Pulmonary Rehabilitation and Exercise in Pulmonary Arterial Hypertension: An Underutilized Intervention

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Recommended Citation

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Pulmonary arterial hypertension (PAH) is a rare and devastating disease characterized by progressive increases in pulmonary arterial pressure and pulmonary vascular resistance which eventually leads to right ventricular failure and death (Badesch et al., 2009; Farber and Lo Scalzo, 2004). It may due to various etiologies such as left heart disease, parenchymal lung disease, chronic thromboembolic disease, hematologic disorders or it may be idiopathic in nature (Table 1) (Simonneau et al., 2013) though the clinical picture of these patients is similar. Patients with all forms of pulmonary hypertension present with progressive dyspnea and increasing exercise intolerance. More specifically WHO Group I pulmonary arterial hypertension (PAH) is due to direct injury to the pulmonary vessels and vascular bed. At present there is no cure for PAH, however over the past decade targeted pharmaceutical options have become available. In addition to these pharmaceutical options, exercise and pulmonary rehabilitation have also been shown to increase exercise capacity, WHO functional class, peak oxygen consumption and resting heart rate (Grunig et al., 2011; 2012a). It has also been seen that exercise may improve health related quality of life (HRQoL) (Chan et al., 2013), depression (Verma et al., 2014) and fatigue (Talwar et al., 2014). It is imperative that clinicians use pulmonary rehabilitation and exercise as an adjuvant therapy in the treatment of these patients as its benefits are clearly evident. This review will attempt to emphasize the importance of structured exercise training in the pulmonary hypertension population despite etiology and disease severity.

INTRODUCTION

Pulmonary hypertension is a rare and devastating disease characterized by progressive increases in pulmonary arterial pressure and pulmonary vascular resistance which eventually leads to right ventricular failure and death. Early thought process was that exercise and increased physical activity may be detrimental to PAH patients however many small cohort trials have proven otherwise. In addition to the many pharmaceutical options, exercise and pulmonary rehabilitation have also been shown to increase exercise capacity as well as various aspects of psychosomatic health. As pulmonary and exercise rehabilitation become more widely used as an adjuvant therapy patient outcomes improve and physicians should consider this in the therapeutic algorithm along with pharmacotherapy.

Keywords: Pulmonary arterial hypertension, Pulmonary rehabilitation, Six minute walk test, Exercise intolerance, Pulmonary hypertension, Health related quality of life

PULMONARY REHABILITATION

Over the last few decades pulmonary rehabilitation has advanced as a therapeutic option in patients of chronic respiratory disease. Evidence based medicine has determined that pulmonary
Pulmonary rehabilitation in pulmonary arterial hypertension

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Pulmonary rehabilitation can reduce dyspnea, increase exercise tolerance and improve health-related quality of life in all settings of chronic respiratory disease (Spruit et al., 2013). Pulmonary rehabilitation is defined as a comprehensive intervention based on a thorough patient assessment followed by patient-tailored therapies, which include, but are not limited to, exercise training, education, and behavior change, designed to improve the physical and psychological condition of people with chronic respiratory disease and to promote the long-term adherence of health-enhancing behaviors (Spruit et al., 2013).

Use of exercise training and pulmonary rehabilitation in cardiopulmonary diseases such as congestive heart failure (CHF) and chronic obstructive pulmonary disease (COPD) have shown patient outcome improvement and are well described (Hambrecht et al., 1998; Ries et al., 2007). However, in the setting of pulmonary hypertension only small cohorts of studies exist. The use of exercise training and pulmonary rehabilitation in a pulmonary hypertension population is still underutilized though studies have shown its beneficial effect (de Man et al., 2009; Martinez-Quintana et al., 2010).

Table 1. World Health Organization’s classification of pulmonary hypertension (Simonneau et al., 2013)

<table>
<thead>
<tr>
<th>Group I - Pulmonary Arterial hypertension (PAH)</th>
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<tbody>
<tr>
<td>Idiopathic PAH</td>
</tr>
<tr>
<td>Heritable PAH (BMPR2, ALK1, ENG, SMAD9, CAV1, KCNK3, Unknown)</td>
</tr>
<tr>
<td>Drug and toxin induced</td>
</tr>
<tr>
<td>Associated with (i) Connective tissue disease, (ii) HIV infection, (iii) Portal hypertension, (iv) Congenital heart disease, (v) Schistosomiasis</td>
</tr>
<tr>
<td>Pulmonary veno-occlusive disease and/or pulmonary capillary hemangiomatosis</td>
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<td>Persistent pulmonary hypertension of the newborn</td>
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<table>
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<tr>
<th>Group II - Pulmonary hypertension due to left heart disease</th>
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<tbody>
<tr>
<td>Left ventricular systolic dysfunction</td>
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<tr>
<td>Left ventricular diastolic dysfunction</td>
</tr>
<tr>
<td>Valvular disease</td>
</tr>
<tr>
<td>Congenital/ acquired left heart inflow/outflow tract obstruction and congenital cardiomyopathies</td>
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<tr>
<th>Group III - Pulmonary hypertension due to lung diseases and/or hypoxia</th>
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<tbody>
<tr>
<td>Chronic obstructive pulmonary disease</td>
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<tr>
<td>Interstitial lung disease</td>
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<tr>
<td>Other pulmonary diseases with mixed restrictive and obstructive pattern</td>
</tr>
<tr>
<td>Sleep disordered breathing</td>
</tr>
<tr>
<td>Alveolar hypoventilation disorders</td>
</tr>
<tr>
<td>Chronic exposure to high altitudes</td>
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<tr>
<td>Developmental lung disease</td>
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<tr>
<th>Group IV - Chronic thromboembolic pulmonary hypertension (CTEPH)</th>
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<th>Group V - Pulmonary hypertension with unclear multifactorial mechanisms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hematologic disorders: chronic hemolytic anemia, myeloproliferative disorders, splenectomy</td>
</tr>
<tr>
<td>Systemic disorders: sarcoidosis, pulmonary histiocytosis, lymphangioleiomyomatosis</td>
</tr>
<tr>
<td>Metabolic disorders: glycogen storage disease, Gaucher’s disease, hypothyroidism</td>
</tr>
<tr>
<td>Others: tumoral obstruction, fibrosing mediastinitis, chronic renal failure, segmental pulmonary hypertension</td>
</tr>
</tbody>
</table>

PULMONARY ARTERIAL HYPERTENSION AND EXERCISE IMPAIRMENT

Pulmonary arterial hypertension is characterized by a progressive increase in pulmonary vascular resistance, leading to right ventricular failure and eventual death (Simonneau et al., 2013). Prior to pharmacological intervention traditional therapies were utilized which included anticoagulants, diuretics, and supplemental oxygen. However, after the pathophysiologic process became well known disease-targeted therapies such as phosphodiesterase inhibitors, endothelin receptor antagonists and prostacyclins have become more widely used (Rubin, 2013). Combinations of these agents are used to decrease pulmonary vascular resistance, decrease pulmonary artery pressures and increase exercise tolerance (Pugh et al., 2013). Combination therapy with disease specific drugs has shown symptomatic relief as well as been shown to increase exercise capacity and possibly survival rate in pulmonary hypertension (Ghofrani et al., 2002; Hoeper et al., 2004). Though these medications address quantitative measures of pulmonary vasculature and hemodynamics, exercise capacity and quality of life continue to decline despite optimized treatment.

http://dx.doi.org/10.12965/jer.150190
Exercise capacity is reduced in the setting of pulmonary hypertension which is associated with depression and anxiety disorders as demonstrated by Lowe et al. (2004). Until a better understanding of exercise training in the setting of pulmonary hypertension was established the early thought process was that exercise and increased physical activity may be detrimental to these patients and may actually increase pulmonary pressures expediting the proliferative process (Badesch et al., 2004).

The pathogenesis of pulmonary hypertension is multifactorial. WHO group I pulmonary hypertension, also referred to as pulmonary arterial hypertension is characterized by abnormalities initiating in the small arteries of the pulmonary vasculature, including enhanced cell proliferation and reduced apoptosis, endothelial dysfunction, thrombosis in situ, inflammation, and plexiform arteriopathy, leading to vascular remodeling and excessive vasoconstriction (McLaughlin et al., 2009). This may due to connective tissue diseases, congenital heart disease, HIV infection, portal hypertension, heritable or it may be idiopathic in nature. In the setting of PAH these changes cause resistance for the blood flowing through the pulmonary arteries. This leads to increased pulmonary artery pressures and increased pulmonary vascular resistance pressures eventually leading to right heart failure due to increased afterload, decreased contractility, and reduced cardiac output (Waxman, 2012). The combination of reduced cardiac output and increased pulmonary vascular resistance creates a situation where exertion causes dyspnea and negatively affects quality of life.

Exertional dyspnea being a common symptom across the pulmonary hypertension population provides a method to assess disease severity and progress. Dyspnea may be measured subjectively by using instruments such as the Borg Dyspnea Scale and Medical Research Council Dyspnea Scale (Bestall et al., 1999; Borg, 1970). The severity of exertion is helpful to assess severity and prognosis in pulmonary hypertension. The 6-min walk test and the measure distance is used to assess exercise capacity in patients diagnosed with pulmonary disease and is used as the determinant and primary endpoint in many exercise training clinical trials (ATS Committee on Proficiency Standards for Clinical Pulmonary Function Laboratories, 2002). Exercise tolerance can also be measured using cardiopulmonary exercise testing (American Thoracic, Society/ American College of Chest Physicians, 2003).

CLINICAL TRIALS OF EXERCISE TRAINING AND PULMONARY REHABILITATION

Though early thought process advised against increased exercise and a change in thinking brought upon clinical trials of exercise respiratory training in the pulmonary hypertension population (Table 2). In Mereles et al., (2006) conducted the first prospective randomized study to evaluate the effects of exercise on chronically ill pulmonary hypertension patients, stable on medication. Thirty patients with either pulmonary arterial hypertension or chronic thromboembolic pulmonary hypertension were randomly assigned to either a control or training group. Primary end points were the change in 6-min walk distance from baseline to week 15 in the scores of the Short Form Health Survey quality-of-life questionnaire (SF-36). Secondary endpoints were also assessed. At week 15, patients in the primary and secondary training groups had an improved 6-min walking distance. Exercise training was shown to be well tolerated and improved scores of quality of life, WHO functional class and peak oxygen consumption. Systolic pulmonary artery pressure values at rest did not change significantly after 15 weeks of exercise and respiratory training (Mereles et al., 2006).

Though the ground work for exercise training in pulmonary hypertension was laid out, furthering testing was required. De Man et al. (2009) conducted a trial of exercise on only patients with WHO Group I PAH. Nineteen clinically stable PAH patients with moderate disease underwent exercise training program for 12 weeks in which maximal capacity endurance capacity and quadriceps function were assessed and serial quadriceps muscle biopsies were obtained. 6-min walk distance and peak exercise capacity did not change after training. However, endurance capacity improved significantly after training. Training also enhanced aerobic capacity of the quadriceps, by increasing capillarization (de Man et al., 2009).

A study by Fox et al. (2011) of 22 PAH or CTEPH patients demonstrated significant increase in 6 MWD as well as Peak VO2. The rehabilitation program consisted of 24 1-h sessions of exercise training/rehabilitation over the course of 12 weeks. Peak work rate during cardiopulmonary exercise test also increased in the rehabilitation group with borderline significance. Echocardiography and blood N-terminal pro-brain natriuretic peptide levels were unchanged. No adverse events occurred due to the rehabilitation program demonstrating safety.

With evidence of the positive benefits of exercise training in the setting of pulmonary hypertension further clinical trials were conducted. Grunig et al. (2011) demonstrated the benefits of exercise training in a connective tissue disease population with pulmonary hypertension (Grunig et al., 2012b). Patients with RHC confirmed connective tissue disease related pulmonary hypertension received in-hospital exercise training for 3 weeks which was con-
continued for another 12 weeks at home. Patients improved the mean distance walked in 6 min, resting heart rate, peak oxygen consumption, and oxygen saturation. In addition systolic pulmonary artery pressure, and diastolic systemic blood pressure improved significantly after 3 weeks of exercise training. The 1- and 2-yr overall-survival rates were found to be 100%, the 3-yr survival rate 73% (Grunig et al., 2012a).

Another study by Grunig et al. (2011) aimed to assess disease progression and survival in patients with severe chronic pulmonary hypertension. Fifty eight patients with severe pulmonary hypertension on stable medication received exercise and respiratory training in hospital for 3 weeks which was later continued at home. They were prospectively followed for 24±12 months. Primary endpoints were time to clinical worsening and survival. Adverse events and changes in the 6-min walking test, QoL, WHO functional class and gas exchange were secondary endpoints and were evaluated at baseline and at weeks 3 and 15. The results showed that all patients tolerated the exercise training well without any severe adverse events. In week 15, 6-min walking test results were significantly improved compared to baseline as well as quality of life scores, WHO functional class, peak oxygen consumption, heart rate at rest and maximal workload. Survival at 1 and 2 yr was 100 and 95%, respectively (Grunig et al., 2011).

Fifteen events occurred during the follow-up. It was concluded that exercise and respiratory training as add-on to medical treatment may improve exercise capacity and QoL, with a good long term safety profile.

**NON PHYSICAL EFFECTS OF EXERCISE AND PULMONARY REHABILITATION**

As pulmonary hypertension’s most debilitating manifestation is exercise intolerance most trials focus on improvement in this characteristic as well as hemodynamics. However, in addition to physical improvement, there is a great deal of improvement in aspects of psychosomatic domains. It is well known that patients with pulmonary hypertension experience depression (Verma et al., 2014) and report increased levels of fatigue severity (Talwar et al., 2014). It is also well known that these patients have a severely impaired health related quality of life (Taichman et al., 2005). These aspects of psychosomatic medicine may also be addressed with the use of structured pulmonary rehabilitation and exercise programs in this population. A study by Kim et al. (2013) showed the effects that dyspnea has on depression and health related quality of life (HRQoL). Pulmonary rehabilitation programs have also showed improvement in fatigue severity as well as depression (Talwar et al.,...
A more recent study by Chan et al. (2013) went ahead to determine the effects of intense treadmill exercise on cardio-respiratory function as well as quality of life in patients with pulmonary hypertension. Patients were randomized to a 10-week disease-specific education only group or education/exercise combined group. The exercise program consisted of 24-30 sessions of treadmill walking for 30-45 min per session. Outcome variables included changes in 6-min walk test (6 MWT) distance, time to exercise intolerance, peak work rate from a cardiopulmonary treadmill test, and quality-of-life measures, including the SF-36 and Cambridge Pulmonary Hypertension Outcome Review (CAMPHOR) (Chan et al., 2013). The results following 10 weeks of intervention, demonstrated an improvement in 6 MWT distance, increased time to exercise intolerance, and peak WR. Additionally, the exercise group showed increased quality of life score according to the SF-36 as well as the CAMPHOR scale (Chan et al., 2013).

CONCLUSIONS

Pharmacotherapy for pulmonary arterial hypertension has advanced over the past decade and has provided clinicians with a disease specific approach which has led to the improvement in prognosis of PAH patients. However, patients continue to experience significant exertional symptoms accompanied by reduced functional capacity and diminished health-related quality of life. From the trials that have been conducted, the evidence supports the benefits and safety of exercise training and rehabilitation programs in the pulmonary hypertension populations. Initial exercise training and pulmonary rehabilitation programs were assessed in a stable, treated, systolic heart failure, and COPD and showed improved functional capacity and reduced exercise intolerance, symptoms of dyspnea, and fatigue.

Trials conducted in a pulmonary hypertension population showed improvement in exercise endurance and muscle strength as well as quality of life with the implementation of exercise training and cardiopulmonary rehabilitation as part of their medical care without a good safety profile. In conjunction with pharmacotherapy all patients with confirmed pulmonary arterial hypertension should be treated with a pulmonary rehabilitation and exercise training program.

CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

REFERENCES


