Bronchial Carcinoid: Case Report and Review of Literature

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Abstract

Bronchopulmonary carcinoid tumors which are slow growing neuroendocrine tumors arising from Enterochromaffin (EC) cells, accounts for 10% of all carcinoid tumors. Mentioned here is a case report of 46 year old male with incidental finding on CXR of left hemithorax inhomogenous opacity and features suggestive of volume loss. CECT showed a well defined 2.5 x 1.5 cm circular intraluminal growth occluding the left main bronchus. Bronchoscopy revealed a red, shiny, smooth walled growth 1 cm beyond the carina in the left main bronchus, attached to the roof of the bronchial wall, causing ball valve mechanism. Histopathology and Cytology of needle aspiration from tumor confirmed the diagnosis of bronchial carcinoid. To avoid misdiagnosis or delay in diagnosis of intrabronchial carcinoid a high index of suspicion should be maintained and radiological, brochoscopical and pathological investigations for accurate and early diagnosis must be undertaken.

Keywords: Bronchial Carcinoid Tumor, Diagnosis, Treatment of Bronchial Carcinoid

1. Introduction

Enterochromaffin (EC) cells, or "Kulchitsky cells", are a type of enteroendocrine and neuroendocrine cells occurring in the epithelia lining the lumen of the digestive tract and the respiratory tract that release serotonin. Carcinoid tumors are neuroendocrine tumors arising from Enterochromaffin (EC) cells1. The Gastrointestinal (GI) tract is the most common site for carcinoid tumors. Of all carcinoids, bronchopulmonary carcinoids represent only 10% cases, of which 80-90% arise within the segmental or subsegmental bronchus and about 10-20% are found in the periphery. Only about 1% to 6% of all lung tumors are carcinoid tumors. Typical carcinoids are more common and atypical carcinoids account for only 10% of all pulmonary carcinoids. Symptom free incidental detection of pulmonary carcinoids are a common occurrence as 25-39% of these patients are asymptomatic and carcinoid syndrome occurs in about 2% of cases only2. Early diagnosis is extremely important as patients benefit from surgical resection and prognosis is very good.

We present a case of bronchial carcinoid detected following evaluation of incidental findings on Chest Radiograph (CXR). Patient was asked to get Contrast Enhanced Computed Tomography (CECT) Thorax, and Bronchoscopic Needle Aspiration of the tumor was done to confirm the diagnosis.

2. Case Report

46 year old male patient was referred to pulmonary medicine OPD for CXR changes (Figure 1.) detected incidentally during evaluation for fracture tibia post road traffic accident. He had no associated chest trauma. On further enquiry he stated that he perceives MMRC grade
1 breathlessness on exertion since 4 months. He had no other chest or constitutional symptoms. There was no significant past or family history. Patient has history of smoking 20 cigarettes per day for 15 years and significant history of alcohol consumption. On admission patient was afebrile and had pulse rate of 88 per minute, respiratory rate of 12 per minute and blood pressure of 110/70 mm Hg and normal general examination, no pallor, no clubbing or lymphadenopathy. His chest examination revealed signs of volume loss, decreased movements, dull percussion note, decreased breath sounds over left hemithorax with bronchial breathing over left interscapular and infrascapular areas. Laboratory investigations (complete hemogram, renal and liver function tests) were unremarkable. CXR (Figure 1.) showed left hemithorax inhomogenous opacity with left shift of mediastinum and features of volume loss. Right hemithorax and bony-soft tissue shadows appeared normal. He was further evaluated with CECT chest (Figure 2(b)) which showed collapse of left lung with minimum aeration noted proximally. A well defined 2.5 x 1.5 cm (Figure 2(a)) circular growth was observed extending from the left main bronchial wall intraluminally occluding the airway. Right lung showed well developed compensatory hyperinflation, causing herniation of right lung to left. There was no evidence of any lymphadenopathy. Bronchoscopy revealed deviated and dilated right lung bronchial openings with normal wall structure. Left main bronchus was occluded due to the mass 1 cm beyond the carina with red, shiny, smooth walled growth attached to the roof of the bronchus (Figure 3). On deep breathing and during coughing the growth moved causing a ball valve mechanism and allowing only minimum airflow to the left lung. Reclining position of the patient showed movement of secretions from the left main bronchus in to the right bronchial branches. There was no bleeding on touch. Needle aspiration was performed following which there was bleeding at the site which was quickly controlled by cold saline wash. Cytology (Figure 4.) of the specimen showed few foci of aggregates of small round cells with monomorphic hyperchromatic nuclei surrounded by scanty cytoplasm. Patient has been referred for excision of the tumor.

**Figure 1.** Chest radiograph showing left lung collapse (left hemithorax homogenous opacity with volume loss).

**Figure 2.** (a) Mediastinal window during phase of contrast shows a well defined 2.5 x 1.5 cm round bordered intraluminal growth in the the left main bronchus occluding the airway. (b) Lung window shows near complete collapse of left lung with minimum aeration post obstruction. Right lung showed well developed compensatory hyperinflation changes only, causing herniation to left.
3. Discussion

Carcinoid was first described in the late 19th century by Lubarsh, who found multiple tumors in the small bowel of 2 patients at autopsy and Siegfried Oberndorfer, a German pathologist at the University of Munich, coined the term karzinoide, or “carcinoma-like”, to describe the unique feature of behaving like a benign tumor despite having a malignant appearance microscopically. Its endocrine-related properties were later described by Gosset and Masson in 1914. Hamperl in 1937 reported the first description of pulmonary carcinoid.

The overall incidence of carcinoid tumors is estimated to be 1 to 2 cases per 100,000 individuals in the United States and is similar in England, Scotland, Spain, Italy, and Japan. This incidence varies with gender, age, and race Carcinoids generally have a higher incidence in African-Americans. A recent analysis of 13,715 carcinoid tumors from the Surveillance, Epidemiology, and End Result (SEER) Program of the National Cancer Institute found that 67.5% of all carcinoid tumors originate in the GI tract and 25.3% originate in the lungs or bronchi. Depending upon the degree of differentiation and lymphnode metastasis Typical Carcinoids have the best prognosis, with a 10-year survival rate of more than 80%. The 5-year survival rate in Atypical Carcinoid without lymph node metastases is 80%, and for Atypical Carcinoid with lymph node metastases is 60%. High vigilance is to be maintained in Typical Carcinoid since recurrence and distant metastasis can develop even after 10 years. Local resection of metastatic lesion in liver also has a 5 year mean survival rate of 78%. The average age of occurrence of Typical Carcinoid tumors is 40-50 years, but Typical Carcinoid tumors have been reported in virtually every age group. Atypical Carcinoid tumors appear in slightly older people than Typical Carcinoid tumors. Carcinoid tumors occur in equal numbers in males and females.

3.1 Risk Factors

Smoking is a major risk factor for many malignancy and bronchial carcinoids are no different. Although the association between bronchial carcinoids and smoking is unclear, between one-third and two-thirds of all patients have been smokers with higher prevalence of smoking being reported in patients with Atypical Carcinoids. No other known carcinogens or exposure to environmental agents has been implicated in carcinogenesis.

Patients with the autosomal dominant syndrome of multiple endocrine neoplasia type 1 (MEN 1) have a high frequency of endocrine malignancies, and foregut carcinoids (i.e., thymus, lung, stomach, or duodenum) arise in approximately 2 percent of cases. There are reports of familial pulmonary carcinoids not associated with the MEN syndrome.

3.2 Clinical Characteristics

The patients of carcinoid most commonly presents with haemoptysis, cough, recurrent pulmonary infections, fever, chest discomfort, unilateral wheezing and shortness of breath. 25 to 40% of patients are asymptomatic. Pulmonary carcinoids may release corticotropin,
resulting in Cushing’s syndrome or GH-releasing hormone (GH-RH) resulting in acromegaly22. Carcinoid syndrome, which occurs when the vasoactive substances escape into the systemic circulation escaping the hepatic degradation, is seen in less than 5% of patients with pulmonary carcinoid23. Metastases occurs usually to mediastinal lymph nodes. The second common site of metastasis is the liver. Rarely the carcinoid tumor may spread to bone, or skin. Overall incidence of metastasis have been identified in fewer than 15% of cases23–25.

In author, presented a case of carcinoid in a 39-year old man with persistent wheezing, episodes of haemoptysis and dry cough who was misdiagnosed as bronchial asthma and was treated with inhaled beta2-agonists and steroids for about 10 months without cure. In author presented a case report of bronchial carcinoid in a 28 year old man who was misdiagnosed as bronchial asthma. Our patient had right lower limb tibial fracture due to road traffic accident and his CXR was done as a screening investigation. Based on the incidental findings on his CXR (Figure 1.), he was initially considered to have a post tuberculous destroyed lung. As he had no past history of tuberculosis he was further evaluated with CECT on which the endobronchial growth was detected. The above cases suggests that absence of characteristic symptoms may lead to misdiagnosis and delay in diagnosis of intrabronchial carcinoid. Thus patients with respiratory symptoms inspite of optimum medical treatment should undergo radiological, bronchoscopical and pathological investigations for accurate and early diagnosis.

3.3 Histology

The first classification of neuroendocrine malignancies was proposed in 1972, which defined Atypical Carcinoid according to histologic criteria, including the number of mitoses per high–power field, the presence of necrosis, increased cellularity with disorganization, nuclear pleomorphism, hyperchromatism, and an abnormal nuclear–to–cytoplasmic ratio. In 1991, In author proposed 4 categories of neuroendocrine lung tumors: Typical Carcinoid, Atypical Carcinoid, Large Cell Neuroendocrine Carcinoma (LCNEC), and Small Cell Neuroendocrine Carcinoma (SCNEC) which was also supported by World Health Organization in 2004 and 2015.

The World Health Organisation/The International Association for the Study of Lung Cancer (WHO/IASLC) classifies Carcinoids according to histological differentiation:

- Typical Carcinoid (<2 mitoses/2 mm2 and no necroses).
- Atypical Carcinoid (2–10 mitoses/2 mm2 and/or confirmed necroses).

3.4 Serum Markers

The diagnostic test of choice for suspected carcinoid tumor is the measurement of urinary levels of the major serotonin metabolite 5-HIAA in a 24-hour urine sample (normal range 2–8 mg/24 hours). Approximately 50% of patients with carcinoids have elevated levels of urinary 5-HIAA, whether or not they have carcinoid syndrome.

An elevated urinary 5-HIAA level has a diagnostic sensitivity of 70% and a specificity of 88% to 100%.

Chromogranin A (CGA), is a 49-kD protein, another tumor marker present in the neurosecretory vesicles of neuroendocrine tumor cells. It is detectable in the plasma of patients with endocrine neoplasms. Because it does not rely on serotonin secretion, serum CGA is a more sensitive and broadly applicable marker than urinary 5-HIAA. Serum CGA levels is measured in patients with metastatic small bowel and appendiceal carcinoid tumors, and also bronchial and rectal carcinoid tumors in whom urinary 5-HIAA levels are less likely to be elevated.

3.5 Radiology

Bronchial carcinoids are visualised on chest radiographs or Computed Tomography (CT). Bronchial Typical and Atypical Carcinoids have similar radiologic features, which depend largely on tumor location. About 80% of bronchial carcinoids arise centrally in the main, lobar and segmental bronchi. Radiological findings include hilar or perihilar masses, endobronchial nodules, findings related to bronchial obstruction, and peripheral nodules. Hilar or perihilar masses are usually well-defined, round or ovoid lesions and may be slightly lobulated at radiography and CT. The tumors range from 2 to 5 cm in size. In approximately 20% of cases, a bronchial carcinoid presents as a solitary pulmonary nodule in the lung periphery, distal to segmental bronchus. These lesions are usually round or ovoid with smooth or lobulated borders. Typical Carcinoids in the periphery are slow growing and should be considered in the differential diagnosis of slow-growing solitary pulmonary nodules. Atypical Carcinoids mostly occur in the lung periphery and are
usually large38,39. Carcinoids are characteristically highly vascular. All carcinoids does not show enhancement and enhancement alone cannot differentiate bronchial carcinoid from bronchogenic carcinoma or does not excludes the diagnosis.

CT using contrast medium is the best method for identifying extrabronchial extends of tissue and mediastinal lymph node enlargement in central tumors. Because of their hypervascularization, carcinoid tumors absorb contrast medium and are often seen as well defined, obstructing tumors40 Up to 20% of Typical Carcinoids are accompanied by hilar or mediastinal lymphadenopathy, which is mostly caused by a reactive inflammatory reaction41. Endobronchial ultrasound guided Transbronchial Needle Aspiration (EPBUS-TBNA) or mediastinoscopy undertaken for the purpose of staging according to the TNM classification for lung cancer42. The histological differentiation grade follows the classification of WHO/IASLC43. Typical Carcinoids are mostly diagnosed at stage I, Atypical Carcinoids mostly at stage II (N1, [hilar] lymph node involvement) or stage III (N2, [mediastinal] lymph node involvement). Hepatic metastases (stage IV) can be detected by using three-phase CT or, alternatively, by using ultrasonography.

3.6 Bronchoscopy
Fiberoptic Bronchoscopy is the best method of choice for tissue diagnosis. About three-fourths of bronchial carcinoids are centrally located and approachable to biopsy during bronchoscopy. The bronchoscopic appearance is a typically pink to red vascular mass with intact overlying bronchial epithelium. Carcinoids are generally attached to the bronchus by a broad base but can be polypoid and create a ball-valve effect44.

Bronchial carcinoids are vascular, and there has been concern for bleeding in the past, particularly after flexible bronchoscopy with biopsy. Inspite of vascular nature, the incidence of serious bleeding complications during bronchoscopic biopsy is very low (<1%)43. The administration of a diluted epinephrine solution before and after biopsy of a suspected endobronchial carcinoid may have reduced the risk of severe bleeding44.

3.7 Treatment
3.7.1 Surgery
All pulmonary carcinoid tumors should be treated as malignancies. Surgery is the only curative approach, and, in view of lack of prospective studies, it is the therapeutical mode of choice in all recommendations45,46. As it is with any other tumor, the most important objective is a microscopically tumor-free resection margin. Following resection of the tumor, 5-year survival rate of 94% was observed in 1109 patients with Typical Carcinoids by The European Society of Thoracic Surgeons Euroendocrine Tumors Working Group42,45 and 3 year survival rate of 67% was observed in the United States by a database analysis of 441 patients with AC47. Another retrospective analysis of 84 patients conducted by43 showed a 5 year survival rate of 91% in patients with Typical Carcinoids and 90% in patients with Atypical Carcinoids43. It is recommended that patients having AC should undergo radical lymphnode resection due to expected high chances of metastases to adjacent lymphnodes. However, patients with Typical Carcinoids are also advised draining lymphnode resection or biopsy since chances of metastasis is present47.

Lung saving surgeries are preferred. Patients having peripheral tumors can be considered for wedge resection or lobectomy48. Certain central tumors need sleeve excision with angioplasty and bronchoplasty, thus further reducing both morbidity and mortality48. Endoscopic resection of endobronchial tumors and endoscopic laser resection can be performed in selected cases. In49, used bronchoscopic laser therapy for endobronchial TC in 11 patients, out of which 6 patients were proven disease free following surgical resection. Rigid bronchoscope is preferred, as it allows multiple uses of instruments at a time and gives good ventilatory support. Most pulmonologists prefer flexible bronchoscope for laser photoesection50.

3.7.2 Treatment of Unresectable Tumors
Various chemotherapy agents have been tried. Somatostatin receptor Analogues (SSA) such as octreotide and lanreotide, which are primarily used for controlling symptoms in carcinoid syndrome, are used along with mTOR inhibitor everolimus (RADIAN T-2 Study) to increase survival rate to 16.4 months from 11.3 months as compared to placebo51–53. The phase II study of Peptide Receptor Radionuclide Therapy (PRRT) using radioactively marked 90Ytrrium (Y) or 177Lutetium (Lu)-SSA (e1) has shown response to treatment in patients with somatostatin refractory tumors44. A pilot study is been conducted for somatostatin receptor ligand 177Lu-DOTATATE55.

Carcinoids are radio resistant, however local lymphnode radiation at atypical carcinoids and unresectable
tumors is being considered as a palliative therapy. Inoperable tumors can be resected bronchoscopically as a palliative approach to relieve obstruction and associated collapse-consolidation. Local resection of metastatic lesion in liver has a 5 year mean survival rate of 78%\textsuperscript{15}.

4. Summary
Carcinoid tumors are malignant neuroendocrine neoplasms arising from Enterochromaffin (EC) cells. Clinical serological as well as radiological studies are unable to differentiate pulmonary neoplasm from bronchial carcinoid. Bronchoscopic needle aspiration or biopsy of carcinoid followed by histological study is the best modality for tissue diagnosis. The survival of patients with carcinoid tumor depends basically on the histologic type and the presence of distant metastasis. Typical bronchial carcinoids rarely metastasize and have an excellent prognosis even when regional lymph nodes are involved; Atypical Carcinoids have a higher likelihood of metastases and a worse prognosis, particularly if mediastinal nodes are involved. Surgical resection is the treatment of choice with good long term survival rate. To avoid misdiagnosis or delay in the diagnosis of intrabronchial carcinoid, the patients with refractory respiratory symptoms inspite of optimum medical treatment should undergo clinical, radiological, bronchoscoical and histopathological investigations for accurate and early diagnosis.

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