HYPOGLYCAEMIC COMA

BY

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Symptoms of hypoglycaemia have only been recognized since the introduction of insulin. Since that time they have been noted in connexion with syndromes other than the over-production and overdosage of insulin, and the symptom-complex has become a well recognized entity. Most of these non-pancreatic causes of hypoglycaemia do not lead to coma. A low blood sugar level may be found in Addison’s disease, Simmonds’ disease and myxoedema and in conditions of chronic hepatic insufficiency, in malnutrition, and following prolonged, severe exercise. In von Gierke’s disease there is a lowered blood sugar level, although the patient rarely complains of symptoms relative to hypoglycaemia.

The pancreatic causes of hypoglycaemia are (a) tumours of the islet tissue, and (b) the condition of hyperinsulinism; the hypoglycaemia in both these conditions may be sufficiently severe to precipitate the patient into coma. During the stabilization and maintenance of insulin therapy such episodes often occur and occasionally may lead to crippling mental or physical sequelae.

Finally, overdosage of insulin will cause hypoglycaemia as exemplified in our case which is reported here.

Case Report

L.W., a white child aged 2 years and 1 month, was admitted to coma to the Transvaal Memorial Hospital for Children on April 30, 1953. The child was a diabetic and was well controlled on 10 to 15 units of N.P.H. insulin each morning. Since her discharge from hospital six weeks before, when her insulin dosage had been stabilized, she had developed and recovered from a mild diarrhoea and upper respiratory tract infection.

The mother stated that she had felt that the child would be cured of her diabetes if she remained sugar-free for a period of time. She had consequently increased the insulin dosage above the prescribed level and had at the same time restricted the child’s diet. The child had been irritable for some time and had tended to become inattentive and ‘dreamy’; the mother had also noted that this was worse before meals. This querulousness and inability to concentrate had become worse in the few days before admission, although it was not attended by signs other than occasional twitching of the left hand for three days before admission. This state of affairs culminated in the parents finding the child unconscious early one morning. Before admission to this hospital the child had been unconscious for 17 hours in another hospital where initially insulin had been given and intravenous glucose administered later, but no control blood sugar estimations done. The child had been energetically treated for ‘shock’, this including the application of hot-water bottles to secure warmth.

On examination the child was deeply comatose with involuntary movements of the left face, arm and leg. The face and hand twitched continuously, whilst the leg was less affected. The urine was sugar-free.

All other systems were clear.

Examination of the cerebrospinal fluid excluded other central nervous system conditions, such as encephalitis. The physical findings, combined with such a definite history of insulin overdosage, were so typical that a diagnosis of hypoglycaemic coma was made, despite the finding of a normal blood sugar level after admission, which it was felt, was due to the intravenous administration of glucose before admission.

A full blood examination on the day of admission gave the following results: Haemoglobin 15 g., erythrocytes 5,000,000, leucocytes 14,500, blood sugar level 202 mg.%, serum potassium level 19·2 mg.%, and serum sodium level 283 mg.%. Four hours after admission the blood sugar level was 324 mg.%, and 12 hours after admission it had fallen to 122 mg.%. On May 1 the cerebrospinal fluid was examined, when no cells were seen, protein was 10 mg.%, chloride 700 mg. % (as NaCl) and sugar 66 mg. %.

On admission an intravenous infusion of isotonic saline was begun and 40 ml. of 50% solution of glucose administered intravenously. On receipt of the laboratory reports insulin was also given. Insulin dosage was guided by the amount of sugar in the urine, tested four-hourly. During the first 60 hours a total of 500 ml. of normal saline and 1,000 ml. of 5% dextrose in normal saline and 2,000 ml. of 5% dextrose in water was given (Fig. 1). Vitamin B1, 1,000 mg., and vitamin C, 100 mg., were added daily to the infusion.

On the third day cortisone therapy was instituted in
HYPOGLYCAEMIC COMA

TABLE 1
 DETAILS OF TREATMENT

<table>
<thead>
<tr>
<th>Day</th>
<th>Insulin (units)</th>
<th>Cortisone (mg.)</th>
<th>Vitamins (units)</th>
<th>Penicillin (units)</th>
<th>Sedation</th>
<th>Fluids (ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10</td>
<td>Nil</td>
<td>Nil</td>
<td>400,000</td>
<td>Nil</td>
<td>500 normal saline</td>
</tr>
<tr>
<td>2</td>
<td>10</td>
<td>Nil</td>
<td>1,000 Vit. B, 100 Vit. C</td>
<td>do.</td>
<td>do.</td>
<td>1,000</td>
</tr>
<tr>
<td>3</td>
<td>5</td>
<td>50</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
<td>2,500</td>
</tr>
<tr>
<td>4</td>
<td>26</td>
<td>50</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
<td>2,000</td>
</tr>
<tr>
<td>5</td>
<td>16</td>
<td>40</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
</tr>
<tr>
<td>6</td>
<td>16</td>
<td>30</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
</tr>
<tr>
<td>7</td>
<td>22</td>
<td>20</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
<td>do.</td>
</tr>
</tbody>
</table>

view of its glycogenic properties, and 50 mg. was given daily for two days and then decreased by 10 mg. daily (Table 1).

From the sixth day the level of consciousness improved and the jerking grew less. The child at this stage responded to painful stimuli. By the eighth day she was fully conscious, but extremely irritable and still manifested occasional twitching of the left arm, hand and leg. It was noted at this time that the patient could not use the left arm and that the left leg was weak.

The child remained in hospital for a further three months during which time a gradual improvement in the left facial palsy and the left leg were noted. The left arm, however, did not improve as much and on her discharge from hospital she still had a considerable paralysis of this limb. During this time the left arm and leg twitched almost daily, but no loss of consciousness was associated.

An air-encephalogram (Fig. 2) was performed on July 27 and Dr. H. Jackson reported that there was dilatation of the right lateral ventricle with displacement to the right. This suggested right cerebral atrophy.

The child has now been followed for 18 months since admission, and this period has seen worsening of the fits. At first they took the form of jerks of the arm, but gradually she started to manifest absences and about 12 months after admission to hospital, akinetic seizures began to make their appearance. Major epilepsy is also present, but these attacks are not as frequent. Attacks of one or other type of epilepsy occur up to 20 times a day, the child falls frequently, and is never without evidences of trauma.

She appears lethargic and the mother has noted that she cerebrates slowly and has learned very few new words since her discharge from hospital. During the early phase of recovery she spoke very little. The mother feels that there is definite mental retardation. The left hemiplegia remains stationary and there is also a degree of dysarthria which shows itself in a thick, slurred speech.

The diabetes remains as before, but she is reasonably well controlled on 10 units crystalline insulin and 10 units P.Z. insulin each morning combined with a free diet.

A Merrill-Palmer intelligence test was given and the following results were obtained:

- Chronological age . . . . 3 years 8 months
- Mental age . . . . . . . . 2 years 3 months
- I.Q. . . . . . . . . . . . . 61

During the test the child was extremely distractable; she was unable to give her full attention to the test. When presented with a test item, she was vague as to
what was required of her. She had a low frustration
tolerance and gave up easily. She worked for a short
time with a test item, and, being faced with apparent
failure, she appeared to become irritated by her inability
to succeed. This was followed by a short temper
tantrum, during which she pushed the items off the table
on to the floor.

She reacted indifferently to praise. When she wanted
to stop working on a test item, no amount of praise or
courage was effective in keeping her at her task.

She was able to name objects, but was unable to explain
their use. Vocabulary was extremely limited, spontane-
ous speech consisting of one or two-word sentences,
only unintelligible. When asked to repeat sentences,
responses consisted of ‘unintelligible baby talk’.

An E.E.G. (Fig. 3) was done on October 13 and Dr.
M. K. Wright reported that there was an area of
diminished amplitude over the right central and parietal
areas, and also a definite spike focus on the convexity of
the left frontal lobe. Such spike foci commonly shift
in young children. The E.E.G. suggested that there
had been rather diffuse cortical damage rather than
a localized lesion.

Discussion

The morbid anatomical findings following hypo-
glycaemic coma may be widespread, and Lawrence,
Meyer and Nevin (1942) have indicated several
different pathological changes. There may be
complete necrosis of nerve cells over areas of the
cerebral cortex and these areas may be remarkably
circumscribed. In addition to the necrotic lesions
there are areas of lesser damage in which cells are not
completely destroyed. There may also be gross
vascular lesions in the cortex and basal ganglia.
Lawrence et al. (1942) compare these changes to the
damage in the central nervous system following severe
anoxaemia due to causes as dissimilar as
poisoning with cyanide, carbon monoxide and ether,
status epilepticus and cardiac arrest during anaes-
thesia. The two cases of Roche (1942), both of
which came to necropsy, had cerebral changes; the
first showed perivascular softening in the thalamus,
caudate and lenticular nuclei, and the second a
generalized softening, especially in the cerebellum.
Hicks (1950), in animal experiments, showed that
the administration of insulin to rats in dosages
sufficient to cause convulsions often led to destruc-
tion of nerve cells of the cerebral cortex and corpus
striatum; the basal nuclei were occasionally affected.
Grayzel (1934) in experiments performed on rabbits
found that in hypoglycaemic animals in which
convulsions occurred brain damage was marked,
whereas in rabbits in which the dose of insulin was
insufficient to cause convulsions minimal or no
microscopic evidence of necrobiosis was found.
The brain damage in these patients is due to the fact
that nerve cells require glucose for their oxidative
processes. The cells, however, carry a very small
reserve of glucose and have to replenish these
stocks from the blood glucose. Thus, in cases in
which the blood sugar level falls too low the cell
metabolism becomes grossly impaired, resulting in
severe damage to or actual death of the cell. The
failure of this essential oxidative process leads to the
final picture which is similar to that found in anoxic
damage to the brain.

Tyler (1941), working with cats, found that the
higher the dose of insulin the greater the brain
damage, and that the giving of phenobarbitone
tended to ameliorate the symptoms. In addition,
cats at a low temperature tended to receive less
damage to the central nervous system. Cerebral
metabolism is depressed by both hypothermia and
by the administration of barbiturates, leading to a
lessened oxygen uptake, and therefore a lessened
need of oxygen, which is obviously desirable in hypoglycaemia. The higher the dosage of insulin in these animals, the more protracted was the fall in blood sugar and the greater was the amount of glucose required to terminate the hypoglycaemia.

The symptoms of hypoglycaemia vary somewhat with the type of insulin used, but are basically the same for both short- and long-acting insulins. They commonly appear several hours after the administration of crystalline insulin, but may occur much sooner if the patient has taken no food or has had to take violent exercise in the interim. The long-acting insulins—globin insulin, N.P.H., P.Z. insulin and the 'lente' insulins—have their maximum effects later, the symptoms are less spectacular and the patient's attention is not drawn to them so readily. He slips quietly and unobtrusively into a stuporous state. The response to the administration of glucose by mouth or by intravenous injection is usually quick. It should, however, be remembered that long-acting insulins continue to act after the blood sugar has been restored to normal and that frequent doses of sugar must be administered.

The difficulty encountered in our case in terminating the coma, despite adequate treatment, has been recorded elsewhere (Klein and Ligterink, 1940). The longer the patient has been in coma the more difficult it is to terminate the condition (Joslin, Root, White and Marble, 1952). Lawrence et al. (1942) state that coma of up to three hours is usually associated with complete recovery, while coma of longer duration is dangerous. Layne and Baker (1939) make the observation that the return of the blood sugar to normal has little effect on the duration of coma. It is important to maintain a normal or rather high blood sugar level during the state of recovery. In our case the observations of Klein and Ligterink (1940), Layne and Baker (1939) and Joslin et al. (1952) are well brought out; this child had a raised blood sugar level on admission to this hospital after having had intravenous glucose in the outside hospital. This is also in accord with the experimental evidence of Tyler (1941), who found that many of the cats used in his research work remained comatose for several days despite a normal blood sugar level.

Hypoglycaemic coma with cerebral damage may lead to progressive mental deterioration and to organic brain damage which may manifest itself clinically as paralysis or epilepsy. Anderson (1940) reports two cases. In one the child's mental age after the onset of coma at 13 years was 5½ years, and in the other child the mental age at 7 did not exceed 4 years. In the first case the child's memory improved, but the second patient had to be placed in an institution. Allan and Crommelin (1942) report a child, aged 6 years, in whom some degree of mental dullness remained months later. In several cases of hypoglycaemic coma reported by Graham (1950) (all adults) the patients had mental changes of a degenerative nature.

Klein and Ligterink (1940) report two cases of hypoglycaemic coma followed by mental retardation. Gardner and Reyersbach (1951) record a case of progressive mental deterioration following brain damage, despite the avoidance of further hypoglycaemia. Murphy and Purtell (1943) in reviewing 26 cases from the literature, eight of which died, record six instances of mental defect, and a further six cases described as personality changes, mental changes, organic psychosis, mental confusion and 'mental invalid'. Two children, one aged 8 and one aged 15, in a series of seven cases reported by Layne and Baker (1939) both showed mental retardation. The older child manifested epilepsy and was ataxic. Several of McQuarrie's (1954) cases of convulsions and coma due to spontaneous hypoglycaemia are mentally retarded and show damage in the central nervous system.

The organic changes may result in paralysis or epileptic seizures, and in all the cases quoted above either organic changes or convulsions occurred. Several cases in the literature, quoted by Murphy and Purtell (1943), were aphasic either permanently or temporarily, but it was not stated whether this was due to mental change or brain damage. In their own case they report mental change and temporary paralysis and the aphasia was obviously due to the mental condition.

The case of Gardner and Reyersbach showed a progressive deterioration in the E.E.G. and that child had convulsions which were extremely difficult to control; in addition, air studies showed definite enlargement of the lateral and third ventricles. In electroencephalograms done on 35 diabetic cases with repeated insulin reactions by Greenblatt, Murray and Root (1946) there were 18 definitely abnormal, nine borderline and eight normal. In our case the E.E.G. was abnormal and an air study showed enlargement of the right lateral ventricle.

The present state of our patient fits in well with the description of mental retardation, frequent epileptiform seizures and hemiplegia referred to by various authors. The convulsions as exhibited in our case are extremely difficult to control and consist of grand mal seizures, petit mal absences and akinetic attacks. The child's response to all forms of medication has been poor. The trigger mechanism, according to the parents, may be either hypoglycaemia or acidosis. Engel, Halberg, Ziegler and
McQuarrie (1952), in very complete studies on two children with epilepsy following hypoglycaemic coma, found that no correlation could be determined between seizures and the blood sugar level. These authors showed that there is, however, less association of the spike-and-wave activity at 2½ per sec. with hypoglycaemia than with hyperglycaemia.

McQuarrie (1954) treated 25 cases of spontaneous hypoglycaemia with cortisone or corticotrophin with very good results. This was also done in our case, and in our opinion should be part of the treatment of similar patients in future, particularly in those cases caused by long-acting insulins.

In the light of the experimental work of Tyler (1941) these comatose patients should perhaps be sedated and subjected to hypothermia. The practical application and degree of hypothermia remain to be determined. It would appear physiological not to warm the patient, and it is possible that the warmth applied to our patient before her transfer to this hospital may have contributed towards the noxious effects of the hypoglycaemic state. In view of recent work on hypothermia during cardiac surgery, this treatment may possibly be tried in future cases. It must be noted that the cats in Tyler’s experiments were not ‘cooled’, they were merely not ‘warmed’. There was a drop in temperature of about 6°C in these animals by the time coma had set in, and this appeared sufficient to ward off a large degree of brain damage.

Summary
A case of hypoglaemic coma is recorded and the pathology discussed. The clinical observation is made that coma may persist for many days despite the restoration and maintenance of a normal blood sugar level. The final state of severe brain damage manifesting itself as convulsive episodes, mental defect and hemiplegia is described.

The literature is briefly discussed.

The suggestion is made that cortisone, sedation and hypothermia may be useful adjuncts in the therapy of this condition.

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References