Case Report

Mitotically active fibrothecoma in association with sclerous peritonitis

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Abstract

Background: Mitotically active fibrothecoma associated with sclerosing peritonitis is an enigmatic and rare condition, described by Clement et al. in 1994, with fewer than 50 pathologically characterized cases. Association of these two lesions is extremely rare and the etiology currently remains unclear; therefore, association with anti-epileptic drugs was suggested.

Case presentation: We report the case of young female, with history of epilepsy, presented with acute severe abdominal pain. Ultrasound revealed a 20 cm mass, possibly arised from ovary, which was associated with septate ascites. The microscopic examination of the mass showed a morphology fibrothecoma with areas of oedema and high mitotique index but without severe atypia.

Conclusion: Mitotically active fibrothecoma associated to sclerosing peritonitis is relatively rare condition. Although its disturbing presentation, the behavior of this rare condition is benign; therefore this entity should be well known by surgeon and pathologist to avoid misdiagnosis and overtreatment of patient.

Abbreviations

hCG: human Chorionic Gonadotropin; CEA: Carcino-Embryonic Antigen; CA-125: Cancer Antigen 125; CD56: Cluster de Différentiation 56; CK AE1/ AE3: Pan cytokeratin AE1/ AE3; CD34: Cluster de Différentiation 34; Ki-67: Proliferating Antigen Ki67

Introduction

Mitotically active fibrothecoma is recently described as a category of ovarian fibromatous tumors, defined by mitotic figures more than 4 per 10 high power fields, without moderate to severe atypia. Association with sclerosing peritonitis is an enigmatic and rare condition, first described in 1994 by Clement et al. with less than 50 pathologically characterized cases [1-3].

Case report

18-year-old woman with history of epilepsy, presented with severe acute abdominal pain. On physical examination, the abdomen was tender with a palpable mass in the left iliac fossea. Ultrasound revealed a 20-cm heterogeneous left pelvic mass as well as free peritoneal fluid. The right ovary and the uterus appeared normal.

Serum analysis revealed that the Human Chorionic Gonadotropin (hCG) test was negative and serum levels of Carcinoeembryonic Antigen (CEA) and Cancer Antigen 125 (CA-125) were normal. The patient underwent laparotomy which showed large left ovarian tumor, associated with sclerosing ascites.

The right ovary, fallopian tubes, and uterus were...
unremarkable. Left salpingo-oophorectomy and intraoperative biopsy of the right ovary were performed.

On gross examination, the outer surface of the left ovary appeared lobulated, measuring 16 cm across its largest diameter (Figure 1). The yellowish sectioned surfaces of the ovarian mass were solid with edematous areas.

Microscopically, the tumor showed variable cellularity (Figure 2) with hypercellular zones alternating with hypocellular foci (Figure 3), it was comprised of fascicles and sheets of plump monomorphous spindle and oval cells, with no atypia. Patchy area of edema was noted. Within the spindle cell population, lutein–like cells were scattered singly or in clusters. Mitotic counts of spindle cells revealed 15 mitotic figures per 10 high–power fields (Figure 4). The right ovarian biopsy was histologically normal.

At immunohistochemistry, tumors cells expressed focally calretin (Figure 5) and inhibin (Figure 6). The proliferated antigen Ki67 was estimated at 20% (Figure 7).

Thus, diagnosis of unilateral mitotically active fibrothecoma associated to sclerosing ascitis was made.

The patient was discharged 7 days later. The clinical outcome was uneven, within 8 months of follow up.
Discussion

Mitotically active fibrothecoma is a recently described category of ovarian stromal tumors, representing less than 10% of fibrothecoma. This entity is defined by mitotic figures more than 4 per 10 high power fields, without moderate or severe atypia [4,5].

The median age of patient is 41 year and most frequent symptoms are abdominal mass, acute pain, ascites as in our case with dysfunctional uterine bleeding [6].

Biologically, two cases are reported in the literature with a high level of CA125 [6].

In gross, the tumor appears as a large ovarian mass, has well-defined capsule and the cut-surface showed typical yellow bosselated appearance.

The average size has been reported to be 9.4 cm [6]. Microscopically, the tumor is hypercellular, usually with area of edema, occasionally imparting a microcystic appearance. Most of the tumor cells are spindle but a minority was round of edema, occasionally imparting a microcystic appearance.

The median mitosis was found to be 6, with a maximum of 19 mitosis under 10 high power fields. Historically, since 1981, as described by Pratt and Scully, the 4 mitotic activities of the cells was given equal importance as nuclear atypia in predicting the aggressiveness of an ovarian fibromatous tumour and classifying as fibrosarcoma. The tumour which exhibit a mitotic activity of ≥ 3/10 HPF is labelled fibromas, while those with ≥ 4/10 HPF mitotic figure were considered malignant in nature [4,7-9].

As time elapsed and more cases were reported, tumours that had high mitotic counts of ≥ 4/10HPF but minimal nuclear atypia were found to be less aggressive and followed a more indolent clinical course. Hence in 2006, Irving et al. proposed the reclassification of fibromatous tumours, introducing mitotically active fibrothecoma for the first time, which fitted into the afore-mentioned criteria. This has been included in the WHO histological classification since 2014 [4,7-9].

Immunohistochemical studies have shown that the tumor cells to be reactive with calretinin, Cluster of differentiation Ki-67 proliferating Antigen (IHC x 400).

Figure 7: Ki-67 proliferating Antigen (IHC x 400).

In rare cases, as the case of our patient, fibrothecoma may be associated with sclerosing peritonitis. The etiology and pathogenesis of this association is not clear; but a relation with anti-epileptic drug has been discussed. Our case consolidate this hypothesis, since our patient, as the case reported by Pati et al, was under anti epilepsy drug [3,10].

On the other hand, some author proposed fibrosing soluble cytokines secreted by tumor cells to be responsible for the peritoneal manifestations [3].

Microscopically, the process of peritonitis consists of fibroblastic and myofibroblastic cell proliferation separated by collagen and fibrin, mesothelial cell proliferation, and occasionally mononuclear inflammatory cells [3].

Currently, the concept is that the tumor is benign in behavior although complications due to sclerosing peritonitis bring down the quality of life [3].

No standard treatment for mitotically fibrothecoma currently exists, although overtreatment should be avoided in women requiring preservation of fecundity. Zong et al. suggested that other risk factors, such as tumor size, growth rate and the evaluation of Ki–67, should be considered [6,11].

Conclusion

Mitotically active fibrothecoma associated with sclerosing peritonitis is relatively rare condition. Although its disturbing presentation with a large tumor size, higher mitotic index and peritoneal involvement, the behavior of this rare condition is benign; therefore this entity should be well known by surgeon and pathologist to avoid misdiagnosis and overtreatment of patient. Standard guideline of treatment is yet to be formulated for this entity.

The authors declare that there are no conflicts of interest associated with this manuscript.

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