

# The role of exercise training frequency in long-term blood pressure control

<b>Submission date</b> 12/06/2025	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 16/06/2025	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
<b>Last Edited</b> 18/06/2025	<b>Condition category</b> Circulatory System	<input type="checkbox"/> Individual participant data

## Plain English summary of protocol

### Background and study aims

High blood pressure is a major risk factor for heart disease. Recent research suggests that performing isometric wall squat exercise training can significantly reduce blood pressure. This study aimed to find out how the frequency of wall squat training affects blood pressure and cardiovascular health, and whether stopping the exercise leads to a reversal of benefits.

### Who could take part?

Adults aged 18–65 years with normal to slightly raised blood pressure (systolic 120–139 mmHg) who were not taking blood pressure medication and had no known cardiovascular, metabolic, or musculoskeletal conditions.

### What did the study involve?

Participants were randomly allocated to one of five groups. All groups completed wall squat training three times per week for the first 4 weeks. In the second 4-week phase, training frequency varied: participants either continued training at three, two or one time per week, stopped training entirely, or were in a control group with no training throughout. Blood pressure and other cardiovascular measurements were taken at baseline, mid-point (week 4), and after 8 weeks.

### What were the possible benefits and risks of taking part?

Participants may have experienced improvements in blood pressure. The exercise is low-risk but may have caused some temporary muscle discomfort. All procedures were designed to be safe and accessible to people with no prior exercise experience.

### Where was the study run from?

The study was conducted at Canterbury Christ Church University (CCCU), within the School of Psychology and Life Sciences.

### When did the study take place and how long did it run for?

September 2019 to March 2025

Who funded the study?

The study was funded internally by Canterbury Christ Church University as part of a postgraduate research degree

Who was the main contact?

Dr Harry Swift, [harry.swift@canterbury.ac.uk](mailto:harry.swift@canterbury.ac.uk)

## Contact information

### Type(s)

Principal Investigator

### Contact name

Dr Harry Swift

### ORCID ID

<https://orcid.org/0000-0003-0499-5706>

### Contact details

North Holmes Road

Canterbury

United Kingdom

CT1 1QU

+44 (0)1227470442

[harry.swift@canterbury.ac.uk](mailto:harry.swift@canterbury.ac.uk)

### Type(s)

Scientific

### Contact name

Dr Damian Coleman

### ORCID ID

<https://orcid.org/0000-0002-7332-8221>

### Contact details

North Holmes Road

Canterbury

United Kingdom

CT1 1QU

+44 (0)1227 767700

[damian.coleman@canterbury.ac.uk](mailto:damian.coleman@canterbury.ac.uk)

### Type(s)

Public

### Contact name

Dr Jim Wiles

### ORCID ID

<https://orcid.org/0000-0002-7790-8063>

**Contact details**

North Holmes Road  
Canterbury  
United Kingdom  
CT1 1QU  
+44 (0)1227 767700  
jim.wiles@canterbury.ac.uk

**Additional identifiers****EudraCT/CTIS number**

Nil known

**IRAS number****ClinicalTrials.gov number**

Nil known

**Secondary identifying numbers**

19/SAS/12C

**Study information****Scientific Title**

Understanding dose-response in isometric exercise training: the role of training frequency in long-term blood pressure control

**Study objectives**

Increasing the frequency of isometric exercise training will result in greater improvements in blood pressure, vascular resistance, and autonomic function in normotensive to pre-hypertensive adults, compared to lower-frequency training or no training.

**Ethics approval required**

Ethics approval required

**Ethics approval(s)**

Approved 19/12/2019, Canterbury Christ Church University ethics board (School of Psychology and Life Sciences, Canterbury Christ Church University, Kent,, Canterbury, CT1 1QU, United Kingdom; +44 (0)1227767700; red.resgov@canterbury.ac.uk), ref: 19/SAS/12C

**Study design**

Single-centre interventional randomized controlled trial

**Primary study design**

Interventional

**Secondary study design**

Randomised controlled trial

**Study setting(s)**

University/medical school/dental school

**Study type(s)**

Efficacy

**Participant information sheet**

Not available in web format, please use contact details to request a participant information sheet

**Health condition(s) or problem(s) studied**

Prevention of hypertension in normotensive to pre-hypertensive adults through isometric exercise training

**Interventions**

All participants are randomly assigned using stratified random allocation to one of five groups. During the first 4 weeks, all intervention groups complete isometric wall squat training three times per week, consisting of four 2-minute bouts at 95% peak heart rate, with 2 minutes rest between bouts. In the final 4 weeks, training frequency is modified: participants either continue training at 3x/week, reduce to 2x/week, reduce to 1x/week, stop training entirely, or remain in the control group with no training throughout.

Training is supervised or remotely monitored to ensure adherence. Assessments are conducted at baseline, mid-point (week 4), and end-point (week 8), measuring resting blood pressure, cardiac output, total peripheral resistance, heart rate variability, and baroreflex sensitivity.

**Intervention Type**

Other

**Primary outcome measure**

Resting systolic blood pressure (SBP) is measured using the Task Force® Monitor during a 5-minute seated continuous recording at baseline, week 4, and week 8

**Secondary outcome measures**

1. Mean arterial pressure (MAP) is calculated from continuous blood pressure data recorded by the Task Force® Monitor during a 5-minute seated period at baseline, week 4, and week 8
2. Diastolic blood pressure (DBP) is measured using the Task Force® Monitor during the same 5-minute seated recording at baseline, week 4, and week 8
3. Total peripheral resistance (TPR) is derived from concurrent blood pressure and cardiac output signals via the Task Force® Monitor during a 5-minute seated measurement at baseline, week 4, and week 8
4. Cardiac output (Q) and stroke volume (SV) are recorded via impedance cardiography using the Task Force® Monitor during a 5-minute seated recording at baseline, week 4, and week 8
5. Heart rate variability (HRV) metrics (LF, HF, LFnu, HFnu, LF/HF ratio) are calculated from ECG data obtained via the Task Force® Monitor during the 5-minute seated period at baseline, week 4, and week 8
6. Baroreflex sensitivity (BRS) is assessed using sequence and spectral methods applied to the 5-minute continuous Task Force® Monitor data at baseline, week 4, and week 8

**Overall study start date**

01/09/2019

**Completion date**

01/03/2025

## **Eligibility**

**Key inclusion criteria**

1. Male and female participants with normal to high-normal sBP (range 120-140 mmHg)
2. University staff and students from Canterbury Christ Church University and their associates via social media and word of mouth
3. Total weekly physical activity level below the 150 minutes of moderate physical activity per week, or specifically  $\leq 600$  MET-minutes, calculated as the sum of all light-intensity activities (light walking = 3.3 METs, light cycling = 3.0 METs, household work = 2.5 METs). Total MET-minutes were computed as  $\text{MET} \times \text{duration (min)}$ . To exclude structured exercise, any reported bout  $\geq 4$  METs lasting  $\geq 10$  minutes (e.g. jogging and weight training) led to exclusion.
4. Free from any injuries or illnesses
5. Not taking any medication (or had previously taken any anti-hypertensive medication)
6. Non-smokers
7. Consumed less than 14 units of alcohol per week
8. Screening was conducted using a standard Physical Activity Readiness Questionnaire
9. Provided written informed consent

**Participant type(s)**

Healthy volunteer

**Age group**

Adult

**Lower age limit**

18 Years

**Upper age limit**

65 Years

**Sex**

Both

**Target number of participants**

118

**Total final enrolment**

100

**Key exclusion criteria**

1. Free from any injuries or illnesses
2. Not taking any medication (or had previously taken any anti-hypertensive medication)
3. Non-smokers
4. Consumed less than 14 units of alcohol per week

**Date of first enrolment**

21/01/2020

**Date of final enrolment**

16/08/2024

## **Locations**

**Countries of recruitment**

England

United Kingdom

**Study participating centre**

**Canterbury Christ Church University**

St. Georges Place

Canterbury

United Kingdom

CT1 1UT

## **Sponsor information**

**Organisation**

Canterbury Christ Church University

**Sponsor details**

North Holmes Road

Canterbury

England

United Kingdom

CT1 1QU

+44 (0)1227 767700

graduatecollege@canterbury.ac.uk

**Sponsor type**

University/education

**Website**

<https://www.canterbury.ac.uk/>

**ROR**

<https://ror.org/0489ggv38>

# Funder(s)

## Funder type

University/education

## Funder Name

Canterbury Christ Church University

## Alternative Name(s)

## Funding Body Type

Private sector organisation

## Funding Body Subtype

Universities (academic only)

## Location

United Kingdom

# Results and Publications

## Publication and dissemination plan

Results from this study will be published in a peer-reviewed journal and a PhD thesis.

## Intention to publish date

03/03/2025

## Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be stored in a publicly available repository.

The dataset is stored in the UK Data Service ReShare repository.

The dataset includes fully anonymised individual-level data on blood pressure, heart rate variability, cardiac output, and vascular resistance across three timepoints (baseline, week 4, week 8), as well as group allocation and basic demographics (age, sex).

Data will be made available indefinitely through the UK Data Service under standard academic licensing terms.

Access will be open to registered users for non-commercial research and teaching purposes, subject to acceptance of a user agreement.

No personally identifiable data are included. Anonymisation involved the removal of names, dates, and indirect identifiers, and aggregation of sensitive subgroups where needed.

Informed consent for data sharing was obtained from participants via approved ethics documentation, and the study was reviewed and approved by Canterbury Christ Church University's ethics committee.

## IPD sharing plan summary

Stored in publicly available repository

**Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Participant information sheet</a>			13/06/2025	No	Yes
<a href="#">Basic results</a>		18/06/2025	18/06/2025	No	No