INCIDENCE OF VIRILISATION IN SEX CORD STROMAL TUMOURS OF OVARY, A 5-YEAR EXPERIENCE IN A TERTIARY CARE GYNAECOLOGICAL CENTRE

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ABSTRACT

BACKGROUND
Androgen secreting ovarian tumours represents less than 5% of all ovarian neoplasms. Hyperandrogenism results in development of acne, hirsutism, androgenic alopecia, menstrual irregularities & virilisation or masculinisation. The objective of this study is to look for incidence of virilisation in sex cord stromal tumours and discussion regarding various causes of hyperandrogenism.

MATERIALS AND METHODS
A retrospective descriptive study of virilising tumours of ovary was conducted in a tertiary care gynaecological centre at Institute of obstetrics and Gynaecology, Chennai for a period of 5 years from February 2012 to February 2017. 4 cases of sex cord stromal tumours presented with signs of virilisation.

RESULTS
A total of 592 ovarian malignancies were identified over a period of 5 yrs. out of which 36 cases were diagnosed as sex cord stromal tumours (6.08%). The most common age group affected are between 41 to 50 yrs. with median age of 45 yrs. Right sided ovary (58%) is more commonly involved than left sided (42%) ovary. Most of the tumours were solid in appearance with average size ranging between 6 cm – 10 cm. The most common symptom is abdominal mass (64%) with 4 cases presented with history of virilisation (11%).

CONCLUSION
Based on present study it may be concluded that patients with symptoms of virilisation require detailed clinical history, physical examination for signs, laboratory investigations & imaging to exclude causes other than ovarian neoplasm.

KEYWORDS
Hyperandrogenism, Ovarian Tumours, Virilisation, Hirsutism.


BACKGROUND
Hyperandrogenism is a common endocrine disorder occurring in women of reproductive age group. Androgen secreting ovarian tumours represents less than 5% of all ovarian neoplasms¹. Davis et al in the year 1900 described first case of ovarian tumour with virilisation,² & over century the number of cases reported in literature has been increased. Ovarian neoplasms like sex cord stromal tumours are associated with excess production of androgens. In other tumours of ovary like epithelial, germ cells or metastatic, the neoplastic nonsteroidogenic cells induce abnormal sex steroid response in ovarian stromal cells which leads to the development of clinical manifestations of virilisation.³ The objective of this study is to look for incidence of virilisation in sex cord stromal tumours and discussion regarding various causes of hyperandrogenism.

MATERIALS & METHODS
A retrospective descriptive study of virilising tumours of ovary was conducted in a tertiary care gynaecological centre in Chennai for a period of 5 years from February 2012 to February 2017. All the relevant data were collected from our record sections & histological slides were reviewed to confirm the diagnosis. During the study period, 36 cases of sex cord stromal tumours were identified out of which 4 cases presented with signs of virilisation.

RESULTS
A total of 592 ovarian malignancies were identified, out of which 36 cases were diagnosed as sex cord stromal tumours with incidence being 6.08%. The most common age group affected are between 41 to 50 yrs. with median age of 45 yrs. (Figure 1). The right sided ovary (58%) was most commonly involved than left sided ovaries (42%) with almost all cases are unilateral. Most of the tumours are solid (58%) in appearance followed by solid & cystic (30%) with only 2 cases cystic (6%) in appearance. The average sizes of the tumours are ranging from 6 cm-10 cm. The most common presenting symptom is abdominal mass (64%) followed by post-menopausal bleeding (11%), menorrhagia (11%), hirsutism (11%) with abdominal mass in 4 cases and 2 cases are incidental finding (3 %) in hysterectomy specimen (Figure 2).

Adult Granulosa cell tumour was the most common sex cord stromal tumour with total of 11(31%) cases followed by...
the coma 8 (22%) cases. The other cases reported were 4 cases of fibroma (10%), 3 cases of massive stromal oedema & granulosa cell tumour with the coma (8%), 2 cases of cystic variant of Adult granulose cell tumour (6%), & 1 case each of fibrothecoma (3%), Juvenile granulosa cell tumour (3%), Sertoli Leydig cell tumour (3%), steroid cell tumour (3%) and Sclerosing stromal cell tumour(3%) (Figure 3).

The details of presenting symptoms, surgical procedure done, histopathological diagnosis & follow up details of all 4 cases of hirsutism with abdominal mass are presented in table 1.

Table 1. Case series of virilisation

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Age Yrs.</th>
<th>Clinical History</th>
<th>Procedure Done</th>
<th>Hormonal Status</th>
<th>HPE</th>
<th>Follow Up (2yrs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>32</td>
<td>Amenorrhea with hirsutism</td>
<td>Staging laparotomy - Right ovary</td>
<td>Total testosterone 4.120 ng/dl, and was persistently elevated in male pattern in dexamethasone suppression test</td>
<td>Sertoli Leydig cell tumour</td>
<td>Asymptomatic during 2 yrs. follow up period</td>
</tr>
<tr>
<td>2</td>
<td>36</td>
<td>Amenorrhea with hirsutism</td>
<td>Left cystectomy</td>
<td>Total testosterone – 5.68 ng/dl</td>
<td>Adult Granulosa cell tumour - cystic variant</td>
<td>Patient died due to decompensated cardiac disease</td>
</tr>
<tr>
<td>3</td>
<td>19</td>
<td>Amenorrhea with hirsutism</td>
<td>Left oophorectomy</td>
<td>Total testosterone – 10.5 ng/dl</td>
<td>Juvenile granulosa cell tumour</td>
<td>At 13 months of follow up patient developed recurrence &amp; died due to disseminated disease.</td>
</tr>
<tr>
<td>4</td>
<td>42</td>
<td>Abdominal pain with hirsutism</td>
<td>Staging laparotomy - Right ovary</td>
<td>Not done as patient was taken up for emergency surgery</td>
<td>Steroid cell tumour</td>
<td>Asymptomatic during 2 yrs. follow up period</td>
</tr>
</tbody>
</table>

Table 2. Common Cause & Rare Lesion

<table>
<thead>
<tr>
<th>Common Cause</th>
<th>Rare Lesion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polycystic ovarian syndrome</td>
<td>Severe insulin resistance syndrome</td>
</tr>
<tr>
<td>Idiopathic hirsutism</td>
<td>Granulosa cell tumour</td>
</tr>
<tr>
<td>Sertoli Leydig cell tumour</td>
<td>Sclerosing stromal cell tumour</td>
</tr>
<tr>
<td>Steroid cell tumours - NOS</td>
<td>Thecoma</td>
</tr>
<tr>
<td>Sclerosing stromal cell tumour</td>
<td></td>
</tr>
<tr>
<td>Leydig cell tumour</td>
<td></td>
</tr>
<tr>
<td>Stromal luteoma</td>
<td></td>
</tr>
<tr>
<td>Gynandroblastoma</td>
<td></td>
</tr>
<tr>
<td>Gonadoblastoma</td>
<td></td>
</tr>
<tr>
<td>Androgen secreting adrenal tumours</td>
<td></td>
</tr>
<tr>
<td>Adrenal gland hyperplasia</td>
<td></td>
</tr>
<tr>
<td>Adenoma &amp; carcinoma</td>
<td></td>
</tr>
</tbody>
</table>
Tumour like conditions:
- Stromal hyperplasia
- Stromal hyperthecosis
- Pregnancy luteomas
- Hyperreactio luteinalis

Rare functioning ovarian tumours:
- Benign cystic teratoma
- Endometrial tumours
- Serous & mucinous cystadenomas
- Brenner tumours
- Leiomyoma
- Metastatic tumours

Cushing’s syndrome
Exogenous drugs.
Congenital adrenal hyperplasia.

Table 2. Causes of hyperandrogenism

<table>
<thead>
<tr>
<th>Causes</th>
<th>Investigations</th>
<th>Additional tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ovarian neoplasms</td>
<td>Increase in testosterone</td>
<td>Imaging – USG, MRI</td>
</tr>
<tr>
<td></td>
<td>Normal/ mild increase in DHEAS</td>
<td></td>
</tr>
<tr>
<td>Adrenal neoplasms</td>
<td>Increase in testosterone</td>
<td>Imaging – USG, MRI</td>
</tr>
<tr>
<td></td>
<td>Increase in DHEAS</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Increase in urine 17 ketosteroids</td>
<td></td>
</tr>
<tr>
<td>Congenital adrenal hyperplasia</td>
<td>Increase in testosterone</td>
<td>ACTH stimulation test is necessary to make diagnosis</td>
</tr>
<tr>
<td></td>
<td>Increase in 17 hydroxyprogesterone</td>
<td></td>
</tr>
<tr>
<td>PCOS</td>
<td>Increase in LH, Prolactin</td>
<td></td>
</tr>
<tr>
<td>Drugs</td>
<td>Normal levels of all hormones.</td>
<td></td>
</tr>
<tr>
<td>Familial/Idiopathic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cushing’s Syndrome</td>
<td>Elevated serum cortisol</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Laboratory investigations in Hyperandrogenism

Figure A. Cystic variant of Adult granulosa cell tumour – cyst wall lined by multi-layered granulosa cells H&E x 10, insert shows Call Exner bodies within stroma. H&E x 4

Figure B. Juvenile granulosa cell tumour – shows follicular pattern of cells with central eosinophilic material and round nucleus without nuclear grooves. H&E x 10

Figure C. Steroid cell tumour – shows round to polygonal cell with granular eosinophilic cytoplasm & hyalinised stroma. H&E x 10.

Figure D. Sertoli Leydig cell tumour – showing Sertoli cells arranged in cords & trabeculae. Inset – Clusters of Leydig cells (arrow) H&E x 10.
androgens are secreted by both adrenal glands & ovaries. The major androgens that are secreted are testosterone, dihydrotestosterone, dehydroepiandrosterone (DHEA), its sulphated form (DHEAS) & androstenedione. Adrenal glands secrete androgens in response to adrenocorticotropic hormone (ACTH) whereas ovaries produce in response to follicle stimulating hormones (FSH) & luteinising hormone (LH). The most important circulating androgen is testosterone & its active metabolite dihydrotestosterone. 80% of testosterone released is bound to sex hormone binding globulin (SHBG), 19% to albumin & remaining 1% is free testosterone which is the active form. These circulating androgens are metabolised in liver, conjugated with glucuronic acids or sulphuric acids and excreted in urine as 17 ketosteroids.4

The underlying cause of androgen excess can be assessed with detailed clinical history, age of onset, duration & severity of symptoms, physical examination & serum hormones.

Clinical Manifestations
The clinical signs & symptoms of hormone producing tumours depend upon age, pre/post-menopausal state and amount of androgen levels in blood. In children it causes heterosexual precocity with acceleration of growth, hirsutism & acne. In prepubertal girls it causes virilisation whereas in adult women it causes menstrual irregularities, clitoromegaly & hirsutism.5

The other symptoms that can occur in virilisation include male pattern balding, deepening of voice, skeletal muscle hypertrophy, breast atrophy & marked growth of facial and body hair.

William et al in his 10 yr study of 462 functioning ovarian tumours, has classified two definite phases of signs and symptoms in virilising tumors.6 The first one is phase of defeminisation followed by phase of masculisation. In early phase, there will be oligomenorrhoea or amenorrhea with involution of breasts and loss of female body contour. The late phase includes acne, hirsutism, clitoral enlargement, increased libido, sterility, enlargement of larynx, deepening of the voice and temporal alopecia.7,8

Causes of Virilisation in Ovarian Pathology
There are many tumours & tumour like conditions that secrete sex steroid resulting in hyperandrogenism & hyperoestrogenism. Ovarian tumours that usually secrete androgens are sex cord stromal tumours like Sertoli Leydig cell tumours, steroid cell tumours not otherwise specified (NOS), Leydig cell tumour (hilar & non-hilar type). The other causes of hyperandrogenism are depicted in table 2.

Sex Cord Stromal Tumours
Sex cord stromal tumours are ovarian tumours that have unique manifestations due to hormone production both estrogenic and androgenic leading to broad spectrum of clinical features. Androgenic tumours are Sertoli Leydig cell tumours, steroid cell tumours, rarely juvenile granulosa cell tumour and cystic variant of adult granulosa cell tumour. All these entities encountered in this present study.

Sertoli Leydig cell tumours (SLCT) are the most common cause of virilising neoplasm of ovary & constitute about 1% of ovarian tumours. Demidov et al in their study of 22 cases of SLCT found that 30% of these tumours are virilising and small percentage cause estrogenic effects.9 A study conducted by Taylor & Norris10 on 30 lipidd cell tumours of ovary shows virilisation in 77% of the cases, 5-23% shows estrogenic activity and 10% associated with Cushing's Syndrome. Granulosa cell tumour usually produces estrogens, but rarely androgenic change may occur. 50% of granulosa cell tumours with cystic gross appearance are associated with androgenic manifestations.11 Till date less than 50 cases of granulosa cell tumours with virilisation have been reported.12

Ovarian Tumours with Functioning Stroma
Some benign or malignant (primary & metastatic) ovarian tumours are hormonally active as they secrete steroid hormones by stromal cells either within or adjacent to tumours. The most common primary ovarian tumour that produce androgens are benign cystic teratoma,13 Serous14 & mucinous cystadenomas,15 Brenner tumour,16 leiomyoma17,18 and metastatic tumours that secrete androgens are Krukenberg tumour.19

Scully RE in his study of 24 cases in ovarian tumours with functioning ovarian stroma classified these cells into 3 types. They are stromal luteal cells, Leydig cells & hilus cell types. Stromal luteal cells produce estrogenic manifestations.

DISCUSSION
In females, androgens are secreted by both adrenal glands & ovaries. The major androgens that are secreted are testosterone, dihydrotestosterone, dehydroepiandrosterone (DHEA), its sulphated form (DHEAS) & androstenedione. Adrenal glands secrete androgens in response to adrenocorticotropic hormone (ACTH) whereas ovaries produce in response to follicle stimulating hormones (FSH) & luteinising hormone (LH). The most important circulating androgen is testosterone & its active metabolite dihydrotestosterone. 80% of testosterone released is bound to sex hormone binding globulin (SHBG), 19% to albumin & remaining 1% is free testosterone which is the active form. These circulating androgens are metabolised in liver, conjugated with glucuronic acids or sulphuric acids and excreted in urine as 17 ketosteroids.4

The underlying cause of androgen excess can be assessed with detailed clinical history, age of onset, duration & severity of symptoms, physical examination & serum hormones.
whereas stromal Leydig cells & hilus cell type produces androgenic manifestations.20

The major source of steroid production in these tumours occurs due to hyperplasia of adjacent stroma either with or without luteinisation. According to Baldwin et al this hyperplasia is due to mechanical effect on the stroma by the tumour which in turn stimulates stromal cells to produce androgens.21 There are certain theories to support this hyperplasia of stromal cells in ovarian tumours with functioning stroma.

Alpha glutathione S transferase has shown to be Immunohistochemical (IHC) marker for delta isomerase, an enzyme active in steroidogenesis. Tiltman et al,22 in their study of 53 epithelial tumours of ovary showed positive staining in luteinized stromal cells of both benign & malignant mucinous tumours. IHC staining of alpha glutathione S transferase in the cytoplasm of ovarian stromal cells of these tumours indicating steroidogenesis.

Hugheson postulated mechanical effect theory, in that the steroidogenesis by these stromal cells is due to the mechanical effect of expanding follicles.23 The other theory for origin of oestrogen in postmenopausal women is due to high circulating levels of pituitary gonadotrophins / HCG leading to stromal steroidogenesis as postulated in the study of Baldwin et al24 & Macdonald et al25. In toplike effect theory the tumour cells produce a substance that simulates the adjacent stromal cells to secrete hormones.

**Tumour like Lesions**

82% of hyperandrogenism in women of reproductive age groups is due to polycystic ovaries,25. It usually begins in puberty & causes hirsutism, menstrual irregularities, and insulin resistance. There are two mechanisms in which PCOS produces androgens-

1. Increased insulin levels in blood inhibits serum human binding globulin (SHBG) which in turn increases free androgens as shown in studies by Plymate et al in human hepatoma cell lines.26
2. Increased LH stimulates ovarian theca cells to produce excess androgens as shown in studies conducted in PCOS theca cell cultures.27

Idiopathic hirsutism is a common cause of virilisation occurring after puberty, which are slowly progressive & are often familial. They are diagnosis of exclusion & occur due to disorder in peripheral androgen activity.28

Pregnancy luteomas are asymptomatic hyperplastic tumour like lesions that are discovered incidentally during surgery. Patients will be hormonally active, and the symptoms disappear after delivery.29

**Adrenal Causes**

Adrenal causes of virilisation include Cushing’s syndrome, congenital adrenal hyperplasia (classical & non-classical), adrenal adenoma & carcinoma. Cushing’s syndrome must be differentiated from polycystic ovarian syndrome as both have same clinical manifestations except for the elevated level of serum cortisol in Cushing’s syndrome & some physical characteristics like central obesity, facial plethora, moon face, and violaceous abdominal striae greater than 1 cm.30,31

Congenital adrenal hyperplasia (CAH) occurs due to deficiency of 21 alpha hydroxylase resulting in accumulation of steroid precursors which are subsequently converted into androgens. There are two types, classical & non-classical. Classical CAH usually presents at birth whereas non-classical CAH does not present until early puberty. Classical type produces clitoral enlargement in females at birth & masculinising features develop as the child grows. In non-classical type there will be premature development of pubic hair, acne & menstrual irregularities.30,32

Adrenal adenomas & carcinomas also produce androgens leading to virilisation.

**Exogenous Androgens**

Many drugs induce hyperandrogenism but the most common drug to induce is synthetic progestins like norgestrel. There are two mechanisms in which synthetic progestins increases testosterone levels. First, they bind directly to androgen receptors thereby inducing androgenic effects; secondly, they decrease serum human binding globulin and increases circulating active testosterone.30, Other drugs which induce hyperandrogenism are danazol & anabolic steroids.

**In investigations**

The laboratory investigations that are usually done for hyperandrogenism are serum testosterone, DHEA & DHEAS, 17 hydroxyprogesterone, low dose dexamethasone suppression test, urinary 17 ketosteroid, serum cortisol & prolactin. The laboratory tests in various causes of hyperandrogenism are depicted in table 3.

**CONCLUSION**

Patients presenting with symptoms of virilisation require detailed clinical history, physical examination for signs, laboratory investigations & imaging. Rapid development of hirsutism, late onset & signs of virilisation should raise suspicion of androgen secreting ovarian neoplasm from adrenal tumours. Serum testosterone level with DHEAS helps in differentiating androgen secreting ovarian or adrenal tumours.

**Abbreviations**

PCOS – Polycystic ovarian syndrome.
ACTH – Adrenocorticotrophic hormone.
FSH – Follicle stimulating hormone.
LH – Luteinising hormone.
DHEA – Dehydroepiandrosterone.
DHEAS – Dehydroepiandrosterone sulfate.
17 OHP – 17 hydroxy progesterone.
IHC – Immunohistochemistry.

**REFERENCES**


