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Prognostic ability of STarT Back Screening Tool combined with work-related factors in patients with low back pain in primary care: a prospective study

Monica Unsgaard-Tøndel,1,2,3 Ottar Vasseljen,2 Tom Ivar Lund Nilsen,2,4 Gard Myhre,3 Hilde Stendal Robinson,5 Ingebrigt Meisingset2,3

ABSTRACT
Objective Primary care screening tools for patients with low back pain may improve outcome by identifying modifiable obstacles for recovery. The STarT Back Screening Tool (SBST) consists of nine biological and psychological items, with less focus on work-related factors. We aimed at testing the prognostic ability of SBST and the effect of adding items for future and present work ability.

Methods Prospective observational study in patients (n=158) attending primary care physical therapy for low back pain. The prognostic ability of SBST and the added prognostic value of two work items; expectation for future work ability and current work ability, were calculated using the explained variance (R²) of the outcomes from univariable and multivariable linear regression and beta values with 95% CIs were used to assess the prognostic value of individual items.

Results The SBST classified 107 (67.7%) patients as low risk and 51 (32.3%) patients as medium/high risk. SBST provided prognostic ability for disability (R²=0.35), pain (R²=0.25) and quality of life R²=0.28). Expectation for return to work predicted outcome in univariable analyses but provided limited additional prognostic ability when added to the SBST. Present work ability provided additional prognostic ability for disability (β=−2.5; 95% CI=−3.6 to −1.4), pain (β=−0.2; 95% CI=−0.5 to −0.002) and quality of life (β=0.02; 95% CI=0.001 to 0.04) in the multivariable analyses. The explained variance (R²) when work ability was added to the SBST was 0.60, 0.49 and 0.47 for disability, pain and quality of life, respectively.

Conclusions Adding one work ability item to the SBST gives additional prognostic information across core outcomes.

Clinical trial number: NCT03626389

INTRODUCTION
Low back pain is a global public health problem with high socioeconomic burden. Low back pain is related to reduced work ability and expected to increase. Primary healthcare practitioners should support people to sustain work and activity by providing multidimensional, precise and patient-centred management. A recent review showed that patients with low back pain desire workplace accommodations and want their employers to be informed about low back pain. In general, improved integration of social and environmental factors in healthcare management is needed. To increase healthy work participation and reduce negative consequences for individuals and society, work ability needs to be addressed in first-line management.

Screening tools could contribute to an integrated multidimensional approach if they are short, easily administered, include the relevant dimensions in the biopsychosocial model, and support treatment allocation. STarT Back Screening Tool (SBST) was developed in UK primary care, to identify patient subgroups for initial treatment by mapping modifiable prognostic physical and psychological factors by means of nine screening items. The predictive power of SBST has previously been evaluated, though there has been considerable heterogeneity between studies and results. Moreover, the SBST does not cover work-related obstacles for recovery.
Another screening tool, the short form Örebro Musculoskeletal Pain Screening Questionnaire (ÖMPSQ), includes the work domain and has been found to be better than SBST for predicting pain and work outcome, while SBST seem to better predict ‘function’ outcome. A recent Cochrane review concluded that individual recovery expectations probably are strongly associated with future work outcome. The authors recommended screening of recovery expectations in order to improve the prognosis and tailoring of low back pain management. In line with this, prospective studies have suggested that work-related risk factors can predict work disability after 3 years in a general working population and that improved perception of work environment were associated with reduced disability in patients with neck and back pain. Moreover, we have previously shown that self-reported work ability, assessed by a single item, was associated with disability, pain and quality of life, which are regarded core outcomes, in patients seeking physiotherapy for low back pain. We therefore hypothesised that the predictive ability of SBST would be improved by adding single items that addressed both current work ability, as well as expectations for future work ability.

On this background, we aimed at examining the prognostic ability of SBST for disability, pain intensity and quality of life after 3 months’ treatment, and to examine if adding two items on future and present work ability to SBST influence the prognostic ability.

**METHODS AND MATERIALS**

**Design and setting**

We used observational data prospectively collected in primary healthcare physiotherapy in Norway in the FYSIOPRIM project. Data were collected at baseline and at 3 months follow-up from patients attending primary care physiotherapy between January 2016 and December 2018. The patients answered baseline questionnaires electronically or at the physiotherapy clinic at the first appointment (baseline) and at 3 months follow-up. Reminders for non-responders at 3 months were sent by email and short message service up to three times. Details of the data collection are provided in the study protocol for the FYSIOPRIM project.

**Participants**

Patients with low back pain as the main problem defined by the physiotherapist was collected from the database of the FYSIOPRIM project. Due to the allocation algorithm in the FYSIOPRIM study, patients classified by therapist with multisite pain as main problem were allocated to another group defined as complex and not included in the present study. Further, we excluded patients with age under 18, pregnancy related disorders, specific neurological disorders (eg, stroke, Parkinson’s disease, multiple sclerosis), recent surgery, chronic obstructive pulmonary disease, fracture and inflammatory arthritis such as rheumatoid arthritis. The patients received usual care physiotherapy with contents based on the physiotherapists’ assessment.

**Variables**

**SBST**

SBST was the prognostic tool examined in this study. The tool has nine screening items on physical and psychological factors related to low back pain and responses are dichotomised into ‘Agree’ and ‘Disagree’. In the current study the medium-risk and high-risk groups were collapsed to one medium/high risk group due to few subjects in the high-risk group. This dichotomisation of the risk groups represents the main difference between the STarT Back risk groups in terms of management volume, were the low-risk group should receive advice and guidance in order to self-manage, while the medium-risk and high-risk groups should in addition receive standardised physiotherapy to address symptoms and function or psychologically informed physiotherapy, respectively.

**Work-related prognostic factors**

We chose the work-related prognostic factors a priori, with the aim to include single work items for screening purposes. Patient’s expectation of future work ability was assessed with item no. 8 (‘In your estimation, what are the chances you will be working your normal duties in 3 months?’) in short form ÖMPSQ. Responses were given on a numerical rating scale from 0 to 10, where 0 indicates not likely and 10 indicates very likely. Additionally, current work ability was measured by a single-item question: ‘describe your current work ability compared with the lifetime best’, and scored on an 11-point numerical rating scale (0–10), where 0 represent ‘completely unable to work’, and 10 represent ‘work ability at its best’. This question is obtained from the work ability index and strong correlation with the seven-item work ability index has been reported.

**Outcomes**

The main outcome in this study was low back pain related disability assessed by Oswestry Disability Index (ODI) (range 0–100, where 0 indicate no disability and 100 indicate 100% disability). Secondary outcomes were pain intensity and health-related quality of life. Together, these three outcomes are defined as core outcome measures in low back pain trials. Pain intensity was assessed by the question ‘How would you rate the pain that you have had during the past week?’, and responded on a numeric pain rating scale ranging from 0 to 10, where 0 indicated no pain and 10 indicated worst imaginable pain. Health-related quality of life was assessed by the Euroqol instrument (EQ-5D-5L). The EQ-5D-5L evaluates the following five dimensions: mobility, self-care, usual activities, pain and/or discomfort and anxiety and/or depression. The response options for each dimension were ‘no problems’, ‘slight problems’, ‘moderate problems’, ‘severe problems’ and ‘extreme problems’. As recommended by the Norwegian guideline, we used the
UK value set to calculate an index value for health status (range −0.285 to 1, where −0.285 means extreme problems on all dimensions, and 1 means perfect health).23 24

**Other baseline variables**

We assessed a range of other baseline factors to provide a comprehensive description of our study sample. Patient-reported background information included age, gender, education (higher education, no/yes), height (cm) and weight (kg) (used to calculate body mass index) and smoking (yes/no) (table 1). Further, physical activity was measured using three questions regarding frequency, duration and intensity of physical activity. According to Nes *et al*, a scale from 0 to 45 was derived from the responses, where a score of 0 was defined as physically inactive.25 Function was assessed using the Patient Specific Functional Scale, where the patients described the most troublesome activity.26 The activity was scored on a Numerical Rating Scale (NRS) from 0 to 10, where 0 indicated not able to perform activity and 10 no problem to perform activity. Patients also reported their employment status and whether they were sick listed. Pain duration was assessed by item number one from the Örebro screening questionnaire17 and the responses were categorised in (1) less than 3 months, (2) 3–12 months and (3) longer

<table>
<thead>
<tr>
<th>Sample size, n (%)</th>
<th>Total sample</th>
<th>Low risk SBST</th>
<th>Medium/high risk SBST</th>
</tr>
</thead>
<tbody>
<tr>
<td>Females, n (%)</td>
<td>n=158</td>
<td>n=107 (67.7)</td>
<td>n=51 (32.3)</td>
</tr>
<tr>
<td>Age, mean (SD)</td>
<td>46 (17.5)</td>
<td>45.7 (18)</td>
<td>46 (16.4)</td>
</tr>
<tr>
<td>Body mass index, median (IQR)</td>
<td>25.2 (22.7–28.6)</td>
<td>24.5 (22.3–27.2)</td>
<td>27.4 (24.3–31.9)</td>
</tr>
<tr>
<td>Higher education, n (%)</td>
<td>90 (57.6)</td>
<td>66 (62.3)</td>
<td>27 (54)</td>
</tr>
<tr>
<td>Current smoker, n (%)</td>
<td>17 (11)</td>
<td>8 (7.6)</td>
<td>9 (18)</td>
</tr>
<tr>
<td>Physically inactive, n (%)</td>
<td>51 (38.4)</td>
<td>33 (35.1)</td>
<td>18 (46.2)</td>
</tr>
<tr>
<td>Patient specific functional scale (0–10), mean (SD)</td>
<td>3.9 (2.2)</td>
<td>4.1 (2.2)</td>
<td>3.4 (2.1)</td>
</tr>
<tr>
<td>Employed (no sick leave benefit), n (%)</td>
<td>72 (46.2)</td>
<td>52 (49.1)</td>
<td>20 (27.8)</td>
</tr>
<tr>
<td>Pain duration, n (%)</td>
<td>40 (25.9)</td>
<td>27 (26)</td>
<td>13 (26)</td>
</tr>
<tr>
<td>&lt;3 months</td>
<td>41 (26.5)</td>
<td>32 (30.8)</td>
<td>9 (18)</td>
</tr>
<tr>
<td>&gt;12 months</td>
<td>73 (47.4)</td>
<td>45 (43.3)</td>
<td>28 (56)</td>
</tr>
<tr>
<td>Uses analgesics, n (%)</td>
<td>74 (48.1)</td>
<td>40 (38.1)</td>
<td>34 (69.4)</td>
</tr>
<tr>
<td>Hopkins Symptom Checklist (1–4)*, median (IQR)</td>
<td>1.6 (1.3–2)</td>
<td>1.4 (1.3–1.8)</td>
<td>2 (1.6–2.4)</td>
</tr>
<tr>
<td>High score, cut-off ≥1.85, n (%)</td>
<td>48 (34.5)</td>
<td>20 (43.5)</td>
<td>26 (56.5)</td>
</tr>
<tr>
<td>Kinesiophobia (0–10), median (IQR)*</td>
<td>3 (0–5)</td>
<td>2 (0–5)</td>
<td>5 (2–8)</td>
</tr>
<tr>
<td>Pain Self-Efficacy Questionnaire 2 item (0–12) median (IQR)</td>
<td>10 (9–12)</td>
<td>11 (10–12)</td>
<td>9 (5–10)</td>
</tr>
<tr>
<td>Moderate-to-severe sleep problems, n (%)</td>
<td>50 (35.5)</td>
<td>25 (25.3)</td>
<td>25 (59.5)</td>
</tr>
<tr>
<td>Quite to very reduced daily activity level, n (%)</td>
<td>75 (47.8)</td>
<td>34 (32.1)</td>
<td>41 (80.4)</td>
</tr>
<tr>
<td>Örebro screening questionnaire short form (0–100)*, mean (SD)</td>
<td>43.8 (15.4)</td>
<td>38.2 (13.1)</td>
<td>55.1 (13.6)</td>
</tr>
<tr>
<td>High risk (&gt;50), n (%)</td>
<td>53 (35.8)</td>
<td>22 (22.2)</td>
<td>31 (63.3)</td>
</tr>
<tr>
<td>Expectations on future work participation (0–10), median (IQR)*</td>
<td>0 (0–5)</td>
<td>0 (0–1)</td>
<td>5 (0–10)</td>
</tr>
<tr>
<td>Work ability (0–10), median (IQR)</td>
<td>7 (3–9)</td>
<td>8 (6–9)</td>
<td>3 (1–6)</td>
</tr>
<tr>
<td>Baseline Oswestry Disability index (0–100), mean (SD)*</td>
<td>22.7 (14.4)</td>
<td>16.5 (9.1)</td>
<td>35.8 (14.8)</td>
</tr>
<tr>
<td>3 months Oswestry Disability index, mean (SD)</td>
<td>16.9 (13.6)</td>
<td>11.2 (8.1)</td>
<td>28.4 (15.2)</td>
</tr>
<tr>
<td>Baseline pain intensity last week (0–10), mean (SD)*</td>
<td>4.9 (2.2)</td>
<td>4.1 (2.1)</td>
<td>6.5 (1.5)</td>
</tr>
<tr>
<td>3 months pain intensity last week (0–10), mean (SD)</td>
<td>3.3 (2.5)</td>
<td>2.4 (2)</td>
<td>5.1 (2.4)</td>
</tr>
<tr>
<td>Baseline health-related quality of life, median (IQR)</td>
<td>0.80 (0.66–0.88)</td>
<td>0.84 (0.77–0.89)</td>
<td>0.64 (0.46–0.75)</td>
</tr>
<tr>
<td>3 months health-related quality of life, median (IQR)</td>
<td>0.89 (0.77–0.94)</td>
<td>0.92 (0.84–0.94)</td>
<td>0.79 (0.53–0.84)</td>
</tr>
</tbody>
</table>

*Low scores indicate low symptom pressure, while high scores indicate high symptom pressure.
than 12 months. Use of analgesics was assessed by the question ‘Have you used pain medication the last week?’ with response option yes or no. Mental distress (anxiety and depression) was assessed by the Hopkins Symptom Checklist - 10 item version (HSCL-10, range 0–4), where higher scores indicated higher mental stress. A cut-off at ≥1.85 on the HSCL-10 was used to define high versus low mental stress. Kinesiophobia was assessed by the question ‘How much fear do you have that these complaints would be increased by physical activity?’. The patients responded on a NRS from 0 to 10, where 0 indicated no fear and 10 indicated very much fear. Pain self-efficacy was assessed by the 2-item version of the Pain Self-Efficacy Questionnaire, and scored on a scale from 0 to 12 where higher score indicated higher pain self-efficacy. Sleep problems were assessed by a single item from the 15D questionnaire, asking patients to define their level of sleep problems on a 5-point Likert scale: ‘I’m able to sleep normally’, ‘I have slight problems with sleeping’, ‘I have moderate problems with sleeping’, ‘I have great problems with sleeping’ and ‘I suffer severe sleeplessness’. Pain interference with daily activity was assessed by the question ‘Due to pain or complaints, how much reduced is your activities of daily life?’. The response options were very much-reduced, quite-reduced, slightly-reduced and not reduced. The Örebro screening questionnaire short form was used to assess risk for long-term work disability on a scale from 0 to 100, where higher score indicated higher risk for long-term work disability.

### Statistical analysis

Descriptive statistics was used to characterise the STarT Back risk groups. Linear regression was used to assess the ability of the STarT Back risk groups and the two work-related items to predict disability, pain intensity and quality of life outcomes at 3 months. We performed the regression analyses with SBST alone, and by adding the other predictors one by one to the model (baseline score of the outcome, expectation of future work ability and current work ability). We used the explained variance (adjusted R²) to compare the different models’ predictive ability. The prognostic ability of the SBST and the two work-related items were evaluated using five regression models. A first, a univariate model (model 1) estimated the prognostic ability of SBST and each of the two work items separately. Model 2 was multivariable analyses including SBST and the baseline values of the outcomes, model 3 included SBST, the baseline values of the outcomes and expectation of future work ability, and model 4 including the SBST, the baseline value of the outcome, expectation of future work ability and current work ability. Finally, in model 5 we conducted a sensitivity analysis similar to model 4, but without baseline values of the outcomes. Baseline ODI values were converted to units of 10 to ease interpretation of the coefficients, meaning a change in outcome should be interpreted per 10-unit change in baseline ODI score. We report the beta coefficients from the linear regression models with 95% CIs. The low risk group in SBST was used as the reference group in all regression models. In addition, we performed sensitivity analyses including pain duration in regression model 4, as duration has been shown to influence the prognostic ability of SBST. All analyses were performed in Stata/IC V.15.1.

### RESULTS

Of the 158 patients included at baseline, 93 (59%) completed follow-up assessment at 3 months (figure 1). Four patients were classified as high risk by the SBST, but these patients were included in the combined medium/high risk group.

Table 1 presents the demographics and baseline characteristics of the total study sample and stratified by the SBST risk groups. Compared with the low risk group (n=107, 67.7%), the medium/high risk group (n=51, 32.3%) had higher symptom loads on most baseline variables and the three outcome measures at baseline. The medium/high risk group was particularly burdened by mental distress, reduced daily activity, more sleep problems, analgesics and psychosocial risk factors for long-term work disability (ie, the Örebro screening questionnaire total score).

Table 2 shows the explained variance (adjusted R²) after adding each of the explanatory variables.

The SBST alone explained 35%, 25% and 28% of the variance in disability, pain intensity and health-related quality of life, respectively. Adding expectation of future work participation to SBST had small effects on explained variance. In model 4, the explained variance of the outcomes after adding current work ability to the SBST increased considerably to 60% for disability, 49% for pain intensity and 47% for health-related quality of life. Tables 3–5 shows associations between the prognostic factors and the outcomes at 3 months for the different
models. Importantly, after including all predictors in the multivariate models, the SBST and work ability was the only variables that consistently and independently predicted outcome after treatment across all the three outcome measures. The estimated associations and explained variances remained largely similar in sensitivity analyses omitting baseline levels of the outcomes from the regression models. Pain duration did not significantly influence the prognostic ability of the SBST or the work-related factors (results not shown).

DISCUSSION

This study examined the prognostic ability of the SBST applied with two risk groups (low risk group and medium/high risk group), and if adding work-related items improved this prediction of core outcomes in patients attending primary care physiotherapy with low back pain as the main problem. The data indicated that SBST independently predicted outcome for disability, pain and quality of life outcome after treatment. Adding one item on current work ability significantly and consistently improved the predictive power of the models for disability, pain and quality of life outcome. Expectation of future work participation did not substantially add prognostic ability to the SBST.

In our study, SBST applied with two risk groups explained 35% of the variance in disability as measured by the ODI at 3 months. That SBST contribute to predict disability after treatment for low back pain is in line with previous research, although with differences in the application of SBST and in follow-up tools and timing: Studies in Germany and UK have shown that SBST applied with the original three risk groups predict future disability measured by Chronic Pain Grade Scale/disability subscale at 12 months, or Roland-Morris Disability Questionnaire at 6 months. Another study applied scores instead of risk group and indicated that overall SBST as well as the psychosocial subscore predict disability assessed by ODI at 6 months. However, the SBST added little to chiropractors’ clinically derived predictions on future disability in Denmark. Systematic reviews suggest that SBST has acceptable performance for discriminating

Table 3  Pain related disability (ODI, 0–100) at 3 months associated with STarT Back Screening Tool (SBST) and work-related factors at baseline

<table>
<thead>
<tr>
<th>Steps</th>
<th>Disability*</th>
<th>Pain intensity†</th>
<th>Health-related quality of life‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>SBST</td>
<td>0.35</td>
<td>0.25</td>
<td>0.28</td>
</tr>
<tr>
<td>SBST + baseline value of outcome</td>
<td>0.45</td>
<td>0.32</td>
<td>0.39</td>
</tr>
<tr>
<td>SBST + expectation of future work participation (0–10)</td>
<td>0.39</td>
<td>0.32</td>
<td>0.33</td>
</tr>
<tr>
<td>SBST + work ability (0–10)</td>
<td>0.60</td>
<td>0.49</td>
<td>0.47</td>
</tr>
<tr>
<td>SBST + all above prognostic factors</td>
<td>0.65</td>
<td>0.50</td>
<td>0.49</td>
</tr>
</tbody>
</table>

*Disability was assessed by the Oswestry Disability Index, range 0–100.
†Pain intensity last week was assessed by a Numerical Rating Scale, range 0–10.
‡Health-related quality of life was assessed by the EQ-5D-5L.
disability outcome, but that the tool add little explanatory information for pain and work absenteeism outcomes. However, there is a lack of agreement between individual studies’ conclusions, probably because previous studies have used different measuring instruments and domains to evaluate outcome, as well as different inclusion criteria and study settings.

SBST and work ability has also been associated with quality of life and daily life outcome in previous studies. Patients’ expectations of future work ability at 2 years, obtained at baseline, was associated with daily life outcome after 6 months in a German study of patients attending physiotherapy. Also in line with our results, a Swedish group found that SBST could predict quality of life as well as work ability in patients with acute or subacute back or neck pain.

Our point of departure was that if screening tools integrate patients’ perspective on the work dimension, it could facilitate and inform the integration of the work domain in management. The screening item from Örebro musculoskeletal pain questionnaire focused on patients’ own expectations on future work ability. The conceptual domains for this expectation item may overlap with both the SBST and the item on current work ability. For instance, in SBST patients are asked to agree or disagree to the following statements: ‘It’s not really safe for a person with a condition like mine to be physically active’ (item 5), and ‘I feel that my back pain is terrible and it’s never going to get any better’ (item 7). Similarly, the question on current work ability could probably be connected to the patient’s expectation of future work ability. This may partly explain why the expectation item did not add prognostic power in the combined prediction model (Model 3).

SBST is short and feasible for clinical use, however, bidimensional and limited to the physical and psychological domains. It has been shown to correlate well with the original as well as the short form. Örebro musculoskeletal pain screening questionnaire. However, another previous study indicated that classification agreement between short form ÖMPSQ and SBST was low and that including work items may add important information to SBST. In the present study, by adding one item on work ability, the predictive ability improved considerably. Assessing work ability can increase clinicians’ awareness and help discriminate patients in need for more extensive mapping of work-related obstacles for recovery, and thus add value to the first-contact decision-making in primary care. Work ability is a complex and multidimensional construct that depend on an interaction between person and work context. Although the work ability item used in the current study does not specify whether it is primarily environmental or individual factors that needs to be targeted, the question may inform preliminary screening of work-related issues, and spur on a patient–therapist dialogue around work. Previous studies have also indicated that self-reported work measures is connected to core outcome: In a Danish study, high self-reported physical work demand was related to sick leave, and higher perceived work ability has previously been related to less disability and pain and higher quality of life in persons with low back pain. Moreover, research has provided evidence that work-related obstacles for recovery do characterise subgroups seeking healthcare for low back pain, and more fine-tuned screening tools for work-related factors (blue flags) have been developed.

Limitations

Patients with complex multisite pain as defined by therapist were excluded from the present study due to the allocation algorithm in the umbrella study. This probably resulted in fewer high-risk patients in our sample.
We thus chose to compare the subgroup with low risk patients to a combined subgroup including both medium- and high-risk patients. When interpreting the results, one should keep in mind that there were few patients classified as high-risk in the collapsed medium/high risk group. Since mono-disciplinary treatment may be best suited for patients with less complex obstacles for recovery, the present sample can be considered representative for the majority of patients attending primary care physiotherapy. Additionally, the optimal cut-off scores for differentiating between the medium and high-risk patients is not straightforward, and risk group allocation may differ between screening instruments. Both patients classified as medium and as high-risk are recommended primary care treatment unless too complex obstacles for recovery. Thus, it may be argued that the cut-off between low and medium risk patients is the most important in this sample. We found, however, no significant differences in patient characteristics at baseline between responders and non-responders at 3 months, indicating limited attrition bias. However, we cannot rule out that responders and non-responders differed for other characteristics.

**Implications**

Our finding that perceived current work ability independently and consistently predict outcome across three core outcome domains, underline the need to address work ability specifically in low back pain management. Adding this work item to the decision support tool SBST may further assist the tailoring of management in primary care and contribute to better integration of the work domain in the initial assessment. In line with this, a recent review indicated that many patients prefer therapists that communicate with their employer, since they experience a social pressure to return to work. Work ability is potentially a modifiable prognostic indicator that can be addressed in the primary care and contribute to better integration of the work domain in the initial assessment. A recent review indicated that many patients prefer therapists that communicate with their employer, since they experience a social pressure to return to work.

### Table 5

<table>
<thead>
<tr>
<th>Health-related quality of life (EQ-5D, −0.285 to 1) at 3 months associated with the STarT Back Screening Tool (SBST) and work-related factors at baseline</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mean difference in health-related quality of life (95% CI)</strong></td>
</tr>
<tr>
<td><strong>Model 1</strong></td>
</tr>
<tr>
<td><strong>SBST</strong></td>
</tr>
<tr>
<td>Low risk</td>
</tr>
<tr>
<td>Medium/high risk</td>
</tr>
<tr>
<td>Work related factors (0-10)</td>
</tr>
<tr>
<td>Expectations, per unit</td>
</tr>
<tr>
<td>Work ability, per unit</td>
</tr>
<tr>
<td>EQ-5D at baseline, per unit</td>
</tr>
<tr>
<td>Explained variance, adjusted R²†</td>
</tr>
</tbody>
</table>

*Model 1=univariable analyses.
†Model 2=SBST + baseline pain intensity.
‡Model 3=model 2 + expectations.
§Model 4=model 3 + work ability.
¶Model 5=model 4 without baseline EQ-5D (sensitivity analysis).

EQ-5D, EuroQol.
Acknowledgements The authors wish to thank the patients and therapists that contributed to the data collection and the Norwegian Fund for Post-Graduate Training in Physiotherapy.

Contributors All authors contributed in the design and discussion of the study, and with comments on the manuscript. IM performed the statistical analysis supported by TILN and OV and MU-T wrote the manuscript draft.

Funding The FYSIOPRIM research programme is funded by the Norwegian Fund for Post-Graduate Training in Physiotherapy. Grant number 360.0.1.

Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not required.

Ethics approval The FYSIOPRIM project was approved by the regional committees for Medical and Health Research Ethics in Norway (REC no. 2013/2030), and the approval covers the assignment in the present study. Informed written consent was obtained from the study participants. The funders role was limited to financial support only. The funder played no role in the design, conduct or reporting of this study.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request.

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