centerline of the shell) are computed with the same correlations as for the HRB (Table 4), however corrected for non-ideal shell flow effects using the Bell-Delaware method. The entrance and exit baffle spacing effects have been neglected for simplicity. The specific geometry of the OBB is listed in Table 5. Additionally, the staggered rotated square layout is used and the tube material is steel.

Note that the number of parallel tubes has increased compared to the HRB from 60 to 120. This was chosen in order to keep the shell length to diameter ratio reasonably low, thus reducing the resulting hot fluid flow length and corresponding pressure drop.

4. Results

In this section, the influence of the flow boiling heat transfer correlations on the predicted heat transfer surface area is reported for both the HRB and the OBB. The results are presented not only specific to the evaporator itself, but also specific to the total boiler area, including the economizer and the superheater for completeness. Firstly, the results of a baseline numerical design at various pressures and ammonia mass fractions are presented for both the HRB and the OBB. This baseline serves as a reference to better understand the heat exchanger size differences when changing operation conditions and using the same flow boiling correlation.
Table 5: Main geometry of the OBB

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of parallel tubes</td>
<td>120</td>
</tr>
<tr>
<td>Inner tube diameter</td>
<td>34 mm</td>
</tr>
<tr>
<td>Outer tube diameter</td>
<td>40.64 mm</td>
</tr>
<tr>
<td>Fin density</td>
<td>236 m$^{-1}$</td>
</tr>
<tr>
<td>Fin height</td>
<td>10 mm</td>
</tr>
<tr>
<td>Fin thickness</td>
<td>1 mm</td>
</tr>
<tr>
<td>Tube pitch</td>
<td>70 mm</td>
</tr>
<tr>
<td>Shell bundle clearance</td>
<td>15 mm</td>
</tr>
<tr>
<td>Shell inside diameter</td>
<td>1014 mm</td>
</tr>
<tr>
<td>Central baffle spacing</td>
<td>507 mm</td>
</tr>
<tr>
<td>Baffle cut ratio</td>
<td>0.20</td>
</tr>
<tr>
<td>Tube to baffle hole clearance</td>
<td>0.8 mm</td>
</tr>
<tr>
<td>Shell to baffle hole clearance</td>
<td>1.6 mm</td>
</tr>
</tbody>
</table>

4.1. Baseline results

Table 6 shows a few baseline results at various pressures and ammonia mass fractions for both the HRB and the OBB, using the pool boiling extension to flow boiling with Táboas et al. [39] and Gungor and Winterton [1].

The heat rate varies from 40 to 57 MW and is largest in the evaporator for both case studies, except that of the HRB at 100 bar and an ammonia mass fraction of 0.9. The heat transfer coefficient of the hot fluid is 6-7 times larger for the OBB compared to the HRB. It means that the heat transfer surface areas are approximately 5 times smaller for the OBB. The cold fluid pressure drop is higher for the HRB than the OBB, because the OBB has twice as many cold fluid circuits and shorter tubes. It also means that the cold fluid mass flux becomes approximately half of that of the HRB. The average heat flux in the evaporator varies from 46 to 66 kW m$^{-2}$ and from 54 to 153 kW m$^{-2}$ for the HRB and the OBB, respectively.

Figure 7 shows the computed temperature profiles of the cold fluid, the hot fluid and the outer tube wall. Again, it indicates that the hot fluid gas-side controls the heat transfer process for the HRB, since the wall temperature is close to that of the cold fluid (Figure 7a). For the OBB (Figure 7b), the wall temperature becomes closer to the average of the cold and the hot fluid throughout
Table 6: HRB and OBB model results

<table>
<thead>
<tr>
<th>Units</th>
<th>HRB</th>
<th>OBB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cold fluid pressure</td>
<td>bar</td>
<td>40</td>
</tr>
<tr>
<td>Bulk ammonia mass fraction</td>
<td>0.5</td>
<td>0.9</td>
</tr>
<tr>
<td>Heat exchange in EC</td>
<td>MW</td>
<td>9.1</td>
</tr>
<tr>
<td>Heat exchange in EV</td>
<td>MW</td>
<td>31.4</td>
</tr>
<tr>
<td>Heat exchange in SH</td>
<td>MW</td>
<td>14.2</td>
</tr>
<tr>
<td>Total heat exchange</td>
<td>MW</td>
<td>54.7</td>
</tr>
<tr>
<td>Cold fluid mass flow</td>
<td>kg s⁻¹</td>
<td>17.4</td>
</tr>
<tr>
<td>Cold fluid mass flux</td>
<td>kg m⁻²s⁻¹</td>
<td>320</td>
</tr>
<tr>
<td>Avg. boiling heat flux</td>
<td>kW m⁻²</td>
<td>58.5</td>
</tr>
<tr>
<td>Average ( h_c ) in EC</td>
<td>Wm⁻²K⁻¹</td>
<td>2430</td>
</tr>
<tr>
<td>Average ( h_h ) in EC</td>
<td>Wm⁻²K⁻¹</td>
<td>14387</td>
</tr>
<tr>
<td>Average ( h_c ) in SH</td>
<td>Wm⁻²K⁻¹</td>
<td>1477</td>
</tr>
<tr>
<td>Average ( h_h ) in SH</td>
<td>Wm⁻²K⁻¹</td>
<td>36.9</td>
</tr>
<tr>
<td>Average ( h_h ) in EV</td>
<td>Wm⁻²K⁻¹</td>
<td>42.4</td>
</tr>
<tr>
<td>Average ( h_h ) in EC</td>
<td>Wm⁻²K⁻¹</td>
<td>48.0</td>
</tr>
<tr>
<td>Heat transfer surface area in EC</td>
<td>m²</td>
<td>15882</td>
</tr>
<tr>
<td>Heat transfer surface area in EV</td>
<td>m²</td>
<td>16154</td>
</tr>
<tr>
<td>Heat transfer surface area in SH</td>
<td>m²</td>
<td>7104</td>
</tr>
<tr>
<td>Total heat transfer surface area</td>
<td>m²</td>
<td>39140</td>
</tr>
<tr>
<td>Cold fluid total pressure drop</td>
<td>bar</td>
<td>1.51</td>
</tr>
<tr>
<td>Hot fluid total pressure drop</td>
<td>bar</td>
<td>0.020</td>
</tr>
</tbody>
</table>

the heat exchanger, thus the heat transfer resistances of the hot and the cold fluids are more even.

Figure 8 presents the heat transfer surface area ratios of the economizer, the evaporator and the superheater at 40 and 100 bar as function of the ammonia mass fraction. The results suggest that the relative size of the evaporator is decreasing with increased ammonia mass fraction for both the HRB and the OBB. Additionally, a higher pressure lowers the relative size of the evaporator. The flow boiling correlations affects only the evaporator size and for that reason, the total boiler area will not show as large area differences as the evaporator may for the different flow boiling correlations.
4.2. Influence of different flow boiling correlations

Figure 9 depicts the predicted evaporator areas for both the HRB and the OBB using the correlations listed in Table 2, for which the areas are normalized by the average predicted area for each conditions (each ammonia mass fraction and pressure).

The results indicate that the area differences are small for the HRB and larger for the OBB. The difference between the highest and lowest predicted areas (maximum area difference) become 6% and 28% for the HRB and the OBB, respectively. The trend with respect to the various correlations
Figure 9: Predicted evaporator areas normalized by the average vs. the different flow boiling correlations (Table 2) for different pressures and ammonia mass fractions; HRB at 40 bar (a), OBB at 40 bar (b), HRB at 100 bar (c), OBB at 100 bar (d).

The results also suggest that the correlation by Rivera and Best [9] always gives the lowest evaporator area, because of its relatively high predicted heat transfer coefficient (Figure 3). The highest area is predicted mostly with the correlation by Khir et al. [10], because of its relatively low heat transfer coefficient (Figure 3 and 4), however for the HRB the correlation by Jung et al. [2] predicts the largest evaporator area at ammonia mass fractions of 0.5 and 0.6.

The influence of using the different flow boiling correlations on the total boiler area is presented in Figure 10 for completeness. The predicted area differences for the total boiler become lower than the evaporator and is related to the evaporator area ratio (Figure 8). The maximum area
For the HRB, the influence of the flow boiling correlations on the required heat transfer surface area is weak during numerical design. However, for the OBB the choice of correlation have larger influence (up to 28 % for the evaporator). If the more elaborate correlations are used such as the classical wide-boiling correlations or the ammonia-water specific pool boiling extensions to flow boiling (excluding the results obtained using the simple correlations by Khir et al. [10] and Rivera and Best [9]), then the maximum difference becomes less than 10 % for the evaporator of the OBB and less than 5 % for the total OBB.
The results indicate that the relatively large differences in the flow boiling heat transfer coefficients as presented in Figure 3 and 4 result in rather small evaporator size differences. The evaporator size differences are almost insignificant for the HRB. For the OBB, the evaporator size differences are less than the typical uncertainties related to the mixture flow boiling heat transfer correlations. These uncertainties may be as high as 40\% and do not necessarily lead to the same uncertainty in the predicted heat transfer area. When excluding the simple Mishra et al. [11] type correlations, the evaporator area difference becomes below 10\% for the OBB. The result is encouraging because it indicates that the use of different elaborate flow boiling correlations for the ammonia-water mixture have minor significance during heat exchanger design.

Figure 11: Relative standard deviation (RSD) of the predicted surface areas of the evaporator and the total boiler; HRB (a) and OBB (b). RSD = \(\frac{1}{\bar{A}} \sqrt{\frac{1}{N-1} \sum_{i=1}^{N} (A_i - \bar{A})^2}\)

Figure 11a and 11b indicate the relative standard deviation (RSD) of the predicted evaporator and total boiler areas. They confirm that the predicted area differences are insignificant for the HRB and small for the OBB when using the correlations listed in Table 2. Additionally, the relative standard deviations do not change significantly with either ammonia mass fraction or pressure. The highest relative standard deviation in the predicted evaporator area is 2\% for the HRB and 9.1\% for the OBB.

5. Discussion
The heat transfer correlations employed in the current work suggest that the nucleate boiling contribution to flow boiling is small compared to the convective contribution to flow boiling. Only at very low mass fluxes, low vapor qualities and high heat fluxes, the nucleate boiling contribution becomes significant for the ammonia-water mixture. For this reason, the different pool boiling mixture correction factors (Figure 2) do not matter much when they are used to predict the mixture flow boiling. In other words, the nucleate boiling is already rather suppressed by the presence of the flow at the current design conditions.

The convective contribution is well predicted with that in a pure fluid correlation with mixture properties [2]. The differences in the predicted heat transfer coefficients in Figure 3 and 4 are thus mainly due to inconsistencies in predicting the convective flow boiling contribution. The classical wide-boiling correlations by Wettermann and Steiner [15] and Jung et al. [2] predict reasonably similar results and are close to the pool boiling extensions to flow boiling, for which the Gungor and Winterton [1] pure fluid correlation was used.

For the above reasons, it is difficult to claim that one method is more appropriate to use than others. The simple Mishra et al. [11] type correlations should be avoided, since these are usually narrowed down to small concentration ranges. It is not important to use an ammonia-water specific pool boiling correlation to predict the nucleate boiling mixture degradation, because the convective boiling contribution dominates for the investigated design conditions. Thus, classical wide-boiling mixture correlations may be used as well.

6. Conclusion

This paper presents a systematic comparison of several flow boiling prediction methods aimed towards the ammonia-water mixtures, in an attempt to quantify their individual influence during heat exchanger design. These prediction methods include: (1) the simple Mishra et al. [11] type correlation, applied/refitted to flow boiling experimental data of ammonia-water mixtures within a narrow composition range, (2) the ammonia-water specific pool boiling correlation extensions to flow boiling using the pure fluid correlation by Gungor and Winterton [1], and (3) the classical wide-boiling correlations used directly for the ammonia-water mixture.
Two case studies related to the use of the Kalina cycle were considered for quantifying the correlations influence on the heat exchanger design: a flue gas based heat recovery boiler (HRB) for a combined cycle power plant and a hot oil based boiler (OBB) for a solar thermal power plant.

From the heat exchanger design simulations, it may be concluded that the different flow boiling correlations resulted in small differences in the predicted heat transfer area of the evaporator, especially when excluding the simple Mishra et al. [11] type correlations. The maximum predicted evaporator area difference compared to the average were 6% and 28% for the HRB and the OBB, respectively. Excluding the simple Mishra et al. [11] type correlations and focusing on the other more elaborate correlations, then the OBB showed below 10% evaporator area difference at maximum. These simulations were performed at various ammonia-mass fractions (0.5 to 0.9) and pressures (40 to 100 bar).

Furthermore, the heat transfer correlations employed in the current work suggest that the nucleate boiling contribution to flow boiling is small compared to the convective contribution to flow boiling for the considered boilers and conditions. Only at very low mass fluxes, low vapor qualities and high heat fluxes, the nucleate boiling contribution becomes significant for the ammonia-water mixtures in the considered range of boiler design conditions.


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