MINIREVIEWS

6974 Applications of time series analysis in epidemiology: Literature review and our experience during COVID-19 pandemic
Tomov L, Chervenkov L, Miteva DG, Batselova H, Velikova T

ORIGINAL ARTICLE

6984 Retrospective Cohort Study
Acute cholangitis: Does malignant biliary obstruction vs choledocholithiasis etiology change the clinical presentation and outcomes?
Tsou YK, Su YT, Lin CH, Liu NJ

Retrospective Study

6995 Usefulness of analyzing endoscopic features in identifying the colorectal serrated sessile lesions with and without dysplasia

7004 Roles of biochemistry data, lifestyle, and inflammation in identifying abnormal renal function in old Chinese
Chen CH, Wang CK, Wang CY, Chang CF, Chu TW

7017 Clinical efficacy and safety of Guipi decoction combined with escitalopram oxalate tablets in patients with depression
Yu J, Xu FQ

7026 Artificial intelligence technology and ultrasound-guided nerve block for analgesia in total knee arthroplasty
Tong SX, Li RS, Wang D, Xie XM, Ruan Y, Huang L

7034 Axenfeld-Reiger syndrome: A search for the missing links
Morya AK, Ramesh PV, Sinha S, Nishant P, Nain N, Ramavath RN, Gone C, Prasad R

Observational Study

7043 Self-management of osteoarthritis while waiting for total knee arthroplasty during the COVID-19 pandemic among older Malaysians
Mahdzir ANK, Mat S, Seow SR, Abdul Rani R, Che Hasan MK, Mohamad Yahaya NH

7053 “In situ bone flap” combined with vascular pedicled mucous flap to reconstruction of skull base defect

7061 Reference values of gait parameters in healthy Chinese university students: A cross-sectional observational study
### Contents

**World Journal of Clinical Cases**  
**Thrice Monthly Volume 11 Number 29 October 16, 2023**

<table>
<thead>
<tr>
<th>Page</th>
<th>Title</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>7075</td>
<td>Effect of T-regulatory cells and interleukin-35, interleukin-10, and transforming growth factor-beta on diffuse large B-cell lymphoma</td>
<td>Wu H, Sun HC, Ouyang GF</td>
</tr>
<tr>
<td>7091</td>
<td>Meta-analysis of the efficacy and safety of daratumumab in the treatment of multiple myeloma</td>
<td>Wang P, Jin SY</td>
</tr>
<tr>
<td></td>
<td><strong>CASE REPORT</strong></td>
<td></td>
</tr>
<tr>
<td>7101</td>
<td>Varicella-zoster virus meningitis with hypoglycorrhachia: A case report</td>
<td>Cao LJ, Zheng YM, Li F, Hao HJ, Gao F</td>
</tr>
<tr>
<td>7107</td>
<td>Unusual presentation of penile giant condyloma acuminatum with spontaneous prepuce perforation: A case report</td>
<td>Hsu FC, Yu DS, Pu TW, Wu MJ, Meng E</td>
</tr>
<tr>
<td>7113</td>
<td>Primary renal lymphoma presenting as renal failure: A case report and review of literature from 1989</td>
<td>Lee SB, Yoon YM, Hong R</td>
</tr>
<tr>
<td>7136</td>
<td>Mucoepidermoid carcinoma of the lung with hemoptysis as initial symptom: A case report</td>
<td>Xie WX, Liu R, Li Z, Zhou PL, Duan LN, Fu DD</td>
</tr>
<tr>
<td>7144</td>
<td>Co-infection of <em>Chlamydia psittaci</em> and <em>Tropheryma whippelii</em>: A case report</td>
<td>Du ZM, Chen P</td>
</tr>
<tr>
<td>7150</td>
<td>Surgical treatment of severe anterior capsular organized hard core cataract: A case report</td>
<td>Wang LW, Fang SF</td>
</tr>
<tr>
<td>7156</td>
<td>First platelet transfusion refractoriness in a patient with acute myelocytic leukemia: A case report</td>
<td>Tu SK, Fan HJ, Shi ZW, Li XL, Li M, Song K</td>
</tr>
<tr>
<td>7162</td>
<td>Rare finding of primary aortoduodenal fistula on single-photon emission computed tomography/computed tomography of gastrointestinal bleeding: A case report</td>
<td>Kuo CL, Chen CF, Su WK, Yang RH, Chang YH</td>
</tr>
<tr>
<td>7170</td>
<td>Rituximab combined with Bruton tyrosine kinase inhibitor to treat elderly diffuse large B-cell lymphoma patients: Two case reports</td>
<td>Zhang CJ, Zhao ML</td>
</tr>
</tbody>
</table>
## Contents

**Thrice Monthly Volume 11 Number 29 October 16, 2023**

<table>
<thead>
<tr>
<th>Page</th>
<th>Title</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>7179</td>
<td>Use of Ilizarov technique for bilateral knees flexion contracture in Juvenile-onset ankylosing spondylitis: A case report</td>
<td>Xia LW, Xu C, Huang JH</td>
</tr>
<tr>
<td>7187</td>
<td>Case of takotsubo cardiomyopathy after surgical treatment of liver hydatid cyst: A case report</td>
<td>Altas Y, Abdullayeva Ü</td>
</tr>
<tr>
<td>7193</td>
<td>Laparoscopic choledocholithotomy and transductal T-tube insertion with indocyanine green fluorescence imaging and laparoscopic ultrasound: A case report</td>
<td>Yoo D</td>
</tr>
<tr>
<td>7200</td>
<td>Hematopoietic stem cell transplantation of aplastic anemia by relative with mutations and normal telomere length: A case report</td>
<td>Yan J, Jin T, Wang L</td>
</tr>
<tr>
<td>7207</td>
<td>Emphysematous thrombophlebitis caused by a misplaced central venous catheter: A case report</td>
<td>Chen N, Chen JJ, Chen T, Zhang W, Fu XY, Xing ZX</td>
</tr>
<tr>
<td>7214</td>
<td>Aggressive angiomyxoma of the epididymis: A case report</td>
<td>Liu XI, Su JH, Fu QZ, Liu Y</td>
</tr>
<tr>
<td>7221</td>
<td>Gastric and intestinal ectopic pancreas: Two case reports</td>
<td>Zhang H, Zhao HY, Zhang FH, Liang W</td>
</tr>
<tr>
<td>7234</td>
<td>Imaging misdiagnosis and clinical analysis of significant hepatic atrophy after bilioenteric anastomosis: A case report</td>
<td>Liang SY, Lu JG, Wang ZD</td>
</tr>
<tr>
<td>7248</td>
<td>Simultaneous thyroglossal duct cyst with parathyroid cyst: A case report</td>
<td>Chen GY, Li T</td>
</tr>
<tr>
<td>7253</td>
<td>Submandibular solid-cystic mass as the first and sole manifestation of occult thyroid papillary carcinoma: A case report</td>
<td>Chen GY, Li T</td>
</tr>
</tbody>
</table>

**LETTER TO THE EDITOR**

<table>
<thead>
<tr>
<th>Page</th>
<th>Title</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>7258</td>
<td>Artificial intelligence and machine learning in motor recovery: A rehabilitation medicine perspective</td>
<td>Swarnakar R, Yadav SL</td>
</tr>
</tbody>
</table>
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RESPONSIBLE EDITORS FOR THIS ISSUE
Production Editor: Hua-Ge Yu; Production Department Director: Xiang Li; Editorial Office Director: Jin-Lei Wang.

NAME OF JOURNAL
World Journal of Clinical Cases

ISSN
ISSN 2307-8960 (online)

LAUNCH DATE
April 16, 2013

FREQUENCY
Thrice Monthly

EDITORS-IN-CHIEF
Bao-Gan Peng, Jerzy Tadeusz Chudek, George Kontogeorgos, Maurizio Serati, Ja Hyeon Ku

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https://www.wjgnet.com/2307-8960/editorialboard.htm

PUBLICATION DATE
October 16, 2023

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ONLINE SUBMISSION
https://www.f6publishing.com

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E-mail: bpgoffice@wjgnet.com https://www.wjgnet.com
Mucoepidermoid carcinoma of the lung with hemoptysis as initial symptom: A case report

Wen-Xing Xie, Rong Liu, Zheng Li, Pei-Ling Zhou, Li-Na Duan, Dan-Dan Fu

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Abstract

BACKGROUND
Mucoepidermoid carcinoma of the lung is a rare malignant tumor, accounting for 0.1%–0.2% of all lung malignancies. It is a primary salivary gland tumor of the lung. Surgical resection is the primary treatment for pulmonary mucoepidermoid carcinoma, for which there has been no standardized treatment strategy. This article reports a case of a young woman with pulmonary mucoepidermoid carcinoma with hemoptysis as the first symptom.

CASE SUMMARY
A 24-year-old female patient presented with "4 d of hemoptysis" as the chief complaint. She had no special history and denied any smoking or drinking history. Physical examination revealed that the vital signs were stable and scattered small wet rales were heard in the left lung. After admission, the lung tumor markers were checked, and no abnormalities were found. After completing the bronchoscopy, a spherical lesion was observed at the main bronchus 1.5 cm away from the protubercle, with obvious pulsation and little blood seepage on the surface, and histopathological biopsy results showed acute and chronic inflammation. She was transferred to the Department of Thoracic Surgery for surgical treatment on the 16th day after admission. After exclusion of surgical conjunctures, the patient underwent resection of the tumor in the left main bronchus with single-pore video-assisted thoracic surgery on the 19th day after admission. The postoperative histopathological biopsy results showed mucoepidermoid carcinoma of the lung. The patient and her family refused to complete genetic testing and she was discharged from the hospital on the 8th day after surgery. During the follow-up period, the patient experienced shortness of breath after feeling active and had no special discomfort.

CONCLUSION
We have documented a case of moderately differentiated mucoepidermoid lung
cancer with hemoptysis as the first symptom to improve clinicians' understanding of the disease and provide a new dimension of thinking for its future diagnosis and treatment.

**Key Words:** Mucoepidermoid carcinoma of the lung; Malignant tumor; Bronchoscopy; Surgical treatment; Case report

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**Core Tip:** Mucoepidermoid carcinoma of the lung is a rare tumor. This paper reports a young female patient who was diagnosed with mucoepidermoid carcinoma of the lung with hemoptysis as the first symptom to improve the clinician's understanding of the disease and to help its clinical diagnosis and treatment.

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**Citation:** Xie WX, Liu R, Li Z, Zhou PL, Duan LN, Fu DD. Mucoepidermoid carcinoma of the lung with hemoptysis as initial symptom: A case report. *World J Clin Cases* 2023; 11(29): 7136-7143

**URL:** https://www.wjgnet.com/2307-8960/full/v11/i29/7136.htm

**DOI:** https://dx.doi.org/10.12998/wjcc.v11.i29.7136

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**INTRODUCTION**

Pulmonary mucoepidermoid carcinoma (PMEC) was first reported by Smetana in 1952 as a rare malignant tumor of the lung[1]. It is the most common primary salivary gland carcinoma of the lung, originating from the small salivary gland in the submucosa of the atmospheric tract[2]. At present, there are few reports on PMEC, and most are case reports. Its clinical symptoms and epidemiological characteristics are not specific or typical. In addition, there is no specific treatment, and surgical excision remains the main treatment. Therefore, it is necessary to increase the attention of clinicians to the disease. This paper reports the case data of a young woman with hemoptysis as the first symptom who was clearly diagnosed with the disease after two lung histopathological biopsies and was discharged from the hospital after surgical treatment, aiming to provide some help for the diagnosis and treatment of the disease and improve the awareness of clinicians about it.

---

**CASE PRESENTATION**

**Chief complaints**
A 24-year-old female patient presented with "4 d of hemoptysis" as the chief complaint.

**History of present illness**
Bright red blood appeared 4 d prior, the amount was approximately 20 mL/d, accompanied by cough and sputum, and the sputum was a small amount of frothy sputum.

**History of past illness**
The patient had a good health previously and had no history of specific diseases.

**Personal and family history**
The patient's personal history and family history were unremarkable.

**Physical examination**
The patient's vital signs were stable, and she had a clear mind and cooperated with the physical examination. The skin mucosa color was normal, and the superficial lymph nodes of the whole body were not enlarged. Bilateral respiratory movement was normal, palpation of both lungs was normal, percussive sound of both lungs was clear, small wet rales were heard in the left lung, and pleural fricative sounds were not heard on either side. No abnormal positive signs were found in the remaining physical examination.

**Laboratory examinations**
D-dimer was 1.17 µg/mL, and hypersensitivity C-reactive protein was 32.88 mg/L. There were no obvious abnormalities in lung tumor markers, routine blood tests, coagulation function, liver function, renal function, or *Mycobacterium tuberculosis* culture.

**Imaging examinations**
Chest computed tomography (CT) revealed left lung emphysema, atelectasis of part of the left lung, and part of the...
bronchus unclear (Figure 1).

Further diagnostic work-up
After admission, the patient was treated by hemostasis and anti-infection therapy, and no hemoptysis occurred. Contrast-enhanced CT of the chest showed that a left main bronchial nodule was seen (Figure 1). Increased soft tissue density shadows appeared in the left hilar area of the lung, and fiberoptic bronchoscopy was recommended. Left pulmonary obstructive emphysema and obstructive atelectasis and obstructive inflammation of the left lung were noted. Further electronic bronchoscopy showed that the main bronchus had a spherical lesion 1.5 cm from the protuberance, with pulsation and slight bleeding on the surface. The considerations were as follows: (1) The lesion in the left main bronchus was suspected to be a tumor; and (2) malignant stenosis of the left main bronchus (Figure 2) was present. The lung tumor markers improved and showed no abnormalities. Cytopathology and DNA ploidy analysis of alveolar lavage fluid (left main bronchoalveolar lavage fluid) showed no definite malignant cells. The results of bronchial tissue biopsy showed that the mucosa presented acute and chronic inflammatory changes, mucinous gland hyperplasia in the lamina mucosa, squamation of the columnar epithelium, and active proliferation of epithelial cells. Some necrotic tissues were also observed, among which small clusters of proliferative glandular epithelial cells were found, the cytoplasm was vacuolar, nucleoli were visible, and no mitotic image was observed. The immunohistochemical results were as follows: Carcinoembryonic antigen (CEA) (+), cytokeratin (CK) (+), CK5/6 (partially +), CK7 (+), P40 (-), TTF1 (-), and NapsinA (-). The automatic immunohistochemical results were as follows: CD117 (-) and Ki67 (5% +). Antacid staining was negative, while periodic acid-Schiff (PAS) was positive (Figure 3).

Based on the immunohistochemistry findings, the patient was transferred to the Department of Thoracic Surgery for surgical treatment on the 16th day after admission. After the exclusion of surgical contraindications, the patient underwent resection of the tumor in the left main bronchus by single-pore video-assisted thoracic surgery on the 19th day of admission (thoracotomy of the left main bronchus and trachea window for tumor resection and anastomosis). Rapid intraoperative pathological examination revealed neoplastic lesions (in the left main bronchus), which were likely to be malignant tumors from the salivating glands. Postoperative histopathological results showed the following: Lung mucoepidermoid carcinoma (in the left main bronchus), without vascular invasion or nerve invasion; group 4 lymph nodes (0/3), group 5 lymph nodes (0/1), group 10 lymph nodes (0/2), and no tumor metastasis; pTNM stage pT1bN0M0. Immunohistochemical results were: CK (+), CK5/6 (partially +), P40 (partially +), P63 (partially +), CK7 (+), CK-H (+), TTF1 (+), NapsinA (+), and CEA (locally +). The automatic immunohistochemical results included CD117 (partially +) and Ki67 (10% +). PAS staining was positive (Figure 4). Head MRI plain + enhanced scans and upper abdominal CT plain + enhanced scans showed no obvious abnormalities. After the operation, the patient was treated with anti-infection, hemostasis, cough suppressant and expectorant, anti-inflammatory treatment, and lung function recovery exercises. After repeat chest CT (Figure 5), the patient was discharged from the hospital on the 8th day after surgery.

MULTIDISCIPLINARY EXPERT CONSULTATION
After considering the pathological results of the patient's two tissue biopsies and imaging data, the pathologist believed that the lesion was a moderately differentiated pulmonary epidermal mucoid carcinoma, and the relevant genetic examination can be further performed to guide subsequent treatment. The radiologist believed that the imaging data of the patient in this case were not consistent with typical adenocarcinoma manifestations, and considering the rarity of the disease, the diagnosis was mainly based on tissue biopsy pathological nodes. The thoracic surgeon considered the patient to be a young woman, and the mass was removed locally. The lung tissue was preserved as much as possible, and close follow-up was performed to determine whether there was metastasis or recurrence. The respiratory surgeon believed that the patient should have a complete bronchoscopy at regular intervals after discharge to clarify the bronchial cavity, and reoperation could have been considered if necessary.

FINAL DIAGNOSIS
The final diagnosis of the presented case was PMEC.

TREATMENT
After surgical treatment, anti-infection, hemostasis, cough suppressant and expectorant; anti-inflammatory treatments; lung function recovery exercises and other treatments were given, the patient improved and was discharged on the 8th day after surgery.

OUTCOME AND FOLLOW-UP
After surgical treatment, the patient was treated with anti-infection, hemostasis, cough suppressant and expectorant, anti-
inflammatory treatments, lung function recovery exercises, etc., the patient improved and was discharged from the hospital on the 8th day after surgery. During the follow-up period, the patient experienced shortness of breath after feeling active and no special discomfort.

**DISCUSSION**

As the most commonly principal salivary gland cancer in the lungs, PMEC originates from the respiratory tract[2]. As a rare malignant lung tumor, it was first reported by Smetana et al[1] as early as 1952, and it was found that it accounted for...
less than 1% of malignant lung tumors\cite{3-5}. Due to its rarity and lack of specific clinical and radiological features, the diagnosis is largely dependent on pathological examination. In addition, PMEC should be distinguished from adenosquamous carcinoma, especially in tiny biopsy specimens obtained by fibrobronchoscopy or lung puncture. It is defined as a tumor composed of mucoepidermoid cells, epidermoid cells, and intermediate cells\cite{6}. At present, there is no unified standard for treatment, and surgical resection is considered the main treatment\cite{7}.

As the most general malignant tumor in the salivary glands, mucoepidermoid carcinoma has an incidence of 0.44/100000 people\cite{8}. Although it can be appeared at every age, it occurs mostly between the ages of 35 and 65, with approximately 60% of cases occurring in women. The large salivary glands are frequently involved and very rarely (less than 1%) it occurs in the lung\cite{9}. Patients with PMEC often have no specific clinical symptoms. Obstructive airway symptoms, mainly cough, dyspnea, or asthma, are usually manifested in tumors located in the central bronchus; 85% of PMECs are
reported to be peripheral to the lung, characterized by cough, chest pain, and lung inflammation. Some asymptomatic patients are found during physical examination[7]. Therefore, there are challenges in the diagnosis of this disease. It is therefore necessary to improve the understanding of clinicians to achieve early detection and early and accurate treatment.

In this case, hemoptysis was the first symptom, and chronic inflammatory changes were considered after the first bronchial histopathologic biopsy. However, the immunohistochemical results suggested CEA (locally +), CK (+), CK5/6 (partially +), and Ki67 (5% +), and PAS staining was positive. Considering the abnormal immunohistochemistry findings, a tumor could not be excluded. PMEC was clearly diagnosed after surgical resection and pathological biopsy. Immunohistochemistry plays an important role in the diagnosis of this disease.

The diagnosis of PMEC mainly depends on histopathological and immunohistochemical examination, and it is histopathologically composed of squamous epithelial cells, mucous cells, and intermediate cells with keratosis defects. PMEC is classified into low-grade and high-grade tumors based on histological features, mitotic frequency, cell atypia, and degree of necrosis[5]. The survival rate of high-grade mucoepidermoid carcinoma was significantly lower than that of low-grade mucoepidermoid carcinoma, and the possibility of metastasis to lymph nodes of high-grade mucoepidermoid carcinoma was found to be 10 times higher than that of low-grade mucoepidermoid carcinoma[10]. In general, PMEC tumors appear as tan or light brown polyloid masses. The central bronchus may exhibit exoplastic tumors that almost completely obstruct the bronchial lumen[11]. The immunohistochemical characteristics of PMEC were retrospectively analyzed and summarized, and the positive rates of P63, CK7, MUC5AC, P40, and CK5/6 were 58/58 (100%), 33/33 (100%), 26/26 (100%), 52/54 (96.3%), and 3/6 (50%), respectively[7]. However, some studies do not support this conclusion. Zhang et al[12] reported that TTF-1 and Napsin A were positive in some PMEC cases, and one paper reported that trastuzumab treatment was effective in metastatic PMEC patients with positive HER2 expression[13]. In addition, some scholars proposed that the Ki-67 index in low-grade cases was lower than that in high-grade cases and proposed that the Ki-67 index might be used as one of the indicators to distinguish PMEC malignancy[14]. At present, most studies are mainly case reports or case series and there are little case data, so the number of studies needs to be expanded for further research.

Studies have shown that the tumor is associated with the t(11;19)(q21-22;p13) translocation, which is associated with the MECT1-MAML2 fusion[15], and some scholars have found that this fusion gene can not only activate HES1 transcription, thus destroying the Notch signaling pathway, but it also activates the protein CREB, thus simulating the activation of cyclic adenosine monophosphate signaling[16,17]. Some scholars proposed that gene fusion could be used as a diagnostic basis for PMEC because it was found to exist in most patients with this disease[7]. In addition, some studies found that gene fusion was often found in low-grade groups[18,19]. At present, most of the genes related to this disease are limited to fusion genes. There are also studies using comprehensive genome amplification to study a small number of high-grade PMECs, and it was found that most patients have at least one gene mutation, and the most common genomic changes occur in CDKN2A and TP53. However, the reliability of this study remains questionable due to the small sample size[20].

More interestingly, EGFR overexpression is present in most PMEC cases, but amplification or mutation of the tyrosine kinase region of the EGFR gene is rarely reported[18]. At present, there is no unified standard for the treatment of PMEC, and the main treatment is surgical excision. The effect of chemoradiotherapy is controversial. There have also been reports of cases effectively treated by chemotherapy, such as apatinib combined with graded stereotactic radiotherapy[14], carboplatin combined with paclitaxel[21], and EGFR-tyrosine kinase inhibitor drug therapy[22]. However, there are individual differences due to the majority of the literature being case reports, and further research is required to define the specific curative effect. Studies show that PMEC is a fairly inert tumor with a relatively optimistic prognosis, which has a better survival rate compared with small cell lung cancer and non-small cell lung cancer. The 5-year survival rate of PMEC is approximately 45%-70%[5]. Mucoepidermoid carcinoma usually involves the proximal bronchus. Therefore, the typical symptoms of mucoepidermoid carcinoma are bronchial obstruction, resulting in such as cough, hemoptysis, asthma, and fever[23]. Salivary gland tumors are rare primary lung lesions. The most common primary salivary gland tumors in the lung mainly include mucoepidermoid carcinoma, adenoid cystic carcinoma, and epithelial-myoepithelial carcinoma. Their morphology, immunophenotype, and molecular characteristics are similar to those in the head and neck or other sites. Because of their rarity, research is often limited, and relevant studies are usually small or limited to individual cases. Fortunately, molecular changes such as MAML2 rearrangement in mucoepidermoid carcinoma of the lung, MYB rearrangement in adenoid cystic carcinoma and clear cell carcinoma, and EWSR1 rearrangement in myoepithelial tumors have been found[2]. These molecular changes help to distinguish salivary gland tumors from other lung tumors to a certain extent and will provide great help for the diagnosis and treatment of this disease.

PMEC is a rare malignant tumor with no specific clinical symptoms. The diagnosis of PMEC mainly relies on pathological and immunohistochemical examination. The diagnosis of the patient in this case was confirmed by two biopsies. Reporting this case is expected to improve the diagnosis rate of this disease to achieve early treatment.

CONCLUSION

PMEC is a rare malignant tumor, and its clinical symptoms often have no specific manifestations. Diagnosis of the disease mainly relies on pathological and immunohistochemical examination. In this case, two tissue biopsies were performed to confirm the disease. When there is unexplained hemoptysis in a young patient, the disease should be considered, and diagnostic work-up should be carried out as soon as possible to confirm the diagnosis.
ACKNOWLEDGEMENTS

First of all, I would like to thank the thoracic surgeons and endoscopy doctors for their help in the treatment of the patient. I would also like to thank the pathologists and imaging department for their relevant information. Last but not least, I would like to thank the patient for her trust in our hospital.

FOOTNOTES

Author contributions: Xie WX and Liu R contributed to manuscript writing and editing and data collection; Li Z and Zhou PL contributed to data analysis; Duan LN and Fu DD contributed to conceptualization and supervision; all authors have read and approved the final manuscript.

Informed consent statement: Informed written consent was obtained from the patient for publication of this report and any accompanying images.

Conflict-of-interest statement: All the authors declare that they have no conflict of interest to disclose.

CARE Checklist (2016) statement: The authors have read the CARE Checklist (2016), and the manuscript was prepared and revised according to the CARE Checklist (2016).

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