Hypertension and Intrapericardial Paraganglioma: An Exceptional Presentation of Multiple Endocrine Neoplasia Type IIA Syndrome

To the Editor:

Paracardiac paragangliomas are very rare. A catecholamine-producing intrapericardial paraganglioma has never been described previously in the context of multiple endocrine neoplasia (MEN) IIA syndrome.1,2 We report the case of a patient with MEN IIA syndrome and secondary hypertension caused by functioning paraganglioma exceptionally located intrapericardially.

A 34-year–old woman presented with recent episodes of headache, palpitation, and flushing. For several months she had arterial hypertension controlled by ramipril; however, she reported high blood pressure spikes (180/90 mm Hg). Moreover, her sister was diagnosed with medullary thyroid cancer. Examination of the patient’s neck demonstrated a small lump in the left thyroid lobe. Aspiration cytology of the nodule (1 cm) revealed a medullary thyroid carcinoma cell pattern. Thyrocalcitonin serum level was high (217.6 pg/mL; normal: 18 pg/mL), whereas calcium, thyroid-stimulating hormone, T3, and T4 levels were normal. A MEN syndrome was suspected, and additional exams were performed. Very high 24-hour urine normetanephrine level (4132 µg/24 hours; normal: 162–527 µg/24 hours) and vanilmandelic acid level (16.2 mg/24 hours; normal: 1–6.7 mg/24 hours) were found. Imaging studies were done to localize the suspected catecholamine-secreting tumor. Computed tomography scan of the chest demonstrated a contrast-enhanced 55-mm intrapericardial mass below the left atrium roof (Figure A and B). Intense123I-metaiodobenzylguanidine uptake concentrated solely in the thoracic mass (Figure C), supporting the diagnosis of intrapericardial functioning paraganglioma without metastases. Mutation of the proto-oncogene RET C618S was documented in the patient and in 5 of her family members (2 children, 1 sister, 1 uncle, and 1 cousin), confirming the diagnosis of MEN type IIA syndrome. Surgical resection of the intrapericardial tumor was carried out. At induction of general anesthesia, the patient had a hypertensive crisis (blood pressure: 210/100 mm Hg) and intraoperatively required multiple doses of phentolamine and labetalol. The tumor was approached by right thoracotomy and was resected without cardiopulmonary bypass. The resected specimen presented as a 45×55-mm pseudocapsulated oval mass, with cut surface showing multiple gray-tan nodules and hemorrhagic areas (Figure D). Pathology revealed a paraganglioma with a resting pattern. Immunohistochemistry was positive for chromogranin A and protein S-100, as well as for tyrosine-hydroxylase, dopamine β-hydroxylase, and met-enkephalin. The tumor was negative for calcitonin and phenylethanolamine N-methyl transferase.

The patient became rapidly well after surgery and no longer had hypertension. On postoperative day 4 her urine normetanephrine level was normal (167 µg/24 hours). After recovery from thoracotomy, total thyroidectomy was performed, and histology documented medullary thyroid carcinoma positive for calcitonin and chromogranin A, confirming the MEN II A syndrome. At follow-up after 9 months from paraganglioma resection the patient is well, without symptoms.

Extra-adrenal tumors producing catecholamines are called paragangliomas. Two histological types are described, chromaffin paragangliomas, usually producing catecholamines and causing hypertension, and nonchromaffin tumors, which are called chemodectomas.3 Extra-adrenal pheochromocytomas most commonly originate in the abdominal superior para-aortic region, between the dia-
phragm and the lower renal poles, frequently in the organ of Zuckerkandl. Only ≈2% of paragangliomas are located within the chest, more frequently in the posterior mediastinum and less frequently within the lungs or in the esophageal wall. Intrapericardial and intracardiac paragangliomas are very rare, usually arising from the left atrial wall or from the interatrial septum, and approximately half of them have been associated with signs or symptoms of excess catecholamine secretion.

Catecholamine-secreting tumors in the MEN II A syndrome are usually located in the adrenal glands. They have never been described previously within the pericardium.

Disclosures

None.

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