INTRODUCTION

Cystic fibrosis (CF), caused by mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene, is the most common autosomal recessive disorder of Caucasians.1,2 Today, the majority of CF cases are diagnosed with newborn screening.3 Despite universal newborn screening, which was achieved in the United States in 2010, increasing numbers of patients are being diagnosed in adulthood.4 With widely available advanced genetic testing and multiple classes of mutations identified, CF is now recognized as a disease with a range of severities, affecting various ages, races, and ethnicities, and is no longer limited to a pediatric disease of Caucasian race.4,5 However, the widening epidemiological background of the disease is posing new diagnostic challenges. Achieving CF diagnosis in the adult population has been notoriously difficult as symptom onset and severity can be variable.6 Moreover, CF patients diagnosed as adults have more subtle symptoms and can have inconclusive sweat chloride test results, which have traditionally been the key to CF diagnosis.7 This diagnostic challenge had been recognized by the US and European CF foundations, and subsequently, consensus guidelines were released in 2008 and revised in 2017.6 The major update in the 2017 consensus guidelines was to lower the threshold for the intermediate sweat chloride levels to 30-59 mmol/L with < 30 mmol/L being considered normal for all ages.

Allergic bronchopulmonary aspergillosis (ABPA) is a frequent complication of CF with an incidence of 8.9% (ranging...
from 3% to 25%). ABPA is a less prevalent complication of asthma with an incidence ranging from 1% to 3%. ABPA results from a hypersensitivity reaction to Aspergillus antigens (most commonly *Aspergillus fumigatus* (AF)) causing a Th2-type inflammatory reaction and airway obstruction. Patients present with wheezing, mucus plugging, hemoptysis, and abnormal radiologic appearance including bronchiectasis. The clinical presentation and diagnosis criteria for ABPA often overlap with the clinical and radiologic features of CF pulmonary exacerbation. Here, we report a case of undiagnosed CF despite a long-term diagnosis of ABPA. This report highlights the evolving CF guidelines, and the greater need to educate general practitioners or primary care providers how to identify patients who need referral to CF centers.

2 | CASE HISTORY

A 58-year-old Caucasian woman, with history of asthma, allergic bronchopulmonary aspergillosis (ABPA), and right upper lung lobectomy secondary to eosinophilic granuloma and osteopenia presented with worsening pulmonary symptoms over the preceding four years. Pertinent family history included asthma in her mother and sister. The patient denied any smoking history.

Her medical history started with childhood asthma that resolved in her late teenage years but recurred at age 35. She had undergone multiple sinus surgeries and septoplasty in the interim years. ABPA was diagnosed at the age of 54 with elevated serum IgE of 711 IU/L and *Aspergillus fumigatus* (AF) recovered from sputum. Her FEV1 was at 64% of predicted levels. She was hospitalized numerous times for Pseudomonas pneumonia and exacerbation of her severe bronchiectasis, each time treated with IV antibiotics (Figure 1A). She also suffered acute pancreatitis with an elevated lipase of 340 at the age of 54.

Due to worsening lung condition and pancreatitis, her primary care doctor ordered a standard 32 CFTR mutation panel. The test revealed one mutation, 394delTT (c.262_263delTT, p.Leu88IleefsX22), and she was diagnosed as a CF carrier. Her sweat chloride was deemed indeterminate at 40 mmol/L based on the 2008 guidelines. Unfortunately, her treatment remained focused on managing severe asthma and ABPA without further workup for CF.

Four years later, at the age of 58, the patient continued to complain of an increasingly congested cough and severe hemoptysis with significant difficulty raising secretions. She was diagnosed with severe bronchiectasis (Figure 1B) and declining lung function. Her FEV1 levels had decreased to 51% despite extensive treatment for asthma and ABPA with systemic steroids. Given her worsening lung condition, the possibility of CF was again raised and a full CFTR gene sequencing was ordered by her primary care physician. The full panel revealed an additional mutation, defining her as compound heterozygous for two rare mutations: 394delTT (c.262_263delTT, p.Leu88IleefsX22) and L967S (c.2900T > C, p.Leu967Ser).

The patient was referred to a CF center for treatment and genetic counseling. Repeat chloride testing at the CF center was 22 mmol/L. Family testing was offered to relatives. Following prescription of a proper airway clearance regimen including dornase alfa and chronic antipseudomonal treatment, her last FEV1 significantly improved from 35% to 53% (Figure 2).

3 | DISCUSSION

This case illustrates how an initial diagnosis of ABPA led to diagnostic confirmation bias resulting in a many-year delay to diagnose CF despite an array of symptoms pointing to CF and continued lung function decline even with full ABPA and asthma therapy. The course of this patients CF was compounded by ABPA, a common complication of CF which is
Our patient's clinical presentation of CF with sinus disease, bronchiectasis, recurrent pulmonary exacerbation with hemoptysis, multi–drug-resistant Pseudomonas, and gastrointestinal manifestations including pancreatitis should have guided the clinicians to obtain sweat chloride testing to help determine the relevance a CF diagnosis. Her first sweat chloride testing at 40 mmol/L is in fact intermediate and should have triggered a referral to an accredited CF center where an extended CFTR gene analysis or CFTR functional analysis would be considered to conclusively diagnose CF. This patient's CF therapy was also significantly delayed due to being incorrectly labeled a CF carrier with one mutation captured on the 32-mutation panel. There are over 1700 recognized mutations, and by only performing a 32-mutation panel for a symptomatic patient, full understanding of their CF diagnosis will be further delayed.11 If a single CFTR variant is identified on limited analysis, then sequencing of the entire CFTR gene should be performed.12

Patients diagnosed with CF as adults often have involvement of fewer organ systems at the time of presentation. This delayed disease manifestation has been attributed to the presence of a residual function mutation combined with a more severe mutation13 and genetic modifiers of the severe genotype.14 Currently, the Clinical and Functional Translation of CFTR project database reports 442 variants.11 Of which, 360 variants are known to be CF-causing and 11 variants are known to be of unknown significance. Our patient had 2 CFTR gene mutations. One allele 394delTT (c.262_263delTT, p.Leu88IlefsX22) has been identified in almost 300 individuals with CF worldwide and is considered a CF-causing variant. The second allele, L967S (c.2900T > C, p.Leu967Ser), is currently classified in the CFTR2 database as a variant of varying clinical consequence. This mutation combination resulted in sufficient CFTR function to delay serious manifestation until our patient reached her late 30s.

As the rate of individuals older than 18 years being diagnosed with CF is increasing,4 the need to educate and update general practitioners must be emphasized to help guide the workup process starting with obtaining relevant history—detailed family history, under-nutrition, chronic sinus disease, and acute or chronic pancreatitis. This need was recognized by the National Institute for Health in the UK who published the National Institute for Health and Care Excellence (NICE) guidance on the diagnosis and management of CF, which encompasses children, young people, and adults and is written for general practitioners.15

Prompt diagnosis of CF will lead to early referral to CF centers and access to specialized medical care. Studies have shown that early intervention in CF management has improved outcomes.16 A study from Denmark has also shown that delayed diagnosis of CF is associated with moderate-to-severe malnutrition at diagnosis.17 Additionally, newly diagnosed adult CF patients who received multidisciplinary, evidence-based care at CF centers achieved significant improvement in FEV1 from the time of diagnosis.18,19 Lastly, the risk factors for older age at diagnosis, diabetes, and lower lung function are predictors of reduced survival.20 Thus, early diagnosis and appropriate treatment improves outcomes for CF patients.

In sum, this case shows the importance of educating general practitioners on the epidemiology of CF and evolving changes in guidelines for CF treatment and diagnosis. It is of particular importance since the CFTR corrector and modulator therapies are dramatically improving the outcomes for
most patients. Had the 2017 CF guidelines and NICE guidelines been followed, this patient could have had earlier access to specialized CF care, which could have improved her quality of life and prevented further deterioration of her lung function.

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CONFLICT OF INTEREST
None declared.

AUTHOR CONTRIBUTIONS
PC, LB, and MG: planned the manuscript. PC: composed the first draft. PC, LB, and MG: edited the final manuscript.

ETHICS STATEMENT
Consent was obtained from the patient prior to publishing this case.

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