Effects of Inpatient Multicomponent Occupational Rehabilitation versus Less Comprehensive Outpatient Rehabilitation on Somatic and Mental Health: Secondary Outcomes of a Randomized Clinical Trial

Lene Aasdahl1 • Kristine Pape1 • Ottar Vasseljen1 • Roar Johnsen1 • Sigmund Gismervik1,2 • Chris Jensen1,3 • Marius Steiro Finland1,4

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Abstract Purpose To evaluate effects on somatic and mental health of a multicomponent inpatient occupational rehabilitation program compared to a less comprehensive outpatient program in individuals on sick leave for musculoskeletal complaints or mental health disorders. Methods A randomized clinical trial with parallel groups. Participants were individuals on sick-leave for 2–12 months with a sick-leave diagnosis within the musculoskeletal, psychological or general and unspecified chapters of ICPC-2. Potential participants were identified in the Social Security System Registry. The multicomponent inpatient program (4 + 4 days) consisted of Acceptance and Commitment Therapy, physical training and work-related problem-solving including creating a return to work plan and a workplace visit if considered relevant. The comparative outpatient program consisted primarily of ACT (6 sessions during 6 weeks). Self-reported health-related quality of life, subjective health complaints, pain and anxiety and depression symptoms were assessed up to 12 months after the program. Results 168 individuals were randomized to the multicomponent inpatient program (n = 92) or the outpatient program (n = 76). Linear mixed models showed no statistically significant differences between the programs, except for slightly more reduced pain after the outpatient program. Conclusions This study presents no support that a 4 + 4 days multicomponent inpatient rehabilitation program is superior to a less comprehensive outpatient program, in improving health outcomes.

Keywords Return to work • Sick leave • Musculoskeletal diseases • Absenteeism • Cognitive therapy

Introduction

Musculoskeletal and mental health disorders are the two leading causes of sickness absence in Norway [1]. Five percent of the gross domestic product is spent on disability and sickness benefits, and this is by far the highest level in the OECD countries [2].

Most occupational rehabilitation programs described in the scientific literature are directed towards specific diagnostic groups, mainly musculoskeletal disorders [3, 4]. Effects reported in the literature are ambiguous. For example, Jensen et al. [5] did not find added effects on return to work or pain reduction of multidisciplinary occupational rehabilitation compared to a brief intervention program for subjects with low back pain. In contrast, Lambeek et al. [6] and Loisel et al. [7] reported that multidisciplinary occupational rehabilitation led to
increased return to work and reduced disability, but with little effect on pain. Others have found increased return to work rates, but no effect on functional status [8]. Studies on the effect of return to work programs for individuals with mental health disorders have also showed increased return to work, but no added reduction in symptoms [9, 10]. However, a recent study showed that work-focused cognitive behavioral therapy for individuals on sick leave with common mental disorders was more effective than usual care in reducing depression and anxiety symptoms, increasing health-related quality of life, as well as increasing or maintaining work participation [11].

In Norway, the occupational rehabilitation services offer both inpatient and outpatient programs to increase work participation and improve health outcomes for patients on sick-leave, and the inclusion of different diagnostic groups in the same rehabilitation programs has been common practice for several years [12]. However, effects of such programs have never been evaluated with a rigorous study design. Recently, we participated in developing a multicomponent occupational rehabilitation program [13]. The program consisted of cognitive behavioral therapy, physical training, creating a return to work plan, and a workplace visit if considered relevant by the participant and rehabilitation team. Physical exercise has been shown to reduce depression [14], seems to reduce pain [15] and is recommended as an adjunctive treatment for anxiety disorders [16]. Different diagnostic groups were included in the program. All activity at the center was framed within a cognitive behavioral therapy approach in the form of Acceptance and Commitment Therapy (ACT) [17]. The ACT model emphasizes accepting both negative and positive experiences, while focusing on a person’s values to guide them towards their goals [17]. In specific diagnostic groups there has been an increasing number of studies on the effect of ACT [18]. Although there are some inconsistencies in the literature [18, 19], studies suggest that ACT may have beneficial effects on chronic pain [20], anxiety [21, 22] and depression [21, 23].

We evaluated the effects of the multicomponent program delivered at the inpatient occupational rehabilitation center by comparing it to a less comprehensive outpatient program consisting mainly of ACT. In a recent study with 12 months of follow-up we found no difference between the programs on number of sickness absence days and return to work (under review). Here, we present results of secondary outcomes related to health as the programs also aimed to improve the participants’ health status and health perception.

We hypothesized that the inpatient program, to a greater extent than the outpatient program, would reduce pain, depression, anxiety and subjective health complaints and increase function and health-related quality of life.

Methods

Study Design and Participants

We conducted a randomized clinical trial with parallel groups, comparing an inpatient multicomponent program (4 + 4 days) with a single-component program (6 sessions during 6 weeks) (hereafter referred to as the inpatient- and outpatient program, respectively) for individuals on sick-leave due to musculoskeletal-, unspecified-, or common mental health disorders. Details about the study design have been published elsewhere [13]. The primary outcome in the main study was sickness absence (under review). The current study assesses effects on somatic and mental health in the inpatient program versus the outpatient program through 12 months follow-up. The study was approved by the Regional Committee for Medical and Health Research Ethics in Central Norway (No.: 2012/1241), and the trial is registered in clinicaltrials.gov (No.: NCT01926574). The results are presented according to the CONSORT statement [24].

Eligible participants were individuals aged 18 to 60 years sick listed 2–12 months with a diagnosis within the musculoskeletal (L), psychological (P) or general and unspecified (A) chapters of ICPC-2 (International Classification of Primary Care, Second edition). Sick leave status at inclusion had to be at least 50 % off work. Exclusion criteria, assessed by a questionnaire and an outpatient screening performed by a physician, a physiotherapist and a psychologist, were: (1) alcohol or drug abuse; (2) serious somatic (e.g. cancer, unstable heart disease) or psychiatric disorders (e.g. high suicidal risk, psychosis, ongoing manic episode); (3) specific disorders requiring specialized treatment; (4) pregnancy; (5) currently participating in another treatment or rehabilitation program; (6) insufficient oral or written Norwegian language skills to participate in group sessions and fill out questionnaires; (7) scheduled for surgery within the next 6 months; and (8) serious problems with functioning in a group setting.

Data was obtained by questionnaires and filled out at six time-points: at screening before inclusion, at the start of the program, at the end of the program, and three, six and 12 months after the inpatient program ended.

Programs

The Inpatient Program

Consisted of group discussions (ACT based) led by team coordinators, individual and group based physical training, mindfulness, psychoeducation on stress and individual meetings with coordinator for work-related problem-
solving and creating a return to work plan. The intervention lasted four full workdays in week 1 and week 4 (8 days in total; 6–7 h each day), separated by 2 weeks at home (week 2 and 3). The two weeks at home included at least two contacts with the team coordinator (in person or by telephone) and a meeting with the employer if regarded relevant and the participant gave permission. A certified ACT instructor supervised the coordinators who mentored the participants before and during (monthly) the intervention. The program took place at Hysnes rehabilitation center, established as a part of St. Olavs Hospital, in central Norway. A more detailed description of the program has been published elsewhere [13].

**The Outpatient Program**

Consisted of group based ACT. The sessions were held at the Department of Physical Medicine and Rehabilitation at St. Olavs Hospital once a week for six weeks, each session lasting 2.5 h. One of two physicians (specialists in Physical Medicine and Rehabilitation) or a psychologist, all three educated in ACT, led the sessions. The participants were encouraged to practise at home between sessions, including a daily 15 min audio-guided mindfulness practice. In addition the participants were offered two individual sessions with a social worker experienced in occupational rehabilitation and trained in ACT to clarify personal values and work-related issues. The intervention also included a motivational group discussion with a physiotherapist on the benefits of physical training. An individual session with both the social worker and ACT therapist present ended the intervention. In this session a summary letter was written to the participant’s general practitioner. A more detailed description of the program has been published elsewhere [13].

**Outcome Measures**

Self-reported data on health and functioning were collected via internet-based questionnaires. The participants received text messages on their mobile telephone when it was time to answer questionnaires and as reminders if they did not respond. If they had not responded after two text-message reminders a project co-worker made a final phone call to remind the participant.

Anxiety and depression were recorded using The Hospital Anxiety and Depression Scale (HADS) [25]. It consists of 14 items, where seven items measure anxiety and seven measure depression symptoms. It is scored on a 4-point Likert scale according to intensity of symptoms the last week. The maximum score is 21 on each subscale. HADS is widely used and has been found to perform well in assessing severity and detecting anxiety and depression, with a cut-off of 8 giving an optimal balance between sensitivity and specificity [26]. HADS was answered at all time-points, except at six months.

Common somatic and mental health problems were recorded using The Subjective Health Complaints Inventory (SHC) [27], which registers complaints in five subscales: musculoskeletal pain, pseudoneurology, gastrointestinal problems, allergy and flu. It consists of 29 questions regarding complaints experienced the last month—each scored on a 4-point Likert scale from 0 “not at all” to 3 “serious”. A severity score can be reported for each subscale or as a total score (score range 0–87) [27]. The questionnaire was answered at the start of the program, and three and 12 months after the program.

To assess pain we used two questions from the Brief Pain Inventory (BPI) [28]. The participants were asked to grade the strongest and average pain during the last week on a 0 (no pain) to 10 (worst imaginable pain) numeric rating scale. The pain questions were answered at all time-points, except at 6 months.

Health-related quality of life was recorded using 15D [29]. It contains 15 dimensions covering physical, mental and social well-being and generates a total score ranging from 1 (no problem on any dimension) to 0 (being dead). It has been suggested that the generic minimal important change is ±0.015 and a large change is ±0.035 [30]. It should be noted that in the Alanne et al. study the cut-off for “slightly better” for pain and depression alone were 0.036 and 0.051, respectively. 15 D was answered at all time-points, except at screening and the end of the program.

Functioning was recorded using COOP/WONKA [31]. It offers a self-reporting assessment of function in six domains. We used four of the domains: physical fitness, feelings, daily activity and social activity. Each domain is scored on a 5-point Likert scale from 1 (no problem/not affected) to 5 (huge problems/considerably affected). Answers were used as a continuous score (range 1–5). It was included at all time-points, except at screening and six months.

Participants were asked to evaluate their general health on a 4-point Likert scale from 1 “poor” to 4 “very good”. The variable was analysed both dichotomized (poor/not very good vs. good/very good) and as a continuous score (range 1–4). The question was answered at all time-points, except at screening.

**Randomization and Blinding**

The present study was part of a larger trial comparing the current (4 + 4 days) and a longer (3.5 weeks) inpatient program, an outpatient program, as well as a treatment as usual control group only followed in sick leave registers.
The current study reports on health outcomes in the 4 + 4 days inpatient program and the comparative outpatient program.

Participants were randomized twice. Firstly, sick listed individuals identified in the Social Security System were randomized to receive an invitation to either the long or short program. Invited participants randomized to the short program completed a short initial questionnaire assessing eligibility. Those eligible were invited for an outpatient screening assessment. If the screening was passed (Fig. 1), the second randomization allocated the subjects to either the inpatient or the outpatient program. The first identified as a potential participant (n=12,007)

Randomization 1

No invitation
Untouched control group (n=4,881)

Received study invitation
Short program (n=3,318)

Accepted (n=275)
Declined (n=799)
No answer (n=1,997)

Letter in return (n=58)
Excluded (n=189)
- Length of sick-leave (n=1)
- Degree of sick-leave (n=13)
- No longer on sick-leave/started to return to work (n=66)
- Serious somatic or psychological disorder (n=6)
- A specific disorder requiring specialized treatment (n=3)
- Pregnancy (n=2)
- Currently participating in another treatment program (n=63)
- Insufficient Norwegian comprehension (n=6)
- Scheduled surgery next 6 months (n=8)
- Problems functioning in a group (n=1)
- Moved to another part of the country (n=1)
- Medical assessment not completed (n=19)

Outpatient pre-screening (n=275)

Randomization 2 (n=168)

Allocated to the inpatient program (n=92)
Completed program (n=74)
- No longer on sick-leave/started to return to work (n=10)
- Logistic problems with childcare (n=2)
- Timing not good (n=1)
- Unknown (n=1)
Withdrawal during program (n=4)
- Health issues (n=1)
- Unknown (n=3)

Answered questionnaires
Screening: 100% (n=92)
Start of program: 85% (n=78)
End of program: 71% (n=65)
3 months after program: 50% (n=46)
6 months after program: 58% (n=53)
12 months after program: 43% (n=40)

Analyzed (n=92)

Allocated to the outpatient program (n=76)
Completed program (n=63)
- No longer on sick-leave (n=2)
- Currently participating in another treatment program (n=1)
- Wanted the multicomponent program (n=1)
- Unknown (n=1)
Withdrawal during program (n=8)
- Not motivated (n=3)
- Travel time (n=1)
- Unknown (n=4)

Answered questionnaires
Screening: 99% (n=75)
Start of program: 78% (n=59)
End of program: 59% (n=45)
3 months after program: 49% (n=37)
6 months after program: 54% (n=41)
12 months after program: 47% (n=36)

Analyzed (n=76)

Fig. 1 Participant flow through the study
randomization was performed by a project co-worker. In the second allocation a flexibly weighted randomization procedure was provided by the Unit of Applied Clinical Research (third-party) at the Norwegian University of Science and Technology (NTNU), to ensure that the rehabilitation center had enough participants to run monthly groups in periods of low recruitment.

It was not possible to blind neither the participants nor the caregivers for treatment. Outcomes were measured using web-based questionnaires that the participants filled out independently on their own. The researchers were not blinded.

Statistics

Sample size was calculated based on the primary outcome, i.e. number of sickness absence days (under review), resulting in 80 persons in each arm. Details about the estimations are published elsewhere [13].

Under the intention to treat principle we used linear (and logistic) mixed-effects models to compare outcome measures of health and function over time for the two rehabilitation programs. In addition to program and time (time points 1–6) we included an interaction term between program and the six time-points in the analyses to assess whether the effects of the programs differed over time. Repeated measurements (at the different time-points) were handled by including a random intercept for person in the models (thereby allowing the participants to start out at different levels) and a random slope for time (allowing individual development over time). The estimates from the analyses (fixed effects) were used to predict health outcomes at different time points for the two programs. We considered \( p \) values (two-tailed) <0.05 to be statistically significant. Precision was assessed using 95\% confidence intervals.

In the main analyses we did not adjust for baseline characteristics, but this was done in sensitivity analyses (gender, age, sick leave diagnosis, work status, education level and type of benefit) to assess the robustness of the results. Supplementary “per protocol” analyses were done by excluding participants that withdrew after randomization (before or during the programs) and/or attended less than 60\% of the sessions of the outpatient program.

Baseline characteristics for responders and non-responders to the 12 month follow-up questionnaire were compared using \( \chi^2 \) test, \( t \) test or Mann–Whitney U test. Median numbers of sickness absence days were compared by Mann–Whitney U test.

All analyses were done using STATA 13.1 (StataCorp. 2013, Stata Statistical Software: Release 13. College Station, TX: StataCorp LP).

Results

In brief, 12,007 potential participants from the regional area were identified in the National Social Security System Registry. Of these, 3,318 were randomized to receive an invitation to the short program and 275 accepted. After screening 107 persons were excluded, withdrew or did not meet for their appointment. The remaining 168 persons were randomized to the inpatient program (\( n = 92 \)) or the outpatient program (\( n = 76 \)). The groups consisted of maximum 9 participants. The flow of participants through the study is illustrated in Fig. 1.

For the inpatient program, 14 people withdrew before they began the program and four quit during the program. For the outpatient program, five people withdrew before the program started and eight during the program. Those who started the outpatient program attended on average 7.9 of the 10 meetings and 59 (83\%) attended at least 60\% of the sessions. For the inpatient program there is no data for number of sessions participants attended, but as it was an inpatient program the participants were assumed compliant if they did not withdraw. All randomized participants were included in the analyses.

The number of people who answered the questionnaires decreased steadily through the study. For the inpatient program 100\% of the participants answered the questionnaire before the screening, 85\% at the start of program, 71\% at the end of the program, 50\% at three months, 58\% at six months and 43\% at 12 months after the program. For the outpatient program the numbers were 99, 78, 59, 49, 54 and 47\%, respectively. One participant in the outpatient program answered none of the questionnaires. At least 3 questionnaires were filled out by 72\% of the participants. A workplace visit was performed for 13\% of the participants randomized to the inpatient program.

Participant Characteristics

The participants were mainly women (79\%), and their mean age was 45 years (SD 9.1) (Table 1). The majority (65\%) of the participants worked full-time prior to their sick leave, 18\% worked part time, 4\% had a graded disability pension and 13\% had no job. About half were on full sick-leave (45\%) and half on graded sick-leave (48\%). A smaller part (7\%) received work assessment allowance, which can be applied for in Norway after being on sick leave for a year. The latter group consisted of individuals who were invited to the study just before their benefit was changed from sick-leave to work assessment allowance. The median number of days on sick-leave the last 12 months before inclusion in the study (i.e. second randomization) was 226 days (interquartile range (IQR) 189–271). Sick-leave diagnoses within the
musculoskeletal chapter in ICPC-2 were most common (52 %), followed by psychological (38 %) and general and unspecified (10 %). The baseline characteristics of the participants in the two programs were fairly similar (Table 1).

### Outcome Measures

#### Comparison of Intervention Groups

Only one of the health measures, strongest pain, showed a statistically significant difference between the programs (Fig. 2 and Table 2). The estimated mean difference in strongest pain from start of the program to 12 months was 1.1 (95 % CI 0.1–2.0, \( p = 0.03 \)) in favor of the outpatient program.

#### Development of Health Outcomes Over Time

Both programs showed increased health-related quality of life from start of the programs to 12 months (Table 2). The other health measures showed no or marginal changes (Table 2 and online supplementary Table 1).

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### Table 1 Baseline characteristics of participants

|                          | Inpatient program (n = 92) | Outpatient program (n = 76) |
|--------------------------|---------------------------|-----------------------------|
| Age mean (SD)            | 45.0 (8.7)                | 45.1 (9.6)                  |
| Women n (%)              | 71 (77 %)                 | 62 (82 %)                   |
| Higher education\(^b\) n (%) | 45 (49 %)                | 31 (41 %)                   |
| Work status n (%)        |                           |                             |
| No work                  | 15 (16 %)                 | 7 (9 %)                     |
| Full time                | 57 (62 %)                 | 52 (68 %)                   |
| Part time                | 15 (16 %)                 | 16 (21 %)                   |
| Graded disability pension| 5 (5 %)                   | 1 (1 %)                     |
| Sick-leave status\(^c\) n (%) |                      |                             |
| Full sick-leave          | 41 (45 %)                 | 35 (46 %)                   |
| Partial sick-leave       | 45 (49 %)                 | 36 (47 %)                   |
| Work assessment allowance| 6 (7 %)                   | 5 (7 %)                     |
| Main diagnoses for sick-leave (ICPC-2)\(^d\) n (%) |                    |                             |
| A-general and unspecified| 9 (10 %)                  | 7 (9 %)                     |
| L-musculoskeletal        | 48 (52 %)                 | 40 (53 %)                   |
| P-psychological          | 35 (38 %)                 | 29 (38 %)                   |
| Length of sick leave at inclusion\(^e\)\(^d\) | | |
| Median days (IQR)        | 224 (189–262)             | 229 (187–275)               |
| HADS mean (SD)           |                           |                             |
| Anxiety (0–21)           | 7.8 (4.4)                 | 7.4 (4.3)                   |
| Depression (0–21)        | 6.7 (4.3)                 | 6.0 (4.1)                   |
| Pain level mean (SD)     |                           |                             |
| Average pain (0–10)      | 4.7 (2.3)                 | 4.6 (2.0)                   |
| Strongest pain (0–10)    | 5.4 (2.5)                 | 5.9 (2.0)                   |
| Quality of life 15D (0–1) |                           |                             |
| Mean (SD)                | 0.79 (0.10)               | 0.79 (0.09)                 |
| Subjective health evaluation n (%) | | |
| Poor                     | 7 (8 %)                   | 10 (13 %)                   |
| Not so good              | 55 (60 %)                 | 39 (51 %)                   |
| Good                     | 15 (16 %)                 | 10 (13 %)                   |
| Very good                | 1 (1 %)                   | 0                           |
| No response              | 14 (15 %)                 | 17 (22 %)                   |

* \(^{a}\) Work status, sick-leave status, diagnosis and length of sick leave recorded at inclusion. Education, HADS and pain recorded at screening. Quality of life and subjective health evaluation recorded at start of program

* \(^{b}\) Higher (tertiary) education (College or university)

* \(^{c}\) Based on data in the medical certificate from the National Social Security System Registry

* \(^{d}\) Number of days on sick leave during the last 12 months prior to inclusion. Measured as calendar days, not adjusted for graded sick leave or part time job

* \(^{e}\) Median days (IQR)
Per Protocol, Sensitivity and Post Hoc Subgroup Analyses

The “per protocol” analyses provided only minor changes in the estimates. The estimated difference between the programs was statistically significant for average and strongest pain, in favour of the outpatient program. The main analyses were repeated adjusting for gender, age, diagnosis, education level, work status and type of benefit received. There were only small changes in the estimates and the adjusted analyses did not change any conclusions about the programs.

We performed subgroup analyses for HADS and average pain according to the two main diagnostic groups (see online supplementary Tables 2 and 3). For the HADS depression subscale there was a somewhat larger reduction in symptoms for participants with a psychological diagnosis. The same was observed for the HADS anxiety subscale for the inpatient program, while for the outpatient program there were only minor differences between the diagnostic groups. For average pain there was little difference between participants with a musculoskeletal- and psychological diagnosis for both programs. When performing the analyses for participants having the highest baseline scores on anxiety, depression and pain the results were similar to the main analyses. The differences between the two programs from start of the programs to 12 months were not statistically significant in any of the subgroup analyses.

Non-Responders

The participants not answering the questionnaire at 12 months were younger than the responders (mean age 43.6 (SD 9.3) vs. 46.7 (SD 8.6), \( p = 0.023 \)). The other baseline values were fairly similar. The median number of sickness absence days during 12 months of follow-up were 87 (IQR 39–146) for the responders and 112 (IQR 44–185) for the non-responders (\( p = 0.252 \)).

Discussion

This randomized clinical trial showed no differences in self-reported health measures between a 4 + 4 days inpatient multicomponent occupational rehabilitation program and a less comprehensive outpatient program consisting mainly of group-based ACT, except for slightly more reduced pain after the outpatient program.

We are not aware of studies comparing inpatient and outpatient return to work programs. No substantial difference on somatic and mental health outcomes between the two rehabilitation programs is in line with some earlier studies on individuals with musculoskeletal complaints.
Table 2  Comparison of estimated health scores between the inpatient and the outpatient program

|                              | Inpatient program | Outpatient program | Estimated difference between programs<sup>b</sup> |
|------------------------------|-------------------|-------------------|-----------------------------------------------|
|                              | Mean          | 95 % CI          | Mean          | 95 % CI          | Mean          | 95 % CI          | p value           |
| **HADS anxiety (0–21)**      |                 |                  |                |                  |                |                  |                   |
| Screening                    | 7.8            | 6.9–8.7          | 7.3            | 6.3–8.3          |                |                  |                   |
| Start of program             | 7.6            | 6.7–8.5          | 6.6            | 5.5–7.6          |                |                  |                   |
| End of program               | 7.1            | 6.2–8.1          | 6.8            | 5.7–7.9          |                |                  |                   |
| 3 months                     | 6.6            | 5.5–7.6          | 5.3            | 4.2–6.5          |                |                  |                   |
| 12 months                    | 6.5            | 5.4–7.6          | 5.7            | 4.5–6.9          | 0.2            | −1.2 to 1.5      | 0.78              |
| **HADS depression (0–21)**   |                 |                  |                |                  |                |                  |                   |
| Screening                    | 6.7            | 5.9–7.6          | 6.0            | 5.1–7.0          |                |                  |                   |
| Start of program             | 6.8            | 5.9–7.7          | 5.8            | 4.9–6.8          |                |                  |                   |
| End of program               | 6.2            | 5.3–7.0          | 5.1            | 4.1–6.1          |                |                  |                   |
| 3 months                     | 5.4            | 4.5–6.4          | 4.3            | 3.2–5.4          |                |                  |                   |
| 12 months                    | 5.9            | 4.8–7.0          | 4.4            | 3.2–5.6          | −0.5           | −2.0 to 1.0      | 0.49              |
| **Average pain (0–10)**      |                 |                  |                |                  |                |                  |                   |
| Screening                    | 4.8            | 4.3–5.2          | 4.6            | 4.2–5.1          |                |                  |                   |
| Start of program             | 4.1            | 3.7–4.6          | 4.6            | 4.1–5.1          |                |                  |                   |
| End of program               | 3.9            | 3.5–4.4          | 4.3            | 3.8–4.9          |                |                  |                   |
| 3 months                     | 3.6            | 3.1–4.1          | 3.8            | 3.2–4.4          |                |                  |                   |
| 12 months                    | 4.0            | 3.4–4.6          | 3.7            | 3.1–4.3          | −0.8           | −1.5 to 0.0      | 0.06              |
| **Strongest pain (0–10)**    |                 |                  |                |                  |                |                  |                   |
| Screening                    | 5.4            | 4.9–5.9          | 6.0            | 5.5–6.5          |                |                  |                   |
| Start of program             | 5.0            | 4.5–5.5          | 6.0            | 5.5–6.6          |                |                  |                   |
| End of program               | 4.9            | 4.4–5.4          | 5.5            | 4.8–6.1          |                |                  |                   |
| 3 months                     | 4.7            | 4.1–5.3          | 4.9            | 4.2–5.5          |                |                  |                   |
| 12 months                    | 5.1            | 4.4–5.8          | 5.1            | 4.3–5.8          | −1.1           | −2.0 to −0.1     | 0.03              |
| **Health-related quality of life (0–1)** |                 |                  |                |                  |                |                  |                   |
| Start of program             | 0.79           | 0.77–0.81        | 0.79           | 0.76–0.81        |                |                  |                   |
| 3 months                     | 0.82           | 0.80–0.85        | 0.83           | 0.80–0.85        |                |                  |                   |
| 6 months                     | 0.82           | 0.80–0.85        | 0.84           | 0.81–0.86        |                |                  |                   |
| 12 months                    | 0.82           | 0.80–0.85        | 0.83           | 0.80–0.86        | −0.02          | −0.05 to 0.02    | 0.41              |
| **SHC total** (0–87)         |                 |                  |                |                  |                |                  |                   |
| Start of program             | 15.2           | 13.4–16.9        | 15.9           | 13.9–17.9        |                |                  |                   |
| 3 months                     | 13.3           | 11.3–15.3        | 13.8           | 11.6–16.1        |                |                  |                   |
| 12 months                    | 14.3           | 12.2–16.5        | 13.9           | 11.6–16.2        | −1.2           | −3.8 to 1.5      | 0.39              |
| **SHC musculoskeletal pain** (0–24) |                 |                  |                |                  |                |                  |                   |
| Start of program             | 6.0            | 5.2–6.8          | 6.7            | 5.7–7.6          |                |                  |                   |
| 3 months                     | 5.2            | 4.3–6.2          | 6.0            | 4.9–7.1          |                |                  |                   |
| 12 months                    | 5.7            | 4.7–6.7          | 5.5            | 4.4–6.6          | −0.9           | −2.1 to 0.4      | 0.17              |
| **SHC pseudoneurology** (0–21) |                 |                  |                |                  |                |                  |                   |
| Start of program             | 5.0            | 4.3–5.7          | 5.1            | 4.3–5.9          |                |                  |                   |
| 3 months                     | 4.4            | 3.6–5.2          | 4.0            | 3.2–4.9          |                |                  |                   |
| 12 months                    | 4.4            | 3.6–5.2          | 4.8            | 4.0–5.6          | 0.3            | −0.8 to 1.4      | 0.58              |
| **SHC gastrointestinal problems (0–21)** |                 |                  |                |                  |                |                  |                   |
| Start of program             | 2.1            | 1.6–2.6          | 2.2            | 1.7–2.8          |                |                  |                   |
| 3 months                     | 1.8            | 1.2–2.3          | 2.1            | 1.4–2.7          |                |                  |                   |
However, we performed subgroup analyses according to the statistical power to detect between group effects. This would likely reduce the possibility of bias due to loss to follow-up. The key element of both the inpatient and outpatient program in the present study was ACT. Differences between the two programs, in addition to the inpatient versus outpatient setting, were that the inpatient program was more extensive and included physical training, creation of a return to work plan and a workplace visit in 13 % of the cases. However, these additional components did not induce additional benefits.

Most occupational rehabilitation programs described in the scientific literature use some sort of cognitive behavioral therapy approach [6, 11]. In ACT the participants are encouraged to accept pain rather than try to control it. It has therefore been argued that pain might not be the best outcome measure for acceptance-based therapies [33]. This also applies to several of the other outcomes in this study like anxiety and depression, as ACT emphasize behavior change and not symptom reduction [34]. This is line with our findings of modest changes for these outcomes. The outpatient program was slightly more effective in reducing one pain variable, but the difference was not clinically significant and due to the number of statistical tests performed this result should be interpreted with caution.

We found an increase in health related quality of life in both groups measured by 15D, estimated to be 0.03 (95 % CI 0.01–0.06) for the inpatient program and 0.05 (95 % CI 0.02–0.07) for the outpatient program. The clinical importance of this change is uncertain, but it is in the area of cut-off suggested as a minimum important change [30]. When this is compared to the rather small changes observed on the other measures this might suggest that the focus of ACT on values and acceptance of negative experiences in life might have changed how the participants perceive their quality of life despite little change in health symptoms.

Few randomized studies have included participants with different diagnoses in the same return to work programs. As we included individuals on sick leave due to musculoskeletal, mental or general/unspecific disorders, some had pain and others not, which was also the case for anxiety and depression symptoms. This would likely reduce the statistical power to detect between group effects. However, we performed subgroup analyses according to the participants’ main sick-leave diagnosis. Participants with a psychological diagnosis had a somewhat larger reduction in depression symptoms than participants with a musculoskeletal diagnosis. However, there was no difference between participants with musculoskeletal and psychological diagnoses in reductions of average pain. As a substantial degree of overlap in symptoms is common in these patients [35, 36] and the diagnostic labelling by the general practitioner may be somewhat arbitrary [37], we performed subgroup analyses for highest baseline scores on anxiety and depression symptoms and average pain. The estimates were fairly similar to the main analyses. It should be noted that the post hoc subgroup analyses were not planned a priori.

The main strength of this randomized study was that all participants were invited from the Social Security System Registry, meaning there was no referral bias. Return to work rehabilitation centers have existed for about 30 years in Norway, but this is the first randomized controlled study investigating effects on somatic or mental health of such programs. The programs were not diagnosis specific and add important knowledge to a field where previous research has focused on diagnosis specific interventions. Also, the study included a broad range of validated health-related measures.

Some limitations should be addressed. Firstly, the response rate for the questionnaires were low at 3, 6 and 12 months. At the start and the end of the programs, more people answered the questionnaires in the inpatient program than in the outpatient program. The participants in the inpatient program answered the questionnaire at the rehabilitation center, while the outpatient participants did it at home. During follow-up, questionnaires were answered at home for both groups and the numbers of missing questionnaires were similar. We therefore assume that the structural differences in collecting questionnaire data account for the differences in responses between the two groups at the start and end of the intervention. For analyses we used linear mixed models which are less sensitive to missing values in outcome data. Still, these models rely on the assumption of “missing at random”, and we cannot disregard the possibility of bias due to loss to follow-up.
However, we consider it unlikely that such bias should influence the two groups differentially and thereby the main results of the study. This assumption is strengthened by register-based sick leave data showing a similar number of sick leave days during 12 months of follow-up between participants responding/not responding to the questionnaire at 12 months.

In the current study there was no usual care control group. Therefore, we cannot distinguish between the effects of rehabilitation and time. It should also be noted that the power calculation for the study was done with regard to the primary outcome (sickness absence) and not the secondary outcomes presented in this article.

With regards to external validity it should be noted that from the over 3000 invitations sent, only 275 individuals accepted the invitation. A possible explanation might be that they had to be prepared to be away from their family for 2 weeks if randomized to the inpatient program. With only about 8% of the invited accepting the invitation the generalizability of the results is a challenge. However, it should be noted that only a small portion of people on sick leave in Norway are referred to occupational rehabilitation centers. By inviting participants this broadly we were able to reach all individuals on sick-leave with these diagnoses without referral bias induced by the general practitioner.

Conclusions

There was no substantial difference between the programs on somatic and mental health; hence, this study presents no support that a 4 + 4 days inpatient multicomponent rehabilitation program is superior to a less comprehensive outpatient program. Whether a longer lasting inpatient program will have greater effects on somatic and mental health will be investigated in an upcoming study.

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Compliance with Ethical Standards

Conflict of interest

Marius Steiro Finland has been employed at Hysnes Rehabilitation Center, St. Olavs Hospital. Sigmund Gismervik is employed at Department of Physical Medicine and Rehabilitation, St. Olavs Hospital. The other authors declare that they have no conflict of interest.

Ethical approval

All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. Informed consent was obtained from all individual participants included in the study.

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References

1. NAV statistics. NAV sickness absence statistics: NAV. 2014. https://www.nav.no/no/Nav+-+og+samfunn/Statistikk/Sykefravar+-+statistikk/Sykefravar/Sykefravar-Statistikknotater.
2. OECD. Mental health and work. Norway: OECD Publishing; 2013.
3. Schandelmaier S, Ebrahim S, Burkhartde SC, de Boer WE, Zambrunn T, Guyatt GH, et al. Return to work coordination programmes for work disability: a meta-analysis of randomised controlled trials. PLoS One. 2012;7(11):e49760.
4. Staal JB, de Rijk A, Houkes I, Heymans MW. Clinical interventions to reduce work disability in workers with musculoskeletal disorders or mental health problems. In: Loisel P, Anema JR, editors. Handbook of work disability Prevention and management. New York: Springer; 2013. p. 317–34.
5. Jensen C, Jensen OK, Christiansen DH, Nielsen CV. One-year follow-up in employees sick-listed because of low back pain: randomized clinical trial comparing multidisciplinary and brief intervention. Spine. 2011;36(15):1180–9.
6. Lambeek LC, van Mechelen W, Knol DL, Loisel P, Anema JR. Randomised controlled trial of integrated care to reduce disability from chronic low back pain in working and private life. BMJ. 2010;340:c1035.
7. Loisel P, Abenhaim L, Durand P, Edaule JM, Suissa S, Gosselin L, et al. A population-based, randomized clinical trial on back pain management. Spine. 1997;22(4):2911–8.
8. Vermeulen SJ, Anema JR, Schellart AJ, Knol DL, van Mechelen W, van der Beek AJ. A participatory return-to-work intervention for temporary agency workers and unemployed workers sick-listed due to musculoskeletal disorders: results of a randomized controlled trial. J Occup Rehabil. 2011;21(3):313–24.
9. Blonk RWB, Brenninkmeijer V, Lagerveld SE, Houtman ILD. Return to work: a comparison of two cognitive behavioural interventions in cases of work-related psychological complaints among the self-employed. Work Stress. 2006;20(2):129–44.
10. van der Klink JJ, Blonk RW, Schene AH, van Dijk FJ. Reducing long term sickness absence by an activating intervention in adjustment disorders: a cluster randomised controlled design. Occup Environ Med. 2003;60(6):429–37.
11. Reme SE, Grasdal AL, Lovvik C, Lie SA, Overland S. Work-focused cognitive-behavioural therapy and individual job support to increase work participation in common mental disorders: a randomised controlled multicentre trial. Occup Environ Med. 2015;72:745–52.
12. Braathen TN, Veiersted KB, Heggenes J. Improved work ability and return to work following vocational multidisciplinary rehabilitation of subjects on long-term sick leave. J Rehabil Med. 2007;39(6):493–9.

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13. Fimland MS, Vasseljen O, Gismervik S, Rise MB, Halsteinli V, Jacobsen HB, et al. Occupational rehabilitation programs for musculoskeletal pain and common mental health disorders: study protocol of a randomized controlled trial. BMC Public Health. 2014;14(1):368.

14. Josefsson T, Lindwall M, Archer T. Physical exercise intervention in depressive disorders: meta-analysis and systematic review. Scand J Med Sci Sports. 2014;24(2):259–72.

15. Hayden JA, van Tulder MW, Malmivaara A, Koes BW. Exercise therapy for non-specific low back pain. Cochrane Database Syst Rev. 2005;3:CD000335.

16. Jayakody K, Gunadasa S, Hosker C. Exercise for anxiety disorders: systematic review. Br J Sports Med. 2014;48(3):187–96.

17. Hayes SC, Strosahl K, Wilson KG. Acceptance and commitment therapy: an experiential approach to behavior change, vol. 1. New York: Guilford Press; 1999. p. 304.

18. Ost LG. The efficacy of Acceptance and Commitment Therapy: an updated systematic review and meta-analysis. Behav Res Ther. 2014;61:105–21.

19. Ruiz FJ. Acceptance and commitment therapy versus traditional cognitive behavioral therapy: A systematic review and meta-analysis of current empirical evidence. Int J Psychol Psychother. 2012;12(3):333–58.

20. Wetherell JL, Afari N, Rutledge T, Sorrell JT, Stoddard JA, Petkus AJ, et al. A randomized, controlled trial of acceptance and commitment therapy and cognitive-behavioral therapy for chronic pain. Pain. 2011;152(9):2098–107.

21. Forman EM, Herbert JD, Moitra E, Yeomans PD, Geller PA. A randomized controlled effectiveness trial of acceptance and commitment therapy and cognitive therapy for anxiety and depression. Behav Modif. 2007;31(6):772–99.

22. Roemer L, Orsillo SM, Salters-Pedneault K. Efficacy of an acceptance-based behavior therapy for generalized anxiety disorder: evaluation in a randomized controlled trial. J Consult Clin Psychol. 2008;76(6):1083–9.

23. Folke F, Parling T, Melin L. Acceptance and commitment therapy for depression: a preliminary randomized clinical trial for unemployed on long-term sick leave. Cogn Behav Pract. 2012;19(4):583–94.

24. Schulz KF, Altman DG, Moher D. CONSORT 2010 statement: updated guidelines for reporting parallel group randomised trials. BMC Med. 2010;8(1):18.

25. Zigmond AS, Snaith RP. The hospital anxiety and depression scale. Acta Psychiatr Scand. 1983;67(6):361–70.

26. Bjelland I, Dahl AA, Haug TT, Neckelmann D. The validity of the Hospital Anxiety and Depression Scale. An updated literature review. J Psychosom Res. 2002;52(2):69–77.

27. Eriksen HR, Ilhøeak C, Ursin H. A scoring system for subjective health complaints (SHC). Scand J Public Health. 1999;27(1):63–72.

28. Cleeland CS, Ryan KM. Pain assessment: global use of the Brief Pain Inventory. Ann Acad Med Singapore. 1994;23(2):129–38.

29. Sintonen H. The 15D instrument of health-related quality of life: properties and applications. Ann Med. 2001;33(5):328–36.

30. Alanne S, Roine RP, Rasanen P, Vainiola T, Sintonen H. Estimating the minimum important change in the 15D scores. Qual Life Res. 2015;24(3):599–606.

31. Nelson E, Wasson J, Kirk J, Keller A, Clark D, Dietrich A, et al. Assessment of function in routine clinical practice: description of the COOP Chart method and preliminary findings. J Chronic Dis. 1987;40(Suppl 1):55–69S.

32. Marchand GH, Myhre K, Leivseth G, Sandvik L, Lau B, Bautz-Holter E, et al. Change in pain, disability and influence of fear-avoidance in a work-focused intervention on neck and back pain: a randomized controlled trial. BMC Musculoskelet Disord. 2015;16(1):94.

33. Veehof MM, Oskam MJ, Schreurs KM, Bohlmeijer ET. Acceptance-based interventions for the treatment of chronic pain: a systematic review and meta-analysis. Pain. 2011;152(3):533–42.

34. Hann KE, McCracken LM. A systematic review of randomized controlled trials of Acceptance and Commitment Therapy for adults with chronic pain: Outcome domains, design quality, and efficacy. J Context Behav Sci. 2014;3(4):217–27.

35. Reme SE, Tangen T, Moe T, Eriksen HR. Prevalence of psychiatric disorders in sick listed chronic low back pain patients. Eur J Pain. 2011;15:1075–80.

36. Von Korff M, Crane P, Lane M, Miglioretti DL, Simon G, Saunders K, et al. Chronic spinal pain and physical-mental comorbidity in the United States: results from the national comorbidity survey replication. Pain. 2005;113(3):331–9.

37. Maeland S, Werner EL, Rosendal M, Jonsdottir IH, Magnussen LH, Ursin H, et al. Diagnoses of patients with severe subjective health complaints in Scandinavia: a cross sectional study. ISRN Public Health. 2012. doi:10.5402/2012/851097.