

Guided exercise intervention for improving physical fitness and weight control in overweight and obese males and females

Submission date
29/09/2009

Recruitment status
No longer recruiting

☐ Prospectively registered

☐ Protocol

Registration date
09/11/2009

Overall study status
Completed

☐ Statistical analysis plan

☐ Results

Last Edited
09/11/2009

Condition category
Nutritional, Metabolic, Endocrine

☐ Individual participant data

☐ Record updated in last year

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

Prof Sulin Cheng

Contact details

Department of Health Sciences
PO Box 35
Jyväskylä
Finland
401001

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers

EWI-study; K-S shp:n Dnro 4U/2009

Study information

Scientific Title

Exercise and weight control intervention for improving physical fitness - fitness line algorithm validation: a randomised controlled trial

Acronym

EWI-study

Study objectives

1. Guided exercise is more effective for improving physical fitness and weight control than unguided exercise. The most sensitive changes will be seen in maximal oxygen uptake (max VO₂) and gut microbiota after a short-term exercise program while the differences will be more pronounced in heart rate variability (HRV) and body composition after long-term exercise program.
2. There will be high dropout after a short-term intervention. Overweight and obese people need guidance and monitoring of their physical condition to motivate them towards continuing exercise and weight control. To establish "habitual exercise" requires long term guidance and commitment.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics Committee of Central Finland Health Care District, Jyväskylä, Finland, approved on the 17th June 2009 (ref: K-S shp:n Dnro 4U/2009)

Study design

Randomised controlled trial

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Other

Study type(s)

Treatment

Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

Health condition(s) or problem(s) studied

Overweight/obesity

Interventions

Phase I (6-week intervention):

Group 1 - exercise follows guidance from a wrist computer: subjects will follow the daily guidance of the device to exercise (Nordic break walking) 30 minutes to 1 hour each time and gradually increase the intensity and duration. The intensity of exercise will be checked every 2nd week and the guidance will be modified accordingly.

Group 2 - general training advice but no device: subjects will follow the instruction by a researcher to exercise (Nordic break walking) 30 minutes to 1 hour a time and gradually increase the intensity and duration.

Group 3 - weight control follows guidance from a wrist computer: subjects follow the weight guidance from device to change their weight either by eating or exercise. The weight (body composition) will be checked every 2nd week and the guidance will be modified accordingly.

Group 4 - weight control gets general advice but no device: subjects follow the instruction by a nutritionist to change their weight either by eating or exercise.

After the first phase of the study, subjects will not be informed that they will be followed-up for phase II of the study. Thus it can be ensured that the maintenance rate is not interfered with by the participants knowing that they will be followed later.

In study phase II (1-year intervention), all of the subjects will be invited back 6 months after the onset of the phase I study. A wrist computer will be provided to them and subjects will follow the instruction to set the exercise intensity/duration and target weight on the basis of their fitness level and weight to exercise. Those people who are not maintaining their leisure time physical activity (LTPA) will be randomly assigned to two groups. One group will have and one will not have 6 months measurements. Phone call tracking interview will be contacted every 3 months interval to evaluate the maintenance rate.

Intervention Type

Other

Phase

Not Applicable

Primary outcome measure

Efficacy of guided exercise on fitness and weight control: physical fitness and body composition (HRV, gut microbiota, Max VO₂, fat mass, lean mass) -

1. Daily physical activity and a diet diary will be assessed at baseline, every week during the intervention, at follow-up point 2, and every week during the long term intervention until follow-up point 3
2. Anthropometry and blood pressure will be measured at baseline, half-way through the long-term intervention and at follow-up points 1, 2 and 3. Blood samples will also be taken at this point.
3. Body composition, and the 10-metre walking test will be measured at baseline and every two weeks through the short term intervention until follow-up point 1, and at follow-up point 2, halfway-way through the long term intervention and at follow-up point 3. The treadmill test will also be taken at all of these points apart from follow-up point 3.
4. Faecal samples, basal metabolic rate and max VO₂ will be measured at baseline and follow-up points 1, 2 and 3

Secondary outcome measures

Maintenance of exercise and weight control intervention:

1. How many people will maintain their physical activity and weight control after a short

intervention?

2. What are the reasons for people stopping exercise and weight control after a short intervention?

3. Does long-term guided exercise and weight control intervention help to maintain peoples LTPA and weight control after intervention?

Overall study start date

20/06/2009

Completion date

31/12/2012

Eligibility

Key inclusion criteria

1. Women and men aged 20 - 50 years
2. Physically inactive (regular exercise less than or equal to two times per week and less than or equal to 45 minutes per session)
3. Body mass index (BMI) greater than 25 kg/m² 38 kg/m²

Participant type(s)

Patient

Age group

Adult

Sex

Both

Target number of participants

100 women and 100 men

Key exclusion criteria

1. BMI greater than 38 kg/m²
2. Serious cardiovascular or musculoskeletal problems
3. Diagnosed type I diabetes
4. Change in weight more than 5 kg during last 6 months

Date of first enrolment

20/06/2009

Date of final enrolment

31/12/2012

Locations

Countries of recruitment

Finland

Study participating centre
Department of Health Sciences
Jyväskylä
Finland
401001

Sponsor information

Organisation
Suunto Oy (Finland)

Sponsor details
c/o Veikko Koivumaa
Valimotie 7
Vantaa
Finland
01510

Sponsor type
Industry

Website
<http://www.suunto.com/suunto/main/index.jsp>

ROR
<https://ror.org/04n2d2p10>

Funder(s)

Funder type
Industry

Funder Name
Suunto Oy (Finland)

Funder Name
University of Jyväskylä (Finland) - Wellness Program

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date**Individual participant data (IPD) sharing plan****IPD sharing plan summary**

Not provided at time of registration