Hypertension episodes of adrenal cortical oncocytoma mimicking pheochromocytoma attacks.

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Abstract

Catecholamine-secreting tumors which are derived from chromaffin cells in adrenal medulla are called pheochromocytoma. The ratio of pheochromocytoma in all hypertensive patients is less than 0.2%. Adrenal oncocytomas can mimic the clinical features of pheochromocytoma. We aimed to present the findings of 2 patients with adrenocortical oncocytoma resembling pheochromocytoma.

Keywords: Adrenal oncocytoma, Pheochromocytoma, Hypertension.

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Introduction

Adrenocortical oncocytomas have rarely been reported on in the medical literature, and most of them have been nonfunctional and benign. Although spontaneous extensor tendon rupture often occurs in association with rheumatoid arthritis, extensor tendon rupture associated with osteoarthritis of the distal radioulnar joint has been rarely reported.

Case 1

A 43-year-old female patient was admitted to emergency department with high blood pressure in September 2014 and to neurology intensive care unit with right hemiparesis, loss of vision in November 2014. She had been discharged with antihypertensive therapy when radiological imaging and laboratory tests were both normal. Despite regular antihypertensive medication her blood pressure was irregular and she was hospitalized at nephrology department in October 2015. She had hypertension otherwise physical examination was normal. She was explored for high blood pressure, and high serum norepinephrine, 24-hour urine normetanephrine levels. Serum ACTH levels were suppressed and cortisol level was slightly increased. Magnetic resonance imaging of pituitary gland was normal. Adrenal magnetic resonance imaging revealed 52 × 38 mm adrenal adenoma (Figure 1). She had been operated. Pathology of adrenal adenoma was consistent with adrenocortical oncocytic neoplasm (Figure 2). Immunohistochemical method revealed positivity for melan A and negativity for chromogranin, synaptophysin, FSH, ER and ACTH (Figure 3). After surrenalectomy patient’s blood pressure did not reduce and she had menorrhagia. Gynecological examination and endometrial biopsy revealed endometrial adenocarcinoma. She had total abdominal hysterectomy and bilateral salpingoooforectomy. She is still having outpatient follow-up.

Case 2

A 62-year-old male patient had type-2 diabetes for nine years and hypertension for twelve years. He had unregulated blood pressure with routine antihypertensive drugs and was investigated at another hospital. MRI of abdomen revealed an adrenal adenoma at a diameter 40 × 30 mm on his left adrenal gland. He was operated on July 2015. But pathological
examination did not reveal the diagnosis. After operation he still had hypertension. And he arrived to our nephrology department. He had hypertension and physical examination was otherwise normal. Laboratory showed slightly decreased potassium level. But his serum ACTH, cortisol, renin and aldosterone levels were both normal. 24 hour urine samples showed increased normetanephrine (Table 1). Specimens of adrenal gland were reevaluated at our pathology department and adrenal adenoma was consistent with adrenal cortical oncocytic adenoma and amyloidosis (Figure 4).

Discussion

Adrenal oncocytomas are very rare. There is female predominance with a median age of 46 years. Our patients were forty three and sixty two years old respectively. These tumors are more common on the left side [1-5]. Most of these tumors are nonfunctional. Tumor size ranged from 3-15 cm; with a median of 8 cm. Oncocytomas are generally benign tumors. The Weiss system is most popular for distinguishing benign from malignant adrenocortical neoplasm⁶. Grossly these tumors are rounded, well-circumscribed and encapsulated. A cut section shows tan brown color with areas of hemorrhagic necrosis. Histologically, these tumors contain cells arranged in solid, tubular, papillary, or trabecular patterns [3,4]. Cells are strongly eosinophilic and granular, due to the presence of numerous mitochondria [4].

Two adrenal oncocytoma patients had resistant hypertension in postoperative period and their clinical, imaging, and laboratory findings resembled pheochromocytoma. In the literature there was presented adrenal cortical oncoytomas mimicking pheochromocytoma. At case 1, serum ACTH level was suppressed and serum cortisol level was slightly increased (only 5 µg/dL). Her appearance was consistent with neither Cushing’s disease nor Cushing’s syndrome. Other pituitary gland hormones were in normal range, MRI of pituitary gland was normal. In preoperative period she had hypertension attacks and serum norepinephrine levels were increased but this situation persisted despite development of adrenal insufficiency after left surrenalectomy. On the other hand in post-operative period 24-hour urine normetanephrine levels were higher than the upper range. Post-operative sintigraphic examination for pheochromocytoma was negative. In case 2 he had hypertension, serum norepinephrine levels were slightly increased and his serum ACTH/cortisol levels were within normal levels in preoperative period. In postoperative period he still had hypertension and 24 hour urine normetanephrine levels were slightly increased. Pathological diagnosis of adrenal adenomas (Cases 1 and 2) was consistent with adrenocortical non-secretory oncocytic neoplasm. Because immunohistochemical method revealed positivity for melan A and negativity for chromogranin, synaptophysin, FSH, ER and ACTH. After surrenalectomy patients were treated with antihypertensives but hypertensive attacks did not developed during follow up period. While amyloidosis in oncocytoma and renal specimen was positive in case 2 but negative in case 1. Case 1 was also operated for endometrial adenocarcinoma. Pathologically these metachronous tumors were different each other [5,6].

Immunohistochemical studies are helpful in distinguishing such cases as pheochromocytomas are positive for chromogranin A and other neuroendocrine markers [7]. Electron microscopic studies will reveal dense-core membrane-bound granules in the cells of pheochromocytomas [8].

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<th>Table 1. Pre-operative and post-operative parameters of patients.</th>
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<td>Parameters</td>
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<tr>
<td>BUN/Creatinine</td>
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Conclusion

In summary we presented two cases with hypertension and adrenal cortical oncocytoma mimicking pheochromocytoma. Amyloid deposition was found positive in one of them. After removal of these nonfunctional oncocytoma hypertension was continued less severely in our cases.

References


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