THE EFFECT OF HISTAMINE ON SKIN AND DEEP TEMPERATURES IN MAN WITH PARTICULAR REFERENCE TO LIVER TEMPERATURE

By W. GRAF

(From the Second Medical Service of St. Erik's Hospital, Stockholm, Sweden)

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Among the numerous investigations on the effects of histamine on regional temperatures and heat regulation the following reports deserve mention as they deal with problems related to the present investigation:

Dale and Laidlaw (1) reported that a large dose of histamine lowered the body temperature of dogs and guinea-pigs. Harmer and Harris (2) administered a moderate dose of histamine to humans, resulting in an increase of the skin temperature and a rise of rectal temperature. These authors suggested that the opposite effect on the rectal temperature would have seemed more likely. Thiesen and Snell (3), in a series of peptic ulcer cases, found that subsequent to a histamine dose gastric temperature fell whereas oral and rectal temperatures rose. Deutsch, Spitzy and Wohlrab (4) stated that after a dose of histamine the gastric temperature fell to the same extent in achlorhydric cases as in normals. The conclusion was drawn that gastric temperature and secretion are not related to each other. Henning, Demling and Kinzelmeier (5) considered measurement of the gastric temperature a possible method of studying, indirectly, blood flow to the stomach mucosa. They also investigated the reactions of the gastric temperature to certain stimuli (6, 7). The lowering of gastric temperature induced with histamine was interpreted as being due to hyperemia with slow blood flow. Spang, Obrecht and Ey (8) obtained the corresponding effect of histamine on gastric temperature but did not consider it justifiable to look upon the gastric temperature as representative for blood flow fluctuations. They also stated that the temperature changes in the stomach do not differ from the pattern of other inner organs. Masuda, Ohara and Katsura (9, 10), in a series of experiments on the temperature of the gastrointestinal tract, concluded that gastric temperature and its changes after a histamine dose are related to the blood flow but not to the gastric secretory activity. Benjamin, Wagner, Zeit, Pisciotta, and Ausman (11), in a study of gastric temperature, claimed that in achlorhydria "intragastric temperature was a straight line tracing." These data are entirely contradictory to the findings of other authors (4). Rossi-Espagnet and Torlontano (12) postulated that the decrease in stomach temperature induced with histamine resulted from cooling of the blood circulating in the dilated skin vessels but offered no direct evidence in support of this opinion. Various authors working with animal experiments have previously expressed this view (13, 14).

METHODS

The procedure applied was described in a previous paper (15). Thermocouple units of copper/constantan or chrome-nickel/constantan were used as measuring units with a common reference junction, and the apparatus was built to compensate for variations in room temperature. The constantan wires measured 0.15 mm. in diameter; the size of the copper wires was 0.08 mm. The size of the junction was 0.5 × 0.3 mm. Junctions and wires were insulated in a nylon tube, the outer diameter of which was 1.55 mm. It was repeatedly checked by means of ice and a heated medium, respectively, so that heat conduction along the thermocouple wires did not distort the results. The measuring units were introduced into the stomach (pyloric region), in the rectal ampulla (immediately above the sphincter), and in the liver. The thermocouple intended for the liver was sterilized for 18 to 24 hours in a 2 per cent solution of benzethonium chloride (NNR) and introduced through a Vim-Silverman-Boecker needle immediately following liver biopsy. The position of the thermocouple in the liver was in several instances controlled by X-ray in two planes. Skin temperature was measured by a thermocouple lightly applied to the center of manubrium. In some cases skin temperature was also measured at the anterior surface of the foreleg. In a few instances the temperature was measured in the right or left hepatic vein, the thermocouple being introduced through a cardiac catheter under fluoroscopic control.

The 40 male patients examined in the present investigation were all admitted to the hospital with alleged or
previously confirmed liver disease. The livers were diagnosed on biopsy as being normal, fatty or cirrhotic but no characteristic differences in temperature pattern between normal and diseased livers have thus far been recognized.

The temperature-measuring equipment was repeatedly checked against a certified mercury thermometer in a water bath of constant temperature, adjusted to different levels between 36 and 38°C. The accuracy of the measuring procedure was found to be ± 0.05°C. As long as the thermocouples mutually yielded entirely uniform figures, small divergences (0.05 to 0.1°C) from the mercury standard were allowed. The room temperature was constant within 0.5°C during most of the experiments, and within 1.0°C in a few. It differed slightly with the seasons of the year (between 18 and 24°C); in the investigations reported, room temperature variations and changes were not found to exert any influence on the organ temperature patterns examined.

Temperature readings commenced at 9:00 to 9:30 A.M. All subjects had fasted for nine hours and each received a sedative (see below) 30 minutes prior to the liver biopsy. At the hour mentioned, some of the patients offered a relative, or apparent, temperature equilibrium with only small deflections in the temperature curves of the different organs. Other subjects, however, showed at the same hour a tendency for slow fall of temperature in all organs. In these cases it was necessary to wait until the temperature seemed stable enough for experimental procedures. Thus some of the experiments were carried out before noon and others after. In the series of experiments with histamine, reproduced in Figure 1, the smaller and larger dose was given alternatively during morning and afternoon hours in order to reduce as far as possible the influence of spontaneous diurnal temperature shifts during the period in question (9:00 A.M. to 2:00 P.M.).

Drugs used

Histamine. A 0.1 per cent solution of 4-β-aminoethyl-imidazol dihydrochloride was used in doses of ½ to 1 ml. given subcutaneously.

Amyobarbital. A 0.1 Gm. dose was given orally in each case 30 minutes prior to the liver biopsy.

![FIG. 1. EFFECT OF HISTAMINE ON SKIN AND DEEP TEMPERATURES](image)

Mean values from 12 cases (see also Table I). Ordinate: change in centigrades from temperature at histamine injection. Abscissa: time in minutes from injection. Absolute temperatures at 0 time:

<table>
<thead>
<tr>
<th></th>
<th>0.5 mg histamine</th>
<th>1.0 mg histamine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rectum</td>
<td>37.23* ± 0.067</td>
<td>37.21† ± 0.070</td>
</tr>
<tr>
<td>Stomach</td>
<td>37.11 ± 0.044</td>
<td>37.12 ± 0.071</td>
</tr>
<tr>
<td>Liver</td>
<td>36.80* ± 0.076</td>
<td>36.82† ± 0.081</td>
</tr>
</tbody>
</table>

* Difference 0.43 ± 0.10°C, P =< 0.001.
† Difference 0.39 ± 0.11°C, P < 0.01.
Promethazine. (N-(β-methyl-β-dimethylaminoethyl) -
phenothiazine chloride) was used as Lergigan®, brand
of Recip Ltd., Stockholm, in doses of 25 mg., given
intramuscularly.

Dilatol®. A brand of Troponwerke, Köln, (1-(p-oxy-
phenyl)-2-(1'-methyl-3'-phenylproplamo) -propanol-
hydrochloride), was given in doses of 5 to 10 mg. sub-
cutaneously.

RESULTS

Organ temperatures during the initial phase of
measurement

In those cases where the organ temperatures
studied displayed a relatively stable level during
the first 30 minutes after applying the thermo-
couples the experimental arrangements were
started without further delay. In cases, however,
which showed mobile temperature levels in the
shape of a slow fall, the experiments had to be
postponed, as mentioned above, until the tempera-
tures leveled off, which occurred within 1 to 2
hours. This fall in temperature did not repre-
sent the normal trend of the diurnal rhythm dur-
ing these hours. In the present material 15 cases
were selected to illustrate the initial temperature
fall described by Renbourn and Taylor (16). In
these 15 cases the temperature at the start of the
measurements was set as zero value and the sub-
sequent changes from this value during 90
minutes are reproduced in Figure 2 (mean values,
n = 15). At 40 minutes the rectal temperature
had a value of $-0.13 \pm 0.015^\circ C$, whereas the
 corresponding figure for the stomach was $-0.25$
$\pm 0.029^\circ C$. The difference between the two or-
gans ($0.12 \pm 0.032^\circ C$) has a P value < 0.01
and is thus statistically significant, implying that
the lag of the rectal temperature is a real one.

After 90 minutes, or slightly more, a relative
stabilization occurred. It is possible that the
initial temperature drop in these cases was due to
an increased "invisible perspiration" as a sign of
stress, but the skin temperatures corresponding to
the organ temperatures in Figure 2 did not show
any characteristic tendency. In any case, the ex-
perimental measures were postponed until the
temperature curves seemed to level off. In many
cases, however, the curve appeared relatively flat
from the beginning and the equilibration period
was rather short. In no instance were the ex-
periments started until 30 minutes after applying
the measuring equipment in order to let the pa-
tient relax after the liver puncture. The meas-
urements were then continued for 3 to 7 hours, i.e.,
ending at from noon to 4 P.M.

Mutual interrelationship between organ tempera-
tures under study

In a majority of cases the rectal temperature
was highest, the gastric and liver temperatures
running a few tenths of a degree centigrade lower.
Occasionally, the liver was higher and also the
stomach temperature was higher than the two
others, but these cases were fairly exceptional.
The relationships between the temperature curves
of the three organs were relatively constant in one
and the same subject but transient crossings some-
times occurred. Inverse relationships between
the curves, as well as the mentioned crossings,
seemed to lack implication to liver disease.

The effect of histamine

The effect of 0.5 and 1.0 mg. histamine dihy-
drochloride on the temperature curves of skin
(manubrium), rectum, liver and stomach is shown
in Figure 1. The centigrade level at the moment
of the injection has been set as zero value and
the subsequent deflections have been followed.
The temperature changes from pre-injection level
are also given in Table I. The smaller dose elicits
TABLE I

Effect of 1 mg. histamine administered subcutaneously in terms of changes from temperature levels before the injection.

Means and standard errors of 12 cases

<table>
<thead>
<tr>
<th>Minutes after Injection</th>
<th>Temperature change in °C</th>
<th>Liver</th>
<th>Stomach</th>
<th>Rectal</th>
<th>Skin</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>-0.10 ± 0.018</td>
<td>-0.09 ± 0.026</td>
<td>-0.01 ± 0.012</td>
<td>+0.20 ± 0.089</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>-0.16 ± 0.016*</td>
<td>-0.16 ± 0.030</td>
<td>-0.04 ± 0.015*</td>
<td>+0.34 ± 0.117</td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>-0.20 ± 0.019</td>
<td>-0.19 ± 0.021†</td>
<td>-0.09 ± 0.024†</td>
<td>+0.38 ± 0.116</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>-0.23 ± 0.020</td>
<td>-0.22 ± 0.023</td>
<td>-0.12 ± 0.026</td>
<td>+0.63 ± 0.156</td>
<td></td>
</tr>
<tr>
<td>25</td>
<td>-0.20 ± 0.028</td>
<td>-0.20 ± 0.023</td>
<td>-0.12 ± 0.022</td>
<td>+0.55 ± 0.181</td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>-0.18 ± 0.028</td>
<td>-0.18 ± 0.027</td>
<td>-0.13 ± 0.023</td>
<td>+0.56 ± 0.194</td>
<td></td>
</tr>
<tr>
<td>35</td>
<td>-0.15 ± 0.032</td>
<td>-0.14 ± 0.032</td>
<td>-0.13 ± 0.024</td>
<td>+0.58 ± 0.225</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>-0.12 ± 0.032</td>
<td>-0.11 ± 0.035</td>
<td>-0.10 ± 0.023</td>
<td>+0.54 ± 0.244</td>
<td></td>
</tr>
<tr>
<td>45</td>
<td>-0.13 ± 0.042</td>
<td>-0.10 ± 0.039</td>
<td>-0.10 ± 0.029</td>
<td>+0.37 ± 0.232</td>
<td></td>
</tr>
<tr>
<td>50</td>
<td>-0.11 ± 0.032</td>
<td>-0.08 ± 0.051</td>
<td>-0.10 ± 0.026</td>
<td>+0.39 ± 0.206</td>
<td></td>
</tr>
<tr>
<td>55</td>
<td>-0.10 ± 0.043</td>
<td>-0.06 ± 0.054</td>
<td>-0.08 ± 0.043</td>
<td>+0.50 ± 0.248</td>
<td></td>
</tr>
<tr>
<td>60</td>
<td>-0.09 ± 0.034</td>
<td>-0.02 ± 0.047</td>
<td>-0.07 ± 0.037</td>
<td>+0.13 ± 0.124</td>
<td></td>
</tr>
</tbody>
</table>

* Difference between liver and rectum at 10 minutes = 0.12 ± 0.02° C. (P < 0.001).
† Difference between stomach and rectum at 15 minutes = 0.10 ± 0.032° C. (P = 0.01).

an average skin temperature rise of 0.35°C, with a concomitant fall of the "deep" temperatures. These display a very uniform pattern, the rectal temperature showing, however, slower fluctuations than the gastric and liver temperatures. No statistical difference was found to exist between the two latter temperature curves. After 45 minutes, all temperatures were back in the vicinity of the pre-injection level. Later on the curves were less representative, being influenced by the diurnal rhythm shift. The experiments were performed on fasting subjects, alternating during morning or afternoon hours. With the larger dose the rise of skin temperature and fall of rectal, hepatic and gastric temperatures were more pronounced, amounting to + 0.6 and −0.2°C, respectively. In this series, the parallelism between liver and stomach temperatures is even more conspicuous and so is the somewhat tardy character of reaction of rectal temperature. At 20 minutes after

**Fig. 3. Effect of 1.0 mg. Histamine**

Ordinate: organ temperatures and liver blood flow, respectively. Abscissa: time.

The liver blood flow has been determined with bromosulphalein technique according to Bradley, Ingelfinger, Bradley and Curry (42).
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FIG. 4. EFFECT OF HISTAMINE BEFORE AND DURING WARMING OF PATIENT

Ordinate: extreme left scale refers to skin temperature and has been lifted in relation to inner scale, which refers to inner organs. Abscissa: time.

Histamine was administered before and after the patient was put under a heating cradle. The temperature-lowering effect was almost entirely abolished.

In two cases of achlorhydria, repeated histamine doses regularly had the same effect on the temp-
temperature curves as in cases with a normal or increased hydrochloric acid secretion. On account of the limited number of observations the exact figures are not reported.

**Effect of anti-histaminic substance on histamine action**

In four cases 0.025 gram of promethazine (Lergigan®) was given 30 minutes prior to a 1 mg. histamine dose. Promethazine did not seem to influence the temperatures measured in any way. A 0.1 to 0.4°C decrease of stomach and liver temperatures (quite congruent) occurred 5 to 20 minutes after the histamine injection, and in three of the four cases a corresponding skin temperature peak appeared (in the fourth case the histamine-induced vasodilatation evidently did not affect the skin area of manubrium used for measuring).

It was concluded that the effect of histamine on the temperatures of body surface and inner organs is not notably influenced by promethazine.

**Comparison of histamine effect to vasodilatation otherwise elicited**

In Figure 5 the effect of 5 mg. Dilatol® is reproduced (one case selected from a series of eight subjects). The same rise of skin temperature and the corresponding opposite behavior of deep temperatures are obtained as with histamine. Dilatol®, however, in this case gave an effect of longer duration. In four out of eight cases, Dilatol® failed to give a characteristic effect, the individual variations in vascular sensitivity to this drug being seemingly greater than to histamine.

**DISCUSSION**

The equipment used in the present investigation might seem to have a fairly great methodological error when compared to certain procedures described in the literature where equipment is repeatedly stated to have an accuracy of 0.005 to 0.01°C. In the present investigation such extremely sensitive instruments are not necessary since spontaneous temperature fluctuations occur unceasingly, having at all the measurement spots a magnitude of 0.05 to 0.1°C per 5 minutes. These fluctuations are met with even during rest and completely basal conditions (16). According to Menzel (17) and Sollberger (18), several more or less regular endogenous temperature fluctuations occur, having a more frequent periodicity than the well-known diurnal rhythm, and having their different wave lengths superimposed on the latter. Hence, an absolute steady state does not seem to exist. A temperature change of less than 0.05°C per 5 minutes, therefore, is seldom significant, and an accuracy of less than ±0.05°C does not constitute an absolute condition when investigating the slow changes which have been studied here.

As mentioned above, Renbourn and Taylor (16) have dealt with the equilibration period when starting temperature measurements, and state, without any attempt to explain the phenomenon, that in general a fall in rectal temperature occurred during this period. During 40 minutes it amounted to −0.18°C. (S.D. ± 0.3°C.) in Renbourn's quoted series (32 subjects). This figure corresponds fairly well to observations in the present investigation (rectal temperature at 40 minutes = −0.13°C. (S.E. ± 0.015°C.).

**Fig. 5. Rise of Skin Temperature and Fall of Deep Temperatures After Administration of 5 mg. Dilatol®**

Ordinate: as in Figure 4. Abscissa: as in Figure 4. See text.
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General pattern of the organ temperatures measured

Behavior of gastric temperature. The temperature curves obtained from the stomach decidedly support the opinion that the existence of normal secretion or achlorhydria does not exert any influence on the stomach temperature and its reactions to histamine. Unlike Benjamin, Wagner, Zeit, Pisciotta, and Ausman (11), the author has found, in conformity with Deutsch, Spitz and Wohlrab (4), that in achlorhydria the fall in temperature elicited with histamine is the same as in cases with normal secretion. The diversity of results with regard to gastric temperature in achlorhydric subjects could not be easily explained, but on the whole an entirely constant temperature level in the stomach (see Reference 11, Figure 3, page 568) seems very unlikely when all other organs show considerable variations.

Behavior of rectal temperature. Figures 1 and 2 show that rectal temperature does not display the same degree of reaction as do gastric and liver temperatures. The rectal temperature pattern shows a slower reaction and the amplitude of its deflections is smaller. The conclusion is therefore justifiable that rectal temperature is not a sensitive index of the rapid temperature fluctuations occurring in the body. Long ago it was suggested that rectal temperature was representative of the critical body temperature but some authors, e.g., Eichna, Berger, Rader and Becker (19), have stressed its inferiority, and Gerbrandy, Snell and Cranston (20) point out that sublingual as well as esophageal temperatures are far better indices of critical tissue temperature.

Mead and Bonmarito (21) performed measurements of rectal temperature gradients at different heights and found that the gastric temperature rose and fell more quickly than rectal temperatures.

Liver temperature. It has been reported that the temperature of the human liver was lower most of the time than the rectal temperature, but that a temporary reversal of the curves could occur, based upon measurings on eight subjects (15). Although this observation seemed inconsistent with some reports based on animal experiments (22-25), it is in agreement with Eichna, Berger, Rader and Becker (19), Hor-
were higher than the rectal temperature of the same subjects. Reference can also be made to Mather, Nahas and Hemingway (31) who mention that "it has been recognized for some time that the rectal temperature in the normal animal is somewhat above that found in the great vessels." Trying to find the key to the considerable rectal heat, Bazett, in a personal communication to Rubin, Horvath and Mellette (32), suggested that the fecal bacterial activity would partly be responsible, but the latter authors found no evidence of this.

Furthermore, it is interesting to observe that according to Klaften (33) and Netter (34) intrauterine temperature seems to lie higher still; according to Bergman (35) it exceeds rectal temperature by 0.3 to 1.0° C., whereby it seems possible that pelvic temperatures in general are high in relation to abdominal ones.

Differences in measuring technique may partly account for contradictory results referred to and certainly it is important that the rectal temperature should be measured at a constant height since it seems to be lower when measured higher up in the rectum (16, 21).

The effect of histamine. Some of the authors quoted above have even yielded results contradictory to those described in this work. Harmer and Harris (2), as well as Thiessen and Snell (3), did not obtain the regular and uniform lowering of deep temperatures but an increase of rectal temperature and a dissociation of deep temperatures. Concerning the first of these reports, the material examined was rather small, and in the second the rectal temperature was measured with a mercury thermometer, which could scarcely be considered satisfactory.

Several authors dealing with the subject have neglected the increased heat loss by cutaneous vasodilatation as an explanation of the lowered "deep" body temperature induced with histamine and have merely suggested changes in blood flow, e.g., in the stomach, as an interpretation of the hypothermic effect of histamine (3, 6, 9, 10).

Results from animal experiments can hardly be accepted as valid for humans. Nonetheless, some of the studies on the mechanism by which histamine exerts its influence on "body" temperature of animals are of interest. Gyermek (13) found that the rectal temperature of rats exposed to an environmental temperature of 20° C. fell from 36.8 to 34.1° C. after injection with histamine. When the room temperature was changed to 30° C. the rectal temperature of the animals rose 1.1° C. The corresponding relations between room temperature and the thermal effect of histamine were observed in rats by Deutsch, Spitzyn and Wohlrab (4) and by Fabinyi-Szebehely and Szebehely (14) in rats and mice. These authors thus draw the conclusion that the temperature-lowering effect on the body of histamine at ordinary or low room temperatures is due to increased heat loss through the dilated skin vessels, whereas Packman, Rossi and Harrison (36) suggest that a decreased heat production may partly be responsible for the effect in question.

It is reasonable and logical to assume that in humans, too, the lowering of central body temperatures elicited with histamine is secondary to the vasodilatation in the skin and increased loss of heat from the body surface. The reactions of humans seem to be identical with that of the animals in the quoted experiments, viz., when the environmental temperature is increased and heat loss thus prevented, the fall in temperature is eliminated or decreased. In Figure 4 a minute temperature fall occurs, presumably due to the fact that the head and neck are not enclosed in the cradle or covered. As reported by Bohnenkamp and Ernst (37), the face and neck have high values for heat radiation (calculations based on Stefan-Bolzmann's law) and consequently a considerable loss of heat can occur here.

Further evidence that histamine acts by means of increased heat loss is the fact that its effect can be imitated with other drugs capable of giving a skin vasodilatation, not only Dilatol® as in Figure 5, but also adenosine triphosphoric acid (author's experiments, not reported here). Dilatol® has been studied by Külz and Schneider (38), who found that in rabbits, cats and dogs small doses elicited a strong vasodilatation in skin and muscles. Hensel, Ruef and Golenhofen (39), studying the effect of Dilatol® on blood flow in the muscle and skin of eight human subjects, noticed a marked increase in blood flow in the muscles but practically none in the skin. Measurements in Hensel's investigations were, however, performed on the soles of the feet and presumably the blood vessels of the upper half of the body are more apt to react to this agent. Consequently, the tempera-
ture-lowering effect of Dilatol® demonstrated in Figure 5 may be due to cutaneous vasodilatation as well.

If the temperature-lowering effect of histamine were due to decreased heat production, it would be reasonable to suppose that a decrease of the basal metabolic rate occurred simultaneously. This, however, is not the case, either in animals (13), or in humans, where, e.g., von Euler and Liljestrand (40) obtained a 7 per cent increase of BMR at ordinary room temperature.

In the experiments reported in the present investigation, the liver and hepatic vein temperatures rose and fell simultaneously with the gastric temperature and no evidence was found that the temperature fall in the liver was less or slower when compared with that of the stomach. Kosaka (41), measuring temperatures in the portal and hepatic veins of dogs, found that when the animals were submitted to cooling the fall in temperature in the hepatic vein was much less than in the rectum, allowing the conclusion that heat production in the liver rapidly increased. The results reported above do not support the hypothesis of such a mechanism, as both liver parenchyma and liver vein temperatures reacted in the same way and to the same extent as did the other organ temperatures.

SUMMARY

1. Rectal, gastric, liver and skin temperatures in man have been measured with thermocouples.

2. The lowering effect of histamine was the same on liver and hepatic vein and gastric temperatures, but less marked on the rectal temperature. It corresponded in time to a rise in skin temperature and both effects were proportional to the histamine doses used.

3. The effect of histamine on gastric temperature was the same in achlorhydric patients as in individuals with normal hydrochloric acid secretion.

4. The effect of histamine could be imitated with other agents eliciting cutaneous vasodilatation. Only when a skin temperature rise occurred did the inner organs display a fall in temperature.

5. If cutaneous vasodilatation is induced by heating a subsequent histamine dose has but little effect on the temperatures.

6. It is concluded that the temperature-lowering effect of histamine is merely due to a transient increase of heat radiation from the skin. This interpretation corresponds to results from animal experiments reported by various authors.

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