Amiodarone Induced Organizing Pneumonia: A Masquerader of Community Acquired Pneumonia

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Abstract

Amiodarone is a widely used antiarrhythmic with well-known adverse effects, being pulmonary toxicity one of the most serious, occurring even with low doses. Amiodarone-induced organizing pneumonia (AIOP) is an infrequent histopathological presentation but a few cases have been reported in the literature.

A 78-year-old woman with atrial fibrillation doing amiodarone 200 mg once daily for two years was admitted to the Emergency Department with progressive dyspnea to rest, productive purulent cough, flulike symptoms and pleuritic chest pain. Blood tests revealed a hypoxemic respiratory failure and a systemic inflammatory response, and the chest radiography showed bilateral, multifocal pulmonary infiltrates. Microbiologic studies of urine, blood and sputum were sterile. The patient was initially treated for community acquired pneumonia, with clinical and radiological worsening despite multiple extended spectrum antibiotics. The complementary study excluded other etiologies, and a transthoracic pulmonary biopsy was performed, revealing histology consistent with organizing pneumonia.

AIOP is typically presented as a community acquired pneumonia that does not respond to antibiotics. The diagnosis depends on a high clinical suspicion, and compatible clinical and radiological pattern. Although it is not the most frequent, the presented case enforces the assumption that toxicity is present even with lower doses such as 200 mg once daily for 2 years.

Keywords: Amiodarone; Pulmonary toxicity; Organizing pneumonia; Treatment

Introduction

Amiodarone is one of the most common antiarrhythmic medications prescribed worldwide [1] and its adverse reactions are very frequent (75% of patients) [2] and extensively studied: corneal deposits in over 90% of the patients (and also development of cataract and optic neuropathy), interaction with other cardiac drugs, symptomatic bradycardia, hepatotoxicity, gastrointestinal disturbances, dermatitis, bluish discoloration of the skin, bone marrow suppression, coagulopathies, peripheral neuropathies, hyperthyroidism and hypothyroidism, being thyroid dysfunction the predominant one [1,3-5].

After the introduction of amiodarone as an antiarrhythmic in 1967, pulmonary toxicity was first described in 1980 [6] and has been reported in some series as the third most common reaction [3], being among the most severe [5]. It is present in 5-10% of patients taking amiodarone [5-8] and it can occur at any dose, even with low dose therapy [9]. The risk for toxicity increases with higher plasma concentrations [10]. However, Amiodarone pulmonary toxicity (APT) is better correlated with the total cumulative dose, rather than with the daily dose or plasma concentration of amiodarone [10,11]. Toxicity can occur at any time after treatment has been initiated. However, it has been described that those who have received a daily dose of 400 mg or more, for more than two months, or a lower dose, commonly 200 mg daily, for more than 2 years are at greater risk [11,12]. The initial loading dose is also believed to be an important contributor as it significantly increases the cumulative dose [4].

Amiodarone and its metabolites can produce lung damage directly by a cytotoxic effect, and indirectly by an immunologic reaction [11,13]. The most typical clinical and histopathological presentation is an interstitial pneumonitis, but less frequent presentations have also been described as such organizing pneumonia and diffuse alveolar damage [14-17].

Report

A 78-year-old woman with a history of hypertensive cardiopathy and atrial fibrillation was admitted with a progressive dyspnea to rest, productive purulent cough, nocturnal wheezing, flu like symptoms and pleuritic chest pain, over one week. She also complained with dry cough and dyspnea on exertion for the past 6 months. She was on losartan/hydrochlorothiazide 50/12.5 mg od, acetylsalicylic acid 100 mg od, and she had also been taking amiodarone 200 mg od for the last 2 years (cumulative dose 146 g - initial loading dose unknown).

On examination she presented with hemodynamic stability (arterial blood pressure 122/71 mmHg; heart rate 89 bpm), polypnea to minimum effort with a respiratory rate of 24 cpm, and bilateral inspiratory crackles, predominant in the right hemi thorax. There was no neurologic impairment. Blood tests revealed hypoxemic respiratory failure (pO2/FiO2 232) and systemic inflammatory response. The chest radiography displayed a bilateral, multifocal pulmonary infiltrates (Figure 1). Microbiologic studies of urine, blood and sputum were sterile. She was initially treated for community acquired pneumonia, with clinical and radiological worsening despite multiple extended spectrum antibiotics. High-resolution computed tomography (HRCT) showed diffuse parenchymal consolidation, with predominance in inferior lobes and posterior right superior lobe; and diffuse pericentimetric ganglion formations and in subcarinal region (Figures 2-4). Bronchial biopsies excluded local malignancy and the pathology of bronchoalveolar lavage (BAL) showed inflammatory infiltrates without granulomas or malignant cells. BAL immunology study was negative for alveolitis with 1.86 × 10^6 lymphocytes/mL.

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10^6 cells, with lymphocyte predominance of 41%, 18% neutrophils, and 12% eosinophils, with a low CD4/CD8 of 0.157, making the hypothesis of interstitial pneumonitis remote. BAL microbiological, mycobacteriological and mycological studies were negative, as well as HIV-1/2, therefore infectious etiology was extremely unlikely. ACE was negative, CD4/CD8 was low and there was no hypercalciuria, excluding sarcoidosis. It was performed a transthoracic pulmonary biopsy, which revealed an increased thickness of the septae, and a fibro-edematic nodule (appearing in one of the represented air spaces), corresponding to an intra-alveolar polyp, consistent with organizing pneumonia. Granulomatose lesions were absent.

Giving the unfavorable clinical course, and the extensive investigation excluding other pathological entities, namely infectious causes, it was considered the diagnosis of AIOP.

Thyroid function was normal. Fundoscopy was performed, showing bilateral whorl-like pattern of corneal epithelial deposits, consistent with amiodarone-induced vortex keratopathy, also called cornea verticillata, markers of amiodarone exposure [18].

Amiodarone was stopped and systemic corticoid therapy initiated with prednisolone 1 mg/Kg/day. There was an impressive clinical response with complete resolution of symptoms and respiratory failure in ten days. Imagiological reevaluation with HRCT at D12 showed significant reduction in bronchopneumonic consolidations and a discrete reduction in mediastinal ganglia. Respiratory functional tests were performed only at D15 of prednisolone, revealing obstruction of small airways, but no restriction pattern and normal DLCO corrected to lung volumes. Prednisolone was tapered gradually, and discontinued 10 months later, with complete clinical and imagiological resolution (Figures 1-7).

Discussion

Organizative pneumonia is commonly presented as community acquired pneumonia, with progressive dyspnea, cough, flu like symptoms, and a typical radiographic pattern mostly with alveolar, multifocal, irregular consolidations [19], being irresponsible to several antibiotic courses, as occurred in this case.

About 50% of occurrences correspond to a cryptogenic form, with the remaining having its origin in connected tissue diseases, drugs (amiodarone, carbamazepine, cephalosporin, nitrofurantoin, phenytoin, etc.) and infections (bacterial such as mycoplasma pneumonia, legionella, pseudomonas aeruginosa, streptococcus pneumonia, staphylococcus aureus, but also viral: HIV, Herpes simplex, influenza; fungi: Cryptococcus neoformans, Pneumocystis jiroveci; and parasites: plasmodium vivax) [19,20].

The diagnosis depends on a high clinical suspicion, and compatible
clinical and radiological pattern. A restrictive pattern in PFTs and a reduced DLCO (>15-20%) strengthens the diagnosis [5,11,20,21]. It may be confirmed with typical histopathological features from lung biopsy, rather obtained by video assisted thorascoscopy (VATS) or open thoracotomy, but it is acceptable and usually more reasonable to perform transbroncquic biopsy, especially in those with typical clinical and radiological features[19,20]. In this case only transthoracic biopsy was performed but the typical clinical presentation, the exposure to amiodarone and the exclusion of other etiologies makes our hypothesis likely.

Risk factors for APT have been suggested such as daily dose, cumulative dose, duration of therapy, patient age and the presence of pre-existing lung disease [3,5,11], but this is not consensual and more investigation is needed in order to define which risk factors are preponderant in development of AIPT to identify the patients at greater risk needing closest monitoring. D.K. Ernawati et al. tested the first four in a large database, and only patient age and duration of amiodarone therapy were confirmed as significant risk factors [3].

AIPT is always a diagnosis of exclusion [5], and there is no way of proving a cause-effect. The exposure risk factor and the typical clinical and imagiological patterns are supporting the diagnosis. PFT usually reveals low lung volumes, a restrictive pattern, and a reduced DLCO. The lack of these patterns in this patient may be explained by the delay in the availability of PFT that were deferred for two weeks later under corticosteroids. An obstruction of the small airways was documented in this case, a defect that has also been described [5,11,20,21].

Treatment decision is made considering disease severity. Stopping amiodarone is mandatory, and it may be sufficient in milder cases, with monitoring of lung function. It is recommended the use of oral corticosteroids in persistent or gradually worsening disease, which usually provide a rapid improvement of symptoms, as it did in our patient [5,11,21].

In case of fulminant disease, intravenous glucocorticoids should be used, with metilprednisolone 125-250 mg each 6 hrs or 750-1000 mg/ day for 3 to 5 days. In the other cases it is suggested an initial dose of prednisiolone of 0.75-1mg/Kg/day, (to a maximum of 100 mg/day). This dose should be maintained for 4 to 8 weeks, and then gradually tapered to 0.5-0, 75 mg/Kg/day for 4 to 6 weeks. After 3 to 6 months, the dose of oral prednisolone should be gradually tapered to zero if the patient remains stable or improved [20,22]. The patient should be routinely monitored with chest radiography and PFT every 2 to 3 months while under glucocorticoid therapy [20]. Relapses are common [19], but it has not occurred in our case 10 months later. It usually occurs when glucocorticoids are tapered, after 1 to 3 months [19,20]. Most of these patients will improve, when going back on prednisolone and the occurrence of relapses does not seem to affect the long term prognosis [22]. After the suspension, the patient should be monitored in the following year with a chest radiograph every 3 months [20].

A monitoring protocol has not been stablished worldwide, but some authors have suggested complete pulmonary function testing (PFT), including a diffusing capacity for carbon dioxide (DLCO) and chest radiography, before initiation of amiodarone therapy, every 3 months in the first year, yearly thereafter, and whenever pulmonary toxicity is suspected [2,21].

Conclusion

AIOP is typically presented as community acquired pneumonia that does not respond to extended spectrum antibiotics. The diagnosis depends on a high clinical suspicion, and compatible clinical and radiological pattern.

Although it is not the most frequent, our case supports that toxicity is present even with lower doses, such as 200 mg daily for 2 years. We must be aware of the multiple adverse reactions of amiodarone, and

every physician should keep in mind that monitoring these patients is crucial. It is important to alert patients about the potential pulmonary toxicity, and the importance of reporting any respiratory symptom.

A monitoring protocol has not been approved yet, but it has been suggested that patients taking amiodarone should be tested with chest radiography and PFT: before starting treatment, every 3 months in the first year, yearly thereafter, or anytime they experience symptoms. A validated strategy is urgent to better anticipate these harmful effects.

Key messages
- AIOP typically presents as a community acquired pneumonia, with typical radiologic pattern and usually a restrictive pattern with reduced DLCO in PFT.
- It is important to have an elevated suspicion index.
- It is always an exclusion diagnose after ruling out other life threatening pulmonary diseases.
- Toxicity is present even with lower doses, such as 200 mg daily for 2 years.
- It is important to alert patients doing amiodarone about the potential pulmonary toxicity and the importance of reporting any respiratory symptom.
- Patients taking amiodarone should be submitted to chest radiography and PFT: before starting the treatment, every 3 months in the first year, yearly thereafter, or anytime they experience symptoms.

Consent
All authors declare that written informed consent was obtained from the patient for publication of this case report.

Ethical Approval
Not applicable.

Competing Interests
Authors have declared that no competing interests exist.

References