Case report of sertoli-leydig cell tumor of bilateral ovaries in a woman with 46XYkaryotype

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Abstract
Sertoli-Leydig Cell Tumors are rare constituting less than 0.5% of ovarian neoplasms and usually unilateral. Average age of presentation is 25 years. We report the case of a 23-year-old unmarried woman who presented with primary amenorrhea and 8 months history of intermittent pain over left lower abdomen. Transvaginal scan revealed rudimentary uterus with left ovarian cyst with mural nodules. Cytogenetic study revealed 46XY karyotype. Histopathological examination of specimen received after surgery showed well-differentiated Sertoli-Leydig Cell Tumor in sections from bilateral ovaries and Serous Surface Papilloma in a section from left ovary. Diagnosis is based on morphological features. Prognosis depends on stage and degree of differentiation.

Keywords: Primary amenorrhea, Sertoli, Leydig, 46XY

Introduction
Sertoli-Leydig Cell Tumor (SLCT) is a rare ovarian tumor belonging to the group of sex-cord stromal tumors. These constitute less than 0.5% of ovarian tumors. The average age of presentation is 25 years. Most tumors are unilateral [1]. The characteristic features of these tumors are the presence of testicular structures that produce androgens that lead to symptoms of virilization depending on the quantity of androgen production. These tumors are also characterized by the degree of differentiation of structures in them, the presence of which determines whether the tumors are benign or malignant [2]. We here present a case report of Sertoli-Leydig Cell Tumor having the rarity of bilateral presentation and male karyotype in a 23-year-old phenotypically female patient.

Case Report
A 23-year-old unmarried woman presented to the Gynecology OPD with primary amenorrhea and a history of intermittent pain over left lower abdomen for 8 months. Physical examination revealed hirsutism and normal stature. Cytogenetic study revealed 46 XY karyotype (male genome). Transvaginal scan showed rudimentary uterus and left ovarian cyst with mural nodules. She was diagnosed as a case of primary amenorrhea with gonadal dysgenesis in the Gynecology Department and she thereafter underwent bilateral gonadectomy with left sided cystectomy. On gross examination of the specimen sent for histopathological examination, the right ovary showed a smooth grayish-white solid mass of size (2X1)cm and left ovary showed a smooth grayish-white solid-cystic mass of size (5X4)cm. Microscopically, section from both the ovaries showed solid tubules of dark blue sertoli cells and the stroma contained leydig cells with pale, vacuolated to eosinophilic cytoplasm.
This indicated a well-differentiated form of SLCT. Serous Surface Papilloma was seen in a section from left ovary. Also, the finding of 46XY karyotype in phenotypically female patient appears to be consistent with the pure gonadal dysgenesis syndrome known as Swyer syndrome.

Sertoli-Leydig Cell Tumor Left Ovary (Left image- 10x; H&E) (Right image- 40x; H&E)

Sertoli-Leydig Cell Tumor Right Ovary (Left image- 10x; H&E) (Right image- 40x; H&E)

Serous Surface Papilloma Left Ovary (Left image- 10x; H&E) (Right image- 40x; H&E)

Discussion

Sertoli-Leydig Cell Tumor (SLCT) is a rare ovarian tumor belonging to the group of sex-cord stromal tumors. These constitute less than 0.5% of ovarian tumors and are usually unilateral[1]. Our case is thus rare in having a bilateral presentation of the same. According to WHO classification, there are five subtypes of Sertoli-Leydig Cell Tumors according to the neoplastic Sertoli and Leydig cells exhibiting varying degrees of differentiation, which include well-differentiated, moderately differentiated, poorly differentiated, retiform and with heterologous elements. The stage and degree of differentiation are the most
important prognostic factors in these tumors. In 1985, Young and Scully reviewed 207 cases; of which all well-differentiated tumors were benign, 11% of tumors with intermediate differentiation, 59% of tumors with poor differentiation, and 19% of those with heterogeneous elements were malignant [1]. Clinically, they present with signs related to androgen production like hirsutism and rarely estrogen [3], or symptoms of mass-occupying lesion as pelvic-abdominal mass and/or pain [1, 4, 5]. Testosterone and androstenedione levels are elevated in approximately 80% of patients with ovarian Sertoli-Leydig Cell Tumors with virilizing manifestations [6, 7].

Our case also has the unusual finding of a 46XY karyotype with the patient being phenotypically female, a finding consistent with Swyer syndrome. Swyer syndrome is a pure gonadal dysgenesis syndrome with 46 XY karyotype, primary amenorrhea, presence of female internal genital tract and bilateral streak gonads in a phenotypic female [8]. There is a testicular differentiation abnormality in Swyer syndrome [9]. They have also elevated gonadotropins and hypoplastic gonads without germ cells [10]. The diagnosis is usually made at adolescence when the primary amenorrhea is investigated [8]. Gonadoblastoma is the most common tumor that develops in Swyer syndrome [6, 7]. We could not comment on the streak gonads in our case due to the presence of mass in both the gonads.

Sertoli-Leydig Cell Tumors are mostly unilateral and diagnosed in stage I, so young patients are treated with conservative surgery [11]. For patients with poor prognostic factors, adjuvant chemotherapy is considered. The BEP regimen i.e., bleomycin, etoposide, and cisplatin regimen does not affect the fertility status of the patient and is thus considered safe. Sertoli-Leydig Cell Tumor should always be taken into consideration in a young female patient with symptoms of virilization and an ovarian mass. The histopathology of the tumor decides the line of management; poorly differentiated tumors require aggressive management because of the high chances of them being malignant and intermediatedifferentiated tumors need an individualized approach [12].

Conclusion
This is a rare case of bilateral Sertoli-Leydig Cell Tumor. Primary amenorrhea with pain abdomen led to further investigations, which revealed left ovarian cyst with mural nodules and rudimentary uterus on transvaginal ultrasound and 46XY karyotype by cytogenetic study. Microscopic examination of sections from the specimen obtained by bilateral gonadectomy with left sided cystectomy revealed well-differentiated Sertoli-Leydig Cell Tumor in bilateral gonads. Presence of 46XY karyotype, primary amenorrhea, normal stature and rudimentary uterus were consistent with Swyer syndrome but we could not comment on the presence of streak gonads due to the presence of mass in bilateral gonads.

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