A pituitary tumour presenting with rhinolalia and galactorrhoea

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DESCRIPTION
A 37-year-old lady presented to her general practitioner (GP) with a history of intermittent galactorrhoea and amenorrhoea. She reported intermittent galactorrhoea to have been present following her last pregnancy 4 years previously. Earlier in the year she had been seen and treated for nasal symptoms which had caused hypernasal speech (rhinolalia). For this she was receiving fluticasone nasal spray.

The patient had a background of hypertension, well controlled on amiloride 10 mg once daily and lisinopril 20 mg once daily. There was no relevant family history.

Following an episode of menorrhagia, the patient had presented to her GP and routine blood tests were performed which included a serum prolactin of 3428 μ/l (25–629 μ/l). The GP requested an MRI scan which revealed a very large (4.4×4.6×2.7 cm) and extensive macroadenoma involving the central skull base (figure 1). There was significant suprasellar extension compressing the chiasm. Although there was no convincing lateral extension into the cavernous sinuses, there was filling of the sphenoid air sinus with the tumour and extension into both the left and right nasal cavities. On the left the extension was significant with displacement of the turbinates.

She was referred to our centre. All the other endocrinology was within normal limits. The prolactin level was confirmed with dilution to exclude a hook effect and trial of low-dose cabergoline was given which suppressed the prolactin to 23 μ/l in keeping with disinhibition. Although felt to be in keeping with a tumour originating from the pituitary, because of the very unusual appearance of this tumour, the nasal component was removed first (figure 2). This confirmed a null cell pituitary adenoma with a Ki67 index <1%. She subsequently underwent trans-sphenoidal surgery with clearance of the remaining tumour and no post-operative complications.

Learning points
▸ In a patient with inappropriate galactorrhoea–check the prolactin levels.
▸ Consider other differential diagnosis for causes of hyperprolactinoma, for example, excluding medication use, renal failure, hypothyroidism and parasellar tumours causing compression or displacement of the pituitary stalk in patients with symptomatic non-physiological hyperprolactinaemia.
▸ Head MRI scanning is essential.
▸ Trial of cabergoline therapy may help shrink tumour avoiding surgical intervention.

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REFERENCES