Do I need to be an endocrinologist to diagnose Cushing’s disease? Around October 1999, while playing cricket after work, I went down for no apparent reason. The same event happened a month later. Being mainly concerned about unexplained falls, I sought help from neurological colleagues. Two neurologists examined me and both reassured me: “Your exam is normal and there is nothing to warrant any further investigations.” I checked my vitamin B12, which was 81.0 μg/ml (normal range 200–900). A combination of B12 deficiency and proximal myopathy could explain my falls. Having put on 4 kg over the past year, I sought advice from an internist. More concerning was that the central weight gain was starting to affect my appearance. The internist ascribed it to middle age spread and having checked my glucose and lipids, referred me to our nutritionist. The nutritionist having seen the results gave a good diet and exercise plan. Following this diet and exercise plan, I developed pain in my left foot and it got so severe that I presented to our radiologist with an x-ray film. On first look, he dismissed it as normal but on closer look identified a hairline fracture in the fifth metatarsal. I was referred to an orthopaedic surgeon who advised me to avoid brisk walks for two weeks. I questioned both of them—why did I break my bone? I was told that it may happen with strenuous workouts. Now I started to notice a change in my sleep, mood, and thoughts. I began to wake up about 3–4 am for no good reason and it was difficult to go back to sleep. Though unable to have a good night’s sleep of seven hours, which used to be a “must” for me, I was not in full of energy during the day but would develop a headache at the end of a busy clinic. I sought advice from my psychiatrist colleague. “It appears like mania or hypomania but B12 is low,” he replied. I forgot it, or if you like you may try some sedatives”. For headache, no recommendation.

I asked my staff nurse to check my blood pressure in the busy clinic and it was 180/100 mm Hg. Not surprisingly, I got more and more worried and one day while doing an endocrine clinic, decided to investigate myself.

While asking for routine analyses, I decided to include thyroid function tests and also cortisol. My cortisol (1600 hours) was 588 nmol/l (140–690), then looking back over the past month, it occurred to me that all of these features could be explained by one syndrome, Cushing’s, but for the low B12. I submitted a sample for 24 hour urine cortisol and reviewed the literature for B12 and cortisol. The 24 hour urine cortisol was 1644 nmol/day (110–436).

I emailed the case to my mentors, Professors A B Grossman and R J M Ross, also mentioning that I had found reports of B12 deficiency producing circadian rhythm abnormalities and dexamethasone non-suppression.1 I was advised to replace B12 and then repeat the tests. The urine cortisol was 466 nmol/day and overnight dexamethasone suppression test 66 nmol. However, after completing an empiric six injection course over one month, I repeated the same tests and the 24 hour urine low dose cortisol was 1911 nmol/day (110–436) and low dose dexamethasone suppression 295 nmol (<360). Subsequent investigations and imaging showed pituitary dependent Cushing’s disease. I underwent transphenoidal selective ad- enomectomy by my neurosurgeon colleague in my own centre. Postoperatively, I had undetectable cortisol and normal pituitary function.

This case illustrates the complexity involved in the diagnosis of Cushing’s disease. It also illustrates two points that physicians should be aware of when consulted by other physicians: “treat physicians as patients when they come for advice for medical problems” and “physicians could also develop any illness and should be diagnosed”. The question about B12 deficiency remains unanswered.

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References

Authors’ reply
We welcome the comments made by Sinharay. They certainly highlight some of the controversies surrounding this condition that we raised in the case presented. We certainly agree that all patients presenting with thyrotoxic periodic paralysis should have their thyroid status checked as, especially in white patients, signs and symptoms of thyrotoxicosis may be subtle and therefore we highlighted this is the summary points. This patient did in fact present in the evening. We were not involved in the patient’s initial management, but we would like to criticise the initial treatment by on-call doctors to use oral and intravenous potassium supplements as the diagnose of thyrotoxicosis had not been made (and was not suspected) and therefore we highlighted this is the summary points. This patient did in fact present in the evening. We were not involved in the patient’s initial management, but we would like to criticise the initial treatment by on-call doctors to use oral and intravenous potassium supplements as the diagnose of thyrotoxicosis had not been made (and was not suspected) and therefore we highlighted this is the summary points.

We welcome Sinharay’s comments on the puzzling male predominance of TPP in a condi- tion that has a female preponderance and that any explanations at present are speculative.

Sinharay disagrees with our comment that when patients are euthyroid they are not at risk of hypokalaemic paralysis. This is contrary to previous experience.1 However, we do agree that patients are at risk of hypokalaemic paralysis if thyrotoxicosis reoccurs. As there is a risk of relapse of hypokalaemic paralysis with recurrence of thyrotoxicosis we agree that there is a case to be made for early definitive treatment.

References

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Thyrotoxic periodic paralysis in western countries

R Sinharay

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