Weak outcome predictors of multimodal rehabilitation at one-year follow-up in patients with chronic pain—a practice based evidence study from two SQRP centres

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Abstract

Background: For patients with chronic pain, the heterogeneity of clinical presentations makes it difficult to identify patients who would benefit from multimodal rehabilitation programs (MMRP). Yet, there is limited knowledge regarding the predictors of MMRP’s outcomes. This study identifies predictors of outcome of MMRPs at a 12-month follow-up (FU-12) based on data from the Swedish Quality Registry for Pain Rehabilitation (SQRP).

Methods: Patients with chronic pain from two clinical departments in Sweden completed the SQRP questionnaires—background, pain characteristics, psychological symptoms, function, activity/participation, health and quality of life—on three occasions: 1) during their first visit; 2) immediately after the completion of their MMRP; and 3) 12 months after completing the MMRP (n = 227). During the FU-12, the patients also retrospectively reported their global impressions of any changes in their perception of pain and their ability to handle their life situation in general.

Results: Significant improvements were found for pain, psychological symptoms, activity/participation, health, and quality of life aspects with low/medium strong effects. A general pattern was observed from the analyses of the changes from baseline to FU-12; the largest improvements in outcomes were significantly associated with poor situations according to their respective baseline scores. Although significant regressors of the investigated outcomes were found, the significant predictors were weak and explained a minor part of the variation in outcomes (15–25%). At the FU-12, 53.6% of the patients reported that their pain had decreased and 80.1% reported that their life situation in general had improved. These improvements were associated with high education, low pain intensity, high health level, and work importance (only pain perception). The explained variations were low (9–11%).

Conclusions: Representing patients in real-world clinical settings, this study confirmed systematic reviews that outcomes of MMRP are associated with broad positive effects. A mix of background and baseline variables influenced the outcomes investigated, but the explained variations in outcomes were low. There is still a need to develop standardized and relatively simple outcomes that can be used to evaluate MMRP in trials, in clinical evaluations at group level, and for individual patients.

Keywords: Multidisciplinary, Multimodal, Outcome, Pain, Rehabilitation

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Background

Approximately 20% of the adult population lives with at least moderate chronic pain [1]. Patients with chronic pain describe wide consequences such as intense and disturbing pain, psychological distress and insomnia, reduced work ability and sick-leave, ill-health, and low quality of life [2–6]. According to Years Lived with Disability (YLDs), a measure of non-fatal health outcomes, pain conditions caused 21% of all YLDs globally ahead of 287 other conditions [7]. The five leading conditions of YLDs in Sweden were low back pain, major depressive disorder, falls, neck pain, and other musculoskeletal disorders.

Acute and chronic pain are influenced by and interact with physical, psychological, social, and contextual factors [8–10], thus a bio-psycho-social framework should be considered in clinical practice [11, 12]. Multimodal (i.e., multidisciplinary) rehabilitation programs (MMRPs) are complex interventions based on such a framework and usually continue over several weeks with both general and patient specific goals. Common components of MMRPs are education, supervised physical activity, training in simulated environments, cognitive behavioural therapy (CBT), and work-related efforts. These components can act independently and interdependently resulting in a combination of effects explained by known and unknown mechanisms and these effects are intended to be greater than the sum of its components [13, 14]. The components are coordinated and delivered by a team of different professionals, but an important prerequisite is that the patient is an active participant [15–18].

MMRPs represent important evidence-based progress in treatment for patients with chronic pain [16–21]. Although there is some evidence that MMRPs are effective, the effect sizes are generally small to moderate. Randomized controlled trials (RCT) and systematic reviews (SR) may reflect artificial situations with regard to unrepresentative selection of patients, risk for researcher allegiance, and problems with practical applications of the interventions investigated [22, 23]. Thus, a complementary necessary step is to investigate whether the evidence reported from RCTs and SRs also holds for a consecutive non-selected flow of patients in real-world practice settings. This research methodology—practice based evidence (PBE)—is based on prospective observational cohort study designs and has also been applied in rehabilitation research [24].

Patients with chronic pain exhibit heterogeneous clinical presentations and the question arises if certain patient characteristics are indicative of outcomes after MMRP. Answering these questions are important as MMRPs are time consuming and expensive even when most of the activities for the patients are group based. Little is known about outcome predictors of MMRPs [18, 25–30]. It is likely that a combination of predictor variables can be identified [30]. Some of the available predictive studies are based on RCTs, but these RCTs have a small sample of patients and may have limited relevance to MMRP in clinical practice [31–33]. Prediction studies based on clinical samples are needed to reliably identify who actually benefits from inclusion and participation in MMRPs.

This PBE study identifies predictors of outcome of MMRPs for patients with chronic pain based on data from the Swedish Quality Registry for Pain Rehabilitation (SQRP) (www.ucr.uu.se/nrs/) for two multidisciplinary pain centres in Sweden. Six outcome domains were identified (pain intensity, physical functioning, emotional functioning, coping, health and ratings of global improvements) mainly according to the recommendations of the IMMPACT-group [12, 34].

Methods

Subjects

This study collected data using the SQRP on 464 patients between 18 and 65 years old who had chronic mainly musculoskeletal pain. These patients were referred to the Pain and Rehabilitation Centre at University Hospital of Linköping or the Pain Rehabilitation Clinic at University Hospital of Umeå between 2008 and 2012. In total, 464 patients with chronic pain were referred to these two facilities to participate in a MMRP. The present study is primarily based on patients who answered the SQRP both at baseline (before the MMRP) and at the 12-month follow-up (n = 227).

MMRP at the two sites

At the Umeå site, the referred patients were assessed by teams consisting of a specialist physician in rehabilitation medicine or by specialists in training under supervision of a senior colleague, a physiotherapist, a social worker, and a psychologist and occupational therapist if needed. If the team considered that the patient needed MMRP and fulfilled the inclusion criteria (disabling chronic pain, on sick leave or experiencing major interference in daily life, no further investigations needed) they were accepted to participate in the program. The MMRP was conducted in groups of six to nine participants and was based on CBT principles and included physiotherapy, ergonomics, training in coping strategies, and education in pain management. The patients were encouraged to take an active part in goal setting. At the Linköping site, medical assessments were performed by senior physicians, primarily from specialists in rehabilitation medicine or similar specialities, or by specialists in training under the supervision of a senior colleague. The majority of the participants were also assessed by a psychologist, an occupational therapist, and a physiotherapist. The Linköping program followed the same main principles as the Umeå program—group treatment
from a CBT perspective, group physiotherapy, interventions targeting improved ergonomics, and occupational therapy. At both sites the MMRPs lasted between 6 and 8 weeks for at least 20 h per week of group-based activities. In addition, lectures in basic pain science and pain management were offered for both patients (both sites) as well as for relatives, friends, and colleagues (only Linköping). Both programs included work related advice and support, and individually tailored sessions with team members were available. Further individual sessions might also be required for a few weeks following the program.

The following inclusion criteria for the MMRP were used: (i) disabling chronic pain (on sick leave or experiencing major interference in daily life due to chronic pain); (ii) age between 18 and 65 years; (iii) no further medical investigations needed; (iv) written consent to participate and attend to the MMRP; (v) and agreement not to participate in other parallel treatments. The following exclusion criteria were used: (i) ongoing major somatic or psychiatric disease; (ii) a history of significant substance abuse; and (iii) state of acute crisis. General exclusion criteria included severe psychiatric morbidity, abuse of alcohol and/or drugs, diseases that did not allow physical exercise, and specific pain conditions with other treatments options available (red flags).

Methods

The SQRP is recognized as a national registry by the Swedish Association of Local Authorities and Regions. The SQRP is mainly based on questionnaires. The patients completed the SQRP questionnaires on three occasions: 1) at the assessment during the first visit to the clinical department (pre-MMRP or baseline); 2) immediately after MMRP (post-MMRP); and 3) at the 12-month follow-up (FU-12). Demographic data are only collected before MMRP. In the present study, the predictions concerned the FU-12.

The main purpose of the SQRP is to present the results of MMRP at a group level to the participating clinical departments. Based on these data, health care providers and researchers can develop a process that will encourage continued improvement of rehabilitation programs. In a review of the overall quality of approximately 60 nationwide registries, the Boston Consulting Group considered the SQRP as one of the top ten national registries in Sweden [35].

The SQRP includes descriptive variables of the patients’ background, pain characteristics, other symptoms such as depression and anxiety, function/activity/participation, and quality of life. Generally validated Swedish language versions of the instruments were used.

Data from the SQRP

Listed below are the variables and instruments of the SQRP that this study used. These are also summarized in Additional file 1: Table S1.

Background data
- Age (years)
- Gender
- Education (elementary school, upper secondary school, University or college or other). This variable was dichotomized into University vs the other alternatives.
- Country of birth (Sweden, Another Nordic country, Europe except Nordic country, or Country outside Europe). This variable was dichotomized into Nordic countries vs. outside Nordic countries.
- Currently working/studying (Yes or No) denoted Work/study-now.
- Three items concerned the attitude about the future. Those working/studying full time were instructed to skip the first two items below; subjects who were seeking a job were instructed to imagine being employed when answering the following items:
  - What do you think it will be like to return to work/studies or to extend working hours if you are not working full time? (Likert scale: 1 (very easy) to 5 (very difficult). This variable was denoted as Own prognosis-RTW.
  - How confident are you about your chances to be fully restored? (Likert scale: 1 (totally convinced) to 5 (not at all convinced). This variable is denoted as Chances-restored.
- What significance does the work have for you in addition to the importance of income? (Five alternatives: 1) Very important, 2) Important, 3) Partially important, 4) Hardly no importance, or 5) No importance). This variable was denoted as Work-importance.

Characteristics of pain
- How many times have you visited a physician for your pain complaints the previous year? 0–1 times; 2–3 times; or ≥4 times). This variable was dichotomized into ≥4 times vs. the other alternatives and is denoted as Dr-visits.
- When did you first experience the pain you are currently troubled by? (days). This variable was denoted as Pain-duration.
- If persistent pain exists, how long? (days). This variable was denoted as Pain-duration-persistent.
- Average pain intensity the last week (a numeric rating scale 0–10 with numbers for guidance and the end points also had verbal descriptions...
Objects (0 = no pain and 10 = worst possible pain). This is denoted as NRS-7days.

- Using 36 predefined anatomical areas (18 on the front and 18 on the back of the body), the subjects marked where they experienced pain: 1) head/face, 2) neck, 3) shoulder, 4) upper arm, 5) elbow, 6) forearm, 7) hand, 8) anterior aspect of chest, 9) lateral aspect of chest, 10) belly, 11) sexual organs, 12) upper back, 13) low back, 14) hip/gluteal area, 15) thigh, 16) knee, 17) shank, and 18) foot. The number of areas associated with pain were counted (between 0 and 36) and this variable was labelled the Pain Region Index (PRI).

**Multidimensional Pain Inventory (MPI)**

The West Haven-Yale Multidimensional Pain Inventory—(WHY) MPI—is a 61-item self-report questionnaire that measures psychosocial, cognitive, and behavioural effects of chronic pain [36, 37]. Part 1 consists of five scales: Pain severity (MPI-Pain-severity); Interference—pain-related interference in everyday life (MPI-PainInterfer); Perceived Life Control (MPI-LifeCon); Affective Distress (MPI-Distress); and Social Support—perceived support from a spouse or significant others (MPI-SocSupp). Part 2 was not used in this study. Part 3 measures to what extent the patients engaged in various activities using four scales. These scales can be combined into a composite scale—the General Activity Index (MPI-GAI)—which was used in the present study [38].

**Hospital Anxiety and Depression Scale (HADS)**

The HADS is a self-assessment questionnaire that measures anxiety and depression using 14 items [39]. HADS comprises seven items in each of the depression and anxiety scales (HAD-D—depression and HAD-A—anxiety). The subscale scores range between 0 and 21, with the lower score indicating the least depression and anxiety possible. HADS is frequently used both in clinical practice and in research and has good psychometric characteristics [39, 40]. It is also validated in its Swedish translation [41].

**The Chronic Pain Acceptance Questionnaire (CPAQ)**

CPAQ measures acceptance behaviours and attitudes towards pain. The process of acceptance refers to the concept of coping with pain; this process changes the focus from controlling pain and related psychological experiences to a wider focus that includes being open to pain without trying to alter the experience. Acceptance includes flexible engagement in goal-directed activities, which include pain as a part of the engagement, without struggling to control pain. CPAQ is a 20-item scale with two subscales: activity engagement (score range: 0–66; denoted CPAQ-AE) and pain willingness (score range: 0–54; denoted CPAQ-PW) [42]. All items are rated on a scale from 0 (never true) to 6 (always true). The CPAQ has shown to be reliable and valid both in the English and Swedish versions [42, 43].

**The Tampa Scale for Kinesiophobia (Tampa)**

Tampa measures fear of movement [44]. The items are rated on a four-point Likert scale from “strongly disagree” to “strongly agree”. The total score has a range from 17 to 68; scores higher than 36 for women and higher than 38 for men indicate high pain-related fear [45]. Tampa has shown to be a reliable assessment tool in chronic pain populations [44, 46]. A study including Dutch, Canadian, and Swedish samples with several different pain types demonstrated that the factor structure was stable across pain diagnoses and nationalities [45].

**Life Satisfaction Questionnaire (LISAT-11)**

LISAT-11 captures the patient’s estimations of satisfaction with life as a whole (LISAT-life) as well as satisfaction in ten specific domains [47]. Each item has six possible answers: 1 = very dissatisfying; 2 = dissatisfying; 3 = fairly dissatisfying; 4 = fairly satisfying; 5 = satisfying; and 6 = very satisfying. This study used LISAT-life together with the following specific items: satisfaction with vocation (LISAT-vocation); satisfaction with economy (LISAT-economy); satisfaction with activities of daily living (ADL) (LISAT-ADL); satisfaction with somatic health (LISAT-somhealth); and satisfaction with psychological health (LISAT-psychhealth).

**The Short Form Health Survey (SF36)**

SF36 intends to represent multidimensional health concepts and measurements of the full range of health states, including levels of well-being and personal evaluations of health [48]. The instrument has eight dimensions (reported using a standardized scale from 0 to 100) [48]. In the present study the scale concerning physical functioning (SF36-PF) was selected. Based on the eight scales, a physical summary component (SF36-PSC) and a mental (psychological) summary component (SF36-MSC) are calculated. Hence, this study used these two summary components together with SF36-PF.

**The European Quality of Life instrument (EQ-5D)**

EQ-5D captures a patient’s perceived state of health [49–51]. The first part of the instrument defines five dimensions: mobility, self-care, usual activities, pain/discomfort, and anxiety/depression. Using these five dimensions, the instrument calculates an index (EQ5D-Index). The EQ5D also measures self-estimation of today’s health according to a 100-point scale, a thermometer-like scale (EQ5D-VAS) with defined end points (high values indicate good health and low values indicate bad health).
Ratings of global improvement
- Retrospective rating of global improvement in life situation (Retro-pain)
- Retrospective rating of global improvement in life situation (Retro-life situation)

Statistics
All statistics were performed using the statistical package IBM SPSS Statistics (version 20.0) and SIMCA-P+ (version 13.0; Umetrics Inc., Umeå, Sweden); in all tests, a probability of <0.05 (two-tailed) was accepted as the criteria for significance. In the tables and text, the mean value ± one standard deviation (±1SD) of the investigated variables are given.

Effect sizes of the rehabilitation program were calculated using the template developed by Lakens [52]. In this paper, Cohen’s $d_w$ was seen as the most appropriate way to present the effects as this uses the average standard deviation of the two measurement points [53]. Roughly, one can interpret the Cohen’s effect sizes using the following criteria: $d = 0.2$ is small, $d = 0.5$ is medium, and $d = 0.8$ is large [54]. Bonferroni correction was used in the detailed analyses over time for the different outcomes. Results were deemed significant at the $p = 0.00083$ level.

Advanced multivariate analyses (MVDA) are essential in PBE studies. Classical statistical methods such as multiple linear regression (MLR) and logistic regression (LR) can quantify the level of relations of individual factors but disregard interrelationships among different factors and thereby ignore system-wide aspects (e.g., when a group of variables correlates with the investigated dependent outcome) [55]. Classical methods assume variable independence when interpreting results [56]. Eriksson and co-workers discussed the risks with considering one separate variable at a time (COST)[57]. They point out that the COST approach has serious drawbacks; the information in multivariate data often remains hidden and there is a risk for spurious results [57]. To handle these drawbacks, we used advanced MVDA (i.e., Principal Component Analysis, PCA) for the multivariate correlation analyses of all investigated variables and Partial Least Square Regression (PLS) for the multivariate regressions using SIMCA-P+. These techniques do not require normal distribution [58].

PCA was used to extract and display systematic variation in a data matrix. All variables were log transformed before the statistical analyses if necessary. A cross validation technique was used to identify nontrivial components (p). Variables loading on the same component p were correlated, and variables with high loadings but with
opposing signs were negatively correlated. Variables with high absolute loadings—i.e., 95% jack-knife uncertainty confidence interval non equal to zero—were considered significant. Hence, the most important variables were those with high absolute loadings. The obtained components are per definition not correlated and are arranged in decreasing order with respect to explained variation. \( R^2 \) describes the goodness of fit—the fraction of sum of squares of all the variables explained by a principal component [59]. \( Q^2 \) describes the goodness of prediction—the fraction of the total variation of the variables that can be predicted by a principal component using cross validation methods [59]. Outliers were identified using two methods: 1) score plots in combination with Hotelling’s \( T^2 \) and 2) distance to model in X-space. No extreme outliers were detected.

**PLS** was used for the multivariate regression analyses [59]. The VIP variable (variable influence on projection) indicates the relevance of each X-variable pooled over all dimensions and Y-variables—the group of variables that best explain Y. VIP \( \geq 1.0 \) was considered significant if VIP had 95% jack-knife uncertainty confidence interval non equal to zero. The PLS analyses were made in two steps. First, all potential regressor variables were entered. If this regression was significant, regressors with VIP > 0.80 were selected and used in the final PLS. Coefficients (PLS scaled and centred regression coefficients, Coeff) were used to note the direction of the relationship (positive or negative). SQRP uses predetermined rules when handling single missing items of a scale or a subscale; details about this is reported in Additional file 1: Table S6. SIMCA-P+, in contrast to traditional statistical packages e.g. SPSS, uses the NIPALS algorithm when compensating for missing data (i.e. in the present study the scales and subscales of a certain subject).

MLR or LR could have been an alternative, but they assume that the regressor (X) variables are independent. If multicollinearity (i.e., high correlations) occurs among the X-variables, the regression coefficients become unstable and their interpretability breaks down. MLR and LR also assumes that a high subject-to-variables ratio is present (e.g., > 5), an assumption not required for PLS. In fact, PLS can handle subject-to-variables ratios < 1. Moreover, PLS, in contrast to MLR and LR, can handle several Y-variables simultaneously.

### Results

#### Drop out analysis

When comparing those who did not and who did complete the questionnaire at FU-12, only one just barely significant difference was found—the pain intensity variable NRS-7days. Those who did not complete the questionnaire had a mean score of 7.02 (SD 1.06) in comparison to those who did (M = 6.61, SD = 1.89) (t(437) = 2.09, \( p = 0.04 \)). Hence, no statistically significant differences were found for demographic variables, psychological well-being, quality of life, acceptance, kinesiophobia, general health, or psychological functioning as measured by the SF-36.

### Background data and characteristics of pain

Background data is reported in Table 1. The majority of the patients were women (>80%). A significant proportion (18.5–36.8%) expressed pessimistic views concerning aspects of return to work (RTW) (Table 1).

The number of anatomical regions with pain (PRI) was 14.7 ± 7.9 out of 36 anatomical regions. Mean pain duration was 2550 ± 2609 days; corresponding figures for persistent pain were 1856 ± 2218 days. Health care seeking was prevalent; 58.9% had visited a physician four or more times during the previous year.

#### Effects of MMRP

For the selected outcomes and domains, significant improvements were found, except for LISAT-life (Table 2). The effect sizes were small to medium [60] (Table 2). The largest effect sizes were found for coping aspects followed by pain intensity aspects. The area of emotional functioning generally showed small effect sizes (Table 2). Detailed tables including all variables—also including registrations immediately post MMRP—are shown in Additional file 1: Tables S2–S5.

#### Predictions of changes in outcomes at 12-month follow-up

The difference (i.e., changes) between pre and FU-12 values for the selected outcomes were calculated and transformed if necessary so that an improvement had a positive value.

#### Regression of changes in pain intensity aspects

In the regression of changes in the two pain intensity aspects (\( R^2 = 0.25, Q^2 = 0.16 \)), the strongest regressors among the background and baseline variables were NRS-7days, MPI-Pain-severity, MPI-SocSupp, Work/study-now, and

| Table 1 Background data for the patients (n = 227) |
|-----------------------------------------------|
| Variable                                      |
| Age (years; mean(SD))                        | 38.1 (10.1) |
| Women                                        | 81.6%      |
| Highest Education (University)               | 24.1%      |
| Country of origin (Nordic countries)         | 91.8%      |
| Currently work or studies (Work/study-now)   | 28.9%      |
| Own prognosis-RTW (very difficult)           | 36.8%      |
| RTW-when (never)                             | 18.5%      |
| Chances-restored (not at all)                | 32.3%      |
| Work-importance (very important)             | 43.0%      |

Own prognosis-RTW = Prognosis of return to work/studies; RTW-when = When return to work/studies; Chances-restored = Chances to be fully restored; Work-importance = Significance of work (except economic)
LISAT-psychehealth (Table 3, model 1). Hence, improvements in pain intensity aspects were associated with the following background and baseline variables: high pain intensities, perceiving social support, working/studying, and satisfaction with psychological health. When excluding the pain intensity variables at baseline, the explained variation decreased as expected (Table 3, model 2). In addition, education (relative high level), low satisfaction with somatic health (LISAT11-somhealth), and relatively short persistent pain duration to some extent were then associated with improvements in pain intensity aspects.

Regression of changes in emotional functioning
In the regression of changes in three psychological variables (MPI-Distress, HAD-D, and HAD-A) ($R^2 = 0.14, Q^2 = 0.10$), the strongest regressors of the background and baseline variables were HAD-A, HAD-D, MPI-Distress, and SF36-MCS (Table 3, model 1). Hence, improvements in psychological status were associated with high psychological and mental distress at baseline. When the three psychological variables at baseline were omitted as regressors, the explained variation decreased (Table 3, model 2). In addition, low life control (MPI-LifeCon) and low satisfaction with life in general (LISAT-life) to some extent was associated with improvements in the three psychological variables.

Regression of changes in physical functioning
In the regression of changes in physical functioning (i.e., MPI-PainInterfer and SF36-PF; $R^2 = 0.19, Q^2 = 0.13$), the
Table 3  Regressions of changes in pain intensity, psychological distress, physical functioning and health (Model 1). In model 2 are excluded the baseline variable used for calculating the changes

|                       | Model 1 | Model 1 | Model 1 | Model 1 | Model 1 |
|-----------------------|---------|---------|---------|---------|---------|
|                       | Changes in pain intensity | Changes in psychological distress | Changes in physical functioning | Changes in health aspects |
| (n = 227)             | VIP     | Sign of Coeff | (n = 227) | VIP     | Sign of Coeff | (n = 227) | VIP     | Sign of Coeff | (n = 202) |
| NRS-7days             | 1.74 +  |           | 1.49 +  | 1.73 -  | EQSD-VAS  | 2.01 -    |
| MPI- pain-severity    | 1.27 +  |           | 1.24 +  | 1.36 -  | EQSD-index| 1.52 -    |
| MPI – social support  | 1.24 +  |           | 1.24 +  |          |          |           |
| Work/study-now        | 1.14 +  |           |          |          |          |           |
| LISAT – psychhealth   | 1.09 +  |           |          |          |          |           |
| R²                    | 0.25    |           | 0.15    | 0.17    | 0.20     |
| Q²                    | 0.16    |           | 0.12    | 0.13    | 0.12     |

|                       | Model 2 | Model 2 | Model 2 | Model 2 |
|-----------------------|---------|---------|---------|---------|
|                       | Changes in pain intensity | Changes in psychological distress | Changes in physical functioning | Changes in health aspects |
| (n = 227)             | VIP     | Sign of Coeff | (n = 227) | VIP     | Sign of Coeff | (n = 227) | VIP     | Sign of Coeff | (n = 197) |
| MPI- SocSupp          | 1.36 +  |           | 1.38 -  | 1.48 -  | Pain-duration-persistent | 1.21 -    |
| Work/study-now        | 1.30 +  |           | 1.10 -  | 1.38 -  | Age | 1.09 -    |
| LISAT-psychhealth     | 1.22 +  |           | 1.02 -  | 1.07 -  | Pain duration | 1.06 -    |
| Education             | 1.08 +  |           |          |          | RTW-when | 1.04 -    |
| LISAT-somhealth       | 1.01 -  |           |          |          | Own prognosis-RTW | 1.03 -    |
| Pain-duration-persistent | 1.01 - |           |          |          | CPAQ-AE | 1.02 -    |
| R²                    | 0.14    |           | 0.07    | 0.15    | 0.05     |
| Q²                    | 0.10    |           | 0.06    | 0.06    | 0.03     |

Background and baseline variables (cf. Additional file 1: Table S1) were used as regressors. The significant variables are shown and for each of these are reported VIP (VIP > 1.0 is significant) and the sign of the coefficient (+ or -). The two bottom rows of each model report $R^2$ and $Q^2$. 
three significant regressors were SF36-PF, RTW-when, and Own prognosis-RTW (Table 3, model 1). Hence, improvements in physical functioning were associated with the following circumstances at baseline: low physical functioning (SF36-PF), perceptions of relatively soon return to work (i.e., RTW-when), and relatively easy to return to work (i.e., Own prognosis—RTW). When excluding the two physical functioning variables at baseline as regressors, the explained variation decreased (Table 3, model 2). To some extent low physical health (SF36-PSC) at baseline was associated with improvements in physical functioning.

Regression of changes in coping aspects
It was not possible to significantly regress the changes in the four coping aspects (i.e., the 4 Y-variables: MPI-LifeCon, CPAQ-AE, CPAQ-PW, and Tampa) using the background and baseline variables as regressors.

Regression of changes in health aspects
In the significant regression of changes in the health variables (i.e., the two EQ5D variables and LISAT-life; \( R^2 = 0.19 \), \( Q^2 = 0.09 \)), the two significant regressors were EQ5D-VAS and EQ5D-index (Table 3, model 1). Hence, improvements in health aspects were associated with low reported health at baseline.

When omitting the health variables at baseline, the explained variation decreased (Table 3, model 2). Short pain durations and low age were then associated with improvements in health aspects.

Retrospective evaluations of pain and in ability to handle life situation in general
At FU-12, 53.6% retrospectively reported improvements for pain and 80.1% for general life situation (Table 2). For Retro-pain, the following background and baseline variables were multivariately important (\( R^2 = 0.11 \), \( Q^2 = 0.04 \)): Education (VIP = 1.84+), MPI-Pain severity (VIP = 1.42), EQ5D-index (VIP = 1.07+), and Work-importance (VIP = 1.07). Hence, improvements were associated with relatively high education, low pain intensity (MPI-pain-severity), high health (EQ-5D-index), and perceiving work as important for reasons other than economic. For Retro-life situation, the important variables at baseline (\( R^2 = 0.09 \), \( Q^2 = 0.05 \)) were MPI-Pain severity (VIP = 1.77), EQ5D-index (VIP = 1.72+), NRS-7days (VIP = 1.25), and education (VIP = 1.20+). Hence, improvements in life situation were associated with low pain intensities (NRS-7days and MPI-Pain-severity), high health (EQ5D-index), and relatively high education at baseline.

PCA of outcome variables
To understand to what extent the changes in outcomes of the different domains together with the two retrospective items above were independent, a PCA was made for the situation at FU-12. Only one significant component (\( R^2 = 0.34 \), \( Q^2 = 0.24 \)) was achieved according to the cross-validation rules in SIMCA-P+. The four most important variables — i.e., highest absolute loadings — were changes in MPI-PainInterfer (loading = 0.33), MPI-LifeCon (loading = 0.31), MPI-Pain severity (loading = 0.30), and MPI-Distress (loading = 0.29). Hence, these four variables showed the most prominent variability across subjects.

Regression of changes in the four MPI variables
When regressing the changes in the four most important MPI variables according to the PCA, the most important background/baseline variables were MPI-LifeCon (VIP = 2.05), MPI-Distress (VIP = 1.46), and Work-importance (VIP = 1.08) (\( R^2 = 0.16 \), \( Q^2 = 0.09 \), \( n = 227 \)). Hence, improvements in these four MPI variables were associated at baseline with low life control, high psychological distress, and perceiving work as important for reasons other than economic.

Discussion
Major results of the present study are listed below.
- The majority of subjects reported benefits of participating in MMRP according both to the repeated measurements (i.e., changes from baseline to FU-12) and to the retrospective evaluations.
- Pain intensity and emotional aspects showed the strongest improvements (i.e., effect sizes) according to the repeated measurements.
- The analyses of the repeated measurements showed that the largest improvements in outcomes were significantly associated with poor situations according to their respective baseline scores.
- Although significant regressors of the investigated domains of outcomes were found, the significant predictors were weak and only explained a small part of the variation in outcomes.

Effectiveness of MMRP
This PBE study confirms the generally positive results for MMRPs according to SRs [16–18], health economics studies, and 10-year follow-ups [19–21] (Table 2). Both the repeated measurements (i.e., baseline vs. FU-12) and the retrospective items (Table 2) showed improvements for this cohort of patients.

Retrospective evaluations at FU-12
The majority of patients retrospectively reported improvements according to both pain and in their ability to handle their life situation in general (Table 2). Fortunately and in agreement with other studies [61, 62], relatively small proportions (3–7%) of the patients reported worse situations at FU-12 (Table 2). Qualitative explorations have been
suggested as one way to develop interventions for patients who do not benefit from MMRP [63].

A mix of background and baseline factors were important regressors. The two retrospective items had several regressors in common. That is, low pain intensity, high education, and high health perception were related to improvements. Similar results have been reported in other studies using retrospective evaluations [33, 61]. In contrast, a SR of MMRP for fibromyalgia patients summarized that the global treatment effect was predicted by the MPI dysfunctional profile (i.e., high pain, disability, and high solicitous responses from significant others) and/or worse baseline status [28]. Hence, it is presently unclear if a relatively good or bad situation with respect to, for example, pain intensity is associated with improvements when evaluated retrospectively.

Changes in outcomes according to the repeated measurements

In the repeated measurements (i.e., changes from baseline to FU-12), the most prominent effects according to Cohen’s d were for emotional and pain aspects (Table 2). Effect sizes of 0.51–0.61 for the two pain intensity variables at FU-12 were found (Table 2) and in fact both variables showed increasing effect sizes over time (Additional file 1: Tables S2 and S4). These results were in contrast to some SRs, which reported no evidence for efficacy with respect to pain intensity [16, 17]. Many pain patients consider pain intensity as the most important aspect to improve in treatments. Several of the included RCTs in SRs of MMRP do not include pain intensity due to the fact that the interventions are not focussed on the pain itself but rather on its consequences [16, 17]. Some authors have suggested that it might be detrimental for the patient to focus on attempting to control, reduce, or cure pain, since this might shift emphasis away from the aspects that are important for the patient’s health and quality of life, such as daily functioning and emotional well-being [64, 65]. Although these lines of arguments appear reasonable, it may ethically problematic if both clinical practice and research ignore the reports of the patients regarding pain intensity. Moreover, patients with high pain intensities require greater reductions in pain intensity than patients with lower pain intensity levels in order to obtain clinically important improvements [66]. In addition to focusing on pain intensity, it might be wise to focus on the other outcomes emphasised by IMMPACT when communicating with patients. As an outcome of MMRP, pain intensity might be associated with problems when interpreting the results of MMRPs (e.g., a successful intervention leads to less fear of movement) as the patient may be more prone to participate in strenuous activities, which may lead to increased pain intensity in a more active patient. However, changes in pain intensity, psychological distress, and coping aspects were positively intercorrelated in this study, findings that do not support such an argument. It is presently unclear if a more prominent focus on pain intensity within the bio-psycho-social context of MMRP can increase the effect sizes or will actually diminish them.

A general pattern was observed where changes in outcomes were significantly predicted by their respective baseline scores (Table 3, model 1). For example, high pain intensity at baseline was associated with more prominent changes in pain intensity aspects. Such results have been observed in other studies and could be due to the fact that a more severe initial status gives more room for improvement [61]. Another related explanation could be regression to the mean [33]. Recently, Bonstra et al. briefly summarized the literature concerning regressors of repeated measures of outcomes and better outcomes were reported for patients with high scores on depression, high fear avoidance beliefs, high perceived stress, optimistic attitudes, low need to socialize, younger age, or high educational level [61]. Hence, the present results (Table 3, model 1) and a study of chronic widespread pain generally agree with that review [33]. In contrast to this literature, a SR concerning fibromyalgia reported that poor outcomes were associated with depression at baseline [28]. Differences across studies with respect to important predictors may be due to factors such as number of potential prognostic factors, cohort characteristics, chosen outcomes, and/or content of MMRP [18, 29, 67, 68].

For two of the domains of outcomes—pain intensity and physical functioning (Table 3, model 1)—additional background and baseline variables were significant. Hence, the regression of the pain intensity changes was also important for satisfaction with psychological health, working/studying, and relatively high social support at baseline. For changes in physical functioning, additional significant regressors were optimistic attitudes towards RTW (i.e., RTW-when and Own-prognosis–RTW).

Based on the regressions of the changes in the repeated measurements (Table 3, model 1), it seems important to choose patients with a relatively severe clinical picture in order to be able to achieve substantial improvements. It is reasonable to assume that this is relevant only up to a certain level of severity. However, even though it was possible to significantly regress most of the domains of outcomes, the explained variations ($R^2$) were low (15–25%). Similar results have been reported by other researchers [61]. Hence, other—not investigated—factors are important and the present results concerning significant background and baseline variables cannot be applied directly in clinical practice with any precision.

Is high or low pain intensity indicative of positive outcomes?

The importance of the level of pain intensity for positive outcome differed between the repeated measurements
and the retrospective evaluations. Improvements in pain intensity aspects according to the repeated measures were associated with high pain intensity at baseline (Table 3, model 1), while in the retrospective evaluations the improvements were associated with low pain intensity at baseline. There are several problems with retrospective items such as recall time, “telescoping”, desirability, and memory aspects [69–71]. When using retrospective items to evaluate changes in a treatment context, additional problems may exist (e.g., overly optimistic assessments) [72]. Hence, the results concerning predictors for the retrospective items may have been associated with bias. On the other hand, repeated evaluations using self-report questionnaires in intervention studies may be problematic [73]. The change that the patient undergoes because of the intervention may affect the interpretations of the questions when presented at follow-ups. MMRPs aim to influence the perception and coping of pain, psychological distress, and various behaviours. Hence, there may be a risk that the attitude towards the items of the questionnaire changed as a consequence of MMRP. This could mean that the follow-up tests cannot simply be compared with the test before MMRP. In order to deepen the knowledge concerning efficacy of MMRPs, objective registrations (e.g., sick-leave registrations, actigraphic recordings, and qualitative interviews [63]) can be important additional sources.

**Evaluations of outcomes of MMRP**

There is a need to develop clinically applicable and standardized ways to evaluate multiple outcomes of MMRPs in individual patients in trials and in PBE studies. This involves both the statistical methods (cf. Statistics) and strategies for handling multiple outcomes. This study used different approaches for evaluation of outcomes. The number of outcomes in MMRPs are generally high; one SR including 46 RCTs found in median nine outcomes and that these generally were not divided into primary and secondary outcomes [16]. In a recent SR, the outcomes were evaluated separately [74], which may be problematic if the outcomes are intercorrelated and in fact they often are as found in the present study. The present study used 14 outcomes divided into six domains that were mainly recommendations from IMMPACT [12, 28, 34]. Different definitions of a positive outcome of a MMRP trial have been presented such as the majority of outcomes had to be significantly better than for the control intervention [16, 17]. The authors of another SR predetermined primary and secondary outcomes and what was necessary to classify an intervention as positive before reviewing the RCTs [75].

The analysis of several outcomes may raise an issue of multiple comparisons and Bonferroni corrections are frequently applied [76] and recommended in pain trials [77]. However, this is a conservative approach when the number of tests increases [76, 78, 79], and this approach reduces the chances to detect real treatment effects. Furthermore, such corrections were designed for corrections of independent comparisons [78]. The latter is generally not present when evaluating the outcome of MMRPs. IMMPACT has presented hierarchical or “gatekeeping” procedures that do not require adjustment for multiplicity [77] but a natural hierarchy of the outcomes. Outcomes may be combined into a single composite outcome [80], but this may be problematic with respect to missing cases and when the components of the composite endpoint are measured on different scales such as non-commensurate outcomes [80].

Multivariate methods that are able to handle non-commensurate outcomes in one analysis have been presented [80]. The presently applied PLS regression can also handle non-commensurate outcomes in one analysis. We used a more comprehensive approach when regressing four outcomes simultaneously. These four outcomes were chosen based on the advanced multivariate correlation analysis (PCA). In agreement with the regressions of the repeated measures for different domains, we found that a relatively severe clinical presentation at baseline was associated with improvements.

**Limitations**

There are several limitations to consider. No control group was included and outcomes can be due both to the effects of MMRP and to other factors such as natural course. On the other hand, these patients on average had their pain for a considerable period (on average more than 2500 days), a period that reduces but does not eliminate the risk that effects are related to the natural course. Patients participating in MMRPs have often tried different unimodal treatments with little or no effects and thus represent a selection of patients. The present cohort of patients also represents a selection of patients with the most complicated clinical pictures as they were referred to a university hospital. Hence, the generalizability of the present results is limited and the results of MMRP in primary care may be different. Not all the MMRP participants answered the questionnaire at FU-12. The only significant difference at baseline was a small difference in pain intensity. But bias cannot be ruled out since the non-responders may have had worse effects of MMRP than the responders. It was not possible to analyse our results in relation to the contents and it was not possible to control for small changes in content over time. There are no internationally accepted definitions of MMRP [61, 81] and no taxonomies for the contents of these programs exist [17, 61], both issues that hinder further improvements. The general goals of MMRP may not be relevant to—or seen as relevant
by—every patient. In the present study, the importance of the general goals was not investigated and handled in the regressions. Another limitation is that we have no records regarding the level of patient attendance on the programs.

**Conclusion**

This study—representing the consecutive non-selected flow of patients in real-world clinical practice settings using SQRP—confirmed systematic reviews that outcomes of MMRP are associated with broad positive effects. A mix of background and baseline variables affected the outcomes investigated, but the explained variations in outcomes were low. In the future there is a need to develop a standardized and relatively simple outcome that can be used to evaluate a complex intervention such as MMRP in trials, in clinical evaluations at group level, and for individual patients.

**Additional file**

Additional file 1: Table S1. Demographic and outcome variables before MMRP (denoted Pre) and 12-month follow-up (FU-12) used in the study. Table S2. Comparison of results on pain, psychological variables and QoL. Table S3. Comparison of results on the SF-36 at pre, post, and follow-up. Table S4. Comparison of results on MPI at pre, post, and follow-up. Table S5. Comparison of results on LISAT at pre, post, and follow-up. Table S6. Handling of single missing items of the different instruments. (DOCK 29 kb)

**Abbreviations**

CBT: Cognitive behavioural therapy; CPAQ: Chronic Pain Acceptance Questionnaire; CPAQ-AE: The Chronic Pain Acceptance Questionnaire subscale activity engagement; CPAQ-PW: The Chronic Pain Acceptance Questionnaire subscale willingness; Dr-visits: Number of visits to a physician previous year; EQ5D-index: The European Quality of Life instrument index; EQSD-VAS: The European Quality of Life instrument health scale; FU-12: 12-month follow-up; HADS: Hospital Anxiety and Depression Scale; HADS-A: Hospital Anxiety and Depression Scale subscale anxiety; HADS-D: Hospital Anxiety and Depression Scale subscale depression; ICF: International Classification of Functioning, Disability and Health; LISAT: Life Satisfaction Questionnaire; LISAT-ADL: The Life Satisfaction Questionnaire subscale Activity of daily living; LISAT-psychhealth: The Life Satisfaction Questionnaire subscale psychological health; LISAT-somhealth: The Life Satisfaction Questionnaire subscale somatic health; LR: Logistic regression; MLE: Multiple linear regression; MMRP: Multimodal rehabilitation programs; MPI: Multidimensional Pain Inventory; MPI-GAI: The MPI subscale General Activity Index; MPI-LifeCon: the MPI subscale Perceived Life Control; MPI-PainInterfer: the MPI subscale perceived interference in everyday life; MPI-SocSupp: The MPI subscale Social support; MVDA: Advanced multivariate analyses; NRS-7days: Average pain intensity the last week; NRS: Numeric pain intensity scale; PLS: Partial Least Square Regression; PPI: Pain region index; RCT: Randomized controlled trials; Retro-life situation: Retrospective global evaluation of change in ability to handle life situation in general; Retro-pain: Retrospective global evaluation of change in pain; RTW: Return to work; SD: Standard deviation; SF36: The Short Form Health Survey; SF36-MSC: The Short Form Health Survey mental summary component; SF36-PF: The Short Form Health Survey subscale physical functioning; SF36-PSC: The Short Form Health Survey subscale psychological functioning; SQRP: Swedish Quality Registry for Pain Rehabilitation; SR: Systematic reviews; Tampa: The Tampa Scale for Kinesiophobia; YLD: Years Lived with Disability

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**Availability of data and materials**

Data will not be shared since this was not included in the consent form signed by the patients or included in the ethical application. Moreover, it cannot be excluded that it will be possible to identify individual patients indirect if the dataset is published.

**Authors’ contributions**

BG, PG, GS, B-MS and PE were involved in study conception and study design. BG performed the data analyses and drafted the manuscript. All authors (BG, PM, GS, B-MS and PE) discussed the results, commented on the manuscript, and approved the current version of the manuscript.

**Competing interests**

The authors declare that they have no competing interests.

**Consent for publication**

Not applicable.

**Ethical approval and consent to participate**

The study was granted ethical clearance by the Umeå University Ethics Committee (Dnr: 2013/192-31). After receiving written information about the study, all subjects signed a consent form that was in accordance with the Declaration of Helsinki.

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