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ÖZ

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INTRODUCTION

Allergic bronchopulmonary aspergillosis (ABPA) is a hypersensitivity reaction usually against *Aspergillus fumigatus* in patients with asthma or cystic fibrosis [1]. It is reported rarely in patients without asthma [2]. ABPA appears in 1 to 2% of patients with asthma and 2 to 15% of patients with cystic fibrosis during the course of the disease [3,4]. A recent meta-analysis demonstrated a prevalence of ABPA in asthma of 12.9% [5]. If ABPA is unrecognized or poorly treated, airway destruction, bronchiectasis, and/or pulmonary fibrosis result in significant morbidity and mortality [1]. Because of the indolent nature of the disease, the first step of diagnosis is suspicion [6]. Radiologic appearance may be misdiagnosed as tuberculosis and malignancy. The aim of this case report is to increase knowledge about ABPA among physicians and emphasize that it can occur with atypical findings.

CASE REPORT

A 31-year-old female presented with chronic cough. She had been diagnosed as smear negative pulmonary tuberculosis eight months ago and treated. Because of lack of response to the treatment, and the suspicious pulmonary mass-like lesion, she had been referred to our clinic. She had no symptoms except for dry cough. She didn’t have any atopic history except usage of asthma drug for one month, two years ago. On physical examination, crackles heard on the left anterior lung field. Chest X-ray showed bilaterally hilar enlargement, nodular opacity on the left upper zone, and infiltrations on the right upper zone and left paracardiac region. High density mass-like lesion with infiltration on the left upper lobe and paracardiac region, and a tubular density on the right upper lobe were observed on chest computed tomography (CT) (Figure 1). There was no abnormality on bronchoscopic examination except for tenacious mucoid secretions from both bronchial trees. To differentiate lesions from vascular anomaly, CT angiography of the thorax was performed. On CT angiography, characteristics of lesions have changed, atelectasis and bronchiectasis appeared on the left upper lobe, and there was a glove finger appearance on the right upper lobe (Figure 2). These findings with eosinophilia (2452/mm³) suggested ABPA although there was no asthma diagnosis. Total IgE level was 28,700 IU/mL. Skin prick test and specific IgE for *A. fumigatus* were positive. In sputum culture, *Aspergillus niger* has grown. Pulmonary function tests were performed and showed reversible mild airway obstruction. The patient was diagnosed as ABPA and treatment with prednisolon 40 mg per day and itraconazole 400 mg per day was started. Sweat chloride value was 40 mmol/L. CFTR gene mutation was studied and no mutation was found. After 15 days, cough disappeared, chest X-ray findings mostly reve-

Figure 1. High density mass-like lesion with infiltration on the left upper lobe and paracardiac region, and tubular density on the right upper lobe.
aled (Figure 3) and total IgE level decreased to 5000 IU/mL. During follow up, IgE level was about 1000 IU/mL. Itraconazole was stopped at second month because of side effects. Steroid doses were gradually tapered, and stopped at sixth month.

**DISCUSSION**

In asthma patients, ABPA should be kept in mind if the disease is uncontrolled despite appropriate treatment. ABPA can rarely be seen in non-asthmatic patients or both asthma and ABPA can be diagnosed at the same time [7]. If the patient has no asthma diagnosis, it is likely that these cases are often misdiagnosed initially as bronchogenic carcinoma or pulmonary tuberculosis [7]. Prasad et al. reported that, 91% of patients had been misdiagnosed as pulmonary tuberculosis before the diagnosis of ABPA [8]. Our patient was given asthma medicine two years ago, but on admission, she didn’t have asthma symptoms and signs and she was misdiagnosed as tuberculosis.

ABPA cases usually present with recurrent asthma attacks, expectoration of brownish mucus plugs, hemoptysis or systemic symptoms including fever, loss of appetit, and malaise. Our patient had atypical clinical history, because her unique complaint was dry cough. The finding of tenacious mucus plugs on bronchoscopic examination can be seen in ABPA. *A. niger* has grown in sputum cultur for fungus. This is accepted as a minor criterion for diagnosis. Because of positive results against *A. fumigatus* in skin prick test and serum specific IgE test, we accepted the cause as *A. fumigatus*.

Pulmonary CT angiography was performed because of the suspected vascular anomaly. The findings of atelectasis and central bronchiectasis and the changing infiltrates with eosinophilia suggested ABPA despite the patient was non-asthmatic.

Eight major criteria were defined for ABPA diagnosis:

1. Asthma,
2. Immediate skin test reactivity to *A. fumigatus*,
3. Elevated serum IgE levels,
4. Transient or fixed pulmonary infiltrates,
5. Central bronchiectasis,
6. Peripheral blood eosinophilia,
7. Elevated specific IgE and/or IgG for *A. fumigatus*,
8. Presence of serum precipitins against *Aspergillus* antigen. However, it is not certain that at least how many criteria are needed for the diagnosis[3]. Our patient had six criteria. She didn’t have typical asthma findings, however there are non-asthmatic ABPA cases in the literature, even rare. Additionally, we couldn’t test the precipitins against *Aspergillus* antigen because of inavailability.

A normal serum IgE level excludes ABPA[7]. Total serum IgE levels are above 1000 ng/mL (420 kU/L) and in acute cases can be above 20,000 ng/mL. The level of total serum IgE was as high as 28,700 IU/mL in our patient. Despite treatment, IgE levels can remain high. In this situation, it should be kept over a lower limit which defined for each patient. Because of it is the most important parameter for the management of the patients, total IgE should be monitored in the first year and then annually[7]. The total IgE level of our patient did not decrease to normal limit, but drawn a plateau over 1000 IU/mL.

ABPA can mimic pulmonary mass rarely[9]. High-attenuation density within these masses can help narrow the differential diagnosis[9]. This entity has been rarely described in the literature mainly as case reports[10,11]. Agarwal et al. reported 18.7% incidence of HAM out of 155 ABPA patients[12]. Currently, the finding of hyperdense mucus on chest CT is considered a characteristic, but not pathognomonic, finding for ABPA[13]. In our case, when evaluated retrospectively, the high density tubular lesion on first CT examination was defined as HAM. The term of HAM is used for mucus plugs that is seen more intense than skeletal muscle on chest CT.

In conclusion, ABPA is not always appear with severe asthma or typical clinical signs. Early diagnosis and treatment are of great importance in terms of prevention of complications and end stage lung disease. Therefore, in patients without asthma, eosinophilia and changing infiltrations on roentgenogram with HAM on CT should suggest ABPA and total IgE levels should be studied.

**REFERENCES**