THE MAXIMUM AND MINIMUM VALUES OF THE HEAT Q TRANSMITTED FROM METAL TO BOILING WATER UNDER ATMOSPHERIC PRESSURE*

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Abstract — The quantity of heat transmitted from a metal surface to boiling water increases as the temperature difference ΔT is increased, but after the ΔT has reached a certain limit, quantity Q decreases with further increase in ΔT. This turning point is the maximum value of heat transmitted. The existence of this point was actually observed in the experiment. Under atmospheric pressure, ΔT corresponding to the maximum value of heat transfer for water at 100°C falls between 20 degC to 40 degC, and Q is between 1080 000 and 1800 000 kcal/m² h (i.e. between 2000 and 3000 kg m² h, if expressed in constant evaporation rate at 100°C); this figure is larger than the maximum value of heat transfer as was previously considered. Also, the minimum value of heat transfer was obtained and in the Q-ΔT curve for the high temperature region, the burn-out effect is discussed.

INTRODUCTION

The improving of evaporation rate (i.e. the quantity evaporated per unit time per unit area) for an evaporator is important, because it reduces the size of the evaporator of a fixed capacity as well as shortens the time required for the generation of steam. If the evaporation rate, hence the heat-transfer quantity Q, has a maximum value, the determination of this value is certainly necessary if one is to control the evaporation rate at all. This article is mainly a report of the experimental determination of the maximum value of heat transfer, although the related minimum value of heat transfer was also obtained experimentally.

As the boiling phenomenon is so involved and complicated, the analysis for boiling heat transfer is an almost impossible problem. Even with the powerful dimensional analysis, the research to date was not very satisfactory, besides, the reported work available in the literature was all experimental.

Now,

\[ Q = \alpha \Delta T \]  

where Q is the heat transmitted from the metal surface per unit area per unit time to the water; \( \alpha \) is the heat-transfer coefficient, and ΔT is the temperature difference between the surface and the water. Equation (1) has been used for obtaining the relationship between ΔT, \( \alpha \) and Q. Among the researchers on boiling heat transfer are Austin [1], M. Jakob and W. Linke [2].

For the purpose of comparison, the work of previous workers is indicated by a, b, c, in Fig. 1. Austin [1] proposed that \( \alpha \) asymptotically approaches \( \alpha = 7000 \) kcal/m² h degC (0.194 cal/cm² *s degC), and the quantity Q increases with ΔT without limit.

In the process of boiling, the water is fully agitated by the generated steam bubbles, and the degree of agitation first increases with increase in ΔT, but because of the lower heat-transfer rate of steam (\( \alpha \)) of the heat-transfer
rate of water), $\alpha$ is not a monotonically increasing function of $\Delta T$. When the boiling is mild, the agitation by the steam bubbling has more effect on the heat transfer, so $\alpha$ and $Q$ both increase as the $\Delta T$ is increased. Whereas, if the generation of steam becomes too fast, most of the metal surface is covered by the steam bubbles. As a result, there is no more water which is in direct contact with the metal surface to be agitated. Therefore, the negative effect (lowering of $\alpha$) takes place and it becomes a matter of heat transfer between metal surface and steam. Thus, contrary to the conclusion of the previous workers, the value of $\alpha$ at 100°C or 200°C $\Delta T$ is conceivably limited (in the order of 1 kcal/m² h degC). Therefore, in the $\Delta T-\alpha$ curve, the ordinate first increases with increase in $\Delta T$ (Fig. 2) to a critical point, then it must decrease for further increase in $\Delta T$. Since $Q$ is the product of $\alpha$ and $\Delta T$, it should not decrease when $\alpha$ first starts to decrease. Differentiating equation (1) and letting $dQ = 0$, one gets $\alpha \Delta T = -d\alpha/d(\Delta T)$; thus at the point $b$, $Q$ also starts to decrease (Fig. 2), and that is the maximum value of heat transfer, $Q_{\text{max}}$. If $\Delta T$ continues to increase, the radiation from the metal surface becomes pronounced and the values of $\alpha$ and $Q$ can go very high, so that the $\alpha$- and $Q$-curve should be concave upward again. The relational function of $\alpha$ or $Q$ on $\Delta T$ is as presented in Fig. 2, where $c$ is the point for the minimum value of $Q$. Nevertheless, the $bc$ part of the curve (as will be discussed later) is so unstable that it is hard to obtain in practice.
FURTHER DISCUSSION ON THE $Q$-$\Delta T$
AND $\alpha$-$\Delta T$ CURVES (FIG. 2)

Some special equipment might be used to maintain the metal surface at a constant
temperature although, in general, equilibrium temperature is attained when the heat trans-
mitted to the water is equal to the heat supplied by the heat source. Let $Q$ be the heat supplied
and the heat transmitted to water at equilibrium state; the temperature of the metal surface at
this value of $Q$ (point P in Fig. 2) is then the sum of the corresponding $\Delta T$ and the temperature
of water. At this point, if the supplied heat is increased by a small amount, $P$ is moved closer
to $b$ and attains another equilibrium state. Suppose, $P$ reaches $b$ now, and if we supply a
little bit more heat to increase $\Delta T$, then $P$ goes beyond $b$ and $Q$ starts decreasing. Therefore,
the excess heat (the difference between supplied heat and the $Q$ from Fig. 2) accumulates in the
metal material so $\Delta T$ is further increased. From there, $Q$ is becoming less and less as $\Delta T$
continuously increases. $P$ then moves rapidly following the $bcd$ curve, and finally reaches the
equilibrium state at $d$, where it has the same value of $Q$ as at point $b$ (Fig. 2). In the experiment, one
supplies the heat to the heat-transfer area, gradually increasing from zero, and records the
surface temperature; at point $b$, the surface temperature will suddenly change; from this,
one notices that the maximum point has been reached. As will be discussed, the value of
$Q_{\text{max}}$ at point $b$ varies with different curvature

do heat-transfer surfaces, but in general, it is
over 30 cal/cm² s. If we assume that $Q_{\text{max}} =$
30 cal/cm² s and calculate $\alpha$ for the heat transfer
between the metal surface and the generated steam, we find that the temperature at point $d$
is in the order of magnitude of the melting point of platinum. As a consequence, for most metals,
when they go beyond $b$, they follow the curve $bcd$ rapidly and melt before they reach $d$.
The $\Delta T$ corresponding to $b$ for water is below
40°C; accordingly the temperature of the metal surface at point $b$ is lower than 140°C. Ordinary
metals are solid at this temperature. As will be discussed later, the experiments were run with
metal wires of iron, nickel, nichrome and platinum. These wires actually melt at point $b$
just like the fuse of low melting point (170°C) does. Only when using an extremely thin
platinum wire for the heat-transfer surface would the point $d$ really exist, because the wire
has a small diameter, hence a small value of $Q_{\text{max}}$. Now, if we reduce the value of $Q$ little by
little, the curve would not reverse from $d$ to $b$
but it goes along $dc$ (as a result of decreasing
$\Delta T$), and suddenly jumps to point $e$ from point
c, for the same reason as for the jump from $b$ to
d. We call the region from $d$ to $c$ the spheroidal
state.

The purpose of this investigation is to study experimentally the above-mentioned phenomenon. But, owing to the large value of $Q_{\text{max}}$, we have to select a suitable type of heat-transfer
surface. We were not able to supply necessary heat for a flat surface which was used by
Austin [1] and by Jakob [2]. When we used a
thin metal plate as the transfer surface, contacting water on one side of the plate and heating
directly with a Bunsen lamp from the other
side in order to supply the heat, we could supply only a few per cent of the $Q_{\text{max}}$. For
various reasons, we have used the following
two methods: (a) A thin metal wire for the heating element as well as the heat-transfer
surface; the wire is set in the water and supplied with the necessary heat by running electric
current through the wire. The experiments
were run with different metals of various diameters. Subcooled water was also used in obtaining $b$. The minimum value of heat transfer was obtained with a thin platinum wire. (b) With a special equipment set-up, the same experiments were repeated on two circular flat plates of 10-mm and 6·48-mm diameters.

EXPERIMENT I—THE $Q$, $x$ AND $AT$ RELATIONS FOR THE SURFACE OF METAL WIRES

In Fig. 3(a), $D$ is the Pyrex glass vessel of 15 cm $\times$ 15 cm $\times$ 24 cm for distilled water. $D$ was set up in a sand bath and was heated with gas or with insulated electric resistance wires in the sand to keep the water at boiling temperature under atmospheric pressure. $ab$ is a metal test wire of 20-cm length; its ends are welded to $aA$ and $bB$ respectively. These silver or nickel plated copper wires are 3-mm in diameter. $A$, $B$ are connected to copper wires of larger diameter which carry the electric current to supply the necessary heat to $ab$. The current going through the wires is about 10 A d.c.; so if $aA$ and $bB$ were bare, or if a metal vessel was used for $D$, the purity of water inside the vessel might have been affected because of the electrolysis. In order to prevent it, $aA$ and $bB$ are cased by glass coating. Alternating current had been tried with the consideration that it might eliminate the electrolysis, but it was not suitable due to the difficulties encountered in the measurement of temperatures, current, voltage, etc.

The electric flow scheme is shown in Fig. 3(b). $CDEF$ forms the Wheatstone bridge, and the test wire $ab$ is fixed in between $CD$. The wires other than for $ab$ are rather large in diameter so that only a few ohms from the resistance in $ab$ were counted for the resistance between section $CD$. $R_1$, $R_2$, $R_3$, are manganin wires. The resistance of $R_2$ and $R_3$ is many hundred ohms, so that most of the current from the source goes to $CDE$, and less than 1 per cent of the total will go through $R_2$ and $R_3$. $R_1$ and $R_2$, are put in an oil bath to avoid the temperature variation due to the heat generation from the

Fig. 3. Equipment for metal wire experiments.
resistors. $R_1$ carries the same amount of current which flows through $ab$. In Fig. 3(c), we have shown that $R_1$ is a rheostat which was constructed with some 40 thin manganin wires stretched between a copper frame and immersed in an oil bath. The resistance is adjusted to the same magnitude as the resistance of the test wire $ab$ during the experiment. Because of the strong current which is flowing, the temperature of oil during an experiment may rise more than 20 degC, but since the manganin wires were used, the variation in $R_1$ can be assumed negligible in this experiment. $S$ is a 30-Ω variac; $G$ is a galvanometer; $N$ is a set of battery cells; $H$ is a d.c. generator; $V$ is a voltmeter; and $I$ is an ammeter.

**THE EXPERIMENTS AND THE RESULTS OF THE DETERMINATION OF THE MAXIMUM VALUE OF $Q$**

Some calibration should be provided before the temperature of $ab$ can be read from the Wheatstone bridge during the experiment; Immerse $ab$ in the oil bath, and vary the oil temperature between 50°C and 150°C (this temperature was measured by mercury thermometer to 1/5 degC precision), take the reading of variac $S$ and plot it against oil temperature when the weak current supplied by $N$ is at equilibrium with the reading in $G$. Since the current used in this calibration is so weak that the temperature of the wire can be taken as the temperature of the oil bath, the calibration curve can be approximated as the reading of $S$ vs. the temperature of the wire. When performing the experiment, the temperature of the test wire $ab$ is therefore obtained from the reading together with the calibration curve.

The calibration with oil may destroy the surface of the metal wires, thus introducing a bad effect in the main experiment. To avoid this, we have run only one point each for a certain kind of metal wire for such calibration with oil, then checked the same calibration with water instead at the temperature from 50°C to 100°C; for temperatures above 100°C, we can extrapolate the result of the calibration.

$ab$ is fixed in the container $D$ (Fig. 3(a)), and after the water in $D$ is heated by Bunsen flame to the required temperature (100°C), a current is supplied by $H$ (Fig. 3(b)), and by adjusting $R_4$. Beginning with 1 A, the current is gradually increased by 0.5 to 1 A, and the corresponding $I$, $V$, $S$, readings are recorded. When $Q$ is close to the maximum value, the variac $S$ is used in addition to $R_4$ (Fig. 3(b)) to keep $G$ at a equilibrium position. When $Q_{\text{max}}$ is reached, part of the wire $ab$ melts out and $G$ will sense a violent shock. At this moment, the readings of $I$, $V$, $S$, are carefully taken, and from this, $\Delta T$ corresponding to the $Q_{\text{max}}$ is obtained. But, to prevent the burn-out of the metal wire, an automatic shut-off device has been applied. During experiments, the electrolysis occasionally introduced some error in the temperature readings. Therefore, in order to shorten the period of one run, the readings in the lower current region were roughly taken. Let $d$ be the diameter of the metal wire in cm, $L$ be the length in cm, the reading of the ammeter and the voltmeter be $I$ and $V$ respectively, the value of $Q$ is then calculated by the following equation:

$$Q = \frac{1}{\pi d L} \cdot \frac{IV}{4184} \text{ cal cm}^2 \text{ s.} \quad (2)$$

The temperature difference between water and the surface of the metal wire $\Delta T$ was obtained from the reading of $S$ and the mercury thermometer; however, there are two points which must not be overlooked:

1. If there were any difference between the temperature of the surface of wire $T_s$ and that of the center of the metal wire $T_0$, it would introduce an error in the value of $\Delta T$. To correct it, we assume that the heat generation from the metal wire due to the current is uniform throughout the wire, and $(T_0 - T_s)$ can be calculated by the following equation (as given in any general heat-transfer book [3]):

$$T_0 - T_s = Wr^2/4\lambda \quad (3)$$
where \( W \) is the quantity of heat generated by the metal wire per unit time per unit volume of the wire, \( r \) is the radius and \( \lambda \) is the thermal conductivity of this metal wire. Since the heat transmitted from the surface of a unit length of wire to the water is \( 2\pi r Q \), which must be equal to the heat generated from the corresponding surface, \( \pi r^2 W \), then, from equation (3):

\[
T_0 - T_r = \frac{Q r}{2\lambda}.
\] (4)

This \( (T_0 - T_r) \) is within 1 degC for platinum and nickel wire, and is less than 2–3 degC for nichrome wire. If we would make more precise correction, we should also consider the variation of the electric resistance due to the temperature difference of the surface and the interior of the metal wire; i.e., the surface of the wire may carry more current because of its lower temperature and its lower resistance. With this consideration, \( (T_0 - T_r) \) could be obtained by solving a simply formulated Bessel function. When it is done, the value of \( (T_0 - T_r) \) is actually less than the one obtained from equation (4), and is within \( \frac{1}{2} \) degC. In the present experiments, we can never neglect the correction made with equation (4), because of the imperfect reproducibility of the experimental results and of item (2) to be discussed. Although the correction can be easily done, it may not positively assure the improvement of the final result. This will be better explained in item (2).

(2) The surface temperature of the metal wire is by no means uniform. This is due to the non-uniform, local generation of steam from the surface. When the vaporization is mild, the bubbles are seen first from the surface where there are scratches or impurities on the surface, then gradually the same bubbling is observed from other locations of the surface after a few seconds. Once the vaporization gets vigorous, most of the surface is covered with steam bubbles, but still there is some part of the surface which is in direct contact with water instead of with steam bubbles. Therefore, the surface temperature will never be everywhere the same, and the surface temperature we recorded during the experiment is therefore only an average value. The degree of non-uniformity of this surface temperature is not yet certain; however, it may be in the order of magnitude of \( \Delta T \), if we consider the temperature for the surface where it is in direct contact with the water to be 100°C. But, actually the temperature of this part of surface is higher than the boiling point of the surrounding water, and therefore the degree of temperature non-uniformity of the metal surface is less than \( \Delta T \) [4]. In view of all these considerations, the few degree centigrade non-uniformity in temperature is not very surprising. Inaccurate measurements may also introduce this amount of error.

The experiments were run with platinum, nickel, nichrome, iron and fuse wires. The results with iron and fuse wire were similar to those with others, except that their surfaces were more seriously affected by electrolysis. Thus, we have omitted the data for iron and fuse. The experimental results were not quite reproducible, therefore, the average value from several runs of the same size and same kind of metal has been taken. The dimension of metals

![Fig. 4. Platinum wire, d = 0.14 mm: water temperature = 100°C.](image-url)
and the number of runs is summarized in Table 1.

Table 2 and Table 3 are lists of some data for illustration. Table 2 is for Run No. 6 with nickel wire of \( d = 0.14 \text{ mm} \) and Table 3 is for Run No. 3 with nichrome wire of \( d = 0.575 \text{ mm} \). The experimental data and the calculated \( Q \) and \( \alpha \) are listed in these tables. The numbers in the last lines of Table 2 and Table 3 are the values for the moment when the metal wires burned out, but in the columns for \( \alpha \), the maximum values appeared in the bold letter before the last line in Tables 2 and 3. Figures 4-9 are the plots of \( Q \) vs. \( T \) (the temperature of the metal wires) from the experimental data for various metal wires. The curves have been drawn in such a way that a straight line is connected between every two adjacent experimental points. The temperature for water has been assumed at 100°C for all cases, and the last point in every graph represents the point when the metal wire burned out. The results as shown are rather scattered among each individual set of data; however, a representative curve is drawn in Fig. 10 for each metal wire. Figure 11 is the \( \alpha \) vs. \( T \) curve.

### Table 1

<table>
<thead>
<tr>
<th>Metal</th>
<th>( d ) (mm)</th>
<th>( L ) (mm)</th>
<th>No. of Runs</th>
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<tr>
<td>Platinum wire</td>
<td>0.14</td>
<td>200</td>
<td>8</td>
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<tr>
<td>Nickel wire</td>
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<td>200</td>
<td>10</td>
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<tr>
<td>Nickel wire</td>
<td>0.40</td>
<td>200</td>
<td>20</td>
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<tr>
<td>Nichrome wire</td>
<td>0.535</td>
<td>200</td>
<td>7</td>
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<tr>
<td>Nichrome wire</td>
<td>0.575</td>
<td>200</td>
<td>9</td>
</tr>
</tbody>
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### Table 2. Experimental results for Run No. 6, nickel wire of \( d = 0.14 \text{ mm} \)

<table>
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<tr>
<th>A</th>
<th>V</th>
<th>W</th>
<th>cal</th>
<th>( Q ) (cal/cm(^2) s)</th>
<th>( \Delta T ) (cal/cm(^2) s degC)</th>
<th>( \alpha ) (cal/cm(^2) s degC)</th>
</tr>
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<tbody>
<tr>
<td>1</td>
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<td>0.3</td>
<td>0.0715</td>
<td>0.0284</td>
<td>0.08</td>
<td>0.0355</td>
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<td>4</td>
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<td>0.303</td>
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<td>0.0645</td>
</tr>
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<td>1.6</td>
<td>12.8</td>
<td>3.05</td>
<td>1.214</td>
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<td>1.896</td>
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<tr>
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<td>11.07</td>
<td>4.41</td>
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<td>20</td>
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<td>82.0</td>
<td>19.52</td>
<td>7.775</td>
<td>14.3</td>
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<tr>
<td>25</td>
<td>5.2</td>
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<td>39.96</td>
<td>12.35</td>
<td>17.0</td>
<td>0.726</td>
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<tr>
<td>30</td>
<td>6.1</td>
<td>183</td>
<td>43.57</td>
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<td>7.9</td>
<td>300</td>
<td>71.5</td>
<td>28.46</td>
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<td>0.949</td>
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<td>39.4</td>
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<td>323</td>
<td>76.9</td>
<td>30.6</td>
<td>32.5</td>
<td>0.942</td>
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### Table 3. Experimental results for Run No. 3, nichrome wire of \( d = 0.575 \text{ mm} \)

<table>
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<tr>
<th>A</th>
<th>V</th>
<th>W</th>
<th>cal</th>
<th>( Q ) (cal/cm(^2) s)</th>
<th>( \Delta T ) (cal/cm(^2) s degC)</th>
<th>( \alpha ) (cal/cm(^2) s degC)</th>
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<tbody>
<tr>
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<td>0.8</td>
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<td>0.0527</td>
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<td>1.383</td>
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<td>0.173</td>
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<td>10</td>
<td>8.25</td>
<td>82.5</td>
<td>19.6</td>
<td>5.44</td>
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<td>12.8</td>
<td>192</td>
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<td>18.8</td>
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<tr>
<td>20</td>
<td>17.0</td>
<td>340</td>
<td>81.0</td>
<td>22.42</td>
<td>25.7</td>
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<td>22</td>
<td>18.7</td>
<td>411.4</td>
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<td>24</td>
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<td>25</td>
<td>21.4</td>
<td>535</td>
<td>127.4</td>
<td>35.29</td>
<td>38.0</td>
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<tr>
<td>26</td>
<td>22.3</td>
<td>580</td>
<td>138</td>
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<td>26.7</td>
<td>23.0</td>
<td>614</td>
<td>146</td>
<td>40.48</td>
<td>46.5</td>
<td>0.860</td>
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</table>
where $\alpha$ is calculated from experimental data. The $\alpha$-curves for metals wire of $d = 0.14$ mm were omitted, because of inconsistent results.

**INTERPRETATION OF RESULTS**

As has been previously mentioned, different positions on the surface of a metal wire may have different $ab$ curves (Fig. 2), because the temperature varies from position to position on the surface; however, when any of these points reaches $b$, the wire burns out. Experimentally, sometimes the wire burned out before it reached point $b$. Suppose on the average, the wire will burn out at $b_1$; then the difference between $b$ and $b_1$ will depend on the diameter of the wire—the smaller the diameter of the wire the smaller will be the difference, so the value $Q_{\text{max}}$ and the corresponding $\Delta T$ are smaller for wire of smaller diameter. In Fig. 10, the burn-out point observed for $d = 0.14$ mm is at the point where the $Q$-curve begins to show a change of slope, though the burn-out points as experimentally detected lie below the $Q_{\text{max}}$.
the calculated in most cases is beyond the maximum point in the curves (Fig. 11). The result varies somewhat with the material used. However, if we take into account the possible experimental error involved, we conclude that the result may be independent of the material used.

Figure 12 is the result of the experiments with nickel wire of \(d = 0.19\) mm and with water temperature at 70°C, 80°C, and 90°C respectively. \(Q_{\text{max}}\) is rather great when the temperature of water is low. The experiments with water at these temperatures were more difficult to control, and there might have been introduced an error of about 10 degC in the value of \(\Delta T\). Figure 13 presents the values for \(x\) as obtained from Fig. 12.

**The experiments and the results of the determination of the minimum value of \(Q\)**

The equipment is the same as the previous one; however, in order to lower the temperature at \(d\) (Fig. 2), the platinum wire of 0.14-mm diameter was used because it has a smaller value of \(Q_{\text{max}}\) and is more durable at higher temperature. When the state moves from \(b\) to \(d\) in Fig. 2, the electric resistance of the platinum wire \(R\) increases rapidly due to the rise of the temperature. When the resistance \(R_4\) in the circuit \(CDER_4H\) (Fig. 3) is greater than \(R\), the
current \( I \) is not lowered much by the increase of \( R \); hence, the quantity of heat \( I^2R \) is very much increased. In this case, point \( d \) of Fig. 2 is at a position higher than point \( b \) and at a temperature so high that even the platinum wire is melted. However, if we reduce the value of \( R_4 \), the difference between the \( Q \) at \( d \) and the \( Q \) at \( b \) is lessened. For this reason, we have used a smaller value of \( R_4 \) in order to prevent the burn-out of the platinum wire. (Recall that we have used large values of \( R_4 \) in the determination of \( Q_{\text{max}} \). The change to smaller \( R_4 \) is to obtain a more adjustable range of the current.) Even so, the point tends to jump from lower position to a higher position as shown by the arrow in Fig. 14. Experimentally, the minimum value of

**Fig. 10.** Average value of \( Q \) for various metal wires; water temperature = 100°C.

**Fig. 11.** Average value of \( \sigma \) for various metal wires; water temperature = 100°C.
FIG. 12. $Q$ curve for water temperature below 100°C.

FIG. 13. $x$ curve for water temperature below 100°C; platinum wire, $d = 0.19$ mm.

FIG. 14. The determination of $Q_{\text{min}}$ for the platinum wire of $d = 0.14$ mm.
$Q$ was determined by reducing the current from the high temperature regime beyond the point b. In the vicinity of the minimum value of $Q$, the platinum wire turns from shining color to red or dark-red spot, and when it passes the point $Q_{\text{min}}$, the temperature rapidly decreases and the galvanometer fluctuates, the brightness of the wire disappears completely and the wire returns to the initial state. Experiments were run with 12 platinum wires and the results are shown in Fig. 14. The resistance vs. temperature curves for platinum wire at higher temperature were extrapolated from the data obtained for the calibration with oil at temperatures below 200°C. The resistance vs. temperature curve for a platinum wire changes every time after the wire is used once. The reason for this may be due to the annealing effect or the expansion of the platinum wire under tension during the strongly heating process, or may be due to the effect of electrolysis. It is not yet completely explored. We presume that all these have caused some 10 deg C of error in $\Delta T$, especially at higher temperatures.

During the tempering process, the metal is changed from high temperature along the $Q$ vs. $\Delta T$ curve. When it reaches $Q_{\text{min}}$, it instantaneously passes $Q_{\text{max}}$ and liberates a large amount of steam. The wire finally becomes black. The $Q$ vs. $\Delta T$ curves can be used to determine the temperature change of the material. Therefore, if we study the case of water in motion or use other liquids instead of water to obtain the similar $Q$ vs. $\Delta T$ curves for each case so as to investigate the mechanism of tempering of metals, we may be able to get some very interesting results.

**EXPERIMENT WITH A FLAT PLATE AS HEATING ELEMENT**

A horizontal flat plate with the water boiling on the top surface was used. The $Q$ vs. $\Delta T$ curve must vary with the size of the flat surface used. Facing the difficulties of being not able to supply sufficient heat to the large flat surface, we had to limit ourselves to two circular surfaces of small diameter, 10 mm and 6.58 mm, in order to obtain the $Q_{\text{max}}$ for the flat plate. Figure 15 is the apparatus set-up: A is a copper solid cylinder with a smaller neck in B, where ab is the heat-transfer surface. C is a cap made of ebonite or of porcelain in the inside and carbonized corks in the outside parts. They were used as the heat insulators. But, if one uses the porcelain and carbonized corks, care must be taken to avoid absorbing water. The resistance wire D was wound on the electrically insulated surface of A and the whole system was insulated with asbestos. The heat generated in D is transmitted to the good conductor A and then to the surface ab by means of B. In this way, we have been able to supply the necessary heat to the surface ab. There might be some heat loss through C, but it is very small compared to the magnitude of the heat transmitted to the water through ab, because the conductivity of C is only $\frac{1}{1000}$ of the conductivity of B. This fact was experimentally tested by attaching thermocouples (copper–constantan) to two positions of B, and to a point in the water close to the surface ab. The axial temperature
distribution as shown in Fig. 16 is a straight line. Thus, the radial heat loss from $ab$ to the water was calculated from $T_1$ and $T_2$ (Figs. 15 and 16) and the thermal conductivity of copper. The thermal conductivity of copper was taken as 320 kcal/m h degC = 0.89 cal/cm s degC.

The copper element $AB$ can hold 10 cal of heat, so that we have to reduce the frequency of the current variation. But, it was found the frequency of current variation did not affect the experimental results very much. The temperature of $ab$ was found by extrapolating the line between $x_1$ and $x_2$ (see Fig. 16).

The heating current, $I$, was recorded in addition to $T_1$ and $T_2$. When $Q$ reaches the maximum value, $T_1$ and $T_2$ change very rapidly, and with our measuring devices, we could only record $T_1$ and $I$. $T_2$ is then obtained from $T_1$, $I$, and Fig. 16. With the $T_1$ and $T_2$ we can calculate the value of $Q_{\text{max}}$ and the corresponding $\Delta T$. The results are shown in Fig. 1, where for a smaller heating surface, we have a larger value of $Q$ at a certain $\Delta T$ reading, such that $Q_{\text{max}}$ is also greater for smaller heating surface. To go a step further, we have designed a device to study the effect of the flow pattern on the $Q_{\text{max}}$. The 10-mm circular surface, with a hollow, ebonite cylinder of the same diameter as the copper surface (top of Fig. 1) was used to control the flow of water adjacent to the copper surface. Notice that there is an adjustable thin plate which divides the ebonite cylinder into two parts. The experiments were repeated with various $S$, which is the distance between the dividing plate $N$ and the surface $ab$. When inserting the dividing plate $N$, the steam bubbles arise along the sides of $N$ and the circulation of water is laminar even with vigorous boiling. Experiment for each value of $S$ was repeated 4–5 times and the average results are presented in Fig. 1. Lifting the distance $S$ 1 mm each time speeds up the flow of water, so that $Q$ has a greater value at a certain $\Delta T$, and so $Q_{\text{max}}$ is also greater. But, when $S$ becomes infinity, i.e., without the dividing plate $N$, the flow becomes very turbulent and the $Q_{\text{max}}$ becomes small again. Figure 17 is the $\alpha$-curve as calculated from the above results, which is similar to Fig. 11 for the case of metal wires.
CONCLUSIONS AND SOME AUXILIARY OBSERVATIONS IN THE EXPERIMENTS

There exists a maximum value of $Q$ for the $Q$ vs. $\Delta T$ curve, which is independent on the material of the metal heating surface. The values of $Q_{\text{max}}$ are rather great compared to the values of $\Delta T$, which are about 20–40 degC as has been determined experimentally. The $Q_{\text{max}}$ is about 30 cal/cm$^2$·s to 50 cal/cm$^2$·s for all dimensions of different heating surfaces, except for the tiny metal wire, provided there is no disturbance to the natural convection of the water. This value for $Q_{\text{max}}$ is far greater than the value which was conventionally accepted. In ordinary evaporators, the value for $Q$ is less than 3 cal/cm$^2$·s even at the position where it is closest to the furnace. Therefore, under normal conditions, the heating surface will never reach the spheroidal state, except possibly when there is scaling on the surface of the evaporator. The problem of scale and the cases for local pressure difference will be treated in the later study.

In the experiments with metal wires some sparks along the surface of the wire were observed. As has been mentioned previously, when $Q$ is small, the bubbling is mild [1], but when $\Delta T$ is about 12 degC, no matter if the temperature of water is 100°C or lower, the generation of steam bubbles becomes suddenly violent, so that all the heating surfaces are covered, and $Q$ abruptly increases at this temperature in the $Q$ vs. $\Delta T$ curve.

What is the temperature of water in contact with the surface, when $\Delta T$ is 20 degC or 30 degC? Granting that there should not be a sudden change on the interface of solid and liquid, we conclude that the water is being superheated. Partridge [5] has experimentally studied the mechanism of steam generation from the dry spot, $mm$ on a surface (Fig. 18). The temperature at this spot on the metal surface is much higher than that of water; therefore, we should say that the $\Delta T$ is actually the average of the temperature of a few higher temperature spots and most of the 100°C parts.

Of course, superheating is required before the pressure can overcome the surface tension of water to generate steam bubbles in the water. The present author has been attempting for years to measure the degree of superheating, but has unfortunately failed. In 1932, Jakob [4] first succeeded in the measurement and he confirmed that it is superheated somewhere near the heating surface. But, this measurement was at the position about 1 mm from the surface, and only 1 degC superheating was reported. It remains as an area for someone to measure the degree of superheating at a location closer to the surface.

![Fig. 18](image)

In the present experiments, the temperature of water should be able to keep at 100°C with heating on the outside of the water container and with the inside heating element. But, as $Q$ becomes large, the water temperature sometimes dropped 0.2–0.5 degC; and the experiment was forced to pause (to find the possible cause of discrepancy). The author thinks that it might be due to the steam bubbles which encourage the vaporization and subcooling of the adjacent water and there is a heat flow from the bulk water to the water adjacent to the steam bubbles; therefore, the temperature of the bulk water dropped.

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Résumé—La quantité de chaleur transmise à partir d'une surface métallique à de l'eau en ébullition croît lorsque la différence de température $\Delta T$ augmente; mais après que $\Delta T$ ait atteint une certaine limite, la quantité $Q$ décroît lorsque $\Delta T$ continue à augmenter. Ce point critique est la valeur maximale de chaleur transmise. L'existence de ce point a été observée actuellement dans l'expérience. Sous la pression atmosphérique, le $\Delta T$ correspondant à la valeur maximale du transport de chaleur pour l'eau à 100°C décroit entre 20° et 40°C, et $Q$ se trouve entre 1 080 000 et 1 800 000 kcal/m² h (c'est-à-dire entre 2 000 et 3 000 kg/m² h, si on l'exprime en vitesse d'évaporation constante à 100°C); ce chiffre est plus grand que la valeur maximale du transport de chaleur qui avait été estimée auparavant. La valeur minimale du transport de chaleur a été aussi obtenue, et l'on discute l'effet de coup de feu sur la courbe $Q-\Delta T$ dans la région des températures élevées.

Zusammenfassung—Die Wärmemenge, die von einer Metalleoberfläche an siedendes Wasser übertragen wird, nimmt mit wachsender Temperaturdifferenz $\Delta T$ zu; hat dieses $\Delta T$ eine bestimmte Grenze erreicht, so nimmt die Wärmemenge $Q$ ab bei weiterhin ansteigendem $\Delta T$. Dieser Umschlagpunkt ist der Maximalwert der übertragbaren Wärme. Das Vorhandensein dieses Punktes wurde tatsächlich experimentell beobachtet. Bei Atmosphärendruck liegt das der maximalen Wärmestromdichte für Wasser von 100°C entsprechende $\Delta T$ zwischen 20 und 40 Grad und zwischen 1 080 000 und 1 800 000 kcal/m² h, d.h. zwischen 2 000 und 3 000 kg/m² h, wenn es als konstante Dampfmassenstromdichte bei 100°C ausgedrückt wird. Diese Beträge sind größer als die früher angenommene maximale Wärmestromdichte. Die minimale Wärmestromdichte wurde ebenfalls bestimmt und in der $Q-\Delta T$ Kurve der Burnout-Effekt für den Bereich hoher Temperaturen diskutiert.

Аннотация—Количество тепла, перенесенного от металлической поверхности к нипящей воде, увеличивается с увеличением разности температур $\Delta T$, но как только $\Delta T$ достигает определенного предела, количество тепла $Q$ при дальнейшем увеличении $\Delta T$ уменьшается. Экстремальной точкой является максимальное значение перенесенного тепла. Существование такой точки наблюдалось экспериментально. При атмосферном давлении $\Delta T$, соответствующей максимальному значению коэффициента теплообмена для воды при 100°C, составляет 20°C-40°C, и $Q = 1 080 000-1 800 000$ ккал/Мч (т.е. 2 000-3 000 ккал/Мч для постоянной скорости испарения при 100°C). Это значение больше максимального значения коэффициента теплообмена, которое рассматривалось раньше. Получено также минимальное значение коэффициента теплообмена. Рассмотрено влияние загрязнений при больших температурах на вид кривой $Q-\Delta T$. 