Effect of specific resistance training on forearm pain and work disability in industrial technicians: cluster randomised controlled trial

Lars L Andersen,1 Markus D Jakobsen,1 Mogens T Pedersen,2 Ole S Mortensen,3 Gisela Sjøgaard,4 Mette K Zebis1,4

ABSTRACT

Objectives: To determine the effect of specific resistance training on forearm pain and work disability in industrial technicians.

Design and setting: Two-armed cluster randomised controlled trial of 20 weeks performed at two industrial production units in Copenhagen, Denmark.

Participants: Working-age industrial technicians both with and without pain and disability.

Interventions: The training group (n=282) performed specific resistance training for the shoulder, neck and arm muscles three times a week. The control group (n=255) was advised to continue normal physical activity.

Outcome: All participants rated forearm pain intensity (Visual Analogue Scale, 0–100 mm) once a week (primary outcome) and replied to a questionnaire on work disability (Disability of the Arm Shoulder and Hand, 0–100) at baseline and follow-up (secondary outcome).

Results: Questionnaires were sent to 854 workers of which 30 (n=282) and 27 (n=255) clusters were randomised to training and control, respectively. Of these, 211 and 237 participants, respectively, responded to the follow-up questionnaire. Intention-to-treat analyses including both individuals with and without pain showed that from baseline to follow-up, pain intensity and work disability decreased more in the training group than in the control group (4–5 on a scale of 0–100, p<0.01–0.001). Among those with pain >30 mm Visual Analogue Scale at baseline (n=54), the OR for complete recovery at follow-up in the training group compared with the control group was 4.6 (95% CI 1.2 to 17.9). Among those with work disability >30 at baseline (n=113), the OR for complete recovery at follow-up in the training group compared with the control group was 6.0 (95% CI 1.8 to 19.8).

Conclusion: Specific resistance training of the shoulder, neck and arm reduces forearm pain and work disability among industrial technicians.

Trial registration number: NCT01071980.

INTRODUCTION

Musculoskeletal disorders are associated with considerable work disability, healthcare costs and sickness absence.1–3 Almost all employees depend on well-functioning upper extremities for daily work tasks. Frequent repetitive movements of the arm or wrist have been associated with development of forearm pain4 and lateral epicondylitis (tennis elbow).5 A recent systematic review quantified the dose–response relationship between work exposure and disorders and reported that repetitive movements for more
than 2 h a day increased the risk for lateral epicondylitis.\(^6\) Data from 2003 showed that the 1-year incidence of forearm pain among newly employed workers across different occupations was 8\(\%\)\(^4\) and the weekly prevalence of moderate-to-severe forearm pain among computer users was 4\(\%\).\(^7\) The aetiology of forearm pain is related to both physical and psychosocial distress.\(^8\) \(^9\) As many occupations inherently involve hand–arm exposure complementing interventions to support existing workplace, ergonomic adjustments should be considered.

General health benefits of regular physical exercise are well known.\(^10\) As most adults spend many hours at work several days a week, the workplace provides a feasible setting for health promotion. While previous studies have documented the effectiveness of workplace interventions with physical exercise for preventing and relieving neck/shoulder and back pain,\(^11\) none of these studies concerned forearm pain. Randomised controlled trials have shown mixed results on the effectiveness of specific resistance training for specific forearm disorders in patients, for example, lateral epicondylitis.\(^12\) \(^13\) We have shown in one study that upper body resistance training can to some extent reduce pain symptoms of the elbow and wrist among office workers.\(^14\) However, the effect of such interventions on work disability is unknown. Thus, randomised controlled trials investigating the effect of workplace physical exercise interventions on forearm pain and work disability are needed. In workplace interventions involving the majority of employees, randomisation at the cluster level (eg, department) is preferred to avoid contamination of interventions.\(^15\)

This cluster randomised controlled trial determines the effect of specific resistance training on forearm pain and work disability in industrial technicians. We hypothesise that specific resistance training reduces pain and work disability.

**METHODS**

**Study design**

We performed a cluster randomised controlled trial in Copenhagen, Denmark, from January (recruitment) to June 2009 (follow-up). All employed technicians from two large industrial production units from the public and private sector were invited to participate. The private sector company specialised in creating biotechnological products by using enzymes and the public sector company specialised in production of vaccines and control of infectious diseases. At both companies, the work involved repetitive tasks, such as pipetting, preparing vial samples for analysis and data processing on a computer including mouse work, all tasks that require precision in work and may result in extended periods of time spent in static working postures.

Pain and disability were not specific inclusion criteria. The rationale for this was that musculoskeletal pain typically is episodic. Including all employees allowed us to study the effect of the intervention at the company level, as well as the treatment and preventive effect, respectively, among subgroups with and without pain at baseline. The main outcome on neck/shoulder pain is described elsewhere.\(^16\) Figure 1 shows the flow of participants through the trial.

In brief, we sent an internet-based questionnaire to 854 prospective participants of which 669 replied. One hundred and four declined to participate or did not reply to the question concerning participation. Exclusion criteria—which led to exclusion of 28 participants—were pregnancy and serious health conditions, such as previous trauma or injuries, life-threatening diseases and cardiovascular disease. Thereby 537 participants were included in the study and randomly assigned at the cluster level to training (n=282 in 30 clusters) or control (n=255 in 27 clusters) (table 1).

All participants gave their written informed consent to participate in this study, which conformed to the Declaration of Helsinki and was approved by the local ethical committee (HC2008103). Trial registration: http://ClinicalTrials.gov (NCT01071980).

**The cluster randomisation procedure**

Eligible volunteers were randomised at the cluster level to training or control. The statistician performing the cluster randomisation had no access to the baseline database. Departments were stratified into 14 strata according to the following nested criteria: company, type of work task and size of department. Strata were labelled alphabetically, and clusters were numbered consecutively within strata. This resulted in a total of 57 clusters with sizes ranging from 1 to 40 participants. A statistician blinded to the identity of the strata, and clusters assigned the clusters within each stratum by simple random allocation to either the training or the control. The consecutive numbers of the clusters within each stratum were written on pieces of paper and drawn from an opaque, tossed plastic bag. To minimise imbalance over several strata with odd numbers of clusters, these strata were paired, and clusters were alternately allocated to either training or control, the first cluster being allocated to either training or control depending on the flip of a coin. Thus, all clusters had the same chance of being allocated to the training group.

**Intervention**

The training group performed specific resistance training for the shoulder, neck and arm with dumbbells (wrist extension, shoulder lateral raise, shoulder front raise, shoulder shrugs, reverse flies) three times of 20 min/week for 20 weeks. Training loads were progressively increased from moderate loadings of 15–20 repetitions maximum (RM)—ie, the number of repetitions that could be performed to momentary fatigue) during the initial weeks to relatively heavier loadings of 8–12 (RM) during the final weeks. The exercises were performed in a slowly controlled manner avoiding sudden jerk and acceleration. Experienced instructors introduced the training programme in
groups of 5–15 colleagues. After the introductory session, the participants had the opportunity to train individually or in self-organised groups. The instructors supervised every other training session throughout the 20 weeks to ensure correct technique and sufficient progression of the training. The supervised sessions were scheduled with assistance from the respective departments to best fit into daily working routines. The remaining training sessions were openly planned, meaning that the participants could train whenever it matched their daily work schedule. The training facilities were placed as near as possible to the workstations. In practice, training locations were established in corridors, storerooms and conference rooms (11 locations in total). All locations were equipped with a training poster, a clock, chairs, two pairs of lifting straps and dumbbells (pairs of 1–25 kg). Participants of the control group were advised to continue their normal physical activity as usual.

Adherence
Participants of the training group logged their training in a diary during each training session. Adherence was defined as the total number of training sessions as a percentage of the maximal possible number of training sessions during the 20 weeks (ie, 60 sessions).

Forearm pain and work disability
The outcomes were based on questionnaire replies. Therefore, blinding of ‘outcome assessors’ does not apply for the present study. In principle, the participants were outcome assessors as we used subjective rating scales. Participants could not be blinded, that is, after the randomisation, each participant was informed about his/her respective group allocation. However, the statistical analyses remained blinded by assigning each group an arbitrary number until after the pre-planned analyses were run.

Primary outcome
Once a week, participants of both groups rated forearm pain intensity at its worst during the previous 7 days on a 100 mm Visual Analogue Scale (VAS), where 0 mm is ‘no pain at all’ and 100 mm is ‘worst possible pain’.17

Secondary outcome
Participants rated work disability at baseline and follow-up by the work module of the Disability of the Arm Shoulder and Hand (DASH) questionnaire, which has previously been validated among industrial workers18: “In the past week did you have any difficulty:” (1) “using your usual technique for your work?”, (2) “doing your usual work because of arm, shoulder or hand pain?”.
Participants replied on a 5-point Likert scale from ‘no difficulty’ to ‘unable’. For comparability with VAS pain scores, the work disability score was normalised on a scale of 0 to 100, where 100 represents the highest level of disability.19

Kaergaard and coworkers20 found that for four complaint questions using 10-point scales, a sum of 12 (ie, average 3 for each of the four questions) was significantly associated with clinical findings. Because we used scales from 0 to 100, we defined pain and disability cases accordingly as those scoring more than 30 on a scale of 0 to 100.

**Sample size**
A priori power analyses showed that a sample size of 120 participants in each group would provide a power of 80% to detect a 15% change in pain. At an estimated dropout or loss to follow-up of 20%, the minimally required number of participants in each group should be 150 in a design with simple randomisation. Calculations based on neck pain intensity from a previous cluster randomised study21 showed that the intra-cluster correlation (ρ) was close to zero. However, we made a conservative estimate and assumed that ρ would be equal to 0.1 and the average cluster size (m) would be equal to 10. Using the formula $1 + (m−1)ρ$,15 we would then need 1.9 times the sample size compared with not having used simple randomisation, that is, 285 participants in each group.

**Statistics**
We performed all analyses in accordance with the intention-to-treat principle and used a generalised linear mixed model (GLIMMIX procedure of SAS) controlled for gender to determine differences in the main outcomes between the two groups from baseline to follow-up. We added a random cluster effect to the model to account for possible intra-cluster correlations. We did not impute missing data as all methods of data imputation have limitations.22 The GLIMMIX procedure inherently accounts for missing data.

Work disability was registered twice, at baseline and follow-up. Pain intensity was registered weekly, and therefore, each participant could have up to 20 registrations. To more accurately report the effect of the intervention on pain, we used linear regression for each participant to estimate the trend over time and thereby determined the change from baseline to follow-up by multiplying the slope with the number of weeks. For the linear regression on pain intensity, we included participants with at least two observations at least 10 weeks apart. The individual changes from baseline to follow-up were then entered in the GLIMMIX procedure. These analyses were pre-planned.

**Treatment and prevention of symptoms**
Using logistic regression analysis controlled for gender and age (Logistic procedure of SAS), we also calculated ORs for complete recovery at follow-up (defined as pain intensity or disability score of <10) among cases at baseline (defined as pain intensity or disability score of

---

**Table 1** Demographics, clinical and work-related characteristics of the participants at baseline

<table>
<thead>
<tr>
<th></th>
<th>Randomised Control</th>
<th>Randomised Training</th>
<th>Pain cases Control</th>
<th>Pain cases Training</th>
<th>Disability cases Control</th>
<th>Disability cases Training</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants</td>
<td>255</td>
<td>282</td>
<td>22</td>
<td>32</td>
<td>44</td>
<td>69</td>
</tr>
<tr>
<td>Demographics</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age, year</td>
<td>43 (9)</td>
<td>44 (11)</td>
<td>45 (9)</td>
<td>45 (11)</td>
<td>43 (9)</td>
<td>45 (10)</td>
</tr>
<tr>
<td>Height, cm</td>
<td>170 (8)</td>
<td>168 (7)</td>
<td>171 (8)</td>
<td>166 (7)</td>
<td>170 (6)</td>
<td>168 (7)</td>
</tr>
<tr>
<td>Weight, kg</td>
<td>73 (14)</td>
<td>70 (14)</td>
<td>72 (12)</td>
<td>69 (17)</td>
<td>71 (14)</td>
<td>69 (14)</td>
</tr>
<tr>
<td>Body mass index, kg/m²</td>
<td>25 (5)</td>
<td>25 (4)</td>
<td>25 (4)</td>
<td>25 (5)</td>
<td>25 (5)</td>
<td>24 (4)</td>
</tr>
<tr>
<td>Women (%)</td>
<td>80%</td>
<td>90%</td>
<td>86%</td>
<td>97%</td>
<td>93%</td>
<td>90%</td>
</tr>
<tr>
<td>Clinical</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Forearm pain intensity (0–100 mm VAS)</td>
<td>10 (16)</td>
<td>12 (18)</td>
<td>52 (19)</td>
<td>52 (17)</td>
<td>23 (28)</td>
<td>18 (24)</td>
</tr>
<tr>
<td>Work disability (0–100)</td>
<td>15 (21)</td>
<td>19 (22)</td>
<td>36 (26)</td>
<td>26 (24)</td>
<td>54 (19)</td>
<td>50 (18)</td>
</tr>
<tr>
<td>Percentage of participants spending more than half of total work time</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Working with the hands twisted or flexed</td>
<td>27%</td>
<td>34%</td>
<td>45%</td>
<td>44%</td>
<td>36%</td>
<td>41%</td>
</tr>
<tr>
<td>Doing the same finger movements several times a minute</td>
<td>58%</td>
<td>65%</td>
<td>73%</td>
<td>56%</td>
<td>75%</td>
<td>80%</td>
</tr>
<tr>
<td>Doing the same arm movements several times a minute</td>
<td>35%</td>
<td>39%</td>
<td>50%</td>
<td>44%</td>
<td>52%</td>
<td>54%</td>
</tr>
<tr>
<td>Working in a static posture</td>
<td>48%</td>
<td>51%</td>
<td>64%</td>
<td>44%</td>
<td>73%</td>
<td>68%</td>
</tr>
<tr>
<td>Other work-related characteristics</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weekly working hours</td>
<td>35 (8)</td>
<td>35 (8)</td>
<td>36 (8)</td>
<td>35 (9)</td>
<td>36 (5)</td>
<td>34 (9)</td>
</tr>
<tr>
<td>Years working as technician</td>
<td>15 (11)</td>
<td>16 (12)</td>
<td>18 (11)</td>
<td>21 (14)</td>
<td>16 (11)</td>
<td>18 (13)</td>
</tr>
</tbody>
</table>

Data are shown for control and training of all randomised participants, pain cases (forearm pain $\geq 30$, scale 0–100 VAS) and disability cases (Disability of the Arm Shoulder and Hand score $\geq 30$, scale 0–100). VAS, Visual Analogue Scale.
30 or more). Similarly, we calculated OR for becoming a case (score of 30 or more) at follow-up among those without symptoms (score of <10) at baseline. These analyses were exploratory.

We used the SAS statistical software V.9.2 for the analyses (SAS institute) and accepted an α level of 5% as statistically significant. We report baseline results as means (SD) and changes from baseline to follow-up as means (95% CIs) unless otherwise stated.

RESULTS
At baseline, there were no significant differences between the two groups regarding demographics, clinical and work-related characteristics (table 1). For the two groups combined, forearm pain and work disability were moderately related (Spearman’s r = 0.41, p < 0.0001).

During the 20-week period, participants from the training and control groups, respectively, had 12 (SD: 6) and 14 (SD: 5) weekly pain registrations. Based on the training diary registrations, participants from the training group followed the programme on average 36 times during the 20-week period, corresponding to an average adherence of 60%. Training load of the specific forearm exercise—wrist extension—was approximately doubled during the training period and increased from 2.3 (1.1) kg during the initial 4 weeks to 4.3 (1.4) kg during the final 4 weeks. Similar improvements were observed for the shoulder exercises.

Intention-to-treat effect
Figure 2 shows the change in forearm pain from baseline to follow-up among the total study population. A priori hypothesis testing of main effects showed a significant group by time effect for forearm pain intensity (p < 0.001) and work disability (p < 0.01). Intention-to-treat analyses showed that from baseline to follow-up, pain intensity decreased significantly more in the training group than in the control group, with a between-group difference of 5.2 mm VAS (95% CI 2.3 to 8.1 mm VAS). Likewise, work disability decreased significantly more in the training group, with a between-group difference of 4.4 (95% CI 1.4 to 7.5).

Treatment effect among employees with pain and disability
Among employees with pain >30 mm VAS at baseline (n = 54), 60% and 31% from the training and control groups, respectively, recovered completely (VAS score of <10 mm). Pain decreased in the training group (−21.1 mm, 95% CI −32 to −11) but not in the control group (−4.5 mm, 95% CI −16 to 7). The between-group difference in pain from baseline to follow-up was significant (16.7 mm, 95% CI 0.8 to 32.5). The OR for complete recovery at follow-up in the training group compared with the control group was 4.6 (95% CI 1.2 to 17.9).

Among employees with work disability (DASH) >30 at baseline (n = 113), 77% and 36% from the training and control groups, respectively, recovered completely (disability score of <10). Work disability decreased both in the training group (−36, 95% CI −43 to −29) and in the control group (−26, 95% CI −33 to −18). The between-group difference in work disability from baseline to follow-up was significant (11, 95% CI 2.5 to 19), that is, the training group improved significantly more than the control group. The OR for complete recovery at follow-up in the training group compared with the control group was 6.0 (95% CI 1.8 to 18.9).

Preventive effect among symptom-free employees
Among those without pain at baseline (<10 mm VAS, n = 286), there was a small but statistically significant increase in pain in the control group (1.6 mm, 95% CI 0.5 to 2.7) and not in the training group (0.4 mm, 95% CI −0.7 to 1.6). This change was not significantly different between the groups (1.1 mm, 95% CI −0.5 to 2.8).

Among those without work disability at baseline (DASH <10, n = 271), there was a small but statistically significant increase in work disability in the control group (2.2, 95% CI 0.7 to 3.6) and not in the training group (1.8, 95% CI −0.01 to 3.5). This change was not significantly different between the groups (0.4, 95% CI −1.6 to 2.4).

Adverse events
Adverse events were minor and transient. Fifteen participants of the training group contacted the physical therapist of the project because of worsening of pain during training and ascribed these symptoms to previous
injuries being provoked during training. With adjustment of the training programme—for example, reducing intensity or range of motion—these participants were able to complete the 20-week programme. Another four participants in the training group withdrew from the study due to worsening of pain during training. These participants did not consult the physical therapist prior to withdrawal.

**DISCUSSION**

Our study documents positive effects of specific upper limb resistance training on forearm pain and work disability in industrial technicians.

The baseline data of our study show that many industrial technicians work with their hands flexed or twisted and perform repeated finger and arm movements for more than half of the work time (table 1). Both physical and psychosocial factors—including prolonged static muscle activity, insufficient variation in movements and a high degree of precision—are risk factors for development of musculoskeletal disorders. Repetitive movements of the arm or wrist are specifically associated with development of forearm pain. Comparable to the definitions of Kaergaard and coworkers, we defined pain and disability cases as those scoring more than 30 on a scale of 0–100. According to this case definition, the prevalence of forearm pain and work disability among the volunteers of the present study was 10% and 21%, respectively. However, those who agreed to participate in the present study generally had more pain symptoms than those who declined. Thus, the present study population may not be representative of industrial technicians in general. Kryger and coworkers reported—a based on clinical examination—a prevalence of 4% of moderate-to-severe forearm pain among computer workers. Although methodological differences exist between our study and Kryger and coworkers, the prevalence of forearm pain may be higher among laboratory technicians than among computer workers in their study. In this regard, Bjorksten and coworkers showed a higher prevalence of shoulder/hand disorders among laboratory technicians compared with other state employees.

The intention-to-treat analysis, including all participants, showed statistically significant effects of specific resistance training on forearm pain and work disability. As our study included many employees with very low levels of pain and disability, the average changes were small (4–5 on a scale of 0–100). The rationale for including participants both with and without pain was that musculoskeletal pain typically is episodic. Including all employees allowed us to study the effect of the intervention at the company level, as well as the treatment and preventive effect, respectively, among subgroups with and without pain at baseline. Thus, the results may be important from a public health perspective as it reflects the overall impact of such interventions at a company level.

We also performed subgroup analyses among employees with and without symptoms at baseline to determine a possible treatment and preventive effect, respectively. Among those without symptoms at baseline, there was only minor development of pain and disability in the control group during the 20-week period, which diminishes the statistical power for detecting a possible preventive effect. We have previously shown a preventive effect of a 1-year physical exercise intervention on development of shoulder pain among office workers. Thus, follow-up periods longer than 20 weeks may be necessary to determine a possible preventive effect of resistance training for musculoskeletal pain symptoms. Future studies investigating the preventive effect of physical exercise may consider using follow-up periods of several years. On the other hand, adherence to a specific physical exercise programme is likely to diminish over time.

Among the technicians in the training group with pain >30 mm VAS at baseline, pain decreased 20 mm from baseline to follow-up. An essential part of interpreting these results concerns their clinical relevance. A great heterogeneity between studies exists in the definition of clinically meaningful changes in pain. However, Farrar and coworkers found that a change in pain intensity of 1.7 points on a 0–10 scale discriminates well between patients with pain who on the patient global impression of change scale improved much or very much and patients who did not improve or worsened. Similarly, Dworkin and coworkers reviewed that a change in pain intensity of 2 points is moderately clinically meaningful and a change of 1 point is the minimally clinically relevant difference. Thus, among those with pain at baseline, the results of our study can be considered clinically relevant. Similar positive results were obtained for work disability. The odds for complete relief of forearm pain and disability were four to sixfolds higher in the training group compared with the control group. Thus, specific resistance training of the upper extremities effectively reduces forearm pain and work disability among employed technicians with symptoms. Randomised controlled trials have shown mixed results on the effectiveness of specific resistance training for specific arm disorders in patients, for example, lateral epicondylitis. The study by Svernlov and Adolfsson showed that eccentric strengthening exercise performed through a 12-week period was superior to conventional low-intensity stretching exercise. By contrast, Martinez-Silvestrini and coworkers found that a 6-week period of strengthening exercise was not more effective than stretching.

In our study, based on the weekly pain registrations, the time-wise change in forearm pain occurred in a roughly linear fashion and became statistically significant after 10 weeks. Likewise, Coury and coworkers reviewed that physical exercise interventions for musculoskeletal disorders lasting <10 weeks rarely find significant results. Thus, workplace interventions with physical
exercise should last at least 10 weeks to effectively reduce musculoskeletal pain symptoms. Additionally, our results indicate a possible preventative effect for participants without pain at baseline since those in the training group did neither increase pain nor disability while such changes occurred for the control group. However, in the present study group, differences did not attain statistical significance, which may be due to the limited intervention time period.

The pathophysiology of forearm disorders is multifactorial, and some evidence exists for an occupational association. In this regard, prolonged muscle fibre activation and local hypoxia during repetitive arm/hand work tasks may contribute to development of pain symptoms. Using the needle biopsy technique and histochemistry, we have previously found increased proportion of very large slow twitch fibres with poor capillarisation—type I megafibres—in chronically painful neck muscles. Whether such findings hold true for painful forearm muscles remain unknown. Boyer and Hastings reported in a review that surgeons commonly observe degenerative changes of the forearm muscles when treating lateral epicondylitis. Regardless of the underlying pathophysiology of forearm pain, we observed reductions of pain and work disability in response to resistance training. Muscle strength increased as evidenced by a doubling of training loads during the 20-week period. As gains in strength lowers the relative exposure of the muscles during work tasks, strength gains may have contributed to the observed reduction of pain and work disability. Adaptations of the connective tissue in response to the high forces exerted during resistance training may also add to the observed improvements. The present strengthening exercises involved both concentric and eccentric muscle contractions, that is, raising and lowering the weight in a controlled manner. Studies have shown promising pain reducing effects along with increased peritendinous type I collagen synthesis in response to eccentric training. Future studies should investigate the multitude of potential pain reducing mechanisms of different types of resistance training for musculoskeletal disorders.

Our study has both strengths and limitations. The cluster randomised controlled design with high statistical power strengthens our study. As we included both public and private sector companies of which most of the invited employees agreed to participate, the external validity of our findings is high. However, the inclusion and exclusion criteria limit the generalisability to technicians with non-specific forearm pain. Although we did not include a clinical examination for diagnosing specific forearm disorders, our results stress the importance of specific resistance training for reducing the overall level of self-reported pain and work disability at the company level. Another limitation is the possible influence of placebo in behavioural interventions. Because we used subjective rating scales and were not able to blind participants, the results may be biased by placebo effects. Another limitation is the loss to follow-up of more than 20% of the participants in the training group. However, using the GLIMMIX procedure to account for missing values at follow-up strengthens the validity of our estimates.

In conclusion, specific resistance training of the shoulder, neck and arm reduces forearm pain and work disability among industrial technicians.

Contributors All authors contributed to conception/design, acquisition of data or analyses and interpretation of data. LLA drafted the article and all co-authors revised it critically for important intellectual content. All authors approved the final version to be published. Thanks to statistician Ole Olsen for performing the randomisation of clusters.

Funding This work was supported by the Danish Working Environment Research Fund (Grant 8-2007-3). The contribution in terms of manpower allowing employees to train during work time for 1 h/week for 20 weeks was given by the workplaces involved.

Competing interests ICMJE conflicts of interest form for each author of this manuscript.

Ethics approval The ethics approval was approved by Local ethical committee.

Provenance and peer review Exploratory analyses from the study is under preparation by the research group. As such, the data set is not yet ready to be shared.

REFERENCES

Implementation of exercise at the workplace

STROBE checklist for “Gouty arthritis, systolic dysfunction and heart failure: results from a 30-year prospective study”, Krishnan E

<table>
<thead>
<tr>
<th>Section/Topic</th>
<th>Item #</th>
<th>Recommendation</th>
<th>Reported on page #</th>
</tr>
</thead>
<tbody>
<tr>
<td>Title and abstract</td>
<td>1</td>
<td>(a) Indicate the study's design with a commonly used term in the title or the abstract</td>
<td>1-2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(b) Provide in the abstract an informative and balanced summary of what was done and what was found</td>
<td>2</td>
</tr>
<tr>
<td>Introduction</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Background/rationale</td>
<td>2</td>
<td>Explain the scientific background and rationale for the investigation being reported</td>
<td>4</td>
</tr>
<tr>
<td>Objectives</td>
<td>3</td>
<td>State specific objectives, including any prespecified hypotheses</td>
<td>4</td>
</tr>
<tr>
<td>Methods</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study design</td>
<td>4</td>
<td>Present key elements of study design early in the paper</td>
<td>4</td>
</tr>
<tr>
<td>Setting</td>
<td>5</td>
<td>Describe the setting, locations, and relevant dates, including periods of recruitment, exposure, follow-up, and data collection</td>
<td>5</td>
</tr>
<tr>
<td>Participants</td>
<td>6</td>
<td>(a) Give the eligibility criteria, and the sources and methods of selection of participants. Describe methods of follow-up</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(b) For matched studies, give matching criteria and number of exposed and unexposed</td>
<td></td>
</tr>
<tr>
<td>Variables</td>
<td>7</td>
<td>Clearly define all outcomes, exposures, predictors, potential confounders, and effect modifiers. Give diagnostic criteria, if applicable</td>
<td>5</td>
</tr>
<tr>
<td>Data sources/</td>
<td>8*</td>
<td>For each variable of interest, give sources of data and details of methods of assessment (measurement). Describe comparability of assessment methods if there is more than one group</td>
<td>5-6</td>
</tr>
<tr>
<td>measurement</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bias</td>
<td>9</td>
<td>Describe any efforts to address potential sources of bias</td>
<td>5-6</td>
</tr>
<tr>
<td>Study size</td>
<td>10</td>
<td>Explain how the study size was arrived at</td>
<td>5-7</td>
</tr>
<tr>
<td>Quantitative variables</td>
<td>11</td>
<td>Explain how quantitative variables were handled in the analyses. If applicable, describe which groupings were chosen and why</td>
<td>5-7</td>
</tr>
<tr>
<td>Statistical methods</td>
<td>12</td>
<td>(a) Describe all statistical methods, including those used to control for confounding</td>
<td>6-7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(b) Describe any methods used to examine subgroups and interactions</td>
<td>6-7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(c) Explain how missing data were addressed</td>
<td>6-7</td>
</tr>
</tbody>
</table>
(d) If applicable, explain how loss to follow-up was addressed 6-7
(e) Describe any sensitivity analyses

| Results | Participants 13* | (a) Report numbers of individuals at each stage of study—eg numbers potentially eligible, examined for eligibility, confirmed eligible, included in the study, completing follow-up, and analysed 7-9
(b) Give reasons for non-participation at each stage  
(c) Consider use of a flow diagram

| Descriptive data 14* | (a) Give characteristics of study participants (eg demographic, clinical, social) and information on exposures and potential confounders  
(b) Indicate number of participants with missing data for each variable of interest 8
(c) Summarise follow-up time (eg, average and total amount)  

| Outcome data 15* | Report numbers of outcome events or summary measures over time 7-10

| Main results 16 | (a) Give unadjusted estimates and, if applicable, confounder-adjusted estimates and their precision (eg, 95% confidence interval). Make clear which confounders were adjusted for and why they were included 8-9
(b) Report category boundaries when continuous variables were categorized 7-9
(c) If relevant, consider translating estimates of relative risk into absolute risk for a meaningful time period 7-9

| Other analyses | Report other analyses done—eg analyses of subgroups and interactions, and sensitivity analyses 9

| Discussion | Key results 18 | Summarise key results with reference to study objectives 3,10

| Limitations | Interpretation 20 | Give a cautious overall interpretation of results considering objectives, limitations, multiplicity of analyses, results from similar studies, and other relevant evidence 10-12

| Generalisability 21 | Discuss the generalisability (external validity) of the study results 10

| Other information | Funding 22 | Give the source of funding and the role of the funders for the present study and, if applicable, for the original study on which the present article is based 3

*Give information separately for cases and controls in case-control studies and, if applicable, for exposed and unexposed groups in cohort and cross-sectional studies.
Dear Dr. Richard Sands,

We have now revised the manuscript (bmjopen-2011-000412) entitled "Effect of specific resistance training on forearm pain and work disability in industrial technicians: Randomized controlled trial" according to the review comments.

Comment from the managing editor:
Further to the reviewers' comments below, we would also like the reporting of the trial to be more transparent and thorough.

In particular, how does it overlap with ref 15, which is in press at a BMC journal? Is that just reporting the baseline results? There should now be a reference for this if it has been accepted (at least a DOI?).

OUR REPLY: Ref 15 provides information on the baseline population and results from the intervention on neck/shoulder pain. Thus besides some of the baseline results there are no overlap between ref 15 and the present results. Ref 15 has now been accepted and published, and the ref has been updated in the reference list of the present manuscript.

Comment from the managing editor:
Although BMC journals are open access, it's unhelpful to refer readers there for all the details of randomisation etc. Please revise the paper in line with the CONSORT statement for cluster randomised trials (eg on method of random allocation, allocation concealment, and blinding). It's at http://www.consort-statement.org/extensions/designs/cluster-trials/

OUR REPLY: We have now provided this information in a new section, "The cluster randomization procedure" in the Methods.

Comment from the managing editor:
The completed CONSORT checklist is the generic one, and doesn't cover cluster RCTs. It should also only state what is explicitly reported in the paper - for example it states that various aspects are reported on p5, whereas this is actually the BMC reference.

OUR REPLY: We have now checked this more carefully and used the checklist for cluster RCT’s. The checklist is uploaded as a supplementary file.

Comment from the managing editor:
Finally, please structure the abstract in line with CONSORT.

OUR REPLY: We have now structured the abstract in line with CONSORT.

Comment from the managing editor:
The contributorship statement says that all authors drafted the protocol. However, authorship should meet all three criteria as laid out by the ICMJE: http://www.icmje.org/ethical_1author.html

OUR REPLY: We have now checked this more carefully and changed accordingly.

*******************************************************************************

Reviewer: Anna Sjörs
Senior Developer
Institute of Stress Medicine
Göteborg, Sweden
The study design and execution seem sound. There are, however, some important details missing in the methods section. Please provide details on the randomization, the procedure for exercise intervention, and the adherence to the protocol (see below).

Comment from reviewer Dr. Anna Sjörs:
An inherent problem with this study is the focus on reducing pain and disability in a population with already low pain and disability scores. If the exercise program is, eventually, intended for treatment of pain in patients with a diagnosed pain condition, the current study population is not representative of actual patients that would receive the treatment. One cannot generalize these findings to a population with more severe pain. The authors argue that the exercise intervention may still be important from a public health perspective. I would like to see a more thorough discussion to justify this statement. If, on the other hand, the main focus is to prevent development of pain in healthy workers, I believe that 20 weeks is a rather short intervention time period, which the authors have addressed briefly in the discussion.

OUR REPLY: It is true that this working population has a relative low mean pain rating. This is why we have conducted a sub-analysis among those with a pain rating greater than 30 mm VAS at baseline. This group approaches pain patients and shows to have particular benefit from the training by having an odds ratio (OR) for complete recovery at follow-up in the training group compared to the control group of 4.6 (95% CI 1.2 to 17.9). This finding is even more convincing regarding the analysis on a sub-group regarding disability. These analyses underline the relevance of this study also for patient groups.

Article summary

Comment from reviewer Dr. Sjörs:
1. Please change arm/hand to forearm pain since hand pain was not the focus of this study.
   OUR REPLY: This has now been changed

Introduction

Comment from reviewer Dr. Sjörs:
2. The numbers given on incidence and prevalence of forearm pain are from 2003 and may not apply to the current study population. Please provide more recent data or soften the statement to indicate that the percentages may have changed.
   OUR REPLY: We have now softened this statement.

Methods

Comment from reviewer Dr. Sjörs:
3. Information on the number and percentages of subjects with pain and disability > 3 could be given in table 1.
   OUR REPLY: We have now added columns to Table 1 showing baseline characteristics of pain-cases and disability-cases for the control and training groups, respectively.

Comment from reviewer Dr. Sjörs:
4. Please provide details on the randomization procedure.
   OUR REPLY: We have now provided this information in a new section, “The cluster randomization procedure” in the Methods.

Comment from reviewer Dr. Sjörs:
5. Please provide more details on how the training was performed. Individually or in groups? Were the participants able to choose location and time of day?
OUR REPLY: We have now provided detailed information of this in the section, “intervention” in the Methods.

Comment from reviewer Dr. Sjörs:
6. Please also give information on adherence to the protocol. How was this monitored?
OUR REPLY: This was monitored by weekly logbook registrations. This information has now been added to the manuscript.

Comment from reviewer Dr. Sjörs:
7. The choice of cut-off for pain cases seems arbitrary and needs to be justified.
OUR REPLY: Kaergaard and coworkers found that for four complaint questions on a 10-point scale a sum of 12 (i.e. average 3 for each of the four questions) was significantly associated with clinical findings. Because we used scales from 0-100 we then used a cut-off at 30. We have now explained this in “Forearm pain and work disability” in the Methods section.

Comment from reviewer Dr. Sjörs:
8. Likewise, justify the cut-off for disability cases.
OUR REPLY: We used the same definition and rationale for disability cases as for pain cases.

Comment from reviewer Dr. Sjörs:
9. How were the individual regressions of pain over time used in the analyses? Were they entered in the GLIMMIX analyses? I cannot find the results of these regressions.
OUR REPLY: we used linear regression for each participant to estimate the trend over time, and thereby determined the change from baseline to follow-up by multiplying the slope with the number of weeks. The individual changes from baseline to follow-up were then entered in the GLIMMIX procedure. This has now been specified in the Statistics section.

Results

Comment from reviewer Dr. Sjörs:
10. Please specify the number of pain cases in each group and not just the total number of 54 pain cases (could be presented in table 1, as suggested above).
OUR REPLY: This information has now been added to Table 1.

Comment from reviewer Dr. Sjörs:
11. Specify the number of disability cases in each group accordingly.
OUR REPLY: This information has now been added to Table 1.

Comment from reviewer Dr. Sjörs:
12. Please report the mean pain and disability scores among cases at the start of the study.
OUR REPLY: This information has now been added to Table 1.

Comment from reviewer Dr. Sjörs:
13. As more of a comment, I would like to point out that Figure 2 is somewhat misleading as the VAS on the Y-axis actually ranges from 0 to 100. Changes in pain scores appear larger than they actually are.
OUR REPLY: True. We have now mentioned in the legend of Figure 2 that the scale ranged from 0 to 100.

Discussion

Comment from reviewer Dr. Sjörs:
14. Please clarify if the prevalence of pain and disability (10 % and 21 %) concerns the total population or one of the subgroups.
OUR REPLY: This concerns the control and training group together. This has now been specified and discussed in relation to the generalisability.

Comment from reviewer Dr. Sjörs:
15. The choice of and discussion of the limit of clinical relevance is somewhat confusing. The sample size calculation uses the limit 15 % change in pain (no reference) as a relevant change. Does this refer to a 15 % difference between groups in pain improvement or to a 15 % change within group? This should be clarified as the comparison between groups in pain improvement is essential, given the study design. In the discussion, however, a reduction of 20 mm (within the training group) is considered clinically relevant (without relating it to the concurrent change in the control group). Since the mean pain intensities among cases before and after the intervention and percentage of change in pain are not reported, it is not possible to relate the results to the given limit of 15 %.
OUR REPLY: The sample size calculation was performed prior to the study and was based on a 15% change as the least relevant change. In hindsight, this calculation should have been based on actual changes between groups rather than percentage change. However, to allow sufficient power even in a cluster randomization design with cluster size below 10 we doubled the number of participants in both groups resulting in approx. 30 clusters and in each group with a total sample size of more than 500. We have now added the calculations for the sample size regarding cluster considerations in the manuscript, and also in more detail the aspect of clinical relevance in the Discussion.

References

Comment from reviewer Dr. Sjörs:
For the discussion of clinical relevance I recommend the following article:
OUR REPLY: We have now included this relevant reference in the manuscript.

**********************************************************************
Reviewer: A.J. van der Beek, PhD
Professor of Occupational Epidemiology
Department of Public and Occupational Health
EMGO Institute, VU University Medical Centre
Van der Boechorststraat 7
NL-1081 BT Amsterdam
The Netherlands

Comment from reviewer Professor van der Beek:
It is recommended to add four columns to table 1, i.e. for participants with and without pain greater than 30 mm VAS at baseline in the training group and for participants with and without pain greater than 30 mm VAS at baseline in the control group. The reader should, in my opinion, have more insight in the
characteristics of these important subgroups, in particular since the study presents relevant results for those with pain at baseline.

OUR REPLY: We have now added columns to Table 1 showing baseline characteristics of pain-cases and disability-cases for the control and training groups, respectively.

Comment from reviewer Professor van der Beek:
I would appreciate a little more information about the actual work being performed by these, most often female, technicians. The title mentions industrial technicians, whereas the Discussion speaks about laboratory technicians. I suggest to add a few sentences to the first paragraph of the Methods.

OUR REPLY: This information has now been added to the first paragraph of the Methods.

Comment from reviewer Professor van der Beek:
Furthermore, more information should be given about the number and size of the clusters. How many participants were in the largest cluster and how many in the smallest?

OUR REPLY: We have now provided this information in a new section, "The cluster randomization procedure" in the Methods.

Comment from reviewer Professor van der Beek:
Finally, it is not fully clear how representative the participants were for the actual working population in the companies at stake. Out of the 854 workers, 211 + 237 replied to the follow-up questionnaire. Although this not bad at all for studies in the occupational setting, it is recommended to pay more attention to differences between subjects who were included in the final analysis and those who did not reply/declined participation/withdrew.

OUR REPLY: Results and discussion on differences and similarities between those who declined and agreed to participate in the study, respectively, have been published in the BMC article. In brief, those who agreed to participate had more pain symptoms. We have now mentioned this in discussion of the present manuscript.

Comment from reviewer Professor van der Beek:
Also, I expect that several participants have one or more missing measurements in the weekly pain ratings. In the first paragraph of the Results it should be mentioned how many measurements were missing for how many participants. And, most importantly, to what extent did this and the loss to follow-up influence the results?

OUR REPLY: The average number of weekly pain ratings is now provided in the Results section. The loss to follow-up is a limitation, which we have emphasized in the limitations section prior to the conclusion.

Comment from reviewer Professor van der Beek:
- It should be mentioned in the legend of Figure 2 that the scale ranged from 0 to 100!
- page 2, line 0: 'Arm/hand is' should be 'Arm/hand pain is'.
- page 4, line 23: 'involves' should be 'involve'.
- page 8, line 50: 'performs' should be 'perform'.
- page 9, line 31 and 35: 'preventative' should (in my opinion) be 'preventive'.
- several times: 'compared with' should be 'compared to' or 'in comparison with'.

OUR REPLY: This has now been changed

********************************************************************************
Reviewer: Anneli Ojajärvi
Senior Specialist
Comment from reviewer Dr. Ojajärvi:
In methods, the authors could explain briefly how they performed the cluster randomized controlled trial.
OUR REPLY: This has now been explained in more detail in the manuscript.

Comment from reviewer Dr. Ojajärvi:
In statistics, the authors have used a generalized linear mixed model controlled only for gender, not age, to determined differences in the main outcomes between the two groups from baseline to follow-up.
OUR REPLY: True. As shown in Table 1 there were more men in the control group than the training group. Age was similar. Therefore we only controlled for gender.