A case of bleomycin-induced lung toxicity

Cuneyt Tetikkurt¹, Deniz Ongel¹, Seza Tetikkurt²

¹ Department of Pulmonary Medicine, Cerrahpasa Medical Faculty, Istanbul University
² Department of Pathology, Bagcılar Training and Research Hospital, University of Health Sciences, Istanbul, Turkey

Abstract

A 64-year-old female was admitted for dry cough, dyspnea, fever, loss of appetite, and weight loss. Past medical history revealed scoliosis, cholecystectomy, and Hodgkin lymphoma. ABG values were: pH: 7.42, pCO₂: 40.2 mm Hg, pO₂: 61.4 mm Hg. Chest CT showed cystic lesions, emphysema, ground glass, and reticular opacities. ABG values worsened under 8L/min nasal oxygen. The patient underwent bilevel positive airway pressure (BiPAP) and methylprednisolone 60 mg/day bid was commenced. The final diagnosis was respiratory insufficiency due to bleomycin toxicity. The patient deceased on the sixth day after transfer to the intensive care unit. Bleomycin is an effective chemotherapeutic agent used for Hodgkin lymphoma treatment. It causes significant lung toxicity in half of the patients. Clinicians should always remember that bleomycin toxicity may lead to fatal complications in patients with comorbid conditions. We present this case to remark the possible consequences of bleomycin toxicity and the precautions taken to preclude bleomycin-induced pulmonary complications are discussed.

Corresponding author: Prof. Cuneyt Tetikkurt, Department of Pulmonary Medicine, Cerrahpasa Medical Faculty, Istanbul University, Tanzimat Sok. Serkan Apt. No:3/16, 34728 Caddebostan, Istanbul, Turkey.
Tel. +90.216.3601977 - Mobile: +90.532.3810900 - Fax: +90.212.5870217.
E-mail: tetikkurt@gmail.com

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Introduction

Bleomycin is a chemotherapeutic agent that was isolated from a strain of Streptomyces verticillus [1,2]. It is frequently used to treat Hodgkin’s lymphoma and germinative tumors. The paramount drawback of bleomycin is its high potential to cause pulmonary toxicity that may progress to pulmonary fibrosis occurs in approximately 10% of the patients [2-6]. Other less common side effects include hypersensitivity and organizing pneumonitis [5-7]. The pathogenesis of bleomycin-induced lung injury is not entirely known. Likely components of bleomycin lung injury include oxidative damage, deficiency of bleomycin hydrolase, genetic predisposition, and inflammatory cytokines. Chronic fibrotic response of bleomycin induced lung injury is due to the loss of bleomycin hydrolase activity [8]. Influence of genetic factors for bleomycin-induced lung injury is suggested by the variable clinical profile of bleomycin sensitivity among patients [9-11]. Diagnosis is determined by integration of symptoms, clinical and radiologic findings. The clinical picture may be complicated and enigmatic that requires a thorough evaluation for differential diagnosis.

The potential of bleomycin lung toxicity is well known but the extent of this complication is not well authenticated. Oxidative damage, deficiency of bleomycin hydrolase, genetic predisposition and inflammatory cytokines are the hallmark pathogenetic mechanisms of bleomycin toxicity. This case report will focus on the potential avertible factors associated with the bleomycin-induced lung toxicity and the patient profile that would forestall the severe or fatal complications of bleomycin-induced lung injury by disputing the clinical precautions that would prevent the potential pulmonary side effects of bleomycin.

Case Report

A 64-year-old female patient was admitted for the evaluation of dry cough, dyspnea, fever, loss of appetite, and weight loss of six kilograms during the previous two months. Personal history revealed scoliosis, laparoscopic cholecystectomy, and Hodgkin lymphoma that was treated with doxorubicin, bleomycin, vinblastine, dacarbazine (ABVD) and radiotherapy six months ago. The BLMH A1450G genotype distribution was homozygous wild-type A/G genotype. She was a current smoker with a 50 p/ year smoking history. At initial admission she was tachypneic with a respiratory rate of 22/min. Blood pressure was 140/70 mm Hg. Bilateral fine rales were present over the middle and lower lung zones. ECG showed sinus tachycardia of 112/min. WBC: 6.9X10³/mm³, RBC: 3.6X10¹²/mm³, Hgb: 10.6 mg/dl, Htc: 32.7 PLT: 296X10³/mm³, Lymph: 0.8X10³/mm³, Eos: %3.9. Creatinine: 0.7mg/dl, AST: 10 IU/L, ALT: 14.9 IU/L, LDH: 165 IU/L, Na: 142 mmol/L, K: 4.3 mmol/L, Ca: 8.2 mg/dl, and CRP: 2.4 mg/dl. Arterial blood gas (ABG) values on admission were: pH: 7.42, pCO₂: 40.2 mm Hg, and pO₂: 61.4 mm Hg. Chest x-ray showed interstitial pattern and ground glass opacities at the middle and lower lung zones. ECG showed sinus tachycardia of 112/min. WBC: 6.9X10³/mm³, RBC: 3.6X10¹²/mm³, Hgb: 10.6 mg/dl, Htc: 32.7 PLT: 296X10³/mm³, Lymph: 0.8X10³/mm³, Eos: %3.9. Creatinine: 0.7mg/dl, AST: 10 IU/L, ALT: 14.9 IU/L, LDH: 165 IU/L, Na: 142 mmol/L, K: 4.3 mmol/L, Ca: 8.2 mg/dl, and CRP: 2.4 mg/dl. Arterial blood gas (ABG) values on admission were: pH: 7.42, pCO₂: 40.2 mm Hg, and pO₂: 61.4 mm Hg. Chest x-ray showed interstitial pattern and ground glass opacities at the middle and lower lung zones. ECG showed sinus tachycardia of 112/min. WBC: 6.9X10³/mm³, RBC: 3.6X10¹²/mm³, Hgb: 10.6 mg/dl, Htc: 32.7 PLT: 296X10³/mm³, Lymph: 0.8X10³/mm³, Eos: %3.9. Creatinine: 0.7mg/dl, AST: 10 IU/L, ALT: 14.9 IU/L, LDH: 165 IU/L, Na: 142 mmol/L, K: 4.3 mmol/L, Ca: 8.2 mg/dl, and CRP: 2.4 mg/dl. Arterial blood gas (ABG) values on admission were: pH: 7.42, pCO₂: 40.2 mm Hg, and pO₂: 61.4 mm Hg. Chest x-ray showed interstitial pattern and ground glass opacities at the middle and lower lung zones. ECG showed sinus tachycardia of 112/min. WBC: 6.9X10³/mm³, RBC: 3.6X10¹²/mm³, Hgb: 10.6 mg/dl, Htc: 32.7 PLT: 296X10³/mm³, Lymph: 0.8X10³/mm³, Eos: %3.9. Creatinine: 0.7mg/dl, AST: 10 IU/L, ALT: 14.9 IU/L, LDH: 165 IU/L, Na: 142 mmol/L, K: 4.3 mmol/L, Ca: 8.2 mg/dl, and CRP: 2.4 mg/dl. Arterial blood gas (ABG) values on admission were: pH: 7.42, pCO₂: 40.2 mm Hg, and pO₂: 61.4 mm Hg. Chest x-ray showed interstitial pattern and ground glass opacities at the middle and lower lung zones.
lung zones (Figure 1). CT revealed diffuse cystic lesions, ground glass opacities, traction bronchiectasis, and reticular opacities in both lungs (Figures 2 and 3). Sputum smear, cytology, and culture for bacteria, mycobacteria, and fungus were negative. Anti-histone antibody was 4.6 units. Following ABG values in the second day were as follows: pH: 7.38, pO2: 55.2 mm Hg, and pCO2: 38.5 mm Hg while the patient received nasal 8L/min oxygen. BiPAP was started and intravenous methylprednisolone 60 mg/day bid was commenced. The ABG values on the third day were as follows: pH: 7.41, pCO2: 49.3 mm Hg, pO2: 49.6 mm Hg and pH: 7.40, pO2: 50.1 mm Hg, pCO2: 49.3 mm Hg. The patient was transferred to the intensive care unit and mechanical ventilation was started. Final diagnosis was respiratory insufficiency due to bleomycin lung toxicity. The patient deceased on the sixth day of her admission in the intensive care unit.

Discussion

Bleomycin is a chemotherapeutic agent that was isolated from a strain of Streptomyces verticillus in 1966 [1]. The drug is frequently used to treat a variety of malignancies, predominantly germ cell tumors and Hodgkin lymphoma since it does not lead to myelosuppression. The major limitation of bleomycin treatment is the potential for life-threatening interstitial pulmonary fibrosis in up to ten percent of the patients receiving the drug [2-4]. Other less common forms of lung injury such as organizing pneumonia and hypersensitivity pneumonitis may also occur [3]. Risk factors for bleomycin-induced lung injury include cumulative dose, low glomerular filtration rate (GFR), older age, supplemental oxygen exposure, bolus drug delivery, and previous lung disease [5-7,12]. Of these, cumulative dose and reduced renal function are the most well-established risk factors [5,13]. Lung toxicity occurs sporadically at doses less than 450 IU but at doses above this range the incidence rises steeply [5,6]. Smoking is another risk factor for bleomycin-induced pulmonary toxicity [14]. Other suggested risk factors include dose rate, combination with other drugs, e.g. cisplatin, radiotherapy and the use of growth factors [2-4, 15,16]. Since bleomycin itself is a major potential risk factor for lung injury, combined bleomycin and radiotherapy has further contributed to the bleomycin-induced lung injury in our patient.

Our patient was 64 years-old, had a heavy smoking history with a silent chronic obstructive pulmonary disease, received radiotherapy together with a high dose bleomycin. The patient also carried a genetic variation in bleomycin hydrolase gene that was identified after bleomycin lung toxicity had developed. The final diagnosis was pulmonary fibrosis caused by bleomycin that lead to respiratory insufficiency and patient’s death. The clinical endpoint of this patient gives rise to the conclusion that patients who undergo bleomycin treatment must be thoroughly evaluated in regard to their pretreatment pulmonary status. Patient history should be exact, must include smoking history and previous lung diseases in a detailed manner. Besides chest x-ray, these patients should undergo a computed tomographic evaluation of the thorax to denote the lung parenchyma for chronic obstructive pulmonary disease, cystic lesions, fibrotic changes, old scars and

Figure 1. Chest x-ray revealing interstitial pattern and ground glass opacities at the middle and lower lung zones.

Figure 2. Chest CT showing bilateral diffuse cystic emphysema, ground-glass opacities, traction bronchiectasis, and interstitial fibrosis.

Figure 3. Chest CT showing bilateral diffuse cystic emphysema, ground-glass opacities, traction bronchiectasis, fibrotic bands, and interstitial fibrosis.
any other probable occult parenchymal lesions or airway diseases. Pulmonary function tests, diffusion capacity and exercise testing should also be performed in equivocal situations. To avoid the combined potential toxicity of radiotherapy and bleomycin induced lung toxicity, these two treatments should be administered separately with a maximum available time interval.

The lung toxicity of bleomycin is well known. The only way to prevent the potential and fatal hazards of this useful drug lies in an elaborate and a meticulous patient evaluation, which is the primary task of the clinician. These patients should be consulted with a pulmonologist for the possible existence of any previous or occult lung disease. Recent data suggests that the overall risk of a clinically apparent severe bleomycin-induced lung injury is low and pulmonary toxicity is reversible without long-term sequela in most of the patients [17,18]. Danish Testicular Cancer database and Lauritzen revealed that only a minority of bleomycin treated patients including those who discontinued bleomycin due to a decrease in carbon monoxide diffusing capacity (DLCO), suffered long-term restrictive disease or obstructive lung disease at five years of follow-up [19,20]. Depending on the clinical criteria and the cumulative bleomycin dose, rates of bleomycin-induced pulmonary toxicity in adults receiving ABVD for Hodgkin lymphoma range from 10 to 53%, and rates of fatal pulmonary toxicity are 4 to 5% [13,21], while long-term functional respiratory impairment is observed in approximately 15 to 18% of the patients [22-25]. The mechanism and pathogenesis of bleomycin-induced lung injury is not fully understood. Oxidative damage, bleomycin hydrolyase deficiency, inflammatory cytokines, genetic susceptibility and most importantly the patient related factors as our patient's case, are the hallmark of this syndrome. Whether the risk is minimum or not, the clinicians should pay utmost attention to the jeopardy of bleomycin toxicity, if at all. A comprehensive and an inclusive identification protocol must be used to identify the patients that carry a probable risk factor for pulmonary bleomycin toxicity.

Conclusions

Our patient had many potential factors that constitute a severe risk for the bleomycin-induced lung toxicity including genetic variation in bleomycin hydrolase gene, old age, smoking history, occult chronic obstructive pulmonary disease, high bleomycin dose and combined radiotherapy. Computed tomography performed at admission revealed chronic obstructive pulmonary lung disease related parenchymal lesions. For patients carrying one or more risk factors for bleomycin-induced pulmonary toxicity, treatment with this agent should be withheld and bleomycin must be replaced by another drug if possible, to prevent the future pulmonary complications of bleomycin toxicity that may be fatal. If treatment with bleomycin is a requisite for the patient, a meticulous clinical search for the presence of a silent or an occult lung disease must be done, while a close and a detailed follow-up should be performed in such patients to cease treatment on time to prevent bleomycin-induced lung toxicity. Asymptomatic patients should also be screened for inherent risk factors that may cause bleomycin-induced pulmonary side effects. We suggest that individuals who are to be treated with bleomycin should undergo a comprehensive and an algorithmic screening surveillance to detect associated risk factors or occult lung disease and to determine the probability of bleomycin-induced lung injury to preclude the severe toxic pulmonary effects of bleomycin treatment. Clinicians should be aware of the bleomycin-induced lung toxicity and identify the possible peril factors before life-threatening treatment complications develop, especially in patients that carry an identifiable risk factor for bleomycin treatment.

References


