The contribution of inspiratory muscles function to exercise limitation in heart failure: pathophysiological mechanisms

Contribuição da musculatura inspiratória na limitação ao exercício na insuficiência cardíaca: mecanismos fisiopatológicos

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Abstract

Background: Heart failure induces histological, metabolic and functional adaptations in the inspiratory muscles. This inspiratory muscle weakness, which occurs in 30% to 50% of the heart failure patients, is associated with reduction in the functional capacity, reduction in the quality of life and with a poor prognosis in these individuals. Objectives: The objective of this review was to discuss the pathophysiological mechanisms that may explain the role of the inspiratory muscles in the exercise limitation with focus in the reflexes that control the ventilation and the circulation during the exercise. Method: We performed searches in the PUBMED database using the terms "inspiratory muscles", "inspiratory muscle training", "metaboreflex" and "chemoreflex" and including studies published since 1980. Results: Inspiratory muscle weakness is associated with exercise intolerance and with an exaggerated inspiratory chemoreflex and metaboreflex in heart failure. The inspiratory metaboreflex may be attenuated by the inspiratory muscle training or by the aerobic exercise training improving the exercise performance. Conclusions: Patients with heart failure may present changes in the inspiratory muscle function associated with inspiratory chemoreflex and metaboreflex hyperactivity, which exacerbate the exercise intolerance.

Keywords: inspiratory muscle training; rehabilitation; chemoreflex; metaboreflex; heart failure.

Resumo

Contextualização: A insuficiência cardíaca (IC) acarreta alterações histológicas, metabólicas e funcionais dos músculos inspiratórios. A fraqueza dos músculos inspiratórios, que ocorre em 30% a 50% dos pacientes com IC, associa-se com a redução da capacidade funcional, prejuízos para a qualidade de vida e piora no prognóstico desses indivíduos. Objetivos: Discutir os mecanismos fisiopatológicos que potencialmente explicam o papel da musculatura inspiratória na limitação ao exercício, abordando-se os reflexos que controlam a ventilação e a circulação durante o exercício. Método: Foram realizadas pesquisas na base de dados PUBMED, utilizando os termos inspiratory muscles, inspiratory muscle training, metaboreflex e chemoreflex e incluindo estudos publicados desde 1980. Resultados: A fraqueza muscular inspiratória está relacionada com intolerância ao exercício e com exacerbação do quimiorreflexo e metaboreflexo na IC. O metaboreflexo inspiratório pode ser atenuado pelo treinamento muscular inspiratório ou pelo treinamento aeróbico, melhorando o desempenho ao exercício. Conclusões: Os pacientes com IC podem apresentar alterações da função muscular inspiratória associadas com hiperatividade quimiorreceptor e metaboreceptor inspiratória, as quais podem agravar a intolerância ao exercício.

Palavras-chave: treinamento inspiratório; reabilitação; quimiorreflexo; metaboreflexo; insuficiência cardíaca.

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Introduction

It has been proposed for several years that impairments in the peripheral skeletal muscle function contribute to the pathophysiology of heart failure (HF). During this period, it has been shown that skeletal muscle abnormalities limit the aerobic capacity during the exercise, which results in metabolites accumulation in the muscles leading to sensitize muscle receptors called metaboreceptors. The metaboreceptors activation induces a reflex response, which results in hyperventilation exacerbating the dyspnea sensation. Moreover, it pronoves peripheral vasconstriction with reduction of blood flow which contributes for the reduction of the tolerance to the exercise on HF.

More recently, studies have shown that inspiratory muscles can also limit physical performance to exercise in healthy subjects and in athletes. Similarly, changes in the inspiratory muscles play an important role in the pathophysiology of exercise limitation in HF. One of these changes is the inspiratory muscle weakness, arbitrarily defined as a maximal inspiratory pressure (PI) less than 70% of the predicted for age and sex. Patient with inspiratory muscle weakness may have a reduced functional capacity determined by the peak oxygen consumption (VO2peak). In addition, inspiratory muscle strength and resistance are directly associated to VO2peak in patients with recent acute myocardial infarction, reinforcing the idea that the inspiratory muscles may contribute for the reduction of functional capacity. A recent study shows that inspiratory muscle strength is also associated with the central hemodynamic measurements at rest, including cardiac output, mean arterial pulmonary pressure and pulmonary vascular resistance in patients with HF. However, no correlation between inspiratory muscle strength and left ventricular ejection fraction was found. Finally, inspiratory muscle strength, assessed by PImax, has an independent impact on the prognosis of patients with HF.

Functional changes of inspiratory muscles may contribute to the activation of cardiovascular reflexes, which aggravate the exercise limitation in HF. Among these reflexes, we highlight the exacerbation of metaboreflex induced by the work of the inspiratory muscles and the increase of the ventilatory responses mediated by the chemoreflex during the exercise. In the present review, we will discuss the role of the inspiratory chemoreflex and metaboreflex in the exercise limitation in HF.

Chemoreflex

The appropriate tissue oxygen supply is maintained by adjustments of the circulation and the ventilation mediated by the complex interaction of multiple systems, with emphasis on the arterial baroreflex, central and peripheral chemoreflex, ergoreflex and pulmonary stretch reflex. Chemoreflexes are the main mechanisms of control and management of the ventilatory responses to changes in the arterial oxygen and carbon dioxide concentrations. The central chemoreceptors, located on the ventral surface of medulla (bulb), respond primarily to the variations of partial pressure of CO2 (PCO2) in the arterial blood. The peripheral chemoreceptors, located in the common carotid artery and in the aorta artery with afferent input to the respiratory center located at the bulb and at the nucleus of the solitary tract, respond primarily to the variations of partial pressure of O2 (PO2) in the arterial blood. Thus, both variations in PO2 and PCO2 result in changes in pulmonary ventilation. In addition, the activation of the central and peripheral chemoreflexes increases the sympathetic nervous activity, which results in the increase of heart rate and blood pressure. In contrast, the increase of the ventilation sensitizes pulmonary stretch receptors located in the bronchi, bronchioles, and lungs, which detect excessive stretching of the lungs, sending information to the dorsal group of respiratory neurons resulting in negative feedback, preventing therefore additional gains of inspiratory volume. Similarly, the increase in blood pressure induced by the chemoreflex can be counterbalanced by the activation of the arterial baroreflex. The elevation of the arterial pressure distends the baroreceptors that transmit afferent signals to the central nervous system, resulting in negative feedback signals to the circulation. Thus, there is an increase of the efferent vagal activity and a reduction of the sympathetic activity, reducing the arterial pressure. Therefore, the reflex control of the circulation and ventilation requires the integration among several systems of control, which are represented in Figure 1.

The chemoreflex sensitivity is abnormally increased in patients with HF, resulting in an exaggerated increase in pulmonary ventilation and muscle sympathetic nervous activity. This exaggerated increase in chemoreflex sensitivity is associated with the reduced negative feedback system of the arterial baroreflex. This demonstrates a impairment in the ventilation and circulation adjustment systems that contribute to the pathophysiology of the HF, since the increase in sympathetic activation results in an adrenergic vasoconstriction and an increase of the left and right ventricular afterload, being the sympathetic hyperactivation an important predictor of mortality in HF. In addition, the exacerbation of the chemoreflex is related to reductions in functional capacity. Therefore, patients with HF and exacerbated chemoreflex might present a greater limitation of exercise performance.
The ventilatory responses to the exercise can be modulated by the chemoreceptors. Studies suggest that both peripheral and central chemoreflexes contribute to the occurrence of hyperventilation during exercise in HF. Indeed, patients with abnominal elevation of the ventilation/carbon dioxide production slope (V̇E/V̇CO2) (>34) present increased peripheral and central chemoreflex sensitivity41. However, the inhibition of the peripheral chemoreflex, via dihydrocodeine, can reduce the V̇E/V̇CO2 slope during progressive exercise42. Hyperventilation during exercise can also be directly mediated by skeletal metaboreflex or the central chemoreflex could be indirectly induced43, since the metaboreflex is a strong predictor of central chemoreflex activity in patients with HF42. It is important to note that both central chemoreflex and ventilatory response induced by muscle skeletal metaboreflex correlated significantly with V̇E/V̇CO2 slope during progressive exercise in HF43.

Another changed ventilatory response to exercise, known as oscillatory ventilation or periodic breathing14,22, can be related to abnormal responses of chemoreflex. A study conducted in patients with HF showed that the peripheral chemoreflex sensitivity contributes in the genesis of the oscillatory ventilation at rest14, since the inhibition of the peripheral chemoreflex normalizes the ventilation in these patients. The inspiratory muscular training may also attenuate the oscillatory ventilation during exercise in patients with HF and inspiratory muscle weakness. In fact, the inspiratory muscle training at 30% PI max reduces the oscillations of ventilation at 57%, concomitant to the increase of 115% of the inspiratory muscle strength14,47. Taking into consideration that oscillatory ventilation may be originated by peripheral chemoreflex, we tested the hypothesis that the chemoreflex is exacerbated in patients with HF and inspiratory muscle weakness. This study demonstrated an inverse association between the peripheral chemoreflex and PI max (r=-0.57, p=0.01), i.e., the inspiratory muscle weakness is related to the exacerbation of the peripheral chemoreflex as shown in Figure 2. Indeed, the group with inspiratory muscle weakness presented a greater peripheral chemoreflex compared with patients with the inspiratory muscle strength preserved24. Thus, the reduction of the inspiratory muscle strength can aggravate the exacerbation of chemoreflex sensitivity in patients with HF, which could potentially be related to reductions in functional capacity and to exercise intolerance36,41.

**Inspiratory muscle metaboreflex in healthy subjects**

The activation of the metaboreflex by the contraction of the skeletal muscles is well-known. However, only in the last decade it was described that the metaboreflex could also be activated by the respiratory muscles working5-7,46. Initially,
it was demonstrated that physical exercise (intensity > 85% of the maximal oxygen consumption \(\dot{V}O_2\text{max}\)) induces diaphragmatic muscle fatigue\(^9\) even in elite athletes\(^{10}\). Then, it was observed that the increase in the inspiratory muscle work (via inspiratory resistance) increases the levels of noradrenaline reducing the blood flow in the leg during maximal exercise on a bicycle\(^{11}\). These findings led to the hypothesis of the existence of an “inspiratory metaboreflex” activated during physical exercise. A theory confirmed in a subsequent study verified the existence of redistribution of blood flow of the active peripheral muscles to the diaphragm, corresponding to more than 14-16% of cardiac output\(^{12}\). Moreover, in an experimental protocol of induction of the inspiratory metaboreflex, through the intense inspiratory effort (inspiratory resistance = 60% of PI\(_{\text{max}}\)) and sustained (ratio between inspiratory time and the total duration of the respiratory cycle [TI/TTot] = 0.70), showed that inspiratory muscle fatigue increases muscle sympathetic nerve activity\(^{13}\) and reduces the muscle blood flow for the inactive leg\(^{14}\) due to the adrenergic vasoconstriction. This response seems to be mediated by the metabolic stimulation of small afferent fibers types III and IV from the respiratory muscles, especially from the diaphragm\(^{15}\). The effects of redistribution of the blood flow during the fatiguing inspiratory work showed the existence of an inspiratory metaboreflex, represented in Figure 3.\(^{16}\) Thus, the activation of the inspiratory metaboreflex during physical exercise that induces inspiratory muscle fatigue\(^{17}\) may limit physical performance\(^{12,55}\) due to the reduction of blood flow to the active skeletal muscles\(^{18}\), exacerbating the fatigue of the peripheral muscles\(^{8}\). On the other hand the reduction of inspiratory muscle work through mechanical ventilation in healthy subjects increases the exercise duration in 14%\(^{6}\) and attenuates quadriceps fatigue during exercise\(^{8}\), probably because it inhibits the inspiratory metaboreflex. Moreover, the conditioning of the inspiratory muscles could minimize the effects of the activation of the inspiratory metaboreflex. Studies show that inspiratory muscle training, performed at 50% of PI\(_{\text{max}}\) attenuates the pressure response\(^{54}\) and reduces the fatigue of the plantar flexors muscles during the activation of the inspiratory metaboreflex in healthy subjects\(^8\). These effects of inspiratory muscle training could contribute to improve the exercise performance. In fact, inspiratory muscle training prolongs the duration of the exercise in bicycle, and also reduces the lactate concentration at the end of progressive exercise test in active subjects\(^{55}\). Additionally, regular aerobic training also attenuates inspiratory metaboreflex. A recent study showed that the inspiratory metaboreflex is attenuated in trained subjects when compared with sedentary healthy subjects\(^{56}\). In fact, fatiguing inspiratory muscle work induced peripheral vasoconstriction with reduced blood flow in the calf of sedentary subjects, but remain almost unchanged in aerobically trained subjects (Figure 4)\(^{56}\). This effect of aerobic training on the inspiratory metaboreflex is probably related to the increased oxidative capacity\(^{57}\), the strength of inspiratory muscles\(^{58}\) and of the fatigue resistance of the diaphragm\(^{59}\). Alternatively, the administration of transcutaneous electric nervous stimulation (TENS) in the cervicothoracic region attenuates the muscle skeletal metaboreflex, probably due to the blockage of the sympathetic nervous activity in young and old subjects, as recently demonstrated\(^{60}\). Thus, the application of TENS in the cervicothoracic region could possibly attenuate the inspiratory metaboreflex impact on physical performance, but this hypothesis still need to be further investigated.

**Figure 3.** Illustration of the inspiratory metaboreflex activated during the fatiguing inspiratory muscle work due to metabolite accumulation that increase afferent phrenic discharge resulting in the increase of the sympathetic activity and peripheral vasoconstriction, exacerbating the fatigue of the active peripheral muscles.

**Inspiratory muscle metaboreflex in subjects with heart failure.**

The inspiratory metaboreflex may limit the exercise performance mainly in HF patients with inspiratory muscle weakness\(^4\). In fact, the weakness of inspiratory muscle often observed in patients with HF\(^{16,61-63}\) may be related to exercise intolerance. In addition, the reduction in inspiratory muscle oxygenation mediated by acute physical exercise in patients with HF\(^{64}\) could increase the ventilatory work and exacerbate the inspiratory muscle metaboreflex limiting the exercise tolerance due to the reduction of the perfusion of the locomotor muscles, as have been previously mentioned.
Thus, strategies that reduce the inspiratory muscle work and attenuate the inspiratory metaboreflex could improve the exercise tolerance. In fact, the reduction of inspiratory muscle work through noninvasive ventilation may improve physical performance in patients with HF. Recently, it has been shown that such intervention improves the oxygenation of the peripheral muscle microcirculation, which may be compatible with the improvement of local blood flow\(^6\). These findings have been currently confirmed by Olson et al.\(^6\), whom found that the reduction in inspiratory muscle work increased the leg blood flow and the cardiac output only in patients with HF, remaining unchanged in healthy controls subjects. It means that patients with HF are more susceptible to exercise intolerance due to the effects of the metaboreflex activation by the inspiratory muscles work, so that the inspiratory muscle training attenuates the inspiratory metaboreflex in patients with HF and with the inspiratory muscles weakness (Figure 5)\(^15\). These effects of inspiratory muscle training may contribute to the improvement of the cardiorespiratory responses, functional capacity and tolerance to the exercise in patients with HF\(^14\). A recent study confirmed the findings of improvement in the exercise tolerance after inspiratory muscle training\(^6\), being the improvement on functional capacity occurs even in HF patients with left ventricle assist device\(^6\). Therefore, the conditioning of inspiratory muscles via inspiratory muscle training could attenuate the inspiratory metaboreflex with possible benefits for the improvement of the exercise tolerance in patients with HF. Thus, interventions that attenuate the activation of the inspiratory metaboreflex may be beneficial for patients that present exercise intolerance, including those with chronic obstructive pulmonary disease. In animal models of HF, it has also been demonstrate that inspiratory muscle training improves hemodynamic function, respiratory mechanics and cardiovascular autonomic control\(^6\).

Some studies suggest that noninvasive ventilation could temporarily attenuate the activation of the inspiratory metaboreflex due to the reduction of the inspiratory muscle work in patients with chronic obstructive pulmonary disease. In the study of Borghi-Silva et al.\(^7\), the bi-level positive airway pressure ventilation improved the \(O_2\) peripheral saturation and reduced quadriceps muscle fatigue during the isokinetic exercise. These data had been supported by a study that demonstrated that the reduction of the inspiratory muscle work, via inhalation of mixture containing helium at 79\%, counterbalanced with oxygen, improved the exercise tolerance in patients with chronic obstructive pulmonary disease that performed an exercise protocol with constant load and high intensity\(^7\). Furthermore, the use of positive pressure support increases the resistance to exercise in these patients\(^7\). Thus, non-invasive ventilatory support could help maintain the physical training of patients that present severe exercise limitations related to chronic obstructive pulmonary disease, probably due to attenuation of the inspiratory metaboreflex\(^7\).

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**Figure 4.** Leg blood flow response, assessed by Doppler ultrasonography, during the induction of the inspiratory metaboreflex at 60% \(P_{Imax}\). Endurance trained subjects are represented by close circles and sedentary subjects are represented by open circles.

**Figure 5.** Leg vascular resistance measured via venous occlusion plethysmography during the induction of the inspiratory metaboreflex at 60% \(P_{Imax}\). The values of the leg vascular resistance pre inspiratory muscle training are represented by close circles. The values of the leg vascular resistance post inspiratory muscle training are represented by open circles.
Implications for the physical therapist practice

The inspiratory muscle training improves functional capacity and quality of life in patients with HF and inspiratory muscle weakness. These findings were confirmed in a recent study that investigated the effects of inspiratory muscle training in HF and found, in their sample, a vast majority of patients with inspiratory muscle weakness.

In addition, the association of inspiratory muscle training with aerobic training promotes additional effects on VO_{2peak} in patients with inspiratory muscle weakness. Randomized controlled clinical trials with a representative sample size could investigate isolated effects, and compared effects between aerobic training and inspiratory muscle training in patients with inspiratory muscle weakness.

Therefore, it is recommended that the physical therapist routinely evaluate inspiratory muscle strength in patients with HF in order to identify patients with muscle weakness, which could benefit from inspiratory muscular training.

Conclusion

Patients with HF may present changes in inspiratory muscle function associated with chemoreflex and inspiratory metaboreflex hyperactivity, which can exacerbate the exercise intolerance.

References