A rare ectopic ACTH secreting atypical pulmonary carcinoid- case report

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Abstract

Pulmonary carcinoids are rare pulmonary neoplasms which account for 1-2% of all lung neoplasms. Pulmonary carcinoid tumors are a rare cause of ectopic ACTH secretion. The incidence of Cushing's syndrome in pulmonary carcinoid tumor is approximately 1%. A case of 25-year-old young male with typical cushingoid appearance presented with headache on and off and episodes of hypertension since 1year. Thoracic CT revealed a nodular region in left lower lobe. The patient underwent left lower lobectomy and histopathology revealed an atypical carcinoid tumor. On immunohistochemistry, tumor cells were positive for CK PAN9, CD117, S100, TTF-1, synaptophysin, chromogranin and negative for P40, CK7, Her-2 and CD99. This case is presented for its rarity.

Keywords: Pulmonary, Ectopic Cushing’s, Carcinoid tumor

Introduction

Carcinoid tumors are the neuroendocrine tumors arising from enterochromaffin cells (type of enteroendocrine or neuroendocrine cells occurring in epithelia lining of digestive tract and respiratory tract that release serotonin). The gastrointestinal tract is the most common site for carcinoid tumors. Of all carcinoids, bronchopulmonary carcinoids represent only 10% cases.

Pulmonary carcinoids are rare pulmonary neoplasms which account for 1-2% of all lung neoplasms. Bronchial carcinoids represent small portion of all pulmonary tumors and about 1-5% of them are associated with ectopic ACTH secretion, which represents 1-10% of Cushing syndrome cases [1-3]. They are a rare cause of ectopic ACTH secretion.

Cushing syndrome is the result of chronic exposure to increased concentration of cortisol hormone, exogenous or endogenous. The endogenous Cushing syndrome comprises three distinct pathogenic disorders: pituitary, adrenal and ectopic [4]. Ectopic Cushing syndrome (ECS) results from autonomous ACTH production from extra pituitary malignancies with elevated plasma levels of ACTH accounting for 15% of cases of Cushing syndrome [5]. It is generally associated with central obesity, metabolic syndrome and hypertension [6].

Neuropsychological disturbances are also frequently observed including depression, emotional irritability, sleep disturbances and cognitive deficits that can be the first manifestation perceived by family members of patients affected by Cushing’s syndrome.

Typical carcinoids are more common and atypical carcinoids are the rarer ones among all pulmonary carcinoids. Treatment is based on decreasing the cortisol levels in blood. Surgery is considered as the mainstay of treatment for ectopic ACTH secreting tumors.

Case Report

A 25-year-old young male presented with headache on and off and episodes of hypertension since 1year. On physical examination, the patient presented with Cushingoid features (moon facies, extensive purple striae on abdomen and arms, marked central obesity and peripheral edema. Blood pressure was high (158/90 mm Hg), cardiac and pulmonary auscultations were normal and there were no abdominal palpable masses.
Routine laboratory results showed-
- Hypercholesterolemia (total cholesterol-270 mg/dL, HDL cholesterol, 82 mg/dL, LDL cholesterol 171 mg/dL and triglycerides 152 mg/dL)
- Elevated plasma glucose 132mg/dL with HbA1c 6.03%
- Hb-13.5 g/dL
- Hypokalemia (K⁺ = 2.6mmol/L) with no other electrolyte disturbances.
- Renal function tests and liver function tests were normal.

As clinical suspicion of endogenous hypercortisolism was high, screening for cortisol levels was done. Tests revealed elevated cortisol levels(45.8µg/dL) and plasma ACTH levels.

<table>
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<th>Morning</th>
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<td>11.26am</td>
<td>255 pg/ml</td>
</tr>
<tr>
<td>11.30am</td>
<td>268 pg/ml</td>
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<tr>
<td>11.32am</td>
<td>272 pg/ml</td>
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<tr>
<td>11.35am</td>
<td>292 pg/ml</td>
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To identify a potential pituitary lesion, a pituitary MRI was performed, there was no identifiable tumor. Bilateral Inferior Petrosal Sinus Sampling (IPSS) revealed peripheral source of ACTH.

Plain chest X-ray showed no abnormalities. Bronchoscopy and cytological examination of bronchoalveolar lavage were negative for malignancy.

Chest CT scan revealed one nodular lesion in the left lower lobe of lung measuring 1.1x0.8cm. A whole body Ga-68 DOTANOC PET CT was planned to characterize the lesion as well as to localize metastatic lymph node which revealed centimetric nodular lesion (1.1x0.8cm) with no significant tracer uptake was seen in left lung lower lobe and a possibility of atypical carcinoid/small cell carcinoma was made.

The patient underwent lower lobe lobectomy of left lung. The resected specimen was sent for histopathological examination.

**Pathological findings**

**Gross**- Lower lobe of left lung was received measuring 10x7x4cm. Bronchus was identified on careful examination, just adjacent to it a growth measuring 1x0.5cm was present (Figure 1).
Microscopy- Sections given from nodular area comprised of nests of uniform population of cells. Cells had scanty to moderate granular cytoplasm. Nuclei were round to oval and central having granular chromatin and inconspicuous nucleoli. Occasional mitotic activity was seen. Surrounding lung parenchyma showed occasional tiny satellite nodules. Rest of the lung parenchyma showed congestion of blood vessels and small number of alveolar macrophages. The resected margin of specimen was free from tumor invasion.

Overall picture was suggestive of atypical bronchial carcinoid tumor.

Immunohistochemistry was advised for proper evaluation.

Markers positive in tumor cells were CK PAN9 (dot like), CD117, ACTH, S100, Synaptophysin, TTF-1 and Chromogranin; negative markers were P40, CK7, Her-2, CD99.

Immunomarkers favored neuroendocrinal tumor with features of atypical carcinoid.

Fig-2: Section representing area of a normal lung and the nodular area.

Fig-3: Image representing section from the nodular area.

Fig-4: Image representing section of the bronchial cartilage.
After the surgery, the patient responded well. The plasma ACTH levels decreased to 25pmol/L. Cortisol level decreased to 6µg/dL. Blood pressure decreased to 130/80. Overall, the patient was symptomatically well with good lung function.

Discussion

Carcinoid was first discovered in late 19th century by Lubarsh, who found multiple tumors in the small bowel of two patients at autopsy. In neuroendocrine tumors, 3 grades on histologic features and biologic behavior are currently recognized:

- Grade I or typical carcinoid
- Grade II or atypical carcinoid
- Grade III or small cell carcinoma/large cell carcinoma

Typical carcinoids occur in both sexes and age of onset ranges from childhood to 9th decade. Atypical carcinoid appear in older age groups. Typical carcinoids are well differentiated, can exhibit distinct recognized histologic pattern (insula, trabecular, glandular, mixed or undifferentiated) and rare mitoses. They usually present as perihilar masses and are generally asymptomatic; however some patients may present recurrent pneumonia, cough, hemoptysis and chest pain.

Ectopic secretion of biologically active hormones is not uncommon, in particular corticotrophin and growth hormone and it occurs in fewer of 5% of patients with these tumors. Conversely, atypical carcinoids represent an intermediate grade neuroendocrine tumors; they may exhibit increased nuclear atypia, focal necrosis or high mitotic indices and their clinical course is aggressive with an high incidence of mediastinal lymph node metastases and a 5-year survival rate of 40–75% [7].

The differential diagnosis of Cushing syndrome and in differentiation of pituitary Cushing syndrome from an ectopic ACTH secreting neoplasm can be difficult [8].

The most useful test is inferior petrosal sinus sampling where patients with pituitary lesions show a gradient in ACTH concentration between the affected side sinus and the periphery in contrast to an ectopic ACTH syndrome where there is no gradient [9]. Localization of the source of ectopic ACTH can be problematic [10]. Small peripheral bronchial carcinoids can be easily missed on CT chest due to poor inspiratory effort, abdominal fat [11].

Confirmation of ectopic ACTH production requires demonstration of immunostaining positive for ACTH in the resected tumour. This may be difficult if the source was metastatic malignancy as in this case only a subpopulation of the cells will produce ACTH making it difficult to be demonstrated by staining. More recently, extraction of appropriate mRNA by real time PCR provides a highly specific means of identifying these tumours [12].

Advances in medical and surgical treatments have improved the overall survival rate for patients with ectopic ACTH [13]. Prognosis depends on primary tumor histology [14]. Patients with small cell lung carcinoma (SCLC) had the worst prognosis, usually dying within 12 months of diagnosis (median 6–8 months) [15,16,17]. Patients with bronchial carcinoids have the best prognosis and are usually considered to have low to moderate grade malignancy [18]. They do have a malignant potential, however, and can be associated with metastases.

Conclusion

Ectopic ACTH secreting tumors present some of the most challenging differential diagnosis in endocrinology and require careful clinical, radiological and pathological investigation. These tumors are best managed in a multidisciplinary setting with close relation between the endocrinologist, endocrine surgeon, pathologist and radiologist.

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References


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