Short report

Status epilepticus, hypothermia and metabolic chaos in a man with agenesis of the corpus callosum

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SUMMARY A case of hypothermia and metabolic derangement in a 60-year-old man following a period of status epilepticus is reported. A CT head scan performed subsequently demonstrated agenesis of the corpus callosum. Hypercalcaemia which has not been reported before was a prominent abnormality in the metabolic profile.

In 1969, Shapiro et al described an association in two patients between spontaneous attacks of hypothermia and agenesis of the corpus callosum. This association has been reported subsequently in eight other cases, of which were intensively investigated both during, and in between, attacks of hypothermia. It is accepted that agenesis of the corpus callosum does not, of itself, cause the disturbance of temperature control, but there are always associated developmental abnormalities involving midline structures which presumably interfere with temperature control mechanisms. In the one pathological study adequately reported, fibrosis of the anterior hypothalamus with neuronal loss was described, and experimental studies have suggested that this structure may be concerned with heat-dissipating mechanisms. In these cases, the episodes of hypothermia have varied from a few minutes or hours, to more than one week in duration, and there has been much discussion as to whether they could be epileptic events. In the early reports, this was assumed to be the case, and the name diencephalic epilepsy was applied to such cases. Some seemed to respond to anticonvulsants, but in more recently studied cases, these were ineffective, and other mechanisms were suggested. Profound metabolic and haematological disturbances have been reported in many of the patients usually occurring at the time of the hypothermic episodes. No mechanism for these has been proposed, and investigations after recovery have failed to show persisting abnormalities.

We report a man with agenesis of the corpus callosum who developed a single episode of hypothermia with a variety of metabolic disorders during a period of status epilepticus, after which he recovered. He had never had disturbances of temperature control before, even in association with a previous episode of status epilepticus.

Case report

A 60-year-old man was admitted to hospital following a fall which caused a fracture of his left scapula. He did not injure his head or lose consciousness. He had suffered from generalised convulsions since the age of eleven years, but these were well controlled on a combined preparation of phenobarbitone 50 mg and phenytoin 100 mg thrice daily. His medical records since 1948 revealed only a single episode of status epilepticus in 1979, which necessitated hospital admission for 48 hours. On that occasion, he was found to have an extensor plantar response on the right, and investigations revealed a normal temperature, normal T4 level and a normochromic normocytic anaemia (Hb = 11.6 g/dl) for which no cause was determined. When reviewed subsequently he was fully recovered. He was said to have always been of low intelligence but had worked conscientiously all his life, running his own successful greengrocery business. After six days in hospital, he appeared to be recovering from his fall, and investigations showed satisfactory anticonvulsant and T4 levels. He then developed brief generalised seizures, often associated with rigidity, which became more frequent. Clinical examination demonstrated persistent signs of a right hemiparesis and a right extensor plantar response. Eventually, these
attacks became almost continuous over a 24 hour period, and failed to respond to intravenous diazepam or chloromethiazole. In association with this, his interictal conscious level deteriorated such that he was not responding to painful stimuli.

At this stage he was transferred for neurosurgical
assessment at the Middlesex hospital. On arrival there, fourteen days after the fall, he was noted to be normoten-
ve with a sinus tachycardia, but within an hour he had
deteriorated further and became unresponsive, cold (rectal
temperature = 29°C), pale and hypotensive (systolic blood
pressure = 30–50 mm Hg) with a sinus bradycardia (40–
60 b.p.m.). His respiratory effort was inadequate and he
was intubated and artificially ventilated. On examination at
this stage, the eyes were in the primary position, with no
reflex movements, and with 2 mm pupils showing a poor
response to light. There was twitching of the tongue and
frequent jerking of the right arm and leg. The limbs were
flaccid with no response to painful stimuli and the reflexes
were exaggerated on the right with bilateral extensor plantar
responses. There was no clinical evidence of blood loss.
Investigations on admission showed anaemia (Hb =
9·0 g/dl, PCV = 0·28%, WCC = 4·6 × 10⁹/l) with ureaemia
(urea = 29·5 mmol/l, creatinine = 184 mol/l, sodium =
150 mmol/l and normal potassium). Calcium (corrected) =
2·96 mmol/l and albumin = 19 g/l. Serum phosphate,
alkaline phosphatase, folate, serum iron, total iron binding
capacity and immunoglobulins were normal. The aspartate
aminotransferase level was initially elevated 484 iu/l (nor-
mally less than 40 iu/l), and rapidly fell to normal after
which there was a transient rise in the alkaline phosphatase
level to a maximum of 448 iu/l (normal range = 100–
280 iu/l). Phenobarboline and phenytoin levels were
within the therapeutic range. The initial endocrine inves-
tigations showed that the T4 level had fallen to 41 nmol/l
from the normal level just after his fall but the TSH was
not elevated at 3·8 mU/l. Subsequent investigation of the
hypothalamic–pituitary axis after neurological improve-
ment showed normal T4 and T3 levels (T4 = 74 mmol/l T3
= 21 nmol/l) and a normal TRH test, LH, FSH, LH-RH
test, prolactin, cortisol, testosterone, magnesium, para-
thyroid hormone and oral glucose tolerance test (OGTT).
Basal growth hormone was normal (5·3 mU/l) but failed to
be suppressed during OGTT on two occasions.

Plain radiographs were normal but a CT scan dem-
strated agenesis of the corpus callosum (fig). EEG showed
frank seizure activity with evidence of a left sided origin.
Cerebrospinal fluid examination contained a total protein
of 0·8 g/l but was otherwise normal.

His hypotension responded dramatically to a volume
load of 5 l of colloid (initial CVP—5), his hypotension to
gradual rewarming and his status epilepticus to intravenous
phenytoin and a thiopentone infusion titrated with a
cerebral function monitor. His haemoglobin level stabil-
ised at 6·6 g/l following the colloid load and required a 4
unit transfusion. A bone marrow aspirate was hypocellular,
with suppression particularly of erythropoiesis. Over the
next two days his condition improved and his temperature
rose to normal. His serum calcium (corrected) fell slowly
from its peak of 2·96 mmol/l to a stable value of
2·30 mmol/l accompanied by a gradual rise in the albumin
to 32 g/l. All other haematological and biochemical
abnormalities corrected. However despite intravenous and
nasogastric feeding, he showed loss of muscle bulk, and,
when testing became possible, he had generalised weak-
ness without loss of reflexes. EMG showed small
polyphasic potentials consistent with a diffuse myopathy.

There was no radiographical evidence of metabolic bone
disease. One month after the fall, he was well enough to be
discharged to the referring hospital, and after a period of
further physiotherapy, he was able to go home with a mild
right hemiparesis and moderate proximal myopathy. On
outpatient review one year later, he had no neurological
abnormalities and no haematological or metabolic abnor-
malities. He remains on anticonvulsants.

Discussion

This patient developed hypothermia and widespread
metabolic and haematological disturbances in associa-
tion with a period of status epilepticus follow-
fing a fall. His fits were reasonably well controlled
before and after this episode and he had previously
had an episode of status epilepticus without tempera-
ture or metabolic disturbance. No precipitating
cause for this episode is apparent. As the fits became
more frequent his general condition and conscious
level deteriorated, and it is impossible to know
whether this was due to the frequent fits, to the
drugs which were given to attempt to control the fits,
or to a developing metabolic disturbance. After
transfer to the Middlesex Hospital, he was found to
have a low body temperature and deranged
metabolism. Of particular interest is the initially
high serum calcium in the presence of a low album-
min. The calcium samples were all taken in the fast-
ing state with venous occlusion. The serum mag-
nesium and parathyroid hormone levels were nor-
mal. In none of the previously recorded cases of
hypothermia with agenesis of the corpus callosum
has an abnormal calcium level been observed.
Although at the same time, there was some elevation of the
plasma sodium and urea, which undoubtedly was
due to fluid depletion. Hyponatraemia occurred in
two cases but the sodium was normal in other cases.
Low serum albumin has not been reported before.
Parallel with the metabolic disturbance and recovery,
there was the development of anaemia which appears to have resulted from suppression of
erythropoiesis in the marrow. Anaemia, usually
with pancytopenia, has occurred in three previously
reported cases.¹ ² ³ It is of interest that, in this
patient, a low haemoglobin had been recorded at the
time of status epilepticus three years previously,
with subsequent recovery to normal, but no further
details of this are available. On this occasion, the
haemoglobin fell considerably and was accompanied
by a low white blood cell count and platelet count.
Endocrinological studies in this, as in other patients
in whom hypothermia occurred in association with
agenesis of the corpus callosum, did not show any
striking abnormality.

The first thyroxine level was marginally low at a
time when the albumin level was low and the total
globulin level was not measured. Subsequently, all studies were normal apart from the failure of growth hormone suppression during an oral glucose tolerance test.

The development of a myopathy has not been described previously as part of this syndrome. Its origin is also obscure, but it could perhaps have resulted from the effects of the metabolic disturbances upon muscle. Like everything else, this recovered, although rather more slowly. Finally this case differs from those previously described in that hypothermia appears to have occurred only once in the patient's life. Much attention has been paid in previous reports to the origin of the hypothermia and, in particular, whether it could result from epileptic activity. The majority of writers have thought that this is unlikely, and, in this case, the fact that it had not occurred during a previous episode of generalised status epilepticus is perhaps also against this. However, recovery occurred when the fits were terminated, although ictal activity was still found on EEG and observed clinically during the recovery phase. The temperature control system is thought to be affected as a result of abnormal development of the anterior hypothalamus, which was found in one case where an adequate autopsy was performed. It is suggested that the thermostat is incorrectly set so that heat losing mechanisms are inappropriately deployed. There are no data available which would elucidate the reason for this, or explain why, in some cases, it tends to occur intermittently, with periods of normal temperature control in between. The effects of raising the body temperature from the hypothermic level using pyrogens or air-conditioned beds has been described in some cases. In most patients attempts at rewarming caused inappropriate sweating until the body temperature fell again to hypothermic levels. In our case, inappropriate sweating was not observed and the temperature rose eventuantly to normal using rewarming methods.

The variety of disturbances in this patient emphasises the dependence of many homeostatic mechanisms on intact cerebral function and, in particular, on function of the hypothalamus.

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References