Exercise Training Improved Pulmonary Gas Exchange Abnormalities in Pulmonary Hypertension due to Heart Failure: A Case Report

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Abstract

Heart failure (HF) is the most common cause of pulmonary hypertension (PH), and reduced exercise capacity and exertional dyspnea are the most frequent concerns in patients with PH-HF. Indeed, carbon dioxide end-tidal partial pressure (PETCO₂) during exercise is a well-established noninvasive marker of ventilation/perfusion ratio in PH. We aimed to evaluate the effect of aerobic exercise training on PETCO₂ response during exercise in a 59-year-old woman with PH secondary to idiopathic dilated cardiomyopathy. The patient with chronic fatigue and dyspnea at mild-to-moderate efforts was admitted to a cardiorespiratory rehabilitation program and had her cardiorespiratory response to exercise assessed during a cardiopulmonary exercise testing performed before and after three months of a thrice-weekly aerobic exercise training program. Improvements in aerobic capacity (23.9%) and endurance time (37.5%) and reduction in ventilatory inefficiency (-20.2%) was found after intervention. Post-intervention improvements in PETCO₂ at ventilatory anaerobic threshold (23.3%) and change in PETCO₂ kinetics pattern, with progressive increases from rest to peak of exercise, were also found. Patient also improved breathing pattern and timing of ventilation. This case report demonstrated for the first time that aerobic exercise training might be able to improve PETCO₂ response during exercise in a patient with PH-HF.

Introduction

Heart failure (HF) is the most common cause of pulmonary hypertension (PH) and is associated with increased morbidity and mortality. Exertional dyspnea and reduced exercise capacity are the most frequent concerns in patients with PH due to HF (PH-HF). The key physiological mechanism in PH-related exertional dyspnea is a lung vasculature abnormality due to excessive pulmonary vascular resistance, which attenuates cardiac output response to exercise. Noteworthy, patients with HF often display ventilatory abnormalities likely related to dysregulation in peripheral control (i.e., impaired skeletal muscle ergoreflex and increased peripheral chemoreceptor sensitivity).

Cardiopulmonary exercise testing (CPX) is an useful tool for assessing severity and prognosis in cardiorespiratory disease. Carbon dioxide end-tidal partial pressure (PETCO₂) during CPX is a well-established noninvasive marker of ventilation/perfusion ratio in patients with lung and heart diseases. Indeed, the PETCO₂ decrease from rest to ventilatory anaerobic threshold (VAT) during CPX is associated with pulmonary gas exchange abnormalities in patients with PH.

Keywords

Aerobic Capacity; Exercise Test; Cardiovascular Diseases; Pulmonary Gas Exchange; High Frequency Ventilation.

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Therapeutic strategies are available for treating primary PH; however, the same does not occur for PH-HF, whereas target therapy agents have failed to demonstrate benefit or have even been harmful. Although exercise training is a well-established non-pharmacological therapy for HF, its effect on patients with PH-HF, particularly on PETCO₂ response during exercise, is unknown. Therefore, our aim was to report the effect of aerobic exercise training on PETCO₂ response during exercise in a patient with PH-HF.

Case Report

Patient’s characteristic and evaluation

A 59-year-old sedentary female (weight: 86 kg, height: 162 cm) with PH secondary to idiopathic dilated cardiomyopathy was admitted for cardiorespiratory rehabilitation in the TotalCare clinic, São Paulo, Brazil. Her comorbidities included systemic hypertension, diabetes, and dyslipidemia. No lower limb edema or increased jugular venous pressure, and resting blood pressure and heart rate of 140/100 mmHg and 60 bpm, respectively, were found during physical examination. Patient had regular heart rhythm with mitral regurgitation and clear breath sounds during auscultation. Her symptoms were chronic fatigue and dyspnea at mild-to-moderate efforts. She had never participated in any exercise program and had no history of hospitalization associated with HF.

Normal sinus rhythm and left branch bundle block were showed during resting electrocardiography. Global left ventricular (LV) hypokinesia with mildly impaired overall systolic function (ejection fraction: 37%), LV dilatation, and moderate PH (PASP: 55 mmHg) were showed during echocardiogram. Echocardiography diagnosis of PH was obtained through right ventricular systolic pressure (RVSP). The echocardiographic assessment was evaluated by the maximal tricuspid regurgitation velocity and the systolic gradient between the right ventricle and the right atrium was calculated.

Baseline medications included carvedilol, hydrochlorothiazide, spironolactone, amlodipine, losartan, acetylsalicylic acid, and simvastatin, and no changes occurred during the previous three month before and throughout the study. The study was approved by the local Ethics Committee (CAEE :26442619.9.0005533) and patient signed an informed consent form.

A symptom-limited maximal CPX was carried out on treadmill (Centurion model 300; Micromed, Brazil) using a Balke modified protocol at controlled room temperature (20-23°C), 2 and 3 days before beginning and after ending the exercise program, respectively. Gas exchange and ventilatory variables were measured breath-by-breath throughout the test by a computerized system (Cortex model III B; Leipzig, Germany) and expressed as 30 sec averages, as previously described. Heart rate was continuously recorded by 12-lead ECG (Elite Ergo PC; Micromed, Brazil) during resting, exercise, and recovery phases of CPX. VAT, peak oxygen consumption (VO₂peak), oxygen uptake efficiency slope (OUES), ventilatory equivalent of CO₂ production (Ve/ VCO₂) slope, noninvasive estimate for physiologic dead space to tidal volume ratio (Ve/Vₜ), PETCO₂ pattern, and ventilatory pattern analysis (i.e., breathing frequency (BF), tidal volume (Vₜ), expiratory time (Tₑ), inspiratory time (Tᵢ), total respiratory time (TOT), mean inspiratory flow (Vᵢ/Tᵢ), and mean expiratory flow (Vₑ/Tₑ)) were assessed during CPX. As previously described, VAT was considered the point at which ventilatory equivalent of O₂ (VE/VO₂) reached the minimum value and began to rise without a concomitant rise in VE/VCO₂. VO₂peak was defined as the maximum VO₂ attained at the end of the exercise period, at which time the patient had reached his/her maximum level of exhaustion (Borg’s rating of perceived exertion scale).

Exercise Training Program

A thrice-weekly supervised exercise training program were performed for three months at the TotalCare clinic’s cardiorespiratory rehabilitation center. Exercise sessions included warm-up (5 min), aerobic (50 min), and cool-down (5 min) exercises. Aerobic exercise was a treadmill walking at the heart rate corresponding to the VAT during CPX (± 5 bpm). Heart rate was monitored throughout the session to ensure that patient exercised within the limits of intensity. The participant attended 90% of the programmed exercise sessions (33 sessions) and did not perform any exercise at home or any other exercise modality.

Outcomes

Severe impairment in cardiorespiratory fitness (< 50% of age predicted peak VO₂), reduced gas exchange efficiency, and abnormal PETCO₂ pattern with progressive decreases were observed at baseline (Table 1 and Figure 1A).
Figure 1 – Ventilatory response to cardiopulmonary exercise testing before (pre) and after (post) 3 months of aerobic exercise training. 

A: Carbon dioxide end-tidal partial pressure (PETCO2), B: Carbon dioxide end-tidal partial pressure change (ΔPETCO2) from rest to ventilatory anaerobic threshold (VAT), C: breath frequency (BF), D: tidal volume (VT), E: physiologic dead space to tidal volume ratio (VD/VT), F: total respiratory time (TOT), G: expiratory time (TE), H: inspiratory time (TI), I: mean inspiratory flow (VT/TI), J: mean expiratory flow (VT/TE).
After three month of aerobic exercise training, patient showed an increase in body mass (1 kg), peak VO$_2$ (23.9%), peak O$_2$ pulse (24.1%), and endurance time (33.3%), as well as in VO$_2$ at VAT (38.1%) and OUES (14.5%) (Table 1). Patient also showed lower V$_{\text{T}}$/VCO$_2$ slope levels (Table 1). PETCO$_2$ analysis during CPX showed increased PETCO$_2$ at VAT (23.3%) and a change in PETCO$_2$ kinetics pattern with progressive increases from rest to peak of exercise when compared to pre-intervention (Figure 1A and 1B). Patient also improved breathing pattern (Figure 1C and 1D) and timing of ventilation (Figure 1G, 1H, 1I, and 1J).

**Discussion**

Progressive increase in PETCO$_2$ from resting to VAT (nearly 5-8 mmHg), followed by its maintenance or slight increase up to the onset of respiratory compensation point and its progressive decrease from respiratory compensation point to maximal effort, is expected in healthy subjects during CPX. In contrast, our data showed progressive decreases in PETCO$_2$ from resting to VAT and maximal effort at baseline (Figures 1A and 1B), which is in accordance with a previous study assessing the PETCO$_2$ response in patients with primary PH. Indeed, the abnormal PETCO$_2$ response was directly associated with disease severity.

Physiological mechanisms involved in the altered PETCO$_2$ pattern found at baseline may include: (1) ventilation-perfusion inequalities due to ventilated alveoli hypoperfusion, resulting in increased physiologic dead space (V$_{\text{T}}$/V$_{\text{E}}$) and (2) increased acidosis at lower work rates, resulting in augmentation of ventilation drive induced by higher levels of hydrogen ion (H$^+$). For instance, the increased pulmonary vascular resistance in patients with PH may blunt cardiac output response during exercise, which results in reduced O$_2$ transport to working muscle, thereby increasing the contribution of anaerobic glycolysis to exercise. Indeed, patients with HF demonstrate impaired skeletal muscle metabolism, which reduces mitochondrial oxidative capacity. Thus, it is possible that both pulmonary gas exchange and muscle metabolism abnormalities may be involved in the impaired baseline PETCO$_2$ response during exercise in the patient with PH-HF.

In contrast, to the best of our knowledge, the present report showed for the first time that three months of aerobic exercise training might improve PETCO$_2$ response to exercise in a patient with PH-HF, as shown by the sharp increase in PETCO$_2$ from rest to VAT (Figure 1B), as well as by the increased PETCO$_2$ levels at both VAT and peak of exercise (Table 1). This result is in line with a previous study demonstrating the effectiveness of aerobic exercise training for improving PETCO$_2$ response in coronary artery disease patients.

It is known that exercise training improves muscle oxidative metabolism during exercise in individuals with HF (e.g., increased O$_2$ uptake and arteriovenous O$_2$ difference, lower phosphocreatine depletion, lower increase in adenosine diphosphate, and decreased lactate accumulation). In addition, training-induced improvements in ventilation efficiency during exercise are, at least in part, due to improvements in muscle receptor reflexes and skeletal muscle metabolism in individuals with HF. In this context, it can be suggested that a training-induced improvement in muscle aerobic metabolism may be reduced acidosis-related ventilatory stimulus during CPX, and thus may be associated with the improved PETCO$_2$ response after training in the present study. The improvements in both VO$_2$ at VAT and OUES suggest an increase in muscle aerobic capacity and thus support this hypothesis.

Improvements in gas exchange efficiency and breathing pattern may be induced by the reduction of physiologic dead space ventilation and may also be involved in the improved PETCO$_2$ response after training. In this context, the reduced V$_{\text{T}}$/VCO$_2$ slope, V$_{\text{T}}$/V$_{\text{E}}$ and tachypneic (BF) and shallow pattern of breathing (T$_{\text{E}}$/T$_{\text{I}}$) after training support this hypothesis. In fact, the reduced V$_{\text{T}}$/T$_{\text{I}}$ and V$_{\text{T}}$/T$_{\text{E}}$ after training indicate a decrease in neuromuscular inspiratory and expiratory drive, respectively. It is important to note that breathing pattern has a significant influence in the V$_{\text{T}}$/V$_{\text{E}}$ ratio during exercise. Specifically, a tachypneic breathing pattern during exercise increases V$_{\text{T}}$/V$_{\text{E}}$ ratio and decreases PETCO$_2$ levels. Moreover, higher levels of breathing frequency are closely related to decreases in expiratory time and, as a consequence, in premature cessation and greater PETCO$_2$ increase during progressive exercise. Thus, the decrease in breathing frequency and the increase in expiratory time found after exercise training may also be associated with the improved PETCO$_2$ response during follow-up in the present study. Interestingly, peripheral chemoreflex control was normalized after exercise training in a rabbit model of heart failure, suggesting that it may be associated with the improvement in ventilation efficiency.

It is noteworthy that the patient also reported an improvement in exertional dyspnea sense after training,
which is in line with the improved exercise capacity, as shown by the training-induced increase in exercise time during CPX. In conjunction, these improvements may result in an increased quality of life.

The echocardiography used for assessing PH is a limitation of the present study. However, we used sequential assessments to confirm the diagnosis. The assessment of only one patient during the follow-up is also a limitation that should be acknowledged.

In summary, this case report showed for the first time that the impaired PETCO₂ response to exercise in PH-HF might be improved by three months of a thrice-weekly aerobic exercise training program. Given that PETCO₂ is a noninvasive marker of ventilation/perfusion ratio and that its abnormal response during CPX is associated with pulmonary gas exchange abnormalities in patients with PH, the present results suggest that aerobic exercise training may be an important tool for improving pulmonary gas exchange abnormalities in patients with PH-HF.

**Author Contributions**

Conception and design of the research: do Prado DM, Fonseca de Campos JP, Miranda TP, Teixeira AB, Staroste M. Acquisition of data: do Prado DM, Fonseca de Campos JP, Miranda TP, Teixeira AB, Staroste M. Analysis and interpretation of the data: do Prado DM, Rocco EA, Ciolac EG. Writing of the manuscript: do Prado DM, Ciolac EG. Critical revision of the manuscript for intellectual content: do Prado DM, Rocco EA, Ciolac EG.

**Potential Conflict of Interest**

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

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**Study Association**

This study is not associated with any thesis or dissertation work.

**Ethics Approval and Consent to Participate**

This study was approved by the Ethics Committee of the Hospital Samaritano under the protocol number 26442619.9.0005533. All the procedures in this study were in accordance with the 1975 Helsinki Declaration, updated in 2013. Informed consent was obtained from all participants included in the study.

**References**