

Survival and course of lung function in the presence or absence of antifibrotic treatment in patients with idiopathic pulmonary fibrosis: long-term results of the INSIGHTS-IPF registry

Running title: IPF long-term outcomes on antifibrotic treatment
9.23 Interstitial Lung Disease

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55 **Abstract**

56 *Rationale.* There is a paucity of observational data on antifibrotic therapy for idiopathic
57 pulmonary fibrosis (IPF).

58 *Objective.* We aimed to assess the course of disease of IPF patients with and without
59 antifibrotic therapy under real-life conditions.

60 *Methods.* We analysed data from a non-interventional, prospective cohort study of
61 consecutively enrolled IPF patients from 20 ILD expert centres in Germany. Data quality was
62 ensured by automated plausibility checks, on-site monitoring, and source data verification.
63 Propensity scores were applied to account for known differences in baseline characteristics
64 between patients with and without antifibrotic therapy.

65 *Results.* Among the 588 patients suitable for analysis, the mean age was 69.8 ± 9.1 years, and
66 81.0% were males. The mean duration of disease since diagnosis was 1.8 ± 3.4 years. The mean
67 % predicted value at baseline for forced vital capacity (FVC) and diffusion capacity (DLCO) were
68 68.6 ± 18.8 and 37.8 ± 18.5 , respectively.

69 During a mean follow-up of 1.2 ± 0.7 years, 194 (33.0%) patients died. The one-year and two-
70 year survival rates were 87% vs. 46% and 62% vs. 21%, respectively, for patients with vs.
71 without antifibrotic therapy. The risk of death was 37% lower in patients with antifibrotic
72 therapy (HR=0.63, 95%CI: 0.45; 0.87; p=0.005). The results were robust (and remained
73 statistically significant) on multivariable analysis. Overall decline of FVC and DLco was slow and
74 did not differ significantly between patients with or without antifibrotic therapy.

75 *Conclusions.* Survival was significantly higher in IPF patients with antifibrotic therapy, but the
76 course of lung function parameters was similar in patients with and without antifibrotic
77 therapy. This suggests that in clinical practice premature mortality of IPF patients eventually
78 occurs despite stable measurements for FVC and DLco.

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81

82 **Key words:** Lung fibrosis, outcomes, survival, adjustment, observational, pirfenidone,
83 nintedanib, antifibrotic therapy

84

85 **Background**

86
87 Idiopathic pulmonary fibrosis (IPF) is a severe respiratory disease characterised by progressive
88 scarring of the lung, leading to respiratory failure and death within 3-5 years from diagnosis.¹
89 Effective treatments are still limited. The antifibrotic treatments pirfenidone and nintedanib
90 have been shown to slow disease progression as measured by annual rate of decline in forced
91 vital capacity (FVC),² but their effect on lung function and survival under clinical practice
92 conditions warrants further exploration.

93 As randomised controlled studies on antifibrotic treatments have limitations in terms of their
94 generalizability due to patient selection/exclusion and duration of follow-up, observational
95 data in unselected IPF patients are needed to provide a more comprehensive picture. A
96 number of registries have been initiated in various countries to provide such real-life data,³⁻⁸
97 but their follow-up is limited to 1-2 years only.

98 The database of the INSIGHTS-IPF registry, one of the largest IPF registries worldwide, offers
99 the opportunity to analyse the course of disease and long-term effectiveness of antifibrotic
100 therapy in IPF. The aims of the present analysis were (1) to describe and compare cohorts of
101 patients with and without antifibrotic therapy, (2) to assess the correlation between
102 antifibrotic drug use and lung function, and (3) to test the correlation between antifibrotic
103 drug use and survival.

104

105 **Methods**

106 *Design and parameters.* The INSIGHTS-IPF (“Investigating significant health trends in idiopathic
107 pulmonary fibrosis”) registry is a nationwide, investigator-initiated observational study. The
108 registry has been continuously enrolling consecutive incident and prevalent patients in routine
109 clinical care across 20 pulmonary specialist centres in Germany since November 2012. Patients
110 ≥18 years of age with a study-site diagnosis of IPF according to the 2011 ATS/ERS/JRS/ALAT IPF
111 guideline⁹ after provision of written informed consent can be enrolled, with no explicit
112 exclusion criteria. The registry’s structure, methodology, and regulatory aspects, as well as a
113 detailed description of the baseline characteristics of the patient cohort, have been reported
114 previously.¹⁰⁻¹² The study has been approved by the ethics committee at the Technical
115 University of Dresden and various local ethical committees. All patients provided informed

116 consent before their data were documented in the registry. The ClinTrials.gov identifier is
117 NCT01695408.

118 Data were collected at enrolment (baseline) and at subsequent 6- to 12-month intervals. At
119 each follow-up visit, all clinical events, including hospitalisation and acute exacerbations (as
120 judged by the treating physician), as well as deaths that occurred during the study period,
121 were recorded by each site. At each visit, if available, a range of routine pulmonary function
122 tests were documented, including forced vital capacity (FVC), diffusing capacity of the lung for
123 carbon monoxide (DLCO), the forced expiratory volume in 1 s (FEV1), and six-minute walk
124 distance (6MWD). The gender, age, and physiology (GAP) index was calculated based on
125 available data.¹³

126 The treating physician was requested to judge the overall clinical course of IPF at baseline and
127 each follow-up visit by the categories: stable disease, slow progression, rapid progression, no
128 judgement possible. Physiologic changes between baseline and 2-year follow-up were
129 categorized as stable if FVC did not change or was improved by $\geq 5\%$; as a moderate decrease if
130 decreased by $>5-10\%$; or as a significant decrease if decreased by $>10\%$.

131 *Quality measures.* All data were collected using a standardised internet-based case report form
132 (eCRF) with secure electronic data transfer to the central database. Quality measures included
133 automated plausibility checks at data entry, statistical checks on data quality (focusing on
134 missing values and outliers) as well as on-site monitoring and source data verification
135 performed in the majority of centres (over 70%).

136 *Data analysis.* Data were summarised by descriptive statistics including means and standard
137 deviations and absolute and relative frequencies at baseline and each subsequent follow-up
138 assessment. Data analysis comprised the period between the first documentation in the
139 registry in December 2012 until the data cut-off point in December 2018. The analyses follow
140 the intention to treat principle, which means that each patient with at least one dose of
141 antifibrotic therapy is assigned to the treatment group.

142 The entire observation period was considered for each patient in the registry in order to
143 compare outcomes, in terms of mortality and pulmonary function test results, between
144 patients who were treated with antifibrotic therapy and those who were not. Patients in the
145 registry who had never been treated with an antifibrotic therapy were assigned to the control
146 group. The first observation in that group was the registry enrolment visit. Patients who
147 started an antifibrotic therapy before enrolment into the registry (start of more than 10 days

148 before, e.g. as participant in a clinical study) were excluded because of the non-availability of
149 clinical data at treatment start. The data were divided into individual treatment episodes for
150 patients who started pirfenidone and/or nintedanib during the observation period. For these
151 patients, the first observation was the initial treatment visit. If a patient was treated with
152 pirfenidone and nintedanib (in sequence) during follow-up, then two treatment episodes were
153 assigned (one for each drug) for the pulmonary function and 6MWD tests at the corresponding
154 time point. In contrast, the risk of mortality was analysed for the last available antifibrotic
155 treatment episode in patients who were treated with pirfenidone followed by nintedanib, or
156 vice versa, during follow-up. All patients with a follow-up period of at least 3 months were
157 included in the analyses. In addition, a follow-up interval of 2 years was considered. The
158 primary analysis for lung function tests and 6MWD is based on the observed values in the
159 registry. Since the number of missing values in lung function tests (FVC: baseline 4.5%, follow-
160 up 20.7%; DLCO: baseline 16.2%, follow-up 31.9%) and 6MWD (baseline 14.1%, follow-up
161 57.3%) were substantial, we applied the technique of multiple imputation for those variables
162 to estimate the missing values as sensitivity analyses. Patients with a missing lung function test
163 tended to be on a less severe disease course compared to patients with available lung function
164 test. Preliminary analyses showed that mortality, age, and comorbidities were associated with
165 the absence of the considered variables. Therefore, the first sensitivity analysis used an
166 imputation model including the predictor variables age, sex, number of comorbidities, IPF
167 duration, mortality, antifibrotic therapy, and the lung function and 6MWD results from the
168 prior visit. The number of imputations was set to 10. As a second sensitivity analysis, the last
169 observation carried forward method for lung function and 6MWD was used as well. The third
170 sensitivity analysis used the imputation of missing values by the worst possible value (FVC,
171 DLCO, and 6MWD of 0) for patients who died.

172 *Propensity score.* INSIGHTS-IPF is an observational study and thus allocation to treatment was
173 not randomly assigned. Consequently, various patient characteristics at baseline may be
174 imbalanced, possibly leading to biased results and conclusions. The standard approach to deal
175 with this problem is to model the probability of treatment assignment by the physician
176 (propensity score) based on the clinical characteristics at treatment start in order to balance
177 the characteristics of the two considered groups of patients.^{14,15,16} The propensity score was
178 estimated by a logistic regression model that included the covariates sex, age, smoking status,
179 number of comorbid diseases, IPF disease duration, FVC % predicted, 6MWD, concomitant
180 therapy with steroids, and the global assessment of the disease course by the physician at

181 baseline. A weight value (inverse probability of treatment weighting, IPTW) was calculated for
182 each patient based on the propensity score.¹⁷ All statistical comparisons between patients
183 with and without antifibrotic therapy were weighted to balance the two groups regarding the
184 clinical characteristics at treatment start.

185 In the primary analysis, the course of the pulmonary function (FVC% and DLCO% predicted)
186 and 6MWD tests were analysed by weighted linear mixed models to account for the possibility
187 of two treatment episodes for a single patient (additional cluster variable) and the longitudinal
188 study design based on the observed values. An interaction term treatment x time was included
189 into the weighted linear mixed models to test for differences in change in the three considered
190 parameters by treatment. Secondary analyses of lung function and 6MWD included the
191 imputed data, which employed two imputation methods: last-observation-carried-forward and
192 worst-case imputation. The risk of mortality was analysed by a multivariable Cox proportional
193 hazard model weighted by the propensity score. The proportional-hazards assumption was
194 tested on the basis of Schoenfeld residuals after fitting the Cox regression model.

195 Data were analysed with STATA 12.1 (StataCorp LP. Stata Statistical Software: Release 12.
196 College Station, TX, USA).

197

198 **Results**

199 A total of 588 patients were deemed suitable for the present analysis. The mean age of the
200 study population was 69.8 years, with a large male preponderance (81.0%). The mean duration
201 of symptoms before the baseline visit was 3.5 ± 4.2 years and the mean time between
202 diagnosis and study enrolment was 1.8 ± 3.4 years. Fifty eight percent of the patients had
203 disease duration of less than 12 months and 47% of less than 6 months. The mean Borg
204 Dyspnea score was 2.2 ± 2.4 , and the GAP index stages were as follows: Stage I in 20.4% of the
205 patients, Stage II in 49.9% of the patients, and Stage III in 29.7% of the patients. In terms of
206 lung function parameters at baseline, the mean predicted FVC was $68.6\% \pm 18.8$ and the mean
207 predicted DLCO was $37.8\% \pm 18.5$. Health-related quality of life as measured on the 100-point
208 visual analogue scale was 59.6 ± 23.6 . As current therapy at baseline, prednisone was reported
209 in 23.6% and N-acetylcysteine in 25.5% of patients.

210 The mean follow-up time was 1.2 ± 0.7 years (maximum of two years) for the total sample, 1.2
211 ± 0.5 years for patients under antifibrotic therapy, and 1.0 ± 0.7 years for patients who had

212 never been treated with antifibrotic therapy. A total of 334 treatment episodes under
213 antifibrotic therapy (168 pirfenidone, 166 nintedanib) were reported for 298 patients in our
214 registry, resulting in 36 patients (12 %) with two episodes. Among these, pirfenidone was the
215 first antifibrotic drug in 29 patients. Seven patients switched from pirfenidone to nintedanib
216 within 3 months after discontinuation of pirfenidone; the other 22 patients started nintedanib
217 on average 13 months after discontinuation of pirfenidone (Table 1).

218 Generalized linear mixed models were used to analyse the pulmonary function and 6MWD
219 tests. These models included all antifibrotic therapy treatment episodes, and were based on
220 the observed values. During the 2 years of follow-up, mean predicted FVC% remained almost
221 stable (Figure 1A, β for change in follow-up=-0.42, 95%CI: -1.44 to 0.60, $p=0.416$), with no
222 significant differences between the two groups (β for time x therapy= -0.65, 95%CI: -1.82 to
223 0.52, $p=0.274$). Predicted DLCO showed a similar course in both groups (Figure 1B), with no
224 significant decline in DLCO (β for change in follow-up =-1.05, 95%CI: -2.40 to 0.30, $p=0.127$) in
225 follow-up and no significant differences between the two groups (β for time x therapy= -0.40,
226 95%CI: -2.56 to 1.77, $p=0.721$). Results for the 6MWD test were available in 89% of patients at
227 baseline; however, this measurement was compromised by a high rate of missing data during
228 follow-up. There was no statistically significant difference in the course of 6MWD results over
229 time (β for change in follow-up = -14.8, 95%CI: -25.6, 4.1, $p=0.076$), considering the observed
230 values. The primary analysis was repeated in patients with disease duration of less than or
231 equal to 12 months at enrolment (prevalent patients, Figure 1: second column). A slightly
232 better course of FVC %, DLCO %, and 6MWD was observed in patients with antifibrotic
233 therapy; however, the difference was not statistically significant. The sensitivity analyses using
234 imputed data and data obtained by the LOCF approach resulted in comparable results to those
235 of the primary analysis. If an FVC % of 0 was imputed in patients who died during follow-up,
236 patients never on antifibrotic therapy tended to have a slightly, but not significantly, stronger
237 FVC decline. The decline in DLCO was worse in patients with antifibrotic treatment, although
238 when imputation of the worst individual value was implemented, there were no significant
239 differences between groups.

240 The risk of mortality was analysed for the last available treatment episode in patients who
241 were treated with pirfenidone ($n=139$) and nintedanib ($n=159$) in follow-up. A total of 194
242 (33.0%) patients died during follow-up. A total of 79 (41%) patients died of IPF related reasons
243 (20% by respiratory failure, and 8% by respiratory infection/pneumonia), followed by

244 complicating comorbidity (8%) and other causes not related to IPF (9%). The reason of death
245 was unknown for 71 (37%) patients.

246 Overall mortality was substantially lower in patients treated with antifibrotic therapy. The risk
247 of death for any reason was 37% lower in patients with antifibrotic therapy compared with
248 those without such therapy (HR = 0.63, 95%CI: 0.45; 0.87; p=0.005, Figure 1). This result was
249 robust (and remained statistically significant) on multivariable analysis, as reported in [Table 2](#).
250 Analysis for both antifibrotic drugs approved for treating IPF, nintedanib and pirfenidone,
251 revealed no statistically significant difference in overall mortality between the two drugs (HR
252 for pirfenidone versus nintedanib = 1.39, 95%CI: 0.87 – 2.22, p=0.164).

253 In patients treated with antifibrotics the risk of IPF-related death was not (statistically
254 significantly) lower compared to patients without such therapy (HR = 0.75, 95%CI: 0.45; 1.25;
255 p=0.266), while the risk of death for unknown reason was 56% lower in patients with
256 antifibrotics (HR = 0.44, 95%CI: 0.26; 0.75; p=0.003). Due to the lower numbers of events in
257 this sub-group analysis this result should be interpreted with caution.

258 We tested the hypothesis whether survival differs between patients with stable FVC (i.e. 10%
259 decline or less during follow-up) compared to patients with worsening of FVC of more than
260 10% during follow-up, regardless of therapy. The risk of mortality was slightly higher in such
261 patients with disease progression compared to stable IPF patients (HR = 1.34, 95%CI: 0.89 –
262 2.02, p=0.163). This result was confirmed while adjusting for the effect for antifibrotic
263 treatment.

264 The risk of mortality was additionally analyzed in patients with disease duration of less than 12
265 months prior to study enrollment. The risk of death in the subsample of incident patients was
266 64% lower in patients treated with antifibrotic therapy compared to controls (HR = 0.44,
267 95%CI: 0.25; 0.78; p=0.003). The result was confirmed in multivariable analysis.

268

269 **Discussion**

270

271 The present analysis of the large and contemporary INSIGHTS-IPF registry indicates that
272 patients on antifibrotic therapy appear to survive significantly longer than IPF patients without
273 antifibrotic therapy. The lower overall mortality risk in the patients treated with antifibrotic
274 medication was mainly driven by patients with unknown reason of death. The statistically non-

275 significant relationship between antifibrotic therapy and IPF-related deaths might be due to
276 the low number of recorded IPF-related deaths (79.4% of deaths).

277 Compared with the recently published observational data from the EurIPF registry, patients in
278 INSIGHTS-IPF were nearly identical in terms of TLC % predicted (70.0% vs 71.2%), FVC %
279 predicted (68.4% versus 68.3%), and FEV₁ % predicted (110% versus 111%), while DLCO %
280 predicted was lower in our study (42.1% versus 37.8%).⁹ A subset of IPF patients with long-
281 term follow up within the EurIPF registry were analysed by Kaplan-Meier analysis (without
282 propensity score matching) in correlation with the date of first IPF diagnosis. The analysis of
283 this subset found that median survival on antifibrotic drugs was 123.1 months (censored cases
284 inclusive, range 84–162 months), compared with a median survival of 68.3 months in patients
285 treated with any other medication including immunosuppressive therapies (censored cases
286 inclusive, range 54–83 months). Functional follow-up data from the EurIPF registry were not
287 reported. Another difference between our data and those of the EurIPF registry, besides the
288 larger number of patients and the statistics applied in our cohort, is the fact that pirfenidone
289 was used in the vast majority (83%) of the EurIPF registry cohort while in our study population
290 nintedanib and pirfenidone were almost equally distributed, slightly favouring nintedanib
291 (53.3%).

292 Interestingly, we observed a similar, stable course of lung function parameters (FVC and DLCO)
293 over time in both groups, with and without antifibrotic therapy, while overall mortality was
294 considerably higher in the group not treated with antifibrotics. At first glance, our data could
295 provide basis for a hypothesis that stable physiological measurements like FVC and DLco alone
296 may not provide a safeguard against premature mortality in IPF. Lung function measurements
297 every 6 to 12 months is common practice and thus employed in our registry. However, such
298 measurements may be less sensitive to detect differences in the course of IPF compared to
299 highly standardized serial measurements at shorter intervals which are commonly applied in
300 clinical trials. Moreover, missing lung function data may have contributed to blunt differences
301 of the slope of FVC and DLco decline between patients with and without antifibrotic therapy.
302 In this context, it is noteworthy that hospital-based FVC measurements, compared with
303 unsupervised daily home measurements, have been suggested to be less sensitive in detecting
304 progression of fibrosis and in predicting subsequent prognosis.¹⁰ However, a recent clinical
305 treatment trial using daily home spirometry for the primary endpoint also revealed potential
306 technical and practical obstacles associated with this methodology.²⁰

307 The phenomenon of emphysema blunting the decline of FVC in both groups may have
308 contributed to this observation, but the prevalence of emphysema as reported by the
309 investigators was low in both groups. The higher preponderance of steroid-treated patients in
310 the group not treated with antifibrotics may also be considered to potentially contribute to a
311 higher mortality in this group. However, the mean prednisone dosage in our study - given to a
312 quarter of patients in our - study was 14 mg/d. In the INPULSIS study the maximum dose was
313 15mg/day and in the ASCEND study, prednisone was only allowed if given for another
314 indication.^{11, 12} Nonetheless, we cannot exclude that unbalanced steroid-medication has
315 contributed to the observed difference. Finally, antioxidant drugs (NAC) were less commonly
316 used in the antifibrotic therapy arm. The impact of these drugs on prognosis is still under
317 discussion and thus a bias cannot be fully excluded.^{13, 14} In consideration of all the limitations
318 our data should be taken as a signal of caution that stability of FVC and DLco may not always
319 protect from premature mortality in the absence of antifibrotic therapy in a fatal disease like
320 IPF. The common practice, still widely used, of withholding antifibrotic therapy from
321 physiologically stable IPF patients may therefore set these patients on a path of increased risk
322 of dying.¹⁵

323 Another important aspect of our study is the fact that all patients were enrolled solely based
324 on investigator judgement. The patients enrolled were, therefore, a cohort which included all
325 the imponderabilities of diagnosis in this complex disease which occur in daily practice. The
326 observed difference in survival in favour of antifibrotic therapy is, therefore, an important
327 argument for the clinical application of these drugs, even though a causative argument cannot
328 be made from our study. This observation is, therefore in accordance with recent clinical trials
329 showing that antifibrotic therapies are effective in progressive fibrotic interstitial lung diseases
330 other than IPF.^{20,26,27}

331 Our data do not identify a cause for the difference in overall mortality between patients with
332 and without antifibrotic therapy. However, one can speculate that acute exacerbation may
333 have contributed substantially to this difference.

334 A number of limitations need to be taken into consideration when interpreting the findings.
335 The major limitation of this study is that patients with existing (prevalent) and newly
336 diagnosed (incident) IPF were documented which may potentially cause lead time bias
337 regarding mortality. This is especially important since time to diagnosis was approximately one
338 year longer in the never-treated population, which could indicate a “healthy survivor effect”.¹⁶

339 Further, there was no randomization between the group of patients who had never been
340 treated with an antifibrotic therapy and patients who were treated with an antifibrotic drug.
341 To account for bias by indication, we calculated a propensity score to estimate the probability
342 of being treated with an antifibrotic drug in our registry based on clinical characteristics.
343 However, there may exist unmeasured variables that cannot be included in the propensity
344 score model that may have impacted the association between antifibrotic therapy and
345 mortality. Furthermore, accompanying therapies such as anti-oxidant or anti-acid therapy may
346 have impacted the results of our analysis. We also had to account for a high proportion of
347 missing values in the pulmonary function tests and in the 6MWD test in the follow-up data,
348 which could have affected our results. The fact that only ILD specialty centers participated in
349 the INSIGHTS-IPF Registry may limit the generalizability of our study.

350 In conclusion, we were able to demonstrate a significant lower all-cause mortality in IPF
351 patients treated with antifibrotic drugs when compared to a matched cohort of IPF patients
352 not treated with antifibrotic drugs. Moreover, our analysis provides the basis for a hypothesis
353 that stability of lung function parameters over time, especially FVC and DLco, in untreated IPF
354 patients may be misleading as our data indicate that stability of these parameters probably do
355 not protect from premature death.

356 **Declarations**

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361

362 **Ethics approval and consent to participate**

363 The study materials were approved by the Ethics Committee of the Medical
364 Faculty, Technical University of Dresden (EK 255082012), and by further local
365 ethic committees as per local requirements.

366

367 **Consent for publication**

368 Not applicable

369

370 **Availability of data and material**

371 The datasets used and/or analysed during the current study are available from the
372 corresponding author on reasonable request.

373

374 **Competing interests**

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Table 1. Characteristics of patients, in the total analysed cohort and by presence or absence of antifibrotic treatment

	Total n=588	Never been treated with antifibrotic therapy n=290	Treated with antifibrotic therapy n=298
Male sex; n(%)	476 (81.0)	230 (79.3)	246 (82.6)
Age; mean (SD), p50	69.8 (9.1); 72	70.3 (9.4); 73	69.2 (8.8); 71
Body Mass Index in kg/m ² ; mean (SD), p50	27.6 (4.1); 27.2	26.9 (4.1); 26.3	28.2 (4.0); 27.7
underweight (BMI<18.5); n (%)	1 (0.2)	1 (0.3)	0 (0.0)
normal weight (18.5 ≤ BMI ≤ 25); n (%)	153 (26.0)	93 (32.1)	60 (20.1)
overweight(25 < BMI ≤ 30); n (%)	291 (49.5)	133 (45.9)	158 (53.0)
obesity (BMI>30); n (%)	143 (24.3)	63 (21.7)	80 (26.9)
Never smoked; n(%)	205 (34.9)	96 (33.1)	109 (36.6)
Ex-smoker; n(%)	372 (63.3)	189 (65.2)	183 (61.4)
Number of comorbidities; mean ± SD	1.7 (1.5); 2	1.8 (1.5); 2	1.7 (1.4); 2
Symptom duration; mean (SD), p50	3.5 (4.2); 2.2	3.5 (4.7); 2.2	3.4 (3.8); 2.1
Age at symptom onset; mean (SD), p50	66.1 (10.5); 68.0	66.4 (11.3); 69.0	65.9 (9.8); 67.7
Age at diagnosis; mean (SD), p50	68.0 (10.0); 70.0	68.1 (10.7); 70.6	68.0 (9.2); 69.9
6-minute walk distance; mean (SD), p50	278.5 (193.9); 330	257.6 (188.7); 300	297.8 (197.1); 360
Borg index; mean (SD), p50	2.2 (2.4); 1	2.2 (2.5); 1	2.2 (2.2); 1
<i>Current therapy</i>			
Prednisone, n (%)	139 (23.6)	86 (29.7)	53 (17.8)
Other steroids, n (%)	11 (1.9)	2 (0.7)	9 (3.0)
Azathioprine, n (%)	14 (2.4)	10 (3.5)	4 (1.3)
Cyclophosphamide, n (%)	1 (0.2)	0 (0.0)	1 (0.3)
Mycophenolate mofetil, n (%)	1 (0.2)	1 (0.3)	0 (0.0)
N-Acetylcysteine, n (%)	150 (25.5)	101 (34.8)	49 (16.4)
Antifibrotic therapy, n (%)	298 (50.7)	0 (0.0)	298 (100.0)
Patients on oxygen therapy	157 (26.7)	86 (29.7)	71 (23.3)
Environmental exposure	199 (33.8)	86 (29.7)	113 (37.9)
Gastro-oesophageal reflux	162 (27.6)	81 (27.9)	81 (27.2)
Family history of ILD	27 (4.6)	20 (6.9)	7 (2.4)
Exposure to drugs	21 (3.6)	9 (3.1)	12 (4.0)
GAP index, n(%)			

Stage I	115 (20.4)	56 (20.4)	59 (20.3)
Stage II	282 (49.9)	128 (46.6)	154 (53.1)
Stage III	168 (29.7)	91 (33.1)	77 (26.6)

Lung function test

Total Lung Capacity, % predicted; mean (SD), p50	71.0 (20.5); 70.5	71.5 (25.7); 69.7	70.5 (14.2); 71.1
Inspiratory Vital Capacity, % predicted; mean (SD), p50	73.2 (20.4); 74.1	70.8 (22.2); 71.8	75.4 (18.4); 76.4
FVC, % predicted; mean (SD), p50	68.6 (18.8); 70.2	66.8 (19.8); 67.9	70.4 (17.5); 71.5
FEV ₁ , % predicted; mean (SD), p50	76.1 (19.7); 76.8	74.1 (20.7); 74.4	77.9 (18.6); 78.4
FEV ₁ : FVC, % predicted; mean (SD), p50	110.9 (11.7); 111.2	111.6 (12.2); 111.9	110.3 (11.2); 110.8
DLCO, % predicted; mean (SD), p50	37.8 (18.5); 35.5	37.6 (20.2); 35.5	38.0 (16.9); 35.2
Health-related quality of life, EQ5D; mean (SD), p50	59.6 (23.6); 60	58.0 (24.1); 60	61.2 (23.1); 65

P50 = median; SD = standard deviation

412 **Table 2. Risk of mortality estimated by a multivariable Cox regression model**

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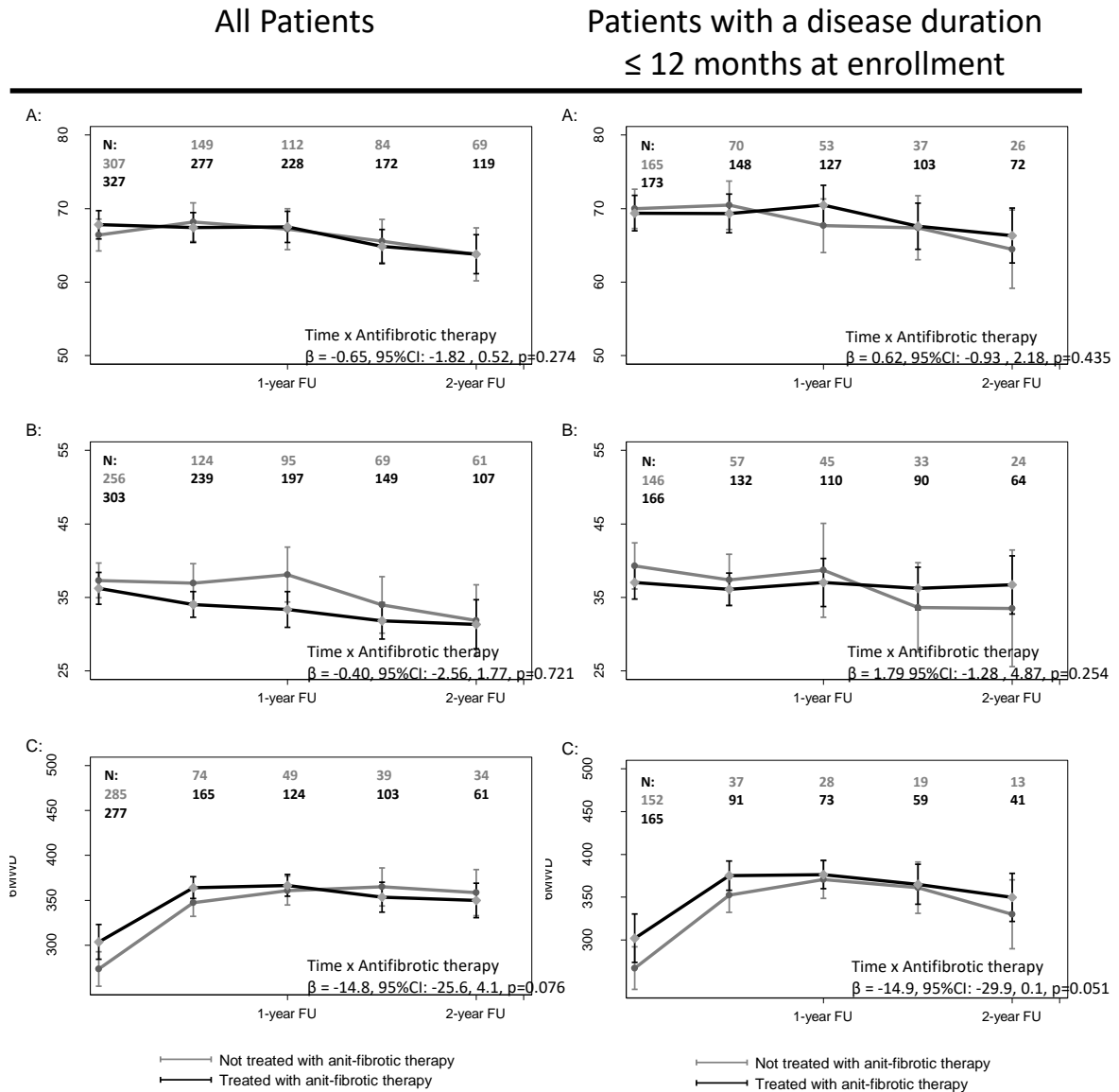
	HR	P value	95% CI
Antifibrotic therapy	0.66	0.016	0.47 ; 0.93
Age	1.09	<0.001	1.07 ; 1.12
Female sex	0.71	0.116	0.47 ; 1.09
IPF disease duration	0.96	0.005	0.94 ; 0.99
Any comorbid disease	1.05	0.821	0.69 ; 1.60
FVC % predicted	0.96	<0.001	0.95 ; 0.98
Overall physician's judgement of clinical course of IPF:			
stable disease	1.00		
slow progression	1.41	0.102	0.93 ; 2.12
rapid progression	2.69	0.002	1.45 ; 4.97

Hazard Ratio (HR) for 1 year change in age and IPF disease duration, HR for 1% change in FVC% predicted.

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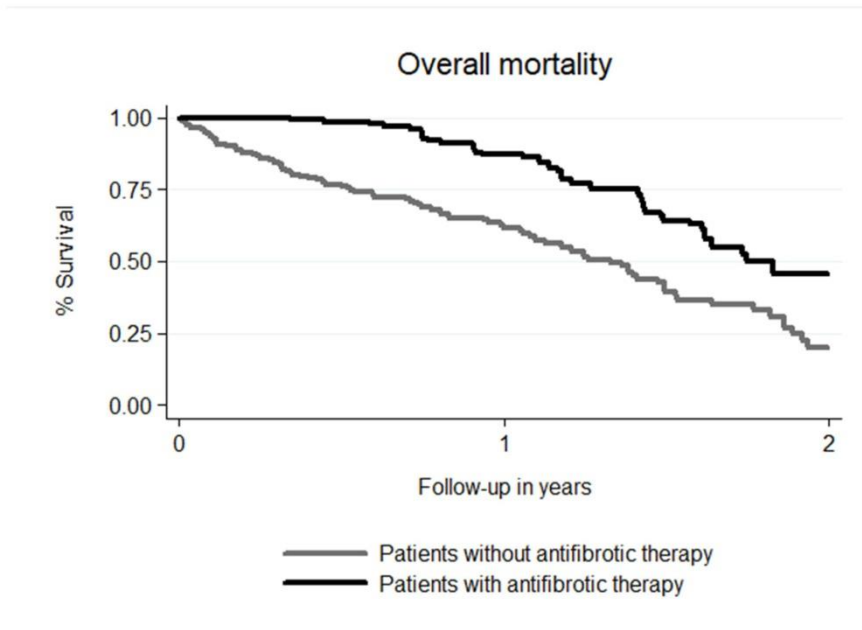
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417 **Figure 1: Change in FVC % predicted (A), DLCO % predicted (B), and 6-minute walking**
 418 **distance (6MWD; C) over the 2-year follow-up (β (interaction term time x therapy) =**
 419 **estimated difference in change during 2-year follow-up in the considered parameter**
 420 **between patients with and without antifibrotic therapy)**
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425 **Figure 2: Risk of mortality within 2 years by antifibrotic treatment (by propensity score**
 426 **weighted Kaplan-Meier survival curves).**
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Number of patients at risk			
With antifibrotic therapy	281	129	57
No antifibrotic therapy	252	139	93

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431 **References**

432

433 1. Raghu G, Rochwerg B, Zhang Y, Garcia CA, Azuma A, Behr J, et al. An Official
434 ATS/ERS/JRS/ALAT Clinical Practice Guideline: Treatment of Idiopathic Pulmonary Fibrosis. An
435 Update of the 2011 Clinical Practice Guideline. *Am J Respir Crit Care Med*. 2015; **192**(2): e3-19.

436 2. Rochwerg B, Neupane B, Zhang Y, Garcia CC, Raghu G, Richeldi L, et al. Treatment of
437 idiopathic pulmonary fibrosis: a network meta-analysis. *BMC Med*. 2016; **14**: 18.

438 3. Wuyts WA, Dahlqvist C, Slabbynck H, Schlessers M, Gusbin N, Compere C, et al. Baseline
439 clinical characteristics, comorbidities and prescribed medication in a real-world population of
440 patients with idiopathic pulmonary fibrosis: the PROOF registry. *BMJ Open Resp Res*. 2018;
441 **5**(1): e000331.

442 4. Jo HE, Glaspole I, Goh N, Hopkins PMA, Moodley Y, Reynolds PN, et al. Implications of
443 the diagnostic criteria of idiopathic pulmonary fibrosis in clinical practice: Analysis from the
444 Australian Idiopathic Pulmonary Fibrosis Registry. *Respirology* 2018; **Oct 17**. doi:
445 **10.1111/resp.13427**.

446 5. Pesonen I, Carlson L, Murgia N, Kaarteenaho R, Skold CM, Myllarniemi M, et al. Delay
447 and inequalities in the treatment of idiopathic pulmonary fibrosis: the case of two Nordic
448 countries. *Multidiscip Respir Med*. 2018; **13**: 14.

449 6. Bouros D, Daniil Z, Papakosta D, Antoniou KM, Markopoulou K, Kolilekas L, et al.
450 Design, Rationale, Methodology, and Aims of a Greek Prospective Idiopathic Pulmonary
451 Fibrosis Registry: Investigating Idiopathic Pulmonary Fibrosis in Greece (INDULGE IPF).
452 *Respiration*. 2018; **96**(1): 41-7.

453 7. O'Brien EC, Durham MT, Gamerman V, Garfinkel S, Anstrom KJ, Palmer SM, et al.
454 Rationale for and design of the Idiopathic Pulmonary Fibrosis-PROspective Outcomes (IPF-PRO)
455 registry. *BMJ Open Resp Res*. 2016; **3**(1): e000108.

456 8. Doubkova M, Svancara J, Svoboda M, Sterclova M, Bartos V, Plackova M, et al. EMPIRE
457 Registry, Czech Part: Impact of demographics, pulmonary function and HRCT on survival and
458 clinical course in idiopathic pulmonary fibrosis. *The clinical respiratory journal*. 2018; **12**(4):
459 1526-35.

460 9. Guenther A, Krauss E, Tello S, Wagner J, Paul B, Kuhn S, et al. The European IPF registry
461 (eurIPFreg): baseline characteristics and survival of patients with idiopathic pulmonary fibrosis.
462 *Respir Res*. 2018; **19**(1): 141.

463 10. Russell AM, Adamali H, Molyneaux PL, Lukey PT, Marshall RP, Renzoni EA, et al. Daily
464 Home Spirometry: An Effective Tool for Detecting Progression in Idiopathic Pulmonary Fibrosis.
465 *Am J Respir Crit Care Med*. 2016; **194**(8): 989-97.

466 11. Richeldi L, du Bois RM, Raghu G, Azuma A, Brown KK, Costabel U, et al. Efficacy and
467 safety of nintedanib in idiopathic pulmonary fibrosis. *N Engl J Med*. 2014; **370**(22): 2071-82.

468 12. King TE, Jr., Bradford WZ, Castro-Bernardini S, Fagan EA, Glaspole I, Glassberg MK, et
469 al. A phase 3 trial of pirfenidone in patients with idiopathic pulmonary fibrosis. *N Engl J Med*.
470 2014; **370**(22): 2083-92.

- 471 13. Behr J, Bendstrup E, Crestani B, Gunther A, Olschewski H, Skold CM, et al. Safety and
472 tolerability of acetylcysteine and pirfenidone combination therapy in idiopathic pulmonary
473 fibrosis: a randomised, double-blind, placebo-controlled, phase 2 trial. *Lancet Resp Med*. 2016;
474 **4**(6): 445-53.
- 475 14. Oldham JM, Ma SF, Martinez FJ, Anstrom KJ, Raghu G, Schwartz DA, et al. TOLLIP,
476 MUC5B, and the Response to N-Acetylcysteine among Individuals with Idiopathic Pulmonary
477 Fibrosis. *American Journal of Respiratory and Critical Care Medicine*. 2015; **192**(12): 1475-82.
- 478 15. Maher TM, Swigris JJ, Kreuter M, Wijsenbeek M, Cassidy N, Ireland L, et al. Identifying
479 Barriers to Idiopathic Pulmonary Fibrosis Treatment: A Survey of Patient and Physician Views.
480 *Respiration*. 2018; **96**(6): 514-24.
- 481 16. Delgado-Rodriguez M, Llorca J. Bias. *J Epidemiol Community Health*. 2004; **58**(8): 635-
482 41.
483
484