Persistent chorea as a manifestation of thyrotoxicosis

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Summary: We report a case of persistent chorea as a manifestation of thyrotoxicosis. The chorea was severe and persisted after the patient was rendered euthyroid. Dopamine antagonists only partially suppressed the involuntary movements during the first few months. It was eventually controlled with haloperidol, but whenever she discontinues the treatment the chorea has returned during the 16 months since she first presented.

Introduction

Choreoathetosis is a recognized manifestation of thyrotoxicosis. It was first described by Sir William Gowers in 1893. There have been several papers in the past reporting cases of chorea due to thyrotoxicosis. In all these reports, except one, chorea was quickly reversible by propranolol, dopamine antagonists or anti-thyroid drugs. There is one case in the literature which describes chorea which dated from an attack of thyrotoxicosis 30 years earlier.

This paper reports a case of severe choreiform movements, with generalized hypotonia associated with thyrotoxicosis. The involuntary movements continued after the patient was rendered euthyroid. This case suggests that the neurological disorder following hyperthyroidism, which usually is transient, may become persistent.

Case report

A 15 year old girl was admitted to hospital as an emergency with violent involuntary movements. These movements had started 3 years earlier and gradually became worse, involving the whole body. She was dropping things at home and school. Shortly before admission the movements became increasingly violent and she collapsed at a friend's party and was unable to stand again.

Until the age of 11 she was short and fat, but then grew rapidly, becoming tall and thin. She grew 15 inches in 2 years and lost weight in spite of a voracious appetite. She also complained of being hot, sweaty and thirsty.

On examination she had violent choreiform movements of her arms, legs, neck and trunk. They were so severe that she had to be nursed in a hammock bed. There was profound generalized hypotonia with dysarthria and dysphagia. She had a tachycardia of 110 per minute, with blood pressure 145/70 mm Hg. There was a soft systolic murmur at the left sternal edge. She had a diffuse goitre. There were no ocular signs of thyrotoxicosis.

The haematological investigation revealed hypochromic microcytic anaemia due to iron deficiency for which no cause was found. Serum electrolytes, renal function, liver function tests and copper studies were normal. No abnormality was found on skull X-ray or computed tomographic scan of the brain. Baseline serum growth hormone and its response to glucose load was normal.

Thyroid function tests showed that she was thyrotoxic with serum free thyroxine (T₄) 86.5 pmol/l (normal range 11–25), serum free triiodothyronine (T₃) 18.7 pmol/l (normal range 3–9), thyroid-stimulating hormone 0.05 mU/l (normal range 0.3–5.0). Thyroglobulin antibodies were positive 1:160 and microsomal antibodies positive 1:100. Streptococcal origin of chorea was excluded by negative throat swabs and normal anti-streptolysin and antihyaluronidase antibodies which were performed on two separate occasions. Anti-DNA antibody tests were also negative. Other bacteriological investigations were also normal and so were the viral complement fixation tests and spirochaetal serology.

As soon as the diagnosis of thyrotoxicosis was made she was started on propranolol 40 mg t.d.s. and carbimazole 20 mg t.d.s. Symptomatic treatment for the chorea was started with chlorpromazine 25 mg t.d.s. The dose was increased and later tetrabenazine 25 mg t.d.s. was added, the dose of which was also increased to 50 mg t.d.s. The thyro-
References

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