Postmenopausal Bleeding with an Accidental Discovery of an Ovarian Strumal Carcinoid Tumor: A Case Report

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Authors’ contributions

This work was carried out in collaboration among all authors. Author EMI selected the case and prepared the images. Authors SSEA and FIE-S reviewed the patient detailed data and shared in collecting the literature. Author HAFH Shared in collecting the literature and writing the draft. Author AAFH selected the needed literature and wrote the manuscript. All authors read and approved the final manuscript.

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ABSTRACT

Aim: Ovarian strumal carcinoid is a tumor formed of thyroid tissue and carcinoid elements. It represents less than 2% of all ovarian tumors and less than 5% of mature teratomas. Physicians in many locations may not be aware about this pathology.

Presentation of Case: A multiparous 78-year-old woman presented by postmenopausal vaginal bleeding. The abdomen& pelvic sonar detected atrophied endometrium and a right adnexal mass with mixed echogenicity and increased blood flow, a finding that was confirmed later with a pelvic MRI which revealed a well-defined mass of abnormal signal at the right adnexa showing cystic

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changes contacting anteriorly the related intestinal loops and the urinary bladder, while contacting posteriorly the recto-sigmoid. It measured 8x6x6 cm. Biopsy revealed atypical epithelial proliferation. CA125 was 90KU/L (normal range, 0-35KU/L) while the levels of CEA, CA19-9 and alpha fetoprotein were normal. After discussion with the family, exploratory hysterectomy and bilateral salpingo-oophorectomy, along with omentectomy, peritoneal washing and peritoneal sampling were performed. Grossly, the right ovary was largely occupied by an ovoid mass (9cmx6cmx2 cm). The tumor was staged as IA according to AJCC 2010. Histological examination of the adnexal mass revealed admixture of benign thyroid tissue and tumor tissue formed of trabeculae & nests of monotonous round cells. Such cells showed positivity for thyroglobulin, synaptophysin, chromogranin and low positivity for ki67 and so a diagnosis of strumal carcinoid tumor was documented. The patient was set to a close follow up. The last follow up was at August 2020 and was satisfactory.

**Discussion and conclusion:** Strumal carcinoids are rare. Pelvic sonar and CA125 are not specific diagnostic tools. They should be considered in the differential diagnosis of any ovarian mass. The disease usually presents at an early stage.

**Keywords:** Ovarian teratoma; strumal carcinoid; Carcinoid syndrome.

**1. INTRODUCTION**

Mature cystic teratomas (MCT) are the most common among all ovarian neoplasms, representing 30-40% of the cases [1]. Rarely MCT, undergo malignant transformation (0.94-2%) [2]. Struma ovarii is a variant of MCT, with a predominant thyroid element. Confirmatory diagnosis is through histopathology [3]. Malignant transformation in struma ovarii is uncommon and mostly in the form of follicular thyroid carcinoma [4].

**2. PRESENTATION OF THE CASE**

A multiparous 78-year-old woman presented by vaginal bleeding without any criteria of a carcinoid syndrome. The woman had an uncomplicated medical and gynecological history with regular menstrual cycle. The general examination was uneventful. Abdominal and pelvic examination revealed a firm right adnexal mass of limited mobility. The abdomen& pelvic sonar detected an atrophied endometrium and a right adnexal mass with mixed echogenicity and increased blood flow. The biopsy of which (may 2018) revealed atypical epithelial proliferation. MRI of abdomen and pelvis revealed a well-defined mass of abnormal signal at the right adnexa showing cystic changes contacting anteriorly the related intestinal loops and the urinary bladder, while contacting posteriorly the recto-sigmoid. Superiorly, it was contacting the related right iliac vessels. It measured 8x6x6 cm. Biopsy revealed atypical epithelial proliferation. Unfortunately, we found in the archive the reports of the sonography and the MRI but not the films. Ca125 was 90ku/l (normal range, 0-35ku/l) while the levels of CEA, CA 19-9 and alpha fetoprotein were normal. CT scan of chest was free. After discussion with the patient and her family, exploratory hysterectomy and bilateral salpingo-oophorectomy, along with omentectomy, peritoneal washing and peritoneal sampling were performed. Fortunately, the postoperative period was uneventful, and the patient was discharged on the fifth postoperative day.

Microscopic examination revealed a compressed normal ovarian tissue with underlying tumor tissue. The tumor was composed partly of thyroid tissue with thyroid follicles filled with colloid and partly of tumor cell clusters composed of monomorphic cells of granular cytoplasm. The nuclear chromatin was finely granular and mitosis was hardly detected. The tumor cells were arranged in trabecular structures or in solid nests (Fig. 1). Moreover, immunohistochemistry revealed positivity for thyroglobulin, synaptophysin, chromogranin and low positivity for ki67 (Fig. 2). The endometrium was atrophic.

The patient was set to a close follow up with gynecological examination and ultrasound imaging every 3-6 months and CT scan of abdomen& pelvis every 6-12 months in the first year following surgery. The follow up during the
following years was recommended to be less frequent. The last follow was satisfactory and was in August 2020.

The Patient Gave Her Consent To Publish The Case.

The Institutional Review Board Approved The Work.

3. DISCUSSION

Carcinoid tumors are usually a slow-growing type of neuroendocrine tumors. It is worth to note that neuroendocrine tumors originate mainly from gastroenteric tissue, pancreatic islet cells, neuroendocrine cells within the respiratory epithelium, and parafollicular cells distributed within the thyroid. They occasionally have secretory characteristics, with a rather unusual and complex disease spectrum with various manifestations, such as flushing, diarrhea, hirsutism, or constipation [5]. Our case did not present with any of these symptoms. Her postmenopausal vaginal bleeding may be due to the atrophic endometrium [6]. Primary carcinoid tumors of the ovary account for <0.1% of all ovarian malignancies, and only 5% of carcinoid tumors are of the ovarian origin. The ovarian carcinoid tumors are typically divided into 4 categories based on the histopathologic patterns: insular, trabecular, mucinous, and strumal [7]. On searching through the PubMed for strumal carcinoid tumors, 100 publications were retrieved. However, none of them were related to our locality. The rate of malignant transformation in MCT worldwide is much higher in postmenopausal ages [8] which coincides with the age of our case. In less than 10% of the published stromal carcinoids, there was a contralateral neoplasm of the ovary [9]. We did not find a contralateral ovarian pathology. Our case was stage I similar to most literature [10]. Coexisting of graves' disease and functioning struma ovarii is a rare condition [11]. One of the limitations of this case report is that no thyroid function tests were documented.

Fig. 1. Histological features of the ovarian tumor
A-admixture of benign thyroid tissue (left) and tumor tissue (right) (H&E; x10)
B-the tumor cells are arranged in acinar and trabecular structures (H&E; x10)
4. CONCLUSION

Strumal carcinoid rarely occurs. It is most probably even rarer in our locality. Pelvic sonar and CA125 are not specific diagnostic tools. Strumal carcinoid should be considered in the differential diagnosis of an ovarian mass in postmenopausal patients.

CONSENT

The patient gave her consent to publish the case.

ETHICAL APPROVAL

The institutional review board approved the work.

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COMPETING INTERESTS

Authors have declared that no competing interests exist.

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