Missed Diagnosis

Missed Addisonian crisis in surgical wards

M. Small, A.C. MacCuish and J.A. Thomson

University Department of Medicine, and Diabetic Unit, Royal Infirmary, 10 Alexandra Parade, Glasgow G31 2ER, UK.

Summary: Three women were admitted to the surgical wards with acute gastro-intestinal symptoms and despite good clinical (pigmentation, hypotension) and biochemical (hyponatraemia, acidosis) evidence of Addisonian crisis had unnecessary investigations with delay in the diagnosis and appropriate management of this medical emergency. Clinicians should take careful note of the typical electrolyte upset seen in Addisonian crisis.

Introduction

Although Addison’s disease is an uncommon condition with a prevalence in the United Kingdom of 39/million, the clinical and biochemical presentation of acute adrenal insufficiency with predominant gastrointestinal symptomatology is well known to undergraduate medical students. We report 3 patients who, over a 3 year period, were admitted to the surgical wards in an Addisonian crisis where a significant delay was noted in making the correct diagnosis.

Case reports

Case 1

A 49 year old post-menopausal female was admitted to the general surgical ward in 1985 with a one week history of epigastric pain and vomiting and a six week history of anorexia with weight loss of approximately 8 pounds. In her past medical history she was diagnosed to have mild type 2 (non-insulin dependent) diabetes in 1984 and follow-up at the diabetic clinic had revealed reasonable glycaemic control on dietary measures alone. In 1981 she had presented to the medical wards with haemoptysis, at which time bronchoscopy was negative. At that time she was noted to have facial plethora, central obesity and to be mildly hypertensive, blood pressure (BP) 150/100 mmHg, but urinary free cortisol levels were normal. At the time of her current admission she was obese and unwell with a BP of 70/0 mmHg, pulse rate 120/min and epigastric tenderness. Chest X-ray, abdominal X-ray and serum amylase were normal, the plasma glucose was 12 mmol/l and urea and electrolytes are listed in Table I. Urinalysis showed moderate ketonuria. The patient was thought to have a peptic ulcer and was treated with intravenous cimetidine and fluids. Following 500 ml of 0.9% saline and 500 ml plasma the BP rose to 100/60 mmHg and the patient was given 9 litres of fluid over the subsequent 36 hours. The patient improved following her rehydration, the ketonuria disappeared and gastroscopy, barium enema and ultrasound of the abdomen failed to reveal a cause for her symptoms. Seven days after admission the patient was well and was allowed home. Four days later however she was re-admitted with the same symptoms, haemodynamic and electrolyte upset (Table I). The arterial pH of 7.30 with a base excess of -14 confirmed metabolic acidosis. The blood glucose was 6.5 mmol/l and again there was moderate ketonuria. Abdominal X-rays were again negative. The patient was treated with intravenous saline and sodium bicarbonate and 2 days later a medical opinion was sought. The hypotension plus electrolyte disturbance suggested a diagnosis of Addison’s disease which was later confirmed.

Case 2

A 40 year old woman was admitted to the surgical ward with a 6 month history of anorexia, nausea and extreme lethargy and in the three months before admission lost over 1 stone in weight. The patient also gave a 10
month history of oligomenorrhoea. Apart from pig-
mentation examination was unremarkable. The
patient was thought to have a peptic ulcer, and in view
of her dehydration and hyponatraemia (Table I) was
given 7 litres of 0.9% saline over the following 48
hours. She improved clinically following re-hydration
although the BP had fallen and remained constant at
90/50 mmHg. Barium meal and gastroscopy were both
normal. On the day of her planned discharge, 8 days
into her hospital admission, her urea and electrolytes
were again found to be abnormal (Table I). The
surgical house officer on further enquiry obtained the
history that the patient’s suntan had not faded follow-
ing her summer vacation and in fact for cosmetic
reasons, she had tried to bleach away the heavy
pigmentation in both palmar creases. The house
officer suspected Addison’s disease which was later
confirmed. The patient was transferred to the medical
ward, and, following the introduction of cortico-
steroids and further intravenous fluids made an
uneventful recovery.

Case 3

A 17 year old girl was admitted to the surgical wards
with a 2 day history of constant lower abdominal pain,
vomiting and sore throat. On examination she was
noted to have cervical lymphadenopathy, a pulse rate
of 130/min and BP of 94/46 mmHg. The abdomen was
mildly tender in the right iliac fossa. The urea and
electrolytes confirmed dehydration but no comment
was made on the other electrolyte disturbance (Table
I). The patient was thought to have a viral illness and
was treated conservatively with parenteral antimi-
etics and analgesics and over the following 2 days
received 7 litres of intravenous fluids, mainly in the
form of 0.9% saline. She made an uneventful recovery
to be discharged 4 days later.

Five months later she was admitted to the medical
wards with a similar history of abdominal pain and
vomiting and identical haemodynamic upset. She was
noted to be pigmented and stated that she had been
able to maintain her suntan from 9 months previously
and that this was unusual. Her friends had also
commented on her pigmentation. The urea and elec-
trolytes were similar to her previous admission (Table
I) and venous gases confirmed a metabolic acidoses
with a pH of 7.28 and base excess − 8. The diagnosis
of Addison’s disease was confirmed and the patient made
an uneventful recovery following intravenous saline and
corticosteroid administration.

In each of the above 3 cases the diagnosis of Addison’s
disease was made on undetectable levels of cortisol
which failed to increase following short and depot
tetracosactrin (Synacthen) test. No patient had
evidence of tuberculosis or adrenal calcification on X-
ray and subsequently adrenal antibodies were detected
in all 3 patients. During the period of rehydration and
prior to the patients being given corticosteroid therapy,
each patient was noted to have mild hypercal-
caemia with adjusted calcium values ranging between
2.7–2.9 mmol/l.

Discussion

These 3 patients were each admitted to the surgical wards
in an Addisonian crisis, because of acute
gastrointestinal symptoms, despite the well recognized
clinical and biochemical features of Addison’s disease
being present in each case. Failure to make the correct
diagnosis led to unnecessary gastrointestinal inves-
tigation and indeed two of the patients were dischar-
ged from hospital without the diagnosis of Addison’s
disease even being contemplated, following clinical
recovery after treatment with saline rehydration.
Although the first case required re-admission 4 days
later, in case 3 it appears likely that a viral illness had
precipitated a relative cortisol deficiency and that the
patient had sufficient cortisol reserve to remain symp-
tom-free for a further 5 months before being admitted.

<table>
<thead>
<tr>
<th>Case 1</th>
<th>Na⁺ (135–145)</th>
<th>K⁺ (3.5–5.0)</th>
<th>HCO₃⁻ (23–30)</th>
<th>Urea (2.5–8.0)</th>
<th>Anion gap (&lt;13)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st admission</td>
<td>122</td>
<td>4.8</td>
<td>14</td>
<td>9.7</td>
<td>13</td>
</tr>
<tr>
<td>2nd admission</td>
<td>126</td>
<td>5.3</td>
<td>13</td>
<td>9.9</td>
<td>22</td>
</tr>
<tr>
<td>Case 2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>On admission</td>
<td>133</td>
<td>—</td>
<td>21</td>
<td>11.1</td>
<td>13</td>
</tr>
<tr>
<td>Day 8</td>
<td>132</td>
<td>5.1</td>
<td>24</td>
<td>12.6</td>
<td>13</td>
</tr>
<tr>
<td>Case 3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st admission</td>
<td>124</td>
<td>5</td>
<td>13</td>
<td>11.5</td>
<td>18</td>
</tr>
<tr>
<td>2nd admission</td>
<td>116</td>
<td>5.9</td>
<td>13</td>
<td>11.6</td>
<td>18</td>
</tr>
</tbody>
</table>

Values are mmol/l
in a similar Addisonian crisis. There is approximately a five-fold excess prevalence of Addison’s disease in diabetes compared to the general population.¹⁻³ In diabetic patients the development of Addison’s disease is usually associated with insulin-dependent diabetes and if such diabetic patients are treated with diet or oral hypoglycaemic agents they often progress to ketosis prone insulin-dependent diabetes;⁴ thus close follow-up of such patients is required.

Our cases also illustrate two other biochemical abnormalities seen in Addisonian crisis, namely metabolic acidosis and hypercalcaemia. The metabolic acidosis is poorly described but it is said that bicarbonate levels of 15–20 mmol/l are commonly seen in Addisonian crisis.⁵ Our 2 patients who complained of abdominal pain (cases 1 and 3) had a more severe acidosis with bicarbonate levels of 13 mmol/l which occurred despite vomiting with loss of acid from the stomach. The production of the acidosis is related to a renal tubular acidosis with reduced ammonia production due to the hypercalcaemia.⁶ In addition the hypotension and hypovolaemia may have resulted in tissue hypoxia with an increase in unmeasured acids, as suggested by the increased anion gap (see Table I).

The first patient presented a more complex metabolic upset – she was an obese, diet-controlled diabetic with anorexia, vomiting, moderate ketonuria, and acidosis despite a relatively normal glucose level. The normoglycaemia and extremely low sodium concentration were out of keeping with the diagnosis of classical diabetic ketoacidosis and should have suggested a further metabolic problem. In diabetic ketoacidosis sodium levels are commonly low¹ although it would be unusual to find a sodium level of 120 mmol/l. In addition, obese diabetics have elevated insulin levels and are resistant to ketosis and it seems likely that the ketonuria in this case was related more to starvation. The improvement, without insulin administration, would support this suggestion. Any diabetic presenting with an acidosis and relatively normal glucose levels should however receive intravenous insulin and hypertonic glucose to correct the acidosis in addition to any other appropriate intravenous fluid therapy.

Transient hypercalcaemia is a well recognized, although uncommon, feature of acute adrenal insufficiency⁸ which is due to the effect of glucocorticoid deficiency leading to decreased calcium excretion and probably increased bone resorption.¹⁰ The findings of hypercalcaemia in our cases suggest that it is more common than the 6% of cases noted in a previous study.⁸

One further point which requires emphasis is that the degree of fluid replacement required in our patients appeared to be out of keeping with the elevation in the baseline urea levels. In Addison’s disease volume depletion seldom exceeds 10% of the total body fluid.¹ The large replacement volumes needed to correct the postural hypotension and hyponatraemia therefore suggest a more severe metabolic decompensation.

In 1885¹¹ Addison described an ‘irritability of the stomach and a peculiar change of colour of the skin’. In patients who present in medical as well as surgical wards with these clinical phenomena, and exhibit the characteristic biochemical abnormalities of acute adrenal insufficiency, there should be no undue delay in the diagnosis of this medical emergency.

Acknowledgement

We wish to thank Mrs M. Tucker for typing the manuscript.

References

Missed Addisonian crisis in surgical wards.

M. Small, A. C. MacCuish and J. A. Thomson

doi: 10.1136/pgmj.63.739.367