Objective: Although pulmonary aspergillosis (PA) is often seen in patients with immunodeficiency, it also identifies a number of clinical diseases that cause morbidity and mortality, which can also occur in patients with immunodeficiency. The aim of this study was to investigate the disease patterns, demographic and clinical features, treatment and prognosis of pulmonary aspergillosis in immunocompetent patients.

Method: Our study is a descriptive cross-sectional study in which the data of immunocompetent patients diagnosed with PA in our hospital between January 2010 and December 2019 were evaluated retrospectively. Patients over the age of 18 with a diagnosis of PA were included in the study. Demographic and clinical features, radiological findings, diagnosis and treatment methods and prognoses of the patients were recorded.

Results: 8 of the patients were female, 3 were male and the mean age was 61 (53 ± 69). Nine patients were diagnosed with chronic pulmonary aspergillosis and two patients were diagnosed with IPA. Four of the patient diagnosed with KPA were chronic cavitary pulmonary aspergillosis, one was chronic fibrosing pulmonary aspergillosis, two were aspergilloma and two were aspergillus nodules. The most common comorbidity was diabetes mellitus (45.4%) and in total, 72.7% of patients had chronic lung disease such as COPD-asthma, tuberculosis, bronchiectasis. Cough, sputum and hemoptysis (90%, 72.7%, 45.4%, respectively) were the most common symptoms. The most common radiological finding on thorax tomography was crescent sign-cavity (72.7%). The diagnostic method used was mostly bronchoalveolar lavage and wedge resection/ lobectomy (45.4%, respectively). It was observed that eight (72.7%) patients were given voriconazole and three (27.2%) other patients surgical procedure performed and one patient was lost after surgery.

Conclusions: Although pulmonary aspergillosis is more frequently seen in immunocompromised patients with malignancy, solid organ and hematological transplants, it has now begun to be seen in immune competent patients. The majority of our patients were middle-aged, women and smokers, people with comorbidity such as DM and chronic lung disease. Therefore, we would like to point out that pulmonary aspergillosis should be kept in mind among differential diagnoses, especially in patients with immunocompetent with comorbidity such as chronic lung disease and DM.

Keywords: Pulmonary Aspergillosis, immunocompetent patients, disease patterns, demographic, clinical features, prognosis
İmmün kompetan Hastalarda Pulmoner Aspergillozis

Öz

Amaç: Pulmoner aspergillozis (PA) sıklıkla immün yetmezlikli hastalarda görülse de immün kompetan hastalarda da ortaya çıkabildi. Amaçımız; immün kompetan hastalarda pulmoner aspergillozisin hastalıktan paternlerini, demografik ve klinik özellikleri, tedavi yöntemlerini ve prognozunu araştırmaktı.

Yöntemler: Çalışmamız Ocak 2010 ve Aralık 2019 tarihleri arasında hastanemizde PA tanısı alan immün kompetan hastaların retrospektif olarak değerlendirildiği tanımlayıcı kesitsel bir çalışmadır. 18 yaş üstü PA tanısı alan hastaların demografik verileri, klinik özellikleri, radyolojik bulguları, tanı ve tedavi yöntemleri ile prognozları retrospektif olarak incelendi.

Sonuçlar: Hastaların 8’si kadın 3’si erkek ve yaş ortalaması 61 (53±69)' idi. Hastaların 9’u Kronik pulmoner aspergilloz (KPA) ve ikisi invaziv PA tanısı aldı. KPA tanısı alan hastaların 4’ü kronik kaviter pulmoner aspergillozis, 1’i kronik fibroz pulmoner aspergilloz, 2’si aspergilloma, diğer 2’si aspergilloz nodülü idi. En sık saptanan komorbidite diabetes mellitus (%45,4) olup hastaların %72,7’sinde koah-astım, tüberküloz, bronşektazi gibi kronik kaviter pulmoner aspergilloz, %45,4 en sık saptanan semptomlardı. Toraks tomografisinde en sık saptanan radyolojik bulgünün ise hilal belirtisi-kavite (%72,7) olduğu belirlendi. Uygulanan tanı yöntemleri sırasıyla bronkoalveolar lavaj (BAL) ve wedge rezeksiyon/ lobektomi (%45,4) olarak saptandı. Tedavide sekiz (%72,7) hastaya vorikonazol verildiği, diğer üç (%27,2) hastaya cerrahi işlem yapıldığı ve bir hastanın ise cerrahi sonrası kaybedildiği görüldü.

Tartışma: Pulmoner aspergilloz daha sik olarak bağışıklığı zayıflamış kanser, solid organ ve hematolojik nakil hastalarda görülmekle birlikte günümüzde immün kompetan hastalarda da görülmeye başlamıştır. Hastalarımızın çoğunluğunu; orta yaş, kadın, sigara içen, altta yatan DM ve kronik aşıklığı hastaları oluşturmaktadır. Bu nedenle özellikle kronik aşıklığı ve DM gibi altta yatan hastalığı olan immün kompetan hastalarda pulmoner aspergillozun ayırıcı tanılar arasında akılda tutulması gerekiyor.

Anahtar kelimeler: Pulmoner aspergillozis, immün kompetan hasta, hastalık paternleri, demografik, klinik özellikleri, prognoz.

INTRODUCTION

Aspergillus spp. It is a widely isolated fungus species both in the external environment (soil, plant residues) and in many indoor environments including hospitals. Aspergillus causes various clinical syndromes. Host immunity is of great importance in the symptoms, clinical course and disease severity of aspergillus infections. The interaction between the pathogen and host immune dysfunction or hyperactivity determines which clinical syndrome is more likely to develop. Nowadays, invasive aspergillosis (IA) can often be seen in immunocompromised neutropic patients as well as increasingly in non-neutropic hosts (such as organ transplant patients, critically ill patients, and patients receiving steroids). Görük et al. reported that IA developed in 15% of febrile neutropenic patients with hematological malignancies. IA can be seen in immunocompetent hosts that a re heavily exposed to fungal spores. Chronic pulmonary aspergillosis (CPA) is a type of disease that causes severe lung damage and is mostly seen in patients with mild immunodeficiency or an underlying lung chronic lung disease (COPD, previous or concurrent tuberculosis (TB) or non-TB mycobacterial disease).

METHOD

This is a descriptive cross-sectional study in which the data of immunocompetent patients diagnosed with pulmonary aspergillosis in our hospital between January 2010 and December 2019 were evaluated retrospectively. Our aim was to determine the disease patterns, demographic and clinical features, and prognosis of pulmonary aspergillosis in immunocompetent patients. Patients over the age of 18 with a diagnosis of PA were included in the study. Data of patients were obtained from their electronic files. Demographic, clinical, radiological features, comorbidities, diagnosis and treatment methods of the patients were recorded. Ethics committee
Approval was obtained for our study with the date of 06.10.2020 and no KA 20/368.

**Study population and case definitions**

Pulmonary aspergillosis diagnosis was established according to diagnostic criteria of international guidelines: The diagnosis of "proven IPA (invasive pulmonary aspergillosis)" was made by demonstrating hiphya compatible with aspergillus in samples taken by lung biopsy or needle aspiration and/or identify of any upper and lower respiratory secretion from the culture.

We did not have any patients meeting the criteria of “possible IPA” and “possible IPA”2.

CPA; mostly consists of a series of disease patents seen in patients with underlying lung disease (3). These clinical conditions are defined as chronic cavity pulmonary aspergillosis (CCPA) - untreated and if progress occurs CFAP (chronic fibrous pulmonary aspergillosis), aspergilloma, Aspergillus nodules (AN). The diagnosis of CPA was established according to the following criteria of The European Society for Clinical Microbiology and Infectious Diseases (ESCMID), the European Respiratory Society (ERS) and the European Confederation of Medical Mycology (ECMM) 2017: (i) one or more nodules with or without fungal ball on thorax CT ii) direct exclusion of Aspergillus spp by biopsy culture or microscopy or exclusion of an immunological response to Aspergillus spp (iii) all present for at least 3 months6,7.

In CCPA; there are one or several thick-walled cavities (when dense fungus hyphae intertwine with collections) with or without fungus balls that develop in a few months8. If CCPA is left untreated, it can progress, and intense fibrotic destruction occurs in lung tissue. This condition is classified as chronic fibrosing pulmonary aspergillosis (CFPA) leading to respiratory failure9. Aspergilloma is a fungus ball composed of aspergillus hyphae, fibrin, mucus, inflammatory cells degenerate blood and epithelial elements in the pre-existing cavity10. Aspergillus nodules are generally defined as solid nodules detected incidentally on CT, which may be single or multiple, especially those that can be confused with conditions such as lung cancer11.

**Study case exclusion criteria**

1- Aspergillus spp. colonization (without aspergillus infection) 2- Severe immunsuppressive patients with solid organ transplantation, hematopoietic stem cell transplantation and receiving chemotherapy for malignancy. 3- In addition, we did not have any patients with allergic bronchopulmonary aspergillosis (ABPA) with follow-up during the screening period for the study.

The antifungal treatment response of our patients was evaluated according to the criteria in the ESCMID / ECMM / ERS clinical guidelines6,7.

**Microbiological processes**

All upper and lower respiratory tract samples were cultured in sabouraud dextrose agar with and without antibiotics after direct microscopic examination. These were incubated in incubators at 25 °C and 37 °C for 3 weeks. Identification of a colony with mold growth was carried out at species level according to macroscopic and microscopic morphology12. Serum galactomannan (GM) antigen level was determined with Platelia Aspergillus Galactomannan EIA kits. For GM, a rate 0.5 ng / mL was considered positive13.

**Statistical Analysis**

Statistical analysis of the data was done using SPSS 25.0 package program. Categorical measurements were summarized as number and percentage, whereas continuous measurements were summarized as mean and standard deviation (median and minimum-maximum, where necessary).
Table I: Diagnostic criteria for invasive fungal disease

| Host Criteria                                                                 |
|-------------------------------------------------------------------------------|
| Neutropenia (> 10 days)                                                      |
| Persistent fever that does not respond to antibiotics (> 96 hours)           |
| Body temperature > 38 °C or > 36 °C in the presence of preparative factors (neutropenia, chemotherapy) |
| GVHD                                                                          |
| Corticosteroid use (> 3 weeks; within the last 60 days)                       |

| Microbiological Criteria                                                    |
|----------------------------------------------------------------------------|
| Histopathology / microscopy (+)                                              |
| Culture (+) (BAL / sputum / sinus aspiration)                               |
| Galactomannan antigen (+) (BAL / BOS / > 2 blood samples)                    |

| Clinical Criteria                                                          |
|----------------------------------------------------------------------------|
| Major Clinical Criteria (Computed Tomography)                               |
| Halo sign                                                                   |
| Crescent sign, consolidation and cavity                                     |

| Minor Clinical Criteria                                                    |
|----------------------------------------------------------------------------|
| Respiratory infection (cough, pleuritic chest pain, hemoptysis)             |
| Pleural frotman                                                             |
| Pleural effusion                                                           |
| Infiltration in addition to major radiological criteria                     |

GCVH: Graft versus host disease, BAL: Bronchoalveolar lavage

Table II: IPA (Invasive pulmonary aspergillosis) diagnose

| Proven IPA                                                                 |
|----------------------------------------------------------------------------|
| Histopathology / cytopathology (+) Example: Biopsy / needle aspiration     |
| Culture (+) Example: Sterile body area (excluding urine and mucosa)        |

| Strongly Probable IPA                                                      |
|----------------------------------------------------------------------------|
| >1 host + 1 microbiological criterion + 1 major (or 2 minor) clinical criterion |

| Possible IPA                                                              |
|----------------------------------------------------------------------------|
| >1 host criterion + 1 microbiological or 1 major (or 2 minor) clinical criterion |

A total of 11 patients: 9 (81.1%) were diagnosed with CPA [4 (56.3%) CCPA, 1 (9.1%) CFPA 2 (18.1%) aspergilloma, 2 (18.1%) aspergillus nodule] and the other 2 (18.1%) were diagnosed as IPA. Eight (72.7%) were female, 3 (27.3%) were male and the mean age was 61 (53-69). Eight patients (72.7%) were smoker. The most common comorbidity was DM 5 (45.4%) patients, and it was also found in 8 (72.7%) patients in total (27.2%, 27.2%, 18.1%), respectively, with chronic respiratory disease. Cough 10 (90%) and sputum 8 (72.7%) were the most common symptoms. Hemothysis was also remarkably 5 (45.4%). Three patients with malignancy (skin-basal cell carcinoma, thyroid follicular carcinoma) did not receive chemotherapy.
Table III: Patient diagnoses, demographic and clinical features

| Patient diagnoses                  | N: 11 (%) |
|-----------------------------------|-----------|
| KPA                               | 9 (81.1)  |
| KKPA                              | 4 (36.3)  |
| KFPA                              | 1 (9.1)   |
| Aspergillum                        | 2 (18.1)  |
| Asperillus nodule (AN)             | 2 (18.1)  |
| IPA                               | 2 (18.1)  |
| Gender M / F                       | 8 (72.7) / 3 (27.3) |
| Mean age                           | 61 (53±69) |
| Smoker                             | 8 (72.7)  |
| Ex-smoker                          | 1 (9.1)   |
| Non-smoker                         | 2 (18.1)  |

Table IV: Patient thorax computed tomography (CT) findings, diagnosis and treatment method

| Thorax CT Findings                  | N: 11 (%) |
|-------------------------------------|-----------|
| Crescent sign, cavity               | 8 (72.7)  |
| One-sided nodule                    | 2 (18.2)  |
| Bilateral nodules                   | 1 (9.1)   |
| Halo sign                           | -         |

Diagnostic Method

| BAL (Bronchoalveolar lavage)         | 5 (45.4)  |
| Wegde resection                     | 5 (45.4)  |
| Bronchial biopsy                    | 1 (9.1)   |
| Transthoracic biopsy                | 1 (9.09)  |

Treatment Method

| 12 weeks                            | 6 (63.6)  |
| 8 weeks                             | 2 (18.2)  |
| Mortality                           | 1 (9.1)   |

Radiological findings, diagnosis and treatment methods of the patients are shown in Table IV. It was found that the disease was most commonly seen in thorax CT in 8 (72.7%) patients as crescent-cavity and 2 (18.2%) in the form of nodules. The diagnosis was most commonly made with bronchoalveolar lavage (BAL) in 5 (45.4%) patients, wedge resection / lobectomy in 5 (45.4%) and transthoracic needle biopsy in another patient. It was found that 8 (72.7%) patients had voriconazole as an antifungal drug and 3 (27.2%) patients underwent surgery. During follow-up, one (9.1%) patient was found to be lost.

Thorax CT images of the patients with pulmonary aspergillosis according to the disease type are shown in Table V.
Table V: Thorax CT images of the patients with pulmonary aspergillosis according to the disease type

| Disease types                           | Description                                                                                                                                                                                                 |
|----------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Aspergilloma                           | 60 years old woman patient. There is a history of previous pulmonary TB, with increased cough, sputum and recurrent hemoptysis for 4 months. In Thorax CT, it is seen that a cavity with a diameter of about 3 cm in the left lung upper lobe apicoposterior segment begins to form and consolidation begins to separate with a thin air ring (crescent symptom). Aspergillus hyphae was detected in pathology with wedge resection. |
| Chronic cavitary pulmonary aspergillosis (CCPA) | 55 years old male patient. Comorbidity: DM, RA, COPD He has had cough, weakness, and recurrent hemoptysis for 5 months. In thorax CT, there was a cavitary lesion with a fungus ball in the right lung upper lobe apicoposterior segment, with an axial diameter of 97 mm in the thin irregular wall, and emphysema in the left lung. Right upper lobectomy was performed due to massive hemoptysis and aspergillus hyphae was detected in pathology. |
| Invasive pulmonary aspergillosis (IPA): | 57 years old man. Comorbidity: COPD, CKD and DM. He applied with hemoptysis in thorax CT, there was a thick and irregular walled cavitary lesion with a sign of fungus and crescent in the posterior segment of the right upper lobe of the right lung. *Aspergillus fumigatus* grew in bronchoscopic lavage culture. BL galactomannan positive, as well as aspergillus hyphae were detected in the right upper lobectomy pathology. |
| Aspergillus nodule                      | 59 years old male has DM and smoker had dry cough for 5 months. In thorax CT, there were nodular lesions with 23x18 mm spiculo contours in the right lung upper lobe anterior segment. As a result of diagnostic wedge resection, aspergillus hyphae was detected in pathology. |

**DISCUSSION**

In this study, diagnosed patients with immunocompetent pulmonary aspergillosis were analyzed. All of the 11 patients 9 were KPA (4 KKPA, 1 KFPA. 2 aspergilloma, 2 aspergillus nodules) and the other 2 patients were IPA. CPA is a neglected pulmonary fungal infection that is estimated to affect more than 3 million people worldwide^{14,15}. It has been found that
patients with CPA are generally middle-aged and male. Our patient group was middle-aged (mean age 61), but 82.7% were women. The high female gender may be related to the single center and patient population of our study.

CCPA is the most common form of CPA, these cavities expand over time. In our study also the most common diagnosis was CCPA (36.3%) and cavities had large diameters. In our study, which was compatible with the literature, the most common diagnosis was KKPA (36.3%). The main risk factors are underlying diseases, especially causing structural damage to the lung; TB, non-TB mycobacteria (NTM) lung diseases, chronic obstructive pulmonary disease (COPD), emphysema, bronchiectasis, asthma, pneumothorax, sarcodosis, pneumonia and other chronic pulmonary diseases. Smith et al. revealed that comorbidity was the most common in tuberculosis and other common conditions in COPD and emphysema in 126 non-immunocompromised patients. As in the literature, it was determined that most of our patients had chronic lung diseases such as tuberculosis, COPD, asthma and bronchiectasis. In addition, 81.1% of patients smoked. Smoking rate of women is high in our country (13.3%). This may have contributed to the formation of chronic lung diseases in patients.

In previous years, it has been shown that the necrotic form of KPA may also occur in patients with mild immune deficiency due to connective tissue diseases such as DM, alcoholism, chronic liver and kidney disease, corticosteroid therapy, malnutrition or rheumatoid arthritis. In our study, the most common comorbidity in accordance with the literature was DM. In addition, these patients had other comorbidities such as chronic renal failure and COPD. We think that these underlying diseases may have contributed to the emergence of a fungal infection such as pulmonary aspergillosis by adversely affecting the immune system and defense mechanisms of the host.

Aspergillus and aspergillosis nodules are less severe forms of CPA. Aspergillus is a fungal ball that commonly occurs in the TB cavity, the underlying cavitary disease. In a study evaluating 544 patients with post-tuberculosis cavity, 11% had radiological findings of aspergilloma. Two of our 3 patients who were diagnosed with aspergilloma had a history of previous tuberculosis.

It is known that IPA is also seen in non-neutropenic immunocompetent patients without classical risk factors. Patients with severe COPD and critically ill patients have the highest risk in this group. There are several reasons for increasing sensitivity to IPA in COPD patients. These; changes in the lung structure, long-term use of corticosteroid therapy, frequent hospitalizations, antibiotic therapy, invasive applications, impaired mucociliary clearance and accompanying comorbid diseases such as diabetes mellitus and malnutrition. In a multicenter study excluding patients with classic risk factors from the USA, 77% of critically ill patients with IA (invasive aspergillus) were found to have taken high-dose steroids during their hospitalization. Both of our patients with IPA had severe COPD, DM, frequent hospital admissions, and long-term use of high doses of steroid.

Symptoms of CPA are nonspecific so diagnosis may be delayed. The most common symptom of the disease is productive cough; others are weight loss and dyspnea. It has been shown that hemoptysis frequently occurs in patients with aspergilloma and can be life-threatening. The most common symptoms consistent with the literature in our patients was cough, sputum and hemoptysis.

CPA appears with radiographic findings such as crescent sign-cavitation (with or without a fungus ball) pleural thickening, peri cavitary infiltrates in thorax CT. Typical finding for CCPA is consolidation in the upper lobes that
progresses to cavitation over time and causes volume loss. Simple aspergilloma occurs as a solid opacity in the upper lobe or as a cavity-creating appearance when the mass separates from the wall in the cavity. In our patients, the most common radiological finding in thorax CT was crescent sign-cavity (72.7%).

**Diagnosis for CPA:** clinical, radiological findings and compliance with diagnostic criteria such as demonstrating aspergillus infection (tissue biopsy and / or microbiological culture). Diagnosis of IPA is also quite difficult in the patient group without typical risk factors. The gold standard for diagnosis is biopsy of lung tissue and / or culture of positive Aspergillus spp. Infection has been proven in half of our patients with reproduction in BAL culture and in the other half by wedge resection / lobectomy and pathological aspergillus hyphae. The reason for the frequent use of surgery in diagnosis is that this method was used for diagnosis and treatment because almost half of our patients applied with serious hemoptysis.

The goal of treatment in CPA is to improve symptoms, reduce and / or prevent hemoptysis and lung fibrosis. Voriconazole or itraconazole is recommended as the first-line treatment option. Voriconazole is primarily recommended in IPA because it decreases mortality. The recommended duration of treatment is 12 weeks, depending on the clinical response, longer treatment may be required. 72.7% of our patients received voriconazole treatment for 12 weeks.

Even when CPA is treated, a 5-year mortality is around 20-50%. Hemoptysis is a complication that can be fatal in patients with CPA. Surgery may be considered in severe hemoptysis or treatment failure, but the rate of postoperative complications is high. In our study, 3 (27.2%) patients with moderate and severe hemoptysis were operated for diagnosis and treatment, and only one was lost postoperatively. Our other patients improved clinically and radiologically. Our results were determined in accordance with the studies in the literature.

Our study has some limitations. The study was single-centered, so the number of patients is relatively small and does not represent the general population. In addition, the reason why our study is a retrospective study was made according to the diagnostic criteria of international guidelines using the examinations performed on the patients. Our number of patients may be low due to the lack of appropriate diagnostic methods. When related literature was reviewed from our country, it was published in the form of case reports. Our study has the highest number of patients from our country, and we think it will contribute to the literature. There is a need to plan forward-looking, multicenter studies.

As a result, in this study, we have revealed the clinical patterns of pulmonary aspergillosis in immune composite patients. The majority of patients are middle-aged, female and smokers. The most common symptoms were productive cough and hemoptysis. All patients had comorbidities most frequently, such as DM and underlying chronic lung disease. The prognosis was quite good. We think that pulmonary aspergillosis, which we know to be seen in people who suppress the immune system, should be kept in mind among the options of diagnosis in immune competent individuals.

**Ethics Committee Approval:** Ethics committee approval was obtained for our study with the date of 06.10.2020 and no KA 20/368.

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