EDITORIAL

A first step against idiopathic pulmonary fibrosis

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Idiopathic pulmonary fibrosis (IPF) is a chronic devastating disease affecting 50 to 300 individuals per million with a median survival time of ~3 yrs [1]. The recent publication of evidence-based guidelines for the diagnosis and management of patients with IPF did not provide patients and caregivers with any reason for optimism [2]. The only recommended procedures in this document were: long-term oxygen therapy in patients with clinically significant hypoxaemia; lung transplantation in selected patients; treatment of asymptomatic gastro-oesophageal reflux (although there is not even one randomised trial showing that this treatment improves lung function decline or survival in IPF patients); pulmonary rehabilitation; and corticosteroid use in patients with acute exacerbation of IPF [3]. Almost all treatment procedures evaluated in that statement were given either a weak or a strong recommendation against use [2].

However, the litany of the negative therapeutic trials in IPF ended with the recent publication of the results of three phase III trials evaluating the efficacy of pirfenidone [4–6]. The results of these trials, including >1,000 patients, support a protective effect of pirfenidone to limit the decline of lung function in patients with IPF [7]. This effect was the basis for the recent approval of pirfenidone (Esbriet®; InterMune Inc., Brisbane, CA, USA) in the European Union (since February 28, 2011). Pirfenidone has been already available in Japan since 2008 (Pirespa®; Shionogi and Co. Ltd, Osaka, Japan) and a generic form was recently made available in India (Pirfenex®; Cipla Ltd, Mumbai, India) [8]. Although a panel of experts voted for approval, the US Food and Drug Administration have not yet granted approval for pirfenidone and have requested that a further phase III clinical trial be performed. Pirfenidone is indicated in Europe in adults for the treatment of mild-to-moderate IPF [9]; however, there is already no consensus about what is mild IPF or moderate IPF compared to severe IPF [3]. A definition of these degrees of IPF severity is needed in order to provide all clinicians with the necessary common vocabulary. Unless we have this vocabulary, we should probably consider that pirfenidone use be limited to patients who meet the inclusion criteria of the CAPACITY (Clinical Studies Assessing Pirfenidone in IPF: Research of Efficacy and Safety Outcomes) trials, i.e. forced vital capacity ≥50% predicted and a diffusing capacity of the lung for carbon monoxide ≥35% pred [5].

The European approval signifies a new era for patients and doctors dealing with IPF in the European Union. First, it offers hope to patients and their families as this active antifibrotic molecule will be available very soon, and it gives doctors a weapon with which to combat this challenging disease. Secondly, it demonstrates that there is still room for non-targeted drugs in modern medicine. Indeed, although pirfenidone has been consistently shown to exert antifibrotic properties in animal models of renal, respiratory, cardiac or hepatic diseases [10–12], the exact mechanism of action of pirfenidone is still poorly understood and might, in fact, be multi-targeted as antioxidant, anti-transforming growth factor and anti-platelet derived growth factor properties have been demonstrated [12]. Further studies are needed to determine more precisely the antifibrotic pathways targeted by pirfenidone as this could help identify new antifibrotic molecules.

Thirdly, as soon as pirfenidone is made available to patients, the design of therapeutic trials in IPF patients will have to adapt to this new situation. Indeed, long-term, placebo-controlled studies will be ethically unsustainable in a progressive disease such as IPF. Both clinicians and pharmaceutical companies will have to incorporate this new information into the way therapeutic trials are developed in the future, at least in the countries where pirfenidone will be available.

Fourthly, IPF is only one of the fibrotic lung diseases that patients and clinicians face every day. For example, connective tissue diseases-associated lung fibrosis is very common and sometimes leads the prognosis, such as in patients with systemic sclerosis or rheumatoid arthritis [13]. Asbestosis and chronic hypersensitivity pneumonitis are environment-induced lung fibrotic diseases with a poor prognosis and limited therapeutic possibilities [14, 15]. Idiopathic fibrotic nonspecific interstitial pneumonia is also a progressive fibrotic lung disease with a significant mortality, although it has a better prognosis than IPF [16]. Altogether, these fibrotic lung diseases have no approved treatment. The evaluation of a possible protective effect of pirfenidone is warranted before this drug is used in these patients. Indeed, we recently learnt that a protective effect in IPF does not necessarily translate into a protective effect in all fibrotic lung diseases, as illustrated by the negative trial of pirfenidone in Hermansky-Pudlak syndrome [17].

Finally, it is clear that pirfenidone is only the first step in the war against lung fibrosis. Pirfenidone has slowed the decline of...
lung function but it has not prevented acute exacerbations in the recent trials [6, 7]. Furthermore, whether pirfenidone has the capacity to target the vascular remodelling which is responsible for the development of sometimes severe pulmonary hypertension in IPF patients is unknown [18]. Although this is a great time for patients with IPF, there is still room for antifibrotic drugs to come before the whole spectrum of fibrotic lung diseases gets covered.

STATEMENT OF INTEREST

B. Crestani received a fee for speaking from InterMune Inc.

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