



The therapy of idiopathic pulmonary fibrosis: what is next?

Vivien Somogyi^{1,2}, Nazia Chaudhuri³, Sebastiano Emanuele Torrisi^{1,4}, Nicolas Kahn¹, Veronika Müller² and Michael Kreuter¹

Affiliations: ¹Center for Interstitial and Rare Lung Diseases, Pneumology, Thoraxklinik, University of Heidelberg, German Center for Lung Research (DZL), Heidelberg, Germany. ²Dept of Pulmonology, Semmelweis University, Budapest, Hungary. ³Manchester University NHS Foundation Trust, Wythenshawe Hospital, Manchester, UK. ⁴Regional Referral Centre for Rare Lung Diseases, University Hospital "Policlinico", Dept of Clinical and Experimental Medicine, University of Catania, Catania, Italy.

Correspondence: Michael Kreuter, Center for Interstitial and Rare Lung Diseases, Thoraxklinik, University of Heidelberg, German Center for Lung Research (DZL), Heidelberg, Germany. E-mail: kreuter@uni-heidelberg.de

@ERSpublications

There is still no cure for IPF. Clinical trials focus on new therapeutic targets, improvements in nonpharmacological therapies and treatment of comorbidities and acute exacerbations. All future therapies should aim to reduce the burden of disease. http://bit.ly/2XdxqAP

Cite this article as: Somogyi V, Chaudhuri N, Torrisi SE, et al. The therapy of idiopathic pulmonary fibrosis: what is next? Eur Respir Rev 2019; 28: 190021 [https://doi.org/10.1183/16000617.0021-2019].

ABSTRACT Idiopathic pulmonary fibrosis (IPF) is a chronic, progressive, fibrosing interstitial lung disease, characterised by progressive scarring of the lung and associated with a high burden of disease and early death. The pathophysiological understanding, clinical diagnostics and therapy of IPF have significantly evolved in recent years. While the recent introduction of the two antifibrotic drugs pirfenidone and nintedanib led to a significant reduction in lung function decline, there is still no cure for IPF; thus, new therapeutic approaches are needed. Currently, several clinical phase I–III trials are focusing on novel therapeutic targets. Furthermore, new approaches in nonpharmacological treatments in palliative care, pulmonary rehabilitation, lung transplantation, management of comorbidities and acute exacerbations aim to improve symptom control and quality of life. Here we summarise new therapeutic attempts and potential future approaches to treat this devastating disease.

Introduction

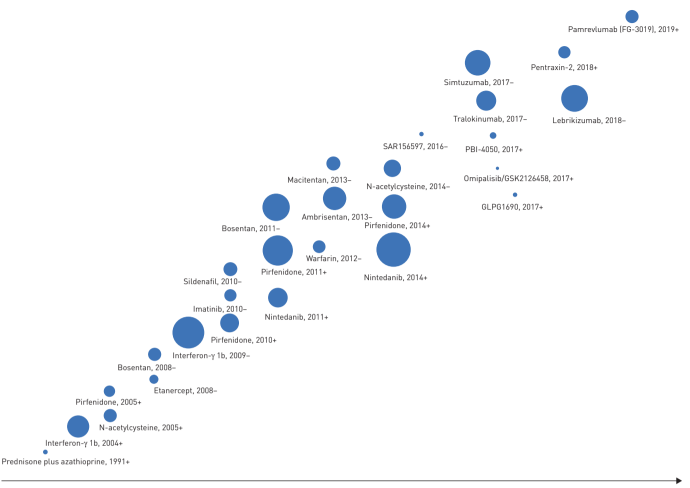
Idiopathic pulmonary fibrosis (IPF) is a chronic, fibrosing idiopathic interstitial lung disease (ILD) characterised by progressive scarring of the lung parenchyma associated with a steady worsening of respiratory symptoms, and decline of pulmonary function, ultimately leading to death [1, 2].

Despite constant efforts, many therapeutic approaches failed until antifibrotic therapy fundamentally altered the therapeutic approach in IPF (figure 1). Currently, two antifibrotic therapies are available: pirfenidone and nintedanib. Pirfenidone has proven anti-inflammatory, antioxidant and antifibrotic effects. After an early feasibility study [4], prospective studies demonstrated that pirfenidone reduces the decline of forced vital capacity (FVC) in patients with IPF [5–8]. *Post hoc* analysis of the pivotal international phase III studies described a significant reduction of the risk in FVC decline, death, disease progression, 6-min walk distance (6MWD) and dyspnoea in the pirfenidone-treated patients compared to placebo [9]. Further *post hoc* analyses described a significant reduction in respiratory-related hospitalisation [10, 11], and consistent effects of pirfenidone in all subgroups analysed, *e.g.* with regards

Publication of this peer-reviewed article was sponsored by Boehringer Ingelheim, Germany (principal sponsor *European Respiratory Review* issue 153).

Received: 27 Feb 2019 | Accepted after revision: 16 May 2019

Copyright ©ERS 2019. This article is open access and distributed under the terms of the Creative Commons Attribution Non-Commercial Licence 4.0.



Time

FIGURE 1 Evolution of treatment in idiopathic pulmonary fibrosis (IPF). Schematic representation of clinical trials of therapy performed for IPF in the past three decades. Circle sizes are an approximate representation of the sample sizes of the clinical trials. +: study ended with positive outcome; -: study ended with negative outcome. Reproduced and modified from [3] with permission.

to FVC decline, 6MWD, health-related quality of life (HRQL) [9, 12]. Furthermore, pirfenidone is associated with adequate safety and tolerability [13]. Nintedanib is a tyrosine kinase inhibitor where *in vitro* analyses have demonstrated inhibitory effects on fibroblast proliferation and differentiation [14, 15]. Phase II and III studies described significant effects on reducing lung functional decline. Additionally, the key secondary end-point, time to the first acute exacerbation, favoured the use of nintedanib [16, 17]. *Post hoc* analyses showed consistent effects of nintedanib reducing FVC decline regardless of the underlying radiological pattern, presence of emphysema or other important baseline characteristics [18, 19], and was effective in patients with more advanced disease [20]. Nintedanib has an adequate safety and tolerability profile [16, 17].

ABCDE and future approaches in IPF

To ensure an optimal patient based approach in IPF, management may be guided by an ABCDE algorithm: Assessment of the needs of patients and caregivers; Backing up the patient with educational resources, self-management strategies and pulmonary rehabilitation; Comfort-care (symptom-based therapies) and treatment of Comorbidities; Disease-modifying treatments; and End-of-life care [21]. Yet, despite significant progress in the treatment of IPF, there is still no cure, and thus new concepts in treatment are needed. These strategies may include therapeutic biomarkers guiding therapy, the use of combination therapies, novel drugs targeting new pathways in fibrosis, therapies aimed at targeting the lung microbiome, novel nonpharmacological therapies with developments in lung transplantation, new ways to approach palliative care, improved treatment of comorbidities and an important unmet need in treatment of acute exacerbation of IPF (AE-IPF).

Biomarkers guiding therapy

A prerequisite for future personalised therapy for IPF is the development of biomarkers that can guide diagnostic, therapeutic and prognostic approaches in managing IPF. Several potential peripheral blood, molecular and radiological biomarkers have been identified and are summarised in table 1.

Diagnostic biomarkers

Accurate diagnosis in IPF remains a challenge and the current gold standard of performing a lung biopsy according to guideline recommendations [1, 2] is not without significant risk [55], and thus often not possible. Thus, the identification of specific biomarkers that can facilitate the accurate diagnosis of IPF is of great interest. A number of peripheral blood biomarkers have been studied that can aid in delineating ILDs from other severe lung diseases. The markers Krebs von den Lungen (KL)-6 [24], chitinase-like protein (YKL40) [41, 56], leucocytes and circulating innate immune cells [57-59], surfactant proteins (SP)-A, -B, -D, among others, have been shown to discriminate IPF from healthy controls [25, 27, 31]. KL-6 may also discriminate ILDs from other benign lung diseases [24, 60]. Another valuable biomarker is the matrix metalloproteinase (MMP)7, where higher levels are related to interstitial lung abnormalities [33]. Furthermore, MMP7 and MMP1 may distinguish between IPF and hypersensitivity pneumonitis [32]. The PROFILE (Prospective Study of Fibrosis in Lung Endpoints) study, the first study to perform a sequential evaluation of >100 serum protein biomarkers, demonstrated that markers of epithelial injury and matrix degradation are important in IPF. The study showed that levels of epidermal growth factor receptor and clusterin were lower and levels of MMP1, MMP7 and SP-D were significantly elevated in IPF patients compared to healthy controls [25]. In addition, PROFILE revealed that oncostatin M and cytokeratin 19 fragment (CYFRA-21-1) were able to distinguish IPF patients from healthy controls [47]. The biomarkers KL-6 and SP-D have been used widely in the diagnostic pathways for IPF in Japan [61]. However, currently, the clinical utility of the abovementioned serum biomarkers is still limited and thus they are not recommended for diagnostic purposes in IPF [2].

Prognostic biomarkers

Progression in IPF is challenging to predict due to the heterogeneity of the population. FVC decline has been shown to be a predictor of mortality in IPF [62] and is the most robust measure adopted in clinical trials. However, baseline FVC does only have limited prognostic value and longitudinal changes in FVC have some weaknesses [63]. Therefore, composite scoring systems incorporating lung physiology, sex and age are deemed more accurate at predicting mortality [64, 65]. It is recognised that biomarkers may augment these models and improve prognostication in IPF. Several biomarkers have been studied that show significant association with prognosis in IPF (table 1) [22, 23, 26, 28–30, 34–40, 42–44]. High MMP7 has shown significant correlations with IPF disease severity as defined as lung function parameters (FVC and diffusion capacity of the lung for carbon monoxide (*DLCO*), and increased breakdown products of MMP activity predicted worse survival in IPF [33].

Furthermore, the PROFILE study showed that C reactive protein degraded by MMP1/8 (CRPM) discriminated IPF from healthy subjects and stable from progressive subjects, predicted progression and indicated poor overall survival [45]. Other predictors of progression and survival in IPF were SP-D and cancer antigen (CA)19-9, -125 [46].

The ongoing INMARK study will assess the effect of nintedanib, on the rates of change of biomarkers of extracellular matrix (ECM) turnover in patients with IPF, their ability as predictors of disease progression and whether the drug alters associations between changes in biomarkers and disease progression [66].

Radiological biomarkers

Several studies have been published in recent years on the potential of high-resolution computed tomography (HRCT) patterns as prospective prognostic biomarkers [67]. A recent study reported about the potential of quantitative computed tomography (CT) tools such as Data-Driven Textural Analysis (DTA) [68]. In this study, automated recognition of IPF on CT resulted in a severity index that correlated with visual and functional changes [68]. Yet, there is still a high need for elaboration of automated, visual CT scoring *via* machine learning development process and for creation of expanded datasets supporting this process [69]. The importance of functional respiratory imaging (FRI) providing quantitative visualisation of different patterns was recently highlighted, where FRI correlated with FVC decline [70]. Furthermore, CT densitometry, especially the area right of the inflexion point in HRCT histogram was reported to correlate with lung function decline [71]. JACOB *et al.* [72] reported recently that a computer scoring based quantification of parenchymal patterns including vessel-related structure scores predicted IPF mortality and functional decline in IPF. Hyperpolarised xenon-129 magnetic resonance imaging could be a potential non-invasive method for estimating gas-exchange impairment in IPF, as this inert gas is able

	Mechanism of action	Outcome of the study	Effect on IPF	
irculating blood biomarkers				
CCL18	Alternative macrophage activation Upregulation of collagen production by lung fibroblasts Higher mortality in patients with serum CCL18 concentrations >150 ng·mL ⁻¹ , higher incidence of disease progression in the group with high serum CCL18 concentrations [22]		Predicts progression and mortality in IPF	
ICAM-1	Adhesion molecule Marker of oxidative stress in the lungs	Predicts poor overall, transplant-free and progression-free survival [23]	Predicts mortality in IPF	
KL-6/MUC1	High molecular weight glycoprotein expressed at ECM surface of type II pneumocytes	Significantly higher level in ILDs [24] Higher levels among patients who died within the study period [25] Lower survival of patients with high KL-6 [26]	Discriminates ILDs from other benign lung diseases Predicts mortality in IPF	
SP-A	Surfactant proteins produced by type II pneumocytes	High level in IPF [27] Independent predictor of survival [28] Associated with the time to death or lung	Discriminate IPF from other ILDs SP-A, SP-D predict mortality i	
		transplantation [29] Predictive effect on those with UIP in HRCT [30]	IPF	
SP-B		Higher level in IPF [31]		
SP-D		High level in IPF [25] Independent predictor of survival time better related to parenchymal involvement [27]		
MMP1	Zinc-dependent proteases involved in the breakdown of ECM components	Distinguish between IPF and HP [32] Elevated levels in IPF [25]	Discriminates IPF from other ILDs	
MMP7	MMP1 the most highly expressed interstitial collagenase degrading	Distinguish between IPF and HP [32] Elevated levels in IPF [25]	Discriminates IPF from other ILDs	
	fibrillar collagens MMP7 the smallest member capable of degrading multiple components of ECM	Related to FVC decline, to higher prevalence of exertional dyspnoea, to ILAs on HRCT and to higher all-cause mortality [33]	Predicts mortality in IPF	
BNP	Natriuretic peptide secreted by cardiac ventricles	Correlation with clinical status, functional exercise testing parameters, functional WHO class II, III [34]	Relates to haemodynamic parameters and prognostic value in patients with left or right heart failure	
VEGF	Growth factor regulating angiogenesis enhancing vascular permeability	Positive correlation with HRCT interstitial score, influence on monthly FVC decline [35]	Reflects severity and predicts progression of IPF	
CD28 downregulation on CD4 ⁺ T52 cells	CD28 co-stimulatory molecule providing signal for activation of naive CD4 lymphocytes	Correlated with decreased FVC and freedom from major adverse events (death or lung transplantation) [36]	Predicts progression, mortalit in IPF	
HSP70 IgG antibodies	HSP70 antibody working against HSP70 autoantigene and activating IL-8 production of monocytes	Associated with decreased FVC and 1-year survival [37]	Predicts progression, mortalit in IPF	
Periostin	Fibroblast activating matrix proteins	Negative correlation with monthly changes in VC, <i>D</i> LCO [38] Increase of honeycombing score on HRCT, predictor of shortened overall survival, time-to-event [39]	Predicts progression, mortalit in IPF	
Osteopontin	Glycoprotein secreted by osteoclasts, macrophages and activated T-cells	Reverse correlation with arterial oxygen tension [40]	Predicts progression in IPF	
YKL40	Chitinase-like protein	Elevated levels in ILDs, correlated with poor survival [24, 41]	Discriminates ILDs from healthy subjects Predicts mortality in IPF, remains predictive after 3-4 years	
BLys	Plasma B lymphocytes stimulating factor	Correlated with pulmonary artery pressures, subjects with higher BLys diminished 1-year survival compared to those with lower BLys [42]	Predicts PH and survival in IPI	

TABLE 1 Peripheral blood and molecular biomarkers in idiopathic pulmonary fibrosis (IPF)

TABLE	1	Continued

	Mechanism of action	Outcome of the study	Effect on IPF	
Circulating fibrocytes	Produce ECM components, mesenchymal markers Potential role in myofibroblast differentiation	High levels correlated with poor survival regardless to preservation of lung function, counts increased further during AE-IPF [43]	Predicts survival in IPF	
CXCL13	Chemokine playing a role in autoimmune processes, mediating B-cell homing to inflammatory foci	High levels correlated with poor FVC and poor major event-free survival (<i>i.e.</i> transplant-free survival) [44]	Predicts progression, mortality in IPF	
EGFR	Epidermal growth factor required for TGF-B1-induced epithelial-mesenchymal transition Crucial in signalling in bronchial epithelium	Lower levels in IPF [25]	Discriminates IPF from healthy subjects	
Clusterin	Known as apolipoprotein J Glycoprotein upregulated by cytotoxic stimuli, maintaining epithelium viability during lung repair	Lower levels in IPF [25]	Discriminates IPF from healthy subjects	
CRPM	C reactive, acute-phase protein degrading by matrix metalloprotease	Higher levels in IPF, could discriminate between stable and progressive subjects and indicated poor overall survival [45]	Discriminates IPF from healthy subjects Predicts progression, mortality in IPF	
CA-19-9	Tumor markers, mucous associated carbohydrate antigens increasing in	High levels highly predictive of progressive fibrosis [46]	Predicts progression, mortality in IPF	
CA-125	metaplastic epithelium in fibrotic lesions Associated with mucous secretion within honeycomb cysts	Rising levels predicted both disease progression and overall survival [46]		
OSM	Glycosylated protein, member of IL-6 family of ligands	Identified baseline prognosis and longitudinal change in individuals with IPF [47]	Discriminates ILDs from healthy subjects Predicts progression, mortality in IPF	
CYFRA-21-1	Intermediate filaments in the cytoskeleton of alveolar and bronchiolar epithelial cells Marker of epithelial cell damage	Identified baseline prognosis and longitudinal change in individuals with IPF [47]	Discriminates ILDs from healthy subjects Predicts progression, mortality in IPF	
olecular biomarkers				
MUC5B	Mucin associated with the development of both familial interstitial pneumonia and sporadic IPF	MUC5B promoter gene polymorphism associated with improved survival independent of clinical factors [48]	Predicts survival in IPF	
uPAR	Plasminogen activator receptor augmenting monocyte adhesion	Elevated serum levels through macrophage overexpression in IPF compared to controls [49]	Discriminates IPF from healthy subjects	
TERT	Reverse transcriptase maintaining telomere integrity	Mutation associated with familial interstitial pneumonias [50] and sporadic, adult-onset IPF [51]	Discriminates familial ILDs and IPF from healthy subjects	
Telomere length	Length of nucleoprotein structures that protect chromosomal ends	Shorter telomere length associated with progression-free survival of IPF [52]	Predicts survival in IPF	
TLR3	Receptor mediating innate immune response to tissue injury, inflammation and viral infection	Polymorphism associated with early lung function decline and death [53]	Predicts progression, mortality in IPF	
α-Defensin	Antimicrobial peptides presenting in granules of neutrophils inhibiting activation of the classical complement pathway	Increased α -defensins localised in the epithelium of the lungs and apoptosis of epithelium in AE-IPF [54]	Predicts AE-IPF	

CCL18: CC chemokine ligand 18; ICAM-1: intercellular adhesion molecule 1; KL: Krebs von den Lungen; MUC: mucin; SP: surfactant protein; MMP: matrix metallopeptidase; BNP: brain natriuretic peptide; VEGF: vascular endothelial growth factor; HSP: heat shock protein; Ig: immunoglobulin; YKL40: chitinase-like protein; BLys: plasma B lymphocytes stimulating factor; CXCL: C-X-C motif chemokine ligand; EGFR: epidermal growth factor receptor; CRPM: C-reactive protein degraded by metalloproteinase-1/8; CA: cancer antigen; OSM: oncostatin M; CYFRA-21-1: cytokeratin 19 fragment; uPAR: urokinase-type plasminogen activator receptor; TERT: telomerase reverse transcriptase; TLR: toll-like receptor; ECM: extracellular matrix; ILD: interstitial lung disease; UIP: usual interstitial pneumonia; HRCT: high-resolution computed tomography; IL: interleukin; HP: hypersensitivity pneumonitis; FVC: forced vital capacity; ILA: interstitial lung abnormality; WHO: World Health Organization; VC: vital capacity; *D*Lc0: diffusing capacity of the lung for carbon monoxide; PH: pulmonary hypertension; AE-IPF: acute exacerbation of idiopathic pulmonary fibrosis; TGF: transforming growth factor. to image the distribution in airspaces as well as in the red blood cells of the vessels and in tissue of interstitial parenchyma [73].

Molecular biomarkers and genetic alterations in IPF

Several genomic biomarkers have been identified that can be predictive of the likelihood of developing IPF and thus may be helpful for diagnostic purposes [49, 54, 74, 75]. The mucin 5B promoter polymorphism (MUC5B) is a strong risk factor for developing sporadic and familial interstitial pneumonias with odds ratios as high as 21.8 (95% CI 5.1-93.5) for developing IPF in those homozygous for the T allele [48]. However, this polymorphism can be present in the normal healthy population at a significant frequency and thus is not specific for the development of IPF. Telomerase reverse transcriptase (TER-C/-T) gene mutations have been reported in familial interstitial pneumonias [50] and in sporadic, adult-onset IPF [51]. Besides TER mutations, short telomere length is also associated with outcome [52]. Toll-like receptor 3 polymorphism, a receptor mediating innate immune response to tissue injury, inflammation and infection showed associations with lung function decline and death [53]. OLDHAM et al. [76] have genotyped single nucleotide polymorphisms between toll-interacting protein (TOLLIP) and MUC5B genes that are associated with IPF susceptibility and survival, and assessed whether the polymorphisms of these genes could interfere with IPF therapies. In their post hoc study, the first significant drug-gene interaction in individuals with IPF was reported, as the TOLLIP rs3750920 TT genotype was associated with a response to N-acetylcysteine. Two studies demonstrated, that immune cell transcriptional profiles may change the response to the lung microbiome [58, 59] and showed association between genomic changes and changes in FVC, mortality and response to antifibrotic therapy [57]. However, personalised medicine is not yet established in IPF with regards to genetic alterations.

Therapeutic biomarkers

In a recent study, biomarkers were tested to interact with pirfenidone treatment and whether it could serve as prognostic, predictive or pharmacodynamic biomarker. There, pirfenidone treatment effects were consistent regardless of baseline biomarker levels, and pirfenidone treatment had no meaningful pharmacodynamic effect on the plasma levels of the prespecified biomarkers [77, 78]. In contrast, a Japanese retrospective analysis has shown prognostic effects of SP-D in an IPF cohort receiving pirfenidone [79]. Taken together, reliable predictive therapeutic biomarkers are still missing and the search for informative biomarkers in IPF must be continued [78].

New approaches in drug treatment in IPF

After several disappointing years of clinical trials of therapies that did not demonstrate efficacy in IPF (figure 1), the anti-fibrotic drugs pirfenidone and nintedanib have been associated with significant slowing of respiratory deterioration in IPF and perhaps prolonged survival [8, 17]. However, the response to antifibrotic treatment is heterogenous and may be limited by side-effects, necessitating the constant need to establish novel therapeutic approaches, including combination therapies and the development of novel compounds.

Combination therapies

The pathogenesis of IPF is a complex interplay of genetic and environmental factors activating numerous profibrotic pathways in multiple cell types [80]. Thus, it is hypothesised that targeting multiple pathways with current established therapies may show synergistic efficacy in IPF, whilst the combined adverse effects are of concern. The INJOURNEY trial [81] addressed this concern in its primary end-point investigating the safety and tolerability of combination therapy of nintedanib with add-on pirfenidone versus nintedanib alone in 105 IPF patients, over a 12-week period. The total number of adverse effects were similar in the two groups; however, nausea and vomiting occurred with greater frequency in the combination group. Despite this, adherence rates were similar in both groups, thus concluding that nintedanib plus pirfenidone therapy had a feasible safety and tolerability profile in IPF. While data have to be interpreted with caution, as the study was not appropriately powered to assess efficacy, there were promising exploratory effects on lung function decline in the combination therapy compared to nintedanib alone (12-week FVC decline of -13.3 mL versus -40.9 mL, respectively) [81]. In a further study, the safety of nintedanib added to pre-existing pirfenidone treatment in 89 IPF patients showed no new safety signals to the known safety profile of either therapy alone [82]. Previous concerns regarding drug-drug interactions between pirfenidone and nintedanib affecting pharmacokinetics and bioavailability [5] have been refuted [83]. While these data are promising with regards to tolerance and safety, an important next step would be to perform larger controlled studies to investigate the efficacy of these combination therapies. Other combination therapies in IPF have been less promising, with a phase II trial investigated the tolerability and safety of pirfenidone with add-on acetylcysteine, based on its antioxidative effects, yielding negative results [84].

Pulmonary hypertension frequently complicates severe fibrosis and is associated with diminished survival in IPF patients [84, 85]. Multiple clinical trials of monotherapies with PAH drugs targeting either pulmonary hypertension–IPF or IPF have yielded negative results [84–88]. However, a subgroup analysis of the STEP-IPF (Sildenafil Trial of Exercise Performance in Idiopathic Pulmonary Fibrosis) trial of sildenafil, a phosphodiesterase-5 inhibitor, reported some positive secondary outcomes including improvements in oxygenation, *DLCO* and, especially, HRQL [86]. These findings led to a combination study of nintedanib and sildenafil *versus* nintedanib alone, the INSTAGE study, in a severe cohort of IPF with *DLCO* <35% predicted [87]. Unfortunately, the study failed to meet its primary end-point of HRQL at 12 weeks. Yet, some exploratory outcomes reported greater differences in HRQL, FVC decline and death at 24 weeks between the two groups. However, these observations are not statistically robust and need to be taken with caution. Subgroup analyses are still pending. An ongoing phase IIb trial (NCT02951429) currently examines the effect of combined sildenafil and pirfenidone therapy in patients with advanced IPF and risk of pulmonary hypertension [88].

Novel therapies

Our understanding of the complex pathogenesis of IPF continues to grow [80]. The repetitive alveolar epithelial cell injury is increasingly recognised as a crucial mediator of the fibrotic process with a complex interplay between host and environment. The activation of multiple pathways leading to fibroblast migration, proliferation and myofibroblast differentiation has identified numerous potential molecular targets of novel therapeutic agents currently being explored in early clinical trials. Here we summarise the key studies exploring these novel therapies in IPF (table 2).

Trials with positive outcomes

PRM-151 (pentraxin-2 analogues)

Pentraxin (PTX)-2, also known as serum amyloid P, is a circulating protein that binds to monocytes and inhibits the differentiation of monocytes into pro-fibrotic fibrocytes and transforming growth factor (TGF)- β producing macrophages, thus promoting epithelial healing and resolution of fibrosis [89–91]. Low PTX-2 levels have been observed in patients with IPF [92] and a recombinant human PTX-2 analogue (PRM-151) has been shown to ameliorate fibrosis in a bleomycin- and TGF- β -overexpressing animal model of fibrosis [93]. A phase I trial (NCT01254409) assessing safety and pharmacokinetics of PRM-151 in patients with IPF has shown a nonsignificant, but improving effect on FVC and 6MWD during the treatment period [94]. A further phase II study demonstrated significant effects in reducing pulmonary function decline and stability in 6MWD over 24 weeks compared to placebo with an acceptable safety profile [95]. The launch of a phase III trial for PRM-151 in IPF has been announced, using FVC as a primary end-point and 6MWD as a key secondary end-point.

Anti-connective tissue growth factor antibodies

Connective tissue growth factor (CTGF or CCN2) is normally expressed in low levels in healthy individuals. However, when expressed in excess it leads to upregulation of TGF- β , deposition of extracellular matrix (ECM) and inhibition of ECM degradation through the inhibition of metalloproteinases [96]. All these actions of CTGF are major profibrotic factors in the development of IPF. Elevated CTGF levels are measured in the bronchoalveolar lavage (BAL) of IPF patients [97, 98]. In the PRAISE study (NCT01890265), the CTGF antagonist pamrevlumab (FG-3019) was determined to have a significant effect on the reduction of the lung function decline of 160 IPF patients, yet full peer-reviewed data are still awaited [99]. The initiation of phase III trials has currently been announced.

PBI-4050

PBI-4050 is an analogue of a medium-chain fatty acid showing affinity towards G-protein receptors targeting and inhibiting multiple pathways involved in pulmonary fibrosis including inhibition of endoplasmic reticulum stress and reactive oxygen species production, epithelial-mesenchymal transition and fibrocyte/fibroblast recruitment, migration, proliferation and differentiation [100]. A phase II trial of 40 IPF patients treated with PBI-4050 alone or in combination with antifibrotic drugs showed no safety concerns [101]. While there was slowing or stability in FVC over a 12-week treatment period with PBI-4050 alone and in the combination with nintedanib, a statistically significant greater decline in FVC was observed for the combination of PBI-4050 and pirfenidone. This was due to reductions in pharmacokinetic levels of PBI-4050 in combination with pirfenidone, suggesting a possible drug-drug interaction. Additional studies of PBI-4050 alone or in combination with nintedanib are currently being planned.

	Mechanism of action	Clinical trial identifier	Study description	Primary outcome measures	Phase of development	Treatment duration
PRM-151	Recombinant form of human SAP	NCT02550873	Randomised, double-blind, placebo controlled	Change from baseline in FVC % pred	II	28 weeks
Simtuzumab	Anti-LOX antibody	NCT01769196	Randomised, double-blind,	The effect of simtuzumab (GS-6624) on	11	148 weeks
Tipelukast	Leukotriene antagonists	NCT02503657	placebo-controlled Randomised, double-blind, placebo controlled	progression-free survival Change from baseline FVC at 26 weeks	II	26 weeks
Tralokinumab	Anti IL-13 antibody	NCT01629667	Randomised dose-ranging	Change from baseline FVC % pred at week 52	II	52 weeks
SAR156597	Anti IL-4 and IL-13 antibody	NCT01529853	Randomised, double-blind, placebo-controlled	Safety/tolerability: number of participants with adverse events	II	6 weeks
Lebrikizumab	Anti IL-13 antibody	NCT01872689	Randomised, double-blind, placebo-controlled	Annualised rate of decrease in FVC % pred over 52 weeks	II	52 weeks
BG00011	Anti-integrin antibody	NCT03573505	Randomised, double-blind, placebo-controlled	Yearly rate of change in FVC	II	52 weeks
Pamrevlumab (FG-3019)	Anti-CTGF antibody	NCT01890265	Randomised, double-blind, placebo-controlled	Change from baseline in FVC % pred at week 48	II	48 weeks
PBI-4050	GPR84 antagonist/ GPR40 agonist	NCT02538536	Open-label, single arm, exploratory, observational study	Number of subjects with abnormal laboratory values and/or adverse events that are related to treatment	II	20 weeks
KD025	Selective inhibitor of ROCK2	NCT02688647	Randomised, phase 2, open-label	Change in FVC in baseline to 24 weeks	II	24 weeks
CC-90001	Kinase inhibitor targeting JNKs	NCT03142191	Randomised, double-blind, placebo-controlled	Percentage point change in FVC % pred	II	24 weeks
GLPG1690	Autotaxin-LPA inhibitor	NCT02738801	Randomised, double-blind, parallel group, placebo-controlled	Safety, tolerability, pharmacokinetic and pharmacodynamic properties of GLPG1690	II	12 weeks
Omipalisib/ GSK2126458	Inhibitor of PI3K/Akt pathway	NCT01725139	Randomised, double-blind, placebo-controlled	To explore a number of doses of GSK2126458 for engagement of pharmacology after short-term dosing	Ι	7–10 days
Sirolimus	mTOR inhibitor	NCT01462006	Double-blind placebo-controlled pilot study	Change in peripheral blood concentration of CXCR4 ⁺ fibrocytes; number of subjects with drug side-effects	NA	22 weeks
Rituximab	Antibody targeting CD20	NCT01969409	Randomised, double-blind, placebo-controlled	Titres of anti-HEp-2 autoantibodies, by indirect immunofluorescence assays over 9 months	II	36 weeks
Co-trimoxazole or doxycycline	Antimicrobial drugs	NCT02759120	Randomised, un-blinded, phase III	Time to first non-elective, respiratory hospitalisation or all-cause mortality	111	9 months

TABLE 2 Current phase II-III trials in idiopathic pulmonary fibrosis (IPF)

SAP: serum amyloid P; FVC: forced vital capacity; LOX: lysyl oxidase; IL: interleukin; CTGF: connective tissue growth factor; GPR: G protein-coupled receptor; ROCK: ρ-associated coiled-coil containing protein kinase; JNK: Jun N-terminal kinase; LPA: lysophosphatidic acid; PI3K/Akt: phosphoinositide 3-kinase/protein kinase B. mTOR: mammalian target of rapamycin; CXCR: C-X-C chemokine receptor; HEp: human epithelial cell.

Autotaxin-LPA inhibitors

Autotaxin enzymes play a pivotal role in epithelial cell apoptosis and endothelial cell damage through the release of bioactive lysophosphatidic acid (LPA) [102]. Levels of LPA and autotaxin are increased in the BAL and exhaled breath condensates of IPF patients [103–105], and thus suggest the role of the autotaxin pathway in fibrosis. In a phase IIa study (NCT02738801, FLORA trial) the safety, tolerability, pharmacokinetic and pharmacodynamic profile of GLPG1690, a selective autotaxin and LPA inhibitor was analysed over a 12-week period. GLPG1690 was well tolerated by IPF patients with a similar safety profile, as placebo and preliminary efficacy analyses demonstrated encouraging results towards halting FVC decline [106]. Currently, international phase III trials to assess the efficacy of GLPG1690 in IPF are underway [107].

Trials with negative outcomes

Anti-lysyl oxidase antibodies

Lysyl oxidase-like 2 (LOXL2) plays an essential role in the cross-linking of collagen and elastin in the production of ECM [108] and elevated levels of LOXL2 are found in IPF [109]. A monoclonal antibody against LOXL2, simtuzumab, was examined in a recent study, where treatment was not associated with beneficial effects on progression-free survival in IPF [110].

Anti-interleukin antibodies

Interleukin (IL)-13 is a T-helper type 2 cell cytokine that has been implicated in promoting lung fibrosis in experimental models [111, 112]. Both IL-13 and IL-4 have an important role in the epithelial-fibroblast cross-talk. A number of monoclonal antibodies targeting IL-13 (the human IgG4 tralokinumab, SAR156597, an inhibitor of both IL-13 and IL-4, and lebrikizumab) have failed to show efficacy in IPF [113].

Ongoing trials

Leukotriene antagonists

Leukotrienes, due to their increased levels in IPF, may be used as therapeutic targets in the future [114, 115]. The leukotriene B4 antagonist tipelukast is currently explored in a phase II trial (NCT02503657) designed for efficacy, safety and tolerability end-points in moderate to severe IPF.

Protein kinase inhibitors

Protein kinases belong to the family of phosphorylation enzymes and inhibitors of protein kinases aim to block the action of these kinases. Several protein kinase inhibitors have been tested in IPF already, as acquired apoptosis resistance of myofibroblasts has been shown to be influenced by protein kinases [116]. A recent phase I study showed an appropriate tolerability of a selective protein kinase inhibitor of the Rho-associated coiled-coil containing protein kinase (ROCK)2. The trial is currently in its second phase II (NCT02688647), analysing further possibilities of ROCK2 [117]. A current phase II (NCT03142191) trial is evaluating CC-90001, a second-generation Jun N-terminal kinase (JNK) inhibitor for efficacy and safety after a first-generation JNK inhibitor (CC-930) showed effects on biomarker plasma levels [118].

Anti-integrin antibodies

Integrins as transmembrane receptors play a pivotal role in ECM adhesion, and integrin $\alpha\nu\beta6$ is involved in different fibrosing processes in the lungs. Based on this function, it may serve as a potential prognostic biomarker in ILDs [119]. A partial inhibition of integrin $\alpha\nu\beta6$ in rodents blocked the development of pulmonary fibrosing processes without aggravating inflammatory processes [120]. The safety and tolerability of a humanised monoclonal antibody (BG00011) against this integrin has been analysed in a phase II trial (NCT01371305). The study has been completed recently, data are still awaited.

PI3K/Akt pathway inhibitors

The phosphoinositide 3-kinase/protein kinase B (PI3K/Akt) is a main signalling pathway for cell proliferation, differentiation, adhesion and survival [121, 122]. The inhibition of these isoforms may be associated with halting fibrosing processes [123, 124]. This idea was supported by a phase I study (NCT01725139) of omipalisib, a strong inhibitor of the PI3K/Akt pathway (GSK2126458), a potential PI3K/mammalian target of rapamycin (mTOR) inhibitor [125]. In a recently published study, omipalisib showed an acceptable tolerability profile in an IPF cohort [126]. In another phase II trial (NCT01462006), sirolimus, a drug targeting mTOR is currently under examination as a possible inhibitor of fibrotic activation in IPF.

CD-20 B-lymphocyte targeting drugs

B cell abnormalities such as plasma B lymphocytes stimulating factors (BLys) are present in sera, BAL and pulmonary parenchyma of IPF patients [37, 127, 128]. The CD20 surface molecule of B lymphocytes

could potentially be targeted by rituximab [42, 129]. Rituximab is currently being assessed in a phase II study of IPF (NCT01969409); however, results are pending. Furthermore, a phase II trial investigated rituximab *versus* combined plasma exchange, and standard of care with corticosteroids as a potential treatment option in AE-IPF was performed. While peer-reviewed results are pending, an interim report stated that 14.3% of the patients showed respiratory and 42.9% haemodynamic deterioration under combined plasma exchange, rituximab and corticosteroid therapy. 42.9% of patients survived to 60 days or to transplantation and 10% showed serious adverse events, and 40% showed non-serious adverse events (NCT01266317) [130].

Targeting the microbiome

Several lines of evidence imply that in IPF progression, acute respiratory deterioration including AE-IPF and mortality are associated with altered microbiome with increased bacterial load or abundance of possibly pathogenic microorganisms [59, 131]. Recent data suggest that lung dysbiosis is associated with inflammation and aberrant lung repair [131]. Thus, targeting the microbiome, *e.g.* by antibiotic drugs might be a possible therapy to slow progression of the disease. Co-trimoxazole, mainly thought of as an antibiotic, but perhaps also associated with immunomodulatory effects, has been studied previously in a randomised trial in IPF. While in the intention-to-treat analysis, no treatment effects were evident, in a per-protocol analysis, improvements in HRQL and reduction in mortality in IPF patients who remained on antibiotic therapy were seen [132]. The impact of co-trimoxazole or doxycycline on the time to first non-elective respiratory hospitalisation and/or on all-cause mortality is being analysed in a phase III study (NCT02759120) [133].

Developments in lung transplantation

Our understanding of the pathogenesis of IPF has improved significantly in recent years and new therapeutic options are emerging in this area; however, the disease is still considered to be incurable. Lung transplantation is a viable treatment option in IPF patients who continue to decline despite current licenced therapies. However, due to the relative contraindications to transplantation, such as the presence of multiple comorbidities in an ageing population that is IPF, lung transplantation may only be appropriate for a minority of patients with IPF [134]. Previously, the International Society for Heart and Lung Transplantation recommended lung transplantation for patients aged <65 years, but the changes made in 2014 have increased this to 70 years at least in some centres [135]. In this context, the United Network for Organ Sharing registry reported comparable post-transplant outcomes in patients aged >70 years to those aged 60–69 years [136]. Therefore, age has to be viewed in the context of the overall condition of the patient. Taking into account the dreadful outcomes, *e.g.* of AE-IPF, an early referral of patients with IPF for lung transplant evaluation is mandatory in those patients that are eligible for it [137].

The challenges attributed to meeting supply with demand and having sufficient organ donation vary from country to country, thus limiting lung donor availability, and unfortunately, in some countries many IPF patients still die waiting on the lung transplant list [138-140]. Accordingly, the appropriate triage of IPF patients according to disease severity and setting-up priorities within the waiting list are of particular importance. In 2005 the United States introduced the lung allocation score (LAS), a scoring system that prioritises lung transplantation according to severity of disease and estimated post-transplant survival. This system has also been adopted by all European transplant countries. The performance of the LAS has increased the number of lung transplantation performed in patients with IPF leading to IPF becoming the most common indication for lung transplantation in the US, and therefore leading to the reduction of waiting list time [141, 142]. Advances in lung transplantation technologies can impact on transplant survival. Nowadays, single, bilateral lobar and heart-lung transplantation is available in IPF [143]. There is little evidence separating single versus bilateral lung transplantation, although the number of bilateral transplants is increasing (single 39% versus bilateral 61%) [143]. Both methods have their advantages and disadvantages. Bilateral lung transplantation has shown increased long-term survival (65.2 months, interquartile range (IQR) 21.4-91.3 months versus single 50.4 months, IQR 17.0-87.5 months) [144] and less chronic lung allograft dysfunction, while single lung transplantation is a quicker and simpler method offering less cardiac manipulation and better peri-operative outcomes [143]. In a younger age group with lower LAS score, bilateral lung transplantation appears advantageous [145]. However, other factors can influence post-transplant survival. Pre-transplant IPF disease severity, use of extracorporeal membrane oxygenation (ECMO) and the presence of pulmonary hypertension or depression, as well as poorer quality of life and higher physiological distress 6 months post-transplant can result in poorer post-transplant survival [146, 147]. ECMO can be indicated in suitably robust IPF patients with severe respiratory failure who deteriorate rapidly despite maximal medical therapy [148]. A recent study demonstrated that patients undergoing ambulatory pre-transplant ECMO therapy had better post-transplantation outcomes than those undergoing lung transplantation from mechanical ventilation [149, 150].

Open questions remain regarding the continuation of antifibrotic therapy prior to lung transplantation. Data from small cohort studies suggest that antifibrotic drugs can be safely continued until lung transplantation [151, 152] and pirfenidone and nintedanib may also decrease the waiting-list mortality due to their disease progression attenuating effects [153]. Genetic markers such as shorter telomere length can also impact the post-transplantation prognosis by increasing the risk of post-transplant infections such as viral cytomegalovirus and thereby increasing the risks of allograft rejection. Genetic phenotyping of transplanted patients is therefore of importance.

Pulmonary rehabilitation

For years, pulmonary rehabilitation has been considered to have limited effect on patients with severe, chronic lung diseases such as IPF [1, 154], primarily due to the low number of articles on rehabilitation, small patient numbers and the lack of standardisation of the length and intensity of pulmonary rehabilitation programmes, making comparisons difficult. Following extensive growing research in this area, pulmonary rehabilitation is now recognised as an integral and essential component of management of patients with IPF and thus is a major feature of recommendations in national and international guidelines [2, 155]. A systematic Cochrane review exploring the impact of pulmonary rehabilitation in IPF reported that improvements of HRQL could be seen almost immediately after starting pulmonary rehabilitation while no adverse events have been reported [141]. However, no clear conclusions can be drawn in severe ILD and data on long-term effects of pulmonary rehabilitation in this advanced cohort are still sparse [156]. Current evidence on pulmonary rehabilitation in IPF shows significant short-term effects on improving exercise capacity (6MWD) and HRQL, while long-term effects are not maintained [157]. In addition, pulmonary rehabilitation has been shown to be useful in patients referred to lung transplantation [158]. Pulmonary rehabilitation should also include nutritional support [1], non-exercise components such as education [159], psychological [160] and symptom management [161], which are all of great importance in IPF.

Ongoing clinical trials are focusing on the outcomes of pulmonary rehabilitation in IPF. A study tested the 6MWD as the primary outcome and quality-of-life changes as secondary outcomes between active and inactive patient groups after a 12-week pulmonary rehabilitation programme (NCT03542318). The active group presented significantly better functional and health status after pulmonary rehabilitation [162]. Another ongoing study is testing the same outcomes comparing nintedanib *versus* nintedanib plus pulmonary rehabilitation patient groups (NCT03717012). A further ongoing trial aimed to determine the short- and long-term effects of oxygen supplementation during pulmonary rehabilitation in IPF (NCT03326089).

New ways to approach palliative intervention

Palliative interventions encompass a broad therapeutic spectrum including education and support of patients and caregivers, early management of symptoms with the goal of improving or maintaining quality of life of patients and end-of-life planning, in a disease which is invariably progressive and life-limiting [21]. Symptoms such as cough, breathlessness, anxiety and depression are common in IPF [163] and have a major impact on the quality of life of patients [164]. It has been proven that lower daily physical activity is associated with worse survival rates [165]. Palliative and supportive interventions in IPF with nonpharmacological and pharmacological interventions aim to improve these major symptoms, and are discussed briefly here.

Breathlessness can be an emotionally frightening and disabling symptom affecting mobility and quality of life and a breathing, thinking and functioning clinical model has been proposed to tackle this symptom [166]. Pulmonary rehabilitation is just one of many interventions that has been used to improve breathlessness. A multicentre prospective study of pulmonary rehabilitation has demonstrated improvements in the St George's respiratory questionnaire, 6-min walk test and breathlessness scores (Medical Research Council and Borg) [167]. Theoretically, oxygen therapy may be useful in improving dyspnoea at rest and during exertion and improving HRQL in IPF; however, studies demonstrating this benefit are limited.

International guidelines recommend the use of long-term oxygen therapy (LTOT) in IPF [1] with low-quality evidence. Oxygen therapy can increase self-confidence and active participation in various daily activities [168]; however, some limitations are described in daily life [169] and patients may develop a fear of being without oxygen after a longer period on LTOT [168]. Currently, there is no evidence of oxygen therapy impacting survival of ILD patients [170]. Conversely, ambulatory oxygen therapy (AmbOx) has been shown to be associated with improvements in HRQL in hypoxic ILD patients [171] and prolonged endurance time at ergometry. However, there was no impact on dyspnoea [172]. Its effect on exercise capacity varies in different studies [170, 172]. Therefore, AmbOx may represent an effective therapeutic option in this disease group.

The evidence for pharmacological therapies such as opiates and benzodiazepines for breathlessness management in IPF is scarce. A recent study reported that low-dose diamorphine reduced breathlessness without causing a fall in oxygen saturation in an elderly cohort with end-stage IPF [173]. As oral morphine therapy reduced dyspnoea and did not cause respiratory depression, low-dose opioids may be effective and safe in the palliative management of IPF [174]. Historically, benzodiazepines have been used for managing dyspnoea, but a Cochrane review of 214 participants with varied chronic diseases showed no benefit of benzodiazepines in ameliorating breathlessness. However, conclusions are limited due to the heterogeneity of studies with a limited number of participants [175].

Cough is present in >80% of IPF patients and an independent predictor of progression in IPF [176]. The pathophysiology and mechanism of cough in IPF is complex and poorly understood and treatment includes the management of comorbidities that can influence cough, such as gastro-oesophageal reflux disease (GORD) [176]. Opioid therapies significantly ameliorate cough and are associated with an improved quality of life [177]. However, the therapeutic impact of opioids on reducing cough is extrapolated from its positive findings in chronic cough patients, and there is limited evidence regarding the safety and efficacy of opioids for the management of cough in the terminal stages and palliative treatment of IPF. Drugs specifically targeting cough in IPF have included thalidomide. Due to its immunomodulatory and anti-inflammatory effects, thalidomide was evaluated in a double-blinded, placebo-controlled trial in patients with IPF and cough. This 24-week cross-over trial in 23 IPF patients revealed that thalidomide significantly reduced cough visual analogue scores and cough-specific quality of life measures compared to placebo [178, 179]. However, thalidomide-treated patients experienced more adverse events compared to placebo (77% versus 22%). The occurrence of these adverse events and the small sample size has limited its use in cough, and thus larger trials are required to further prove it safety and efficacy. Anecdotally, some IPF patients feel their cough is improved with antifibrotic therapy. In an open-label 12-week study of cough in IPF, the antifibrotic pirfenidone has been shown to reduce objective cough measurements with 24 h cough counts, as well as subjective improvements in cough visual analogue scores. However, this study was not a placebo-controlled study, thus reflecting the limitations of the study [180].

Future novel agents include a pilot study of nebulised sodium cromoglicate (PA101) in IPF reporting to be effective in patients with chronic cough [181]. Currently, an ongoing phase 2b trial was initiated to assess the potential role of inhaled cromolyn sodium (RVT-1601, formerly PA101B) in the therapy of persistent chronic cough in IPF (NCT03864328).

Treatment of comorbidities

While IPF is the main reason for death in patients suffering from this chronic disease, the cause of death in 30–40% of these patients is related to other conditions, *i.e.* comorbidities [182]. Cardiovascular and thromboembolic disorders, GORD, depression, sleep disorders, pulmonary hypertension, emphysema, diabetes and lung cancer are the most common comorbidities in IPF [183]. This association might be explained partially by the elderly population, which is mainly affected by IPF [184]. Yet, many comorbidities worsen the prognosis of the disease [185, 186], reflected by a recent comorbidome (figure 2) illustrating the association of frequencies and groups of diseases to prognosis.

Several studies have focused on the relationship between comorbidities and the burden of disease [85, 187-190]. The German INSIGHTS IPF registry has determined that an increased number of comorbidities decreases HRQL [191]. Recently, the novel comorbidity assessment TORVAN model has shown that comorbidities may refer to a lower survival rate based on clinical and physiological parameters as in the GAP index [192]. While no effects of antifibrotic drugs on comorbidities have been reported so far, several reports focused on the effects of comedications or treatments of comorbidities in IPF. Three studies have reported an association between worsened prognosis and the use of anticoagulants, mainly warfarin, for reasons other than IPF [188, 193, 194]. Recent studies suggested that a new strategy of anticoagulation may have positive effects. Namely, the onset of direct thrombin inhibitors may even show antifibrotic effects that open new avenues in antithrombotic therapy in IPF [195-198]. GORD is a frequent comorbidity in IPF. Based on possible favourable effects of anti-acid therapy on meaningful end-points in IPF, the recent international IPF guideline recommends antacid therapies for all IPF patients. However, owing to newer conflicting data, several national guidelines do not support this recommendation [199-202]. Another approach to GORD in IPF is laparoscopic surgical treatment. Here, a phase II study (WRAP-IPF) showed clinically meaningful impact on lung function and further improvements of other important end-points like respiratory hospitalisations in the group of patients after surgical intervention [203]. Statins reduce cholesterol levels to reduce the risk of cardiovascular morbidity [204, 205]. In addition to their anti-inflammatory effects, they may have a protective effect against smoking and slow down the deterioration of age-associated lung function [206]. In a post hoc analysis, IPF patients, who

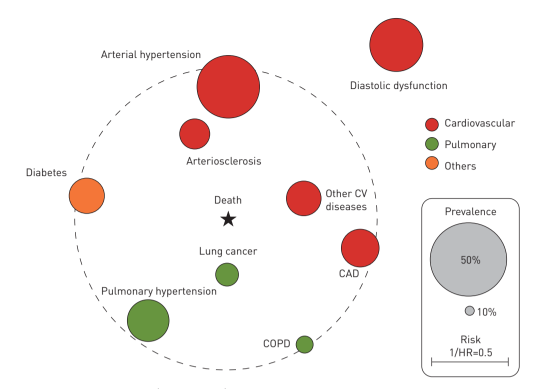


FIGURE 2 Graphic expression (comorbidome) of comorbidities with >10% prevalence in the entire cohort, and those comorbidities with the strongest association with mortality (hazard ratio (HR) >1, 95% CI >1; p<0.05) [166]. The area of the circle relates to the prevalence of the disease. The proximity to the centre (mortality) expresses the strength of the association between the disease and risk of death. This was scaled from the inverse of the HR (1/HR). All bubbles associated with a statistically significant increase in mortality are fully inside the dotted orbit (1/HR <1). Bubble colours represent organ systems or disease clusters. CV: cardiovascular; CAD: coronary artery disease; COPD: chronic obstructive pulmonary disease. Reproduced and modified from [186] with permission.

were on statins, had a better prognosis, lower hospitalisation rates, and mortality compared to those who did not receive statin therapy [207, 208]. Also, newest data show some effects of ACE-inhibitors on outcomes in IPF [209].

Urgent unmet need for AE-IPF

AE-IPF is a frequent and severe complication of IPF associated with poor prognosis, especially high hospital mortality [210]. AE-IPF can occur at any time, but is more common in advanced stages of the disease [211]. Idiopathic and triggered, *e.g.* infection-associated, AE-IPF have a similar clinical appearance and outcome [212].

Due to the lack of robust evidence, there is no accepted guideline for the treatment of acute exacerbations, thus practice for AE-IPF varies in different centres [213]. Recently, it was demonstrated in a large international survey that corticosteroids combined with broad-spectrum antibiotics including macrolides are widely used. Yet, doses, duration and route of steroids vary significantly between centres, while only a minority never use steroids. Retrospective data suggest that ECMO has no meaningful effects on the prognosis of patients who are not eligible for lung transplantation [214]. Preventive strategies such as vaccination and antifibrotics are extensively adopted in AE-IPF without clinical evidence [213, 215]. Future research is required to develop strategies for diagnosis, prevention and treatment of AE-IPF.

Conclusions

In recent years, our understanding of the pathophysiology of IPF and potential therapies for IPF have significantly evolved, yet there is still no halt to disease progression and no cure. Future therapies should aim to stabilise the disease, ameliorate symptoms and improve quality of life with the ultimate goal of reducing the burden of disease. Our long-term goal must be to reverse this devastating disease and discover novel ways to cure it, in addition to finding new effective therapies for a major unmet need and the deadliest complication of IPF, acute exacerbations. As such, currently several clinical trials are under way and focusing on new therapeutic approaches. Furthermore, the research field in IPF is expanding to

focus on approaches to improve nonpharmacological therapies to improve quality of life such as rehabilitation, symptom management, enhanced management of comorbidities and a better understanding of the effects of comedications. These advances in research pave the way for an interesting few decades in IPF diagnosis and management.

Conflict of interest: None declared.

References

- 1 Raghu G, Collard HR, Egan JJ, et al. An official ATS/ERS/JRS/ALAT statement: idiopathic pulmonary fibrosis: evidence-based guidelines for diagnosis and management. Am J Respir Crit Care Med 2011; 183: 788–824.
- 2 Raghu G, Remy-Jardin M, Myers JL, et al. Diagnosis of idiopathic pulmonary fibrosis. An official ATS/ERS/JRS/ ALAT clinical practice guideline. Am J Respir Crit Care Med 2018; 198: e44–e68.
- 3 Raghu G, Idiopathic pulmonary fibrosis: lessons from clinical trials over the past 25 years. *Eur Respir J* 2017; 50: 1701209.
- 4 Raghu G, Johnson WC, Lockhart D, *et al.* Treatment of idiopathic pulmonary fibrosis with a new antifibrotic agent, pirfenidone: results of a prospective, open-label phase II study. *Am J Respir Crit Care Med* 1999; 159: 1061–1069.
- 5 Azuma A, Nukiwa T, Tsuboi E, *et al.* Double-blind, placebo-controlled trial of pirfenidone in patients with idiopathic pulmonary fibrosis. *Am J Respir Crit Care Med* 2005; 171: 1040–1047.
- 6 Taniguchi H, Ebina M, Kondoh Y, et al. Pirfenidone in idiopathic pulmonary fibrosis. Eur Respir J 2010; 35: 821–829.
- 7 Noble PW, Albera C, Bradford WZ, *et al.* Pirfenidone in patients with idiopathic pulmonary fibrosis (CAPACITY): two randomised trials. *Lancet* 2011; 377: 1760–1769.
- 8 King TE Jr, Bradford WZ, Castro-Bernardini S, et al. A phase 3 trial of pirfenidone in patients with idiopathic pulmonary fibrosis. N Engl J Med 2014; 370: 2083–2092.
- 9 Noble PW, Albera C, Bradford WZ, *et al.* Pirfenidone for idiopathic pulmonary fibrosis: analysis of pooled data from three multinational phase 3 trials. *Eur Respir J* 2016; 47: 243–253.
- 10 Nathan SD, Albera C, Bradford WZ, et al. Effect of pirfenidone on mortality: pooled analyses and meta-analyses of clinical trials in idiopathic pulmonary fibrosis. Lancet Respir Med 2017; 5: 33–41.
- 11 Ley B, Swigris J, Day BM, et al. Pirfenidone reduces respiratory-related hospitalizations in idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2017; 196: 756–761.
- 12 Albera C, Costabel U, Fagan EA, *et al.* Efficacy of pirfenidone in patients with idiopathic pulmonary fibrosis with more preserved lung function. *Eur Respir J* 2016; 48: 843–851.
- 13 Costabel U, Bendstrup E, Cottin V, *et al.* Pirfenidone in idiopathic pulmonary fibrosis: expert panel discussion on the management of drug-related adverse events. *Adv Ther* 2014; 31: 375–391.
- 14 Wollin L, Maillet I, Quesniaux V, *et al.* Antifibrotic and anti-inflammatory activity of the tyrosine kinase inhibitor nintedanib in experimental models of lung fibrosis. *J Pharmacol Exp Ther* 2014; 349: 209–220.
- 15 Hilberg F, Roth GJ, Krssak M, *et al.* BIBF 1120: triple angiokinase inhibitor with sustained receptor blockade and good antitumor efficacy. *Cancer Res* 2008; 68: 4774–4782.
- 16 Richeldi L, Costabel U, Selman M, et al. Efficacy of a tyrosine kinase inhibitor in idiopathic pulmonary fibrosis. N Engl J Med 2011; 365: 1079–1087.
- 17 Richeldi L, du Bois RM, Raghu G, *et al.* Efficacy and safety of nintedanib in idiopathic pulmonary fibrosis. *N Engl J Med* 2014; 370: 2071–2082.
- 18 Brown KK, Flaherty KR, Cottin V, et al. Lung function outcomes in the INPULSIS* trials of nintedanib in idiopathic pulmonary fibrosis. Respir Med 2019; 146: 42–48.
- 19 Raghu G, Wells AU, Nicholson AG, et al. Effect of nintedanib in subgroups of idiopathic pulmonary fibrosis by diagnostic criteria. Am J Respir Crit Care Med 2017; 195: 78–85.
- 20 Wuyts WA, Kolb M, Stowasser S, *et al.* First data on efficacy and safety of nintedanib in patients with idiopathic pulmonary fibrosis and forced vital capacity of ≤50% of predicted value. *Lung* 2016; 194: 739–743.
- 21 Kreuter M, Bendstrup E, Russell AM, et al. Palliative care in interstitial lung disease: living well. Lancet Respir Med 2017; 5: 968–980.
- 22 Prasse A, Probst C, Bargagli E, *et al.* Serum CC-chemokine ligand 18 concentration predicts outcome in idiopathic pulmonary fibrosis. *Am J Respir Crit Care Med* 2009; 179: 717–723.
- 23 Richards TJ, Kaminski N, Baribaud F, et al. Peripheral blood proteins predict mortality in idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2012; 185: 67–76.
- 24 Horimasu Y, Hattori N, Ishikawa N, et al. Different MUC1 gene polymorphisms in German and Japanese ethnicities affect serum KL-6 levels. *Respir Med* 2012; 106: 1756–1764.
- 25 Saini G, Jenkins G, Mckeever T, et al. The PROFILE study: a prospective study of fibrosis in lung endpoints to discover and qualify biomarkers for use in clinical trials. Am J Respir Crit Care Med 2012; 185: A5169.
- 26 Yokoyama A, Kondo K, Nakajima M, et al. Prognostic value of circulating KL-6 in idiopathic pulmonary fibrosis. Respirology 2006; 11: 164–168.
- 27 Takahashi H, Fujishima T, Koba H, et al. Serum surfactant proteins A and D as prognostic factors in idiopathic pulmonary fibrosis and their relationship to disease extent. Am J Respir Crit Care Med 2000; 162: 1109–1114.
- 28 Kinder BW, Brown KK, McCormack FX, et al. Serum surfactant protein-A is a strong predictor of early mortality in idiopathic pulmonary fibrosis. Chest 2009; 135: 1557–1563.
- 29 Greene KE, Ye S, Mason RJ, et al. Serum surfactant protein-A levels predict development of ARDS in at-risk patients. Chest 1999; 116: 90S-91S.
- 30 Nagata N, Kitasato Y, Wakamatsu K, *et al.* Prognostic value of immunohistochemical surfactant protein A expression in regenerative/hyperplastic alveolar epithelial cells in idiopathic interstitial pneumonias. *Diagn Pathol* 2011; 6: 25.
- 31 Kahn N, Rossler AK, Hornemann K, *et al.* C-proSP-B: a possible biomarker for pulmonary diseases? *Respiration* 2018; 96: 117–126.
- 32 Vij R, Noth I. Peripheral blood biomarkers in idiopathic pulmonary fibrosis. Transl Res 2012; 159: 218–227.

- 33 Armstrong HF, Podolanczuk AJ, Barr RG, et al. Serum matrix metalloproteinase-7, respiratory symptoms, and mortality in community-dwelling adults. MESA (Multi-Ethnic Study of Atherosclerosis). Am J Respir Crit Care Med 2017; 196: 1311–1317.
- 34 Song JW, Song JK, Kim DS. Echocardiography and brain natriuretic peptide as prognostic indicators in idiopathic pulmonary fibrosis. *Respir Med* 2009; 103: 180-186.
- 35 Ando M, Miyazaki E, Ito T, et al. Significance of serum vascular endothelial growth factor level in patients with idiopathic pulmonary fibrosis. Lung 2010; 188: 247–252.
- 36 Gilani SR, Vuga LJ, Lindell KO, et al. CD28 down-regulation on circulating CD4 T-cells is associated with poor prognoses of patients with idiopathic pulmonary fibrosis. PLoS One 2010; 5: e8959.
- 37 Kahloon RA, Xue J, Bhargava A, et al. Patients with idiopathic pulmonary fibrosis with antibodies to heat shock protein 70 have poor prognoses. Am J Respir Crit Care Med 2013; 187: 768–775.
- 38 Okamoto M, Hoshino T, Kitasato Y, *et al.* Periostin, a matrix protein, is a novel biomarker for idiopathic interstitial pneumonias. *Eur Respir J* 2011; 37: 1119–1127.
- 39 Tajiri M, Okamoto M, Fujimoto K, et al. Serum level of periostin can predict long-term outcome of idiopathic pulmonary fibrosis. Respir Investig 2015; 53: 73–81.
- 40 Kadota J, Mizunoe S, Mito K, *et al.* High plasma concentrations of osteopontin in patients with interstitial pneumonia. *Respir Med* 2005; 99: 111–117.
- 41 Korthagen NM, van Moorsel CH, Zanen P, *et al.* Evaluation of circulating YKL-40 levels in idiopathic interstitial pneumonias. *Lung* 2014; 192: 975–980.
- 42 Xue J, Kass DJ, Bon J, et al. Plasma B lymphocyte stimulator and B cell differentiation in idiopathic pulmonary fibrosis patients. J Immunol 2013; 191: 2089–2095.
- 43 Moeller A, Gilpin SE, Ask K, *et al.* Circulating fibrocytes are an indicator of poor prognosis in idiopathic pulmonary fibrosis. *Am J Respir Crit Care Med* 2009; 179: 588–594.
- 44 Vuga LJ, Tedrow JR, Pandit KV, et al. C-X-C motif chemokine 13 (CXCL13) is a prognostic biomarker of idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2014; 189: 966–974.
- 45 Jenkins RG, Simpson JK, Saini G, *et al.* Longitudinal change in collagen degradation biomarkers in idiopathic pulmonary fibrosis: an analysis from the prospective, multicentre PROFILE study. *Lancet Respir Med* 2015; 3: 462–472.
- 46 Maher TM, *et al.* The PROFILE (Prospective Observation of Fibrosis in the Lung Clinical Endpoints) study. *Am J Respir Crit Care Med* 2017; 195: A6790.
- 47 Simpson JK, Maher TM, Bentley J, *et al.* CYFRA-21-1 as a biomarker with prognostic potential in idiopathic pulmonary fibrosis: an analysis of the PROFILE cohort. *Am J Respir Crit Care Med* 2017; 195: A6791.
- 48 Seibold MA, Wise AL, Speer MC, et al. A common MUC5B promoter polymorphism and pulmonary fibrosis. N Engl J Med 2011; 364: 1503–1512.
- 49 Desai B, Mattson J, Paintal H, et al. Differential expression of monocyte/macrophage-selective markers in human idiopathic pulmonary fibrosis. Exp Lung Res 2011; 37: 227–238.
- 50 Armanios MY, Chen JJ, Cogan JD, *et al.* Telomerase mutations in families with idiopathic pulmonary fibrosis. *N Engl J Med* 2007; 356: 1317–1326.
- 51 Tsakiri KD, Cronkhite JT, Kuan PJ, *et al.* Adult-onset pulmonary fibrosis caused by mutations in telomerase. *Proc Natl Acad Sci USA* 2007; 104: 7552–7557.
- 52 Stuart BD, Lee JS, Kozlitina J, *et al.* Effect of telomere length on survival in patients with idiopathic pulmonary fibrosis: an observational cohort study with independent validation. *Lancet Respir Med* 2014; 2: 557–565.
- 53 O'Dwyer DN, Armstrong ME, Trujillo G, *et al.* The Toll-like receptor 3 L412F polymorphism and disease progression in idiopathic pulmonary fibrosis. *Am J Respir Crit Care Med* 2013; 188: 1442–1450.
- 54 Konishi K, Gibson KF, Lindell KO, *et al.* Gene expression profiles of acute exacerbations of idiopathic pulmonary fibrosis. *Am J Respir Crit Care Med* 2009; 180: 167–175.
- 55 Durheim MT, Kim S, Gulack BC, et al. Mortality and respiratory failure after thoracoscopic lung biopsy for interstitial lung disease. Ann Thorac Surg 2017; 104: 465–470.
- 56 Korthagen NM, van Moorsel CH, Barlo NP, *et al.* Serum and BALF YKL-40 levels are predictors of survival in idiopathic pulmonary fibrosis. *Respir Med* 2011; 105: 106–113.
- 57 Herazo-Maya JD, Sun J, Molyneaux PL, *et al.* Validation of a 52-gene risk profile for outcome prediction in patients with idiopathic pulmonary fibrosis: an international, multicentre, cohort study. *Lancet Respir Med* 2017; 5: 857–868.
- 58 Huang Y, Ma SF, Espindola MS, *et al.* Microbes are associated with host innate immune response in idiopathic pulmonary fibrosis. *Am J Respir Crit Care Med* 2017; 196: 208–219.
- 59 Molyneaux PL, Willis-Owen SAG, Cox MJ, et al. Host-microbial interactions in idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2017; 195: 1640-1650.
- 60 Satoh H, Kurishima K, Ishikawa H, *et al.* Increased levels of KL-6 and subsequent mortality in patients with interstitial lung diseases. *J Intern Med* 2006; 260: 429–434.
- 61 Homma S, Bando M, Azuma A, *et al.* Japanese guideline for the treatment of idiopathic pulmonary fibrosis. *Respir Investig* 2018; 56: 268–291.
- 62 du Bois RM, Weycker D, Albera C, *et al.* Forced vital capacity in patients with idiopathic pulmonary fibrosis: test properties and minimal clinically important difference. *Am J Respir Crit Care Med* 2011; 184: 1382–1389.
- 63 King TE Jr, Tooze JA, Schwarz MI, *et al.* Predicting survival in idiopathic pulmonary fibrosis: scoring system and survival model. *Am J Respir Crit Care Med* 2001; 164: 1171–1181.
- 64 Lee SH, Park JS, Kim SY, *et al.* Comparison of CPI and GAP models in patients with idiopathic pulmonary fibrosis: a nationwide cohort study. *Sci Rep* 2018; 8: 4784.
- 65 Torrisi SE, Ley B, Kreuter M, *et al.* The added value of comorbidities in predicting survival in idiopathic pulmonary fibrosis: a multicentre observational study. *Eur Respir J* 2019; 53: 1801587.
- 66 Maher TM, Stowasser S, Nishioka Y, *et al.* Investigating the effects of nintedanib on biomarkers of extracellular matrix turnover in patients with IPF: design of the randomised placebo-controlled INMARK^{*} trial. *BMJ Open Respir Res* 2018; 5: e000325.
- 67 Wu X, Kim GH, Salisbury ML, *et al.* Computed tomographic biomarkers in idiopathic pulmonary fibrosis. The future of quantitative analysis. *Am J Respir Crit Care Med* 2019; 199: 12–21.

- 68 Humphries SM, Yagihashi K, Huckleberry J, *et al.* Idiopathic pulmonary fibrosis: data-driven textural analysis of extent of fibrosis at baseline and 15-month follow-up. *Radiology* 2017; 285: 270–278.
- 69 Walsh SLF. Imaging biomarkers and staging in IPF. Curr Opin Pulm Med 2018; 24: 445-452.
- 70 Clukers J, Lanclus M, Mignot B, et al. Quantitative CT analysis using functional imaging is superior in describing disease progression in idiopathic pulmonary fibrosis compared to forced vital capacity. Respir Res 2018; 19: 213.
- 71 Loeh B, Brylski LT, von der Beck D, *et al.* Lung CT densitometry in idiopathic pulmonary fibrosis for the prediction of natural course, severity and mortality. *Chest* 2019; 155: 972–981.
- 72 Jacob J, Bartholmai BJ, Rajagopalan S, et al. Predicting outcomes in idiopathic pulmonary fibrosis using automated computed tomographic analysis. Am J Respir Crit Care Med 2018; 198: 767–776.
- 73 Mammarappallil JG, Rankine L, Wild JM, *et al.* New developments in imaging idiopathic pulmonary fibrosis with hyperpolarized xenon magnetic resonance imaging. *J Thorac Imaging* 2019; 34: 136–150.
- 74 Hambly N, Shimbori C, Kolb M. Molecular classification of idiopathic pulmonary fibrosis: personalized medicine, genetics and biomarkers. *Respirology* 2015; 20: 1010–1022.
- 75 Chiba H, Otsuka M, Takahashi H. Significance of molecular biomarkers in idiopathic pulmonary fibrosis: a mini review. *Respir Investig* 2018; 56: 384–391.
- 76 Oldham JM, Ma SF, Martinez FJ, et al. TOLLIP, MUC5B, and the response to N-acetylcysteine among individuals with idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2015; 192: 1475–1482.
- 77 Neighbors M, Cabanski CR, Ramalingam TR, *et al.* Prognostic and predictive biomarkers for patients with idiopathic pulmonary fibrosis treated with pirfenidone: *post-hoc* assessment of the CAPACITY and ASCEND trials. *Lancet Respir Med* 2018; 6: 615–626.
- 78 Kreuter M, Maher T. Gazing into the crystal ball: can treatment response be predicted in IPF? *Lancet Respir Med* 2018; 6: 570–572.
- 79 Ikeda K, Shiratori M, Chiba H, et al. Serum surfactant protein D predicts the outcome of patients with idiopathic pulmonary fibrosis treated with pirfenidone. *Respir Med* 2017; 131: 184–191.
- 80 Lederer DJ, Martinez FJ. Idiopathic pulmonary fibrosis. N Engl J Med 2018; 378: 1811–1823.
- 81 Vancheri C, Kreuter M, Richeldi L, *et al.* Nintedanib with add-on pirfenidone in idiopathic pulmonary fibrosis. Results of the INJOURNEY trial. *Am J Respir Crit Care Med* 2018; 197: 356–363.
- 82 Flaherty KR, Fell CD, Huggins JT, *et al.* Safety of nintedanib added to pirfenidone treatment for idiopathic pulmonary fibrosis. *Eur Respir J* 2018; 52: 1800230.
- 83 Richeldi L, Fletcher S, Adamali H *et al.* No relevant pharmacokinetic drug–drug interaction between nintedanib and pirfenidone. *Eur Respir J* 2019; 53: 1801060.
- 84 Behr J, Bendstrup E, Crestani B, *et al.* Safety and tolerability of acetylcysteine and pirfenidone combination therapy in idiopathic pulmonary fibrosis: a randomised, double-blind, placebo-controlled, phase 2 trial. *Lancet Respir Med* 2016; 4: 445–453.
- 85 Lettieri CJ, Nathan SD, Barnett SD, et al. Prevalence and outcomes of pulmonary arterial hypertension in advanced idiopathic pulmonary fibrosis. Chest 2006; 129: 746–752.
- 86 Swigris JJ, Han M, Vij R, et al. The UCSD shortness of breath questionnaire has longitudinal construct validity in idiopathic pulmonary fibrosis. Respir Med 2012; 106: 1447–1455.
- 87 Kolb M, Raghu G, Wells AU, et al. Nintedanib plus sildenafil in patients with idiopathic pulmonary fibrosis. N Engl J Med 2018; 379: 1722–1731.
- 88 Behr J, Nathan SD, Harari S, et al. Sildenafil added to pirfenidone in patients with advanced idiopathic pulmonary fibrosis and risk of pulmonary hypertension: a phase IIb, randomised, double-blind, placebo-controlled study – rationale and study design. Respir Med 2018; 138: 13–20.
- 89 Pilling D, Gomer RH Regulatory pathways for fibrocyte differentiation. In: Bucala R, Ed. Fibrocytes: New Insights into Tissue Repair and Systemic Fibroses. Singapore, World Scientific Publishing, 2007; pp. 37–60.
- 90 Pilling D, Buckley CD, Salmon M, *et al.* Inhibition of fibrocyte differentiation by serum amyloid P. J Immunol 2003; 171: 5537–5546.
- 91 Pilling D, Roife D, Wang M, *et al.* Reduction of bleomycin-induced pulmonary fibrosis by serum amyloid P. *J Immunol* 2007; 179: 4035–4044.
- 92 Dillingh MR, van den Blink B, Moerland M, et al. Recombinant human serum amyloid P in healthy volunteers and patients with pulmonary fibrosis. Pulm Pharmacol Ther 2013; 26: 672–676.
- 93 Murray LA, Rosada R, Moreira AP, et al. Serum amyloid P therapeutically attenuates murine bleomycin-induced pulmonary fibrosis via its effects on macrophages. PLoS One 2010; 5: e9683.
- 94 van den Blink B, Dillingh MR, Ginns LC, *et al.* Recombinant human pentraxin-2 therapy in patients with idiopathic pulmonary fibrosis: safety, pharmacokinetics and exploratory efficacy. *Eur Respir J* 2016; 47: 889–897.
- 95 Raghu G, van den Blink B, Hamblin MJ, et al. Effect of recombinant human pentraxin 2 vs placebo on change in forced vital capacity in patients with idiopathic pulmonary fibrosis: a randomized clinical trial. JAMA 2018; 319: 2299–2307.
- 96 Kono M, Nakamura Y, Suda T, *et al.* Plasma CCN2 (connective tissue growth factor; CTGF) is a potential biomarker in idiopathic pulmonary fibrosis (IPF). *Clin Chim Acta* 2011; 412: 2211–2215.
- 97 Bonniaud P, Martin G, Margetts PJ, *et al.* Connective tissue growth factor is crucial to inducing a profibrotic environment in "fibrosis-resistant" BALB/c mouse lungs. *Am J Respir Cell Mol Biol* 2004; 31: 510–516.
- 98 Wang X, Wu G, Gou L, et al. A novel single-chain-Fv antibody against connective tissue growth factor attenuates bleomycin-induced pulmonary fibrosis in mice. Respirology 2011; 16: 500–507.
- 99 Gorina E, Richeldi L, Raghu G, *et al.* PRAISE, a randomized, placebo-controlled, double-blind phase 2 clinical trial of pamrevlumab (FG-3019) in IPF patients. *Eur Respir J* 2017; 50: OA3400.
- 100 Gagnon L, Leduc M, Thibodeau JF, *et al.* A newly discovered antifibrotic pathway regulated by two fatty acid receptors: GPR40 and GPR84. *Am J Pathol* 2018; 188: 1132–1148.
- 101 Khalil N, Manganas H, Ryerson CJ, et al. Phase 2 clinical trial of PBI-4050 in patients with idiopathic pulmonary fibrosis. Eur Respir J 2019; 53: 1800663.
- 102 Aoki J, Inoue A, Okudaira S. Two pathways for lysophosphatidic acid production. *Biochim Biophys Acta* 2008; 1781: 513–518.
- 103 Tager AM, LaCamera P, Shea BS, *et al.* The lysophosphatidic acid receptor LPA1 links pulmonary fibrosis to lung injury by mediating fibroblast recruitment and vascular leak. *Nat Med* 2008; 14: 45–54.

- 104 Oikonomou N, Mouratis MA, Tzouvelekis A, *et al.* Pulmonary autotaxin expression contributes to the pathogenesis of pulmonary fibrosis. *Am J Respir Cell Mol Biol* 2012; 47: 566–574.
- 105 Montesi SB, Mathai SK, Brenner LN, *et al.* Docosatetraenoyl LPA is elevated in exhaled breath condensate in idiopathic pulmonary fibrosis. *BMC Pulm Med* 2014; 14: 5.
- 106 Maher TM, van der Aar EM, Van de Steen O, *et al.* Safety, tolerability, pharmacokinetics, and pharmacodynamics of GLPG1690, a novel autotaxin inhibitor, to treat idiopathic pulmonary fibrosis (FLORA): a phase 2a randomised placebo-controlled trial. *Lancet Respir Med* 2018; 6: 627–635.
- 107 Maher T, Kreuter M, Lederer D, *et al.* Rationale, design and objectives of two phase III, randomised, placebo-controlled studies of GLPG1690, a novel autotaxin inhibitor, in idiopathic pulmonary fibrosis (ISABELA 1 and 2). *BMJ Open Respir Res* 2019; 6: e000422.
- 108 Moon HJ, Finney J, Ronnebaum T, et al. Human lysyl oxidase-like 2. Bioorg Chem 2014; 57: 231-241.
- 109 Chien JW, Richards TJ, Gibson KF, et al. Serum lysyl oxidase-like 2 levels and idiopathic pulmonary fibrosis
- disease progression. *Eur Respir J* 2014; 43: 1430–1438.
 Raghu G, Brown KK, Collard HR, *et al.* Efficacy of simtuzumab *versus* placebo in patients with idiopathic pulmonary fibrosis: a randomised, double-blind, controlled, phase 2 trial. *Lancet Respir Med* 2017; 5: 22–32.
- 111 Zhu Z, Homer RJ, Wang Z, *et al.* Pulmonary expression of interleukin-13 causes inflammation, mucus hypersecretion, subepithelial fibrosis, physiologic abnormalities, and eotaxin production. *J Clin Invest* 1999; 103: 779–788.
- 112 Belperio JA, Dy M, Burdick MD, *et al.* Interaction of IL-13 and C10 in the pathogenesis of bleomycin-induced pulmonary fibrosis. *Am J Respir Cell Mol Biol* 2002; 27: 419–427.
- 113 Parker JM, Glaspole IN, Lancaster LH, et al. A phase 2 randomized controlled study of tralokinumab in subjects with idiopathic pulmonary fibrosis. Am J Respir Crit Care Med 2018; 197: 94–103.
- 114 Peters-Golden M, Henderson WR Jr. Leukotrienes. N Engl J Med 2007; 357: 1841-1854.
- 115 Izumo T, Kondo M, Nagai A. Effects of a leukotriene B4 receptor antagonist on bleomycin-induced pulmonary fibrosis. *Eur Respir J* 2009; 34: 1444–1451.
- 116 Garneau-Tsodikova S, Thannickal VJ. Protein kinase inhibitors in the treatment of pulmonary fibrosis. *Curr Med Chem* 2008; 15: 2632–2640.
- 117 Zanin-Zhorov A, Weiss JM, Nyuydzefe MS, et al. Selective oral ROCK2 inhibitor down-regulates IL-21 and IL-17 secretion in human T cells via STAT3-dependent mechanism. Proc Natl Acad Sci USA 2014; 111: 16814–16819.
- 118 van der Velden JL, Ye Y, Nolin JD, *et al.* JNK inhibition reduces lung remodeling and pulmonary fibrotic systemic markers. *Clin Transl Med* 2016; 5: 36.
- 119 Saini G, Porte J, Weinreb PH, et al. ανβ6 integrin may be a potential prognostic biomarker in interstitial lung disease. Eur Respir J 2015; 46: 486–494.
- 120 Horan GS, Wood S, Ona V, *et al.* Partial inhibition of integrin ανβ6 prevents pulmonary fibrosis without exacerbating inflammation. *Am J Respir Crit Care Med* 2008; 177: 56–65.
- 121 Diehl N, Schaal H. Make yourself at home: viral hijacking of the PI3K/Akt signaling pathway. *Viruses* 2013; 5: 3192–3212.
- 122 Diab S, Fidanzi C, Léger DY, et al. Berberis libanotica extract targets NF-κB/COX-2, PI3K/Akt and mitochondrial/caspase signalling to induce human erythroleukemia cell apoptosis. Int J Oncol 2015; 47: 220–230.
- 123 Zhang XL, Xing RG, Chen L, *et al.* PI3K/Akt signaling is involved in the pathogenesis of bleomycin-induced pulmonary fibrosis *via* regulation of epithelial-mesenchymal transition. *Mol Med Rep* 2016; 14: 5699–5706.
- 124 Zhang L, Li Y, Liang C, et al. CCN5 overexpression inhibits profibrotic phenotypes via the PI3K/Akt signaling pathway in lung fibroblasts isolated from patients with idiopathic pulmonary fibrosis and in an *in vivo* model of lung fibrosis. Int J Mol Med 2014; 33: 478–486.
- 125 Mercer PF, Woodcock HV, Eley JD, *et al.* Exploration of a potent PI3 kinase/mTOR inhibitor as a novel anti-fibrotic agent in IPF. *Thorax* 2016; 71: 701–711.
- 126 Lukey PT, Harrison SA, Yang S, *et al.* A randomised, placebo-controlled study of omipalisib (PI3K/mTOR) in idiopathic pulmonary fibrosis. *Eur Respir J* 2019; 53: 1801992.
- 127 Dall'Aglio PP, Pesci A, Bertorelli G, et al. Study of immune complexes in bronchoalveolar lavage fluids. Respiration 1988; 54: Suppl. 1, 36–41.
- 128 Dobashi N, Fujita J, Murota M, *et al.* Elevation of anti-cytokeratin 18 antibody and circulating cytokeratin 18: anti-cytokeratin 18 antibody immune complexes in sera of patients with idiopathic pulmonary fibrosis. *Lung* 2000; 178: 171–179.
- 129 Marchal-Sommé J, Uzunhan Y, Marchand-Adam S, *et al.* Cutting edge: nonproliferating mature immune cells form a novel type of organized lymphoid structure in idiopathic pulmonary fibrosis. *J Immunol* 2006; 176: 5735–5739.
- 130 Donahoe M, Valentine VG, Chien N, et al. Autoantibody-targeted treatments for acute exacerbations of idiopathic pulmonary fibrosis. PLoS One 2015; 10: e0127771.
- 131 O'Dwyer DN, Ashley SL, Gurczynski SJ, et al. Lung microbiota contribute to pulmonary inflammation and disease progression in pulmonary fibrosis. Am J Respir Crit Care Med 2019; 199: 1127–1138.
- 132 Shulgina L, Cahn AP, Chilvers ER, *et al.* Treating idiopathic pulmonary fibrosis with the addition of co-trimoxazole: a randomised controlled trial. *Thorax* 2013; 68: 155–162.
- 133 Hammond M, Clark AB, Cahn AP, *et al.* The Efficacy and Mechanism Evaluation of Treating Idiopathic Pulmonary fibrosis with the Addition of Co-trimoxazole (EME-TIPAC): study protocol for a randomised controlled trial. *Trials* 2018; 19: 89.
- 134 Glanville AR, Estenne M. Indications, patient selection and timing of referral for lung transplantation. *Eur Respir J* 2003; 22: 845–852.
- Weill D, Benden C, Corris PA, et al. A consensus document for the selection of lung transplant candidates: 2014
 an update from the Pulmonary Transplantation Council of the International Society for Heart and Lung Transplantation. J Heart Lung Transplant 2015; 34: 1–15.
- 136 Kilic A, Merlo CA, Conte JV, *et al.* Lung transplantation in patients 70 years old or older: have outcomes changed after implementation of the lung allocation score? *J Thorac Cardiovasc Surg* 2012; 144: 1133–1138.
- 137 Lamas DJ, Kawut SM, Bagiella E, et al. Delayed access and survival in idiopathic pulmonary fibrosis: a cohort study. Am J Respir Crit Care Med 2011; 184: 842–847.

- 138 Paik HC, Haam SJ, Lee DY, *et al.* The fate of patients on the waiting list for lung transplantation in Korea. *Transplant Proc* 2012; 44: 865–869.
- 139 Adamali HI, Judge EP, Healy D, et al. International collaboration: a retrospective study examining the survival of Irish citizens following lung transplantation in both the UK and Ireland. BMJ Open 2012; 2: e000605.
- 140 Kourliouros A, Hogg R, Mehew J, *et al.* Patient outcomes from time of listing for lung transplantation in the UK: are there disease-specific differences? *Thorax* 2019; 74: 60–68.
- 141 Annual Data Report of the US Organ Procurement and Transplantation Network (OPTN) and the Scientific Registry of Transplant Recipients (SRTR). Introduction. *Am J Transplant* 2013; 13: Suppl. 1, 8–10.
- 142 US Department of Health and Human Services. Organ Procurement and Transplantation Network Data. http:// optn.transplant.hrsa.gov/Data Date last updated: June 16, 2019.
- 143 The International Society for Heart and Lung Transplantation. International Thoracic Organ Transplant (TTX) Registry data slides: adult lung transplantation slideset. 2018. https://ishltregistries.org/registries/slides.asp Last date accessed: January 25, 2019.
- 144 Schaffer JM, Singh SK, Reitz BA, *et al.* Single- *vs* double-lung transplantation in patients with chronic obstructive pulmonary disease and idiopathic pulmonary fibrosis since the implementation of lung allocation based on medical need. *JAMA* 2015; 313: 936–948.
- 145 Salamo O, Roghaee S, Schweitzer MD, *et al.* White donor, younger donor and double lung transplant are associated with better survival in sarcoidosis patients. *Sci Rep* 2018; 8: 6968.
- 146 Bleisch B, Schuurmans MM, Klaghofer R, *et al.* Health-related quality of life and stress-related post-transplant trajectories of lung transplant recipients: a three-year follow-up of the Swiss Transplant Cohort Study. *Swiss Med Wkly* 2019; 149: doi: 10.4414/smw.2019.20019.
- 147 Singh VK, Patricia George M, Gries CJ. Pulmonary hypertension is associated with increased post-lung transplant mortality risk in patients with chronic obstructive pulmonary disease. *J Heart Lung Transplant* 2015; 34: 424–429.
- 148 George PM, Patterson CM, Reed AK, et al. Lung transplantation for idiopathic pulmonary fibrosis. Lancet Respir Med 2019; 7: 271–282.
- 149 Biscotti M, Gannon WD, Agerstrand C, et al. Awake extracorporeal membrane oxygenation as bridge to lung transplantation: a 9-year experience. Ann Thorac Surg 2017; 104: 412–419.
- 150 Fuehner T, Kuehn C, Hadem J, et al. Extracorporeal membrane oxygenation in awake patients as bridge to lung transplantation. Am J Respir Crit Care Med 2012; 185: 763–768.
- 151 Mortensen A, Cherrier L, Walia R. Effect of pirfenidone on wound healing in lung transplant patients. Multidiscip Respir Med 2018; 13: 16.
- 152 Leuschner G, Stocker F, Veit T, *et al.* Outcome of lung transplantation in idiopathic pulmonary fibrosis with previous anti-fibrotic therapy. *J Heart Lung Transplant* 2017; S1053-2498(17)31886-7.
- 153 Delanote I, Wuyts WA, Yserbyt J, *et al.* Safety and efficacy of bridging to lung transplantation with antifibrotic drugs in idiopathic pulmonary fibrosis: a case series. *BMC Pulm Med* 2016; 16: 156.
- 154 Nici L, ZuWallack R, Wouters E, *et al.* On pulmonary rehabilitation and the flight of the bumblebee: the ATS/ ERS Statement on Pulmonary Rehabilitation. *Eur Respir J* 2006; 28: 461–462.
- Barratt SL, Creamer A, Hayton C, et al. Idiopathic pulmonary fibrosis (IPF): an overview. J Clin Med 2018; 7: E201.
- 156 Dowman L, Hill CJ, Holland AE. Pulmonary rehabilitation for interstitial lung disease. *Cochrane Database Syst Rev* 2014; 10: CD006322.
- 157 Cheng L, Tan B, Yin Y, *et al.* Short- and long-term effects of pulmonary rehabilitation for idiopathic pulmonary fibrosis: a systematic review and meta-analysis. *Clin Rehabil* 2018; 32: 1299–1307.
- 158 da Fontoura FF, Berton DC, Watte G, et al. Pulmonary rehabilitation in patients with advanced idiopathic pulmonary fibrosis referred for lung transplantation. J Cardiopulm Rehabil Prev 2018; 38: 131–134.
- 159 Holland AE, Fiore JF, Goh N, *et al.* Be honest and help me prepare for the future: what people with interstitial lung disease want from education in pulmonary rehabilitation. *Chron Respir Dis* 2015; 12: 93–101.
- 160 Ryerson CJ, Cayou C, Topp F, *et al.* Pulmonary rehabilitation improves long-term outcomes in interstitial lung disease: a prospective cohort study. *Respir Med* 2014; 108: 203–210.
- 161 Bajwah S, Ross JR, Wells AU, *et al.* Palliative care for patients with advanced fibrotic lung disease: a randomised controlled phase II and feasibility trial of a community case conference intervention. *Thorax* 2015; 70: 830–839.
- 162 Sciriha A, Lungaro-Mifsud S, Fsadni P, et al. Pulmonary rehabilitation in patients with interstitial lung disease: the effects of a 12-week programme. Respir Med 2019; 146: 49–56.
- 163 Garibaldi BT, Danoff SK. Symptom-based management of the idiopathic interstitial pneumonia. *Respirology* 2016; 21: 1357–1365.
- 164 van Manen MJ, Geelhoed JJ, Tak NC, et al. Optimizing quality of life in patients with idiopathic pulmonary fibrosis. Ther Adv Respir Dis 2017; 11: 157–169.
- 165 Nishiyama O, Yamazaki R, Sano H, et al. Physical activity in daily life in patients with idiopathic pulmonary fibrosis. Respir Investig 2018; 56: 57–63.
- 166 Spathis A, Booth S, Moffat C, *et al.* The breathing, thinking, functioning clinical model: a proposal to facilitate evidence-based breathlessness management in chronic respiratory disease. *NPJ Prim Care Respir Med* 2017; 27: 27.
- 167 Tonelli R, Cocconcelli E, Lanini B, *et al.* Effectiveness of pulmonary rehabilitation in patients with interstitial lung disease of different etiology: a multicenter prospective study. *BMC Pulm Med* 2017; 17: 130.
- 168 Duck A, Spencer LG, Bailey S, et al. Perceptions, experiences and needs of patients with idiopathic pulmonary fibrosis. J Adv Nurs 2015; 71: 1055-1065.
- 169 Belkin A, Albright K, Swigris JJ. A qualitative study of informal caregivers' perspectives on the effects of idiopathic pulmonary fibrosis. BMJ Open Respir Res 2014; 1: e000007.
- 170 Bell EC, Cox NS, Goh N, *et al.* Oxygen therapy for interstitial lung disease: a systematic review. *Eur Respir Rev* 2017; 26: 160080.
- 171 Visca D, Mori L, Tsipouri V, et al. Effect of ambulatory oxygen on quality of life for patients with fibrotic lung disease (AmbOx): a prospective, open-label, mixed-method, crossover randomised controlled trial. Lancet Respir Med 2018; 6: 759–770.
- 172 Sharp C, Adamali H, Millar AB. Ambulatory and short-burst oxygen for interstitial lung disease. *Cochrane Database Syst Rev* 2016; 7: CD011716.

- 173 Allen S, Raut S, Woollard J, *et al.* Low dose diamorphine reduces breathlessness without causing a fall in oxygen saturation in elderly patients with end-stage idiopathic pulmonary fibrosis. *Palliat Med* 2005; 19: 128–130.
- 174 Simon ST, Higginson IJ, Booth S, et al. Benzodiazepines for the relief of breathlessness in advanced malignant and non-malignant diseases in adults. *Cochrane Database Syst Rev* 2016; 10: CD007354.
- 175 Ryerson CJ, Abbritti M, Ley B, et al. Cough predicts prognosis in idiopathic pulmonary fibrosis. Respirology 2011; 16: 969–975.
- 176 van Manen MJ, Birring SS, Vancheri C, et al. Cough in idiopathic pulmonary fibrosis. Eur Respir Rev 2016; 25: 278–286.
- 177 Kohberg C, Andersen CU, Bendstrup E. Opioids: an unexplored option for treatment of dyspnea in IPF. *Eur Clin Respir J* 2016; 3: 30629.
- 178 Horton MR, Santopietro V, Mathew L, et al. Thalidomide for the treatment of cough in idiopathic pulmonary fibrosis: a randomized trial. Ann Intern Med 2012; 157: 398-406.
- 179 Horton MR, Danoff SK, Lechtzin N. Thalidomide inhibits the intractable cough of idiopathic pulmonary fibrosis. *Thorax* 2008; 63: 749.
- 180 van Manen MJG, Birring SS, Vancheri C, et al. Effect of pirfenidone on cough in patients with idiopathic pulmonary fibrosis. Eur Respir J 2017; 50: 1701157.
- 181 Birring SS, Wijsenbeek MS, Agrawal S, et al. A novel formulation of inhaled sodium cromoglicate (PA101) in idiopathic pulmonary fibrosis and chronic cough: a randomised, double-blind, proof-of-concept, phase 2 trial. Lancet Respir Med 2017; 5: 806–815.
- 182 Kreuter M, Ehlers-Tenenbaum S, Schaaf M, *et al.* Treatment and outcome of lung cancer in idiopathic interstitial pneumonias. *Sarcoidosis Vasc Diffuse Lung Dis* 2015; 31: 266–274.
- 183 Kreuter M, Baroke E, Ehlers-Tenenbaum S, et al. Other comorbidities in idiopathic pulmonary fibrosis. In: Costabel U, Crestani B, Wells AU, eds. Idiopathic Pulmonary Fibrosis (ERS Monograph). Sheffield, European Respiratory Society, 2015; pp. 186–195.
- 184 Collard HR, Ward AJ, Lanes S, et al. Burden of illness in idiopathic pulmonary fibrosis. J Med Econ 2012; 15: 829–835.
- 185 King CS, Nathan SD. Idiopathic pulmonary fibrosis: effects and optimal management of comorbidities. Lancet Respir Med 2017; 5: 72–84.
- 186 Kreuter M, Ehlers-Tenenbaum S, Palmowski K, *et al.* Impact of comorbidities on mortality in patients with idiopathic pulmonary fibrosis. *PLoS One* 2016; 11: e0151425.
- 187 Nathan SD, Basavaraj A, Reichner C, et al. Prevalence and impact of coronary artery disease in idiopathic pulmonary fibrosis. Respir Med 2010; 104: 1035–1041.
- 188 Hyldgaard C, Hilberg O, Bendstrup E. How does comorbidity influence survival in idiopathic pulmonary fibrosis? *Respir Med* 2014; 108: 647–653.
- 189 Nadrous HF, Pellikka PA, Krowka MJ, et al. The impact of pulmonary hypertension on survival in patients with idiopathic pulmonary fibrosis. Chest 2005; 128: 6 Suppl., 616S–617S.
- 190 Tomassetti S, Gurioli C, Ryu JH, *et al.* The impact of lung cancer on survival of idiopathic pulmonary fibrosis. *Chest* 2015; 147: 157–164.
- 191 Kreuter M, Swigris J, Pittrow D, *et al.* The clinical course of idiopathic pulmonary fibrosis and its association to quality of life over time: longitudinal data from the INSIGHTS-IPF registry. *Respir Res* 2019; 20: 59.
- 192 Torrisi SE, Ley B, Kreuter M, *et al.* The added value of comorbidities in predicting survival in idiopathic pulmonary fibrosis: a multicenter observational study. *Eur Respir J* 2019; 53: 1801587.
- 193 Tomassetti S, Ruy JH, Gurioli C, *et al.* The effect of anticoagulant therapy for idiopathic pulmonary fibrosis in real life practice. *Sarcoidosis Vasc Diffuse Lung Dis* 2013; 30: 121–127.
- 194 Kreuter M, Wijsenbeek MS, Vasakova M, et al. Unfavourable effects of medically indicated oral anticoagulants on survival in idiopathic pulmonary fibrosis. Eur Respir J 2016; 47: 1776–1784.
- 195 Crooks MG, Hart SP. Coagulation and anticoagulation in idiopathic pulmonary fibrosis. *Eur Respir Rev* 2015; 24: 392–399.
- 196 Bogatkevich GS, Ludwicka-Bradley A, Silver RM. Dabigatran, a direct thrombin inhibitor, demonstrates antifibrotic effects on lung fibroblasts. Arthritis Rheum 2009; 60: 3455–3464.
- 197 Bogatkevich GS, Ludwicka-Bradley A, Nietert PJ, et al. Antiinflammatory and antifibrotic effects of the oral direct thrombin inhibitor dabigatran etexilate in a murine model of interstitial lung disease. Arthritis Rheum 2011; 63: 1416–1425.
- 198 Scotton CJ, Krupiczojc MA, Königshoff M, et al. Increased local expression of coagulation factor X contributes to the fibrotic response in human and murine lung injury. J Clin Invest 2009; 119: 2550–2563.
- 199 Kreuter M, Wuyts W, Renzoni E, et al. Antacid therapy and disease outcomes in idiopathic pulmonary fibrosis: a pooled analysis. Lancet Respir Med 2016; 4: 381–389.
- 200 Kreuter M, Spagnolo P, Wuyts W, *et al.* Antacid therapy and disease progression in patients with idiopathic pulmonary fibrosis who received pirfenidone. *Respiration* 2017; 93: 415–423.
- 201 Lee JS, Collard HR, Anstrom KJ, et al. Anti-acid treatment and disease progression in idiopathic pulmonary fibrosis: an analysis of data from three randomised controlled trials. *Lancet Respir Med* 2013; 1: 369–376.
- 202 Lee CM, Lee DH, Ahn BK, et al. Protective effect of proton pump inhibitor for survival in patients with gastroesophageal reflux disease and idiopathic pulmonary fibrosis. J Neurogastroenterol Motil 2016; 22: 444–451.
- 203 Raghu G, Pellegrini CA, Yow E, et al. Laparoscopic anti-reflux surgery for the treatment of idiopathic pulmonary fibrosis (WRAP-IPF): a multicentre, randomised, controlled phase 2 trial. Lancet Respir Med 2018; 6: 707–714.
- 204 Bansilal S, Castellano JM, Fuster V. Global burden of CVD: focus on secondary prevention of cardiovascular disease. Int J Cardiol 2015; 201: Suppl. 1, S1–S7.
- 205 Briel M, Vale N, Schwartz GG, *et al.* Updated evidence on early statin therapy for acute coronary syndromes: meta-analysis of 18 randomized trials involving over 14,000 patients. *Int J Cardiol* 2012; 158: 93–100.
- 206 Alexeeff SE, Litonjua AA, Sparrow D, et al. Štatin use reduces decline in lung function: VA Normative Aging Study. Am J Respir Crit Care Med 2007; 176: 742–747.
- 207 Vedel-Krogh S, Nielsen SF, Nordestgaard BG. Statin use is associated with reduced mortality in patients with interstitial lung disease. *PLoS One* 2015; 10: e0140571.

- 208 Kreuter M, Bonella F, Maher TM, *et al.* Effect of statins on disease-related outcomes in patients with idiopathic pulmonary fibrosis. *Thorax* 2017; 72: 148–153.
- 209 Kreuter M, Lederer DJ, Molina-Molina M, et al. Association of angiotensin modulators with the course of idiopathic pulmonary fibrosis. Chest 2019. In press doi: 10.1016/j.chest.2019.04.015.
- 210 Novelli L, Ruggiero R, De Giacomi F, *et al.* Corticosteroid and cyclophosphamide in acute exacerbation of idiopathic pulmonary fibrosis: a single center experience and literature review. *Sarcoidosis Vasc Diffuse Lung Dis* 2016; 33: 385–391.
- 211 Raghu G, Richeldi L. Current approaches to the management of idiopathic pulmonary fibrosis. *Respir Med* 2017; 129: 24–30.
- 212 Collard HR, Ryerson CJ, Corte TJ, et al. Acute exacerbation of idiopathic pulmonary fibrosis. An International Working Group Report. Am J Respir Crit Care Med 2016; 194: 265–275.
- 213 Kreuter M, Polke M, Walsh S, *et al.* A global perspective on acute exacerbation of idiopathic pulmonary fibrosis (AE-IPF): results from an international survey. *Eur Respir J* 2018; 52: Suppl. 62, OA542.
- 214 Trudzinski FC, Kaestner F, Schäfers HJ, *et al.* Outcome of patients with interstitial lung disease treated with extracorporeal membrane oxygenation for acute respiratory failure. *Am J Respir Crit Care Med* 2016; 193: 527–533.
- 215 Polke M, Behr J, Kabitz HJ, *et al.* Diagnosis and therapy of acute exacerbations of idiopathic pulmonary fibrosis (AE-IPF) in Germany. *Eur Respir J* 2018; 52: Suppl. 62, PA2903.