**CASE REPORT**

**Catamenial hemoptysis and pneumothoraces in a patient with cystic fibrosis**

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Hemoptysis or pneumothorax that recurs with the onset of menses is strongly suggestive of thoracic endometriosis syndrome (TES). TES is a rare disorder, with relatively few cases reported in the literature. A 32-year-old woman with cystic fibrosis, who over a period of several months had experienced recurrent catamenial hemoptysis and pneumothoraces, including an episode of life-threatening hemoptysis that coincided with menstruation, is presented. Thoracic computed tomography and magnetic resonance imaging scans, as well as a bronchoscopic evaluation that demonstrated endobronchial lesions that disappeared after menses, support the diagnosis of TES in the present patient. The patient was treated empirically with danazol and subsequently underwent a successful double-lung transplantation. Danazol was discontinued postoperatively, and she was started on an oral contraceptive. Eighteen months post-transplant, she has not experienced a recurrence of her catamenial symptoms, despite having resumed a regular menstrual cycle.

**Key Words:** Catamenial; Cystic fibrosis; Hemoptysis

**CASE PRESENTATION**

Over several months, a 32-year-old woman with CF (AF508, ΔI507 genotype) experienced several episodes of mild, streaky hemoptysis and two episodes of spontaneous left pneumothorax, which coincided with the onset of menses. She was nulligravid, had no known previous history of pelvic endometriosis nor any history of pelvic pain, dysmenorrhea or uterine manipulation. In September 2004 she had developed sudden massive hemoptysis on the second day of menses. Immediately following this episode, she had collapsed and required resuscitation. The volume of blood expectorated was estimated to be approximately 500 mL and was associated with an acute fall in hemoglobin concentration of 18 g/L. Before this event, she was relatively well; her baseline lung function had remained stable (forced expiratory volume in 1 s 32% of predicted), and there were no symptoms to suggest an antecedent infectious exacerbation. She was nonetheless treated with antibiotics and had recovered. With the onset of her next menstrual cycle, she experienced recurrent streaky hemoptysis. Bronchial arterial angiography demonstrated the presence of an extensive collateral circulation, and several ectatic arteries were embolized. A computed tomography (CT) scan obtained during menses demonstrated the presence of pulmonary parenchymal lesions, consistent with hemorrhage, that resolved completely on a follow-up CT scan obtained midway through her menstrual cycle (Figure 1). Thoracic magnetic resonance imaging, obtained during menses, demonstrated...
hyperintense lesions (consistent with hemorrhage or endometrial tissue) that had localized to her left apical pleura. Bronchoscopy, performed premenstrually, showed at least two friable purplish lesions in her segmental bronchi, which were no longer evident after menses. Bronchial washings did not show endometrial cells. At the patient’s request, neither biopsies nor brushings were performed. Given the strong clinical suspicion of TES, she was started empirically on danazol and became amenorrheic. She experienced another episode of life-threatening hemoptysis approximately three months after starting danazol therapy coincident with an infectious exacerbation of CF, and subsequently underwent double-lung transplantation. Histological evaluation of her native lungs did not demonstrate the presence of endometrial tissue. Danazol treatment was discontinued postoperatively, and the patient was started on an oral contraceptive. Twenty-four months post-transplant, she had no further episodes of hemoptysis or pneumothoraces, despite having resumed menses.

DISCUSSION
To our knowledge, this is the first report of catamenial hemoptysis and pneumothoraces occurring in a patient with CF. In the present patient, the clinical history, as well as the dynamic radiographic and bronchoscopic appearance, strongly suggest a diagnosis of TES. Although we were unable to exclude an infectious exacerbation of the underlying CF, the lack of clinical symptoms to suggest an infectious etiology, as well as the recurrent catamenial nature of both the hemoptysis and the pneumothoraces, did not favor this diagnosis. Nonetheless, she was treated empirically with antibiotics. Furthermore, she was known to be colonized with Aspergillus fumigatus, but had experienced recurrent hemoptysis, despite initiation of treatment with itraconazole. Although a definitive histopathological diagnosis was not obtained, it should be noted that in approximately two-thirds of the cases of TES reported in the literature, diagnosis was established based on historical and clinical criteria alone (3). Furthermore, treatment with danazol has been demonstrated to result in macroscale regression of ectopic endometriomas, which may have made pathological confirmation more difficult in the present case (4). The fact that this patient experienced both catamenial hemoptysis and pneumothoraces suggested that ectopic endometriomas had localized to both pleura and pulmonary parenchyma, which has also been previously described (2,5). In previous reports (1), the volume of blood expectorated in the setting of catamenial hemoptysis was usually less than 200 mL, and life-threatening hemoptysis has hitherto not been described. We propose that the massive nature of the hemoptysis experienced by our patient may be a consequence of the bronchial arterial collateral circulation (visualized angiographically) that had developed due to her underlying bronchiectasis. Interestingly, although it is generally accepted that the most likely etiology of catamenial hemoptysis is TES, there is a single case report (3) in the literature of a pulmonary arteriovenous malformation presenting with cyclical bleeding coincident with menses. This may suggest that in some disease processes, at least, the pulmonary vasculature may be sensitive to hormonal changes, and it is therefore possible that such a mechanism was affecting the ectatic bronchial circulation in our patient. However, this possibility would not be expected to explain the occurrence of catamenial pneumothoraces.

The optimal treatment of TES is uncertain, with surgical resection and ovarian suppression being the most widely reported (1,6). Danazol is reported to be effective, although treatment failures have been described (7). Other treatments reported to be successful include gonadotropin-releasing hormone antagonists, oral contraceptives, estrogen subdermal implants or the use of clomiphene citrate therapy (1,8). More recently, the use of endobronchial laser ablation has been described for lesions that are detected bronchoscopically (9,10). In some cases, at least, there may also be spontaneous resolution or improvement in symptoms, even in the absence of therapy (11).

Bronchoscopic findings in patients with catamenial hemoptysis and presumed TES are variable. Multiple bilateral purplish-red submucosal lesions are most commonly described, but appearances ranging from a single tiny red mucosal lesion (9) to diffuse mucosal hyperemia (8) have been reported. The

Figure 1) Corresponding slices of thoracic computed tomography scans taken during menses (A) and mid-cycle (B). Images taken during menses demonstrated the presence of pulmonary parenchymal lesions (white arrows) consistent with hemorrhage, which were no longer evident on a scan taken approximately two weeks later when the patient was mid-cycle.
yield of bronchial biopsy in establishing the diagnosis of TES is curiously low, although cytological analysis of samples obtained with brush biopsy may be more sensitive. However, as in our patient, follow-up bronchoscopic examination performed postmenstrually or in mid-cycle typically demonstrates resolution of previously visualized lesions (8). Similarly, radiographic appearance on chest CT scans can be variable, but may include ill-defined parenchymal opacities, nodular lesions or ground glass opacifications; these lesions are often dynamic and change with the menstrual cycle.

Controversy also exists in terms of the presumed etiological mechanisms by which ectopic endometrial tissue arises in the thorax; three theories have been proposed (1,12). The theory of coelomic metaplasia maintains that uterine endometrium and pleural mesothelium share the same embryological origin, and undefined pathogenic stimuli may result in the differentiation of pleural-based precursor cells into endometrial cells. However, this does not explain the occurrence of intrapulmonary endometriosis. The second theory is that of the transdiaphragmatic passage of endometrial cells through diaphragmatic defects; this too could explain pleural deposits of endometrium but would not be expected to result in intraparenchymal pulmonary deposits. The third theory suggests that endometrial cells may embo-lize from the pelvis to other extraterine or extrapelvic sites by using the lymphatics or the bloodstream. Alternatively, ectopic endometrium that arises in the pleural space as a consequence of another mechanism (such as transdiaphragmatic passage of endometrium) could embo-lize via regional lymphatics. This mechanism could explain the development of multiple sites of ectopic endometrium, including intraparenchymal lung lesions. However, none of these theories can in isolation explain all of the clinical manifestations of TES, and the etiology may be multifactorial. In the present patient, the risk of recurrence of pulmonary endometriosis in the lung allograft is unknown.

REFERENCES