Haemochromatosis and aldosterone deficiency presenting with *Yersinia pseudotuberculosis* septicaemia

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Summary: A 50 year old man presented with a pyrexial illness following a holiday abroad. *Yersinia pseudotuberculosis* was isolated from blood culture. Response to appropriate antibiotic therapy was prompt and complete, but full recovery was complicated by an episode of hyperkalaemia, hyponatraemia and the passing of large volumes of dilute urine. Three years previously he had developed diabetes mellitus and hypogonadotropic hypogonadism. Investigation on recovery showed underlying haemochromatosis and aldosterone deficiency.

Introduction

*Yersinia pseudotuberculosis* is a Gram-negative coc-cobacillus ubiquitous within the animal kingdom. Human infection can occur by ingestion of food contaminated by animal excreta, drinking infected milk or water, or by direct contact with an infected animal.¹ In man it produces two distinct clinical entities, acute mesenteric adenitis and septicaemia.²⁻⁴ Disordered iron metabolism is probably the important common factor in those illnesses that predispose to yersinia septicaemia.⁵⁻⁶ We report a case of severe *Yersinia pseudotuberculosis* septicaemia in a man with hitherto undiagnosed haemochromatosis. A rare manifestation of haemochromatosis is aldosterone deficiency due to iron deposition in the zona glomerulosa of the adrenal cortex.⁷ We believe our patient has this complication which became manifest during the acute illness as polyuria, hyponatraemia and hyperkalaemia.

Case report

After 4 days holiday in Rhodes, a 50 year old Caucasian male developed a dull and constant lower abdominal ache, pain in both leg and arm muscles, headache, nausea, vomiting and fever. He was treated with unspecified intravenous antibiotics but following his return to England he continued with increasing nausea, intermittent fever, night sweats, shivering and complained of the recent onset of pleuritic chest pain.

On examination he looked ill and icteric. His abdomen was generally tender but without guarding or rebound. An appendicectomy scar was evident. He ran a low grade temperature which peaked at 39°C forty-eight hours after admission.

Three years previously, at another hospital, the patient was found to have diabetes mellitus requiring insulin. Investigation for impotence 7 months after the onset of his diabetes had shown hypogonadotropic hypogonadism with low plasma testosterone, follicle stimulating hormone and luteinising hormone levels. A normal increment in plasma cortisol levels had been demonstrated in response to insulin-induced hypoglycaemia and thyroid function tests were normal.

Investigations on admission were as follows: haemoglobin 13.3 g/dl, total white cell count 20 × 10⁹/l, 82% neutrophils, erythrocyte sedimentation rate 65 mm/h, bilirubin 20 µmol/l, alkaline phosphatase 7 King Armstrong units, aspartate transaminase (AST) 109 units, alanine transaminase (ALT) 85 units, albumin 28 g/l, globulin 45 g/l, plasma potassium 4.7 mmol/l, sodium 127 mmol/l, chloride 93 mmol/l, bicarbonate 29 mmol/l, urea 6.8 mmol/l, creatine 88 µmol/l, amylase 50.5 units. Chest X-ray showed right basal consolidation. Abdominal X-ray showed a moderate sized liver shadow. Treatment with erythromycin was started for a possible atypical pneumonia. Two days after admission his clinical condition deteriorated. He remained febrile at 39°C with a markedly tender liver and abdominal tenderness to very light palpation. Moderate diarrhoea was present with 13 loose stools in the first 4 days of admission. The illness was following a ‘typhoid-
like' course and despite its rapid onset, only 4 days after his arrival in Rhodes, a diagnosis of typhoid fever was considered. A Widal test taken at this time (day 4) subsequently showed no evidence of *Salmonella typhi* or *paratyphi* infection.

Seven days after admission a Gram-negative bacillus was isolated after 48 hours incubation from one only out of several sets of blood cultures. This was identified as *Yersinia pseudotuberculosis* on the basis of its mobility at 22°C and lack of mobility at 37°C and by API. The latter is a standard identification system for Enterobacteriaceae and other Gram-negative rods using a series of miniaturized biochemical tests and a data base. Subsequent serology from the national reference centre at Leicester Public Health Laboratory showed an antibody to type IA of 160 at 7 days, rising to 1280 by day 8 and falling to 640 by day 22. Attempts at culture from faeces failed in spite of cold enrichment. Coincidental with the patient’s improvement it was noticed that he was passing large volumes of dilute urine, specific gravity 1005–1010, with hyperkalaemia (highest 7.1 mmol/l) and hyponatraemia (lowest 120 mmol/l). Hyponatraemia had been present at admission and was considered to be a non-specific feature of severe illness. Blood glucose ranged from 4.4–13.3 mmol/l. The plasma and urinary electrolyte pattern suggested adrenal failure. A 5-hour depot tetracosactrin test was carried out and pending the results treatment with hydrocortisone and fludrocortisone was begun. The plasma potassium was 3.6 mmol/l and sodium 130 mmol/l on day 14. The tetracosactrin test yielded normal base line plasma cortisol (517 nmol/l, reference range 200–750 nmol/l) and an appropriate 5 hours response (1171 nmol/l).

On recovery from the acute *yersinia* infection investigations for haemochromatosis showed serum ferritin levels markedly elevated at 4550 and 4760 ng/ml (male reference range 25–240). Liver biopsy showed massive accumulation of stainable iron in hepatocytes and Kupffer cells. The portal tracts were expanded by fibrous tissue but there was no portal-portal linking and the appearances fell short of cirrhosis. The patient was maintained on insulin and 200 μg/day of fludrocortisone and some weeks after recovery, adrenal function was investigated. Basal plasma cortisol was 390 nmol/l and increased to 1360 nmol/l five hours after 1 mg depot tetracosactrin intramuscularly, a normal response. Having discontinued the fludrocortisone for 6 days, an angiotensin infusion test was carried out and showed no significant increment in plasma aldosterone above the basal level in response to angiotensin (Table I).

### Discussion

*Yersinia* septicaemia may present as a subacute localized form with hepatosplenic abscesses or follow a ‘typhoidal course’ with a rapid rise in temperature, rigors, abdominal, joint and limb pain, hepatosplenomegaly, exudation in serous cavities and a continuing or intermittent temperature.

Table I Plasma aldosterone response to infusion of angiotensin II 5 ng/kg/min

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<tr>
<th>Time (minutes)</th>
<th>Plasma aldosterone* (pmol/l)</th>
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<tbody>
<tr>
<td>0</td>
<td>440/500</td>
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<tr>
<td>30</td>
<td>320</td>
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<td>60</td>
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<td>120</td>
<td>470</td>
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*Reference range after overnight recumbency, 100–500 pmol/l.

Although *Y. pseudotuberculosis* and *Y. enterocolitica* grow rapidly in a defined iron sufficient medium, they fail to release detectable siderophore activity in an iron-deficient medium. Both of these organisms can use common exogenous siderophores produced by other micro-organisms and are therefore able to thrive in the gut. Body iron is, however, mostly bound in the form of haemoglobin or ferritin. In conditions associated with iron overload, with saturation of serum transferrin, the pool of iron is bound non-specifically to other proteins and is more available to bacteria. This more ready supply of iron will encourage *yersinia* septicaemia.

Our patient exhibited classic features of *Y. pseudotuberculosis* septicaemia with pre-existing
haemochromatosis and diabetes mellitus, a thyroidal-like illness with marked abdominal tenderness, full recovery with intensive and early treatment with appropriate antibiotics, and isolation of type 1A, responsible for the large majority of human infections. The predilection for iron to be deposited in the zona glomerulosa has long been known. Only recently, however, has aldosterone deficiency causing a clinical problem been described. In our patient the water, sodium and potassium derangement were compatible with adrenocortical hypofunction but normal glucocorticoid release from the adrenal was demonstrated. However, although the basal plasma aldosterone level was in the normal range, indeed at the upper end, it did not increment as it should have done in response to angiotensin infusion. We conclude that although aldosterone production was adequate for unstressed life, the demands of the very severe illness could not be met. This resulted in considerable loss of sodium and water, and potassium retention.

Acknowledgements

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References

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