Acute antihypertensive effect of self-selected exercise intensity in older women with hypertension: a crossover trial

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Introduction

Hypertension (HTN) affects more than 1 billion people worldwide,1 and its prevalence among older adults is ~70%.2 Prevalence of HTN is higher among women than men.3 HTN is a major risk factor for coronary artery disease, heart failure, stroke, kidney disease, and mortality.1,2 Even in older adults with HTN who are treated with antihypertensive medication(s), high systolic blood pressure (BP) levels are associated with cardiovascular events and all-cause mortality.3 Therefore, efforts should be made to decrease BP in older adults with HTN, in order to attenuate the risk of cardiovascular events and mortality.4,5

Physical exercise is a cornerstone of nonpharmacological therapies for individuals with HTN, and is widely recommended by guidelines for HTN.6 Systematic
reviews and meta-analyses of randomized controlled trials (RCT) have reported a clear effect of aerobic exercise training for reducing both office and ambulatory BP levels, especially in individuals with HTN. A recent network meta-analysis of 391 RCTs confirmed these findings reporting that among individuals with HTN (ie, systolic BP ≥140 mmHg), the systolic BP-lowering effect of aerobic exercise was ~9 mmHg, which is similar to the effect of antihypertensive medications. Taken together, these data strongly endorse aerobic exercise as an antihypertensive therapy.

Exercise prescription in clinical guidelines for the management of HTN have focused on the frequency–intensity–time–and type principle, recommending that hypertensive patients should perform at least 30 minutes of moderate-intensity aerobic exercise (ie, 40%–59% of heart rate [HR] reserve) most days of the week. This “exercise dose” elicits both an acute and a chronic BP-lowering effect in individuals with HTN. The acute exercise BP-lowering effect, called postexercise hypotension (PEH), is characterized by reduced BP values in a postexercise period compared to preexercise and/or by reduced BP values in a postexercise period compared to a control condition. In older adults with HTN, 45 minutes of moderate-intensity aerobic exercise at 50% of maximal oxygen uptake (VO$_{2\max}$) can elicit ambulatory PEH that lasts up to 22 hours, including both awake and asleep periods. The maintenance of lower BP levels through ambulatory PEH has clinical implications, as these reductions can help prevent cardiovascular events. Therefore, studies characterizing ambulatory PEH, including the magnitude and duration of its presence, following a specific exercise approach may provide valuable information on its efficacy to improve acute BP control in older individuals with HTN.

Although ambulatory PEH is well documented in older individuals with HTN following formal exercise prescriptions (ie, HR- or VO$_2$-based) of moderate-intensity aerobic exercise under direct supervision with well-controlled settings, this is not a realistic scenario for most of these patients. Investigations about the acute effect of feasible exercise approaches on ambulatory BP in “real-world” settings are needed in older individuals with HTN. In this context, self-selected exercise (SSE)-intensity has been suggested as an alternative to prescribed exercise intensity in an effort to improve physical activity levels of inactive populations. SSE is characterized by individuals choosing their own exercise intensity according to their preferences.

Older individuals typically self-select an exercise intensity close to their ventilatory threshold, which is within the recommended range proposed for patients with HTN (ie, ≥40% of HR or VO$_2$ reserve). In addition, SSE-intensity has psychological advantages compared to prescribed exercise intensity, such as higher perceived autonomy, self-efficacy, and affect and lower perceived exertion. When SSE-intensity is performed in an outdoor environment, individuals display higher externally focused attention and affect, lower perceived exertion, and greater enjoyment compared to treadmill exercise.

Therefore, this study aimed to investigate the acute effect of an SSE-intensity session on ambulatory BP in inactive older women with HTN. In addition, HR, affect (ie, feeling of pleasure/displeasure), and perceived exertion during the SSE-intensity session were examined. We hypothesized that an SSE-intensity session would elicit an ambulatory BP-lowering effect, due to participants choosing an intensity within the recommended range proposed for patients with HTN (ie, ≥40% of HR reserve). Furthermore, we hypothesized that participants would report the SSE-intensity as light–moderate and pleasant.

**Methods**

**Trial design**

This randomized, controlled, crossover clinical trial was designed to assess the effect of a single SSE-intensity session on ambulatory BP in older women with HTN, and is reported in accordance with the CONSORT statement guidelines. Each participant performed the following procedures during a 4-week period: initial health screening and body composition assessment, assessment of physical activity levels with a pedometer over a 7-day period, maximal graded exercise test, familiarization with ambulatory BP monitoring procedures, familiarization with SSE-intensity session, a single SSE-intensity session and a control session separated by a 7- to 10-day period. The study was conducted from August 2016 to May 2017 at the Department of Physical Education, Federal University of Rio Grande do Norte, Brazil. The Institutional Ethics Committee approved the study (protocol 1526048). Participants were informed about all procedures of the study and provided written informed consent.

**Participants**

A total of 23 older women with HTN participated in this study. However, only 20 were included in the final
analysis (see Figure 1). Participants were recruited from the university cardiology outpatient service of the Federal University of Rio Grande do Norte, Brazil. Inclusion criteria were age 60–75 years, previous diagnosis of HTN according to the Brazilian guidelines on HTN, currently taking antihypertensive medication, no participation in exercise-training programs in last 6 months, "inactive", ie, <7,500 steps/day, no known cardiovascular, metabolic, renal, or respiratory disease, and no musculoskeletal conditions limiting ability to exercise at moderate and vigorous intensity (ie, knee arthritis or pain). Exclusion criteria were: pain or injury limiting ability to exercise during study procedures and changes in antihypertensive medication(s) during the study.

Procedures
Screening visit
Participants were screened at their first visit using the Physical Activity Readiness Questionnaire, medical history, medication use, and physical activity levels (short-version International Physical Activity Questionnaire). Clinical examination provided body weight (kg), height (m), resting HR (bpm), and resting BP (mmHg) measurements. Body mass index (BMI) was calculated as weight (kg) divided by height (m) squared. Resting HR was measured after 10 minutes of rest in a supine position in a calm and controlled room (24°C–26°C degrees) using an HR monitor (Polar Electro®, Kempele, Finland). Office BP was measured after the resting HR measurement in a seated position using an oscillometric device (Omrón® HEM-780-E, Kyoto, Japan).

Figure 1 Flow diagram of the study.
Abbreviation: ABPM, ambulatory blood pressure monitoring.
in triplicate with 1-minute intervals between each measure. The average value of the last two measures was recorded.\textsuperscript{23} Afterward, the participant’s body composition was assessed by dual-energy X-ray absorptiometry (GE Healthcare\textsuperscript{2} Lunar Prodigy Advance, Madison, EUA). At the end of the screening visit, participants were fitted with a wrist pedometer (Omron\textsuperscript{26} HJ-321, Kyoto, Japan) to confirm their physical activity status as assessed by the number of steps accumulated per day over a 1-week period.

**Physical activity level**

Pedometers were individually adjusted for the participants based on their stride length, weight, and height according to the manufacturer’s instructions. A total number of steps per day <7,500 was used as a criterion to classify the participants as “inactive”.\textsuperscript{24} Moreover, the number of steps per day performed at a moderate cadence (ie, 60 steps per minute)\textsuperscript{27} for at least 10 minutes continuously was recorded. This variable was captured to certify that the participants walked less than 30 minutes per day at a moderate cadence (ie, <1,800 steps) continuously or in bouts of at least 10 minutes. One week after the first visit, the participants returned to the laboratory to return the wrist pedometer and to perform a maximal graded exercise test.

**Maximal graded exercise test**

The participants performed a maximal graded exercise test on a motorized treadmill (Movement\textsuperscript{5} RT350, Pompeia, Brazil) to determine their maximal HR (HR\textsubscript{max}), as well as to exclude symptoms and electrocardiographical alterations suggestive of heart disease. Participants were continuously monitored with 12-lead electrocardiography by a cardiologist during the maximal graded exercise test. Following a 2-minute warm-up, an incremental ramp protocol was performed by the participants, starting at a speed of 2.5 km/h without grade and increasing 0.3 km/h and 1% grade every minute until volitional exhaustion. The participants’ rating of perceived exertion (RPE) was assessed every minute using Borg’s (6–20) RPE scale.\textsuperscript{28} BP was measured every minute and immediately following volitional exhaustion. The graded exercise test was considered maximal when the participant reached an RPE >17 and HR within ±10 bpm of age-predicted HR\textsubscript{max} (220-age).\textsuperscript{29} For participants using a β-blocker, only RPE was used as a criterion of volitional exhaustion. Participants performed the maximal graded exercise test under their regular antihypertensive medication(s). After the exercise test, participants took a shower and had an ambulatory BP monitoring device (Cardios\textsuperscript{®} CardioMapa, São Paulo, Brazil) placed on their nondominant arm to familiarize them with 24-hour ambulatory BP monitoring.

**Ambulatory blood pressure monitoring**

Ambulatory BP was assessed three times during the study: during the familiarization session, following the SSE-intensity session, and following the control session. The familiarization session was conducted in an effort to avoid an alert reaction, which can happen when wearing an ambulatory BP monitoring device for the first time. This response is more common in women than in men.\textsuperscript{30} This familiarization procedure increased the validity of the ambulatory BP response measurement following the SSE-intensity and control sessions. After each experimental session, participants were instructed to take a shower before being fit with the ambulatory BP device (Cardios\textsuperscript{®} CardioMapa, São Paulo, Brazil) on their nondominant arm according to the Brazilian guidelines for ambulatory BP monitoring.\textsuperscript{31} The device was programmed to measure BP every 15 minutes during awake and every 30 minutes during asleep periods. A minimum of 16 and eight BP measurements during awake and asleep periods, respectively, had to be successfully recorded to be included in the final analysis.\textsuperscript{31} Participants came back to the laboratory the following morning after each session so that the BP information could be downloaded to a computer. Data were recorded during a 20-hour period with the awake and asleep periods defined according to each participant. The average values for each period were considered for data analysis. In addition, BP load was calculated during the 20-hour period. This index is defined as the percentage of readings that exceeds the normal BP values in both awake and asleep periods. In the awake period, systolic BP ≥135 and diastolic BP ≥85 mmHg were considered abnormal. In the asleep period, systolic BP ≥120 and diastolic BP ≥70 mmHg were considered abnormal.\textsuperscript{31} All procedures related to ambulatory BP monitoring were explained to the participants at the screening visit, the familiarization session, and immediately before the experimental sessions, based on the Brazilian guidelines for ambulatory BP monitoring.\textsuperscript{31}

**Experimental sessions**

The SSE-intensity and control sessions were performed in a randomized order, separated by 7–10 days, on a standard outdoor 400 m track. All procedures were performed in the
morning (7–8 am). Participants were instructed to abstain from moderate-vigorous physical activity and caffeinated products on the days of the experimental sessions. Additionally, participants were instructed to abstain from alcohol, maintain their eating habits and sleeping pattern during the 24-hour period prior to the experimental sessions. Participants’ resting BP was measured before the experimental sessions following the same procedures as that of the screening visit. They were then fitted with a HR monitor (Polar Electro®, Kempele, Finland) to monitor HR responses during both sessions. For the control session, participants remained seated for 40 minutes at the same location where the exercise session was performed. Before the SSE-intensity session, participants were instructed to regulate their own pace using the following statement: “We would like that you exercise for 30 minutes at a pace of your own choosing. You can freely change the pace at any time during the exercise”. Following a stretching routine, participants warmed up for 5 minutes to adjust their pace before starting the 30 minutes of SSE-intensity session. After, participants cooled down for 5 minutes walking at a slow pace. Whole-body perceived exertion using Borg’s (6–20) RPE scale was reported by participants every 10 minutes. They were instructed on use of the RPE scale and Feeling Scale at the screening visit, the familiarization visit and immediately before the SSE-intensity session. RPE was defined as subjective intensity of effort, strain, and/or fatigue felt during exercise. Low and high perceptual anchors for Borg’s RPE scale were established during the maximal graded exercise test. A rating of 6 (low anchor, “very, very light”) was assigned to the lowest exercise intensity, while a rating of 20 (high anchor, “very, very hard”) was assigned to the highest exercise intensity. The Feeling Scale is an 11-point bipolar scale ranging from +5 to −5 recommended by the American College of Sports Medicine to assess affect (ie, feelings of pleasure/displeasure) during exercise. This scale presents verbal anchors: −5= very bad; −3= bad; −1= fairly bad; 0= neutral; +1 fairly good; +3= good; and +5= very good. The participants received standard instructions explaining the Feeling Scale according to Hardy and Rejeski.

Following both experimental sessions, participants were instructed to shower before being fitted with the ambulatory BP device, as previously described. In addition, they were fitted with wrist pedometer to monitor the number of steps they accumulated over the ambulatory BP monitoring period. When they returned in the following day to remove the ambulatory BP monitoring device, they filled out a modified Pittsburgh Sleep Quality Index (PSQI) to assess the quality and pattern of sleep in the night prior. Two trained investigators (IBBC and DS) conducted the experimental sessions.

Sample size and randomization
Sample size was calculated using the preliminary data of the primary outcome of the present study (ie, awake systolic ambulatory BP; results obtained from 10 participants); ie, reduction of 3.6 mmHg (95% CI, 1.0 to 6.2 mmHg) between the SSE-intensity (117.7±8.4 mmHg) and control (121.3±7.9 mmHg) sessions. Based on this preliminary result, 22 participants were needed to power the study, considering 1-β of 80% and an alpha of 5%. After all baseline assessments, a computer-based simple randomization (www.graphpad.com) was carried out to determine the order of the experimental sessions. Due to logistics, only the participants were blinded to the order of the experimental sessions.

Statistical analysis
Data normality was tested using the Shapiro–Wilk test, skewness and kurtosis (score: z = −1.96 to +1.96). Results are expressed as mean ± SD for the parametric data and median and 25th–75th percentile for the nonparametric data. A paired t-test was used to compare the mean values of resting BP, BP load, number of steps, sleep quality score, and ambulatory BP between the sessions. Cohen’s d was used to determine the effect size (ES) of the mean difference. The Hopkins’s criteria to interpret the magnitude of the ES were followed: <0.2 is trivial, 0.2–0.5 is small, 0.6–1.1 is moderate, 1.2–1.9 is large, and 2.0–4.0 is very large. A generalized estimating equation (GEE) was used to assess the condition (2 factors: SSE-intensity session vs control session) by time (3 factors: average values of BP during 1 to 6-hour, 7 to 13-hour, and 14 to 20-hour periods) interaction on the ambulatory BP, adjusted for resting BP values. The GEE model for each outcome was based on the goodness of fit. The normality of the residuals was verified by normal Q-Q plot. For all analyses, a two-tailed p<0.05 was considered statistically significant. Data analysis was performed using IBM SPSS Statistics version 25 (IBM®, Chicago, USA).

Results
Table 1 outlines the characteristics of the participants. The mean length of time of since menopause and HTN diagnosis
was 16.9±7 and 10.8±9.1 years, respectively. The temperature (28 °C, 95% CI 27–28 °C) and humidity (70%, 95% CI 70–80%) of the outdoor track was similar between the SSE-intensity and control sessions (P>0.05). Resting systolic (117.2±11.8 vs 119.1±12.1 mmHg; P=0.30) and diastolic (70.0±7.8 vs 71.1±8.4 mmHg; P=0.38) BP values immediately before the SSE-intensity and control sessions were similar. In addition, the participants accumulated the same number of steps (5,835±2,627 vs 5,880±1,969 steps/day; P=0.91) in the day of the SSE-intensity and control sessions and experienced similar sleep quality (3.5, 95% CI 1–6 vs 4, 95% CI 1–5.8 arbitrary units; P=0.76) in the day prior to the sessions.

Table 2 shows the results of the SSE-intensity session. On average, the participants exercised at 5.1±1.1 km/h and spent ~90% of the exercise time at moderate-to-vigorous intensity. The participants reported the SSE-intensity session as light to moderate (ie, RPE between 10–12) and pleasant (ie, basic affect between 3–4). There were no adverse events during the exercise sessions.

Table 3 shows the results of the ambulatory BP monitoring following the SSE-intensity and control sessions. Average values of systolic BP in the total 20-hour and awake periods were lower following the SSE-intensity session compared to the control session.
(P<0.05). No differences were observed in the average values of systolic BP during the asleep period and diastolic BP during the 20-hour, awake, and asleep periods between the SSE-intensity and control sessions (P>0.05). The systolic BP load was lower in the SSE-intensity session compared to the control session (P<0.05). No differences were found for diastolic BP load between the SSE-intensity and control sessions (P>0.05).

Table 3 Ambulatory blood pressure monitoring following the self-selected exercise-intensity and control sessions in inactive older women with hypertension

<table>
<thead>
<tr>
<th>20-hour</th>
<th>Control</th>
<th>SSE</th>
<th>Δ (95% CI)</th>
<th>P</th>
<th>ES</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBP (mmHg)</td>
<td>119.2±6.9</td>
<td>115.8±7.0</td>
<td>−3.4 (−5.9 to −0.9)</td>
<td>0.010</td>
<td>0.6</td>
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<tr>
<td>DBP (mmHg)</td>
<td>67.0±6.1</td>
<td>66.0±6.8</td>
<td>−1.0 (−2.8 to 0.7)</td>
<td>0.219</td>
<td>0.3</td>
</tr>
<tr>
<td><strong>Awake</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>121.6±7.5</td>
<td>117.6±8.0</td>
<td>−4.0 (−6.4 to −1.6)</td>
<td>0.003</td>
<td>0.8</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>69.7±6.9</td>
<td>68.1±8.3</td>
<td>−1.6 (−3.5 to 0.3)</td>
<td>0.101</td>
<td>0.4</td>
</tr>
<tr>
<td><strong>Asleep</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>114.7±7.8</td>
<td>111.8±6.8</td>
<td>−2.9 (−6.6 to 0.8)</td>
<td>0.114</td>
<td>0.4</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>61.6±6.1</td>
<td>61.3±6.4</td>
<td>−0.3 (−3.1 to 2.4)</td>
<td>0.812</td>
<td>0.1</td>
</tr>
<tr>
<td><strong>BP load</strong></td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>SBP (%)</td>
<td>17.9 (6.7–28.7)</td>
<td>11.7 (2.6–19.9)</td>
<td>−6.5 (−12.0 to −1.1)</td>
<td>0.014</td>
<td>0.6</td>
</tr>
<tr>
<td>DBP (%)</td>
<td>10.3 (1.7–19.9)</td>
<td>7.2 (1.7–11.9)</td>
<td>−0.6 (−5.1 to 3.9)</td>
<td>0.327</td>
<td>0.1</td>
</tr>
</tbody>
</table>

Notes: Values expressed as mean ± SD or median (25th-75th percentile). Bold values indicate significant difference (P<0.05).
Abbreviations: SBP, systolic blood pressure; DBP, diastolic blood pressure; SSE, self-selected exercise ES, effect size.

Figure 2 shows the ambulatory BP behavior over the 20-hour period following the SSE-intensity and control sessions. There was significant condition by time interaction for systolic BP (W2) =7.0; P=0.030. Post hoc analysis revealed a decrease of 6 mmHg (95% CI 2.7–9.3 mmHg) in the first 6-hour period following the SSE-intensity session compared to the control session (P=0.004; Figure 2, panel A). For DBP, there was no significant condition by time interaction – W(2) =2.8; P=0.252 (Figure 2, panel B). Table S1 shows the comparison of the absolute values of ambulatory BP over the 20-hour period (ie, 1–6 hours, 7–13 hours, and 14–20 hours).

Discussion
To the best of our knowledge, this is the first study to investigate the effect of a single SSE-intensity session on ambulatory BP in inactive older women with HTN. The main findings of this study were: (i) the SSE-intensity session elicited a clinically meaningful reduction of 6 mmHg in the ambulatory systolic BP in the first 6-hour period postexercise. This contributed to the reduction in systolic BP seen during the awake (4 mmHg) and the total 20-hour periods (3.4 mmHg) when compared to the control session; (ii) the participants met the recommended intensity for aerobic exercise (ie, ≥40% of HR reserve) according to the guidelines for HTN and reported the SSE-intensity session as light and pleasant.

Our findings regarding the acute effect of the SSE-intensity session on ambulatory BP are in accordance with previous studies that have observed a greater magnitude of ambulatory PEH in the first hours following a HR- or VO2-based aerobic exercise session in individuals with HTN. Santos et al have observed a greater systolic ambulatory PEH in the first 5 hours following 45 minutes of both a light (50% of HRmax; 7.7±2.4 mmHg) and moderate (75% of HRmax; 9.4±2.8 mmHg) aerobic exercise session in middle-aged individuals with resistant HTN over 19 hours of ambulatory BP monitoring. Ash et al have demonstrated a 10-hour systolic (5.6±1.8 mmHg) and diastolic (3.6±1.3 mmHg) BP-lowering effect only in the awake period following 30 minutes of an aerobic exercise session at 60% of VO2peak in middle-aged individuals with pre- to established stage 1 HTN over 19 hours of ambulatory BP monitoring. Our results suggest that the systolic ambulatory BP-lowering effect following the SSE-intensity session in the first 6 hours improved the acute BP control of the participants; ie, reduced the average values of systolic BP during the awake and 20-hour periods and the systolic BP load.

All participants in the study took antihypertensive medication(s) and presented controlled office (<140/90
mmHg) and ambulatory (<130/80 mmHg) BP values. Recent data\textsuperscript{4,40} have shown that more intensive systolic BP control (ie, <120 mmHg) reduces the risk of fatal and non-fatal cardiovascular events and all-cause mortality compared with traditional target of systolic BP (ie, <140 mmHg). Despite a well-controlled BP status of the participants, the SSE-intensity session elicited a systolic ambulatory PEH that lasted up to 6 hours. Therefore, the ambulatory PEH observed in our study may have clinical significance in improving the acute BP control of these inactive older women with diagnosed HTN. It should be noted that there is a dose-response relationship between resting BP levels and the magnitude and duration of PEH.\textsuperscript{11} Thus, it seems reasonable to think that a SSE-intensity session could elicit a greater and longer ambulatory PEH in older adults with HTN who present higher BP levels than those included in the present study. Future studies should address this question.

A control condition without a previous exercise session is crucial for the assessment of PEH occurrence as it is supposed to only take sources of BP variation unrelated to exercise and the normal circadian variation of BP into account.\textsuperscript{39,41} In addition, the ambulatory BP monitoring during a control condition demonstrates the cardiovascular load during daily activities, including sedentary behaviors in different positions (ie, seated and supine), posture changes with orthostatic stress, walking and other physical activities. Thus, the ambulatory systolic BP-lowering effect of ~4 mmHg observed in the awake and 20-hour periods and the reduction of ~6% in the systolic BP load

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**Figure 2** Ambulatory BP monitoring hour-to-hour following the self-selected exercise-intensity (SSE) and control sessions in inactive older women with hypertension. **Notes:** (A, B) represent systolic and diastolic blood pressure responses, respectively. Values expressed as mean ± SD. *P*<0.05 (difference from control session compared to the same period).

**Abbreviations:** BP, blood pressure; SSE, self-selected exercise.
following the SSE-intensity session, as compared to the control condition, may play a role in attenuating the overload on the cardiovascular system imposed by sedentary behaviors and light-intensity activity (ie, <4 METs). It should be noted that older adults spend more than 95% of their awake time in sedentary behaviors and light physical activities.42

Our data support the idea that a SSE-intensity session elicits a systolic ambulatory PEH, which was similar to previous studies that have used traditional HR- or VO2-based exercise prescription approaches.11 Overall, the studies have reported that middle-aged and older individuals self-select an exercise intensity close to ventilatory threshold16–18 and within the recommended range proposed for apparently healthy individuals and patients with HTN (ie moderate to vigorous; ≥40% of HR or VO2 reserve).6 In our study, the participants exercised at a brisk walking pace (5.1±1.1 km/h; ~105 steps/minute), spent ~90% of the exercise session at moderate to vigorous intensity, and presented a mean HR response equivalent to 58.8% of HR reserve. Therefore, it seems clear that the acute ambulatory BP-lowering effect observed in our study is associated with the exercise intensity level achieved by the participants during the SSE-intensity session. This finding is consistent with a previous study that observed an ambulatory PEH following a VO2-based moderate intensity aerobic exercise session in older individuals with HTN.15

Despite the participants having presented HR responses consistent with moderate to vigorous exercise intensity, they reported a RPE consistent with light physical activity (ie, RPE from 10 to 11) during the SSE-intensity session. Exercise at a self-selected pace elicits a high perceived autonomy and pleasant feelings, which may have contributed to attenuate a perceived exertion consistent with a moderate-intensity aerobic exercise.20 Moreover, the various stimuli present in an outdoor setting seems to reduce internal attention focus (eg, increased HR, breath, pain) and increase more externally focused attention.21 Thus, individuals can report a lower RPE than expected for a moderate to vigorous exercise intensity. This aspect seems to be important for older adults, especially those with a low self-efficacy for exercise, as this is an important factor associated with lack of engagement in regular physical activity by this population.43

The participants reported the SSE-intensity session as pleasant. This can be partially explained by the magnitude of the metabolic stress imposed on the participants during the exercise. Overall, the individuals self-select an exercise intensity close to ventilatory threshold.16–18 Lind et al18 reported that SSE-intensity is characterized by a stable and positive affective state, which is often maintained close to, but below, the ventilatory threshold. Therefore, SSE-intensity seems to be an interesting approach for increasing physical activity levels in inactive older women with HTN. In this context, it is important to highlight that even among hypertensive patients taking antihypertensive medication(s) and with well-controlled BP levels, those who are physically active have a lower cardiovascular risk than their inactive peers.44

From a practical perspective, SSE-intensity seems to be a simple exercise approach for improving acute BP control of inactive older adults with HTN through the maintenance of systolic ambulatory PEH. In addition, it seems more feasible than the traditional HR- or VO2-based aerobic exercise prescription in community-based settings where there may be no formal supervision. Therefore, it seems reasonable that clinicians and healthcare providers consider providing instructions for SSE-intensity for older adults with HTN, especially for those who are inactive and unfit.

Despite our novel findings, some limitations of this study must be mentioned. First, our sample was composed of medicated hypertensive older women with well-controlled office and ambulatory BP status. Our findings may not be generalizable to other hypertensive populations. Second, only HR responses were monitored as a physiological marker of the exercise intensity, therefore, it was not possible to characterize the SSE-intensity session according to VO2 and metabolic domains (below, at or above the ventilatory threshold). Third, despite the instruction for the participants to abstain from alcohol and to maintain their eating habits in the 24-hour period before the experimental sessions, the compliance with this instruction was not assessed. However, it should be noted that important confounding factors were monitored in order to improve the internal validity of this study, such as the familiarization with the ambulatory BP monitoring device, the measurement of the quality and patterns of sleep in the night prior to the experimental sessions, and the assessment of the physical activity levels by pedometer over the 20 hours of ambulatory BP monitoring following the experimental sessions.

Conclusion
A single SSE-intensity session lasting 30 minutes elicited a reduction in ambulatory systolic BP in inactive older...
women with HTN in the first 6-hour period post exercise, which contributed to the reduction of systolic BP during the awake and 20-hour periods. This SSE-intensity approach met the recommended intensity for aerobic exercise (ie, moderate to vigorous) according to the guidelines for HTN. In addition, it was perceived as light and pleasant. Therefore, SSE-intensity should be considered as a simple alternative to encourage inactive older women with HTN become more physically active. Future studies should investigate whether SSE-intensity sessions can elicit a chronic ambulatory BP-lowering effect and improve health-related physical fitness in older adults with HTN.

Acknowledgments
Master’s degree scholarship granted by Coordination for the Improvement of Higher Education Personnel (CAPES, Brazil) to IBBC. All procedures were approved by the Ethics Committee of the Federal University of Rio Grande do Norte (protocol 1526048) and were carried out in line with the Declaration of Helsinki. The authors thank the cardiologist Bruno Erick de Barros Lucena to conduct the graded exercise test with the participants of the study.

Disclosure
The authors report no conflicts of interest in this work.

References

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### Supplementary material

**Table S1** Ambulatory blood pressure monitoring per period following self-selected exercise-intensity and control sessions in inactive older women with hypertension

<table>
<thead>
<tr>
<th></th>
<th>Awake (mmHg)</th>
<th></th>
<th>Asleep (mmHg)</th>
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<tr>
<td></td>
<td>1–6 hours</td>
<td>7–13 hours</td>
<td>14–20 hours</td>
<td></td>
</tr>
<tr>
<td><strong>SBP</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>121.4±7.5</td>
<td>121.6±7.9</td>
<td>115.4±7.5</td>
<td></td>
</tr>
<tr>
<td>SSE</td>
<td>115.4±7.5*</td>
<td>119.5±9.4</td>
<td>112.5±6.8</td>
<td></td>
</tr>
<tr>
<td>∆ (95% CI)</td>
<td>6.0 (2.7–9.3)</td>
<td>2.1 (−0.5 to 4.7)</td>
<td>2.5 (−1.1 to 6.1)</td>
<td></td>
</tr>
<tr>
<td><strong>DBP</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>69.6±7.3</td>
<td>69.6±6.7</td>
<td>62.2±5.8</td>
<td></td>
</tr>
<tr>
<td>SSE</td>
<td>67.1±8.2</td>
<td>68.9±8.7</td>
<td>62.0±6.3</td>
<td></td>
</tr>
<tr>
<td>∆ (95% CI)</td>
<td>2.5 (0.2–4.7)</td>
<td>0.7 (−1.4 to 2.8)</td>
<td>0.1 (−2.4 to 2.6)</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:** Values expressed as means ± SD and 95% CIs. *P*<0.05 compared to the same period of the control session.

**Abbreviations:** SBP, systolic blood pressure; DBP, diastolic blood pressure; SSE, self-selected exercise.