A Rare Abdomino-Pelvic Tumor: Paraganglioma

Um Caso Raro de Tumor Abdominopélvico: Paraganglioma

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ABSTRACT
Paragangliomas are rare tumors, with a reported incidence of 2–8 per million. They are chromaffin cell tumors that develop from the neural crest cells and may be divided in tumors derived from the parasympathetic or sympathetic ganglia. We report a case a of a 32-year-old nulliparous woman, referred to our Infertility Clinic. Abdomino-pelvic ultrasound identified a large abdominopelvic tumor, without ovarian origin (both ovaries were identified and had normal morphology). Magnetic Resonance Imaging suggested a right adnexal multicystic, vascularized mass close to iliac vessels and questioning an ovarian origin. At exploratory laparotomy, a 10 cm encapsulated and vascularized mass was found beginning just below right renal artery and extending to the level of the broad ligament. This mass was totally excised and histopathology was consistent with Paraganglioma.

Keywords: Abdominal Neoplasms; Paraganglioma; Pelvic Neoplasms.

INTRODUCTION
Paragangliomas are rare tumors, with a reported incidence of 2–8 per million.¹ They are chromaffin cell tumors that develop from the neural crest cells and may be divided in tumors derived from the parasympathetic or sympathetic ganglia. These tumors comprise 10–18% of all chromaf fin tumor.² Parasympathetic ganglia-derived tumors are found almost exclusively in the neck and skull base and arise within the carotid body and globus jugulotympanic. The best known and most common sympathetic ganglia-derived tumors grow within the adrenal medulla and are known as pheochromocytoma. In contrast, sympathetic paragangliomas, also known as extraadrenal pheochromocytomas, arise outside the adrenal gland and can be found anywhere along the sympathetic chain from the base of the skull and neck (5% of cases) to the bladder and prostate gland (10%). Of those found along the aorta, 10% are in the thorax and 75% are in the abdomen, with most of the latter located in the organ of Zuckerkandl (centered around the root of the inferior mesenteric artery).

The authors report a case of a large inferior para-aortic paraganglioma in a 32-year-old woman.

CASE REPORT
A 32-year-old healthy nulliparous woman with a history of primary infertility of 2 years followed at our Infertility Clinic was submitted to routine pelvic ultrasound study which revealed a uterus with 3 small fibroids, normal right ovary and a large abdominopelvic mass of solid and cystic nature, measuring 64.4 x 52.6 x 57.1 mm (Fig.s 1 and 2).

Magnetic Resonance Imaging (MRI) revealed a multicystic, vascularized adnexal mass with 60 x 35 x 104 mm located close to the iliac vessels and questioning a right ovarian origin. Although the patient had a normal body mass index her first physical examination did not reveal any abdomino pelvic mass. Evaluation of blood pressure was also normal. Blood tests showed normal values, including tumor markers levels such as carcinoembryonic antigen, carbohydrate antigen 125, and α-feto-protein. Based on magnetic resonance imaging report the patient was offered an exploratory laparotomy.

At surgery a 10 cm encapsulated and vascularized solid and cystic mass was found beginning just below right renal artery and extending to the level of the broad ligament. Both ovaries were morphologically normal. The tumor was completely excised (Fig. 3).
Histopathology revealed a solid tumor of 11.5 x 6.5 x 5.5 cm, nests of cells with a trabecular and alveolar pattern consistent with a Paraganglioma (Figs. 4 and 5).

**DISCUSSION**

Paraganglioma is the generic term applied to tumors of paraganglia regardless of location. Paragangliomas of the adrenal medulla, the most common site of paragangliomas, are known as pheochromocytomas while those located outside the adrenal gland are known as extra-adrenal pheochromocytomas/paragangliomas. Extra-adrenal paraganglia are divided into two categories: those related to the parasympathetic system and those connected with the sympathetic system. The former, usually nonchromaffin, are found mainly along the supradiaphragmatic branches of the vagus and glossopharyngeal nerves and are believed to have a chemoreceptor function. The latter are chromaffin, associated with the peripheral sympathetic nervous system, and produce catecholamines in response to sympathetic neural stimulation. They lie along the thoracolumbar paravertebral region, from high in the neck, near the superior cervical ganglion to the abdomen and pelvis, predominantly in the posterior mediastinum and retroperitoneum. Rarely, paragangliomas have been described in unusual sites, such as the gallbladder, mesentery, kidney, prostate and ovary. The paraganglioma in our patient was located in the area of Zuckerkandl organ, extending to the pelvic cavity.

Most paragangliomas are diagnosed in the third to fifth decade of life. Extra-adrenal paragangliomas present with abdominal pain and/or palpable abdominal mass however, they may be asymptomatic even when associated with large masses, as our patient. About 15% of patients are asymptomatic and the diagnosis is usually done in the context of diagnostic imaging procedures for other purposes (incidentalomas). Other symptoms include nausea, vomiting, diarrhea, abdominal distension and weight loss.

The majority of abdominal pheochromocytomas and paragangliomas are benign; malignant pheochromocytomas account for nearly 10% of all pheochromocytomas.

Clinical, biochemical and radiological features are inadequate to predict malignancy which can be established by the presence of distant metastases mainly to the liver, lymph nodes, lung and/or bone, either at diagnosis or during follow-up. Local invasion and various histopathological features can be suggestive; however, these features are not widely accepted and there is a need for the development of more sensitive and specific diagnostic means. This said, the lack of firm predictors of malignancy, coupled with the variable course of this rare disease, make life-long follow-up of patients with chromaffin-cell tumors mandatory.

Contrast-enhanced abdominal computed tomography (CT) scan is useful for diagnosis; however, there are no specific features for this condition. Abdominal MRI and scintigraphy performed with radiotracer-labeled metaiodobenzylguanidine (MIBG) are essential in the identification and characterization of paragangliomas.
Figure 4 - Histopathology: Paraganglioma (trabecular pattern)

Figure 5 - Histopathology: Paraganglioma (alveolar pattern)

(distinguishing functional from non-functional tumors). MRI has the highest sensitivity in detection of extra-adrenal paragangliomas and pheochromocytomas. The primary treatment for paragangliomas is complete surgical resection. Conventional treatment consists of open exploration and resection of the mass. Annual biochemical testing (plasma catecholamines, plasma-free metanephrines, urinary catecholamines, urinary vanillylmandelic acid, urinary total metanephrines, and urinary fractionated metanephrines), CT scans and/or MRI and MIBG scans are essential in the assessment for metastatic disease, tumor recurrence or delayed appearance of multiple primary tumors.

CONCLUSION

Paraganglioma of the abdomino-pelvic cavity is an extremely rare tumor. Clinical and imaging data of patients with extra-adrenal, intra-abdominal paragangliomas are variable. Many of them may be asymptomatic even when the lesion is quite large.

It may appear as an incidental finding on gynecological ultrasound examination, especially in a tertiary women’s hospital such as ours.

These cases highlight how paraganglioma may be mistaken for gynecological masses. A careful approach during sonography, depending on the operator, may in some cases better exclude the ovarian origin of adnexal masses than MRI. Although it is important not to forget that MRI is important in differential diagnosis of adnexal masses and has the highest sensitivity in detection of extra-adrenal paragangliomas and pheochromocytomas.

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CONFLICTS OF INTEREST

The authors declare that there are no conflicts of interest.

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