Pulmonary rehabilitation for cystic fibrosis: A narrative review of current literature

Evgenia I. Kalamara¹, Evangelos T. Ballas², Georgia Pitsiou¹,³, Guergana Petrova²,⁴

¹Respiratory Failure Unit, General Hospital of Thessaloniki “G. Papanikolaou”, Thessaloniki, Greece; ²Medical University Sofia, Bulgaria; ³Medical school Aristotle university of Thessaloniki; ⁴Pediatric Clinic, University Hospital Alexandrovska, Sofia, Bulgaria

Abstract

Pulmonary rehabilitation is a key component in cystic fibrosis care. This review summarizes the recent evidence in the area of pulmonary rehabilitation for cystic fibrosis in the form of questions and answers regarding interventions, indications, benefits and risks of pulmonary rehabilitation. Pulmonary rehabilitation includes airway clearance techniques, exercise training, education and behaviour change and can improve patients’ exercise capacity, muscle strength, quality of life and nutritional status. Airway clearance techniques have beneficial effects for clearing mucous. Over the past years, evidence for the beneficial effects of exercise training on exercise capacity and overall lung health is growing. In cystic fibrosis, multiple factors result in reduced exercise capacity. All modalities of pulmonary rehabilitation should be offered to patients with cystic fibrosis, as the benefits in most cases outweigh the risks, though the optimal regimens need to be yet defined.

Introduction

Cystic fibrosis (CF) is an autosomal recessive disease and one of the most common life-shortening condition affecting approximately 70,000 individuals worldwide [1], caused by mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene. The CFTR protein is expressed in epithelial cells and has several functions, primarily serving as an ion channel. It regulates liquid volume on epithelial surfaces through chloride secretion and inhibition of sodium absorption.

Airway disease is hypothesized to be mainly a result of reduced airway surface liquid volume which subsequently leads to mucociliary dysfunction, increase of bacterial load, and chronic inflammation. The abnormal composition and secretion of mucus affects multiple organs and causes various complications, thus rendering CF a systemic disease. Therefore, CF management requires a multidisciplinary approach, preferably in the setting of a specialized center. The treatment should be individualized to the needs of each patient, and should aim at the management of airway infection and the development of an active lifestyle, in order to maintain good nutrition and exercise capacity [2].

Pulmonary rehabilitation is an intervention customized to each patient and includes airway clearance techniques, exercise training, education and behaviour change. Its target is the amelioration of the physical and psychological status of people with chronic respiratory disease and the promotion of long-term adoption of health-improving behaviour. It is acknowledged as an important adjunct to the treatment of many chronic respiratory diseases [3]. In patients with CF, pulmonary rehabilitation traditionally focused on airway clearance techniques. Nowadays the management also extends to other areas. Systematic exercise training and personalized rehabilitation programs are offered to CF patients [4], while new technologies are being gradually incorporated to the rehabilitation process during the past few years.

This review was conducted in order to map the recent research done in the area of pulmonary rehabilitation for cystic fibrosis and answer questions about its main aspects, as an aid for respiratory medicine physicians, general practitioners, physiotherapists or other clinicians involved in the care of CF patients.
Methods

This literature review took place between March 2019 and September 2020. Our protocol was drafted using the Preferred Reporting Items for Systematic Reviews and Meta-analysis Protocols, with appropriate modifications for the purposes of a narrative review [5]. For this review all relevant information was identified using Embase, Cochrane Library, Science Direct, Scopus, PubMed databases and Google Scholar.

A literature search was performed. Search words were cystic fibrosis, pulmonary rehabilitation, exercise capacity, lung function, quality of life, physiotherapy, airway clearance, exercise training. Relevant peer-reviewed journal papers were included if they were published between the period of 2000 to 2020 and written in English. The papers could include subjects of various ages (children, adolescents, adults) and severity of disease. Papers were excluded if they did not fit into the conceptual framework of the study, for example papers focused on chronic respiratory conditions other than cystic fibrosis.

The selected literature included 23 Randomized control trials, 8 Cochrane Systematic Reviews, 7 guidelines, 7 clinical studies, 12 reviews, 1 systematic review, 2 other original articles.

What interventions does pulmonary rehabilitation include?

Airway clearance techniques

An integral part of pulmonary rehabilitation is the application of airway clearance techniques, that are used to enhance the mucociliary clearance system, in order to transport secretions proximally up the airways [6].

These include conventional chest physiotherapy (mainly involves postural drainage, percussion and vibration), active cycle of breathing techniques (ACBT), which consist of breathing control exercises, thoracic expansion exercises and forced expirations, positive expiratory pressure (PEP) therapy, which is defined as breathing against a positive expiratory pressure using a mask or mouthpiece, autogenic drainage (three-level breathing sequence beginning at low lung volumes, followed by breathing at mid-lung volumes, followed by deep breathing, huff and coughing- it uses controlled breathing to achieve the highest possible airflow in different bronchi generations), mechanical percussion, high-frequency chest compression and non-invasive ventilation (NIV).

Notably, PEP therapy can be oscillating or non oscillating. Oscillating PEP combines oscillation of airflow with PEP in order to loosen secretions [7,8]. Aerobic exercise can also be considered as an airway clearance technique, as it reduces mechanical impedance of sputum and enhances expiratory flow rates [9].

Exercise training

In pulmonary rehabilitation, out-patient, in-patient and home-based programs can be utilized. Exercise is a basic component of rehabilitation programs, and it includes lower and upper extremity training, inspiratory muscle training, as well as chest physical therapy techniques. Usually, the proposed schedule consists of a minimum of two sessions per week, preferably three or more, for 4, 6, 9 or 12 weeks (minimum of 12 supervised sessions recommended). The duration varies from 30 to 60 min and exercise can be continuous or intermittent and training can target endurance and/or strength. Patients should be able to continue effective training once supervised sessions have ended and regular physical activity five times a week for 30 min each time is encouraged regardless of rehabilitation, as healthy living advice [10,11].

Notably, the American College of Sports Medicine recommend moderate-intensity aerobic (endurance) physical activity for a minimum of 30 min on five days each week or vigorous-intensity aerobic physical activity for a minimum of 20 min on three days each week in order to promote and maintain health in adults [12].

As far as home-based programs are concerned, several protocols have been used, based on aerobic and muscle strength training. Happ et al. utilized an at-home bicycle exercise regimen performed three times a week, consisting of two exercise routines of different intensity performed in an alternating sequence during each session. Schneiderman-Walker et al. also used a three times weekly program that required a minimum of 20 min of aerobic exercise at a heart rate of approximately 150 beats/min. While there is still no standardization regarding self-regulated, home-based exercise, further evaluation is needed as such interventions are fairly easily conducted and low-cost [13-15].

Kriemler et al. compared the effects of 6 months training (aerobic or anaerobic) versus no training and found similar improvements in FEV1, FVC and exercise capacity. However, anaerobic training lowered static hyperinflation, while aerobic training did not. The authors assumed that improvements in FVC with strength training might be because of a decrease in hyperinflation in this group [16]. Based on these facts, aerobic and anaerobic training can be regarded as similarly effective.

A randomized control trial by Selvadurai et al. compared aerobic and resistance training in children with cystic fibrosis (CF) admitted to hospital with an intercurrent pulmonary infection with a control group. The study demonstrated that children who received aerobic training had significantly better peak aerobic capacity, activity levels, and quality of life than children who received the resistance training program. Children who received resistance training had better weight gain (total mass, as well as fat-free mass), lung function, and leg strength than children who received aerobic training. Therefore it is implied that a combination of aerobic and anaerobic training may be the optimal training modality for CF patients [17].

It is noteworthy that not all training modalities are suitable for all patients, as in the case of severe pulmonary impairment. Gruber et al. used an individualized training program with supplemental oxygen over 6 weeks in patients with severe disease and found comparable improvements in exercise capacity with conventional training in patients with less severe disease [18]. This study is in favor of individualized training interventions.

Concerning the intensity of training, most studies that managed to improve exercise capacity used intensities at about 50% VO2max or 60-85% of maximum heart rate [16,19-21]. A study of Whitley et al assessed the immediate effects of different exercise intensities on pulmonary function and diffusion capacity. Moderate exercise improved diffusion, while vigorous exercise caused airflow restrictions [22]. Taking all these into account, the optimal exercise training protocol for CF patients is yet to be defined.

Education, behavioural change promotion and adjunctive therapies

In addition to these, in pulmonary rehabilitation programs psychosocial assessment and intervention are offered, as depression and anxiety are common in patients with chronic lung disease. Evaluation of disability and education of both patient and family...
are part of this type of intervention [11]. Moreover, nutritional interventions are appropriate, considering the high probability of malnourishment and decreased muscle mass among patients with chronic respiratory conditions [11]. Adjunctive therapies, such as bronchodilators, oxygen therapy, non-invasive ventilation (NIV) and neuromuscular stimulation are also of considerable help during a rehabilitation program. Last but not least, patients referred to pulmonary rehabilitation should be assessed about their smoking status and smoking cessation services should be available [11].

Integration of new technologies in pulmonary rehabilitation programs

It is noteworthy that for the last couple of years, there has been special interest in new technologies, including video games, social media and web-based platforms and the potential for their application in the rehabilitation process.

In the CF population, a randomized controlled trial used the Nintendo Wii platform to deliver a 6-week home training program, with participants followed up for 12 months after the intervention. Exercise capacity, muscular strength and quality of life were improved in the short-term. The effects of training on muscle performance and quality of life were sustained over 12 months. However, adherence was 95% at 6 weeks but substantially reduced in the long term, with 65% of the subjects not using the active video game at all at 12 months [23]. Bishay et al. compared a fitness tracker with a personalized exercise prescription and social media platform to exercise prescription alone, although no significant difference in patient outcomes was noted between the two groups [24].

An RCT by Salolini et al. involving children and adolescents with CF compared a traditional stationary cycle training intervention to a training intervention using the interactive Xbox Kinect platform. Heart rate, SpO2, dyspnea, fatigue and subject satisfaction were evaluated. The study concluded that training with Xbox Kinect provided a cardiovascular demand similar to a stationary cycle, while it caused less dyspnea and fatigue and was more enjoyable than the stationary cycle. Therefore, such a modality has the potential to be used as an exercise intervention in young patients with CF. Similarly, Holmes et al. reported that training using the Xbox Kinect may be a suitable alternative to conventional exercise modalities for adults with CF [25,26]. O’Donovan et al. concluded that active video games are a useful source of light to moderate intensity physical activity in children with cystic fibrosis [27]. There are also other examples of studies including social media or web-based platforms as an aid to exercise [28,29] and several relative studies that are on-going [30,31].

These new modalities have potentially beneficial effects on fitness and training adherence, however further research is needed to study the long-term effects of such interventions [32].

What are the benefits of pulmonary rehabilitation in general?

Pulmonary rehabilitation is known to improve exercise capacity, muscle strength, dyspnea and health status compared with usual care. Self-reported measures of activities of daily living (ADL) and psychological status also improve. In addition to these, pulmonary rehabilitation modestly ameliorates physical activity levels and body weight/nutritional status [11].

What are the reasons for reduced exercise capacity in patients with cystic fibrosis?

Thick bronchial secretions, altered respiratory mechanics and gas exchange and decreased lung function are hallmark features of the disease. Severe airflow limitation and dynamic hyperinflation results in increased dead space ventilation and exertional dyspnea. In certain patients, concomitant cardiovascular abnormalities may play a role. Another important factor in the impairment to exercise is the involvement of peripheral muscles. Defective muscle metabolism, malnutrition, electrolyte disturbances, physical myocardial infarction, severe pulmonary hypertension, unstable diabetes, severe exercise-induced hypoxemia, abdominal aortic aneurysm >5.5 cm deemed inoperable, severe locomotor impairment, severe peripheral vascular disease. Stable cardiovascular disease is not a contraindication. Active cigarette smoking is, in certain cases, considered as a relative contraindication, although patients with COPD should be referred regardless of their smoking status [10,11].

As far as CF patients are concerned, pulmonary rehabilitation is a key element of care. Airway clearance techniques are frequently described as a cornerstone of CF treatment and should be performed across the lifespan. Exercise is also recommended, and patients should be offered an individualized program, according to their capability and preferences. Regular physical activity should be encouraged and should include weightbearing exercise in order to optimize bone density, while strength training programs should be prescribed to optimize muscle mass. Nutritional and psychological interventions are also acknowledged as an integral part of the standard of care in CF [7,33].

Even in complex CF cases, physiotherapy management is advised to be continued, with proper alterations and modifications, rather than discontinued, when possible. For instance, in case of hemoptysis the regimens are altered to minimize the risk of re-bleeding and in case of pneumothorax physiotherapy is continued when feasible but, minimizing the amount of positive pressure generated inside the patient lungs [34]. The American Diabetes Association Clinical Care Guidelines for CF-related diabetes recommend that patients should perform moderate aerobic exercise for at least 150 min per week, but monitoring of blood glucose levels before activity, consumption of extra carbohydrates or alterations of insulin dosage may be required [35].

Pregnant women with CF are encouraged to maintain a regular airway clearance routine, though postural drainage should be avoided. Exercise programs should be appropriate for the cardiorespiratory and musculoskeletal changes during pregnancy, and adequate hydration during exercise should be maintained [36].

What are the indications and contraindications of pulmonary rehabilitation?

Presence of long-term debilitating symptoms, dyspnea Medical Research Council (MRC) score of more than 2, motivation of patient to follow the rehabilitation program, rehabilitation prior to volume reduction surgery or lung transplantation are some of the criteria for patient selection for pulmonary rehabilitation in general. However, rehabilitation is not usually appropriate when the patient is considered unable to follow the proposed program or when serious comorbidities are present, such as angina pectoris, recent
inactivity, systemic inflammation and oxidative stress, oral corticosteroid use and CF-specific gene defects can be implicated in the peripheral muscle dysfunction. In certain cases, diaphragm strength is decreased despite a normal muscle mass, which can be possibly attributed to functional changes due to hyperinflation [37].

### What are the benefits of airway clearance techniques in CF patients?

Concerning airway clearance, bibliography clearly recommends its application when compared to no airway clearance or cough alone [11,38-40]. A 2015 Cochrane review found a significant increase in the amount of sputum expectorated in the patient groups that applied airway clearance compared to spontaneous cough or not using any airway clearance technique, concluding that methods of clearing the airways have short-term benefits for moving mucous [39]. Most studies do not show significant differences in the lung function of CF patients after chest physiotherapy [8,39,41-43]. Evidence from the Cochrane systematic reviews support that no one airway clearance regimen is better than another [8,31].

### What are the benefits of exercise training in CF patients?

Evidence suggests that CF patients with better physical fitness have better quality of life [7]. Exercise programs can improve fitness, exercise capacity, thoracic mobility, quality of life, maintain bone mineral density and lower the rate of decline in pulmonary function [13,19,44-51]. Moreover, a training effect, as measured by a decrease in lactate levels and heart rate can be achieved [52]. Both aerobic or anaerobic physical training has a positive effect on primary outcomes in CF patients (exercise capacity, strength and lung function) [53].

**Pulmonary function and exercise capacity**

A prospective, longitudinal study by Schneiderman et al. that included 212 patients with CF recruited over a 9-year period, concluded that patients with CF with increasing activity levels had a reduced rate of FEV1 decline compared to those who did not become more active over the study period [54]. Kriemler et al. and Hebestreit et al. both studied the effects of physical training over a 6-month period and found improved exercise capacity and lung function in the intervention group compared to the control group [16,55]. Santana-Sosa et al. reported amelioration of muscle strength and VO2 peak after an 8-week exercise program [20,21]. A retrospective cohort study of 2014 concluded that regular exercise is associated with a reduced decline in FEV1 and body mass index (BMI) in adults with CF. It is noteworthy that exercise capacity seems to be linked to the prognosis of CF patients. VO2 peak, peak work rate, ventilatory equivalent for oxygen and carbon dioxide have been found as predictors of death or lung transplantation at 10-year follow-up [56].

**Health-related quality of life, need for hospitalization, body posture and strength**

VO2 peak correlates with HRQL in CF patients and when improved it is associated with better nutritional status and health perception [57]. Hebestreit et al. concluded that a combined aerobic and strength training ameliorates exercise capacity, lung function and HRQL [19]. Another RCT by Klijn et al. has shown that anaerobic training has a beneficial effect on aerobic and anaerobic performance, as well as on quality of life [50].

Urquhart et al. reported that supervised, outpatient exercise and physiotherapy are associated with improvements in QOL and exercise tolerance, a reduction in intravenous antibiotic days, and a trend towards reducing lung function decline in children with CF [58]. Perez et al. detected that VO2 peak was the only variable significantly associated with time to hospitalization, therefore greater aerobic fitness is associated with a lower risk of hospitalization [59].

An RCT by Schindel et al. studied the effect of exercise training on posture, and resulted that after 3 months the intervention group showed a decrease in cervical and lumbar lordosis, thoracic kyphosis, lateral chest distance and abdominal protrusion [60]. Rovedder et al. reported increased upper limb strength over a 3-month combined aerobic and strength training program [14].

### Are there risks related to exercise? And if so, do the benefits outweigh the risks in the CF population?

A patient may experience dyspnea, productive cough and fatigue during exercise. Moreover, some patients with bronchiectasis or asthma may suffer exercise-induced bronchoconstriction. Exercise-induced hypoxemia (drop in oxygen saturation by more than 4 or below 90%) may also be observed, especially in patients with advanced lung disease. An estimated 20-25% of all patients with CF may present exercise-induced hypoxemia [36]. Patients with exercise-induced hypoxemia, as well as patients treated with macrolides are also at higher risk of developing cardiac arrhythmias during exercise. In general, approximately 5-10% of all patients present cardiac arrhythmias with exercise. Furthermore, exercise can potentially trigger a pneumothorax or an episode of hemoptysis, although the evidence is scarce. Musculoskeletal injuries are always possible with exercise, just as in the healthy population [61].

As far as CF patients are concerned, electrolyte losses, injury to spleen or oesophageal hemorrhage in the presence of portal hypertension, hypoglycemia in the setting of CF-related diabetes mellitus and worsening of preexisting arthritis are potential risks during exercise. In addition to these, when pulmonary rehabilitation programs are organized, special attention should be given to hygiene aspects to avoid cross-infection.

The number of different complications described may be unsettling, however the frequency of true adverse events is small and these mainly appear in patients with advanced disease. Individual assessment of each patient is always advised, but in most cases the benefits of exercise clearly outweigh the potential harms [61-63].

### Discussion

Pulmonary rehabilitation is a key component in CF care. Traditionally, airway clearance techniques are considered a cornerstone of CF treatment and should be performed across the lifespan in CF [7]. Evidence supports that airway clearance...
regimens have beneficial effects for moving mucous and are similarly effective [8,39].

Regular exercise training is increasingly being recommended for CF patients. Further research is necessary to accurately assess the benefits of exercise training in the CF population. Moreover, there is a lack of studies investigating the effects of physical exercise training on other significant outcomes, such as bone health, diabetic control and pulmonary exacerbations. However, evidence for the beneficial effects of exercise training on lung health and exercise capacity is growing. Exercise improves aerobic capacity and probably lowers the rate of decline in pulmonary function. Exercise capacity is also correlated with the prognosis of CF patients [56]. Thus, there is no reason to discourage the integration of exercise in all CF patients care [32]. This is also supported by evidence that show a relatively small frequency of true adverse events of exercise. Personalized assessment of patients is needed, but in most cases the benefits of exercise outweigh the potential harms [61,64].

The optimal training components (type, frequency, duration, intensity) need to be determined. Taking into account the existing literature, exercise training programs need to be tailored to the patient. For non-complex cases, perhaps a program of moderate-intensity exercise, 5 times per week, for 30 min per session is a reasonable regimen [12,19-22,65]. There is an increasing interest in the integration of new technologies in the pulmonary rehabilitation process (video game activities, social media and web-based platforms). These new modalities have potentially beneficial effects on fitness and training adherence, however further research is needed to study the long-term effects of such interventions [32].

### Limitations

This article is a narrative review, therefore it represents a broad overview of a topic-related research area. Data extraction relies on description of study findings rather than on continuous or categorical statistical values.

### Conclusions

Pulmonary rehabilitation is a multidisciplinary, individualized intervention that incorporates exercise training, airway clearance, education and behaviour change and a key component in CF patients care, as it can improve exercise capacity, muscle strength, quality of life and nutritional status of patients. Both airway clearance techniques and exercise training are invaluable parts of pulmonary rehabilitation. Airway clearance regimens have beneficial effects for moving mucous, and they seem to be similarly effective. On the other hand, exercise improves aerobic capacity (commonly affected in CF due to multifactorial pathology) and probably lowers the rate of decline in pulmonary function in CF patients. The optimal training components (type, frequency, duration, intensity) need to be determined, but it seems that a combination of aerobic and anaerobic training may be the optimal training modality for CF patients. When planning a training program, personalized assessment is needed, but in most cases the benefits of exercise outweigh the potential harms, so there is no reason to discourage the integration of exercise in all CF patients care.

In the past few years, new technologies (video game activities, social media and web-based platforms) can also be integrated in the rehabilitation process. Further research is needed to study the long-term effects of such interventions, however new means offer new possibilities for CF care. For instance, tele-rehab could be evolved and become a safe standard of care, especially during the COVID-19 era.

### References

5. PRISMA. Transparent reporting of systematic reviews and meta-analyses.


57. Hebestreit H, Schmid K, Kieser S, et al. Quality of life is asso-