Reducing shoulder complaints in employees with high occupational shoulder exposures: study protocol for a cluster-randomised controlled study (The Shoulder-Café Study)

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Method Article

Keywords: Exercise, Intervention, Mechanical exposure, Occupation, Randomised controlled trial, Shoulder, Training programme

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Abstract

Background: In Denmark, exercise therapy in combination with work modifications is the first choice treatment for persons with shoulder complaints and high occupational shoulder exposures. To obtain this treatment they must visit several healthcare providers, which makes usual care fragmented and uncoordinated. Therefore, we developed a new intervention which unifies the expertise that is needed. The main hypotheses are that a group-based Shoulder-Café intervention will reduce (I) shoulder complaints and (II) occupational shoulder exposures more effectively than an individual-based Shoulder-Guidance intervention (active control – enhanced usual care).

Methods: A cluster-randomised trial is conducted including 120 employees with high occupational shoulder exposures. Companies (clusters) are randomised to either Shoulder-Café or Shoulder-Guidance with a 1:1 allocation ratio. Participants are 18–65 years old and have an Oxford Shoulder Score (OSS) ≤ 40. Both interventions include a home-based shoulder exercise programme, assessment of shoulder exposures by technical measurements and self-report, and general information on how to reduce shoulder exposures. The Shoulder-Café course also includes three café meetings with physiotherapist-supervised exercises, clinical shoulder evaluation, education on shoulder anatomy, workplace-oriented counselling, and an opportunity for a workplace visit by a health and safety consultant. The primary outcomes are the OSS at 6 month follow-up (hypothesis I), and the mean number of minutes/day with the arm elevated > 60° shortly after end of intervention (hypothesis II). We will use mixed model analysis that allows for company clustering, and data will be analysed according to the intention-to-treat principle.

Discussion: Persons with shoulder complaints and high occupational shoulder exposures are an obvious target group for secondary prevention efforts. We developed the Shoulder-Café to reduce shoulder complaints and shoulder exposures while unifying the expertise that is needed to evaluate and treat shoulder complaints. If the intervention is effective, it would warrant widespread implementation.

Trial registration: The trial was registered at Clinicaltrials.gov on 18 May 2017 (ID: NCT03159910).

Keywords: Exercise, Intervention, Mechanical exposure, Occupation, Randomised controlled trial, Shoulder, Training programme.

Background

Shoulder complaints prevail in the working age population and constitute a common reason to consult a general practitioner [1]. In the general population, the prevalence of self-reported shoulder complaints is estimated to be 16–26% [1, 2], and in occupations with high mechanical shoulder exposures, shoulder disorders are especially frequent [3, 4]. High occupational shoulder exposures are associated with an approximately doubled risk of surgery for subacromial impingement syndrome [5-7], and when combined with shoulder complaints, a more than five-fold increase in risk of later surgery has been reported [8]. Based on these findings, persons with shoulder complaints and high occupational shoulder exposures seem an obvious target group for secondary prevention efforts.
The Danish Health Authority recommends exercise therapy as first choice treatment for shoulder complaints related to subacromial impingement syndrome [9, 10]. In case of shoulder complaints in combination with high occupational shoulder exposures, the Danish Health Authority also recommends work modifications [10]. Today, employees with shoulder complaints must visit several healthcare providers to meet this recommendation, which is why they are often repeatedly seen by general practitioners, physiotherapists in private practice and municipalities, job centres, departments of orthopaedic surgery, and departments of occupational medicine [11]. This makes usual care fragmented and uncoordinated as experienced by the patients [12]. To unify the necessary expertise to evaluate and treat shoulder complaints, a café intervention was recently developed and pilot-tested in Central Denmark Region [12]. The café concept was based on an intervention study of patients after lumbar spinal fusion, where a Back-café (three café meetings plus one exercise instructions by a physiotherapist, and featuring the opportunity to exchange experiences) scored better in daily function than group-based physiotherapist-supervised exercises and individual-based video training [13]. This indicated positive effects of a café concept per se.

We further developed the pilot-tested café intervention [12] to target employees with shoulder complaints and high occupational shoulder exposures. Our café intervention, the Shoulder-Café, unifies clinical examination of the shoulders, patient education, supervised and home-based shoulder exercises, advice from a health and safety consultant on work modifications, and assessment of shoulder exposures at work.

This trial compares a group-based Shoulder-Café intervention with an individual-based Shoulder-Guidance intervention (active control – enhanced usual care). The main hypotheses are that the Shoulder-Café will reduce I) shoulder complaints and II) occupational shoulder exposures more effectively than the Shoulder-Guidance. In connection with hypothesis I, we also expect positive effects on fear avoidance beliefs, patients’ global impression of change, and a series of supplementary outcomes.

**Methods**

**Design and setting**

The design is a cluster-randomised controlled trial with two parallel groups: Shoulder-Café and Shoulder-Guidance. We chose cluster-randomisation at company level to prevent contamination between groups. T₀ is the start of the intervention. With regard to hypothesis I, baseline data is collected shortly before T₀ and follow-up data is collected 6 and 12 months after T₀. With regard to hypothesis II, baseline data is collected shortly after T₀ and follow-up data is collected shortly after end of intervention (EOI, around 3 months after T₀). The setting is Central Denmark Region. A stakeholder group with members from trade unions, municipal rehabilitation centres, general practice, and the Health Planning Agency in Central Denmark Region has been established to facilitate the completion of the project and subsequent implementation of the Shoulder-Café if the results favour this intervention. This study protocol is written
in accordance with the SPIRIT checklist [14] (Additional files 1a and 1b) in conjunction with the TIDieR checklist [15].

**Trial population**

The trial population consists of employees from occupations with high mechanical shoulder exposures who experience shoulder complaints. Relevant occupations are identified by means of a Danish Job Exposure Matrix (The Shoulder JEM), which is based on five experts’ ratings and covers all occupations in Denmark [16]. We selected occupations which fulfilled at least one of the following criteria: upper arm elevation > 90° ≥ 1 hour/day, highly repetitive work ≥ 0.5 hours/day, moderately repetitive work ≥ 4 hours/day, and a force score ≥ 3 (range 1–5) [5, 8]. Kitchen assistants with moderate exposures are also included to ensure adequate representation of women. Companies are recruited in batches according to their geographical location. To achieve adequate patient enrolment we will gradually widen the geographical distribution of companies within Central Denmark Region and include more occupational groups. The selected occupations are grouped according to industry: service (cleaning, kitchen and laundry assistants, hairdressers, and gardeners/pavers), manufacture (dairy, bread, and wood industry workers) and construction (electricians, carpenters, plumbers, bricklayers, house painters, welders, blacksmiths, and insulation workers). In a batch mode, we contact relevant companies in Central Denmark Region with at least 10 employees identified in The Central Business Register (https://datacvr.virk.dk/data/index.php?q=forside&language=en-gb). If a company accepts participation, employees from the relevant occupations are asked to fill in an electronic or postal screening questionnaire which – together with telephone screening - determines eligibility. The companies will distribute the questionnaires because according to the Danish Data Protection Act they are not allowed to give us a list with all possible participants. Thus, we cannot calculate the exact percentage that participated.

Based on the screening questionnaire, employees are invited to participate in the telephone screening if they meet the following inclusion criteria: 18–65 years old, employed in one of the selected occupations, and an Oxford Shoulder Score (OSS) ≤ 40 [17, 18]. The OSS, which exists in a Danish version [19], consists of 12 items, each referring to the past 4 weeks, with a total score ranging from 0 (worst) to 48 (best). We set the screening criterion at an OSS ≤ 40 to ensure that the included employees have shoulder complaints. The cut-off level was based on the pilot café intervention [12], where around 20% had an OSS ≤ 40, and is supported by mean scores of 42–47 in asymptomatic populations [20, 21]. Employees are excluded if they do not provide sufficient contact information or decline further participation. Based on the telephone screening, the following additional exclusion criteria are applied: no current shoulder complaints, sickness absence expected to continue into the intervention period, weekly working hours < 20, previous shoulder surgery, previous breast cancer operation, other health conditions expected to affect participation (e.g. rheumatoid arthritis, pregnancy), and inability to communicate in Danish. Employees may also decline further participation at this step. An additional exclusion criterion is failure to complete the baseline questionnaire (electronic or postal) before T₀. The time between completion of the screening questionnaire and the telephone screening is expected to be around 5 weeks,
and the subsequent time before enrolment is expected to be around 4 weeks. Figure 1 presents the expected flow of participants through the study.

**Randomisation**

Companies (clusters) are randomly allocated to Shoulder-Café or Shoulder-Guidance with a 1:1 allocation ratio using computer-generated random number assignment. Randomisation is stratified by industry (service, manufacture, construction) using blocking within strata with randomly permuted block sizes of 2, 4, and 6. A research assistant prepares closed envelopes with printed randomisation numbers and the corresponding intervention inside. Companies are contacted batch wise. When all relevant employees from a company have completed screening, the principal investigator (JT) opens the envelope and invites eligible employees from the company to their first Shoulder-Café or Shoulder-Guidance attendance. The randomisation result is not revealed to the participants until they have signed the informed consent (obtained by JT) and completed the baseline questionnaire. The baseline questionnaire includes self-reported typical occupational shoulder exposures (see “Other assessments” below), while baseline assessment of occupational shoulder exposures with respect to hypothesis II takes place after the randomisation result has been revealed.

**Interventions**

The Shoulder-Café is designed as a complex intervention [22] with interacting components unified into a group intervention, whereas the Shoulder-Guidance is a simpler individual intervention. Consecutively, around 60 employees are scheduled to attend one of around 12 Shoulder-Café courses. Concurrently, around 60 employees are scheduled to attend a Shoulder-Guidance course. Each course lasts around 3 months with variations depending on practical issues, e.g. care givers’ time schedules. Physical attendance will take place at six geographically dispersed municipal health centres. A description of the Shoulder-Café and Shoulder-Guidance is presented in Table 1.

The following elements are identical in the Shoulder-Café and the Shoulder-Guidance:

- A home-based shoulder exercise programme with instructions for individual tailoring, described in a pamphlet (Additional file 2). Exercises for treating shoulder complaints have shown promising results [23-26], but the optimal type, intensity, frequency, and duration of these exercises are not clear [27-31]. Our exercise programme was constructed by JT in cooperation with three physiotherapists from the orthopaedic shoulder department at Silkeborg Regional Hospital (SRH). Based on studies showing effect of exercise programmes [23-26, 32], easily learned exercises were selected taking into account elements known to motivate exercise adherence (e.g. a limited number of exercises) [33]. The programme consists of four exercises: one posture corrective exercise and three resistance exercises, performed bilaterally with an elastic band (Thera-band©). The three resistance exercises, each with three levels, consist of two exercises for the scapula stabilising muscles (wall slide and low row/high row) and one for the rotator cuff muscles (external rotation). Participants are recommended to start with the exercises at level one, and to perform 3 sets of up to 15 repetitions 3–
4 times per week during the intervention period and preferably also thereafter. When a participant is able to perform 3 sets of 15 repetitions of an exercise without aggravating pain (lasting > 1 hour after exercise), he/she is encouraged to progress to the next level of that particular exercise.

- General information on occupational shoulder exposures and how to reduce them, described in a pamphlet (Additional file 3). The pamphlet, developed by AD in collaboration with PF, SWS, and SDC, focuses on work with elevated arms, repetitive shoulder movements, and forceful shoulder exertions. It is based on previous assessments of occupational shoulder exposures [16], exposure-response relationships with shoulder disorders [5-8], and years of experience from work as occupational health physicians (PF and SWS) and as a health and safety consultant (SDC).

- Assessment of occupational shoulder exposures based on:
  - Technical measurements of postures and movements performed using an Axivity (AX3) accelerometer [34], processed to yield minutes/day with the arms elevated > 30°, > 60°, and > 90°, and median angular velocity (°/s) (as a measure of repetition) during work. Axivity measurements are performed on the more affected shoulder (right shoulder in case of similar symptoms). The participants are instructed to wear the accelerometer for at least 1 and preferably 5 working days and to register working hours (start and stop times), main tasks, and whether it was a typical working day in a work diary. Data from one measurement day of ≥ 4 hours per person is considered enough for characterisation at the group level.
  - Self-reported estimates of the average level of forceful shoulder exertions for each working day using the Borg CR-10 [35].

Exposure assessment is performed shortly after the 1st café meeting / intervention contact and shortly after EOI (see Table 1). All participants receive individual written feedback on their shoulder exposures after these two exposure assessment periods (Additional file 4).

**Shoulder-Café**

A Shoulder-Café course includes 3 café meetings spaced around 6 weeks apart. The principal investigator (JT) will attend all 1st and 3rd café meetings. Each café meeting lasts for about 2 hours and includes 15–30 minutes of small talk and exchange of experiences over a cup of coffee / tea to secure social networking and interpersonal relationships. In addition, a Shoulder-Café course contains:

- Individually tailored shoulder exercises (in accordance with the exercise pamphlet, additional file 2), supervised by physiotherapists from the six municipality health centres. At each café meeting, the attending physiotherapist spends 1 hour demonstrating the exercises, correcting participants performing the exercises, and answering questions in relation to the exercises. To secure fidelity, the physiotherapists have attended a training session led by JT prior to the 1st café meeting and follow a pre-defined guideline (Additional file 5).

- A clinical shoulder evaluation of each participant performed at the 1st café meeting by a physiotherapist according to a pre-specified form (Additional file 6) and protocol. The protocol is...
Based on the Danish guideline for diagnosing patients with shoulder complaints [9] and was developed by JT in cooperation with 3 physiotherapists from the orthopaedic shoulder department at SRH, an orthopaedic surgeon (TK), and two occupational health physicians (PF and SWS). The aim of the examination is to clinically characterise the participants. If, as an exception, a participant is identified with a “red flag” (e.g. progressive non-mechanical pain or weight loss) [36], he/she is advised to contact his/her general practitioner and a statement regarding advice against exercise is recorded; the participant will still be included in the intention-to-treat analyses. The three physiotherapists, who take turns performing the examinations, had been physiotherapists for 12–18 years, had special training in clinical evaluation of shoulder complaints, and had worked 3–7 years in the orthopaedic shoulder department at SRH at the start of the interventions.

- Education about shoulder anatomy (Additional file 7) for 45 minutes at the 1st café meeting is provided by the above-mentioned experienced physiotherapists. The goal is to educate participants in the taking of appropriate action to reduce their shoulder complaints.

- Workplace-oriented counselling focussing on reducing shoulder exposures. The counselling is given by a health and safety consultant (SDC), who had been a physiotherapist for 18 years and had been working as health and safety consultant for 14 years at the start of the He has 45 minutes at his disposal at the 2nd café meeting (Additional file 8), where he also answers questions about the individual feedback on shoulder exposures (additional file 4). The counselling is based on theories from “The motivational conversation” [37], “Stages of change” [38], and “The health belief model” [38] in order to increase the participants’ motivation for self-generated changes. There is also time to discuss organisational and other factors which might be barriers for work modifications. Previous experience indicates that health and safety advice is less likely to be implemented if the advice is too general or will take long time to implement [39]. Therefore, our focus is on feasible and specific work modifications that can be implemented within a short time frame, i.e., modifications that are cheap, uncomplicated, and fit workplace conditions. Advice on more far-reaching modifications may be given, too. A workplace visit by the health and safety consultant is an option when necessary to find ways to reduce the shoulder exposures. Plans of action that are based on a workplace visit are often focussed and clearly outlined, which increases their chances of being implemented [39]. The workplace visits are attended by the health and safety consultant, the participant, a working environment representative, and, if possible, the employer/supervisor. Initially, 1–3 tasks are prioritised which entail high shoulder exposures and are difficult to perform while having shoulder complaints. Again, the focus is on specific work modifications that are feasible within a short time frame. The advice is documented by the health and safety consultant and categorised as ways to reduce high task exposures (technical solutions) and ways to reduce the duration of tasks with high exposures (organisational solutions) for the individual participant. After the workplace visit, the health and safety consultant sends a summary of the advice to the employee, the working environment representative, and the employer/supervisor. We have resources for a maximum of 50 1-hour workplace visits.
The physiotherapists, who supervise the exercises and perform the clinical examinations, and the health and safety consultant are financially compensated by the project.

**Shoulder-Guidance**

The Shoulder-Guidance includes an initial 20–30-minute individual appointment, staffed by a physiotherapist student or a project physiotherapist; the remaining parts of the guidance are delivered as postal letters or emails.

**Outcome measures**

Table 2 provides the time schedule of the trial and the timing of assessments of primary, secondary, and supplementary outcomes as well as assessments of baseline characteristics and measures of adherence and adverse events.

**Primary outcomes**

In relation to hypothesis I

The primary outcome is the OSS at 6 month follow-up. We chose a patient-reported outcome [40] which directly measures the participants’ shoulder complaints. The OSS has been translated and cross-culturally adapted to Danish [19] and is a valid, reliable, and responsive shoulder-specific measure [17, 41-44]. It is one of the recommended first choice instruments in patients with shoulder disorders [45]. The OSS was developed for patients undergoing shoulder surgery [17], but has also been used in patients who have not been operated on [43, 44] and asymptomatic persons [20, 21]. Follow-up after 6 months was chosen to allow the potential effects on shoulder pain and disability to evolve.

In relation to hypothesis II

The primary outcome is work with the arm elevated > 60° (minutes/day) according to Axivity measurements shortly after EOI. This outcome was chosen based on the available evidence on work with elevated arms [3, 5-7] and because we think that this measure will be more responsive to change than minutes/day with the arm elevated > 90°, which occurs to a limited extent in some of the included occupations. The timing was chosen because we expect that most work modifications will occur within the intervention period and because we want to use the 2nd measurement feedback to motivate the participants for further work modifications.

**Secondary outcomes**

In relation to hypothesis I

Listed in order of priority, the secondary outcomes are:
• The OSS at 12 month follow-up. We added this time point because increasing effects of a training intervention 12 months after T₀ has been reported previously [24].

• The Fear-Avoidance Beliefs Questionnaire – Physical Activity (FABQ-PA) [46] at 6 month follow-up in a version modified for the shoulder [47]. FABQ-PA contains four items about shoulder pain in relation to physical activity [46-48]. The score ranges from 0–24 with higher scores reflecting a higher tendency for fear-avoidance beliefs [46]. According to the fear-avoidance model, pain-related fear may cause people to avoid physical activities (including shoulder exercises), and reduction of an exaggerated reaction pattern of this kind may be part of the intervention's mechanism of action [48-50].

• Patients’ global impression of change (PGIC) [51] at 6 month follow-up, which reflects the participants’ general impression of change with regard to their shoulder condition rated on a 7-point Likert scale ranging from 1 (much better) to 7 (much worse) (https://www.sciencedirect.com/science/article/pii/S2287888215300684). Our a priori definition of improvement is the range 1 “Much better”, 2 “Better”, and 3 “A little better”.

• The FABQ-PA [46] at 12 month follow-up.

In relation to hypothesis II

Listed in order of priority, the secondary outcomes are:

• Minutes/day working with the arm elevated > 90° according to Axivity measurements shortly after EOI.

• Mean median angular velocity (°/s) according to Axivity measurements shortly after EOI.

• Average forceful shoulder exertions assessed by the Borg CR-10 scale [35] shortly after EOI.

• Minutes/day working with the arm elevated > 30° according to Axivity measurements shortly after EOI.

Supplementary outcomes

In relation to hypothesis I

Intensity of shoulder pain at rest and during activity measured on a numerical rating scale (NRS, ranging from 0 to 10), quick version of the Disabilities of the Arm, Shoulder and Hand (quick DASH) and work module [52], health-related quality of life using the EQ5D-3L [53]), work ability using the Work Ability Score [54, 55], PGIC at 12 months follow-up, overall satisfaction with the intervention at 6 and 12 months, and the degree to which the participant felt sufficiently informed about 1) how to handle shoulder complaints, 2) how to perform shoulder exercises, 3) how to reduce occupational shoulder exposures at 6 months follow-up (5 point scale).

In relation to hypothesis II
Work modifications measured at 6 month follow-up.

**Supplementary outcome measures will be selected from these variables.**

**Other assessments**

Other baseline assessments were smoking status, body mass index, duration of shoulder complaints, psychosocial work exposures (job demands, job control, and social support based on the Karasek-Theorell model) [56], occupational mechanical shoulder exposures (self-reported upper arm elevation, repetitive shoulder movements, forceful shoulder exertions, and use of vibrating tools). In addition job title, weekly working hours, and system of wage payment were assessed at baseline and at 12 month follow-up and work status was assessed as 12 month follow-up. At 6 and 12 months follow-up, all participants were also asked how often exercise was performed.

**Adherence**

Adherence to the home-based exercise programme is monitored using an exercise diary and a BandCizer© sensor mounted on the elastic band (Thera-band©). The BandCizer© records the exercise dose quantified as time under tension [57-59]. Adherence to the exposure assessment will be described as the percentage of the participants that has ≥ 1 work day with ≥ 4 hours of Axivity data and/or a Borg CR-10 rating in the 1st and in the 2nd exposure assessment period. For the Shoulder-Café group, adherence to café meetings will also be described (Table 2).

**Co-interventions and adverse events**

The questionnaires at 6 and 12 month follow-up will ask about co-interventions and adverse events (Table 2).

**Data collection and data management**

All questionnaires will be collected by the principal investigator (JT). Companies will be reminded by email and phone if few or no screening questionnaires have been returned after 1–2 months. Participants who do not return the follow-up questionnaires will be reminded to do so by email and finally by postal letter. Data from paper screening questionnaires will be scanned by PostNord [60]. Data from electronic screening, baseline, and follow-up questionnaires will be directly captured in REDCap (version 7.4.17, Vanderbilt University), while data from paper versions of baseline and follow-up questionnaires and from exercise diaries will be manually entered into REDCap. Data from the BandCizer© will be processed to yield date, number of training sessions, number of exercise sets, number of repetitions, time under tension for each repetition, and total time under tension for each training session. Variables based on data from the BandCizer© will be entered into REDCap. Axivity data (Axivity Ltd, Newcastle, United Kingdom) will be downloaded using OmGui open-source software (OmGui Version 1.0.0.28; Open Movement, Newcastle University, Newcastle upon Tyne, United Kingdom) and saved in raw format files. MatLab (Build
8.6.0.267246 (R2015b) 64 bit) and STATA 15 (StataCorp LP, College Station, TX, US) will be used for data processing and statistical analyses. Data cleaning will be documented in Stata do files.

Blinding

Blinding of participants and care providers is not possible due to the character of the interventions. To prevent this from influencing the answers on the OSS and other patient-reported outcomes, all participants receive an active intervention. With respect to shoulder exposures, the outcome assessor (AD) will be blinded to intervention arm. We have developed a statistical analysis plan (SAP) to minimise the risk of analysis bias (Additional file 9).

Sample size

We aim to be able to show a difference between the groups of at least 5 points in the OSS [61, 62] at six month follow-up. With an expected SD of 8 points [24], an intraclass correlation coefficient of 0.05 [63, 64], and a mean cluster-size of four, the study size needs to be \( \geq 96 \) (2 x 48) with a two-sided significance level of 0.05 and a power of 0.80. We aim to include 60 employees in each group to ensure that 50 employees in each group complete the study. Power calculations were carried out with Stata 15 (StataCorp LP, College Station, TX, USA; power twomeans with cluster option).

Statistical methods

All analyses will be performed according to the intention-to-treat principle. Regarding hypothesis I, a mixed model analysis of the OSS will be performed including "intervention" (Shoulder-Café and Shoulder-Guidance), "time" (6 and 12 month follow-up), "intervention x time", baseline OSS, sex, age, and industry (service, manufacture, construction) as fixed effects, adjusting for random effects of participant and company (cluster). FABQ-PA will be analysed likewise, but will be adjusted for baseline FABQ-PA instead of baseline OSS. In the analysis of PGIC at 6 months the outcome will be dichotomised as described above. We will use a risk difference model if around 50% of the participants improve. If a considerably smaller percentage (< 20%) improves, we will employ a relative risk model using improved as the outcome, while if a considerably larger percentage (> 80%) improves, we will employ a relative risk model using not improved as the outcome. The analysis of PGIC will be adjusted for sex, age, and industry and use robust standard errors to take into account clustering at company level.

Regarding hypothesis II, a mixed model analysis of the primary outcome (minutes/day working with the arm elevated > 60°) will be performed including "intervention" (Shoulder-Café and Shoulder-Guidance), baseline minutes/day working with the arm elevated > 60°, sex, age, and industry (service, manufacture, construction) as fixed effects, adjusting for random effects of company (cluster). The analyses for the secondary outcomes will be performed likewise, but will be adjusted for the respective baseline values instead of the baseline number of minutes/day working with the arm elevated > 60°.

If no more than two questions in the OSS are left unanswered, single mean imputation will be used [18], otherwise the total score will be left missing. Axivity measures are considered missing in case of < 4
hours of measurement data during 1 working day. Loss to follow-up will be addressed by sensitivity analyses comparing realistic scenarios; subgroup analyses are not intended. Additional information is described in the SAP (Additional file 9).

**Harms and data monitoring**

The intervention is based on non-invasive methods and is not expected to cause any adverse events other than possible temporary muscle tenderness after shoulder exercises. Therefore, no data monitoring committee has been established and no stopping rules defined. Any unexpected serious adverse event will be reported to the Committee on Health Research Ethics in Central Denmark Region within seven days after the principal investigator has become aware of the event.

**Publication policy**

Hypotheses I and II will be addressed in separate publications. The main publication regarding hypothesis I will be prepared first and the main publication regarding hypothesis II shortly thereafter. We intend to publish positive, negative, and inconclusive results. Authorship will be determined in accordance with the recommendations of the International Committee of Medical Journal Editors. Furthermore, we plan to disseminate the results to key stakeholders through the projects’ stakeholder group. The authors do not have any publication restrictions.

**Satellite studies**

Two prospective cohort studies are planned based on the cluster-randomised trial. One study, with the OSS as the primary outcome, will investigate the relative influence of shoulder exercises and reduced occupational shoulder exposures on shoulder complaints. Another study will investigate the intensity of shoulder pain at rest and during activity (NRS) monitored week-by-week with short message service as a predictor of subsequent weekly exercise dose, and the potential influence of fear avoidance beliefs on this relationship. Further, a process evaluation [65, 66] is nested in the trial to assist later contextualisation of the outcomes. The findings from this may point to areas that warrant further consideration or development prior to a potential wider implementation of the Shoulder-Café intervention. The process evaluation employs semi-structured interviews [67] with six participants from the Shoulder-Café (n = 2) and Shoulder-Guidance (n = 4) conducted 1 month after EOI and 12 observations [68] of Shoulder-Café (n = 9) and Shoulder-Guidance (n = 3) sessions. All interviews and observations are supervised by a senior project participant (MTH).

**Discussion**

Several studies have found that exercise is effective in reducing shoulder complaints [23-29, 31, 69, 70], but optimal ways to exercise remain to be established. Few studies have evaluated interventions that have addressed occupational shoulder exposures in order to prevent or reduce shoulder complaints [71-73]. The disappointing results of these studies may be related to the fact that for the most part they were
completed in office environments and health care settings, where shoulder exposures are at most moderate to begin with [71-73]. Only one study that we are aware of included participants with high shoulder exposures, but did not document whether the intervention reduced the exposures [74]. The combination of shoulder exercises and workplace-oriented advice using a café concept is a novel approach.

The strengths of this study are the randomised controlled design, cluster randomisation at company level to prevent contamination between groups, use of validated patient-reported outcomes to assess shoulder complaints, and technical measurements of shoulder postures and movements.

Stigmatisation of employees with shoulder complaints is avoided as the intervention takes place outside the company and after working hours. This enables participants to decide whether they want to inform their workplace about their participation.

A limitation of the study is the inability to blind participants to the intervention, but both groups receive an active intervention in order to reduce the risk of biased outcome reporting. Baseline assessment of occupational shoulder exposures takes place after the randomisation result has been revealed. However, Axivity accelerometers are mounted on all participants at their 1st intervention appointment and we use technical measurements performed on several working days. This should guard against differential participation and differential misclassification of occupational shoulder exposures. Additionally, participants and non-participants will be compared with respect to self-reported occupational shoulder exposures according to the baseline questionnaire.

A further limitation is that it is not possible to differentiate between the separate effects of exercise, work modification, diagnostic clarification, education, and workplace-oriented counselling on the participants’ shoulder complaints, but the analyses in relation to hypothesis II and one of the planned satellite studies will reveal to which extent reduced occupational shoulder exposures may have played a part. To give a further indication of the relative influence of the intervention elements, we will ask the participants at 6 month follow-up to which degree they feel that the intervention provided them with sufficient knowledge about (1) how to handle shoulder complaints, (2) how to exercise, and (3) how to reduce their shoulder exposures. The process evaluation may aid in this evaluation. If shoulder exposures are reduced by handing over high-load tasks to colleagues, the problem may only be relocated. On the other hand, the possibility of exposure modification in periods with increased pain may be in all employees’ favour.

If the results turn out positive, we believe that the Shoulder-Café intervention has the potential to be implemented on a larger scale. The pilot study café intervention is already implemented in three municipalities in Central Denmark Region, and the project has a stakeholder group to back up the process. Further, it should be possible to develop the intervention to involve other musculoskeletal regions, which has already been requested by one of the participating municipalities.

**Abbreviations**
CI: Confidence interval

EOI: End of intervention

FABQ – PA: Fear Avoidance Beliefs Questionnaire – Physical Activity

OSS: Oxford Shoulder Score

PGIC: Patients’ Global Impression of Change

SAP: Statistical analysis plan

SRH: Silkeborg Regional Hospital

T₀: Start of intervention

Declarations

Trial Status

Protocol version 1.0: Issue date: 22 January 2019. Recruitment of participants started in May 2017 and is ongoing. Recruitment of participants is expected to end no later than June 2019.

Ethics approval and consent to participate

The trial protocol, the informed consent forms and other requested documents have been reviewed and approved by The Danish Data Protection Agency on 7 September 2016 (case number: 1-16-02-498-16) and The Committee on Health Research Ethics in Central Denmark Region approved the trial on 20 March 2017 (case number: 1-10-72-271-16). Written consent is obtained from all participants. The trial was registered at Clinicaltrials.gov on 18 May 2017 (ID: NCT03159910). Important protocol modifications will be communicated to these agencies.

Consent for publication

Written informed consent was obtained from the person appearing in the exercise pamphlet (additional file 2). A copy of this consent is available for review by the Editor of this journal.

Availability of data and materials

Not applicable since this manuscript is a study protocol.

Competing interests

None of the authors have any competing interests.

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Authors' contributions

The trial was planned by all authors. JT is responsible for participant recruitment and data collection and has drafted this manuscript in close collaboration with SWS. All authors have revised the manuscript for important intellectual content and have read and approved the final manuscript. All authors will have access to the final trial dataset.

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Tables

Table 1 Content and time schedule of the Shoulder-Café and the Shoulder-Guidance.
| **Shoulder-Café** | **Shoulder-Guidance (active control – enhanced usual care)** |
|-------------------|-------------------------------------------------------------|
| **1st café meeting (T₀):** | **1st intervention contact - individual appointment (T₀):** |
| · Distribution of home-based exercise pamphlet, BandCizer®, Axivity accelerometers*, diaries, and elastic bands | · Distribution of home-based exercise pamphlet, BandCizer®, Axivity accelerometers*, diaries, and elastic bands |
| · Presentation of participants and networking with the group |  |
| · Supervised exercises with individual tailoring according to the exercise pamphlet |  |
| · Clinical evaluation of the participants’ shoulders |  |
| · Education about shoulder anatomy |  |
| **At home:** | **At home:** |
| · Home-based exercises and exercise diary | · Home-based exercises and exercise diary |
| **At work:** | **At work:** |
| · Shoulder exposure assessment and work diary | · Shoulder exposure assessment and work diary |
| **2nd café meeting (~1.5 month after T₀):** | **2nd intervention contact – postal letter or email (~1.5 month after T₀):** |
| · Written feedback on the 1st exposure assessment | · Written feedback on the 1st exposure assessment |
| · Written general advice on reduction of occupational shoulder exposures | · Written general advice on reduction of occupational shoulder exposures |
| · Supervised exercises with individual tailoring according to the pamphlet |  |
| · Education about shoulder exposures |  |
| · Advice on work modifications and possibility to ask questions about the 1st exposure assessment |  |
| · Offer of a workplace visit to find ways to reduce the exposures |  |
| · Networking with the group |  |
| **At home:** | **At home:** |
| · Home-based exercises and exercise diary | · Home-based exercises and exercise diary |
3rd café meeting (end of intervention ~3 months after $T_0$):
- Distribution* of Axivity accelerometers and work diaries
- Supervised exercises with individual tailoring according to the pamphlet
- Networking with the group

3rd intervention contact – postal letter (end of intervention ~3 months after $T_0$):
- Distribution of Axivity accelerometers and work diaries

At work:
- Shoulder exposure assessment and work diary

Postal letter or email:
- Written feedback on the exposure assessment shortly after end of intervention

6 month follow-up (~6 months after $T_0$):
- Electronic or postal questionnaire

6 month follow-up (~6 months after $T_0$):
- Electronic or postal questionnaire

12 month follow-up (~12 months after $T_0$):
- Electronic or postal questionnaire

12 month follow-up (~12 months after $T_0$):
- Electronic or postal questionnaire

* The Axivity accelerometer is mounted, unless the participant is going on holiday or expects atypical work, e.g. due to course participation.

**Table 2:** Schedule for study procedures. For each batch of companies, the two interventions (Shoulder-Café and Shoulder-Guidance) start and end simultaneously.
| Time point                  | Pre-intervention | Intervention | Follow-up |
|----------------------------|------------------|--------------|-----------|
|                             |                  | 1st day after | Shortly after | Shortly after | 6 months after | 12 months after |
|                             |                  | (T₀) T₀       | EOI 3 months T₀ | EOI T₀ | T₀ | T₀ |

**ENROLMENT**

- Company randomisation *  
- Informed consent  
- Revealing randomisation result to participants **

**INTERVENTION**

- Shoulder-Café  
- Shoulder-Guidance

**OUTCOME ASSESSMENTS**

**Primary outcomes**

- **Hypothesis I**
  - OSS at 6 month follow-up

- **Hypothesis II**
  - Arm elevation > 60°

**Secondary Outcomes**

- **Hypothesis I**
  - OSS at 12 month follow-up
  - FABQ-PA at 6 month follow-up
  - PGIC at 6 month follow-up
  - FABQ-PA at 12 month follow-up

- **Hypothesis II**
  - Arm elevation > 90°
  - Repetitive shoulder movements  
  - Forceful shoulder exertions  
  - Arm elevation > 30°

**Supplementary outcomes**

- Hypothesis I
  - NRS at rest
  - NRS during activity
  - Quick DASH + work module
  - EQ5D-3L
  - Work Ability Score
  - PGIC at 12 month follow-up

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*Note: x indicates the time point of assessment.*
Satisfaction questionnaire  
Felt informed about how to handle shoulder complaints, how to perform shoulder exercises, and how to reduce occupational shoulder exposures (questionnaire)  

Hypothesis II  
Work modification  

OTHER ASSESSMENTS  
Smoking status  
Body mass index  
Duration of shoulder complaints  
Psychosocial work exposures  
Occupational mechanical shoulder exposures (self-reported)  
Job title  
Weekly working hours  
System of wage payment  
How often exercise was performed  
Work status  
Typical working day  

ADHERENCE  
Café meetings ****  
Exercise diary  
Work diary  

CO INTERVENTIONS  
Steroid injection  
Shoulder surgery  
Seen by doctor because of shoulder complaints  
Shoulder treatment by physiotherapist outside the project  
Shoulder treatment by chiropractor  
Pain medication in last 4 weeks  

ADVERSE EVENTS
* Randomisation of a batch of companies takes place around 2 weeks before the start of an intervention. ** The randomisation result is revealed to the participants after the baseline questionnaire is filled in. *** The baseline questionnaire is scheduled to be filled in a few days before the 1st intervention day, but may be filled in on the 1st day (see **). **** For the intervention group, only.

Abbreviations: quick DASH = Disabilities of the Arm, Shoulder and Hand – quick version, EOI = end of intervention, EQ5D-3L = EuroQol 5D-3L, FABQ-PA = Fear Avoidance Beliefs Questionnaire – Physical Activity, NRS = numerical rating scale (range 0–10), OSS = Oxford Shoulder Score, PGIC = Patients’ Global Impression of Change, $T_0$ = start of intervention.

**Figures**
Figure 1

Expected flow of participants through the study.

Supplementary Files

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