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Psychometric Validation of the BODY-Q in Danish Patients Undergoing Weight Loss and Body Contouring Surgery

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**Background:** A well-developed patient-reported outcome instrument is needed for use in Danish bariatric and body contouring patients. The BODY-Q is designed to measure changes in important patient outcomes over the entire patient journey, from obesity to post-body contouring surgery. The current study aims to psychometrically validate the BODY-Q for use in Danish patients.

**Methods:** The process consisted of 3 stages: translation and linguistic validation, field-test, and data analysis. The translation was performed in accordance with the International Society for Pharmacoeconomics and Outcomes Research and World Health Organization guidelines, and field-test data were collected in 4 departments in 2 different hospitals. Field-test data were analyzed using Rasch Measurement Theory.

**Results:** A total of 495 patients completed the Danish BODY-Q field-test 1–4 times, leading to a total of 681 assessments with an overall response rate at 76%. Cronbach α values were ≥ 0.90, and person separation index values were in general high. The Rasch Measurement Theory analysis provided broad support for the reliability and validity of the Danish version of the BODY-Q scales. Item fit was outside the criteria for 34 of 138 items, and of these, 21 had a significant chi-square P value after Bonferroni adjustment. Most items (128 of 138) had ordered thresholds, indicating that response options worked as intended.

**Conclusion:** The Danish version of the BODY-Q is a reliable and valid patient-reported outcome instrument for use in Danish bariatric and body contouring patients. (Plast Reconstr Surg Glob Open 2017;5:e1529; doi: 10.1097/GOX.0000000000001529; Published online 20 October 2017.)

INTRODUCTION

In Denmark, between 2003 and 2015, there were approximately 16,400 bariatric procedures performed.¹² Many patients who have bariatric surgery develop excess skin as a consequence of massive weight loss. Body contouring with removal of excess skin is often needed to fully restore a patient’s body image and health-related quality of life (HR-QOL).³

To be able to measure change in HR-QOL along the patient weight loss journey, a well-developed patient-reported outcome (PRO) instrument is needed. Previous literature concerning PRO instruments for bariatric and/or body-contouring surgery has identified the need for a new and comprehensive PRO instrument.⁴⁻⁹ In a recent review, Gilmartin et al.¹⁰ also call for a new PRO instrument designed for the patient group and point to the newly published BODY-Q as a possible candidate tool.¹¹
The Moorehead-Ardelt Quality of Life Questionnaire is the PRO instrument that is included in the Danish Bariatric Surgery Database to document change in HR-QOL for Danish patients having bariatric and body-contouring surgery. We have previously described the limitations of the Moorehead-Ardelt Quality of Life Questionnaire in the Danish system and the need for a more appropriate PRO instrument for use in monitoring outcomes following bariatric and body contouring surgery.

The BODY-Q is a new PRO instrument designed for weight loss and body-contouring treatments. The BODY-Q was specifically designed to measure changes in important patient outcomes over the entire patient journey, from obesity to post-body contouring surgery. The BODY-Q consists of 18 independently functioning scales grouped into 3 main domains: appearance, HR-QOL, and experience of health-care. There is also an obesity-specific symptom checklist; however, it is important to notice that the symptom checklist is not scored as a scale, but rather as a set of independent items measuring symptoms.

INTERNATIONAL RECOMMENDED GUIDELINES FOR ITEM GENERATION, ITEM REDUCTION, AND PSYCHOMETRIC EVALUATION WERE FOLLOWED IN THE DEVELOPMENT OF THE BODY-Q. A STRENGTH OF THE BODY-Q IS THE USE OF RASCH MEASUREMENT THEORY (RMT), WHICH CONTRIBUDES TO DEVELOPING INTERVAL LEVEL MEASUREMENT SCALES.

The BODY-Q was developed and field-tested in the United States, Canada, and United Kingdom, which limit its immediate generalization to other non–English-speaking countries. When adapting a PRO instrument for use in another country and/or culture, it is essential to start with an appropriate translation and linguistic validation study, followed by psychometric validation. Due to the lack of a comprehensive and robust PRO instrument for patients who undergo bariatric and body-contouring treatments in Denmark, we translated and evaluated the BODY-Q in a large sample of Danish patients having bariatric and/or body contouring. In this article, we describe the findings from the Danish field-test and psychometric validation study.

METHODS AND MATERIALS

The study was approved by the Danish Data Protection Agency before commencement. Ethics approval was applied for at the Regional Scientific Ethical Committee for Southern Denmark, who found no reason for approval, as our study was an interview and questionnaire survey.

The process consisted of 3 stages described further underneath: translation and linguistic validation, field-test, and data analysis.

Stage I: Translation, Cultural Adaption and Linguistic Validation

The translation and linguistic validation was performed according to recommendations of the International Society for Pharmacoeconomics and Outcomes Research and the World Health Organization and is described in detail elsewhere. Briefly, 2 independent forward translations were produced by a professional translator and a clinician, both of whom had Danish as their mother tongue and were fluent in English. The harmonized forward translation was provided to a professional translator who had English as their mother tongue and was fluent in Danish who performed a back translation. The back translation was then compared with the original English version. Discrepancies were discussed with the instrument’s developers and revised in an iterative manner until a final version was produced. An expert panel meeting was held to ensure the BODY-Q included all clinically relevant issues from the perspective of Danish clinicians. We then conducted 2 rounds of cognitive interviews with a total of 22 patients. These interviews were used to determine if any aspect of the BODY-Q was poorly worded or asked about issues not relevant to patients in Denmark. Feedback from patients and proof reading by 2 clinicians was used to revise and finalize the Danish version of the BODY-Q.

Stage II: Field-Test

The BODY-Q and relevant demographic and clinical questions were developed into an online REDCap (i.e., Research Electronic Data Capture) survey (http://project-redcap.org). Researchers at Odense University Hospital are granted access to REDCap through the Odense Patient data Explorative Network. In the period June 2015 to June 2016, 4 groups of patients were recruited from Odense University Hospital (Department of Endocrinology and Department of Plastic Surgery) and Hospital of Southwest Jutland (Bariatric Center/Department of Endocrinology and Department of Plastic Surgery):

1. Prebariatric surgery: This group included patients who were referred for obesity treatment and/or bariatric surgery.
2. Postbariatric surgery: This group included patients who had bariatric surgery. Participants were asked to complete the BODY-Q survey at the following intervals: 4–5, 12, and 24 months postsurgery.
4. Postbody contouring: Patients who had undergone body contouring following massive weight loss. Participants had undergone at least one of the following procedures: abdominoplasty, upper arm lift, thigh lift, buttocks lift, and/or breast lift. Patients were asked to complete the BODY-Q survey 3 and 12 months after body contouring.

Patients scheduled for an outpatient appointment were sent an information letter about the study, with a link to access and complete the BODY-Q survey in REDCap. The Hospital of Southwest Jutland, patients were sent mobile text reminders before their scheduled outpatient clinic appointment. At both hospitals, patients attending an outpatient appointment who had not completed the BODY-Q survey before their appointment were invited to do so using a tablet in the outpatient clinic. Patients were, according to agreement with the Danish Data Protection Agency, asked electronically to provide informed consent.
6. Differential Item Functioning: Differential item functioning (DIF) refers to the stability of an instrument, that is, whether an item is responded to differently by subgroups within a population. To examine DIF, the BODY-Q dataset from the original study sample (Canada, the United States, and the United Kingdom) was included alongside the Danish dataset and analyzed in RUMM2030. We examined DIF by country to determine if the Danish sample differed from the original study sample. Items with potential DIF were located by significant chi-square values after Bonferroni adjustment. If DIF was located, the sample was split by DIF and then looked at to see if it made any difference.

7. Correlation to original scoring: To examine whether the original BODY-Q scoring key can be used, we correlated the logit scores for each scale’s set of items with for the Danish study sample and original study sample (Canada, the United States, and the United Kingdom).

**RESULTS**

A total of 495 patients completed the Danish BODY-Q field test between 1 and 4 times. Including data for patients who filled out the experience of care scales twice (i.e., once in relation to their experience of care in weight loss clinic and once for their experience in relation to their experience of care in the bariatric surgery clinic), there were a total of 681 assessments. Characteristics for each patient group are outlined in Table 1. The overall response rate was 76%. For the bariatric group, the response rate was 84%, and for the body-contouring group, it was 72%. Sixteen percentage of responses were prebariatric surgery, 33% were postbariatric surgery, 34% were prebody contouring, and 17% were postbody contouring. The percentage of missing responses at item level was 1% in total.

Overall, the RMT analysis provided broad support for the reliability and validity of the Danish version of the BODY-Q scales. Appendix 1 provides the detailed RMT statistics for the 138 BODY-Q items, organized by scale, and within scale, by the item order based on the Rasch model in the original study (Appendix, Supplemental Digital Content 1, [http://links.lww.com/PRS/G572](http://links.lww.com/PRS/G572)). For item fit statistics, item fit was outside the criteria of -2.5 to +2.5 for 34 of 138 items, and of these, 21 had a significant chi-square P value after Bonferroni adjustment.

Most items (128 of 138) had ordered thresholds. Of the 10 disordered items, all except one item in the Physical Function scale [Bending over (e.g., to tie your shoes)] was in a patient experience scale, including the Doctor scale (item 100, 101, 102, 103, 105, and 106) and Information scale (item 129, 130, and 132).

Table 2 provides the scale performance findings. In terms of scale reliability, Cronbach α values were ≥ 0.90 for all scales. Appearance scales and QOL scales evidence high reliability, with all PSI values (with and without extremes) ≥ 0.74 [exception Hips/Outer Thighs = 0.37 (without extremes)]. The PSI values for the patient experience scales were lower and ranged from 0.32 to 0.70 (with extremes) and 0.69–0.86 (without extremes).

Item residual correlations were above 0.30 (range, 0.31–0.52) for 10 pairs of items within 8 scales. In subtest analysis, the correlated items were found to have minor influence on scale reliability with a difference in PSI value ≤ 0.73.
Figure 1 shows examples of the person-item threshold distribution for the social (1a) and psychological (1b) function scales. Person locations are shown in the upper half, whereas item locations are located in the lower half. The figures confirm that the scales are able to provide information for all levels of the constructs measured (Figs. 1, 2).

DIF was examined by comparing the original study population (Canada, the United States, and the United Kingdom) with the Danish study population. The original study sample consisted of 965 assessments provided by 734 participants. We adjusted the sample size to 500 for the analysis, and DIF was significant for 27 of 138 items. However, when these items were split on the variable (country) with DIF, and the new person locations were correlated with the original person locations, DIF was found to have a negligible impact. For 17 of 18 scales, the Pearson correlation was $\geq 0.99$ for the calculated score. The correlation for the remaining scale (office staff) was 0.97. Our findings confirm that the original scoring key can be used in the Danish version.

The physical symptom checklist was completed 572 times. Table 3 shows frequencies for each symptom. The most frequent symptom reported to be there all the time was joint pain followed by short of breath with mild exercise and back pain.
Fig. 1. Person-item threshold distribution for the social wellbeing scale.

Fig. 2. Person-item threshold distribution for the psychological wellbeing scale.

<table>
<thead>
<tr>
<th>Symptom</th>
<th>All the Time, N (%)</th>
<th>Often, N (%)</th>
<th>Sometimes, N (%)</th>
<th>Never, N (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feeling tired during the day</td>
<td>58 (10)</td>
<td>140 (25)</td>
<td>215 (38)</td>
<td>39 (7)</td>
</tr>
<tr>
<td>Back pain</td>
<td>65 (11)</td>
<td>102 (18)</td>
<td>188 (32)</td>
<td>102 (18)</td>
</tr>
<tr>
<td>Joint pain</td>
<td>76 (13)</td>
<td>105 (18)</td>
<td>159 (28)</td>
<td>114 (20)</td>
</tr>
<tr>
<td>Leg pain or discomfort</td>
<td>46 (8)</td>
<td>102 (18)</td>
<td>156 (27)</td>
<td>147 (26)</td>
</tr>
<tr>
<td>Feeling of balance</td>
<td>27 (5)</td>
<td>57 (10)</td>
<td>194 (34)</td>
<td>174 (30)</td>
</tr>
<tr>
<td>Feeling weak</td>
<td>32 (6)</td>
<td>70 (12)</td>
<td>184 (32)</td>
<td>166 (29)</td>
</tr>
<tr>
<td>Short of breath with mild exercise</td>
<td>66 (12)</td>
<td>57 (10)</td>
<td>144 (25)</td>
<td>184 (32)</td>
</tr>
<tr>
<td>Swollen feet</td>
<td>40 (7)</td>
<td>58 (10)</td>
<td>152 (27)</td>
<td>292 (35)</td>
</tr>
<tr>
<td>Skin rash or infection</td>
<td>28 (5)</td>
<td>55 (10)</td>
<td>133 (23)</td>
<td>236 (41)</td>
</tr>
<tr>
<td>Too much perspiration</td>
<td>34 (6)</td>
<td>50 (10)</td>
<td>134 (23)</td>
<td>223 (39)</td>
</tr>
</tbody>
</table>
DISCUSSION

The findings from our study provide broad support that the BODY-Q scales were acceptable, reliable, and valid in a Danish sample of bariatric and body-contouring patients. Based on the high response rate obtained for the online questionnaire survey and the low amount of missing data at item level (despite the large number of items tested), the BODY-Q was judged to be acceptable to patients. Patients completed all relevant scales, and a subset of patients even completed the experience of care scales twice. Elsewhere, we report that patients in the cognitive interview phase of the translation process found the BODY-Q to be acceptable and that they were happy to fill it out and even relieved to have the opportunity to express how they felt.

Within the RMT analysis, we observed some degree of misfit that we need to explore. Item fit was outside the criteria for 34 of 138 items, and of these, 21 had a significant chi-square value after Bonferroni adjustment. This only indicates that the observed responses to these items did not fit the Rasch model perfectly and that the misfit needs to be explored further. Previously a way to adjust for this was to exclude items; however, never literature suggests that reasons should be explored and explained instead. As we found all items clinically relevant and important, we did not wish to exclude any items, and working through the affected items, we found good explanations to the statistical results. For example, 1 item was about the size of the body, which can be interpreted as both weight and subjective size. Another example was items asking about how toned specific parts of the body were. In Danish toned is not the most commonly used word, and as cognitive interviews suggested, it was translated into well trained instead, which might have influenced results. For thresholds, 128 of 138 items were ordered, indicating that response options worked as intended. Most disordered items were in the Doctor and Information scale, and when comparing the item order of these scales with the original BODY-Q item order, it was in broad agreement, which provides evidence of validity.

In terms of reliability, Cronbach α values were ≥0.90 for all scales, and PSI values for all scales measuring appearance and quality of life concerns were high. Lower reliability was noted for the experience of care scales, which may warrant further examination. This finding is likely due to the finding that a high proportion of patients in the Danish sample reported scores at the ceiling on these 4 scales, particularly according to the doctor scale. Ceiling effects are important since they limit the ability to measure change and thereby measurement accuracy. The observed ceiling effects in the experience of care scales might be explained by cultural differences, a too small sample size—or simply reflection of patients’ satisfaction with treatment. These findings again could relate to the fact that the original sample is more heterogeneous, as participants were included from a range of weight loss and plastic surgery healthcare providers from 4 countries (England, Scotland, the United States, and Canada). Saiga et al. recently published a Japanese translation and psychometric analysis of the BREAST-Q and also found ceiling effect in the experience of care scales. They suggest that the ceiling effect could be associated with study design, which could also be the situation in our design, where patients were asked to fill out the BODY-Q as preparation for their follow-up visit in the outpatient clinic.

In item residual correlations, a cutoff point at 0.3 is often used, though it needs to be used carefully while the number of items potentially can influence. When looking at the scales and items, some amount of dependency must be expected; however, it has the potential to falsely inflate reliability. To investigate this further, we did a subtest analysis showing a difference in PSI value ≤0.73. If the reliability decreases in the subtest, it should be viewed as a more valid description of data. Based on this, the correlated items that we found should only have minor influence on scale reliability.

There are several limitations to our study. First, the sample was composed mainly of women, which was similar to the development sample and reflects the fact that more women come forward for bariatric and body-contouring surgery. Second, our sample does not include patients from all regions of Denmark. However, our study included 4 different departments in 2 hospitals, and as Denmark is a small country with a total population of approximately 5.7 million people, our sample must be considered representative. Finally, we did not examine test–retest reliability or responsiveness. These psychometric aspects could be the focus of future research.

CONCLUSIONS

Our study confirms that the Danish version of the BODY-Q is a valid and reliable PRO instrument for use in Danish bariatric and body-contouring patients. Furthermore, our study and analysis underlines the importance of doing a thorough translation and linguistic validation followed by field-testing and psychometric evaluation. Finally, we have demonstrated that the Danish version of the BODY-Q fulfills the need of a valid and reliable PRO instrument for the bariatric and body-contouring population in Denmark.

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