Fatal Asymmetric Interstitial Lung Disease after Erlotinib for Lung Cancer

Shaohua Ren\textsuperscript{a}  Yuan Li\textsuperscript{a}  Weiwen Li\textsuperscript{a}  Zhongwei Zhao\textsuperscript{b}  Chunxian Jin\textsuperscript{b}  Dengke Zhang\textsuperscript{b}

Departments of \textsuperscript{a}Respiratory Medicine and \textsuperscript{b}Radiology, Lishui Central Hospital, Lishui, PR China

**Abstract**

Pulmonary toxicity is a known complication of erlotinib, one of the epidermal growth factor receptor tyrosine kinase inhibitors. It consists of diverse entities such as interstitial pneumonitis, bronchiolitis obliterans with organizing pneumonia and pulmonary fibrosis. In our report, an unusual case of an asymmetric interstitial lung disease was described. A 68-year-old female presented with resistant cough, hemoptysis and a right lung atelectasis on chest X-ray. She underwent selective bronchial artery embolization successfully after pharmaceutical therapy failed to stop hemoptysis. Flexible bronchoscope revealed that the opening of the right main bronchus was blocked completely by a neoplasm with a distance < 2 cm to the carina and the sample of bronchoscopic biopsy confirmed the diagnosis of lung adenocarcinoma (cT3N2M0). Dyspnea and asymmetric interstitial lung disease in the nontumorous lung were noted on the 6th day.

**Key Words**

Lung cancer \cdot Selective bronchial artery embolization \cdot Erlotinib \cdot Interstitial lung disease

**Established Facts**

- Drug-associated interstitial lung disease is a rare, but serious and often fatal adverse reaction to erlotinib, one of the oral epidermal growth factor receptor tyrosine kinase inhibitors.
- Lung injury may be perpetuated by ventilation strategies that do not limit lung volumes and airway pressures. Autopositive end-expiratory pressure is one of the risk factors for barotrauma.

**Novel Insights**

- We introduce a rare condition of asymmetric interstitial lung disease in a patient treated with erlotinib for lung adenocarcinoma.
- The additional affliction of the right lung was related to the dual effects of the obstructing tumor at the right main bronchus and selective bronchial artery embolization, which resulted in overexpansion, hypoventilation and hence underperfusion.
of erlotinib therapy (150 mg daily) which had been efficacious in its anticancer effect. Discontinuing erlotinib use and treatment with corticosteroids could not relieve her symptoms. The patient deteriorated rapidly and died of progressive respiratory failure. We explored the mechanisms of asymmetric interstitial lung disease.

Introduction

Interstitial lung disease (ILD) is a rare but fatal complication of erlotinib which was approved recently for the treatment of advanced non-small-cell lung cancer. The distribution of disease is bilateral and symmetrical in typical ILD. We present a patient who developed an unusual type of asymmetric ILD after erlotinib treatment for her lung adenocarcinoma.

Case Report

A 68-year-old female was admitted with a history of resistant cough and hemoptysis for 2 months. Physical examination revealed normal vital signs. Chest auscultation was significant for no breathing sound in the right lung, and jugular venous distention was not evident. Blood routine, biochemical tests and electrocardiogram were all within a normal range. An echocardiogram showed normal left ventricular systolic and diastolic functions. Chest X-ray (CXR) displayed a right lung atelectasis (fig. 1a). She underwent selective right bronchial artery embolization (BAE) successfully for her refractory hemoptysis as pharmaceutical therapy had failed to stop bleeding (fig. 1b, c). Flexible bronchoscope showed a neoplasm obstructing the entire opening of the right main bronchus with a distance <2 cm from the carina (fig. 1d). Bronchoscopic biopsy histology led to the diagnosis of adenocarcinoma (cT3N2M0).

The patient began therapy with erlotinib, 150 mg daily, and within 5 days her condition improved dramatically. On the 6th day, she had a good appetite and normal temperature without any signs of infection, but complained of dyspnea. Velcro rales could be heard at the bottom of the left lung. The blood routine was within a normal range. Chest CT demonstrated resolution of the atelectasis and the right lung had reexpanded well, but there was a large area of ground-glass attenuation (GGA), mainly in the left

![Fig. 1.](image-url)
ILD Is a Rare but Fatal Complication of Erlotinib

lung (fig. 2a). Erlotinib-associated ILD was considered and the drug was discontinued immediately (it had been taken for 6 days in total). The patient was started on 250 mg of methylprednisolone administered intravenously, every 12 h. Despite all the treatments, including corticosteroids, oxygen supply and jet ventilation, her progressive dyspnea with hypoxemia caused a rapid deterioration of her condition. The arterial blood gas under a nonrebreathing mask O2 supply (FiO2 = 100%) was: pH 7.450, PaO2 54.5 mm Hg, PaCO2 24.1 mm Hg and SaO2 80%. Chest CT taken 4 days later revealed a diffuse distribution of GGA with a mosaic pattern and consolidation in the left lung, and GGA became more obvious at the base of the lower lobe of the right lung (fig. 2b). All of these abnormalities progressed rapidly on CXRs in the following days (fig. 2c, d). She refused mechanical ventilation, was discharged and died at home.

Discussion

The patient did not undergo chemotherapy, radiotherapy or receive any other agent that could potentially damage lung tissues. There were no signs of infection or cardiogenic pulmonary edema. Considering the temporal relationship between the use of erlotinib and the onset of symptoms, it is reasonable to conclude that the fatal ILD in the patient was induced by erlotinib [1, 2].

Erlotinib, an epidermal growth factor receptor tyrosine kinase inhibitor, is an effective antitumor agent for the treatment of non-small-cell lung cancer, especially in Asians. Lately, it is recommended as a first-line treatment for patients with advanced epidermal growth factor receptor mutation-positive non-small-cell lung cancer [3]. ILD is a rare but potentially life-threatening complication of erlotinib therapy with an overall incidence about 0.2–6.5% in different studies [4, 5].

It is intriguing that the involvement of GGA was confined in the unilateral (or very asymmetric) left lung. Asymmetric ILD is not commonly seen, but does exist rarely [6–11]. Padley et al. [6] described an asymmetric form of acute respiratory distress syndrome (ARDS) following pulmonary resection. The authors found that density increased more in the nonoperated lung than in the operated lung in the patients who developed ARDS following lobectomy. This was considered as an asym-
metric form of ARDS rather than compensatory hyper-expansion of the residual lung on the operated side. Kon-doh et al. [7] also found that in 5/236 patients with interstitial pneumonia who underwent a surgical lung biopsy, the extent of the parenchymal involvement was significantly greater on the nonoperated side (p = 0.0251) in all patients. The etiology of asymmetric ARDS or the extent of the parenchymal involvement was mainly related to intraoperative respiratory management, i.e. single-lung ventilation, which brought on high pressure or barotrauma and oxygen toxicity in the nonoperated lung [6, 7]. Asymmetric ILD is also seen under nonsurgery conditions. Acid reflux and nonacid reflux in gastroesophageal reflux may proceed to the development of idiopathic pulmonary fibrosis (IPF) [8]. In a recent study, Tchera- kian et al. [9] found the rates of gastroesophageal reflux and acute exacerbations were significantly higher in the patients with asymmetrical IPF than in those with symmetrical IPF. Most of the patients with asymmetrical IPF (62.5%) showed a predominance of fibrosis in the right lung. Furthermore, the side of the fibrosis, whether right or left, matched with the patient’s preferred posture for falling asleep in 94% of the asymmetrical IPF cases. Repeated microaspirations associated with gastroesophageal reflux may be locoregional factors in the development of asymmetrical IPF and acute exacerbations. Unilateral usual interstitial pneumonia and unilateral IPF were occasionally reported in the patients with slow-growing pulmonary artery sarcoma or congenital absence of the right interlobar pulmonary artery, respectively. Chronic pulmonary ischemia associated with systemic collateral vessels was hypothesized for the pathogenesis of the diseases [10, 11].

As in those cases under surgery conditions, the additional affliction of the right lung in our case was due to the apparent check-valve effect of the obstructing tumor at the opening of right main bronchus which resulted in overexpansion, hypoventilation and hence underperfusion of the right lung. GGA at the bottom of the right lung was as a result of relatively higher pressure and ventilation at the base in a respiratory cycle.

Apart from the above, BAE may also play a role in the protection of the right lung by reducing the lung blood supply. Blockade of bronchial blood flow has been observed as minimizing pulmonary injury in sheep [12]. In humans, although the bronchial arteries make up only 1% of cardiac output, the function of BAE is more than providing nutrition for the walls of the bronchi, pulmonary arteries and veins. The bronchopulmonary arterial anastomoses can respond in the form of hypertrophy and perform a vital, adaptable function; more blood can be directed through the anastomoses toward the pulmonary artery to poorly ventilated or atelectatic areas in order to feed the parenchyma where its own capillaries would not carry oxygenated blood [13]. Furthermore, the diameter of the bronchial arteries might increase in the patient because of the presence of hemoptysis [14]. BAE carried out then might not only reduce the blood supply to the bronchial tumor, but also to the rest of the right lung, where a collateral circulation between the bronchial and pulmonary circulation was developed anatomically after chronic obstruction of the pulmonary arterial system by the lung tumor. However, the exact mechanism of how the reduction of lung blood supply by BAE would minimize pulmonary injury is not yet known.

In a recent study, Ter et al. [15] found that the development of the fatal ILD might be associated with high erlotinib and metabolite levels. It is possible that BAE also provides unintended protection of the right lung in our case. BAE reduced the blood supply to the bronchial tumor, joined together with the check-valve effect of the obstructing tumor, somehow decreasing the blood perfusion and lowering drug distribution in the whole right lung accordingly, and thus finally producing less toxicity in the right lung.

**Financial Disclosure and Conflicts of Interest**

The authors have nothing to disclose.

---

References

ILD Is a Rare but Fatal Complication of Erlotinib


