A Case of Granulosa Cell Tumour with Atypical Presentation

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Summary:
Introduction: Granulosa cell tumour is one of the rare variations of ovarian tumour. As the Granulosa cells secrete Estradiol, the patient with granulosa cell tumour usually present with features of precocious puberty. But our patient present with androgenic features, which inspires us to report this atypical case.

Materials and methods: This young patient admitted in BBMH, as a diagnosed case of ovarian tumour with features of virilisation. Evaluation of patient with history taking, clinical examination and investigations are done, and information is noted accordingly.

Conclusion: Features of virilisation are a nightmare for a lady of 16 years old. So during management of this patient proper counselling is very crucial and during surgery ovarian reserve should be maintained as far as possible considering her future quality of life and obstetric outcome.

Keywords: Granulosa cell tumour, virilization.

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Introduction:
Granulosa Cell tumour [GCT] is one of the rare varieties of sex cord stromal tumours. It contributes 5% of ovarian tumors1. Mean age of presentation is 13 years. Most patients present with sexual precocity due to excessive estrogen production and rarely produce Androgen2. Our patient presented with these rare features of tumour. Granulosa cell tumour was first described by Scully in 19773. In contrast to adult type juvenile one is more aggressive with high mitotic index.

Case Report:
Miss X, a 16-year-old unmarried girl of middle class family got admitted with the complaints of secondary amenorrhoea and abnormal facial hair growth for last 1 year. According to the patient, she had history of spontaneous menarche at the age of 12 years and of regular cycle. She developed amenorrhoea for last one year and feeling heaviness in lower abdomen for last 6 months. On general examination, she had average body built, height 5 feet 3 inches, weight 52 kgs, hirsutism with male pattern and excessive course hairs on upper arms and lower limbs. On mons pubis the hair distribution is of normal female pattern. Other features of virilisation were absent except mild regression of breast. On abdominal examination mild tenderness was present over lower abdomen. Vaginal inspection revealed clitoromegaly [2.5cm]. On digital rectal examination, there was a lump 8x8cm size, mobile, non-tender, firm on posterior fornix. Ultrasono scan revealed a solido -cystic mass 10x7 cm at left adnexal area with small pelvic collection. Enocrinological studies of serum FSH, LH, ACTH were normal except serum Testosterone level, which was raised [15.6nmol/l]. On laparotomy, small amount of peritoneal fluid was found, aspirated and sent for cytology. A mass 10x8cm, party cystic partly solid, well capsulated was identified in left adnexal area. Opposite ovary and tube were healthy. There were no peritoneal seedlings or palpable pelvic or para aortic lymph nodes. Left sided salpingo-ophorectomy was done. Histopathogy revealed Granulosa cell tumour. Cytology of peritoneal fluid failed to identify any malignant cell. For abnormal hair growth she was advised dermatological consultation. Following surgery from next cycle, she resumed menstruation. Serum testosterone level regressed to normal within two months. After one year follow up she is having regular menstruations and has not developed any unwanted symptom.

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Discussion:
Though ovarian tumours present late stage, hormonally active ovarian neoplasm like GCT are diagnosed early due to the effect of hormones on target tissues. This is a rare ovarian tumour with two distinct clinicopathological sub-type like Adult and Juvenile. Adult variety is common accounts to 95%.[4]

Generally the adult variety occurs in peri [40-45 years] and post menopausal women with peak incidence at 50-55 years. The Juvenile one occurs in pre-pubertal age group. GCT as a hormone producing tumour [Estrogen] commonly present with sexual precocity. Other presenting features are – pain abdomen, abdominal distension, menstrual abnormalities like- menorrhagia, inter-menstrual bleeding, post-menopausal bleeding. Endocrine manifestation are related with hyper-secretion of Estrogen from granulosa cells. 15% of the tumours are hormonally inert.[7]

One of the rare presentation of GCT is amenorrhoea with features of virilization. The luteinizing form produces androgen leading to virilisation. To date less than 50 cases of virilizing GCT have been reported in literature.[8] Our patient presented with this feature, the unique point which drew our attention to report the case.

Sunil kumarkota et al reported one case in which a 16-year-old girl presented with primary amenorrhoea and virilization over 3 years period.[9] Another case report by Arunnayak et al in which a 19 years old unmarried girl presented with secondary amenorrhoea and virilizing features like our patient.[10] She was stage 1A and unilateral salpingo-oophorectomy was done. Another patient of 17 years, was reported by Reddy et al in the year 2014, presented with secondary amenorrhoea with pelvic mass.[11] During reproductive age the typical picture of Androgen secretion is oligo-menorrhoea followed by amenorrhoea, defeminisation and progressive masculinization.

On physical examination, mass in lower abdomen could be palpable, usually unilateral upto 12 cm. On cut section the tumour is multi loculated, cystic with yellow white solid area. On histo-pathological examination, there are five histological patterns like – micro, macro-follicular, insular, trabecular and spindle/sarcomatoid. Call Exner

Fig.-1: a. Excessive facial hair growth, b. Clitoromegaly, c. Gross anatomy of tumour, d. Histopathological picture of tumour
Endometrial carcinoma in 5-13% of cases is seen in GCT. Diffuse or multifollicular pattern with microcytes containing eosinophilic secretion and coffee bean nuclei are the commonest diagnostic point. In our case the gross and histology revealed these described features.

The serum tumour markers raised in GCT are Estradiol, Inhibin, Anti-mullerian hormone and CA-125 in patients with sex-cord stromal tumour, pelvic and paraaortic lymphadenectomy may not be included in the staging surgery of the patient. Ultimate diagnosis is done by histopathological examination. Due to Estrogen stimulation, there is chance of development of endometrial histopathology must be included in the management protocol. The prognostic factors in GCT includes – staging, intra-peritoneal disease, tumour size, patient age, histologic grade of differentiation, mitotic activity and nuclear atypia. Survival rate after 10 years for stage 1,2,3 and 4 are 87%, 75%, 20% and 0% respectively.

The role of adjuvant therapy is controversial. But it is recommended in advanced stage disease. Limited data are available regarding the chemotherapeutic agents in stage 3 and 4 disease. BEP therapy have been tried successfully, Radiotherapy is reserved for recurrent disease in selected group of patients. As these tumours are known for relapse, long term follow-up is necessary for early detection.

Conclusion:
Our patient presented with some unusual characteristics of GCT. She did not present with precautious puberty, rather presented with secondary amenorrhoea. Abnormal hormonal secretion consisted mainly of Androgen, with sign of virilisation, which is a nightmare for any young adult female. But medical science is magic at some point; it removes all her agony by a simple incision and changes her life.

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