Effectiveness of implementing a best practice primary healthcare model for low back pain (BetterBack) compared with current routine care in the Swedish context: an internal pilot study informed protocol for an effectiveness-implementation hybrid type 2 trial

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ABSTRACT

Introduction Low back pain (LBP) is a major health problem commonly requiring healthcare. In Sweden, there is a call from healthcare practitioners (HCPs) for the development, implementation and evaluation of a best practice primary healthcare model for LBP.

Aims (1) To improve and understand the mechanisms underlying changes in HCP confidence, attitudes and beliefs for providing best practice coherent primary healthcare for patients with LBP; (2) to improve and understand the mechanisms underlying illness beliefs, self-care enablement, pain, disability and quality of life in patients with LBP; and (3) to evaluate a multifaceted and sustained implementation strategy and the cost-effectiveness of the BetterBack model of care (MOC) for LBP from the perspective of the Swedish primary healthcare context.

Methods This study is an effectiveness-implementation hybrid type 2 trial testing the hypothesised superiority of the BetterBack MOC compared with current routine care. The trial involves simultaneous testing of MOC effects at the HCP, patient and implementation process levels. This involves a prospective cohort study investigating implementation at the HCP level and a patient-blinded, pragmatic, cluster, randomised controlled trial with longitudinal follow-up at 3, 6 and 12 months post baseline for effectiveness at the patient level. A parallel process and economic analysis from a healthcare sector perspective will also be performed. Patients will be allocated to routine care (control group) or the BetterBack MOC (intervention group) according to a stepped cluster dogleg structure with two assessments in routine care. Experimental conditions will be compared and causal mediation analysis investigated. Qualitative HCP and patient experiences of the BetterBack MOC will also be investigated.

Dissemination The findings will be published in peer-reviewed journals and presented at national and international conferences. Further national dissemination and implementation in Sweden and associated national quality register data collection are potential future developments of the project.

Strengths and limitations of this study

▸ This will be the first study of effectiveness and implementation of a best practice model of care in low back pain primary care in Sweden.
▸ An international consensus framework is used for the development, implementation and evaluation of the BetterBack model of care.
▸ The main trial’s a priori methodology has been informed and refined by an internal pilot phase.

Background

Low back pain (LBP) is a prevalent and burdensome condition in Sweden and globally. LBP can be described by its location and by its intensity, duration, frequency and influence on activity. The natural course of LBP is often self-limiting, but a large majority experience pain recurrence and 20% may experience persistent symptoms. LBP is commonly categorised as non-specific, where a pathoanatomical cause cannot be confirmed through diagnostic assessment. Approximately <1%-4% of LBP cases in primary healthcare may show signs underlying malignancy, fracture, infection or cauda equina syndrome requiring medical intervention. Furthermore, neuropathic pain may be present in 5%-15% of cases. Medical imaging studies display a high prevalence of varying spinal morphology and degenerative findings in...
both symptomatic and non-symptomatic younger and older adults. This suggests that LBP is more typically a result of benign biological and psychological dysfunctions, as well as social contextual factors influencing the pain experience.

In Sweden, previous studies by our research group suggest the healthcare process for patients with LBP tends to be fragmented, with many healthcare practitioners (HCPs) giving conflicting information and providing interventions of varying effectiveness. Our studies have shown that only a third of patients on sick leave for musculoskeletal disorders receive evidence-based rehabilitation interventions in primary care. Furthermore our research has also demonstrated that there are still interventions that physiotherapists in primary care consider to be relevant in clinical practice despite the absence of evidence or consensus about the effects. Our preliminary data suggest that when patients with LBP are referred to specialist clinics, up to 48% have not received adequate evidence-based rehabilitation in primary care. There is therefore a strong case for change to address what care should be delivered for LBP and how to deliver it in the Swedish primary healthcare setting.

The development of best practice clinical guidelines aims to provide HCPs with recommendations based on strength of available evidence as well as professional consensus for the intervention’s risk and benefits for the patients. Best practice clinical guidelines for LBP are lacking in Sweden but have recently been developed by the Danish Health and Medicines Authority and the English National Institute for Health and Care Excellence. These national guidelines provide a thorough assessment of current evidence and can be used in Sweden to form the basis for locally adapted recommendations. Common to LBP, central recommendations from best practice clinical guidelines for arthritis are also education and exercise therapy aimed at improving patient self-care. Guideline-informed models of care (MOC) such as ‘Better Management of Patients with Osteoarthritis (BOA)’ in Sweden and ‘Good Life with Osteoarthritis’ in Denmark (GLA:D) have been successfully implemented with broad national HCP use. Furthermore, improvements in patient-reported pain, physical function and decreased use of pain medication after receiving these MOCs have been reported. A similar best practice MOC for LBP could potentially improve HCP evidence-based practice and patient-rated outcomes in the Swedish primary healthcare setting.

Recently an international consensus framework has been established to support the development, implementation and evaluation of musculoskeletal MOC. MOC intends to act on behavioural change mechanisms to attain specific behavioural targets. In order to achieve effective and efficient implementation of an MOC in primary healthcare, it is important to apply knowledge from implementation science. Implementation science is the scientific study of uptake of research findings and evidence-based practices into routine practice to improve the quality and effectiveness of healthcare and services. Implementation strategies focus on minimising barriers and maximising enablers that impact on the implementation and use of evidence-based practices. It has been suggested that a multifaceted strategy involving simultaneous use of several implementation strategies may be more effective than single-faceted strategies, but the evidence base is inconclusive. A recent systematic review however suggests that the most important aspects of successful implementation strategies are an increased frequency and duration of the implementation intervention and a sustained strategy.

There is therefore a clear rationale for evaluating the extent to which and how a best practice MOC for LBP (BetterBack) implemented with a sustained multifaceted strategy is potentially effective in the Swedish primary care context. The costs in relation to effects are important to consider in order to deliver healthcare efficiently. This article describes a protocol for a Better-Back MOC effectiveness and implementation process evaluation. The protocol conforms to the Standard Protocol Items: Recommendations for Interventional Trials guidelines, with a checklist provided in online supplementary file 1.

AIMS

The overall aim is to investigate the effectiveness and implementation process of the BetterBack MOC for LBP in a Swedish primary healthcare context. The specific trial objectives are to (1) improve and understand the mechanisms underlying changes in HCP confidence, attitudes and beliefs for providing best practice primary healthcare for patients with LBP, (2) improve and understand the mechanisms underlying change in illness beliefs, self-care enablement, pain, disability and quality of life in patients with LBP, and (3) evaluate a multifaceted and sustained implementation strategy and cost-effectiveness of the BetterBack MOC for LBP in the Swedish primary healthcare context.

HYPOTHESES

1. HCP-reported confidence, attitudes and beliefs for providing primary healthcare for LBP will show statistically significant improvement after a sustained multifaceted implementation of the BetterBack MOC compared with baseline before implementation. Intentional and volitional HCP-rated determinants of implementation behaviour regarding the BetterBack MOC will mediate improved confidence,
attitudes and beliefs in a causal effects model. This will correlate with more coherent care according to best practice recommendations.

2. The sustained multifaceted implementation of the BetterBack® MOC will result in more statistically significant and greater clinically important improvement compared with current routine care for LBP regarding patient-reported measures for illness beliefs, self-care enablement, pain, disability and quality of life. Improvements in illness beliefs and adequate patient enablement of self-care will mediate the effect on these outcomes.

3. A sustained multifaceted implementation of the BetterBack® MOC compared with current routine care will result in fewer patients with persisting LBP, fewer requiring specialist care, increased adherence to best practice recommendations and more statistically significant incremental cost-effectiveness ratio (ICER) based on cost per EuroQoL 5-Dimension Questionnaire (EQ-5D) quality-adjusted life years (QALY) gained.

METHODS

Study design

The WHO Trial Registration Data Set is presented in table 1. This study is an effectiveness-implementation hybrid type 2 trial testing the hypothesised superiority of the BetterBack® MOC compared with current routine care.29 The design involves an effectiveness evaluation of the BetterBack® MOC at the HCP and patient level, as well as a process evaluation of a sustained multifaceted implementation strategy conducted simultaneously. Evaluations are focused at the HCP and patient levels because the MOC is targeted at changing HCP behaviour, who then in turn implements behavioural change strategies at a patient level. This trial design was chosen for its potential to provide more valid effectiveness estimates based on pragmatic implementation conditions. This is in contrast to best or worst case implementation conditions common in traditional efficacy or effectiveness trials.30 31 Another advantage of the hybrid design is its potential to accelerate the translation of the MOC to real-world practice. This is in contrast to a time lag between efficacy, effectiveness and then dissemination steps in traditional research.29 The trial design is outlined in figure 1.

As outlined in table 2, the design at the HCP level involves data collection in the cohort before and prospectively after implementation of the BetterBack® MOC. At the patient level, data are collected in a single-blinded, pragmatic, randomised controlled, stepped cluster format with longitudinal follow-up at 3, 6 and 12 months post baseline. Randomisation at the patient level is not possible due to potential carry-over effects of the HCP transitioning back and forth between providing routine care or the BetterBack® MOC for different patients. Instead cluster randomisation is conducted at the start of the study, where patients are allocated thereafter to routine care (control group) or the BetterBack® MOC (intervention group) depending on the clinic’s allocation. Patients remain in their allocated group throughout the study.

A stepped cluster structure instead of a parallel structure of MOC implementation is applied due to the logistics involved in implementation in different geographical areas. The specific stepped cluster structure applied in the context of our study is classified as a dogleg with two assessments in routine care.30 31 The term ‘dog leg’ has been used by methodologists because the stepped structure resembles the form of a dog hind leg.32 As displayed in table 2, this involves the first cluster being assessed after the implementation of the BetterBack® MOC. The second cluster is assessed after a period of current routine care (control), and assessed again after the implementation of the BetterBack® MOC. The third cluster receives current routine care (control) throughout the trial. However, studying the implementation of the BetterBack® MOC in cluster 3 is planned to occur as a final step at the end of the study.

An advantage of using the dogleg structure with two assessments in routine care is that it allows for an internal pilot phase of initial implementation of the BetterBack® MOC in cluster 1 compared with clusters receiving current routine care. Another advantage is that data generated will still contribute to the final analyses to maintain trial efficiency.32 33 One objective for an internal pilot is to confirm the HCP acceptability of the intervention and trial within the first cluster.32 33 A progression criteria for continuing the trial requires that HCPs who have completed the BetterBack® education workshop rate on average a maximum of 2.5 out of 5 on the following determinants of implementation behaviour question: I expect that the application of BetterBack® MOC will be useful (1=agree completely to 5=do not agree at all).

Another objective of the internal pilot is to monitor patient recruitment in all three clusters during the first 2 months to provide information on the optimal cross forward time for cluster 2. In the dogleg design it is possible to vary the time point of cluster 2 to cross forward from the control to intervention condition if the patient recruitment process in either cluster 1 or 3 is more or less than expected in the internal pilot (see table 2). In the event that cluster 1 recruits less than expected and cluster 2 or 3 recruits more than expected, then cluster 2 will cross forward to the intervention condition immediately after the internal pilot. If cluster 1 recruits more than expected and cluster 2 or 3 recruited less than expected during the internal pilot phase, then cluster 2 will cross forward to the intervention condition later in the trial to allow adequate current routine care data collection. Clusters were expected to recruit and gather data for at least 20 patients with LBP per month in the internal pilot. A final objective with the internal pilot phase is to assess baseline variation and change over 3 months for implementation process and patient primary outcome...
Table 1  WHO Trial Registration Data Set

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<td>Contact for public queries</td>
<td>Allan Abbott, MPhysio, PhD (+46 (0)13 282 495) (<a href="mailto:allan.abbott@liu.SE">allan.abbott@liu.SE</a>)</td>
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measures to inform if our a priori sample size calculation needed to be revised in the continuation of the trial.

Study setting
The Östergötland public healthcare region has a total population of 453,596 inhabitants with approximately 5000 patients per year accessing primary care physiotherapy due to LBP. In the public healthcare region of Östergötland, a large majority of consultations for LBP are via direct access to the 15 primary care physiotherapy rehabilitation clinics. A smaller percentage of consultations are via referral to these rehabilitation clinics from the 36 primary healthcare general practices in the region. Therefore the focus of this study is on the physiotherapeutic rehabilitation process for LBP in primary care. The rehabilitation clinics form three clusters in Östergötland healthcare region. These clusters are based on municipal geographical area and organisational structure of the rehabilitation clinics, which help to minimise contamination between separate clusters of clinics (figure 2). Cluster west comprised 5 clinics with 27 physiotherapists, cluster central comprised 6 clinics with 44 physiotherapists, and cluster east comprised 6 clinics with 41 physiotherapists.

Eligibility criteria
Registered physiotherapists practising in the allocated clinics and regularly working with patients with LBP will be included in the study. These physiotherapists will assess the eligibility of consecutive patients before and after the implementation of the BetterBack☺ MOC based on the following criteria:

► Inclusion criteria: men and women 18–65 years; fluent in Swedish; and accessing public primary care due to a first-time or recurrent episode of acute, subacute or chronic-phase benign LBP with or without radiculopathy.

► Exclusion criteria: current diagnosis of malignancy, spinal fracture, infection, cauda equina syndrome, ankylosing spondylitis or systemic rheumatic disease, previous malignancy during the past 5 years; spinal surgery during the last 2 years; current pregnancy or previous pregnancy up to 3 months before consideration of inclusion; patients who fulfil the criteria for multimodal/multiprofessional rehabilitation for complex long-standing pain; and severe psychiatric diagnosis.
Table 2  Study design and schedule of enrolment, interventions and assessments

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0, control condition; 1, intervention condition; grey-shaded cells, internal pilot; T, assessment time. Period where 2-week crossover from control to intervention can occur dependent on patient recruitment rates identified in the internal pilot study. BIPQ, Brief Illness Perception Questionnaire; DIBQ, Determinants of Implementation Behaviour Questionnaire; EQ-5D, EuroQol 5-Dimension Questionnaire; HCP, healthcare practitioner; MOC, model of care; NRS, Numeric Rating Scale; ODI, Oswestry Disability Index; PABS-PT, Pain Attitudes and Beliefs Scale for physical therapists; PCS, Practitioner Confidence Scale; PEI, Patient Enablement Index; PGIC, Patient Global Rating of Change; PROMS, patient-reported outcome measures.
Interventions
Control condition: current routine physiotherapeutic care for LBP in primary healthcare

Patients attending rehabilitation clinic clusters that have not yet completed the implementation of the BetterBack© MOC will receive treatment as usual according to current routine care clinical pathways (figure 3). A clinical pathway specified in Östergötland public healthcare region requires that for patients accessing primary care due to LBP, a triage is to be performed by licensed HCPs (physiotherapists, nurses or general practitioners (GPs)), to triage for specific pathology of serious nature. These approximately 1%–4% of patients with suspected specific pathology of serious nature are then to be examined by GPs and referred for specific intervention in secondary or tertiary healthcare. The majority of patients with LBP who on initial triage are assessed as having benign LBP are then scheduled for physiotherapy consultation and implementation of an LBP management plan. If the patient has persistent functional impairment and activity limitation despite 2–3 months of primary care intervention, the clinical pathway specifies inclusion criteria for specialist care referral pathways (figure 3).

Figure 2 Municipal resident population and number of physiotherapy rehabilitation clinics and therapists in the west, central and east organisational clusters in Östergötland healthcare region.

Figure 3 Current routine care clinical pathway for LBP in Östergötland healthcare region. The primary care physiotherapy process outlined by the red square is the focus area for the implementation of the BetterBack© model of care for LBP. GP, general practitioner; LBP, low back pain.
Intervention condition: the BetterBack MOC for LBP

Development, design and implementation of the BetterBack MOC for LBP

A framework for the development of musculoskeletal MOC was used to guide the development of the BetterBack MOC for LBP. The high prevalence and burden of LBP, and the lack of evidence-based clinical practice guidelines and a call for a best practice MOC requested by physiotherapy clinic managers in the Östergötland healthcare region have been identified in the primary care of LBP. Therefore, a case for change has been justified to improve current physiotherapeutic health service delivery for the primary care of LBP. The content and structure of the BetterBack MOC were developed by engaging a work group of physiotherapy clinicians (clinical champions) from each primary care cluster in the Östergötland public healthcare region and physiotherapy academics at Linköping University. A Template for Intervention Description and Replication checklist is described in online supplementary file 2. To identify which key areas of contemporary care were of relevance for the BetterBack MOC, the following tasks were performed by the work group:

1. Discussion and outline of the current routine care clinical pathway for LBP and areas needing improvement: the work group concluded that the BetterBack MOC needed to focus on the following:
   - WHO/WHERE: the primary care physiotherapy process for the management of patients with LBP in Östergötland healthcare region outlined by the red square in figure 3.
   - Analysis and discussion of existing international best practice clinical guidelines: the following thorough and up-to-date systematic critical literature reviews and international clinical guidelines were analysed and discussed by the work group: refs 13–15.
   - 3. Adaptation of best practice clinical guidelines to the Swedish context: the development of evidence-based recommendations was based on the Swedish National Board of Health and Welfare methods for guideline construction. The overall grade of evidence together with a consensus position based on professional experience and patient net benefit versus harms and costs are the key aspects on which the work group has formulated local recommendations to reflect their strength.
   - The recommendations have been externally reviewed by local physicians and international experts from the University of Southern Denmark. A summary of the Östergötland healthcare region physiotherapeutic clinical practice guideline recommendations for primary care management of LBP with or without radiucopathy as well as the support tools used in the BetterBack MOC is provided in online supplementary file 3.

4. Considering potential barriers to the uptake of evidence-based recommendations by HCP, the work group identified and discussed targeted HCP behavioural change priorities of relevance for the BetterBack MOC. The work group discussion led to a rationale for the BetterBack MOC content and implementation described in table 3:

   - WHY: The main HCP target behaviour was the adoption of the BetterBack MOC to influence HCP delivery of care coherent with best practice recommendations.
   - WHAT: This would require the contents of the MOC to change impeding barrier behaviours such as low confidence in skills/capabilities for improving LBP patient management, a biomedical treatment orientation rather than a biopsychosocial orientation, and low awareness or beliefs of the negative consequences of the MOC.
   - HOW: BetterBack MOC content used to overcome the modifiable barriers includes support tools aimed at further education and enablement of HCP clinical reasoning in providing LBP assessment and treatment coherent with the Swedish adaptation of best practice clinical guidelines. The support tools include assessment proformas with associated instruction manual, clinical reasoning flow charts linking assessment findings to relevant treatment interventions, patient education brochures and group education material on LBP self-care, as well as a functional restoration programme (online supplementary file 3).
   - WHEN/HOW MUCH/TAILORING: The functional restoration programme and patient education components used, and their individual and group-based delivery and dosing, are individualised based on the HCP clinical reasoning of the type and grade of patients’ functional impairments and activity limitations (online supplementary file 3).
   - PROCEDURE: Figure 4 displays a flow diagram showing the steps involved for HCPs in delivering the contents of the BetterBack MOC.

The Behaviour Change Wheel (BCW) was used by the work group as a logic model to theorise the process of how the BetterBack MOC content applied at the guideline policy level could guide theory-informed intervention functions using specific behaviour change techniques.

To help investigate possible mediators of behavioural change interventions in the BetterBack MOC, the Theoretical Domains Framework (TDF) was integrated into the BCW. The TDF comprised 14 theoretical domains/determinants of behavioural change which could potentially influence behaviour change technique effect on the central source of behaviour. The central source of behaviour in the behaviour change wheel is described by the COM-B model. In the COM-B model, a person’s capability (physical and psychological) and opportunity (social and physical) can influence on motivation (automatic and reflective), enacting behaviours that can then alter capability, motivation and opportunity. The BCW and TDF are displayed in figure 5.

5. The following sustained multifaceted implementation strategy for the BetterBack MOC was developed:
<table>
<thead>
<tr>
<th>Target behaviour</th>
<th>Rationale based on barriers to be addressed</th>
<th>BetterBack® MOC content to overcome the modifiable barriers</th>
<th>BCT (^{44})</th>
<th>Functions</th>
<th>Mechanism of action</th>
<th>COM-B</th>
<th>TDF</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>BCT (^{44})</td>
<td>Functions</td>
<td>Mechanism of action</td>
<td>COM-B</td>
<td>TDF</td>
</tr>
<tr>
<td>Improved HCP confidence and biopsychosocial orientation in treating LBP through adoption of BetterBack® model of care</td>
<td>1. Low confidence in skills/capabilities for improving LBP patient management. 2. Use of a biomedical treatment orientation rather than a biopsychosocial orientation. 3. Low awareness of the model. 4. Beliefs of negative consequences of the model.</td>
<td>1. Multifaceted implementation strategy—workshop education. Evidence-based model of care and clinical implementation tools (see online supplementary files 1 and 2).</td>
<td>1.2 Problem-solving</td>
<td>Enablement</td>
<td>Psychological capability</td>
<td>Behavioural regulation</td>
<td>Psychological capability</td>
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<td>1.4 Action planning</td>
<td>Enablement</td>
<td>Psychological capability</td>
<td>Goals</td>
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<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>2. Feedback on behaviour</td>
<td>Training</td>
<td>Reflective motivation</td>
<td>Behavioural regulation</td>
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<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>3. Social support</td>
<td>Enablement</td>
<td>Social opportunity</td>
<td>Social influences</td>
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<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>4. Instruction on how to perform behaviour</td>
<td>Education</td>
<td>Psychological capability</td>
<td>Knowledge</td>
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<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>5. Information about social and environmental consequences</td>
<td>Persuasion</td>
<td>Social opportunity</td>
<td>Social influences</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>6.1 Demonstration of behaviour</td>
<td>Modelling</td>
<td>Psychological capability</td>
<td>Social influences</td>
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<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>6.2 Social comparison</td>
<td>Persuasion</td>
<td>Social opportunity</td>
<td>Social influences</td>
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<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>6.3 Information about other’s approval</td>
<td>Persuasion</td>
<td>Social opportunity</td>
<td>Social influences</td>
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<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>8.1 Behavioural practice/rehearsal</td>
<td>Training</td>
<td>Physical capability</td>
<td>Physical skills</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>8.7 Graded task</td>
<td>Training</td>
<td>Physical capability</td>
<td>Physical skills</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>9.1 Credible source</td>
<td>Persuasion</td>
<td>Reflective motivation</td>
<td>Reinforcement</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>9.2 Pros and cons</td>
<td>Persuasion</td>
<td>Reflective motivation</td>
<td>Beliefs about consequences</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>9.3 Comparative imagining of future outcomes</td>
<td>Enablement</td>
<td>Reflective motivation</td>
<td>Beliefs about consequences</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>13.2 Framing/reframing</td>
<td>Enablement</td>
<td>Psychological capability</td>
<td>Cognitive and interpersonal skills</td>
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<td></td>
<td></td>
<td>BetterBack® MOC content to overcome the modifiable barriers</td>
<td>15.1 Verbal persuasion about capability</td>
<td>Enablement</td>
<td>Psychological capability</td>
<td>Beliefs about capabilities</td>
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### Table 3  Continued

<table>
<thead>
<tr>
<th>Target behaviour</th>
<th>Rationale based on barriers to be addressed</th>
<th>BetterBack© MOC content to overcome the modifiable barriers</th>
<th>Mechanism of action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decreased patient LBP and disability, as well as improved patient enablement of self-care.</td>
<td>1. Maladaptive beliefs on the cause and course of LBP (illness perception)=low outcome expectation, anxiety, catastrophising, fear avoidance, illness beliefs. 2. Low belief in ability to control pain, low belief in ability to perform activities, low baseline physical activity.</td>
<td>1. BetterBack© part 1: individualised information at initial and follow-up visits. 2. BetterBack© part 1: patient education brochure. 3. BetterBack© part 2: group education.</td>
<td>Com-B</td>
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<td>Lay language pedagogical explanation of function impairment and activity limitation-related assessment findings and matched goal-directed treatment. Lay language education on the spine’s structure and function, natural course of benign LBP and advice on self-care. Pain physiology, biomechanics, psychological coping strategies and behavioural regulation.</td>
<td>Psychological capability</td>
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<td>5.1 Information about health consequences 9.1 Credible source 4.1 Instruction on how to perform behaviour 5.1 Information about health consequences 1.2 Problem-solving 3.1 Social support 4.1 Instruction on how to perform behaviour 4.3 Reattribution 5.1 Information about health consequences 6.1 Demonstration of behaviour 6.2 Social comparison 8.1 Behavioural practice/rehearsal 8.2 Behaviour substitution</td>
<td>Knowledge</td>
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<td>Education Persuasion Education Education Education Enablement Enablement Education Modelling Persuasion Training Enablement Persuasion Enablement Education</td>
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<td>Psychological capability Psychological capability Psychological capability Psychological capability Psychological capability Social opportunity Social opportunity Social opportunity Physical capability Physical capability Psychological capability Behavioural regulation</td>
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<td>Education Education Education Education Education Education Education Education Education Education Education Education Education</td>
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<td>Information about health consequences Demonstration of behaviour Social comparison Behavioural practice/rehearsal Behaviour substitution Credible source Comparative imagining of future outcomes Incentive (CME diploma) Social comparison Framing/reframing</td>
<td>Social influences</td>
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<td>Education Modelling Persuasion Training Enablement Persuasion Enablement Persuasion Enablement Enablement Persuasion Enablement</td>
<td>Social influences</td>
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<td>Psychological capability Psychological capability Social opportunity Physical capability Cognitive and interpersonal skills</td>
<td>Knowledge</td>
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<td>Psychological capability Psychological capability Behavioural regulation Reflective motivation Beliefs about consequences Reflective motivation Emotion Reflective motivation Memory, attention and decision processes</td>
<td>Knowledge</td>
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<td>Behavioural regulation Reflective motivation Reinforcement Reflective motivation Reinforcement Reflective motivation Physical skills Cognitive and interpersonal skills</td>
<td>Knowledge</td>
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<th>Mechanism of action</th>
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</thead>
<tbody>
<tr>
<td><strong>Mode</strong></td>
<td><strong>Content</strong></td>
<td><strong>BCT</strong></td>
<td><strong>Functions</strong></td>
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<tr>
<td></td>
<td>1.1 Goal-setting</td>
<td>Enablement</td>
<td>Reflective motivation</td>
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<td></td>
<td>1.5 Review behaviour goal(s)</td>
<td>Enablement</td>
<td>Reflective motivation</td>
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<td></td>
<td>2.2 Feedback on behaviour</td>
<td>Training</td>
<td>Reflective motivation</td>
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<td>6.1 Demonstration of behaviour</td>
<td>Modelling</td>
<td>Psychological capability</td>
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<td>7.1 Prompts/cues</td>
<td>Environmental restructuring</td>
<td>Automatic motivation</td>
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<td>8.1 Behavioural practice/ rehearsal</td>
<td>Training</td>
<td>Physical capability</td>
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<td>8.7 Graded task</td>
<td>Training</td>
<td>Physical capability</td>
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<td>12.6 Body changes</td>
<td>Training</td>
<td>Reflective motivation</td>
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<td></td>
<td>15.1 Verbal persuasion about capability</td>
<td>Enablement</td>
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<td>1.8 Behavioural contract</td>
<td>Incentivisation</td>
<td>Reflective motivation</td>
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<td>2.3 Self-monitoring of behaviour (training diary)</td>
<td>Training</td>
<td>Reflective motivation</td>
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<td></td>
<td>2.2 Feedback on behaviour</td>
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</tbody>
</table>

BCT, behavioural change technique; CME, continued medical education; COM-B, Capability, Opportunity, Motivation and Behaviour Model; HCP, healthcare practitioner; LBP, low back pain; MOC, model of care; TDF, Theoretical Domains Framework.
An implementation forum including rehabilitation unit managers and clinical researchers was formed. The implementation forum collaborated on forming overarching goals, timeline and logistics facilitating and sustaining the implementation of the BetterBack MOC in the primary care rehabilitation clinic clusters in the Östergötland public healthcare region.

An MOC support team was formed. This comprised experienced clinicians (clinical champions) from each rehabilitation unit together with clinical researchers facilitating local implementation and sustainability of the BetterBack MOC at the rehabilitation units.

A package of education and training that the support team can use to assist the use of the BetterBack MOC by HCP was developed.

- Physiotherapists in the three geographical clusters of public primary care rehabilitation clinics in Östergötland will be offered to participate in a 13.5-hour (2 days) continued medical education workshop. The workshop is designed by the

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**Figure 4** Steps involved for healthcare practitioners in delivering the contents of the BetterBack model of care. ICD-10, International Classification of Diseases-10.

**Figure 5** The Behavioural Change Wheel and the Theoretical Domains Framework (TDF).
support team with at least two clinical researchers and one experienced clinician from the rehabilitation unit cluster present in the support team’s delivery of the workshop for each cluster. The HCP education provided in the workshop format is described in online supplementary file 4.

- Key components of the educational programme are the following:
  - Education and persuasion about evidence-based recommendations for LBP care and the BetterBack® MOC through an experiential learning process applying problem-based case studies and clinical reasoning tools.
  - Training and modelling of the practical use of the BetterBack® education and physical intervention programmes aiming at self-care, as well as function and activity restoration.
  - Access to a website describing the BetterBack® MOC. A chat forum will give an opportunity for clinicians to ask questions and share different experiences of the new strategy managed by the support team.
  - To consolidate the BetterBack® MOC use at the local clinics, the local support team member will provide continued maintenance of education at their clinics and even educate new staff.

6. Once HCP behaviour change has occurred, it is anticipated that HCP use of the BetterBack® MOC may influence patient outcomes. A rationale for causal mediation effects can be proposed based on the Common Sense Model of Self-Regulation.42 This suggests a potential effect of the BetterBack® MOC on improved patient-reported pain, physical function and quality of life may be mediated by improved patient illness beliefs, such as cognitive and emotional illness representations, as well as adequate coping through self-care enablement.43 The patient target behaviours are therefore focused on the understanding of the mechanisms and natural course of benign LBP and the enablement of self-care. This requires content of the MOC to change patients’ impeding barrier behaviours such as maladaptive illness beliefs on the cause and persistent course of LBP (low outcome expectation, anxiety, catastrophising, fear avoidance and negative illness beliefs), low self-care enablement and low baseline physical activity.43 The content for the patient education and functional restoration programme included in the BetterBack® MOC therefore reflects these aspects and is shown in online supplementary file 3. These are also characterised according to the BCW, behavioural change technique taxonomy44 and TDF in table 3.

OUTCOMES
Implementation process
Primary outcome measure
- Practitioner Confidence Scale (PCS)45 mean change from baseline to 3 months post baseline. Practitioner-reported confidence is the primary HCP behavioural change goal for the HCP education and training workshop in the multifaceted implementation of the BetterBack® MOC. The 3-month time frame allows for the development and consolidation of HCP behavioural change after application in repeated patient cases.

Secondary outcome measures
- PCS45 mean immediate change from baseline to directly after the HCP education and training workshop, as well as mean long-term change from baseline to 12 months post baseline. This secondary outcome is important for the understanding of longitudinal HCP behavioural change.
- Pain Attitudes and Beliefs Scale for physical therapists (PABS-PT)46 mean change from baseline to directly after the HCP education and training workshop, as well as at 3 and 12 months post baseline.

Implementation outcomes
Primary outcome measure
- Proportional difference between control and intervention groups for incidence of participating patients receiving specialist care for LBP between baseline and 12 months after baseline. Incidence proportion, analogous to cumulative incidence or risk, is calculated by taking the number of patients receiving specialist care of LBP and dividing it by the total number of patients recruited to the study. The main goal of both the control and intervention conditions in primary care for benign first-time or recurrent debut of LBP is to improve patient-reported outcomes without the need of secondary or tertiary healthcare processes.

Secondary outcomes measures
- Mean difference between control and intervention groups for change between baseline and final clinical visit regarding grade of patients’ functional impairment and activity limitation according to the International Classification of Functioning, Disability and Health (ICF) brief core set for LBP.47
- The proportion of patients who receive the BetterBack® MOC and registration of healthcare codes coherent with the Swedish best practice clinical recommendations.

Patient outcomes
Primary outcome measure
- Numeric Rating Scale for lower back-related pain intensity (NRS-LBP) during the latest week.48 The mean difference between control and intervention groups in change between baseline and 3 months post
baseline will be analysed. Pain intensity is the primary functional impairment that patients with LBP contact primary healthcare for and has been recommended by international consensus to be included as a core outcome domain for clinical trials in non-specific LBP.49 International consensus even recommends patient-reported NRS change over 6 months as a core metric for pain management interventions.50

► Oswestry Disability Index V.2.1 (ODI).51 The mean difference between control and intervention groups in change between baseline and 6 months post baseline will be analysed. Disability, analogous to decreased physical functioning and activity limitation, has been recommended by international consensus to be included as a core outcome domain for clinical trials in non-specific LBP.49 International consensus even recommends patient-reported ODI change over 6 months as a core metric for functional restoration.50

Secondary outcome measures

► NRS-LBP48 and ODI50 mean difference between control and intervention groups in short-term change from baseline to 3 months post baseline and mean long-term change from baseline to 12 months post baseline. These secondary outcomes are important for the understanding of longitudinal patient-rated changes in pain intensity and disability after primary care intervention.

► The European Quality of Life Questionnaire (EQ-5D).52 The mean difference between control and intervention groups in change between baseline and 3, 6 and 12 months post baseline will be analysed. Health-related quality of life has been recommended by international consensus to be included as a core outcome domain for clinical trials in non-specific LBP.49 International consensus even recommends patient-reported EQ-5D change over 6 months as a core metric for pain management interventions.50

► The Brief Illness Perception Questionnaire (BIPQ).53 The mean difference between control and intervention groups in change between baseline and 3, 6 and 12 months post baseline will be analysed. Illness perception has been shown to predict longitudinal pain and disability outcomes in several LBP studies.54-58

► Patient Enablement Index (PEI),59 Patient Global Rating of Change (PGIC)60 and Patient Satisfaction (PS)61 mean difference between control and intervention groups at 3, 6 and 12 months post baseline will be analysed.

Participant timeline

The trial timeline is shown in table 2. The intervention schedule started with the development of evidence-based recommendations and the BetterBack® MOC, which occurred during June 2016–February 2017. The enrolment schedule started with cluster enrolment and randomisation in March 2017. This resulted in the first allocated cluster 1 (west) entering internal pilot of implementing the BetterBack® MOC HCP education and training workshop, which occurred in March 2017. This was followed up with a 2-month internal pilot of patient enrolment schedule occurring in all three clusters during April–May 2017. In order to finalise a sample size calculation for the main trial, baseline data collected during the internal pilot are compared with follow-up data 3 months after baseline for the primary outcome measure questionnaires to analyse initial HCP and patient effects of the implementation of BetterBack® MOC in cluster 1 compared with the control conditions in clusters 2 and 3. In the transition to the main trial, patient enrolment and baseline assessments will then continue to occur until January 2018. The eventual time of crossing forward of cluster 2 into the implementation of the BetterBack® MOC is determined by the internal pilot trial results. Participants in the trial will be followed up longitudinally at 3, 6 and 12 months after baseline measures. The schedule for assessments is also outlined in table 2.

Sample size

An initial sample size estimation in the planning stage of the study assumed at least a small Cohen’s d effect size (d=0.35) for the HCP behavioural change primary and secondary outcomes. This is based on previous literature showing small-moderate HCP behavioural change effects sizes using similar interventions to increase the uptake of evidence-based management of LBP in primary care.62 63 Considering also a one-tailed P=0.05 for the benefit of the multifaceted implementation of the BetterBack® MOC, an 80% statistical power and a 20% loss to follow-up, a sample size of n=63 HCP is needed for a matched pairs t-test statistics comparing baseline and follow-up means. We assume a possible carry-over of a similar effect size (d=0.35) on patient behavioural change primary and secondary outcomes. Considering also a one-tailed P=0.05 for the benefit of the multifaceted implementation of BetterBack® MOC compared with usual care and an 80% statistical power, the number of patients required for an individually randomised simple parallel group design would be n=204. Adjusting for the design effect due to cluster randomising, an intraclass correlation of 0.01 and a cluster autocorrelation of 0.80, a dogleg design with two assessments in routine care and 100 patients in each cluster section would require at least n=402 patients over 2.41 clusters according to algorithms described by Hooper and Bourke.90 In a balanced recruitment schedule, this equates to 14 patients per month per cluster for a total of 3 clusters. Allowing for potential unbalanced recruitment flow and a potential dropout in the longitudinal outcomes at 3, 6 and 12 months post baseline, each cluster will aim for up to 20 patients per month, equating to a potential total study of n=600.

Recruitment

In an effort to curb recruitment difficulties, strategies to promote adequate enrolment of participants into the study will be used. We anticipate less problems with
recruitment into the prospective cohort study design investigating the multifaceted implementation of the BetterBack© MOC at the HCP level. This is due to the study having been endorsed by clinical department managers calling all HCPs working with patients with LBP at their clinics to participate. However, recruitment of patients into the cluster randomised controlled trial is dependent on the feasibility of recruitment processes adapted to the context of each individual clinic and the compliance of HCPs to administer recruitment of consecutive patients. A strategy to optimise the administration of patient recruitment will involve the author KS regularly visiting participating clinics to inform HCPs of the study protocol and help streamline practical administration of the protocol in the context of the individual clinics. KS will also monitor weekly recruitment rates from the clinics and provide motivational feedback on recruitment flow to clinical department managers and designated clinical champions who will provide additional motivational feedback to HCP. In accordance with the Consolidated Standards of Reporting Trials, a flow diagram displaying participant enrolment, allocation, follow-up and analysis will be constructed.64 Reasons for exclusion, declined participation, protocol violations and loss to follow-up will be monitored by KS.65

Allocation and blinding
Random concealed allocation of clusters was performed by a blinded researcher randomly selecting from three sequentially numbered, opaque, sealed envelopes. The method resulted in the following order: 1=cluster west, 2=cluster central and 3=cluster east. KS informed the clinics in the different clusters of their allocation to the control or intervention study condition. Due to the nature of the study and intervention, HCPs conducting patient measurements and treatment cannot be blinded to group allocation. Risk of bias is minimal as the primary and secondary outcomes are patient self-reported questionnaires. Patients will be blinded to group allocation. The researcher responsible for statistical analysis will not be blinded to group allocation, but an independent statistician will review statistical analysis.

Data collection
Data will be collected through quantitative questionnaires and qualitative focus group and semistructured interviews. In the case of non-response to questionnaires, a questionnaire will be resent via post a total of three times. In case of continued non-response, this will be complemented with a telephone call as a final effort for data collection.

Implementation process
► The PCS contains four items reported on a 5-point Likert scale, where a total score of 4 represents greatest self-confidence and 20 represents lowest self-confidence for managing patients with LBP. The structural validity in terms of internal consistency of the items has been shown to be good with a Cronbach’s α coefficient=0.73 in a single factor model for self-confidence.45 The questionnaire has been forward-translated by our research group from English to Swedish.
► The PABS-PT consists of two factors where higher scores represent more treatment orientation regarding that factor. One factor with 10 items measures the biomedical treatment orientation (score 0–60) and one with 9 items measures the biopsychosocial treatment orientation (score 0–54).46 Each item is rated on a 6-point Likert scale ranging from 1=’totally disagree’ to 6=’totally agree’. The internal consistency of the biomedical factor has been shown to be good with a Cronbach’s α range of between 0.77 and 0.84. Furthermore, the biopsychosocial factor has been shown to be adequate with a Cronbach’s α range of between 0.62 and 0.68.65 Construct validity and responsiveness to educational interventions have been shown to be positive along with the test–retest reliability with reported intraclass correlation coefficient (ICC) on the biomedical factor of 0.81 and on the biopsychosocial factor of 0.65.66 The questionnaire has been forward-translated from English to Swedish in a previously published study.66

► The Determinants of Implementation Behaviour Questionnaire (DIBQ) was originally constructed based on the domains of the TDF.41 67 Confirmatory factor analysis resulted in a modified 95-item questionnaire assessing 18 domains with sufficient discriminant validity. Internal consistency of the items for the 18 domains was good, ranging from 0.68 to 0.93 for the Cronbach’s α coefficient.68 The questionnaire has been forward-translated by our research group from English to Swedish. After face validity consensus in our research group regarding relevant domains for the implementation of BetterBack© MOC, the questionnaire was shortened to the following domains: knowledge, skills, beliefs about capabilities, beliefs about consequences, intentions, innovation, organisation, patient, social influence and behavioural regulation, totalling to 57 items. Questions were adapted to the context of HCP-reported determinants of an ‘expected’ implementation of BetterBack© MOC for measurement directly after the HCP education and training workshop. HCP-reported determinants retained original wording for the questionnaires at 3 and 12 months after the implementation of BetterBack© MOC. The response scale used for each DIBQ question in our study is a 5-point Likert scale ranging from 1=’totally agree’ to 5=’totally disagree’.
pain clinic, orthopaedic or neurosurgical care, and the number receiving surgery.

- Clinical reasoning and process evaluation tool (CRPE tool): grade of patients’ functional impairment and activity limitation according to the ICF brief core set for LBP is assessed by the physiotherapist at baseline and final clinical contact, where light, moderate, severe and very severe impairment/limitation are coded 0–4, respectively. A total score for baseline and follow-up measures is calculated from the sum of the functional impairment divided by the number of functional impairments, and a similar total score is calculated for activity limitations. A worsening of functional impairments and activity limitations measured at follow-up with the CRPE will be considered in the analysis of adverse events. Swedish Classification of Health Interventions (KVÅ) codes for assessment and treatment interventions will be assessed to analyse coherence with the Swedish best practice clinical recommendations. International Classification of Diseases-10 diagnosis codes will also be recorded.

- The Keele STarT Back Screening Tool is reported by patients at baseline providing a stratification of prognostic risk of persistent pain. The overall score ranging from 0 to 9 is used to separate the low-risk patients from the medium-risk subgroups, where patients who achieve a score of 0–3 are classified into the low-risk subgroup and those with scores of 4–9 into the medium-risk subgroup. To identify the high-risk subgroup, the last five items must score 4 or 5.

- Focus groups performing qualitative Strengths, Weaknesses, Opportunities and Threats (SWOT) analyses will be conducted by HCPs between 3 and 6 months after implementation.

- Semistructured interviews with 10 HCPs at 3 months after implementation will be conducted to investigate determinants of implementation behaviour and if other determinants need to be added to the DIBQ. The interviews will be deductively analysed according to the TDF and BCW frameworks.

- Semistructured interviews investigating patient experience of receiving care for LBP will be performed on 10 patients. These patients will have received care after implementation of the BetterBack© MOC.

- Economic costs of developing the BetterBack© MOC as well as performing the implementation strategy (staff time, HCP training and printed resources).

Patient outcome measures

- NRS-LBP intensity during the latest week is an 11-point scale consisting of integers from 0 through 10, with 0 representing ‘No pain’ and 10 representing ‘Worst imaginable pain’. Previous research in an LBP cohort has shown a test–retest reliability ICC of 0.61, a common SD of 1.64 points, SE of measure of 1.02 and minimal clinically important difference (MCID) in LBP after treatment of 2.72,73

- ODI V.2.1 assesses patients’ current LBP-related limitation in performing activities such as personal care, lifting, walking, sitting, standing, sleeping, sex life, social life and travelling. The ODI consists of 10 items with response scales from 0 to 5, where higher values represent greater disability. The ODI is analysed as a 0–100 percentage variable, where lower scores represent lower levels of LBP disability. A reduction of 10 points is considered the MCID in LBP after treatment.50 In Scandinavian conditions, the coefficient of variation, ICC and internal consistency of the ODI is 12%, 0.88–0.91 and 0.94, respectively.74,76 Good concurrent validity has also been shown.75

- The EQ-5D measures generic health-related quality of life and is computed into a 0–1.00 scale from worst to best possible health state by using the Swedish value sets.77 A reduction of 0.08 points is considered the MCID in LBP after treatment.78 The mean change after treatment for LBP has been reported to be 0.12 (SD ±0.30).79

- The BIPQ analyses cognitive illness representations (consequences, outcome expectancy, personal control, treatment control and knowledge), emotional representations (concern and emotions) as well as illness comprehensibility. An overall score of 0–80 represents the degree to which the LBP is perceived as threatening or benign, where a higher score reflects a more threatening view of the illness.32 The BIPQ has been shown to be valid and reliable in a Scandinavian sample of patients with subacute and chronic LBP. The BIPQ has a Cronbach’s α of 0.72 and a test–retest ICC of 0.86, an ICC range for individual items from 0.64 to 0.88, an SE of measurement of 0.63 and minimal detectable change of 1.75.80

- The PEI has a score range between 0 and 12, with a higher score intended to reflect higher patient self-care enablement.39

- PGIC asks the patient to rate the degree of change in LBP-related problems from the beginning of treatment to the present. This is measured with a balanced 11-point numerical scale. A reduction of 2 points is considered the MCID in LBP after treatment.60

- PS is measured with a single-item patient-reported question. The question asks ‘Over the course of treatment for this episode of LBP or leg pain, how satisfied were you with the care provided by your health care provider? Were you very satisfied (1), somewhat satisfied (2), neither satisfied nor dissatisfied (3), somewhat dissatisfied (4), or very dissatisfied (5)?’.61

- Economic costs of health service utilisation.

Data management

All paper-based questionnaire data will remain confidential and will be kept in a lockable filing cabinet in the research group office. A password-protected coded database only accessible to the research team will be kept on a data storage device in the research department. The research team will regularly monitor the integrity of trial
data. Trial conduct will be audited on a weekly basis by the research team.

Statistical analysis
Statistical significance will be assessed with an alpha level of 0.05. All results will be reported as estimates of mean ±SD and also effect size (eg, mean difference) with 95% CIs. An intention-to-treat (ITT) principle applying multiple imputation will be used. A sensitivity analysis will compare per-protocol and ITT databases. A sensitivity analysis will also be used to assess the significance of a washout period by comparing the complete database against the same database without data collected during the 2 weeks in conjunction with the BetterBack® implementation in each cluster.

Implementation process and outcome analysis
Analysis of variance statistics comparing baseline and follow-up means will be used for implementation process and outcome measures. Causal mediation analysis will be used to analyse indirect mediational effects of multiple putative determinants of implementation behaviour measured with the DIBQ directly after the HCP education and training workshop (intention stage) or at 3 or 12 months (volition stages) on the effect of baseline PCS or PABS-PT or 3-month or 12-month follow-up measurement of PCS or PABS-PT. If the HCP education and training workshop does not have a causal effect on improved prospective outcomes, we will analyse where the causal pathway breaks down. Causal mediation analysis will be performed using the program PROCESS63 within IBM SPSS Statistics V.23 (figure 6).

Patient outcome measures for the control and intervention groups will be compared using multilevel analyses of repeated measurements and experiment condition as fixed effects and participants and clusters as random effects with IBM SPSS. Fixed-effect interactions between the experimental condition and The Keele StArT Back Screening Tool will also be assessed. Patient population-specific MCID will be assessed for primary and secondary outcomes based on an anchor method where PGIC serves as an anchor. Applying a 1-1-1 multilevel mediation procedure with all effects random in MPLUS, the products of (1) the independent variable (experimental condition: control or intervention) to the mediator (change in BIPQ, PEI), and (2) the mediator to the dependent variable (change in NRS, ODI or secondary outcome scores pretreatment to post-treatment) when the independent variable is taken into account, will be tested for mediation (figure 7).

Economic analysis
The reference case analysis is based on a healthcare sector perspective. The EQ-5D will be used to calculate the ratio of costs to QALY saved for patients. The ICERs for the multifaceted implementation strategy and the usual care condition will be calculated and plotted on a cost-effectiveness plane. This is based on the Swedish guideline-priced direct costs of health service utilisation, organisational costs of developing the BetterBack® MOC, as well as performing the implementation strategy and overall intervention clinical outcome effectiveness. The ICER will also be calculated per patient avoiding specialist care. To estimate a distribution of costs and health measures and CIs for ICER, bootstrapping will be used.

Data monitoring
All outcome questionnaires are formatted for use of scan processing software for automated data entry into the Statistical Package for the Social Sciences package. KS, who is not blinded to treatment allocation, will perform regular data checks during data entry and provide feedback when necessary to HCPs regarding data omissions. KS will also double-check data entry to detect and correct input errors, and range checks will be undertaken prior to data analysis.

Ethics and dissemination
The ethics application including consent forms in Swedish is available on request to the authors. There are no known risks for participants. Voluntarily participating HCPs will complete questionnaires. All participating patients are informed orally and in writing about the study on the first visit at participating primary healthcare clinics. They are informed that participation is voluntary and that they can at any time withdraw their participation. The HCP intervention will not be affected by the patient’s decision to participate or not participate in the study. Data collection will not be performed for those not participating. A signed patient consent form will be collected from patients by the HCP before baseline measures are collected and intervention is commenced according to the study protocol. All collected data will be entered into a database accessible to the authors. A code list will be created where each participant will be represented by a code so that the database will be anonymous. The code list with personal data will be stored separately in locked filing cabinets at Linköping University to protect confidentiality before, during and after the study. Data analyses and reporting will be performed using the de-identified database. The authors plan to disseminate the findings through manuscript publications in scientific journals and presentation at conferences.

Patient and public involvement
The adaptation of best practice clinical guidelines to the Swedish context, the construction of the BetterBack® MOC, as well as the development of the research question, study design and outcomes measures involved interpretation of literature and professional experience of the patient net benefit versus harms and costs. Specific investigations of priorities, experience and preferences of the patients in the Östergötland healthcare region were not performed in this development phase. No patient advisors or other public are involved in the study. HCPs
working with patients with LBP at their clinics ask consecutive patients to participate in the study and adhere to the prescribed intervention. Patients have no other involvement in recruitment and conduct of the study. Semistructured interviews on 10 patients randomly selected will investigate the priorities, experience, burden and preferences of the intervention. Patients’ satisfaction regarding the intervention is assessed by the patients themselves through a questionnaire. The dissemination of the study findings to participating patients will occur through popular science summary publication.

**Internal pilot trial results**
The initial implementation of the BetterBack MOC in cluster 1 allowed for an internal pilot to determine the HCP acceptability of the intervention and trial within the first cluster. A progression criteria for continuing to the main trial required that HCPs who have completed

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**Figure 6** Causal mediation model to analyse indirect mediational effects of multiple putative determinants of implementation behaviour measured with the DIBQ directly after the healthcare practitioner education/training workshop (intention stage) or at 3 or 12 months (volition stages) for the effect of baseline PCS or PABS-PT on 3-month or 12-month follow-up measurement of PCS or PABS-PT (c). DIBQ, Determinants of Implementation Behaviour Questionnaire; PABS-PT, Pain Attitudes and Beliefs Scale for physical therapists; PCS, Practitioner Confidence Scale.
the BetterBack® education and training workshop rate on average a maximum of 2.5 out of 5 on the following determinants of implementation behaviour question: I expect that the application of BetterBack® MOC will be useful (1=agree completely to 5=do not agree at all). The 27 HCPs participating in the internal pilot in cluster 1 responded to the question with a mean value of 1.7 (SD 0.8), which subsequently fulfilled the HCP progression criteria.

The resulting internal pilot patient flow for April and May was n=28 and n=28 for cluster 1 west (intervention), n=5 and n=12 for cluster 2 central (control), as well as n=14 and n=22 for cluster 3 east (control) consecutively. This informed the decision to move the cluster 2 transition from control to intervention condition to occur later in the schedule, planned for September 2017, to allow for more control condition patient recruitment and data collection. The flow of patient recruitment and the process of 3-month follow-up in the internal pilot were used to inform the optimal time point of patient-reported primary outcome for the main trial. Our initial planning was to measure patient-reported primary outcome at 6 months post baseline based on the definition of persistence/chronicity of symptoms being often defined in the literature to be of 3 and up to 6 months in duration.82 Our internal pilot study had a 3-month follow-up rate of 80% resulting after up to three reminders sent to many of these patients. This informed of a likely risk of non-response at later follow-up time points. Furthermore, feedback from participating HCPs even reported a larger clinical interest in 3-month patient follow-up data. Therefore the internal pilot informed the choice to revise our patient-reported primary outcomes to 3 months post baseline with subsequent amendments of the trial registration on ClinicalTrials.gov: NCT03147300.

Our internal pilot study was also used to assess baseline variation and change over 3 months in HCP and patient-reported primary outcome measures in the control and intervention arms to aid calibration of the sample size calculation. Multilevel analyses of repeated measurements and experiment condition as fixed effects and participants and clusters as random effects revealed an intracluster correlation of <0.01 for all primary outcomes measures. A small effect size in favour of the intervention condition was shown for HCP-reported PCS (d=0.33) directly after implementation but increased to a moderate effect size after 3 months (d=0.51). Patient-reported NRS showed a small effect size (d=0.28). Therefore, the internal pilot data supported our a priori sample size calculation for the main trial regarding PCS and NRS. However no effect size difference was observed between experimental conditions for ODI. It is possible that when statistical power improves when the trial progresses, potential differences in ODI may be detectable between experimental conditions.

CONCLUSION
The effectiveness-implementation hybrid type 2 trial with dogleg stepped cluster structure allowed for the use of an internal pilot to inform feasibility and optimise method efficiency for the progression of the trial.
Contributors AA and BÖ formulated the trial’s original aims and hypothesis. AA, KS and BÖ developed intervention materials. AA, KS, PE, PN and ÖB designed the study methodology. AA, PN and BÖ procured funding for the trial. AA, KS, PE, PN and ÖB have reviewed and finalised the protocol.

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Competing interests None declared.

Patient consent Detail has been removed from this case description/these case descriptions to ensure anonymity. The editors and reviewers have seen the detailed information available and are satisfied that the information backs up the case the authors are making.

Ethics approval Ethical clearance for the study (Dnr: 2017-35/31) has been attained through the Regional Ethics Committee in Linköping.

Provenance and peer review Not commissioned; externally peer reviewed.

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