Abstract

A total of 3 to 5% of all ovarian malignancies include malignant ovarian germ cell tumors (MOGCTs). They are subdivided into germinomatous and non-germinomatous tumors. Common types of non-germinomatous tumors include yolk sac and immature teratoma. Ovarian yolk sac tumors (YST) are the second most frequent histological subtype of MOGCTs, after ovarian dysgerminoma. They account for 20% of MOGCTs and are frequent especially in childhood and in early adulthood. We report the case of a yolk sac tumor of the ovary in a 13-year-old female.

Keywords: Alpha-fetoprotein, Ovarian germ cell tumors, Yolk sac tumor.

INTRODUCTION

Yolk sac tumors (YSTs) are defined as tumors that resemble the yolk sac, allantois, and extraembryonic mesenchyme and also known as endodermal sinus tumors. Yolk sac tumors are rare and a subtype of germ cell tumor. It accounts for about 10% of malignant germ cell tumors. This tumor presents as a rapidly growing mass in younger women.

CASE REPORT

A 13-year-old female came with chief complaints of lower abdominal pain and distension since 5 months. She had associated loss of weight and appetite. Per abdominal examination revealed a firm mass of about 15 × 16 cm in the lower abdomen, which was mobile freely horizontally; vertical mobility was restricted.

INVESTIGATIONS

- Complete hemogram: hemoglobin 9.2 g/dL; other parameters were normal; random blood sugar, liver function test, and renal function test were normal.
- Serum alpha-fetoprotein (AFP) was elevated (3,029 ng/mL).
- Other tumor markers were negative.
- Follicle-stimulating hormone and luteinizing hormone were normal.
- Magnetic resonance imaging: Well-encapsulated lesion measuring 16 × 15 cm, uterine size appropriate for age.

Impression: Dysgerminoma; karyotyping = normal female karyotype; histopathology report showed features of YST (Fig. 1). Biopsy of left ovary, biopsy of right ovary, and omental biopsy were negative for malignancy; cytology was negative for malignancy.

Treatment Given

Treatment provided was left ovarian cystectomy with left salpingectomy, biopsy of right ovary with omental biopsy.

Intraoperative Findings

The patient had a right ovarian well-encapsulated tumor, measuring 15 × 16 cm. Uterus was infantile; left ovary and tube were normal (Figs 2 to 4).

DISCUSSION

Yolk sacs are defined as tumors that resemble the yolk sac, allantois, and extraembryonic mesenchyme and also known as endodermal sinus tumors.

Yolk sac tumors are commonly seen in males and females, involving the testis, ovary, and other sites, such as...
Yolk sac tumors of the testis are observed in two forms, i.e., pure YST in young children and mixed type in adults.\textsuperscript{1,2} Yolk sac tumors are common in Asian children than in other races. In adults, these tumors are more common in white individuals than in other races.\textsuperscript{3,4}

The etiology of YST is unknown. Hypermethylation of the $RUNX3$ gene promoter and overexpression of GATA-4, a transcription factor that regulates differentiation and function of yolk sac endoderm, may play important roles in the pathogenesis of YSTs.\textsuperscript{3,4} But these hypotheses have not been proved.

Yolk sac tumors are solid gray-white with a gelatinous, mucoid appearance. Necrosis, cystic changes, and hemorrhage are often seen. Grossly, the overall appearance of a YST is heterogeneous.

Many other forms of these YSTs include endodermal sinus papillary, solid, glandular, festoon, myxomatous, sarcomatoid, macrocytic, polyvesicular, hepatoid, and parietal. These various forms do not predict prognosis, but may pose difficulties in distinguishing yolk sac YSTs from other tumors or other subtypes.\textsuperscript{5}

Yolk sac tumors of the endodermal sinus type consist of a central vessel rimmed by fibrous tissue, surrounded by malignant epithelial cells in a cystic space which is lined by flattened tumor cells. This endodermal sinus-like structure is called a Schiller-Duval body and is pathognomonic of YSTs. Schiller-Duval bodies are found only in 50 to 75\% of tumors.

Yolk sac tumor cells express AFP, which can be detected on tissue sections by immunohistochemistry as well as by biochemical estimation.\textsuperscript{6} Alpha-fetoprotein is not specific for YSTs.\textsuperscript{7} In 40 to 80\% of cases of YSTs, placental alkaline phosphatase is present, which is a marker of several different subtypes of germ cell tumors.

Other markers of YSTs have been reported recently. The prominent one is glypican 3. It is a membrane-bound heparan sulfate proteoglycan. Glypican 3 is more sensitive but less specific than AFP, as glypican 3 can be detected in some immature teratomas, in addition to liver and chorioepithelioma.\textsuperscript{8,9}

The missing of short arm of chromosome 1 (1p36) and the long arm of chromosome 6 (6q) as well as presence of the long arm of chromosomes 1 and 20 has been reported in YSTs. A few papers have demonstrated the molecular signature of these tumors based on expression from microarray studies.\textsuperscript{10}

About 80 to 90\% YSTs in children belong to stage I. The presence of a YST component in stage I tumors has been associated with decreased likelihood of occult metastasis.

Age is not a prognostic tool and does not predict and help in diagnosing YSTs. The amount of AFP elevation, in addition to other serum tumor markers in mixed germ cell tumors, has been shown to correlate with prognosis. For serum AFP, levels <1,000 ng/mL show a good prognosis. Levels >10,000 ng/mL show a poor prognosis. Tumors in between these levels are associated with an
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CONCLUSION

Differential diagnosis of yolk sac tumor includes embryonal carcinoma, seminoma, juvenile granulosa cell tumor, and dysgerminoma. Staging of the disease and increase in AFP along with other tumor markers can help in predicting the prognosis.

REFERENCES