24-item Roland-Morris Disability Questionnaire was preferred out of six functional status questionnaires for post-lumbar disc surgery

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Abstract

Objective: Measurement properties of questionnaires should be based on samples of populations on whom these measurements will be used. The purpose of this study is to establish an evidence based recommendation regarding the use of functional status questionnaires in patients following a lumbar disc surgery by a direct comparison of the reproducibility and responsiveness.

Study Design and Setting: The measurement properties of six functional status questionnaires were assessed: 1) Roland-Morris Disability Questionnaire (RDQ-24), 2) Modified Roland-Morris Disability Questionnaire (MRDQ), 3) short Roland-Morris Disability Questionnaire (RM-18), 4) Physical Functioning scale, 5) Role Limitations-Physical scale of the SF-36, and 6) The Main Complaint (MC). Subjects (n = 97) that still suffered residual complaints 6 weeks following a lumbar disc surgery completed the questionnaires before and 3 months after treatment. In a direct comparison the A) The test-retest reproducibility (Intraclass Correlation Coefficients [ICC] and the Standard Error of Measurement [SEM]) and B) 3 parameters of responsiveness (Minimal Detectable Change [MDC], Standardised Response Mean [SRM], and the Area Under the receiver operator characteristic Curve [AUC]) were assessed.

Results: This study suggests the superiority of the 3 versions of the RDQ compared to the 3 other questionnaires. Comparing the 3 versions of the RDQ reveals no substantial differences thereby indicating that the 2 modified version of the RDQ hold no better measurement properties in this specific population.

Conclusion: The use of the RDQ-24 for this specific post-surgery population is suggested. The optimal cut-off point of the RDQ-24 that minimizes the overall classification error was found to be 3.5 with a sensitivity of 94.6% and a specificity of 88.2%.

Keywords: Functional status; Low back pain; Lumbar disc surgery; Roland-Disability Questionnaire; SF-36; Measurement properties

1. Introduction

A recent systematic review that assessed the effectiveness of various rehabilitation programs for patients after lumbar disc surgery concluded that is was unclear what the exact content of active post-surgery rehabilitation should be [1]. The aim of rehabilitation programs in general is to improve the functional status of patients. Restriction of function is a patient-referenced concept that is different for each individual. Questionnaires have been developed for measuring functional status, but they were designed for patients with nonspecific low back pain. One widely used questionnaire is the Roland-Morris Disability questionnaire (RDQ) [2]. Several modifications to the RDQ have been suggested, but these modifications seem to provide only modest improvement in measurement properties that were in general considered to be satisfactory [3]. An international group of experts [4] suggested the use of the original version because in addition to the satisfactory measurement properties the RDQ has been widely used in many studies (and countries) for nonspecific low back pain.

However, measurement properties of questionnaires should be based on samples of populations on whom these measurements will be used in clinical practice. Davidson and Keating [5] argue that differences in measurement properties of two versions of the RDQ between their study and the study by Patrick et al. [6] can be due to differences in study populations. Whereas Davidson and Keating included patients that were seeking help from a physical therapist, Patrick et al. included patients with sciatica secondary to a herniated lumbar intervertebral disk. The measurement properties of the various versions of the RDQ for patients after a lumbar disc surgery have not been assessed. In addition to the various versions of the RDQ, the measurement
properties of other, widely used questionnaires, such as the Physical Functioning subscale and the Role Limitations-Physical subscale of the SF-36 [7] and the Main Complaint (MC) [8], are unknown in this specific population. The SF-36 subscales and the MC were chosen because these are relatively brief instruments. Several authors have advocated the direct comparisons of evaluative functional status questionnaires in a single patient group [5,9,10].

The purpose of this study is to establish an evidence-based recommendation regarding the use of functional status questionnaires in patients after lumbar disc surgery by a direct comparison of the reproducibility and responsiveness of the included functional status questionnaires. Reproducibility is defined as the ability to measure attributes in a consistent manner when administered on several occasions to stable subjects [11]. In other words, the question to be answered for evaluative instruments is whether the instrument produces similar results on repeated administration when no real change in health status has occurred within this time frame [12]. Lack of reproducibility can be due to random measurement error [11] and real within-subject variance [13,14]. Both components together lead to measurement fluctuations in health status in the absence of real change. These fluctuations have been termed “background noise” [12].

Responsiveness is the extent to which different results are obtained on repeated administration of the same instrument when a real change in health status has occurred [13,15–17]. There are various ways of expressing the responsiveness, and there is no consensus on the most appropriate strategy for quantifying responsiveness. In this article, responsiveness is defined as the ability to detect clinically important changes [11,18]. To explore the responsiveness from this perspective, an explicit external criterion is used to define whether a patient has deteriorated, has not changed, or has improved such that the improvement can be considered clinically relevant.

2. Methods

2.1. Study population

The study population consisted of participants (n = 105) of a randomized controlled trial concerning the effectiveness on the rehabilitation after lumbar disc surgery [19,20]. Patients were checked for eligibility for this trial by the neurosurgeon during the routine 6-week post-surgery consultation. Patients were eligible if they were between 18 and 65 years of age, if this was their first-time lumbar disc surgery, and if they still were restricted in the activities of daily life or had not yet (fully) resumed work 6 weeks after surgery. Patients with a confirmed and relevant co-morbidity that possibly affected the spine, such as morbus Bechterew, were excluded. Patient characteristics were collected at baseline. Follow-up measurement took place immediately after the treatment period (3 months after randomization). The medical ethics committee of the Maastricht University Hospital approved the research protocol.

2.2. Questionnaires

2.2.1. Functional status questionnaires

The following functional status questionnaires, completed by the patients, were taken into account for the purpose of this study. The same research assistant was present during both measurements (at baseline and after 3 months) in case patients had difficulties completing the questionnaires. Because these analyses are based on data obtained from the trial, the choice of the competing measures was determined by the design of the trial. In the trial, the original Roland-Morris Disability questionnaire (RDQ-24) was selected as main outcome measure because Beurskens [21] reported higher point estimates of change and an international group of experts [4] suggested the use of the original RDQ. Two modifications of the RDQ were administered to analyze whether these modifications had better measurement properties. We also assessed whether other questionnaires measuring functional status, as listed below, had better measurement properties as compared with the RDQ-24.

1. The original (RDQ-24) [2] contains 24 yes/no items. Patients are asked whether the statements apply to them that day (the last 24 hours). The RDQ-24 score is calculated by adding up the number of “yes” items, ranging from 0 (no disability) to 24 (maximum disability). Because in this population we expected sciatica to be prevalent, we changed the terminal phrase of each statement from “because of my back pain” to “because of my back or leg problem” as suggested by Patrick [6].

2. The Modified Roland-Morris Disability questionnaire (MRDQ) contains 23 items. Patrick [6] suggested that the responsiveness of the RDQ-24 could be increased by removing five potentially redundant items and adding four additional items relating to sexual function, daily work, expressions of concern to others, and the need to rub or hold areas that hurt.

3. The short Roland-Morris Disability questionnaire (RM-18) contains 18 items because Stratford [22] concluded that six items were redundant and measurement properties were equivalent to the RDQ-24. (Both modifications [MRDQ and RM-18] used the phrasing “because of my back or leg problem.” Scoring methods were also identical to the RDQ-24.)

4. The 10-item Physical Functioning scale of the SF-36 (version 1) [7] (SF-36 PhF) is used to measure activity limitations experienced at this moment. Every item has three response options. After transformation of the scores, the range is from 0 (maximum limitations) to 100 (no limitations).

5. The four-item Role Limitations-Physical scale of the SF-36 (version 1) [7] measures activity limitations experienced the last 4 weeks. Every item has two
response options. After transformation of the scores, the range is from 0 (maximum limitations) to 100 (no limitations).

6. The Main Complaint (MC) [8] measures the limitation that patients experience while performing activities selected at baseline in a standardized way. Patients selected three activities they performed frequently, that they perceived as important in their daily life, and that were hampered by their back or leg complaints. Patients rated the severity of these main complaints on a 100-mm visual analogue scale. For this study, only the first main complaint was used.

2.3. External criterion

In this study, the explicit external criterion exploring the responsiveness was the seven-point global perceived effect (GPE) scale (1 = completely recovered, 7 = worse than ever) that was administered at the 3-month follow-up. If a patient indicated “complete recovery” or “much improved,” the patient was coded as “improved,” which was regarded as clinically important. Patients who indicated “slightly improved,” “no change,” or “slightly worsened” were coded as “unchanged.” Patients who rated their complaints as “much worsened” or “worse than ever” were coded as “deteriorated.” Patients who indicated “complete recovery” or “much improved,” therefore, were classified as clinically important. Given this small probability, it is likely that a patient whose score exceeds the MDC has changed.

2.4. Data analysis

2.4.1. Reproducibility

Patients who reported to be unchanged were included in the test-retest analysis, assuming that these patients had no clinically relevant improvement. Because the large number of statements that patients had to respond to and the 3-month interval, it was assumed that patients were not able to recall their initial responses.

Test-retest reproducibility was assessed using the following two methods.

1. In a two-way random model, the intraclass correlation coefficients (ICC) for agreement [23] were calculated for each questionnaire as the ratio of variance between subjects and the total variance. These variance components were computed with ANOVA for random effects. The ICC ranges from 0 to 1.

2. The standard error of measurement (SEM) was calculated to express measurement error in the same units as the original questionnaire. The SEM is defined as the square root of the within-subject variance consisting of variance between measures (to account for systematic error between measurements) and the residual variance. The 95% confidence interval (CI) was calculated as described by Brennan [24].

2.4.2. Responsiveness

The responsiveness of each questionnaire was investigated in the following three ways:

1. The SEM (based on “unchanged” subjects) was used to calculate the minimal detectable change (MDC). The MDC is calculated as $1.96 \times \frac{\sigma}{\sqrt{n}}$, where $\sigma$ is the standard deviation and $n$ is the sample size. Many authors state that the MDC expresses the minimal magnitude of change, expressed in scale points, required to be 95% confident (hence the 1.96) that the observed change between the two measures (hence the $\frac{\sigma}{\sqrt{n}}$) reflects real change and not just measurement error. Because only stable patients are assessed, one does not know the likelihood that a patient has truly changed. Therefore, a more accurate interpretation of the MDC is that it expresses the magnitude of change, with a chance of less than 5%, that a patient being stable is truly stable. Given this small probability, it is likely that a patient whose score exceeds the MDC has changed. Because the explicit external criterion that is used labels patients as deteriorated, not changed or as improved such that the improvement can be considered clinically relevant, in this study the MDC can be interpreted as clinically important change.

2. Another method to investigate responsiveness is relating the magnitude of change to the variability in score [25]. The effect size statistic is calculated by taking the mean change found in a variable (the signal) and dividing it by the standard deviation (SD) of that variable (the noise). In the clinimetric literature, there is no consensus with regard to what SD to take. Some authors propose the SD of the baseline scores [18,26]; the statistic is then referred to as effect size. Others have suggested taking the SD of the mean change score of the same group [5,10,25,27]. This statistic often is referred to as standardized response mean (SRM). In this study, we calculated the SRM because we believe that measuring change is a function of the SD of the change score. SRMs were calculated for “unchanged” and “improved” subjects separately.

3. A third method of evaluating responsiveness assesses the ability of an instrument to discriminate between clinically relevant (improved) and clinically irrelevant (unchanged). If the functional questionnaires are considered as diagnostic tests for improvement, then these instruments can be described in terms of sensitivity and specificity for detecting change as established by the gold standard [13]. The receiver operator characteristic (ROC) curve is a graph of true positive (sensitivity) versus false positive (1-specificity) for each of several cut-off points in score change. The area under the ROC curve reflects the ability of the test to discriminate between subjects who have improved from subjects who are unchanged. A value of 1 for the AUC represents perfect (100%) accuracy, whereas a value of 0.5 represents chance alone [13]. For every questionnaire, a cut-off point is calculated for which sensitivity and specificity jointly minimize the total error in misclassification.
A prerequisite for measuring improvement is a prior score that does not equal the best possible score on a particular scale; the same holds true for measuring deterioration. In other words, prior scores at the floor or at the ceiling of the scale affect the ability of the scale to detect a meaningful change. Davidson and Keating [5] have coined the term “scale width” to indicate the capacity of a scale to have initial scores that are far enough onto the scale to allow detection of change in scores over time. Because we defined the MDC as the minimal required change, we examined scale width in terms of not more than 15% of the respondents within 1 MDC from the theoretical minimum or maximum of a particular scale. Scale width as defined here assumes a single measurement. Averaging measurements over occasions can reduce scale width. For all statistics, SPSS 10.1 for Windows was used.

To summarize the results, we tried to define criteria regarding the various measurement properties investigated. The scale width was only measurement property for which we found criteria in the literature. For scale width, the 15% criterion was suggested by McHorney et al. [28]. If less than 15% of the responders had initial scores within 1 MDC from one anchor of the scale, this was labeled “good”; scale width was labeled “negative” if initial scores exceeded the 15% rule. The ability of measuring improvement and deterioration were assessed separately. Our interpretation of the standardized response means (SRMs) was broadly based on the criteria for the effect size as described by Cohen [25]: ≤0.20 is regarded as “negative,” >0.20 and ≤0.50 as “doubtful,” >0.5 and <0.8 as “good,” and values of 0.80 or greater as “very good.” These values apply to the SRM for “improved” subjects, whereas for “unchanged” subjects SRM values had to be <0.50 to be rated “good,” and values of ≥0.50 were rated as “negative.” For the ICC, we consider values ≤0.60 as “negative,” >0.60 and ≤0.80 as “doubtful,” >0.80 and <0.90 as “good,” and 0.90 or greater as “very good.” For the AUC, the same criteria were used. For the SEM, we calculated the corresponding percentage of SEM related to the total score of the scale. Criteria that we used were: ≤5% was “very good,” >5% and ≤10% was “good,” >10% and <20% was “doubtful,” and values of 20% were considered as “negative”. Because the MDC is based on the SEM, no criteria for the MDC were defined.

3. Results

Of the 105 patients included in the trial, eight dropped out before the 3 months post-treatment measurement. Table 1 presents the information on the characteristics of the 97 participants with complete data on pre-treatment and post-treatment measurements.

The mean scores at pre-treatment and the mean scores at post-treatment for subjects in each of the seven categories of GPE, for the total group, and for the two clustered groups that were used in the analyses are presented in Table 2.

On the seven-point GPE, five patients rated themselves as “completely recovered,” and 51 patients rated themselves as “much improved.” Only seven patients indicated deterioration (“much worsened” or “worse than ever”). Due to their small number, they were excluded from the analysis. These results show that the direction and the magnitude in differences between baseline measurement and post-treatment measurement are as expected: The change over time for each questionnaire declined with the decreasing categories of GPE. Moreover, the results showed that there were no statistically significant differences between the change scores of the “slightly improved” group and the “no change” group. However, there were statistically significant differences between the change scores of the “slightly improved” group and the “much improved” group.

Table 3 presents all ICCs. The ICCs ranged from 0.14 (95% confidence interval [CI] 0.0–0.40) for the MC to 0.78 (95% CI 0.57–0.89) for the MRDQ. The SEM, the MDC, and the between-subject and within-subject variance components for all questionnaires are also presented in Table 3.

Table 4 shows that for the subjects that were “unchanged” (n = 34), the mean change scores were small, indicating no difference between pre-treatment and post-treatment measurement except on the MC and on the SF-36 RLPh, where scores improved by 22.2 mm (SD 24.1 mm) and 14.7 (SD 32.6), respectively, between initial measurement and post-treatment.

Table 4 shows that the mean changes of the questionnaires differed between the patients in the “improved” and “unchanged” group. SRMs in the “improved” group exceed the SRMs in the “unchanged” group. The SRMs in the improved group could all be labeled as “very good.” The RDQ24 and the MC showed the largest SRMs (2.02), whereas the SRM of the SF-36 RLPh was the smallest with 1.40. The areas under the ROC curve (AUC) showed that there were no differences between the three versions of the RDQ as indicated by the largely overlapping 95% CIs (Fig. 1). The three remaining questionnaires had smaller AUC, especially the SF-36 RLPh and the MC.

Assessing the scale width (Table 5) revealed that the three versions of the RDQ were comparable with regard to detecting improvement. For detecting deterioration, MRDQ

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Characteristics of patients (n = 97)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variable</td>
<td></td>
</tr>
<tr>
<td>Mean age in years (SD)</td>
<td>43.3 (8.8)</td>
</tr>
<tr>
<td>Sex (% female)</td>
<td>41.2%</td>
</tr>
<tr>
<td>Duration of this episode before operation</td>
<td></td>
</tr>
<tr>
<td>2–6 months</td>
<td>51.5%</td>
</tr>
<tr>
<td>7–12 months</td>
<td>39.2%</td>
</tr>
<tr>
<td>13 months or more</td>
<td>9.3%</td>
</tr>
<tr>
<td>Previous low back or leg pain (%)</td>
<td>81.4%</td>
</tr>
<tr>
<td>Work status</td>
<td></td>
</tr>
<tr>
<td>Employed</td>
<td>80.4%</td>
</tr>
<tr>
<td>Unemployed</td>
<td>5.2%</td>
</tr>
<tr>
<td>Not in labor force</td>
<td>14.4%</td>
</tr>
</tbody>
</table>
Table 2

Mean scores and standard deviations at pre-treatment (T0) and at post-treatment (T1) for subjects per category of GPE for the total group and for collapsed categories as used in analysis: “improved” and “unchanged”

<table>
<thead>
<tr>
<th>Category of GPE</th>
<th>T0</th>
<th>T1</th>
<th>T0</th>
<th>T1</th>
<th>T0</th>
<th>T1</th>
<th>T0</th>
<th>T1</th>
<th>T0</th>
<th>T1</th>
<th>T0</th>
<th>T1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Completely recovered (n = 5)</td>
<td>15.4 (2.1)</td>
<td>1.0 (2.2)</td>
<td>14.6 (2.1)</td>
<td>0.8 (2.2)</td>
<td>14.0 (2.3)</td>
<td>0.8 (2.2)</td>
<td>59.0 (14.7)</td>
<td>92.0 (7.6)</td>
<td>5.0 (11.2)</td>
<td>66.2 (20.3)</td>
<td>13.8 (12.8)</td>
<td></td>
</tr>
<tr>
<td>Much improved (n = 51)</td>
<td>12.8 (4.3)</td>
<td>4.6 (4.7)</td>
<td>12.7 (4.3)</td>
<td>4.6 (4.7)</td>
<td>10.9 (3.6)</td>
<td>4.0 (4.2)</td>
<td>56.5 (16.4)</td>
<td>81.4 (12.7)</td>
<td>5.9 (12.8)</td>
<td>66.6 (12.6)</td>
<td>13.8 (22.0)</td>
<td></td>
</tr>
<tr>
<td>Slightly improved (n = 23)</td>
<td>15.3 (3.6)</td>
<td>13.9 (4.2)</td>
<td>15.7 (3.8)</td>
<td>14.2 (4.2)</td>
<td>13.1 (3.1)</td>
<td>12.3 (3.5)</td>
<td>57.6 (18.5)</td>
<td>48.0 (21.0)</td>
<td>12.7 (14.1)</td>
<td>74.4 (13.9)</td>
<td>5.5 (23.5)</td>
<td></td>
</tr>
<tr>
<td>No change (n = 9)</td>
<td>15.7 (3.6)</td>
<td>15.9 (4.1)</td>
<td>15.7 (3.8)</td>
<td>16.0 (4.2)</td>
<td>13.1 (3.1)</td>
<td>12.5 (3.5)</td>
<td>50.0 (21.0)</td>
<td>47.1 (21.0)</td>
<td>11.7 (14.1)</td>
<td>90.0 (14.1)</td>
<td>5.5 (23.5)</td>
<td></td>
</tr>
<tr>
<td>Much worsened (n = 6)</td>
<td>14.3 (3.6)</td>
<td>15.8 (4.2)</td>
<td>14.5 (3.8)</td>
<td>16.0 (4.6)</td>
<td>12.2 (3.4)</td>
<td>13.5 (3.3)</td>
<td>45.8 (16.9)</td>
<td>35.8 (21.3)</td>
<td>16.7 (20.4)</td>
<td>63.0 (16.1)</td>
<td>24.8 (18.6)</td>
<td></td>
</tr>
<tr>
<td>Worse than ever (n = 1)</td>
<td>22.0</td>
<td>23.0</td>
<td>22.0</td>
<td>23.0</td>
<td>18.0</td>
<td>18.0</td>
<td>5.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>75.0</td>
<td>82.0</td>
</tr>
</tbody>
</table>

Total group (n = 97)             | 14.0 (4.2) | 8.7 (6.3) | 14.3 (3.9) | 7.6 (4.5) | 14.6 (3.8) | 8.8 (5.0) | 47.3 (18.7) | 50.0 (17.1) | 6.2 (14.0) | 74.5 (19.2) | 43.4 (29.8) |

Categories used in analysis

| Improved (n = 56)               | 13.0 (2.1) | 4.3 (4.7) | 12.9 (4.2) | 4.3 (4.7) | 11.2 (3.6) | 3.8 (4.2) | 56.7 (16.1) | 82.3 (12.7) | 5.8 (12.6) | 62.1 (14.1) | 26.6 (19.7) |
| Unchanged (n = 34)              | 15.4 (3.7) | 14.3 (3.9) | 15.7 (3.7) | 14.6 (3.8) | 13.4 (3.1) | 12.5 (3.3) | 46.5 (17.3) | 45.8 (16.4) | 5.1 (14.8) | 69.9 (13.9) | 65.0 (25.5) |

Abbreviations:

GPE, global perceived effect; RDQ-24, Roland-Morris Disability Questionnaire; MRDQ, Modified Roland-Morris Disability Questionnaire; RDQ-18, Roland-Morris Disability Questionnaire 18 items; SF-36 PhF, Physical Functioning scale SF-36; SF-36 RLPh, Role Limitations-Physical scale SF-36; MC, Main Complaint.

a Standard deviations shown in brackets.

4. Discussion

In this study, the reproducibility and responsiveness of the six questionnaires measuring functional status in patients after a lumbar disc surgery were assessed in a direct comparison. The reproducibility was assessed in the test-retest analysis including “unchanged” patients. The ICC values of the three versions of the RDQ were comparable, ranging from 0.74 (RDQ-24) to 0.78 (MRDQ). The magnitude of these ICCs was labeled as “doubtful.” The ICCs of the other three questionnaires were all labeled as “negative.” The values of the SEM showed that the three versions of the RDQ were similar and superior to the other three functional status questionnaires. This indicates that none of the questionnaires included in this project was superior to the RDQ-24.

Responsiveness was defined as the ability to detect clinically important changes. For all three versions of the RDQ, the MDCs were smaller than the other three questionnaires, meaning that these are more sensitive in detecting a clinically important change. The MDC (expressed in percentages of the scale range) for the MC and the SF-36 RLPh were 63.6% and 69.3%, respectively. The SRM was calculated and the AUC were assessed to evaluate responsiveness in terms of sensitivity to change and specificity to change. The results of both strategies led to the same conclusion: The three versions of the RDQ performed better in discriminating between “improved” and “unchanged” as compared with both the subscales of the SF-36 or the MC. The SRM of the main complaint for improvement was comparable to the SRM of the RDQ-24; however, also in “unchanged” patients, the SRM of the MC was large. Most likely the MC is measuring something other than or in addition to the outcome of interest. Finally, because of floor and ceiling effects, the MC and the SF-36 RLPh might have serious difficulties in detecting deterioration, whereas the MRDQ only slightly exceeds the 15% rule for detecting deterioration.

Results from this study suggest that the three versions of the RDQ are superior to the two subscales of the SF-36 and the MC regarding the reproducibility and the responsiveness. Comparing the three versions of the RDQ reveals only small differences. The MRDQ has a slightly higher ICC value. The SEMs, and consequently the MDCs, are comparable, but the MRDQ slightly exceeds the 15% rule at the upper end of the scale. The RDQ-24 is slightly better in discriminating between “unchanged” and “improved” as indicated by the values of the SRMs and the AUCs.

The measurement properties were assessed in this specific post-surgery population because measurement properties are not fixed numbers but are highly dependent on the patient group, including diagnosis and stage and the timing of data.
collection. However, our study shows that the ICC and the MDC of the original RDQ (RDQ-24) in this population are comparable with studies including patients with chronic low back pain [3]. There have been several proposals for modifications of the original RDQ. These modifications seem to provide only modest improvement in measurement properties that were in general considered to be satisfactory [3], and an international group of experts [4] suggested the use of the original RDQ version because, in addition to the satisfactory measurement properties, the original RDQ has been widely used in many countries for nonspecific low back pain. Results from our study show that both modifications of the original RDQ. These modifications have been widely used in many countries for nonspecific low back pain. Results from our study show that both modifications of the original RDQ. These modifications resulted in concordance with Beurskens [10]. There was support for including “slightly improved” in the “unchanged” group because there were statistically significant differences between the change scores of the “slightly improved” group and the “much improved” group. Moreover, there were no statistically significant differences between the change scores.

4.1. External criteria for measuring change

In this study, we were interested in change in functional status. We used global perceived effect as the external criteria. Our clustering into “improved” and “unchanged” groups was in concordance with Beurskens [10]. There was support for including “slightly improved” in the “unchanged” group because there were statistically significant differences between the change scores of the “slightly improved” group and the “much improved” group. Moreover, there were no statistically significant differences between the change scores.
Fig. 1. ROC curves and areas under the curve.

Table 5

<table>
<thead>
<tr>
<th>Questionnaires</th>
<th>Area</th>
<th>Std. Error</th>
<th>95% CI</th>
<th>Cut-off</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDQ-24</td>
<td>0.949</td>
<td>0.024</td>
<td>0.902-0.995</td>
<td>3.5</td>
</tr>
<tr>
<td>MRDQ</td>
<td>0.944</td>
<td>0.024</td>
<td>0.898-0.991</td>
<td>3.5</td>
</tr>
<tr>
<td>RDQ-18</td>
<td>0.948</td>
<td>0.025</td>
<td>0.899-0.977</td>
<td>2.5</td>
</tr>
<tr>
<td>SF-36 PhF</td>
<td>0.810</td>
<td>0.047</td>
<td>0.719-0.902</td>
<td>12.5</td>
</tr>
<tr>
<td>SF-36 RLPh</td>
<td>0.773</td>
<td>0.051</td>
<td>0.673-0.873</td>
<td>37.5</td>
</tr>
<tr>
<td>MC</td>
<td>0.752</td>
<td>0.056</td>
<td>0.643-0.861</td>
<td>29.5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Questionnaires</th>
<th>Percentage of subjects with initial scores that hamper detection of improvement</th>
<th>Percentage of subjects with initial scores that hamper detection of deterioration</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDQ-24</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>MRDQ</td>
<td>0</td>
<td>17</td>
</tr>
<tr>
<td>RDQ-18</td>
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<td>0</td>
</tr>
<tr>
<td>SF-36 PhF</td>
<td>12</td>
<td>10</td>
</tr>
<tr>
<td>SF-36 RLPh</td>
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<td>99</td>
</tr>
<tr>
<td>MC</td>
<td>8</td>
<td>83</td>
</tr>
</tbody>
</table>

Abbreviations: RDQ-24, Roland-Morris Disability Questionnaire; MRDQ, Modified Roland-Morris Disability Questionnaire; RDQ-18, Roland-Morris Disability Questionnaire, 18 items; SF-36 PhF, Physical Functioning scale SF-36; SF-36 RLPh, Role Limitations-Physical scale SF-36; MC, Main Complaint.

of the “slightly improved” group and the “no change” group, both of which were included in the “unchanged” group. Still, regarding the use of this external criterion, some comments have to be made. First, there is the issue of correlated error because every patient completes both the included questionnaires and the external criteria. Furthermore, Norman et al. [29] question the validity of these single-item global rating of unknown reproducibility as the standard for evaluating a multi-item tool that presumably yields measurements of superior reproducibility compared with the global rating. Their concern is that global ratings may be influenced by recall bias, for which there is some evidence [30]. Moreover, global perceived effect, as used in the current study, is an all-encompassing measure for improvement that includes
For assessing the reproducibility, we used two methods: the ICC and SEM. The ICC is generally accepted in the medical literature as the preferred method for quantifying reliability [11,18,37,39]. We calculated the ICC for agreement (as opposed to the ICC for consistency) [23] because we view differences between the two measurements in absolute scores on a questionnaire, regardless of the reason, as disagreement. However, the variance of interest with regard to the ICC is the between-subjects measures, whereas in longitudinal changes the magnitude of the within-subject variance over time is relevant [15,40,41]. Furthermore, the ICC, expressed as a dimensionless number between 0 and 1, is hardly interpretable in terms of scale points. In addition, Beckerman et al. [40] have demonstrated that reliability coefficients such as the ICC are not appropriate for gaining insight into the methodologic quality of instruments measuring change over time within a subject. Because the ICC and the related MDC are better suited for that purpose [40], these measures are presented in this study. However, the SEM and the related MDC have many faces, and even the taxonomy varies considerably [42]. Because the design of this study follows a two-way random effects model, the SEM was calculated as the square root of the within-subject variance consisting of the variance between measures and the residual variance (see Table 3). In this way, systematic disagreements between the two measurements are accounted for.

### 4.2. Various methods for assessing reproducibility

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### 4.3. Various methods for evaluating responsiveness

Three strategies have been used to assess responsiveness. We considered the MDC, based on “unchanged” subjects as a measure for responsiveness, because it defines the minimal change needed for labeling the change (with 95% confidence) as a clinically relevant change. In the present study, the MRDQ (21.7%) and the RDQ-24 (22.5%) seemed to be most sensitive to detect clinically important change. The two other strategies (SRM and ROC curves) have their advantages. The advantage of the SRM is that it is easier to calculate, although there is the controversy with regard to what SD to use. The ROC curves visualize the relation between the true-positive and false-positive rates at different cut-off points of change scores. Moreover, the ROC curve identifies the optimal cut-off point for the desired combination of sensitivity and specificity.

In conclusion, the measurement properties of the three versions of the RDQ were similar and superior to the other three functional status questionnaires, thereby indicating that none of the other included questionnaires holds better measurement properties than the RDQ-24. Based on the results of this study, we suggest the use of the RDQ-24 for this specific post-surgery population. The optimal cut-off point of the RDQ-24 that minimizes the overall classification error was found to be 3.5, with a sensitivity of 94.6% and a specificity of 88.2%.

### References


