Case report and review of the literature

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Abstract

Background: The present of malignant transformation in struma ovarii is exceedingly rare. Malignant struma ovarii is usually asymptomatic and infrequently diagnosed preoperatively. Because of its rarity, there is no consensus about diagnosis and management in the literature.

Case presentation: A 40-year-old female presented for her obstetric examination with an incidental finding of a pelvic mass. Patient was asymptomatic at presentation. A follow-up ultrasound confirmed the presence of a 3-cm mass in the left adnexa. Patient underwent a cytoreductive surgery (hysterectomy, bilateral salpingectomy and oophorectomy, omentectomy, appendectomy, and pelvic lymphadenectomy). Histopathology revealed a malignant struma ovarii with a focus of papillary thyroid carcinoma and the omentum metastasis. The patient with stage FIGO IIIc received 6 cycles of paclitaxel/carboplatin regimen after surgery. The patient subsequently had a thyroid scan that was normal with normal thyroid function. At a follow-up of 12 months, she is alive, in good clinical condition, and disease-free.

Conclusions: Because of the rarity of these tumors and their lack of firm prognostic factors, treatment decisions should be made individually, based on pathologic and clinical parameters.

Keywords: Struma ovarii, Papillary thyroid cancer, Metastases

Background

The majority of germ cell tumors are mature cystic teratomas, which account for approximately 15–20 % of all ovarian tumors [1]. Of these, 15 % contain thyroid tissue [2, 3]. Struma ovarii is diagnosed when thyroid tissue comprises more than 50 % of the teratoma [4, 5]. Struma ovarii accounts for only 2 % of all mature teratomas, and less than 5 % of struma ovarii present malignant transformation [4, 6, 7]. Owing to malignant struma ovarii (MSO) rarity, there has been some controversy about the diagnosis and treatment, and prognosis is difficult to evaluate.

We report an unusual case of papillary thyroid cancer in MSO, with metastasis to the omentum. Here, we presented a review of literature including the clinicopathologic features, differential diagnosis, and management.

Case presentation

A 40-year-old female presented with an incidental finding of a pelvic mass during her obstetric examination before 6 years. A palpable mass was noted in the left-sided pelvis on physical examination. The patient did not have special symptoms and ascites. Abdominal ultrasonography (Voluson S8, General Electric Company, USA) revealed a 3.0 × 3.1 cm mixed echogenicity mass in the left adnexa, suggestive of an ovarian teratoma. The data obtained by routine blood, urine, and thyroid function tests, as well as tumor markers, were in the normal range. The patient had no history of thyroid...
dysfunction and no family history of ovarian or endocrine
tumors. The patient had no relevant past interventions.
Malignancy could not be excluded.

The patient underwent laparoscopic ovarian cyst
divesting surgery. At exploration, the uterus, right ovary,
and fallopian tube were normal. The left ovary was en-
larged, and a benign-looking white and smooth ovarian
cystic mass was noted. Upon sectioning, the ovarian mass
was partially cystic and grossly resembled papillary tumor.
The peritoneal washing was negative on cytological exam-
ination. An intra-operative frozen and permanent paraffin
section was proved to be a malignant struma ovarii with
multiple foci of papillary thyroid carcinoma (Fig. 1a). In
the papillary thyroid carcinoma focus, papillary thyroid
tissue with increased mitotic activity and ground glass
nuclei, intra-nuclear inclusions were detected. Also, a few
psammoma bodies were identified. The tumor was re-
stricted within the capsule, and vascular invasion was not
identified. Immunohistochemically, tumor cells were
strongly positive for thyroglobulin (Tg), thyroid transcrip-
tion factor-1 (TTF-1), galetin-3, cytokeratin-19 (CK19),
and human bone marrow endothelial cell-1 (HBME-1)
(Fig. 2). Based on all these findings, the diagnosis was pap-
illary thyroid cancer arising within malignant struma
ovarii. Thus, 1 week later, the patient underwent a cytore-
ductive surgery (hysterectomy, bilateral salpingectomy
and oophorectomy, omentectomy, appendectomy, and
pelvic lymphadenectomy). The pathologic examination
showed no metastatic neoplasm was found, except on the
omentum material (Fig. 1b). After surgery, the patient
with stage FIGO IIIc received 6 cycles systemic chemo-
therapy of paclitaxel (175 mg/m²; over 3 h intravenous in-
fusion) plus carboplatin (area under the curve [AUC] = 5;
over 1 h intravenous infusion), given on day 1 of a 21-day
cycle. Patient subsequently had a thyroid scan that was
normal with normal thyroid function.

Discussion

Struma ovarii is a teratoma in which thyroid tissue is
present exclusively or forms a grossly recognizable com-
ponent of a more complex teratoma [8]. The rate of ma-
lignant transformation in struma ovarii is extremely low.
The mean age at diagnosis of MSO was 43 years old [6, 9].
Most cases of MSO are subclinical. Clinical hyperthyroid-
ism manifestations are seen in 5–8 % of the cases [10].
Moreover, insufficient concordance between morphologic
features and clinical outcome in MSO is striking, making
the behavior of these tumors particularly unpredictable
[11]. It is reported that BRAF V600E gene mutations, a
common pathogenesis for all papillary thyroid cancers, re-
gardless of body location, were present in two thirds of
MSO with papillary features [12].

The predominant sites of metastasis were adjacent pelvic
structures, including the contralateral ovary; hematologic
dissemination includes metastases of the lung, bone, liver,
and brain, but metastasis is rare in patients with MSO. In
this case, the patient had omentum metastasis. The reason
is that the tumor can spread via regional lymphatics to pel-
vic and paraaortic lymph nodes and can directly infiltrate
the omentum [11, 13].

Since it is a rare type of germ cell tumor encountered,
there are no uniform diagnostic criteria for MSO. MSO
is most often diagnosed postoperatively. The histopatho-
logical diagnosis adheres to the same criteria used for
thyroid carcinoma: “ground glass” overlapping nuclei
and nuclear grooves, or mitotic activity and vascular in-
vasion [4]. Eighty-five percent of papillary carcinomas
with the characteristics are “ground glass” nuclei. All
pathologic patterns of thyroid gland malignancy may be
found in struma ovarii, and papillary carcinoma is the
most common [14–16]. Immunohistochemical staining
with Tg, HBME-1, and galectin-3, often positive in pap-
illary thyroid carcinoma, can also help confirm the

Fig. 1 H&E staining of malignant struma ovarii tissues. H&E staining (a) showed fibrous cores of tissue lined by neoplastic cells which
contain large overlapping “ground-glass” nuclei characteristic of thyroid papillary carcinoma. b Metastasis to omentum is shown (H&E,
original magnifications ×400)
diagnosis and are necessary for differential diagnosis from carcinoid and granulosa cell tumors [17, 18].

The absence of a primary lesion in the thyroid is necessary to exclude metastatic thyroid carcinoma to the ovary [4, 11]. The differential diagnosis from thyroid-type cancers from the thyroid and ovary depends on immunohistochemical stains for CK19 and TTF-1, supporting the independent existence of two cancers. Besides family history and a physical examination followed by an ultrasound of the thyroid gland, it is reported that unique characteristics on ovarian magnetic resonance imaging (MRI) might help to differentiate the two diseases.

The optimal treatment regimen for MSO is currently debated due to the difficulty to distinguish between benign and malign struma ovarii. Most authors advocate for an aggressive treatment based on TAH/BSO, lymph-node dissection and omentectomy, followed by adjuvant therapy, including external radiotherapy, chemotherapy, and thyroid suppression, regardless of the presence of distant metastases at time of diagnosis.

MSO might be over-treated, and this might impact long-term the quality of life and fertility of affected women. Thus, conservative treatment such as unilateral oophorectomy should be the choice for patients who wish to fertility preservation, if pelvic imaging and surgical exploration do not reveal extra-ovarian disease [9, 19].

In addition, on the basis of the similarities between MSO and thyroid carcinoma, near-total thyroidectomy followed by radioactive iodine (I\textsuperscript{131}) should be first-line therapy for patient with a higher risk of recurrence or with palindromia [12]. Treatment with thyroxine was not documented consistently enough to determine its impact on recurrence rates.

Patients with MSO had an excellent disease-specific survival rate, regardless of the management strategy employed. In 2015, Adam et al. published a study that included 68 patients with MSO [9]. The data indicated that overall survival rates for all patients were 94.3 % at 10 years and 84.9 % at 20 years, which was in agreement with the finding by Robboy et al. published in 2009 [20]. After TAH/BSO and chemotherapy, our patient is currently disease-free for the past 1 year. But long-term follow-up with thyroglobulin levels is necessary due to reports of increasing recurrence rates.

Conclusions

Based on the rarity of these tumors and their lack of firm prognostic factors, treatment decisions should be made individually, based on pathologic and clinical parameters.

Consent

Written informed consent was obtained from the owner of the patient for the publication of this case presentation and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

Abbreviations

MSO: malignant struma ovarii; Tg: thyroglobulin; TTF-1: thyroid transcription factor-1; CK19: cytokeratin-19; HBME-1: human bone marrow endothelial cell-1; TP: paclitaxel/carboplatin; MRI: magnetic resonance imaging.

Competing interest

The authors declare that they have no competing interests.

Authors’ contributions

YZ and CW collected the clinical data and drafted the manuscript. YS and SQX helped to draft the manuscript. SJU carried out the pathological diagnosis and immunohistochemical staining. YZ and RH assessed the ultrasound imaging. GNZ conceived the study and reviewed the manuscript. YZ and CW contributed equally to this article. All authors read and approved the final manuscript.
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