

# LJMU Research Online

Fecchio, RY, de Sousa, J, Oliveira-Silva, L, da Silva Junior, N, de Abreu, A, da Silva, G, Drager, L, Low, DA and Forjaz, C

Effects of dynamic, isometric and combined resistance training on blood pressure and its mechanisms in hypertensive men.

https://researchonline.ljmu.ac.uk/id/eprint/18703/

#### Article

**Citation** (please note it is advisable to refer to the publisher's version if you intend to cite from this work)

Fecchio, RY, de Sousa, J, Oliveira-Silva, L, da Silva Junior, N, de Abreu, A, da Silva, G, Drager, L, Low, DA ORCID logoORCID: https://orcid.org/0000-0001-7677-8634 and Forjaz, C (2023) Effects of dynamic, isometric and combined resistance training on blood pressure and its mechanisms in

LJMU has developed LJMU Research Online for users to access the research output of the University more effectively. Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Users may download and/or print one copy of any article(s) in LJMU Research Online to facilitate their private study or for non-commercial research. You may not engage in further distribution of the material or use it for any profit-making activities or any commercial gain.

The version presented here may differ from the published version or from the version of the record. Please see the repository URL above for details on accessing the published version and note that access may require a subscription.

For more information please contact <a href="mailto:researchonline@ljmu.ac.uk">researchonline@ljmu.ac.uk</a>

1 **Tittle:** Effects of dynamic, isometric and combined resistance training on blood pressure 2 and its mechanisms in hypertensive men. 3 **Authors**: Rafael Y Fecchio<sup>1</sup>; Julio CS de Sousa<sup>1</sup>; Laura Oliveira-Silva<sup>1</sup>; Natan D da Silva 4 Junior<sup>1</sup>; Andrea P de Abreu<sup>2</sup>; Giovânio V da Silva<sup>2</sup>; Luciano F Drager<sup>2</sup>; David A Low<sup>3</sup>; 5 Cláudia LM Forjaz<sup>1</sup> 6 7 1 Exercise Hemodynamic Laboratory, School of Physical Education and Sport, 8 9 University of São Paulo, São Paulo, São Paulo, Brazil. 10 2 Hypertension Unit, Renal Division of Hospital das Clínicas, Medical School, University 11 of São Paulo, São Paulo, São Paulo, Brazil. 3 Research Institute of Sport and Exercise Sciences, Faculty of Science, Liverpool John 12 Moores University, Liverpool, Merseyside, United Kingdom. 13 14 15 **Conflicts of Interest** 16 We declare that the authors do not have any conflict of interest. 17 **FUNDING** 18 This study was supported by the National Council for Scientific and Technological 19 Development (CNPQ, process 304436/2018-6), the São Paulo Research Foundation 20 21 (FAPESP, process 2018/12390-1, 2018/19151-2, 2018/23653-3 and 2019/02649-0) and the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior - Brasil (CAPES, 22 process 0001). 23 24 25 Address for correspondence: Cláudia Lúcia de Moraes Forjaz 26 Av. Prof. Mello Moraes, 65, Butantã, São Paulo/SP - 05508-030 - Brazil 27

Phone: +55+11 30913136; FAX: +55+11 30913136 Email: cforjaz@usp.br

Word count: 4.661 number of tables: 3; number of figures: 3

28

29

30

#### **ABSTRACT**

32

Although dynamic resistance training (DRT) and isometric handgrip training (IHT) may 33 decrease blood pressure (BP) in hypertensives, the effects of these types of training have 34 35 not been directly compared, and a possible additive effect of combining IHT to DRT (combined resistance training - CRT), has not been investigated. Thus, this study 36 compared the effects of DRT, IHT and CRT on BP, systemic hemodynamics, vascular 37 function, and cardiovascular autonomic modulation. Sixty-two middle-aged men with 38 treated hypertension were randomly allocated among four groups: DRT (8 exercises, 50% 39 of 1RM, 3 sets until moderate fatigue), IHT (30% of MCV, 4 sets of 2 min), CRT (DRT 40 + IHT) and control (CON – stretching). In all groups, the interventions were administered 41 42 3 times/week for 10 weeks. Pre- and post-interventions, BP, systemic hemodynamics, 43 vascular function and cardiovascular autonomic modulation were assessed. ANOVAs and ANCOVAs adjusted for pre-intervention values were employed for analysis. Systolic BP 44 decreased similarly with DRT and CRT (125±11 vs. 119±12 and 128±12 vs 119±12 45 mmHg, respectively; all P<0.05), while peak blood flow during reactive hyperaemia (a 46 marker of microvascular function) increased similarly in these groups (774±377 vs. 47 1067±461 and 654±321 vs. 954±464 mL/min, respectively, all P<0.05). DRT and CRT 48 did not change systemic hemodynamics, flow-mediated dilation, and cardiovascular 49 autonomic modulation. Additionally, none of the variables were changed by IHT. In 50 conclusion, DRT, but not IHT, improved BP and microvascular function in treated 51 52 hypertensive men. CRT did not have any additional effect in comparison with DRT alone.

53

- **Keywords**: hypertension; strength training; vascular function; autonomic modulation;
- 55 hemodynamics

#### INTRODUCTION

Hypertension is one of the major modifiable risk factors for cardiovascular disease<sup>1</sup>, causing around 8 million deaths per year, mainly due to stroke, myocardial infarction and sudden death<sup>2</sup>. Blood pressure (BP) control among individuals with hypertension remains sub-optimal (i.e., 43.5%)<sup>3</sup>, and complementary non-pharmacological interventions, such as exercise training, are recommended to improve BP control<sup>3,4</sup>. Recently, resistance training has been considered for hypertension treatment with dynamic resistance training (DRT) recommended by both the American and European guidelines<sup>3,4</sup>, while isometric handgrip training (IHT) is advised only by the American guidelines<sup>3</sup>.

Meta-analytic data demonstrated that DRT reduces systolic/diastolic blood pressures (SBP/DBP) by -6.11 (95%CI: -10.23 to -1.99) / -2.75 (95%CI: -4.27 to -1.22) mmHg in treated hypertensives<sup>5</sup>. Such effects may be related to vascular adaptations induced by training since studies have reported improved resistance vessel function in healthy<sup>6</sup> and pre-hypertensive<sup>7</sup> individuals after DRT, which still needs to be evidenced in hypertensives. Concerning IHT, a recent meta-analysis<sup>8</sup> indicated that it decreases SBP/DBP by -6.00 (95%CI: -7.75 to -4.26 / -2.75 (95%CI: -3;78 to -1.72) mmHg, which might be related to the training effects improving cardiac vagal modulation and vasomotor sympathetic modulation<sup>9</sup>.

Current literature has suggested IHT in hypertension management based on its potential of higher adherence given its short duration (11 min per session) and execution with portable device<sup>10</sup>. However, its use as a stand-alone exercise therapy has drawbacks. Differently from DRT that promotes generalized musculoskeletal and metabolic benefits<sup>11</sup>, IHT has musculoskeletal effects confined to the small muscle mass exercised

and only minor impact on overall health. Given that, IHT is recommended in addition, and not in place of conventional exercise modes, such as DRT<sup>10</sup>. However, by the best of our knowledge, no previous study investigated the possible additive effect of associating IHT to DRT on BP control.

Based on this background, it is possible to hypothesise that the addition of IHT to DRT, in a combined resistance training (CRT), besides improving general health status, may also induce a greater BP decrease in hypertensives as such protocol would combine the DRT vascular effects<sup>6,7</sup> and the IHT autonomic effects<sup>9</sup>.

Therefore, the current clinical trial was designed to assess and compare the effects of DRT alone, IHT alone and CRT on BP, systemic hemodynamics, markers of vascular function, and cardiovascular autonomic modulation in treated hypertensives. The hypotheses were: i) DRT alone would decrease BP and improve vascular function; ii) IHT alone would equally decrease BP compared with DRT and would improve cardiovascular autonomic modulation; and iii) CRT would induce a greater BP-lowering effect than both DRT and IHT, promoting both vascular and autonomic improvements.

### **METHODS**

# **Subjects**

This study was registered at the Brazilian Clinical Trials [(RBR-4fgknb) at <a href="http://www.ensaiosclinicos.gov.br">http://www.ensaiosclinicos.gov.br</a>, and all procedures were approved by the Ethics Committee of the School of Physical Education and Sport, University of São Paulo (process 2.870.688). All participants were informed of the benefits and risks of the

investigation prior providing written consent before enrolment. Experimental procedures were performed at the School of Physical Education and Sport of University of São Paulo.

Preliminary medical evaluation was performed at the Hospital das Clínicas of the Medical School of the University of São Paulo.

Middle-aged (30 to 65 years old) hypertensive men were recruited from advertisements posted at the University of Sao Paulo's media. The study was conducted with men to avoid the influence of menstrual cycle and menopause status on BP and its mechanisms<sup>12</sup>.

The inclusion criteria were: i) be receiving anti-hypertensive pharmacological treatment with drugs and doses maintained for at least the last 4 months; and ii) not be physically active (i.e. not accumulating more than 150 min per week of leisure physical activity, not performing exercise training more than 2 times per week, and had not performed resistance training in the previous 6 months). The exclusion criteria were: i) taking drugs that directly act on cardiac autonomic modulation (i.e. nondihydropyridine calcium channel blockers or beta-adrenergic receptor antagonists); ii) presence of secondary hypertension; iii) presence of hypertension-induced target organ damage; iv) presence of other cardiovascular disease despite hypertension; v) presence of symptoms or electrocardiographic alterations during a graded maximal exercise test; vi) body mass index  $\geq 35 \text{ kg/m}^2$ ; vii) presence of diabetic complications or insulin use; viii) presence of musculoskeletal problems that impair resistance training execution; and ix) SBP/DBP  $\geq$  160/105 mmHg that are the maximal BP values recommended for beginning exercise by the Brazilian Hypertension Guidelines<sup>13</sup>.

Inclusion and exclusion criteria were checked through preliminary procedures. In an initial visit, the participants answered an anamnesis, fulfilled a questionnaire, and underwent anthropometric and BP evaluations. The anamnesis involved questions about health history, regular medication use, and physical activity routines. The International Physical Activity Questionnaire was completed<sup>14</sup>. Weight and height were measured (Welmy® W300A, São Paulo, Brazil) and body mass index calculated. Auscultatory BP was measured in triplicate on both arms with the participants in the seated position for at least 5 min. This BP evaluation was repeated in another visit and the six values obtained for each arm were averaged with the highest value between the arms being considered as the BP level of each participant. In another visit, medical evaluations were conducted, including clinical examination and collection of urine and blood samples to exclude secondary hypertension and target-organ lesion. For that, the basic laboratorial evaluation recommended by the Brazilian Hypertension Guidelines<sup>13</sup> were followed and included the analyses of plasma potassium, uric acid, and creatinine; fasting plasma glycose, triglycerides, and total, HDL- and LDL-cholesterol concentrations; conventional urine analyses; and the estimation of glomerular filtration rate. Finally, a graded maximal exercise test was performed on a cycle ergometer (Lode Medical Technology, Corival, Groningen, Netherlands) with electrocardiogram (Welch Allyn, Cardioperfect ST2001 model, Netherlands) evaluated by a physician.

129

130

131

132

133

134

135

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

The participants who fulfilled the study criteria underwent two familiarization sessions to the exercises employed in the study as already done in previous research<sup>15</sup>. In these sessions, they executed 2 sets of 20 repetitions with the lowest workload allowed by each equipment (Edge Line, Movement Fitness, Sao Paulo, Brazil) in 8 dynamic resistance exercises (bench press, leg press, lat pull down, left leg extension, right leg extension, arms curl, left leg curl and right leg curl) followed by the execution of 4 sets of 2 min isometric handgrip exercise at 5% of maximal voluntary contraction (MVC). On another day, they did 1 repetition maximum (1RM) tests in all aforementioned exercises

following standardized protocol<sup>16</sup> as already done in previous studies<sup>17,18</sup>. Afterwards, participants performed a standardized evaluation of handgrip MVC with left and right hands<sup>19</sup>.

### **Procedures**

This study was a four-parallel-arm randomized controlled trial designed to evaluate and compare the effects of DRT, IHT and CRT. The pre-specified primary outcome was BP, and the secondary outcomes were muscle strength, systemic hemodynamics, vascular function, and cardiovascular autonomic modulation.

The participants were randomly allocated among four groups: DRT, IHT, CRT and control (CON), with a 1:1:1:1 allocation ratio. Randomization was performed after the pre-intervention evaluations by an independent researcher (i.e. not involved directly in the recruitment and data collection) using the block method through sealed envelopes (i.e. sorting among the four options in each envelop). In all four groups, the intervention period lasted 10 weeks and the intervention sessions were conducted 3 times per week. Each session was individually supervised by an exercise specialist and conducted at the institution's gym facility. The outcomes were assessed in experimental sessions conducted pre- and post-interventions, with the post-evaluations being conducted after a minimal interval of 48h in relation to the last intervention session.

Prior to the experimental sessions, the participants received the following instructions: i) not to ingest vitaminic supplements in the previous 72h; ii) not to perform exercise in the previous 48h; iii) not to consume alcoholic beverages in the previous 24h; iv) not to smoke in the previous 8h; v) to keep their usual daily activities and sleep habits in the previous day; vi) to use their regular medications as usual; and vii) to come to the

session after fasting for at least 8h. The experimental sessions started between 07:00-07:30 a.m. and the laboratory temperature was maintained between 20-22°C.

During the experimental sessions, assessments started after 10 min of seated rest. Firstly, continuous signals of electrocardiogram, photoplethysmographic BP and respiration were recorded for 10 min for cardiovascular autonomic modulation evaluation. Then, auscultatory BP, cardiac output (CO) and heart rate (HR) were measured in triplicate for systemic hemodynamic evaluation. Afterwards, for vascular evaluation, the participants moved to the supine position, and after a 10-min interval, images and doppler flow signals of the brachial artery were recorded initially for 1 min without any stimulus (baseline) and then for 3 min after 5 min of forearm vascular occlusion (post-occlusion).

#### **Interventions**

The DRT group executed the 8 dynamic resistance exercises previously mentioned on specialized equipment (Edge Line, Movement Fitness, Sao Paulo, Brazil). In each exercise, the participants executed 3 sets of repetitions until moderate fatigue (defined by a visual reduction on movement velocity) and kept a 90-s interval between sets and exercises. The intensity was initially set at 50% of 1RM and was increased by 2-5% and 5-10% for upper- and lower-limb exercises, respectively, when the participants could perform more than 15 repetitions without moderate fatigue in two consecutive sets <sup>20</sup>. This DRT protocol followed the hypertension guidelines<sup>3,4</sup>.

The IHT group executed the isometric handgrip exercise on a specific device (ZonaPlus, Zona Health, Boise, Idaho, USA). In each session, the participants executed 4 sets of 2-min isometric contractions at 30% of MVC, alternating the hands (i.e. 2 sets per hand) and maintaining a 60-s interval between the sets. MVC was measured at the

beginning of each training session. After each session, the device provided a score quantifying the performance of the handgrip squeeze, and values  $\geq 80$  indicated effective training. This IHT protocol followed hypertension guidelines<sup>3</sup>.

The CRT group executed, in each training session, the same protocol as the DRT group followed by the same protocol performed by the IHT group.

The CON group executed 30-min stretching sessions. In each session, the participants executed 20 to 25 exercises and in each exercise, they executed 2 to 3 attempts keeping the highest degree of stretching without pain for 20-30 s. This active control intervention was proposed for this study to assure a similar interaction of the participants with the research team and to multiple BP measurements, since it is known that adaptation to these factors that would happen in the training groups (DRT, IHT and CRT) can decrease BP.

Adherence to each intervention was calculated as the percentage of the 30 offered sessions actually performed by each participant (i.e. sessions performed / 30 x 100).

## Measurements

Auscultatory BP was measured by a trained evaluator using a calibrated aneroid sphygmomanometer (Mikatos, Missouri, Sao Paulo, Brazil). Measurements were done on the dominant arm employing an adequate cuff size. SBP and DBP were respectively defined as phases I and V of Korotkoff sounds. Mean BP (MBP) was calculated as: MBP =  $DBP + [1/3 \times (SBP - DBP)]^{21,22}$ .

For systemic hemodynamic evaluation, CO was assessed by the indirect Fick method through CO<sub>2</sub> rebreathing technique<sup>23</sup> using a gas analyser (Medical Graphics

Corporation, CPX/Ultima, Minnesota, USA) and a bag containing hypercapnic gas (8-10%  $CO_2$ ). Firstly, the participants spontaneously breathed the ambient air for the measurement of  $CO_2$  production and the estimation of  $CO_2$  arterial content from end-tidal  $CO_2$  pressure. Then, via a two-way valve, participants started to inhale the hypercapnic gas until  $CO_2$  achieved an equilibrium and  $CO_2$  venous content could be estimated. Then, CO was calculated as:  $CO = VCO_2$  / ( $CO_2$  venous content –  $CO_2$  arterial content). Systemic vascular resistance (SVR) was calculated from: SVR = MBP / CO. Stroke volume (SV) was calculated from: SV = CO / HR.

225

226

227

228

229

230

231

232

233

234

235

236

237

238

239

240

241

242

243

244

245

246

247

248

249

Vascular function evaluation was assessed through a linear array probe attached to a high-resolution ultrasound machine (General Eletric Medical Systems, LOGIQ 7, California, USA) following guidelines<sup>24,25</sup>. Assessments were performed at the brachial artery of the dominant arm, ~5 cm proximal to the antecubital fossa, and using an insonation angle of 60°. Firstly, vascular images and doppler flow signal were continuously recorded for 1 min as baseline. From these records, arterial diameter was automatically detected, and blood flow velocity was quantified (Quipu, Cardiovascular Suite, Pisa, Italy). Blood flow (BF) was calculated as: BF = arterial cross-sectional area x blood flow velocity. Vascular conductance (VC) was calculated as: VC = BF / MBP. For vascular function assessment, a vascular occlusion period was initiated immediately after the baseline assessment using a cuff positioned at the forearm that was inflated to 250 mmHg for 5 min. When the cuff pressure was released, recordings of vascular images and doppler signals were taken for 3 min. Microvascular function (i.e. resistance vessels function) was assessed by the peak BF (i.e. highest absolute value) achieved during the reactive hyperaemia following cuff deflation<sup>25</sup>. Arterial endothelial function was assessed by flow-mediated dilation (FMD)<sup>24</sup> calculated by arterial diameter change from the baseline to the post-occlusion period as: FMD (%) = [(peak arterial diameter - baseline)] arterial diameter) / baseline diameter] x 100. The stimulus underlying FMD was evaluated by peak shear rate calculated at the post-occlusion period as: peak shear rate = 4 x peak BF velocity / arterial diameter.

Cardiovascular autonomic modulation evaluation followed the respective Task Force guidelines<sup>26</sup>. Briefly, HR was continuously measured through three-lead electrocardiogram (EMG System of Brazil, EMG 030110/00B), Sao Paulo, Brazil), beat-by-beat BP was monitored using finger photoplethysmography (Finapress Measurement System, Finometer, Arnhem, Netherland) and respiratory movements were measured via elastic thoracic belt (Pneumotrace 2, UFI, Morro Bay, USA). These signals were continuously acquired and recorded through a data acquisition system (Dataq Instruments, DI-720, Akron, Ohio, USA) with a sampling rate of 500Hz. Temporal sequences of R-R intervals, SBP and respiration were generated and analysed at the frequency domain through the autoregressive model using the Heart Scope II Software (A.M.P.S. LLC, Version 1.3.0.3, New York, USA). Cardiac sympathovagal balance was defined by the ratio between the low- and high-frequency bands of R-R interval variability (LF/HF<sub>R-R</sub>). Sympathetic vasomotor modulation was defined by the low-frequency band of SBP variability (LF<sub>SBP</sub>). Baroreflex sensitivity (BRS) was evaluated by the transfer function method<sup>27</sup>.

### **Statistical analysis**

The minimal sample size estimated for this study was 60 participants (i.e. 15 per study arm). This number was calculated for the primary outcome (SBP), considering an effect size (d) of -0.41<sup>28</sup>, a statistical power of 0.90, an alpha value of 0.05 and a correlation among repeated measures of 0.68<sup>29</sup>.

Data normality was checked by Shapiro-Wilks test and outliers identified through box plots. Non-normal data was transformed by natural logarithm to meet assumptions of the subsequent inferential analysis. The efficacy of interventions on the study's outcomes were analysed by two-way mixed ANOVAs considering group as a between factor (DRT vs. IHT vs. CRT vs. CON) and time (pre- vs. post-intervention) as a within factor. When significant main effects or interactions were observed, pairwise comparisons were done by Newman-Keuls post-hoc tests. Additionally, changes ( $\Delta$  = post-intervention – pre-intervention) adjusted for pre-intervention values were compared between the groups by ANCOVAs, and Bonferroni post-hoc tests were applied for pairwise comparisons when a significant effect was observed.

Data is presented as mean  $\pm$  standard deviation, and significance level was set at P value < 0.05 for all analyses.

#### **RESULTS**

Data recruitment took place from September 2018 to November 2021. Due to coronavirus 2019 disease, the study's procedures were interrupted or restrained from March 2020 to September 2020 and from March 2021 to June 2021.

The clinical trial flowchart is shown in Figure 1. Two hundred and nineteen participants were contacted, 106 performed the initial visit, 96 provided written consent and 70 were randomly allocated into the study's groups. The clinical trial ended after the assignment of 70 participants considering the minimal sample size required (i.e. 60 participants) and a dropout rate of 15.0% <sup>30</sup>. Indeed, there were 8 (11.4%) dropouts during the intervention period and 62 participants concluded the entire experimental protocol. Due to technical issues, data for the autonomic modulation evaluation was missed for two

participants (DRT: n=1 and CON: n=1). As the study was designed to evaluate and compare the efficacy of DRT, IHT and CRT, only data from the subjects who finished the experimental protocol were analysed. Groups characteristics were similar at the beginning of the study as shown in Table 1.

Adherences to the intervention sessions were high and similar among the groups (DRT:  $89\pm7\%$ ; IHT:  $90\pm9\%$ ; CRT:  $90\pm7\%$ ; CON:  $88\pm9$ , p = 0.917). During the interventions, participants from CRT executed dynamic and isometric exercises with similar intensities and volumes as DRT and IHT, respectively (data not shown).

None of the interventions changed isometric handgrip MVC of the left nor the right arm (left: +1±6, +3±5, +2±9, and -1±4; and right: +1±6, +3±4, +2±5, and -1±6 kg for DRT, IHT, CRT and CON, respectively, all p > 0.05). On the other hand, DRT and CRT significantly increased 1RM strength (all p<sub>group x time</sub> <0.05) in all exercises (bench press: +11±11 and +11±10 kg; leg press: +33±24 and +32±26 kg; lat pull down: +12±9 and +11±7 kg; left leg extension: +10±11 and +11±10 kg; right leg extension: +10±12 and +11±10 kg; arms curl: +12±8 and +7±12 kg; left leg curl: +8±5 and +7±4 kg; and right leg curl: +8±4 and +6±5 kg for DRT and CRT, respectively), while no change was observed for the IHT and the CON groups (bench press: -1±4 and +3±7 kg; leg press: -4±16 and +5±17 kg; lat pull down: 0±4 and +1±10 kg; left leg extension: -3±9 and +2±8 kg; right leg extension: -4±10 and +1±7 kg; arms curl: -2±3 and -1±5 kg; left leg curl: 0±4 and 0±3 kg; and right leg curl: 0±4 and 1±4 kg for IHT and CON, respectively).

SBP decreased significantly from pre- to post-intervention after the DRT and the CRT and did not change after the IHT and the CON ( $p_{group\ x\ time}=0.003$ , Table 2). Additionally, SBP changes adjusted to pre-intervention values observed with DRT and CRT were significantly different from CON (p=0.002, Figure 2). DBP did not change

significantly in any group (p > 0.05, Table 2) and changes in DBP adjusted for preintervention values were similar among the groups (p = 0.096, Figure 2).

SVR, CO, SV and HR did not change significantly in any group (all p > 0.05, Table 2) and changes in these variables adjusted for pre-intervention values were similar among the groups (all p > 0.05, Figure 2).

Baseline BF and VC as well as FMD did not change significantly in any group (all p > 0.05, Table 3) and changes in these variables adjusted for pre-intervention values were similar among the groups (all p >0.05, Figure 3). There was significant main effect of time for peak shear rate ( $p_{time} = 0.011$ ), demonstrating that peak shear rate increased significantly and similarly from pre- to post-intervention in all groups, including CON. Accordingly, changes in peak shear rate adjusted for pre-intervention values were similar between the groups (p = 0.083). On the other hand, peak BF increased significantly from pre- to post-intervention after DRT and CRT and did not change after IHT and CON ( $p_{group \ x \ time} = 0.007$ ). Additionally, peak BF changes adjusted to pre-intervention values observed with DRT and CRT were significantly different from CON (p = 0.008).

Regarding autonomic modulation responses, there were no significant main effects nor interactions (group vs. time) for LF/HF<sub>R-R</sub>, nor LF<sub>SBP</sub> (all p > 0.05, Table 3). Accordingly, changes between the groups adjusted for pre-intervention values were similar for these variables (all p >0.05, Figure 3). There was a significant main effect of time for BRS ( $p_{time}$  = 0.046), showing that BRS increased significantly and similarly from pre- to post-intervention in all groups, including CON. Accordingly, changes in BRS adjusted for pre-intervention values were similar among the groups (p = 0.306).

The current study has two main findings. First, DRT, but not IHT, decreased BP and improved microvascular function in treated hypertensive men. Second, the addition of IHT to DRT, in the CRT, did not promote any additive effect in comparison to DRT alone on either BP, systemic hemodynamics, vascular function or autonomic modulation.

DRT produced a net reduction (i.e. DRT vs CON, Figure 2) of -8.4 [95%CI: -15.9 to -0.8] mmHg in SBP, which is in accordance with the study hypothesis and within the range of reduction reported in a previous meta-analysis for SBP in treated hypertensives after DRT (-6.1; 95%CI: -10.2 to -2.0 mmHg)<sup>5</sup>. Moreover, the BP-reduction observed is comparable to the net effect reported for aerobic training (-8.3; 95%CI: -10.7 to -6.0 mmHg)<sup>31</sup>, and for the main anti-hypertensive drug classes used in monotherapy (-8.8; 95%IC: -9.6 to -8.0 mmHg)<sup>32</sup>. This BP-lowering effect induced by DRT might have clinical relevance given that a 5 mmHg decrease in SBP has been shown to reduce the risk of major cardiovascular events by about 9% <sup>33</sup>. Indeed, 75% (n=12) of the participants in the DRT group presented this clinically meaningful reduction in SBP (Supplementary Figure 1).

The BP-lowering effect induced by DRT was accompanied by an increase in peak BF during hyperaemia, which reflects an improvement in microvascular function<sup>25</sup>. As BP is mainly regulated by resistance vessels, such improvement in microvascular function may be responsible, at least in part, for the reduction in SBP induced by DRT. By our knowledge, this is the first study to demonstrate that DRT improves microvascular function in treated hypertensives. This adaptation was probably triggered by mechanism deflagrated during each exercise execution. Along this line, skeletal muscle activity produces vasodilatory factors (e.g., adenosine, CO<sub>2</sub>, lactate/H<sup>+</sup>, and K<sup>+</sup>)<sup>34</sup>, but during the concentric phase of dynamic resistance exercise, blood flow is restricted<sup>35</sup>. However, during the rest periods between the exercise repetitions and sets, blood flow increases,

producing shear stress and vasodilation, which reveals ischemia/reperfusion cycles<sup>36</sup> that may chronically improve microvascular function<sup>37</sup>. Additionally, such microvascular function improvement after DRT might have clinical relevance once microvascular dysfunction is typical in hypertension<sup>7</sup>, and an attenuated reactive hyperaemia is associated with higher risk of major cardiovascular events<sup>38</sup>. On the other hand, DRT did not improve arterial endothelial function evaluated by FMD. Likewise, a previous study<sup>6</sup> with healthy individuals with preserved endothelial function also found unchanged FMD and increased peak BF after DRT. Thus, the absence of FMD changes after DRT in the current study might be related, at least in part, to the apparently preserved baseline FMD presented by the participants; which might be due to the fact that almost all the sample was taking angiotensin II receptor blockers or angiotensin-converting enzyme inhibitors that already improve FMD<sup>39</sup>.

Contrary to the study's hypothesis, IHT did not reduce SBP nor DBP. Indeed, although meta-analytic data indicates that IHT reduces BP in general population<sup>8</sup>, a recent evidence-based Consensus Document<sup>40</sup> concluded that such hypotensive effect is greater in normotensive than hypertensive individuals; suggesting that target population may explain, at least in part, this current result. Therefore, as the BP-lowering is clinically important in hypertension, more research is required to actually elucidate whether IHT can decrease BP in this specific population, i.e. treated hypertensives.

The present results also do not support an effect of IHT on cardiovascular autonomic modulation. Although, prior data<sup>9</sup> reported improvements in cardiovascular autonomic markers after IHT in hypertensives, a meta-analysis<sup>41</sup> published during this study execution concluded that IHT does not modify cardiac autonomic modulation in hypertensives. Therefore, the current results support that IHT does not improve cardiovascular autonomic control in treated hypertensives.

CRT produced a net reduction (i.e. DRT vs CON, Figure 2) of -10.7 [95%CI: -18.3 to -3.0] mmHg in SBP, with 60% (n=9) of the participants of this group presenting a clinically meaningful (> 5 mmHg) reduction in SBP (Supplementary Figure 1). In addition, CRT increased peak BF during reactive hyperaemia. These responses, however, were similar to DRT, demonstrating that CRT effects were driven by DRT and IHT had no additive effect. Interestingly, a previous meta-analysis<sup>31</sup> also reported no additive effect of the combination between DRT and aerobic exercise training in BP reduction. Therefore, obtaining an additive BP-lowering effect through the addition of different exercise modes seems to be challenging.

The current study has important clinical implications. The findings support DRT as a valuable additional non-pharmacological intervention for hypertension management since it reduced BP and improved microvascular function even in hypertensive patients already taking pharmacologic treatment. On the other hand, the results raise caution regarding the replace of conventional exercise modes by IHT for hypertension management given the observed lack of efficacy. Lastly, the present results do not also support the association of IHT to DRT given the absence of additive effects in comparison to DRT alone.

It is important to mention the limitations of the current study. Participants were non-active middle-aged men without cardiovascular disease. Thus, caution is needed when extrapolating the current results to individuals with other characteristics, such as elderly, women and patients with cardiovascular disease. Few participants (n=6, 9% of final sample) had been infected by SARS-CoV-2 before the study enrolment, but none of them had to be hospitalized, and their prevalence was similar among the study's groups. As in many clinical trials, although adequately powered for the primary outcome (SBP:  $\beta = 0.921$ ), analysis for secondary outcomes can be underpowered. Finally, the results

regarding the comparisons among the training protocols (DRT, IHT and CRT) are restricted to the specific protocols employed in the present study. It is possible to speculate that the divergent responses between DRT and IHT might be explained, at least in part, by the different amount of muscle mass involved in each protocol, since DRT enrolled a whole-body training and the vascular adaptations induced by training are greater in regions directly mobilized during the exercise sessions<sup>37,42</sup>. Nevertheless, the protocols employed in the present study were designed based on the recommendations of the hypertension guidelines<sup>3,4,13</sup> but the employment of other protocols might reveal different results.

In conclusion, DRT, but not IHT, reduced BP and improved microvascular function in treated hypertensive men. The addition of IHT to DRT, in a CRT protocol, did not produce additive effects when compared to DRT alone.

### **ACKNOWLEDGEMENTS**

The authors want to acknowledge the volunteers of the current study. This study was supported by the National Council for Scientific and Technological Development (CNPQ, process 304436/2018-6), the São Paulo Research Foundation (FAPESP, process 2018/12390-1, 2018/18327-0, 2018/23653-3 and 2019/02649-0) and the Coordination for the Improvement of Higher Education Personnel (CAPES, process 0001). We declare that authors or any organization with which we are associated do not have any conflict of interest.

#### SUPPLEMENTARY MATERIAL

Supplementary information is available at Hypertension Research's website

## **REFERENCES**

448	1	Brouwers S, Sudano I, Kokubo Y, Sulaica EM. Arterial hypertension. Lancet
449		2021; <b>398</b> : 249–261. doi:10.1016/S0140-6736(21)00221-X
450	2	Lawes CM, Hoorn S Vander, Rodgers A. Global burden of blood-pressure-
451		related disease, 2001. Lancet 2008; <b>371</b> : 1513–8. doi:10.1016/S0140-
452		6736(08)60655-8
453	3	Whelton PK, Carey RM, Aronow WS, Ovbiagele B, Casey DE, Smith SC et al.
454		$2017\ ACC\ /\ AHA\ /\ AAPA\ /\ ABC\ /\ ACPM\ /\ AGS\ /\ APhA\ /\ ASH\ /\ ASPC\ /\ NMA$
455		/ PCNA Guideline for the Prevention , Detection , Evaluation , and Management
456		of High Blood Pressure in Adults A Report of the American College of
457		Cardiology / American Heart Association T. <i>Hypertension</i> 2018; <b>71</b> : 1269–1324.
458		doi:10.1161/HYP.0000000000000066
459	4	Williams B, Mancia G, Spiering W, Rosei E, Azizi M, Burnier M. 2018
460		ESC/ESH Guidelines for the management of arterial hypertension. J Hypertens
461		2018; <b>36</b> : 1953–2041. doi:10.1093/eurheartj/ehy339
462	5	Oliver-Martínez PA, Ramos-Campo DJ, Martínez-Aranda LM, Martínez-
463		Rodríguez A, Rubio-Arias J. Chronic effects and optimal dosage of strength
464		training on SBP and DBP: a systematic review with meta-analysis. J Hypertens
465		2020; <b>38</b> : 1909–1918. doi:10.1097/HJH.0000000000002459
466	6	Rakobowchuk M, Mcgowan CL, de Groot PC, Hartman JW, Phillips SM,
467		MacDonald MJ. Endothelial function of young healthy males following whole
468		body resistance training. J Appl Physiol 2005; 98: 2185–2190.
469		doi:10.1152/japplphysiol.01290.2004.

4/0	/	Beck D1, Martin JS, Casey DP, Braith RW. Exercise training improves
471		endothelial function in resistance arteries of young prehypertensives. J Hum
472		Hypertens 2014; <b>28</b> : 303–9. doi:10.1038/jhh.2013.109
473	8	López-Valenciano A, Ruiz-Pérez I, Ayala F, Sánchez-Meca J, Vera-Garcia F.
474		Updated systematic review and meta-analysis on the role of isometric resistance
475		training for resting blood pressure management in adults ~. J Hypertens 2019; 37:
476		1320–1333. doi:10.1097/HJH.0000000000002022
477	9	Taylor AC, McCartney N, Kamath M V., Wiley RL. Isometric training lowers
478		resting blood pressure and modulates autonomic control. Med Sci Sports Exerc
479		2003; <b>35</b> : 251–256. doi:10.1249/01.MSS.0000048725.15026.B5
480	10	Millar P, Paashuis A, McCartney N. Isometric Handgrip Effects on
481		Hypertension. Curr Hypertens Rev 2009; 5: 54-60.
482		doi:10.2174/157340209787314351
483	11	Westcott WL. Resistance Training is Medicine: Effects of strength training on
484		health. Am Coll Sport Med 2012; 11: 209–216.
485		doi:10.1249/JSR.0b013e31825dabb8
486	12	Coylewright M, Reckelhoff JF, Ouyang P. Menopause and hypertension: An age-
487		old debate. <i>Hypertension</i> 2008; <b>51</b> : 952–959.
488		doi:10.1161/HYPERTENSIONAHA.107.105742
489	13	Barroso W, Rodrigues C, Bortolotto L, Gomes M, Brandão A, Feitosa A.
490		Brazilian Guidelines of Hypertension - 2020. <i>Arq Bras Cardiol</i> 2021; <b>116</b> : 516–
491		658.
492	14	Craig CL Marshall AL Siöström M Bauman AE Booth ML Ainsworth BE et

493		al. International physical activity questionnaire: 12-Country reliability and
494		validity. <i>Med Sci Sports Exerc</i> 2003; <b>35</b> : 1381–1395.
495		doi:10.1249/01.MSS.0000078924.61453.FB
496	15	Queiroz ACC, Sousa JCS, Silva ND, Tobaldini E, Ortega KC, De Oliveira EM et
497		al. Captopril does not Potentiate Post-Exercise Hypotension: A Randomized
498		Crossover Study. Int J Sports Med 2017; 38: 270–277. doi:10.1055/s-0042-
499		123044
500	16	Maud PJ, Foster C. Physiological Assessment of Human Fitness. 1st ed.
501		Champaign, IL: Human Kinetics doi:10.1097/00005768-199607000-00027
502	17	Queiroz ACC, Sousa JCS, Cavalli AAP, Silva Jr ND, Costa LAR, Tobaldini E et
503		al. Post-resistance exercise hemodynamic and autonomic responses: Comparison
504		between normotensive and hypertensive men. Scand J Med Sci Sport 2015; 25:
505		486–494. doi:10.1111/sms.12280
506	18	Rezk CC, Marrache RCB, Tinucci T, Mion D, Forjaz CLM. Post-resistance
507		exercise hypotension, hemodynamics, and heart rate variability: Influence of
508		exercise intensity. Eur J Appl Physiol 2006; <b>98</b> : 105–112. doi:10.1007/s00421-
509		006-0257-y
510	19	Carlson DJ, Inder J, Palanisamy SKA, McFarlane JR, Dieberg G, Smart NA. The
511		efficacy of isometric resistance training utilizing handgrip exercise for blood
512		pressure management: A randomized trial. Medicine (Baltimore) 2016; 95:
513		e5791. doi:10.1097/MD.000000000005791
514	20	Ratamess NA, Alvar BA, Evetoch TK, Housh TJ, Kibler W Ben, Kraemer WJ et
515		al. Progression Models in Resistance Training for Healthy Adults. <i>Med Sci Sport</i>

516		Exerc 2009; <b>41</b> : 687–708. doi:10.1249/MSS.0b013e3181915670
517	21	Malachias M, Souza W, Plavnik F, Rodrigues C, Sociedade Brasileira de
518		Cardiologia. 7ª Diretriz Brasileira de Hipertensão Arterial. Arq Bras Cardiol
519		2016; <b>107</b> : 1–83.www.arquivosonline.com.br
520	22	Sharman JE, Lagerche A. Exercise blood pressure: Clinical relevance and correct
521		measurement. J Hum Hypertens 2015; <b>29</b> : 351–358. doi:10.1038/jhh.2014.84
522	23	Collier CR. Determination of mixed venous CO2 tensions by rebreathing. <i>J Appl</i>
523		Physiol 1956; 9: 25–29.
524	24	Thijssen DHJ, Bruno RM, van Mil ACCM, Holder SM, Faita F, Greyling A et al.
525		Expert consensus and evidence-based recommendations for the assessment of
526		flow-mediated dilation in humans. Eur Heart J 2019; : 1–14.
527		doi:10.1093/eurheartj/ehz350
528	25	Limberg JK, Casey DP, Trinity JD, Nicholson WT, Wray DW, Tschakovsky ME
529		et al. Assessment of resistance vessel function in human skeletal muscle:
530		guidelines for experimental design, Doppler ultrasound, and pharmacology. Am J
531		Physiol Hear Circ Physiol 2020; 318: H301–H325.
532		doi:10.1152/ajpheart.00649.2019
533	26	European Society of Cardiology; North American Society of Pacing and
534		Electrophysiology. Heart rate variability. Standards of measurement,
535		physiological interpretation, and clinical use. Task Force of the European Society
536		of Cardiology and the North American Society of Pacing and Electrophysiology.
537		Eur Heart J 1996; <b>17</b> : 354–381. doi:10.1161/01.CIR.93.5.1043
538	27	Robbe HW, Mulder LJ, Rüddel H, Langewitz WA, Veldman JB, Mulder G.

539		Assessment of Baroreceptor Reflex Sensitivity by Means of Spectral Analysis.
540		Hypertension 1987; <b>10</b> : 538–543.
541	28	MacDonald H V., Johnson BT, Huedo-Medina TB, Livingston J, Forsyth KC,
542		Kraemer WJ et al. Dynamic Resistance Training as Stand-Alone
543		Antihypertensive Lifestyle Therapy: A Meta-Analysis. J Am Heart Assoc 2016;
544		<b>5</b> : e003231. doi:10.1161/JAHA.116.003231
545	29	Bottini B, Carr A, Prisant L, Rhoades R. Variability and similarity of manual
546		office and automated blood pressures. <i>J Clin Pharmacol</i> 1992; <b>32</b> : 614–619.
547	30	Maher CG, Sherrington C, Herbert RD, Moseley AM, Elkins M. Reliability of
548		the PEDro scale for rating quality of randomized controlled trials. Phys Ther
549		2003; <b>83</b> : 713–721. doi:10.1093/ptj/83.8.713
550	31	Cornelissen V, Smart N. Exercise Training for Blood Pressure: A Systematic
551		Review and Meta-analysis. J Am Heart Assoc 2013; 2: e004473.
552		doi:10.1161/JAHA.112.004473
553	32	Naci H, Salcher-Konrad M, Dias S, Blum MR, Sahoo SA, Nunan D et al. How
554		does exercise treatment compare with antihypertensive medications? A network
555		meta-analysis of 391 randomised controlled trials assessing exercise and
556		medication effects on systolic blood pressure. Br J Sports Med 2018; 0: 859–869.
557		doi:10.1136/bjsports-2018-099921
558	33	Adler A, Agodoa L, Algra A, Asselbergs FW, Beckett NS, Berge E et al.
559		Pharmacological blood pressure lowering for primary and secondary prevention
560		of cardiovascular disease across different levels of blood pressure: an individual
561		participant-level data meta-analysis. <i>Lancet</i> 2021; <b>397</b> : 1625–1636.

562		doi:10.1016/S0140-6736(21)00590-0
563	34	Sarelius I, Pohl U. Control of muscle blood flow during exercise: local factors
564		and integrative mechanisms. Acta Physiol 2010; 199: 349–365.
565		doi:10.1111/j.1748-1716.2010.02129.x
566	35	Asmussen E. Similarities and dissimilarities between static and dynamic exercise.
567		Circ Res 1981; <b>48</b> : I3-10. doi:10.1249/01.MSS.0000115224.88514.3A
568	36	Mohrman DE, Heller LJ. Cardiovascular Physiology. 8th ed. New York (NY):
569		McGraw-Hill Education
570	37	Green DJ, Hopman MTE, Padilla J, Laughlin MH, Thijssen DHJ. Vascular
571		Adaptation to Exercise in Humans: Role of Hemodynamic Stimuli. <i>Physiol Rev</i>
572		2017; <b>97</b> : 495–528. doi:10.1152/physrev.00014.2016
573	38	Anderson TJ, Charbonneau F, Title LM, Buithieu J, Rose MS, Conradson H et al.
574		Microvascular function predicts cardiovascular events in primary prevention:
575		Long-term results from the firefighters and their endothelium (FATE) study.
576		Circulation 2011; <b>123</b> : 163–169. doi:10.1161/CIRCULATIONAHA.110.953653
577	39	Shahin Y, Khan JA, Samuel N, Chetter I. Angiotensin converting enzyme
578		inhibitors effect on endothelial dysfunction: A meta-analysis of randomised
579		controlled trials. Atherosclerosis 2011; <b>216</b> : 7–16.
580		doi:10.1016/j.atherosclerosis.2011.02.044
581	40	Hanssen H, Boardman H, Deiseroth A, Moholdt T, Simonenko M, Kränkel N et
582		al. Personalized exercise prescription in the prevention and treatment of arterial
583		hypertension: a Consensus Document from the European Association of
584		Preventive Cardiology (EAPC) and the ESC Council on Hypertension. Eur J

585		Prev Cardiol 2022; <b>29</b> : 205–215. doi:10.1093/eurjpc/zwaa141
586	41	Almeida JPA de S, Bessa M, Lopes LTP, Gonçalves A, Roever L, Zanetti HR.
587		Isometric handgrip exercise training reduces resting systolic blood pressure but
588		does not interfere with diastolic blood pressure and heart rate variability in
589		hypertensive subjects: a systematic review and meta-analysis of randomized
590		clinical trials. <i>Hypertens Res</i> 2021; <b>44</b> : 1205–1212. doi:10.1038/s41440-021-
591		00681-7
592	42	McGowan CL, Visocchi A, Faulkner M, Verduyn R, Rakobowchuk M, Levy AS
593		et al. Isometric handgrip training improves local flow-mediated dilation in
594		medicated hypertensives. Eur J Appl Physiol 2007; 99: 227–234.
595		doi:10.1007/s00421-006-0337-z
596		
597		

#### FIGURE CAPTIONS

- FIGURE 1 Flow diagram of the current trial. N, number of participants; BMI, body mass index; COVID-19, coronavirus disease 2019; HIV, human immunodeficiency virus; BP,
- blood pressure; EXP, experimental session; MI, myocardial infarction; DRT, dynamic
- resistance training; IHT, isometric handgrip training; CRT, combined resistance training;
- 603 CON, control.

604

598

- FIGURE 2 Between-groups comparisons of changes (post-intervention preintervention) adjusted for pre-intervention values for the following variables: systolic blood pressure (SBP – panel a), diastolic blood pressure (DBP – panel b), systemic vascular resistance (SVR – panel c), cardiac output (CO – panel d), stroke volume (SV – panel f) and heart rate (HR – panel g). DRT, dynamic resistance training; IHT, isometric
- 610 handgrip training; CRT, combined resistance training; CON, control. Analysis: One-way
- 611 ANCOVA adjusted for pre-intervention values.

612

FIGURE 3 Between-groups comparisons of changes (post-intervention – pre-613 intervention) adjusted for pre-intervention values for the following variables: ratio 614 between low- and high-frequency bands of R-R interval variability (LF/HF<sub>R-R</sub> – panel a), 615 low-frequency band of systolic blood pressure variability (LF<sub>SBP</sub> – panel b), baroreflex 616 sensitivity (BRS – panel c), baseline vascular conductance (VC – panel d), baseline blood 617 flow (BF - panel e), peak blood flow (panel f), peak shear rate (panel g) and flow-618 mediated dilation (FMD - panel h). DRT, dynamic resistance training; IHT, isometric 619 620 handgrip training; CRT, combined resistance training; CON, control; nl, natural 621 logarithm. Analysis: One-way ANCOVA adjusted for pre-intervention values.

622

Table 1. Sample characteristics obtained at preliminary procedures

	DRT	IHT	CRT	CON	P
N	16	15	15	16	
Age (years old)	54±7	55±7	50±11	52±10	0.457
COVID-19 without hospitalization – n (%)	2 (13)	1 (7)	2 (13)	1 (6)	0.862
Physical activity levels (minutes / week)	41±43	57±55	35±41	57±50	0.476
Anthropometric					
Height (m)	1.75±0.06	1.74±0.08	1.77±0.09	1.76±0.06	0.617
Weight (kg)	91±12	86±15	91±18	88±11	0.642
BMI (kg/m²)	29.8±3.5	28.1±3.5	28.8±4.0	28.4±3.5	0.591
Blood pressure					
SBP (mmHg)	130±12	131±13	134±12	127±10	0.505
DBP (mmHg)	88 <u>±</u> 9	88±7	88±8	85±7	0.621
Pharmacological treatment					
Anti-hypertensive treatment duration (months)	118±91	105±87	95±78	114±80	0.883
Anti-hypertensive monotherapy – n (%)	9 (56)	8 (53)	6 (40)	9 (56)	0.810
Anti-hypertensive polytherapy – n (%)	7 (44)	7 (47)	9 (60)	7 (44)	0.810
ARB – n (%)	12 (75)	11 (73)	11 (73)	10 (63)	0.894
ACEi – n (%)	2 (13)	1 (7)	4 (27)	4 (25)	0.440
CCB – n (%)	5 (31)	5 (33)	7 (47)	5 (31)	0.812
DIU – n (%)	6 (38)	6 (40)	5 (33)	4 (25)	0.854
Statins – n (%)	1 (6)	3 (20)	3 (20)	1 (6)	0.510

Data: mean±standard deviation or number (percentage). DRT, dynamic resistance training; IHT, isometric handgrip training; CRT, combined resistance training; C, control; COVID-19, coronavirus disease 2019; BMI, body mass index; SBP, systolic blood pressure; DBP, diastolic blood pressure; ARB, angiotensin receptor blocker; ACEi, angiotensin-converting enzyme inhibitor; CCB, calcium channel blocker; DIU, diuretic. Physical activity levels were evaluated by the International Physical Activity Questionnaire. Analysis = One-way ANOVA for continuous data and Fisher's exact test for categorial data.

Table 2. Blood pressure and systemic hemodynamics parameters measured pre- and post-interventions in the 4 experimental groups: dynamic resistance training (DRT); isometric handgrip training (IRT); combined resistance training (CRT) and control (CON).

	DRT	IHT	CTR	CON	
SBP (mmHg	g)				P group = 0.511
PRE	125±11	128±13	128±12	127±14	P time = $0.000$
POST	119±12*	125±14	119±12*	129±16	P group x time = $0.003$
DBP (mmH	<b>g</b> )				P group = 0.764
PRE	85±10	87±8	87±6	86±9	P time = $0.642$
POST	84±10	86±10	84±8	89±10	P group x time = $0.091$
CO (L/min)					P group = $0.107$
PRE	$5.6 \pm 1.0$	$5.0\pm0.9$	5.1±1.0	$4.8\pm0.8$	P time = $0.158$
POST	$5.2 \pm 1.0$	5.3±1.1	$4.7\pm0.9$	$4.6\pm0.6$	P group x time = $0.201$
SVR (U)					P group = 0.133
PRE	18±4	21±4	21±4	21±4	P time = $0.449$
POST	19±5	20±5	21±4	23±3	P group x time = $0.306$
SV (mL)					P group = 0.995
PRE	82±17	77±15	81±24	83±17	P time = $0.101$
POST	76±17	83±16	76±16	77±15	P group x time = $0.066$
HR (bpm)					P group = 0.060
PRE	69±11	66±11	65±13	60±7	P time = $0.908$
POST	70±7	65±9	64±12	61±6	P group x time = $0.379$

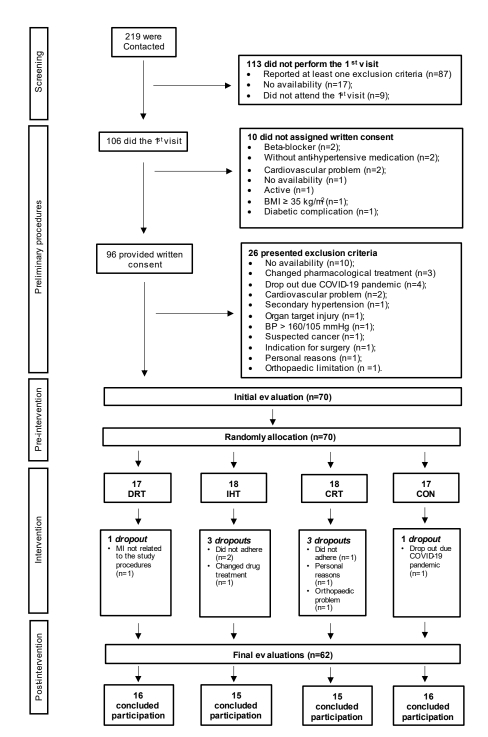
Data: mean  $\pm$  standard deviation. SBP, systolic blood pressure; DBP, diastolic blood pressure; CO, cardiac output; SVR, systemic vascular resistance; SV, stroke volume; HR, heart rate. Analysis: Two-way mixed ANOVA. \*Significantly different from pre-intervention (P<0.05).

Table 3. Vascular function and cardiovascular autonomic modulation parameters measured pre- and post-interventions in the 4 experimental groups: dynamic resistance training (DRT); isometric handgrip training (IRT); combined resistance training (CRT) and control (CON).

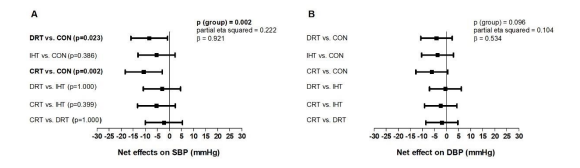
(IRT), combined resistar	DRT	IHT	CTR	CON				
VASCULAR FUNCTION CONTROL OF THE CON								
Baseline VC (mL.min	P group = 0.489							
PRE	$1.16\pm0.70$	$1.09\pm0.64$	$0.93\pm0.46$	$0.98 \pm 0.60$	P time = $0.137$			
POST	$1.34\pm0.63$	$1.19\pm0.73$	$1.10\pm0.50$	$1.03\pm0.55$	P group x time =			
					0.940			
Baseline BF (mL/min)	)				P group = 0.614			
PRE	110±59	$107 \pm 55$	$90\pm40$	96±57	P time = $0.205$			
POST	121±50	$114 \pm 70$	105±51	102±53	P group x time =			
					0.968			
/ - / - /								
Peak BF (mL/min)	774.277	501 : <b>3</b> 00	CEA : 221	929 - 259	P group = 0.161			
PRE	774±377	581±298	654±321	828±358	P time = 0.000			
POST	1067±461*	714±336	954±464*	786±223	P group x time =			
					0.007			
Peak shear rate (s <sup>-1</sup> )					P group = 0.161			
PRE	723±289	564±206	656±253	849±412	P time = 0.011			
POST	819±309*	688±266*	788±353*	851±314*	P group x time =			
1031	017±307	000±200	700-333	031±314	0.510			
					0.310			
FMD (%)					P group = $0.711$			
PRE	6.0±3.3	$6.6\pm4.2$	5.6±2.6	$6.2\pm4.0$	P time = $0.588$			
POST	$6.6\pm2.9$	$7.1\pm4.2$	6.2±2.2	5.5±2.7	P group x time =			
					0.642			
CARDIOVASCULAR	R AUTONOM	IIC MODUL	ATION					
nl LF/HF <sub>R-R</sub>					P group = 0.110			
PRE	$0.76\pm0.86$	$0.33\pm0.83$	$0.06\pm1.03$	$0.38\pm0.82$	P time = $0.065$			
POST	$0.45\pm1.05$	$0.48\pm0.51$	$-0.30\pm1.02$	$0.05\pm1.12$	P group x time =			
					0.320			
LLE (2)					D 0.210			
nl LF <sub>SBP</sub> (ms <sup>2</sup> ) PRE	1.07   1.09	1.70±1.27	1.63±1.05	1.52±1.01	P group = 0.310 P time = 0.692			
POST	1.97±1.08 2.10±1.00	1.70±1.27 1.55±1.03	1.03±1.05 1.22±1.38	1.52±1.01 1.69±1.00				
rusi	∠.10±1.00	1.33±1.03	1.22±1.38	1.09±1.00	P group x time = 0.596			
					0.370			
nl BRS					P group = $0.124$			
(mmHg/bpm)					1 510up - 0.121			
PRE	1.41±0.55	1.54±0.51	$1.92\pm0.47$	1.76±0.58	P time = 0.046			
POST	1.56±0.48*	1.90±0.45*	1.91±0.56*	1.79±0.76*	P group x time =			
					0.161			

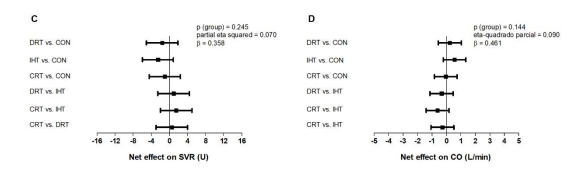
Data: mean $\pm$ standard deviation. DRT = dynamic resistance training; IRT = isometric handgrip training; CRT = combined resistance training; CON = control; BF = blood flow; VC = vascular conductance; FMD = flow-mediated dilation; nl = natural logarithm; LF/HF<sub>R-R</sub> = ratio between low- and high-frequency bands of R-R interval variability; LF<sub>SBP</sub> = low-frequency band of systolic blood pressure variability; BRS = baroreflex sensitivity. Analysis: Two-way mixed ANOVA. \*Significantly different from pre-intervention (P<0.05).

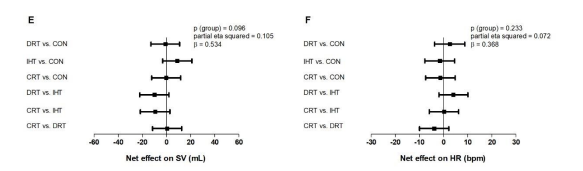
## **FIGURE 1**



### **FIGURE 2**







### **FIGURE 3**

