Multidisciplinary pain management program for patients with chronic musculoskeletal pain in Japan: a cohort study

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Introduction: Multidisciplinary pain management is a useful method for treating chronic musculoskeletal pain. Currently, few facilities in Japan offer multidisciplinary pain treatment, especially in the inpatient setting. We implemented a multidisciplinary pain management program based on International Association for the Study of Pain recommendations. This study described our initial efforts in implementing the program, and reported 3- and 6-month follow-up results.

Materials and methods: Our pain management team included orthopedic surgeons, psychiatrists, nurses, physical therapists, clinical psychologists, pharmacists, and nutritionists. The 3-week inpatient pain management program comprised exercise therapy, psychotherapy, and patient education. We evaluated patients using the Brief Pain Inventory (BPI), Pain Catastrophizing Scale (PCS), Pain Disability-Assessment Scale (PDAS), Hospital Anxiety and Depression Scale (HADS), Pain Self-Efficacy Questionnaire (PSEQ), EuroQol Five Dimensions (EQ-5D), and physical examinations (flexibility, muscle endurance, walking ability, and physical fitness). Statistical analyses were performed using paired t-tests and Wilcoxon matched-pairs signed-rank sum tests with Bonferroni correction after the Friedman test.

Results: Data for 23 patients were analyzed before and immediately after the program. Statistically significant improvements were seen in BPI, PCS, PDAS, HADS, PSEQ, EQ-5D, flexibility, muscle endurance, walking ability, and physical fitness. Eight patients were also assessed 3 and 6 months after the program. PCS (rumination and helplessness) scores and flexibility showed significant improvement at 3 and 6 months. Significant improvement was seen in PDAS and HADS (anxiety) scores and muscle endurance at 6 months, and in PSEQ scores immediately and at 3 and 6 months.

Conclusion: Our inpatient pain management program can improve patients’ physical function and ability to cope with chronic musculoskeletal pain, which supports improved quality of life. Our program is currently being expanded to better assist patients with chronic musculoskeletal pain.

Keywords: chronic musculoskeletal pain, multidisciplinary pain treatment, biopsychosocial model, inpatient pain management program

Plain language summary
Multidisciplinary pain management is a useful method for treating chronic musculoskeletal pain. Currently, few facilities in Japan offer multidisciplinary pain treatment, especially in the inpatient setting. We implemented a multidisciplinary pain management program based on International Association for the Study of Pain recommendations. This study described our initial efforts in implementing the program and reported 3- and
6-month follow-up results. Our pain management team included orthopedic surgeons, psychiatrists, nurses, physical therapists, clinical psychologists, pharmacists, and nutritionists. The 3-week inpatient pain management program involved exercise therapy, psychotherapy, and patient education. We evaluated patients using the Brief Pain Inventory (BPI), Pain Catastrophizing Scale (PCS), Pain Disability-Assessment Scale (PDAS), Hospital Anxiety and Depression Scale (HADS), Pain Self-Efficacy Questionnaire (PSEQ), EuroQol Five Dimensions (EQ-5D), and physical examinations (flexibility, muscle endurance, walking ability, and physical fitness). We analyzed results for 23 patients before and immediately after the program. There were statistically significant improvements in all measures immediately after the program. Eight patients were also assessed 3 and 6 months after the program. PCS (ruminating and helplessness) scores and flexibility showed significant improvement at 3 and 6 months. Significant improvement was seen in PDAS and HADS (anxiety) scores and muscle endurance at 6 months, and in PSEQ scores immediately and at 3 and 6 months. Our inpatient pain management program improves patients’ physical function and ability to cope with chronic musculoskeletal pain, supporting improved quality of life. Our program is currently being expanded to better assist patients with chronic musculoskeletal pain.

Introduction
Musculoskeletal pain is a common problem that affects millions of people worldwide. Chronic musculoskeletal pain reduces people’s quality of life (QOL) and may hinder normal social life. A multidisciplinary approach for pain management is useful for treating chronic musculoskeletal pain, and has been used in the USA since the 1950s. A biopsychosocial model of well-being is an important concept in multidisciplinary pain management. This is a general model or approach based on the assumption that biological, psychological, and social factors play significant roles in affecting human function during disease or illness. Multidisciplinary pain treatment requires special facilities. However, few facilities in Japan are able to provide multidisciplinary pain treatment, particularly in an inpatient setting. We implemented an inpatient multidisciplinary pain management program in our hospital based on the biopsychosocial model recommended by the International Association for the Study of Pain (IASP). 3–7 We previously reported our inpatient multidisciplinary pain management program was being expanded to improve support for patients with chronic musculoskeletal pain. 2 However, our previous results reflected an evaluation immediately after the program. Therefore, it was unclear whether our inpatient multidisciplinary pain management program could lead to long-term improvement for patients with chronic musculoskeletal pain.

This article aimed to describe our initial efforts in implementing the multidisciplinary pain management program for Japanese patients, which used a biopsychosocial model for pain self-management. We also aimed to report results from the 3- and 6-month follow-up.

Materials and methods

Patients
We treated 102 patients with chronic musculoskeletal pain from April 2015 to March 2018. Twenty-three patients completed our inpatient multidisciplinary pain management program. Data for these 23 patients were analyzed before and immediately after the program. Eight patients were also able to be assessed at 3 and 6 months after the program. The remaining 15 patients dropped out of this study. Two patients recovered completely and did not need to continue regular hospital visits, eight patients lived far from our hospital and could not continue regular hospital visits, four patients did not agree to regular hospital visits, and one patient had developed lung cancer.

Multidisciplinary pain management program
The pain management center at Hoshi General Hospital has a team of orthopedic surgeons, psychiatrists, nurses, physical therapists, clinical psychologists, pharmacists, and nutritionists. The center was established in April 2015 and is technically supported by the Department of Pain Medicine at Fukushima Medical University School of Medicine. First, patients are screened by orthopedic surgeons and physical therapists for mechanical and biological changes in the body. Psychiatrists and clinical psychologists then evaluate patients’ psychosocial and social factors, and radiologists assess their cortical function. A flowchart of our multidisciplinary pain treatment system is shown in Figure 1. The inclusion criteria for our 3-week inpatient multidisciplinary pain management program were: (1) patients who had difficulty working or attending school because of chronic musculoskeletal pain, (2) patients confined to life at home but that wished
to return to work or school, (3) patients who accepted our program, and (4) patients who were able to pay the hospital expenses for our inpatient program. Exclusion criteria were: (1) older adults who were unable to answer the questionnaires; (2) patients with dementia or intellectual disabilities; and (3) patients who would have difficulty participating in the program (determined by multidisciplinary conference).

The program incorporates exercise therapy, psychotherapy, and patient education. Specific details of the program are as follows. (1) Exercise therapy, which comprises physical fitness and individual training by physical therapists, including aerobic exercise (walking, underwater exercise) and strengthening and stretching of muscles. A sit-up exercise was used to strengthen trunk flexor muscles and an extension exercise was used to strengthen trunk extensor muscles. Exercise therapy took place over 6–7 hrs/week, giving a total of 20 hrs. (2) Psychotherapy and cognitive behavioral therapy were used to develop patients’ assertiveness (three 60–120 min sessions). Patients also received relaxation training (two 60–120 min sessions), and role-playing to increase healthy behaviors and decrease pain behavior (two 60–120 min sessions). All sessions were led by psychologists. (3) Patient education, which was delivered by various specialists. These sessions covered pain mechanisms (orthopedic surgeon), exercise and pacing (physical therapist), assertiveness and relaxation training (psychologist), side effects of drugs (pharmacist) and daily life habits associated with nutrition (nutritionist). Each education session lasted 30–60 mins, and patients received 20 sessions in total.

The program aims to help patients return to a functional daily life by acquiring new habits and coping methods for their pain. A characteristic of our program is that family members or significant others also participated in some education and psychotherapy sessions. Our previous report detailed preliminary results for our 3-week inpatient multidisciplinary pain management program.

Specialists’ roles

The roles of each of the seven specialist groups have been described in detail in our previous report. An additional role performed by clinical psychologists was classifying patients into three subgroups: dysfunctional (DYS-type), interpersonally distressed (ID-type), and adaptive coper (AC-type) using the West Haven-Yale Multidimensional Pain Inventory (WHYMPI). The WHYMPI is a multi-dimensional assessment instrument for patients with chronic pain. These three subgroups have specific symptom patterns that could be evaluated by cluster analysis of the nine main WHYMPI scales. Characteristics of the DYS-type subgroup were high pain severity, low function, high depression, and low life control. The ID-type subgroup was characterized by a low level of support from their partner and social environment. Finally, characteristics of the AC-type subgroup were relatively low pain.

Figure 1 Flowchart for the evaluation and diagnosis of chronic musculoskeletal pain at Hoshi General Hospital. First, orthopedic surgeons evaluate physical, neurological, and imaging findings and diagnose the presence of any specific musculoskeletal disorders. Second, physical therapists evaluate physical function. Third, psychiatrists diagnose any psychological disorders associated with chronic musculoskeletal pain, including psychiatric disorders. Fourth, clinical psychologists evaluate the psychosocial factors associated with pain. Finally, we perform brain magnetic resonance imaging and cerebral blood-flow scintigraphy to evaluate cortical function.

Abbreviations: MPI, Multidimensional Pain Inventory; CAARS, Conners’ Adult ADHD Rating Scales.
intensity and relatively low interference from pain (functional interference). We classified patients into these subgroups because subgroups were assigned during initial pain psychology interviews, and pain self-management was taught depending on subgroup assignment.

Evaluations of pain, associated factors, and physical function

We evaluated inpatients with chronic musculoskeletal pain using the Brief Pain Inventory (BPI)\(^9\) to assess pain severity, and several other scales to assess pain-related psychosocial factors. These scales included the Pain Catastrophizing Scale (PCS) to evaluate the degree of rumination, magnification, and helplessness;\(^10\) the Pain Disability Assessment Scale (PDAS);\(^11\) the Hospital Anxiety and Depression Scale (HADS) to assess anxiety and depression;\(^12\) and the Pain Self-Efficacy Questionnaire (PSEQ).\(^13\) We also administered the EuroQol five dimensions questionnaire (EQ-5D)\(^14\) to evaluate patients’ QOL. Patients’ physical function was measured using the finger-floor-distance test to evaluate the degree of flexibility, the 30-second sit-to-stand test to assess muscle endurance, the 2-step test to evaluate walking ability, and 6 mins of walking to assess physical fitness.

Statistical analysis

We compared patients’ pain, associated factors, and physical abilities before and immediately after participation in the 3-week inpatient program. Statistical analyses for each item were performed using paired \(t\)-tests. We evaluated the same measures at 3 and 6 months after completion of the program. Statistical analyses were performed using Wilcoxon matched-pairs signed-rank sum test with Bonferroni correction after the Friedman test. We considered \(p\)-values less than 0.05 to be statistically significant. Statistical analyses were performed using IBM SPSS version 25 (IBM Corp., Armonk, NY, USA).

Ethics approval and consent to participate

This study was approved by the ethics committees of the participating institutions: Fukushima Medical University (Reference number: 2429) and Hoshi General Hospital (Reference number: 27-3). All patients provided written informed consent before this study started.

This study was conducted in accordance with the Declaration of Helsinki.

Results

In total, 23 patients (eight males, 15 females) aged 20–79 years (mean age 52.2 years) participated in our inpatient program from April 2015 to March 2018. Participants’ clinical characteristics, chief complaints, structural disorders, and psychiatric diagnoses are shown in Table 1. Analysis of psychiatric diagnoses showed that six patients (one male, five females) had personality disorders and 15 patients (seven males, eight females) had pervasive developmental disorder, attention deficit hyperactivity disorder, or somatoform disorders (Table 1). Only five cases (all females) were without a personality or developmental disorder diagnosis (Table 1). Eighteen patients (78%) (six males, 12 females) showed abnormal findings on cerebral blood flow scintigraphy.

Results before and immediately after the program (23 patients)

A comparison of results for the 23 patients before and immediately after the program showed statistically significant improvements in pain and associated factors (Table 2) and physical functions (Table 3).

Results before, immediately after, and 3 and 6 months after the program (eight patients)

Eight patients (four males, four females) aged 20–69 years (mean age 42.9 years) were also evaluated at 3 and 6 months after the program (marked by bold font in Table 1). Statistically significant improvement was seen in pain and associated factors (Figure 2) and physical functions (Figure 3).

Discussion

Our inpatient multidisciplinary pain management program was based on a biopsychosocial model guided by IASP recommendations. The present study showed that pain intensity (measured with the BPI) and pain-associated elements including psychosocial factors (measured with the PCS, PDAS, HADS, and PSEQ), QOL (measured with the EQ-5D), flexibility, muscle endurance, and physical fitness were significantly improved after completing the program. In addition, pain relief and improvement in psychosocial factors (PCS rumination and helplessness, PDAS, HADS-Anxiety, and PSEQ) and physical functioning (flexibility and muscle endurance) were maintained for at least 6 months after completing the program. There were minimal detectable changes for pain-associated...
| Case       | Chief complaint                        | Structural disorder                                      | Psychiatric diagnosis                                                                 |
|------------|---------------------------------------|----------------------------------------------------------|----------------------------------------------------------------------------------------|
| 1. 41-year-old man | Low back pain                        | None                                                     | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 2. 53-year-old woman | Posterior cervical pain, bilateral omalgia, low back pain | None                                                     | Dependent personality disorder, somatoform disorders                                     |
| 3. 69-year-old woman | Posterior cervical pain, bilateral omalgia, low back pain | Spondylosis                                               | Compulsive personality disorder, somatoform disorders                                      |
| 4. 55-year-old woman | Posterior cervical pain, bilateral upper extremity numbness | Ossification of posterior longitudinal ligament (after surgery) | Narcissistic personality disorder                                                     |
| 5. 20-year-old man | Low back pain                        | None                                                     | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 6. 56-year-old woman | Posterior cervical pain, bilateral omalgia, low back pain, bilateral gonalgia | Spondylosis, knee osteoarthritis                          | Attention deficit hyperactivity disorder, dependent personality disorder, somatoform disorders |
| 7. 31-year-old woman | Posterior cervical pain, bilateral omalgia, low back pain | None                                                     | None                                                                                   |
| 8. 47-year-old woman | Low back pain                        | None                                                     | Attention deficit hyperactivity disorder, histrionicism personality disorder, somatoform disorders |
| 9. 56-year-old woman | Low back pain, left-leg pain          | None                                                     | None                                                                                   |
| 10. 51-year-old man | Low back pain                        | None                                                     | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 11. 55-year-old man | Posterior cervical pain, bilateral omalgia, back pain, bilateral arm pain, low back pain, bilateral leg pain | None                                                     | Pervasive developmental disorder, autism spectrum disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 12. 58-year-old woman | Posterior cervical pain, bilateral omalgia, low back pain, right coxalgia, bilateral gonalgia, bilateral leg pain | Right-hip osteoarthritis, 4th lumbar degenerative spondylolisthesis | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 13. 75-year-old woman | Low back pain, bilateral leg pain    | Spondylosis                                               | None                                                                                   |
| 14. 73-year-old woman | Low back pain, bilateral leg pain    | Spondylosis, scoliosis                                  | None                                                                                   |
| 15. 68-year-old woman | Back pain, low back pain, bilateral leg pain | Spondylosis, ossification of yellow ligament (after surgery) | None                                                                                   |

(Continued)
Table 1 (Continued).

| Case         | Chief complaint                                | Structural disorder | Psychiatric diagnosis                                                                 |
|--------------|------------------------------------------------|---------------------|----------------------------------------------------------------------------------------|
| 16. 34-year-old man | Low back pain                                | None                | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 17. 79-year-old man  | Posterior cervical pain, left-arm pain, low back pain, bilateral leg pain | Spondylosis         | Pervasive developmental disorder, attention deficit hyperactivity disorder, compulsive personality disorder, somatoform disorders |
| 18. 42-year-old woman | Bilateral sole numbness, cool feeling, and burning sensation | None                | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 19. 42-year-old woman | Low back pain                                | None                | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 20. 53-year-old woman | Low back pain                                | Spondylosis         | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 21. 40-year-old man  | Low back pain, bilateral sacroiliac joint pain | None                | None                                                                                    |
| 22. 38-year-old woman | Low back pain                                | None                | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |
| 23. 38-year-old man  | Low back pain, left lower and lateral leg pain and numbness | None                | Pervasive developmental disorder, attention deficit hyperactivity disorder, somatoform disorders |

Notes: Eight patients (four males, four females) aged 20–69 years (mean age 42.9 years) were also evaluated at 3 and 6 months after the program (marked by bold font in Table 1). Statistically significant improvement was seen in pain and associated factors (Figure 2) and physical functions (Figure 3).
elements (Table 2). PCS total scores ranged from 32.7–24.5 points. The cut-off value for the PCS total score is 30 points. Among our participants, the PCS total score was >30 points before program and <30 points immediately after program, which showed more improvement in PCS total score than the minimal detectable change. PDAS scores ranged from 29.4–18.8 points. The cut-off value for the PDAS is 10 points; the PDAS was >10 points before and immediately after program, which suggested the PDAS was not improved to normal condition in the program. However, participants’ PDAS scores still showed significant improvement immediately after the program. HADS scores of 8–10 points indicate suspicion of anxiety/depression and scores >11 points are considered to indicate the presence of anxiety/depression. In this study, participants’ HADS scores improved more than the minimal detectable change. The minimal clinically important difference for the PSEQ was more than 9%, which suggested participants’ PSEQ scores improved more than the minimal detectable change. Finally, the BPI and EQ-5D have no clear standards; however, our results suggested pain severity (BPI) and quality of life (EQ-5D) were improved immediately after the program.

### Multidisciplinary pain treatment for chronic musculoskeletal pain

The pathology of chronic musculoskeletal pain has intricately related biological and psychosocial components. As the duration of musculoskeletal pain is lengthened, the casual relationship between pain and pain-related factors becomes obscure. Furthermore, pain behavior and social factors related to pain may become more involved. The IASP indicates that changes in physical and psychosocial

| Table 2 Changes in pain and associated factors (n=23) |
|-----------------------------------------------|
| **Before program** | **Immediately after program** | **p-value** | **r** |
| BPI | 24.5±10.15 | 20.0±8.90 | 0.0001 | 0.73 |
| PCS (rumination) | 15.1±5.19 | 12.7±5.21 | 0.004 | 0.58 |
| PCS (magnification) | 6.3±4.09 | 4.4±3.53 | 0.001 | 0.67 |
| PCS (helplessness) | 11.5±5.58 | 7.5±5.67 | 0.001 | 0.68 |
| PCS (total) | 32.7±14.05 | 24.5±13.68 | 0.001 | 0.71 |
| PDAS | 29.4±12.29 | 18.8±12.56 | 0.001 | 0.64 |
| HADS (anxiety) | 8.7±4.86 | 6.2±4.02 | 0.001 | 0.66 |
| HADS (depression) | 9.8±4.81 | 6.5±4.35 | 0.001 | 0.65 |
| PSEQ | 22.1±11.90 | 33.2±13.93 | 0.001 | 0.69 |
| EQ-SD | 0.53±0.15 | 0.64±0.1745 | 0.009 | 0.53 |

*Notes: p-values less than 0.05 were considered statistically significant. r shows the effect size between before and immediately after the program. The cut-off value for the PCS total score is 30 points. The PCS total score was >30 points before the program and <30 points immediately after program. The result suggested the PCS total score improved more than the minimal detectable change. The PDAS cut-off value is 10 points. The PDAS was >10 points before and immediately after program. The result suggested PDAS scores might not have improved to normal condition; however, the PDAS showed significant improvement immediately after the program. For the HADS, 8–10 points indicates suspicion of anxiety/depression and >11 points indicates anxiety/depression. The result suggested HADS scores improved more than the minimal detectable change. The minimal clinically important difference for the PSEQ is >9%. The result suggested PSEQ scores improved more than the minimal detectable change. The BPI and EQ-5D have no clear standards; however, these results suggested pain severity (BPI) and quality of life (EQ-5D) improved immediately after the program.

**Abbreviations:** SD, standard deviation; BPI, Brief Pain Inventory; PCS, Pain Catastrophizing Scale; PDAS, Pain Disability Assessment Scale; HADS, Hospital Anxiety and Depression Scale; PSEQ, Pain Self-Efficacy Questionnaire; EQ-5D, EuroQol Five Dimensions Questionnaire.

| Table 3 Changes in physical functions (n=23) |
|-----------------------------------------------|
| **Before program** | **Immediately after program** | **p-value** | **r** |
| Finger floor distance (flexibility) | 15.8±16.30 | 8.1±13.73 | 0.005 | 0.59 |
| 30-second sit-to-stand test (muscle endurance) | 14.0±7.46 | 18.9±8.87 | 0.0001 | 0.77 |
| 2-step test (walking ability) | 1.3±0.27 | 1.4±0.23 | 0.01 | 0.53 |
| 6 mins walking (physical fitness) | 424.9±127.64 | 477.0±133.96 | 0.005 | 0.59 |

*Notes: p-values less than 0.05 were considered statistically significant. r shows the effect size between before and immediately after the program.

**Abbreviation:** SD, standard deviation.
factors from chronic musculoskeletal pain include muscle weakness associated with reduction in daily activity, somniphaty, malnutrition, drug dependence, dependence on family, isolation from family or society, decline in job performance, and economic burden. These factors may prolong pain duration and decrease the effects of therapy. Therefore, usual treatment delivered by a single department may not be sufficient for people with chronic pain.

The IASP recommends multidisciplinary pain treatment after distinguishing acute pain from chronic pain and evaluating various facets of a patient’s pain. A multidisciplinary approach has been applied in Europe and the USA since Bonica highlighted the necessity of such an approach in the 1950s. Multidisciplinary approaches are delivered by a team of professionals and may include education for patients, patient-centered cognitive behavioral therapy, exercise therapy, and pain-coping training. Professional groups involved include doctors from various medical departments (eg orthopedic surgeons, psychiatrists, anesthesiologists, physicians, neurologists, dentists), nurses, physical therapists, occupational therapists, clinical psychologists, pharmacists, nutritionists, and social workers. Each professional considers the patient’s pathology after open discussions in conference, and contributes to a comprehensive plan that focuses on the patient’s life, including sleep and nutritional support.

**Characteristics of our inpatient multidisciplinary pain management program**

An innovation of our multidisciplinary pain management program was the inpatient component.3

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Figure 2 Changes in pain-associated factors. Statistically significant improvement was seen in: PCS (rumination) at 3 (p=0.03) and 6 (p=0.003) months after the program; PCS (helplessness) at 3 (p=0.012) and 6 (p=0.04) months; PDAS at 6 months (p=0.04); HADS (anxiety) at 6 months (p=0.016); and PSEQ immediately after (p=0.004), at 3 months (p=0.012), and at 6 months (p=0.03). *p<0.05.

**Abbreviations:** BPI, Brief Pain Inventory; PCS, Pain Catastrophizing Scale; PDAS, Pain Disability Assessment Scale; HADS, Hospital Anxiety and Depression Scale; PSEQ, Pain Self-Efficacy Questionnaire; EQ-5D, EuroQol Five Dimensions Questionnaire.
Cognitive behavioral therapy

The recognition or perception of pain is considered as important as pain sensation. The essence of cognitive behavioral therapy in pain control is teaching perspectives and methods that help to improve pain self-management (particularly for problems related to external accidental stress), and develop new behaviors via operant conditioning that will improve social skills learning. In our facility, we classified patients into three subgroups (DYS-type, ID-type, and AC-type) using the WHYMPI. The subgroups were assigned during initial pain psychology interviews, and pain self-management was taught depending on subgroup assignment (type of pain). DYS-type patients can be effectively managed through operant conditioning, and patients in this group were rewarded if they attained the goals we set. For patients categorized as ID-type, it was important to help their families understand the concept of chronic musculoskeletal pain and how to change their way of attending to the patient. As patients in the AC-type subgroup tend to have difficulty adapting to some situations, we prioritized teaching this group about pacing during activities through exercise. We supervised all treatment for patients and family members. Our program aimed to help patients escape from passivity, dependency, and the tendency to be controlled by their external environment, so that they were able to manage their pain by themselves. The treatment goal was improving patients’ QOL despite their chronic pain.

Exercise therapy

Exercise and stretching with pacing improves flexibility, muscle endurance, walking ability, and physical fitness. It also activates dopaminergic/serotonergic and norepinephrinergic

**Figure 3** Changes in physical ability. Statistically significant improvement was seen in finger-floor distance (FFD) (flexibility) at 3 (\(p=0.04\)) and 6 (\(p=0.04\)) months after the program, and the 30-second sit-to-stand test (muscle endurance) at 6 months (\(p=0.012\)). \(^*p<0.05.\)
neurotransmission in the descending spinal pain-inhibition pathways. Changes in whole locomotorium, such as muscle atrophy, muscle degeneration, shortening of connective tissue, osteoporosis, and psychosocial factors are associated with increased chronic musculoskeletal pain. In turn, chronic musculoskeletal pain leads to disuse of the locomotorium to avoid pain. Some cases fall into kinesiophobia, where patients tend not to move their body because of the fear of pain worsening. Pain typically increases when moving in disuse conditions, meaning the fear of moving may be reinforced. Moreover, patients may develop avoidance reactions and excess wariness. We can explain these states using a fear-avoidance model. The goal of exercise therapy in our program was to allow patients to escape from this type of negative feedback loop, improve their flexibility and physical strength, promote healthy living and improved QOL, and support the recovery of physical ability. Exercise therapy can be effective on its own; however, the effects are greater if it is combined with cognitive behavioral therapy.

Outpatient management after inpatient multidisciplinary/interdisciplinary pain management programs for chronic musculoskeletal pain in other facilities in worldwide

Several studies have reported on follow-up after inpatient multidisciplinary or interdisciplinary pain management programs in other countries. We compared the outcomes from our program with those from other facilities. Table 4 shows the follow-up outcomes after inpatient programs in other countries. The Pain Treatment Center at the Red Cross Hospital in Kassel, Germany offers a 1-week inpatient multidisciplinary self-management program for patients with fibromyalgia. That program involves a team of rheumatologists, physiotherapists, occupational therapists, nurses, dieticians, and social workers. Patients are followed for 3 weeks after the program. The program had no effect on psychological distress, functional and symptomatic consequences, and self-efficacy for patients with fibromyalgia; however, there was a small short-term effect on skills and behavior important for managing and participating in healthcare (Table 4). The Pain Centre of the University Medical Centre Groningen in the Netherlands offers 4-week inpatient multidisciplinary cognitive behavioral treatment for patients with chronic pain. Treatment is delivered by a team including neurologists, physicians, physical therapists, and psychologists. Patients are followed for 1 year (Table 4). The Department of Rheumatology and Physical Medicine, University Hospital of Zurich, Switzerland offers a 4-week inpatient interdisciplinary pain program for patients with chronic pain, which involves a team of physicians, physical therapists, psychiatrists, and psychologists. That program showed a strong association between change in pain severity and physical functioning and change in baseline affective health and coping levels during the first outpatient management period. The program follows patients until 5-months after discharge (Table 4). A 4-week inpatient cognitive behavioral pain program for patients with chronic pain is offered at INPUT Pain Management, St Thomas’ Hospital, London (UK). That program involves a team psychologists, physiotherapists, nurses, occupational therapists, and anesthetists. Patients are followed for 6 months. Although that program has similarities to ours, it differs in length (4 weeks) and does not include nutritionists (nutritional education), clinical psychologists, or pharmacists. The program outcomes (QOL, pain intensity, psychological function, and physical function) were also similar to our results immediately after program, and were well maintained at the 6-month follow-up (Table 4). Finally, an inpatient program (at least 12 days of treatment) is offered for patients with chronic spinal back pain at the Loreley Hospital of Conservative Orthopaedics and Center for...
Table 4 Outcomes at follow-up after inpatient programs in other countries

| Program Location                          | Outcome | Outcome immediately after program                                                                 | Outcome at follow-up          | Follow-up rate (%) |
|------------------------------------------|---------|---------------------------------------------------------------------------------------------------|------------------------------|--------------------|
| Pain Treatment Center, Red Cross Hospital (Kassel, Germany) | NRS     | Significant post treatment improvement in pain intensity, disability, pain acceptance, catastrophizing, and depression. | 3 months: maintained well | Not listed          |
|                                          | MPSS    |                                                                                                  |                              |                    |
|                                          | PDI     |                                                                                                  |                              |                    |
|                                          | ADS-K   |                                                                                                  |                              |                    |
|                                          | CPAQ    |                                                                                                  |                              |                    |
|                                          | PCS-H   |                                                                                                  |                              |                    |
| HRC Bethesda Children’s Hospital (Budapest, Hungary) | VAS     | Short- and long-term improvements in clinical outcomes and resource use among pediatric patients with severe chronic pain. | 3 months: maintained well | 62.5% 5/8          |
|                                          | PPDI    |                                                                                                  | 2 years: maintained well      | 75% 6/8            |
|                                          | CDI     |                                                                                                  |                              |                    |
|                                          | CBCQ    |                                                                                                  |                              |                    |
|                                          | WCI     |                                                                                                  |                              |                    |
|                                          | LTS     |                                                                                                  |                              |                    |
| James A. Haley Veterans’ Hospital (Tampa, FL, USA) | NRS     | Significant improvements in pain intensity, pain-related fear, mobility, pain-related negative affect, vitality, implausible symptoms, and sleep. | 3 months: maintained well | Not listed          |
|                                          | POQ-VA  |                                                                                                  |                              |                    |
|                                          | SIS     |                                                                                                  |                              |                    |
|                                          | CSQ     |                                                                                                  |                              |                    |
|                                          | SPQ     |                                                                                                  |                              |                    |
| Lillehammer Hospital for Rheumatic Diseases (Lillehammer, Norway) | CHQ20   | Self-management program had no effect on psychological distress, functional and symptomatic consequences and self-efficacy, but a small short-term effect on skills/behavior important for managing and participating in healthcare. | 3 weeks: maintained well | 80.3% 118/147      |
|                                          | EC-17   |                                                                                                  |                              |                    |
|                                          | ASES    |                                                                                                  |                              |                    |
|                                          | FIQ     |                                                                                                  |                              |                    |
|                                          | VAS     |                                                                                                  |                              |                    |
| Pain Centre of the University Medical Centre Groningen (Groningen, the Netherlands) | VAS     | Significant improvements in pain, fatigue, walking distance, muscle strength, anxiety, depression, somatization, negative self-efficacy, and catastrophizing in the intervention period. | 6 months: maintained well | 76% 26/32          |
|                                          | 6 min walking test |                                                                                                 | 12 months: maintained well |                    |
|                                          | SCL-90  |                                                                                                  |                              |                    |
|                                          | RAND-36 |                                                                                                  |                              |                    |
|                                          | PCL     |                                                                                                  |                              |                    |
|                                          | TSK     |                                                                                                  |                              |                    |

(Continued)
Table 4 (Continued).

| Department of Rheumatology and Physical Medicine, University Hospital of Zurich (Zurich, Switzerland) | Outcome |
|---|---|
| | Outcome immediately after program |
| | Outcome at follow-up |
| | Follow-up rate (%) |
| MPI | Pain intensity (MPI) and depression (HADS) worsened slightly during 5 months. Long-term significant improvements for the other outcomes. | 5 months: Pain intensity and depression worsened slightly, but the other outcomes remained almost stable (follow-up at other facilities). | 71.8% |
| HADS | | | 308/429 |
| CSQ | | | |

| INPUT Pain Management, St Thomas’ Hospital (London, UK) | Outcome measures: QOL, physical performance, pain intensity, and distress, depression severity and confidence; significant improvements immediately after treatment. | 6 months: maintained well | 71.1% |
| Dysfunction score (SIP) (%) | | | 118/166 |
| Pain intensity score | | | |
| Pain distress score | | | |
| Depression score (BDI) | | | |
| Self-efficacy score (PSEQ) | | | |
| Length of 10 min walk (meters) | | | |
| Time of 20 meters walk (seconds) | | | |
| Number of stairs in 2 mins | | | |
| Number of sit-ups to tolerance | | | |

| Loreley Hospital of Conservative Orthopaedics and Center for Musculoskeletal Medicine, (Loreley, Germany) | Significant improvements in pain intensity, QOL, and function. | 6 months: maintained well | 53.4% |
| NRS | | | 133/249 |
| CPI | 1 year: maintained well | 43.8% | 109/249 |
| ODI | | | | |

Abbreviations: NRS, Numerical Rating Scale; MPSS, Mainz Pain Staging System; PDI, Pain Disability Index; ADS-K, Allgemeine Depressions Skala-Kurz version; CPAQ, Chronic Pain Acceptance Questionnaire; PCS, Pain Catastrophizing Scale; PCS-H, Pain Catastrophizing Scale-helplessness; VAS, Visual Analog Scale; PPDI, Pediatric Pain Disability Inventory; CDI, Children Depression Inventory; CBCQ, Child Behavior Checklist Questionnaire; WCL, the Ways of Coping Inventory; LTS, Likert-type scale; POQ-VA, the Pain Outcome Questionnaire-Department of Veterans Affairs; SIS, Symptom Implausibility Scale; CSQ, Coping Strategy Questionnaire; SP, Sleep Problems Questionnaire; CHQ20, General Health Questionnaire 20, EC-17, Effective Musculoskeletal Consumer Scale; ASES, Arthritis Self-Efficacy Scale; FIQ, Fibromyalgia Impact Questionnaire; SCL-90, Symptom Check List-90; PCL, Pain Cognition List; TSK, Tampa Scale for Kinesiophobia; MPI, Multidimensional Pain Inventory; SF-36, Short-Form 36 Questionnaire (SF-36); HADS, Hospital Anxiety and Depression Scale; SIP, Sickness Impact Profile; BDI, Beck Depression Inventory; PSEQ, Pain Self-Efficacy Questionnaire; CPI, Von Korff Questionnaire; ODI, Oswestry Low Back Pain Disability Index; QOL, quality of life.
Our inpatient pain management program may be able to improve patients’ ability to cope with their chronic musculoskeletal pain, as well as their QOL and physical abilities. Our program is currently being expanded to better assist patients with chronic musculoskeletal pain, from immediately after the program to 6-month follow-up.

Conclusion

Our inpatient pain management program may be able to improve patients’ ability to cope with their chronic musculoskeletal pain, as well as their QOL and physical abilities. Our program is currently being expanded to better assist patients with chronic musculoskeletal pain, from immediately after the program to 6-month follow-up.

Limitations

Controversial points of our program

In addition to our program’s advantages, it also had a few weak or controversial points. First, although our program is open to all, we need to carefully consider each patient’s rehabilitation. Second, because it may be difficult to sufficiently change how patients think and behave in a short period, patients may need to be followed-up after discharge. Finally, a large number of medical personnel are involved in our inpatient program, and their incomes may not be adequate given the insurance system for chronic pain in Japan. These points need to be addressed before our inpatient program can be generalized.

Limitations of this study

The present study had certain important limitations. First, there was no control group. We compared the treatment outcomes of our program with similar programs in other countries; however, further studies including control groups are required. Second, our study population was small, and larger populations are needed in subsequent studies. Third, the follow-up period was relatively short, and more studies with long-term follow-up are needed to evaluate long-term outcomes.

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Author contributions

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data, took part in drafting the article or revising it critically for important intellectual content, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest in this work.

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