First report of pulmonary sclerosing pneumocytoma with malignant transformation in both cuboidal surface cells and stromal round cells: a case report

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Abstract

Background: Pulmonary sclerosing pneumocytoma (PSP) is a rare benign tumor. Although lymph node metastasis has been reported, it is still considered benign. No malignant transformation has been reported. This is the first case of malignant transformation of both cuboidal surface cells and stromal round cells.

Case presentation: A 64-year-old male had been complaining of intermittent hemoptysis several times per day for eight months. Chest computed tomography scan showed parenchymal infiltration with cystic lesion in the right lower lobe accompanied by enlarged right hilar lymph nodes. Lobectomy and systemic lymph node dissection was performed. On grossly pathological examination, the lesion was 50 mm from the bronchial stump. It was a mixture of both cystic and solid components and 30 mm × 20 mm in size with unclear border. Microscopically, the cuboidal surface cells transformed to adenocarcinoma. The stromal round cells also had a malignant transformation. The Ki-67 proliferation index in malignant cuboidal surface cells and stromal round cells were 70 and 55%, respectively. Furthermore, E-cadherin was negative in primary tumor but positive in metastatic lymph node, which suggested that the mesenchymal to epithelial transition may play an important role in lymph node metastasis.

Conclusions: To our knowledge, we present the first case of malignant transformation of both cuboidal surface cells and stromal round cells in PSP. The process of mesenchymal to epithelial transition may play an important role in lymph node metastasis.

Keywords: Pulmonary sclerosing pneumocytoma, Malignant transformation, Mesenchymal to epithelial transition, Stromal round cell, Cuboidal surface cell

Background

Pulmonary sclerosing pneumocytoma (PSP) is a rare benign tumor which has been described as sclerosing hemangioma [1]. It was previously considered as a vascular neoplasm, and now as a derivative from the primitive respiratory epithelium [2]. It is predominant in females, most commonly seen in middle aged females [3, 4]. Patients are always asymptomatic and computed tomography (CT) and X-ray of chest shows solitary, well circumscribed masses. The key pathological features of PSP involve two types of cells, cuboidal surface cells and stromal round cells, which are both neoplastic. Immunohistochemistry (IHC) studies show that thyroid transcription factor-1 (TTF-1) and epithelial membrane antigen (EMA) are both positive [2]. Pancytokeratin (CKpan) and Napsin A are both positive in cuboidal surface cells, while negative in stromal round cells [5]. Though lymph node metastasis has been reported, PSP is still considered benign [6, 7]. We report a unique case of PSP with malignant transformation in both cuboidal surface...
cells and stromal round cells, which has not been reported before.

Case presentation
A 64-year-old male had been complaining of intermittent hemoptysis several times per day for eight months. He had no fever, chest pain, shortness of breath, dizziness or amaurosis. He had no relevant medical history especially no history of cancer. He had no smoking history. The patient was admitted to The First Affiliated Hospital, School of Medicine, Zhejiang University due to symptoms getting worse. Chest computed tomography scan on July 1st, 2018 showed parenchymal infiltration with cystic lesion in the right lower lobe accompanied by enlarged right hilar lymph nodes (Fig. 1). Transbronchial lung biopsy under bronchofibroscopy was free of tumor cells. A primary surgical resection was recommended by surgeons. Lobectomy and systemic lymph node dissection was done on July 4th, 2018. The patient is now well after he recovered from surgery. So far there were no signs of tumor recurrence or metastasis.

Upon grossly pathological examination, the lesion was located in the right lower lobe, 50 mm from the bronchial stump. It was gray-tan to yellow on the section, with foci of hemorrhage. The lesion was a mixture of both cystic and solid components and was 30 mm *20 mm in size with unclear border. The solid component was in the middle of the lesion and was 17 mm*17 mm in size, surrounded by honeycomb cystic components.

Microscopically, the structure of the solid component of the tumor was similar to a typical PSP. It was composed of areas of cuboidal surface cells and stromal round cells. The tumor showed a hemorrhage pattern (Fig. 2). Bronchial adenomatous hyperplasia and cystic dilatation were noticed in surrounding areas. TTF-1 and EMA were positive in both cuboidal surface cells and stromal round cells (Fig. 2) while CKpan and Napsin A were only positive in cuboidal surface cells.

In the case reported, while most of the surface cells being similar to a typical PSP in some areas of the tumor, a few transformed to adenocarcinoma. The nuclei were columnar and containing hyperchromatic nuclear chromatin. In addition, the surface cells replaced the alveolar lining and invaded the fibrous stroma and vascular walls with TTF-1, EMA, Napsin A and CKpan all positive. The Ki-67 proliferation index was 70% (Fig. 3). We also noticed atypical adenomatous hyperplasia (AAH) of cuboidal cells in the transition area (Fig. 3). Cuboidal surface cells proliferated along preexisting alveolar walls with mild to moderate cellular atypia. A typical hobnail appearance was also seen in the atypical cuboidal surface cells. Substantial gaps along the surface of basement membrane in the transition area were also evident of AAH.

A few stromal round cells had small, well-defined borders and central bland nuclei without nucleoli similar to that in a typical PSP. However, mild to moderate atypical stromal round cells proliferation was seen in the transition region (Fig. 4). Binuclearization and intranuclear eosinophilic inclusions were common in the transition area in our case. Furthermore, abundant cytoplasm, nuclear polymorphism, prominent nucleoli and irregular mitosis were observed in malignant stromal round cells, adjoining the transition areas (Fig. 4). Vascular invasion

Fig. 1 Chest computed tomography scan showed that parenchymal infiltrate with cystic lesion in the right lower lobe of lung
Fig. 2 (a, b) Tumor nodule showed a typical pulmonary sclerosing pneumocytoma of hemorrhage growth pattern comprising of large blood-filled spaces lined by surface cells (h&i). (c, d) Cuboidal surface cell were positive for pancytokeratin (CKpan), round cells were negative for CKpan. (e, f) Cuboidal surface cell were positive for Napsin A, round cells were negative. (g, h) Both cuboidal surface and stromal round cells were positive for thyroid transcription factor-1 (TTF-1). (i, j) Both cuboidal surface and stromal round cells were positive for epithelial membrane antigen (EMA).
and pulmonary parenchyma involvements were also found in malignant lamellarlike stromal round cells. TTF-1, P63 and EMA were all positive. Only a small amount stromal round cells were positive for CKpan. However, stromal round cells were negative for beta-catenin and E-cadherin. The Ki-67 proliferation index in these areas was 55%, which was significantly increased compared to typical PSP areas (Fig. 4). Both stromal round cells and surface cells were negative for Progesterone receptor, CD20, CD3, S-100, Melana, HMB45, Myogenin, MyoD1, CgA and Syn. Further molecular investigation using a polymerase chain reaction panel showed that no EGFR, ALK or ROS1 mutation was detected.

In this case, we also found mediastinal lymph nodes involvement. The architecture of lymph nodes was replaced...
by abnormal proliferated stromal round cells with either vacuolated or eosinophilic cytoplasm (Fig. 5). IHC showed that these cells were positive for TTF-1, partial positive for CKpan and E-cadherin, but negative for beta-catenin. However, the E-cadherin was negative in malignant stromal round cells in the primary tumor (Fig. 4).

**Discussion and conclusion**

PSP is considered as a rare benign tumor [1]. In searches of PubMed and Embase database, there are 24 cases of PSP with lymph node metastasis and recurrence (Table 1) [2, 6–25]. Five cases have mediastinal lymph node metastasis, four have distant metastasis, and only one have recurrence of PSP. However, no case about malignant transformation of PSP has been reported. One case reports overgrown stromal round cells and bone metastasis, accompanied by increased cellularity and necrotic areas, but a Ki-67 index of less than 5% [22]. Another case reports PSP with metastatic spread to stomach with Ki-67 indeice in primary tumor and metastatic gastric lesion of 17.6 and 19.4%, respectively [19]. The, however, no malignant pathomorphological change has been reported. In lyoda’s research, cases with recurrence has a Ki-67 index of 0.4% [26]. These results show no significantly increased proliferation of cells even in patients with recurrence or metastasis. In our case, the Ki-67 proliferation index of the malignant cuboidal surface cells and the stromal round cells are 70 and 55%, respectively. The high proliferative activity and pathomorphological change in both cuboidal surface cells and stromal round cells suggest that PSP transformed to a malignant tumor. Liu reports a case of coexistence of PSP and primary adenocarcinoma in the same tumor [27], which is different from our...
In our case, the two well-established epithelial markers, E-cadherin and CKpan [28], are both positive in the metastatic lymph nodes with similar levels (black arrows in Fig. 5), although they should be negative in metastatic lymph nodes (composed of stromal round cells). The epithelial marker expression in metastatic lymph nodes suggests the mesenchymal-epithelial transition (MET) during lymph node metastasis. Previous studies show that MET process is able to promote distal metastasis in breast cancer [29], especially for establishing macrometastasis [30–32], which, combined with our results, suggests that the MET process may play an important role in lymph node metastasis of PSP. Although based on previous studies, lymph node involvement doesn’t affect long-term survival rate [6, 23], patients with malignant PSP may still need close follow-up.

In summary, we report the first case of malignant transformation in both cuboidal surface cells and stromal round cells, which suggests the malignant potential of PSP. The fact that E-cadherin is negative in primary tumor but positive in metastatic lymph nodes suggests that the process of MET plays an important role in lymph node metastasis of PSP.

**Abbreviations**

AAH: Typical adenomatous hyperplasia; CKpan: Pancytokeratin; CT: Computed tomography; EMA: Epithelial membrane antigen; IHC: Immunohistochemistry; MET: Mesenchymal-epithelial transition; PSP: Pulmonary sclerosing pneumocytoma; TTF-1: Thyroid transcription factor-1

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**Authors’ contributions**
TX collected the data, reviewed the literature, drafted and edited the manuscript. TXD conceived the study, participated in experiment design and data acquisition and edited the manuscript. All authors read and approved the final manuscript.
Available data and materials
The datasets used in this study are available from the corresponding author on reasonable requests.

Ethics approval and consent to participate
The study received ethics approval from the Commission for Scientific Research in the First Affiliated Hospital, School of Medicine, Zhejiang University. The patient provided written informed consent.

Consent for publication
Written informed consent was obtained from the patient and his healthy sibling for the publication.

Competing interests
The authors declare that they have no competing interests.

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