Prospective Study

Short health scale: A valid measure of health-related quality of life in Korean-speaking patients with inflammatory bowel disease

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AIM
To evaluate the short health scale (SHS), a new, simple, four-part visual analogue scale questionnaire that is designed to assess the impact of inflammatory bowel disease (IBD) on health-related quality of life (HRQOL), in Korean-speaking patients with IBD.

METHODS
The SHS was completed by 256 patients with Crohn’s disease (CD) and ulcerative colitis (UC). Individual SHS items were correlated with inflammatory bowel disease questionnaire (IBDQ) dimensions and with disease activity to assess validity. Test-retest reliability, responsiveness and patient or disease characteristics with probable association with high SHS scores were analyzed.
INTRODUCTION

Crohn’s disease (CD) and ulcerative colitis (UC) are the two most common variants of inflammatory bowel disease (IBD); both are chronic relapsing disorders that significantly affect daily life\(^1\)\(^,\)\(^2\). The biological burden of IBD had been measured using disease activity scales such as the Crohn’s disease activity index (CDAI) in CD\(^3\). However, these do not reflect the well-being of patients with chronic illness. In contrast, an important patient-reported outcome, the health-related quality of life (HRQOL), refers to the subjective perception of illness and disease impact on daily life and general well-being; the HRQOL is therefore an essential part of health assessment in patients with IBD in both clinical practice and clinical trials\(^{4,5}\). Early HRQOL scales for IBD involved extensive questions relating to specific symptoms, social function, limitation of activities, and mental health, and were time-consuming to complete and evaluate\(^\text{6}\). Since then, shorter scales have been developed, such as the short inflammatory bowel disease questionnaire (IBDQ)\(^7\), and short health scale (SHS)\(^8\). The SHS consists of four simple 100-mm visual analog scales assessing factors traditionally associated with HRQOL (symptom burden, activities of daily living, disease related worry, and sense of general well-being). The questions were designed to be openended, so that patients could score any or all aspects of their life that they felt were important to them when completing the questionnaire. The SHS was validated in Swedish-speaking\(^9\), Norwegian-speaking\(^10\), and English-speaking\(^11\) patients with IBD.

Although the incidence of IBD in Asia has increased rapidly in recent years\(^12\)\(^-\)\(^14\), only a few studies have investigated HRQOL in Korean patients with IBD\(^15\). Thus, we aimed to evaluate the SHS in Korean-speaking patients with IBD.

RESULTS

Of 256 patients with IBD, 139 (54.3%) had UC and 117 (45.7%) had CD. The correlation coefficients between SHS questions about “symptom burden”, “activities of daily living”, and “disease-related worry” and their corresponding dimensions in the IBDQ ranged from 0.62 to 0.71, compared with correlation coefficients ranging from -0.45 to -0.61 for their non-corresponding dimensions. There was a stepwise increase in SHS scores, with increasing disease activity in both CD and UC (all \(P\) values < 0.001). Reliability was confirmed with test-retest correlations ranging from 0.68 to 0.90 (all \(P\) values < 0.001). Responsiveness was confirmed with the patients who remained in remission. Their SHS scores remained unchanged, except for the SHS dimension “disease-related worry”. In the multivariate analysis, female sex was associated with worse “general well-being” (OR = 2.28, 95%CI: 1.02-5.08) along with worse disease activity.

CONCLUSION

The SHS is a valid and reliable measure of HRQOL in Korean-speaking patients with IBD.

Key words: Quality of life; Inflammatory bowel disease; Ulcerative colitis; Crohn’s disease; Disease activity

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Core tip: The short health scale (SHS) is a new, simple, four-part visual analog scale questionnaire that is designed to assess the impact of inflammatory bowel disease (IBD) on health-related quality of life (HRQOL). In Korean-speaking IBD patients, total SHS scores correlated with total IBDQ scores in both Crohn’s disease (CD) and ulcerative colitis (UC). There was a stepwise increase in SHS scores with increasing disease activity in both CD and UC. Reliability was confirmed with test-retest correlations. Thus, SHS is a valid and reliable measure of HRQOL in Korean-speaking patients with IBD.

Materials and methods

Patients

A total of 272 patients with IBD, who visited the Kangbuk Samsung Hospital and Soonchunhyang University Hospital, Bucheon between March 2016 and August 2016 were invited to participate in this study. Of these, 265 patients agreed (97.4% response rate). After excluding 9 patients whose questionnaires did not have evaluable SHS data, 256 patients were finally enrolled. This study was approved by the Institutional Review Board of Kangbuk Samsung Hospital, South Korea.

Questionnaires

QOL was measured using the SHS. The questionnaire was designed to be self-administered and patients were asked to place a mark on the 10-cm visual analog scale that they thought was appropriate to their condition (Figure 1). Scores were presented for each of four dimensions including symptom burden, social function, disease-related worry, and sense of general well-being; the scores were then added for a total score. One gastroenterologist (SKP) translated the English SHS to Korean. QOL was also determined using the 32-item IBDQ that collects data on four dimensions, including bowel symptoms, social function,
emotional function, and systemic symptoms. We used the Korean version of the IBDQ, which has been validated for IBD[15]. Disease activity at the time of questionnaire completion was assessed by consulting physicians without knowledge of the questionnaire results. To assess disease activity, the CDAI was used for CD[3], and the Mayo score for UC[16]. CD in remission was defined as a CDAI < 150, mild disease as 150-220, moderate as 220-450, and severe as > 450. UC in remission was defined as a Mayo score 0-2, mild disease as 3-5, moderate as 6-10, and severe as 11-12.

Evaluation of SHS
Validity: We assessed validity by correlating both individual SHS items and total SHS score with IBDQ dimensions and total score. The score for each SHS question should be closely associated with other HRQOL measures reflecting the same dimension of health (convergent validity), whereas the association with variables that measure other health dimensions should be less (discriminant validity). The SHS item “symptom burden” was expected to be closely associated with the IBDQ dimension “bowel symptoms”, the SHS item “social function” with the IBDQ dimension “social function”, the SHS item “disease-related worry” with the IBDQ dimension “emotional function”, and the SHS item “general well-being” with the IBDQ dimension “systemic symptoms” or “emotional function”. We also compared SHS scores according to disease activity, as it can be assumed that disease activity would influence QOL (known-groups comparison/predictive validity).

Reliability: Test-retest reliability was determined from results 2 to 8 wk apart for SHS questions in patients in remission.

Responsiveness: Changes in SHS scores in patients who remained in remission and who experienced a change in disease activity, from remission to mild to moderate activity or vice versa, were measured for responsiveness. To evaluate responsiveness, the patients were offered a second appointment 3-6 mo later, or earlier in the event of deterioration.

Influence of patient or disease characteristics on SHS: Patient or disease characteristics with probable association with high SHS scores were analyzed. High score was defined as ≥ 5 cm for each SHS question.

Statistical analysis
Continuous data were presented as medians and interquartile ranges and categorical variables were expressed as percentages. Correlations between continuous data were analyzed using Spearman’s rank correlation coefficient (rs). For test-retest reliability, total score was analyzed using the Bland-Altman plot. Differences in unpaired and paired groups were assessed using the Kruskal-Wallis test and Wilcoxon signed-rank test, respectively. To investigate the influence of patient or disease characteristics on SHS, multivariate logistic regression analysis was performed. In the multivariate analysis, variables with probable association with high scores for each of the four SHS dimensions were included. These variables were age (< 40 years, ≥ 40 years), sex (female, male), education (beyond high school, junior school), smoking status (never, previous, or current smoker), family history (yes, no), disease type (CD, UC), disease duration (≥ 5 years, < 5 years), anti-tumor necrosis factor (TNF) or azathioprine use (yes or no), history of operation (yes or no), and disease activity based on CDAI for CD and Mayo score for UC (remission, mild, moderate, severe). For each variable, the odds ratio (OR) and 95%CI were provided. Two-sided P values < 0.05 were considered to be statistically significant. Statistical calculations were performed using SPSS 18.0 (SPSS Inc., Chicago, IL, United States).

RESULTS
Patients
Demographic and disease characteristics are shown in Table 1. Among 256 patients who were finally enrolled, median age was 37 and 64% were male. Of these,
In UC, correlation coefficients between the SHS questions “symptom burden”, “activities of daily living”, and “disease-related worry” and their corresponding dimensions in the IBDQ were also higher (0.61-0.71) than for their non-corresponding dimensions (0.45-0.59). The remaining SHS question score, “general well-being”, showed closest correlation with the IBDQ score for emotional function, similar to that for the SHS question score for “worry”, with a correlation coefficient of 0.67 for CD and 0.62 for UC.

Correlations between the four SHS dimensions and disease activity scores are shown in Figure 2.

There was a stepwise increase in all SHS dimensions, with increasing disease activity. For both CD and UC 139 (54.3%) had UC and 117 (45.7%) had CD. The median duration of disease was 4.8 years.

**Evaluation of SHS Validity:** Table 2 shows correlations between the four SHS dimensions and their corresponding IBDQ items. To establish convergent validity and discriminant validity, each SHS question score should be more closely associated with corresponding than with non-corresponding dimensions of the IBDQ questionnaires. In CD, the correlation coefficients between the SHS questions “symptom burden”, “activities of daily living”, and “disease-related worry” and their corresponding dimensions in the IBDQ ranged from 0.64 to 0.71, compared with correlation coefficients ranging from -0.45 to -0.63 for their non-corresponding dimensions. In UC, correlation coefficients between the SHS questions “symptom burden”, “activities of daily living”, and “disease-related worry” and their corresponding dimensions in the IBDQ were also higher (0.61-0.71) than for their non-corresponding dimensions (0.45-0.59). The remaining SHS question score, “general well-being”, showed closest correlation with the IBDQ score for emotional function, similar to that for the SHS question score for “worry”, with a correlation coefficient of 0.67 for CD and 0.62 for UC.

Correlations between the four SHS dimensions and disease activity scores are shown in Figure 2. There was a stepwise increase in all SHS dimensions, with increasing disease activity. For both CD and UC...
patients in remission, median “disease-related worry” scores were 20 (range 0-100), and were higher than for other dimensions.

**Reliability:** To assess reliability, test-retest results for 36 patients with IBD (20 with UC, 16 with CD) in remission were investigated. The Spearman rank correlation coefficients for test-retest scores for the SHS questions “symptom burden”, “activities of daily living”, “disease-related worry”, and “general well-being” were 0.70 ($P < 0.01$), 0.63 ($P < 0.01$), 0.79 ($P < 0.01$), and 0.60 ($P < 0.01$), respectively. The correlation coefficient for the SHS total score was 0.74 ($P < 0.01$). In the Bland-Altman plot, mean difference was -2.81 (95%CI: -15.55-9.94) (Supplement Figure 1).

**Responsiveness:** Table 3 shows responsiveness in 88 patients with IBD. Seventy-five patients remained in stable remission and were reexamined after a median 4 mo. Their SHS scores remained unchanged, except for the SHS dimension “disease-related worry”, which decreased during follow-up (17.5-7.5, $P < 0.05$). A change in disease activity occurred in 21 patients; 15 were in remission at baseline, but changed to mild to moderate disease activity during the follow-up period; 6 patients with mild to moderate activity at baseline repeated the questionnaires when in remission at the median 4-mo follow-up. There was no significant change in scores in the four SHS dimensions between the two test occasions.

**Influence of other patient and disease characteristics on SHS**

In the multivariate analysis, disease activity was associated with high scores for each SHS question after adjusting for patient- or disease-related variables (Table 4). In addition, a 3.53-fold increase (95%CI: 1.18-10.56) in the adjusted odds for high score on “activities of daily living” was observed in young patients. In the SHS dimension “disease-related worry”, the adjusted odds were 4.74 (95%CI: 1.06-21.2), 2.98 (95%CI: 1.43-6.20), and 2.56 (95%CI: 1.06-6.22) among patients with more than high school education, disease duration $\geq$ 5 years, and anti-TNF use, respectively. Female sex was associated with worse “general well-being” (OR = 2.28, 95%CI: 1.02-5.08) along with worse disease activity.

**DISCUSSION**

In this study, we verified appropriate psychometric properties for the SHS in Korean patients with IBD, as were previously shown in Western patients with IBD. We assessed the SHS for validity, reliability, and responsiveness, and all were confirmed except responsiveness in patients whose disease activity had changed. Associated patient and disease factors such as age, sex, and education were identified in our study.
The SHS is a simple and standardized four-item questionnaire that is traditionally associated with HRQOL\[8,9\]. It is quick and easy to administer, and our Korean patient population appeared to have little difficulty completing the survey. Validity was confirmed by correlation between the four SHS dimensions and their corresponding IBDQ items. In both CD and UC, the SHS questions for “symptom burden” and “activities of daily living” showed a close association with their corresponding dimensions in the IBDQ, and correlation coefficients were higher than for their non-corresponding dimensions, thus establishing convergent and discriminant validity. Previous studies examined either patient ratings of IBD concerns\[8,17\] or IBDQ emotional function\[11\], and both showed high correlations with the SHS “worry” dimension. In this study, we compared the “worry” dimension with IBDQ emotional function and confirmed convergent and discriminant validity. However, for “general well-being”, previous studies examined different dimensions, i.e., either IBDQ emotional function\[8\] or IBDQ systemic symptoms\[11\]. In this study, we found close associations between the SHS “well-being” dimension for both IBDQ emotional function and systemic symptoms in CD, and emotional function in UC. This is consistent with results in Swedish-speaking patients\[8\], and different from those in English-speaking patients\[11\]. This indicated that well-being is the most comprehensive among various dimensions associated with HRQOL, which is formed by the interaction of biological, psychological, social, and economic variables in patients with IBD. Validity was also supported by the significantly worse scores in all SHS questions in patients with mild to severe disease, compared with those in remission.

The SHS good reliability, although it was assessed in a small group of patients in remission, using surveys 2 to 6 wk apart. The correlations between the two measurements were high in both CD and UC, and corresponded with the results of previous studies\[8-11\]. Although patients in stable remission showed no significant change in score, the SHS was not responsive to changes in disease activity. Patients who changed from remission to active disease or vice versa between the two follow-up visits had changes in symptom, worry, well-being, and total scores, but the results did not reach statistical significance, possibly due to the small number of patients. However, patients in stable remission showed no significant change in score, except in the worry dimension. “Disease-related worry” scores were relatively high in patients in remission, compared with scores for other dimensions, and improved after a median 4 mo. It is possible that worry will decrease when a patient is in remission for a certain period of time.

In this study, we also investigated patient and disease factors that might affect QOL. Only a study in Norwegian patients reported factors associated with SHS, and unemployment was adversely associated with SHS social function and general well-being in UC patients\[10\]. Interestingly, we found that younger age, higher education, and female sex were adversely associated with SHS activity, worry, and well-being dimensions, respectively. In the worry dimension, longer disease duration and anti-TNF use were also adversely associated. As we expected, moderate to severe disease activity was associated with worse scores in all SHS questions, after adjusting for patient- or disease-related variables, especially in the “symptom burden” dimension.

This is the first study to test and validate the SHS in Asian patients. In addition, associated patient and disease factors in the SHS were identified in our study. The open-ended nature of the four SHS questions helps clinicians determine which component of health is affected, and enables decision-making for therapeutic intervention in bowel inflammation or the need for psychological support. The SHS is comprehensive and simple to complete, and can quickly provide information for use in both clinical practice and clinical trials.

Our study had several limitations. First, most subjects were outpatients in remission. Evaluation

| Table 4 Influence of other demographic and disease factors on high short health scale score |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|
| Age < 40 (vs ≥ 40)              | 2.76 (0.79-9.56) | 3.53 (1.18-10.56) | 1.06 (0.47-3.28) | 1.67 (0.70-3.94) |
| Sex, Female (vs male)           | 0.85 (0.25-2.90) | 3.70 (0.98-7.45)  | 1.79 (0.83-3.85) | 2.28 (1.02-5.08) |
| Education, beyond high school (vs junior school) | 0.33 (0.02-5.51) | 1.73 (0.25-11.95) | 4.74 (1.06-21.2) | 1.31 (0.22-7.75) |
| Smoking (vs non or ex-smoking)  | 3.79 (0.86-16.43) | 3.43 (0.92-12.77) | 1.02 (0.33-3.16) | 2.78 (0.94-8.21) |
| Family history (vs no)          | 0.34 (0.06-1.92) | 1.93 (0.20-16.62) | 0.50 (0.14-1.80) | 0.41 (0.10-1.59) |
| Disease, CD (vs UC)             | 0.54 (0.15-1.91) | 0.52 (1.17-1.54)  | 0.81 (0.35-1.89) | 1.41 (0.58-3.43) |
| Disease duration ≥ 5 yr (vs < 5 yr) | 1.33 (0.45-3.89) | 2.12 (0.81-5.52)  | 2.98 (1.43-6.20) | 1.62 (0.77-3.45) |
| Anti-TNF use (vs no)            | 0.81 (0.18-5.59) | 1.69 (0.53-5.44)  | 2.56 (1.06-6.22) | 1.01 (0.48-2.68) |
| Azathioprine use (vs no)        | 0.55 (0.16-1.91) | 1.39 (0.49-3.94)  | 1.39 (0.62-3.08) | 1.69 (0.72-3.96) |
| Bowel operation history (vs no) | 1.19 (0.28-4.94) | 1.80 (0.50-6.47)  | 0.91 (0.40-2.06) | 1.09 (0.44-2.68) |
| Disease activity, moderate to severe (vs mild or remission) | 16.6 (4.19-66.18) | 8.02 (2.43-26.43) | 12.56 (3.66-43.09) | 6.03 (1.98-18.40) |

SHS: Short health scale; CD: Crohn’s disease; UC: Ulcerative colitis; TNF: Tumor necrosis factor.
of a population with a higher proportion of patients with moderate to severe disease is needed. Second, the number of patients who entered the study in remission and subsequently had a relapse during follow-up was small, and limited the confirmation of good responsiveness. Third, only Koreans were included, and further studies of other Asians should be performed to determine SHS reliability as an HRQOL instrument in Asia.

In conclusion, the SHS is a valid and reliable measure of HRQOL in Korean-speaking IBD patients. The SHS can be used in clinical trials and clinical practice to identify the main problems affecting QOL in IBD patients.

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COMMENTS
Background
The short health scale (SHS) was validated in Swedish-speaking, Norwegian-speaking, and English-speaking patients with inflammatory bowel disease (IBD). Although the incidence of IBD in Asia has increased rapidly in recent years, only a few studies have investigated health-related quality of life (HRQOL) in Korean patients with IBD. Thus, we evaluate the SHS in Korean-speaking patients with IBD.

Research fronts
An important patient-reported outcome, the HRQOL, refers to the subjective perception of illness and disease impact on daily life and general well-being; the HRQOL is therefore an essential part of health assessment in patients with IBD in both clinical practice and clinical trials.

Innovations and breakthroughs
This is the first study to test and validate the SHS in Asian patients. We verified appropriate psychometric properties for the SHS in Korean patients with IBD, as were previously shown in Western patients with IBD. We assessed the SHS for validity, reliability, and responsiveness, and all were confirmed except responsiveness in patients whose disease activity had changed. Associated patient and disease factors such as age, sex, and education were identified in our study.

Applications
The open-ended nature of the four SHS questions helps clinicians determine which component of health is affected, and enables decision-making for therapeutic intervention in bowel inflammation or the need for psychological support. The SHS is comprehensive and simple to complete, and can quickly provide information for use in both clinical practice and clinical trials.

Peer-review
Both the total score for SHS and IBDQ in Crohn’s disease and ulcerative colitis as well as reliability should be also presented as Bland-Altman plot.

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