ABSTRACT

Introduction: The structural and mechanical adaptations of the right ventricular (RV) myocytes in response to hypertension associated with low-intensity endurance training (LIET) have not been studied in experimental models. Objective: To determine the effects of LIET on the structural and mechanical properties of RV myocytes in spontaneously hypertensive rats (SHR). Methods: Male SHR and normotensive Wistar rats (age: 16 weeks) were allocated to groups (n=7): WIS (Wistar Controls); SHR-C (SHR Controls) and SHR-T (SHR Trained; 60 min/day, 50-60% of maximal exercise capacity, 5 days/week for 8 weeks). Systolic arterial pressure (SAP), isolated RV myocyte dimensions, contractility, intracellular Ca\(^{2+}\) transient ([Ca\(^{2+}\)]\(_t\)), and ventricular Ca\(^{2+}\) regulatory proteins were measured. The statistical analysis was performed by one-way ANOVA followed by the Tukey post hoc test (α=5%). Results: LIET reduced the SAP in SHR animals (SHR-C, 164 ± 2 mmHg vs. SHR-T, 152 ± 4 mmHg; P<0.05); Hypertension increased cell length (WIS, 156.8 ± 2.7 µm; SHR-C, 166 ± 3.1 µm; P<0.05) but did not affect cell width or volume (P>0.05). LIET did not change the cell dimensions in the SHR-T. Neither hypertension nor LIET affected myocyte contractility or the expression of Ca\(^{2+}\) regulatory proteins in the RV of the SHR-C and SHR-T groups. Hypertension did not affect the amplitude of the [Ca\(^{2+}\)]\(_t\) transient or the time to half resting level (P>0.05), but increased the time to peak (WIS, 58 ± 1 ms vs. SHR-C, 79 ± 2 ms; P<0.05). LIET increased the amplitude of the [Ca\(^{2+}\)]\(_t\) transient (WIS, 2.28 ± 0.07 F/F\(_0\) vs. SHR-C, 2.48 ± 0.08 F/F\(_0\); vs. SHR-T, 2.87 ± 0.08 F/F\(_0\); P<0.05), but did not alter the times to peak or to half resting level. Conclusion: LIET had no effect on the structural and mechanical properties of RV myocytes in the SHRs, although it increased the amplitude of the [Ca\(^{2+}\)]\(_t\) transient and reduced the SAP.

Keywords: Exercise; Rats; Hypertension; Heart.

RESUMO

Introdução: As adaptações estruturais e mecânicas de miócitos do ventrículo direito (VD) em resposta à hipertensão associada ao treinamento aeróbio de baixa intensidade (TABI) não foram estudadas em modelos experimentais de rato. Objetivo: Determinar os efeitos do TABI sobre as propriedades estruturais e mecânicas de miócitos do VD em ratos espontaneamente hipertensos (SHR). Métodos: Ratios SHR e ratos Wistar machos e normotensos (idade: 16 semanas) foram distribuídos em grupos (n = 7): WIS (Wistar controle); SHR-C (SHR controle) e SHR-T (SHR treinados; 60 min/dia, 50-60% da capacidade máxima de exercício, 5 dias/semana por 8 semanas). Procedeu-se à medição de pressão arterial sistólica (PAS), dimensões dos miócitos isolados do VD, contratilidade, transiente intracelular de Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_t\)) e de proteínas reguladoras de Ca\(^{2+}\). A análise estatística foi realizada por ANOVA one-way, seguida por teste de Tukey post hoc (α = 5%). Resultados: O TABI reduziu a PAS nos animais SHR (SHR-C, 164 ± 2 mmHg vs. SHR-T, 152 ± 4 mmHg; P < 0,05). A hipertensão aumentou o comprimento celular (WIS, 156,8 ± 2,7 µm; SHR-C, 166,6 ± 3,1 µm; P < 0,05), mas não afetou a largura ou o volume (P > 0,05), O TABI não alterou as dimensões celulares nos SHR-T. Nem a hipertensão nem o TABI afetaram a contratilidade dos miócitos ou a expressão das proteínas reguladoras do Ca\(^{2+}\) no VD dos grupos SHR-C e SHR-T. A hipertensão não afetou a amplitude do transiente de [Ca\(^{2+}\)], e o tempo até a metade do nível de repouso (P > 0,05), mas aumentou o tempo até o pico (WIS, 58 ± 1 ms vs. SHR-C, 79 ± 2 ms; P < 0,05). O TABI aumentou a amplitude do transiente de [Ca\(^{2+}\)]; (WIS, 2,28 ± 0,07 F/F\(_0\) vs. SHR-C, 2,48 ± 0,08 F/F\(_0\); vs. SHR-T, 2,87 ± 0,08 F/F\(_0\); P < 0,05), mas não alterou os tempos até o pico e a metade do nível de repouso. Conclusão: O TABI não teve efeito sobre as propriedades estruturais e mecânicas de miócitos do VD de SHR, embora tenha aumentado a amplitude do transiente de [Ca\(^{2+}\)], e reduzido a PAS.

Nível de evidência I, Estudos terapêuticos – Investigação dos resultados do tratamento.

Descritores: Exercício; Ratos; Hipertensão; Coração.

RESUMEN

Introducción: Las adaptaciones estructurales y mecánicas de los miocitos del ventrículo derecho (VD) en respuesta a la hipertensión asociada al entrenamiento de resistencia de baja intensidad (ERBI) no se han estudiado en modelos experimentales de ratas. Objetivo: Determinar los efectos del ERBI sobre las propiedades estructurales y mecánicas de los miocitos del VD en ratas espontáneamente hipertensas (SHR). Métodos: Ratas SHR y ratas Wistar machos y normotensas (edad: 16 semanas) fueron distribuidos en grupos (n = 7): WIS (Wistar control); SHR-C (SHR control) y SHR-T (SHR entrenados; 60 min/día, 50-60% de la capacidad máxima de ejercicio, 5 días/semana por 8 semanas). Se realizó la medición de presión arterial sistólica (PAS), dimensiones de los miocitos aislados del VD, contratilidad, transitorio intracelular de Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_t\)) y proteínas reguladoras de Ca\(^{2+}\). La estadística se realizó por ANOVA one-way, seguida por prueba de Tukey post hoc (α = 5%). Resultados: El ERBI redujo la PAS en SHR (SHR-C, 164 ± 2 mmHg vs. SHR-T, 152 ± 4 mmHg; P < 0,05). La hipertensión aumentó el largo del miocito (WIS, 156,8 ± 2,7 µm; SHR-C, 166,6 ± 3,1 µm; P < 0,05), pero no afectó el largo o el volumen (P > 0,05). El ERBI no afectó la amplitud del transitorio de [Ca\(^{2+}\)]; (WIS, 2,28 ± 0,07 F/F\(_0\) vs. SHR-C, 2,48 ± 0,08 F/F\(_0\); vs. SHR-T, 2,87 ± 0,08 F/F\(_0\); P < 0,05), pero no alteró los tiempos hasta el pico y la mitad del nivel de reposo. Conclusión: El ERBI no tuvo efecto sobre las propiedades estructurales y mecánicas de los miocitos del VD en SHR, aunque aumentó la amplitud del transitorio de [Ca\(^{2+}\)], y redujo la PAS.

Nivel de evidencia I, Estudios terapéuticos – Investigación de los resultados del tratamiento.
INTRODUCTION

Hypertension is the leading risk factor for cardiovascular disease and a precursor of heart failure thus producing major concern to public health authorities. Hypertension impacts negatively the myocardial structure and function leading to systolic and diastolic impairment, mainly in the left ventricle. The spontaneously hypertensive rat (SHR) is a well-established model of genetic hypertension. Young (4 months) SHR show a compensated cardiac state and in spite of presence of severe myocardial hypertrophy, the left ventricle pump function is preserved until the age of 12 months. However, severe cardiac dysfunctions develop thereafter and presence of heart failure is reported at the age between 18 and 24 months. In the compensated phase of hypertension in SHR, it has been reported that the left ventricular (LV) myocyte shortening increases, whereas the action potential duration and the time course of intracellular global Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_i\)) transient, cell shortening and relaxation are prolonged.  

Preservation of cardiac functioning can be obtained by several interventions in hypertension, like the anti-hypertensive therapy, a healthy diet, smoking cessation, being exercise training an important hypotensive non-pharmacological therapeutic strategy. The recommended low-intensity exercise training (LIET) improves LVotropic e isotropic performance, β-adrenergic responsiveness, phosphorylation of [Ca\(^{2+}\)]\(_i\) regulatory proteins, and attenuates systolic dysfunction in the compensatory phase of hypertension in female SHRs. Most of these studies have been concentrated in the left ventricle myocardium while little is known about the effects of hypertension and LIET in the right ventricle.

It is important to point out that overall cardiac performance depends on the simultaneous and synchronous activity of both ventricles. Right ventricle function is also an important prognostic marker in conditions that have traditionally been regarded as primarily LV pathologic conditions, such as congestive heart failure and acute myocardial infarction. Right ventricle works as a barometer of cardiovascular symptoms, once total exercise time until fatigue (TTF). The training intensity throughout the training period was progressively increased, which reached 1 h/day, 0% grade, at 50–60% of MRS on the third week. The MRS test was repeated at the end of the 4\(^{th}\) week of training in the animals of the SHR-T group to update the training intensity. Two days after the last training session, the MRS test was repeated in all animals to evaluate their total exercise time until fatigue (TTF). The animals from the WIS and SHR-C groups were handled every day (5-10 min, 0% grade, 0.3 km/h, 3 days/week).

The body weight (BW) of all rats was measured every week, and the resting heart rate (RHR) and the systolic arterial pressure (SAP) were measured every week. A tail-cuff method (PowerLab 4/30, ADInstruments, USA). Two days after the last MRS test, the rats were weighed and then sacrificed by cervical dislocation. The hearts were removed and mounted on a Langendorff system and superfused for ~5 min with a modified Hepes-Tyrode solution with the following composition (in mM): 116 NaCl, 1.43 MgCl\(_2\), 5.4 KCl, 0.75 CaCl\(_2\), 5 Hepes, 10 glucose, 20 taurine and 10 creatine, pH 7.3 at 37°C. The perfusion solution was changed to

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Descriptores: Ejercicio; Ratas; Hipertensión; Corazón.

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a calcium-free solution containing EGTA (0.1 mM) for 6 min, and after that, hearts were perfused for 15–20 min with a solution containing 1 mg/ml collagenase type II ( Worthington, USA). The digested heart was removed from the apparatus, the ventricles and the right ventricle were carefully isolated and weighed. The right ventricle was cut into small pieces and placed in conical flasks with collagenase-containing solution supplemented with 1% bovine serum albumin ( Sigma-Aldrich, Saint Louis, MO, USA). The cells were dispersed by agitating the flasks for periods of 5 min at 37°C, and were separated from the non-dispersed tissue by filtration and stored at −5°C until needed. Only the calcium-tolerant, quiescent, rod-shaped cardiomyocytes showing clear cross striations were studied, within 2–3 h after isolation.

The isolated cardiomyocytes were placed in a chamber mounted on the stage of an inverted microscope (Nikon Eclipse, TS100, USA). The chamber was perfused with Hepes-Tyrode solution at room temperature. Steady-state 1 Hz contractions were elicited via platinum bath electrodes (Myopacer, Field Stimulator, Ionoptix, USA) with 5 ms voltage pulses with an intensity of 20 V. The cells were visualized on a PC monitor with a NTSC camera (Mycom, Ionoptix, USA). This image was used to measure cell shortening in response to electrical stimulation using a video motion edge detector (IonWizard, Ionoptix, USA). Cell shortening (expressed as a percentage of resting cell length) and maximal velocity of contraction and relaxation were calculated. The cell image was used to determine the measurements of resting cell length and midpoint width, which were used to calculate the cell volume as previously described.²⁴

Isolated cardiomyocytes from the right ventricle were loaded with fluo-4 AM, membrane-permeant form of the Ca²⁺ indicator at 5 μM (Molecular Probes, Eugene, OR, USA) for 20 min at room temperature and subsequently washed with the extracellular Hepes-Tyrode solution to remove dye excess, [Ca²⁺], transients were elicited by field-stimulating through a pair of platinum electrodes with a 0.2 ms supra-threshold voltage square pulses. Cells were stimulated at 1 Hz to produce steady-state conditions. A Meta LSM 510 scanning system (Zeiss GmbH, Germany) with a 63x oil immersion objective was used for confocal fluorescence imaging. Fluo-4 was excited at 488 nm (Argon laser), and the emission intensity was measured at > 510 nm. The digital image was processed using the Matlab® platform. The Ca²⁺ levels were reported as F/F₀, where F is the maximal fluorescence intensity average measured during the systolic phase of the [Ca²⁺] transients, and F₀ is the minimal fluorescence intensity average measured between 1 Hz contractions during the diastolic phase of the [Ca²⁺] transients. The time to peak of the [Ca²⁺], transient and time from peak transient to half resting level of [Ca²⁺], were determined.

The RV samples harvested from the myocardium after enzymatic perfusion were homogenized in a buffer containing 50 mM potassium phosphate (pH 7.0), 0.3 M sucrose, 0.5 mM DTT, 1 mM EDTA (pH 8.0), 0.3 mM PMSF, 10 mM NaF and phosphatase inhibitors cocktail (1:100, Sigma-Aldrich, Saint Louis, MO, USA). The cells were dispersed by agitating the flasks for periods of 5 min at 37°C, and were separated from the non-dispersed tissue by filtration and stored at ~5°C until needed. Only the calcium-tolerant, quiescent, rod-shaped cardiomyocytes showing clear cross striations were studied, within 2–3 h after isolation.

Hypertension induced a increase in RV cell length (Figure 1A), but did not affect cell width and volume. (Figures 1B, 1C) Liet did not modify RV cell length, width and volume in hypertensive rats. (Figures 1A, 1B, 1C)

**Cardiomyocyte contractility and [Ca²⁺], transients**

Hypertension and Liet did not modify the contractile properties of the RV myocytes in WIS, SHR-C and SHR-T groups. (Figures 2A, 2B, 2C) Figure 3 shows typical line-scan images recorded from field-stimulated cardiomyocytes that were loaded with the fluorescent Ca²⁺ indicator fluo 4-AM along with representative records of cardiomyocytes contractions. As shown in Figure 4A, hypertension did not affect [Ca²⁺], transient amplitude and cell shortening, respectively. Liet increased the [Ca²⁺], transient amplitude of RV myocytes from SHR-T group compared to SHR-C and WIS groups (Figure 4A), although it has not affected cell shortening (Figure 2A).
Notably, no differences were observed in the baseline fluorescence intensity average that was measured between contractions at the diastolic phase of \([Ca^{2+}]_i\) transients between the three groups. The time to peak of \([Ca^{2+}]_i\) transient (Figure 4B) of RV cells from SHR-C group was higher as compared to those from WIS group. LIET did not modify this parameter in cells from SHR-T group (Figure 4B). Hypertension and LIET did not affect the time from peak to half-resting level of \([Ca^{2+}]_i\) (Figure 4C). SHR-T group had faster time from peak to half-resting level of \([Ca^{2+}]_i\) when compared to the WIS group. (Figure 4C)

Neither hypertension nor LIET modified the expression of calcium regulatory proteins in the right ventricle of the WIS, SHR-C and SHR-T groups. (Figures 5A, 5B, 5C e 5D)

DISCUSSION

We observed that LIET was able to improve the exercise capacity and a tendency to decrease RHR. This result indicates that the exercise training protocol was efficient to produce cardiac adaptations. The resting bradycardia has been considered to be a hallmark of exercise training adaptation in experimental animals.\(^{25}\)

More important, LIET was efficient in decreasing the SAP in SHR animals. The hypotensive effect of regular exercise in hypertensive individuals is well-documented.\(^{1,10,11}\) The exercise intensity influences its pressure-lowering effect, inasmuch as larger reductions are detected at lower exercise training intensities.\(^{12,26}\)

We also observed that hypertension increased the RVW/TL ratio along with RV myocyte length, but our LIET did not counteract these structural adaptations. It is described that aerobic exercise-induced cardiac hypertrophy is more pronounced in the left ventricle, while ventricular dilation is common in both ventricles.\(^{27}\) Our results suggest some degree of hypertrophy in the right ventricle caused by hypertension, which may contribute to chamber enlargement. Pathological LV hypertrophy is associated with cardiac dysfunction, reduced \([Ca^{2+}]_i\), transients and contractility, increased interstitial fibrosis, decreased vascularization, and re-expression of fetal genes.\(^{2,8,28,29}\) On the other hand, the right ventricle can be affected by conditions that have been regarded as LV pathologic conditions, like the increased filling pressures during hypertension,\(^{19}\) in addition to the increased sympathetic activity in both ventricles.\(^{19}\) Our
data cannot confirm a established pathological hypertrophy in the RV myocytes from SHRs as we observed only an increase in the time to peak of the \([\text{Ca}^{2+}]_i\) transient. The other parameters of the \([\text{Ca}^{2+}]_i\) transient were normal, as well as cellular contractility and the expression of the \([\text{Ca}^{2+}]_i\) transient regulatory proteins, although we did not measure the histological content of this chamber.

Our results demonstrate that LIET induced an increase in the amplitude of RV myocyte \([\text{Ca}^{2+}]_i\), transient in SHR, but did not affect cellular mechanical properties. Because the \([\text{Ca}^{2+}]_i\) transient controls cardiomyocyte contractile activity, exercise training-induced changes in cell contractility are mediated by changes in the \([\text{Ca}^{2+}]_i\) transient. However, we did not observe any modifications in RV cell contractility in SHR-T group. Normally, the related changes in the rates of rise and decay of \([\text{Ca}^{2+}]_i\), transient and the rates of contraction and relaxation, indicate that changes in contractility and in the \([\text{Ca}^{2+}]_i\) transient are closely linked\(^3\), as we observed in LV myocytes from SHR and normotensive rats\(^2\) submitted to LIET.

The cardiomyocyte contraction-relaxation and the \([\text{Ca}^{2+}]_i\) transient are controlled by proteins that regulate calcium handling.\(^3\) We demonstrated here that LIET did not modify the expression of PLB\(_{er16}\), SERCA2a and NCX in the RV of SHR-T group. Previously, our group showed that hypertension did not affect the expression of PLB\(_{er16}\), PLB\(_{ser16}\), SERCA2a, or NCX in the LV of SHR animals. However, LIET increased the expression of SERCA2a and PLB\(_{er16}\) and reduced the PLB\(_{er}/\text{SERCA2a}\) ratio without changes in NCX and PLB\(_{er}\) levels in the LV of both hypertensive and normotensive rats\(^6\). In normotensive rats, we have already shown that LIET was not capable of modifying these proteins in the RV\(^2\).

In this study, RV myocytes of SHR demonstrated tiny responses to LIET. We have to consider that RV is connected serially to the LV and thus is obligated to pump the same effective stroke volume. Nevertheless, with a thinner and more compliant wall than the LV, it has a distinct physiology related to the low hydraulic impedance characteristics of the pulmonary vascular bed. The output of left and right ventricles are similar, under normal conditions, but the RV output requires approximately 25% of stroke work which implicates in less energy cost because of the low pressure pulmonary system and to the unique characteristics of the RV pressure–volume relationship\(^5\). It is worthy to note that these RV specific characteristics could potentially explain the absence of more pronounced adaptations in response to LIET at the cellular and molecular levels.

Moreover, the absence greater structural and mechanical changes in the RV myocytes from SHR submitted to LIET can be explained by the age of these animals since they were in the compensated phase of cardiac remodeling, where the myocardium of both control and exercised animals are able to cope with exercise and hypertension stresses in a positive way. Another possibility is that structural and mechanical adaptive responses of the RV myocytes of SHR to LIET depend on exercise protocols with longer duration and higher intensity. It is noteworthy that different aerobic exercise types or resistance training may lead to different adaptations.

The analysis of RV structural and mechanical properties at the cellular and molecular levels is important to characterize the functional and morphological remodeling in response to endurance training during hypertension. Advances in the comprehension of such cardiac remodeling is of clinical relevance in the assessment of exercise performance and may help to understand the differences between physiological and pathological forms of RV remodeling. Additional studies are necessary to understand the effects of different types of exercise training on the structure and function of the right ventricle in this model of hypertension.
CONCLUSION

The compensation phase of hypertension in SHR leads to mild eccentric hypertrophy in the RV, probably due to increased cell length. Although RV myocyte mechanical properties are not affected by either hypertension or LIET, the time course of the \([\text{Ca}^{2+}]_i\) transient prolongs in response to hypertension, while LIET increases the amplitude of the \([\text{Ca}^{2+}]_i\) transient. Despite that, neither hypertension nor LIET modify the RV \(\text{Ca}^{2+}\) regulatory proteins.

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REFERENCES


All authors declare no potential conflict of interest related to this article.

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