The vital@work study. The (cost-) effectiveness of a lifestyle intervention in order to improve older workers' vitality.

No registrations found.

Ethical review Positive opinion

Status Pending

Health condition type -

Study type Interventional

Summary

ID

NL-OMON24340

Source

NTR

Brief title

The vital@work study

Health condition

Vitality (vitaliteit), aging (work force) (vergrijzing), Lifestyle (leefstijl/ BRAVO), Early retirement (vroeg pensioen), productivity (productiviteit)

Sponsors and support

Primary sponsor: VU University Medical Center, EMGO Institute

Source(s) of monetary or material Support: Sichting Instituut GAK (SIG)

Intervention

Outcome measures

Primary outcome

Vitality and lifestyle behaviour (Physical activity, dietary behaviour, alcohol consumption, smoking habits).

Secondary outcome

Work engagement and Productivity, General health status, quality of life, sick leave and costeffectiveness.

Study description

Background summary

The Vital@work Study. The (cost-) effectiveness of a lifestyle intervention in order to improve older worker's vitality.

J.E. Strijk, K.I. Proper, A.J. van der Beek, W. van Mechelen, MD

Background

As a consequence of an aging workforce, workers need to prolong their working life. vitality is of great importance for the endured employability of aging workers. Since having high energy levels is one of the main components of vitality, lifestyle could contribute to aging workers' vitality by positively affecting aging workers activity. As a consequence, a lifestyle intervention which aims to improve lifestyle is considered to be an effective instrument to keep aging workers vital. The aim of this study is to develop and evaluate a lifestyle intervention program to promote health in order to improve older workers' (45 years and older) vitality.

Methods

The intervention will be evaluated using a RCT design with 2 arms. The older workers in the intervention group (n=230) will be receiving tailored lifestyle intervention during 6 months. The older workers in the control group (n=230) will receive usual care. Measurement will take place at baseline and after 6 and 12 months after implementing the intervention. Primary outcome measures are vitality and lifestyle behaviour (physical activity, dietary behaviour, alcohol consumption, smoking habits). Secondary outcome measures are work engagement, productivity, general health status, quality of life, sick leave and cost-effectiveness.

Results

No results are available yet. This project will start January 2008 and the follow up measurements will continue until the end of 2010.

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Study objective

Our hypothesis is that older workers in the intervention group will improve lifestyle and thereby vitality at the short term (6 months) and at the long term (12 months), whereas in the control group lifestyle and vitality will remain the same as at baseline.

Study design

The first follow-up after the baseline measurement (T=0 weeks) will take place at 6 months (T=24 weeks) after implementing the intervention. The participants will be asked to fill in the same questionnaires as for the baseline measurement. This procedure will be repeated for the second follow-up at twelve months (T=52 weeks) after the start of the intervention.

Intervention

Detailed information about the format and content of the intervention cannot be given yet, as it will be developed during the first phase of the project using the invertention mapping protocol. The intervention will be aimed to achieve behavioural changes as to a healthier lifestyle among older workers to improve workers' vitality. Based on current literature and experience, the intervention may include the following components: Using social ecological models and implementation intentions, respondents will receive a tailored intervention by means of counselling/ face-to-face/ telephone/internet contacts. The control group will receive care as usual.

Contacts

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Eligibility criteria

Inclusion criteria

- 1. Aged 45 or older
- 2. Contract of employment of at least 20 hours a week
- 3. Contract of employment till the end of the measurements
- 4. Informed consent

Exclusion criteria

1. Disease which makes physical activity impossible

Study design

Design

Study type: Interventional

Intervention model: Parallel

Allocation: Randomized controlled trial

Masking: Single blinded (masking used)

Control: Active

Recruitment

NL

Recruitment status: Pending

Start date (anticipated): 01-12-2007

Enrollment: 450

Type: Anticipated

Ethics review

Positive opinion

Date: 03-03-2008

Application type: First submission

Study registrations

Followed up by the following (possibly more current) registration

No registrations found.

Other (possibly less up-to-date) registrations in this register

No registrations found.

In other registers

Register ID

NTR-new NL1195 NTR-old NTR1240

Other Research Committee VUmc: WC2007-009

ISRCTN wordt niet meer aangevraagd

Study results

Summary results

N/A