Annals of the Rheumatic Diseases, 1989; 48, 430–431

Case report

Acute adrenal insufficiency as a manifestation of the anticardiolipin syndrome?

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Summary A 34 year old man with a past history of deep vein thrombosis, positive lupus serology, and high titres of anticardiolipin antibodies developed acute adrenal insufficiency. Association of anticardiolipin antibodies with adrenal insufficiency has not been previously reported.

Key words: anticardiolipin antibodies, systemic lupus erythematosus.

Anticardiolipin antibodies have been associated with recurrent venous and arterial thrombosis, thrombocytopenia, fetal losses, and neurological disease. We report a patient with 'atypical lupus' and raised anticardiolipin antibody titres who developed acute adrenal insufficiency.

Case report

A 34 year old white man was admitted to hospital in January 1984 with a 24 hour history of cramp-like lower abdominal pain and fever. He had returned a few days earlier from a two week cruise in the Caribbean. At the age of 17 he had suffered from a deep femoral vein thrombosis confirmed by venography. A routine blood test when he was 19 had shown the presence of antinuclear antibodies in a low titre.

The abdominal pain had started a few hours after playing squash. He did not complain of nausea, vomiting, or diarrhoea. Physical examination on admission was unremarkable except for a temperature of 38°C, skin hyperpigmentation, attributed to his recent trip, and right lower abdominal tenderness without guarding. Laboratory findings showed haemoglobin 146 g/l, white blood cell count 10.1×10⁹/l with a normal differential, platelets 208×10⁹/l, and Westergren sedimentation rate of 53 mm/h. Blood urea and electrolytes were normal. Blood, stool, and urine cultures were negative. The patient subjectively improved and was discharged nine days later without a specific diagnosis.

He was readmitted to hospital a week later feeling extremely weak and still complaining of abdominal pain. He was dehydrated and had lost 4.5 kg. His temperature was 37.5°C and blood pressure 70/30 mmHg. Laboratory investigation showed haemoglobin 125 g/l, white blood cell count 1.9×10⁹/l (67% neutrophils), and platelet count 154×10⁹/l. Blood urea was 14.3 mmol/l, serum potassium 6.1 mmol/l, sodium 117 mmol/l, and creatinine 150 μmol/l. Repeated blood, stool, and urine cultures were negative. Partial thromboplastin time was 101.5 s (control 33.6 s), prothrombin time 13.9 s (control 11.1 s), and thrombin time 34 s (control 17 s). The addition of normal plasma did not correct these abnormalities. Factor II was reduced to 0.35. High titres of fibrin degradation products determined twice with a 12 hour interval were present (>1280 μg/l, normal <80 μg/l). Fibrinogen was 3.96 g/l (normal range (NR) 1.6–4.2 g/l). Antinuclear antibodies (homogeneous pattern) were present (1/1280). Veneral Disease Research Laboratory (VDRL) test and direct Coombs' test were positive. Anti-DNA, antithyroid, anti-adrenal, and antismooth muscle antibodies were absent. Serum cortisol was 49.7 nmol/l (NR 190–700 nmol/l).

Accepted for publication 23 July 1988.

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Acute adrenal insufficiency and the anticardiolipin syndrome

A diagnosis of primary adrenal insufficiency was made, and the patient was treated with steroids. His clinical status rapidly improved. The diagnosis was confirmed by a three day corticotrophin stimulation test, which showed decreased urinary cortisol concentration (max: 88 nmol/24 h, NR 200–745). He has remained completely asymptomatic, treated with a maintenance dosage of cortisone acetate 37.5 mg and fludrocortisone acetate 0.05 mg daily. Analysis of stored serum samples obtained in June 1984 (four months after his hospitalisation) showed high levels of anticardiolipin antibodies (enzyme linked immunosorbent assay (ELISA): 1:86 optical density (OD) units, normal <0.4 OD units).

Discussion

Idiopathic or autoimmune Addison's disease accounts for more than 85% of all cases of primary adrenal insufficiency. Anti-adrenal antibodies are detected in two thirds of the patients, and other tissue autoantibodies are frequently seen. Interestingly, antinuclear antibodies are rarely present and, so far, only two cases of primary adrenal insufficiency have been reported in association with systemic lupus erythematosus. In both cases vasculitis restricted to the adrenal glands, and not autoimmunity, was suggested as the underlying pathophysiological mechanism.

Our patient had clinical and serological manifestations typical of those previously described in patients with anticardiolipin antibodies. He had a history of venous thrombosis at a young age and numerous abnormalities were demonstrated in his serum (antinuclear antibodies, leucopenia, 'lupus anticoagulant', low factor II, false positive VDRL test, and positive Coombs' test). Like many patients with antibodies against cardiolipin he did not satisfy the revised criteria for a diagnosis of systemic lupus erythematosus.

Although we cannot rule out an autoimmune aetiology to his adrenal failure, the absence of anti-adrenal antibodies, the negative family history, and the fact that autoimmune adrenal insufficiency has never been reported in typical systemic lupus erythematosus militate against this hypothesis. We believe that bilateral adrenal vein thrombosis should be considered as a possible aetiology in this case in view of the underlying hypercoagulable state (anticardiolipin antibodies) and the presence of high titres of fibrin degradation products during the acute episode. The absence of septicaemia, the normal platelet count, and fibrinogen concentration argue against the suggestion that increased fibrin degradation products may be explained by disseminated intravascular coagulation.

Adrenal vein thrombosis is a rare cause of adrenal insufficiency, which has been reported mainly in necropsies. Diagnosis of adrenal vein thrombosis is by adrenal venogram. This was not carried out in our patient as the diagnosis was not suspected at the time.

No anticoagulant treatment or prophylaxis was prescribed for our patient, and he has remained symptom free despite the persistence of serological abnormalities.

References

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doi: 10.1136/ard.48.5.430

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