Bipolar affective disorder, stress fractures, fungal dermopathy and ‘tree frog fingers’

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A 51-year-old woman was referred to the Department of Endocrinology following her visit to a dermatological out-patient clinic. She gave an eight-year history of medical problems: following the death of her father she had consulted her general practitioner with weight loss, anxiety, psychomotor retardation and suicidal ideation. After these symptoms had failed to improve on treatment with a tricyclic antidepressant she was admitted to the local psychiatric hospital. There she received eight courses of electroconvulsive therapy and was discharged on prophylactic lithium and tricyclic antidepressant therapy. Three years later the (premenopausal) patient suffered a stress fracture of her second left metatarsal, followed by stress fractures of her third and fourth left metatarsals one year later. A year later she noticed a puffy swelling of the pulp of her right thumb. Over the following two years, the left thumb, right ring and index fingers were all affected by similar changes and she became aware of a tendency to bruise easily.

In the year preceding her presentation, the patient suffered a further stress fracture (third right metatarsal) and a recurrence of psychiatric symptoms four weeks after she had stopped her lithium and clomipramine therapy. Her sleep requirement dropped to three hours, her attention was poor and she became emotionally labile. Her maniacal symptoms precipitated a further psychiatric admission and the commencement of carbamazepine therapy. After her discharge, the patient’s long-standing tinea pedis became so troublesome that she consulted a dermatologist.

Examination revealed no markedly plethoric facies, centripetal obesity or hirsutes. Pale striae were noted in both flanks. Body Mass Index was 28.3. There was extensive tinea pedis. The hand changes are illustrated in the figures. Skin thickness was reduced to less than 2 mm. Her pulse was 72 beats/min, blood pressure 170/110 mmHg. Otherwise cardiovascular and chest and abdominal examination were unremarkable. Neurological examination revealed no abnormalities. The dermatologist advised topical treatment of the intertriginous fungal infections. He was, however, intrigued by the frog-finger-like appearance of both her thumbs, her right ring and index fingers and suggested some simple screening tests, following which the patient was referred for endocrinological investigation.

Questions

1. What diagnosis did the dermatologist suspect?
2. Which screening tests were performed to support the diagnosis?
3. Which further investigations were required before treatment could be undertaken?
Answers

QUESTION 1
Cushing’s syndrome is the only unifying diagnosis in this patient presenting with psychopathy, osteoporosis, dermopathy (easy bruising and skin fragility, reduced skin thickness, tinea pedis and tree frog fingers’), mild obesity and hypertension. Like hypercortisolism, thyrotoxicosis may initially present with psychiatric symptoms, and it is a cause of osteoporosis. It may also present as a dermopathy (pretibial myxoedema, infiltrative ophthalmopathy) and is associated with distal finger changes (thyroid acropathy). The digital changes, however, resemble clubbing and do not affect the pulp of the finger tip.

QUESTION 2
Unfortunately a random cortisol estimation is not a very useful diagnostic test in this setting. The initial screening test performed was a 24-h urine free cortisol (UFC). This was found to be raised at 640 nmol/24 h (reference range 25–280 nmol/24 h). Whilst measurement of UFC is a good screening procedure (elevated in 90% of patients with Cushing’s syndrome) alternatives would have included a cortisol day curve (midnight cortisol normally less than 50% of 09.00 h value) and a 1 mg overnight dexamethasone suppression test. All of these tests have their drawbacks, particularly in the setting of psychiatric illness, stress, drug treatment, hormone replacement therapy and oral contraception, obesity and alcoholism. If any of the screening tests are abnormal or borderline and there is a high clinical index of suspicion, a low-dose dexamethasone suppression test should be performed (0.5 mg 6 hourly for 48 h). In the presented case, 09.00 h serum cortisol failed to suppress during this test (cortisol 825 nmol/l after 24 h and 1055 nmol/l after 48 h of dexamethasone 0.5 mg 6 hourly). This result established the diagnosis of hypercortisolism. In a high-dose dexamethasone suppression test (2 mg 6 hourly for 48 h), 09.00 h cortisol showed significant suppression, thus suggesting a diagnosis of (pituitary) Cushing’s disease rather than that of an adrenal adenoma or ectopic adrenocorticotropic (ACTH) production.

QUESTION 3
After haematological and biochemical complications of Cushing’s disease had been excluded (leucocytosis, lymphopenia, thrombocytosis, electrolyte disturbance, hyperglycaemia), the presence of hypercortisolism was confirmed once carbamazepine and hormone replacement therapy had been discontinued. The mean value of a serum cortisol day curve was 457 nmol/l with a midnight cortisol of 413 nmol/l (patient asleep). Low and high dose dexamethasone suppression corroborated the outcome of the earlier tests. Further investigations were aimed at identifying the cause of the patient’s hypercortisolisma. Biochemical tests included a corticotropin-releasing factor (CRF) test, which revealed a blunted ACTH and cortisol response with values rising from 23 to 39 ng/l and 474 to 678 nmol/l, respectively. This supported a diagnosis of (pituitary) Cushing’s disease. Bilateral inferior petrosal sinus catheter sampling gave no indication of the site of a possible ACTH-producing pituitary adenoma (ACTH rise after CRF stimulation on the right from 22 to 85 ng/l, on the left from 22 to 98 ng/l). Radiological investigations included normal chest and skull radiographs. Computed tomography of the pituitary showed a bulky gland filling the fossa, with a slightly convex upper surface. Magnetic resonance imaging showed a pituitary fossa floor sloping downwards anterolaterally to the right side with a suggestion of a low signal lesion in the right pituitary lobe. A slightly bulky gland seemed to deviate the pituitary stalk to the left. In view of the history of stress fractures DXA-densitometry was performed, revealing bone density to be 67–79% below the mean of females of the patient’s age.

A diagnosis was made of Cushing’s disease caused by a pituitary microadenoma, possibly on the right. The patient was referred for transsphenoidal hypophysectomy. The immunohistochemical examination of the excised specimen revealed a chromophobe adenoma with positive immunostaining for ACTH and negative staining for other pituitary hormones. Some areas in the anterior pituitary showed Crookes’ hyaline change. This is an ultrastructural feature associated with ACTH-producing pituitary adenomas, consisting of bundles of fine perinuclear filaments.

Despite the histological proof of successful surgery, postoperative 09.00 h cortisol remained detectable (472 ng/l six days post-op), and a second operation was thought to be necessary. Exploration of the sella revealed a tissue segment of 1–2 mm diameter adherent to the cavernous sinus which was dually excised. After this surgical revision, cortisol values became undetectable and the patient was considered cured.

Discussion
Cushing’s syndrome affects most systems of the human body. The full-blown clinical picture is characterised by progressive specific-site adiposity, myopathy, dermopathy, hirsutism, stress fractures, psychopathy, glucose intolerance, hypercholesterolaemia, hypertension, atherosclerosis and immunosuppression. The psychopathy, as illustrated in the presented case, may become manifest as depression, mania or psychosis and can precede the onset of the classical physical signs by several years. Whilst the classical syndrome outlined above is relatively easy to recognise in clinical practice, more subtle manifestations of Cushing’s syndrome are difficult to spot and characterise, particularly in the setting of concomitant medical conditions such as obesity, alcoholism and depression. In the case presented here, the diagnosis was suspected in a dermatological out-patient department on the basis of a recently described sign.1

Dermatological complications of persistent hypercortisolism include fungal infections
whose spread and persistence are thought to be caused by cortisol-mediated reduction of IL-1 secretion from monocytes/macrophages and IL-2 release from activated lymphocytes resulting in a suppression of T-lymphocyte proliferation. The most characteristic skin changes of Cushing’s syndrome, however, are the result of an abnormal collagen metabolism. Easy bruising and violaceous striae formation are considered to be due to reduced collagen–platelet aggregation and cortisol-induced protein catabolism in blood vessel walls. Collagen catabolism and increased turnover also cause abnormalities of bone architecture leading to stress fractures. It has recently been shown that various serum parameters of collagen turnover such as type I procollagen and aminoterminal propeptide of type III procollagen are increased in Cushing’s syndrome.3

In the seminal report of profound acquired digital collagen atrophy (tree frog fingers), histological examination of the digital pulp showed focally decreased amounts of normal collagen with obvious dermal thinning. The dermal elastin fibres were noted to be of normal morphology but clumped as a consequence of the loss of collagen. These histological findings were broadly similar to more generalised cutaneous changes reported in previous studies of the effect of hypercortisolism on skin structure.4 As demonstrated in the digital photographs, the patient presented here exhibited a profound reduction of cutaneous elasticity in her affected fingers.

The case reported by Groves et al described similar changes in a patient with Cushing’s syndrome due to a benign adrenal adenoma.5 The patient described here represents the first case of marked digital cutis laxa due to Cushing’s syndrome caused by a pituitary adenoma (Cushing’s disease). Our report highlights the diagnostic challenge that Cushing’s syndrome continues to present. Dermatological changes serve to discriminate between true hypercortisolism and differential diagnoses including obesity, polycystic ovary syndrome and depression. Like skin-fold thickness,6 profound digital collagen atrophy with reduced elasticity should be looked for as a sign of Cushing’s syndrome.

### Final diagnosis
Cushing’s disease caused by a pituitary microadenoma.

### Keywords:
Cushing’s disease; dermopathy; collagen

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