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HEAT TRANSFER COEFFICIENTS OF CONDENSING VAPORS

BY

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A THESIS SUBMITTED TO THE FACULTY OF THE DEPARTMENT OF CHEMICAL ENGINEERING OF NEWARK COLLEGE OF ENGINEERING

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#### ABSTRACT

The film coefficients of heat transfer for vapors condensing on a single horizontal tube have been measured for methyl, i-propyl, and m-butyl alcohols. The equipment used was specifically designed for these studies, and represents an improvement over that used by other investigators. The data collected were analyzed using a modified Wilson method which is based on a rigorous theoretical analysis.

It has been found that the condensing film coefficients for methyl and i-propyl alcohols wary with the heat duty. This is to be expected from the analysis mentioned above. No variation was found for n-butyl alcohol.

The observed values of the heat transfer coefficients have been compared with the values calculated with the Musselt and Bromley equations. The Bromley equation is a modification of the Musselt equation and takes into account the effect of the heat capacity of condensate. Excellent agreement between the observed coefficients and those predicted by the Musselt equation were obtained with methyl and i-propyl alcohols. Good agreement was found with n-butyl alcohol. In all cases the Bromley equation predicted higher values of the condensing film coefficient than did the Musselt equation.

Close agreement between the observed and theoretical coefficients is a result of the refined theoretical and experimental methods used.

1

## APPROVAL OF THESIS

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FOR

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APPROVED:

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#### INTRODUCTION

Investigators have generally used one of two methods for measuring the film coefficients of heat transfer for vapors condensing on single horizontal tubes. These are the embedded thermocouple method as used by Kirkbride (14), Othmer and Berman (18), and Wallace and Davison (24); and the Wilson method (26). Both of these methods have inherent disadvantages.

Rhodes and Younger (23) pointed out that the average wall temperature as measured with thermocouples can be computed only by making certain assumptions that may not be valid. Baker and Mueller (1) later proved that there is no point on a tube wall at which a thermocouple can be located to obtain a truly representative tube wall surface temperature.

Rhodes and Younger (23) showed that some of Wilson's assumptions were not correct. Based on this earlier work, Chu, Flitcraft, and Holeman (6) proposed a unique modification of the Wilson method using a rigorous theoretical analysis. This modified technique has greatly improved the Wilson method.

Chu and associates' proposed technique has been applied to only a limited extent. As a result, this investigation was initiated to test this modified Wilson method using another homologous series of organic compounds. Methyl, i-propyl, and n-butyl alcohols have been selected for these studies. The physical properties of these alcohols have been widely investigated and are more reliable than those of more complex organic compounds. Thus, the predicted condensing film coefficients, which are to be compared with the observed coefficients, can be calculated with a reasonable assurance of reliability.

The predicted coefficients will be calculated using both the Musselt and Bromley equations. The Bromley equation (4) is a modification of the Musselt equation and includes a correction for the effect of the heat capacity of condensate. The Bromley equation is to be compared with Nusselt's equation to determine the magnitude of this correction.

The equipment used to measure the heat transfer coefficients was specifically designed for these studies and represents an improvement over that used by the other investigators. 2

#### THEORY

The Musselt equation (15) (17) is generally used to predict the coefficients of hest transfer for the filmwise condensation of pure vapors on a colder surface. Applied to a single, horizontal, cylindrical tube, the equation is

$$h_0 = 0.725 (k_f^3 \rho_f^2 g \lambda / D_0 u_f \Delta t)^{0.25}$$
 I

which is based on the assumption that streamline flow exists throughout a continuous condensate film, and that gravity alone causes the condensate to flow over the smooth surface. The possible acceleration effect of the vapor velocity upon the film thickness is also neglected.

Many investigators (14) (16) (24) have measured the condensing film coefficients of pure organic vapors with the aid of embedded thermocouples. Inasmuch as the condensate film thickness around the perifery of a cylindrical tube varies, it is to be expected that the film temperature would also vary. Therefore, the measurement of surface temperatures with thermocouples may result in considerable error. Baker and Mueller (1) proved that there is no point on the surface of a cylindrical tube at which a thermocouple can be located to obtain a representative surface temperature.

Investigators (3) (23) have utilized the Wilson method to avoid the difficulties associated with direct temperature measurements. The Wilson method is represented by the following equation

$$\mathbf{R} = \sum \mathbf{R}_{1} = \mathbf{R}_{V} + \mathbf{R}_{V} + \mathbf{a} / \mathbf{V}^{0 \cdot 8}$$
 II

where  $R_{\psi}$  is the thermal resistance of the condensate film at a finite cooling water flow rate, and "a" is a constant. This equation is based on the assumption that changes in cooling water rate have no effect upon  $R_{\psi}$ . Rhodes and Younger (23) pointed out that  $R_{\psi}$  varied with water rate and postulated that

$$R_{\psi} = R_{\psi\phi} + b/\psi^{0,\delta} \qquad III$$

Beatty and Katz (3) applied this method to their work with finned tubes, with a modification to allow for the effect of cooling water temperature upon the water film resistance.

Chu, Fliteraft, and Heleman (6) stated that Equation III was empirical, since there is no theoretical justification for setting up R<sub>y</sub> as a function of R<sub>ye</sub> and the cooling water flow rate. They proposed a modification of the Wilson method based on a rigorous theoretical analysis. It was pointed out in their paper and by other investigators (7) (23) that the group  $(k_f^3 P_f^2 g \lambda //4_f)^{0.25}$  appears to remain constant with varying temperature for most organic solvents. Therefore, for steady state heat transfer through a condensing vapor, the following equation has been derived (6).

$$h_{\phi} = c q^{-1/3} \qquad IV$$

This equation will result in a straight line with a negative slope equal to one-third on a log-log plot. Thus, Chu and associates (6) have deduced that the condensing film coefficient varies with the heat transferred.

The values of the group  $(k_f^3 \beta_f^2 g_{\lambda} / M_f)^{0.25}$  for methyl, i-propyl,

and n-butyl alcohols are listed in Table 1. Also included in this listing are the values of this group for other organic compounds as reported by Chu and associates (7).

The overall thermal resistance from the condensing vaper to the cooling water side of a condenser tube is represented by

$$\frac{1}{U_0 A_0} = \frac{1}{b_0 A_0} + \frac{x}{k_0 A_{sv}} + \frac{D_1 U_{s2}}{150 A_1 (1+0.011t) V} + \frac{1}{V_0 A_0}$$

From Equation IV it is evident that  $h_0$  is constant at constant values of q. Also, the thermal resistance of the tube wall is negligible compared to the other terms of Equation V. Therefore, the first two terms of Equation V are constant at constant values of q. A plot of  $1/V_0A_0$  against  $1/(1 \ 0.011t)V^{0.6}$  should result in a straight line at equal values of q. The intercept of this line, equal to  $1/h_0A_0$   $x/k_0A_{gV}$ , can be used to calculate the condensing film coefficient,  $h_0$ , since

$$\frac{1}{V_0 A_0} = \frac{T}{q} = \frac{1}{h_0 A_0} + \frac{X}{k_0 A_{av}}$$

and

$$\frac{1}{h_0} = \frac{1}{U_0} - \frac{x}{k_0} \frac{A_0}{k_0}$$
 VII

Several sets of experimental data can be obtained by varying the cooling water flow rate and the pressure in the test condenser vapor space. For any one set, the overall water to vapor temperature difference would be kept constant; and from one set to another the temperature difference changed by readjusting the pressure in the vapor space. Two or more points, at constant q, can be gotten from the above sets of data. These points, when replotted as  $1/U_{c}A_{c}$  against

# TABLE 1

RATIO OF  $(k_f^3 P_f^2 s h/\mu_f)^{0.25}$  at different temperatures

COMPOUNDS	UPPER TEMP.	LONER TEMP.	RATIO
Methyl alcohol	72	63	1.02
1-Propyl alcohol	80	69	1.13
n-Butyl alcohol	<b>9</b> 8	86	1.09 ~
Sthyl acetate	60	ho	1.01
Benzene	60	30	1.03
Toluene	90	40	1.09
Trichlorosthylens	60	ьо	1.02
Mitromethane	80	50	1.02
Bronobensens	100	50	1.08
n-Hexyl alcohol	100	50	1.07
Perchloroethylene	100	50	1.03
Carbon tetrachloride	60	30	1.11

.

 $1/(1+6.011t) = \sqrt{0.8}$ , yield a straight line. The value of h<sub>0</sub> at the constant value of q is then calculated from the intercept of this line.

As previously stated, the Musselt equation is used to predict condensing film coefficients. Browley (4) has pointed out that in the basic derivation by Musselt the effect of the heat capacity of condensate was assumed negligible. This assumption is valid at low pressures. However, at high pressures and large tube wall to saturated vapor temperature differences, sensible heat may significantly affect heat transfer.

This consideration led to the theoretical development of a modified Husselt equation. The equation for vapors condensing on a single horizontal tube is  $h_{0} = 0.728 \left[ \frac{ke^{3} f_{e}(f_{e}-f_{e})g_{A}(1+0.k-A)^{2}}{D_{0}\mathcal{U}_{f}At} \right]^{0.25}$ VIII

Bromley's equation should predict higher values of h than the Masselt equation.

#### DESCRIPTION OF APPARATUS

The equipment used in this work, as shown in Figures 1 and 2, consisted basically of a kettle, a single tube horizontal condenser, a cooling water circulation system, and a vacuum pump.

The jacketed kettle was of stainless steel construction and had a five gallon capacity. Steam, at an initial pressure of 90 psig., was introduced into the jacket through a 3 to 15 psig. throttling valve.

The test condenser was a 0.375 inch outside diameter brass tube 2h inches long having a wall thickness of 0.035 inches. The effective outside surface area was 0.196 sq. ft., and the thermal conductivity was 60 Btu/(Hr.) (sq. ft.) (°F/ft.). A 2.5 inch schedule h0 stainless steel pipe, flanged at both ends, was used as the condenser jacket.

Vapor from the kettle entered the jacket through three 0.5 inch diameter distributors. Condensate returned to the kettle by gravity flow through two downcomers and liquid seal traps. The temperature of the uncondensed waper taken overhead was measured by a -1 to 101°C or 99 to 201°C thermometer graduated in 0.1°C increments. A glass secondary condenser was used to assure that excess vapor was passing the test condenser at all times. Condensate returned to the kettle through a liquid seal trap.

The condenser jacket contained a special thermocouple installation used for measuring bulk condensate temperatures. A finely threaded fitting permitted adjustment of the position of the thermocouple junction.







EXPERIMENTAL APPARATUS

CONDENSER JACKET

10

The system was evacuated and maintained at low pressures by a Cenco-Hypervac & pump of 1.44 cubic ft. per minute free air displacement. The pressure was controlled at the desired level by a cartesian manostat and measured with a U-tube mercury manometer.

The kettle - condenser - vacuum system was checked for leaks. It was considered to be air-tight when no noticeable increase in absolute pressure occurred over a one-half hour period, after evacuation to an absolute pressure level of one inch of mercury.

Water was circulated by a centrifugal pump rated at 25 gallons per minute at 80 ft. head and 1.0 specific gravity. Flow rates were measured with a calibrated Fisher-Porter rotameter rated a 13.6 gallens per minute.

Two 55 gallon drums were used as holdup and constant head supply for the pump. Fresh water flowed to the pump from the supply drum. Heated water from the condenser could be discharged to the second drum and then to the sever. The secondary drum was equipped with a steam coil which allowed pre-heating of the cooling water.

Condenser inlet water temperatures were measured with a 0 to  $50^{\circ}$ C thermometer graduated in 0.1°C increments. The outlet water temperature measurements were made with a thermocouple installation. A pair of 0 to 6°C Beckmann thermometers, each of which could be read to 0.002°C, was used to determine the water temperature rise in the test condenser.

The entire kettle-condenser system was well insulated. Magnesia

lagging was used on all pipe lines. The condenser jacket was enclosed in a plywood box which was filled with powdered vermiculite. This method of insulation minimized heat losses.

Eighteen gage, copper-constantan thermocouples were used. These thermocouples were factory calibrated at  $\pm 0.8$  °C up to a temperature of 375 °C.

The alcohols used were reagent grade containing a maximum of 0.5 per cent impurities. The boiling ranges were less than  $1^{\circ}$ C.

#### EXPERIMENTAL PROCEDURE

The test alcohol was charged to the kettle after the system had been cleaned. Before heating with steam, the cooling water was turned on and the system was allowed to reach a steady state condition. At this point a series of reference readings was taken on the Beckmann thermometers. The rotaneter reading and water temperature were also taken. This blank correction on the Beckmann thermometers was used to determine the true water temperature rise in the condenser tube, as discussed in the appendix section "Reference Correction of Beckmann Thermometers".

The system was then evacuated, bypassing the manostat to permit more rapid evacuation. When the absolute pressure decreased to the vapor pressure of the alcohol in the kettle, the pump was turned off and the bypass closed. Steam was then admitted to the kettle jacket. When the system pressure reached the desired level, the manostat was put in operation and the vacuum pump restarted. This method of startup minimized the pessibility of non-condensable gases remaining in the system.

Five complete sets of readings were taken at one minute intervals after the system reached equilibrium. The average of these readings was used in the calculations. A set of data consisted of a rotameter reading, absolute inlet and outlet water temperatures, vapor temperature, bulk condensate temperature, kettle pressure, and a simultaneous reading of the Beckmann thermometers. The secondary condenser was constantly checked to assure that excess vapors were present in the test condenser at all times.

After the completion of a run, the water rate was changed. If necessary, the pressure level was readjusted to maintain a constant overall water to vapor temperature difference.

Twenty to thirty minute time intervals were required for the system to return to equilibrium after conditions were changed.

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#### EXPERIMENTAL RESULTS AND TREATMENT OF DATA

The experimental results for methyl, i-propyl, and n-butyl alcohols are presented in Tables 2, 3, and 4, which include the calculated results of  $(1 \times 10^3) / (1+0.011t)V^{0.8}$ . The data are plotted in Figures 3 to 11. Figures 3, 6, and 9 show the variation of the water flow rate and temperature rise in the test condenser. Figures 4, 7, and 10 are a form of the Wilson plot and represent the heat transferred as a function of  $1/V^{0.8}$  at varying overall vapor to cooling water temperature differences.

In determining the heat transfer coefficients,  $h_{0}$ , a series of straight lines were drawn parallel to the abscisses of Figures 4, 7, and 10, intersecting two or more of the constant vapor - water temperature difference lines. These horizontal lines represent constant values of the heat transferred, q. At each intersection, the overall temperature difference, and  $(1 \times 10^3) / (1+0.011t)v^{0.8}$  were read.  $\Delta T/q$  was calculated and plotted against  $(1 \times 10^3) / (1+0.011t)v^{0.8}$ for each value of q. These plots, shown in Figures 5, 8, and 11, yielded straight lines which were extrapolated to the ordinates. The intercepts represent  $\Delta T/q$  or  $1/V_{0}A_{0}$  at infinite water flow. The values of  $h_{0}$ , the condensing film coefficients, are calculated from these intercepts.

TABLE 2 TABULATED RESULTS - METHYL ALCOHOL

							<b>.</b> .	
RUN NO.	INLET WATER TEMP. °C	WATER TEMP. RISE C	VAPOR TEMP. °C	OVERALL WATER- VAPOR AT. C	WATER Flow W Lb/hr	HEAT DUTY Q BTU/HR	$\frac{1\times10^{3}y^{8}}{100.011t}$	
12345	5.50	1.759	64.29	57.94	2610	8260	1.24	
	5.80	2.280	64.60	57.65	1950	8010	1.58	
	5.23	1.536	64.93	59.04	3310	9150	1.05	
	6.15	3.108	65.50	57.77	1320	7390	2.11	
	6.72	3.964	64.36	55.62	980	7000	2.65	
6	5.50	1.957	64.59	58.13	2270	8000	1.29	
7	5.90	2.730	66.14	58.86	1580	7760	1.86	
8	7.80	4.491	65.25	55.20	830	6695	2.94	
9	7.30	3.449	65.26	56.21	1160	7210	2.30	
10	7.15	2.744	55.80	47.19	1250	6175	2.19	
11 12 13 14 15	5.96 5.70 5.60 5.70	1.883 1.462 1.120 1.357 1.802	55.81 55.69 55.91 55.86 55.93	48.89 49.27 49.70 49.65 49.36	2040 2640 3310 2950 2290	6920 6950 6670 7210 7430	1.51 1.23 1.044 1.14 1.39	
16	6.13	2.410	56.06	48.72	1550	6720	1.87	
17	7.25	3.975	56.98	47.71	850	6080	2.96	
18	6.20	1.540	51.65	44.70	2295	6360	1.38	
19	6.10	1.338	51.72	45.00	2670	6430	1.225	
20	6.04	1.166	51.83	45.28	2960	6220	1.13	
21	6.63	1.893	52.63	45.06	1845	6290	1.62	
22	6.20	1.146	52.09	45.52	3260	6720	1.033	
23	6.46	1.866	52.25	44.42	1910	6410	1.59	
24	6.90	2.202	52.43	43.73	1645	6520	1.12	
25	7.12	2.650	52.73	42.60	1250	5970	2.20	
26	7.65	3.477	52.96	41.02	920	5750	2.78	
27	6.40	1.024	45.71	39.10	3350	6175	1.019	
28	6.37	1.155	45.65	38.87	2920	6070	1.138	
29	6.40	1.262	45.79	38.86	2630	5980	1.23	
30	6.50	1.386	45.75	38.56	2350	5860	1.35	
31	6.80	1.543	46.36	38.49	2035	5650	1.50	
32	7.02	1.897	46.63	38.08	1630	5560	1.79	
33	7.43	2.249	47.28	37.66	1315	5325	2.09	
34	8.20	3.022	48.03	36.32	930	5060	2.73	
35	6.35	1.015	46.23	39.32	3330	6080	1.035	

TABLE 2 (Con't)TABULATED RESULTS - METHYL ALCOHOL

RUN NO.	INLET WATER TEMP. C	WATER TEMP. RISE C	VAPOR TEMP <sup>o</sup> c	OVERALL WATER- VAPOR AT. C	WATER Flow W Lb/hr	HEAT DUTY Q BTU/HR	1x10 <sup>3</sup> y <sup>8</sup> 170.011t
36 37 38 39 40	6.50 6.55 6.55 6.60 6.75	1.307 1.440 1.633 1.696 2.137	60.76 60.76 60.75 60.80	53.61 53.49 53.39 53.30 52.98	3330 2970 2590 2345 1950	7800 7700 7630 7165 7490	1.005 1.118 1.243 1.342 1.550
41 42 43 44 45	6.90 7.20 7.40 7.60 7.80	2.478 2.983 3.357 3.897 4.330	61.87 61.89 62.09 62.56 62.69	53.73 53.20 53.11 53.01 52.72	1660 1300 1150 970 840	7410 6985 6950 6805 6545	1.757 2.11 2.32 2.62 2.92





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	A 7000 Btu/hr	· · · · · · · · · · · · · · · ·
	• 6600 Btu/hr	
,010	▲ 5750 Btu/hr	
· · · · · · · · · · · · · · · · · · ·	· · · · · · · · · · · · · · · · · · ·	
	0,4 0,8 1,2 1,6 2,0 2,4 2,8 3,2	······································

TABLE 3 TABULATED RESULTS - ISOPROPYL ALCOHOL

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RUN NO.	INLET WATER TEMP. °C	WATER TEMP. Rise °C	VAPOR TEMP <sup>O</sup> C	OVERALL WATER- VAPOR AT.ºC	WATER Flow W Lb/hr	HEAT DUTY Q BTU/HR	<u>1×10<sup>3</sup>y8</u> 1/0.011t
46	7 • 15	2.080	83.13	74.94	1320	4940	2.10
47	6 • 80	1.405	82.80	75.30	2010	5080	1.523
48	6 • 70	1.169	82.77	75.49	2370	4980	1.338
49	6 • 70	1.032	82.30	75.08	2670	4970	1.200
50	7 • 35	2.653	82.37	73.69	980	4675	2.63
51 52 53 55 55 55	7.00 7.73 5.97 5.90 6.60	2.286 3.605 .876 1.038 2.444	82.36 82.35 82.40 82.40 71.03	74.22 72.82 75.99 75.98 63.21	1155 680 3255 2 <b>7</b> 20 990	4760 4420 5120 5080 4360	2.33 3.48 1.05 1.30 2.65
56	6.90	2.952	71.78	63.40	810	4300	3.08
57	7.32	3.572	72.35	63.24	650	4180	3.64
58	5.98	1.475	69.24	62.52	1700	4510	1.775
59	5.75	1.277	69.71	63.32	1980	4550	1.575
60	5.60	1.093	69.57	63.42	2300	4530	1.388
61	5.50	•973	69.70	63.72	26 <b>7</b> 5	4680	1.282
62	5.50	•861	69.43	63.50	3010	4660	1.123
63	5.60	•803	69.18	63.18	3240	4680	1.052
64	6.52	2•221	70.43	62.80	1100	4440	2.42
65	6.70	1•822	65.29	57.68	1245	4985	2.205
66	7.00	2.327	66.22	58.06	975	4090	2.68
67	7.40	2.576	67.71	59.02	830	3840	3.00
68	7.90	3.183	68.71	59.22	660	3780	3.59
69	6.51	1.421	65.48	58.26	1650	4220	1.788
70	6.30	1.159	64.05	57.17	1990	4150	1.547
71	6.25	•998	64.62	57.87	2310	4150	1.382
72	6.10	•820	64.41	57.90	2785	4110	1.190
73	6.00	•714	64.74	58.38	3220	4140	1.060
74	5.90	•913	59.09	52.71	2300	3780	1.385
75	6.00	1•095	59.46	52.91	1930	3800	1.590
76	5.80	•774	58.28	52.09	2705	3760	1.220
77	5.75	•636	57.76	51.69	3240	3710	1.060
78	6.24	1•329	59.98	53.07	1540	3680	1.89
79	6.50	1•582	60.44	53.15	1270	3620	2.20
80	6.87	1•945	61.09	53.25	1070	3750	2.50
81	7.20	2•439	61.51	53.07	830	3640	3•15
82	8.40	3•735	62.44	52.17	505	3400	4•46
83	7.63	2•883	61.58	52.51	660	3420	3•58





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TABLE4TABULATED RESULTSN-BUTYL ALCOHOL

RUN NO.	INLET WATER TEMP.	WATER TEMP. RISE OC	VAPOR TEMP. °C	OVERALL WATER- VAPOR AT, C	WATER Flow W Lb/hr	HEAT DUTY Q BTU/HR	1x10 <sup>3</sup> y <sup>8</sup> 1/0.011t
84	7.50	2.251	95.14	86.51	1400	5665	2.00
85	8.10	3.378	95.86	86.07	930	5660	2.72
86	7.30	1.958	95.45	87.17	1680	5920	1.74
87	7.20	1.685	94.01	85.97	1980	6010	1.53
88	7.02	1.425	94.01	86.28	2345	6020	1.34
89	7.00	1.255	94.06	86.43	2680	6050	1.20
90	7.18	1.030 9	93.45	85.75	3235	6000	1.04
91	9.00	4.399	97.25	86.05	700	5540	3.34
92	8.90	4.358	97.14	86.06	720	5650	3.26
93	7.72	2.484	99.27	90.31	1365	6100	2.03
94	8.25	3.570	99.53	88.69	955	6140	2.62
95	8.88	4.427	100.23	89.14	750	5980	3.16
96	7.50	1.057	98.65	90.62	3240	6170	1.03
97	7.58	1.331	98.55	90.30	2595	6220	1.23
98	8.04	2.250	98.61	89.44	1515	6140	1.86
99	7.80	1.688	99.10	90.46	2040	6210	1.47
100	7.96	1.623	104.07	95.30	2265	6630	1.36
101	8.11	1.866	104.10	95.06	1940	6520	1.53
102	8.37	2.304	104.40	94.88	1595	6610	1.78
103	8.80	2.635	105.78	95.66	1400	6630	1.96
104	9•30	3.782	106.60	95.41	960	6540	2.59
105	9•50	4.253	107.07	95.44	830	6350	2.89
106	6•48	5.371	103.85	94.68	650	6290	3.64
107	8•00	1.169	103.65	95.07	3210	6750	1.03
108	8•00	1.299	103.80	95.15	2840	6640	1.13
109	8.00	1.420	108.67	99.96	2740	7010	1.17
110	7.98	1.220	108.38	100.40	3210	7050	1.03
111	8.30	1.714	109.35	100.19	2270	7000	1.35
112	8.35	1.901	109.79	100.49	1980	6790	1.50
113	8.50	2.422	110.45	100.74	1620	7060	1.75
114	8.80	2.992	110.44	100.49	1310	7050	2.06
115	6.67	5. <b>517</b>	108.60	99.17	665	6610	3.63
116	8.23	1.237	108.78	99.93	3210	7150	1.03
117	9.30	4.605	111.31	99.71	800	6640	3.03
118	8.80	3.406	111.07	100.57	1120	6860	2.31

	IA						
RUN NO.	INLET WATER TEMP. °C	WATER TEMP. RISE °C	VAPOR TEMP. <sup>O</sup> C	OVERALL WATER- VAPOR AT. C	WATER Flow W Lb/hr	HEAT DUTY Q BTU/HR	1x10 <sup>3</sup> y <sup>8</sup> 170.011t
119	8.80	3.658	116.25	105.62	1115	7340	2.33
120	8.40	2.137	116.01	106.54	1995	7670	1.49
121	8.60	2.659	116.14	106.21	1580	7560	1.78
122	8.80	3.180	116.71	106.29	1310	7500	2.05
123	9.30	4.496	117.37	105.82	990	7370	2.69
124	7.86	5•768	116.79	106.00	665	6910	3•53
125	7.43	5•479	116.27	106.10	705	6950	3•37
126	8.81	5•056	116.54	105.20	775	7060	3•06

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	FIGURE 11 - RELATION BETWEEN OVER-ALL
	THERMAL RESISTANCE AND WATER FLOW RATE
.028	N-BUTYL ALCOHOL
· · · ·	
.026	
0	
\$ .025	
	0 7150 Btu/hr
,024	△ ?000 Btu/hr □ 6780 Bts/hr
	▲ 6350 Btu/hr ● 6030 Btu/hr
023	<b>♦ 5960 Btu/hr</b>
.028	
··•	0.4 $0.8$ $1.2$ $1.6$ $2.0$ $2.4$ $2.8$ $3.2$ $3.6$ $4.0$
* •	(1/0.0111)¥0.8
	······································

#### DISCUSSION

It has been found that the value of  $h_0$  varies with q for methyl and i-propyl alcohols. No variation was found for n-butyl alcohol. The values of  $h_0$  for methyl alcohol ranged from h85 to 527 with q varying from 7750 to 5750. For i-propyl alcohol,  $h_0$  varied from 221 to 250 with q changing from 4700 to 3800. The value of  $h_0$  for n-butyl alcohol was 208 with q ranging from 7150 to 5960.

The h<sub>e</sub> values were determined from lines drawn through two or three points. Most of the values for methyl alcohol were obtained from three points and most of those for the other alcohols from two points. It is desirable to obtain as many points as possible. However, this is not always practicable. As is evident from Figures h, 7, and 10, the number of points determined is dependent upon the slopes and spacing of the curves. The slopes of the curves are a function of each individual alcohol; the spacing is an experimental variable.

The data of Chu, et. al.; (7) indicate that the slopes of the above mentioned curves become very steep at high cooling water flow rates. Pump head limitations in this work prevented the experimental determination of the steep end of these curves. Investigation at high water rates would have provided additional points for the determination of  $h_{e^*}$ 

Reducing the spacing between curves to obtain more points was

impracticable due to data overlap. This overlap is best demonstrated on Figures  $3_5$   $6_5$  and  $9_2$ .

It would have been desirable to make investigations over wider ranges of q. Limitations of the pressure controller made it impossible to obtain lower values of this parameter.

Since many of the values of  $h_0$  were determined from only two points, it is necessary to discuss the accuracy of these pairs of points. For the most part, these points were obtained from the head of one curve and the tail of another. (Figures 4, 7, and 10.) The head, or high cooling water rate end, was usually well defined by a large number of data points. The tail, or low water pate end, was usually defined by fewer points, tending to make this end of the curve less reliable. Also, as the overall temperature difference decreased, the curves became flatter. This was to be expected since from a theoretical consideration, q will equal zero when the overall temperature difference is zero, no matter what the rate of cooling water flow. A slight change in slope of these flat curves can make a large difference in the intercept selected to calculate  $h_0$ . Therefore, the points obtained from low water rates and/or low overall temperature differences are least reliable.

The water flow rate variation was less than one per cent, except at low flows where it was occasionally as high as three per cent. It is believed, therefore, that the value of q is correct to within  $\pm 3$  per cent. The Beckmann thermometers could be read to within 0.002°C. Since the minimum cooling water temperature rise was about 0.800°C the maximum error involved in reading these thermometers is  $\pm 0.5$  per cent. The average variation of the overall water to vapor temperature difference is  $\pm 1$  per cent. This variation was kept to a minimum by adjusting vapor temperature as water temperature changed. The overall error of the observed condensing film coefficients is estimated to be 5 to 10 per cent.

As shown in a previous section, the relation of  $h_0$  and q is such that a log-leg plot of  $h_0$  and q should give a straight line with a slope of minus one-third. This plot is shown in Figure 12 and the data listed in Table 5. The data on methyl and i-propyl alcohols confirm this theoretical relationship. The slope for methyl alcohol was -0.278 and for i-propyl alcohol was -0.405. The deviation of these slopes from -0.333 is within the accuracy of the graphical techniques involved. No variation of  $h_0$  and q was found for n-butyl alcohol. The reasons for this are unknown at this time.

The value of  $h_0$  at the lowest measured value of q for i-propyl alcohol is inconsistent with the other determinations. As discussed above, this inconsistency is probably a result of the inaccuracies inherent in the determination of  $h_0$  at low values of  $q_0$ .

As discussed in the "Theory", Browley re-derived the Musselt equation, taking into account the effect of the heat capacity of condensate. The observed results of the condensing film coefficients of heat transfer have been compared to those predicted by both the

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Russelt and Bromley equations. These results are presented in Table 5.

The agreement between the observed values of  $h_0$  and those predicted by the Nusselt equation is excellent for methyl and i-propyl alcohols. The ratie of  $h_{obs}/h_{calc}$  ranges from 0.975 to 1.020 for methyl alcohol and from 0.975 to 0.992 for i-propyl alcohol, with a ratie of 0.863 for the one inconsistent data point for i-propyl alcohol. The agreement between the observed and the predicted values of n-butyl alcohol is good, the ratio ranging from 0.879 to 0.934. For the most part, for the alcohols studied, the observed values of  $h_0$  were lower than the theoretical values.

The condensing film coefficients predicted by the Bromley equation are higher than those predicted by the Nusselt equation. Bromley's predictions were 2% higher for methyl alcohol, h to 5 per cent higher for i-propyl alcohol, and 6 to 7 per cent higher for n-butyl alcohol. This equation, although supposedly better than Nusselt's at high pressures, apparently overpredicts coefficients at low and reduced pressures. It can be concluded from these observations that the Nusselt equation satisfactorily predicts condensing film coefficients at low pressures where the effect of the heat capacity is negligible.

The values of  $h_{0}$  were calculated at constant values of q. Since the variations of latent heats were small over the range investigated in this work, the tube loadings and film thicknesses for any one alcohol remained essentially constant for these values of  $h_{0}$ . Also, the acceleration effects of the vapor on the condensate film were minimized by using a large condenser jacket. These are further reasons for the agreement between the observed and theoretical results predicted by the Musselt equation.

This study shows that the ratio of  $h_{obs}/h_{calc}$  decreases as the molecular weight of the alcohol increases. The average of the ratios for the three alcohols in order of increasing molecular weight are 0.997, 0.950, and 0.910. It should be noted that the work of Chu, et. al., (7) indicates a ratio of 0.420 for n-hexyl alcohol.

It has been suggested in the past that sub-cooling of condensate occurs and effects the experimental results. In order to measure the effect of sub-cooling, if any, a thermocouple was installed to measure the condensate temperature. These temperature data are listed in Tables 8, 9, and 10.

In all cases the measured condensate temperatures were about 5 to  $10^{\circ}$ C lower than the saturated vapor temperatures. Since this measured bulk condensate temperature is higher than any average calculated with the equation recommended by Mc Adams (15),  $t_{f} = t_{sv} -0.75 (t_{sv}-t_{s})$ , it is felt that sub-cooling did not effect the experimental results of this work. Furthermore, excess vapors were present in the test condenser at all times.

Another possible source of error is the presence of noncondensable gases in the condensing vapor. The experimental technique, as described in a previous section, precludes this possibility. The fouling of a condenser tube surface effects the condensing film coefficients. Rhodes and Younger (23) found that the apparent thermal resistance of a fouled tube can be 10 to 20 per cent higher than that of a cleaned tube. This may partially account for the fact that the ratios of  $h_{obs}/h_{calc}$  were usually less than unity.

The predicted condensing film coefficients are always affected by the physical property data used. The properties of the alcohols used in this work were obtained from the <u>International Critical</u> <u>Tables</u> (10) and from other, more recent, literature (2) (8) (9) (15) (21). There is considerable inconsistency in these data, especially for thermal conductivity.

TABLE 5         COMPARISON BETWEEN OBSERVED AND CALCULATED HEAT TRANSFER         COEFFICIENTS OF CONDENSING VAPOR									
q	Temp. Diff. Across Cond.	Temp. <u>ho</u> Diff. Observed Theoretica Across Nusselt Bron Cond.		Temp. <u>ho</u> Diff. Observed Theoret Across Nusselt B		ho ho Dbserved Theoretical Nusselt Bromley			
BTU/HR	°F	<u>(a)</u>	(b)	(c)	(a)(b)	<u>(a)/(c)</u>			
		MET	THYL AL	COHOL					
7750 7400 6600 6200 5750	103 96 87 87 80 69	485 511 521 511 520 527	493 502 515 513 525 540	504 512 525 524 535 549	•984 1•020 1•012 •997 •991 •975	•962 •998 •992 •975 •972 •960			
		ISOI	PROPYL	ALCOHOL					
4700 4550 4100 3800	135 114 105 96	236 242 250 221	242 248 252 256	255 260 262 266	•975 •975 •992 •863	•925 •931 •954 •831			
		NORMAI	L BU <b>TYL</b>	ALCOHOL					
7150 7000 6780 6600 6350 6030 5960	192 181 181 172 172 162 156	210 210 211 207 208 208 208 204	225 228 228 229 229 230 232	242 243 243 244 244 244 244 245 246	•934 •921 •926 •904 •908 •904 •904 •879	.868 .864 .868 .848 .852 .849 .829			

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FIGURE 12

#### CONCLUSIONS

The technique of Chu, Flitcraft, and Holeman, for measuring the film coefficients of condensing vapors, has been checked with methyl, i-propyl, and n-butyl alcohols. Chu and associates concluded that

$$h_0 = c q^{-1/3}$$

The work on methyl and i-propyl alcohols presented in this paper substantiate their conclusion. No variation of h<sub>o</sub> and q was found for n-butyl alcohol.

The theoretical condensing film coefficients calculated from the Musselt and Bromley equations were compared with the observed values. The Musselt equation predicted values of h<sub>o</sub> closer to those observed than did the Bromley equation. Excellent agreement of theoretical and observed coefficients was found for methyl and i-propyl alcohols, and good agreement was found for n-butyl alcohol. In all cases the Bromley equation predicted higher values of h<sub>o</sub> than did the Musselt equation. It can be concluded from these observations that the Musselt equation satisfactorily predicts condensing film coefficients at low pressures where the effect of the heat capacity of condensate is negligible.

The close agreement between the theoretical and observed coefficients is attributed to 1) the rigorous theoretical analysis of the Wilson method, as presented by Chu and associates, eliminating variables not previously considered and 2) the use of equipment, designed specifically for this work, which represents an improvement over apparatus used by other investigators.

## RECOMMENDATIONS

The following recommendations are forwarded:

- 1. Continue investigations with the aliphatic alcohols. Determine the relationship, if any, between the ratio of  $h_{obs}/h_{oalc}$  and molecular weight.
- 2. Investigate other homologous series of organic compounds.
- 3. Conduct investigations over a wider range of cooling water flow rates. Initiate studies at positive pressures.

# NOMENCLATURE

•

a, b, c	=	constants
A, Ai, Aav	=	external, inside, and average surface area of a tube perpendicular to the flow of heat, sq. ft.
c <sub>p</sub>	=	heat capacity of condensate, Btu/lb°F.
D <sub>1</sub> , D <sub>0</sub>		inside, outside diameter of tube, ft.
8	=	gravitational constant, $4.17 \times 10^8$ ft./(hr) <sup>2</sup>
ho, hobs, hcalc	-	film coefficient, observed film coefficient and calculated film coefficient of condensate outside of a tube, Btu/(hr) (°F) (sq.ft.)
k <sub>f</sub>	-	thermal conductivity of condensate film, Btu/(hr) (sq.ft.) (°F/ft.)
ׄ	*	thermal conductivity of tube wall, Btu/(hr.) (sq.ft.) (°F/ft.)
q, Q	-	rate of heat transfer, Btu/hr.
R		thermal resistance, $({}^{\circ}F)$ (hr.)/Btu. R. for tube wall, R. for condensing vapor, R. for condensate at infinite rate of flow of water, and R for total resistance ( $\pm 1/U$ )
t	=	temperature, "F or "C. t for water bulk, t <sub>f</sub> for condensate film, t <sub>s</sub> for outside tube surface, t <sub>sv</sub> for saturated vapor.
Δ <b>\$</b>	:	temperature difference across condensate, °F.
<b>AT</b>	=	overall (water bulk to saturated vapor) temperature difference, °F.
Ue	=	overall heat transfer coefficient based on outside tube surface area Btu/ (hr.) (°F) (sq.ft.)
¥	2	average velocity of flow, ft./sec. based on a water density of 62.3 lb./cu. ft. ( $\forall = lb/hr$ . in the calculation procedures).
x	-	thickness of tube wall, ft.
λ		latent heat of vaporization, Btu/lb.

Pr Pr	-	condensate film, vapor density, 1b./cu. ft.
μ <sub>f</sub>	•	absolute viscosity of condensate film, lb./(hr.) (ft.)

APPENDII

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Т	ABLE	6		
ROTAMETER CALIBR	ATION	L - (	DRIGINAL	DATA

POINT NO.	ROTAMETER READING %FMD	TIME MIN.	WEIGHT DELIVERED LBS.	WATER TEMP. °C	WATER Flow Lb/hr
1	9.5	5.78	60.0	10.3	623
2	15.0	2.425	40.0	11.0	990
3	20.0	3.66	80.0	10.5	1311
4	25.0	2.91	80.0	10.0	1650
5	30.0	2.43	80.0	9•9	1975
6	35.0	2.08	80.0	10.0	2305
7	40.0	1.815	80.0	9.0	2645
8	45.0	1.630	80.0	8.5	2945
9	50.0	1.452	80.0	8.4	3300

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#### REFERENCE CORRECTION OF BECKMANN THERMOMETERS

Two Beckmann thermometers were used to measure the cooling water temperature rise. No attempt was made to adjust the mercury columns to the same scale reading. However, both mercury levels were adjusted to the lower ends of the scales at the prevailing cooling water temperatures for maximum utilization of the scales.

Since the mercury columns in the Beckmann thermometers were not preset to the same scale reading, the indicated temperature difference at zero heat duty had to be determined. This was accomplished by pumping water through the condenser tube at a constant rate until the system reached equilibrium. The temperature difference was then read. This reading, or "blank", was algebraically added to the indicated temperature rise at a finite heat duty to obtain the true increase.

It was found that there was a small, but measurable, rise in water temperature due to friction in the condenser tube. This temperature rise increased with increasing water flow rate. A calibration curve was developed to determine the magnitude of this effect. This curve is shown in Figure 14. It is evident that this temperature rise would have a significant effect upon temperature differences measured under test conditions. Therefore, a correction for the friction effect, as shown in Figure 14, was employed.

It is convenient to illustrate the use of the correction factors

with an example. Assume that a "blank" was established for the Beckmann thermometers at a rotameter reading of 50 per cent. At this water rate, a temperature difference of  $0.500^{\circ}$ C was indicated. Under test conditions at the same flow rate, say the temperature difference was  $1.700^{\circ}$ C. The actual temperature difference in the condenser tube would then be 1.700 - 0.500 or  $1.200^{\circ}$ C. As shown by Figure 1h, the increase due to friction alone was  $0.262^{\circ}$ C. Therefore, the true temperature difference resulting from the condensing vapor is  $1.200 - 0.262^{\circ}$ C or  $0.938^{\circ}$ C.

All differential temperature data presented in this work were corrected in the above manner.

TABLE	7
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TEMPERATURE RISE DUE TO FRICTION CONDENSER TUBE BLANK										
R U N NO .	ROTA- Meter RDG	INLET WATER TEMP.	INLET BECK- MANN	OUTLET BECK- Mann	△T BECK- Mann	RELA- TIVE AT				
	¢.	°C	TEMP.	TEMP.	°c	<u>°c</u>				
Cl	50.0	6.20	2.102	0.412	1.690	0.262				
<b>C</b> 2	40.00	6430	2.217	0.615	1.602	0.174				
сз	30.0	6.75	2.597	1.066	1.531	0.103				
С4	19.0	7.40	3.228	1.757	1.471	0.043				
C5	9.0	9.20	5.091	3.659	1.432	0.004				
<b>C</b> 6	14.9	8.00	3.889	2.434	1.455	0.027				

	VARIATION OF	THE RELATIV	E WATER	
		FLOW RATE		
0.36				
0.32				
0.28				
	-			Ø
0.24				
0,20				
0,16		•		
0.12				· · ·
		0		
0,08				
0.04	مر			
0.00	0 <sup>-</sup> 10 20	50	40	50
ROTAN	TER READING, S	FULL METER	differential	

RUN NO.	ROTA- Meter	INLET TEMPE	WATER Rature	OUTLET TEMPER	WATER Ature	VAPOR TEMP.	FILM	TEMP.
	KDG	ABSO- LUTE	BECK- MANN °C	ABSO- Lute MV	BECK- MANN °C	°c	MV	°c
C 1 2 C 3	45.0 39.5 29.7 50.0 50.0	5.60 5.50 5.80 5.20 5.23	2.301 2.153 2.517 2.451 1.954	0.266 0.300 0.195	0.627 2.191 3.146 0.804 1.631	64.29 64.60 64.93	2.42 2.44 2.44	58.8 59.3 58.4
с4567	28.4 20.1 15.0 34.5 24.0	6.20 6.15 6.72 5.50 5.90	2.440 2.860 3.437 2.235 2.615	0.310 0.370 0.240 0.296	0.823 4.398 5.853 2.535 3.754	65.50 64.36 64.59 66.14	2.46 2.44 2.40 2.45	59•8 59•3 58•4 59•5
с с 8 9 с	13.7 31.3 12.7 17.7 45.2	7 • 70 6 • 70 7 • 80 7 • 30 6 • 00	3.468 2.399 3.575 3.032 0.887	0.450 0.384	1.621 0.465 6.222 4.618 1.158	65.25 65.26	2.50 2.48	60.6 60.2
10 C 11 12 13	19.0 37.0 31.0 40.0 50.0	7 • 15 5 • 70 5 • 96 5 • 70 5 • 70	1.926 1.468 1.608 1.400 1.393	0.347 0.258 0.242 0.244	5.118 0.423 2.488 1.795 1.428	55.80 55.81 55.69 55.91	2.07 2.07 2.06 2.07	50.9 50.9 50.7 50.9
C 14 15 16 17	26.3 44.7 34.7 23.5 13.0	6.10 5.60 5.70 6.13 7.25	2.166 1.844 1.885 2.315 3.416	0.222 0.248 0.288 0.392	0.552 1.453 1.919 3.127 2.424	55.86 55.93 56.06 56.98	2.05 2.06 2.08 2.14	50.4 50.7 51.1 52.5
C 18 19 20 21	27.2 34.8 40.4 46.3 28.0	6.50 6.20 6.10 6.04 6.63	2.531 2.302 2.236 2.214 2.715	0.260 0.250 0.246 0.295	0.846 2.107 1.796 1.552 2.918	51.65 51.72 51.83 52.63	1.90 1.92 1.92 1.95	46.9 47.4 47.4 48.1
с 22 23 24 с	40.0 49.3 29.0 25.0 14.2	6.40 6.20 6.46 6.90 7.65	2.310 2.269 2.463 2.885 3.629	0.236 0.272 0.302	0.471 1.496 2.544 3.350 2.031	52.09 52.25 52.43	1.93 1.94 1.95	47•7 47•9 48•3

.

TABLE 8ORIGINAL DATE - METHYL ALCOHOL

RUN ROTA- NO. METER		INLET WATER Temperature		OUTLET TEMPER	OUTLET WATER TEMPERATURE		FILM	TEMP.
	x 20	ABSO- LUTE	BECK- MANN °C	ABSO- Lute MV	BECK- MANN °C	°c	<u> </u>	°c
25 26	19.0 14.1	7.12	3.108 3.657	0.346 0.400	4.140 5.536	52 <b>.7</b> 3 52 <b>.</b> 96	1.98 2.01	48.8 49.5
27 28	50.6 44.2	6.40 6.37	2.422	0•240 0•246	1.664	45.71 45.65	1.62 1.63	40.2 40.4
29 30	39.8 35.7	6.40 6.50	2.365	0.248 0.258	1.936	45•79 45•75	1.65 1.65	40.9 40.9
31 32	30.9 24.8	6.80 7.02	2.750 2.973	0.280 0.300	2.714 3.328	46.36 46.63	1.72 1.74	42.6 43.1
33 34 35 C	20.0 14.2 50.3 50.0	7.43 8.20 6.35 6.50	3.336 4.133 2.360 2.352	0.333 0.400 0.250	4.088 5.663 1.640 0.692	47.28 48.03 46.23	1.77 1.81 1.71	43.8 44.8 42.4
36	50.3	6.50	2.417	0.256	2.061	60.76	2.27	55.4
37 38 39 40 41	45.0 39.3 35.6 29.6 25.2	6.55 6.55 6.60 6.75 6.90	2.432 2.408 2.468 2.579 2.709	0.264 0.270 0.276 0.300 0.320	2.256 2.473 2.624 3.217 3.714	60.76 60.76 60.75 60.80 61.87	2.27 2.27 2.28 2.32	55•4 55•4 55•4 55•7 56•6
42 C	19.8 13.7	7.20	3.038	0.352	4.577 1.621	61.89	2•33	56.8
43 44 45	17.5 14.8 12.9	7.40 7.60 7.80	3.172 3.367 3.540	0.394 0.412 0.440	4.667 5.413 6.025	62.09 62.56 62.69	2.36 2.39 2.40	57.5 58.2 58.4

TABLE 8 (con<sup>1</sup>t) Original date - Methyl Alcohol

	ORIGINAL DATA - ISOPROPYL ALCOHOL								
RUN NO.	ROTA- METER	INLET Tempe	WATER Rature	OUT LET TEMPER	WATER	VAPOR TEMP.	FILM	TEMP,	
	xDG	ABSO- LUTE °C	BECK- MANN °C	ABSO- LUTE MY	BECK- MANN °C	°c	MV	<u>°c</u>	
сс 467 47	37.4 19.0 20.1 30.5 35.9	6.65 7.20 7.15 6.80 6.70	2.445 2.968 2.920 2.567 2.494	0•320 0•280 0•264	0.452 1.077 3.104 2.029 1.682	83.13 82.80 82.77	3.15 3.15 3.12	75•3 75•3 74•7	
c 49 50 51	46.2 23.5 40.4 15.0 17.6	6.80 7.10 6.70 7.35 7.00	2.576 2.803 2.444 3.042 2.703	0.262 0.352 0.342	0.552 0.948 1.501 3.880 3.163	82.30 82.37 82.36	3.10 3.13 3.13	74•2 74•9 74•9	
52 53 53 54 c	10.9 40.1 49.2 41.2 19.4	7•73 6•30 5•97 5•90 6•65	3.456 1.925 1.658 1.591 2.287	0.410 0.235 0.240	5.260 0.080 0.611 0.774 0.576	82.35 82.40 82.40	3.15 3.12 3.10	75•3 74•7 74•2	
55 56 57 58 c	15.1 12.4 10.0 25.8 19.7	6.60 6.90 7.32 5.98 6.17	2.206 2.495 2.956 1.576 1.768	0.316 0.350 0.390 0.250	2.957 3.764 4.850 1.305 0.017	71.03 71.78 72.35 69.24	2.69 2.74 2.75 2.60	65.0 66.1 66.4 62.9	
59 60 61 62 63	30 <b>.1</b> 35.0 40.5 45.5 49.0	5.75 5.60 5.50 5.50 5.60	1.379 1.261 1.232 1.243 1.326	0.230 0.220 0.210 0.210 0.210	0.846 0.510 0.319 0.176 0.170	69.71 69.57 69.70 69.43 69.18	2.60 2.60 2.58 2.58 2.58	62.9 62.9 62.9 62.5 62.5	
64 C 65 66	17.0 40.1 19.0 19.0 14.9	6.52 6.70 6.30 6.70 7.00	2.152 1.925 2.270 2.296 2.636	0.300 0.290 0.340	2.643 0.080 0.543 2.391 3.254	70 <b>.</b> 43 65.29 66.22	2.65 2.45 2.50	64.1 59.5 60.6	
67 68 69 70 71	12.7 10.2 25.0 30.3 35.1	7.40 7.90 6.51 6.30 6.25	3.027 3.576 2.159 1.932 1.869	0.350 0.400 0.270 0.250 0.250	3.902 5.065 1.821 1.301 1.060	67.71 68.71 65.48 64.05 64.62	2.55 2.60 2.45 2.38 2.40	61.8 62.9 59.5 57.9 58.4	

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 TABLE 9

 IGINAL DATA - ISOPROPYL ALCOHOL

	<del></del>							
RUN NO.	ROTA- METER RDG	INLET WATER TEMPERATURE		OUTLET WATER Temperature		VAPOR TEMP.	FILM	TEMP.
		ABSO- LUTE OC	BECK- MANN °C	ABSO- Lute MV	BECK- MANN <sup>O</sup> C	°c	MV	°c
72 73	42.2	6.10 6.00	1.748	0.230 0.220	0.705	64.41 64.74	2.40 2.40	58.4 58.4
74 75	35.0 29.3	5.90 5.90 6.00	1.475	0.230 0.240	2.372	59.09 59.46	2.20 2.20	53•9 53•9
76 77 78 79 c	41.0 49.0 23.4 19.3 18.9	5.80 5.75 6.24 6.50 6.70	1.366 1.363 1.752 1.994 2.251	0.210 0.210 0.250 0.275	2.077 1.867 3.138 3.653 0.495	58.28 57.76 59.98 60.44	2.16 2.15 2.24 2.26	53.0 52.7 54.8 55.2
80 81 82 83	16.3 12.7 7.8 10.2	6.87 7.20 8.40 7.63	2.464 2.823 4.098 3.225	0.300 0.340 0.445 0.375	2.665 3.531 6.117 4.384	61.09 61.51 62.44 61.58	2.31 2.32 2.36 2.33	56.3 56.6 57.5 56.8

TABLE 9 (Con't) ORIGINAL DATE - ISOPROPYL ALCOHOL

	OR	IGINAL	TAB DATA -	LE 10 NORMAL	BUTYL	ALCOHOL		~~
RUN NO.	ROTA- Meter	INLET TEMPE	WATER	OUTLET TEMPER	WATER	VAPOR TEMP.	FILM	TEMP.
	R	ABSO- LUTE	BECK-	ABSO- LUTE NV	BECK- MANN °C	°c	NY_	<u>°c</u>
с 84 85 86 87	21.3 21.3 14.2 25.5 30.1	7.45 7.50 8.10 7.30 7.20	2.884 2.933 3.514 2.761 2.660	0•35 0•40 0•325 0•305	1.219 3.519 5.258 3.031 2.629	95•14 95•86 95•45 94•01	3•47 3•53 3•54 3•53	82.4 83.8 84.0 83.8
88 89 90 c 91	35.6 40.6 48.9 20.0 10.7	7.02 7.00 7.18 7.70 9.00	2.505 2.498 2.691 3.128 4.378	0.29 0.28 0.27 0.48	2.176 1.960 1.857 0.571 6.256	94.01 94.06 93.45 97.25	3•53 3•53 3•52 3•56	83.8 83.8 83.6 84.4
92 93 94 95 0	11.1 20.75 14.6 11.5 40.8	8.90 7.72 8.25 8.88 7.50	4.293 3.107 3.647 4.269 2.927	0.48 0.37 0.43 0.48	6.129 3.031 4.683 6.172 0.211	97.14 99.27 99.53 100.23	3.56 3.55 3.62 3.67	84.4 84.2 85.8 86.9
96 97 98 99 c	49.0 39.3 23.0 31.0 38.0	7 • 50 7 • 58 8 • 04 7 • 80 7 • 90	3.006 3.022 3.438 3.180 3.224	0.30 0.31 0.365 0.33	1.276 1.649 3.092 2.224 0.059	98.65 98.55 98.61 99.10	3•55 3•56 3•65 3•60	84.2 84.4 86.4 85.4
100 101 102 C 103	34.4 29.5 24.2 21.3 21.3	7.96 8.11 8.37 8.80 8.80	3.351 3.506 3.728 4.136 4.149	0.34 0.35 0.395 0.42	1.836 2.267 2.958 1.078 3.699	104.07 104.10 104.40 105.78	3.88 3.88 3.90 4.00	91.6 91.6 92.0 94.0
104 105 106 107 108	14.75 12.7 10.0 48.5 43.0	9.30 9.50 6.48 8.00 8.00	4.638 4.924 1.794 3.488 3.440	0.48 0.51 0.42 0.32 0.32	5.364 6.128 4.123 1.377 1.507	106.60 107.07 103.85 103.65 103.80	4.05 4.11 4.01 3.92 3.90	95.1 96.4 94.2 92.4 92.0
109 110 111 112 113	41.5 48.5 34.5 30.1 24.6	8.00 7.98 8.30 8.35 8.50	3.444 3.444 3.675 3.715 3.868	0•335 0•32 0•355 0•37 0•40	1.644 1.383 2.223 2.481 3.195	108.67 108.38 109.35 109.79 110.45	4.10 4.07 4.13 4.15 4.19	96.2 95.5 96.8 97.2 98.1

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ORIGINAL DATE - NORMAL BUTYL ALCOHOL								
R UN NO.	ROTA- Meter	INLET WATER TEMPERATURE		OUTLET WATER TEMPERATURE		VAPOR TEMP.	FILM TEMP.	
	RDG	ABSO- LUTE	BECK- MANN °C	ABSO- LUTE MV	BECK- MANN °C	°c	MV	°c
114 115 116	20.0 10.2 48.5	8.80 6.67 8.23	4.149 1.965 3.692	0.435 0.45 0.35	4.062 4.440 1.649	110.44 108.60 108.78	4.23 4.25 4.12	99•0 99•4 96•6
117	30.1	8.40 9.30	3.512	0.525	6.213	111.31	4.22	98.8
118 119	17.1 17.0	8.80 8.80 8.40	4.168	0.45 0.45	4.467 4.719	111.07 116.25	4.20 4.48	98.3 104.4
120 121	30.3 24.0	8.40 8.60	3.766 3.951	0.38 0.415	2.780 3.524	116.01 116.14	4.44 4.50	103.5 104.8
122 123 124 C	20.0 13.9 10.2 27.0	8.80 9.30 7.86 8.60	4.137 4.673 3.213	0.45 0.52 0.52	4.252 6.130 5.953 0.840	116.71 117.37 116.79	4.55 4.62 4.58	105.9 107.4 106.5
125	10.8	7.43	2.763	0.45	5.223	116.27	4.43	103.3
126	11.9	8.81	4.151	0.52	6.185	116.54	4.44	103.5

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TABLE 10 (con<sup>t</sup>t) RIGINAL DATE - NORMAL BUTYL ALCOHOL

## SAMPLE CALCULATIONS

I. RUN NO. 2 - METHYL ALCOHOL

1. Rate of Heat Transfer, q

Water flow rate = 1950 lb./hr. Water temperature rise = 2.280°C Heat capacity of water = 1.0 Btu/(lb) (°F) q = 1950 (1.0) (2.280) (1.8) = 8010 Btu/hr.

Inlet water temperature = 7.60°C Water temperature rise = 2.280°C

Bulk temp. (°C) = 5.807  $\frac{2.280}{2}$  = 6.94°C

Bulk temp. (°F) = (6.94) (1.8) + 32 = 44.6°F

3. <u>Overall Water to Vapor Temperature Difference</u>, <u>°C</u> Water bulk temperature = 6.94 °C Saturated vapor temperature = 64.60 °C

Overall temp. difference = 64.60 - 6.94 = 57.66 °C

$$4. \frac{1 \times 10^3}{(1+0.011t)^{0.0}}$$

Water bulk temperature = 14.6 F

Water flow rate, V = 1950 lb./hr.

$$\frac{1 \times 10^3}{(1+0.011t)} v^{0.8} = \frac{1 \times 10^3}{[1+0.011(144.6)](125)} = 1.58$$

## II. METHYL ALCOHOL AT 9 = 7400 Btu/hr

## 1. Observed Heat Transfer Coefficient, h.

From Figure 4 at q = 7400 Btu/hr

AT.ºC	$\frac{1 \times 10^3}{(1+0.0115)v^{0.8}}$	∆ <b>T, <sup>e</sup>f</b>	∆ <b>I/q</b> ( <sup>e</sup> F) (hr)/(Btu)
57.38	2.12	103.3	0.01396
53.25	1.53	95.9	0.01296
48.81	0.92	87.8	0.01186

 $\Delta T/q$  vs  $\frac{1 \times 10^3}{(1+0.011t)}$  v<sup>0.8</sup> is plotted in Figure 5.

at  $\frac{1 \times 10^3}{(1+0.011t)v^{0.8}} = 0$ ,  $\frac{\Delta T}{q} = 0.01025$ 

Tube wall thickness, x = 0.00292 ft. Outside tube surface area = 0.1962 sq.ft. Inside tube surface area = 0.1596 sq.ft. Tube thermal conductivity, k = 60 Btu/(hr) (sq.ft.) ( $^{\circ}F/ft$ )

Average tube surface area,  $A_{av} = \frac{0.1962 - 0.1596}{\ln \frac{0.1962}{0.1596}} = 0.1772$ 

 $\frac{x}{k A_{gv}} = \frac{0.00292}{60(0.1772)} = 0.00027$   $\frac{1}{U_{g}A_{0}} = \frac{\Delta T}{q} = \frac{1}{h_{0}A_{0}} + \frac{x}{k A_{gv}}$   $h_{0} = \frac{1}{0.1962(0.01025 - 0.00027)} = 511 \text{ Btu/(hr) (sq.ft.)(°F)}$ 

2. Theoretical Heat Transfer Coefficient, h.

At infinite water flow rate, water bulk temp = inlet water temp = 7.0°C (Figure h at T = 53.25°C) Saturated vapor temp., t<sub>mw</sub> = 61.5°C Thermal resistance from water to vapor = 0.01025 Thermal resistance of tube wall = 0.00027 Water to saturated vapor temp. diff. = 61.5- 7.0 = 54.5°C Tube surface temp., t = 7.0 + 5h.5 [0100027] = 8.h°c Film temp.,  $t_{r} = t_{ou} - 0.75 (t_{ou} - t_{o})$ - 61.5 -0.75 (61.5- 8.4) - 21.7°C Temp. drop across condensate film = 61.5 -8.4 = 53.1°C At t, = 21.7°C Thermal conductivity, k<sub>r</sub> = 0.1235 Btu/(hr)(sq.ft.)(<sup>9</sup>F/ft) Liquid density,  $\beta_e = h9.2$  lb/cu.ft. Latent heat,  $\lambda = 503$  Btu/lb. Viscosity,  $\mathcal{M} = 1.40 \text{ lb}/(\text{hr})(\text{ft.})$ Heat capacity, ep = 0.601 Btu/(1b) (<sup>6</sup>F) Gravitational constant,  $g = h.17 \times 10^8$  ft/hr<sup>2</sup> Outside tube diameter, D. = 0.03125 ft. Temp. drop across condensate film = 95.7°F

(a) h<sub>0</sub> by Musselt equation  $h_{0} = 0.725 \sqrt{\frac{k_{f}^{3} \beta_{f}^{2} g \lambda}{D_{0} \mathcal{U}_{f} \Delta t}}$ 

;

$$h_{0} = 0.725 \sqrt{\frac{(0.1235)^{3} (h9.2)^{2} (h.17x10^{8}) (503)}{0.03125 (1.40) (95.7)}}$$
  
= 502 Btu/(hr) (sq.ft.) (°F)

(b) h by Broaley Equation

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$$h_{0} = 0.728 \sqrt{\frac{k_{f}^{3} / (P_{f} - P_{f}) g (1 + 0.4 \frac{\Delta 4 CB^{2}}{\lambda})}{D_{0} / (f \Delta 4}}}$$

$$h_{0} = 0.728 \sqrt{\frac{(0.1235)^{3} (49.2)^{2} (4.17 \pm 10^{8}) [1 + 0.4 \frac{(95.7)(0.601)}{503}]^{2}}{0.03125 (1.40) (95.7)}}$$

$$= 512 \quad Btu/(hr) (sq.ft.) (^{0}F)$$

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