Prognostic Factors for Physical Functioning After Multidisciplinary Rehabilitation in Patients With Chronic Musculoskeletal Pain

A Systematic Review and Meta-Analysis

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Objectives: This systematic review aimed to identify and evaluate prognostic factors for long-term (≥6 mo) physical functioning in patients with chronic musculoskeletal pain following multidisciplinary rehabilitation (MDR).

Materials and Methods: Electronic searches conducted in MEDLINE, PsycINFO, EMBASE, CINAHL, Web of Science, and Cochrane CENTRAL revealed 25 original research reports, published 1983-2016 (n = 9436). Potential prognostic factors relating to initial pain and physical and psychological functioning were synthesized qualitatively and quantitatively in random effects meta-analyses. The level of evidence (LoE) was evaluated with Grading Recommendations Assessment, Development and Evaluation (GRADE).

Results: Pain-related factors (intensity and chronicity) were not associated with function/disability at long-term follow-up, odds ratio (OR) = 0.84; 95% confidence interval (CI), 0.65-1.07 and OR = 0.97; 95% CI, 0.93-1.00, respectively (moderate LoE). A better function at follow-up was predicted by Physical factors; higher levels of initial self-reported functioning, OR = 1.07; 95% CI, 1.02-1.13 (low LoE), and Psychological factors; low initial levels of emotional distress, OR = 0.77; 95% CI, 0.65-0.92, low levels of cognitive and behavioral risk factors, OR = 0.85; 95% CI, 0.77-0.93 and high levels of protective cognitive and behavioral factors, OR = 1.49; 95% CI, 1.17-1.90 (moderate LoE).

Discussion: While pain intensity and long-term chronicity did not predict physical functioning in chronic pain patients after MDR, poor pretreatment physical and psychological functioning influenced the prognosis negatively. Thus, treatment should further target and optimize these modifiable factors and an increased focus on positive, psychological protective factors may perhaps provide an opening for yet untapped clinical gains.

Key Words: chronic musculoskeletal pain, GRADE, interdisciplinary rehabilitation, meta-analysis, prognostic factors, treatment outcome

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Chronic musculoskeletal pain (ie, pain duration >3 mo) such as chronic neck/shoulder and back pain, or generalized widespread pain, is a major health and socio-economic burden. Although etiology, localization, and diagnoses might differ, chronic pain itself could be considered a disease in its own right.¹ About a quarter of the adult population live with chronic pain of significant intensity,²,³ which may result in poor health including psychological distress, reduced quality of life, impaired physical functioning, reduced work ability, and increased sick leave.⁴

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From a therapeutic perspective, chronic musculoskeletal pain is a complex, multifaceted condition. A biopsychosocial approach is necessary for understanding and treating chronic pain—as a result, a comprehensive, multimodal and interdisciplinary, pain management method, here referred to as multidisciplinary rehabilitation (MDR, also known as interdisciplinary rehabilitation, multimodal rehabilitation, and multimodal pain therapy) is advised for this patient group. On the basis of a cognitive-behavioral therapy approach, it incorporates education, physical activity and exercise, coping skills, and occupational therapy sessions in a multimodal rehabilitation program. MDR is administered by multidisciplinary teams, which commonly include physicians, psychologists, physiotherapists, occupational therapists, social workers, and other health professionals. The team’s collaboration in assessment and shared goal-setting is an essential component, adding value beyond the effects of the multiple modalities provided in pain treatment. Existing data shows that MDR is effective compared with single-treatment or treatment-as-usual programs, but the effects are at best moderate and need further study.

Studying effectiveness and effect moderators of MDR in patients with chronic musculoskeletal pain has been recognized as a major challenge. The complexity of the various pain conditions and the complexity of the intervention itself, accompanied by the lack of a standardized, internationally accepted definition of the treatment, hinder comparative clinical trials and meta-analyses, which delays evidence on how outcomes for this patient group can be optimized. It is, however, believed that outcomes would improve if treatments could be better customized to a patient’s profile, that is the characteristics of their initial biopsychosocial status.

Prognostic factor research aims to identify factors associated with clinical outcomes to provide data on the likely health outcomes among people with a given health condition. Riley et al. state that prognostic factors can help “inform clinical and therapeutic decisions (either directly or as part of prognostic models for individualised risk prediction)... and help identify targets for new interventions that aim to modify the course of a disease or health condition.” Predictive factor (or predictor) is a term related to the term prognostic factor, and these are sometimes used synonymously. Predictive factor, however, is used more in the context of measures of response to a given therapy among others. In the present study, we have chosen to use the term prognostic factor or indicator consistently.

Although many clinical studies have performed these analyses to identify factors of importance for future outcomes in patients with chronic musculoskeletal pain, the body of evidence of prognostic factors is still insufficient to predict MDR outcomes. Rather than looking at any study in isolation, systematic reviews can provide an overview of a whole body of evidence of possibly important prognostic factors available at pretreatment level, across a number of outcomes that are targeted by the MDR. In this first part of the systematic reviews, we have focused on physical functioning as the main outcome.

Improving physical functioning and decreasing pain’s interference with functioning are of great clinical importance, and these are therefore important targets of MDR. Knowledge of early prognostic indicators of outcome is therefore of great clinical importance as well.

**OBJECTIVE**

The aim of this systematic review was to identify, evaluate, and meta-synthesize published data on prognostic factors, related to baseline information on pain and physical and emotional functioning, for physical functioning at least 6 months post MDR in patients with chronic musculoskeletal pain.

**MATERIALS AND METHODS**

This systematic review used a random effects meta-analysis of published original research reports with a longitudinal follow-up of early prognostic factors preceding MDR. The review was conducted by an interdisciplinary research team. It conforms to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement with particulars of the Meta-analysis Of Observational Studies in Epidemiology (MOOSE) guidelines in consideration. A protocol with the defined objectives, eligibility criteria, and planned methods of the complete review project was registered in the International Prospective Register of Systematic Reviews (PROSPERO, id:CRD42016025339) at an early stage of the study, and a study protocol reporting the review process was published ahead of the study.

**Data Sources and Search Procedure**

Articles published in English between 1980 and April 2017 were identified through systematic electronic searches of 6 reference databases: MEDLINE and PsyINFO (via Ovid), EMBASE (via Elsevier), CINAHL (via EBSCO), Web of Science (via Thomson Reuters), and the Cochrane Central Register of Controlled Trials (CENTRAL). With the support of
a research librarian, we developed a comprehensive search strategy combining 4 search parameters; “Chronic Pain”—“Multidisciplinary Rehabilitation”—“Treatment Outcome”—“Prediction,” for inclusion of all common diagnoses of chronic musculoskeletal pain conditions targeted in MDR comprising a follow-up of clinically important outcomes and explorative approaches to all prognostic factors possibly studied. An a priori decision was made to only search for published work. Consistent with the explorative objective, the search was unrestricted except for 2 limitations; publication language and publication date. To identify additional studies, a manual search of reference lists of obtained and relevant articles was conducted. The complete search strategy is described in detail in Supplemental Appendix 1 (Supplemental Digital Content 1, http://links.lww.com/CJP/A533).

Study Selection

Inclusion criteria were: (1) studies with a longitudinal design, either observational (cohort, case-control) or experimental/clinical trials (randomized controlled trial), (2) studies that investigated prognostic factors of treatment outcome, (3) in adults aged 18 to 67 years (ie, the working-age population), with a chronic musculoskeletal pain condition; defining chronic as a duration of >3 months and delimitating musculoskeletal pain conditions to common nonspecific musculoskeletal pain diagnoses such as back pain, neck pain, and generalized pain syndromes (including fibromyalgia and general widespread pain) but not those emanating from, for example, malignancies, systemic or inflammatory diseases (eg, rheumatoid arthritis), or degenerative joint diseases (eg, osteoarthritis-related joint pain), (4) studies on patients that had taken part in multidisciplinary/interdisciplinary/multimodal rehabilitation following the biopsychosocial model and coordinated by ≥3 different health professionals. MDR could be of any duration/intensity and rehabilitation approach, in inpatient or outpatient settings, (5) studies on interventions that targeted core outcome domains as recommended by the Initiative on Methods, Measurement, and Pain Assessment in Clinical Trials (IMMPACT), and reported results on either pain, physical functioning, work ability, or health-related quality of life (QoL) and emotional functioning.30 Outcome measures were allowed to vary as long as they could be grouped under the domains of interest, (6) studies with a follow-up of 6 months or longer were set as a minimum time criterion for analysis of clinically relevant long-term outcomes, and (7) only original research reports in peer-reviewed journals, published in English, and in full text were eligible.

Studies were excluded if: (1) they lacked a presentation of data from baseline to a follow-up of at least 6 months in the prediction analyses, or (2) they investigated the process of change as a prognostic factor, that is the actual changes occurring during treatment as prognostic indicators of outcome. Eligibility criteria were defined as PICOT (ie, Population Intervention/Variable of Interest Comparator Outcome and Time).25

The study selection procedure was performed in the Covidence online systematic review platform31 and a PRISMA flow diagram32 was used to document the flow of included and excluded studies, along with the reasons for exclusion (Fig. 1). The selection process was performed in 4 steps: (1) screening of titles, (2) screening of abstracts, (3) screening of full texts for PICO eligibility, and (4) screening of full texts for relevance according to study objective.

A first raw screening of titles was performed by one reviewer. During the following selection steps, every article was appraised by 2 reviewers independently. The articles were randomly assigned to the reviewer teams. Every step was first piloted to validate the interpretation of the criteria. Interrater agreement throughout the review process was evaluated and agreement ranged from 72% to 83% (Cohen κ = 0.342 to 0.648). Disagreements were resolved through discussions with the full review team.

In the current study, further selection was made for papers evaluating the outcome “Physical functioning.” Typically, measures commonly used in clinics assess either the ability for various sorts of functioning, or conversely the inability for functioning, that is disability—thus reflecting opposite perspectives of the same construct (physical functioning). Moreover, only prognostic factors related to initial pain and physical and emotional functioning were included for analysis in the present paper. Sociodemographic factors will be presented elsewhere (Fig. 1).

Quality Assessment

Articles deemed relevant from the full text screening were assessed for internal validity with The Quality in Prognostic Studies (QUIPS)-tool.33 Potential threats to validity were assessed within the 6 domains: (1) study participation, (2) study attrition, (3) prognostic factor measurement, (4) outcome measurement, (5) study confounding, and (6) statistical analysis and reporting. Similar to Cochrane’s risk of bias (RoB) assessment, but with emphasis on evaluating critical methodological criteria for bias in prognostic studies as recommended by the Cochrane Prognosis Methods Group. All articles were assessed independently by 2 reviewers: 1 senior reviewer assessed all studies, which were then divided between 2 other researchers in accordance with the randomization scheme. The process was piloted a priori for interrater agreement. The percent agreement ranged between 48% and 81% and the Prevalence and Bias Adjusted Kappa-Ordinal Scale (PABAK-OS) across RoB domains varied between 0.227 and 0.719. Consensus on overall study quality, was based on thoughtful scrutiny for all 6 domains together. We classified a study to have a low RoB when at least 5 of the domains had low RoB and none of the domains had high RoB, a Moderate RoB when the study had a maximum of 2 moderate RoB and the rest low RoB, and a high RoB study when one or more domains had high RoB or there were 3 domains or more with moderate RoB. The full RoB assessment was also analyzed across every RoB-domain separately to identify specific problematic areas pertaining to a specific outcome. The analyses of overall study quality were later also incorporated in the Grading of Recommendations Assessment, Development and Evaluation (GRADE) summary, under the factor “study limitations.”

Data Extraction and Data Syntheses

From each included study, data were collected on: (1) participant and sample characteristics, (2) intervention characteristics, (3) treatment-related characteristics, and (4) outcomes of interest. The synthesis of the outcomes for each prognostic factor, was rated as low/moderate/high RoB. The synthesis of the between-studies risk of bias (ROB), for overall study quality, was based on thoughtful scrutiny for every outcome as we avoided making a simple summary score. Every outcome was assessed in 2 ways: (1) by classifying each study into 3 levels of RoB based on the ratings of all 6 domains together. We classified a study to have a low RoB when at least 5 of the domains had low RoB and none of the domains had high RoB, to a Moderate RoB when the study had a maximum of 2 moderate RoB and the rest low RoB, and a high RoB study when one or more domains had high RoB or there were 3 domains or more with moderate RoB. (2) RoB was also analyzed across every RoB-domain separately to identify specific problematic areas pertaining to a specific outcome. The analyses of overall study quality were later also incorporated in the Grading of Recommendations Assessment, Development and Evaluation (GRADE) summary, under the factor “study limitations.”
characteristics, (3) independent variables (potential prognostic factors) and assessment methods, (4) dependent variables (outcome domains) relating to physical functioning (primary outcome in the present study), work ability, health-related QoL, pain, emotional functioning and their assessment methods, (5) research design, kind of study, study phase and follow-up time, and (6) statistical outcomes, conclusions and further statistical data. Data were extracted to a digital coding protocol by 2 reviewers (W.J.A.G., E.T.) independently, and compared for data accuracy and consensus before analysis.34

Descriptive analysis was then performed on this database. When coding was completed, all reported variables (potential prognostic factors), n ≥ 200, were presented to the review team for a consensus-reaching grouping process, by which similar variables were collated into coherent domains, with related prognostic factors, to be used in further analyses. Variables that were too disparate to be included in any domain were specified in the original synthesis file, for transparency of the grouping process. When all found prognostic factors and domains were set, the analyses for the current study with the primary outcome physical functioning was initiated, parting the remaining outcome domains for later analysis.

A narrative synthesis of the relation between each potential prognostic factor and the outcome physical functioning was performed, in which the direction (positive, negative, or absence of association) was stated. Depending on how data were presented in the original studies, results were, if necessary, reversed to fit the chosen reporting direction of synthesis, that is for “positive outcome,” for example low levels of disability and high levels of physical functioning.

A quantitative synthesis was also performed. When at least 2 studies provided data on the same prognostic factor, a subsequent meta-analysis was aimed for, based on our a priori decision. All outcome data required for the meta-analyses were extracted from the coding protocol and

FIGURE 1. PRISMA flow chart of study selection. HRQoL indicates health-related quality of life; PRISMA, Preferred Reporting Items for Systematic Review and Meta-Analysis.
complemented with details from the articles by the 2 reviewers together, and then double-checked once more. To quantify the strength of the relationship between identified prognostic factors and corresponding outcomes, the statistical outcomes (effect sizes) from single studies were converted into a common index to permit pooling across studies.\textsuperscript{33} The odds ratio (OR) was set as the common index used in our analyses, an effect size frequently used in prognostic studies. Web-based calculators\textsuperscript{36,37} were used to compute and transform any relevant data that were not reported as ORs, that is continuous and correlational data, into ORs and their 95\% confidence intervals (CIs). The complete methodology for these procedures is described in Lipsey.\textsuperscript{38} In the software Review Manager,\textsuperscript{39} variance weighted pooled ORs were then computed in a random effects model for each prognostic factor, using the generic inverse variance method, which permits a wide selection of data formats in the analyses.\textsuperscript{40} For every meta-analysis, inverse variance method, which permits a wide selection of effects model for each prognostic factor, using the generic weighted pooled ORs were then computed in a random study phase.\textsuperscript{40} If there was not a majority for low RoB, we downgraded for study limitation. Judgment of inconsistency, influencing the estimates of prognosis, was based on an evaluation of all analyses (narrative, quantitative, sensitivity, and the I\textsuperscript{2} statistics). Indirectness, generalizability, was assessed through an estimation of our included material. Imprecision was not deemed possible to judge in our study. Publication bias was assessed through funnel plots and a comparison of effects included in narrative and quantitative syntheses.

RESULTS

Results of the Literature Search

Electronic searches identified 3355 candidate studies, and 491 full text articles were retrieved. A total of 187 studies met the PICOT eligibility criteria and were subsequently screened once more for relevance. Of these, 105 studies met our relevance criteria and were included in the present review for further analysis, within the prespecified outcomes—physical functioning, pain, work, and QoL. During the data extraction and the process of narrative synthesis, additional studies were excluded for the following reasons: lack of sufficient data on the prognostic factors of interest (10), data provided only on change factors instead of baseline factors (10), mixed group analyses, that is prediction analyses of MDR-treated groups and control groups together (5), or double reporting of data (1), wrong outcome (1), or other (3). As a result, 75 studies remained eligible for analysis, and from these, the 25 studies that reported on prognostic factors for the outcome Physical functioning were selected for analysis and included in the present report (Fig. 1).

Description of Studies Included in Analysis

The 25 included studies consisted of 24 cohort studies\textsuperscript{41–43,48–68} and 1 randomized controlled trial.\textsuperscript{69} Nineteen studies were primary prognostic factor studies while 3 examined prognosis as their secondary aim, and 2 examined validation/study methodology. Follow-up time ranged from 6 to 18 months with a loss to follow-up between 0\% and 51\% (median = 14\%). In total, 9436 participants were included in the studies, with sample sizes ranging from 39 to 3106 participants for the single studies (mean n = 377, median n = 143). The studies were published between 1983 and 2016. Nineteen studies were conducted in Europe (Sweden 6, Germany 4, the Netherlands 5, Norway 1, Finland 1, Denmark 1, Switzerland 1), 5 in North America (USA 3, Canada 2), and 1 in New Zealand (Table 1). Studies included patients with an average age ranging from 38 to 54 and the percentage of females in their samples ranged from 35\% to 100\%. Studied diagnoses were chronic (low) back pain (n = 12), chronic pain (n = 9), fibromyalgia (n = 2), generalized widespread pain (n = 1), and whiplash-associated disorder (n = 1). The participants’ average pain duration ranged from 3 months to > 10 years; the majority


| References | Country | Study Design* (Phase I-III)† | Type of Study | Prognostic Factor Measurement Instruments | Outcomes Measurement Instruments | Follow-up Period (mo) |
|------------|---------|-------------------------------|---------------|------------------------------------------|---------------------------------|----------------------|
| Angst et al41 | Switzerland | (i) II | SF-36 BP & PF, HADS, sports activities h/wk, CSQ | SF-36 PF | 6 |
| Bendix et al48 | Denmark | (i) II | Biering-Sørensen tests of abdominal and back muscle endurance, aerobic capacity l/min, mobility and ADL-scores/disability index | ADL-scores/disability index | 12 |
| Bergström et al49 | III | MPI-S, profiles: DYS, AC, and ID | MPI-Interference | 12, 18 |
| Bergström et al50 | III | BAS | MPI-Interference | 12 |
| Breman et al51 | II | VAS, HADS | MPI-Interference | 6 |
| Checchanowski et al52 | USA | III | VAS, RMDQ, CES-D, CSQ, RSQ | SF-36 PF | 6 |
| de Rooij et al53 | The Netherlands | (i) III | NRS, HADS subscale: anxiety, BDI-II, SCL-90, IPQ-R, CSQ, PCI subscale: resting (avoidance behavior), DGSS, TSK | MPI-Interference | 6 |
| Dobkin et al54 | Canada | (i) II | FIQ, MPQ-VAS, ASES, CES-D | FIQ | 6 |
| Farin et al55 | Germany | (i) III | BAS, NRS, timeline, treatment motivation | SF-12 PC, ODI | 6 |
| Gerdlle et al56 | II | NRS, MPI, pain duration/persistance, SF-36 PF, PCS & MCS, HADS, CPAQ, TSK, EXPECT RTW | MPI-Interference, SF-36 PF | 12 |
| Glattacker et al57 | Germany | (i) II | VAS, pain duration, SF-36, ODI, IRQ, RKP-B | SF-36 PF and R, ODI, FCi | 6 |
| Harkapa et al58 | Finland | (i) III | VAS, medical optimism, LoC-beliefs | FCi | 12 |
| Lénastry & Olszynski59 | Canada | (i) III | VAS, MPI, pain duration, HADS, SF-36, ODI | PDI | 15 |
| Lillegård et al60 | Norway | (i) II | VAS, HADS | Functional Health Status (COOP/WONCA) | 12 |
| Ludwig & Bergsten et al61 | Sweden | III | TSK | DRI | 6 |
| McCready et al62 | USA | (i) III | VAS | MVAS | 12 |
| Moradi et al63 | Germany | (i) II | Biering-Sørensen test, Villiger test, Oesch test | PDI, FFBH-R | 6 |
| Moss-Morris et al64 | New Zealand | (i) III | SF-36 PCS | SF-36 PCS | 6 |
| Persson et al65 | Sweden | (i) III | DRI, MPI subscales, COPM | COPM | 12 |
| Ruscheweyh et al66 | Germany | (i) III | NRS, pain duration | PDI | 6 |
| Trief & Yuan 67 | USA | (i) III | MMPI | Activity level | 8-12 |
| van Hooff et al68 | The Netherlands | (i) II | VAS, ODI, ZSDS, PSEQ, PCS, TSK | ODI | 12 |
| Vendar et al69 | The Netherlands | III | MPI | QBPD, MISE | 6 |
| Vendring et al70 | The Netherlands | III | MPI | QBPD, MISE | 6 |
| Verkerk et al71 | The Netherlands | (i) II | VAS, pain duration, QBPD, SF-36 PCS & MCS, SCL-90, TSK | QBPD | 12 |

* (i), hypothesis generating; (ii), extensive exploratory, (iii), confirmation of hypothesis.
† RCT study.
§ Not included in synthesis.

ADL-score/Disability index indicates Low Back Pain Rating scale; ASES, Arthritis Self-Efficacy Scale; B200 Isostation, Physical performance test-back extension strength; BAS, Body Awareness Scale; BDI, Beck Depression Inventory; BDI-II, Beck Depression Inventory (II); Biering-Sørensen test, physical performance test—back muscle strength; BP, Bodily Pain; BRQ, Beliefs about Rehabilitation Questionnaire; CES-D, Center for Epidemiological Studies—Depression Scale; COOP/WONCA, Functional Health Status measurement (Darmouth COOP Functional Health Assessment Charts/World Organization of Family Doctors); COPM, Canadian Occupational Performance Measure; CPAQ, Chronic Pain Acceptance Questionnaire; CSQ, Coping Strategies Questionnaire; DGSS, Dutch General Self-Efficacy Scale; DR1, Disability Rating Index; EXPECT RTW, perceptions of prognosis on return to work; FABQ, Fear-Avoidance Beliefs Questionnaire; FCI, Functional Capacity Index (based on RMDQ); FFBH-R, Hannover Functional Ability Questionnaire (German); FIQ, Fibromyalgia Impact Questionnaire; HADS, Hospital Anxiety and Depression Scale; Health expectations, not specified instrument; Health Optimism, Health Optimism Brief Scale; IPQ-R, Illness Perceptions Questionnaire-Revised; KKG, Control beliefs Concerning Illness and Health (German); LoC, Locus of Control beliefs, from Health+Pain Locus of Control Scale; MISE, Maximal Isometric Strength Extension (Trunk muscle performance test); MMPI, Minnesota Multiphasic Personality Inventory; MMPI-2, PSY-5 scale, The MMPI-2 Personality Psychopathology Five; MPI, Multidimensional Pain Inventory; MPI-Interference, subscale of MPI; MPI-S, MPI-Swedish version. Here classified into profiles: Dysfunctional (DYS), Adaptive Coper (AC), and Interpersonally Distressed (ID); MPQ, McGill Pain Questionnaire; MVAS, The Million Visual Analogue Scale; NHP, Nottingham Health Profile, subscale: physical ability; NRS, Numeric Rating Scale; ODI, Oswestry Disability Index; Oesch test, physical performance test-arms strength; Pain duration, measured by self-report/questionnaires; PCI, Pain Coping Inventory; PCS, Physical Component Summary; PCS, Pain Catastrophizing Scale; PDI, Pain Disability Index; PF, Physical Functioning; PSEQ, Pain Self-Efficacy Questionnaire; QBPD, Quebec Back Pain Disability Scale; RMDQ, Roland Morris Disability Questionnaire; RP, Role-Physical; RSQ, Relationship Scale Questionnaire; SCL-90, Symptom Checklist-90; SF-12, 12-Item Short Form Health Survey; Physical Health Summary Scales (PCS); SF-36, 36-Item Short Form Health Survey—Stages of change, Pain Stages of Change Questionnaire; TSK, Tampa Scale for Kinesiophobia; VAS, Visual Analogue Scale; Villiger test, physical performance test-Step test; ZSDS, Zung Self-Rating Depression Scale.
of them had had chronic (persistent) pain for several years. Participants were recruited or referred from primary care, secondary care, or insurance providers.

Interventions were described using the following nomenclature; “multidisciplinary/multimodal/ interdisciplinary” (19), “functional restoration program” (5), and “work hardening program” (1). The intervention duration varied mainly between 2 and 8 weeks, although some interventions were performed in 2 phases, in which a longer follow-up period with continued rehabilitation time was offered for as long as a year. Twelve of 25 studies reported an MDR intervention time of 4 to 8 weeks, and 7 studies reported a longer duration; either >8 weeks or >8 weeks when both phases were added together. The majority of studies reported an average total of 100 hours, although this could be delivered as full time treatment over the period of a couple of weeks or more spread out over a couple of months (Table 2).

Outcome Measures
Both generic and disease-specific measures for physical functioning were used. The outcomes relating to physical functioning were assessed either with measures of physical functioning or measures of disability, or a combination of both. Outcome measures used to assess physical functioning, included ADL scores, the Coop Functional Health Assessment Charts (COOP/WONCA), Functional back capacity (FFbH-R), the Maximal Isometric Strength Extension (MISE), the Functional Capacity Index (FCI), and scales from the 36-Item Short Form Health Survey (SF-36); Physical Functioning (PF), Role-Physical (RP), Physical Component Summary (PCS) and respectively, the 12-Item Short Form Health Survey (SF-12). For disability, measures included the Roland-Morris Disability Questionnaire (RMDQ), the Oswestry Disability Inventory (ODI), the Disability Rating Index (DRI), the Quebec Back Pain Disability Scale (QBPDFS), the Pain Disability Index (PDI), the Fibromyalgia Impact Questionnaire (FIQ), and the Multi-dimensional Pain Inventory (MPI)-Interference scale. Most of the measures were based on self-reports, that is Patient Reported Outcome Measures (PROM), whereas some were performance-based and assessed by the MDR team.

Prognostic Factors
A total of 87 baseline factors were identified, which were operationalized into domains. Three domains and their related potential prognostic factors were included for synthesis; Pain-related factors, Physical function-related factors, and Psychological factors, in analogy to the assessment topics of the IMMPACT.

(1) Pain-related factors: pain intensity and pain duration. Assessment measures included Numeric Rating Scale (NRS), Visual Analogue Scale (VAS), and the SF-36—Bodily pain (SF36—BP).
(2) Physical function-related factors: performance-based function (e.g. muscle strength, mobility, aerobic capacity, and self-rated function, expressed in terms of physical ability or disability). Function-related factors were assessed with the same measures as the primary outcome (e.g., PDI, ODI, SF-36).
(3) Psychological factors: psychological measures were sorted under higher order factors “emotional distress” and “cognitive-behavioral factors” to ensure relatively homogenous categories.70

- Emotional distress, for example, anxiety and depression. Assessment measures included the Hospital Anxiety and Depression Scale (HADS), the Beck Depression Inventory (BDI), the Center for Epidemiological Studies-Depression Scale (CES-D), the Minnesota Multiphasic Personality Inventory (MMPI), the Symptom Checklist-90 (SCL-90), and the SF-36 Mental component scale (SF-36 MCS).
- Cognitive-behavioral factors, either with a positive direction, for example, health optimism, personal control, and self-efficacy or with a negative direction, risk, for example, catastrophizing, fear of movement, avoidance behavior, and external locus of control. Assessment measures included the Health Optimism Scale, the Health Locus of Control Scale, the Dutch General Self-efficacy Scale (DGSS), Pain Self-Efficacy Questionnaire (PSEQ), the Pain Coping Inventory (PCI), the Coping Strategy Questionnaire (CSQ), the Tampa Scale of Kinesiophobia (TSK), the Revised Illness Perceptions Questionnaire (IPQ-R), the Beliefs about Rehabilitation Questionnaire (BRQ), the Minnesota Multiphasic Personality Inventory-2 (MMPI-2), the Multidimensional Pain Inventory (MPI), subscale Life control—some of which evaluate both risk and protective factors.

Sociodemographic-related, Medical-related, and Work-related factors were identified as well, but will be reported in a separate report, due to the large amount of diverse factors provided in these domains.

Methodological Quality
The within-studies RoB is presented as a total percent of included studies for the 6 assessed domains of validity (Fig. 2). The domains study attrition and study confounding emerged with the highest RoB (ie, low quality), mainly due to insufficient reporting on these topics in the paper. Ratings for individual studies are displayed in Table 3.

PROGNOSTIC FACTORS FOR PHYSICAL FUNCTIONING—NARRATIVE AND QUANTITATIVE ANALYSES

Pain-related Factors
Pain Intensity
The association between baseline pain intensity and physical functioning after MDR was assessed in 16 studies,41–43,48,51–55,57,59,62,63,65,68,69 including a total of 8191 participants.

The narrative analyses indicated inconclusive results. Eight studies32,43,52,53,55,57,59,68,69 reported no association between pain intensity at baseline and outcome. Four studies54,57,59,63 showed that lower levels predicted positive outcomes while 2 studies41,51 showed that high pain levels at baseline predicted positive results at follow-up. Two studies had conflicting results, depending on pain location46 or type of analysis (uni/multivariate)62 (Table 4).

Five studies (4 low, 1 high RoB) provided continuous data for inclusion in a meta-analysis (n = 2676). Results of the meta-analysis showed that initial pain intensity was not associated with improvement in physical function at follow-up, OR = 0.84; 95% CI, 0.65-1.07; P = 0.16 (Fig. 3A).
### TABLE 2. Description of Participants and Intervention

| References | Diagnosis       | Participants (n) | Age Mean (SD), Median (IQR) (y) | % Female | Duration of Pain/Disability Mean (SD); Median (IQR) | Intervention Profile | Intervention Time | Intervention Details |
|------------|-----------------|------------------|---------------------------------|----------|----------------------------------------------------|----------------------|-------------------|---------------------|
| Angst et al41 | Whiplash        | 175              | Mean 37.4 (11.7)                | 79       | Mean, 13.3 mo (10.7)                               | Interdisciplinary, multimodal program | 4 wk              | Inpatient care. The program included physiotherapy individually and in small groups, medical training therapy (MTT, graded exercise), passive therapy modules, occupational therapy, creative therapy, neuropsychological treatment with group information about pain, individualized cognitive behavioral therapy and a test psychological setting. |
| Bendix et al48 | CLBP            | 621*             | Median, 40                      | ≥ 6 mo   | Functional restoration program with the goal of restoring the patient’s health physically and psychologically | 3 wk daily sessions, 8 h, and 3 wk 1 d/ wk, in total 135 h | Outpatient care. A combination of physical and ergonomic training, psychological pain management, patient education, and counseling about return to work. |
| Bergstrom et al49 | CBP           | 156              | Mean, 42.5 (9.5)                | 48       | Mean, 38 mo (SD, 63.4)                             | Work hardening program | 4 wk              | Outpatient care in groups consisting of 10-12 participants. Rehabilitation was based on interdisciplinary collaboration and the patient as an active team member. Basic Body Awareness Therapy, swimming pool exercises, rhythm and movement exercises, and relaxation exercises were modalities run by physiotherapists. Other sessions in the program were coping, ergonomics, and education about pain mechanisms and its consequences. The patients also had individual activities and contacts. On the basis of the patient’s needs, contact was established with key persons such as the patient’s primary care physician and representatives from the Swedish Social Insurance Agency and their employer, as one main goal of the MMRP was decreased sick leave and return to work. These persons were invited together with the patient and significant others to a final team meeting at the end of the program. |
| Bergström et al50 | Chronic pain    | 39               | Mean, 41                        | 80       | ≥ 6 mo                                             | Multimodal program based on cognitive-behavioral principles and focused on pain management and education about pain and its consequences and bodily and psychological reactions to pain. | 5 wk              | Treatment was performed both in group and in individual sessions. There was a daily combination of physical treatment, including qigong, body awareness, pool exercise, and sessions with a cognitive-behavioral approach. The program had a nonpharmacological profile. |
| Bremander et al52 | Chronic pain    | 97               | Mean, 44.6 (9.7)                | 88       | ≥ 3 mo                                             | Functional restoration program aimed at improving pain management skills and physical and psychological functioning, with the main focus on coping with daily life to improve HRQoL. | 3 wk inpatient care and 6 mo outpatient care | (Continued) |
| References          | Population Diagnosis | Participants (n) | Age Mean (SD), Median (IQR) (y) | % Female | Duration of Pain/Disability Mean (SD); Median (IQR) | Intervention Profile | Intervention Time | Intervention Details |
|---------------------|----------------------|------------------|---------------------------------|----------|-----------------------------------------------------|----------------------|-------------------|---------------------|
| Ciechanowski et al51| Chronic pain         | 111              | Mean, 44.7 (10.7)               | 55       | Mean, 6.3 y (7.8)                                   | Multidisciplinary program aimed at improving patient pain-management skills and physical and psychological functioning | 3 wk                | Outpatient care. The program contained physical and occupational therapy, individual cognitive-behavioral psychotherapy, vocational counseling, group pain education and coping-skills training, and the tapering of opioid and sedative-hypnotic medications when indicated. It also included a strong emphasis on fostering active pain self-management skills and reducing reliance on health care providers and passive pain management strategies. |
| de Rooij et al52    | CWP                  | 138              | Mean, 45.0 (10.3)               | 95       | NR                                                  | Multidisciplinary program                                         | 7 wk with group treatment, 2×3.5 h/wk. Individual treatment was offered for 4-6 mo, with variable frequency | 3 mo with 2-4 sessions per week, 2-4 h per session | The program included cognitive behavioral-therapy, the acquisition of pain management skills (eg, goal setting, structuring of daily activities, pacing strategies, ergonomics), physical training (eg, exercise), relaxation training, education about neuro-physiology and medication management, and assertiveness training. The treatment was tailored to the patients personal goals and was performed in groups and on an individual basis. The multidisciplinary team involved rehabilitation physicians, physiotherapists, occupational therapists, psychologists, and social workers. In a small group outpatient care setting, 6-8 sessions of physotherapy, occupational therapy, nursing education and intervention, and cognitive-behavior therapy. These sessions were held as closed groups (ie, did not admit new members once they started) with each treatment modality delivered by a different health professional. |
| Dobkin et al53      | Fibromyalgia         | 53               | Mean, 53.6 (14.5)               | 100      | NR                                                  | Interdisciplinary, multimodal program to educate patients about FM, prepare them to manage symptoms, improve sleep and coping skills, teach stress management, and to develop a fitness program that progressed slowly over time | 3 mo with 2-4 sessions per week, 2-4 h per session | 3 mo with a mean length of 20.6 (4.5) rehabilitation days | The multimodal programs included educational, somatic, psychotherapeutic, social, and occupation-related therapy. Examples of individual treatment elements are information (eg, providing information on chronic back pain and rehabilitation goals in educational group sessions), training based on a biopsychosocial disease model (eg, discussing dysfunctional health beliefs), occupational therapy, physical therapy, exercise therapy, and psychotherapeutic treatment to modify maladaptive illness behavior and learn techniques for relaxing and coping with stress. Multicenter study, 4 inpatient and 7 outpatient rehabilitation centers. The patient generally had 4-5 therapy sessions a day on workdays and 2-4 at weekends. Two outpatient rehabilitation centers. |
| Farin et al54       | CLBP                 | 688 but only 468 answered the 6 mo follow-up | Mean, 51.0 (11.2)               | 57       | Chronification (%), % <1, 13.0 1-2, 11.1 3-5, 18.6 6-10, 16.3 > 10, 40.2 | Multidisciplinary program                                         | 3 wk with a mean length of 20.6 (4.5) rehabilitation days | The multimodal rehabilitation program was conducted in groups of 6-9 participants and included physiotherapy, ergonomics, training in coping strategies, and education in pain science and pain management. Work-related advice for workdays and weekends was provided. |
| Gerdle et al55      | Chronic pain         | 464 but only 227 answered the 12 mo follow-up | Mean, 38.1 (10.1)               | 81.6     | Mean, 6.98 y (7.15)                                 | Multimodal program based on CBT principles                        | 6-8 wk, at least 20 h/wk | Two outpatient rehabilitation centers. The patient generally had 4-5 therapy sessions a day on workdays and 2-4 at weekends. Two outpatient rehabilitation centers. |

(Continued)
### TABLE 2. (continued)

| References               | Population Diagnosis | Participants (n) | Age Mean (SD), Median (IQR) (y) | % Female | Duration of Pain/Disability Mean (SD); Median (IQR) | Intervention Profile | Intervention Time | Intervention Details |
|--------------------------|----------------------|------------------|----------------------------------|----------|-----------------------------------------------|---------------------|------------------|---------------------|
| Glattacker et al43       | CLBP                 | 105              | Mean, 54 (11)                    | 37       | 49.5% > 10 y, and 3.8% <1 y                    | Interdisciplinary, multimodal program | 3 wk             | The inpatient care program included patient education, physical therapy, health education programs, occupational therapy, and psychological treatment, mainly in groups. However, altering illness beliefs was not an explicit or standardized component of the rehabilitation programs |
| Harkapaa et al56         | CLBP                 | 175              | Mean, 42.1                       | 48       | NR                                            | Multimodal back treatment program modified to emphasize the role of intensive physical training and work hardening methods | A 3-d preprogram, 5-wk home training period and 4-wk intensive, inpatient program | The main goal was to increase the daily functioning of the patient by improving physical functioning, by overcoming the fear of pain related to different activities and by increasing feelings of control and mastery. During the preprogram the rationale and methods were explained. The home training was a self-care program which consisted mainly of stretching and light physical exercises, aimed at preparing the patient for the intensive program. The 4-wk intensive program consisted of physical exercises, general work hardening methods, back school, relaxation training, cognitive-behavioral group therapy and socio-economic counseling. 7-8 h daily training sessions. |
| Lemstra & Olszynski69     | Fibromyalgia         | 43               | Mean, 49.7 (9.6)                 | 86       | Mean, 121.7 mo                                | Multidisciplinary program | 6 wk             | The intervention consisted of 18 group exercise therapy sessions supervised by a physical and exercise therapist, 2 group pain and stress management lectures by a psychologist, 1 group education lecture by a rheumatologist and 1 group dietary lecture, and 2 massage therapy sessions. There was no vocational or return to work component. The primary components of the intervention were submaximal general exercise, education, lifestyle changes, and self-management. Active participation was maximized with supervised visits, phone calls with every absence, and scheduled attempts to determine knowledge retention. The patients were involved in developing their own management plan, developing realistic short-term expectations, and identifying barriers to recovery and management. |
| Lillefjell et al57        | Chronic pain         | 143              | Mean, 45.7 (8.9)                 | 74       | NR                                            | Multidisciplinary program | 5 wk intensive, 4 d a week + 52 wk follow-up, -1-3 d a week, in total 57 wk of treatment | Mapping of the participants resources, the intensive training period of 6 h/d for 5 wk consisted of individual and group based training to improve functional capacity. Group-based education/training. Indoor and outdoor activities every day. Individual exercise program, eg, endurance, strength, relaxation. During follow-up training the functional capacity continues along with individual counselling and plan for work reentry. Additional exercise was also offered. |
| Lüning-Bergsten et al58   | CBP                  | 265              | Females: median, 45 (37-51)      | 49       | Sick leave days 2 y earlier, females: median, 275 (150-485) | Multidisciplinary program | 4 wk full-time | The inpatient program included physical training, education in anatomy, physiology, ergonomics, pain management, relaxation techniques, and physical work. | (Continued) |
### TABLE 2. (continued)

| References          | Population Diagnosis | Participants (n) | Age Mean (SD), Median (IQR) (y) | Duration of Pain/Disability Mean (SD); Median (IQR) | Intervention Profile | Intervention Time | Intervention Details |
|---------------------|----------------------|------------------|---------------------------------|------------------------------------------------------|----------------------|-------------------|---------------------|
| McGeary et al59     | CDOD                 | 3106             | males: median, 44 (37-50)       | males: 242 (85-425)                                   | Functional restoration program | NR, (usually 3-5 wk) | This program used quantitatively directed exercise progression under the supervision of both physical and occupational therapists and participation, which included individual counseling, group therapy, stress management, vocational reintegration, and future fitness management. Outpatient probably |
| Moradi et al60      | CLBP                 | 162              | Mean, 46 (11)                   | Mean, 2.3 y (0.8)                                    | Multidisciplinary program | 3 wk, with 5 d/ wk and 8-h sessions, total of 120 h | This inpatient program integrated physical exercises, ergonomic training, psychotherapy, patient education, behavioral therapy, and workplace-based interventions on an individual basis and in group sessions |
| Moss-Morris et al61 | Chronic pain         | 76               | Mean, 42.4 (9.49)               | Mean, 7.05 y (6.88)                                  | Multidisciplinary program | 4 wk, 5 d a week, 7 h/d | The emphasis of the outpatient care program was on reactivation and included components of graded goal directed exercise, relaxation, pain education, goal setting as well as information and therapeutic suggestions on specific issues such as sleep and mood management. Specific sessions in the psychoeducation component of the program address “ways of thinking,” “stress,” and “fear and avoidance” using cognitive restructuring techniques that focus on anxious or catastrophic thinking that inhibits reactivation |
| Persson et al62     | Chronic pain         | 555              | Mean, 40 (9.5)                  | Median, 217 wk (120-343)                             | Multidisciplinary program aiming at improving pain management strategies, with an overall goal to increase participation in society at large | 5 wk, and 2 d of follow-up 2 mo after discharge | The team offered education about pain and pain-related topics, as well as homework. In addition to the rehabilitation plan, all participants specified their most important everyday occupational problems, further targeted during the activity training. The group-based treatment enabled participants to share useful pain strategies with each other |

(Continued)
TABLE 2. (continued)

| References            | Population       | Diagnosis  | Participants | Mean Age (SD), Median (IQR) | % Female | Duration of Pain/Disability Mean (SD); Median (IQR) | Intervention Profile | Intervention Time | Intervention Details                                                                 |
|-----------------------|------------------|------------|--------------|-----------------------------|----------|------------------------------------------------------|----------------------|------------------|--------------------------------------------------------------------------------------|
| Ruscheweyh et al63    | Chronic pain     | 65         | Mean, 49.3 (12.3) | 74                      | Mean, 8.0 y (8.5) | Multidisciplinary program | 4 wk, outpatient, > 100 therapy hours | 6 wk             | The program consists of medical therapy, psychological therapy, physical therapy, art therapy, and patient education |
| Trief & Yuan64        | CLBP             | 132        | NR           | 51                      | Mean, 4 y       | Multidisciplinary program | 2-wk program (10 d), including 100 h including pretreatment and 2 follow-ups | 4 wk, daily        | This inpatient program contained both physical and occupational therapy. |
| Van Hooff et al65     | CLBP             | 524        | Mean, 45 (9.6) | 58                      | Mean, 13 y (10.8) | Multidisciplinary program |                                                 |                  | Intensive inpatient program with combined physical and psychological (CPP) program and included a cognitive behavioral approach in collaboration with the spine surgeons |
| Vendrig et al66       | CBP              | 120        | Mean, 41.3 (9.0) | 35                      | Mean, 47.6 mo (37.6) | Functional restoration program with the aim of restoring a normal pattern of daily functioning, including a complete return to work. On the basis of the functional restoration approach | Outpatient treatment was given daily and consisted of group sessions, which included back school, discussion of deep-rooted beliefs about symptoms and disabilities, and education on stress management. The physical training occurred according to operant learning principles (graded activity) and activities such as swimming and squash were also part of the program. The occupational therapist assisted the patient in the process of returning to work. The clinical psychologist provided group sessions in which an eclectic approach was adopted to identify and modify maladaptive behaviors, enhance adequate coping skills, and improve emotional awareness |
| Vendrig et al66       | CBP              | 120        | Mean, 41.3 (9.0) | 35                      | Mean, 47.6 mo (37.6) | Functional restoration program, aimed at achieving a normal pattern of functioning, including return to regular work. Decrease of pain or improvement of pain coping were not the direct aims of the program | Outpatient treatment was given daily and consisted of group sessions, which included back school, discussion of deep-rooted beliefs about symptoms and disabilities, and education on stress management. The physical training occurred according to operant learning principles (graded activity) and activities such as swimming and squash were also part of the program. The occupational therapist assisted the patient in the process of returning to work. The clinical psychologist provided group sessions in which an eclectic approach was adopted to identify and modify maladaptive behaviors, enhance adequate coping skills, and improve emotional awareness |
| Verkerk et al68       | CNLBP            | 1760       | Mean, 40.1 (10.6) | 74                      | Mean, 7.7 y (8.8) | Multidisciplinary program | 2 mo, 16 sessions of 3 h (a total of 48 h) + 3 mo self-supporting activity |                  | The outpatient rehabilitation program was coached by a multidisciplinary team (physical therapist, physician, health scientist and psychologist). Behavioral principles were applied to encourage patients to adopt adequate normal behavioral movement aimed at physical recovery |

*Intervention group.
†Not included in synthesis.
CBP indicates chronic back pain; CDOD, chronic occupational musculoskeletal disorders; CLBP, chronic low back pain; CNLBP, chronic nonspecific low back pain; CWP, chronic widespread pain; IQR, interquartile range; NR, not reported.

Pain Duration
The association between pain duration before MDR and physical functioning was assessed in 8 studies43,54,55,62,63,65,68,69 including a total of 3800 participants. Four of 8 studies54,55,62,69 reported no association with outcome, 2 studies showed a negative association,63,65 and 2 studies43,68 reported conflicting results on multiple outcome measures, showing either no association or a negative association in favor of short duration (Table 4).
Five studies (3 low, 1 moderate, 1 high RoB) were included in a meta-analysis (n=2978). The pooled OR (95% CI) showed no association with physical functioning; that is, the results...
indicate pain duration at baseline is not a prognostic indicator for outcome, OR = 0.97; 95% CI, 0.93-1.00; P = 0.08 (Fig. 3B).

Sensitivity Analyses and LoE (GRADE)

The sensitivity analyses for both pain intensity and pain duration showed that our results remained robust when examining the influence of study quality, follow-up time, measurement instruments, uni/multivariate analyses, and when compared with a fixed-effects model. The GRADE analyses of pain intensity as well as pain duration showed that, due to downgrading as a result of “inconsistency of the results,” there is evidence of moderate quality that baseline pain level and pain duration cannot predict physical functioning at ≥ 6-month follow-up of MDR (Table 7).

Physical Function–related Factors

The association between baseline and follow-up physical functioning was assessed in 15 studies (n = 4868). Physical function was assessed either by patients’ actual performance of physical tests (and evaluated by therapists) — or by patients’ own reporting of their function, activities, or disability, that is completing questionnaires (PROMs). The factors were divided into 2 groups and analyzed separately due to the qualitative differences of the assessment methods (Table 5).

Performance-based Physical Factors

Two studies investigated 6 performance-based physical factors (n = 783). The tests evaluated isometric endurance, mobility, and aerobic capacity as prognostic factors. The narrative analyses indicated no prognostic value for outcomes related to physical function, both studies reported no significant association. Both studies were rated as having high RoB. Because of limited data, a meta-analysis was not appropriate.

Self-reported Function, Activities/Disability

Fourteen studies examined the association between self-reported physical functioning and outcome (n = 4706). The narrative analyses of self-assessed physical function revealed inconclusive results. Higher levels of function at baseline were significantly associated with a positive outcome in 6 studies, while low levels of function associated with a positive outcome were reported in one study and no significant association was reported in another one study. However, 6 studies presented inconclusive results depending on measures used, either showing an inconsistency between a positive association and no association (3 studies) or between a negative association and no association (3 studies).

Eight studies (5 low, 2 moderate, 1 high RoB) were included in a meta-analysis (n = 3444). The pooled OR (95% CI) showed that high baseline function was associated with positive outcome, OR = 1.07; 95% CI, 1.02-1.13; P = 0.01 (Fig. 4).

Sensitivity Analyses and LoE (GRADE)

The results of self-reported physical function remained robust when excluding high RoB studies, and were independent of a fixed or random model. However, when analyzing the 3 studies with shorter follow-up times, there was no longer any significant association between physical function at baseline and outcome. Moreover, in studies with univariate analysis only, the associations disappeared as well.

The Grade synthesis showed there was no evidence (−) of prognostic value of performance-based physical function and that there was low evidence (+++) of a small effect of self-rated initial high physical functioning as prognostic for good physical functioning at follow-up after MDR (Table 7). Downgrading was due to “study limitations” and “inconsistency of the results.” For performance-based physical function, the initial GRADE LoE was set at +++ due to unclear study phases.

Psychological Factors

Seventeen studies investigated baseline psychological factors. Of these, most were categorized as either emotional factors or cognitive behavioral factors. For the purpose of analyses, cognitive-behavioral factors were divided into protective factors or risk factors. A few remaining factors, mostly relating to personality traits, were considered too compound or dissimilar and were therefore not synthesized in this context.

Emotional Factors

Fifteen studies investigated emotional factors relating to mood/distress, for example, depression and anxiety and their association to physical functioning at follow-up. The narrative analyses showed inconclusive results concerning their prognostic value. Six studies investigating emotional factors relating to mood/distress, for example, depression and anxiety and their association to physical functioning at follow-up.
did not demonstrate any significant associations; 6 studies showed differing results between anxiety and depression, 2 studies showed that low levels of depression/anxiety at baseline could predict positive results at follow-up, while 1 study showed some degree of initial anxiety/depression was associated with a positive outcome. Anxiety and depression were analyzed both separately and in combination with each other (Table 6). Eight studies (5 low, 3 high RoB) with continuous data were included in a meta-analysis (n = 3483). The pooled OR

**TABLE 3. Risk of Bias (RoB) Ratings of the Included Studies, Assessed With the Quality in Prognostic Studies (QUIPS)-tool**

| References | Study Participation | Study Attrition | Prognostic Factor Measurement | Outcome Measurement | Study Confounding | Statistical Analysis and Reporting |
|------------|---------------------|-----------------|------------------------------|---------------------|------------------|-----------------------------------|
| Angst et al | Low                 | Moderate        | Low                          | Low                 | Low              | Low                               |
| Bendix et al | Moderate            | High            | Low                          | Moderate            | Low              | Low                               |
| Bergstrom et al | Low | Low | Low                           | Low                 | Moderate         | Low                               |
| Bergström et al | Low  | Moderate | Low                         | Moderate            | Moderate         | Moderate                          |
| Bremanster et al | Low  | Moderate | Low                         | Low                 | Low              | Low                               |
| Ciechanowski et al | Moderate | Low | Low                           | Low                 | Low              | Low                               |
| de Rooij et al | Low                 | Moderate        | Low                          | Low                 | Low              | Low                               |
| Dobkin et al | High               | Moderate        | Moderate                      | Moderate            | Moderate         | Moderate                          |
| Farish et al | Moderate            | Moderate        | Low                          | Low                 | Low              | Low                               |
| Gerdie et al | Moderate            | High            | Low                          | Low                 | Moderate         | Low                               |
| Glattacker et al | Moderate | Moderate | Low                         | Low                 | Low              | Low                               |
| Harkapaa et al | Moderate     | Moderate | Low                         | Low                 | Low              | Low                               |
| Lennstra & Ouyama et al | Low | Moderate | Low                         | Moderate            | Moderate         | Low                               |
| Lillefjell et al | Moderate         | High            | Moderate                      | Low                 | Moderate         | Low                               |
| Lünning-Bergsten et al | Low | Moderate | Low                         | Moderate            | Moderate         | Low                               |
| McGeary et al | Low                 | Moderate        | Low                          | Low                 | Low              | Low                               |
| Moradi et al | Moderate            | High            | Low                          | Moderate            | High             | Moderate                          |
| Moss-Morris et al | Low  | Moderate | Low                         | Low                 | High             | Moderate                          |
| Persson et al | Moderate            | Low             | Low                          | Low                 | Low              | Low                               |
| Ruscheweyh et al | Low  | High   | Low                         | Low                 | Moderate         | Low                               |
| Trief & Yuan | Moderate            | Moderate        | Moderate                      | Moderate            | Moderate         | High                              |
| Van Hooff et al | Low                 | Low             | Low                          | Low                 | Low              | Low                               |
| Vending et al | Moderate            | Low             | Low                          | Moderate            | High             | Moderate                          |
| Vending et al | Moderate            | Low             | Low                          | Low                 | Low              | Low                               |
| Verkerk et al | Low                 | Moderate        | Low                          | Low                 | Low              | Low                               |

Categorization of RoB on study level and between studies:

- **Low**: all domains are classified as having Low RoB, or up to one Moderate RoB.
- **Moderate**: mainly low RoB-domains and up to two moderate RoB.
- **High**: ≥ one domain with high RoB or ≥ 3 moderate RoB.
### TABLE 4. Narrative Analyses of Pain-related Factors

| References          | Risk of Bias | Instrument | Direction | MA* | Pain intensity | Pain Duration |
|---------------------|--------------|------------|-----------|-----|----------------|---------------|
| Angst et al41       | Low          | SF 36-BP   | +         |     |                |               |
| Bendix et al46      | High         | VASBack pain | -         |     |                |               |
|                     |              | VASLeg pain |           |     |                |               |
| Bremander et al52   | Low          | VAS        | 0 No excl |     |                |               |
| Ciechanowski et al51| Low          | VAS        | + Yes     |     |                |               |
| de Rooij et al52    | Low          | NRS        | 0 Yes     |     |                |               |
| Dobkin et al53      | High         | MPQ-VAS    | 0         |     |                |               |
| Farin et al54       | Moderate     | VAS        | - a       |     |                |               |
|                     |              | VAS        | - b       |     |                |               |
| Gerdle et al55      | High         | NRS        | 0 g       |     |                |               |
|                     |              | MPIpain severity | 0 f     |     |                |               |
| Glattacker et al56  | Low          | VAS        | 0 a       | -  | Yes            |               |
|                     |              | VAS        | 0 b       |  0 |                |               |
|                     |              | VAS        | 0 c       |  0 |                |               |
| Lemstra & Olszynski60| High         | VAS        | 0         |     |                |               |
| Lillefjell et al57  | High         | VAS        | - Yes     |     |                |               |
| McGeary et al59     | Low          | VAS        | -         |     |                |               |
| Persson et al52     | Low          | MPIpain severity | 0 a     |     |                |               |
|                     |              |             |           |     |                |               |
|                      |              |             |           |     |                |               |
| Ruscheweyh et al52  | High         | NRS        | 0         | -  | Yes            |               |
| van Hooff et al55   | Low          | VAS        | 0         | Yes | 0 j            | Yes           |
| Verkerk et al56     | Low          | VAS        | 0         | Yes |                |               |

* + indicates favors high levels of pain intensity/duration at baseline; 0, no association between pain intensity/duration and positive outcome; −, favors low levels of pain intensity/duration at baseline.

** Bold indicates multivariate analyses.**

* Included in meta-analyses.

Details on multiple outcome measures: *OIDI, 1SF-12PCS, 2SF-36/MPI-Interference, 3SF-36 PF, 4SF-36 RP, 5ODI, 6COPM-performance, 7COPM-satisfaction, relative change, absolute change.

BP indicates Bodily Pain; COPM, Canadian Occupational Performance Measure; subscales Performance and Satisfaction; MPI, Multidimensional Pain inventory; MPI-Pain severity, subscale of MPI; MPQ-VAS, McGill Pain Questionnaire; Visual Analogue Scale for pain rating; NRS, Numeric Rating Scale; ODI, Oswestry Disability Index; Pain duration was measured by selfreport/questionnaires; PCS, Physical Health Summary Scales; PF, Physical Functioning; RP, Role-Physical; SF-12, 12-Item Short Form Health Survey; SF-36, 36-Item Short Form Health Survey; VAS, Visual Analogue Scale.
Cognitive and Behavioral Factors—Protective Factors

Nine studies (n = 2288) examined various cognitive and behavioral factors relating to self-efficacy, control beliefs, and health optimism; factors commonly attributed to strengthening a person’s resilience, that is with protective effects. The narrative analyses showed diverse results. Three studies found no association from 6 examined protective factors and 1 study showed a negative association, indicating low levels of 1 factor was associated with a positive outcome.

Four studies (3 low, 1 moderate RoB) were included in a meta-analysis (n = 1392). The pooled OR (95% CI) showed, contrary to the narrative analysis, an association between low levels of cognitive and behavioral risk factors and a positive outcome, OR = 0.77; 95% CI, 0.65-0.92; P = 0.003 (Fig. 5A).

Results identified 20 items with no association and 9 in favor of low levels for a positive outcome.

Six studies (2 low, 3 moderate, and 1 high RoB) were included in a meta-analysis (n = 1173). The pooled OR (95% CI) showed, contrary to the narrative analysis, an association between low levels of cognitive and behavioral risk factors and a positive outcome, OR = 0.85; 95% CI, 0.77-0.93; P = 0.0008 (Fig. 5C).

Cognitive and Behavioral Factors—Risk Factors

Eleven studies (n = 4068) examined the association between various “negative” cognitive and behavioral factors and outcome, that is potential risk factors. These were related to illness and self-efficacy beliefs, fear-avoidance beliefs and behavior, catastrophizing, and dimensions of somatic discomfort/somatization.

The narrative analyses of cognitive and behavioral risk factors indicated a majority of nonsignificant associations.
| Reference | Research Design | Performance-based physical function | Self-reported function |
|-----------|----------------|-------------------------------------|------------------------|
| Anjet et al. | Low | Mobility, muscle endurance, back muscle endurance, aerobic capacity | SF-36 PT, Sports training |
| Bondix et al. | High | ADL, Oswestry Disability Index, Sport activities | + |
| Ciechanowski et al. | Low | RMDQ | +, Yes |
| de Reuš et al. | Low | MRI, MRI productivity | - |
| Dobbin et al. | High | FD | + |
| Gordis et al. | High | SF-36 PF, SF-36 RP, ODI | +, Yes |
| Glattarter et al. | Moderate | SF-36 PF, SF-36 RP | +, Yes |
| Harbapaa et al. | Moderate | FCI | - |
| Lemme & Olayemi | High | FCI | - |
| Lisejürgen et al. | High | VAS, VAS phys. capacity | + |
| Montani et al. | High | Storti Sorensen test, VAS | SF-36 PCS |
| Veen-Merrie et al. | High | COPM | + |
| Persson et al. | Low | COPM | + |
| van Noord et al. | Low | ODI | +, Yes |
| Verbak et al. | Low | SF-36 PCS, SPPED | + |

+ indicates favors high levels of function at baseline; 0, no association between level of function and positive outcome; − favors low levels of function at baseline.

Bold indicates multivariate analyses.

*Included in meta-analyses.

Details on multiple outcome measures: ADL-score/Disability index, Low Back Pain Rating scale; B200 Isostation, physical performance test—back extension strength; Biering-Sørensen test, physical performance test—back muscle performance test; COPP/WONCA, Functional Health Status measurement (Darmouth COOP Functional Health Assessment Chart/World Organization of Family Doctors); COPM, Canadian Occupational Performance Measure; DRI, Disability Rating Index; FCI, Functional capacity index (based on RMDQ); FFbH-R, Hannover Functional Ability Questionnaire (German); FIQ, Fibromyalgia Impact Questionnaire; MBSE, Maximal Isometric Strength Extension (Trunk muscle performance test); MPI, Multidimensional Pain inventory; MPI-general activity, subscale of MPI; MPI-Interference, subscale of MPI; MPQ, McGill Pain Questionnaire; MVAS, The Million Visual Analogue Scale; NA, not available; NHP, Nottingham Health Profile, subscale: physical ability; NRS, Numeric Rating Scale; ODI, Oswestry Disability Index; Oesch test, physical performance test—arms strength; PCS, Physical Health Summary Scales; PDI, Pain Disability Index; PF, Physical Functioning; QBPDS, Quebec Back Pain Disability Scale; RMDQ, Roland Morris-Disability Questionnaire; RP, Role-Physical; SF-12, 12-Item Short Form Health Survey; SF-36, 36-Item Short Form Health Survey; VAS, Visual Analogue Scale; Villiger test, Physical performance test—Step test.
data and when compared with a fixed effects model. All in all, sensitivity analyses of the psychological factors clearly showed that the results were robust.

In summary, based on a GRADE analysis of these results including sensitivity analyses, the results showed that (a) there is moderate quality evidence that low initial emotional distress predicts a positive outcome on physical functioning at follow-up after MDR, (b) there is moderate quality evidence that high levels of protective cognitive behavioral factors predict a positive outcome of physical functioning at follow-up after MDR, and (c) there is moderate quality evidence that low levels of cognitive behavioral risk factors predict a positive outcome (Table 7). Downgrading was due to “study limitations” (a, c) and suspected “publication bias” (b).

**DISCUSSION**

**Summary of the Results**

To synthesize the evidence on prognostic factors for long-term (≥6 mo) physical functioning in patients with chronic musculoskeletal pain after MDR treatment, we examined 25 studies (n = 9436) that included a total of 87 potential prognostic factors relating to initial pain and physical and psychological functioning.

The key finding of this review confirmed that pretreatment psychological factors as well as physical function/disability are important prognostic indicators of functional outcome after MDR while common pain variables did not appear to provide evidence on prognosis.

Regarding psychological factors, results showed a moderate LoE that low levels of emotional distress, high levels of cognitive and behavioral protective factors, and low levels of cognitive and behavioral risk factors predicted a better physical functioning in long-term follow-up. Moreover, results showed a low LoE that high levels of self-reported physical function predicted better physical functioning. Our results also indicated, with moderate levels of evidence, that pain severity and pain duration did not predict physical functioning after MDR in patients with chronic musculoskeletal pain at least 6 mo after treatment.

**Comparison With Previous Reviews**

**Pain Factors**

Our study found that pain severity and pain duration did not have any prognostic value (moderate LoE), indicating that pretreatment information on pain per se is not informative for the further clinical course, at least not where physical function is concerned. The review of van der Hulst et al,20 also reported that pain duration lacked prognostic value. But contrary to our study, they found evidence that higher pain intensity was associated with worse outcome. However, this conclusion was based on only 2 articles, one of which is included in our study.48 While the other study included findings on a dissimilar subgroup of population, intervention, and outcome. On the other hand, the review of de Rooij et al19 reported the opposite, that is high pain intensity being associated with a better outcome, though this conclusion was based on only 1 study. In previous reviews22,24 that have investigated prognostic ability in earlier phases of pain chronicity (acute and subacute), pain variables presented with evidence of a negative impact on outcome. In our results, however, pain ratings were not significantly related to the outcome, in this case physical functioning, although the direction of the association was in accordance to these previous results, maybe indicating a less prognostic value over time.

**Physical Factors**

In the synthesis we differentiated between objectively measured performance-based and self-assessed physical functioning. The assessment of performance-based function was only investigated in 2 studies, and showed no association, which is in line with van der Hulst et al.20 Moreover, the study of Wessels et al,21 which investigated the association of changes in physical performance factors with improvement in disability, also reported that there was no association with outcome. Further research is needed to elucidate the topic, to investigate whether more objectively measured dimensions
### TABLE 6. Narrative Analyses of Psychological Factors

| References            | RoB  | Emotional distress | CB protective | CB Risk |
|-----------------------|------|--------------------|---------------|---------|
|                        |      | Instrument | Direction | MA*     | Instrument | Direction | MA*     | Instrument | Direction | MA*     |
| Angst et al<sup>11</sup> | Low  | HADS A    | 0         |          |          |          |          | CSQ      | 0         |          |
|                        |      | HADS D    | +         |          |          |          |          |          |           |          |
| Bremander et al<sup>12</sup> | Low  | HADS A    | +         |          |          |          |          | CSQ      | 0         | Yes      |
|                        |      | HADS D    | +         |          |          |          |          |          |           |          |
| Ciechanowski et al<sup>13</sup> | Low  | CES-D    | 0         | Yes      |          |          |          | CSQ      | 0         | Yes      |
| de Rooij et al<sup>15</sup> | Low  | HADS A    | -         |          |          | IPQR: IC | 0         | CSQ      | 0         |          |
|                        |      | BDI-II   | 0         | Yes      |          | IPQR: TC | 0         | PCI      | 0         |          |
|                        |      |          |           |          | IPQR: PC | +         | IPQR:ER  | 0         |          |
|                        |      |          |           |          |          | IPQR:TL | 0         |          |           |          |
|                        |      |          |           |          | SCL90    | 0         | IPQR: TLC| 0         |          |
| Dobkin et al<sup>19</sup> | High | CES-D    | 0         |          |          | ASES     | 0         | TSK      | 0         |          |
| Farin et al<sup>14</sup> | Moderate |          | | | LoC:Internal | 0<sup>6</sup> | | LoC:External | 0<sup>6</sup> | |
|                        |      |          |           |          | IPQR:IC  | 0<sup>6</sup> | | FABQ<sup>a</sup> | 0<sup>6</sup> | |
| Gardle et al<sup>16</sup> | High  | HADS A    | 0         |          | CPAQ AE  | 0         |          | TSK      | 0         |          |
|                        |      | HADS D    | 0         |          | CPAQ PW  | 0         |          |          |           |          |
|                        |      | MPI      | 0         |          | MPI-LC   | 0         |          |          |           |          |
|                        |      |           |           |          | SF36-MCS | 0         |          |          |           |          |
| Glaistacher et al<sup>17</sup> | Moderate | SF-36 MH      | 0<sup>14,5</sup> | | IPQR: IC  | 0<sup>6</sup> | | IPQR: ER  | 0<sup>6</sup> | Yes |

(Continued)
### TABLE 6. (continued)

| *References* | *Multivariate Analyses* | *Baseline* | *Outcome* | *COPM-performance* | *COPM-satisfaction* | *Activity level* | *QBPDS* | *Horse* | *COPM-performance* | *COPM-satisfaction* | *Activity level* | *QBPDS* | *Baseline* | *Outcome* | *COPM-performance* | *COPM-satisfaction* | *Activity level* | *QBPDS* |
|--------------|-------------------------|------------|-----------|---------------------|---------------------|---------------------|---------|---------|---------------------|---------------------|---------------------|---------|------------|-----------|---------------------|---------------------|---------------------|---------|---------|
| + indicates favors high levels at baseline; 0, no association between psychological factor and positive outcome; −, favors low levels at baseline. Bold indicates multivariate analyses.

*Included in meta-analyses.
†Excluded from MA, due to dichotomized prognostic factor- outlier.

Details on multiple outcome measures:  aSF-12 PC, bODI, cSF-36 PF, dSF-36-RP, eODI, fCOPM-performance, gCOPM-satisfaction, hActivity level, iQBPDS, jMISE trunk muscle performance, krelative change, labsolute change.

ASES indicates Arthritis Self-Efficacy Scale; BDI, Beck Depression Inventory; BDI-II, Beck Depression Inventory (II); BRQ, Beliefs about Rehabilitation Questionnaire; CES-D, Center for Epidemiological Studies-Depression Scale; COPM, Canadian Occupational Performance Measure; CPAQ, Chronic Pain Acceptance Questionnaire; CSQ, Coping Strategies Questionnaire; DGSS, Dutch General Self-Efficacy Scale; DRI, Disability Rating Index; EXPECT RTW, Perceptions of prognosis on Return to Work; FABQ, Fear-Avoidance Beliefs Questionnaire; FCI, Functional Capacity index (based on RMDQ); FFI-H-R, Hannover Functional Ability Questionnaire (German); FFIQ, Fibromyalgia Impact Questionnaire; HADS-A, Hospital Anxiety and Depression Scale subscale Anxiety; HADS-D, Hospital Anxiety and Depression Scale subscale Depression; Health expectations, single question, not specified instrument; Health Optimism, Health Optimism brief Scale; IPQ-R, Illness Perceptions Questionnaire-Revised; KKG, Control beliefs Concerning Illness and Health (German); LoC, Locus of Control beliefs, from Health+Pain Locus of Control Scales; MCS, Mental Component Summary; MH, Mental Health; MMPI, Minnesota Multiphasic Personality Inventory; MPI, Multidimensional Pain Inventory; MPI-Affective distress subscale of MPI; MPI-Life control subscale of MPI; ODI, Oswestry Disability Index; PCI, Pain Coping Inventory; PCS, Physical Health Summary Scales; PCS, Pain Catastrophizing Scale; PDI, Pain Disability Index; PF, Physical Functioning; PSEQ, Pain Self-Efficacy Questionnaire; QBPSD, Quebec Back Pain Disability Scale; RMDQ, Roland Morris-Disability Questionnaire; RP, Role-Physical, RSQ, Relationship Scale Questionnaire; SCL-90, Symptom Checklist-90; SF-12, 12-Item Short Form Health Survey; SF-36, 36-Item Short Form Health Survey; Stages of change, Pain stages of change questionnaire; TSK, Tampa Scale for Kinesiophobia; ZSDS, Zung Self-Rating Depression Scale. |
of physical functioning could have a prognostic value for outcome. On the other hand, self-assessed physical functioning emerged as a major outcome topic, and proved valuable in predicting outcome. We found, with low levels of evidence, that self-assessed physical function predicts physical functioning 6 mo after MDR. Our meta-analysis strengthened the results from the qualitative analyses of van der Hulst et al,\textsuperscript{20} where it was found that self-assessed physical functioning could predict physical functioning. Also, as the findings were reproduced in a mixed-diagnosis chronic pain population—instead of a more homogenous chronic low back pain population—the generalizability of the findings increased. However, the reasons for the inconsistency in reported direction of the association (either favoring higher or lower baseline status), which were also noted by van der Hulst and colleagues, need to be further examined.

FIGURE 5. Psychological factors: A, Forest plot of comparison between baseline emotional distress and association with positive outcome. B, Forest plot of comparison between baseline levels of cognitive behavioral protective factors and association with positive outcome. C, Forest plot of comparison between baseline levels of cognitive behavioral risk factors and association with positive outcome. The assessment measures for outcome and prognostic factor (PF) reported, type of analyses, and whether estimates (ORs) were combined from plural measures are presented in the footnotes. 5A: (1) Outcome: Activity level; PF: MMPI-Anx Pt; univariate; (2) Outcome: ODI; PF: Zung Self-rated Depression scale; univariate; (3) Outcome: COOP-WONCA; PF: HADS-A and HADS-D; multivariate. Combined OR; (4) Outcome: RMDQ; PF: CES-D; univariate correlation; (5) Outcomes: QBPDS and MISE; PF: Anxiety: MMPI-2 Pt, ANX and PBS and Depression: MMPI-2 D and DEP; multivariate. Combined OR; (6) Outcome: MPI Interference; PF: HADS-A and SCL-90 psychological functioning; multivariate and BDI-II, univariate. Combined OR; (7) Outcomes: QBPDS, relative and absolute recovery; PF: SF-36MCS; multivariate. Combined OR; (8) Outcome: MPI Interference; PF: MPI Affective distress; multivariate. 5B: (1) Outcome: MPI Interference; PF: DGSS, illness coherence, IPQ Personal control, IPQ Treatment control; uni- and multiv. Combined OR; (2) Outcome: FCI; PF: Health optimism; multivariate. (3) Outcomes: Coping Performance and Satisfaction; PF: MPI Life control; multivariate. Combined OR; (4) Outcome: ODI; PF: PSEQ self-efficacy; univariate. SC; (5) Outcome: RMDQ; PF: CSQ; univariate; (2) Outcomes: QBPDs, MISE; PF: MMPI-2 Hs, MMPI-2 HEA; multivariate. Combined OR; (3) Outcome: ODI; PF: TSK and PCS. Combined OR; (4) Outcome: FCI; PF: Other LoC; multivariate; (5) Outcome: ODI and SF-RP; PF: IPQ-R timeline acute-chronic, BRQ identity, BRQ process expectation; multivariate. Combined OR; (6) Outcome: MPI Interference; PF: IPQ-R; Timeline, Conseq., Emotional repr., Timeline cycl., PSQ, PCS, TSK; uni- and multiv. Combined OR. BDI indicates Beck Depression Inventory; CI, confidence interval; CopM, Canadian Occupational Performance Measure; COOP-WONCA, Coop Functional Health Assessment Charts HADS, Hospital Anxiety and Depression Scale; MISE, Maximal Isometric Strength Extension; MPI, Multidimensional Pain Inventory; ODI, Oswestry Disability Index; OR, odds ratio; PF, Physical Functioning; QBPDs, Quebec Back Pain Disability Scale; RMDQ, Roland-Morris Disability Questionnaire; RP, Role-Physical; SF-36, 36-Item Short Form Health Survey.
### TABLE 7. Summary of Findings and Overall Quality as Assessed With GRADE

| Domain                  | Potential Prognostic Factor | Total Number of Participants (No. Studies) | Total Number of Participants (No. Studies) | Estimated Effect Size (95% Confidence Interval)* | Phase | GRADE Factors | Study Limitations | Inconsistency | Indirectness | Imprecision | Publication Bias | Moderate/Large Effect Size | Dose Effect | Overall Quality (Level of Evidence) |
|-------------------------|-----------------------------|-------------------------------------------|-------------------------------------------|-----------------------------------------------|-------|---------------|------------------|---------------|-------------|-------------|-----------------|-------------------------------|-------------|--------------------------------------|
| Pain                    | Pain level                  | 8191 (16)                                 | 2676 (5)                                  | OR, 0.84 (0.65-1.07)                          | +++  | +             | 0                | —             | 0           | 0           | 0               | 0               | 0            | Moderate quality (+++)          |
|                         | Pain duration               | 3800 (8)                                  | 2978 (5)                                  | OR, 0.97 (0.93-1.00)                          | ++   | +             | 0                | —             | 0           | 0           | 0               | 0               | 0            | Moderate quality (+++)          |
| Physical                | Performance-based function | 783 (2)                                   | NA (0)                                    | NA                                            | +++  | —             | 0                | —             | 0           | 0           | 0               | 0               | 0            | Very low quality (−)            |
|                         | Self-reported function      | 4706 (14)                                 | 3444 (8)                                  | OR, 1.07 (1.02-1.13)                          | ++   | —             | 0                | —             | 0           | 0           | 0               | 0               | 0            | Low quality (++)                |
| Psychological           | Emotional functioning       | 4358 (15)                                 | 3483 (8)                                  | OR, 0.77 (0.65-0.92)                          | ++   | —             | 0                | 0             | 0           | 0           | 0               | 0               | 0            | Moderate quality (+++)          |
|                         | Cognitive and behavioral    | 2288 (9)                                  | 1392 (4)                                  | OR, 1.49 (1.17-1.90)                          | ++   | 0             | 0                | 0             | 0           | 0           | —               | 0               | 0            | Moderate quality (+++)          |
|                         | protective factors          |                                           |                                           |                                               |      |               |                  |               |             |             |                 |                 |              |                                      |
|                         | Cognitive and behavioral    | 4068 (11)                                 | 1173 (6)                                  | OR, 0.85 (0.77-0.93)                          | ++   | —             | 0                | 0             | 0           | 0           | 0               | 0               | 0            | Moderate quality (+++)          |
|                         | risk factors                |                                           |                                           |                                               |      |               |                  |               |             |             |                 |                 |              |                                      |

Significant estimates in bold style.

GRADE indicates Grading of Recommendations Assessment, Development and Evaluation; NA, not available; OR, odds ratio.
Psychological Factors

We found high levels of emotional distress predicted poor outcome, which is in line with previous assumptions and reports,19,70,71 however, there is a lack of consistent evidence.20 This is the first time it has been shown in a meta-analysis based on >3000 participants, and our results confirm the importance of patients’ emotional functioning for treatment outcome.

Cognitive and behavioral factors are implied to have an impact on treatment outcome19,20,70 and this was also confirmed by our results. These essential factors of the pain experience may both strengthen the ability to deal with chronic pain as well as hinder patients’ adaptation. The narrative analyses of cognitive behavioral risk factors indicated a majority of nonsignificant associations but the meta-analysis revealed them to be significant prognostic factors for a negative outcome. While addressing these factors is at the core of pain management in MDR, our results show that high levels on cognitive and behavioral risk factors are related to poorer functional outcome. This implies that our current best evidence practice may not be addressing the coping problems of these patients satisfactorily. Indeed, Morley et al72 pointed out that results of cognitive-behavioral therapy pain management programs are modest at best, and these results have led to calls for improvements in treatment models.73,74

High levels of cognitive and behavioral protective factors predicted a better level of physical functioning in long-term follow-up. The results confirm the importance of factors attributed to a person’s resilience in determining outcome. Indeed, in a recent publication, the importance of factors related to a positive affect has been lifted forward as one way to improve treatments for chronic pain.75 As psychological risk and protective cognitive and behavioral factors are not mutually exclusive, MDR treatment should focus on both lowering the psychological risk factors and enhancing the protective psychological factors.

The prognostic ability of the psychological factors with a negative bearing, emotional distress (OR = 0.77), and cognitive and behavioral risk factors (OR = 0.85), respectively, was somewhat lower compared with the prognostic ability of the psychological protective factors (OR = 1.49). This could be due to treatment effects, as in most MDR treatment programs the negative psychological factors are not mutually exclusive, MDR treatment should focus on both lowering the psychological risk factors and enhancing the protective psychological factors.

On the whole, as today’s management of chronic pain still gains only moderate effects, and the evidence to guide optimal treatment tailoring is limited, the importance of identifying prognostic indicators is of major clinical relevance. A prerequisite is that we are able to identify who is at risk of poor outcomes and who is most likely to benefit. Until now, no previous meta-analysis review studies have been conducted on this topic and, to our knowledge, this study is the first well-powered systematic review to summarize the available literature on prognostic factors specifically for this major patient group.

Methodological Considerations

The strength of this systematic review is that it synthesizes factors of importance for physical functioning, one of the main targeted outcomes of MDR, rather than exploring a single prognostic factor impact or a selected part of the chronic pain-population, for example based on diagnosis. The study takes its standing point from a pragmatic perspective, hypothesizing that some factors probably exist that are common for the chronic pain population in general, irrespective of initial pain diagnosis, that is generic factors of importance for treatment outcome. From a methodological point of view, a body of evidence derived from longitudinal and pragmatic cohort studies enables high confidence in the field of prognosis, in comparison to more selected experimental randomized controlled trial studies.47 On the other hand, attrition and confounding can limit the internal validity of observational studies. The way of creating high-level evidence by unifying these observational studies with systematic synthesis methods is therefore a strength of this study.

The interdisciplinary review team with expertise in all fields relating to the aim of the study enabled a precise study selection, which led to great confidence in the identification of both the population of interest and the intervention of interest. The team was generally in agreement during the study selection process, despite the heterogeneity of retrieved studies. Good inter-rater agreement was strived for in all selection steps and RoB ratings, by introducing every phase with a pilot.

Omitting gray literature is likely to have introduced some information bias; however, it would be too time consuming to also collect and deal with this type of spread-out information, which is often not reported in enough detail. Including only articles in English is a potential source for information bias as well; however, it was a necessity for maintaining the strictness and specificity during the scrutiny of the study selection process. In addition, some reporting biases, for example publication bias or selective reporting of outcomes or analyses, cannot be ruled out. Significant results have a greater chance of being made available. Still, we found many studies presenting nonsignificant results. We believe this was partly due to our broad review scope and an exploratory search strategy, which permitted a vast amount of material, independent of primarily targeted prognostic factors in the original research publications. We put great effort into using these, often nonsignificant, variables in our syntheses, either narratively or quantitatively if data were provided. This has hopefully led to adding power and reducing possible asymmetry. As the relatively small number of studies reporting on each comparison precluded a detailed and meaningful analysis of funnel plots for publication bias, we attempted to visually analyze the narrative tables for symmetry of significant versus nonsignificant reporting. For some results, for example, the synthesis of protective psychological factors, the effect emerged stronger in the meta-analysis, which could likely be a result of missing nonsignificant data.

The risk of selection bias may have been introduced in the initial screening of titles, which was performed by one reviewer instead of 2. However, it was necessary to reduce the recall volume resulting from the broad and sensitive search strategy, and this stage therefore dealt only with identifying titles that precluded inclusion. The following screening process had a robust arrangement with randomization of studies and independent teams constituted by a senior and junior researcher.

Other sources for limitations of the study results may arise if narrative and quantitative syntheses are based on incompatible study heterogeneity or low study quality. We aimed to provide a well-powered overview of potential prognostic indicators of various MDR outcomes—as a result heterogenous studies were included with regard to
types of pain conditions/regions and clinical settings. This was based on the premise that common prognostic factors for “the chronic pain disease itself” probably exist. Although unique in its kind, some loss of specificity is therefore a consequence and limit to this review. To the best of our ability, great effort was put into a sensible study selection and a coherent collating of our found predictors and outcomes, in the sense of minimizing incompatible (noncomparable) factors. We are thus confident that the study populations and study interventions constituted a sample in accordance with the pragmatic, wide selection of individuals with chronic pain that would normally participate in MDR. The same applies for the grouping of factors and outcomes, which were measured with various instruments; however, all with the intention of capturing dimensions of the same construct. Incompatible measures or measures with measurement properties considered to be too vague were not included in analyses. In the present study, the OR was used as the common index in the meta-analysis, although the OR has sometimes been criticized for its difficulty in interpretation. We stated in our study protocol that we will present associations between prognostic factors and outcome by means of OR, and this could enhance comparisons with future MAs. A random effects model was chosen for the statistical analyses, as it assumes and deals better with the anticipated heterogeneity.

Heterogeneity, measured by $I^2$, was generally high for almost all comparisons (range: 48% to 94%). Although $I^2$ indicated high heterogeneity, our attempts to investigate the source for these differences did not reveal any systematic reasons for the variance. Sensitivity analyses proved our results were in general robust. The direction of the associations remained stable and did not result in any major change of variation in the effect, except for the factor physical functioning. The effect estimates remained stable when comparing studies based on statistical analyses (univariate vs. multivariate) and study quality (low vs. high), and follow-up time (shorter vs. longer), although the statistical significance level occasionally decreased to nonsignificant for the emotional distress and cognitive and behavioral risk factors. Sometimes the effects of the prognostic factors seemed to be strengthened over time, when comparing shorter versus longer follow-up time (eg protective cognitive and behavioral factors), but the limited number of included studies in each meta-analysis did not permit further detailed moderator analyses of follow-up time or further aspects of clinical diversity.

Although our sensitivity analysis of potential factors influencing the stability of our results was generally stable, we cannot exclude true heterogeneity. With more unexplained variance across studies, some caution in the interpretation of the results was required and we therefore downgraded all pain and physical function domains in the GRADE, due to “inconsistency.”

Study quality, that is poor methodological quality may also impose limitations to the validity of study results, for example, low power, low attrition rates, or inadequate analyses are likely to affect the estimates and widen the 95% CIs in smaller studies. Our included studies were to a large extent of good methodological quality, with at least two thirds having low or moderate RoB. Still, “study limitations” was the most common reason for downgrading the GRADE. The measures for both outcomes and prognostic factors were mainly of “good” quality and statistical analyses were relevant but attrition and dealing with confounding were the weakest domains—which can seriously impact the results in prognostic factor studies. The assessment of study quality relies to a great extent to the level of relevant reporting. Often study quality was downgraded due to unclear detailing on, for example, study participation and attrition, which might not have been actual sources for bias. Moreover, for some RoB domains, the PABAK-OS was found to be unacceptably low. However, it was easy to obtain consensus on the overall RoB scores during the consensus discussions. On a general note, it was apparent that reporting has improved over the past decades, possibly as a result of the devise of reporting guidelines, for example the STROBE checklist. All in all, we believe our results have external validity and can be generalized within the context of the population and intervention of interest—still keeping in mind that our findings may apply to this specific outcome “Physical functioning” and possibly not to the other dependent variables that will be analyzed in subsequent reviews.

CONCLUSIONS

Physical functioning at 6 months or longer after MDR was not predicted by initial pain level or pain duration (chronicity), contrary to previous indications, and therefore should not be used for assumptions of treatment prognosis. Better physical functioning was predicted by high levels of initial self-assessed physical functioning. Furthermore, a better outcome was predicted by low levels of emotional distress and low levels of cognitive and behavioral risk factors, indicating that treatment should further target and optimize these modifiable factors. Finally, high levels of protective cognitive and behavioral factors were strong prognostic indicators of better physical functioning at 6 months or more after MDR, and an increased focus on positive, psychological protective factors may perhaps provide an opening for yet untapped clinical gains. The prognostic ability of the investigated factors may have been confirmed, but substantial heterogeneity between the studies was present and the effect sizes were in general fairly low, explaining only a limited part of the variance of outcome. Further research is naturally warranted to identify more important prognostic factors. Ultimately, this body of evidence can contribute to the development of clinical prediction models, which, in turn, will generate a basis for the future optimization of multidisciplinary biopsychosocial rehabilitation in chronic pain.

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