SERTOLI-LEYDIG CELL TUMOR: A RARE OVARIAN TUMOR
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ABSTRACT: Sertoli- Leydig cell tumor or “Androblastoma” or “Arrhenoblastoma” is a rare virilizing tumor of ovary, mostly with masculinizing features. Many but certainly not all tumors are hormonally active. Although classified under malignant tumors the degree of malignancy is less than that of ovarian carcinoma in general. Unilateral oophorectomy was done and the patient recovered well with diminution of masculinizing features.

KEYWORDS: Clitoromegaly, Hirsutism, Sertoli- Leydig cell tumor, Testosterone.

INTRODUCTION: Sertoli-Leydig cell tumor (SLCT) is a rare ovarian tumor that belongs to the group of sex-cord stromal tumors. These constitute less than 0.5% of ovarian tumors. Most tumors are unilateral, seen during the second and third decades of life and are of less well differentiated variety. Many patients have symptoms of virilization due to the presence of androgen element in tumor. The degree of differentiation is age related.¹ Intermediate differentiated tumors comprises of 54% of tumors (data reference 2).

CASE: A 20 year old nulliparous unmarried woman came to Government general hospital with complaints of progressive oligomenorrhoea of one and half years duration. She had been amenorrhic for the past one and half year. She had noticed excessive growth of hair on face, chest and limbs and hoarseness of voice for one year, recession of breast (and increased sweating and fatigue for one year). Headache for 10 years which is occipital in nature with nausea and reeling sensation and relieved on taking some analgesic. There is no history of anorexia, weight loss/gain. Her medical and family history is uneventful.

Her general physical examination was normal except for the presence of hirsutism FERRIMAN-GALLWAY score of 13, and clitoromegaly. Per rectal examination revealed normal
findings. An ultrasound examination of the pelvis showed 5x4.2cms solid mass in left adnexa, and MRI abdomen showed 4.3x3.6cms solid space occupying lesion in left ovary. Hormone profile in blood indicated excessive androgenic activity in the form of elevated testosterone level. (513ng/dl; normal 20-80ng/dl), serum and rostenidione (7.66ng/ml; normal 3-3.5ng/ml), DHEA (81.7ng/ml; normal 1.8-12.5ng/ml), however DHEAS levels were normal. MRI brain, visual fields, and fundus examination were normal.

Based on these investigations a diagnosis of virilizing (androgen producing) tumor of ovary was made. The patient underwent exploratory laparotomy and left salpingo-oophorectomy. The size of the left ovary is 7x5cms and on cut section showed a well encapsulated 5x3cms mass lesion. Rest of the abdomen appeared normal. Biopsy revealed Sertoli-Leydig cell tumor of intermediate grade differentiation. Biopsy from right ovary and omentum were normal.

Patient was kept on cyclical oral contraceptive pills and she resumed regular cycles. Post-operatively hirsutism, clitoromegaly, decreased. Hoarseness decreased over a course of 6months. Patient is under regular follow-up. She got married and delivered a healthy male baby after two years of surgery. Need for radiotherapy excluded by radiotherapist.
COMMENT: Menstrual irregularities are the most common hormone related manifestation. If virilizing signs are present menstrual irregularities are universal. Androgen excess result in defeminization. Breast atrophy, loss of sub cutaneous deposits and female contour are common.

In Women with androgen secreting sertoli-leydig cell tumors growth of facial and body hair, development of acne and external genitalia occur in about 33-38% of cases. Deepening of voice is noted in 75% of virilized patients. Zaloudek and Norris have the largest series in which the incidence of virilizing signs is reported.

Clitoral hypertrophy, defined as clitoral length more than 2cm is seen in approximately half of virilized patients. After tumor resection many androgenic manifestations resolve over time. Virilizing symptoms of clitoromegaly, hirsutism, and deepening of voice regress less reliably. Abdominal pain, discomfort and swelling noted in about 15%.

Ultrasound remains the best imaging modality, but normal ultrasound findings in presence of clinical and laboratory evidence of androgen excess cannot exclude the diagnosis. Computerized tomography is helpful in differentiating it from adrenal tumors. Study of tumor markers such as inhibin and AFP are useful to study the response to therapy in patients with advanced or recurrent SL tumors.

Hormonal studies are useful to differentiate between ovarian and adrenal tumors. Patients who demonstrate signs of virilization will have elevated plasma testosterone. Urinary 17-alpha ketosteroids are usually normal or slightly elevated in contrast to adrenal tumors.

Ovarian and adrenal venography with selective venous sampling is considered as “gold standard” for localizing androgen secreting tumors. 97% of tumors are diagnosed at stage 1 due to early androgenic manifestations. Well differentiated tumors are benign. Moderately and poorly differentiated tumors have malignant potential ranging from 3-34%. SLCT having retiform pattern are associated with 20% malignancy rate.

Young woman desirous of childbearing should be treated with unilateral salpingo-oophorectomy. In older woman TAH with bilateral salpingo-oophorectomy is the treatment of choice.

CONCLUSION: Whenever a young girl presents with menstrual disturbances and hirsutism with elevated androstenedione levels, Sertoli-Leydig cell tumor should be suspected and unilateral salpingo-oophorectomy often gives good results.

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