IDIOPATHIC NEONATAL HYPERGLYCAEMIA

BY

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Attention has recently been drawn to a temporary diabetic state in newborn infants with complete recovery in weeks or months (Hutchison, Keay, and Kerr, 1962; Sweetnam and Sykes, 1962). Infants with this condition exhibit little or no ketosis and have low insulin requirements compared with true diabetes mellitus presenting at this age (Guest, 1949; Gans, 1954; Hofman-Bang, 1954). The condition is a separate entity from the hyperglycaemia seen in association with infections and cerebral abnormalities. For these reasons we suggest the use of the term ‘idiopathic neonatal hyperglycaemia’.

The paucity of published material on this subject since 1850 suggests that such cases are rare, but we have learnt of several treated by colleagues in the past 12 months, and some cases may pass unrecognized. We consider that the condition is more common than has previously been supposed.

Early recognition and careful treatment are necessary in the more severe cases if sequelae are to be avoided. We therefore feel justified in reporting two cases, together with a brief review of the published material.

Case Reports

Case 1. A male first child of healthy parents; there is no family history of diabetes mellitus. He was born on July 29, 1962, after a normal pregnancy by breech delivery, following surgical induction of labour at 42 weeks gestation. Delivery was not difficult; the placenta was described as ‘very infarcted’.

His birth weight was 4 lb. 6 oz. (1·98 kg.) and he made good progress, feeding well from the breast until the 7th day, when he developed mild diarrhoea and vomiting. There was no weight loss until the 10th day, when he lost 2 oz. (57 g.). Routine examination of the urine showed a reducing substance, identified chromatographically as glucose. Breast milk was insufficient and artificial feeding was introduced. He was transferred to the South London Hospital at the age of 12 days.

On admission he weighed 4 lb. 8 oz. (2·08 kg.) and showed the characteristic appearance of dysmaturity.

He was vigorous and not dehydrated. Blood sugar (two and a half hours after a feed) was 465 mg./100 ml.; urine sugar 2%o; acetone nil. The glucose tolerance test at 3 weeks gave the following results: fasting 115 mg./100 ml.; ⅓ hour 153 mg.; 1 hour 200 mg.; ⅔ hour 175 mg.; and 2 hours 200 mg./100 ml.

Serum electrolytes were normal. No pathogens were isolated from rectal swabs. Urinary 17-ketosteroids were 0·3 mg./24 hours; and 17-ketogenic steroids were 0·73 mg./24 hours. There was no serum insulin-like activity at age 6 weeks.

He continued to have slightly loose stools, which improved on oral neomycin. Half cream milk was given, and he gained weight until the age of 5 weeks, at which time he weighed 6 lb. (2·73 kg.) (637 g. above birth weight). This rate of progress was not maintained and did not improve when sugar was omitted from his feeds, the calorie requirement being made up with milk powder. Occasional pyrexial episodes responded to oral fluids, and were attributed to polyuria.

At the age of 9 weeks he weighed only 5 lb. 15 oz. (2·7 kg.) and insulin treatment was started. In view of the importance of avoiding hypoglycaemia (Hutchison et al., 1962; Brown and Wallis, 1963), treatment was begun with small doses of soluble insulin, 0·2 units after each feed, i.e. 1·2 units a day. This resulted in a weight gain of 15 oz. (425 g.) in one week. There was little further weight gain until the dose of insulin had been increased to 1 unit after each feed, i.e. 6 units a day. A weaning diet and full cream milk were introduced at 16 weeks.

At the age of 17 weeks, 4 consecutive insulin doses were omitted. His weight fell by 10 oz. (283·5 g.) and his temperature rose to 102· F. (39· C.). The blood sugar was 1,028 mg./100 ml. and alkali reserve 15·3 mEq/l. He was given soluble insulin, 1 unit, and rehydrated (N/2 saline subcutaneously and a milk feed). Five hours later the blood sugar had fallen to 334 mg./100 ml.

Subsequent progress was interrupted only by minor upper respiratory infections and on one occasion by vomiting, which appeared to be associated with teething.

At the age of 8 months he weighed 13 lb. 10 oz (6·2 kg.) and an intravenous tolbutamide test was performed (Fig. 1). This showed a significant fall in blood sugar suggesting the presence of functioning islet-cell tissue.

At the age of 15 months he was having 8 units of insulin a day in three divided doses and was below the 3rd
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...and he handled toys well. Insulin was increased to 16 units in two doses and this resulted in a weight gain of 2 lb. 5 oz. (1,055 g.) in one month.

COMMENT. Treatment has been directed throughout to maintain a satisfactory weight increase and to avoid hypoglycaemia, with its attendant risk of irreversible brain damage. From assessment at the age of 15 months, it seems that while we have achieved the second objective, we have fallen short of the first. It remains to be seen whether he will reach normal stature.

Though the complete recovery implicit in the diagnosis has not yet been confirmed in this child, we feel confident in including him in this group of cases for the following reasons.

1. The low birth weight and post-maturity in common with 12 previous cases.

2. The degree of response to insulin shown by the weight gain following the introduction of relatively small doses of insulin, and the fall of blood sugar from 1,028 to 534 mg./100 ml. following 1 unit of insulin.

3. The absence of ketosis at the onset of the disease and the relatively minor degree of ketosis associated with a high blood sugar when insulin was withheld.

Fig. 1.—Case 1. Intravenous tolbutamide test (15 mg. kg.) at 8 months, showing fall of blood sugar after 1 hour.

Fig. 2.—Case 1. Progress chart showing weight curve, insulin therapy, and blood sugars.
(4). The tolbutamide test at 8 months suggests recovering pancreatic function.

There is no evidence at present of any complications of hyperglycaemia and the serum cholesterol is normal (155 mg./100 ml.), but a closer control of the blood sugar level must be attempted for as long as he needs insulin. Fig. 2 shows the progress up to 4½ months.

Case 2. This fourth child of a healthy Ghanaian family was born normally on December 25, 1962 at the London Hospital at 38 weeks gestation, there having been mild hydramnios. She weighed 7 lb. (3.17 kg.) and seemed shocked and pale at birth but required no resuscitation. After 9 hours she became oedematous and cold (rectal temperature 93°F. (33·9°C.). The hypothermia was satisfactorily corrected, and in the next three days a diuresis occurred, resulting in loss of oedema fluid and 1 lb. (453 g.) in weight. She then appeared healthy, but failed to gain, despite an adequate intake from the breast. On the twelfth and thirteenth days complementary feeds were given by tube, and this was followed by dramatic weight loss to 5 lb. (2·26 kg.) associated with polyuria.

On examination she was lively, thirsty, bright-eyed, but grossly dehydrated, though still passing urine.

Investigations showed a blood sugar of 890 mg./100 ml. (true glucose 840 mg./100 ml.) and heavy glycosuria, but no acidosis or ketonuria.

Treatment was instigated with intravenous half-normal saline and intramuscular soluble insulin 5 units.

Two hours later the blood sugar was 720 mg./100 ml., and after four hours it was 130 mg./100 ml. At this time generalized convulsions developed. These were temporarily relieved by 11 g. glucose intravenously, but occurred frequently and were not relieved by calcium, sedation, or further glucose; they ceased spontaneously after 36 hours, by which time small milk feeds were being tolerated. The blood sugar varied from 300 to 600 mg./100 ml., the serum calcium and other electrolytes remained within normal limits, and in 24 hours the hydration was restored to normal. (The serum magnesium level was not measured.) Three small doses of insulin (total 0·8 units) were given during this time in an attempt to reduce the osmotic diuresis, and thereby improve the hydration and lessen the electrolyte loss. The EEG remained normal throughout, but the infant did not exhibit normal neurological responses for 7 days.

She sucked well, however, and breast feeding was soon re-established, but she failed to gain during three days without insulin. Soluble insulin 0·1 units every 4 hours was therefore given with feeds, increasing over the next few weeks to an average of 6 units a day in order to maintain a satisfactory weight gain. Two episodes of ketosis with minimal acidosis, one lasting 12 hours, the other almost three days, responded to extra insulin and M/6 sodium lactate solution by mouth. No attempt was made to eliminate glycosuria or hyperglycaemia, which persisted to 10 weeks of age. The blood sugar having gradually fallen to the region of 100 mg./100 ml., insulin was withdrawn at this stage for the purpose of investiga-

tion and found to be no longer necessary. She now weighed 9 lb. 1 oz. (4·1 kg.).

Plasma insulin levels were estimated at 10½ weeks, when insulin had been withdrawn for three days. A very high fasting level was found (165 μU./ml.) with a further rise after intravenous tolbutamide and after oral glucose (Fig. 3); hypoglycaemia did not develop during a prolonged starvation period (Fig. 4). Insulin antibodies were demonstrated in the plasma.

At 12 weeks a glucose tolerance test was normal and the plasma insulin levels were also recorded (Fig. 5).
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The features of idiopathic neonatal hyperglycaemia are sufficiently characteristic to constitute a distinct clinical picture. There is hyperglycaemia with consequent polyuria, polydipsia, and dehydration, but without proportionate ketosis; and this is followed by eventual recovery to normal glucose tolerance.

The 13 published cases of idiopathic transient hyperglycaemia, together with our two cases, are summarized in the Table. They all presented by the age of 6 weeks, and all but three were diagnosed in the first 4 weeks of life. Two had severe associated disease (Cases 2 and 3) which, it may be argued, provoked the diabetic-like state; however, gangrene (Case 2) or abscess (Case 3) may equally well have been the result of hyperglycaemia.

Six other cases of neonatal hyperglycaemia have been recorded, all of which died. Four of these we have excluded because they showed pancreatic abnormalities at post-mortem examination (Limmer and Miller, 1935; Lewis and Eisenberg, 1935; Devine, 1938; Cuno, 1911). The other two may have fitted the category of idiopathic hyperglycaemia and may have proved temporary had the infants survived; one died before insulin was discovered (Kitselle, 1852), and the other shortly after the onset of treatment (Hickish, 1956).
The case described by Jeune and Riedweg (1960) had associated gastro-enteritis due to a specific type of Esch. coli, severe acidosis, and stupor with blood-stained cerebrospinal fluid. It is difficult to postulate that hyperglycaemia of two days' duration was the primary factor in this child’s illness.

Since 1850 there have been only three reports of true diabetes mellitus with such early onset, i.e. 11, 21, and 39 days, respectively (Guest, 1949; Hofman-Bang, 1954; and Gans, 1954). These cases differ strikingly from those with idiopathic hyperglycaemia. Two had severe ketosis, and the third was diagnosed by the mother, who, having other diabetic children, found glucose in the urine wrung out from the nappin before symptoms appeared. A strong family history was present in two cases.

Aetiology. Definite dysmaturity was present in all but 4 cases, there being a very low birth weight for gestation period in 11. Of the remaining 4 cases, 2 (Cases 5 and 7) weighed only 6 lb. 2 oz. (2,777 g.) at term, and one of these (Case 7) was described as post-mature.

Both sexes were equally affected, and a family history of diabetes mellitus was recorded in two (Cases 1 and 13).

Cerebral damage appears to play little part in the causation of this syndrome. It was considered necessary to examine the cerebrospinal fluid in only 4 cases: Keidan’s case had xanthochromic, blood-stained fluid with the protein rising to 1,000 mg./100 ml.; Nawrocka-Kanska’s case showed a raised protein and xanthochromia; Engleson mentioned the sugar but did not record the other findings, and one of the cases described by Hutchison et al. had a protein level of 80 mg./100 ml. Keidan (1955) was unable to explain the significance of the cerebrospinal fluid changes, but Nawrocka-Kanska (1952) considered the cerebral haemorrhage to be the fundamental cause. However, neither of the two infants described here exhibited signs of cerebral damage before or during the development of hyperglycaemia, nor was this a feature in the other 10 cases. If cerebral damage were a significant aetiological factor, one would expect symptoms of hyperglycaemia to be common in babies with cerebral birth trauma, and this is not so. The transient glycosuria secondary to cerebral lesions is usually due to a lowered renal threshold.

**Insulin Studies.** There are no previous insulin studies in these infants. Plasma insulin activity has been satisfactorily demonstrated in the normal newborn and in infants of diabetic mothers by the rat diaphragm technique (Baird and Farquhar, 1962; Stimmmer, Braize, and O’Brien, 1964).

Dr. Keith Taylor kindly undertook estimations of serum insulin activity in Case 1 (M.A.) before the start of insulin therapy and in two healthy infants, one of whom was dysmature. He reports, ‘no insulin was detectable in M.A.’s serum by the rat diaphragm method. The serum was a post-prandial specimen, and serum insulin was by contrast readily detectable in the serum of the two healthy infants, the amount present being comparable to that seen in normal adults. It cannot of course be certain in

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**Table: Idiopathic Hyperglycaemia**

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Reference</th>
<th>Age at Onset (days)</th>
<th>Sex</th>
<th>Birth Weight (kg.)</th>
<th>Maturity (wk.)</th>
<th>Highest Blood Sugar (mg. 100 ml.)</th>
<th>Ketonuria</th>
<th>Associated Condition</th>
<th>Insulin Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Ramsey (1927)</td>
<td>30</td>
<td>M</td>
<td>2:2</td>
<td>40</td>
<td>Nil</td>
<td>Nil</td>
<td>Gangrene</td>
<td>7 days</td>
</tr>
<tr>
<td>2</td>
<td>Lawrence and McCance (1931)</td>
<td>18</td>
<td>F</td>
<td>3:75</td>
<td>40</td>
<td>600</td>
<td>Nil</td>
<td>Abscess surrounding</td>
<td>2 weeks</td>
</tr>
<tr>
<td>3</td>
<td>Strandqvist (1932)</td>
<td>35</td>
<td>M</td>
<td>2:2</td>
<td>40</td>
<td>420</td>
<td>Nil</td>
<td>Pyrexia</td>
<td>6 weeks</td>
</tr>
<tr>
<td>4</td>
<td>Arey (1953)</td>
<td>13</td>
<td>M</td>
<td>2:2</td>
<td>40</td>
<td>555</td>
<td>Nil</td>
<td>'Toil'</td>
<td>Not given</td>
</tr>
<tr>
<td>5</td>
<td>Keidan (1955)</td>
<td>22</td>
<td>F</td>
<td>2:78</td>
<td>40</td>
<td>240</td>
<td>Once</td>
<td>Vomiting, &amp; cerebral</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Nawrocka-Kanska (1952)</td>
<td>12</td>
<td>M</td>
<td>2:7</td>
<td>Not recorded</td>
<td>268</td>
<td>Yes</td>
<td>Haemorrhage</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Engleson and Zetterqvist (1957)</td>
<td>5</td>
<td>M</td>
<td>2:78</td>
<td>40+</td>
<td>720</td>
<td>Nil</td>
<td></td>
<td>3 months</td>
</tr>
<tr>
<td>8</td>
<td>Engleson and Zetterqvist (1957)</td>
<td>11</td>
<td>M</td>
<td>2:0</td>
<td>40</td>
<td>400</td>
<td>Nil</td>
<td></td>
<td>27 weeks</td>
</tr>
<tr>
<td>9</td>
<td>Hutchinson et al. (1962)</td>
<td>7</td>
<td>F</td>
<td>1:93</td>
<td>40</td>
<td>1,292</td>
<td>Nil</td>
<td>Ethisterone effect</td>
<td>41 months</td>
</tr>
<tr>
<td>10</td>
<td>Hutchinson et al. (1962)</td>
<td>17</td>
<td>F</td>
<td>2:4</td>
<td>40</td>
<td>660</td>
<td>Nil</td>
<td></td>
<td>18 months</td>
</tr>
<tr>
<td>11</td>
<td>Hutchinson et al. (1962)</td>
<td>12</td>
<td>F</td>
<td>2:13</td>
<td>38</td>
<td>800</td>
<td>Nil</td>
<td></td>
<td>22 days</td>
</tr>
<tr>
<td>12</td>
<td>Hutchinson et al. (1962)</td>
<td>42</td>
<td>F</td>
<td>1:84</td>
<td>43</td>
<td>750</td>
<td>Nil</td>
<td></td>
<td>2 weeks</td>
</tr>
<tr>
<td>13</td>
<td>Sweetnam and Sykes (1962)</td>
<td>14 or sooner</td>
<td>F</td>
<td>1:54</td>
<td>40</td>
<td>1,200</td>
<td>Mild</td>
<td></td>
<td>8 months</td>
</tr>
<tr>
<td>14</td>
<td>Present paper: Case 1</td>
<td>7</td>
<td>M</td>
<td>1:98</td>
<td>42</td>
<td>1,028</td>
<td>Tr.</td>
<td>Mild gastro-enteritis</td>
<td>At least 15 months</td>
</tr>
<tr>
<td>15</td>
<td>Present paper: Case 2</td>
<td>13</td>
<td>F</td>
<td>3:17</td>
<td>38</td>
<td>980</td>
<td>Yes</td>
<td>Oedema</td>
<td>8 weeks</td>
</tr>
</tbody>
</table>
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these experiments whether insulin was absolutely deficient or was present and masked by antagonistic factors in the serum of the hyperglycaemic child.

The studies in Case 2 (E.M.S.) were performed by Dr. Ellis Samols by immuno assay. Very high levels of insulin activity were demonstrated. The fasting level of 165 units three days after withdrawal of insulin treatment, fell to 80 units a week later. Remaining exogenous insulin may have contributed to this and the interpretation of the results was further complicated by the presence of insulin antibodies resulting from previous therapy.

However, Dr. Samols thought it fair to conclude that: `there was a rise in insulin after tolbutamide and there was a rise in insulin after glucose.'

Though our findings do not shed much light on the basic lesion, the following points emerge. The failure to demonstrate insulin activity in the serum of Case 1 at 6 weeks may be taken as evidence either of defective insulin production or of the presence of insulin antagonists. The response to intravenous tolbutamide, i.e. the fall of blood sugar in both infants and the rise of insulin activity in Case 2, suggests either recovery of pancreatic function or a decrease of insulin antagonists; Case 2 had at this stage recovered clinically.

Following the intravenous tolbutamide test in Case 1 the possibility of treatment with tolbutamide was considered. It seemed, however, that if a gradual recovery of islet cell function were taking place, stimulation of insulin release might 'exhaust' the pancreas and hinder recovery. Treatment with tolbutamide failed in one of Hutchison's cases.

TREATMENT AND SEQUELAE. Keidan (1955) drew attention to the low insulin requirements of these babies compared with those in true diabetes mellitus and doubted whether insulin was necessary. However, neither of our cases thrived without it and the effect of withholding it in Case 1 at the age of 4 months clearly demonstrated the need for its continued use. The mild episodes of ketosis in Case 2 were taken as an indication for increasing the dose of insulin to which they responded.

Hutchison et al. were the first to describe permanent sequelae in these infants, but few other authors have reported a comparable follow-up. Hypoglycaemia in the newborn period is known to be a cause of permanent brain damage (Brown and Wallis, 1963; Neligan, Robson, and Watson, 1963). As Sweetnam and Sykes state, hypoglycaemia at any time during the stage of rapid brain growth is likely to be particularly dangerous in this respect. Hutchison et al. considered that hypoglycaemic insulin reactions were responsible for the mental retardation of two of their cases, though they do not accept the possibility in their third defective child.

Whether or not hypoglycaemia resulting from the too liberal use of insulin is the only cause of brain damage it seems imperative to avoid it. We think that treatment should be aimed at the maintenance of a normal rate of growth rather than the elimination of hyperglycaemia and glycosuria. Neither of these two cases developed the cushingoid appearance reported by Hutchison et al. and attributed by these workers to a growth hormone effect, as part of a basic hormonal defect in these infants. We think it is more likely due to the effect of excessive insulin.

Assessment of pancreatic islet cell function by serum insulin assays and intravenous tolbutamide tests at the onset and during the course of the illness would be valuable. We can offer no explanation for the apparently anomalous metabolic situation in which there is a failure of carbohdrate utilization without evidence of accumulation of the products of fat katabolism as seen in true diabetes mellitus. Perhaps this is in some way related to the relative absence of subcutaneous fat in babies suffering intrauterine starvation. Against this is the fact that four of the infants were not thin at birth. We are intrigued by the comparison between these infants and newborns suffering from idiopathic hypoglycaemia, who also frequently show evidence of intrauterine starvation.

Summary

Two cases of idiopathic neonatal hyperglycaemia are presented together with studies of plasma insulin activity, and a brief review of 13 previously reported cases is given. A clinical syndrome affecting mainly immature infants emerges.

The insulin studies suggest islet-cell failure or the presence of insulin antagonists rather than failure of 'end-organ responsiveness'.

Note: Case 2 was shown at the Royal Society of Medicine, Section of Paediatrics Meeting, March 22, 1963, and an abridged account was published in the Proc. roy. Soc. Med. (Mortimer, 1963).

We are indebted to Dr. Mary J. Wilmers and Dr. Richard H. Dobbs for permission to publish these cases and for their valuable help and criticisms. We would like to thank Dr. Keith Taylor, King's College Hospital and Dr. E. Samols, Royal Free Hospital, for the insulin studies, and the House Officers and Nursing Staff who were so conscientious in the care of these babies and in the collection of specimens.

REFERENCES


Addendum.

Case 1, at the age of 27 months, still requires a small dose of insulin. He remains free from ketosis and is making satisfactory progress. Further investigations are planned.