PROJECT TITLE: EXPERIMENTAL INVESTIGATION ON THE EFFECTS OF COOLANT CONCENTRATION ON SUB-COOLED BOILING AND CRUD DEPOSITION ON REACTOR CLADDING AT PROTotypical PWR OPERATING CONDITIONS.

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Proposal Objectives: (1) To obtain experimental data for subcooled pool boiling heat transfer coefficients on Zircaloy clad rods for coolants with varying concentrations of boron and lithium. The sensitivity of crud deposition on bulk subcooling, heat flux, system pressure, and coolant concentration is to be investigated. (2) To obtain a visual record of nucleation site density, bubble size, and nucleation frequency for subcooled boiling in dilute mixtures on Zr-4 clad rods. (3) To develop experimental techniques using neutron depth profiling (NDP) to measure the deposition rate of boron and lithium on Zr-4 rods.
ABSTRACT

Increasing demand for energy necessitates nuclear power units to increase power limits. This implies significant changes in the design of the core of the nuclear power units, therefore providing better performance and safety in operations. A major hindrance to the increase of nuclear reactor performance especially in Pressurized Deionized water Reactors (PWR) is Axial Offset Anomaly (AOA) - the unexpected change in the core axial power distribution during operation from the predicted distribution. This problem is thought to be occur because of precipitation and deposition of lithiated compounds like boric acid (H$_2$BO$_3$) and lithium metaborate (LiBO$_2$) on the fuel rod cladding. Deposited boron absorbs neutrons thereby affecting the total power distribution inside the reactor. AOA is thought to occur when there is sufficient build-up of crud deposits on the cladding during subcooled nucleate boiling.

Predicting AOA is difficult as there is very little information regarding the heat and mass transfer during subcooled nucleate boiling. An experimental investigation was conducted to study the heat transfer characteristics during subcooled nucleate boiling at prototypical PWR conditions. Pool boiling tests were conducted with varying concentrations of lithium metaborate (LiBO$_2$) and boric acid (H$_2$BO$_3$) solutions in deionized water. The experimental data collected includes the effect of coolant concentration, subcooling, system pressure and heat flux on pool the boiling heat transfer coefficient. The analysis of particulate deposits formed on the fuel cladding surface during subcooled nucleate boiling was also performed.

The results indicate that the pool boiling heat transfer coefficient degrades in the presence of boric acid and lithium metaborate compared to pure deionized water due to lesser nucleation. The pool boiling heat transfer coefficients decreased by about 24% for 5000 ppm concentrated boric acid solution and by 27% for 5000 ppm lithium metaborate solution respectively at the saturation temperature for 1000 psi (68.9 bar) coolant pressure. Boiling tests also revealed the formation of fine deposits of boron and lithium on the cladding surface which degraded the heat transfer rates. The boron and lithium metaborate precipitates after a 5 day test at 5000 ppm concentration and 1000 psi (68.9 bar) operating pressure reduced the heat transfer rate 21% and 30%, respectively for the two solutions.

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1 INTRODUCTION

1.1 Overview

To meet the increasing demand for energy, it is essential that the electrical power be generated at greater economy and safer conditions. Many of the existing nuclear pressurized water reactors (PWR) are planning to increase their licensed peak power limits to meet the higher demand for energy. Consequently, to provide better economy and improve the safety of operating nuclear reactors, core designs require higher peaking factors.

The higher power increases the possibility of a portion of core being subjected to subcooled nucleate boiling due to higher temperatures in the core. However, these new demands on reactors contribute to a phenomenon known as the “Axial Offset Anomaly” (AOA), which is the unexpected deviation in axial power distribution in the core from the predicted distribution. AOA is thought to occur when the crud deposits build up on the fuel rod cladding undergoing subcooled nucleate boiling [Frattini et al; 2001]. During subcooled nucleate boiling, several compounds become supersaturated near boiling sites leading to precipitation on the surface of the heater. This effect of subcooled nucleate boiling has a major consequence, especially in case of lithiated compounds of boron such as lithium metaborate (LiBO2) and boric acid (H3BO3) because of its high neutron capture cross section. The high neutron capture cross section of boron in lithiated compounds has a major effect on the local power distribution in a reactor by absorbing the neutrons in the core. The change in the neutron density in the core affects the axial power distribution in the reactor thereby causing “Axial Offset Anomaly”.

To resolve the problem of AOA, it is very important to understand the process affecting AOA. As AOA occurs during subcooled nucleate boiling upon the build up of crud deposits, it is essential to understand the phenomenon of subcooled boiling very well. The crud that builds on the Zircaloy-4 (Zr-4) clad fuel rod surface consists of corrosion products released in the reactor coolant system. The crud deposits are porous and hence affect the nucleation sites on the clad surface which in turn affect the rate of boron precipitation on the surface of fuel rod. The deposition rates are affected by several factors, which include the surface temperature, coolant concentration, and degree of subcooling of the bulk fluid. However, the information available about interactions between subcooled boiling and lithium and boron deposits is very scarce. Hence, an experimental study was conducted to understand the interaction of subcooled boiling and boron and lithium deposition on the surface of zircaloy clad rods.

The experimental study was conducted using specially designed high pressure boiling experimental equipment to measure the subcooled boiling heat transfer coefficient under varying operating conditions under varying operating conditions. The boiling tests were performed at prototypical PWR conditions with an electrical test heater simulating a fuel rod in a core of reactor. The boiling tests were conducted with varying concentrations of boric acid and lithium metaborate solutions with deionized water to measure the effect of coolant concentration on the boiling heat transfer coefficient.
1.2 Working of a PWR

Pressurized water reactor or PWR is a nuclear reactor, where the water is used as a primary coolant as well as secondary coolant. A schematic of working model of the pressurized water reactor is shown in Figure 1-1. The reactor vessel encloses the core, which consist of bundles of fuel rods. The purpose of the primary coolant is to remove the heat generated at the core, which is caused when the energy released in nuclear reactions is transformed by collisions into random molecular motion (heat) [Nero, 1979]. Highly pressurized deionized water is passed through the core in the reactor vessel; the deionized water is then passed through a steam generator where a secondary coolant (water) is allowed to boil, producing steam to run the turbine. Using a secondary coolant helps in making the nuclear reactor safe by preventing any nuclear waste from mixing with steam supplying the turbines. The primary coolant also serves the purpose of neutron moderator by slowing down the neutrons in the reactor vessel. Boric acid is added to the primary coolant to control the power generated in the PWR, as boron is a good absorber of neutrons. The heat transfer from the fuel rods to the primary coolant (deionized water) in the reactor vessel occurs by subcooled boiling. AOA is believed to occur when there is sufficient build up of deposits on the fuel rod under subcooled boiling conditions.
1.3 Organization of Report

This report has been primarily divided into 6 chapters. The first chapter introduces the project. The literature concerning previous research related to AOA is discussed in Chapter 2. The experimental equipment used for obtaining the pool boiling results are explained in detail in Chapter 3 and Chapter 4 gives the procedures employed carrying out the experimental work. Chapter 5 presents the results obtained for pool boiling tests conducted at pressures of 100 psi, 200 psi, 500 psi and 1000 psi. The results of particulate boron and lithium deposits onto the Zr-4 clad are also presented in the Chapter 5. Analysis of results obtained from the pool boiling experiments are developed and presented in Chapter 6. Chapter 7 gives the conclusions generated by this investigation.
2 LITERATURE SURVEY

2.1 Overview

This chapter discusses the previous relevant research work done regarding subcooled pool boiling and effect of particulate deposits on heat transfer.

2.2 Pool Boiling Heat Transfer

“Pool boiling is defined as boiling from heated surface submerged in a large volume of stagnant liquid”[Collier and Thome 1994] If this liquid is at boiling point, it is called ‘saturated pool boiling’, or if the temperature of bulk liquid is below its boiling point, then it is called ‘subcooled pool boiling’. As the surface temperature of the heater exceeds the saturation temperature of the liquid, nucleation begins on heater surface.

The boiling process depends upon the nature of the surface, thermo physical properties of the fluid and vapor bubble dynamics [Sachdeva 2001]. The results obtained from the boiling experiments are usually represented in the form of a “boiling curve” which is a plot of surface heat flux against the heater wall surface temperature (T_w). Boiling curves are sometimes presented with wall superheat instead of wall temperature. Wall superheat is defined as the temperature difference between the wall temperature and bulk fluid temperature (T_b). For boiling tests with coolant at saturation temperature, wall superheat (T_w-T_b) is equivalent to T_w-T_s, where T_s is the saturation temperature. Figure 2-1 shows the pool boiling curve for deionized water at atmospheric pressure. The region AB is the natural convection heat transfer region, where the temperature gradients are set up in a pool.

![Figure 2-1 Pool boiling curve for water at atmospheric pressure [Source: Collier and Thome, 1999]](image)
The onset of nucleate boiling (ONB) is the condition where the wall superheat temperature becomes sufficiently large to cause vapor nucleation at the heater surface. This condition usually occurs close to the meeting point of AB and B’C as shown in Figure 2-1. The B’C in Figure 2-1 indicates the nucleate boiling region where vapor nucleation occurs at the heater surface. The nucleation begins with a few individual bubbles at low heat flux and with increasing heat flux, the vapor structure changes because of bubble coalescence. With further increasing heat flux, vapor patches and columns are formed close to the surface.

The critical heat flux (CHF or point D) indicates the upper limit of nucleate boiling where the interaction between the liquid and vapor streams restrict the liquid supply to the heating surface. The transition boiling region (DE) is marked by the formation of an unstable vapor blanket over the heating surface that releases large patches of vapor at more or less regular intervals. At this point, the nucleation rate becomes so high that the flow of fresh liquid to the heater surface becomes restricted by the vapor film formed, causing a decline in the heat transfer rates. Wetting of the heating surface is only intermittent in this region. The film boiling region (EF) is characterized by formation of a stable vapor film which covers the entire heating surface and vapor is released from the film as regularly spaced bubbles. The heat transfer occurs by conduction and convection through the vapour. Heat transfer occurring through radiation becomes significant as the surface temperature of the heater increases to a very high value.

2.3 Boiling for Mixtures

Kamoshida and Isshiki have investigated the pool nucleate boiling of multi component lithium halide salt solutions at saturation under atmospheric pressure (Kamoshida and Isshiki, 1994). Kamoshida and Isshiki have performed pool boiling tests with binary system solutions of H₂O/LiCl and H₂O/LiBr and ternary system solutions of H₂O/LiCl+LiBr. It was found that the boiling curves of salt solutions, in the higher heat flux regions, have higher degree of superheat than deionized water. Their results also indicated that the mixed salt solutions have higher superheat in the transition region and therefore have lower heat transfer rates than a normal solution. The reduction in heat transfer for mixed solutions was attributed to lower coalescence of bubbles than the other solutions.

The work done on pool boiling indicates that the heat transfer coefficients for mixtures vary significantly from their pure components. For the deionized water-ethylene glycol mixture, system reductions in heat transfer coefficients on the order of 65% were observed on both smooth and the Turbo BIII tubes at a mole fraction of 0.5 (Schnelle, 2002). These experiments were conducted at saturation temperatures near atmospheric pressures. Schnelle noted that the variation in the heat transfer coefficient depends on many factors such as concentration of the solution, level of subcooling, and surface roughness (schnelle, 2002).
2.4 Influence of Particulate Deposits

Two studies have been found indicating that deposits and scaling generated during boiling considerably affect the heat transfer rate. Consequently, nucleate boiling heat transfer may be reduced by scale formation even when the bulk foulant concentration is below its saturation level (Steinhagen and Jamialahmadi 1990). Steinhagen and Jamialahmadi have studied the interaction between scale formation and bubble formation and its effect on heat transfer using calcium sulphate (CaSO₄) solution. It was found that the variation of heat transfer coefficient as a function of time was characterized by a sharp decrease to a minimum, followed next by an increase to a maximum and then a decrease to an asymptotic minimum. Pool boiling tests were performed using a saturated CaSO₄ solution over a period of 130 hours. The heat transfer coefficient decreased by nearly 30% for a constant heat flux of 38.5 kW/m². This change in heat transfer coefficient is attributed to the dissolved and deposited CaSO₄. Particulate deposition occurs due to evaporation at the base of the growing bubbles. Even though nucleation behavior has been studied for pool boiling, there is no generally accepted model to explain bubble growth behavior due to the complexity of boiling heat transfer. With addition of ethylenediaminetetraacetic acid (EDTA) to CaSO₄ solution, the boiling heat transfer coefficients was found higher than pool boiling heat transfer coefficient for saturated CaSO₄ solution under similar test conditions (Steinhagen and Jamialahmadi, 1991). The increase in pool boiling heat transfer coefficient under influence of the EDTA was explained by the decreased calcium deposits on the heater surface, indicating that the deposits of the precipitate has a influence on the heat transfer rates.

Zhao and Tsuruta (2002) studied bubble growth behavior and developed the micro layer theory. The cycle of a bubble consists of two parts, one being the lifetime and other is the waiting time of nucleus activity. “The lifetime of the individual bubble consist of three durations : initial growth duration, final growth duration and the condensation duration before the individual bubble collapses”[ Zhao and Tsuruta 2002].

2.5 Axial Offset Anomaly

The problem of Axial Offset Anomaly is widely reported in many of the operating PWR’s. A report on the performance cycles of the Callaway nuclear power plant indicated an maximum axial offset of about 10% decrease and an axial offset of 6% increase in the power distribution (Konya, et all 1993). This study indicates the deviation from predicted power in cycles 4 and 5 of the Callaway power plant owned by Union Electric (St Louis, Missouri). One trend which was common in cycles 4 and 5 was that the negative deviation in the core axial offset was more significant that the positive deviation from the predicted power generation. Defloor (1993) has explained another significant AOA problem that has occurred in the DOEL Plant in Belgium (Konya, et all, 1993). During the plant’s 11th cycle of operation it was observed that there was a negative deviation from the predicted power and the maximum offset was found to be only about 4%. Further examination of AOA at the DOEL plant has revealed a thick layer of crud deposits on the surface of fuel. Investigations by Union Electric and Westinghouse into the AOA problem at Callaway plant indicated a 25 ppm increase in boron concentration after a power trip during the period when axial offset was observed. This implies that there is a relation between axial offset observed in PWR and boron concentration on the
fuel rod. Even though the performance reports of nuclear power plants indicate a strong relationship between the boron concentration and AOA, there is less research done detailing the how boron concentration affects the axial offset. In order to understand the interaction between coolant concentration and the total power shifts, it is very important to study how the coolant concentration affects the heat transfer rates between fuel rods and the coolant.

The main reason for the AOA problem is believed to be the effect of particulate deposits on subcooled nucleate boiling (Frattini, et. al 2001). Earlier studies indicated that subcooled nucleate boiling causes corrosion deposition in the upper spans of fuel assemblies. Although the main causes of AOA are known in general, it is still not clear how the mechanism occurs. Corrosion deposits tend to occur at the boiling regions. To understand the mechanism of deposition, it is essential to understand the relation between rate of heat transfer and deposition. Even though there have been extensive studies on the effect of some additives like CaSO₄ on heat transfer rate, there has been no significant work done on the effect of boron and lithium metaborate precipitation on heat transfer rate. Hence, an experimental study was proposed to study the effect of coolant solution concentration on heat transfer characteristics at test conditions simulating the fuel rod of a PWR.

2.6 Objectives

The primary objective of this project was to obtain experimental data for subcooled pool boiling heat transfer coefficients on Zircaloy clad rods for coolants with varying concentrations of boron and lithium. The experimental data obtained will include influence of coolant concentration, bulk fluid subcooling, bulk fluid system pressure and heat flux on heat transfer rate. The project also aims at obtaining characteristic information about the deposits of boron and lithium on Zr-4 rods including a visual record of nucleation for subcooled boiling on Zr-4 clad rods.
3 EXPERIMENTAL EQUIPMENT

3.1 Overview

This chapter presents a description of the experimental Equipment used for pool boiling tests. The experimental test facility was designed in such a way that it would simulate the conditions of a fuel rod in the core of a Pressurized Water Reactor (PWR). This was done using a test chamber which could withstand very high pressures typical of a PWR vessel. An electrical test heater was used to simulate the fuel rod of PWR. A specially built ‘DC electrical power supply unit’ was used to power the test heater. A nitrogen cylinder and a compressed gas accumulator were used in combination to increase the coolant pressure to very high values. Pool boiling conditions were assumed for all the experimental tests conducted, because simulating the flow conditions of the primary coolant in a PWR is a very complex task requiring expensive equipment. The details of all the equipment used to perform subcooled boiling tests are discussed in this chapter. However, some subcooled boiling tests were conducted at a pressure of 100 psi, using a preliminary experimental set-up from that of the existing experimental set-up. To differentiate between the test set-up used for lower pressure (100 psi) and higher pressures, the existing pool boiling test equipment is hereafter referred to as “new pool boiling equipment” (higher pressures). This chapter primarily is divided into two parts. The first part describes the preliminary pool boiling test facility and the second part provides a detailed description of the new pool boiling test facility. As some of the equipment used in both preliminary and new pool boiling test setup is the same, they are explained in greater detail under the new experimental set up.

3.2 Preliminary Pool Boiling Test Equipment

The pool-boiling test facility consisted of a high-pressure test chamber, bulk heater, test heater, nitrogen cylinder and a hydro-pneumatic accumulator. A schematic diagram of the pool boiling test facility is shown in Figure 3-1. The experimental investigation necessitates use of a test chamber, which can operate under high pressures. Typically the pressures of the coolant in PWR range from 6.8MPa (1000psi) to 15.5MPa (2200psi) (Loftness, 1964). A unique high pressure vessel with view ports on either side was used for conducting the experimental investigation. The test chamber (pressure vessel) was 40.64 cm in length with outer diameter of 12.065cm and inner diameter of 9.8425cm (Vleet, 1985). The test chamber had two optical ports located on the either side of the chamber facilitating the observation of the bubble formation on the test heater. The clear fused quartz windows were seated in tapered Teflon cushioning. The windows were held tight by clamping bushings with help of eight socket head screws.

The pressure vessel was designed and constructed by Van Vleet (1985) to investigate subcooled and saturated nucleate boiling on thin wires under transient conditions under high pressure conditions. The pressure vessel, made of 316 stainless steel, was used for conducting the experimental investigation. The test chamber (pressure vessel) was 40.64 cm in length with outer diameter of 12.065cm and inner diameter of 9.8425cm (Vleet, 1985). The test chamber had two optical ports located on the either side of the chamber facilitating the observation of the bubble formation on the test heater. The clear fused quartz windows were seated in tapered Teflon cushioning. The windows were held tight by clamping bushings with help of eight socket head screws.

The bulk heater was inserted axially at the center of a flange at one end of the test chamber. The specially built test heater was placed axially at the center of the other flange. The large diameter base and inside mounting of the bulk and test heaters
accommodate high pressures by a compression fitting in the holder base. Copper O-rings [Duniway, 2006] prevent leakage from the test chamber on the bulk heater end. The bulk heater’s function is to raise the temperature of the coolant in the test chamber. A 500 W immersion type cartridge heater manufactured by Watlow Inc [Watlow, 2006] was used for bulk heating of the water for pressures 100 and 200 psi.

![Figure 3-1 Schematic diagram of Pool boiling test Equipment](image)

The test heater had Zr-4 cladding and was obtained from a commercial reactor fuel supplier. Care was taken to prevent any damage to the outer surface of cladding. The heating element of the test heater was made of inconel 718. The test heater had an outer diameter of 9.5 mm (0.374 inch) and a heating element length of 2.5 cm (1 inch). The test heater simulated the conditions of a fuel rod inside a pressurized deionized water nuclear reactor. PTFE O-rings were used for effective sealing between the test heater flange and the pressure vessel. Power supplied to the test and bulk heater was manually controlled with potentiometers to regulate the amount of heat transfer to the fluid inside the test chamber. The test heater and bulk heater were operated on 115 V AC 60 hz power supply.

A combination of a nitrogen cylinder and an accumulator were used to regulate the pressure of the fluid in the test chamber. As the water is incompressible, nitrogen gas is used to indirectly build pressure inside the testing chamber. The gas compressed hydraulic accumulator contained a bladder to prevent the nitrogen from mixing with the deionized water. A pressure regulator connected to nitrogen cylinder controlled the pressure inside the test chamber. A pressure transducer connected to the test chamber
indicated the pressure inside the testing chamber. A relief valve was used to bleed the excessive pressure.

A T-type thermocouple was used to monitor the bulk temperature of the coolant. The voltage across the test heater was measured by a voltmeter connected in parallel. As measuring current requires placing an ammeter in series to the electrical circuit, a one ohm resistor is connected in series to the test heater. The current through the test heater was measured indirectly by monitoring the voltage drop across the resistor. The use of potentiometers helped in better control of electrical power supplied to the test heater which prevented voltage fluctuations.

3.3 New pool boiling test facility

A new electrical test heater was acquired to operate at the higher pressures and higher temperatures. A 100-ampere DC power supply already existing in the lab was used for supplying the power to the heater. Changes were made to the existing pool boiling test setup for operation of the new test heater. A schematic of the new pool boiling test facility is shown in Figure 3-2. Details of each component for the new pool boiling setup are explained later in this chapter.

3.3.1 Test Heater

The new test heater clad with zircaloy drew power from the 23 kW DC power supply unit. Stern Laboratories Inc. (Hamilton, Ontario, Canada) manufactured the heater. Figure 2 is a photograph of the test heater. A specially designed inner sleeve with holes drilled axially for four K-type thermocouples and INCONEL 718 filament was press fit into the Zr-tube. The physical parameters of the heater were not changed from that of previous heater to maintain the same heating configuration. A picture of the test heater with four thermocouples can be seen in Figure 3-3. The physical parameters of the heaters are as follows:

- Zr Sheath O. D =0.374 inches.
- Sheath Overall Length =6.75 inches.
- Maximum Heat Flux = 3 MW/m².
- Heated Length (boiling region) = 1.0 inch.
Figure 3-2 Schematic of new pool boiling test facility.
3.3.2 Electrical DC Power Supply Unit

The electrical DC power supply had a combination of three 25 kVA single phase dry type distribution transformers and six silicon controlled rectifier (SCR) units to supply the required DC output from 3 phase AC input. The power supply system converted 208 Volts AC, 3 phase, 3 wire, and 60 Hz delta connected power supply into a controlled output. “The output ranged from ‘0 to 120’ VDC output at up to 180 amperes” [Hamilton, R C, 2000]. The power supply had operational limits due to the possible damage of the silicon controlled rectifiers (SCR’s) at their lower thermal limit. This unit cannot be operated beyond 130 VDC. The operating range of the power supply unit is represented in Figure 3-4 where the voltage and the current output should be within the envelope given. The load powered by the “electrical DC power supply” was required to have a minimum electrical resistance of 0.42Ω to prevent damage to SCR’s.
The maximum operating current to the load from the “electrical DC power supply” was 180 A. The total heat generation due to electrical losses depended upon the ‘current’ through the resistor and electrical resistance of the resistor. The electrical resistance of the test heater (0.07Ω) does not meet the minimum requirement condition for the operation of the “DC electrical power supply”. Hence, an additional load was added in series to the electrical test heater. The resistor (load) prevented damage to SCR. The power supplied by the “Electrical DC Power Supply” was controlled manually using the keys on ‘Fisher Rosemount DPR950 Controller display’.

### 3.4 Heat Exchanger

The use of ‘electrical DC power supply unit’ was restricted by the electrical resistance of the total load. An additional resistor was added in series to the test heater to ensure that the total electrical resistance of the load is in the operation range of the ‘electrical DC power supply unit’. Due to the high current in the resistor, the heat generated in the resistor (conduction losses) was very high. At a maximum current of 180 A, the heat generated is 18 kW for a resistor with an electrical resistance of 0.55Ω. Hence, a resistor with good heat dissipation characteristics was fabricated. As heat transfer by natural convection is not sufficient, the heat generated in the resistor must be removed by cold water. Inconel tubes served the dual purpose of acting as a resistor in the electric circuit and providing an effective means of heat dissipation. Hence, the name “heat exchanger” is used instead of the conventional “resistor.”

Laboratory tap water was used to cool the inconel tubes. The unique property of inconel is that its electrical resistivity remains fairly constant over a broad range of temperatures. The photograph of the heat exchanger is provided in Figure 3-5.
The heat exchanger design calculations were made to determine the dimensions needed and are presented in the next paragraphs. The length of the inconel tubes was calculated from the electrical resistance required. The resistivity of inconel is known to be 1.03 $\mu\Omega$m at 20$^\circ$C. Even at 100$^\circ$C, its resistivity is 1.04 $\mu\Omega$m. The electrical resistance of a tube is given by the

$$R = \frac{\rho L}{A}$$ \hspace{1cm} (3.1)

where:

- $R$ = resistance of tube in ohms
- $A$ = cross-section of the tube in sq m.
- $\rho$ = electrical resistivity of material (inconel) in $\Omega$m.
- $L$ = length of the tube in meters.

The heat generated had to be removed constantly to avoid the over heating of the inconel tubes. The minimum and maximum allowable electrical resistance of the heat exchanger (resistor) was 0.41$\Omega$ and 0.71$\Omega$, respectively. Considering an electrical resistance of 0.55$\Omega$ (average) for the heat exchanger, the maximum heat generated was 18 kW. Cold tap water was circulated through the inconel tubes to remove the generated heat.
The design characteristics of the heat exchanger tubes were as follows:

- The total electrical resistance of the tubes should lie within the thermal limit of the ‘electrical DC power supply’. Consequently, the electrical resistance of the heater has to be greater than $0.41 \Omega$ and less than $0.71 \Omega$.
- The rate of heat dissipation in the inconel tubes (heat exchanger) is dependent on the flow rate of the water through the inconel tubes.

Turbulent flow was maintained in the inconel tubes to ensure the maximum heat transfer between the tubes and water. Hence, the velocity of the water inside the tubes was high enough such that the Reynolds number was at least 2300 (transition for conduits). A hose of length 15 meters supplied tap water to inconel tubes. As a result, a maximum pressure drop of 40 psi was assumed over total length of hose and tubes. The flow velocity was computed from Darcy’s formula using Moody’s chart for the friction factor.

The following variables were used in the calculations:

- $D_o =$ outer diameter of the tube in meters.
- $D_i =$ inner diameter of the tube in meters.
- $h_t =$ pressure difference between source and outlet of water in meters of hg.
- $P_1 =$ Supply pressure of water at inlet to the heat exchanger.
- $P_2 =$ Outlet Pressure of water in heat exchanger.
- $f =$ friction factor.
- $L =$ Total length of the Inconel tubing.
- $Re =$ Reynolds number
- $\rho =$ Density of the fluid in kg/m$^3$
- $\mu =$ Dynamic viscosity of the fluid.
- $Nu =$ Nusselt Number
- $Pr =$ Prandtl Number
- $K =$ Thermal conductivity of the fluid in W/mk.
- $Q =$ flow rate in m$^3$/sec.
- $A =$ hollow cross section of the Inconel tubes.
- $V =$ velocity of the water flowing through the tube.
- $Q =$ Water flow rate in heat exchanger in m$^3$/sec
- $K_{water} =$ Thermal Conductivity of water at 300K $=0.613$ W/mk.
- $E_{max} =$ Maximum heat generated in Inconel tubes.
- $T_i =$ Inlet Temperature of water entering the heat exchanger.
- $T_o =$ Outlet Temperature of water leaving the heat exchanger.
- $A_{surf} =$ Surface area of the heat exchanger

The flow calculations were performed with an intial assumption of $D_o = 0.25”$ and $L=1m$. The final dimensions of the inconel tubes were obtained by iterating the initial guess values. The iterations were done with tube diameter varying from 0.25 inch to 1 inch.

$$\Delta P = P_1 - P_2$$ (3.2)
The pressure of the tap water supply in the laboratory was 40 psig. The outlet pressure was atmospheric pressure and the pressure head was calculated from equation (3.2). The pressure head difference in metres is \( h_i \). The friction factor \( f \) was initially guessed and corrected with iterations. The velocity of the water inside the tube was calculated using Darcy’s formula using equation (3.3).

\[
V = \sqrt{\frac{2gD_i h_i}{fl}}
\]  

(3.3)

The Reynolds number was computed from equation (3.4) once the flow velocity and physical properties of the fluid were known.

\[
Re = \left( \frac{VD_i \rho}{\mu} \right)
\]  

(3.4)

If Reynolds number was greater than 2300 (critical Reynolds number for internal flow) it was turbulent flow. Moody’s chart gives a graphical relation between the Reynolds number, roughness factor, and friction factor. The roughness value for tubes was assumed as 0.004 based on the surface finish of the tubes. Hence, the friction factor was estimated from Moody’s chart.

The Dittus-Boelter equation (3.5) was used to determine the Nusselt number for turbulent flow in tubes.

\[
Nu = 0.023 Re^{\frac{4}{5}} Pr^n
\]  

(3.5)

where \( n \) is 0.4 for heating (surface temperature \( T_s > T_b \) bulk temperature) and \( Pr \) for water is 5.8466. The water flow rate can be calculated from the velocity by equation (3.6)

\[
Q = A*V
\]  

(3.6)

The coefficient of convective heat transfer \( h \) can be determined from Nusselt’s number using the equation (3.7)

\[
h = \left( \frac{Nu K_{water}}{D_i} \right)
\]  

(3.7)

The maximum heat generated is 18 kW. For the cooling to be efficient, the outlet temperature of water through the inconel tubes should be less than its boiling temperature. The heat lost by the inconel tubes is gained by the flowing tap water. Using the energy balance equation, the outlet temperature of water from the inconel heat exchanger was calculated using

\[
T_o = T_i + \frac{E_{max}}{hA_{surf}}
\]  

(3.8)

Iterations were carried out using the equations (3.1) to (3.8) until the electrical resistance of heat exchanger was higher than 0.41\( \Omega \). A sample calculation is included in the appendix. It was found that an inconel tube diameter of half inch and length of thirty feet
would satisfy the requirements. A tap water flow rate of 2.9gpm was found to be
sufficient to cool the inconel tubing.

To fabricate the heat exchanger, the 30 ft long inconel tube was cut into 6
pieces of 5 feet each. The inconel tubes were laid parallel, ends joined by copper tube
bends as seen in Figure 3.5. Wooden frames supported the tubes and two garden hoses
transported the tap water flow to and from the heat exchanger.

3.5 **Bulk Heater**

A bulk heater was used to control the bulk temperature of the fluid. As the boiling
tests were conducted at higher pressures, the saturation temperatures are also high; hence
more power was needed to raise the temperature of fluid. So the preliminary bulk heater
of 500W capacity was replaced with a specially built 2 kW “Fire” rod immersion type
heater (Watlow, Inc.) suitable for operation at pressures up to 2000 psi. The input power
to the heater was controlled using a conventional rheostat. A photograph of test heater is
given in Figure 3-6.

![Bulk Heater](image)

Figure 3-6 Bulk heater with flange and copper gasket for the preliminary test
equipment.
3.5.5 Electric Circuit

The DC power supply supplied electrical current to the test heater and heat exchanger which were connected in series. The resistance of the test heater was very low at about 0.07\,\Omega. The total resistance of the load was within the operational limits of the DC power supply unit. A special control unit on the DC power supply was used in metering the power input to the heater. A voltmeter in conjunction with the readout of the control unit indicated voltage across the load and total power supplied to the load. The test heater was a sheath return type heater. It is designed in such a way that only the positive terminal of power supply was connected directly to the heater. The negative terminal was connected to the test chamber. The test chamber was electrically grounded. Sheath return type heaters are more commonly used for high current applications.

3.5.2 High Speed Camera

To understand the phases of the bubble growth at the nucleation sites, a high-speed camera was used. Because the bubbles rise and collapse very quickly, it is a tough task to identify the exact growth process of bubbles rising from the heater surface. A high-speed camera can capture images at a relatively fast rate compared to a conventional movie camera. Some high speed cameras can capture images at speeds of 12,000 pictures per second (PPS) where conventional movie cameras capture images at a speed of 30 PPS.

The high-speed camera used here was a ‘HYCAM model # 41-0005’ manufactured by Redlake Corp and was used for observing the bubble growth behaviour. A photograph of the Hycam is provided in Figure 3-7. It was a high speed 16mm motion picture camera with a rotating prism. The maximum speed of the camera was 11,000 half frames per second. The film transport had a maximum film capacity of 400 feet of standard thickness film where spools of 100’, 200’ and 400’ films can be loaded in the camera. The PPS dial in the high-speed camera was used to set the frame rates as per the requirement. The capture speed of the camera was adjusted by rotating the dial in conjunction with the multiplier switch. The minimum speed was 20 frames per second.
3.5.3 Pressure Transducer

A pressure transducer was used to monitor the bulk fluid pressure inside the test chamber. The pressure transducer used for the preliminary pool boiling set up was replaced with a pressure transducer measuring gauge pressure, Model PX35K1-3KGV (Omega Engineering, Inc.). The pressure transducer was calibrated for a range from 0 to 3000 psig and had an operating temperature range from 15 to 70°C. A stainless steel tube connects the pressure transducer and test chamber separating the transducer from the heat generation in the test chamber. A pressure transducer was preferred over a pressure gauge for better accuracy. The calibration data for transducer is provided in Table 3-1. The excitation for the transducer was 10 V DC. The accuracy of the transducer is 0.25% which includes linearity, hysteresis and repeatability. The characteristics this pressure transducer are given in Table 3-2.

The transducer was connected to a strain gauge panel meter to read the pressure. The DP25-S strain gauge panel meter (Omega Engineering, Inc.) used the calibrated
output voltage from the transducer to display the pressure in psi. The DP25-S panel meter had a digital display and its broad scaling capability allowed for the display of most engineering units (Omega Engineering 2002). The strain gauge panel meter also had a facility for taring the output value.

<table>
<thead>
<tr>
<th>Reading</th>
<th>Pressure (Psig)</th>
<th>Transducer Output Data (mV DC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.0</td>
<td>0.000</td>
</tr>
<tr>
<td>2</td>
<td>1500.0</td>
<td>15.031</td>
</tr>
<tr>
<td>3</td>
<td>3000.0</td>
<td>30.004</td>
</tr>
<tr>
<td>4</td>
<td>1500.0</td>
<td>15.055</td>
</tr>
<tr>
<td>5</td>
<td>0.0</td>
<td>0.015</td>
</tr>
</tbody>
</table>

Table 3-1 Calibration data for Pressure transducer (Omega Engineering Inc.)

calibration sheet

<table>
<thead>
<tr>
<th>Item</th>
<th>Parameters</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Balance</td>
<td>0.310 mVdc</td>
</tr>
<tr>
<td>2</td>
<td>Sensitivity</td>
<td>30.004 mVdc</td>
</tr>
<tr>
<td>3</td>
<td>Input Resistance</td>
<td>350.3 ohms</td>
</tr>
<tr>
<td>4</td>
<td>Output Resistance</td>
<td>351.0 ohms</td>
</tr>
</tbody>
</table>

Table 3-2 Pressure transducer characteristics (Omega Engineering, Inc.)
4 EXPERIMENTAL PROCEDURE

4.1 Solution Preparation

Boiling tests were performed with pure deionized water, boric acid solution and lithium metaborate solution with various concentrations. PPM or parts per million is defined as the number of solute particles per million parts of solution. Consequently,

$$PPM = \left( \frac{\text{Mass of solute}}{\text{Mass of solution}} \right) \times 1000000 \quad (4.1)$$

and so one ppm is equivalent to 1 milligram of solute per liter of deionized water (mg/l). The weight of the solute may be calculated from equation (4.1) when the parts per million and weight of the solution are known.

Boiling heat transfer experiments were conducted at 500 PPM, 1000 PPM, 2000 PPM and 5000 PPM concentration levels. Measurement of the weight of the solute (boric acid and lithium metaborate) was done using a weighing scale. This weighing scale could measure a maximum weight of 2.61 kg and had a least count of 0.1g. The boric acid solution was prepared by mixing the measured quantity of boric acid powder into the measured quantity of distilled deionized water. Typically, for every pool boiling test conducted, the coolant required was about a gallon. The weight of solute corresponding to a solvent volume of one gallon is presented in Table 4-1.

<table>
<thead>
<tr>
<th>Solution no.</th>
<th>Coolant Concentration (PPM)</th>
<th>Weight of the solute (g/gallon)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>500</td>
<td>1.9</td>
</tr>
<tr>
<td>2</td>
<td>1000</td>
<td>3.8</td>
</tr>
<tr>
<td>3</td>
<td>2000</td>
<td>7.6</td>
</tr>
<tr>
<td>4</td>
<td>5000</td>
<td>19</td>
</tr>
</tbody>
</table>

Table 4-1 Coolant concentration and weight of the solute

4.2 Boric Acid

Boric acid is a white powder soluble in water. The chemical properties of boric acid are presented below obtained from its Material Safety Data Sheet (MSDS-Fisher Scientific, 2005).

Physical State: Solid (powdered)

pH value: 3.6-4.0 (4% aqueous solution)

Solubility: 4.9g/100g in water @ 20°C

Specific Density: 1.44 (water=1.0)

Molecular Formula: $\text{H}_3\text{BO}_3$
Molecular Weight: 61.83

Boric acid solution in deionized water was used as coolant to determine the pool boiling heat transfer coefficient. The boric acid solution boiling tests were performed with varying concentration of 500 ppm, 1000 ppm, 2000 ppm and 5000 ppm, respectively.

### 4.3 Lithium Metaborate

The physical and chemical properties of lithium metaborate are presented below obtained from its Material Safety Data Sheet (MSDS, Fischer 2005)

- **Physical State:** Crystalline Powder
- **Density:** 1.397 g/cm³ at 20°C
- **Molecular Formula:** LiBO₂
- **Melting Point:** 845°C
- **Molecular Weight:** 49.75

The boiling tests were performed with the same concentrations as for boric acid, namely 500 ppm, 1000 ppm, 2000 ppm and 5000 ppm lithium metaborate solution in deionized water.

### 4.4 Experimental Procedure

The experimental procedure to obtain the boiling heat transfer test data for the new setup is as follows:

1. The pressure vessel, test heater and bulk heater were cleaned with acetone to remove any deposits, settled due to earlier experiments. The cleaning of the surface ensures that all the boiling tests are performed at similar conditions.

2. The flanges of the pressure vessel are firmly fastened with eight bolts. Copper o-ring provides the necessary sealing arrangement. Care was taken to avoid leakage as improper fitting of the o-ring may lead to mechanical failure at high pressures.

3. The solution for the boiling test was prepared. A weight balance was used to measure the quantity of the solute in solution. The test chamber was filled with the prepared solution through the filling port.

4. Check bulk fluid temperature measurement. The temperature of the bulk fluid was measured with a T-type “quick disconnect” thermocouple (Model No. HTQIN 316G-12, Omega Engineering). The temperature reader on measurement panel indicates the temperature of the test heater.

5. Check test heater surface temperature measurement. Four K-type thermocouples, which measured the surface temperature of the electrical test heater, were connected to the thermocouple reader (DP-24 T, Omega Engineering) mounted on the measurement panel.
6. Check bulk fluid pressure transducer. The pressure of the coolant was measured from the transducer connected to the test chamber through a 3/16” stainless steel tube. The output from the transducer was fed into the DP-25 strain gauge panel meter.

7. The pressure of the coolant in the test chamber was then increased to the required level. The required pressure was obtained by pressurizing the gas side of accumulator from a nitrogen cylinder. The pressure was controlled by a regulatory valve on nitrogen cylinder. This results in increase of the pressure of the solution inside the test chamber.

8. After the solution was pressurized to the required pressure, the electrical bulk fluid heater was turned on to heat the solution. The amount of heat supplied to the solution was controlled by regulating the power supplied to bulk fluid heater using a rheostat. The temperature of the bulk fluid (coolant) was maintained at a constant temperature thus sustaining a specific subcooled condition. For example, a 20°C subcooled condition at 1000 psi pressure requires a temperature of 264.8°C which is 20°C below the saturation temperature of 284.8°C.

9. Cold water was circulated through the inconel heat exchanger to remove the heat generated due to its electrical resistance.

10. Power was supplied to the electrical DC power supply by turning on the 3-phase, 240 volts, 100 Amps AC input. The power input to the test heater is slowly increased using the control unit on the DC power supply panel. As the input power to the test heater was increased, the power supply to the bulk heater was decreased keeping the bulk temperature constant.

11. Measure the voltage drop across the electrical test heater using the multimeter. The voltmeter on the ‘electrical DC power supply unit’ panel measured the voltage drop across the total load.

12. Measure the electrical current flowing through the test heater using the clamp type ammeter (Fluke 2005).

13. The power supplied to the test heater was increased in steps and the measurements mentioned in steps 4, 5, 6, 11 and 12 were recorded at each step only after the test heater temperatures reached steady state operation.

14. The test chamber was brought back to ambient conditions after the test was conducted by gradually decreasing the power supplied to the test heater and bulk heater, respectively.

15. Finally, the pressure of the coolant was decreased by bleeding off the nitrogen gas in the accumulator.

4.5 Equipment Maintenance

This section discusses the details concerning equipment failures that occurred during the subcooled pool boiling tests and the precautions that need to be taken for proper functioning of pool boiling experimental equipment.
The test heater hook up wire was burnt causing sparks, disrupting the power circuit of the boiling set up. It was found that improper thermal insulation caused the two power supply wires to come in contact shorting the circuit. As the temperature of the pressure vessel rises to about 250°C at 1000-psi pressure, the power cables should have excellent electrical and thermal insulations. A fiberglass sleeve and high temperature resistant thermal tape was found to adequately insulate the power supply wires.

The quartz window in the optical view port, which was subjected to high pressure and high temperature, has a large thermal gradient across its thickness. The quartz window used in the pool boiling tests failed, even when the operating temperature was below the maximum working temperature of quartz window. The cracking of quartz window has been observed thrice during the operation of tests. The reason for breakage is likely due to the additional stress on the quartz window due to the unequal expansion rates of stainless steel housing and quartz. It was found that the failure could be due to improper cushioning and failure of the gasket. A new gasket (size 1.75” OD X 1.0” ID X 0.0625” thickness) made of “Non Asbestos C4401” was used to replace the failed gasket (Ernest 2005). The ”Non Asbestos C4401” material is a high temperature resistant and high pressure resistant material. These gaskets are manufactured by combining non–asbestos fiber with rubber. The non–asbestos C4401 can operate at a maximum temperature of 399°C. Replacing the gaskets has fixed this problem to a large extent. However, care should be taken to avoid sudden changes in temperatures of the test chamber as it may lead to cracking of the optical windows.

### 4.6 Data Reduction

The measured parameters in the pool boiling test were the bulk fluid temperature in the test chamber, surface temperatures of the test heater, the current flowing through the heater, the pressure inside the test chamber and the total voltage drop across the test heater and the heat exchanger. These measurements were used in the determination of heat flux and temperature difference between test heater surface and bulk fluid. The details of calculations for determination of heat flux are presented below. The variables and constants used in the calculations of heat flux are as follows:

- \( R_f \) = diameter of heater element (filament) = 0.0031111m
- \( R_{CI} \) = copper sleeve radius = 0.004178m
- \( R_o \) = outside diameter of heater rod (cladding) = 0.00475m
- \( Z_{LB} \) = boiling region length = 0.0254 m
- \( Z_{LNC} \) = \( Z_{LB} \) = natural convection length =0.0254 m
- \( Z_{EER} \) = copper sleeve length = 0.0508 m
- \( A_{CU} \) = cross – sectional area of copper sleeve
- \( V \) = voltage drop across heater in volts
- \( I \) = electrical current flowing though the heater in amperes
R = resistance of the test heater = 0.07Ω

\[ T_{CAVE} \] = average temperature of heater surface which is determined from four thermocouple temperatures \( T_{C1}, T_{C2}, T_{C3}, T_{C4} \)

\[ T_{KAVE} = T_{CAVE} + 273.15 \text{ K} \]

\[ T_{BAVE} = \text{bulk fluid temperature (°C)} \]

\[ R_C = \text{thermal contact resistance between copper sleeve and zr-4 clad (m}^2 \cdot \text{K/W)} \]

\[ R_{TOT} = \text{total thermal resistance (m}^2 \cdot \text{K/W)} \]

\[ Q = \text{total input power to the test heater. (W)} \]

\[ q” = \text{heat transfer flux (W/m}^2 \) \]

\[ Q_{LOSS} = \text{heat loss in cladding due to conduction in copper sleeve (W)} \]

\[ K_{COND_CU} = \text{thermal conductivity for copper} = 391.0 \text{ W/(m. K)} \]

\[ K_{COND_ZR} = \text{thermal conductivity of zircalloy (W/m.K)} \]

\[ K_{COND_ZR} = 7.51 + 0.0209T_{KAVE} – (1.45 \times 10^{-5}) T_{KAVE}^2 + (7.67 \times 10^{-9}) T_{KAVE}^3 \]

The electrical power input to the test heater is given by equation (4.2). The current was measured from a special ammeter, which was used to measure high currents by measuring the magnetic field intensity around the power cable.

\[ Q = I^2 R \quad (4.2) \]

The measurements from the four thermocouples located between the cladding and copper sleeve of the test heater were used for computing the average surface temperature of test heater. The four thermocouples were located at a 90° angle to each other from the center of heater. The average temperature of the test heater is given by

\[ T_{CAVE} = \left( \frac{T_{C1} + T_{C2} + T_{C3} + T_{C4}}{4} \right) \quad (4.3) \]

The actual wall temperature of the test heater was determined by applying steady state heat conduction equation. The wall temperature of heater was slightly less than average thermocouple temperature because of the thermal resistance. The wall temperature was computed using

\[ Q = \left( \frac{T_{WALL} - T_{CAVE}}{R_{TOT}} \right) \quad (4.4) \]

\[ T_{WALL} = T_{CAVE} - QR_{TOT} \quad (4.5) \]

The thermal resistance was due to cladding material between the thermocouples and outer wall of test heater. As heat transfer occurs in radial direction, the thermal resistance \( R_{TOT} \) was calculated by using steady state heat equation for a hollow cylinder. The thermal resistance of a hollow cylinder is given by equation (4.6) (Sachdeva, 1988).
\[ R_{\text{thermal}} = \frac{\ln \left( \frac{r_{\text{outer}}}{r_{\text{inner}}} \right)}{2\pi kL} \]  

(4.6)

Rewriting equation (4.6) with test heater parameters, we obtain

\[ R_{\text{TOT}} = \frac{\ln \left( \frac{R_O}{R_T} \right)}{2\pi k_{\text{CONDZR}} Z_{LB}} + R_C \]  

(4.7)

where \( R_C \) is thermal contact resistance which was calculated from natural convection cooling tests. The contact resistance is given by

\[ R_C = \frac{0.0000692}{2\pi R_T Z_{LB}} \]  

(4.8)

Substituting (4.7) equation in equation (4.5) gives the wall temperature of the test heater

\[ T_{\text{WALL}} = T_{\text{CAVE}} - Q \left[ \frac{\ln \left( \frac{R_O}{R_T} \right)}{2\pi k_{\text{CONDZR}} Z_{LB}} + R_C \right] \]  

(4.9)

The heat transfer \( Q \) was corrected for conduction loss through the copper conductor connected to test chamber. The corrected \( Q \) is given by equation (4.10) where \( Q_{\text{CORRECTED}} \) is the actual heat transfer occurring from the test heater to the bulk fluid.

\[ Q_{\text{CORRECTED}} = Q - Q_{\text{LOSS}} \]  

(4.10)

The heat loss was calculated by

\[ Q_{\text{LOSS}} = k_{\text{CONDCU}} A_{\text{CU}} \Delta T_L \]  

(4.11)

The temperature of coolant is the bulk fluid temperature. The heat transfer is directly proportional to the temperature difference between the wall temperature of test heater and the bulk fluid temperature. This temperature difference \( \Delta T_{\text{DEL}} \) is defined by

\[ \Delta T_{\text{DEL}} = T_{\text{WALL}} - T_{\text{BAVE}} \]  

(4.12)

The heat flux across the test heater was represented as follows

\[ q'' = \frac{Q_{\text{CORRECTED}}}{2\pi R_O Z_L} \]  

(4.13)

The coefficient of convective heat transfer was computed using

\[ h = \frac{q''}{\Delta T_{\text{DEL}}} \]  

(4.14)

A FORTRAN program was used to compute the heat flux and coefficient of convective heat transfer incorporating the necessary equations. Dr. Steve Bajorek and Dr. Ken Shultis wrote this program for data reduction for pool boiling heat transfer analysis. Changes were made to the program to suit to the present test heater conditions and is noted to produce two output files. This program is given in the appendix.
4.7 Uncertainty

The thermocouples used to measure the surface temperatures of the electrical test heater had an accuracy of .1°C. The thermocouple used for measuring the bulk temperature of the liquid had an accuracy of 0.2°C. The thermocouple measurements influence the superheat uncertainty. The tolerance of the test heater diameter is .002 mm. The pressure transducer had an accuracy of .25%. The error percentage in the measurements of current and voltage are 3% and 2%, respectively. The maximum deviation in the heat flux due to uncertainty in the measurements is therefore 4.5%. The maximum deviation in the superheat value is 2.5% and the uncertainty in the heat transfer coefficients due to the inaccuracies of the instrumentation is 7.2%.
5 RESULTS

5.1 Boiling Curves

Subcooled boiling solution tests were performed at pressures of 100 psi, 200 psi, 500 psi and 1000 psi. All the pressures indicated in the thesis are in absolute scale. The solution boiling tests were performed at subcooling temperatures of 30°C, 20°C, 10°C and at saturation temperatures. The boiling tests performed using deionized water serve as reference for comparison with the solution boiling tests of boric acid and lithium metaborate.

5.2 Tests at 100 psi (6.9 bar)

Boiling tests were performed using the preliminary experimental set up (with AC electrical power supply) at 100 psi (6.9 bar). The variation in the heat transfer characteristics are observed from boiling curves where each test is numbered for reference.

Boiling tests were conducted with pure deionized water and with varying concentrations of boric acid and lithium metaborate solution. The results for tests done with deionized water are shown in Figure 5-1 showing the variation of heat flux with superheat. The boiling curves are plotted for conditions with coolant at 30°C subcooling, 20°C subcooling, 10°C subcooling, and at saturation conditions. The abscissa in Figure 5-1 represents the temperature difference between test heater surface and the bulk coolant (deionized water) which is also called superheat. The ordinate in Figure 5-1 represents the heat flux generated by the electrical test heater in kW/m².

![Figure 5-1 Boiling curve for deionized water at 100 psi pressure (6.9 bar).](image-url)
The results of tests numbered 651 through 654 for deionized water can be seen in Figure 5-1. From Figure 5-1, observe that boiling curves for all subcooling conditions match well for lower heat flux. It can also be observed that for same heat flux, the degree of superheat is different indicating that the heat transfer coefficient decreases with increase of subcooling. The maximum heat flux is about 250 kW/m² for all the subcooling curves, which is below the critical heat flux for deionized water. The boiling curve for test 651 increases linearly until 70 W/m² heat flux and then rises steeply. This behavior can be explained by a change in phase of the water at surface of heater. The heat transfer between the heater and the coolant occurs by natural convection until the onset of nucleate boiling in the liquid. Beyond the saturation temperature, the heat transfer between heater and coolant occurs through nucleate boiling. The beginning of bubble formation is termed as nucleation. As the temperature of coolant is below its saturation temperature, the process is called subcooled nucleate boiling. The increase in the heat transfer rates is due to subcooled nucleate boiling. Similar phenomenon is observed for boiling curves at different subcooling rates.

![Boiling curve for 500 ppm boric acid solution at 100 psi pressure (6.9 bar)](image)

Figure 5-2 Boiling curve for 500 ppm boric acid solution at 100 psi pressure (6.9 bar)

Figure 5-2 shows results for tests done from #661 to 664 for 500 ppm concentrated boric acid solution. The results follow a similar trend to that of Figure 5-1 except for a small variation in superheat values. The boiling curves for 500 ppm boric acid and deionized water are compared in Figure 5-3. The boiling curves for 500 ppm boric acid tests coincide very well with that of deionized water for lower heat flux values. This indicates that addition of boron at 500ppm concentration does not have significant effect of heat transfer rates in the natural convection region. Upon keen observation, it can be found that the boiling curves for 500 ppm boric acid solution appear to have a slightly higher degree of super heat for similar heat flux values as that obtained with deionized water. The difference in boiling curves for deionized water and 500 ppm boric
acid is evident from Figure 5-3. It can also be noted that the change in superheat values for 500 ppm boric acid tests is small when compared to the actual superheat values for deionized water. However, the small increase in superheat values is due to a change in fluid physical properties which causes less nucleation than deionized water. It can be concluded that boric acid slightly reduces the heat transfer rate at concentration level of 500 ppm and at 100 psi pressure for nucleate boiling.

![Figure 5-3 Comparison of boiling curves for deionized water and 500 ppm boric acid solution at 100 psi (6.9 bar).](image)

The test results with 500 ppm lithium metaborate solution are presented in Figure 5-4. It can be observed that there is a change in the superheat values of 500 ppm lithium metaborate solution with respect to deionized water but it is not significant. However, degradation of the boiling heat transfer coefficients seems to be marginally more in the case of lithium metaborate than boric acid for the same concentration of 500 ppm. This signifies that the lithium metaborate also slightly reduces the heat transfer rate at a concentration of 500 ppm at 100 psia pressure under nucleate boiling. The degradation in heat transfer due to the presence of lithium metaborate at 500 ppm concentration can be observed in Figure 5-5.

However degradation of boiling heat transfer coefficients seems to be marginally more in case of lithium metaborate than boric acid for the same concentration of coolant. For instance, at a heat flux of 164 kW/m², the solution of 500 ppm boric acid (test #661) at 30 °C subcooling resulted in 4.5% decrease in heat transfer coefficient and a reduction of 4.8% in heat transfer coefficient case of 500 ppm lithium metaborate (test #666) at 30 °C subcooling. The reference for comparison of heat transfer coefficients is test #651 for deionized water, which has a boiling heat transfer coefficient of 39.72 kW/m²°C at a similar heat flux value and 30 °C subcooling.
Figure 5-4 Boiling curves for 500 ppm lithium Metaborate solution at 100 psi (6.9 bar)

Figure 5-5 Comparison of deionized water and 500 ppm concentrated lithium metaborate test results.
Figure 5-6 Boiling curve for 2000 ppm boric acid solution at 100 psi pressure (6.9 bar).

Figure 5-7 Boiling curves for 2000 ppm lithium Metaborate solution at 100 psi (6.9 bar).
Figure 5-8 Degradation of boiling heat transfer coefficients with 2000 ppm lithium metaborate solution at 200 psi (6.9 bar).

Figure 5-9 Boiling curve for 5000 ppm boric acid solution at 100 psi pressure (6.9 bar).
Test results with 2000 ppm boric acid solution are presented in Figure 5-6 where considerable change in pool boiling heat transfer coefficients with 2000 ppm boric acid solution occurs. Figure 5-7 shows the results of the boiling tests with 2000-ppm lithium metaborate solution and Figure 5-8 displays the same results in terms of the heat transfer coefficient. For example, with a heat flux of 250 kW/m², the boiling heat transfer coefficient decreased by 20% when compared to deionized water at saturation temperature.

Test results for 5000 ppm boric acid solution are presented in Figure 5-9. As with the results of other boiling curves, it is observed that all four boiling curves coincide at lower heat fluxes indicating that the concentration level of coolant has no effect on the natural convective heat transfer. A comparison of the test results for test 694 and 654 is shown in Figure 5-10 at the 10 °C subcooled condition. It is clear that the temperature gradient between the heater wall and coolant solution is considerably higher in the case of 5000 ppm boric acid solution. The decrease in the heat transfer coefficient is due to the reduced nucleation which was not uniform and unlike that of deionized water leading to lower heat transfer rates.

A reduction of 24.2% is shown in Figure 5-10 in the heat transfer coefficient for the 5000 ppm boric acid solution (heat flux = 153 kW/m²) at saturated conditions compared to the boiling heat transfer coefficient of deionized water (13.08 kW/m²°C, heat flux = 153 kW/m²). Similarly a reduction of 25.5% is seen for the heat transfer coefficient with 5000 ppm lithium metaborate solution (heat flux = 154 kW/m²) as shown in Figure 5-12 when compared with that obtained from deionized water at a similar heat flux. The test results for 5000 ppm lithium metaborate is shown in Figure 5-11.
Figure 5-11 Boiling curves for 5000 ppm lithium Metaborate solution at 100 psi (6.9 bar).

Figure 5-12 Degradation of boiling heat transfer coefficients with 5000 ppm lithium metaborate solution (6.9 bar).
5.3 Tests with New Experimental Equipment

Boiling tests were performed at 100 psi with the new experimental set-up (DC electrical power supply) to compare the results of the preliminary experimental set up. The new test heater has the same dimensional and material properties as the old test heater. The test results for deionized water (tests through #721 to 725) are presented in Figure 5-13. The boiling test results from the new experimental set up match very well with results from the preliminary set-up. Figure 5-13 shows the results obtained from tests through 721 to 725 (new setup) and tests 651 through 655 (preliminary set up). A similar trend was obtained for tests done with 2000 ppm boric acid and 2000 ppm lithium metaborate where the percent difference is only 1.3%. These results verify that the new experimental set-up was operating properly.

Figure 5-13 Comparison results from old and new experimental set up (6.9 bar).

5.4 Tests at 200 psi pressure

Boiling tests performed at a pressure of 200 psi (13.3bar) showed similar results to those of tests done at lower pressure. The tests done are performed using the new experimental setup. Figure 5-14 shows results indicating boiling curves for varying degree of subcooling for deionized water. It can be clearly observed from the results that at lower heat flux, the boiling curves coincide very well. This can be explained due to lack of nucleation and this region indicates heat transfer takes place through natural convection. The boiling curves begin to rise steeply after a point indicating the onset of nucleate boiling. Because of nucleation, the rate of heat transfer increases rapidly. The boiling curves rise steadily with an increase of heat flux.
The boiling tests have revealed that the rate of heat transfer decreases with an increase of subcooling as can be seen in Figure 5-14. Also observe that the onset of nucleate boiling (ONB) is dependent on the degree of subcooling. For instance, the onset of nucleate boiling seems to occur at a heat flux of 45 kW/m² for saturated conditions whereas ONB occurs at the heat flux of 101 kW/m² for 30°C subcooled conditions. The maximum heat flux for all degrees of subcooling is well below the critical heat flux value of 1 MW/m² of deionized water (Incropera, 2001).

Boiling tests were performed with varying concentrations of boric acid in the coolant. Tests were performed with concentrations of 500 ppm, 1000 ppm, 2000 ppm and 5000 ppm, similar to tests at 100 psi.

Figure 5-15 shows the heat transfer coefficients for 500 ppm concentrated boric acid solution. Figure 5-20, 5-25, and 5-30 represent the results of the pool boiling tests for 1000 ppm, 2000 ppm and 5000 ppm concentrations of boric acid, respectively. Figure 5-17, Figure 5-22, Figure 5-28, and Figure 5-33 show the results of tests for 500 ppm, 1000 ppm, 2000 ppm and 5000 ppm concentration level of lithium metaborate, respectively.
Figure 5-15 Boiling Curve for 500 ppm boric acid at 200 psi (13.8 bar)

Figure 5-16 Boiling curves for deionized water and 500 ppm boric acid at 200 psi (13.8 bar)
Figure 5-17 Boiling curves for 500 ppm LiBO₂ Solution, 200 psi (13.8 bar)

Figure 5-18 Boiling curves for deionized water and 500 ppm LiBO₂ solution, 200 psi (13.8 bar)
Figure 5-19 Boiling curves for deionized water and 500 ppm LiBO₂ solution, 10°C subcooling and 200 psi (13.8 bar)

Figure 5-21 indicates the decrease in boiling heat transfer coefficient due to the presence of boron at 1000 ppm concentration, where for a heat flux of 218 kW/m², the heat transfer coefficient is 8.2 kW/m²°C compared to a value of 9.1 kW/m²°C for deionized water indicating almost a 9% drop. It can be observed from Figure 5-26 and Figure 5-27 that the presence of boric acid presence has a significant effect on the heat transfer rate. Additionally, the degradation of heat transfer coefficient continues with increasing concentration of boric acid from 500 ppm to 2000 ppm and is explained by the observed change in nucleation. For deionized water, bubbles were found to rise uniformly along the heater surface. For boric acid tests 825 and 826, the bubbles departed the heated surface in irregular clumps. With increasing heat flux, it was observed that the number of bubbles increased and departed the heated surface in a sporadic manner. This variation in bubble behavior patterns shows that the presence of boron reduces nucleation consequently reducing the heat transfer coefficient.

For a 2000 ppm concentration of boric acid, the heat transfer coefficient (heat flux = 221 W/m², saturated) decreased 19.9%. Figure 5-31 shows the variation of heat transfer coefficient in presence of boron at 5000 ppm concentration. Comparing Figure 5-31 and Figure 5-26, observe that the heat transfer coefficient does not change significantly. This indicates boric acid concentration changes from 2000 ppm to 5000 ppm decrease the heat transfer coefficient much less than occurs from 0 to 2000 ppm.

Observations from the boiling test results indicate that the boiling heat transfer rates for 200 psi (13.8 bar) follow a similar trend as the results obtained for 100 psi (6.9 bar). Observations from Figure 5-18 and Figure 5-19 indicates degradation of the boiling heat transfer coefficient for 500 ppm lithium metaborate relative to deionized water. A decrease of the heat transfer coefficient by 6.5% occurs at 10°C subcooling. Figure 5-23,
Figure 5-24, Figure 5-29, Figure 5-33, and Figure 5-34 show the influence of lithium metaborate solution compared to deionized water at 200 psi (13.8 bar). Tests with lithium metaborate at a pressure of 200 psi (13.8 bar) showed a similar trend as the tests conducted with a bulk liquid pressure of 100 psi (6.9 bar). The tests results indicate a reduction of the heat transfer coefficient by 23% (heat flux = 221 kW/m², saturated) for the 5000 ppm lithium metaborate solution. Comparing the heat transfer coefficients from Figure 5-30 and Figure 5-33, observe that lithium metaborate has greater influence than boric acid. In general, it is concluded from these boiling tests that the reduction in heat transfer coefficient is higher in presence of lithium metaborate solution than boric acid solution.

Figure 5-20 Boiling curves for 1000 ppm boric acid solution at 200 psi (16.9 bar).
Figure 5-21 Degradation of boiling heat transfer coefficients due to 1000 ppm boric acid concentration at 10°C bulk fluid subcooling, 200 psi (13.8 bar).

Figure 5-22 Boiling curves for deionized water and 1000 ppm LiBO₂ solution, 200 psi (13.8 bar).
Figure 5-23 Effect of lithium metaborate concentration on heat transfer coefficients at 30 °C subcooling and 200 psi pressure (13.8 bar).

Figure 5-24 Effect of lithium metaborate concentration on heat transfer coefficients at 20 °C subcooling and 200 psi (13.8 bar).
Figure 5-25 Boiling curves for tests with 2000 ppm concentration boric acid solution at 200 psi (13.8 bar).

Figure 5-26 Boiling curves for deionized water and 2000 ppm concentration boric acid solution at 200 psi (13.8 bar).
Figure 5-27 Effect of boric acid concentration on heat transfer at 20 C subcooling and 200 psi pressure (13.8 bar).

Figure 5-28 Boiling curves for 2000 ppm LiBO₂ solution, 200 psi (13.8 bar).
Figure 5-29 Effect of lithium metaborate concentration on heat transfer at 20°C subcooling and 200 psi pressure (13.8 bar).

Figure 5-30 Boiling curves for tests with 5000 ppm concentration boric acid solution at 200 psi (13.8 bar).
Figure 5-31 Effect of boric acid concentration (5000 ppm) on heat transfer at 20 C subcooling and 200 psi pressure (13.8 bar).

Figure 5-32 Effect of boric acid concentration (5000 ppm) on heat transfer at 10 C subcooling and 200 psi pressure (13.8 bar).
Figure 5-33 Boiling curves for 5000 ppm LiBO₂ solution, 200 psi (13.8 bar).

Figure 5-34 Effect of lithium metaborate concentration (5000 ppm) on heat transfer at 30 C subcooling and 200 psi pressure (13.8 bar).
5.5 **Boiling Curves at 500 psi**

Boiling tests were conducted at a higher pressure of 500 psi. The boiling curves for pure deionized water at 500 psi are presented below in Figure 5-36. For lower heat fluxes, the boiling curves match well indicating that natural convection heat transfer is dominate.

5.5.1 Tests with Boric Acid Solution

Figure 5-37 shows the results with 500 ppm boric acid. The variation in heat transfer coefficient is observed in Figure 5-38. As with the lower pressures previously discussed, observe that the 500 ppm boric acid solution at pressure of 500 psi has less influence on boiling the heat transfer coefficient compared to solutions with higher concentrations of boric acid.
Figure 5-36 Boiling curves for deionized water at 500 psi (34.5 bar).

The test results of 1000 ppm boric acid and 2000 ppm boric acid are shown in Figure 5-39 and Figure 5-41, respectively. Figure 5-42 indicates the decrease in heat transfer coefficient in the presence of boric acid at 2000 ppm concentration with respect to deionized water. From Figure 5-42, the degradation in boiling heat transfer coefficient is 21%, when compared with deionized water (tests numbered 930 and 904). Figure 5-40 shows the variation in boiling curves for the 1000 ppm boric acid solution test compared to deionized water at subcooling 10°C (231.4°C). Figure 5-44 shows boiling tests numbered 935 and 939 where the degree of subcooling ranges from 30°C to 0°C with 5000 ppm concentrated boric acid solution.

Figure 5-38, Figure 5-43 and Figure 5-45 represent the change in the boiling curves for 500 ppm, 2000 ppm and 5000 ppm concentration boric acid solutions, respectively, with reference to the boiling curves of deionized water at similar subcooling conditions. From Figure 5-45, observe that the superheat for the 5000 ppm boric acid solution is considerably greater than that of deionized water at a saturation temperature of 284.4°C (34.5 bar) for subcooling temperatures of 10°C, 20°C, and 30°C. This indicates
that the temperature gradient between the heater surface and bulk fluid is increased due to resistance offered by the boric acid solution. The nucleation pattern for 5000 ppm boric acid solution was not uniform and unlike that of deionized water which likely explains the reason for the decrease in heat transfer rates - poor coalescence of bubbles in the presence of boric acid leading to lesser nucleation.

Figure 5-37 Boiling curves for tests with 500 ppm concentration boric acid solution at 500 psi (34.5 bar).
Figure 5-38 Boiling curves for deionized water and 500 ppm concentration boric acid solution at 500 psi (34.5 bar).

Figure 5-39 Boiling curves for tests with 1000 ppm concentration boric acid solution at 500 psi (34.5 bar).
Figure 5-40 Effect of boric acid concentration on heat transfer at 10 C subcooling and 500 psi pressure (34.5 bar).

Figure 5-41 Boiling curves for tests with 2000 ppm concentration boric acid solution at 500 psi (34.5 bar).
Figure 5-42 degradation of boiling heat transfer coefficients due to 2000 ppm boric acid solution (test 930) at 500 psi (34.5 bar).

Figure 5-43 Boiling curves for deionized water and 2000 ppm concentration boric acid solution at 500 psi (34.5 bar).
5.5.2 Tests with Lithium Metaborate

The Figure 5-46 shows the results from experiments 915 through 918 for the 500 ppm lithium metaborate concentration. The boiling curves vary in subcooling ranging...
from 30 °C to saturation and indicate that the heat transfer coefficient is less affected for a concentration level of 500 ppm. Figure 5-48, Figure 5-50, and Figure 5-53, show the boiling test results with 1000 ppm, 2000 ppm, and 5000 ppm concentration of lithium metaborate, respectively. These figures indicate that superheat is maximum for the concentration level of 2000 ppm compared to the other concentrations. Figure 5-51 shows the variation in the boiling curve for saturated temperature compared to that of deionized water.

Figure 5-52 shows the degradation of the heat transfer coefficient for same test. Observe that the heat transfer coefficient is reduced by 24.4% (heat flux = 169 kW/m², saturated) in the presence of lithium metaborate at 2000 ppm concentration compared to deionized water. It is also interesting to note that at 5000 ppm concentration, the degradation of heat transfer is nearly the same as that of 2000 ppm concentration. Figure 5-54 shows the degradation in heat transfer coefficient for 5000 ppm concentrated lithium metaborate solution at saturated conditions compared to deionized water. For instance, the heat transfer coefficient is decreased by 23% for 5000 ppm concentration of lithium metaborate for test numbered 946 (heat flux = 169 kW/m²). This trend indicates that the increase of lithium in the coolant beyond a certain concentration does not have a significant effect on the heat transfer rate. This certain limiting lithium coolant concentration is 2000 ppm - beyond this concentration, the nucleation pattern is unchanged.
Figure 5-46 Boiling curves for tests with 500 ppm lithium metaborate solution at pressure of 500 psi (34.5 bar).

Figure 5-47 Effect of boric acid concentration on heat transfer at 20°C Subcooling condition and 500 psi pressure (34.5 bar).
Figure 5-48 Boiling curves for tests with 1000 ppm lithium metaborate solution at pressure of 500 psi (34.5 psi).

Figure 5-49 Effect of lithium metaborate concentration on heat transfer at saturated condition and 500 psi pressure (34.5 psi).
Figure 5-50 Boiling curves for tests with 2000 ppm lithium metaborate solution at pressure of 500 psi (34.5 bar).

Figure 5-51 Effect of lithium metaborate concentration on heat transfer at saturated condition and 500 psi pressure (34.5 bar).
Variation in boiling heat transfer coefficients

Figure 5-52 degradation of boiling heat transfer coefficients in presence of lithium metaborate (2000 ppm) at 500 psi (34.5 bar).

Figure 5-53 Boiling curves for tests with 5000 ppm lithium metaborate solution at pressure of 500 psi (34.5 bar).
Figure 5-54 degradation of boiling heat transfer coefficients in presence of lithium metaborate (5000 ppm) at 500 psi (34.5 bar).

5.6 Boiling Curves at 1000 psi

Boiling tests done at 1000 psi are very significant as it approaches the pressure typical of PWR conditions. Figure 5-55 shows the boiling curves for deionized water at 1000 psi for tests numbered 951 through 954. The boiling curves show a similar trend for all the subcooled conditions and saturated conditions as those of lower pressures. The boiling curves coincide well for lower heat fluxes indicating that heat transfer by natural convection at this higher is maintained. It can also be observed that the heat transfer coefficient decreases slightly when compared to tests done with deionized water at lower pressures. Marginal degradation of the heat transfer coefficients is explained by the nucleation pattern. It was observed that at higher test pressures, the nucleation was restricted. Due to higher pressures, the cavities which act as nucleation sites, are completely filled with liquid therefore causing resistance for the formation of new bubbles. It may be recalled that cavities, which are partially wetted, have more chance of being nucleation sites than cavities that are completely wetted [Collier and Thome 1994].
Repeatability is an important concern during boiling tests. The reproducibility of results depends on following the same defined procedure for conducting the boiling tests. Boiling tests were performed in regular intervals for the same conditions in order to check the repeatability of results. Repeated pool boiling tests were performed with deionized water at 10°C subcooling. The results of test numbered 953, 983, 985 and 986 have a boiling heat transfer coefficient of 9.05 kW/m²°C, 9.5 kW/m²°C, 8.75 kW/m²°C, and 9.1 kW/m²°C for a heat flux of 238 kW/m². The maximum boiling heat transfer coefficient was found to be 4.3% more than the average heat transfer coefficient, while the minimum was 3.8% less than the average boiling heat transfer coefficient. The deviation in results is due to the uncertainty in the measurements of temperature, voltage and current.

5.6.1 Tests with Boric Acid Solution

Figure 5-56 shows the boiling curves for tests numbered 958 through 961 for 500 ppm boric acid solution. It can be observed from Figure 5-57 that the influence of the boric acid solution at 500 ppm concentration is not significant (less than ??? percent). Figure 5-58 shows the boiling test results for all subcooled and saturated conditions with the 1000 ppm boric acid solution. Observe that the temperature difference between the wall and bulk fluid is higher than that of deionized water indicating that the nucleation affects the heat transfer coefficients. Figure 5-60 shows boiling curves obtained from tests numbered 975 through 978 for boric acid at a concentration of 2000 ppm. Figure 5-61 and Figure 5-62 show the change in boiling curves of the above mentioned tests with that of deionized water. Note that the addition of boric acid to water has a degrading
effect on the heat transfer coefficient. Figure 5-63 displays the results of tests done with 5000 ppm concentrated boric acid.

Figure 5-64 shows the degradation in heat transfer coefficient for test numbered 992. The boiling heat transfer coefficient decreases by 23% (for \( q = 236 \text{ kW/m}^2 \)) for the 5000 ppm concentrated boric acid solution at saturated temperature when compared to heat transfer coefficient for deionized water. The test results have thus indicated that the boiling heat transfer coefficient is significantly affected by the presence of boron at higher concentrations. The likely reason for the decrease in heat transfer coefficient is due to the observed decrease in nucleation sites. The boron particles in the solution seem to reduce the coalescence in the bubbles thereby reducing the heat transfer rate.

![500 ppm Boric Acid, 1000 psi (68.9 bar)](image)

Figure 5-56 Boiling curves for tests with 500 ppm boric acid solution at pressure of 1000 psi (68.9 bar).
500 ppm Boric Acid, 1000 psi (68.9 bar)

Figure 5-57 Effect of boric acid concentration on heat transfer at 20°C subcooling and 1000 psi pressure (68.9 bar).

1000 ppm Boric Acid, 1000 psi (68.9 bar)

Figure 5-58 Boiling curves for tests with 1000 ppm boric acid solution at pressure of 1000 psi (68.9 bar).
Figure 5-59 Effect of boric acid concentration on heat transfer at 30° C subcooling and 1000 psi pressure (68.9 bar).

Figure 5-60. Boiling curves for tests with 2000 ppm boric acid solution at pressure of 1000 psi (68.9 bar).
Deionized Water and 2000 ppm Boric Acid, 1000 psi (68.9 bar)

Figure 5-61 Boiling curves for deionized water and 2000 ppm concentration boric acid solution at 1000 psi (68.9 bar).

Deionized Water & 2000 ppm Boric Acid, 1000 psi (68.9 bar)

Figure 5-62 Effect of boric acid concentration (2000 ppm) on heat transfer coefficients at pressure of 1000 psi (68.9 bar).
Figure 5-63 Boiling curves for tests with 5000 ppm boric acid solution at pressure of 1000 psi (68.9 bar).

Figure 5-64 Degradation of boiling heat transfer coefficients for 5000 ppm concentrated boric acid solution at 1000 psi pressure (68.9 bar) and saturated condition.
5.6.2 Boiling Curves for Lithium Metaborate

Pool boiling test results for 500 ppm concentration lithium metaborate solution are presented in Figure 5-65. Figure 5-66 indicates the variation in superheat values for the 500 ppm lithium metaborate (LiBO$_2$) solution relative to deionized water. The heat transfer coefficient is noted to decrease by 4.5% (heat flux = 176 kW/m$^2$, saturated). Figure 5-67 shows the results of the boiling tests with 1000 ppm lithium metaborate solution (tests numbered 971 through 974). Here, the superheat values for the boiling curves are higher for tests with 1000 ppm lithium metaborate compared to that of the 500 ppm lithium metaborate solution. This denotes a decline in the boiling heat transfer rate for 1000 ppm lithium metaborate over the 500 lithium metaborate solution. For a heat flux value of 149 kW/m$^2$, the heat transfer coefficient decreases from 5.7 kW/m$^2$ C$^{-1}$ to 5.3 kW/m$^2$ C$^{-1}$ indicating a decrease of about 6% in the heat transfer coefficient for the 1000 ppm lithium metaborate when compared with deionized water at 10°C the subcooled condition.

![Boiling curves for tests with 500 ppm lithium metaborate solution at pressure of 1000 psi (68.9 bar).](image-url)
Figure 5-66 Effect of lithium metaborate concentration on heat transfer coefficients at 10 °C subcooling and 1000 psi pressure (68.9 bar).

Figure 5-67 Boiling curves for tests with 1000 ppm lithium metaborate solution at pressure of 1000 psi (68.9 bar).
Figure 5-68 Degradation of boiling heat transfer coefficients due to 1000 ppm lithium metaborate at 10°C subcooling (68.9 bar).

Figure 5-69 Boiling curves for tests with 2000 ppm lithium metaborate solution at pressure of 1000 psi (68.9 bar).
The boiling test results with 2000 ppm concentrated lithium metaborate solution are presented in Figure 5-69. Figure 5-70 shows the boiling test results for the 2000 ppm lithium metaborate solution. Figure 5-70 and Figure 5-71 show the decrease in boiling heat transfer coefficients for the 2000 ppm concentration of lithium metaborate solution in comparison to the boiling heat transfer coefficient of deionized water. The increase in the superheat values is a direct indication of a decline in the heat transfer coefficient for the same value of heat flux. Figure 5-72 shows the degradation of heat transfer coefficient with the 2000 ppm concentration of lithium metaborate solution. The boiling transfer coefficient for deionized water is 9.39 kW/m²°C (heat flux = 195.4 kW/m², saturated). The boiling heat transfer coefficient is reduced by a value of 2.7 kW/m²°C for the 2000 ppm lithium metaborate solution when compared to the boiling heat transfer coefficient of deionized water at (heat flux = 167 kW/m²). The reduction in heat transfer coefficient is 20 % (heat flux = 167 kW/m², saturated).

![Graph](image.png)

Figure 5-70 Boiling curves for deionized water and 2000 ppm concentration lithium metaborate solution at 1000 psi (68.9 bar).
Deionized water & 2000 ppm Lithium Metaborate, 1000 psi (68.9 bar)

Figure 5-71 Effect of lithium metaborate concentration (2000 ppm) on heat transfer coefficients at 1000 psi pressure (68.9 bar).

variation in boiling heat transfer coefficients, 1000 psi

Figure 5-72 degradation in heat transfer coefficients with influence of 2000 ppm lithium metaborate solution at saturated temperature and 1000 psi pressure (68.9 bar).
Figure 5-73 shows the boiling curves for tests numbered 993 through 997 for all the subcooled conditions and the saturated condition. The boiling curves for the 5000 ppm lithium metaborate solution are shown in Figure 5-73 and the deviation between the saturated boiling curve with the 5000 ppm concentration solution in comparison to the boiling test done with saturated deionized water are given in Figure 5-74. Figure 5-75 shows the degradation of the boiling heat transfer coefficient for the 5000 ppm lithium metaborate solution at the 10 °C subcooled condition. It can be observed from Figure 5-75 that the boiling heat transfer coefficient decreases to value of 10.4 kW/m²°C for a heat flux of 210 W/m². The corresponding heat transfer coefficient for deionized water is 13.9 kW/m²°C giving a reduction equal to 26 % when compared to the 5000 ppm lithium metaborate solution.

![Boiling curves for tests with 5000 ppm lithium metaborate solution at pressure of 1000 psi (68.9 bar).](image)

Figure 5-73 Boiling curves for tests with 5000 ppm lithium metaborate solution at pressure of 1000 psi (68.9 bar).
Figure 5-74 Effect of lithium metaborate concentration on heat transfer coefficients at saturation temperature and 1000 psi pressure (68.9 bar).

Figure 5-75 Degradation in heat transfer coefficients with influence of 5000 ppm lithium metaborate solution at saturated temperature and 1000 psi pressure (68.9 bar).
5.7  **Formation of Deposits**

Clearly concluded from the pool boiling test results is that the presence of boric acid and lithium metaborate in the coolant affects the heat transfer characteristics associated with subcooled nucleate boiling. As discussed earlier, the main reason for the occurrence of axial offset anomaly is subcooled nucleate boiling inside the core of a PWR and is observed in a PWR only after years of continuous operation. As it is not pragmatic to run the boiling tests for such a long period of time, boiling tests were run continuously with higher concentrations of boron and lithium in the coolant for five days to simulate the conditions favorable for deposition of particulates.

5.7.1  **Test Procedure**

The boiling test for deposits was conducted with 5000 ppm boric acid and lithium metaborate solutions at 1000 psi pressure and 10°C subcooling. The procedure employed for each test is described below:

1. Power was supplied to the bulk heater using a rheostat control. The coolant was heated and maintained at a steady state temperature of 274.2°C, which is 10°C below the saturation temperature at 1000 psi.

2. The power supplied to the test heater was increased until a heat flux of 250 kW/m² was reached and maintained thereafter by adjustment.

3. The coolant pressure was recorded at 20 minute intervals.

4. The current through the test heater was recorded using a clamp-type ammeter at 20 minute intervals.

5. The temperature of the test heater from the four K-type thermocouples and the bulk coolant temperature were recorded every 20 minutes.

6. The entire experimental set up was keenly monitored for any sudden changes in temperature, pressure and for mechanical and failures.

7. After the test was run for 5 days, the heat flux to the deionized water was slowly decreased to zero by decreasing the power input to test heater.

8. Similarly the power supplied to the bulk heater was decreased to zero in a steady process.

9. The test chamber was not disturbed in any manner so as to not disturb the test heater as it might have affected the particulate deposits.

10. After the coolant temperature reached the ambient temperature, the pressure in the test chamber was very gradually decreased to ambient pressure. Decreasing the pressure of coolant was done gradually to avoid sudden movement of coolant within test chamber which could have dislodged the particulate deposits.

11. After the coolant reached ambient conditions, it was discharged slowly through a bleed valve.
12. The test heater flange was unfastened from the test chamber for observation of the deposits. Care was to be taken while transporting the test heater as to not disturb the particulates deposited on the test heater.

13. The test heater surface and particulate deposits were then carefully observed using a microscope in conjunction with a camera to record the images.

14. After the visualization process, the deposits are then scraped into a tiny crucible for weighing. Care was taken to not damage the surface of the test heater. The net weight of deposits was calculated where the difference in weight of crucible before after collection yielded the net weight of particulate deposits. Because of the test heater surface irregularities, some particles settled in surface cavities.

15. The remaining deposits were then washed thoroughly with a measured quantity of acetone. The washed liquid was carefully collected in a small beaker. The advantage of using acetone for weight measurement was its unique volatile property. After the washed liquid was collected, it was kept open to the atmosphere. Due to its extreme volatile nature, acetone evaporated leaving behind the deposits. The beaker was then weighed. The difference in weights gave the weight of the deposits.

16. The above step was repeated until the net weight of deposits for a washing attempt was zero. This method was very effective as acetone has good cleaning characteristics.

17. The total weight of the deposits was obtained by summations of all the net weights obtained by direct measurement and by the acetone cleaning method.

5.7.2 Boric Acid Deposits

The variation of the heat transfer coefficient during the boiling test indicates that boron deposits have considerable effect on heat transfer characteristics. The boiling curve for the "boron deposition test" is shown in Figure 5-76. The change in heat transfer coefficient with time is plotted in Figure 5-77.

The 5000 ppm boric acid deposition test was conducted with a constant heat flux of 250 W/m². Observe from Figure 5-77 that the boiling heat transfer coefficient declines with increasing time. Initially, the heat transfer coefficient increased slightly until it reached a steady state. The heat transfer rate seemed to be constant for the first four hours of test and declined gradually thereafter over the remainder of the test. The decline in heat transfer rate is due to deposition of boric acid precipitates onto the test heater surface. For this deposition test, the boiling heat transfer coefficient decreases from 8.9 kW/m²°C to 7.0 kW/m²°C showing a reduction of 21% over a period of 136 hours. It can also be observed that after a running time of 80 hours, there is an increase in heat transfer coefficient. While this increase in boiling heat transfer coefficient is unexplained, it may be due particulate deposits being dislodged from the test heater surface.
Figure 5-76 Boiling curve for 5000 ppm boric acid solution at 1000 psi (68.9 bar).

Figure 5-77 Effect of boron deposits on heat transfer coefficient (68.9 bar).
The test heater was observed using a Canon high resolution camera with 400X magnification. This camera had an optical lens fitted to it and was mounted on a stand. As the surface of the test heater is cylindrical, only a portion of surface can be clearly viewed through the microscope at a time. Images were taken at the center of the boiling region, and at the beginning and end of the boiling region, to get information concerning the particulate deposit material density. Figure 5-78 shows a photomicrograph of test heater at the end of the boiling region on the test heater. It may be recalled that the actual length of heater filament is only one inch and away from the flange end. The boron deposits (white in color) can be clearly observed in Figure 5-78. The deposited particles seem to be scattered along the surface of the heater in the valleys on the surface. The test heater surface of the can also be observed at 400X magnification in Figure 5-78 where the imperfections of the surface are clearly visible. Figure 5-79 shows the deposited particle size distribution associated with Figure 5-78. The average size of the particle was found to be 2.62 microns (1micron =10^{-6} m) which is based on particle area. Figure 5-80 and Figure 5-82 show the photomicrographs of the test heater at the boiling region. It is evident that the deposits are higher in concentration which confirms the reason for degradation in heat transfer rate. Figure 5-81 and Figure 5-83 show the particle size distribution characteristics associated with Figure 5-80 and Figure 5-82, respectively. It can be observed that the maximum number of particles lie in size range of 5.96 sq microns. The particle size is presented by average area covered by the particle as it affects the deposits by covering the nucleation sites. It appears that most of the particulates are separate and scattered over heater surface. However some of the particulates seem to be form clusters. The average area of the particulate is 8.2 square microns at the boiling region.

![Figure 5-78 Photomicrograph of test heater adjacent to boiling region at 400X magnification for 5000 ppm boric acid deposition test.](image)
Boron deposits particle distribution at end of boiling region

Figure 5-79 Particle size distribution of the deposited particles at end of boiling region (total particle count = 260) for the 5000 ppm boric acid deposition test.

Figure 5-80 photomicrograph of boron deposits at boiling region on test heater for the 5000 ppm boric acid deposition test.
Figure 5-81 Particle size distribution for boric acid deposits for Figure 5-80, 5000 ppm boric acid deposition test (68.9 bar).

Figure 5-82 Photomicrograph of boron deposits on test heater at 400X magnification for the 5000 ppm boric acid deposition test (68.9 bar).
Figure 5-83 Particle size distribution for 5000 ppm boric acid deposits for Figure 5-82 (total particle count = 723).

The previously shown photomicrographs clearly show that boron particles settled on the surface of the test heater. Figure 5-80 shows the deposited boron particles on the surface of the test heater at the boiling region - the surface area over the 1.0 inch test heater filament. This indicates that boron deposition occurs due to subcooled nucleate boiling. It is also to be noted that the surface roughness plays a significant role in deposition of boron particles. Figure 5-82 indicates that density of deposited particulates is higher in the valleys of the surfaces. The possible reason for higher deposition in the valleys is due to higher nucleation at valleys in the surface. Previous studies have indicated that nucleation is higher in the cavities and troughs of the hot surface. The particle size of the deposits was calculated using Sigmascan. Sigmascan Pro (Systat Software) is an image analysis software tool. Using the threshold intensity feature of Sigmascan, a number of particles can be counted. This feature is based on the principle of difference in intensity of pixels in a image. The images obtained from the Canon high resolution camera setup have been calibrated and each pixel on the image corresponds to 2.98 square microns. Using this pixel conversion factor, the size of the particles and associated statistical data has been obtained.

Figure 5-84 shows a photomicrograph of the test heater surface outside the one-inch long boiling region. The numbers of deposited particulates are remarkably less than deposit rates at the boiling region. The lesser density of deposits strongly suggests that the deposition rates are directly dependent on nucleation rates. Figure 5-85 shows the particulate distribution associated with Figure 5-84. The number of particles with area in the 2.98 sq micron bracket is eighteen, whereas in Figure 5-84 there are 162 particles in the boiling region. Recapping for the 5000 ppm boric acid solution deposition test, a
particulate deposit weight of 12.2mg was removed from the heated test surface and that the heat transfer coefficient was degraded by 22% at 10°C subcooling.

Figure 5-84 Photomicrograph of test heater outside the boiling region (400X magnification) for the 5000 ppm boric acid deposition test (68.9 bar).

Figure 5-85 particle size distribution for boric acid deposits for Figure 5-84 (total particle count = 65) for 5000 ppm boron deposition test.
5.7.3 Lithium Metaborate

Boiling test for lithium metaborate deposition was conducted with a 5000 ppm concentration solution. The procedure employed was similar to that of the boric acid boiling particulate deposition test described in the previous section. The deposition test conducted with 5000 ppm lithium metaborate solution has resulted in failure of the test heater due to apparent high surface temperature achieved. Figure 5-86 Photograph of the failed test heater for the 5000 ppm lithium metaborate particulate deposition test showing the damaged Zr-4 cladding, and although not evident from the photograph, a large amount of particulate lithium deposits.

![Figure 5-86 Photograph of the failed test heater for the 5000 ppm lithium metaborate particulate deposition test (68.9 bar).](image)

The boiling test for deposition of the 5000 ppm concentration lithium metaborate solution was conducted over a time of 5 days similar to that of the boric acid deposition test. Figure 5-87 shows the degradation of the boiling heat transfer coefficient with time where heat transfer coefficient is not affected significantly for the first 300 minutes of time but thereafter decreases at a uniform rate from 7.8 kW/m²°C to 5.5 kW/m²°C (nearly 30% reduction) over a period of 140 hours. The degradation in heat transfer coefficient is due to the apparent constant deposition rate of lithium metaborate.

The test heater was observed using a Zeiss LSM5 Pascal microscope located in the Department of Biology at Kansas State University. The LSM 5 is a laser scanning confocal microscope capable of exciting fluorescent markers with laser lines of 458/488/514/543/633 nm. Figure 5-88 shows a photomicrograph of the test heater after the boiling test with the 5000 ppm concentrated lithium metaborate solution.
Figure 5-87 Degradation of heat transfer coefficient with boiling test time for 5000 ppm concentration Lithium Metaborate solution

Figure 5-88 Photomicrograph of test heater with deposits of precipitate of 5000 ppm lithium metaborate at 600X magnification (68.9 bar).

Figure 5-88 shows the lithium metaborate deposits on the test heater surface where the concentration of deposits is high at the center of the boiling region surface. Figure 5-89 shows a part of the image captured by the microscope of the boiling region at a magnification of 600X. The microscope scans the specimen line by line to complete an entire picture of the specimen. Figure 5-89 shows the image of test heater surface captured in one scan. The deposits appear to be clusters of tiny particulates. Unlike boron deposits, the lithium metabaorate deposits are smaller in size but higher in density.
Lithium metaborate particulates appear to deposit in random clumps as seen in Figure 5-91.

Figure 5-90 shows the particulate size distribution for lithium metaborate deposits. The size of the particulates was calculated by analyzing the photomicrographs in a similar method to that of the boric acid photomicrographs. The results reveal that most of the particulates lie in the size range of 0-3 sq microns. It can also be observed that the size of particles is relatively smaller than that of the boric acid deposits. The average size of particulate material was found to be 5.8 sq microns. However, particulates with area greater than 15 microns are higher when compared to boric acid deposits indicating that some of the particulates combine to form a clusters. Figure 5-91 shows another image taken from the Zeiss LSM5 Pascal microscope at the boiling region with a lesser magnification of 100X where the deposits of lithium metaborate precipitate are observed to have settled on the surface of test heater. This image shows again that the deposits are formed in clusters of small individual particles. It may be recalled that the bubble formation occurred in odd clumps for the 5000 ppm lithium metaborate solution. The reason for these odd clumps in nucleation is explained by the distribution pattern for the lithium metaborate deposits.

Figure 5-89 Photomicrograph of test heater with 5000 ppm concentration solution lithium metaborate deposits at magnification of 600X (68.9 bar).
Figure 5-90 Particulate size distribution of lithium metaborate deposits on test heater for total particle count of 1326.

Figure 5-92 shows a photomicrograph of the test heater taken away from the boiling surface region of the test heater. The density of deposits is less compared to the density of deposits at boiling region. However, there is minimal nucleation occurring at the transition region between the boiling and non boiling region due to heat transfer along the test heater by conduction. The reduction in heat transfer in the presence of lithium metaborate can be explained by the observed decrease in nucleation rates apparently due to the particulate deposition.

5.7.4 Nucleation

A high speed camera was used to capture the nucleation at the test heater surface. The videos were taken using 16mm Kodak Hawk Eye Surveillance film processed at the Yale film Studio in California (Yale Film and Video 2006). The boiling images were captured for tests with deionized water and different concentrations of boric acid and lithium metaborate. The processed films were later viewed using a Howell 16mm film projector Model 2580 (Howell 1985. Observations from viewing the nucleation films revealed the variation in bubble behavior. It was found that nucleation was smooth for deionized water with a significant number of bubbles. Figure 5-94 shows the bubble formation pattern for deionized water with a relatively high heat flux of 240 kW/m². Figure 5-93 shows the image captured on the film taken for a boiling test with deionized water at 10° C subcooling, 1000 psi bulk fluid pressure, and 150 kW/m² heat flux. It has been observed with increasing heat flux that the bubble density increases along with the combining rate of the bubbles upon leaving the heated surface. In other words, the bubbles seem to be collapsing in lesser time when compared with the bubble collapsing time for 100 psi. Figure 5-93 and Figure 5-94 were captured at a speed of 500 frames per
second. Figure 5-94 shows columns of bubbles indicating that the nucleation rate is very high and coalescence of the bubbles leads to the formation of columns of bubbles which appear to be leaving the heater surface uniformly in all directions.

Figure 5-91 Photomicrograph of test heater after the 5000 ppm concentration lithium metaborate boiling test at magnification of 100X (68.9 bar).

Figure 5-92 Photomicrograph of test heater after the 5000 ppm lithium metaborate boiling deposition test at edge of boiling region at magnification of 600X (68.9 bar).
Figure 5-93 Visual image of nucleation for deionized water at 1000 psi (68.9 bar).

Figure 5-94 Image of bubbles above test heater surface at 240 kW/m² heat flux for deionized water at 20X magnification (pressure = ?)
Images were captured at a higher speed (frame rate) using the HYCAM high speed camera. Figure 5-95 shows an image taken at 6000 half frames/sec speed for a boiling test with deionized water where the heat flux is 240 kW/m². Figure 5-95 has the same test conditions as Figure 5-94, but much higher frame rate or speed. It is observed that this higher flux, the nucleation rate is so high that at the slower frame rate, the bubbles seem to leave the surface in clusters, but in fact, they are individual bubbles leaving heater surface with an average bubble size measured to be $\frac{1}{16}$th inch in diameter. Also, observation from Figure 5-95 reveals that the bubbles are mostly spherical in shape and seem to be nucleating uniformly.

Similar to high-speed images taken with deionized water, nucleation high-speed film images were captured for the boiling test with 5000 ppm concentration boric acid solution at 1000 psi (68.9 bar). Observations of the high-speed film reveals that bubble density is less than with deionized water. The delay time between the collapsing of an old bubble and the growth of a new bubble was found to be 0.042 seconds whereas the value for deionized water was found to be 0.036 seconds. The delay time for 5000 ppm lithium metaborate was 0.045 seconds. Figure 5-96 shows the nucleation image captured at 3000 fps for 5000 ppm concentration boric acid. Note that the number of bubbles is relatively few compared to that of deionized water at the same conditions indicating the influence of boric acid deposition on the surface of the test heater.

![Figure 5-95 bubbles captured at speed of 3000 fps, 10°C subcooled and 1000 psi for at 20X magnification](image)

Figure 5-97 shows the nucleation image captured at 3000 fps for the boiling test with 5000 ppm concentration lithium metaborate solution. It is clearly evident from the figure that the number of bubbles is less than those of deionized water showing a reduced bubble density. The bubble density for lithium metaborate can be extrapolated based on images taken at the higher frame rate speed. The bubble density for lithium metaborate solution was found to be approximately 1406 bubbles per sq cm (viewing area) based on
the data collected from the nucleation images. The values can be considered for comparison. Similar approximations have revealed the bubble density was 1968 per sq cm for 5000 ppm boric acid and 1406 per sq cm for deionized water. These approximations are based on a heat flux value of 200 kW/m². The analysis of images obtained from high speed camera indicate that nucleation is more uniform for deionized water and therefore leads to a better boiling heat transfer rate. In comparison, the bubble behavior of boric acid seems to indicate bubbles of a smaller size and less uniform indicating that boric acid deposits affect the nucleation by reducing the nucleation sites which thereby lead to a reduction in heat transfer coefficient. Results for lithium metaborate revealed a much smaller number of bubbles, but the relative size of the bubbles was bigger compared to that of deionized water. Furthermore, for the lithium metaborate solution, the generation of bubbles was random in nature and seemed to be occurring in clumps. This could be possibly be explained by behavior of lithium metaborate deposits to form clusters thereby completely covering the nucleation sites. The deposition images of lithium metaborate revealed that the deposition occurred in patches on the heater surface.

Figure 5-97 shows bubbles leaving the surface of test heater for the boiling test with 5000 ppm lithium metaborate acid solution. Figure 5-97 was taken at 6000 half frames per minute speed which implies that the picture was captured in a time frame of little less than 1/100th of a second. It can be observed in Figure 5-97 that there are a fewer number of bubbles indicating that the nucleation rate has significantly degraded in the presence of lithium metaborate.

![Figure 5-96 Picture of nucleation for 5000 ppm boric acid solution taken at 3000 fps and at 20X magnification.](image-url)
Figure 5-97 Picture of nucleation for 5000 ppm lithium metaborate solution taken at 3000 fps and at 20X magnification.
6 ANALYSIS OF RESULTS

The factors affecting the heat transfer coefficient under the experimental nucleate pool boiling conditions are discussed in this chapter. The boiling test results obtained are evaluated and the effect of various parameters affecting the boiling heat transfer coefficient are discussed.

6.1 Effect of Coolant Concentration

Evaluation of the boiling test results have revealed that the presence of boric acid and lithium metaborate in deionized water affect the boiling heat transfer coefficient by causing a resistance to the heat transfer between the heater and the bulk liquid. The details are discussed below.

6.1.1 Effect of Boric Acid

The evaluation of boiling tests performed at bulk fluid pressures of 100 psi, 200 psi, 500 psi and 1000 psi reveal that the boiling heat transfer coefficient decreases with increasing boric acid concentration in the bulk fluid (coolant). Increased resistance to uniform nucleation caused by the boric acid solution likely explains the decrease in heat transfer rate. The nucleation that occurs during the boiling process leads to the formation of boric acid deposits on the heater surface thereby creating a layer of resistance between the heater and bulk fluid. The photomicrographs taken after the boiling test with 5000 ppm concentration boric acid confirm the presence of boron particulates on the test heater. The effect of varying the boric acid concentration in deionized water on the heat transfer coefficient is shown in Figure 6-1 where the bulk fluid pressure is 1000 psi in conjunction with a saturation temperature of 284.4°C. Even though there is a decrease in pool boiling heat transfer coefficient for 500 ppm concentration boric acid over that of deionized water, the decrease is not significant. The heat transfer coefficient was reduced by 4.9% at a heat flux of 236 kW/m² when compared to the pool boiling heat transfer coefficient of 14.7 kW/m²°C for deionized water. However, for 1000 ppm concentration of boric acid, the decrease in heat transfer coefficient is considerable. For a heat flux of 236 kW/m², the heat transfer coefficient decreased by 9% to a value of 13.4 kW/m².

The pool boiling heat transfer coefficients for 2000 ppm and 5000 ppm concentration solutions of boric acid at a heat flux of 236 kW/m² are 11.9 kW/m² and 11.4 kW/m², respectively. It can be observed that the pool boiling heat transfer coefficient is lowest for coolant with a 5000 ppm concentration of boric acid. Figure 6-2 shows the degradation of the pool boiling heat transfer coefficient with increasing boric acid concentration. Also note from Figure 6-2 that the boiling heat transfer coefficient decreases steadily up to a concentration of about 2000 ppm. The rate of decrease in the heat transfer coefficient reduces with concentrations of boric acid greater than 2000 ppm indicating that further addition of boric acid to the solution does not significantly affect the heat transfer rate.
Figure 6-1 Degradation of pool boiling heat transfer coefficients with variation of coolant concentration (boric acid as additive).

Figure 6-2 Degradation of pool boiling heat transfer coefficients of boric acid solution relative to deionized water.
6.1.2 Effect of Lithium Metaborate

The evaluation of the boiling curves obtained from tests with different concentrations of lithium metaborate indicates a similar trend as with boric acid. The pool boiling heat transfer coefficients decrease with increasing concentration of lithium metaborate in deionized water. However, the reduction rate of the pool boiling heat transfer coefficient for lithium metaborate is slightly different from that of boric acid. Changing the bulk fluid pressure does not have a significant effect on heat transfer rate and is shown in Figure 6-3 under different concentrations of lithium metaborate solution. Observed from Figure 6-3 that the pool boiling heat transfer coefficient decreases with increasing concentration of the lithium metaborate solution. It is also interesting to note that the change in heat transfer coefficient is not significant for those boiling tests with 2000 ppm and 5000 ppm concentrations. This indicates that upon further addition of lithium to deionized water, the variation in boiling heat transfer coefficient will not be significant.

Figure 6-4 shows the percentage change in the pool boiling heat transfer coefficient for varying concentration of lithium metaborate for the boiling tests done at a coolant pressure of 1000 psi and at saturation temperature of 284.4 °C. The boiling heat transfer coefficient for deionized water at a heat flux of 236 kW/m² is 14.6 kW/m²°C at the saturation temperature of the bulk fluid. The pool boiling heat transfer coefficient shows a reduction of 6.6%, 12%, 20.2% and 24.9% for 500 ppm, 1000 ppm, 2000 ppm and 5000 ppm concentration of lithium metaborate solution, respectively where the bulk fluid is saturated temperature. In comparison with boric acid, it was found that lithium metaborate has a greater influence on the pool boiling heat transfer rate than boric acid as seen from Figure 6-4 and Figure 6-2. This is consistent with the observed nucleation and deposition patterns for associated with the boric acid and lithium metaborate solutions. Recall that the lithium metaborate deposits occurred in clusters and had a higher deposition rate than the boric acid deposits which explains the higher degradation of pool boiling heat transfer. Figure 6-5 shows a comparison of the effects of boron and lithium on the pool boiling heat transfer rate where it can be observed that the addition of lithium metaborate has resulted in a higher degradation of heat transfer rate relative to the addition of boric acid.

6.2 Effect of Subcooling

With increasing subcooling of bulk fluid, the pool boiling heat transfer coefficient decrease as seen in Figure 5-1 and Figure 5-14. Additionally, Figure 5-36 and Figure 5-55 show that the boiling heat transfer coefficient decreases with increasing level of subcooling from 10 °C to 30 °C. The boiling heat transfer coefficient was higher for tests done at saturated temperature because the onset of nucleate boiling (ONB) occurs at a lower heat flux for those boiling tests conducted at saturation pressure. The evaluation of boiling the tests reveal that the pool boiling heat transfer rate are highest when the temperature gradient between hot surface and coolant is a minimum. Figure 6-7 shows the degradation in the pool boiling heat transfer coefficient with increasing the bulk fluid...
subcooling. The ordinate in the Figure 6-7 indicates the heat transfer ratio \( \frac{\alpha}{\alpha_{\text{saturation}}} \), which is the ratio of

![Variation in heat transfer coefficients, 1000 psi](image)

**Figure 6-3** Degradation of pool boiling heat transfer coefficients with variation of coolant concentration (lithium metaborate as additive).

![Effect of lithium metaborate concentration on pool boiling heat transfer coefficients](image)

**Figure 6-4** Degradation of pool boiling heat transfer coefficients of lithium metaborate solution relative to deionized water.
Effect of coolant concentration on pool boiling heat transfer coefficients

![Graph showing degradation of pool boiling heat transfer coefficients with concentration of coolant.](image)

Figure 6-5 Comparison of the effect of boron and lithium concentration on pool boiling heat transfer rates.

Effect of bulk subcooling on pool boiling heat transfer (1000 psi)

![Graph showing effect of bulk subcooling on pool boiling heat transfer.](image)

Figure 6-6 Effect of bulk subcooling on pool boiling heat transfer.
the boiling heat transfer coefficient at any subcooling to the boiling heat transfer coefficient at saturation temperature. Observe that with increasing degree of subcooling, the heat transfer coefficient decreases.

6.3 **Effect of Pressure**

The increase of pressure does not have a significant effect on the heat transfer coefficient. However, note from Figure 6-7 that increasing bulk fluid pressure, while maintaining a fixed heat flux, reduces the nucleation rate consequently decreasing the pool boiling heat transfer rate. This is due to the greater external fluid pressure on the bubbles rising from the heater surface. In other words, the heat flux required for nucleation to occur increases with increasing pressure. The reduction in heat transfer coefficient can be also be explained by the possibility of active nucleation cavities being completely filled with coolant thereby limiting bubble formation in the cavity. The likelihood of a new bubble nucleating is high in semi–wet cavities and low in fully–wet cavities. Tests with varying concentrations of boric acid and lithium metaborate solutions show slight degradation of the heat transfer coefficient with increasing pressure. For a change in system pressure from 200 psi to 1000 psi, the boiling heat transfer coefficient decreases slightly from 4.7 kW/m²°C to 4.3 kW/m²°C for a 5000 ppm concentration boric acid solution. For boiling tests with a 5000 ppm concentration of lithium metaborate, the heat transfer coefficient showed a slight degradation from 4.6 kW/m²°C to 4.1 kW/m²°C for a change of coolant fluid pressure from 200 psi to 1000 psi, respectively.

Figure 6-7 Effect of pressure on pool boiling heat transfer coefficient.
6.4 Analysis of Particulate Deposits

The photomicrographs of the boric acid deposits reveal that the particulate deposits were scattered over the entire surface above the heating element (nucleation region of the test heater). The total weight of the boric acid deposits was found to be 12mg for a boiling test with 5000 ppm boric acid solution at 1000 psi pressure and 10°C bulk subcooling. The analysis of this boric acid deposition test indicates that significant deposits are formed on the test heater which leads to a decrease in heat transfer coefficient over time. The average number of deposition particles was found to be 742 per sq mm for the 5000 ppm boric acid solution test. The increase in pool boiling heat transfer coefficient during the first few hours of the test (Figure 5-77) result from the deposits serving as heat transfer “fins” augmenting the heat transfer. Afterwards, the heat transfer coefficient decreases uniformly with the exception of 80 hours where a fraction of the boron deposits have been apparently dislodged from the heater surface. The deposition rate however may have reached a saturation or steady state condition after 100 hours as the change in the pool boiling heat transfer coefficient appears insignificant after a time of 110 hours.

Deposits of lithium metaborate reveal that deposits of this material are more dense in particle number than that of lithium metaborate where the deposits seem to form clusters of cohesive particles. The total weight of the deposits from the lithium metaborate test was found to be 22mg for 5000 ppm lithium metaborate concentration at 1000 psi and 10°C subcooling. As seen in Figure 5-91 the lithium metaborate deposits occur in patches which leads to irregular nucleation patterns. The average number of deposition particles was found to be 1021 particles per sq. mm. From the test results, it appears that the lithium metaborate enhances the deposition rate of boron onto the Zr-clad test heater. The degradation of heat transfer coefficient in the presence of lithium metaborate is significantly higher than that of boric acid. It can be concluded that the higher deposition rate in the presence of lithium metaborate is cause for greater reduction of heat transfer rates.

6.5 Analysis of Bubble Growth Behavior

Observations from the visual recordings reveal that nucleation occurs in a uniform manner for deionized water. The bubble size obtained from nucleation images varied from 0.8mm to 2.5mm in diameter for a heat flux of 150 kW/m². The visual recordings of the nucleation obtained for boric acid solution reveal that nucleation is less uniform. The bubble size for boiling tests with 5000 ppm boric acid solution obtained from the nucleation images varied from 0.6mm to 2mm in diameter for a heat flux of 150 kW/m². The analysis of nucleation recordings for tests with lithium metaborate revealed that the nucleation occurred in a non-uniform and sporadic manner. The bubbles occurred in odd clumps indicating a strong presence of lithium metaborate deposits on the test heater surface. The bubble size varied from 0.6mm to 3mm for a heat flux of 150 kW/m². The larger size of bubbles in the case of lithium metaborate is due to the bubbles combining with each other. However, as explained in nucleation results section of this report, the nucleation density is lower with both the boric acid and lithium metaborate coolant solutions than deionized water accounting for degradation of boiling heat transfer rates. It is the variation in nucleation patterns that explains the change in heat transfer rate for
the boric acid and lithium metaborate solutions. As explained in nucleation results, the coolant concentration affects the nucleation process by decreasing the nucleation densities. For 5000 ppm boric acid, the nucleation densities decreased by 20% when compared with deionized water and in presence of 5000 ppm lithium metaborate solution, the number of bubbles decreased by 38%.
7 CONCLUSIONS

This experimental investigation generated a data base for subcooled pool boiling heat transfer coefficients on Zr-4 clad heater rods for coolants with varying concentration of boron and lithium. The generated empirical data includes the effect of bulk subcooling, bulk fluid pressure, heat flux and coolant concentration and is useful for better understanding of axial offset anomaly (AOA) phenomenon in PWR power nuclear reactors.

Analysis of the results obtained from the investigation reveal that the increase of boric acid and lithium metaborate solution concentration results in the degradation of the boiling heat transfer coefficient. The boiling heat transfer coefficient was found to be slightly lower in the presence of lithium than with boron. The degradation in the heat transfer coefficient, relative to that of deionized water, was about 24% for boron and about 27% for lithium at a coolant concentration of 5000 ppm, respectively. The pool boiling tests showed a consistent decline in boiling heat transfer rate with increasing degree of subcooling. The experimental investigation revealed that the system pressure did not have a significant effect on the pool boiling heat transfer coefficient.

The inspection of the electrical test heater clad with Zr-4, revealed the formation of particulate deposits on the surface after undergoing subcooled nucleate boiling for five days with 5000 ppm concentrations of boric acid and lithium metaborate. The formation of deposits on the heater surface confirms that the precipitation of lithiated compounds like lithium metaborate on the test heater occurs during subcooled nucleate boiling. A visual recording of the nucleation was obtained at subcooled boiling conditions with varying concentrations of boron and lithium. The image analysis of the deposits reveal that the boron deposits are scattered and less dense than the deposits of lithium. The greater deposition quantity of lithium also explains the reason for its greater effect on the boiling heat transfer coefficient. The experimental data generated in this investigation contributes to the understanding of AOA.revealing that the deposition of boron and lithium precipitates is a factor in determining the heat transfer rate from the surface of a Zr-4 clad fuel element. The influence that the particulate boron and lithium metaborate precipitates have after a 5 day test at 5000 ppm concentration and 1000 psi operating pressure are 21% and 30%, respectively.
REFERENCES


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APPENDIX A

A.1 Program for Data Reduction in FORTRAN

PROGRAM PBDATA
PARAMETER (pi=3.141593)
PARAMETER (twopi=2*pi)
COMMON/XDATA/idnum,idate,isurf,icomp(5),frac(5),npts,iunits,
& volts(40),amps(40),P(40),Tb(2,40),Tc(4,40),Te(4,40)
COMMON/XOUTP/delt(40),Qloss(40),alpha(40),Twall(40),flux(40),
& TCvar1(40),TCvar2(40)
CHARACTER*20 inpfil, datfil, sdffil
DATA jin/1/,jinp/2/,jdat/3/,jsdf/4/,jban/5/
OPEN(jin,FILE='test.inp',STATUS='OLD')
OPEN(jban,FILE='pbdata.txt',STATUS='UNKNOWN')
C--- Read number of data files to be processed
READ(jin,10)
10     FORMAT( )
     READ(jin,10)
     READ(jin,15) nfiles
15     FORMAT(3X,I3)
     READ(jin,10)
C*************** Begin BIG loop to process each data file
    DO 900 IFILE=1,nfiles
20       FORMAT(3A20)
WRITE(*,25) inpfil
25       FORMAT(/,10X,'Begin data reduction for file ',A20)
C--- open output files
OPEN(jinp,FILE=inpfil,STATUS='OLD')
OPEN(jdat,FILE=datfil,STATUS='UNKNOWN')
OPEN(jsdf,FILE=sdffil,STATUS='UNKNOWN')
C--- process the data -- many possible cases
     IF (idnum.GE.1000) THEN
20       FORMAT(3A20)
         WRITE(*,25) inpfil
25       FORMAT(/,10X,'Begin data reduction for file ',A20)
C--- open output files
         OPEN(jinp,FILE=inpfil,STATUS='OLD')
         OPEN(jdat,FILE=datfil,STATUS='UNKNOWN')
         OPEN(jsdf,FILE=sdffil,STATUS='UNKNOWN')
     END IF
     IF (idnum.GE.1000) THEN
     C********** FLAT DISK TEST SECTIONS
     condcu = 391.0
     ri = 0.0127
     ro = 0.0159
     AI = 0.507E-03
     AO = 0.792E-03
     thick = 0.8E-03
     perim = 2. * twopi * ri + 2. * thick
     AX = twopi * ri * thick
     zlb = ro - ri
     zlnc = 5.
     zl = zlnc * perim / condcu / AX
     D1 = 0.001
     D2 = 0.005
C-----------------
D3 = 0.009
DO 1050 I = 1, npts
    STC = Tc(1,I) + Tc(2,I) + Tc(3,I)
    SX2 = D1*D1 + D2*D2 + D3*D3
    SX  = D1 + D2 + D3
    STCX= Tc(1,I)*D1 + Tc(2,I)*D2 + Tc(3,I)*D3
    S2X = SX * SX
    Twall(I) = (STC*SX2 - SX*STCX) / (3.*SX2 - S2X)
    TBave = (Tb(1,I) + Tb(2,I))/2.
    delt(I) = Twall(I) - TBave
    watts = volts(I) * amps(I)
    Qloss(I) = SQrt(zlnc*perim*condcu*AX)*delt(I)*TANH(zl)
    flux(I) = (watts - Qloss(I)) / AI
    alpha(I)= flux(I)/delt(I)
    Qloss(I)= 100.*Qloss(I)/watts
    TCvar1(I) = 0.
    TCvar2(I) = 0.
1050        CONTINUE
ELSEIF ((isurf.EQ.1).OR.(isurf.EQ.7).OR.(isurf.EQ.8)) THEN
C**********************************************************
C           isurf = 1,7,8
C           Smooth Tube - See section B.2, pg. 101 Shakir
C**********************************************************
ri=  0.00630
rt=  0.00870
ro=  0.01110
C--         This heater had a 3 inch boiling length
zl=  0.07620
znc=0.01270
zl = zl+ znc
condcu= 391.0
C
DO 135 I=1,npts
    watts = volts(I)*amps(I)
    Q = watts
    TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I))/4.
    TBave=(Tb(1,I)+Tb(2,I))/2.
    TLave=(Te(1,I)+Te(2,I))/2.
    Tvar = 0.
    DO 133 J=1,4
        Tvar = Tvar + (TCave - Tc(J,I))**2
    133           CONTINUE
    TCvar1(I)= SQrt(Tvar)
    TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
    Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
    delt(I)= Twall(I) - TBave
    flux(I)= Q/(twopi*ro*zl)
    alpha(I)= flux(I)/delt(I)
    deltl= TCave - TLave
    deltM= TLave - TBave
    Qloss(I)= condcu*pi*(ro*ro-ri*ri)*deltL/(0.5*zl) +
    & alpha(I)*twopi*ro*zlnc*deltM
    Q = Q - Qloss(I)
    Qloss(I)= 100.*Qloss(I)/watts
    Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)

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delt(I) = Twall(I) - TBave
flux(I) = Q/(twopi*ro*zlb)
alpha(I) = flux(I)/delt(I)

ELSEIF (isurf.EQ.2) THEN
  C**************************************************
  C           isurf = 2, High Flux  (Bajorek)
  C**************************************************
  ro= 0.009335
  RC= 0.008382
  rt= 0.007798
  ri= 0.004763
  zlb= 0.0508
  zlnc= 0.041625
  zl= zlb + zlnc
  CONDHF= 242.0
  C
  DO 145 I=1,npts
      watts = volts(I)*amps(I)
      Q = watts
      TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I)+2.*(Te(2,I)+
      &    Te(4,I)))/8.
      TBave=(Tb(1,I)+Tb(2,I))/2.
      TLave=(Te(1,I)+Te(3,I))/2.
      Tvar = 0.
      DO 143 J=1,4
          Tvar = Tvar + (TCave - Tc(J,I))**2
      143           CONTINUE
      TCvar1(I)= SQrt(Tvar)
      TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
      Twall(I)= TCave - Q*(1./(twopi*CONDHF*zlb))*ALOG(ro/rt)
      delt(I)= Twall(I) - TBave
      flux(I)= Q/(twopi*ro*zlb)
      alpha(I) = flux(I)/delt(I)
      deltl= (Te(2,I)+Te(4,I))/2. - TLave
      deltM= TLave - TBave
      Qloss(I)= CONDHF*pi*(ro*ro-ri*ri)*deltl/(0.5*zlnc) +
      &      alpha(I)*twopi*ro*zlnc*deltM
      Q = Q - Qloss(I)
      Qloss(I)= 100.*Qloss(I)/watts
      Twall(I)= TCave - Q*(1./(twopi*CONDHF*zlb))*ALOG(ro/rt)
      delt(I)= Twall(I) - TBave
      flux(I)= Q/(twopi*ro*zlb)
      alpha(I) = flux(I)/delt(I)
  145        CONTINUE
  ELSEIF (isurf.EQ.3) THEN
  C************************************************************
  C        isurf = 3, Finned Tube, 19 Fins per Inch (Bajorek)
  C************************************************************
  ri   = 0.00470
  rt   = 0.00591
  ro   = 0.00800
  RFIN = 0.009535
  zlb  = 0.0508
  zlnc = 0.03175

  C
  DO 145 I=1,npts
      watts = volts(I)*amps(I)
      Q = watts
      TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I)+2.*(Te(2,I)+
      &    Te(4,I)))/8.
      TBave=(Tb(1,I)+Tb(2,I))/2.
      TLave=(Te(1,I)+Te(3,I))/2.
      Tvar = 0.
      DO 143 J=1,4
          Tvar = Tvar + (TCave - Tc(J,I))**2
      143           CONTINUE
      TCvar1(I)= SQrt(Tvar)
      TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
      Twall(I)= TCave - Q*(1./(twopi*CONDHF*zlb))*ALOG(ro/rt)
      delt(I)= Twall(I) - TBave
      flux(I)= Q/(twopi*ro*zlb)
      alpha(I) = flux(I)/delt(I)
      deltl= (Te(2,I)+Te(4,I))/2. - TLave
      deltM= TLave - TBave
      Qloss(I)= CONDHF*pi*(ro*ro-ri*ri)*deltl/(0.5*zlnc) +
      &      alpha(I)*twopi*ro*zlnc*deltM
      Q = Q - Qloss(I)
      Qloss(I)= 100.*Qloss(I)/watts
      Twall(I)= TCave - Q*(1./(twopi*CONDHF*zlb))*ALOG(ro/rt)
      delt(I)= Twall(I) - TBave
      flux(I)= Q/(twopi*ro*zlb)
      alpha(I) = flux(I)/delt(I)
  145        CONTINUE
zle = zlb + zlnc
condcu = 391.0
DO 155 I=1,npts
   watts = volts(I)*amps(I)
   Q = watts
   TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I))/4.
   TBave=(Tb(1,I)+Tb(2,I))/2.
   Tvar = 0.
   DO 153 J=1,4
      Tvar = Tvar + (TCave - Tc(J,I))**2
   153 CONTINUE
   TCvar1(I)= SQrt(Tvar)
   TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
   Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
   delt(I)= Twall(I) - TBave
   flux(I)= Q/(twopi*RFIN*zlb)
   alpha(I)= flux(I)/delt(I)
   deltM= 0.5 * delt(I)
   Qloss(I)= alpha(I)*twopi*ro*zlnc*deltM
   Q = Q - Qloss(I)
   Tvar = Tvar + (TCave - Tc(J,I))**2
   155 CONTINUE
ELSEIF (isurf.EQ.4) THEN
C****************************************************************
C          isurf = 4    (Schnelle)
C          Smooth surface, 0.75 inch diameter, 2.5 inch boiling length
C          Four thermocouples at midpoint of heated length.
C****************************************************************
   ri= 0.47625E-02
   rt= 0.71438E-02
   ro= 0.95250E-02
   zlb= 0.0635
   zlnc=0.0349
   zl = zlb + zlnc
   zlOSS = 0.675 in. = 0.017145 m
   zlOSS = 0.017145
   condcu= 391.0
   condcu = 339.
   DO 165 I=1,npts
      watts = volts(I)*amps(I)
      Q = watts
      TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I))/4.
      TBave=(Tb(1,I)+Tb(2,I))/2.
      Tvar = 0.
      DO 163 J=1,4
         Tvar = Tvar + (TCave - Tc(J,I))**2
      163 CONTINUE
      TCvar1(I)= SQrt(Tvar)
TCvar2(I) = 100.*TCvar1(I)/(TCave-TBave)
Twall(I) = TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
delt(I) = Twall(I) - TBave
flux(I) = Q/(twopi*ro*zl)
alpha(I) = flux(I)/delt(I)
delt = (Te(3,I)+Te(4,I))/2. - (Te(1,I)+Te(2,I))/2.
deltM = ((Te(1,I)+Te(2,I))/2. + TLave)/2. - TBave
Qloss(I) = condcu*pi*(rt*rt-ri*ri)*deltL/zlOSS
Q = Q - Qloss(I)
Qloss(I) = 100.*Qloss(I)/watts
Twall(I) = TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
delt(I) = Twall(I) - TBave
flux(I) = Q/(twopi*ro*zlb)
alpha(I) = flux(I)/delt(I)

165 CONTINUE
ELSEIF (isurf.EQ.5) THEN

C*************************************************************************
C           isurf = 5   Smooth Tube - 0.75 in. OD (Bajorek)
C     Three valid thermocouples at center of tube.
C*************************************************************************

ri= 0.47625E-02
rt= 0.71438E-02
ro= 0.95250E-02
zlb= 0.05080
zlnc=0.01905
zl = 0.03810 + 0.05080
condcu= 391.0
DO 575 I=1,npts
watts = volts(I)*amps(I)
Q = watts
TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I))/3.
TBave=(Tb(1,I)+Tb(2,I))/2.
TLave=(Te(1,I)+Te(2,I)+Te(3,I)+Te(4,I))/4.
Tvar = 0.
DO 573 J=1,4
Tvar = Tvar + (TCave - Tc(J,I))**2
573 CONTINUE

TCvar1(I)= SQrt(Tvar)
TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
delt(I) = Twall(I) - TBave
flux(I) = Q/(twopi*ro*zl)
alpha(I) = flux(I)/delt(I)
deltL = TCave - (Te(1,I)+Te(2,I))/2.
deltM = ((Te(1,I)+Te(2,I))/2. + TLave)/2. - TBave
Qloss(I) = condcu*pi*(ro*ro-ri*ri)*deltL/(0.00635) +
alpha(I)*twopi*ro*zlnc*deltM
Qloss1 = condcu*pi*(ro*ro-ri*ri)*deltL/(0.00635)
Qloss2 = alpha(I)*twopi*ro*zlnc*deltM
Qloss(I) = Qloss1 + Qloss2
Q = Q - Qloss(I)
Qloss(I) = 100.*Qloss(I)/watts
Twall(I) = TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
delt(I) = Twall(I) - TBave
flux(I) = Q/(twopi*ro*zlb)
alpha(I) = flux(I)/delt(I)

CONTINUE

ELSEIF (isurf.EQ.6) THEN

C****************************************************************
C isurf = 6 Smooth Tube - 0.75 in. OD (Bajorek)
C Four valid thermocouples at center of tube.
C****************************************************************

ri= 0.47625E-02
rt= 0.71438E-02
ro= 0.95250E-02
zlb= 0.05080
zln=0.01905
zl = 0.03810 + 0.05080
condcu= 391.0

C
DO 175 I=1,npts
watts = volts(I)*amps(I)
Q = watts
TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I))/4.
TBave=(Tb(1,I)+Tb(2,I))/2.
TLave=(Te(3,I)+Te(4,I))/2.
Tvar = 0.
DO 173 J=1,4
Tvar = Tvar + (TCave - Tc(J,I))**2
173 CONTINUE
TCvar1(I)= SQrt(Tvar)
TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
delt(I)= Twall(I) - TBave
flux(I)= Q/(twopi*ro*zl)
alpha(I)= flux(I)/delt(I)
deltL= TCave - (Te(1,I)+Te(2,I))/2.
deltM= ((Te(1,I)+Te(2,I))/2. + TLave)/2. - TBave
Qloss(I)= condcu*pi*(ro*ro-ri*ri)*deltL/(0.00635) +
& alpha(I)*twopi*ro*zln*c*deltM
Q = Q - Qloss(I)
Qloss(I)= 100.*Qloss(I)/watts
Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
delt(I)= Twall(I) - TBave
flux(I)= Q/(twopi*ro*zlb)
alpha(I)= flux(I)/delt(I)
175 CONTINUE

ELSEIF (isurf.EQ.9) THEN

C****************************************************************
C isurf = 9
C Nuclear clad heater - 17x17 Vantage 5A clad with
C 2-inch long cartridge heater.
C The heater diameter after press fit is 0.245 in.
C****************************************************************

ri= 0.003111
C-- The thermocouples are on a 0.329 in diameter circle.
rt= 0.004178
C-- The outside diameter is 0.374 in
ro= 0.00475
The length of the boiling region is 2.0 in.
zlb = 0.0508

Assume the length of tube in natural convection is 2 in
zlnc = 0.0508
zl = zlb + zlnc

Thermal conductivities for copper and zircalloy
condcu = 391.0
CONDZR = 7.51

DO 185 I = 1, npts
watts = volts(I) * amps(I)
Q = watts
TCave = (Tc(1,I) + Tc(2,I) + Tc(3,I))/3.
TBave = (Tb(1,I) + Tb(2,I))/2.

Since this heater lacks error TCs, assume that TLave is equal to TBave.
TLave = (Te(1,I) + Te(2,I))/2.
Tvar = 0.
DO 183 J = 1, 3
Tvar = Tvar + (TCave - Tc(J,I))**2
183 CONTINUE
TCvar1(I) = SQRT(Tvar)
TCvar2(I) = 100.*TCvar1(I)/(TCave - TBave)

TCwall(I) = TCave - Q*(1./((twopi*condcu*zlb))*ALOG(ro/rt))
RC = 0.0002/(twopi*rt*zlb)
TCwall(I) = TCave - Q*(ALOG(ro/rt)/(twopi*CONDZR*zlb)+RC)
delt(I) = TCwall(I) - TBave
flux(I) = Q/(twopi*ro*zlb)
alpha(I) = flux(I)/delt(I)
deltL = TCave - TLave
deltM = TLave - TBave
Qloss(I) = CONDZR*pi*(ro*ro-ri*ri)*deltL/(0.5*zl) +
alpha(I)*twopi*ro*zlnc*deltM

Q = Q - Qloss(I)
Qloss(I) = 100.*Qloss(I)/watts
TCwall(I) = TCave - Q*(1./((twopi*CONDZR*zlb))*ALOG(ro/rt))
delt(I) = TCwall(I) - TBave
flux(I) = Q/(twopi*ro*zlb)
alpha(I) = flux(I)/delt(I)
185 CONTINUE
ELSEIF (isurf.EQ.10) THEN

C**********************************************************
C       isurf = 10
C       Nuclear clad heater - 17x17 Vantage 5A clad with
C                             1-inch long cartridge heater.
C**********************************************************
ri = 0.003111
C The copper sleeve has radius:
RCI = 0.004178
C The thermocouples are on a 0.329 in diameter circle.
rt = 0.004178
C Assume the TCs are centrally located between the sleeve
C and the heating element.
rt = 0.003645
The outside diameter is 0.374 in.
\[ \text{ro} = 0.00475 \]
The length of the boiling region is 1.0 in.
\[ \text{zlb} = 0.0254 \]
Assume the length of tube in natural convection is 1 in.
\[ \text{zlnc} = 0.0254 \]
\[ \text{zl} = \text{zlb} + \text{zlnc} \]
Thermal conductivities for copper and zircalloy
\[ \text{condcu} = 391.0 \]
\[ \text{RES} = 0.07 \]
\[ \text{RES} \] is the electrical resistance of test heater

```
DO 195 I=1,npts
  watts = amps(I)*volts(I)
  
  Q = watts
  TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I))/4.
  TKAVE= TCave + 273.15
  CONDZR = 7.51 + 0.0209*TKAVE - 1.45E-5*TKAVE**2 +
            7.67E-9*TKAVE**3
  
  TBave=Tb(1,I)
  TLave= TBave
  Tvar = 0.
  DO 193 J=1,4
    Tvar = Tvar + (TCave - Tc(J,I))**2
  193 CONTINUE
  TCvar1(I)= SQrt(Tvar)
  TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
  
  Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
  RC is the contact resistance between the copper sleeve and
  the Zr clad. Based on natural conv. cooling tests.
  RC = 0.0002/(twopi*rt*zlb)
  RC = 1.4516E-4/(twopi*rt*zlb)
  RC = 0.00069/(twopi*rt*zlb)
  Twall(I) = TCave-Q*(ALOG(ro/rt)/(twopi*CONDZR*zlb)+RC)
  delt(I)= Twall(I) - TBave
  flux(I)= Q/(twopi*ro*zl)
  alpha(I)= flux(I)/delt(I)
  deltL= TCave - TLave
  deltM= TLave - TBave
  
  FMIN = 0.1
  FMAX = 1.0
  TSAT = 100.0
  
  RC = max((1.0+(FMIN-FMAX)*(TCave-TSAT)/100.),FMIN)*RC
  FRC = (TSAT-TCave)/50.
  TCHF = TSAT+75.
  FRC = ((TCHF-TCave)/(TCHF-70.))**3
  FRC = min(FMAX,max(FRC,FMIN))
  RC = FRC*RC
  
  FRC = max((1.0+(FMIN-FMAX)*(TCave-TSAT)/100.),FMIN)
  Heat loss in the nuclear clad heater is assumed to be due to
  conduction through the copper sleeve.
```
\[ DTC = 0.0015875 \]
\[ ACU = \pi (RCl \cdot RCl - ri \cdot ri) - \pi \cdot DTC^2 \]

* Conditions at nucleation.

* ZERR = 2.00 * 0.0254

\[ Qloss(I) = \text{condcu} \cdot ACU \cdot deltL / ZERR \]

* The low TC is assumed to be pulled out 2.54 cm based on conditions at nucleation.

\[ Qloss(I) = 0. \]

\[ Qloss(I) = \text{CONDZr} \cdot \pi (ro \cdot ro - ri \cdot ri) \cdot deltL / (0.5 \cdot zl) + \alpha(I) \cdot twopi \cdot ro \cdot zlnc \cdot deltM \]

\[ Q = Q - Qloss(I) \]

\[ RCLAD = \text{ALOG} (ro / RCl) / (twopi \cdot \text{CONDZr} \cdot zlb) \]

* The heater diameter is 0.372 in (0.0094488 m)

\[ ri = 0.47244 \times 10^{-2} \]

\[ rt = 0.63627 \times 10^{-2} \]

\[ RSDO = 0.008001 \]

\[ \text{The outside diameter of the tube is 0.740 in. = 0.018796 m} \]
ro = 0.93980E-02

C-- The heater has a heated length of 2.5 in = 0.0635 m
zlb = 0.06350
zln = 0.02530
zl = zlb + zln

C-- The error TCs are located 0.25 in off of the heater
zLOSS = 0.039065

C-- The thermal conductivity of Alloy C12200 (ASTM B359)

C is k = 196 Btu/ft2-hr-F = 339.2 W/m2-K
condcu = 339.2

RC = 0.03

DO 585 I=1,npts
  watts = volts(I)*amps(I)
  Q = watts
  TCave=(Tc(1,I)+Tc(2,I)+Tc(3,I)+Tc(4,I))/4.
  TBave=(Tb(1,I)+Tb(2,I))/2.
  TLave=(Te(1,I)+Te(2,I))/2.
  Tvar = 0.
  DO 583 J=1,4
    Tvar = Tvar + (TCave - Tc(J,I))**2
  583 CONTINUE
  TCvar1(I)= SQrt(Tvar)
  TCvar2(I)= 100.*TCvar1(I)/(TCave-TBave)
C
  Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
  Twall(I)= TCave - Q*(ALOG(ro/rt)/(twopi*condcu*zlb) + RC)
  delt(I)= Twall(I) - TBave
  flux(I)= Q/(twopi*ro*zlb)
  alpha(I)= flux(I)/delt(I)
  deltL= TCave - TLave
C
  deltM= ((Te(1,I)+Te(2,I))/2. + TLave)/2. - TBave
  Qloss(I)= condcu*pi*(RSDO*RSDO-ri*ri)*deltL/zlOSS
  Q = Q - Qloss(I)
  Qloss(I)= 100.*Qloss(I)/watts
C
  Twall(I)= TCave - Q*(1./(twopi*condcu*zlb))*ALOG(ro/rt)
  Twall(I)= TCave - Q*(ALOG(ro/rt)/(twopi*condcu*zlb) + RC)
  delt(I)= Twall(I) - TBave
  flux(I)= Q/(twopi*ro*zlb)
  alpha(I)= flux(I)/delt(I)
  585 CONTINUE
ELSEIF (isurf.EQ.12) THEN
C *****************************************************************
C *** isurf = 12 ****** Turbo-BIII Enhanced Tube
C
C *** isurf = 12 Turbo-III Tube - 0.740 in. OD (Schnelle)
C Four valid thermocouples at center of tube inserted thru
C sleeve to obtain contact with Turbo-III inner surface.
C Two thermocouples are assumed for estimating heat loss.
C The loss TCs are located at the heater edge, and
C 0.675 inch (0.017145 m) inside the heated region.
C*****************************************************************
C *** The heater diameter is 0.372 in (0.0094488 m)
  ri = 0.47244E-02
C-- The thermocouple dia. is (0.559+0.035)=0.594in = 0.0150876m
  rt = 0.75438E-02
C-- The copper sleeve OD is 0.630 in. (0.016002 m)
\[ RSDO = 0.8001 \times 10^{-2} \]

- The outside diameter of the tube is 0.740 in. = 0.018796 m
- \( r_o = 0.93980 \times 10^{-2} \)

- The heater has a heated length of 2.0 in = 0.0508 m
- \( z_{lb} = 0.05080 \)
- \( z_{ln} = 0.02530 \)
- \( z_l = z_{lb} + z_{ln} \)

- The error TCs are located 0.25 in off of the heater
- \( z_{LOSS} = 0.00635 \)

- The thermal conductivity of Alloy C12200 (ASTM B359)
- \( k = 196 \text{ Btu/ft2-hr-F} = 339.2 \text{ W/m2-K} \)

\[ \text{DO 595 } I=1,npts \]
\[ \text{watts} = \text{volts}(I) \times \text{amps}(I) \]
\[ \text{Q} = \text{watts} \]
\[ \text{TCave} = (\text{Tc}(1,I)+\text{Tc}(2,I)+\text{Tc}(3,I)+\text{Tc}(4,I))/4. \]
\[ \text{TBave} = (\text{Tb}(1,I)+\text{Tb}(2,I))/2. \]
\[ \text{TLave} = (\text{Tc}(1,I)+\text{Tc}(2,I))/2. \]
\[ \text{deltL} = \text{TCave} - \text{TLave} \]
\[ \text{Tvar} = 0. \]
\[ \text{DO 593 } J=1,4 \]
\[ \text{Tvar} = \text{Tvar} + (\text{TCave} - \text{Tc}(J,I))^2 \]
\[ \text{CONTINUE} \]

593

\[ \text{TCvar1}(I) = \sqrt{Tvar} \]
\[ \text{TCvar2}(I) = 100.*\text{TCvar1}(I)/(\text{TCave}-\text{TBave}) \]
\[ \text{Qloss}(I) = \text{condcu} \times \pi \times (RSDO \times RSDO - r_i \times r_i) \times \text{deltL}/z_{LOSS} \]
\[ \text{Q} = \text{Q} - \text{Qloss}(I) \]
\[ \text{Qloss}(I) = 100.*\text{Qloss}(I)/\text{watts} \]
\[ \text{C} \]
\[ \text{Twall}(I) = \text{TCave} - Q \times (1./(\text{two pi} \times \text{condcu} \times z_{lb})) \times \text{ALOg}(r_o/rt) \]
\[ \text{Twall}(I) = \text{TCave} - Q \times (\text{ALOG}(\text{ro}/\text{rt})/(\text{two pi} \times \text{condcu} \times z_{lb})) \]
\[ \text{delt}(I) = \text{Twall}(I) - \text{TBave} \]
\[ \text{flux}(I) = Q/(\text{two pi} \times \text{ro} \times z_{lb}) \]
\[ \text{alpha}(I) = \text{flux}(I)/\text{delt}(I) \]

595

\[ \text{CONTINUE} \]

ELSE

C*****************************************************************************
C   ERROR: not valid surface
C*****************************************************************************

\[ \text{WRITE}(*,*) \text{"**ERROR: invalid surface number", isurf} \]
\[ \text{STOP} \]

ENDIF

C*****************************************************************************
C Print out results
C*****************************************************************************

\[ \text{CALL OUTPUT(jdat)} \]
\[ \text{CALL SDFOUT(jsdf)} \]
\[ \text{WRITE}(*,225) \text{datfil} \]

225 FORMAT(10X,'OUTPUT FILE IS ',A20)
\[ \text{WRITE}(*,226) \text{sdffil} \]

226 FORMAT(10X,'PLOT FILE IS ',A20)
\[ \text{IF(IFILE.EQ.nfiles) CALL BANNER(jban)} \]
\[ \text{CLOSE(jinp)} \]
\[ \text{CLOSE(jdat)} \]
\[ \text{CLOSE(jsdf)} \]

900 CONTINUE

CLOSE(jban)
SUBROUTINE BANNER(N)
C *******************************************************************
C  Prints out a banner and ids of surfaces and substances
C ************************************************************************
WRITE(N,10)
WRITE(N,200)
WRITE(N,500)
RETURN
END
SUBROUTINE INPUT(jin)
C *******************************************************************
C Reads in an input file
C *******************************************************************
COMMON/XDATA/idnum,idate,isurf,icomp(5),frac(5),npts,iunits,
& volts(40),amps(40),P(40),Tb(2,40),Tc(4,40),Te(4,40)
COMMON/SOLUTE/ ppmB, ppmLi

C-- Function to convert degF to degC
FtoC(x)= (x-32.)*(5./9.)
C
C--- READ(jin,1) is a header line that is ignored
WRITE(*,*) 'TEST input subroutine start'
READ(jin,1)
READ(jin,10) idnum,idate,isurf,iunits
C     READ(jin,*) idnum,idate,isurf,iunits
READ(jin,1)
READ(jin,20) (icomp(i),i=1,5)
READ(jin,1)
READ(jin,30) (frac(i),i=1,5)
READ(jin,1)
READ(jin,40) npts,ppmB,ppmLi
READ(jin,1)
C-- readin temperatures
READ(jin,60) (volts(n),amps(n),(Tc(i,n),i=1,4),Tb(1,n),Tb(2,n),
& (Te(i,n),i=1,4),P(n),n=1,npts)
C-- change temps in degF (iunits=1) to degC(iunits=2) (if necessary)
IF (iunits.EQ.1) THEN
   DO 55 n=1,npts
      DO 56 i=1,4
         Tc(i,n) = FtoC(Tc(i,n))
         Te(i,n) = FtoC(Te(i,n))
      56 CONTINUE
   Tc(1,n) = FtoC(Tb(1,n))
   Tc(2,n) = FtoC(Tb(2,n))
  55 CONTINUE
ENDIF
1 FORMAT( )
10 FORMAT(10X,I5,9X,I6,10X,I5,10X,I5)
20 FORMAT(5I15)
30 FORMAT(5E15.5)
WRITE(*,*) 'TEST 09'
40 FORMAT(10X,I5,5X,F10.1,5X,F10.1)
60 FORMAT(F5.1,1X,F5.2,10F6.1,1X,F6.1)
RETURN
END
SUBROUTINE OUTPUT(J)
C ******************************************************************
C  Produces output data files
C ******************************************************************
COMMON/XDATA/idnum,idate,isurf,icomp(5),frac(5),npts,iunits,
& volts(40),amps(40),P(40),Tb(2,40),Tc(4,40),Te(4,40)
COMMON/XOUTP/delt(40),Qloss(40),alpha(40),Twall(40),flux(40),
& TCvar1(40),TCvar2(40)
COMMON/SOLUTE/ ppmB, ppmLi

WRITE(J,5)
  5  FORMAT(10X,'idnum',11X,'date',8X,'surface')
WRITE(J,10) idnum,idate,isurf
10  FORMAT(10X,5X,9X,I6,10X,15)
WRITE(J,20)
20  FORMAT(5(6X,'Component'))
WRITE(J,30) (icomp(i),i=1,5)
30  FORMAT(5I15)
WRITE(J,40)
40  FORMAT(5(7X,'Fraction'))
WRITE(J,50) (frac(i),i=1,5)
50  FORMAT(5E15.5)
WRITE(J,60)
60  FORMAT(11X,'npts')
WRITE(J,70) npts
70  FORMAT(10X,I5)
WRITE(J,80)
80  FORMAT(/,9X,'V',4X,'I',2X,'TC-1',2X,'TC-2',2X,'TC-3',2X,'TC-4',
  &      2X,'TB-1',2X,'TB-2',2X,'TE-1',2X,'TE-2',2X,'TE-3',
  &      2X,'TE-4',4X,'P')
WRITE(J,100) (volts(N),amps(N),(Tc(i,N),i=1,4),Tb(1,N),Tb(2,N),
  &     (Te(i,N),i=1,4),P(N),N=1,npts)
100  FORMAT(4X,F6.2,F5.2,10F6.1,F7.1)
WRITE(J,120)
120  FORMAT(/,2X,'NO',1X,' SUPERHEAT',2X,'HEAT flux',5X,'alpha',
  &      5X,'Qloss',5X,'Twall',4X,'TCvar1',4X,'TCvar2')
WRITE(J,140) (N,delt(N),flux(N),alpha(N),Qloss(N),Twall(N),
  &     TCvar1(N),TCvar2(N),N=1,npts)
140  FORMAT(2X,I2,1X,F10.1,1X,4F10.1,F10.2,F10.1)
IF((ppmB+ppmLi).GT.0.) THEN
WRITE(J,150) ppmB
150     FORMAT(/,' Boron concentration   =',F8.1,' ppm')
WRITE(J,160) ppmLi
160     FORMAT(/,' Lithium concentration =',F8.1,' ppm')
ENDIF
RETURN
END
SUBROUTINE SDFOUT(J)
C ********************************************************************
C  Creates plot data files
C ********************************************************************
COMMON/XDATA/idnum,idate,isurf,icomp(5),frac(5),npts,iunits,
&     volts(40),amps(40),P(40),Tb(2,40),Tc(4,40),Te(4,40)
COMMON/XOUTP/ delt(40),Qloss(40),alpha(40),Twall(40),flux(40),
&     TCvar1(40),TCvar2(40)
C
WRITE(*,15)
ZKW = 1000.
WRITE(J,20) (delt(N),flux(N)/ZKW,alpha(N)/ZKW,Twall(N),
  &     VOLTS(40),amps(40),P(40),Tb(2,40),Tc(4,40),Te(4,40),
  &     (Te(i,N),i=1,4),P(N),N=1,npts)
110  FORMAT(4X,F6.2,F5.2,10F6.1,F7.1)
WRITE(J,120)
120  FORMAT(/,2X,'NO',1X,' SUPERHEAT',2X,'HEAT flux',5X,'alpha',
  &      5X,'Qloss',5X,'Twall',4X,'TCvar1',4X,'TCvar2')
WRITE(J,140) (N,delt(N),flux(N),alpha(N),Qloss(N),Twall(N),
  &     TCvar1(N),TCvar2(N),N=1,npts)
140  FORMAT(2X,I2,1X,F10.1,1X,4F10.1,F10.2,F10.1)
IF((ppmB+ppmLi).GT.0.) THEN
WRITE(J,150) ppmB
150     FORMAT(/,' Boron concentration   =',F8.1,' ppm')
WRITE(J,160) ppmLi
160     FORMAT(/,' Lithium concentration =',F8.1,' ppm')
ENDIF
RETURN
END
& Tb(1,N),P(N),N=1,npts)
WRITE(*,20) (delt(N),flux(N)/ZKW,alpha(N)/ZKW,Twall(N),
& Tb(1,N),P(N),N=1,npts)

15 FORMAT(' Delt(n) flux/1000 alpha/1000',
& ' Twall Tb P')
20 FORMAT(5X,F10.4,5X,F10.2,5X,F10.1,3F10.1)
C
RETURN
END
A.2 Sample Data File.

Input file with data for pool boiling test at 1000 psi pressure for ionized water at 30 subcooling.

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APPENDIX B

B.1 Product Reference

1. Duniway Stockroom Corporation
   Product: Copper –O rings, 2006, Code: SG-600
   Details:
   Duniway Stockroom Corporation
   1305 Space Park Way
   Mountain View, CA 94043
   Ph :800-446-8811

2. Richard Greene Company
   Product: electrical test heater, Code: SN6N-3206
   Manufacturer Details:
   Watlow
   12001 Lackland Road
   St louis, MO 63146-41001
   Ph: 314-878-4600, fax: 314-878-6814

3. McMaster-Carr
   Product : Pte O-rings
   Manufacturer details:
   McMaster-Carr Supply Company
   P.O. Box 4355
   Chicago, IL 60680-4355
   Ph: 630-833-0300 fax: 530-834-9427

4. Omega Engineering Inc
   Product: Pressure Transducer, Model: PX35k1-3kGV
   Product: Pressure Transmitter, Model: DP-25S
   T-type thermocouple, Model : HTQIN 316G-12
   Details:
   Omega Engineering Inc
   One Omega drive,
   PO Box 4047
   Stamford, CT  06907-0047
   Ph: 203-359-1660 fax- 203-359-7700

5. Microgroup
   Product : Inconel Tubes, Grade : 600
   Manufacturer Details:
   Microgroup Inc.
   7 Industrial Park Road
   Medway, MA 02053-1732
6. Linweld
   Product: Nitrogen cylinder, Type: Industrial
   Details:
   Linweld Inc.
   PO Box 682
   Manhattan, KS 66505
   Ph: 785-537-0395

7. Yale Film and Video
   Product: 16mm film processing, Model: Hawkeye Surveillance Film
   Details:
   Yale Film and Video
   10555 Victory Blvd,
   North Hollywood, CA 91606
   Ph 818-508-9253, fax 818-762-0688

8. Machined Glass Specialists Inc
   Product: Fused Quartz Window, Model: custom order.
   Dimensions: 1.75” Dia x 0.75” thick
   Details:
   Machined Glass Specialists, Inc.
   245 Hiawatha Trail
   Springboro, Ohio 45066
   Ph: 937-743-6166, fax 937-743-6168

9. John C Ernest Co Inc
   Product: Gaskets for fused quartz, Model: 583 1.75-1.00-0.0625
   Details:
   John C. Ernest Co. Inc
   21 Gail Ct.
   Sparta, NJ 07871
   Ph: 973-940-1600.  www.sightglass.com

10. Visual Instrumentation Corporation
    Product: high speed camera (purchased 1984), Model: Hycam: 41-0005
    Details:
    Visual Instrumentation Corporation
    1110 West Avenue L-12, Unit 2,
    Lancaster, CA, USA 93534-7039
    Ph: 661-945-7999 fax 661-723-5667
11. Stern Laboratories Inc  
   Product: electrical test heater  
      Solid-state electrical DC power supply  
Details:  
Stern Laboratories, Inc.  
1590 Burlington Street East,  
Hamilton, Ontario, Canada L8H 3L3  
Ph 905-548-5300 fax 905-545-5399

12. Alfa Aesar  
   Product: Lithium metaborate puratonic  
      Boric Acid  
Details:  
www.alfaaesar.com

13. Swagelock  
   Product: union tube fittings  
Details:  
Kansas City Valve & Fitting Co.  
4707 Roe Parkway  
Roeland Park, Kansas 66205

14. Fluke  
   Product: Clamp Ammeter, Model: i-410  
Details:  
www.fluke.com

15. Systat Inc.  
   Product: Sigmascan- Image Analysis Software.  
Details:  
http://www.systat.com/products/SigmaScan/
\textbf{B.2 Sample Heat Exchanger Calculations}

Assuming $\Delta P = 40$ psi.
Assuming internal diameter of inconel tube is 0.5", $D_i = 0.01021\text{m}$
$D_o =$ Outer diameter of inconel tubing = 0.127 m
$\Delta P = 28122.6 \text{N/m}^2$.

Head loss $h_L = \frac{\Delta P}{\rho g} = 2.87 \text{m}$

\(\rho\) = density of the fluid = 996 kg/m$^3$
\(g\) = acceleration due to gravity = 9.81 m/sec$^2$

Using equation (3.3) with initial guess $f = 0.02$ and assuming the length of the tube = 8.5 m, we have

Solid cross-section of the inconel tubing = \(\frac{\pi(D_o^2 - D_i^2)}{4}\) = 0.0000158 sq m.

Resistance of the inconel tubes is $R = \frac{\rho l}{A}$ equation (3.1)

Total Electrical resistance of the inconel tubing = \(\frac{1.03*8.5}{0.0000158}\) = 0.553 $\Omega$.

Using Darcy’s formula equation (3.3) gives

\[ V = \sqrt{\frac{2gD_i h_i}{fL}} = 1.9865 \text{ m}^2/\text{sec} \]

Viscosity of the fluid = $\mu = 0.000855 \text{N/m}^2$

Reynolds number = $Re = \left(\frac{VD_i \rho}{\mu}\right) = 27492$

Reynolds number is greater than 2300, indicating that the flow is turbulent.
Surface area of the inconel tubes = $(\pi)*D_i*L = 0.317238 \text{sqm}$
Corrected friction factor $f$ (from Moody chart) = 0.027

Prandtl number = 5.846 and $n = 4$.
From equation (3.5) the Nusselt number is

\[ Nu = 0.023 \left(Re^{\frac{4}{5}}Pr^n\right) = 165.9 \]

Required flow rate of water to remove the heat generated in the inconel tubes is

\[ Q \text{ (flow rate) = } A*V \]

where $A$ is the area of tube cross section and $V$ is the velocity of the fluid.
\[ A = \frac{\pi D_i^2}{4} = \frac{\pi (0.01021)^2}{4} = 0.000111 \text{m} \]

\[ Q = \text{flow rate} = A \times V = 0.000111 \times 1.9865 = 0.00022 \text{ m}^3/\text{sec} \]

Heat transfer coefficient between inconel tubes and cold water is given by following equation

\[ h = \left( \frac{N_u K_{\text{water}}}{D_i} \right) \]

where \( K_{\text{water}} \) = thermal conductivity of water = 0.613 W/mK

Substituting the values of nusselt number, thermal conductivity of water and internal diameter we have

Heat Transfer Coefficient = 8560 W/m\(^2\)C

Total heat transfer from Inconel tubes to water is \( h \times A_{\text{surf}} \times (\Delta T) \)

where

\( A_{\text{surf}} \) is the interface area through which heat transfer occurs

\( \Delta T = \) temperature gradient.

Assuming a temperature gradient to be 15\(^\circ\)C, the total heat transfer is

\[ (8.56 \text{ kW/m}^2\text{C})(0.000111)(15) = 40.73 \text{ kW}. \]

The maximum heat generated is approximately 20 kW. This iteration was repeated with the improved friction factor and smallest possible dimensions (inside diameter and length) was chosen for the inconel tubes.