Effectiveness of workplace active rest programme on low back pain in office workers: a stepped-wedge cluster randomised controlled trial

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ABSTRACT

Objective This study aimed to investigate the effectiveness of workplace active rest programme (WARP) on chronic low back pain (LBP) among office workers.

Design A closed cohort, stepped-wedge cluster randomised trial was conducted. The total duration of the study was 16 weeks (4 weeks for each step). Sequence allocation was randomised, but no one was blinded.

Setting This study was conducted in three offices in a Japanese electronics company. One office was for the administrative department, the others are for the engineering department.

Participants We recruited 29 office workers with LBP greater than 3 months. LBP due to specific injury or disease was excluded. The median age was 38 years, and 26 (90%) were male. All participants completed the study.

Interventions In the intervention phase, participants performed WARP comprising frequent stand-up and individualised brief exercise/physical activity during work. Physical therapists held an LBP workshop and developed tailor-made programmes before introducing WARP. We instructed participants to perform WARP at five timings during work. Control phase was set before the intervention and participants stayed as usual.

Primary and secondary outcome measures The primary outcome was pain intensity of LBP assessed using the Brief Pain Inventory. The secondary outcomes were work productivity loss measured using the Work Limitations Questionnaire, LBP disability assessed using the Roland-Morris Disability Questionnaire, psychosocial subscale assessed using the STarT Back Screening Tool and physical activity measured using triaxial accelerometers. These outcomes were collected at baseline and at 4-month follow-up evaluation.

Results In the intention-to-treat analysis, WARP did not show any significant effects on pain intensity ($p=0.01$; 95% CI $-0.50$ to 0.52) and on the secondary outcomes. The median adherence to WARP was 28.6% (IQR, 16.8–41.1), which was equal to 1.43 times per day. No adverse effect was observed.

Conclusions The present study was unable to confirm the effectiveness of active rest in improving LBP. Hence, further study needs to investigate its effectiveness.

Strengths and limitations of this study

- This study is the first pragmatic trial conducted in a real-world setting that investigates the feasibility and effectiveness of active rest.
- All participants completed the workplace active rest programme (WARP).
- Adherence to WARP was lower than expected.
- Because recruited office workers had relatively mild low back pain, we were unable to confirm whether WARP is effective among office workers with severe low back pain.

INTRODUCTION

Low back pain (LBP) is a prevalent health problem among office workers1 2 and is the leading cause of decreasing healthy life expectancy worldwide.3 Moreover, LBP results in a large socioeconomic burden due to work productivity loss and medical expenses.4 5 In terms of both individual and social impact, LBP among office workers is a crucial problem which should be tackled.

Office workers are workers who stay in prolonged sitting position during most of their working time.6 7 Prolonged sitting is one of the causes of LBP, which is also due to several factors such as increased disc pressure,8 decreased trunk mobility9 and less posture variation.10 Although previous studies have investigated the effectiveness of ergonomic intervention and back support, these are considered ineffective in improving LBP.11 12 Recently, the use of standing desk has been shown to be effective in improving LBP,13 but it has the following limitations: it requires a lot of space and is costly. Therefore, easy-to-use solutions are required in the workplace.

Active rest (taking a break with exercise/physical activity in the workplace) could possibly improve LBP because it has the
following characteristics: (1) sedentary break by standing up, which can prevent prolonged sitting; and (2) exercise/physical activity, which is recommended in the LBP guidelines. A previous study showed that office-based stretching (10–15 min/session, 3 times/week) was effective in reducing the occurrence of musculoskeletal discomfort when compared with no intervention. However, in our study, we developed a shorter exercise programme involving frequent sessions (a few minutes per session, five times per day, except on weekends) because we aimed to promote frequent standing to break the habit of prolonged sitting. Although a positive effect of active rest on LBP has been shown in a laboratory study, its effectiveness in the real-world setting is still unknown. We hypothesised that there is a difference in effectiveness between laboratory and real-world setting. Thus, the present study aimed to investigate the effectiveness of the workplace active rest programme (WARP) on chronic LBP and work productivity loss among office workers in a real-world setting.

METHODS

Study design

The present study was conducted according to the extension of the Consolidated Standards of Reporting Trials 2010 statement for stepped-wedge cluster randomised controlled trial (SW-CRT). We used a closed cohort SW-CRT involving randomisation of clusters to different sequences. SW-CRT is a crossover design with repeated measurement, in which clusters switch from control to intervention condition. SW-CRT is a suitable study design if we assume that the intervention will do more good than harm, hence making it unethical to withhold the intervention from a control group. Thus, because it is morally acceptable and beneficial for participant recruitment, we introduced the SW-CRT design. Moreover, this is the pragmatic design, which increases statistical power and decreases needed clusters compared with those in parallel CRT. The present clinical trial was registered with University Hospital Medical Information Network (UMIN) Clinical Trials Registry.

As figure 1 shows, we conducted the present study in three offices (clusters) in a Japanese electronics company. We set three sequences, where an office switched from the control condition to the intervention condition one by one. The total duration of the study was 16 weeks (4 weeks for each step). Evaluation was conducted at baseline and at four points during the last week of each step. Due to the study’s closed cohort design, participants assessed in different periods were the same participants.

Patient and public involvement

Office workers with LBP were not involved in developing the research question, but we consulted them about the design of the study (especially the intervention programme) in terms of feasibility and applicability by joining the employees’ health committee. During the trial, they helped us to hold an LBP workshop by arranging a room and equipment. We asked them to assess the burden of the intervention before they joined the study. We already disseminated the results of our study to the participants and reported them at the employees’ health committee.

Participants’ recruitment

We recruited 29 participants from three offices of a Japanese electronics company in July 2018. Three offices were separated from one another. First, participants were approached by the public health nurse working in this company. When they expressed interest in the study, the public health nurse introduced them to us. Subsequently, the researchers explained the study to the participants, and the participants provided informed consent for inclusion in the study.

Office workers were eligible for the present study if they have the following characteristics: (1) full-time workers (all workers worked in the same day shifts), (2) engaged in desk work greater than 4 hours/daily working time (self-reported) and (3) had LBP for greater than 3 months.
The location of LBP was defined as pain between the 12th rib and the inferior gluteal folds. Exclusion criteria were as follows: (1) LBP caused by fracture and trauma injuries, infectious diseases and internal organ disorders and (2) difficulty participating in the study due to medical or surgical disease. Cluster-level eligibility criteria were as follows: (1) an office where most workers were engaged in desk work and (2) supervisors granting permission in the performance of the study. Whereas office A was for an administrative department, office B and office C were for engineering department.

All participants provided written informed consent for inclusion in the study.

Randomisation and blinding
Offices were randomly assigned to one of the three successive sequences (one office per sequence) after all clusters and participants were recruited. A researcher who was not involved in the recruitment performed random allocation using computer-generated random numbers and coded information about offices. To prevent contamination, both clusters and participants were not informed of the time the intervention started and the detailed programme of the intervention until 2 weeks before the intervention started. We also asked the participants exposed to the intervention not to disclose the programme content to other workers. Due to the nature of the present study, participants, intervenient and outcome assessors (self-reported) could not be blinded. The data analyst was also not blinded to group allocation.

Intervention
In the intervention phase, we offered WARP in two parts. First, we held the LBP workshop (group), followed by the introduction of active rest in the workplace. The LBP workshop was held when the group moved from the control phase to the intervention phase.

The purposes of the LBP workshop were as follows: to allow the participants to understand LBP and sedentary behaviour, develop customised exercise programme and explain how WARP is performed after the workshop. The LBP workshop was held at the company’s gymnastics room after work for 90 min by two or three physical therapists (PTs) (PTs with expertise in LBP, at least 3 or more experience years) including the primary researcher (YT). To avoid inconsistency on workshop contents in PTs, we discussed and agreed with its contents before workshop. To perform any intervention to the participants (usual work).

Primary outcome
The primary outcome was LBP intensity. We used the pain intensity subscale of the Brief Pain Inventory (BPI), which is well validated and reliable among patients with non-cancer pain including LBP. BPI consists of four questions rating pain intensity separately at ‘worst’, ‘least’, ‘average’ and ‘now’ during the last 24 hours using 11-point Numerical Rating Scale (NRS), ranging from 0 (no pain) to 10 (worstimaginable pain). Finally, the mean of these four items was used as the BPI score: BPI score=(worst+least+average+now)/4. A Japanese version of BPI has good validity and reliability.

At the time of trial registration, although we had planned to evaluate weekly LBP intensity, we changed to once in 4 weeks evaluation. This is because weekly evaluation was not feasible at this company in terms of responders’ burden in answering questionnaires.
Secondary outcome

The Roland-Morris Disability Questionnaire (RDQ) is a validated 24-item questionnaire that assesses disability due to LBP, such as ‘I change position frequently to try and get my back comfortable’. Each item is scored either 0 or 1, with all scores summed to a total between 0 and 24 (a higher score indicates higher disability level).

The STarT Back Screening Tool is a validated screening tool that predicts future disability level. We used the five-item psychosocial subscale of the STarT Back Screening Tool, including fear of movement, depressive symptom, catastrophic attitude, anxiety and pain distress. Scores ranged from 0 to 5 (a higher score indicates higher possibility of future disability level).

The Work Limitations Questionnaire (WLQ) is a validated 25-item questionnaire that evaluates work productivity loss due to physical/psychological issues. The WLQ is composed of the following four subscales: (1) time management (difficulty in performing job tasks in a timely manner and in scheduling tasks); (2) mental-interpersonal demands (difficulty in performing cognitive job tasks and in interacting with colleagues); (3) physical demands (ability to perform job tasks involving body strength, movement, endurance, coordination and flexibility); and (4) output demands (work quantity and quality reduction and timeliness of completed work). Additionally, ‘not applicable’ was also provided as a response option and treated as a missing value. All subscales scores were converted to percentage, from 0% (least limited) to 100% (most limited). Work productivity loss (%) was calculated from the weighed sum of the four subscale scores using a validated algorithm ranging from 0% to 24.9%. A higher score indicates higher level of work productivity loss.

To measure physical activity and sedentary behaviour, we distributed triaxial accelerometers (Active style Pro HJA-750C, Omron Healthcare) to the participants during each step. Details of the accelerometer measurement procedure were described elsewhere. Participants were instructed to wear triaxial accelerometers on their waist during only working time for 5 days. Data were recorded in 60 s epoch. In addition to the number of steps, time spent in moderate-to-vigorous physical activity (MVPA, ≤3.0 metabolic equivalent; METs), light physical activity (1.5 < MET < 3.0) and sedentary behaviour (MET ≤1.5) was calculated using R V.3.5.2. Days with at least zero count lasting over 60 min.

Adherence

To evaluate adherence to WARP, we asked participants to keep diaries on whether they performed WARP or not in each five timing. Adherence is calculated 100% if they performed WARP at all five timings during the whole intervention phase. Because WARP is a programme at the workplace, we did not include holidays when assessing adherence.

Sample size

We calculated the sample size using formula specific for stepped-wedge design. The primary outcome difference and SD were set as 2.0 and 2.5, respectively. The following assumed parameters were used: cluster size=10, intra-cluster correlation coefficient=0.05, number of step=3, number of baseline measurement=1, measurement after each step=1, two-sided α level=0.05 and β=80%. To detect a 2-point difference in primary outcome, a total of 22 participants were needed. Considering dropout, we estimated 30 participants as the required sample size, and 29 participants actually joined the present study. Although we set cluster size as 10 before recruitment, the actual size of the two clusters was 8. We conservatively performed sample size calculation by changing some parameters. However, the required sample size was not changed (22 participants) even if it is 8 participants. Therefore, this difference would not affect the results of our study.

Statistical analysis

For the characteristics of participants, categorical variables were presented as frequency and percentage and continuous variables as mean±SD. If the distributions of the continuous variables were skewed, data were presented as median (range or IQR).

We performed both intention-to-treat (ITT) and per-protocol analyses to investigate both the effectiveness and the efficacy of WARP. The primary analysis was ITT analysis because this study aimed to investigate the pragmatic effectiveness of WARP in a real-world setting. Regarding ITT analysis, we performed linear mixed effect model for all outcomes, setting the intervention as the fixed effect, individual and office as the random effect, and calendar time as the confounding factor. Regarding per-protocol analysis, we also performed the linear mixed effect model...
for all outcomes after excluding participants whose adherence to WARP was 28.6% (median) or less. Unstandardised coefficients and 95% CIs were calculated.

All statistical analyses were performed using Stata/IC V.15.1 software. P<0.05 was considered statistically significant.

RESULTS
We recruited 29 office workers from three offices in July (figure 2). As planned, office A performed the intervention in the first period (August), office B in the second period (September) and office C in the third period (October). All participants continued WARP until the end (no dropout) of the study. Twenty-eight participants completed the baseline and each follow-up evaluation (T1–T4). Only one participant did not answer the T3 evaluation, but answered other evaluations.

The median age was 38 years, and 26 (90%) were male (table 1). The median pain intensity assessed using BPI was 2.0 (IQR, 0.8–2.2), and the median score on RDQ was 1.0 (0.0–2.0). Only two participants had clinic or alternative care, and only one participant often received analgesic medication. The median proportion of sedentary time was 79.6% (68.1–84.1). The median productivity loss estimated by WLQ was 2.2% (0.8–5.9). Regarding the difference in characteristics of the three offices, participants were younger in office C than in other offices. Pain intensity was lighter in office B than in other offices.

The median adherence to WARP was 28.6% (16.8–41.1), which is equal to 1.43 times per day (figure 3). Participants with higher adherence had relatively higher pain intensity, disability due to LBP and higher work productivity loss (online supplemental table 1) compared with those with lower adherence. Furthermore, low adherence was related to longer duration of WARP (adherence, office A=B=C).

For ITT analysis with adjustment for time effects, pain intensity did not improve better in the intervention phase compared with the control phase (β, 0.01; 95% CI –0.50 to 0.52) (table 2). Regarding secondary outcomes, no significant improvement was observed. For per-protocol analysis with adjustment for time effects (n=14), time management demands and mental-interpersonal demands (WLQ subscale), MVP A improved better in the intervention phase compared with the control phase. RDQ, productivity loss and step significantly improved better in the intervention phase compared with the control phase. Calendar time had significant or marginal significant positive effects on the primary and secondary outcomes. Any adverse effects were not reported in the present study.

Regarding participants’ satisfaction with WARP, 4 (14%) were very satisfied, 10 (34%) were satisfied and 15 (52%) were normal. No one was unsatisfied with WARP. As regards positive comments, some said the following: “I understood my back pain could be improved, and exercise was easy to perform,” “It was nice to know effective stretch,” “I feel my back pain is gradually improved,” “I could be careful for prolonged sitting,” “I want to make use of personalized exercise,” “Back pain was gradually improved,” “I could consider problems and methods for solving back pain,” and “It was nice to undertake an...
exercise instruction from professionals.” As regards negative comments, some said the following: “Not enough follow-up other than questionnaire,” “Regular feedback based on follow-up data can motivate us to perform this program, but actually no feedback in this program,” “There were few people doing exercise around me, so it was hard to do exercise,” and “I wanted to know exercise during sitting.”

Table 1  Characteristics of the participants

|                      | All | Office A | Office B | Office C |
|----------------------|-----|----------|----------|----------|
| n                    | 29  | 8        | 8        | 13       |
| Age, median (IQR)    | 38.0 (28.0–45.0) | 43.5 (37.0–46.5) | 41.5 (29.5–46.0) | 32.0 (27.0–38.0) |
| Sex, n (%)           |     |          |          |          |
| Male                 | 26 (90) | 6 (75)   | 7 (88)   | 13 (100) |
| Female               | 3 (10) | 2 (25)   | 1 (12)   | 0 (0)    |
| BMI, median (IQR)    | 21.9 (20.2–24.6) | 20.9 (19.9–23.8) | 21.5 (20.3–24.3) | 22.6 (21.5–24.6) |
| Lumbar disc herniation, n (%) | 2 (7) | 0 (0) | 1 (12) | 1 (8) |
| Lumbar canal stenosis, n (%) | 2 (7) | 0 (0) | 1 (12) | 1 (8) |
| Pain intensity, median (IQR) | 2.0 (0.8–2.2) | 1.9 (1.1–3.0) | 0.6 (0.0–2.1) | 2.0 (1.2–2.5) |
| RDQ, median (IQR)    | 1.0 (0.0–2.0) | 1.0 (0.0–1.5) | 0.0 (0.0–1.0) | 1.0 (0.0–2.0) |
| STaR T Back, median (IQR) | 0.0 (0.0–1.0) | 0.0 (0.0–0.5) | 0.0 (0.0–0.5) | 0.0 (0.0–1.0) |
| Medicine, n (%)      |     |          |          |          |
| None                 | 23 (79) | 5 (62)   | 7 (88)   | 11 (85)  |
| Rarely               | 3 (10) | 2 (25)   | 0 (0)    | 1 (8)    |
| Sometimes            | 2 (7) | 1 (12)   | 1 (12)   | 0 (0)    |
| Often                | 1 (3) | 0 (0)    | 0 (0)    | 1 (8)    |
| Always               | 0 (0) | 0 (0)    | 0 (0)    | 0 (0)    |
| Seek for clinic care, n (%) | 2 (7) | 0 (0) | 1 (12) | 1 (8) |
| Seek for alternative care, n (%) | 2 (7) | 2 (25) | 0 (0) | 0 (0) |
| Physical activity, median (IQR) | | | | |
| Time spent for sedentary (%) | 79.6 (68.1–84.1) | 74.1 (58.5–80.0) | 78.9 (63.6–84.9) | 81.6 (73.5–85.2) |
| Time spent for LPA (%) | 16.3 (12.6–24.4) | 19.4 (15.5–32.9) | 17.2 (12.4–27.9) | 13.4 (11.0–19.2) |
| Time spent for MVPA (%) | 4.5 (2.9–7.1) | 5.6 (3.5–10.1) | 3.9 (2.7–5.9) | 4.1 (3.0–6.3) |
| Step                 | 4763.4 (3553.1–6228.4) | 4763.4 (3962.9–8457.4) | 4569.5 (3490.1–6228.4) | 4593.9 (3624.5–5636.5) |
| Wearing time (min)   | 708.4 (663.3–757.1) | 682.7 (635.4–744.4) | 757.0 (667.4–847.3) | 707.1 (692.2–743.5) |
| Other musculoskeletal pain, n (%) | | | | |
| Neck                 | 17 (59) | 4 (50)   | 4 (50)   | 9 (69)   |
| Shoulder             | 18 (62) | 4 (50)   | 5 (62)   | 9 (69)   |
| Elbow                | 3 (10) | 0 (0)    | 2 (25)   | 1 (8)    |
| Hand                 | 4 (14) | 1 (12)   | 2 (25)   | 1 (8)    |
| Hip                  | 4 (14) | 1 (12)   | 1 (12)   | 2 (15)   |
| Knee                 | 7 (24) | 2 (25)   | 4 (50)   | 1 (8)    |
| Foot                 | 7 (24) | 3 (38)   | 2 (25)   | 2 (15)   |
| Sleep quality, n (%) |     |          |          |          |
| Good                 | 15 (52) | 5 (62)   | 4 (50)   | 6 (46)   |
| Bad                  | 14 (48) | 3 (38)   | 4 (50)   | 7 (54)   |
| Productivity loss, mean (IQR) | 2.2 (0.8–5.9) | 1.8 (1.2–2.5) | 2.8 (0.4–5.1) | 2.2 (1.3–6.9) |
| Time management, median (IQR) | 0.0 (0.0–15.0) | 0.0 (0.0–5.0) | 0.0 (0.0–10.0) | 0.0 (0.0–15.0) |
| Physical demand, median (IQR) | 0.0 (0.0–10.0) | 2.5 (0.0–25.0) | 0.0 (0.0–0.0) | 0.0 (0.0–10.0) |
| Mental-interpersonal demand, median (IQR) | 8.3 (0.0–16.7) | 5.6 (1.4–9.7) | 11.1 (2.8–18.1) | 11.1 (0.0–22.2) |
| Output demand, median (IQR) | 10.0 (0.0–25.0) | 7.5 (0.0–17.5) | 13.1 (0.0–30.0) | 10.0 (0.0–30.0) |

STaR T Back, STaR T Back Screening Tool; BMI, body mass index; LPA, Low physical activity; MVPA, Moderate-vigorous physical activity; RDQ, Roland-Morris Disability Questionnaire.
In summary, ITT analysis showed that WARP did not have significant positive effects on LBP intensity and other secondary outcomes such as LBP disability or work productivity. The median adherence to WARP was 28.6% (1.43 times per day), which was significantly lower than we expected. Per-protocol analysis revealed that WARP was not associated with LBP outcomes, but WARP had significant positive effects on some subscales of work productivity (time management demands, mental-interpersonal demands) and MVP.

Although a recent systematic review investigated the current evidence on active rest, they concluded that there was low-quality evidence for the conflicting effectiveness in LBP. Studies included in the systematic review were conducted in the laboratory setting or healthy subjects without LBP. Therefore, this is the first randomised controlled trial that investigates the effectiveness of active rest in LBP and work productivity in a real-world setting. However, we were not able to demonstrate the significant positive effect of WARP on LBP. While the present study evaluated the effect of short and frequent office-based exercises (a few minutes per session, five times per day, except weekends) on LBP symptom reduction, a previous study showed the effect of long and less frequent office-based exercises (10–15 min per session, 3 times per week) on LBP symptom reduction. These differences between the two study designs should be considered when interpreting the results of our study.

We have two potential explanations about the negative results of our study. First, it might be due to low adherence to WARP, which could diminish its efficacy. Although we considered some strategies to keep adherence (eg, introducing WARP to all workers other than...
the participants of this study in the same office, ringing the chime to inform them of WARP timing, and tailor-made exercise programme), these might be insufficient to improve adherence. Previous studies suggested supervised exercise and group-based exercise. However, there were no strict supervision or group-based exercises in our study because we tried to investigate the effectiveness of a pragmatic easy-to-use solution. Moreover, lower adherence to workplace exercise was influenced by poorer psychosocial work environment (eg, influence at work, work pace, quantitative demands, interpersonal relations) and lower exercise self-efficacy. A further study should be conducted to perform such strategies to improve adherence, but simplicity and acceptance from the employee and the employer should be considered in terms of practical use. The second potential explanation for the negative results is that the participants in our study had lower level of LBP intensity at baseline, which leads to low motivation for WARP and floor effect. Actually, participants with lower LBP intensity had lower adherence than those with high LBP intensity. We considered the floor effect owing to the mild pain by specifically recruiting workers with back pain (NRS was 3 or higher). However, a time lag between recruitment and baseline assessments due to coordinating the schedule of the LBP workshop might have led to a decrease in pain levels at the time the study was actually conducted. Future studies should focus on the fluctuations of outcome variables between recruitment and baseline assessments.

Regarding per-protocol analysis, unstandardised coefficients of most outcome parameters were significantly positive compared with those of the ITT analysis. A previous study reported that active rest (10 min fitness programme at lunch break) has positive effects on vigour, interpersonal stress and physical activity. Although the results of the per-protocol analysis should be carefully interpreted owing to selection bias and an underpowered analysis, these results indicate that WARP could have positive effects if its adherence was ideally kept.

Several limitations should be considered when interpreting the results of our study. First, adherence to the programme was very low, which might lead to underestimation of the potential efficacy of WARP. Second, severity level of LBP was relatively mild in this population, which might cause floor effect especially for BPI and RDQ. We should have set the inclusion criteria about the severity level of LBP to eliminate the floor effect. Otherwise, if mild LBP is common in the working population compared with primary care, we should focus on the incidence or recurrent incidence of LBP in terms of primary prevention. Finally, owing to the limited number of workplace settings and types included within one company, the results of the study should not be considered generalisable to other workplace settings.

We were unable to conclude that active rest is effective in LBP and productivity loss from the results of the present study. However, the present study provided valuable information for conducting similar research, although the strategies implemented in this study might be insufficient to maintain adherence. In the future, we need to study its effectiveness with high adherence or among workers with higher level of LBP intensity.

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Data, STATA code for statistical analyses and R code for data processing of accelerometers are available upon reasonable request.

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