Unusual Causes of Adnexal Masses in Early Reproductive Period

Erken Üreme Çağında Adneksial Kitlelerin Nadir Sebepleri

ÖZET
Üreme çağında tespit edilen ovaryan tümörler akıl kaçırtıcıdır. Bu yazida üç ayrı tanılı vaka sunuyoruz; Ovaryen immature teratomu (IT), ovari primer peritoneal seröz papiller kansiyomu (PSPC) ve metastatik ovaryan diffuz büyük B hücreli Lymphoması. Ovrene ait IT siklikla solid, artmış alfa fetoprotein değerleriyle birlikte görülüen nadir bir tümördür. PSPC, normal büyüklükte overler, yaygın peritoneal karsinomatosis ve belirgin artmış CA12-5 değerleriyle birlikte olan hastalarda tanıda düşünlmesi gerekir. Lymphoma, CT de çok sayıda büyümüt nodların füzyonu veya tek dönemde genislemiş lenf nodlarını içeren solid kitleler olarak karşımıza çıkar. Lenfomalar en hızlı doubling zamanına sahip olduğundan radyolojik incelenme bir kaç günden itibaren gösterebilir ki bu bir ipucu olabilir. Özellikle nodal kitleler ile birlikte ovaryan kistler ovaryan metastazı olan lenfomayı düşündürür.

Anahtar kelimeler: Ovaryum ve tuba uterina hastalıklar; üreme

ABSTRACT
Ovarian tumors when detected in the reproductive period are challenging. We present three different cases diagnosed as an immature teratoma (IT) of ovary, a primary peritoneal serous papillary carcinoma (PSPC) and a metastatic ovarian diffuse large B-cell Lymphoma in the early reproductive period. IT of the ovary is an uncommon tumor; predominantly solid with increased serum alpha-fetoprotein levels may be associated with a greater likelihood. PSPC should be suggested as a diagnosis in patients with extensive peritoneal carcinomatosis, relatively normal sized ovaries, and a highly elevated serum CA-125 level. Lymphomas on CT may appear solid, include singular enlarged lymph nodes and fusion of multiple enlarges nodes. As lymphomas have fastest doubling time the radiologic evaluation may progress in few days and may be a clue for the diagnosis. Especially, ovarian cysts with nodal masses suggest a lymphoma with ovarian metastasis.

Key words: Adnexal Disease; Reproduction
the right ovarian mass was resected. At the frozen section immature teratoma of the ovary was the first diagnosis. Since the tumor didn’t extend beyond the left ovary, and the patient wanted a fertility sparing surgery, unilateral salpingo-oopherectomy and full staging procedure were performed. The final pathologic diagnosis was Grade II astrocytoma and neuroblastoma arising in immature teratoma with negative lymph nodes, peritoneal washings and biopsies (stage Ia) (Figure 1). With the medical oncology consultation a three cycle of bleomycin, etoposide and cisplatin treatment was decided.

Case 2
A 27 years old woman (G1,P1) was admitted with a complaint of bilateral groin pain for a few months. Pelvic Computed Tomography (CT) showed a 39x30mm mass adjacent to the right ovary (Figure 2). The tumor markers were all in normal ranges. At the laparotomy, the uterus and both ovaries were normal. The mass fixed to the posterior isthmic part of the uterus in cul-de-sac was excised. Intraoperative pathology consultation reported mass as a tumor being at least borderline malignant serous papillary tumor. Then, full staging procedure was performed. The fact that the patient was a young woman of childbearing age, the operation was completed without any further invasive procedures.

Due to the final pathological diagnosis of primary peritoneal papillary serous carcinoma (PSPC) (Figure 3) with residual disease, second-look surgery was performed with a complete debulking surgery (total abdominal hysterectomy, bilateral salpingo-oopherectomy, bilateral pelvic lymph node dissection, infracolic omentectomy, appendectomy and low anterior resection of colon). The tumor was reported to infiltrate the serosal surfaces of the rectal and sigmoidal wall. The patient, with an uneventful recovery was transferred to medical oncology unit for chemotherapy treatment.

Case 3
A 26-year-old female patient (G1,P1) complaining of pelvic pain and distention with 8kg weight loss was admitted to our gynecology clinics. The ultrasonography and the abdominal MRI confirmed bilateral ovarian tumor with another solid 75x83mm mass just over the bifurcation of aorta just beneath the intestinal loops (Figure 4). At the laparotomy, biopsies from ovaries and the mesenteric mass resulted as lymphoma (Figure 5). The surgery was finished without any further process. The patient was transferred to medical oncology unit for chemotherapy treatment.

DISCUSSION
The challenging period for adnexal masses in reproductive period begins if the mass have malignancy criteria. US, is so far the first step to confirm the presence of a mass and to determine its organ of origin. Further radiological methods (CT, MRI) are useful for accurate and differential diagnosis. But the diagnosis of these suspicious pelvic masses ultimately requires an exploratory laparotomy (3).

IT of the ovary is an uncommon tumor; comprising less than 1% of teratomas of the ovary (4). Malignant transformation is uncommon and seen in approximately 2% of cases, usually in older women (most commonly squamous cell carcinoma 75% to 80%). Tumors of neuroepithelial origin are extremely rare in teratomas. We describe a case of ovarian IT with having both astrocytoma and neuroblastoma in the same tumor. IT may be solid or

![Figure 1.](image1)

- 1. Mature ectoderm (H&E X100)
- 2. Immature glial neuroplastic component (H&E X400)
- 3. Grade 2 glial (astrocytic) component (H&E X400)

![Figure 2.](image2)

39x30mm complex mass adjacent to the right ovary posterior to the uterus
cystic with solid components. These solid parts are usually recognized at CT and MR imaging. The existence of solid component can be sign of an immature teratoma, although mature teratomas sometimes have large solid parts. There is no complete diagnostic criterion for immature teratoma, although large, predominantly solid masses with increased serum alpha-fetoprotein levels may be associated with a greater likelihood (5,6).

PSPC have pathologic features closely resemble those of their ovarian counterparts (7). Some characteristics of CT of PSPC are reported as mesenteric or omental involvement, ascites, peritoneal thickening, extensive peritoneal calcification and normal-appearing ovaries (9-12). However some may show no sign of these criteria and have relatively large ovarian masses (8,9). The CA-125 antigen seems to be the most effective tumor marker for a primary peritoneal carcinoma (12). However, it should be noted that not all primary peritoneal carcinomas exhibit increasing levels of CA-125 (13). PSPC should be suggested as a diagnosis in patients with extensive peritoneal carcinomatosis, relatively normal sized ovaries, and a highly elevated serum CA-125 level.

Ovarian involvement by lymphoma may be the early manifestation of a systemic disease (14) or less commonly lymphoma may arise de novo in the ovary (15,16).

Lymphomas on CT may appear solitary, multiple-nodular or diffuse. Solitary mass includes singular enlarged lymph nodes and fusion of multiple enlarges nodes. The most frequently seen multiple-nodular type is diagnosed by enlarged lymph nodes with regional distribution. Both solitary and multiple-nodular types have clear margins. The CT features of diffuse type show diffuse but non-confluent enlarged lymph nodes in the mesenteric and retroperitoneal region with uniform density and cobblestone appearance. As lymphomas have fastest doubling time the radiologic evaluation may progress in few days and may be a clue for the diagnosis. Especially, ovarian cysts with nodal masses suggest a lymphoma with ovarian metastasis (17-21).

MRI may also be a helpful for differential diagnosis. Ferrozzi et al (22) reported that the MRI findings of metastatic ovarian lymphoma included solid bilateral masses, which were low signal intensity on T1 weighted images and mildly high signal intensity on T2 weighted images. Lesions were homogeneous with mild to moderate contrast enhancement with intravenous gadolinium.

Adnexal masses at reproductive age have great importance. Preoperative evaluation has a key role in diagnosis.

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**Figure 3.**
Papillary structures with serous epithelia. (H&E X100)

**Figure 4.**
Bilateral ovarian tumor with another solid 75x83mm mass just over the bifurcation of aorta

**Figure 5.**
1- Lymphoid infiltration, diffuse pattern (H&E X100)
2- Infiltration with centroblast and centrocytes (H&E X400)
3- CD20 staining of malign cells (X400)
4- High proliferation index with Ki-67 staining (X400)
differential diagnosis. All adnexal masses need further whole body physical examination. It should always be remembered that an ovarian tumor may be metastatic. Radiological evaluation especially CT and MRI may be helpful to differentiate the mass.

REFERENCES
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