Sir,

Tuberculosis (TB) is the second most common fatal infectious disease, following human immunodeficiency virus/acquired immunodeficiency syndrome (HIV/AIDS), in the adult population. Socio-economic deprivation, immigration, wars, omission of tuberculosis control programs, and HIV/AIDS epidemics have led to an increase in the incidence of tuberculosis. The World Health Organization (WHO) publishes a global tuberculosis report annually. The 2013 report stated that, in 2012, 8.6 million people developed TB, and 1.3 million people died from the disease, including 320,000 deaths among HIV-positive individuals [1].

As a rare complication of spontaneous pneumothorax, spontaneous hemopneumothorax (SHP) is an emergency condition that can be life threatening because of active bleeding in the pleural space; the condition has a high mortality rate and requires early diagnosis and treatment [2, 3]. A report has suggested that TB will not cause SHP due to thick and extensive pleural adhesions [4].

We present here four cases with SHP secondary to a hypovolemic shock condition due to hemorrhage, together with a literature review due to the rarity of this condition and discuss applicable treatment approaches. Pleural biopsy specimens were stained with Kinyoun and cultivated on Löwenstein-Jensen medium. Cultivation was also performed using a liquid automatized Bactec 460 TB system (Becton-Dickinson).

The first patient was a 34-year-old male admitted to the Emergency Room (ER) because of pain in his right chest and back. With no history of trauma, the patient’s physical examination showed diminished breath sounds in the right hemithorax. His chest X-ray revealed an appearance consistent with hydrothorax with a partial pneumothorax line on the right (Figure 1A). Laboratory findings reported the hemoglobin level of 10.4 g/dL and a hematocrit level of 31.4%. No pathological data was detected in coagulation parameters. Tube thoracotomy (TT) was performed on the right after a preliminary diagnosis of SHP and obtained 1000 cc of fresh blood. Emergency right thoracotomy was performed following 300 ml of hemorrhage during the first hour, a clinical condition associated with hypotensive and hypovolemic shock (80/50 mmHg); a hemoglobin value reduced to 8.5 g/dL and a hematocrit value to 25.2% in the clinical follow up. Exploration showed that the apex of the upper lobe was adherent to the chest wall with a small adhesion, and aberrant vessels with diameters of ~0.4 mm at this site and an actively bleeding varicose pattern between the chest wall and pleura were identified. A suture procedure was carried out, and three units of blood were transfused. TB treatment was initiated two months after the procedure was carried out, and three units of blood were transfused. TB treatment was initiated two months after the

The second patient was a 25-year-old male admitted to the ER with complaints of shortness of breath and sudden pain in the right chest showed diminished breath sounds on the right in chest auscultation. His chest X-ray showed a partial pneumothorax line on the right consistent with hydrothorax (Figure 1B). The patient had no history of trauma. Laboratory findings reported a hemoglobin level of 12.6 g/dL, and a hematocrit level of 36.8%. TT was performed on the right after a preliminary diagnosis of SHP. Air drainage was initially performed and yielded 700 cc of hemorrhagic fluid. Emergency right thoracotomy was performed following 300 ml of hemorrhage during the first hour followed by 200 ml of hemorrhage per hour, a clinical condition associated with hypovolemic hypotension (90/50 mmHg) with a hemoglobin value of 9.4 g/dL and a hematocrit value of 28.2% in his clinical follow up. Exploration found a cavity lesion with a dimension of 4x3x2

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Tuberculosis cases presenting with spontaneous hemopneumothorax and hypotension

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Tuberculosis cases presenting with spontaneous hemopneumothorax and hypotension

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lesion in the apical region of the hemithorax. The bullous lesions were excised, and bridging ligation was performed. The patient was followed up by the CDD upon detection of AFB in his sputum and treatment was initiated thereafter.

The fourth patient was a 43-year-old male with no history of trauma who was admitted to the ER because of breath difficulty and sudden pain in the right chest. He was hypotensive (90/50 mmHg), dyspneic, and tachypneic, with no breath sounds heard in the right hemithorax. His chest radiography was consistent with total pneumothorax in the right and hydrothorax and deletion of the diaphragm contours. The bleeding profile and biochemical values were within normal limits, a hemoglobin level of 9.1 g/dL, a hematocrit level of 29.4%, and a platelet count of 435,000/mm³. TT was performed on the right hemithorax. Intravenous fluid replacement was initiated. Air drainage was initially performed, yielding 1200 cc of fresh blood. Air discharge and hemorrhagic fluid drainage were stopped at the first hour, and no additional surgical interventions were considered due to his hypotensive condition and improved dyspnea. Two units of blood were transfused upon the decrease of the hemoglobin value to 8.3 g/dL and hemat-

cm, ruptured to the pleura in the apical segment of the upper lobe, and segmentectomy was performed. Clinical sample revealed acid fast bacilli (AFB). TB treatment was initiated due to this result by the Chest Diseases Department (CDD) and microbiological evaluation reported tuberculosis after two months.

The third patient was a 28-year-old male admitted to the ER with complaints of pain in the left chest and shortness of breath. With no history of trauma, the patient's thoracic radiograph showed an increased density varying with partial pneumothorax on the left. The bleeding profile and biochemical values were within normal limits and a hemoglobin level of 11.5 g/dL, and a hematocrit level of 32.4%. TT was performed on the left hemithorax. Air drainage was initially performed, yielding 1250 cc of hemorrhagic fluid. Emergency video-assisted thoracoscopic surgery (VATS) was performed in the left hemithorax following 250 ml of hemorrhagic fluid drainage during the first hour due to hypotensive and hypovolemic condition (90/40 mmHg), and a hemoglobin value reduced to 9.1 g/dL and a hematocrit to 27.2%. Exploration found bullous lesions, with a maximum diameter 2×2 cm, in the apical segment of the upper lobe and an actively bleeding adhesion

Figure 1 | The chest X-ray and CT images of the patients
ocrit to 26.4%. The patient was followed up by the CDD upon detection of AFB in his sputum and a high level of adenosine deaminase (ADA: 120 U/L) in the pleural fluid; TB treatment was initiated by the CDD (Figure 1C and 1D).

Although hemopneumothorax is typically a result of trauma, SHP is not associated with a trauma history. Associated with pneumothorax, SHP involves ≥400-ml bleeding in the pleural cavity [4]. SHP is observed mostly in the adolescent age group and may cause life-threatening bleeding [2]. The incidence of SHP varies between 0.5% and 12% [5]. Our clinic treated 184 spontaneous pneumothorax patients between 2008 and 2013, only six of whom (3.2%) were diagnosed with SHP. SHP secondary to TB was found in four patients.

Three bleeding mechanisms were defined in SHP: first, the lysis of adhesions between visceral and parietal pleura; second, the rupture of congenital aberrant vessels between the parietal pleura and bulla; third, the rupture of the vascularized bullae. Some patients were found to have abnormal vascularization between the lung and parietal pleura [3]. TB, like other chronic infections, can cause abnormal vascularization connecting the parietal pleura and chest wall, leading to aneurysms and intrathoracic hemorrhage if the vessels rupture. This appears to be the explanation for the massive bleeding in two of the cases (Cases 1 and 3). Soo-Kim et al. reported their 12-year experience with SHP [4]. They showed that pleural adhesions after TB do not cause SHP because they are thick and extensive. In two of our cases, very little thickening was observed in the pleura, which could be easily decorticated, whereas SHP was caused by cavity perforation in one of our patients.

SHP cannot compensate for even minimal bleeding because the lung is collapsed, which can lead to severe blood loss. Tube thoracostomy should be the first choice in the treatment of SHP. Studies have demonstrated that only tube thoracostomy with conservative treatment is enough for the treatment of SHP in some patients [3]. Fluid and blood replacement, where necessary, should be started at the stabilization of hemodynamics, the patient should be closely followed, and the need for emergency surgical intervention should be considered [6]. Persistent hemothorax ≥200 mL/h for consecutive 3 hours, persistent air leak, impaired lung expansion, and empyema are indications for thoracotomy or VATS [7].

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CONFLICTS OF INTEREST

The authors declare that they have no conflict of interest

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