Birt-Hogg-Dubé (BHD) syndrome is an autosomal dominant inherited disorder that is characterized by skin fibrofolliculomas, renal tumors and multiple lung cysts with or without spontaneous pneumothorax. The disease is caused by germline mutations in the FLNC gene that codes for a protein of unknown function called folliculin. Patients with BHD syndrome do not always have all three manifestations of the skin, kidney and lung. To the best of our knowledge, there has been no case report of the radiologic findings of the lung manifestation in a patient with BHD syndrome in Korea. We report here on two cases of BHD syndrome that presented with spontaneous pneumothorax. The pulmonary abnormalities consisted of multiple thin-walled cysts of various sizes and shapes in both lungs.

**Index words:** Lung  
Cysts  
Pneumothorax

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**Case Reports**

**Case 1**

A 47-year-old woman visited our emergency room for an acute onset of dyspnea. Her chest radiograph (Fig. 1A) showed large pneumothorax in the right lung and a radiolucent area in the left lower lung field. The chest CT scan taken after closed thoracostomy showed multiple cysts of various sizes and shapes in both lungs (Figs. 1B–C). The pulmonary cysts were sharply demarcated with thin walls and they measured 0.3–5.5 cm in diameter. The intervening lung parenchyma between the cysts appeared normal. The initial radiologic impression after the CT scan was lymphangioleiomyomatosis (LAM). The patient had no smoking history. She had a history of thoracoscopic bulla resection for a left pneumothorax 15 years ago. The patient reported that her two sisters and one of her daughters had undergone thoracic surgery for spontaneous pneumothorax. The pa-
tient then underwent wedge resection in the right lung, and this revealed bulla and blebs without evidence of LAM [Fig. 1D].

Eight months after the patient’s surgery, her daughter visited our hospital and presented with recurrent pneumothorax. The chest CT scan of the daughter after closed thoracostomy also revealed multiple thin-walled cysts in both lungs [Figs. 1E, F]. The daughter had a previous history of wedge resection at our hospital a year previously, and her previous lung specimen revealed bulla only [Fig. 1G].

**Case 2**

A 54-year-old woman presented with dyspnea, cough and chest pain. Her chest radiograph showed pneumothorax in the left lung. The chest CT scan after closed thoracostomy revealed multiple thin-walled cysts and bullae in both lungs (Figs. 2A-C). They were round or oval shaped and some of them showed a lobulating contour. The intervening lung parenchyma between the cysts was normal without any evidence of pulmonary nodule or fibrosis. She had a history of closed thoracostomy due to right pneumothorax 35 years ago. She had a 20 pack-year smoking history and had stopped smoking 5 years ago. She had multiple whitish papules measuring 2–3 mm on her cheek, and a punch biopsy was performed. The pathologic result was perifollicular fibrosis, which was consistent with the histologic diagnosis of fibrofolliculoma [Fig. 2D]. Ultrasonography of the kidneys did not show any abnormality.

**Discussion**

Birt-Hogg-Dubé (BHD) syndrome is a rare autosomal dominant inherited disorder that is characterized by skin fibrofolliculomas, renal tumors and multiple lung...

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Fig. 1. Chest radiograph (A) of patient 1 shows a large pneumothorax in the right lung with passive atelectasis of the right lung. The chest CT images (B and C) after closed thoracostomy show multiple cysts of various sizes and shapes in both lungs. The pathologic diagnoses of the pulmonary cysts were bulla and blebs (D).
cysts with or without spontaneous pneumothorax [1–3]. Skin fibrofolliculoma is a benign hamartoma of the hair follicle and this is the hallmark of BHD’s cutaneous manifestation. Fibrofolliculomas usually appear as multiple, dome-shaped, whitish papules in the face [4, 5]. The renal tumors associated with BHD syndrome show various histologic types, ranging from benign oncocytes to malignant renal carcinomas. Patients with BHD syndrome have a 7-fold increased risk of developing renal neoplasm and the most life-threatening complication of BHD syndrome is renal cancer [4, 6]. More than 80% of adult patients with BHD syndrome have multiple lung cysts. Despite the presence of multiple lung cysts, the lung function is usually unaffected [7]. The histology of the pleuropulmonary lesions include intraparenchymal collections of air surrounded by normal parenchyma or a thin fibrous wall, and blebs within the pleura [8]. Patients with BHD syndrome have a 50-fold increased risk of spontaneous pneumothorax [6].

The clinical diagnosis of BHD syndrome requires the fulfillment of one major or two minor criteria. The major criteria include: 1) at least five fibrofolliculomas with at least one histologically confirmed and 2) pathogenic FLCN germline mutation. The minor criteria include: 1) multiple lung cysts that are bilateral basally located, with or without spontaneous pneumothorax, 2) renal cancer with an early onset (< 50 years old) or multifocal or bilateral renal cancer, and 3) a first-degree relative with BHD syndrome [5]. Based on these criteria, our patients could be diagnosed as having BHD syndrome (multiple lung cysts and a familial history of spontaneous pneumothorax in patient 1, and multiple lung cysts and fibrofolliculomas in patient 2).

Patients with BHD syndrome do not always have all three manifestations, including skin, kidney and lung manifestations. Toro et al. reported that skin fibrofolliculomas, pulmonary cysts and renal tumors were found in 98%, 89% and 23% of 198 patients, respectively [7]. According to a report of Tobino and colleagues, pulmonary cysts may develop earlier than the skin and renal lesions; therefore, the detection of pulmonary cysts on chest CT scan may be useful for making an early diagnosis of this syndrome. On the analysis of the thin-section CT findings of pulmonary cysts in 12 patients with genetically diagnosed BHD syndrome, multiple, irregular-shaped cysts of various sizes with a predominance in the lower medial lung zone were characteristic for BHD syndrome. In addition, cysts abutting or including the proximal portions of the lower pulmonary arteries or veins were found in all the patients [9]. These findings can be helpful for making the radiologic differential diagnosis for diseases associated with multiple pulmonary cysts such as lymphangioleiomyomatosis, Langerhans cell histiocytosis and lymphocytic interstitial pneumonitis [10].
In summary, multiple lung cysts with or without spontaneous pneumothorax may be the first manifestation of BHD syndrome. Therefore, clinicians must be aware of this syndrome and they should recommend that the affected patient and family members undergo proper screening for renal tumor.

References

Birt-Hogg-Dubé 증후군에서 나타난 자발성 기흉: 2 증례

배형주∙우옥희∙용환석∙강은영∙김현구2∙최영호2∙신봉경3∙김윤경

Birt-Hogg-Dubé (BHD) 증후군은 피부의 섬유모낭종, 신장종양, 다발성 폐낭종을 나타내는 드문 상염색체 우성 유전질환으로 자발성 기흉을 동반하는 경우가 있다. 이 질환은 기능이 잘 알려지지 않은 folliculin이라는 단백질을 합성하는 FLCN 유전자의 생식계열 변이 때문에 발생한다. BHD 증후군 환자들은 항상 피부, 신장, 폐의 세 가지 이상징후를 모두 나타내는 것은 아니다. 지금까지 대한민국에서 BHD 증후군의 폐 이상징후에 대한 증례보고는 없었다. 이에 우리는 자발성 기흉으로 나타난 BHD 증후군의 두 증례를 보고하고자 한다. BHD 증후군의 폐 이상 소견은 얇은 벽을 가진 다양한 크기와 모양의 양측성 다발성 폐낭종으로 구성되어 있었다.