Tonelli et al. BMC Pulmonary Medicine (2017) 17:130 DOI 10.1186/s12890-017-0476-5

BMC Pulmonary Medicine

RESEARCH ARTICLE

Open Access



Effectiveness of pulmonary rehabilitation in patients with interstitial lung disease of different etiology: a multicenter prospective study

Roberto Tonelli^{1*}, Elisabetta Cocconcelli⁴, Barbara Lanini², Isabella Romagnoli², Fabio Florini³, Ivana Castaniere¹, Dario Andrisani¹, Stefania Cerri¹, Fabrizio Luppi¹, Riccardo Fantini¹, Alessandro Marchioni¹, Bianca Beghè¹, Francesco Gigliotti² and Enrico M. Clini¹

Abstract

Background: Recent evidences show that Pulmonary Rehabilitation (PR) is effective in patients with Interstitial Lung Disease (ILD). It is still unclear whether disease severity and/or etiology might impact on the reported benefits. We designed this prospective study 1) to confirm the efficacy of rehabilitation in a population of patients with ILDs and 2) to investigate whether baseline exercise capacity, disease severity or ILD etiology might affect outcomes.

Methods: Forty-one patients (IPF 63%, age 66.9 ± 11 ys) were enrolled in a standard PR course in two centers. Lung function, incremental and endurance cyclo-ergometry, Six Minutes Walking Distance (6MWD), chronic dyspnea (Medical Research Council scale-MRC) and quality of life (St. George Respiratory Questionnaire-SGRQ) were recorded before and at the end of PR to measure any pre-to-post change. Correlation coefficients between the baseline level of Diffuse Lung Capacity for Carbon monoxide (DLCO), Forced Vital Capacity (FVC), 6MWD, power developed during incremental endurance test, GAP index (in IPF patients only) and etiology (*IPF* or *non-IPF*) with the functional improvement at the 6MWDT (meters), at the incremental and endurance cyclo-ergometry (endurance time) and the HRQoL were assessed.

Results: Out of the 41 patients, 97% (n = 40) completed the PR course. Exercise performance (both at peak load and submaximal effort), symptoms (iso-time dyspnea and leg fatigue), SGRQ and MRC significantly improved after PR (p < .001). Patients with lower baseline 6MWD showed greater improvement in 6MWD (Spearman r score = -.359, p = .034) and symptoms relief at SGRQ (r = -.315, p = .025) regardless of underlying disease.

Conclusion: Present study confirms that comprehensive rehabilitation is feasible and effective in patients with ILD of different severity and etiology. The baseline submaximal exercise capacity inversely correlates with both functional and symptom gains in this heterogeneous population.

Keywords: Pulmonary rehabilitation, Interstitial lung diseases, Endurance test, Endurance time, Functional performance

* Correspondence: roberto.tonelli@me.com

¹Respiratory Diseases Unit-AOU Policlinico, Department of Medical and Surgical Sciences, University of Modena Reggio Emilia, Modena, Italy Full list of author information is available at the end of the article



© The Author(s). 2017 **Open Access** This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

Background

Interstitial lung diseases (ILDs) comprise a heterogeneous group of chronic conditions characterized by lung parenchymal involvement with different degrees of inflammation and fibrosis resulting in impaired gas exchange and restrictive physiology. Clinically, the development of irreversible and progressive parenchymal fibrosis leads to ventilatory constraint and abnormal lung mechanics with limited exercise capacity and dyspnea on exertion [1]. In particular, the impaired level of gas exchange seems to be the major cause leading to exercise intolerance in these patients [2]. Consequently, as ILD progresses, the patient's daily activities decline early following symptoms (shortness of breath, tiredness, muscle fatigue). This reduction in everyday performance begins even before that ventilatory limitation with functional impairment occurs [3]. Furthermore ILD patients experience greater physical and social limitations once ventilatory constraint has established, reducing their functional reserves [2]. Finally, due to the progressive exercise limitation, individual's healthrelated quality of life (HRQoL) is markedly affected [4].

Thus, the proper management of dyspnea represents a critical need for patients suffering from ILD. Pulmonary rehabilitation (PR) programs have been widely assessed and validated in patients with Chronic Obstructive Pulmonary Disease (COPD) [5], for which they have been proved to be effective in reducing respiratory and non-respiratory (i.e. peripheral muscle fatigue) symptoms and improving functional performance status which is consistent with improvement in HRQoL [6]. Nonetheless, there is a growing body of evidence showing that PR consisting of tailored and supervised training on aerobic and resistance exercises, breathing techniques and education sessions focused on self-management of symptoms and physical activity promotion could improve outcomes like dyspnea, functional capacity and quality of life also in patients with ILD [7–11]. However, the ultimate guidelines on diagnosis and management of Idiopathic Pulmonary Fibrosis (IPF) [1] provide a weak recommendation for PR in this particular form of ILD, due to the low quality of evidence on the real achievable gain. Furthermore, there is still no clear evidence regarding which type of ILD form (idiopathic VS non idiopathic) could benefit the most from a PR program.

With reference to this, Huppmann and colleagues prospectively investigated the effect of a 4-week PR program in a wide population of 402 patients with ILDs. The authors reported clinically relevant improvements in both functional exercise capacity and in HRQoL independent on the underlying disease (which were IPF, cryptogenic organizing pneumonia, hypersensitivity pneumonitis, sarcoidosis, or others) [12]. However, in a systematic review by Holland et al., patients with IPF seem to have lower improvement in functional capacity following PR as compared with others suffering from different etiologies of ILD [13].

Even more, the impact of disease severity on PR outcomes seems not to be well clarified in patients with ILDs. Lower lung derangement, less oxyhaemoglobin desaturation and lower level of secondary pulmonary hypertension have been associated with greater improvement in functional capacity in patients with IPF but were not predictors of benefit in those with other ILDs [14]. This might indicate that PR is likely to be effective in IPF at early stage of the disease, whereas different etiology of ILD may benefit regardless of disease severity. Indeed, in a prospective multicenter cohort study, Ryerson and colleagues had showed that patients with ILD (of whom around only one third were IPF) and a low functional capacity at baseline had greater benefit from rehabilitation [3].

In summary despite the beneficial effect of PR in ILD patients has been supported by a growing body of evidence, the ultimate IPF guidelines still provide a weak recommendation for PR [1] and 2) the impact of disease severity and ILD etiology on PR outcomes remains not well understood. The present prospective observational two-center study has been designed 1) to confirm the positive impact of pulmonary rehabilitation delivered both as in- and outpatient program in a population of patients with ILDs of different etiology [15] and 2) to further investigate whether baseline exercise capacity, disease severity or ILD etiology might differently affect clinical outcomes following a standard PR course.

Methods

Patient population and study design

Between January 2013 and January 2015, 41 consecutive patients with ILDs of different etiology were prospectively enrolled as in-patients (n = 30, 73%) or out-patients (n = 11, 27%) to perform a standard comprehensive PR program [6] both at "Villa Pineta" Rehabilitation Hospital in Pavullo n/F (Modena) and at "Don Gnocchi" Institute in Firenze. Patients not fulfilling criteria for PR [6] were excluded. Diagnosis of ILD was performed according to the ATS/ERS international consensus classification of idiopathic interstitial pneumonias [16].

The study had a prospective design. It was approved by the local review board and the Ethics Committee at both Institutions. All the individuals gave their informed consent to participate and to publish data on scientific journals.

The study flowchart is shown in Fig. 1.

Assessment and outcomes

Functional status

Routine pulmonary function tests (forced vital capacity-FVC and forced expiratory volume in 1st second-FEV₁) and diffusive lung capacity (DLCO), obtained with subjects



in a comfortable seated position, were measured according to the American Thoracic Society/European Respiratory Society (ATS/ERS) recommendations [17]. Lung volumes were measured by volume-displacement body plethysmograph (Autobox DL 6200; SensorMedics, Yorba Linda, CA); the normal reference values were those by the European Community for Coal and Steel [18].

Blood gases were measured by an automated analyzer (IL-1650, Instrumentation Laboratory, Milan, Italy for the center of Modena and ABL 800, Radiometer, Copenaghen for the center of Florence) from an individual's arterial sample taken when in resting condition while breathing room air.

Disease severity score of IPF patients was recorded using the GAP-staging system [19], which includes gender, age, forced vital capacity (FVC) and diffusion lung capacity for carbon monoxide (DLCO).

Incremental exercise testing (1-min increments of 10 W) to a symptom-limited maximum was performed on an electronically braked ergometer (Ergo-Metrics 800 s; SensorMedics) for lower limbs pre-post PR. The patients were familiarized with the apparatus days prior to the test; they were encouraged to keep exercising for as long as possible while pedaling at 50 to 60 rpm and were instructed to maintain work levels until they reached a symptom-limit. In addition, the six-minute walked distance (6MWD) was carried out according to the ATS/ERS guidelines [20]. All the subjects underwent

two consecutive tests with the best result taken as representative.

A pulmonologist or a cardiologist and a rehabilitation therapist, unaware of the study purposes, closely supervised each exercise session; during the session, heart rate and peripheral oxygen saturation (SpO_2) were monitored continuously.

Symptoms and health-related quality of life

The perceptions of dyspnea and limb efforts were determined at the beginning and at the end of 6MWD and every 2 min during cycle-ergometry using the modified Borg scale [21]. This scale is a vertical list with labeled categories describing increasing intensities of the perceived symptom (dyspnea or fatigue). Subjects were asked to rank the overall intensity of respiratory discomfort and leg effort by pointing to a score on a large scale from zero (none, i.e. no sensation at all) to 10 (maximum, i.e. the most severe sensation they had ever experienced).

Chronic exertional dyspnea was assessed by using the 5-point (0 to 4) MRC scale [22].

The St. George Respiratory Questionnaire (SGRQ) as validated into the Italian version was used to assess the patient's health-related quality of life (HRQoL) [23]. Total and component (symptoms, activity and impacts) scores were recorded for analysis. This questionnaire was previously used also in patients with ILD [24, 25].

Maximal load and level of symptom achieved during incremental exercise test, endurance time and dyspnea recorded at iso-time during endurance test, distance walked, chronic dyspnea and HRQoL were used as outcome measures, suitable for a pre-to-post comparison.

Pulmonary rehabilitation program

The two centers involved provided a standardized similar and shared PR course as recommended [6, 26]. PR program consisted of: 6-h/week individually exercise training, including endurance training for upper and lower limbs, 2 session of breathing techniques lasting 30 min for four to five times per week and 3 sessions of group education per week. Exercise training consist of aerobic (treadmill, stationary bikes) and resistance training (light weights, resistance bands) and included supervised cycle and supported arm ergometry, and leisure walking used as outcome measures, suitable for a pre-to-post comparison.

Breathing training consisted of breathing techniques (controlled and diaphragmatic breathing), pacing and energy conservation. Supplemental oxygen was delivered to maintain normal level of oxygen saturation. PR was tailored on the patient's functional status and performance. The educational topics included medication and oxygen use, nutrition, panic control and relaxation techniques, as well as psychosocial support and issues of palliation and/or end-of-life related to the disease progression [6]. If needed, patients received psychosocial support. The PR program lasted at least 24 sessions of rehabilitation training and was conducted 6 days a week, once daily the first week and twice daily thereafter. Each rehabilitation session lasted at least 3 h.

Statistical analyses

Per-protocol analysis was conducted by means of STATA 11.2 statistical package (StataCorp, Texas, USA). Data were obtained from prospectively maintained medical records and computerized databases. A p value lower than 0.05 was required for significance.

We used Kolmogorov–Smirnov test in order to assess the distribution of data showing that all data follow a normal distribution. Pre to post comparisons were performed by paired and unpaired *t*-test, χ square test and Wilcoxon rank sum test. Changes from baseline in 6MWDT (meters) and in Endurance Time (minutes) and SGRQ (total score) following PR were considered as indicators of improvement in functional capacity and HRQoL respectively [3, 27]. The statistical power expected for the present study population was calculated with reference to the primary outcomes and the minimum difference appreciable by our sample was assessed at ≥3% of baseline values for major outcomes.

With reference to etiology patient were divided in "IPF" and "non IPF". Baseline DLCO, FVC and GAP

index (for IPF patients only) were considered indicators of lung derangement and disease severity. The distance covered at 6MWD and the power (watt) developed during the incremental endurance test at baseline were taken to assess functional performance.

To asses correlation between baseline exercise capacity, disease severity and ILD etiology with functional and symptoms improvement a bivariate analysis by Spearman rank correlation coefficient was applied.

Results will be described as mean \pm standard deviation (SD), except otherwise indicated.

Results

Forty out of 41 patients referred to PR completed the program; one subject with hypersensitivity pneumonitis (non IPF) dropped out as he developed bacterial pneumonia in hospital, once he had completed 20 rehabilitation sessions. Among patients who completed the program the number of rehabilitation sessions performed ranged from 26 to 32 and did not differ between in and outpatients (p = .15).

Demographics and baseline data on function, symptoms, disease etiology and severity are shown in Table 1. Baseline features were comparable with no statistically significant difference between patients enrolled in the two centers and involved in the in- and outpatient program. Mean DLCO and volumes (overall mean FVC 74.5 ± 21.42% of predicted value) indicated a mild-tomoderate lung restriction. Notably, the overall mean GAP index score in IPF patients was 3.74 ± 1.69 over a maximum of 7. Twelve subjects presented abnormal gas exchange on effort and were prescribed for using oxygen as a supportive therapy. At baseline, the overall mean distance covered at the 6MWDT was 376.8 m \pm 94.6 m and the overall mean endurance power developed during incremental exercise test was 57.5 w ± 23.7 w, which revealed a substantial reduction of exercise capacity both at submaximal and maximal performance test. Among patients with ILD other than IPF, 8 had pulmonary fibrosis associated to collagen vascular disease, 4 had chronic hypersensitivity pneumonitis, 2 had sarcoidosis and 1 had asbestos-related ILD.

In Table 2 outcome measures and their absolute or relative change following PR are displayed: the standard rehabilitation course resulted in a significant improvement in all the pre specified outcomes, being not influenced by PR setting and center of enrollment (see Fig. 2). Distance covered, peak and endurance exercise performance increased after PR, while symptoms score at peak exercise or iso-time decreased substantially. Notably, SGRQ score improved in all the categories. The minimal clinically important difference (MCID) in 6MWD and SGRQ was achieved by 75.6% and 80.3% of patients respectively. Nor IPF diagnosis neither other baseline features significantly influenced the achievement of the MCID in the major

Baseline characteristics	Overall	Center of provenience		р	Setting of PR program		р
		A ^a (n. 20)	B ^b (n. 21)		In (n. 30)	Out (n. 11)	
Age (years)	66.9 ± 10.9	67.8 ± 9.7	65.4 ± 11.1	.74	70.3 ± 15.7	62.4 ± 7.1	0.12
Gender (males: females)	27: 14	13: 7	14: 7	.99	21:9	8: 3	0.99
BMI (kg/m²)	28.1 ± 3.4	27.8 ± 2.4	28.3 ± 1.1	.84	27.6 ± 7.4	29.3 ± 1	0.46
Diagnosis (IPF: non IPF)	26: 15	13: 7	13: 8	.99	18: 12	8: 3	0.72
Smoking history (yes: no)	30: 11	16: 4	14: 7	.48	21:9	9: 2	0.69
GAP index -for IPF only-	3.74 ± 1.69	4.14 ± 2.19	3.24 ± 1.15	.11	4.36 ± 3.19	3.24 ± 1	0.345
Inpatients (in: out)	30: 11	15: 5	15: 6	.99	-	_	-
O ₂ therapy (yes: no)	12: 29	5: 15	7: 14	.73	10: 20	2: 9	.46
O ₂ therapy (continuous: during effort)	8: 4	3: 1	5: 3	.99	7: 3	1: 1	.99
Arterial pO_2 (mmHg)	68 ± 12	70 ± 10	66 ± 7	.14	64 ± 14	71 ± 3	.11
Arterial pCO_2 (mmHg)	37 ± 4	38 ± 5	37 ± 2	.4	37 ± 4	37 ± 1	.99
FVC (% predicted)	74.5 ± 21.4	72.4 ± 16.4	75.8 ± 7.4	.39	70.4 ± 16.4	77.8 ± 8.4	.16
FEV ₁ (% predicted)	78.7 ± 20.7	79.6 ± 19.6	77.4 ± 21.7	.74	72.6 ± 19.6	79.4 ± 12.7	.29
DLCO (% predicted)	45.5 ± 20.9	43.3 ± 21.2	46.9 ± 19.5	.57	42.4 ± 19.2	47.9 ± 12.5	.38

Table 1 Baseline features of study population

^aCenter A = "Don Gnocchi" Institute, Firenze, Italy

^bCenter B = "Villa Pineta" Rehabilitation Hospital in Pavullo n/F, Modena, Italy. Continuous variables are indicated as mean ± standard deviation, non continuous as n

outcomes. Lower 6MWD covered at baseline significantly correlated with improvement in 6MWD (Fig. 3, panel f), endurance time at cycle ergometer (Fig. 5, panel f) and in SGRQ (total score) (Fig. 4, panel f) following PR, whereas power reached at incremental endurance test, FVC, DLCO, GAP index score and etiology did not (see also in Figs. 3, 4 and 5 panels a to e).

Discussion

In this study we evaluated the impact of a standard comprehensive PR program conducted in the real life in patients with ILD of different etiology, functional performance and lung impairment. Overall, findings confirm that these patients are likely to gain large benefits from PR delivered as in and outpatient setting, both in terms of exercise performance, symptoms, and perceived quality of life regardless of underlying disease. In particular, our study extends the concept that patients with both IPF and non-IPF showing lower submaximal exercise capacity at baseline are more likely to benefit in terms of exercise performance and HRQoL.

Multiple aspects deserve discussion.

First, the extent of improvements experienced by the enrolled ILD-patients seems even greater than that reported in previous studies. In particular, the absolute change in the average distance covered at the 6MWDT

|--|

Outcomes	n	Before PR	Post PR	Absolute change	Relative change (%)	р
MRC	40	2.8 ± 0.8	1.7 ± 1.1	- 1.1 ± 0.8	-40.8 ± 35	<.001
SGRQ (Total)	39	50.6 ± 13.9	38.5 ± 13.7	- 12.1 ± 11.1	-23.3 ± 19	<.001
SGRQ (Activity)	39	67.6 ± 14	589 ± 19.6	- 8.8 ± 17.6	-12.5 ± 24.7	0.009
SGRQ (Impact)	39	43.7 ± 19.1	29.9 ± 15.9	- 13.7 ± 14.6	-29.9 ± 41.5	<.001
SGRQ (Symptoms)	39	40.4 ± 20.9	26.7 ± 20.8	- 17 ± 19.8	-36 ± 22.1	<.001
6MWDT (m)	39	376.8 ± 94.6	430.9 ± 96.4	54.1 ± 55.4	16.7 ± 37.8	<.001
Dyspnea 6MWTD (Borg Scale)	39	5.2 ± 2.3	3.8 ± 2.2	-1.4 ± 2.1	-23.4 ± 40.1	.015
Leg fatigue 6MWTD (Borg Scale)	39	3.5 ± 2.7	2 ± 2.2	-1.6 ± 1.5	-48.1 ± 71.5	.006
Cycle dyspnea ^a (Borg scale)	40	6.1 ± 2	4.2 ± 2.7	- 1.8 ± 1.9	-33 ± 35.1	<.001
Cycle leg fatigue ^a (Borg scale)	40	5.9 ± 2.1	3.7 ± 2.7	- 2.2 ± 2	-39.5 ± 106	<.001
Cycle endurance time (min)	40	7.7 ± 3.8	12.5 ± 8.4	4.8 ± 6.9	66 ± 108.1	<.001
Cycle endurance power (watt)	40	57.5 ± 23.7	88.2 ± 57.1	31 ± 53.5	63.4 ± 33.9	.003

Each outcome value is reported as mean value \pm standard deviation (SD); p value are referred to relative changes

^aCycle dyspnea and leg fatigue were assessed at isotime



after PR has been 54.1 \pm 55.4 m (+ 16.7% from baseline, p < .001), compared to the 44.34 m (95% CI 26.04 to 66.64) as reported in a recent meta-analysis [28, 29]; furthermore the average reduction seen in the MRC score for dyspnea has been greater than 1 (- 1.1 \pm 0.8, p < .001). The observed major benefits might be due to the peculiar baseline features of the study population, constituted of ILD patients with mild to moderate impairment of respiratory function and a measurable exercise capacity (although substantially reduced as expected) even at peak intensity. The permissive level of lung function derangement let almost all the patients to

complete the rehabilitation program (only 1 patient left the study protocol for the onset of infectious pneumonia), with increased exercise tolerance both at submaximal and maximal performance. Thus we can postulate that less deranged lung in interstitial diseases may provide greater chance to successfully undertake and complete rehabilitation training, issuing its positive results over a limited baseline functional capacity.

Second, data from our study also show how ILD patients of different nature and severity could equally gain benefits from PR without any significant difference in terms of improvement. In particular the nature of the fibrotic lung







changes does not impact the positive effect of PR. As a matter of fact, regardless of the etiology, patients affected with ILD share common clinical features with distressing dyspnea, exercise intolerance and persistent cough [3]. What might differentiate IPF from other forms of ILD, especially non-idiopathic, is the systemic involvement of the underlying disease that could affect the adherence to exercise training, thus reducing the effectiveness [10]. Therefore, our study has demonstrated that PR could provide benefits irrespectively of the nature of ILD and the ensuing eventual systemic involvement. Researches have widely questioned the impact of disease severity and the subsequent functional limitation on the effect of PR in ILD patients [8]. Ryerson and colleagues showed how a lower



baseline 6MWD could predict larger improvement in distance covered after PR [3]. Similarly, those with relatively higher levels of baseline dyspnea experienced major prolonged symptoms relief [14]. Conversely, other studies show that greater and sustained improvements in functional capacity following exercise training have been described when patients present a higher baseline FVC and less desaturation on the baseline 6MWDT [30], suggesting that PR could be more effective if offered earlier in the disease course. This discordance may be explained by differences in the patient population (e.g. the presence of comorbidities) or study design (cohort vs randomized trial). In our study the different level of lung derangement expressed by levels of FVC and DLCO do not affect PR benefits. This suggests that motivated ILD patients with functional impairment from mild to moderate should be early referred to PR because they are likely to benefit.

Third, our study confirms that patients with lower baseline functional performance may experience the greater clinical benefit in terms of exercise capacity and symptoms relief. These data confirmed what already reported by Ryerson and colleagues [3], adding a significant correlation with lower submaximal baseline functional capacity and greater HRQoL improvement after PR. It seems interesting to stress that the power reached at incremental endurance test performed at baseline did not inversely correlate with functional improvement after PR. In a relatively recent study, Holland and colleagues directly compared the cardiorespiratory responses during 6MWDT and cardiopulmonary exercise test (CPET) in patients with ILD [31]. They showed how desaturation on exertion was higher during 6MWDT than during CPET independently on the physiological load. This seems due to the increased alveolar ventilation reached during cycling, with a subsequent offset of oxyhaemoglobin desaturation through augmentation in partial pressure of oxygen in arterial blood (PaO₂). They conclude that 6MWDT, eliciting high but submaximal functional responses in ILD patients, is a unique tool to derive information across the range of disease severity. Our study enhances this concept suggesting that 6MWDT seems even more sensitive than incremental endurance test in identifying ILD patients with greater functional improving potential. It is worth to note that patients with a higher baseline 6MWD showed less increase in distance covered after PR, probably due to a reduced capacity for further improvement. Moreover a greater 6MWDT was not correlated with lack of improvement in other functional or symptomatic outcomes, thus confirming that physical intervention (rehabilitation) is a critical approach even in patients with a higher level of baseline exercise capacity.

Our study presents several limitations; first of all this was a preliminary trial whose explorative nature resulted in the lack of a sample size computation analysis. A subsequent study with a sample size computed on the basis of the evidence provided by the present study is needed. Secondly the limited size of the study population did not allow detailed subgroup analyses. In addition, the lack of a control group could limit the likely significance of our findings, while the lack of follow-up after PR did not allow to assess the persistence of the reported benefits over time nor comparing it between diagnosis (e.g. IPF vs non-IPF). Furthermore, all the patients enrolled showed a baseline mild to moderate impairment of lung function, without any case of severe lung derangement. This did not let to evaluate the effect of PR model on the more compromised patients and to investigate whether a more severe functional limitation could significantly affect the improvement after PR.

Future research is therefore required to better define the subset of ILD patients that can benefit most from exercise training and investigate whether different and/or more specific modalities should be used to enhance gains following PR in patients with more severe disease.

Conclusions

There is a growing body of evidence to support pulmonary rehabilitation as a fundamental treatment for patients with lung fibrosis that should be early referred to tailored exercise programs. Our study has demonstrated that in ILD patients the level of lung derangement or etiology at baseline do not affect outcomes improvement following rehabilitation. Moreover, a lower derangement in lung function may provide a greater chance to successfully undertake and complete PR. Since the degree of improvement across measured outcomes is remarkable, this suggests that pulmonary rehabilitation should be a first line therapy for managing symptomatic patients with ILD of different nature and mild to moderate severity.

Abbreviations

6MWD: Six Minutes Walking Distance; ATS: American Thoracic Society; CPET: Cardiopulmonary exercise test; DLCO: Diffuse Lung Capacity for Carbon monoxide; ERS: European Respiratory Society; ET: Endurance test; FVC: Forced vital capacity; GAP: Gender age pulmonary function; HRQoL: Health related quality of life; ILD: Interstitial lung disease; IPF: Idiopathic Pulmonary Fibrosis; MCID: Minimal clinically important difference; MRC: Medical Research Council; PAO₂: Partial pressure of oxygen in arterial blood; PR: Pulmonary rehabilitation; SGRQ: St. George Respiratory Questionnaire

Acknowledgements

Not applicable.

Funding

Not applicable

Availability of data and materials

Raw data is publicly available and can be found at the Respiratory Disease Unit of the Azienda Ospedaliero Universitaria Policlinico di Modena, Via del Pozzo 71, 41,124, Modena, Italy.

Authors' contributions

RT and IC designed the study, enrolled the patients, collected data, wrote the manuscript and performed statistical analysis. IR, RF helped in designing the study, enrolled the patients, performed the study procedures, collected data and wrote the manuscript. BL, FF helped in designing the study and collecting data. AM and BB participated in the design of the study and performed the statistical analysis. EMC and FG conceived of the study, and participated in its design and coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The study was approved by the local review board and the Ethics Committee at both Institutions (Ethics Committee of Modena, University Hospital of Modena, via del Pozzo 71, 41,124 Modena, Italy and Ethics Committee of Florence, IRCSS don Gnocchi, Via di Scandicci, 269, 50,143 Firenze, Italy ref. number 64/14). All the individuals gave their written informed consent to participate.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details

¹Respiratory Diseases Unit-AOU Policlinico, Department of Medical and Surgical Sciences, University of Modena Reggio Emilia, Modena, Italy.
²Respiratory Diseases Unit, Don Gnocchi IRCCS, Florence, Italy. ³Rehabilitation Unit, Villa Pineta Hospital, Pavullo n/F, Modena, Italy. ⁴Respiratory Diseases Unit, Cardiovascular Department, University of Padua, Padua, Italy.

Received: 1 February 2017 Accepted: 2 October 2017 Published online: 10 October 2017

References

- Raghu G, 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.
- Katzenstein AL, Myers JL. Idiopathic pulmonary fibrosis: clinical relevance of pathologic classification. Am J Respir Crit Care Med. 1998;157:1301–15.
- Ryerson CJ, Cayou C, Topp F, Hilling L, Camp PG, Wilcox PG, Khalil N, Collard HR, Garvey C. Pulmonary rehabilitation improves long-term outcomes in interstitial lung disease: a prospective cohort study. Respir Med. 2014;108(1):203–10.
- Chang JA, Curtis JR, Patrick DL, Raghu G. Assessment of health- related quality of life in patients with interstitial lung disease. Chest. 1999;116:1175–82.
- From the global strategy for the diagnosis, management and prevention of COPD. Global initiative for chronic obstructive lung disease (GOLD); 2017. Available from: http://www.goldcopd.org.
- Nici L, Donner C, Wouters E, et al. American Thoracic Society/European Respiratory Society statement on pulmonary rehabilitation. Am J Respir Crit Care Med. 2006;173:1390–413.
- Nishiyama O, Kondoh Y, Kimura T, Kato K, Kataoka K, Ogawa T, Watanabe F, Arizono S, Nishimura K, Taniguchi H. Effects of rehabilitation in patients with idiopathic pulmonary fibrosis. Respirology. 2008;13:394–9.
- Ferreira A, Garvey C, Connors GL, Hilling L, Rigler J, Farrell S, Cayon C, Shariat C, Collard HR. Pulmonary rehabilitation in interstitial lung disease. Benefits and predictor of response. Chest. 2009;135:442–7.
- Holland AE, Hill CJ, Conron M, Munro P, McDonald CF. Short term improvement in exercise capacity and symptoms following exercise training in interstitial lung disease. Thorax. 2008;63(6):549–54.
- Naji NA, Connor MC, Donnelly SC, McDonnell TJ. Effectiveness of pulmonary rehabilitation in restrictive lung disease. J Cardpulm Rehabil. 2006;26(4):237–43.
- Salhi B, Troosters T, Behaegel M, Joos G, Derom E. Effects of pulmonary rehabilitation in patients with restrictive lung diseases. Chest. 2009;137(2): 273–279.

- Huppmann P, Sczepanski B, Boensch M, Winterkamp S, Schönheit-Kenn U, Neurohr C, Behr J, Kenn K. Effects of in-patient pulmonary rehabilitation in patients with interstitial lung disease. Eur Respir J. 2013;42:444–53.
- Holland AE, Hill C. Physical training for interstitial lung disease. Cochrane Database Syst Rev. 2008;4:CD006322.
- Holland AE, Hill CJ, Glaspole I, Goh N, McDonald CF. Predictors of benefit following pulmonary rehabilitation for interstitial lung disease. Respir Med. 2012;106:429–35.
- Kenn K, Gloeckl R, Behr J. Pulmonary rehabilitation in patients with idiopathic pulmonary fibrosis – a review. Respiration. 2013;86:89–99.
- American Thoracic Society. European Respiratory Society international multidisciplinary consensus classification of the idiopathic interstitial pneumonias. Am J Respir Crit Care Med. 2002;165:277–304.
- Wanger J, Clausen JL, Coates A, Pedersen OF, Brusasco V, Burgos F, Casaburi R, Crapo R, Enright P, van der Grinten CPM, Gustafsson P, Hankinson J, Jensen R, Johnson D, MacIntyre N, McKay R, Miller MR, Navajas D, Pellegrino R, Viegi G. Standardisation of the measurement of lung volumes. Eur Respir J. 2005;26:511–22.
- Quanjer PH, Tammeling GJ, Cotes JE, Pederson OF, Peslin R, Yernault JC. Lung volumes and forced ventilatory flows. Report working party standardization of lung function tests, European Community for steel and coal. Official statement of the European Respiratory Society. Rewiew. Eur Respir J. 1993;16(suppl 9):5–40.
- Ley B, Ryerson CJ, Vittinghoff E, Ryu JH, Tomassetti S, Lee JS, et al. A multidimensional index and staging system for idiopathic pulmonary fibrosis. Ann Intern Med. 2012;156(10):684–91.
- 20. American Respiratory Society. ATS statement: guidelines for the six-minute walk test. Am J Respir Crit Care Med. 2002;166:111–7.
- Borg GAV. Psychophysical basis of perceived exertion. Med Sci Sports Exerc. 1982;14:377–87.
- 22. Medical Research Council. Committee on reasearch into chronic bronchitis. Instruction for use of the questionnaire on respiratory symptoms: Devon WJ, Holman; United Kingdom; 1966.
- Carone M, Bertolotti G, Anchisi F, Zotti AM. The St George's Respiratory Questionnaire (SGRQ): Italian version. Rassegna Patol App Respir. 1999; 14:31–7.
- 24. Patel AS, Siegert R, Brignall K, et al. S133 The assessment of health related quality of life in interstitial lung disease with the King's brief interstitial lung disease questionnaire (K-BILD). Thorax. 2011;66:A61.
- Nishyiama O, Taniguchi H, Kondoh Y, Kimura T, Ogawa T, Watanabe F, Nishimura K. Health-related quality of life in patients with idiopatic pulmonary fibrosis: what is the main contributing factor? Respir Med. 2005;99:408–14.
- Ries AL, Bauldoff GS, Carlin BW, Casaburi R, Emery CF, Mahler DA, Make B, Rochester CL, Zuwallack R, Herrerias C. Pulmonary rehabilitation: joint ACCP/ AACVPR evidence-based clinical practice guidelines. Chest. 2007;131(5 Suppl):4S–2.
- Swigris JJ, Wamboldt FS, Behr J, et al. The 6 minute walk in idiopathic pulmonary fibrosis: longitudinal changes and minimum important difference. Thorax. 2010;65:173–7.
- Lacasse Y, Goldstein R, Lasseron TJ, et al. Pulmonary rehabilitation for chronic obstructive pulmonary disease. Cochrane Database Syst Rev. 2006;4:CD003793.
- Dowman L, Hill CJ, Holland AE. Pulmonary rehabilitation for interstitial lung disease. Cochrane Database Syst Rev. 2014;(10):CD006322.
- Kozu R, Jenkins S, Senjyu H. Effect of disability level on response to pulmonary rehabilitation in patients with idiopathic pulmonary fibrosis. Respirology. 2011;16:1196–202.
- Holland AE, Dowman L, Fiore LJ, Brazzale D, Hill CJ, McDonald CF. Cardiorespiratory responses to 6-minute walk test in interstitial lung disease: not always a submaximal test. BMC Pulm Med. 2014;14:136.