Primary ovarian carcinoid tumor. Case report

Tumor carcinoide primario de ovario. Reporte de caso

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ABSTRACT
Primary neuroendocrine tumors are rare. They belong to a group of heterogeneous neoplasms that express similar immunohistochemical markers. Carcinoid tumors are the most common neuroendocrine neoplasms. Most of them arise in the gastrointestinal and bronchopulmonary tract. Primary carcinoid tumors of the ovary are rare entities that represent approximately 0.3% of all carcinoid tumors and less than 0.1% of all ovarian neoplasms. They have an evident clinical presentation with symptoms such as skin redness, diarrhea, and bronchospasm. This tumor spreads to other organs through the bloodstream. The primary tumors are usually located in the ovary. The treatment is surgical, except for metastatic cases. The prognosis is usually good. 

Key words: Primary carcinoid tumor, Neuroendocrine tumor, Ovary, Carcinoid.

INTRODUCCION
Neuroendocrine tumors are epithelial neoplasms with predominant differentiation of neuroendocrine cells, which carcinoid tumors comprise the main subtype. Primary ovarian carcinoid tumors are rare, accounting for 0.3% of all carcinoid tumors and less than 0.1% of malignant ovarian tumors. They originate most frequently from the gastrointestinal and bronchopulmonary systems. Those appearing in the ovary may be primary or metastatic.

Most ovarian primary carcinoid tumors are unilateral and classified into four types: insular (the most common), trabecular, mucinous, and mixed. Neuroendocrine tumors release a large amount of serotonin into the systemic circulation and cause manifestations of carcinoid syndrome in the absence of metastasis. Its rarity within the genital tract may lead to late diagnosis unless patient's systemic symptoms are recognized for being related to pelvic pathology. Other types are generally not associated with the carcinoid syndrome. A case of a primary ovarian carcinoid tumor is presented.

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Declaration of ethical aspects
Acknowledgment of authorship: All authors declare that they have contributed to the study, study design, data collection, data analysis, and interpretation, critical review of the intellectual content, and final approval of the submitted manuscript.

Ethical responsibilities: Protection of human subjects: The authors declare that the procedures performed were in accordance with the ethical provisions of the corresponding human research ethics committee, as well as the World Medical Association and the Declaration of Helsinki.

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CLINICAL CASE

A 60-year-old female patient consulted because of exertional dyspnea after walking a few meters, lower limb edema and abdominal pain, which progressively increased in the previous six months, accompanied by abrupt facial redness that progressively worsened (around 15 episodes per day), and later diarrhea (four to five episodes per day), anorexia, night sweats, and abdominal distention, for the past five years. She reported a history of chronic arterial hypertension treated with calcium channel blockers. She denied chest pain, palpitations, paroxysmal nocturnal dyspnea, and other significant medical, surgical, or family history.

On physical examination, temperature was 37.7°C, heart rate 100 beats per minute, respiratory rate 16 breaths per minute, and blood pressure 145/95 mmHg. Heart sounds were rhythmic and without murmurs, and vesicular murmur was audible in both lung fields, with no aggregates. The abdomen was globule, with evidence of hepatomegaly, changing dullness, and a positive wave sign, with a palpable, hard and irregular tumor in the suprapubic area.

Electrocardiogram revealed sinus rhythm with an incomplete right bundle branch block, and plain chest X-ray showed mild cardiomegaly without signs of pericardial calcifications, with lung fields clear. Abdominopelvic ultrasound revealed the presence of a tumor in the right ovary measuring 11 x 8 x 6 centimeters with dominant peripheral vascularization, arterial and venous waves. Magnetic resonance images revealed a solid-cystic pelvic tumor of approximately 12 centimeters, apparently arising from the right ovary, with a scant amount of free fluid in abdominal cavity (Figure 1). No intestinal, liver, or bladder lesions were observed. The upper and lower endoscopy did not show evidence of gastrointestinal pathologies and chest tomography did not show tumors.

In view of the possibility of a neuroendocrine tumor, the following tests were ordered: urinary 5-hydroxyindoleacetic acid (5-HIAA) 65 mg / 24 hours (normal value (NV) 2 to 6 mg / 24 hours), serotonin 1 318 mcg / L (NV 80-400 mcg / L) and chromogranin A 1 130 mcg / L (NV 27 to 94 mcg / L). Gastrin, vasoactive intestinal peptide, glucagon, and calcitonin values were within normal limits. Hematology, renal function, electrolytes, coagulation profile, liver function, and tumor marker values alpha-fetoprotein, human chorionic gonadotropin, carcinoembryonic antigen, and CA-125 were all within normal limits.

Positron emission tomography showed a pelvic tumor with moderate to intense uptake of gallium-labeled octreotide, with no evidence of metastatic intestinal disease, suggesting the possibility of a primary ovarian carcinoid tumor.

Patient underwent laparotomy while being treated with octreotide (50 mcg / h 24 hours before surgery). An ovoid, solid, multinodular, and irregular tumor of approximately 12 centimeters originating from the right ovary, firmly adhered to the cul-de-sac, without apparent infiltration to adjacent structures was found. A frozen biopsy of tumor was performed, and squamous epithelium, mature respiratory epithelium and bone, cartilage, smooth muscle, adipose, and brain tissue (glial and choroidal tissue) were observed.

Given the findings, a total hysterectomy plus bilateral oophorosalpingectomy, peritoneal lavage, omentectomy, pelvic and para-aortic lymphadenectomy were performed. There were no complications during surgery and she was discharged on day four. Chromogranin A (17 mcg / L) and urinary 5-HIAA (2.8 mg / 24 hours) values were within normal limits six weeks after surgery. Symptoms also disappeared. Patient did not return for postoperative follow-up.
Macroscopic study showed a solid, smooth-surfaced, congestive tumor measuring $9 \times 6 \times 5$ centimeters confined to the right ovary, with intact capsule and weighing 152 grams (Figure 3). On section, a grayish, nodular area was observed, with no evidence of cystic changes. Microscopic evaluation showed uniform polygonal cells with eosinophilic granular cytoplasm and regular nuclei with little mitotic activity, arranged in solid sheets with a trabecular and insular pattern. These patterns were lined with layers of cells with central, homogeneous, and round nuclei with small nucleoli without mitosis or atypia (Figure 4). The amount of cytoplasm in the cells was large and strongly acidophilic. Immunostaining showed strong positivity for CK19 along with diffuse immunostaining for CD56 and localized immunopositivity for CK7, NSE, CDX2, and synaptophysin and negative for CK20, CEA, CA125, TTF-1, HNF1 beta, and MiB with Ki67 dispersed in less than 1% in the minor individual cell staining. Ultrastructural study of the cells revealed numerous intracytoplasmic neurosecretory granules. Final diagnosis was primary ovarian neuroendocrine carcinoid tumor.

**DISCUSSION**

Carcinoid tumors are well-differentiated neuroendocrine tumors that arise from enterochromaffin cells and secrete serotonin and other vasoactive substances. Most carcinoid tumors are found in the gastrointestinal tract (55%) and bronchopulmonary region (30%). The small intestine is the most common site (45%), followed by the rectum (20%), appendix (17%), colon (11%), and stomach (7%). Primary ovarian tumors are even rarer, accounting to less than 0.1% of ovarian malignancies and 1% of all carcinoid tumors.

Most carcinoid tumors are relatively slow-growing, but they often metastasize and some can behave aggressively. Approximately 19% of carcinoid tumors present with metastatic disease; the most common site is the liver, regardless of its primary origin. Even with significant hepatomegaly caused by infiltration of the liver parenchyma due to metastatic lesions, liver biochemistry may be within normal limits. Ovarian
Involvement can be primary or metastatic. Primary ovarian carcinoid tumors are generally unilateral and are composed of epithelial elements of gastrointestinal or respiratory origin. They often arise within a cystic teratoma or dermoid tumor, and up to 60% coexist with these tumors.

Primary ovarian carcinoid tumors are accidentally discovered in most women by ultrasound imaging. Patients rarely present with abdominal pain, constipation, hirsutism, or a pelvic mass. Clinical entity is probably multifactorial and mediated by vasoactive hormones, such as serotonin, tachykinins, kallikrein, and prostaglandins. All these substances are metabolized and inactivated by hepatic monoamine oxidases. However, in some cases, presence of liver metastases is not required for the development of the syndrome, since these substances can reach systemic circulation directly when the primary tumor is in the lung or the ovary. In the latter case, vasoactive substances reach the systemic circulation through the inferior vena cava (right ovary) or renal vein (left ovary).

Diagnosis of carcinoid tumors requires multiple imaging modalities and biochemical tests if there are symptoms of carcinoid syndrome. Computed tomography (76% sensitivity, 96% specificity) and ultrasound (93% sensitivity, 95% specificity) are useful for locating the tumor, although they cannot identify whether the lesion is carcinoid. Functional imaging or photon emission tomography can confirm diagnosis, identify metastases, and stage of disease. This requires administration of radiolabeled octreotide, a synthetic analog of somatostatin that is absorbed by carcinoid tumors. These tests have a sensitivity of 94%.

Those with symptoms suggestive of carcinoid syndrome may have elevated 24-hour urine 5-HIAA (the main breakdown product of serotonin) as a screening test. Elevated urinary concentrations confirm presence of the syndrome and guide the use of imaging to locate and stage the tumor. Chromogranin A can also be used as a marker, but its specificity is lower because it is also secreted by pheochromocytomas.

Carcinoid tumors are characterized by growth patterns that suggest neuroendocrine differentiation, including organoid, insular, trabecular, palisade, ribbon, and rosette arrangements. Individual tumor cells have uniform cytologic features with moderate, finely granular eosinophilic cytoplasm and nuclei with a finely granular chromatic pattern. Histological criteria to discriminate between typical and atypical ovarian carcinoid tumors have not been established. Since ovarian carcinoid incidence is very low, with no cases of aggressive histological characteristics reported, there was no prior need to establish criteria for atypical carcinoid tumor of the ovary. Primary ovarian carcinoid tumors composed of more than one pattern may be classified as primary mixed carcinoid tumors, regardless of the predominant pattern.

Trabec-
Primary ovarian carcinoid tumors neurosecretory granules are round to oval and show slight variations in size, thus differing from marked variations seen in insular carcinoid tumors. Immunocytochemical analyses of trabecular carcinoid tumors show a much broader range of neurohormonal polypeptides compared to those in insular carcinoid tumors[13].

Primary ovarian carcinoid tumors must be differentiated from metastatic carcinoid tumors or cord-like pattern Sertoli-Leydig cell tumors. Metastatic carcinoid tumors are bilateral, with peritoneal metastases and poor prognoses; presence of teratomatous elements help to exclude a metastatic lesion[13]. The most common subtypes that metastasize to the ovary are insular and trabecular[7]. Sertoli-Leydig cell tumor shows similar cordon patterns and, unlike trabecular carcinoids, has inhibin-positive tubular formations and absence of neurosecretory granules[13].

Therapeutic options are complex. Treatment with somatostatin analogs accompanied or not by tumor reduction-resection may improve symptoms and the negative hemodynamic changes on cardiac function[8]. Somatostatin analogs inhibit the release of biogenic amines and peptides, including serotonin, and relieve symptoms. Surgery is the standard treatment for resectable tumors, although it is not often possible in patients with metastatic disease. Cytoreductive surgery and chemotherapy are often necessary to prolong survival. Octreotide is useful in patients with symptoms of carcinoid syndrome to relieve symptoms and prevent progression of heart disease[14]. Perioperative treatment should include intravenous infusion with octreotide to reduce the risk of carcinoid crisis. Antihistamines and corticosteroids may be helpful in controlling flushing and bronchospasm. There is no evidence to support the use of adjuvant treatments (hormones, chemotherapy, or radiation)[8].

Prognosis is excellent for stage I ovarian carcinoid tumors with 5-year survival rate of 90%. Metastatic carcinoid tumors (most commonly of small intestine) tend to be bilateral, insular type, with tumor deposits in both ovaries. In this group of patients, prognosis is poor since survival rate is less than 50% at five years. The 3-year mortality rate for patients with carcinoid syndrome and heart disease is 31%, while those without heart disease have approximately twice the survival rate[15].

In conclusion, primary ovarian carcinoid tumor is a very rare neuroendocrine neoplasm, which should be considered in patients with carcinoid syndrome symptoms with location other than the gastrointestinal tract or lungs. Patients with these ovarian tumors and systemic symptoms must be evaluated for endocrinological-cardiac status and undergo early surgery, to avoid progression to cardiac syndrome. Symptoms should be considered to facilitate early diagnosis and treatment of this curable malignancy, with excellent survival rates.

**References**


