Cystic Fibrosis Late Diagnosis: A Case Report

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Abstract

Cystic Fibrosis (CF) is an autosomal recessive disease that affects mucus and sweat producing cells involving multiple organs. CF is usually diagnosed in childhood; however, a considered number of adults are diagnosed every year. Atypical CF can be a milder form of the CF disorder, and individuals with atypical CF can remain undiagnosed for many years. Physicians should suspect of CF in adult individuals when these present recurrent pneumonia or bronchiectasis. Therefore, since early diagnosis of CF can avoid morbidities and unnecessary hospitalizations, the recognition of CF symptoms by clinicians is necessary to avoid late diagnosis.

Keywords: Late diagnosis; Atypical cystic fibrosis; Cystic Fibrosis (CF)

Introduction

Cystic Fibrosis (CF) is a recessive disease involving an autosomal gene, the Cystic Fibrosis Trans-membrane Conductance Regulator (CFTR) gene located at 7q31.2, which regulates the activity of chloride and sodium channels at the surface of the epithelium cell [1-3]. This disease affects the cells that produce mucus and sweat in multiple organs, being the lung the most severely affected and responsible for 90% of the deaths in patients with CF [4].

Defective CFTR protein results in thick and sticky mucus that can obstruct the pathways, since CFTR proteins in normal conditions lets chloride ions to pass through the mucus cells producing water and making the mucus thinner [5]. These obstructions, especially in the lungs, lead to massive neutrophil infiltration that induces the release of elastase that stimulates the lung antiproteases increasing lung destruction [6]. Besides the tissue destruction, neutrophils degranulation contributes to the mucus hyperviscosity since it releases a large amount of nucleic acids and cytosol matrix proteins [7]. Following the lungs, the GIT are severely affected since the mucous plugs obstruct pancreas enzymes and bile flow into the duodenum resulting in digestion abnormalities and malabsorption. Some patients present Distal Intestinal Obstruction Syndrome (DIOS) characterized by ileocecal obstruction of inspissated intestinal contents due to pancreatic insufficiency [8,9]. Although all these symptoms associated to GIT is involved as a multiorgan manifestation, they have not yet been systematically quantified. Tabori et al. [10] showed that patients with a severe course of the disease or with genotypes causing a moderate to severe abdominal involvement have more GI symptoms and that these symptoms are not associated with liver disease (CFLD) or elevated Liver Function Tests (LFTs).

Bronchiectasis, chronic infections associated with pneumonia, hemoptysis, pneumothorax and respiratory failures are complications associated with CFTR defected protein in respiratory system. GIT complications include nutritional deficiencies, including vitamins deficiencies and diabetes, progressive hepatic dysfunction, gallstones, intestinal obstruction, Small Intestine Bacterial Overgrowth (SIBO) and DIOS. Normally, infertility, osteoporosis, electrolyte imbalances, dehydration, fatigue, weakness and really low blood pressure are other complications associated with CF [11].

The aim of this work is to report a 28 years old female that was diagnosed with CF.

Case Report

28 years old female was referred to Emergency Room of the Regional Hospital of Presidente Prudente (HRPP) with worsening dyspnea accompanied by greenish vomit, inappetence, epigastric pain, and paresthesia of the right upper limb. She did not have a history of smoking or alcoholism. She complained of weight loss, holocranial headache and cough with expectoration. There was absence of relevant complaints in the other systems.

The patient history showed that she was born vaginally, weighting 2,780 g, and 51 cm of height. At 15 days of life she presented her first lung manifestation with pleural effusion. At 5 years old she started with a history of upper and lower airway infections episodes that were refractory to the proposed treatments. By 13 years old she underwent her first surgical procedure, a right inferior and middle lobectomy. Three years after her first surgery, she was submitted to a new pulmonary lobe resection, a superior right lobectomy. Four years later, she underwent amputation of the right bronchial stump which presented chronic inflammatory processes with fibrosis, hyalinization and hemorrhagic areas. Figure 1 shows the images of flexible bronchoscopy that was made before and after the amputation of the right bronchial stump.

On examination she presented good general condition, afibrile, acyanotic, aniceric, stained, hydrated and with tachypnea. Her heart rate was 88 beats per minute; her respiratory rate was 28 breaths per minute. Auscultation revealed rhythmic heart rate and vesicular murmur in the left hemithorax with presence of crepitant rales on the left pulmonary base. Her abdomen was flat, normotensive, with hdroaerose noises and without visceromegalias.

Based on the history of the patient, the initial diagnosis was of residual bronchiectasis of the left lung, with the suspicious of cystic fibrosis. She was admitted in the HRPP for the 5th time in the period of 1 year for complaints regarding pulmonary affections, and this time she was diagnosed with opportunistic infection by Mycobacterium abscessus. The patient also underwent pilocarpine sweat iontophoresis and the diagnosis of Cystic Fibrosis was confirmed. The patient was then treated with 600 mg Amikacin, 500 mg Clarithromycin, 2 g Cefepime and 400 mg Ciprofloxacin.
After diagnostic confirmation, the patient was hospitalized five more times in a period of four years as a consequence of infected bronchiectasis. She underwent respiratory physiotherapy and she was treated with 500 mcg/day Beclomethasone and 100 mcg Salbutamol when necessary. In March of this year (2017) she underwent a spirometry that showed a very severe obstructive disorder, that slowed her vital forced capacity to 34% of the expected value. Nowadays she has been using 24 mcg/day Formoterol and presents dyspnea to small efforts. Figure 2 shows images of a recent CT showing the absence of the right lung and the presence of bronchiectasis.

Discussion and Conclusion

Cystic Fibrosis (CF) is a genetic condition with high prevalence among Caucasian populations with an incidence of 1:2500 live births. One in 25 persons is asymptomatic carriers. CF can be diagnosed at birth for early medical and nutrition intervention that can lead to improved outcomes [12, 13]. And although most people with classic CF will be diagnosed through newborn screening or symptoms in early childhood, there are atypical forms of CF that will only be recognized in adults. The diagnosis in the first year of life is typically accompanied of meconium ileus, failure to thrive, pulmonary infections, diarrhea, and steatorrhea. In adults, patients usually present respiratory symptoms or male infertility due to Congenital Bilateral Absence of the Vas Deferens (CBAVD) [14].

The data from CF Foundation showed in 2001 that in an n=100, 9.9% of the new diagnoses in US occur in the age of 18 or older. The tendency is that this rate must increase, since the identification of over 1000 mutations now makes possible the diagnosis in those who have atypical symptoms or equivocal sweat test results [15].

Wilde research has been showing that the individuals who are diagnosed in a late age display better lung function, higher rates of pancreatic sufficiency, fewer complications, and longer life expectancy than those who are diagnosed at birth or as children. They also can live a normal life, being married and working full time [16-18]. However, a study has shown that late diagnosis does not mean good prognosis. In
fact, it was found that patients diagnosed at an early stage generally did better than those diagnosed at a later stage [16].

There are many reasons for late diagnosis in CF. Mild or absent symptoms may delay a patient seeking medical attention, a physician may not suspect of CF in a patient with atypical symptoms, and some tests, such as the sweat test, can present normal or borderline results. Normally, pulmonary infections are the reason an adult is diagnosed with CF, and GIT symptoms rarely lead to diagnosis [16].

Therefore, a physician can suspect of CF in an adult, even though this pathology is taught as an infant disease. Certain pulmonary signs, however, such as recurrent pneumonia, progressive obstruction possibly identified as asthma or chronic obstructive pulmonary disease, chronic sinusitis, or nasal polyposis can lead to suspicious CF [19,20]. Among GIT symptoms in atypical CF, it can be included chronic constipation or diarrhea and chronic or recurrent pancreatitis [21].

Delay in the diagnosis can lead to substantial morbidity and several hospitalizations that might be avoided. There are excellent treatments for the various symptoms of CF, including chest physiotherapy, antibiotic therapy, diet modification, and pancreatic enzyme replacement. Thus, a closer relationship with a clinician can help identify other complications and appropriate treatment can be offered, improving quality of life and life expectancy to those who are affected by CF.

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