Pressure effect on CHF enhancement in pool boiling of nanofluids

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This paper investigates critical heat flux (CHF) in saturated pool boiling for water and TiO₂ nanofluid on a 7 mm diameter vertical copper surface at pressures of 0.1 to 0.8 MPa. The nanofluid was prepared by dispersing 0.002 wt% TiO₂ nanoparticles in deionized water. The CHF of the nanofluid was enhanced about two times over that of water boiling at atmospheric pressure. With the increasing pressure, however, the CHF enhancement with the nanofluid decreases, and almost disappears at 0.8 MPa.

Key Words; critical heat flux; nanofluids; pool boiling; high pressure; in-vessel retention; experiment

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1. Introduction

Nanofluids are suspensions of nanoparticles dispersed in a base liquid. A unique characteristic in nanofluid boiling is that the critical heat flux (CHF) is significantly higher than the CHF of the base liquid, and on account of this characteristic nanofluids have potential applications as coolants of thermal devices with high heat flux components, including nuclear power reactors. The enhancement of the CHF with nanofluids was reported first by You et al. [1] in 2003. They carried out experiments with pool boiling on an upward facing surface with an Al₂O₃-water nanofluid at 0.02 MPa and found that the CHF with the nanofluid was 3.1 times higher than that with water. Since then, many researchers [2-6] have measured the CHF in nanofluid boiling using various kinds of nanoparticles (SiO₂, Al₂O₃, CuO, and TiO₂), and reported that the CHF is markedly enhanced compared with the CHF of water, similarly to the results by You et al.

It has been established that nanoparticles are deposited on heating surfaces during boiling of nanofluids and the general consensus is that this is likely the cause of the CHF enhancement with nanofluids. Kwark et al. [7] measured the CHF of water boiling on a 10mm × 10mm upward facing heating surface on which the Al₂O₃ nanoparticles were previously deposited and reported that the CHF of water increases with the improvement in wettability (with the decrease in the static contact angle) due to the nanoparticle deposition. Kim and Kim [8] boiled different nanofluids with 0.2mm diameter horizontal wires to deposit layers of the nanoparticles on the wires and concluded that the CHF increase shown by the effect of the nanofluids is a result of both the increase in surface wettability and the capillarity in the porous layer of the deposited nanoparticles. Sakashita [9] measured liquid-vapor behaviors close to a heating surface using a conductance probe during pool boiling of water on an upwards-facing heating surface coated with TiO₂ nanoparticles, and specified the macrolayer thicknesses formed
beneath vapor masses. Sakashita [9] established that the macrolayers formed on the TiO$_2$
coated surface are thicker than those on the uncoated (clean) surface, and concluded that
this is most likely a main cause of the CHF enhancement with nanofluids. Kim et al. [10]
measured the CHF of water boiling on a 0.2mm horizontal wire on which a TiO$_2$
nanoparticle coated layer was previously deposited by previous boiling in nanofluid. The
wettability and thickness of the deposited layer were varied by controlling the heat flux
and duration of this previous boiling. Through these measurements, Kim et al. concluded
that capillarity and additional liquid held in the porous layer (which leads to an increase
in the effective macrolayer thickness) are responsible to the CHF increase. In a similar
study, Park et al. [11] measured the CHF of water on a 0.49mm diameter horizontal wire
on which Al$_2$O$_3$ nanoparticle layers with different thicknesses and porosities were
previously deposited by previous boiling in the nanofluid. Park et al observed that with
the increase in the previous boiling time the CHF increases steeply for short previous
boiling times, and after peaking gradually decreases with longer previous boiling times.
Park et al. mentioned that the steep increase in the CHF at short previous boiling times
can be explained by the increase in wettability and the change in hydrodynamic
instability, and that the gradual decrease in the CHF for longer previous boiling times by
a decrease in the porosity of the nanoparticle layer.

As mentioned above, valuable information has been accumulated for the CHF
enhancement with nanofluids; however, most studies are carried out at atmospheric or
lower pressures. Considering the application of nanofluids in emergency core cooling
systems in nuclear power plants, it is important to know the CHF behavior of nanofluids
at higher pressures. For example, Buongiorno et al. [12] proposed the use of nanofluids
to enhance the in-vessel retention (IVR) capability in management strategies of severe
accidents in light-water reactors. In the IVR strategy, the reactor vessel is flooded with
water if a severe accident occurs and the decay heat of the molten core relocated to the
lower head of the reactor vessel is removed through nucleate boiling on the outer wall of
the reactor vessel. Therefore, if the CHF of the water can be expected to be enhanced by
using nanofluids in a severe accident condition, the use of nanofluids would improve the
IVR safety margin of currently operating reactors and enable the IVR strategy to be
applied to advanced reactors with higher core power densities. Under severe accident
conditions, it is expected that the ambient pressure surrounding the reactor vessel would
be higher than the atmospheric pressure, and therefore it is important to examine the
CHF behavior of nanofluids at higher than atmospheric pressure to assess the capability
and applicability of the IVR strategy.

In research on the CHF of nanofluids at pressures above the atmospheric
pressure, Lee et al. [13] carried out experiments with Fe3O4-water and Al2O3-water
nanofluids boiling on 0.4mm diameter horizontal wires at pressures of 0.1, 0.6, and 1.0
MPa, and reported that the CHF enhancement remains at almost the same value (1.4 to
1.6 times the CHF of water) at the three different pressures. However, it is also well
known that the CHF of fine wires varies in a complicated manner dependent on the wire
diameter [14, 15]. Therefore, there is the possibility that the effect of pressure on the
CHF enhancement using nanofluids will be different for different configurations of the
boiling surfaces. Kwark et al. [7] carried out an experiment with water boiling on a 10
mm×10 mm surface with an Al2O3 nanoparticle coated layer previously deposited by
nucleate boiling of an Al2O3-water/ethanol nanofluid. They measured the CHF at
pressures of 0.02, 0.05, 0.1 and 0.2 MPa, and obtained results where the CHF
enhancement due to the nanoparticle coating gradually decreases as the pressure
increases. To the best of the author’s knowledge, no other research has been reported for
the CHF in boiling with nanofluids at higher pressures.

This study carried out an experiment with boiling of TiO2-water nanofluid on a 7
mm diameter copper heating surface. The heating surface was placed vertically to allow
the observation of the boiling behaviors on the heating surface using high speed video under high pressure condition. The CHF in water and with the nanofluid was measured at pressures 0.1 to 0.8 MPa, and the effect of the pressures on the CHF enhancement using nanofluids was examined.

2. Experiment

The experiments were carried out in saturated pool boiling conditions. Figure 1 shows the heating module used in the present experiments. One end of a copper rod placed in a cylindrical stainless steel housing unit served as the heating surface and it was heated with a cartridge heater inserted into the copper rod. The diameter of the heating surface was 7 mm. Three sheathed thermocouples (0.5 mm diameter) were embedded 5 mm, 12 mm, and 19 mm below the heating surface to determine the surface temperature and surface heat flux. The copper rod and the surrounding 1mm thick stainless steel flange (the end surface of the housing unit) were attached by shrink-fitting. The surfaces of the copper and the stainless steel flange were machined to be flush, and then Ni was electroplated over the two surfaces to avoid corrosion and nucleation of bubbles at the interface between the copper surface and the surrounding flange. The thickness of the Ni plating was about 3μm. Figure 2 shows the high pressure cell. The cell is cylindrical with an inner diameter of 50 mm and a height of 150 mm. The heating module shown in Figure1 was inserted in the cell from the side of the cell, and a sapphire window was provided in the opposite side of the cell. The boiling behaviors on the vertical heating surface were observed via the sapphire window using high speed video with a maximum 2000 fps and a microscope with a maximum 320-fold magnification.
The present experiment used TiO\textsubscript{2} nanoparticles (Sigma-Aldrich) with a vendor specified average size of 25 nm. The experiments with the nanofluid were carried out in the following manner: 0.002 wt% of TiO\textsubscript{2} nanoparticles was mixed with de-ionized water (electro conductivity below 0.2 $\mu$S cm\textsuperscript{-1}) and dispersed in an ultrasonic bath for 30 min. Then the nanofluid was poured into the high pressure cell and decompressed with a vacuum pump to make it boil at room temperature. After boiling at room temperature for several minutes for degassing, the liquid was heated to a saturation temperature at a set pressure level by heaters wound around the high pressure cell and maintained at that temperature by adjusting the input to the heaters and the flow rate of the cooling water through the cooling coils. Then the boiling curves were recorded at stepwise increasing heat inputs to the cartridge heater in the copper heating module. When approaching the CHF, the size of the increments in the heat input was reduced, and the heat flux when the surface temperature was raised rapidly was defined as the CHF. The time from the start of boiling to the occurrence of CHF was about 1 to 1.5 hrs.

During each experiment, the bulk temperature of the liquid was maintained within $\pm0.5$ K of the set saturation temperature. The uncertainties in the saturation pressure caused by this temperature uncertainty increased with the saturation pressure and were estimated as $\pm0.01$ MPa at 0.8 MPa, the maximum pressure in the present experiment. The surface temperature and the surface heat flux were calculated from the three thermocouples embedded in the copper rod by assuming steady state one-dimensional heat conduction. Figure 3 shows examples of temperature distributions at three different heat fluxes for the nanofluid boiling at 0.5 MPa. The heat flux of $q=4.56$ MW m\textsuperscript{-2} is about 1% lower than the CHF. As shown in Figure 3, the linearity of temperatures at the three locations is quite good and the deviation from the linearity was less than 1%. The heat loss by the heat conduction through the surrounding stainless steel flange was estimated by solving the 2-D heat conduction equation and the error due to
the heat loss was determined as less than 4% of the heat flux estimated with the three thermocouples.

<Figure 3>

3. Experimental Results

3.1 Appearances of Water Boiling at Higher Pressures and of Heating Surface after the CHF Measurements with Nanofluid

Figure 4 shows the appearance of boiling for water at 0.8 MPa, the highest pressure in the present experiments. Figure 4(a) shows the boiling appearance at $q=1.1$ MW m$^{-2}$ ($q/q_{CHF}=0.24$) and Figure 4(b) is at $q=3.3$ MW m$^{-2}$ ($q/q_{CHF}=0.73$), with the 7 mm diameter heating surface encircled by a broken line. At atmospheric pressure, large coalesced bubbles completely covering the heating surface are formed at heat fluxes above around 0.5 MW m$^{-2}$ (images are not shown). At higher pressures, the bubble growth rates and the diameters at detachment of the bubbles decrease with the increase in the pressure, and a large vapor mass completely covering the heating surface is not formed at $q=1.1$ MW m$^{-2}$, but there are several smaller coalesced bubbles on the heating surface (Figure 4(a)). At $q=3.3$ MW m$^{-2}$, a large coalesced bubble completely covers the heating surface (Figure 4(b)). In the present experiments with a 7 mm diameter heating surface, the large coalesced bubbles completely covering the heating surface were formed at 40% to 50% of the CHF.

<Figure 4>

Figure 5 shows the appearances of the heating surface after the CHF measurements for the boiling of the nanofluid at three different pressures: 0.1, 0.25, and 0.8 MPa. The appearances of the TiO$_2$ coatings appear somewhat different on the three surfaces probably due to the differences in shape and size of the bubbles and the nucleation site densities, while most of the surface is covered with a TiO$_2$ layer. For the
three surfaces, the static contact angle was measured at room temperature under atmospheric pressure by detaching the heating module from the high pressure cell and by placing a water droplet (3 μL) on the surface. The static contact angles at the three surfaces showed similar values, around 0° to 5°.

3.2 Pressure Effect on CHF Enhancement with the Nanofluid

Figure 6 shows boiling curves for water and the nanofluid at two pressures, 0.1 and 0.6 MPa, with the arrows in the figures indicating the CHF. Figure 6 also shows the predicted results for water with the Stephan and Abdelsalam [16] correlation. In the pressure range of the present experiment, there are no obvious differences in the heat transfer coefficients of water and the nanofluid, and the boiling curves agree fairly well with the results with the Stephan and Abdelsalam correlation. However, the enhancement ratios of the CHF with the nanofluid are quite different for 0.1 and 0.6 MPa.

Figure 7 shows the variation in the measured CHF with pressure for water and the nanofluid. The CHF of the boiling nanofluid is larger than that of water over the whole range of pressures, showing that the CHF of water is enhanced with the TiO$_2$ nanofluid. However, the CHF values of water and nanofluid to approach similar values with the increases in pressure.

Figure 7 also plots the predicted results with the widely used CHF correlation in Equation (1) proposed by Zuber [17] and Lienhard and Dhir [18], and the correlation proposed by Kandlikar [19]. The correlation by Zuber and Lienhard and Dhir is given by

$$q_{CHF} = K \cdot \rho_v^{1/2} H_{fg} \left( \sigma g (\rho_l - \rho_v) \right)^{1/4}$$

(1)
where $K$ is a constant which Zuber derived theoretically as $K=0.131$, and Lienhard and Dhir modified to $K=0.149$. Kandlikar proposed a CHF model incorporating the effects of a dynamic receding contact angle and surface orientation as given by

$$q_{CHF} = \rho_v^{1/2} H_{fg} \left( \sigma g (\rho_l - \rho_v) \right)^{1/4} \left[ \frac{1}{\pi} + \frac{\pi}{4} (1 + \cos \beta) \cos \phi \right]\right]^{1/2}$$

where $\beta$ is the dynamic receding contact angle and $\phi$ is the angle of inclination of the surface measured in relation to the upward position ($\phi=90^\circ$ for the present measurements with the vertical surface). The present experiments did not measure the dynamic receding contact angle, $\beta$, and therefore, $\beta=45^\circ$, the value recommended for water/copper systems by Kandlikar, was used for water. For the nanofluid, $\beta$ was assumed to be $0^\circ$, because the static contact angle measured after the CHF measurements was close to $0^\circ$ as mentioned in the preceding section.

For water boiling, the measured CHF at 0.1 MPa agrees well with the values predicted by Equation (1), at higher pressures however, the measured CHF values are larger than the predicted values and the discrepancy increases with the increases in pressure. Equation (2) predicts lower values than the present data for water and nanofluid and the predicted values by Equation (1) over the whole range of pressures (The main reason of the discrepancy for the CHF of water between Equations (1) and (2) comes from the term in the last parenthesis on the right hand side of Equation (2) expressing the effect of the surface orientation. With $\beta=45^\circ$, Equation (2) predicts the CHF for a vertical surface ($\phi=90^\circ$) as 43% lower than the CHF for a horizontal surface ($\phi=0^\circ$)).

The reason why the present data of CHF for water tends to be different from the predictions of Equation (1) with the increase in pressure is unclear. The present data are obtained for a vertical surface of relatively small size (7mm diameter), and this may lead to a pressure dependence different from Equation (1) derived for an infinitely large upward facing surface. Further study would be necessary to elucidate details of this.
Figure 8 plots the CHF enhancement ratio versus pressure (averaged CHF values of two or three data sets were used for water at 0.1, 0.5 and 0.6 MPa.) The CHF of the nanofluid is about double the CHF of water at 0.1 MPa, but the enhancement ratio decreases with increasing pressure falling to 1.08 at 0.8 MPa. These results show that the effect of the CHF enhancement with nanofluid deteriorates with the increase in pressure and almost disappears at 0.8 MPa.

The CHF enhancement mechanism observed with nanofluids has not been clarified; however, according to previous studies at atmospheric or lower pressures, the improved surface wettability due to the deposition of nanoparticles on the heating surfaces is likely to be one of the main causes of the CHF enhancement. In the following, the results of Figure 8 are examined from the viewpoint of surface wettability under high pressure (high temperature) conditions. The research into wettability at high temperature conditions has been carried out by Bernardin et al. [20] and Hazuku et al. [21]. Bernardin et al. measured the contact angles of water droplets on an aluminum plate in the temperature range of 25 to 175 °C (pressure range 0.1 to 0.82 MPa), and obtained the result that the contact angle, \( \theta \), remains at a constant value for surface temperatures below 120 °C, and that above 120 °C it decreases in a linear manner with the temperature derivative \( d\theta/dT = -0.55 \text{deg.} K^{-1} \). Hazuku et al. measured the contact angles of water droplets on aluminium and stainless steel surfaces at surface temperature up to 300 °C (15 MPa) and obtained results similar to Bernardin et al. for surface temperatures below 200 °C. The surface wettability of water boiling in the present experimental conditions was estimated based on the results by Bernardin et al. and shown in Figure 9. Figure 9 plots the changes in the contact angle of water with pressure, under the assumption that the contact angle remains constant to 120 °C (the measured value at
room temperature, 75°, was used), and that it decreases with the gradient

\[ \frac{d\theta}{dT} = -0.55 \text{deg.} \, K^{-1} \] above 120 °C. In Figure 9, the solid symbols (●) are for the contact angle based on the saturation temperature at each pressure, and the open symbols (○) are based on the measured surface temperature at the CHF. It is found that the contact angles at the CHF (the open symbols (○)) decrease greatly as the pressure increases from 0.1 to 0.4 MPa. From the result in Figure 9, it may be surmised that a main reason why the CHF enhancement effect decreases with the increasing pressure could be that the surface wettability is increased with the increase in pressure, and that the effect of the wettability improvement due to the nanoparticle deposition on the CHF is reduced.

<Figure 9>

4. Conclusions

The CHF for pool boiling of water and TiO₂-water nanofluid on a 7 mm diameter vertical copper surface was measured in the pressure range of 0.1 to 0.8 MPa. The conclusions obtained in the present study may be summarized as follows:

(1) The CHF of water and nanofluid increase steeply with the increase in the pressure.

(2) The effect of the CHF enhancement with the nanofluid decreases with the increase in the pressure and almost disappears at 0.8 MPa.

(3) When the CHF enhancement is assumed to be caused by the improvement in surface wettability due to the deposition of nanoparticles on the heating surface, the cause of the result described in (2) may be that the wettability of the heating surface is increased with the increase in pressure.
Nomenclature

$C$  Specific heat [J kg$^{-1}$ K$^{-1}$]

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

$H_{fg}$  Latent heat of evaporation [J kg$^{-1}$]

$K$  Constant in Equation (1)

$p$  Pressure [Pa]

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

$T$  Temperature [°C]

Greek symbols

$\beta$  Dynamic receding contact angle

$\Delta T_{sat}$  Superheat [K]

$\rho$  Density [kg m$^{-3}$]

$\sigma$  Surface tension [N m$^{-1}$]

$\theta$  Contact angle [degree]

$\phi$  Angle of inclination of the surface measured relative to the upward position [°]

Subscripts

$CHF$  Critical heat flux

$l$  Liquid

$v$  Vapour

$w$  Wall
References


Captions of figures

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Figure 9  Change in contact angle with pressure, estimated from the results by Bernardin et al.
Figure 1 Heating module

Figure 2 High pressure cell
Figure 3 Temperature distributions in the copper heating module
(with boiling nanofluid at 0.5 MPa)
Figure 4 Appearance of water boiling at 0.8 MPa at two heat fluxes

(a) $q=1.1 \text{MW/m}^2$  (b) $q=3.3 \text{MW/m}^2$

Figure 5 Appearance of the heating surface after the CHF measurements with the nanofluid at three pressures

(a) 0.1MPa  (b) 0.25MPa  (c) 0.8MPa
Figure 6 Boiling curves for water and nanofluid at 0.1 and 0.6 MPa
Figure 7 CHF vs. pressure for water and nanofluid

Figure 8 CHF enhancement ratio vs. pressure
Figure 9 Change in contact angle with pressure, estimated from the results by Bernardin et al.