

High versus low load training in females

Submission date 17/04/2025	Recruitment status Recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
Registration date 01/05/2025	Overall study status Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
Last Edited 23/04/2025	Condition category Other	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

Performing resistance exercise increases muscle mass and muscle mass is a strong predictor of illness and death in North America's aging population. Recent research has found that both relatively heavy loads and relatively light loads are equally effective at producing muscular adaptations. Additionally, extrinsic factors such as protein provision/diet may influence adaptations. However, the source of protein intake and its interaction with different training loads have yet to be extensively investigated in women. Research to date emphasises that protein types – casein, whey, egg, soy and even other forms of protein from plant sources – are likely no different in their ability to promote training-induced hypertrophy (muscle growth) and strength gains. However, very little of this work is carried out in younger women. Thus, the main aim of this study is to assess the interaction between training load (high vs low) and protein source (casein vs whey) on body composition, muscle hypertrophy and strength gains in women.

Who can participate?

Healthy female volunteers aged 18-35 years

What does the study involve?

The participants will be randomly allocated to either a relatively heavy-lifting group (about 80% one-rep max [1RM], able to perform 8-12 repetitions) or a relatively light-lifting group (about 30-50% 1RM, able to perform 15-25 repetitions). They will perform 10 weeks of resistance exercise training and be given 40 g of whey or casein protein following each training session. The researchers will take various measurements on muscle size, body composition and muscle strength pre- and post-training intervention.

What are the possible benefits and risks of participating?

There are no proposed benefits to participants. However, the findings of this study may contribute to the development of nutritional guidelines that maximise muscle health.

As with any research, there are risks associated with participating, such as during blood sampling or muscle biopsies (samples). The research team has done everything possible to mitigate these risks and will gladly provide further information if requested.

Where is the study run from?

McMaster University in Hamilton, Ontario (Canada)

When is the study expected to run for?

January 2025 to August 2026

Who is funding the study?

Natural Sciences and Engineering Research Council of Canada (NSERC) (Canada)

Who is the main contact?

Dr Stuart Phillips, phillis@mcmaster.ca

Contact information

Type(s)

Scientific, Principal Investigator

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Additional identifiers

EudraCT/CTIS number

Nil known

IRAS number**ClinicalTrials.gov number**

Nil known

Secondary identifying numbers

HiREB #18158

Study information

Scientific Title

The effect of high load versus low load resistance training and protein type on muscle mass and strength in females

Acronym

HER-TRAIN

Study objectives

The working hypothesis is that higher and lower load resistance exercise training (RET) will induce equivalent muscle hypertrophy, measured as muscle mass (D3-creatine), lean mass (dual-energy x-ray absorptiometry [DXA]), and strength (i.e., no effect of load); and protein supplementation will be equally effective regardless of source in promoting strength and hypertrophy.

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 07/04/2025, Hamilton Integrated Research Ethics Board (237 Barton Street East, Hamilton, L8L 2X2, Canada; +1 (0)905 521 2100, Ext 42013; eREBhelpdesk@hhsca.ca), ref: 18158

Study design

Randomized controlled trial

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

University/medical school/dental school

Study type(s)

Treatment

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

Resistance exercise and protein feeding

Interventions

This is a 2 x 2 nested factorial RCT where participants will be randomized to a training intervention (higher or lower load) and then randomized within each arm to receive either whey or casein protein.

Participants will be randomized to either a relatively heavy-lifting group (HL: ~80% 1RM, able to perform 8-12 repetitions/set) or a relatively light-lifting group (LL: ~30-50% 1RM, able to perform 15-25 repetitions/set). They will perform 10 weeks of resistance exercise training (training twice weekly) and be given 40 g of whey or casein protein (also by random assignment) following each training session. The researchers will take various measurements of muscle size, body composition and muscle strength pre- and post-training intervention.

Intervention Type

Other

Primary outcome measure

Skeletal muscle mass assessed by D3-Cr-measured muscle mass at baseline and 10 weeks

Secondary outcome measures

Lean body mass:

1. Dual-energy x-ray absorptiometry (DXA)-measured lean body mass at baseline and 10 weeks.

Muscle strength:

2. One repetition-maximum on a leg press machine and biceps curl measured at baseline, 5 and 10 weeks

3. Unilateral isometric maximum voluntary contraction of knee extension/elbow flexion measured at baseline and 10 weeks

Overall study start date

01/01/2025

Completion date

31/08/2026

Eligibility

Key inclusion criteria

1. Between the ages of 18-35 years (inclusive)
2. Able to maintain a habitual diet and perform RET three times per week throughout the trial
3. Able to begin exercise (assessed by Get Active Questionnaire [GAQ])
4. Understand the study procedures and sign this form providing informed consent to participate in the study

Participant type(s)

Healthy volunteer

Age group

Adult

Lower age limit

18 Years

Upper age limit

35 Years

Sex

Female

Target number of participants

52

Key exclusion criteria

1. Use of tobacco- or cannabis-related products (smoking or vaping)
2. A history of neuromuscular problems or muscle and/or bone-wasting diseases
3. Any acute or chronic illness; cardiac, pulmonary, liver, or kidney abnormalities; insulin- or non-insulin-dependent diabetes or other metabolic disorders (all ascertained through medical questionnaires)
4. Use of medications known to affect protein metabolism (i.e. corticosteroids, non-steroidal anti-inflammatory drugs (prescription use or daily use of over-the-counter medication), or prescription strength acne medications)
5. Have any electronic medical or metal implants
6. Currently pregnant or planning to get pregnant
7. Dairy allergy

Date of first enrolment

05/05/2025

Date of final enrolment

31/05/2026

Locations

Countries of recruitment

Canada

Study participating centre

McMaster University

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Sponsor information

Organisation

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Sponsor type

University/education

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ROR

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Funder(s)

Funder type

Government

Funder Name

Natural Sciences and Engineering Research Council of Canada

Alternative Name(s)

Conseil de Recherches en Sciences Naturelles et en Génie du Canada, NSERC, CRSNG

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

Canada

Results and Publications

Publication and dissemination plan

Planned publication in a peer-reviewed journal.

Intention to publish date

31/10/2026

Individual participant data (IPD) sharing plan

The data-sharing plans for the current study are that all data will be made publicly available upon reasonable request from the PI, Dr Stuart Phillips (phillis@mcmaster.ca).

IPD sharing plan summary

Available on request