Catamenial pneumothorax with bubbling up on the diaphragmatic defects: a case report

Bo Dong, Chun-Li Wu, Yin-liang Sheng, Bin Wu, Guan-Chao Ye, Ya-Fei Liu, Shi-Hao Li, Lu Han and Yu Qi*

Abstract
Background: Catamenial pneumothorax is characterized by spontaneous recurring pneumothorax during menstruation, which is a common clinical manifestation of thoracic endometriosis syndrome. There are still controversies about its pathogenesis.

Case presentation: A 43-year-old woman with a history of endometriosis came to our hospital due to recurring pneumothorax during menstruation. Uniportal Video-assisted Thoracoscopic Surgery (VATS) exploration was performed on the eve of menstruating. We thoroughly explored the diaphragm, visceral and parietal pleura: The lung surface was scattered with yellowish-brown implants; no bullae were found; multiple diaphragmatic defects were found on the dome. And surprisingly, we caught a fascinating phenomenon: Bubbles were slipping into pleural cavity through diaphragmatic defects. We excised the diaphragmatic lesions and wedge resected the right upper lung lesion; cleared the deposits and flushed the thoracic cavity with pure iodophor. Diaphragmatic lesions confirmed the presence of endometriosis, and interestingly enough, microscopically, endometrial cells were shedding with impending menses. After a series of intraoperative operations and postoperative endocrine therapy, the disease did not recur after a period of follow-up.

Conclusion: We have witnessed the typical signs of catamenial pneumothorax at the accurate timing: Not only observed the process of gas migration macroscopically, but also obtained pathological evidence of diaphragmatic periodic perforation microscopically, which is especially precious and confirms the existing theory that retrograde menstruation leads to diaphragmatic endometriosis, and the diaphragmatic fenestration is obtained due to the periodic activities of ectopic endometrium.

Keywords: Case report, Catamenial pneumothorax, Endometriosis, Thoracic endometriosis syndrome TES
**Case presentation**

A 43-year-old woman with a history of endometriosis came to our hospital because of frequent chest stuffiness (attacking every three to six months), which can be traced back to 2 years ago. The X-ray demonstrated a right mild pneumothorax. Subsequently, this recurring symptom closely related to menstruation bothered her deeply. During this time, she tried closed drainage of pleural cavity, however, the frequent recurrence of her symptoms was bothersome. Chest CT scan showed a recurring pneumothorax on the right, no obvious bullae, we decided to explore the source of pneumothorax under uniportal VATS.

We thoroughly explored the diaphragm, visceral and parietal pleura: no bullae were found; multiple diaphragmatic defects were found on the dome (Fig. 1a). And surprisingly, bubbles were slipping through the defects to the thoracic cavity (Fig. 1b); the visceral pleura was scattered with yellowish-brown implants (Fig. 1c). We excised the diaphragmatic lesions and wedge resected the right upper lung lesions (shown in Fig. 1d); cleared the deposits and rinsed pleural cavity with pure iodophor; wiped the parietal pleura with dry gauze to preclude hidden lesions.

Histopathological examination revealed endometriosis in diaphragmatic lesions, periodic shedding of endometrial cells and a large number of blood cells can be seen in the gland cavity, immunohistochemistry showed CD10, ER and PR positive (Fig. 2), right upper lung lesions showed inflammation.

On the second day after operation, she had menstruation. Her postoperative process was uneventful. Subsequently, she received endocrine therapy (Gonadotropin Releasing Hormone agonist GnRH-a; 3.6 mg; once every 28 days, 4 times in total), after a period of follow-up, the pneumothorax did not recur.

---

**Fig. 1** Intraoperative findings; **a** There were multiple defects in the diaphragmatic dome, and the swing of liver can be clearly seen along with breathing movement. **b** With breathing, bubbles were overflowing into the pleural cavity through the fenestrations. **c** Yellowish-brown nodules and fibrinous deposits scattering in the pleural cavity and on the visceral pleura. **d** Tight attachments of the upper right lung that cannot be washed away.
Discussion
Catamenial pneumothorax is characterized by spontaneous recurring pneumothorax during menstruation, which is a common clinical manifestation of thoracic endometriosis syndrome. Surgical treatment has become one of its standardized treatment methods, in previous literature reports, the diaphragmatic defect observed during surgery is the most common sign [4]. The most appropriate time for surgery is recommended during menstrual period, so that the typical signs and pathological findings can be observed more clearly. However, the testimony of gas migration is only an occasional capture in imaging [5]. In our case, we chose to operate on the eve of menstruation, the initial expectation was to confirm the existence of thoracic endometriosis via pathology. By chance, we witnessed the scene of bubbles boiling on the diaphragmatic defects and diaphragmatic decidual ectopic gland cells under the microscope. Until now, there has not been such a typical discovery of this rare case.

The relationship between catamenial pneumothorax and endometriosis has been confirmed, in our case, the endometriosis history and pathological conclusions can also validate the speculation. Its pathogenesis can be explained by the “Determinant of uterineeutopic endometrium “as a clinical manifestation of TES: Eutopic endometrium is the root cause, its own characteristics determine the location of the thoracic lesions, diaphragm is the most common, and so do the lungs and the pleura. Meanwhile, about catamenial pneumothorax, there are mainly three related mechanisms [6]: 1. Anatomical theory: Retrograde menstruation resulting in subdiaphragmatic endometriosis, air migrates to pleural cavity through the genital tract and acquired diaphragmatic perforation during menstruation; 2. Metastasis theory: Ectopic endometrial tissue spreads to the thoracic cavity through blood or lymphatic vessels; 3. Physiological theory: The effects of prostaglandin F2.

During the operation, we found a large number of yellowish-brown nodules, considering that the visceral pleura was also involved. After repeatedly flushing the
pleural cavity with sterile water, we noticed that there was yellowish-brown deposition tightly sticking to upper right lung, but the postoperative pathology revealed inflammatory exudation. We believe that the fibrin deposits and numerous yellow–brown nodules were exudate of the pleura and the previously shed endometrial tissue.

High recurrence rate is the main trouble in the treatment of catamenial pneumothorax. However, for patients with separate diaphragmatic lesion, there are reports in the previous literatures that surgical treatment usually has a very good prognosis [7, 8]. For this patient, in addition to removing the diaphragmatic lesions and reconstructing diaphragm, we also used pure iodophor flushing the thoracic cavity to promote pleural adhesions, which we believed that may be used as another preventive measure to form coagulated iodinated protein protective membrane on the lung surface. Carefully exploring the entire pleural cavity and wiping the parietal pleura with dry gauze to exclude other lesions. After the perioperative period, we added endocrine therapy. Through this series of systematic treatment measures, we believe that the efforts paid off.

Surgical exploration of catamenial pneumothorax is usually found with diaphragmatic fenestrations. However, in our case, the witness of bubbling up on the diaphragmatic defects and the positive pathological results have proved that we accurately grasped the timing of surgery. At this time, we discovered a phenomenon that has not been reported before and further clarified the pathogenesis of catamenial pneumothorax, which is still controversial until now.

Abbreviations
VATS: Video-assisted thoracic surgery; SP: Spontaneous pneumothorax; CT: Computed tomography; TES: Thoracic endometriosis syndrome; GnRH-a: Gonadotropin releasing hormone agonist.

Acknowledgements
We acknowledge the contribution made by the Department of Pathology, the First Affiliated Hospital of Zhengzhou University in the interpretation of the histology.

Authors' contributions
BD: Drafting of the manuscript and treated the patient. YQ: Provided the overall supervision of the case and the production of the manuscript. YLS, BW and CLW: Designed and participated in this surgery. LH, YFL, GCY and SHL: Attend to the patient and figures processing. All authors read and approved the final manuscript.

Availability of data and materials
Data sharing is not applicable to this article as no datasets were generated or analysed during the current study.

Declarations

Ethics approval and consent to participate
Ethical approval was given by the Ethics Committee of the First Affiliated Hospital of Zhengzhou University, Zhengzhou, Henan Province, China.

Consent to publication
For publication of this case report, including the images contained within, informed written consent was obtained from the patient. A copy of the consent form is available for review by the Editor of this journal.

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

Received: 9 December 2020 Accepted: 15 April 2021
Published online: 20 April 2021

References
1. Maurer ER, Schaal JA, Mendez FL. Chronic recurring spontaneous pneumothorax due to endometriosis of the diaphragm. J Am Med Assoc. 1958;168(15):2013–4.
2. Lillington GA, Mitchell S. Catamenial pneumothorax. JAMA. 1972;219(10):1326–32.
3. Baoquan L, Liangjian Z, Qiang W, Hai J, Hezhon C, Zhiyun X. Catamenial pneumothorax associated with multiple diaphragmatic perforations and pneumoperitoneum in a reproductive woman. J Formos Med Assoc. 2014;113(6):385–7.
4. Hsu PK. Tissue graft to cover diaphragmatic defects in a patient with catamenial pneumothorax. Ann Thorac Surg. 2019;107(e143).
5. Roth T, Alfano M, Schussler Q, Magdaleinat P, Regnard JF. Catamenial pneumothorax: chest X-ray sign and thoracoscopic treatment. Ann Thorac Surg. 2002;74(2):563–5.
6. Kawaguchi Y, Fujita T, Hanaoka J. Catamenial pneumothorax with bullae. Ann Thorac Surg. 2015;99(3):1075–8.
7. Takahashi M, Matsukura T, Hirai T, Mino N. Recurrent catamenial hemothorax treated by coverage with polyglycolic acid sheets. J Thorac Cardiovasc Surg. 2013;145(1):300–2.
8. Baoquan L, Liangjian Z, Qiang W, Hai J, Hezhong C, Zhiyun X. Catamenial pneumothorax associated with multiple diaphragmatic perforations and pneumoperitoneum in a reproductive woman. J Formos Med Assoc. 2014;113(6):385–7.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.