A Rare Case Presenting with Symptoms of Familial Pheochromocytoma

Familyal Feokromasitoma Semptomlarıyla Başvuran Nadir Bir Olgu

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ABSTRACT
Pheochromocytoma is a rare tumor in which seen an incidence of 1 per 100.000 in the general population. Pheochromocytoma is a catecholamine producing neuroendocrine tumor arising from adrenal medulla. A 46-year-old man experienced headache, sweating and palpitation referred to our hospital. Familial pheocromocytoma was detected. We presented this case as whole family members had suffered from morbidities and deaths due to this disorder.

Key words: Pheochromocytoma, familial disorder, hypertension

ÖZET

Anahtar kelimeler: Feokromasitoma, ailesel hastalık, hipertansiyon

INTRODUCTION
Pheochromocytoma is a rare tumor in patients with hypertension and general population that a catecholamine producing neuroendocrine tumor arising from adrenal medulla. Thus, these results in discharge of catecholamine, and is the second common cause of secondary endocrine hypertension, as adrenal dependent. Persistent hypertension, headache, and palpitations (especially sinus tachycardia) are the most common symptoms.

Advances in diagnosis and genetics now challenge the traditional rule of 10 for pheochromocytoma (10% bilateral, 10% extra-adrenal, 10% familial, 10% malignant). Prevalence of bilateral adrenal tumors is higher than 10% in some familial phaeochromocytoma syndrome such as multiple endocrine neoplasia type 2 and Von Hippel-Lindau Syndrome. In this patient population misdiagnosed as primary hypertension is frequent. Consequently they will lose the chance of early diagnosis and curative treatment.

We aimed to present this rare case with familial pheochromocytoma, family members had suffered from morbidity and death.

CASE REPORT
A 46-year-old man experienced headache, sweating and palpitation referred to our hospital. His arterial tension was measured 200 mmHg as
systolic, 100 mmHg as diastolic from the right arm. 190 to 80 from the left lower extremity measurements were 210/110 from both sides. We saw that his has a diagnosis of pheochromocytoma from his previous reports. Reports showed the scintigraphic findings consistent with the accumulation of activity of neuroendocrine tumor in the right adrenal region. He underwent surgery, and applied right adrenalectomy in 1988. He has been without complaints until 2009. He admitted to hospital with same symptoms. He lost 10 kg in 6 months.

At that time glucagon challenge test was applied, and his urine epinephrine was 149 mcg/24 h (0-20 mcg/24 h), norepinephrine was 324 mcg/24h (18-80 mcg/24 h), dopamine 147 mcg/24 h(65-400 mcg/24 h), metanephrine 21 mg (<1.3 mg/24 h), vanilmandelic acid (VMA) 7.6 mg/24 h (<6.8 mg/24 h), cortisol 132.5 mcg/dl, serum glucose 230 mg /dl. According to the test results he have received the diagnosis of type 2 diabetes mellitus.

His disease has been silent up to 2014 was admitted us with same symptoms. His urine epinephrine was 2.4 mcg/24 h, norepinephrine was 604 mcg/24 h, normetanephrine 1328mcg/24 h, metanephrine 222 mg/24 h, VMA 7.6 mg/24 h, cortisol 322 mcg/dl. A multilobule, 4.4.8 cm in diameter, and heterogeneous mass was detected in the left adrenal gland by contrast-enhanced CT. This mass was closely adjacent to the renal vein, and was successfully resected. Preoperative doxazosin, and postoperative prednisolone treatments were given. We invited the family members due to suffering from the same symptoms. All of them had a diagnosis of pheochromocytoma. However, familial transmission was not taken into consideration, so far. His grandfather was dead from pheochromocytoma. His brother and two sisters were stricken pheochromocytoma from his stepmother. One of his stepsisters was died due to paraganglioma and pheochromocytoma. Our patient has four brother. All of them have a diagnosis of pheochromocytoma. One of them was operated. Family tree is shown in the figure 1.

Figure1 The family tree for pheochromocytoma of patient.
DISCUSSION

Pheochromocytomas are rare neuroendocrine tumors with a highly variable clinical presenting. The serious and potentially lethal cardiovascular complications of these tumors are due to potent effect of secreted catecholamines. The most common clinical presentation are headaches, sweating, palpitation and hypertension. Hereditary pheochromocytomas occur in multiple endocrine neoplasia type 2 (MEN type 2), Von Hippel Lindau Syndrome, Neurofibromatosis type 1 and the familial paragangliomas. All patients with suspected pheochromocytoma should undergo biochemical testing. Standard biochemical test include measurements of urinary and plasma catecholamines, urinary metanephrines (normetanephrine and metanephrine) and urinary VMA. Tumor localisation should ideally only be initiated once there is unequivocal biochemical evidence for pheochromocytoma, CT scans or T2-weighted MRI for localisation of pheochromocytoma. Genetic testing should be done in all patients with pheochromocytoma. Genetic testing is recommended to those patients who have a positive family history or those who are younger than 50 years, especially children. Germline mutations in five genes have been identified to be responsible for familial pheochromocytoma: The VHL gene, the Ret gene, the NF 1 gene and the genes encoding the B and D subunit of mitochondrial succinate dehydrogenase (SDHB and SDHD). In our case; all these symptoms were observed. Due to clinical findings and family history like pheochromocytoma, we wanted to studied genetic diagnostic analysis. However, genetic analysis can not be performed because of technical difficulties.

CONCLUSION

Pheochromocytoma may relapse even in the late period as shown in our case. Therefore, physician should be sceptical in resistant and complicated hypertension. Methods for genetic diagnosis are rising gold standart techniques to determine the potential risks, malignancy of disease, and for the maintenance of treatment.

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