Impact of exercise training on cardiotoxicity and cardiac health outcomes in women with breast cancer undergoing anthracycline-containing chemotherapy: a study protocol for a randomized controlled trial.

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Abstract

Background: Anthracyclines are chemotherapeutic agents frequently used in breast cancer (BC) treatment. Although improving disease-free and overall survival, the use of anthracyclines is associated with a cumulative risk of cardiac toxicity. Preventive strategies to optimize cardiac health are needed and exercise is proposed as a potential non-pharmacological approach for counteracting of anthracycline-related cardiotoxicity (ARC). Most of the data on the effects of exercise to reduce ACT is from animal studies, with only a few studies with a limited number of patients indicating beneficial effects. To better understand the effectiveness of exercise in mitigation of ARC, clinical, real-world trials claim for a larger sample size and more accurate and valuable clinical biomarkers. In this study, we intend to include a large sample size and investigate cardiac function throughout serial measures of biomarkers and imaging techniques. Methods: This protocol describes a two-armed prospective randomized controlled trial that will explore the cardioprotective effect of a structured exercise program in women with BC undergoing anthracycline-containing chemotherapy (ACT). Ninety adult women with early BC and with a therapeutic decision to receive ACT will be randomly assigned (1:1) to an intervention group or a control group. Patients allocated to the intervention group will perform a supervised exercise program three times per week, consisting of a combination of aerobic and resistance training with progressive intensity and volume, during the time period they received ACT. The control group will receive standard BC care. Primary outcomes related to cardiac (dys)function will be circulating N-terminal pro-brain natriuretic peptide (NT-proBNP) levels, resting left ventricular (LV) longitudinal strain, and resting LV ejection fraction. Secondary outcomes will include the assessment of resting blood pressure, resting heart rate (HR), resting HR variability (HRV), recovery HR, physical function outcomes, self-reported physical activity level, health-related quality of life and fatigue.
Data will be obtained at baseline (t0), after the end of anthracycline-treatment (t2), and three months after t2 (t3). Additionally, N-terminal pro-brain natriuretic peptide levels will be measured 1-24 hours prior to each anthracycline-treatment cycle (t1). Discussion: The implementation of the present study design, using novel clinical biomarkers, will determine the effect of structured exercise interventions at mitigating ARC, with the overall aim of finding means to further improve BC care.

Trial registration: ISRCTN, ISRCTN32617901. Registered on 24 October 2018. Last updated on 11 January 2019, https://doi.org/10.1186/ISRCTN32617901.

Background

Over the last three decades the epidemiology of breast cancer (BC) has been marked by the clear increase in survival rates (1). The accessibility of screening and the discovery of new therapeutic options are among some relevant factors related to the improved management of cancer. However, despite their undeniable clinical importance, anti-cancer treatments are also associated with a frequent induction of side effects. Among those, cardiotoxicity emerges as a major challenge limiting treatment options (2) and contributing to morbidity and mortality in this patient population (3-5).

In the BC setting, cardiotoxicity is typically associated with exposure to traditional cytotoxic therapies, and particularly to the use of anthracyclines (6). Anthracyclines are important and effective chemotherapeutic agents, frequently administered in curative and palliative regimens for BC, although their clinical use is limited by cardiac dysfunction, usually seen as cardiotoxicity (7). The American Society of Echocardiography and European Association of Cardiovascular Imaging define cardiotoxicity as a decrease in left ventricular ejection fraction (LVEF) of >10 % to a value of <53 % (8). However, it has been proposed that anthracycline-related cardiotoxicity (ARC) is a continuous (9) and dose-dependent phenomenon (10) that starts with an acute myocardial damage (11) which
can be detected by the elevation of circulating cardiac biomarkers (12) and by the impairment in left ventricular (LV) longitudinal strain (13), in turn preceding the commonly reported LV ejection fraction progressive decline (9). Particularly when underestimated, not or insufficiently prevented and untreated, ARC may ultimately lead to overt heart failure (9).

Unfortunately, and in addition to the manifestations associated with treatment with anthracyclines, BC survivors often exhibit a phenotype characterized by the presence of risk factors for cardiovascular disease (CVD) development including: advanced age, obesity, prior CVD, poor cardiorespiratory fitness, and inappropriate lifestyles (smoking, alcoholism and a sedentary lifestyle) (14, 15). It is therefore not surprising that when compared to healthy individuals of a similar age, this patient population have higher prevalence and mortality of CVD (4). Considering the high risk of cardiac dysfunction, the study of preventive strategies are an emerging and unmet need, as they may act to mitigate the cardiac damage associated with the use of cardiotoxic agents and could be associated with improvements in overall cardiovascular health.

In adults with CVD without cancer, exercise training is recognised as an important approach in cardiac rehabilitation (16). Particularly, two meta-analyses have shown favourable effects of exercise on attenuation of N-terminal pro-brain natriuretic peptide (NT-proBNP) levels (17) and mitigation of LV remodeling (18) in patients with heart failure. Also, in the oncological setting, it has become clear that exercise is a safe and effective supportive therapy in the management of several treatment-related side effects and in improving overall physical fitness (19, 20). Furthermore, the Task Force for cancer treatments and cardiovascular toxicity of the European Society of Cardiology suggests in a recent position paper the possible utility of aerobic exercise as a promising strategy to attenuate ARC (21). However, this position remains to be complementarily analysed as the
evidence so far is mainly from studies in animal models (22, 23), with only a few studies with a limited number of patients have evaluated the cardioprotective role of exercise in cardiac function of women with breast cancer receiving cardiotoxic treatments. Further clinical studies involving a large sample size and accurate clinical biomarkers are thus warranted to better understand the effectiveness of exercise in mitigation of ARC.

Hypothesis
The primary aim of this study is to ascertain whether a structured exercise program mitigates ARC, measured by the level change of circulating biomarkers (NT-proBNP) and cardiac (dys)function endpoints (LV global longitudinal strain and LV ejection fraction). The secondary aim is to evaluate the effectiveness of the intervention in the regulation of some cardiac health parameters: resting blood pressure, resting heart rate (HR), resting HR (HRV) and recovery HR. As exploratory objectives, we will assess physical function (cardiorespiratory capacity, upper limb strength and lower limb functionality), self-reported physical activity level, health related to quality of life (HR-QOL) and fatigue. We hypothesize that exercise may limit the degradation of cardiac function and structure and benefit cardiac outcomes. We also believe that patients in the intervention group will improve overall physical fitness, HR-QOL, decrease their perception of fatigue and increase physical activity levels.

Methods/design

Study Design
This is a study protocol for a two-armed prospective randomized controlled trial that will explore the cardioprotective effect of a structured exercise program, compared to standard care, in adult women undergoing anthracycline-containing chemotherapy (AC-CT) for early BC. The study design and protocol adhere to the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) guidelines (Additional file 1). The study
design is outlined in Figure 1.

**Ethical approval**

This study will be conducted in compliance with the Declaration of Helsinki Ethical Principles (1975) and it received approval by the Ethics Committee of the Centro Hospitalar de Vila Nova de Gaia/Espinho (CHVNG/E; Vila Nova de Gaia, Portugal) (reference number: 145/2018-1). The study is registered in the International Standard Randomised Controlled Trial Number (ISRCTN32617901). Any protocol amendments will be submitted to the CHVNG/E for ethical approval and updated on the ISRCTN.

**Participant recruitment**

We intend to recruit 90 adult women with early invasive BC, scheduled to receive AC-CT and followed-up in the Medical Oncology Department of the CHVNG/E. Participants will be recruited considering the eligibility criteria presented in Table 1. Recruitment will take place in two distinct phases. In a first instance, potential participants will be identified in the multidisciplinary consultation involving medical oncologists, surgeons and radioncologists. After this preliminary phase, the eligibility of each patient will be confirmed by the oncologist during a medical consultation. The oncologist will present the study to the patients considered eligible, explaining, offering the inclusion and providing written informed consent. Written informed consent will be obtained from all patients and they will be informed that they are under no obligation to participate and they may withdraw their consent at any time. The withdrawal from the study or non-participation, will have no consequences for medical follow-up and care. Where possible, the reasons for withdrawal from the study will be recorded. All the participants will be followed from the acceptance period (t0) and after 3 months of the end of the AC-CT (t3). If recruitment is not achieving the target sample size, we will extend the recruitment for additional hospitals.
Randomization

After confirmation of eligibility and baseline assessments (t0), patients will be randomized through an Internet software (www.sealedenvelope.com), with a 1:1 ratio between a supervised exercise group (intervention group) and a usual care group (control group), using a permutated block design with random block sizes (4, 6, 8) with stratification by two dichotomous variables, known as risk factors for ARC:

Age (Under/50 years or older).

Receive anti-HER2 therapy (Yes/No).

This process will be performed by an external individual who is blinded to the study and who will place the sequence in a numbered, opaque, sealed envelope. The allocation of participants will then be reported to an oncologist (AJ) who will subsequently inform the patients about the assignment group.

Study arms

Intervention group

Patients allocated to the intervention group will perform a supervised exercise program specifically developed for BC patients, based in the guidelines of the American College of Sports Medicine (27) and in a close cooperation between physical sports researchers (PA, DE, AA) and medical staff (oncologists, surgeons, radiologists, physiatrist, and physiotherapist) of the CHVNG/E. The exercise program comprises 3 weekly sessions guided in small groups (<5 patients) in an appropriately equipped room of the CHVNG/E, supervised by the main author (PA) and a physiotherapist. Each session will involve an initial warm-up (5 min), followed by resistance and aerobic training (60 min), and ending with a cooldown phase (5 min). The program will be started after 1-2 days of the first AC-CT session and will be conducted over the respective treatment of each patient. It should be noted that the proposed exercise intervention will never be intended to replace or
interfere with the current standard BC care.

**Aerobic training:** It will include the combination of treadmill, stationary bike, and stepping. This phase will be monitored through HR (each participant will wear a heart rate monitor during exercise training sessions) and RPE measure by a 0-10 point modified Borg Scale (minimal effort = 0; maximum effort = 10) (29). During the first two weeks, the participants will perform 20 minutes (divided equally among the three exercise modes) of aerobic training in a light intensity [<50% of measured HR reserve (based on maximum HR reached in the cardiorespiratory test), reporting 2-4 ('easy' to 'somewhat easy') on modified Borg scale]. After this period, 3 minutes will be added every two weeks until a volume of 30 minutes of aerobic training is reached. At this stage, participants will be encouraged to perform moderate-to-high intensity training [65%-80% of measured HR reserve, reporting 5-8 ('somewhat hard' to 'hard') on modified Borg scale] until the end of the intervention.

Participants will be reminded weekly (through email and phone) of their exercise training schedule and the importance of adherence to achieve the established objectives.

**Resistance training:** It will include upper body (shoulder press, chest press, lat pulldown, biceps curls, and triceps extension) and lower body (squat, calf raise, leg press, leg extension, and leg curl) weight-training exercises. All the exercises will be performed at the maximum possible joint range of motion, using resistance machines and free weights. Rating perceived exertion (RPE) will be measured using a 0-10-point OMNI-Resistance Exercise Scale (OMNI-RES, minimal effort = 0; maximum effort = 10) (28). During the first week, participants will perform 2 sets and 10 repetitions of each exercise without additional resistance or with the lowest available [reporting 2-4 ('easy' to 'somewhat easy') on the OMNI-RES]. After this phase, if no adverse events or symptoms were reported for a specific exercise, resistance will be added so that each participant could be
able to perform 3 sets with 12 maximal repetitions (12-RM) of each exercise. When the participants can complete 3 sets and more than 12-RM at the set weight in 3 consecutive sessions, then the resistance will be increased between 5%-10%.

Usual care group

Patients allocated to the usual care group will receive the standard BC care. The CG will not receive any specific advice regarding physical activity and will not be asked to be inactive. In compensation for the participation in this study, it will be offered the possibility of performing the same exercise program after the final assessments are completed.

Study assessments

The schedule of the study outcome assessments is outlined in Figure 2. Study assessments will be scheduled upfront and participants will be regularly reminded (through email and phone) to ensure a complete follow-up. Primary, secondary and exploratory outcomes will be measured in all participants at three different moments:

- t₀ (baseline assessments): Between 0-14 days prior to the first chemotherapy session.
- t₂ (post-treatment assessments): Between 1-5 days after the end of AC-CT.
- t₃ (follow-up assessments): After 3 months of t₂.

In addition, for analysis of circulating NT-proBNP, blood samples will be collected between 1-24 hours before each AC-CT cycle (t₁: during-treatment assessments). Patients will be instructed and remembered to avoid drinking alcoholic and caffeine-containing beverages, to abstain from smoking for 12 hours prior and to avoid vigorous physical activities 24 hours prior to all examinations.

Study outcomes

Primary outcomes

Circulating NT-proBNP levels
Resting LV global longitudinal strain

Resting LV ejection fraction

**Secondary outcomes**

Cardiac health outcomes:

Resting blood pressure

Resting HR

Resting HRV

Recovery HR

Physical function

Cardiorespiratory fitness

Upper limb strength

Lower limb functionality

Self-reported physical activity level

Health-related quality of life

Fatigue

**Assessment of the primary outcomes**

Resting LV ejection fraction and resting LV longitudinal strain

Resting LV ejection fraction will be calculated using the biplane method of disks (modified Simpson’s rule) from the apical four- and two-chamber view (30). For resting LV global longitudinal strain assessment, two-dimensional grey-scale images will be acquired in the apical four-, two- and three-chamber views, with a frame rate of 60 to 100 fps. Three cardiac cycles will be digitally stored and Velocity Vector Imaging (VVI) software (Siemens Medical Solutions United States of America Inc) will be used in the analysis.

Echocardiographic acquisitions will be performed by a single experienced cardiologist
blinded to the patient assignment group.

**Circulating NT-proBNP levels**

Nonfasting venous blood samples will be drawn by a nurse oncologist. The assessment of N-terminal pro-brain natriuretic peptide (NT-proBNP) levels will be conducted in the local clinical analysis laboratories of the CHVNG/E, which are certified by the United Kingdom National External Quality Assessment Service. This professional staff will be blinded to the patient assignment group.

**Assessment of the secondary outcomes**

**Resting blood pressure and resting HR**

Resting blood pressure (systolic and diastolic blood pressure) and resting HR will be measured using a standard automated device Philips SureSignsVM6 (Philips Medical System, Andover, United States of America). Two measurements will be carried out. The first measurement will be preceded by 5 minutes resting period and a second reading will be taken after 3 minutes. If necessary, additional records shall be obtained until two consecutive stable measurements (differences <5 mmHg for blood pressure and <7 bpm for HR) are obtained. The average of the two stable measurements will be considered for the analysis. This procedure will be carried out by a study investigator (ALA) not blinded to the patient assignment group.

**Resting HRV**

HRV is a non-invasive method to analyse cardiac autonomic function through the measurement of successive heart beats variations (RR). Resting HRV will be analysed using a HR monitor Polar V800 (Polar Electro Oy, Kempele, Finland) with a Polar H7 chest strap. During the RR recording, patients will be seated in a comfortable position. They shall be required to breathe spontaneously, to avoid any movements and to maintain neutral thoughts during the time of data acquisition. The first 5 minutes will be excluded
(stabilization period) and the remaining 5 minutes will be used to calculate the time-domain (standard deviation of successive normal RR [SDNN], and root mean square of successive normal RR [RMSSD]) and frequency-domain indices (low-frequency spectral component [LF], and high-frequency spectral component [HF]). In all the cases, the RR recordings will be exported to the Kubios v2 HRV software (Biosignal Analysis and Medical Imaging Group at the Department of Applied Physics, University of Kuopio, Kuopio, Finland). Occasional, artefact noise shall be automatically replaced with the interpolated adjacent RR interval values (filter power < low). This procedure will be carry out by the first author (PA) not blinded to the patient assignment group.

Recovery HR

Recovery HR will be determined as the absolute difference between the HR at peak effort during the cardiorespiratory exercise test (CRET) and the HR at 60-seconds, and 120-seconds post-exercise. HR values will be derived from a continuous record obtained via CRET (Mortara X-Scribe, Mortara, United States of America). This procedure will be carried out by study investigators (EV, MT), blinded to the patient assignment group.

Cardiorespiratory fitness

Cardiorespiratory fitness will be evaluated by means of a symptom-limited CRET on a treadmill (Mortara X-Scribe, Mortara, United States of America), using a modified version of the Bruce protocol (31). Expired gases will be continuously collected throughout exercise and analysed for ventilatory volume (VE) and for oxygen (O2) and carbon dioxide (CO2) content, using dedicated analysers. Standard spirometry [forced expiratory volume in 1 second (FEV1)] and forced vital capacity (FVC) will also be undertaken before the test. Equipment calibration and measurements will be done in accordance to the recommendations of the American Thoracic Society and American College of Chest Physicians (32). The following parameters will be calculated and considered for analysis:
peak oxygen consumption (peak $\text{V}O_2$, measured in millilitre per kilogram per minute), peak respiratory exchange ratio (RER), defined by the ratio of CO2 production to O2 consumption at peak effort, oxygen consumption at the anaerobic threshold (AT), defined as the point at which CO2 production increases disproportionately in relation to O2 consumption, obtained from a graph plotting O2 consumption against CO2 production, and total exercise duration (measured in seconds). The maximum HR achieved will also be recorded. The CRET will be conducted on an independent day to the remaining outcomes assessments out by study investigators (EV, MT) blinded to the patient assignment group. Patients will be in a fasted state and will not be asked to discontinue current medication before the test.

**Upper limb strength**

Upper limb strength will be evaluated by the maximal voluntary grip strength (measured in kilograms), using a digital handgrip dynamometer (Saehan Corporation, Masan, South Korea – model SH5003). Each subject will perform six trials, three in each arm, with an alternating bilateral sequence. The results will be given by the average of the three trials, respectively for operated and non-operated limb. This procedure will be carried out by the first author (PA), not blinded to the patient assignment group.

**Lower limb functionality**

Lower limb functionality will be evaluated by the sit-to-stand test using a straight-backed chair (40-centimetre high). It will be required that each subject, keeping plantar support flat on the floor and arms crossed at the chest, sit and stand as many times as possible for 30 seconds. The score of the test will be determined by the number of repetitions done respecting the above procedure. This procedure will be carried out by the first author (PA), not blinded to the patient assignment group.

**Self-reported physical activity level**
International Physical Activity Questionnaire-Short (IPAQ-SF) will be used to calculate the metabolic equivalent (MET) minutes per week spent in walking, moderate and vigorous activities. Sedentary behaviour will be determined based on time spent sitting per day (minutes). Considering the obtained scoring, participants will be categorised as a low, moderate, or high physical activity level. In this study will be used the Portuguese language of the IPAQ-SF (33). Scoring will be analysed by the first author (PA), not blinded to the patient assignment group.

Health-related to quality of life and fatigue
The European Organization for Research and Treatment in Cancer (EORTC) Quality of Life C-30 (QOL-C30) is a self-administered, validated questionnaire to assess HR-QOL in cancer patients (34). It is composed of nine multi-item scales: five functional scales (physical, role, cognitive, emotional, and social), three symptom scales (fatigue, pain, and nausea and vomiting), and a global health and quality-of-life scale. Additionally, there are five single items of commonly reported symptoms by cancer patients (dyspnea, sleep disturbance, appetite loss, constipation and diarrhea), and an item that evaluates the perceived financial impact of the disease. In this study, the third version of this questionnaire will be used, in the Portuguese language (35). Analyses will include the five functional scales, fatigue scale and the global health and quality-of-life scale. The scoring of the several scales will be carried out by the first author (PA), not blinded to the patient assignment group.

Demographic, anthropometric and clinical data
Demographic, anthropometric and clinical data will be recorded during the enrolment process (−t₀). Demographic data includes age, sex, and education. Anthropometric data includes weight, height, and body mass index. Clinical data includes disease, treatment information, past medical history and current medication. These data will be extracted
from the patients’ electronic medical files by two study coordinators (ALA, AJ) not blinded to the patient assignment group.

Safety

The safety of the intervention will be assessed by weekly tracking and monitoring the number of adverse events according to the National Cancer Institute Common Terminology Criteria for Adverse Events version 4.0 (https://ctep.cancer.gov/protocoldevelopment/electronic_applications/ctc.htm). A meeting between the study investigators will be held every weekly to review and discuss the reported adverse events. All serious adverse events will be immediately reported to the CHVNG/E ethics commission, to all study members and will be reported in the study results. Adverse events will be evaluated by the study investigators who will make the decision to stop the study early if there is an increased risk of clinical relevance.

Data management

Study data will be managed by two study investigators (PA, AJ) using a predesigned criterion to data collection form [Microsoft Office Excel version 2016 (Microsoft Corporation, Redmond, WA, United States of America] and Statistical Package for the Social Sciences files version 23.0 (IBM Corporation, Armonk, NY, United States of America)] with double-entry. Regularly data checks will be performed to ensure data quality. To ensure patients’ anonymity, they will be identified by codes and only the authors involved in the trial will have access to the full identification. The total number of patients who meet the study eligibility will be recorded, as will the number of patients who will agree or not agree to participate in the study, the number of patients who assigned to each study arm, the number of patients who participated in all sessions, the attendance of each patient in the intervention sessions, the number of patients who provided follow-up data, the number of patients included in the final analysis and the
Sample size calculation

Sample size was carried out by a power calculation based on resting LV ejection fraction outcome, using a non-commercial statistical power analysis program (G*Power Version 3.1.9.2). Based on an effect size of 0.6 in resting LV ejection fraction presented in a previous study (26), to ensure a statistical power of 80% and a significance level of 0.05, through a t-test for two independent groups, the recruitment of 72 participants is required. Predicting a 20% dropout rate (19), we estimate that a total of 90 patients will be needed (n= 45 in each arm). There are no early planned stopping rules. Adverse events will be evaluated by the study investigators who will make the decision to stop the study early if there is an increased risk of clinical relevance.

Statistical analyses

Statistical data analysis will be performed using Statistical Package for the Social Sciences. The statistical significance will be set at a p< 0.05. An intention-to-treat and per protocol approaches will be used both for all analyses. Data analysis will start with standard descriptive methods to describe the data (means and standard deviations will be calculated for continuous variables and absolute and relative frequencies for categorical variables).

The comparison of the continuous variables between the two study groups will be made using a one-way analysis of covariance (ANCOVA), adjusted for the effect of the baseline values (covariate). A linear two-way mixed ANCOVA model with repeated measures (t1, t2 and t3) will be performed to test the difference over time between the two study groups and interaction (Group × Time), on primary and secondary outcomes, with the same covariate as the one-way ANCOVA. Bonferroni’s post-hoc procedure will be performed to locate the pairwise differences. Normality will be verified by the Kolmogorov-Smirnov
test and the homogeneity of the variance will be validated by the Levene's test. Effect size will be calculated to estimate variance between moments through partial eta-squared. The cut-off values were interpreted as 0.02 for small effect size, 0.13 for moderate and 0.26 for large (36). The chi-squared test will be used to check the existence of a relationship between categorical variables. Effect size will be calculated using Cramer’s V test and their interpretation will be based on the following cut-off values: 0.10 for a small effect, 0.30 for a medium effect and 0.50 for a large effect. (36)

No interim analyses will be conducted.

**Blinding**

This study will involve the prescription of exercise sessions. To carry out a rigorous exercise prescription and to ensure an adequate follow-up of each patient, participants, physical trainer (PA), medical oncologists (AJ) and nurse oncologists (ALA) will not be blinded to group assignment. Due to the lack of resources, only the evaluators that will make the acquisition of echocardiographic outcomes (resting LV ejection fraction and LV global longitudinal strain), circulating NT-proBNP levels, cardiorespiratory fitness and recovery HR data, will be blinded to the group assignment.

**Limitations**

There are some limitations to this study, which should be noted. Firstly, we will only include patients followed from CHVNG/E. Secondly, due to the impossibility of blinding patients and some of the involved authors about the study group assignment, the open design of this study may influence the assessments retention rate of participants who were allocated the control group. Thirdly, we will stratify our sample considering the age (under/over 50 years old) and use of trastuzumab (yes/no). However, there are other risk factors associated with ARC, including: total cumulative anthracycline dose, pre-existing cardiac disease and treatment with mediastinal radiation, that should also be considered.
Although these hinderances should be acknowledged, we believe the findings from the present study will provide important data which will be of relevance to the contemporary literature in this subject.

**Dissemination**

Findings of this study will be involved in a doctoral thesis of the main author, will be submitted to a peer-reviewed journal for publication and presented at relevant conferences and disseminated to the public by the partnership with CHVNG/E, Universidade da Beira Interior (Covilhã, Portugal) and community stakeholders.

**Discussion**

CVD is one of the leading causes of death among women with BC (3-5) and a foremost concern in the clinical practice of oncology (2). According to Gernaat et al (4) BC survivors have an absolute risk of dying from CVD that ranges from 1.6% to 10.4%. The recurrent use of cardiotoxic therapies and the presence of risk-factors for the development and worsening of cardiovascular health are the main factors related to this phenomenon. The implementation of preventive strategies aiming to optimize cardiovascular care in BC survivors is therefore an emerging need. Until now, these include: early identification of potential risk-factors, the management of anthracycline cumulative dose (and possible use of ancillary therapies), or the use of heart failure drugs (beta-blockers, angiotensin converting enzyme inhibitors, and angiotensin receptor antagonists) (21). However, these approaches were mainly established to reduce the toxicity of therapeutic agents, and ensuring their efficacy, but not providing a general preventive cardiovascular approach. It is thus pivotal to investigate the effects of non-pharmacological approaches prevention strategies in order to counteract ARC and related complications in BC patients.

Currently, exercise is recognised as a safe and effective supportive intervention to improve physical function and HR-QOL in BC survivors during (19) or after treatment (20).
Furthermore, exercise has also been proposed as a potential tool to mitigate ARC in humans (21, 37), although this remains uncertain since this is manly supported by data from animal studies and/or using less accurate biomarkers (22, 23). In fact, there is limited data on whether the benefits of exercise also include protection from anthracycline-related cardiac damage in women with BC. To the best of our knowledge, Kirkham et al (24) were the first to test this hypothesis by analysing the efficacy of a single aerobic bout performed 24 hours prior to each treatment. The authors observed positive effects on systemic outcomes (cardiac output, resting HR, body weight, and psychological symptoms), but did not observe relevant changes in echocardiographic outcomes (LV mass, LV ejection fraction, strain imaging and the E/A ratio) or circulating cardiac biomarkers (troponin t and NT-proBNP). In a recent non-randomized trial, Howden et al (26) verified that performing a 2-weekly supervised exercise program plus a weekly unsupervised aerobic session attenuated the reduction in peak $\dot{V}o_2$ and the increase of troponin levels in women with BC undergoing AC-CT, when compared to a usual care group. However, these authors did not also observe any relevant changes in the measured echocardiographic outcomes. So far, the overall clinical significance of exercise in preventing cardiac dysfunction associated with treatment with anthracyclines remains to be clarified in controlled clinical settings.

The present study will explore the cardioprotective role of exercise and potential mitigation of ARC, as well as establish the possible effect of exercise on different health outcomes in women with BC. A total of 90 adult women with early BC and with therapeutic decision to receive AC-CT from the CHVNG/E will be enrolled. Recruitment will take place between 1 November 2018 and 31 November 2020 (expected date of recruitment completion). Follow-up is expected to be completed by 30 June 2021.

We anticipate that the results of this study will add new knowledge to what the literature
currently offers, clarifying the effects of a supervised exercise program on different established markers and more accurate biomarkers of cardiotoxicity in women with BC undergoing AC-CT. Furthermore, we expect that the findings from this study may help in future policies related to cancer care management and contribute to the ascertainment of the role of exercise programs during antineoplastic in-hospital treatment.

TRIAL STATUS

Recruitment started on 1st November 2018 and is expected to be completed by 31st November 2020. Last edited: 11th January 2019.

Abbreviations

AC-CT: Anthracycline-Containing Chemotherapy; ARC: Anthracycline-Related Cardiotoxicity; AT: Anaerobic Threshold; BC: Breast Cancer; CRET: Cardiorespiratory Exercise Test; CO2: Carbon Dioxide; EORTC: European Organization for Research and Treatment in Cancer; FVC: Forced Vital Capacity; HF: High-Frequency Spectral Component; HR: Heart Rate; HR-QOL: Health Related to Quality of Life; HRV: Heart Rate Variability; IPAQ-SF: International Physical Activity Questionnaire-Short; LF: Low-Frequency Spectral Component; LV: Left Ventricular; MET: Metabolic Equivalent; NT-proBNP: N-Terminal pro-Brain Natriuretic Peptide; O2: Oxygen; peak Vo2: Peak Oxygen Consumption; QOL-C30: Quality of Life C-30; RER: Respiratory Exchange Ratio; RMSSD: Root Mean Square of Successive Normal RR; RR: Successive Heart Beats Variations; SDNN: Standard Deviation of Successive Normal RR; VE: Ventilatory Volume; VVI: Velocity Vector Imaging.

Declarations

Ethics approval and consent to participate

Ethics approval of this protocol was conferred by the ethics Committee of the Centro CHVNG/E (reference number: 145/2018-1). The trial will be conducted in compliance with
the Declaration of Helsinki Ethical Principles (1975). Informed written consent will be obtained from all participants before enrolment.

Consent for publication

Not applicable.

Availability of data and materials

The data that will support the findings of this study will be available on request from the corresponding author. The request will be analysed by the research team and the ethics committee that ethically approved the study.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

The idea of the above-mentioned trial was conceived and designed by PA and AJ. PA, DE, AA and AJ designed the intervention. CN and PA developed the statistical analysis and sample size calculation. EV and MT designed and will perform the cardiorespiratory assessments. FS designed the echocardiography acquisition. ALA will collect some secondary outcomes and together with AJ will record demographic, anthropometric and clinical data. PA and AJ will be responsible for the management of study data. PA drafted the manuscript, which underwent revision by all other authors. All authors read and approved the publication of this final manuscript.

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Tables

Table 1 Eligibility criteria

| INCLUSION CRITERIA:                                      | EXCLUSION CRITERIA:                                                                 |
|----------------------------------------------------------|------------------------------------------------------------------------------------|
| · Female gender.                                         | · Contraindications to maximal exercise testing.                                   |
| · Contraindications to maximal exercise testing.         | · Decompensated diabetes mellitus.                                                |
| · Aged 18 years or older.                                | · Severe anaemia (<8g / dL) uncorrectable with transfusion and / or iron and / or vitamin deficiency replacement. |
| · Aged 18 years or older.                                | · Pregnancy.                                                                       |
| · Histological diagnosis of stage IA-IlC breast carcinoma.| · Known significant heart disease (myocardial infarction, congestive heart failure, cardiomyopathy). |
| · Scheduled to receive AC-CT.                            | · Usual medication-containing beta-blockers.                                       |
| · Follow-up at the Medical Oncology clinic at the CHVNG / E. |                                                                                   |
| · Consent of the assistant oncologist for the practice of exercise. |                                                                                   |
| · Able to provide informed consent.                      |                                                                                   |
| · Acceptance of randomization to intervention group or control group. |                                                                                   |

Figures
Figure 1

Study flow. ACH: anthracycline-containing chemotherapy; SCC: standard breast cancer care; SEP: structured exercise program.
| TIMEPOINT        | Enrollment | Allocation | Post-allocation | Follow-up |
|------------------|------------|------------|-----------------|-----------|
| −t₀              | 0          | t₀         | t₁              | t₂        | t₃        |

**ENROLMENT:**
- Eligibility screen ✓
- Informed consent ✓
- Demographic data ✓
- Anthropometric data ✓ ✓ ✓ ✓ ✓
- Clinical data ✓ ✓ ✓ ✓ ✓
- Allocation ✓

**INTERVENTIONS:**
- Supervised exercise program
- Usual cancer care

**ASSESSMENTS:**
- Primary outcomes
  - NT-proBNP ✓ ✓ ✓ ✓ ✓
  - Resting LV global longitudinal strain ✓ ✓ ✓ ✓
  - Resting LV ejection fraction ✓ ✓ ✓
- Secondary
| outcomes                       | t0 | t1 | t2 |
|-------------------------------|----|----|----|
| Resting blood pressure        | ✓  | ✓  | ✓  |
| Resting heart rate            | ✓  | ✓  | ✓  |
| Resting heart rate variability| ✓  | ✓  | ✓  |
| Heart rate recovery           | ✓  | ✓  | ✓  |
| Physical Function:            |    |    |    |
| Cardiopulmonary capacity      | ✓  | ✓  | ✓  |
| Upper limb strength           | ✓  | ✓  | ✓  |
| Lower limb functionality      | ✓  | ✓  | ✓  |
| Self-reported physical activity level | ✓ | ✓ | ✓ |
| Health Related to Quality of Life | ✓ | ✓ | ✓ |
| Fatigue                       | ✓  | ✓  | ✓  |

**Figure 2**

The schedule of enrolment, interventions, and assessments. -t0 Enrolment process t0 Baseline, t1 During anthracycline-containing chemotherapy, t2 Post anthracycline-containing chemotherapy (which coincides with the end of the intervention), t3 After 3 months of t2, NT-proBNP N-Terminal pro-Brain Natriuretic Peptide, LV Left ventricular.

**Supplementary Files**

This is a list of supplementary files associated with the primary manuscript. Click to download.

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