Ciliated muconodular papillary tumor of the lung presenting with polymyalgia rheumatica-like symptoms: a case report

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Abstract: Ciliated muconodular papillary tumors (CMPTs) of the lung have been recently characterized as low-grade malignant tumors and may be indistinguishable from adenocarcinoma in situ (AIS) because they are both abundant in mucous and spread along the alveolar walls. Herein, we report a case of CMPT with polymyalgia rheumatica (PMR)-like symptoms, which resolved after resection. After the surgery, antinuclear antibody tests were performed, but no abnormalities were noted. Furthermore, the lung tumor could not be distinguished from AIS, as revealed by a pathological examination. This case demonstrates two key points: the paraneoplastic symptoms of CMPT can indicate PMR, and it is difficult to diagnose peripheral lung tumors as CMPT unless there is a completely resected specimen available. The possibility exists that an increased number of older patients will be diagnosed with CMPT because of the increasing frequency of computed tomography performed in this population. Therefore, it is important for clinicians to obtain completely resected specimens to ensure accurate diagnosis and management of CMPT.

Keywords: Adenocarcinoma in situ (AIS); ciliated muconodular papillary tumor (CMPT); differential diagnosis; paraneoplastic syndromes

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Introduction

Ciliated muconodular papillary tumors (CMPTs) of the lung have been recently characterized as low-grade malignant tumors. It is difficult to distinguish CMPT from adenocarcinoma in situ (AIS) (1-3). Polymyalgia rheumatica (PMR)-like symptoms associated with lung cancer (4,5). CMPT is thought to be associated with benign tumors; however, we herein report a case of CMPT with PMR-like symptoms that resolved after resection, which was indistinguishable from AIS before pulmonary resection.

Case presentation

A 78-year-old man with a history of diabetes mellitus and hypertension had experienced bilateral pain and stiffness in his proximal thigh muscle, which occurred throughout the day, starting first thing in the morning; however, he was not hospitalized. He was referred to our hospital because of a lung tumor, which was identified on computed tomography (CT). Laboratory investigations showed a carcinoembryonic antigen level of 7.5 ng/mL and a C-reactive protein level of 0.4 mg/dL. CT indicated the presence of a lung nodule, 1.9 cm in diameter, with a cavity in the right S3 segment (Figure 1). The nodule had a standardized uptake value of 3.56 on 18F-fluorodeoxyglucose positron emission tomographic imaging. No other nodules were identified. Based on these findings, lung cancer was suspected. However, a diagnosis could not be made based on bronchoscopic biopsy findings, and, therefore, surgical biopsy was performed. After the patient’s health was assessed, thoracoscopic surgery was performed utilizing four ports to biopsy the tumor.
Intraoperative rapid diagnosis with a frozen section of the tumor indicated AIS; hence, we performed right upper lobectomy with lymph node dissection. After the surgery, we consulted another hospital, because histological examination of the resected specimen showed a low nuclear grade, and the tumor did not resemble AIS. The tumor consisted of a mixture of ciliated columnar, mucous, and basal cells in glandular and papillary growth patterns, and basal cells showed positive p40 staining. These features were consistent with those of CMPT; thus, the patient was diagnosed accordingly (Figure 2). After the surgery, his bilateral pain and stiffness in the proximal thigh muscle were relieved. Symptom relief was maintained despite reducing the dose of analgesics. We performed antinuclear antibody tests; however, no abnormality was noted. Thirty months after the surgery, the patient did not have any symptoms and had no tumor recurrence. Nonetheless, he is being monitored carefully because the etiology of the tumor is not understood clearly.

**Discussion**

This case demonstrates two key points: the paraneoplastic symptoms of CMPT can indicate PMR, and it is difficult to diagnose peripheral lung tumor cases as CMPT without obtaining a completely resected specimen.
The diagnosis of PMR is based on the Provisional Classification Criteria for Polymyalgia Rheumatica (Table 1) (6). The current patient had experienced bilateral pain and stiffness in his proximal thigh muscle, which started upon waking up every morning. Laboratory investigations just before the operation showed absence of anti-citrullinated protein antibodies and other antinuclear antibodies. Thus, it was revealed that he had PMR, because he met two criteria and his score was 4. However, after the surgery, his symptoms disappeared. He is being monitored carefully, and his symptoms have not returned.

Secondly, AIS is a precancerous lesion. Moreover, it is difficult to distinguish CMPT from AIS, because they are both abundant in the mucous cells and spread along the alveolar walls. In a previous study, the authors reported that they could judge that the tumor was a low-grade malignant tumor using frozen sections (7). However, we could not determine that the tumor was not AIS. After the surgery, we consulted with another hospital, and two more features were identified consistent with CMPT. First, there were no atypical cells. Second, there were mixed structures constructed with ciliated cells, mucous cells, and basal cells positive for p40. The differential diagnosis for CMPT and AIS may become easier with the accumulation of data from more cases; however, currently, it is difficult to investigate in detail without observing viable pathological specimens after the operation.

In summary, CMPT can present with paraneoplastic symptoms that indicate PMR; therefore, it is difficult to diagnose peripheral lung tumors as CMPT without obtaining a completely resected specimen. Hence, it is important to resect the tumor completely.

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Footnote

Conflicts of Interest: This paper was presented at the 57th Annual Meeting of the Japan Lung Cancer Society at Fukuoka on December 20, 2016.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of written consent is available for review by the Editor-in-Chief of this journal.

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