**RÉSUMÉ**

Struma ovarii chez une femme de 56 ans – rapport de cas

**Introduction.** Struma ovarii représente une tumeur rare, seulement 1% des tumeurs de l’ovaire, avec une incidence de 0,3 à 0,7%. Le diagnostic positif est obtenu par examen microscopique; parfois, des taux sériques élevés d’hormones thyroïdiennes, de CA 125 et d’échographies peuvent suggérer un diagnostic préopératoire.

**Case presentation.** Nous rapportons le cas d’une femme de 56 ans avec une struma ovarii unilatérale et aucune preuve clinique ou paraclinique du diagnostic à venir. L’échographie transvaginale a révélé un utérus polyfibromateux et des annexes apparentment

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**ABSTRACT**

**Introduction.** Struma ovarii represents a rare tumor, only 1% of the ovarian tumors, with an incidence of 0.3-0.7%. The positive diagnosis is obtained by microscopic examination; sometimes elevated serum levels of thyroid hormone, CA 125 and ultrasound aspects can suggest the preoperative diagnosis.

**Case presentation.** We report the case of a 56-year-old woman with unilateral struma ovarii and no clinical or paraclinical evidence of the diagnosis to come. Transvaginal ultrasound revealed polypfibromatous uterus and apparently normal adnexa. Partial hysterectomy with bilateral salpingo-oophorectomy (considering the patients age) through laparotomy was performed under spinal anaesthesia. The histopathological result showed an endometrial polyp, multiple

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**CASE REPORT**

**STRUMA OVARIII IN A 56-YEAR-OLD WOMAN – A CASE REPORT**

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INTRODUCTION

Struma ovarii is a mono-dermal teratoma composed predominantly or solely of thyroid tissue. This type of tumor accounts for 3% of ovarian teratomas. It is encountered in the reproductive age, with most patients being in the fifth decade of life. It is found almost always unilaterally, in only 10% and 15% being bilateral. It is most commonly asymptomatic, seldom clinically discernable as a palpable pelvic mass and even less frequently associated with ascites (1/3) or Meigs syndrome. On some occasions, hyperthyroidism is present. High serum levels of CA – 125 have been reported. At ultrasound examination, it can appear complex and nonspecific. However, low-resistance blood flow or good vascularized solid component in the central portion are important clues for the diagnosis of struma ovarii.

On gross examination, the tumor has a color that can vary from red to brown and green and a size of no more than 10 cm. It consists of predominantly solid, soft tissue.

Like the thyroid tissue it emulates, struma ovarii is made of hundreds of thousands of follicles lined by cuboidal or low columnar epithelium. Cytology is usually with minimal atypia and low mitotic activity. One can encounter variants such as the microfollicular, the pseudotubular or the solid pattern. The latter can be composed of oxyphilic (abundant eosinophilic cytoplasm) or clear cells (pale cytoplasm).

Association with dermoid cysts, mucinous tumors or Brenner tumors has been described. Malignant change in struma ovarii most commonly leads to papillary or follicular carcinoma, but it is still unclear whether the criteria used in the thyroid gland should be applied. Thyroglobulin and TTF1 are essential immunohistochemical markers for a struma diagnosis, with the former being more specific. When suspecting a carcinoid, additional stains such as chromogranin or synaptophysin are useful in differentiating the two entities. Histologic heterogeneity makes the differential diagnosis more difficult. The most important neoplasm a pathologist should exclude are ovarian cystadenoma (when confronted with a cystic struma), steroid cell tumours, carcinoid tumours, Sertoli-Leydig cell tumours, renal clear cell carcinoma, and metastatic melanoma (in case of oxyphilic struma).

Differential diagnoses include: clear cell carcinoma (primary or metastatic from the kidney), primary or secondary hydatid cyst, metastatic tumors from renal sarcoma, endometrioid carcinomas, Sertoli cell tumour, hepatoid yolk sac tumour, malignant melanoma, serous cystadenoma, pregnancy luteomas, metastatic thyroid carcinoma of the ovary, secondary tumors from retroperitoneum, GIST tumors, primary or secondary carcinoid tumors, tumoral or benign appendiceal pathology.

Most often, the prognosis is favourable, with typical struma ovarii being benign. Even in the small percentage of histologically malignant results – 5-10%, the outcome is favourable, with only few patients who die of this disease. The strumal component, abundant ascites, adhesions and defects of the ovarian serosa are correlated with the malignant type. Only half of the malignant tumors associate extension beyond the ovaries.
The recommended treatment is oophorectomy and, in case of malignant struma, extraovarian tumor removal is advised².

Patients are advised to have a long-term follow-up².

The aim of this case report is to show the diagnostic stages and treatment of a rare case of struma ovarii, which represents no more than 1% of all ovarian tumours and 3% of all dermoid tumors¹.

CASE PRESENTATION

A 56-year-old woman was admitted to the Department of Obstetrics and Gynaecology of “St. Pantelimon” Emergency Clinical Hospital, Bucharest, Romania, on November 2018, for persistent vaginal bleeding during menopause. From the patient’s personal history, we noted one birth through C-section in 1994, the occurrence of menopause at the age of 51 years, obesity stage II, fibroadenoma of the left breast with surgical intervention in 2000, hypercholesterolemia, hypertriglyceridemia, smoker (20 cigarettes/day). Dilatation and uterine curettage were performed in May 2018, with the histological result of endometrial polyp.

On admission in our department, the patient was cooperative, the blood pressure was 120/75 mmHg, heart rate 73 beats/minute, with moderate vaginal bleeding. The gynaecological examination with the speculum showed no macroscopic lesions on the cervix and moderate bleeding coming from the uterine cavity. On the bimanual examination, the uterus was firm, with increased volume, irregular contour, mobility was preserved, sensitive at mild palpation, bilateral adnexa were normal. Laboratory exams were in normal range. Transvaginal ultrasound was performed and revealed a polyfibromatous uterus and apparently normal adnexa.

The informed consent was obtained and exploratory surgery through laparotomy was performed under spinal anaesthesia (spinal block/ intradural block/ intrathecal block) and followed by partial hysterectomy.

Figure 1. Thyroid tissue: intraovarian colloid. HE staining, x4.

Figure 2. Ovary with corpus albicans and thyroid follicles. HE staining, x10.

Figure 3. Thyroid follicles in the ovary. HE staining, x20.

Figure 4. Struma ovarii – thyroid follicles. HE staining, x4.
with bilateral salpingo-oophorectomy, considering the age and examinations prior to the surgery.

The postoperative evolution was favourable, without any complications. On day 7, the patient was released from the hospital with good general condition and afebrile.

The histopathological result showed an endometrial polyp with glands composed of unistratificate columnar epithelia, some of them cystically dilated, multiple intrauterine leiomyomas, left ovary which showed lobular display and consisted of thyroid follicles, which looked like normal thyroid tissue. The cells were cuboidal to columnar and dense colloid is seen inside the follicles (Figures 1-4). The right ovary was sclera hyalinized and both salpinges were atrophic.

At the 6 months’ follow-up of the patient no complications occurred and the laboratory investigations were within normal limits.

DISCUSSION

The clinical diagnosis of struma ovarii is difficult due to the rarity of the disease. Most of the times the patient is asymptomatic, with no abnormal paraclinical investigations. This is the case of our patient, who had normal serum levels of the thyroid hormones or CA 125, with normal vaginal ultrasound of the ovaries and no ascites. If the preoperative diagnosis occurs, laparotomy surgery is advised, due to the risk of tumor rupture intra-abdominally with dissemination, in case of a malignant tumor.

Oophorectomy is preferred and in case the malignant struma ovarii is confirmed, second intervention is planned and scheduled for pelvic and para-aortic lymph nodes sampling, peritoneal cytologic washing, partial or total omentectomy. Thyroidectomy and follow-up with Iodine-131 whole body scanner are recommended. In some cases, fertility can be preserved as needed and follow-up with the intervention as explained earlier.

The postoperative management of the patients after surgery is difficult, due to the rarity of the cases. Both guidelines of ovarian cancer and thyroid cancer are used.

CONCLUSIONS

The treatment of choice has to be carefully selected because of the high possibility of clinically misdiagnosing this pathology. If the diagnosis is established, laparotomy is advised due to the better accessibility and manipulation of the tumor by the surgeon. Fertility can be preserved, but caution must be implied. The majority of the patients has a good outcome.

Compliance with Ethics Requirements:

“The authors declare no conflict of interest regarding this article”

“The authors declare that all the procedures and experiments of this study respect the ethical standards in the Helsinki Declaration of 1975, as revised in 2008(5), as well as the national law. Informed consent was obtained from the patient included in the study”

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