Hyperthyroidism with periodic paralysis, acropachy, pre-tibial myxoedema, transient atrial fibrillation and myopathy

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Summary
A case of thyrotoxicosis presenting with periodic paralysis and later complicated by acropachy, pre-tibial myxoedema and transient atrial fibrillation is described. This association has not been reported previously. Possible aetiological links are discussed.

Introduction
Periodic paralysis, idiopathic atrial fibrillation, acropachy, manic depressive illness, skeletal manifestations, pernicious anaemia and pre-tibial myxoedema have all been separately reported in association with thyrotoxicosis. It is very unusual to encounter a patient with many of the complications. In Singapore thyrotoxic periodic paralysis is relatively common (6.0% of all thyrotoxics) while pre-tibial myxoedema is rare (0.7% of all thyrotoxics). The association of the two conditions in a patient has not been described previously. We report in this paper a patient with hyperthyroidism presenting with periodic paralysis who subsequently developed acropachy, transient atrial fibrillation, pre-tibial myxoedema and myopathy.

Case report
A 48-year-old Chinese man presented in February 1973 with sudden onset lower limb weakness after a heavy meal and had had to be carried to a hospital. Examination revealed a thin, anxious man with typical signs of hyperthyroidism: lid retraction of both eyes with no periorbital swelling, warm moist hands with fine finger tremors, nail dystrophy but no clubbing, slightly enlarged thyroid gland with no bruit and no evidence of tracheal compression (Fig. 1). He had evidence of lower limb proximal muscle weakness but not other findings were evident in the locomotor system. Cardiorespiratory system and abdomen were normal.

Laboratory investigations confirmed thyrotoxicosis with a serum tri-iodothyronine (T3) uptake of 175% (normal 75–115), thyroxine (T4) 9·6 μg/dl (normal: 3–7), free thyroxine index 16·8 (normal: 2·25–8·05). His serum potassium was low (3·1 mmol/litre) on admission but became normal with potassium supplements. Anti-thyroglobulin and anti-microsomal antibodies were absent. He was treated with radioiodine (7mCi 131I) and was euthyroid a month later with no further episodes of leg weakness.

Ten months after his initial presentation he was noted to be mildly clubbed. He failed to turn up for

FIG. 1. The patient on initial presentation
review 12 months after admission only to be re-admitted 4 months afterwards because of relapse thyrotoxicosis with increasing weakness in all 4 limbs over a 48-hr period.

Clinical examination revealed an obviously thyrotoxic man. His thyroid gland had not increased in size but a bruit could be heard over the slightly bigger right lobe. He had bilateral exophthalmos with diplopia on looking to the left and upwards. Clubbing of the fingers with tremors were noted. His toes were also clubbed (Figs. 2 and 3). His limbs were generally weak (MRC Grade 3/5) especially over the proximal musculature. Some rough nodular swellings were noted over his shins (Fig. 4). His serum T3 uptake was 170%, T4 8.0 μg/dl and free thyroxine index 13.6. Serum potassium was low at 3.0 mmol/litre. Other biochemical investigations (calcium, phosphate, magnesium, muscle enzymes, immunoglobulin levels) were normal and antithyroid antibodies (anti-thyroglobulin and antimicrosomal) were absent. An electromyogram showed myopathic changes in his proximal muscles with normal motor nerve conduction. X-ray of hands and forearms failed to show any periosteal elevation and biopsy of the nodular areas showed typical lesions of pre-tibial myxoedema.

Oral potassium supplements and carbimazole were started. His diplopia and weakness improved with a mild residual proximal weakness. He was discharged with no improvement of his clubbing or skin lesions.

He became euthyroid a few months later and antithyroid therapy was discontinued 10 months after his relapse. His clubbing, however, was noted to be...
more gross. Repeat chest X-ray and hand X-ray failed to reveal any abnormalities.

He was re-admitted 6 months later in March 1976 with another episode of sudden generalized weakness. He was now in atrial fibrillation with a ventricular response of about 120/min. There were no finger tremors but clubbing was obvious. His thyroid gland was small and firm with no bruit heard. His limbs were very weak (MRC Grade 2/5) especially over the proximal musculature with minimal muscle wasting. The rough areas over his shins were still present.

Biochemical investigations confirmed thyrotoxic periodic paralysis; his serum potassium was 1.9 mmol/litre, T3 uptake 146%, thyroxine 12.4 µg/dl, free thyroxine index 1.84 and electrocardiogram showed atrial fibrillation.

Oral potassium supplements were restarted and he was given a second dose of radioactive iodine (10mCi 131I) after which his muscle power improved with residual mild proximal weakness. His atrial fibrillation was transient and he converted to sinus rhythm two days later with a rate of 85/min.

He remained euthyroid both clinically and biochemically. His skin lesions gradually subsided but his finger clubbing remained and became more gross. Repeated chest X-rays to date have failed to reveal any lung pathology. He was euthyroid when last seen in March 1981.

Discussion

Periodic paralysis is a well-known although not a common complication of thyrotoxicosis, except amongst the Chinese, Japanese and a few other ethnic groups (Cheah, 1978; Minns, Newlin and Day, 1978). The incidence amongst Chinese in Singapore was found to be 6.0%: 19.2% in male Chinese and 0.3% in female Chinese (Cheah, 1974). There have been reports of the familial occurrence of thyrotoxic periodic paralysis (Yueng and Au, 1978) and it has been suggested that the racial predilection to this condition for Chinese, Japanese and other mongoloid races is genetically determined and that patients affected share similar HLA antigens such as A2/BW22 and AW19/B17 (Yeo et al., 1978). It appears that thyrotoxicosis unmasks the defect by an increased release of insulin to carbohydrate ingestion causing the shift of potassium into muscles thereby causing hypokalaemia and resultant muscular paralysis (Cheah, 1974). Our patient's HLA profile was A2/BW22, there was no family history of hypokalaemic periodic paralysis.

The incidence of thyrotoxic pre-tibial myxoedema in Singapore was found to be 0.7%; 1.6% in males and 0.3% in females (Kee and Cheah, 1975). Gimlette (1960) found a higher percentage of its incidence. Pre-tibial myxoedema has also been associated with finger clubbing and exophthalmos.

Our patient is unusual in that he presented with
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thyrotoxic periodic paralysis and later developed pre-tibial myxoedema and clubbing. The precise aetiology of these three conditions remains obscure and it has been suggested that pre-tibial myxoedema and clubbing are due to localized antigen-antibody reaction (Kriss, Pleshakor and Chien, 1964). Kriss and co-workers suggested that local destruction of the thyroid gland caused release of antigen which would stimulate antibody production. The antigen-antibody complex resulting from this could then get fixed to the leg tissue because of dependency and the local inflammatory reaction resulting causes the condition of pre-tibial myxoedema. A similar mechanism could explain the clubbing (acropachy).

Recent work has shown that all clinically thyrotoxic patients with abnormal thyroid function tests show evidence of myopathy on electromyography although not all show clinical myopathy (Puvanendran et al., 1979).

There is no evidence yet to suggest that an immunological reaction is responsible for the muscular complications of thyrotoxicosis although this mechanism cannot be excluded completely. Instead these complications have been thought to be due to the physiological effect of excessive thyroid hormone on the muscles (Ramsay, 1966; Satoyoshi, Murakami and Torin, 1963).

Another unusual aspect of our patient’s condition was that he showed a rather rapid response to his first dose of radio-iodine, being euthyroid after one month, only to relapse over a year later. The reason for this is not known. Perhaps there is an association between this and his multiplicity of complications.

Thyrotoxicosis is a relatively common disease. Why is it that the above complications occur rarely? Genetical susceptibility to their predisposition might be partially responsible as suggested by studies on HLA antigens in patients with thyrotoxic periodic paralysis (Yeo et al., 1978). There are no similar studies on the HLA profile of patients with pre-tibial myxoedema or acropachy, as yet. Perhaps these might tell us more about the aetiology and pathogenesis of these complications and provide an explanation for the multiple complications of thyrotoxicosis that our patient suffered.

References


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