Case Report

A rare case of Cushing’s disease concurrent with papillary thyroid carcinoma

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Abstract

Background: Although a nodular thyroid disease is higher in patients with pituitary adenoma, concurrent thyroid cancer with pituitary tumor is uncommon.

Case Presentation: We report a young woman with discovered papillary thyroid carcinoma after 1-year transsphenoidal surgery for Cushing’s disease. Thyroidectomy was done and patient is well after three years follow-up.

Conclusion: We suggest the patient with functional pituitary adenoma be evaluated for thyroid tumor.

Keywords: Cushing’s disease, Papillary thyroid carcinoma

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Nodular thyroid disease (NTD) is one of the common thyroid disease and is frequent in general population. The most important risk factors are iodine deficiency, goitrogenic foods, radiation, aging, genetics, gender and smoking (1). An increased frequency of NTD has also been reported with functional pituitary adenomas especially with acromegaly and after that is more associated with prolactinoma and Cushing’s disease (2). However, concomitant papillary thyroid carcinoma and pituitary adenomas are not common, especially Cushing’s disease.

Case Presentation

A 33-year-old woman with a history of depression and anti-depressant drugs therapy was referred to the endocrine clinic due to headache, weight gain about 16 kilograms and purple stria on the skin during last year. The patient’s medications included sertraline and quetiapine, which weight gain continued despite quetiapine discontinuation and change to bupropion. The menstrual cycle was normal but the patient complained of hypomenorrhea and occasional galactorrhea since last year. She did not mention history of hirsutism, hypertension and diabetes mellitus. On physical examination, blood pressure was 120/80 mmhg and weight was 101 kg. Given the mentioned symptoms, the patient was evaluated for Cushing’s syndrome. In primary laboratory findings, morning basal plasma cortisol was 29 mic/dl and UFC (urinary free cortisol) 82.7 mic/dl (in normal range of laboratory). Morning fasting cortisol was not suppressed after 1 mg dexamethasone suppression test (9.9 mic/dl). Liddle test (low dose dexamethasone suppression test) was done and the results for serum and urinary cortisol were 10 micgr/dl and 15.5 micgr/dl respectively. The ACTH level was 118 pg/ml and a 14*15 mm pituitary adenoma was found in MRI. The patient’s visual field was normal. Except for prolactin which was high (907 miu/ml: normal up to 495), other pituitary axis functions were normal (table 1).
Cushing disease concurrent with papillary thyroid carcinoma

Table 1: Pre pituitary surgery laboratory finding

| Laboratory Test | Result | Normal range: |
|-----------------|--------|---------------|
| TSH (mIU/L)     | 1.6    | 0.5-5.5       |
| FreeT4 (ng/dl)  | 0.8    | 0.8-1.7       |
| Prolactin (Miu/l) | 907    | 132-498       |
| IGF1 (ng/ml)    | 289    | 107-246       |
| GH base (ng/ml) | 0.9    | 0.06-6        |
| GH 2 hours after 75 gr glucose (ng/ml) | 0.2 | <1 |
| NA (meq/l)      | 136    | 135-145       |
| K (meq/l)       | 4      | 3.5-5.2       |
| WBC (10³/micl)  | 13900 (PMN 4000-75%, lym 25%) | 11000 |
| FBS (mg/dl)     | 95     | 70-99         |
| Anti Tpo (unit/ml) | 1.3 | <30          |

The patient was treated with cabergoline 0.5mg/week for 3 months, though the prolactin level decreased to 10 miu/ml but the pituitary adenoma did not shrink and finally she underwent TSS (Trans spheroidal surgery) with the diagnosis of corticotroph adenoma. In a pathology report, pituitary tumor was confirmed. In IHC study, the GH and ACTH markers were positive in majority of the tumor cells, but TSH, FSH, LH, P53 and prolactin were negative. Synaptophysin and chromogranin A were positive and ki67 was positive in 1% of tumoral cells.

After pituitary surgery prednisolone tablet, 5 mg/d and nasal desmopressin spray was started and is continued until now. The weight gain stopped and the skin stria disappeared. After 1 year follow-up, the patients was well but complained of lumpy filling in neck. The thyroid examination was suspicious for thyroid nodule in right lobe. Morphologic characteristic of the nodule in sonography included a solid hypoecho signal, 10*9 mm in size, ill defined, taller than wide without calcification in right thyroid lobe. In addition, another 6*4 mm nodule with benign appearance was reported in right thyroid lobe. A sonography guided fine needle thyroid nodule aspiration (FNA) was performed. The thyroid cytopathologic diagnosis was atypia (or follicular lesion) of undetermined significance (AUS/FLUS). The FNA of thyroid nodule was repeated 3 months later and the papillary thyroid carcinoma diagnosis was reported in second cytology report. Total thyroidectomy was performed. In final pathology report, the tumor size was 1 cm without extrathyroidal extension and capsular or vascular invasion and there was one follicular adenoma in right lobe. No lymph node was involved. Based upon the clinicopathologic report and low risk of recurrence (papillary thyroid cancer confined to thyroid) the patient did not receive Radiiodine ablation or treatment. Thyroglobulin and ant-thyroglobulin antibodies were undetectable three months after thyroidectomy.

After 3 years of follow-up, the patient is well with her weight and has normal menstrual cycle. She is on levothyroxine 150 micgr in addition to carbonate calcium, vitamin D. The laboratory and radiology findings (neck sonography and brain MRI) are acceptable and the patient does not need to prednisolone or desmopressin.

Discussion

NTD is more frequent in patients with functional pituitary adenomas (2). About 8-16 percent of NTD are malignant. The thyroid PTC can be seen concurrent to some kind of pituitary tumors. Screening for PTC is recommended in patients with...
acromegaly because of increased risk of developing cancer in this group (3-13). Though an increased risk of NTD in patients with CD and prolactinoma has been reported in recent studies (3, 4, 7, 9, 10), but screening for PTC is not recommended yet (2-19).

Synaptophysin is positive in normal anterior and posterior pituitary gland and all pituitary adenomas. Chromogranin A is characteristic for gonadotrope adenoma and is poorly expressed in GH adenomas. For the differentiation between GH secreting tumor and gonadotropes complementary tests should be done. The positivity of cells for ACTH and GH antibodies can be because of the cross reaction of GH Aβ s with ACTH adenoma or less probably , the cross reaction of ACTH Aβ, s with GH secreting adenoma (20).

In a recent study in Turkey, the prevalence of NTD was 36% in prolactinoma, 34% in CD and 60% in patients with acromegaly. Also, the frequency of PTC was higher in CD (11.4%) compared to patients with acromegaly (10.8%) (21). Sheng-Fong et al. reported a 58-year-old man in Taiwan that had advanced PTC and then presented with pituitary apoplexy. The final diagnosis was recurrent ACTH producing adenoma without cushingoid features (22). In another study in 1968 in the USA, 2 cases of Cushing’s syndromes were reviewed which PTC was diagnosed after 2 and 5 years respectively (23). Kanazunorg et al. reported a 75-year-old woman with pituitary, thyroid and left adrenal gland pituitary tumor was preclinical Cushing’s disease accompanied with PTC and adrenal incidentaloma (24).

The similar frequency of PTC seen among the different FPA groups may reflect their common etiopathogenic mechanisms, including cytokines or growth factors. Another possibility is that the genetic and epigenetic factors that cause pituitary adenoma development may also be responsible for PTC. May be a close follow-up of NTD in CD patients beneficial for the early diagnosis of PTC. Further studies on larger cohorts are needed to assess the prevalence of PTC, and to specify the related risk factors.

Our results suggested that patients with Cushing’s disease may be needed to a close follow up for NTD, as the papillary thyroid carcinoma was confirmed in this study.

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Conflict of Interest: The authors declare that they have no conflicts of interest.

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