Rare Interstitial Pneumonia Due to the Use of Aripiprazol: A Case Report

Aripiprazol Kullanımına Bağlı Nadir Görülen İnterstisyel Pnömoni: Olgu Sunumu

Gulistan Alpagat, Ayse Baccioglu, Sumeyra Alan Yalim, Merve Poyraz, Betul Dumanoglu, Ayse Fusun Kalpaklioglu

Abstract

It is accepted that 2–3% of interstitial lung diseases (ILD) are due to drugs, and that 70% of drug-induced lung diseases are due to ILD. We present here the case of a 51-year-old female patient who had been taking aripiprazole for bipolar disorder for 3 years and had been undergoing asthma treatment for 2 years. ILD was considered based on the patient’s clinic, a restrictive pattern in the pulmonary function test (PFT), a decreased carbon monoxide diffusion test, laboratory results, radiological findings and biopsy. The possible etiologies of infection, aspiration, heart failure, pet feeding status and exposure to hazardous respiratory substances were excluded. The suspected drug was discontinued and oral corticosteroid treatment was started. Lung toxicity can develop with drug treatments, and so drug history should be questioned in detail and discontinued immediately in case of any doubt. ILD is rare, and if the drug is not suspected and continued, the disease may become chronic and progress to respiratory failure. We present this case to increase awareness of the disease.

Key words: Interstitial Lung Disease, aripiprazole, interstitial pneumonitis, steroid treatment.

Özet

İnterstisyel akciğer hastalıklarının (IAH) %2-3’unun ilaçlara bağlı olduğu ve ilaca bağlı akciğer hastalıklarının %70’inin IAHdan olduğu kabul edilmektedir. Elli bir yaşında kadın hasta 3 yıldır bipolar bozukluk için aripiprazol kullanmaktadır ve 2 yıldır astım tedavisi görmekteydi. Hastanın kliniği, solunum fonksiyon testinde (SFT) restriktif patern olması, azalmış karbon monoksid difüzyon testi, laboratuvar sonuçları, radyolojik bulgular ve biyopsi ile IAH düşünüldü. Olası etyolojiler: enfeksiyon, aspirasyon, kalp yetmezliği, evcil hayvan besteme durumu ve tehlike solunum maddelerine maruziyet dışlandı. Şüpheli ilaç kesilerek, oral kortikosteroid tedavisi başlandı. İlaç tedavileri ile akciğer toksisitesi gelişebilir. İlaç öyküsü ayrıntılı olarak sorgulanmalı ve şüphe durumunda derhal kesilmişdir. IAH nadir görülür; ilaca şüphelenilmese de devam edilirse hastalık kronikleşebilir ve solunum yetmezliğine ilerleyebilir. Hastalıklarla ilgili farkındalığı artırmak için bu olguyu sunmaya amaçladık.

Anahtar Sözcükler: Interstisyel Akciğer Hastalığı, aripiprazol, interstisyel pnömoni, steroid tedavi.

Department of Immunology and Allergy Disease, Kırıkkale University Faculty of Medicine, Kırıkkale, Turkey

Kırıkkale Üniversitesi Tip Fakültesi, İmmunoloji ve Allerji Hastalıkları, Kırıkkale

Submitted (Başvuru tarihi): 30.04.2021 Accepted (Kabul tarihi): 05.10.2021

Correspondence (İletişim): Gulistan Alpagat, Department of Immunology and Allergy Disease, Kırıkkale University Faculty of Medicine, Kırıkkale, Turkey

e-mail: gulistanalpagat16@gmail.com
Interstitial lung disease (ILD) is a heterogeneous group of diseases, the etiology of which can be influenced by many factors. It is accepted that 2–3% of ILDs are due to drugs, and ILDs account for 70% of drug-induced lung diseases (1). We present here a case that developed non-specific interstitial pneumonia (NSIP) after the regular use of aripiprazole for bipolar disorder.

CASE
A 51-year-old female patient was referred to the respiratory outpatient clinic with severe asthma, and complaints of dry cough and shortness of breath for about 2 years. The symptoms were of moderate severity, preventing such strenuous activities as exercise. She had been under treatment with high dose inhaled corticosteroid (ICS) and long-acting beta2-agonists (LABA) (salmeterol+fluticasone 50 mg/500 mg/day), montelukast-sodium (10 mg/day) and tiotropium (18 mcg/day), however her symptoms in recent months had been unresponsive. She had been taking aripiprazole for bipolar disorder for 3 years and levothyroxine for Hashimoto’s thyroiditis. The patient was an active smoker with 10 pack/years, but no alcohol or other substance use disorders. She had no recent travel history, pet ownership, occupational exposure or tuberculosis history. A physical examination revealed bilateral crakles and digital clubbing. The respiratory rate was 16/min, axillary temperature was 37°C, heart rate was 80/min, systemic blood pressure was 125/75 mmHg, oxygen (O2) and saturation at room air was 96%. Other system examinations were normal.

In laboratory tests, serum C-reactive protein was 4.1 mg/L, erythrocyte sedimentation rate was 41 mm/h, white blood cell was 7,500/mm3, hemoglobin was 11.2 g/dL and platelet count was 211,000/mm3. Serum autoimmune panel (ACE, ANA, anti-dsDNA, c-ANCA, p-ANCA, AMA, ASMA, anti-CCP, RF, ASA, anti-Ro, anti-La) resulted normal twice. Resting blood tests, including glucose, hepatic markers, renal and hepatic function tests were also within the normal range. Chest X-ray showed bilateral non-homogeneous infiltrations in the middle-low lung areas (Figure 1). High resolution computed tomography (HRCT) revealed mediastinal retrocardial lymphadenopathy (LAP) (18x11 mm), mosaic pattern, peripheral reticular density and patchy ground-glass opacities in the bilateral lower lung parenchyma with peribronchial thickening (Figure 2). Sputum acid-fast bacilli (ARB) samples resulted negative three times, as well as a sputum mycobacterium culture. In pulmonary function tests (PFT), a restrictive pattern with FEV1 33% (0.92L), FVC as 36% and FEV1/FVC 98% was noted. The increase in FEV1 reversibility was lower than 10% and 100 mL. A diffusion lung test with carbon-monoxyde (DLCO) was 50% and DLCO/VA was 110%. Cardiac failure was ruled out by a cardiologic consultation, normal echocardiography (ECHO) and a normal serum pro-brain natriuretic peptide (pro-BNP). To diagnose ILD, a fiber optic bronchoscopy (FOB) was performed, and no endobronchial lesion was observed. A bronchoalveolar lavage (BAL) analysis showed a normal cellular profile with a negativity of culture. Diagnostic video-assisted thoracic surgery (VATS) was performed to specify interstitial pneumonitis. A lung biopsy report identified “hypertrophy of smooth muscles in terminal bronchial structures, increase in the distance of the alveolar ductus, subpleural and alveolar septal thickening, increased vascular wall thickness, mononuclear inflammatory cell infiltration, including scattered lymphocytes and plasmocytes in the interstitial area with lymphoid aggregates patches, mild to moderate interstitial inflammation, and relatively homogeneous interstitial fibrosis”. Histochmical staining revealed staining in favor of connective tissue in the focal areas (Figure 3).

A diagnosis of NSIP was made because of restriction in PFT, decrease in DLCO, and histopathological findings of lung biopsy in accordance with radiological findings and clinical correlation. The possible etiologies of infection, aspiration, cardiac failure, pet ownership and negativity in the serum auto-immune panel were excluded. Collagen tissue diseases such as rheumatoid arthritis or scleroderma were absent, based on negative autoimmune blood markers, and rheumatologic consultation. The most common indicators of ILD were absent in the patient’s history, including long-term exposure to such hazardous inhaled materials as asbestos. Considering that the onset of symptoms occurred during aripiprazole use with an indication of bipolar disease, drug-related lung disease was considered.

Aripiprazol was discontinued as a suspicious drug, and an alternative treatment was suggested by psychiatry. For the treatment of NSIP, oral corticosteroid (OCS) was initiated at 0.5 mg/kg with prophylaxis of gastritis and pneumocystis carinii, given the presence of severe dyspnea and low PFT values. An ICS-LABA combination was continued for its steroid-sparing effect. After 2 weeks, the OCS dose was reduced to 16 mg, and discontinued after 11 months through slow tapering. During follow-up, clinical complaints decreased, PFT values increased.
(FEV1 as 47%-1.30L, FVC as 40%, FEV1/FVC as 123%), and radiologic improvement was noted. The patient has been under follow-up for 5 years, and has had four NSIP flares with hospitalization. She has used OCS seven times in 5 years. Even though aripiprazol was discontinued, a mild progression in FEV1 as 35% (0.96L), FVC as 32% and FEV1/FVC as 119% was seen, in contrast to the severe radiologic progression (Figure 4 and 5). Her dyspnea and cough became persistent, and she was prescribed long-term oxygen treatment. Radiologic images of the patient are presented in Figure 6 from before the start of OCS treatment, 6 months after starting and during the 5-year follow-up. The patient has not developed coronavirus disease.

**DISCUSSION**

NSIP occurs with the uniform enlargement of the alveolar septa along with fibrosis and inflammation (2). It affects men and women equally, and the average age of onset is 48. Shortness of breath and dry cough are the most common patient complaints, and a combination of ground-glass opacities, consolidation and irregular lines showing subpleural peripheral distribution are common radiological findings. Although these findings are characteristic, atypical UIP cases may have a similar appearance. The average life expectancy varies between 1.3 and 15 years (3). NSIP may be idiopathic or may develop as a complication of connective tissue disease, hypersensitivity pneumonia, drug-induced interstitial lung disease and diffuse alveolar injury (4).

This was a rare case of NSIP development due to aripiprazole – an atypical anti-psychotic agent that has been widely used for the management of schizophrenia and bipolar diseases. Its mechanism of action includes partial agonistic activity at the dopamine-D2, and serotonin 5-HT1A receptors, and antagonistic activity at the 5-HT2A receptors (5). Even though the drug is generally well-tolerated, it has some common adverse events, such as extrapyramidal symptoms and weight gain. Its best known pulmonary effect is non-cardiogenic pulmonary edema, which can be diagnosed based on its acute setting and typical ECHO findings. Pulmonary edema was ruled out in our patient based on a normal ECHO and normal pro-BNP, and the absence of bilateral central ground-glass opacities in the pulmonary parenchyma.

ILD due to aripiprazole is a rare finding, with only one case reported in literature (6) – being a 36-year-old woman who developed respiratory symptoms after beginning aripiprazole treatment. Drug-related ILD (DI-ILD), however, was only diagnosed 4 years later, compared to...
3 years in our patient. In our case, the time until the onset of respiratory symptoms after arisiprazole was not clear, although the temporal relationship between drug commencement and the development of respiratory symptoms supported the etiologic relationship after the elimination of other reasons. The specific diagnosis of hypersensitivity pneumonitis in the reported case was based on the characteristic findings of ground-glass appearance on HRCT scan with the lymphocytic predominance in the BAL sample. The previous case report’s radiology was also compatible with hypersensitivity pneumonitis, with bilateral multifocal mosaic pattern and ground-glass attenuation. Although the radiologic findings of the two patients were similar, indicating hypersensitivity pneumonitis, NSIP, as a subgroup of ILD, was diagnosed in our patient due to the chronic inflammation pattern on histopathology, and no lymphocytic predominance in BAL.

Figure 4: Peripheral and sub-rough reticular in zone opacities, heterogeneous density increase. (Last X-ray)

Figure 5: Irregularly limited and heterogeneous density increases in the lower bilateral zones, ground-glass appearance and septal thickening progression (last HRCT)

Figure 6: Radiological images from before the start of OCS treatment, 6 months after starting and during the 5-year follow-up (in order of top to bottom)

DI-ILD diagnoses are based on the compatibility of laboratory and imaging data in clinically compatible patients, based on an evaluation of the physical examination, laboratory and radiological findings together. The laboratory analysis is required to exclude infectious diseases, and autoimmune or rheumatological diseases. While some anti-infective and immunomodulatory drugs are known to cause lung damage, infections can be ruled out through culture studies in bronchoalveolar lavage (BAL) examination. PFTs with increased FEV1/FVC and decreased FVC and DLCO can help in the identification of a restrictive or obstructive model, although these results may overlap with other ILDs. HRCT is the most sensitive radiological examination for the diagnosis of ILDs (7-9).

Biopsy is not routinely recommended for diagnosis. Histological examination shows great similarity in idiopathic and drug-related forms. Transbronchial biopsy with FOB may be helpful in ruling out other ILDs, such as sarcoidosis (10). In NSIP, lung biopsies shows interstitial pneumonia, fibrosis or bronchiolitis obliterans, and findings suggestive of eosinophilic or extrinsic allergic alveolitis can be observed in some forms of DI-ILD (11,12).
ILD represents a heterogeneous group of pathologies that may be related to different causes. A low percentage of these lung diseases may be secondary to the administration of drugs or substances. Drug-Induced Interstitial Lung Diseases (DI-ILDs) have been associated with the administration of drugs such as anti-inflammatory and immunomodulators drugs, chemotherapeutics and cardiovascular drugs (13–15). Diagnostic doubt can be resolved through an analysis of the possible cause-effect relationship between the onset of a pulmonary pathology and exposure to a drug, and it is thus necessary to evaluate any recent pharmacological anamnesis and to investigate any previous ones (16). Toxic and immunological mechanisms have been held responsible for the development of DI-ILDs (1,17). Various clinical pictures such as lung parenchymal disease, acute lung injury, pulmonary hemorrhage, malignancy, mediastinal, neuromuscular and pleural/pericardial pathologies can develop after drug use (18). The patient's clinic due to DI-ILDs may vary from mild to severe, and may sometimes be fatal (19). The National Cancer Institute common terminology criteria for adverse events suggest a grading system from mild (1) to fatal (5). At the time of diagnosis, our patient had grade 2 (moderate) dyspnea, being symptomatic, but not interfering with daily activities, and progressed to 3 (severe) as symptomatic with limitation in daily activities, as well as an oxygen need in 5 years. A DI-ILD prognosis may have a favorable outcome after drug withdrawal, although complete recovery is rare. A significant proportion of cases follow a progressive clinical course with mortality-related respiratory failure, and progression of the primary underlying disease (15). As in the reports of previous cases, our case saw a slow progression of lung disease over 5 years, and also her bipolar disease worsened, due possibly to recurrent OCS use and alternative medicines that were not as effective as aripiprazole.

In addition to drug discontinuation and supportive therapy, corticosteroids are useful in the treatment of I LD diseases. It is recommended that oral corticosteroids (OCS) be initiated at a dose of 0.5–1mg/kg/day, continued for 2–6 months and gradually tapering the dose after clinical improvement is achieved. If the symptoms or radiologic findings recur during OCS dose reduction or interruption, the OCS dose should be increased again. Oxygen therapy and inhaled bronchodilator therapy are recommended as supportive therapies (20-23). In our case, after the discontinuation of aripiprazole and the administration of systemic corticosteroid therapy for almost a year, her symptoms, PFTs and radiological findings partially improved. Even though the radiologic and clinical improvement was more apparent in the previously reported case (6), our patient’s clinical and radiologic stage was more severe, and permanent restrictive changes progressed in 5 years to respiratory failure.

There is a lack of information in literature charting the rate of improvement or decline in patients who did not develop fibrosis after the drug was stopped, and so studies involving large case series are needed.

The limitations of this case report include the lack of an exact initiation time of ILD, which was based on the patient’s anamnesis, with the onset of respiratory symptoms after aripiprazole being reported by the patient. Another limitation is the diagnosis of ILD based on non-specific clinical and radiologic findings that could have occurred for other reasons that we were unable to identify.

CONCLUSION

The frequency and facilitating factors of DI-ILD are not fully known. The patient’s medication history should be questioned in detail when investigating the etiology of ILD. In patients with respiratory symptoms that began after starting treatment with a specific drug, one should keep DI-ILD in mind. Initiating treatment without delay is very important for patient prognosis. As in our case, disease progression and the development of respiratory failure cannot be prevented in some patients, despite the currently available treatments. Although lung biopsy is recommended for diagnosis, when a biopsy cannot be performed for any reason, a diagnosis should be made based on clinical and radiological findings, the drug in question should be stopped immediately, and treatment with steroids and supportive drugs should be started.

CONFLICTS OF INTEREST

None declared.

AUTHOR CONTRIBUTIONS


YAZAR KATKILARI


REFERENCES


5. Croxtail JD. Aripiprazole: a review of its use in the management of schizophrenia in adults. CNS Drugs 2012; 26:155-83. [CrossRef]


