Feminizing adrenocortical carcinomas are defined as malignant tumors with estrogens overproduction with or without other adrenocortical hormones. They are rare even in endocrinology as they account for less than 1−2% of all adrenal tumors. They are mainly observed in men (median age = 42 years, range = 19−77 years); they are rare in children and exceptional in women. The prognosis is poor, especially when metastases are present at diagnosis.

In males, the main symptom is gynecomastia, which may or may not be associated with other hypogonadism features. Our aim was to describe a case without gynecomastia and try to understand why this important clinical symptom was absent in our patient.

Feminizing Adrenocortical Carcinoma Without Gynecomastia

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ABSTRACT
Malignant feminizing adrenocortical tumors are exceedingly rare. Their main presentation is gynecomastia. In these estrogen secreting tumors (with or without other adrenocortical hormones) lack of gynecomastia is exceptional as in our case. A 44-year-old man presented with abdominal pain. Radiological examination revealed a tumor measuring 120 × 95 mm in the retroperitoneal area with numerous metastases. Pathological examination pleaded for an adrenal origin with a Weiss’s score of 5. Six months later, the tumor relapsed, and he had a second surgery and was sent for hormone assessment. Clinical examination showed a skinny man with severe fatigue. He had no Cushingoid features. Gynecomastia and galactorrhea were absent. Penile length, testicular volume, and body hair growth were normal. Several cutaneous nodules were present. Biological assessment showed high morning plasma cortisol, which failed to be suppressed by treatment with 2 mg dexamethasone. Plasma estradiol and 17OH progesterone levels were high, but his testosterone levels were low. Radiological exploration showed numerous metastases: pleural, pulmonary, retroperitoneal, and abdominal. He was treated with classical chemotherapy, but he died four months after diagnosis.

A 44-year-old man presented with abdominal pain. Radiological assessment revealed a large tumor measuring 120 × 95 mm in the left retroperitoneal area with numerous metastases. He was operated on without any hormonal assessment. Pathological examination showed an adrenal origin with a Weiss’s score of 5 [Figure 1]. Six months later, he had another surgery for a relapsing tumor and was sent to our department for hormonal evaluation. Clinical examination showed a skinny man with severe fatigue and anorexia. His body mass index (BMI) was 16 kg/m², and his blood pressure was 110/60 mm. He had no Cushingoid features. Gynecomastia and galactorrhea were also absent. Penile length, testicular volume, and body hair growth were normal. However, many cutaneous nodules were present in the thoracic and abdominal areas. Hormonal assessment showed mixed adrenal secretion where estrogens were prevailing. He had high morning and midnight plasma cortisol, which failed to be suppressed with 2 mg dexamethasone, and very high estradiol and 17OH progesterone contrasting with low testosterone [Table1]. Imaging studies revealed a relapsing tumor with numerous pleural, pulmonary, abdominal, and retroperitoneal metastases [Figure 2].

As Mitotane treatment was not available, he had chemotherapy (doxorubicin, carboplatin, and
etoposide). He did not respond well to the treatment and died four months after diagnosis.

**DISCUSSION**

Secreting and non-secreting adrenocortical carcinomas are rare malignant tumors with a dire prognosis, and typically occur in males in the fourth and fifth decades of life. Adrenocortical neoplasms may lead to a variety of endocrine syndromes depending on the type of produced hormones. When estrogens overproduction occurs, the term feminizing adrenal tumors (FATs) is used. Most of these tumors are malignant. Benign tumors, although rare, exist.

Feminizing adrenocortical carcinomas are exceedingly rare in adults and account for 1−2% of adrenocortical carcinomas. They are prevailing in adult males, although they can be observed in children. Female cases are exceptional.

Patients with FATs usually complain of decreased libido and erection or ejaculation problems. They also have gynecomastia with or without other features of hypogonadism. In 1965, the first literature review recorded 52 FATs and demonstrated that patients with FATs have gynecomastia (98%), a palpable abdominal mass (58%), and a testicular atrophy (52%). Patients with FATs usually complain of diminished libido (48%) and breast tenderness (42%).

On biological assessments, estrogens overproduction, with or without an increase in other adrenal hormones, are the main abnormalities. The increase in estrogens production results from peripheral androgens conversion, but also from direct production of the adrenocortical tumor.

Gonadotropin hormones are normal-low or decreased. The response to luteinizing hormone-releasing hormone is generally blunted.

The main characteristic of these adrenocortical tumors is their ability to produce estrogens, although cortisol may be subnormal or high with or without clinical expression. Women can develop androgen excess. Estrogens overproduction occurs through an increase in androgen substrates such as androstenedione. However, it can also result from augmentation in aromatase activity within the tumor leading to an increased synthesis of estrone.

Bilateral breast enlargement results from the imbalance between high estrogen and low free androgens. Gynecomastia is the major clinical expression. However, its absence does not eliminate the diagnosis of FATs. When gynecomastia is missing, one should discuss the resistance of breast tissue to estrogens, very rapid evolution, and severe protein degradation.

FATs have a very poor prognosis. Mortality among adult males with FATs is high as the three-year survival is less than 20% after tumor resection.

**Table 1:** Hormonal evaluation of the patient.

<table>
<thead>
<tr>
<th>Hormones</th>
<th>Patient’s results</th>
<th>Normal range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estradiol</td>
<td>645.32*</td>
<td>55–165 pmol/L</td>
</tr>
<tr>
<td>Cortisol level</td>
<td>784.34</td>
<td>50–550 nmol/L</td>
</tr>
<tr>
<td>8 AM</td>
<td>453.00</td>
<td></td>
</tr>
<tr>
<td>Midnight</td>
<td></td>
<td></td>
</tr>
<tr>
<td>17OH Progesterone</td>
<td>11.7</td>
<td>0.0–1.6 nmol/L</td>
</tr>
<tr>
<td>Androstenedione</td>
<td>8.82</td>
<td>0.3–3.1 ng/mL</td>
</tr>
<tr>
<td>DHEA-S</td>
<td>1337</td>
<td>95–530 µg/dL</td>
</tr>
<tr>
<td>Total testosterone</td>
<td>6</td>
<td>10–40 nmol/L</td>
</tr>
</tbody>
</table>

*DHEA-S: dehydroepiandrosterone sulfate; *mean value.*
especially in patients diagnosed late with multiple abdominal or extra-abdominal metastases.

Common sites of intra-abdominal metastases include the liver, intestine, peritoneum, and retroperitoneal lymph nodes. Extra-abdominal metastases may be observed in the lungs, pleura, bones, and other sites.

Treatment is based mainly on surgery with complete resection of the tumor and its metastases when possible. For recurrent or metastatic FATS, radiation therapy and various chemotherapies can be considered. However, with the exception of Mitotane, there is no other effective medication.\textsuperscript{5} Aromatase inhibitors have been used without success.\textsuperscript{5}

**CONCLUSION**

Feminizing adrenal carcinomas are exceptional and have a dire prognosis. Gynecomastia, which is the classical manifestation, was absent in this patient probably due to the resistance of the breast tissue to estradiol.

**Disclosure**
The authors declared no conflicts of interest.

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**REFERENCES**


**Figure 2:** (a) A large adrenal tumor (orange arrow), (b) pulmonary metastases (red arrows), and (c) abdominal (blue arrow), retroperitoneal lymph nodes (yellow arrow), and pleural effusion (green arrow).