A pulmonary pleomorphic carcinoma is a rare, poorly differentiated, primary malignancy. We report an atypical manifestation of this rare tumor, with the air crescent and CT halo signs.

Index words: Carcinoma  
Lung neoplasms  
Tomography, X-ray computed

Pulmonary pleomorphic carcinoma is rare, accounting for approximately 0.1–0.4% of all lung malignancies. These tumors usually present in symptomatic older male smokers as large peripheral masses, often with chest wall invasion and metastases at diagnosis [1]. We recently experienced an atypical case of pleomorphic carcinoma with the air crescent and CT halo signs in a young woman without a history of smoking. The air meniscus and CT halo signs associated with pleomorphic carcinoma have not been previously reported. Here we present the radiology and pathology correlations as well as a review of the medical literature.

Case Report

A 38-year-old woman was referred to our hospital with an abnormality on chest X-ray. The patient complained of a dry cough over the past few months. In addition, she had neither a history of smoking nor a significant medical history. Upon admission, her physical and laboratory findings were unremarkable. In addition, sputum studies for acid-fast bacilli, Aspergillus organisms, other pathological organisms, and malignant cells were negative. On the initial chest X-ray, a large mass was discovered in the right lower lobe of the lung with the air crescent (Fig. 1A). The chest CT showed a 6 cm sized sharply defined, homogenous mass with lobulation at the periphery of the right lower lobe of the lung (Fig. 1B). After contrast infusion, the mass enhanced inhomogeneously with a focal eccentric area of decreased enhancement. In the lung window setting, the mass was surrounded by ground glass attenuation (Fig. 1C). An interface between the ground glass attenuation and normal parenchyma was relatively sharply delineated. A crescent-shaped air space with a smooth inner surface was seen at the anterior aspect of the mass. The mass obstructed the anterior basal segmental bronchus, resulting in segmental atelectasis. There was no evidence of pleural effusion, chest wall invasion, lymphadenopathy, or distant metastasis. The differential diagnosis before surgery included sclerosing hemangioma with peritumoral hemorrhage, lung cancer arising from a cavity, and cavitory lung cancer. Aspergilloma was ruled out by
Fig. 1. Pleomorphic carcinoma in a 38-year-old woman.
A. Initial chest PA reveals a thick-walled cavitary mass with an air crescent sign in the right lower lobe of the lung.
B. Axial contrast enhanced CT scan shows a 6 cm sized, lobulated, inhomogeneous enhancing mass located at the periphery of the right lower lobe of the lung (white thick arrow). Note the segmental atelectasis at the anterior basal segment, caused by the obstruction of the mass.
C. The CT image obtained using lung window settings shows a crescent-shaped air space (arrowhead) as well as peritumoral areas of ground glass attenuation (double thin black arrows). Note the relatively sharp interface between the ground glass attenuation and normal lung parenchyma.
D. Photomicrograph of the histopathological specimen shows solid tumor with multi-nucleated giant cells (inset) and adenocarcinoma mixed subtype with BAC in the surrounding parenchyma. [H & E, ×12] The peritumoral areas of ground glass attenuation represent the BAC component (arrows) of this tumor.
E. Photomicrograph shows the adenocarcinoma component of the tumor. Note the cluster of tumor cells within the cystic space. The wall of the cystic space is also lined with adenocarcinoma tumor cells (arrows). [H & E, ×12]
the enhancement pattern of the mass on CT. A fiberoptic bronchoscopy and transbronchial washing cytology were performed, but the findings were inconclusive. The patient underwent a right lower lobectomy for both a definite diagnosis and treatment. The macroscopic examination showed a lobulated, yellowish mass with an eccentric cystic space. The microscopic examination revealed a pleomorphic carcinoma consisting of adenocarcinoma and giant cell carcinoma. Giant cell carcinoma was mainly located in the central portion of the mass, which was surrounded by adenocarcinoma with bronchioloalveolar carcinoma (BAC) features (Fig. 1D). The cystic space within the mass was lined by tumor cells, adenocarcinoma, and also contained some islets of adenocarcinoma components (Fig. 1E).

**Discussion**

Pulmonary pleomorphic carcinoma is a subtype of carcinoma with pleomorphic, sarcomatoid, or sarcomatous components. Histologically, it is defined as a non-small cell lung carcinoma with neoplastic spindle and/or giant cells, having at least a 10% spindle cell or giant cell component (4–6). Pleomorphic carcinoma is common in older men and is strongly associated with a history of smoking. It usually presents as a large, peripheral, necrotic mass with upper lobe predilection, invading the chest wall (3, 6).

Recently, two reports on the CT features as well as the radiological and pathological correlations of the tumors have been published. According to Kim et al. (7) in a recent study of 10 cases of pleomorphic carcinoma of the lungs, the tumors preferentially manifest as large peripheral masses with central low attenuation areas, frequently invading the pleura and chest wall. They concluded that low attenuation areas on contrast enhanced CT scans corresponding to areas of myxoid degeneration, necrosis, or hemorrhage on pathology.

In addition, Kim et al. (8) suggested that the CT features might be dominated by the epithelial component of the tumor after review of the CT and pathological findings of 30 cases of pleomorphic carcinoma. According to their study, pleomorphic carcinomas with adenocarcinoma or large cell components tend to be located in the peripheral zone. However, tumors with squamous cell components were located in the central zones. In their study, central necrosis was frequent but cavity formation in the tumor was uncommon for the pleomorphic carcinoma. The frequency of necrosis and tiny cavity formation was higher in the large and giant cell subtypes compared to others. Moreover, peritumoral areas of ground-glass attenuation were characteristics of the large and giant cell subtypes, and are associated with intra-alveolar macrophage aggregation and alveolar wall thickening with inflammatory cell infiltration and mild fibrosis on the pathology specimens.

In our case, the mass was initially thought to be a sclerosing hemangioma, because the mass was a well-defined contrast-enhancing juxtapleural mass in a young woman without a history of smoking. Moreover, the air crescent and CT halo signs, in our case, were characteristics consistent with sclerosing hemangioma, which were explained by repeated peritumoral bleeding with fibrosis and bronchial communication with the check valve phenomenon (9).

The radiology and pathology correlations, in our case, revealed that the air crescent represented the cystic changes associated with adenocarcinoma, without the evidence of communication with the airway. Moreover, the peripherally located bronchioloalveolar carcinoma was associated with ground glass attenuation. Kim et al. (8) reported that peritumoral ground glass attenuation was caused by inflammation. In our case, the ground glass attenuation, manifested by the bronchioloalveolar component of the pleomorphic carcinoma, was clearly demarcated with associated cystic changes; CT findings well known to be consistent with bronchioloalveolar carcinoma. Wang et al. (10) reported a case report of adenocarcinoma with an air crescent sign as observed in our case. They concluded that the emphysematous or cystic change between the tumor-infiltrating bands was caused by the infiltration of adenocarcinoma cells with the paracapitatural effect around the pulmonary parenchyma.

In summary, this unusual case of pleomorphic carcinoma with the air crescent and CT halo signs were associated with the adenocarcinoma and bronchioloalveolar carcinoma component of a pleomorphic carcinoma.

**References**

3. Chang YL, Lee YC, Shih JY, Wu CT. Pulmonary pleomorphic (spindle) cell carcinoma: peculiar clinicopathologic manifestations
different from ordinary non-small cell carcinoma. Lung Cancer 2001;34:91-97

Yeong Yi An, et al: Pleomorphic Carcinoma with the Air–Crescent and CT Halo Signs

Air Crescent Sign과 CT Halo Sign을 보인 폐 다형성 암종: 증례 보고

가톨릭대학교 의과대학 성균관병원 영상의학과
가톨릭대학교 의과대학 인천성모병원 영상의학과
가톨릭대학교 의과대학 성균관병원 병리과
가톨릭대학교 의과대학 성균관병원 흉부외과

안영이, 박현진, 김기준, 유진영, 조규도, 백준현

폐의 다형성 암종은 원발성 폐 종양의 0.1-0.4%를 차지하는 분화도가 좋지 않은 드문 종양이다. 이 종양은 대개 고령의 흡연 남성에서 크기가 큰 말초성 종괴로 나타나며, 진단 당시에 흉벽 침범이나 원격전이가 있는 경우가 많다. 저자들의 증례에서는 흡연력이 없는 수 개월간의 마른기침을 주소로 내원한 38세 여자환자에서 air crescent sign과 CT halo sign을 보인 다형성 암종 1예를 경험하였기에 문헌고찰과 함께 보고한다.