Dose response relationships of insulin hypoglycaemia and gastric acid in man

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SUMMARY Fourteen studies of gastric acid secretion in a basal hour and in the two hours after a single intravenous injection of soluble insulin (0.01 to 0.40 units/kg) were performed in a healthy man. The peak acid output after insulin (measured as the two consecutive 15-minute samples giving the highest acid output) was significantly correlated with the lowest concentration of blood glucose, the fall in blood glucose, the rate of fall of blood glucose, and the maximum fall of blood glucose in any 15 minutes. Peak acid outputs after insulin were similar over the range 0.1 to 0.2 units/kg, and greater than at lower or higher doses.

These results are contrary to the accepted assumption that insulin-stimulated acid secretion is an 'all-or-none' phenomenon. They support instead the hypothesis that insulin hypoglycaemia provides a quantitative glycopenic stimulus producing quantitative vagal acid response. Extreme hypoglycaemia, below about 15 mg/100 ml of blood glucose, inhibits insulin-stimulated acid secretion.

Certain assumptions about the insulin test have been widely held (Roholm, 1930; Hollander, Jemerin, and Weinstein, 1942) in the 40 years since intravenous insulin was found to be an hypoglycaemic stimulus of gastric acid secretion (Simici, Popescu, and Diculesco, 1927). These assumptions are that the acid response is an 'all-or-none' phenomenon which is (1) initiated when the blood sugar falls to a threshold value of 40-50 mg/100 ml; (2) not related to the degree of hypoglycaemia below this threshold; (3) not dependent on the fall in blood sugar; and (4) not related to the rate of fall of blood sugar.

Hollander (1946) later suggested the insulin test as a method of recognizing completeness of vagal denervation of the stomach in man, and his test is used universally, despite lack of agreement on technique and interpretation. However, Clark, Curnow, Murray, Stephens, and Wyllie (1964) suggested that a fall in blood sugar of 30 to 50 mg/100 ml evoked a maximum response regardless of the initial fasting level, whilst Demand, Gross, and Berg (1968) found that the rate of increase of acid output was significantly correlated with the concentration of blood glucose at the time of maximum gastric secretory response.

Almost all investigations of this problem have been done in series of different individuals and it is clear that the relationship between blood sugar and gastric acid needs further examination by tests of different doses of insulin in the same individual.

Method

The subject, a healthy male aged 37 years and without dyspepsia, fasted overnight. The right nostril and throat were sprayed with 3% lignocaine in isotonic saline. A plastic nasogastric tube (Rayx Porges, 14 or 16 mm circumference) was positioned by fluoroscopy in the most dependent part of the body of the stomach, and a plastic cannula inserted in a vein in the forearm. The stomach was emptied and the tube connected to a continuous pump suction, interrupted by manual syringe suction. After a basal hour (four 15-minute periods of collection) soluble insulin
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(Boots, insulin BP) was given as a rapid single intravenous injection over the range 0-01-0-40 units/kg body weight, and eight more 15-minute collections were made. The volume, pH, and titratable acidity of each specimen of gastric juice were measured by methods already described (Baron, 1963). Venous blood was taken before the insulin injection, and 15, 30, 45, 60, 90, and 120 minutes after the insulin to determine the whole blood glucose by a glucose oxidase AutoAnalyzer technique (Wincey and Marks, 1961). Although capillary blood glucose may more closely resemble the glucose concentration reaching the glyco-receptors in the brain responsible for vagal stimulation, venous blood was collected in these studies, so that potassium, cortisol, and growth hormone could be measured as well. These results will be reported separately. Gillespie, Gillespie, and Kay (1969) found no significant difference between arterial and venous blood sugar concentrations during insulin tests.

There were 14 tests with insulin in the range 0-01-0-4 u/kg (Table I). In the one test in which 0-01 u/kg insulin was given, gastric acid was not stimulated at all. The analyses of the relationships between acid secretion and blood glucose changes are based on 13 tests.

Results

In order to make maximum use of the data and to compare the results with the many variables which have been described, extensive analyses have been carried out to examine the relationships between acid secretion and changes in the concentration of blood glucose.

The following variables of acid secretion have been examined: (1) The basal acid output, calculated as the sum of the four basal 15-minute periods. (2) The peak acid output calculated as the sum of the two highest consecutive 15-minute collections, multiplied by two, and expressed as m-equiv/hour (Baron, 1962 and 1963). (3) The maximum acidity, which was the highest concentration of titratable acidity achieved, expressed as m-equiv/litre. (4) The maximum increase in acidity being the maximum acidity minus the mean acidity in the half hour before insulin, and expressed as m-equiv/litre.

Four variables of blood glucose were also examined. These were: (1) the lowest blood glucose, being the lowest value observed after the injection of insulin, and expressed as mg/100 ml. (2) The maximum fall of blood glucose, being the difference between the initial blood glucose and the lowest blood glucose (expressed as mg/100 ml). (3) The rate of maximum fall of blood glucose, being the quotient of maximum fall of blood glucose divided by the time (in minutes) for blood glucose to fall from an initial value to its lowest concentration (expressed as mg/100 ml per minute). (4) The maximum fall of blood glucose in any 15-minute period expressed as mg/100 ml.

Each of these four absolute values of blood glucose has also been expressed as a percentage of the initial, fasting blood glucose, thus giving eight different methods of expressing change in blood glucose.

Basal Secretion

There was no significant correlation between basal acid output and the blood glucose at the end of the basal hour (r = 0.02, p = 0.47).

Secretion after insulin

In the first half hour there was a slight and insignificant decrease in acid output. After this initial half hour of nearly constant secretion, volume, titratable acidity, and acid output increased to a peak and then declined to basal levels within two hours of the injection of the insulin. In one test, which was prolonged, acid output remained low in the third hour, and there was no suggestion of a late response after 120 minutes (Fig. 1). Concentrations of blood glucose decreased to their lowest level at 30 minutes after the insulin injection and rose again towards basal levels within 2 hours. In only two tests was the lowest blood glucose delayed until 45 minutes.

Timing of peak acid output and dose relationship

The peak half hour of acid secretion in the two hours after insulin was evenly distributed during the period 30-120 minutes, and showed no relationship to the dose of insulin, the lowest blood glucose, or the maximum fall in blood glucose.

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<table>
<thead>
<tr>
<th>Insulin Dose (u/kg)</th>
<th>No. of Tests</th>
<th>No. of Tests in which Peak Half-hour Acid Occurred in Each Period (min)</th>
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<tr>
<td></td>
<td></td>
<td>0-30 15-45 30-60 45-75 60-90 75-105 90-120</td>
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<tr>
<td>0:01</td>
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<td>1</td>
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<td>0:4</td>
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<td>1</td>
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Lowest blood glucose (mg/100 ml)

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<tr>
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<th>No. of Tests</th>
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<td>11-20</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>21-30</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>31-40</td>
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<td>1</td>
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Maximum fall in blood glucose (mg/100 ml)

<table>
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<th>No. of Tests</th>
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<td>21-30</td>
<td>1</td>
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<td>31-40</td>
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<tr>
<td>&gt; 40</td>
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All tests

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<th></th>
<th>No. of Tests</th>
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<tbody>
<tr>
<td></td>
<td>3</td>
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</tbody>
</table>

Table I Timing of peak acid output in relation to insulin dose and changes in blood glucose
The usual hypoglycaemic symptoms and signs appeared 20-30 minutes after insulin was injected but lessened with repeated testing.

**Peak Acid Output**

The graph of the relationship of peak acid output after insulin and the lowest blood glucose suggested that acid output increased as blood glucose levels became lower; the slope of the straight line fitted to this data was significantly different from zero (Fig. 2a). However, further inspection of the data shown in Fig. 2a suggested that there might be a systematic biphasic acid secretory response to changes in the lowest blood glucose, with peak acid output rising as blood glucose falls, but declining when lowest blood glucose falls below about 15 mg/100 ml.

Some evidence for these trends was provided by fitting quadratics to the data (Fig. 2b): the residual variation of the data about the quadratics was significantly less than that about the straight line. Although the quadratic is the most convenient curve for testing whether a trend is linear, many other alternative curves would fit the data as well or even better. For example, straight lines can be fitted separately for the two phases of the data, stimulation and inhibition, and these lines fit the data very well (Fig. 3). There is no statistical basis for choosing between these various ways of describing the data, and the choice must rest on...
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the plausibility of the relationships and their descriptive usefulness. With a small number of observations there is a good case for just reporting the actual data and this policy has been adopted with the rest of the results of acid output.

With increase in the maximum fall of blood glucose (the second of the four variables of blood glucose examined), peak acid output after insulin increased, but again there was a situation in which large falls in blood glucose led to inhibition of acid output (Fig. 4a). The graphs for the rate of maximum fall of blood glucose (third variable) (Fig. 4b), and maximum fall of blood glucose in any 15 minutes (fourth variable) (Fig. 4c) showed similar trends. In all but one of the eight measurements of changes in blood glucose, a quadratic

\[
\begin{align*}
\text{Fig. 2b Quadratic correlations } (y &= a + b_1x + b_2x^2) \\
\text{of peak acid output after insulin with lowest blood glucose expressed as mg/100 ml (pb}_2 = 0.0000003) \\
\text{and as a percentage of initial blood glucose (pb}_2 = 0.0000003) \text{ in all 13 tests.}
\end{align*}
\]

\[
\begin{align*}
\text{Fig. 3 Linear correlations of peak and output after insulin with lowest blood glucose over the range in which hypoglycaemia is associated with stimulation of gastric acid secretion } (r = -0.84, P = 0.001; \\
\text{r = 0.91, P = 0.0001}) \text{ and the range in which extreme hypoglycaemia is associated with partial inhibition of acid secretion } (r = 0.75, P = 0.007; r = 0.85, \\
P = 0.03).
\end{align*}
\]
Fig. 4a  Peak and output after insulin and maximum fall in blood glucose.

Fig. 4b  Peak and output after insulin and rate of maximum fall in blood glucose.

Fig. 4c  Peak acid output after insulin and maximum fall in blood glucose in any 15 minutes.
curve provided a better fit to the data than a straight line, thus providing evidence that the trends were not linear; however, it is also clear from the graphs that many of these measurements are highly correlated with each other.

TITRATABLE ACIDITY
The maximum acidity and increase in acidity were related in a very similar fashion to that of peak acid output to measurements of blood glucose.

INSULIN DOSE AND BLOOD GLUCOSE
There were of course statistically significant correlations (Table II) between the dose of insulin and all the different expressions of changes in blood glucose described above. The larger doses of insulin prolonged the hypoglycaemia so that two hours after the injection of insulin blood glucose was also inversely correlated with the dose of insulin.

INSULIN DOSE AND ACID OUTPUT
Peak acid outputs after insulin increased over the range 0-01-0-1 units/kg, were similar over the range 0-1-0-2 units/kg, and then tended to fall (Fig. 5a). The relationship with insulin dose and acid output over the whole two hours was similar (Fig. 5b). Similar curves were obtained if the acid output after insulin was expressed for the periods 0-60, 60-120, 30-90, or 30-120 minutes, or as peak acid output minus basal acid output.

Discussion

These results are not compatible with the accepted assumption (Bachrach, 1953) that gastric acid secretion after intravenous insulin is an 'all-or-none' response to a blood sugar below 50 mg/100 ml, and suggest instead a relationship between insulin dose, hypoglycaemia, and acid secretion by the stomach. The hypoglycaemia-sensitive cephalic centres (Jögi, Ström, and Uvnäs, 1949) may respond to the lowest blood glucose concentration achieved, the absolute fall, the maximum rate of fall, or the maximum fall in any unit time. The hypoglycaemic stimulus might be either an absolute change in blood glucose concentration or a proportion of initial concentration of blood

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Table II Linear correlations of insulin dose and changes of blood glucose in 13 tests

<table>
<thead>
<tr>
<th>Lowest blood glucose (mg/100 ml)</th>
<th>0.7243</th>
<th>0.0026</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lowest blood glucose (percentage of initial blood glucose)</td>
<td>0.7083</td>
<td>0.0034</td>
</tr>
<tr>
<td>Maximum fall of blood glucose (mg/100 ml)</td>
<td>0.7026</td>
<td>0.0037</td>
</tr>
<tr>
<td>Maximum fall of blood glucose (percentage of initial blood glucose)</td>
<td>0.7083</td>
<td>0.0034</td>
</tr>
<tr>
<td>Rate of maximum fall of blood glucose (mg/100 ml per min)</td>
<td>0.6597</td>
<td>0.0041</td>
</tr>
<tr>
<td>Rate of maximum fall of blood glucose (percentage of initial blood glucose)</td>
<td>0.6597</td>
<td>0.0041</td>
</tr>
<tr>
<td>Maximum fall of blood glucose in any 15 minutes (mg/100 ml)</td>
<td>0.7833</td>
<td>0.0008</td>
</tr>
<tr>
<td>Maximum fall of blood glucose in any 15 minutes (percentage of initial blood glucose)</td>
<td>0.8255</td>
<td>0.0003</td>
</tr>
</tbody>
</table>
glucose before the stimulus. Inhibition may occur in association with extreme hypoglycaemia. The present data are compatible with all these possibilities. There are three major phenomena to be considered and they will be discussed separately.

AN HYPOGLYCAEMIC THRESHOLD FOR THE INITIATION OF ACID STIMULATION

The smallest dose of insulin (0.03 u/kg), which elicited a blood glucose response from the subject produced a blood glucose as low as 36 mg/ml. An even smaller dose (0.01 u/kg) produced a blood glucose of 58 mg/100 ml but no acid stimulation. It was not possible from these limited studies to determine a precise hypoglycaemic threshold for initiation of acid secretion, but various estimates have been made ranging from 18-67 mg/100 ml blood sugar (Stempien, Lee, and Dagradi, 1968) to 50-85 mg/100 ml blood sugar (Hiles, 1947). Although most authors accept the threshold hypothesis, there have been suggestions ‘that the vagal stimulation resulted more from a rapid decline in blood sugar than from its actual value at any one time’ (Hiles, 1947), or that ‘a falling blood sugar may be sufficient for stimulation regardless of the rate or depth of that fall’ (Stempien et al, 1968). However, the results in diabetic humans (Kalk and Meyer, 1932) and dogs (Kemp, Herrera, Isaza, and Eisenberg, 1968a) seem conclusive evidence for the hypothesis of a threshold blood glucose for initiation of acid secretion, and against the hypotheses of initiation by fall or rate of fall of blood glucose.

It seems reasonable to conclude from these studies that when the blood glucose falls, acid secretion is initiated at a certain threshold, and that a fall of blood glucose which does not reach this threshold, however rapid and even if the value is as large as several hundred mg/100 ml, will not initiate gastric secretion. Although there does not seem to be any universal threshold for blood sugar in man or dog, the human threshold appears to be between 33 and 73 mg/100 ml (Demand et al, 1968), but may not be the same in the same individual from day to day. This threshold is below the range of blood glucose during normal activities, and there was no correlation between basal acid output and blood glucose in the present series. It is, therefore, unlikely that variations in spontaneous interdigestive basal secretion in man could be due to variations in blood sugar.

INSULIN DOSE, HYPOGLYCAEMIA, AND ACID OUTPUT

The results presented have shown a quantitative relationship between insulin hypoglycaemia and gastric acid output. This relationship has not previously been described, and the literature has therefore been reviewed critically to explain these differences.

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Fig. 6 Acid output and lowest blood sugar, recalculated from the data of dog 77 of Jemerin et al (1942) (r = -0.917; P = 0.014).
**Animal data**

Hollander et al (1942), in their original study, concluded that 'it is impossible, however, to establish any quantitative relation between the magnitude of the insulin dosage or the hypoglycaemia on the one hand, and the volume-rate of secretion or the acidity on the other'. However, Jemerin, Hollander, and Weinstein (1942) claimed 'a rough parallel between the degree of hypoglycaemia and the magnitude of response was obtained in this animal'. This discrepancy may be related to Hollander’s group having measured volume and acidity separately and omitted to calculate their product, acid output. Their data on dog 77 have therefore been recalculated (Fig. 6); there is a significant inverse linear correlation of gastric acid output with the lowest blood sugar concentration, similar to the results in the present paper (Fig. 2a). Similarly, recalculation of the data of Metys and Ronský (1959) suggests an inverse linear correlation (similar to Figs. 2a and 6) of acid output with the lowest blood sugar concentration (n = 6, r = −0.69, p = 0.066).

The most recent and apparently definitive studies of insulin hypoglycaemia in the dog (Davis and Brooks, 1962 and 1963; Davis, Brooks, and Robert, 1965; Brooks, 1967) claimed that the authors had confirmed the old ‘all-or-none’ law, although they failed to consider that acid response after single parenteral injections of drugs in man or dog should be analysed in terms of peak acid output (Marks, Komarov, and Shay, 1960; Baron, 1962 and 1963; Makhlouf, 1968) irrespective of the timing of peak acid output (Baron, 1968 and 1969). Recalculation of the data of Davis and his co-workers suggests that the acid changes are correlated with the blood sugar changes, and thus the acid changes are correlated with the insulin dose.

A dose-response relationship for vagally stimulated gastric acid was suggested for 2-D-deoxyglucose which is thought to stimulate the cephalic centres by the production of intracellular glycopenia, without reduction of glucose concentration in the extracellular fluid (Hirschowitz and Sachs, 1965; Eisenberg, Emas, and Grossman, 1966). Hirschowitz (1966) therefore deduced that insulin would be expected to show a similar dose-acid response relationship if to the observed acid output was added the amount of acid which insulin had inhibited at the same time. Hirschowitz and Sachs (1967) were able to demonstrate a significant correlation between the degree of hypoglycaemia and pepsin output, but not with acid output.

**Human data**

These are summarized in Table III. The results in the present paper suggest that acid secretion once

<table>
<thead>
<tr>
<th>Source</th>
<th>Drug</th>
<th>Dose</th>
<th>Route</th>
<th>Findings</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roholm (1930)</td>
<td>Insulin</td>
<td>12 units</td>
<td>Intravenously</td>
<td>No correlation between the lowest, the fall or the rate of fall of blood sugar and the increase in, or maximum, acidity.</td>
<td>'All-or-none' relationship</td>
</tr>
<tr>
<td>Ihre (1938)</td>
<td>Insulin</td>
<td>16 units</td>
<td>Intravenously</td>
<td>Acid output after 20 units not more than after 16 units. Gastric stimulation not related to hypoglycaemia.</td>
<td>'All-or-none' relationship</td>
</tr>
<tr>
<td>Bachrach (1949) quoted by Bachrach and Bachrach (1967)</td>
<td>Insulin 0-1, 0.2-0.4 u/kg in same patient</td>
<td>Intravenously</td>
<td>Similar acidities after each of these three doses (secretory volume and acid output not reported).</td>
<td>'All-or-none' relationship</td>
<td></td>
</tr>
<tr>
<td>Brooke (1949)</td>
<td>Insulin</td>
<td>10 units</td>
<td>Intravenously</td>
<td>No correlation between acid response and fall or rate of fall of blood sugar</td>
<td>'All-or-none' relationship</td>
</tr>
<tr>
<td>Sharick and Campbell (1951)</td>
<td>Insulin 2-2 u/kg</td>
<td>Subcutaneously</td>
<td>No correlation between lowest blood sugar and acid response.</td>
<td>'All-or-none' relationship</td>
<td></td>
</tr>
<tr>
<td>Clark et al (1964)</td>
<td>Insulin</td>
<td>15 units</td>
<td>Subcutaneously</td>
<td>In Fig. 2 a clear linear correlation can be seen between insulin-stimulated acid (as % histamine-stimulated) and maximum fall of blood sugar.</td>
<td>Maximum acid response evoked by fall in blood sugar of 30-50 mg/100 ml</td>
</tr>
<tr>
<td>Aylett (1965)</td>
<td>Tolbutamide</td>
<td>100 mg</td>
<td>Intravenously</td>
<td>No correlation between blood glucose and gastric secretion.</td>
<td>'All-or-none' relationship</td>
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<tr>
<td>Demand et al (1968)</td>
<td>Insulin</td>
<td>0-09 u/kg/hr</td>
<td>Intravenously</td>
<td>Significant correlation between rate of increase of acid output and the absolute blood glucose concentration at the maximum of secretory stimulation but no correlation with the insulin dose, the absolute or relative blood glucose at the onset of secretion, or the rate of fall of blood glucose.</td>
<td>'All-or-none' relationship</td>
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<tr>
<td>Gillespie et al (1969)</td>
<td>Insulin</td>
<td>20 units</td>
<td>Intravenously</td>
<td>No correlation between blood sugar change and acid response.</td>
<td>'All-or-none' relationship</td>
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</table>

Table III  The relationship between hypoglycaemia and gastric acid in man
initiated after insulin hypoglycaemia is a quantitative vagal acid response in response to a quantitative glycopenic stimulus. It seems clear that previous reports, which have almost consistently claimed an ‘all-or-none’ response, have been almost exclusively based on studies of different subjects or animals. In the few studies where the same animal has been studied on more than one occasion, recalibration of the data produced complete agreement with the interpretation of the human data of the present study.

The graphs of the correlation of acid output with lowest blood sugar (Figs. 2 and 6) are mirrored by the dose-response curves of peak acid output and insulin dose (Fig. 5a). However, the scatter of points along the line in Fig. 5 is considerable, presumably because the reproducibility of the insulin test is poorer than for other gastric near-maximal stimuli. For example, the coefficient of variation of 25 duplicate measurements of peak acid output after 0-15 units/kg insulin was 16% (Hubel, 1966) as compared with 8% after an augmented histamine test (Baron, 1963), or 9% after pentagastrin (Baron, 1969).

The imprecision of this relationship of acid output and insulin dose is understandable if ‘the dose of insulin required to produce a given degree of hypoglycaemia cannot be predicted, because it varies not only in different individuals (man or animal) but also in the same individual from day to day’ (Bachrach, 1953).

Inhibition of gastric secretion after insulin

In the present study there was no significant inhibition in the first half hour after insulin, but acid output was apparently increasingly inhibited at extreme hypoglycaemia, below about 15 mg/100 ml. The literature on this problem is most confusing, because inhibition has been reported both soon and late after insulin, because different commercial insulins contain different amounts of glucagon, and because the inhibition after insulin may not be due to the hypoglycaemia but may be related to some other change in the internal environment of the cephalic centres or the parietal cells of the stomach.

Animal data

Necheles, Olson, and Morris (1940 and 1941) noticed depression of gastric motility and tone after 0-5-0-7 u/kg Squibb insulin in the dog; this inhibition coincided with very low blood sugars (10-28 mg/100 ml), and, as the blood sugar rose to medium and slightly subnormal values, the typical insulin hypermotility appeared. Necheles, Olson, and Scruggs (1942) showed that mild hypoglycaemia from low doses of insulin stimulated gastric secretion from Pavlov pouches; severe hypoglycaemia from higher doses of insulin produced less stimulation; other similar results have been summarized in Table IV.

The simplest explanation for the inhibition of gastric secretion associated with hypoglycaemia was that the low blood glucose directly inhibits parietal cell function as does anoxia (Pickett and van Lierie, 1939) analogous to the depression of intestinal enzyme output (Kneller and Nass, 1949) or colonic motility (Kilnberg and Cornell, 1964) associated with severe hypoglycaemia. Indeed, Jordan and Quintana (1964) were able to inhibit insulin-stimulated gastric secretion by inducing profound hypoglycaemia with a larger dose of insulin; both this hypoglycaemia and the inhibition were still produced if glucagon-free insulin was used, and both were prevented by glucose infusions. Eisenberg, Woodward, Quintana, and Dragstedt (1963) could not prevent inhibition of histamine-stimulated acid by glucose infusion, and therefore suggested that ‘a transient hyperkalsis of the parietal cell temporarily interferes with acid elaboration ...’.

Hirschowitz and his co-workers (Hirschowitz and O’Leary, 1964; Hirschowitz, 1966; Hirschowitz and Sachs, 1967a and b) have systematically studied the inhibitory action on histamine-stimulated secretion of large doses of insulin. This inhibition was obtained with glucagon-free insulin, was dose-dependent, associated with a fall in plasma and gastric juice potassium, and not reversed by prevention or correction of the hypoglycaemia, but was reversed or prevented by intravenous potassium chloride or by intravenous rubidium. The inhibition associated with very high doses of insulin in the rat (Lee and Thompson, 1967) and dog (Kemp, Herrera, Tsukamoto, and Eisenberg, 1968b) coincided with depression of blood sugar and serum potassium.

It seems clear from these studies that insulin could both (vagally) stimulate and (peripherally) inhibit gastric secretion. There are two possible experimental methods of confirming this double action of insulin by studying the two effects independently.

First, ruminants (who have no psychic vagal phase of gastric secretion) show only inhibition of gastric acid after insulin with a significant correlation between blood sugar and titratable acidity. This inhibition was unaffected by vagotomy and was therefore a peripheral effect associated with hypoglycaemia (Hitchcock,
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Karvonen, and Phillipson, 1948). Secondly, a non-hypoglycaemic drug such as 2-D-deoxy-glucose could be used as a vagal stimulant; increasing doses showed no dose-response reversal, and the peak acid output after 2-D-deoxy-glucose was significantly higher than after insulin (Hirschowitz and Sachs, 1965; Eisenberg et al, 1966).

INHIBITION OF GASTRIC SECRETION AFTER INSULIN IN MAN

Initial phase
Mahler (1930) noted a fall in acid secretion in the first 25 minutes after insulin, attributable to a primary inhibitory action of insulin itself. A similar inhibitory phase was also noted by Roholm (1930) after Leo insulin. Olson and Necheles (1953) therefore systematically studied in 15 patients with duodenal ulcer the change in acid after 25 units insulin and found a significant decrease at 30 minutes compared with pre-insulin acid output. A similar significant reduction was found in the volume, acidity, and acid output at 15 and 30 minutes after regular insulin in 33 healthy subjects (Olsen and Necheles, 1955) and it was thought possible that this inhibition could be due to the glucagon in this insulin.

Aylett (1963) has reviewed the effect of glucagon on gastric secretion in man and animal, having shown (Aylett, 1962) that as little as 2 mg glucagon intravenously would significantly inhibit gastric secretion in response to a water meal and that inhibition was present even if the glucagon was given at the same time as glucagon-free insulin. Aylett agreed that the biphasic action of ‘insulin’ upon gastric secretion may have been due to the glucagon contamination, and she suggested that ‘glucagon-free insulin’ should always be used for this test.

The glucagon content of commercial insulin varies considerably, was probably always considerable from the 1920s to the 1940s, and may have been responsible for the many previous reports of an early phase of inhibition of gastric secretion after ‘insulin’. However, in the last twenty years the manufacturing methods in Britain and Denmark have changed, and these insulins, unlike some American insulins (Rowlinson and Lesford, 1951), now contain insignificant amounts of hyperglycaemic factor. In the present studies no significant early inhibitory phase was noted after the Boots insulin, which with other British insulins, as well as Novo insulin, have glucagon concentrations so low (less than 0.01%) as to be negligible, and therefore are suitable for the insulin test, as is Lilly crystalline insulin (0.1% glucagon).

EXTREME HYPOGLYCAEMIA IN MAN
In one patient in the series of Clark et al (1964) a fall in blood glucose of more than 60 mg/100 ml (fasting and lowest blood glucose not stated) was associated with relatively low peak acid output after insulin compared with post-histamine acid output.

There are no other reports in the literature of extreme hypoglycaemia being achieved in man in an insulin test, but the results of the present study in which blood glucose concentrations below 15 mg/100 ml were associated with decreased acid output are entirely compatible with the results in animals subject to extreme hypoglycaemia, irrespective of whether the mechanism of inhibition of parietal cell function is by decrease in the glucose concentration and/or the potassium concentration in the blood.

Criteria for an adequate insulin test in the unoperated subject
Obviously, the results in one individual may not be taken as representative and further tests with different doses of insulin are needed in many different subjects. However, the present results suggest that an insulin dose of about 0.2 u/kg may be optimum in producing sufficient hypoglycaemia (blood glucose below 30 mg/100 ml) to guarantee initiation of gastric secretion in an individual, to ensure a near maximum vagal acid output, and to prevent blood glucose falling so low (below 15 mg/100 ml) that hypoglycaemic inhibition of gastric secretions or dangerous side effects are produced.

Addendum

Isenberg, Stening, Ward, and Grossman (1969) gave intravenous injections of insulin (0-025-0.4 u/kg) to five healthy men. Peak 15-minute acid output was similar after 0.1, 0.2 and 0.4 u/kg and highest after 0.2 u/kg. Blood glucose at 30 minutes were similar after 0.1-0.4 u/kg, but no acid/glucose correlations were made. Stening and Isenberg (1969) gave cats with gastric fistulas 0.03-1 u/kg insulin intravenously and found 0.5 u/kg produced the highest peak 15-minute acid output.

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References