Sex Cord Stromal Tumor with Annular Tubules: A Case Report

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Abstract:

Background: Sex cord tumor with annular tubules (SCTAT) is an uncommon ovarian tumor. One third of the cases are related with hyperestrogenism and Peutz-Jeghers syndrome (PJS). They are mostly seen in reproductive period and only rarely in pediatric patient group. Here, we present a 12 years old patient with SCTAT with a review of related literature.

Case Report: In a patient operated with a pre-diagnosis of acute appendicitis, a mass was found out in the right ovary. An appendectomy and right salpingo-oophorectomy were performed. In cross section of ovary, a nearly 6 × 5 cm regular bordered mass was observed which included millimetric white nodular foci and hemorrhagic changes (Figure 1).

Conclusion: This case is worth discussing since besides being an infrequent entity, SCTAT is also very rare in pediatric age.

Keywords: Ovarian tumors, Sex-cord tumors, Sex cord tumor with annular tubules

Introduction

Sex cord tumors with annular tubules (SCTAT) are first described by Scully [1]. They are rare tumors of the ovary which account for less than 1% of all sex cord stromal tumors [2]. One third of SCTAT are accompanied by Peutz-Jeghers syndrome (PJS) [3,4]. The cases with PJS are slightly younger (average age, 27) than isolated cases (average age, 36) [2]. The tumor is extremely rarely found in pediatric patients [3-7]. Clinical diagnosis of SCTAT is very difficult and diagnosis is mostly made with pathological examination [7].

Case Report

Twelve years old female patient was admitted to emergency clinic with a sudden severe pain localized at right lower abdomen, nausea and vomiting. At abdominal ultrasound, ascites and an appendix of 5.8 cm. diameter were observed at right lower abdomen. Clinical history and examination did not reveal any significant finding. The patient was operated with a prediagnosis of acute abdomen (acute appendicitis?). Right adnex was 8 × 3 cm, torsioned and gangrenous. An appendectomy with right salpingoo-oophorectomy were performed. Macroscopically, at the cross-sections of 7 × 5 × 3 cm, gray-purple ovarian tissue; a nearly 6 × 5 cm regular bordered mass was observed which included millimetric white nodular foci and hemorrhagic changes (Figure 1).

The surrounding ovarian tissue also included hemorrhagic areas. Sections of tuba and appendix did not reveal an abnormal feature. At histopathological examination; in fibrous stroma of the ovarian tissue, a tumoral area was observed which consisted of simple and complex tubular structures which had PAS positive hyaline material in the center surrounded by peripheral palisading cells (Figures 2 and 3).
The cells had small single nuclei and wide pale cytoplasm. One mitotic figure was noted per 10 high power fields (HPF) and microcalcification was not observed. SCTAT, sertoli cell tumor, gonadoblastoma and microfollicular granulosa cell tumor were considered for differential diagnosis. Tumor cells were positively stained for inhibin, calretinin, vimentin and were negative for EMA, PLAP, laminin and CD117 (Figures 4 and 5).

Discussion

Since SCTATs have common features with sertoli and granulosa cell tumors, they are classified as a subgroup of sex cord stromal tumors [2,3,8]. Due to hyperestrogenism, puberty precox in pediatric group, and menstrual irregularities as menometrorrhagia in reproductive age might be the clinical presentations of SCTAT. The tumor that is found incidentally in a case related with PJS may be accompanied with hyperpigmented macules in oral mucosa, gastrointestinal hamartomatous polyps and various tumors in genital system, breast and pancreas [2,3]. In preoperative examination and clinical history of our case, findings related with hyperestrogenism or PJS were not detected.

Sertoli cell tumor, gonadoblastoma and microfollicular granulosa cell tumor should be taken into account for differential diagnosis of SCTAT. Complex tubules are not seen in sertoli cell tumors. In microfollicular granulosa cell tumors, hyaline bodies in lumens of tubular structures are not very dominant, a feature important for differential diagnosis. In gonadoblastomas there are sex cord stromal elements and hyaline bodies, however a germ cell component and abnormal gonadal development (most often phenotypic females who are virilized) are seen contrary to SCTAT. In our case, in addition to histopathological findings, PLAP and CD 117 were negative and germ cells were not present [9].

Clinical behaviour and pathological features of the tumors differ according to the presence or absence of PJS. In cases with PJS, bilateral, multiple and smaller (<3 cm) tumors with benign behaviour are observed. On the contrary, in cases without PJS, unilateral, bigger (as big as 33 cm) pelvic masses that may behave in a malignant behaviour, are detected [9]. On 20% of the cases without PJS, since malignancy and distant metastasis might be observed, these tumors are known to have low malignant potential. They may spread through pelvic, paraaortice and supraclavicular lymph nodes and metastasize to retroperitoneum, parietal and visceral peritoneum, liver, kidney and lung [9,10].

Malignant potential of the tumor may not be recognized microscopically [10]. However, in literature, there are studies was made which revealed that she had no complaint during postop six years.
reporting that more than 3-4 mitosis in 10 HPF and infiltrative growth pattern are related with aggressive behaviour [9]. In the case presented here, there wasn’t any finding in favour of malignancy. In postoperative period, the patient did not come for routine follow up, but a phone contact could be made. Accordingly, it was apparent that no complaint was present in a period of postop 6 years.

In literature search there are found reports including mostly a single case and a very few large study groups. So, currently, there are no widely accepted standard treatment protocols. In unilateral tumors, ipsilateral pelvic and paraaortic lymphadenectomy and unilateral salpingo-oophorectomy are the recommended therapy for young patients [9]. In cases with PJS, even if the tumor is present in only one ovary, wedge resection is recommended for the other ovary. Even though the effect is not clear, chemotherapy and radiotherapy are performed for local recurrence and distant metastasis [7].

Long term follow up is mandatory for SCTATs which have relatively good prognosis despite high recurrence risk [7].

Conclusion

The case presented here which is found incidentally in absence of a PJS is worth attention especially because of its pediatric age.

References


