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Limitations of the Knee Society Score in evaluating outcomes following revision total knee arthroplasty.

Elie Ghanem
The Rothman Institute of Orthopaedics, Thomas Jefferson University Hospital

Ian Pawasarat
The Rothman Institute of Orthopaedics, Thomas Jefferson University Hospital

Adam Lindsay
The Rothman Institute of Orthopaedics, Thomas Jefferson University Hospital

Lauren May
The Rothman Institute of Orthopaedics, Thomas Jefferson University Hospital

Khalid Azzam
The Rothman Institute of Orthopaedics, Thomas Jefferson University Hospital

See next page for additional authors

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Limitations of the Knee Society Score in Evaluating Outcomes Following Revision Total Knee Arthroplasty

By Elie Ghanem, MD, Ian Pawasarat, MA, Adam Lindsay, MD, Lauren May, MD, Khalid Azzam, MD, Ashish Joshi, MD, MPH, and Javad Parvizi, MD, FRCS

Investigation performed at The Rothman Institute of Orthopaedics, Thomas Jefferson University Hospital, Philadelphia, Pennsylvania

Background: Traditionally, the results of revision total knee arthroplasty have been determined with use of surgeon-based measures such as the Knee Society rating system. Recently, outcome and quality-of-life measures have shifted toward a greater emphasis on patient-based evaluation. The aim of our study was to determine the validity and responsiveness of the Knee Society rating system compared with the Short Form-36 health survey (SF-36), Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC), and a four-question 4-point Likert scale satisfaction questionnaire following revision total knee arthroplasty.

Methods: A total of 152 patients underwent revision total knee arthroplasty at our institution, between August 2003 and January 2007, and had a two-year follow-up evaluation after revision surgery. The SF-36, WOMAC, Knee Society rating system, and satisfaction scores were completed preoperatively and postoperatively. Spearman correlation coefficients were calculated to determine the degree of correlation for each outcome scale. The SF-36, WOMAC, and patient satisfaction were correlated with the Knee Society rating system.

Results: Both before and after surgery, the correlation among items of the Knee Society rating system displayed low to negligible levels of association. The Knee Society rating system pain score showed modest levels of convergent construct validity with the WOMAC and SF-36. However, the Knee Society functional score displayed negligible to low correlation with its WOMAC functional counterpart preoperatively. The Knee Society pain and functional scores, respectively, showed marked and moderate association with satisfaction. The change in the Knee Society pain and functional scores had moderate association with the SF-36 and WOMAC counterparts, except low correlation was displayed between the pain scores for the Knee Society rating system and the SF-36. The Knee Society rating system pain score was found to be the most responsive of the measures with a standardized response mean of 1.6, whereas the Knee Society rating system functional score was found to be the least responsive at 0.7.

Conclusions: Currently, there is no so-called gold standard that optimally reflects the status of the knee, as well as the patient, prior to and following revision total knee arthroplasty. Ideally, numerous assessment scales should be administered to the patient in order to accurately reflect the patient characteristics for the purpose of academic study, but from a practical standpoint, this may not be feasible. We encourage further research and development of a simple and concise standardized questionnaire for use before and after revision total knee arthroplasty.

Revision total knee arthroplasty has been shown to be an effective and safe surgical option for the treatment of the failed total knee arthroplasty. With an increase in the number of primary total knee arthroplasties being performed, the total number of revisions is expected to rise. One dilemma that faces the orthopaedic community is how to assess patient outcome following these complex procedures. There are currently numerous assessment tools used worldwide for the evaluation of outcome following total knee arthroplasty: the disease-specific tools (the Western Ontario and McMaster Universities Osteoarthritis Index [WOMAC], the McMaster-Toronto Arthritis Patient Preference Questionnaire, the Knee

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Society rating system, the Oxford knee score, global health tools (Short Form-36 [SF-36] health survey), and cost-toutility outcomes tools (quality-adjusted life years)\textsuperscript{16-18}. Traditionally, the results of revision surgery have been determined with use of surgeon-based measures such as the Knee Society rating system, although this outcomes tool has never been validated\textsuperscript{19}. This issue should soon be addressed as the Knee Society rating system is currently undergoing an update and validation to include objective as well as functional assessments to better reflect patient outcome following knee arthroplasty\textsuperscript{19}.

Patient-based questionnaires including the SF-36, the WOMAC, and a Likert satisfaction scale can evaluate the patient’s perception of quality of life following revision total knee arthroplasty\textsuperscript{21}. Previous studies have examined the correlation among conventional knee scoring systems, general health scores, disease-specific scores, and patient satisfaction after primary total knee arthroplasty\textsuperscript{21,22}. Given the complexity of revision total knee arthroplasty and the paucity of studies that have evaluated outcome measures after revision surgery, a thorough analysis of the different survey systems used would be useful\textsuperscript{21}.

Since the publication of the Knee Society clinical rating system in 1989, it has been broadly accepted as an objective measure of the knee in patients having total knee arthroplasty\textsuperscript{21}. The Knee Society rating system has yet to be validated or shown to be responsive for patients following revision total knee arthroplasty. Although Lingard et al. showed the Knee Society rating system to have adequate convergent construct validity, those authors also stated that the WOMAC and SF-36 are preferable for outcome assessment of total knee arthroplasty as these tools are able to be completed by the patient alone\textsuperscript{21}. The aim of our study was to determine the correlation between commonly used outcome questionnaires, specifically those that are used at our institution following revision total knee arthroplasty. We hypothesized that a marked discrepancy exists between the Knee Society rating system and the patient-centered assessment questionnaires (i.e., SF-36, WOMAC, and the satisfaction scale) in reflecting patient outcome.

**Materials and Methods**

Following institutional review board approval, a thorough review of the revision joint replacement database at our institution was performed to identify patients undergoing revision total knee arthroplasty and extract data for the 304 cases occurring between August 2003 and January 2007. The patients excluded from our final cohort included eighty-five who had a revision because of infection, thirty-five who had a revision for patellar and/or polyethylene exchanges, fifteen who had a revision for conversion of internal fixation or unicompartmental knee replacement to a total knee replacement, and four who had revision for nonprosthetic failure such as retinacular release or extensor mechanism repair. An additional thirteen patients, including nine with failure of the total knee replacement prior to the twenty-four-month follow-up, three who died, and one who was lost to follow-up, were excluded. Our final cohort after exclusion consisted of 152 patients with a mean age of sixty-seven years (range, thirty-six to eighty-nine years) who underwent revision total knee arthroplasty at our institution during the study period and completed preoperative and postoperative assessments. All patients underwent revision total knee arthroplasty because of mechanical failure, including aseptic loosening of total knee arthroplasty components in 106 patients and knee instability in forty-six patients.

There are many possible tools to measure the preoperative as well as postoperative functioning of patients after revision total knee arthroplasty, although at our institution we use the SF-36 for global assessment, the WOMAC for disease-specific evaluation, the Knee Society rating system for the knee-specific measure, and the satisfaction scale for the patient’s impression of the success of the procedure. The SF-36 is a standardized thirty-six-question form used to determine eight dimensions of health (physical functioning, physical role, bodily pain, general health, vitality, social functioning, emotional role, and mental health), which are compiled to form both physical and mental health profiles of the patient, each consisting of 100 points\textsuperscript{25-27}. This questionnaire is suited for, and was constructed for, use in health policy evaluations, general population surveys, clinical research and practice, as well as other applications involving a diverse population\textsuperscript{27}.

The WOMAC is a disease-specific measure of osteoarthritis in the lower extremities, which consists of three dimensions: pain, stiffness, and function. The WOMAC subscales include five questions on pain for a possible total of 20 points, two questions on stiffness for a possible 8 points, and seventeen questions on function for a possible 68 points, for a total score of 96 points that can be normalized to 100 points in order to facilitate comparison with other questionnaires\textsuperscript{27}. The questions surrounding function include items such as “taking off socks/stockings” and “getting in and out of a car.” Those relevant to pain include “walking on a flat surface” and “going up or down stairs.” Questions regarding stiffness include severity of stiffness in the morning as well as after inactivity later in the day.

The Knee Society score is an assessment of the knee that comprises two arms, the functional ability and clinical examination scores, each with a 100-point denominator. The functional score pertains to the functional capacity of the patient, which includes walking distance, stair-climbing ability, and the use of walking aids. This score is compiled by allocating points for stair-climbing ability and walking distance while making deductions for the use of a walking aid. A Knee Society rating system score of 100 represents unlimited walking distance and normal stair-climbing without the use of a walking aid. The clinical examination score focuses on knee motion, stability, alignment of the knee, and pain\textsuperscript{28,29}. This score is composed of 100 points, including 50 points allocated for the evaluation of motion, stability, and alignment and 50 points allocated for pain. The physical examination portion is based on the knee motion, stability, and alignment; a score of 50 indicates a minimum knee movement of 0° to 125°, without an active extension lag, without instability, and possessing normal mechanical axis alignment. Of the 50 points allocated for pain,
points are deducted with increasing frequency and severity of pain. Pain is evaluated in a single question, which may be answered with one of seven responses, indicating varying levels of pain intensity and frequency.

Patient satisfaction with the outcome of the revision total knee arthroplasty was determined with use of a previously validated four-question 4-point Likert scale. This survey includes questions regarding patient satisfaction with pain relief, improvement in function in the home and during recreation, and overall satisfaction with the procedure. The total satisfaction score is calculated as a mean of the responses to the four questions and then is converted to a 100-point scale, with 100 representing the highest level of satisfaction. Patients were considered to be very satisfied if the score was >75, satisfied if the score was >50 and £75, dissatisfied if the score was >25 and £50, and very dissatisfied if it was £25.

Preoperative baseline forms, including the SF-36, WOMAC, and Knee Society rating system, were completed by patients and physicians at a maximum of six weeks prior to revision surgery. At the two-year follow-up visit, the SF-36, the WOMAC, and the satisfaction questionnaire were completed by the patient while their surgeon completed the Knee Society rating system forms after evaluating the patient and the postoperative radiographs. The surgeon was blinded to patient responses during the preoperative and postoperative evaluation.

**Statistical Analysis**
The means and standard deviations were calculated for each outcome measure and compared with use of the t test. Spearman correlation coefficients were calculated to determine the extent of correlation of the different subscales and dimensions of the SF-36 and WOMAC with the clinical and

<p>| TABLE I Mean Preoperative and Postoperative Outcome Scores and Standardized Response Mean |
|-------------------------------------------|---------|---------|---------|---------|</p>
<table>
<thead>
<tr>
<th></th>
<th>Preop.</th>
<th>Postop.</th>
<th>Change</th>
<th>Standardized Response Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Knee Society clinical</td>
<td>34.0 (17.5)</td>
<td>64.2 (21.6)</td>
<td>29.7 (26.8)</td>
<td>1.4</td>
</tr>
<tr>
<td>Knee Society pain</td>
<td>23.0 (26.0)</td>
<td>69.7 (33.5)</td>
<td>44.9 (40.1)</td>
<td>1.6</td>
</tr>
<tr>
<td>WOMAC pain</td>
<td>51.9 (19.2)</td>
<td>24.2 (4.5)</td>
<td>-26.8 (25.4)</td>
<td>-1.3</td>
</tr>
<tr>
<td>SF-36 bodily pain</td>
<td>31.9 (20.4)</td>
<td>62.8 (26.8)</td>
<td>30.4 (31.3)</td>
<td>1.1</td>
</tr>
<tr>
<td>Functioning</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Knee Society functioning</td>
<td>41.6 (18.0)</td>
<td>60.2 (27.2)</td>
<td>16.2 (25.8)</td>
<td>0.7</td>
</tr>
<tr>
<td>WOMAC functioning</td>
<td>50.3 (20.5)</td>
<td>29.2 (21.4)</td>
<td>-20.9 (26.26)</td>
<td>-0.9</td>
</tr>
<tr>
<td>SF-36 physical functioning</td>
<td>39.2 (15.5)</td>
<td>55.6 (21.9)</td>
<td>15.9 (23.64)</td>
<td>0.9</td>
</tr>
</tbody>
</table>

*The values are given as the mean, with the standard deviation in parentheses. The scores reported are standardized to a value of 100 points to facilitate comparison. †WOMAC = Western Ontario and McMaster Universities Osteoarthritis Index, and SF-36 = Short Form-36. The preoperative scores include values of the thirteen patients excluded from the final cohort due to early failure, loss to follow-up, or death.

<table>
<thead>
<tr>
<th>TABLE II Correlations Among Items of the Clinical Score*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Item Analysis on</td>
</tr>
<tr>
<td>Knee Society Rating System</td>
</tr>
<tr>
<td>Pain</td>
</tr>
<tr>
<td>Range of motion</td>
</tr>
<tr>
<td>Anterior-posterior</td>
</tr>
<tr>
<td>Medial-lateral</td>
</tr>
<tr>
<td>Flexion contracture</td>
</tr>
<tr>
<td>Extension lag</td>
</tr>
<tr>
<td>Anatomical alignment</td>
</tr>
</tbody>
</table>

*The preoperative scores are in the white area, and the postoperative scores are in the gray area.
functional Knee Society rating system scores at the preoperative and the two-year postoperative period. A similar analysis was performed for the change in score from baseline to follow-up evaluation for the above-mentioned assessment forms. The strength of correlation was indicated by an r value, with 0 to 0.20 indicating negligible; 0.21 to 0.40, low; 0.41 to 0.60, moderate; 0.61 to 0.80, marked; and 0.81 to 1.00, high correlation. A p value of <0.05 was considered to be significant. Responsiveness was assessed with use of standardized response means, which are calculated by dividing the mean change from the preoperative state by the standard deviation of the change in score.

**Source of Funding**
There was no external source of funding for this study.

**Results**
At the two-year follow-up evaluation, the mean postoperative scores of the SF-36, WOMAC, and Knee Society rating system showed improvement from the baseline levels (Table I). All reported r values below depicting correlations among the Knee Society rating system, WOMAC, SF-36 scores, and satisfaction scores were found to be significant (p < 0.01).

**Item Analysis**
Preoperatively, the level of correlation among the clinical items of the Knee Society rating system were characterized as negligible to low, except for anterior-posterior and medial-lateral stability, which displayed a high level of correlation, and flexion contracture, which displayed a moderate level of correlation (Table II). Postoperatively, similar negligible to low levels of correlation among most Knee Society rating system items were seen, although anterior-posterior and medial-lateral stability again displayed a high degree of association. The Knee Society rating system functional ability scores showed low correlation preoperatively and moderate to marked correlation postoperatively (Table III).

**Validity**
The preoperative and postoperative correlations of the Knee Society rating system, WOMAC, SF-36, and satisfaction scores are shown in Table IV. As there is currently no validated means of assessing the clinical measurements (range of motion, alignment, and stability), it was impossible to test for convergent construct validity for the above parameters. The preoperative Knee Society pain score showed a low degree of correlation with the WOMAC pain score but a moderate level of association with the SF-36 pain score. Interestingly, the postoperative assessments show marked association of the Knee Society rating system pain score with the WOMAC pain score but only moderate correlation with the SF-36 pain score. Satisfaction also displayed a marked association with the postoperative Knee Society pain score (Table IV).

The preoperative functional scores for the Knee Society rating system displayed a negligible degree of association with the WOMAC functional score and a low level of association with the SF-36 functional score. The postoperative functional scores for the Knee Society rating system displayed a marked

| Table III Correlations Among Items of the Knee Society Functional Score* |
|-----------------------------------|---------------------|-----------------|-----------------|
| Item                              | Walking Aid (R value) | Stair Climbing | Walking Distance |
| Walking aid                      | -0.44               | -0.45          |                 |
| Stair climbing                   | -0.37               | 0.63           |                 |
| Walking distance                 | -0.33               | 0.36           |                 |

*The preoperative scores are in the white area, and the postoperative scores are in the gray area.

| Table IV Correlation Among the Knee Society Rating System, WOMAC, SF-36, and Satisfaction Scores* |
|---------------------------------------------------------------------|---------------------|---------------------|---------------------|
| Knee Society Rating System                                        | WOMAC               | SF-36               |
| Pain                   | Function | Clinical | Pain | Function | Pain | Pain | Satisfaction |
| Knee Society pain      | 0.16     | 0.43     | 0.83 | -0.59    | -0.69 | 0.47 | 0.52 | 0.62 |
| Knee Society function  | 0.79     | 0.30     | 0.33 | -0.74    | -0.60 | 0.85 | 0.49 | 0.42 |
| Knee Society clinical  | -0.28    | -0.19    | -0.24 | -0.47    | -0.54 | 0.40 | 0.46 | 0.49 |
| WOMAC function         | -0.37    | -0.10    | -0.29 | 0.64     | 0.83 | -0.81 | -0.62 | -0.60 |
| WOMAC pain             | 0.18     | 0.30     | 0.16 | -0.44    | -0.27 | 0.66 | 0.66 | 0.51 |
| SF-36 function         | 0.45     | 0.21     | 0.36 | -0.57    | -0.65 | 0.46 | 0.42 |

* R values are shown. The preoperative scores are in the white area, and the postoperative scores are in the gray area. WOMAC = Western Ontario and McMaster Universities Osteoarthritis Index, and SF-36 = Short Form-36.
association with the functional WOMAC, a high degree of correlation with the functional SF-36, and a moderate degree of correlation with satisfaction (Table IV).

**Discussion**

Traditionally, the outcome of total knee arthroplasty has been evaluated with use of physician-based scales. Recently, an increasing amount of emphasis has been placed on the use of patient-based measures of surgical outcome that can accurately reflect patient satisfaction and improvement. The determination of which outcome measures to use has been fueled by evidence suggesting the existence of disparities between physician and patient perception of success. Lieberman et al. reported significant differences in patient and physician evaluation of pain and satisfaction after total hip arthroplasty. The general assessment of physical and mental health as well as comorbidities of the patient are important variables that come into play in the evaluation of a patient with a clinically functional knee, and these factors are often lacking in the surgeon-based measurement tools.

In an effort to increase the accuracy and completeness of patient outcome measures, many investigators have chosen to combine various outcome questionnaires when evaluating patients after a revision total knee arthroplasty. However, to date, there is no so-called gold standard for this measurement. While patient self-reported measures of outcome such as the SF-36, the WOMAC, and the four-question 4-point Likert scale for patient satisfaction have previously been validated, the Knee Society clinical rating system has yet to be validated, particularly for patients after revision total knee arthroplasty.

Lingard et al. claimed the face validity of the Knee Society rating system is questionable, as patients were not included in the item-selection process and the number of selected items is limited. In the same study, it was also demonstrated that the physical examination score is subject to misrepresentative scores, as poor correlation among the items of the clinical score on the Knee Society rating system make it possible for two very different patients to receive the same score. For example, a patient who presents with an extremely stiff, pain-free, well-aligned knee will receive a score similar to a patient with mild pain, excellent knee motion, and normal alignment. These are clearly very different

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**TABLE V Correlation of the Change Among the Knee Society Rating System, WOMAC, SF-36, and Satisfaction Scores**

<table>
<thead>
<tr>
<th>Knee Society Rating System</th>
<th>WOMAC</th>
<th>SF-36</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pain</td>
<td>Function</td>
</tr>
<tr>
<td>Knee Society pain</td>
<td>0.81</td>
<td></td>
</tr>
<tr>
<td>Knee Society clinical</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Knee Society function</td>
<td>0.36</td>
<td>0.34</td>
</tr>
<tr>
<td>WOMAC pain</td>
<td>–0.39</td>
<td>0.35</td>
</tr>
<tr>
<td>WOMAC function</td>
<td>0.26</td>
<td>0.20</td>
</tr>
<tr>
<td>SF-36 pain</td>
<td>0.57</td>
<td>0.45</td>
</tr>
<tr>
<td>SF-36 function</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Satisfaction</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*R values are shown. WOMAC = Western Ontario and McMaster Universities Osteoarthritis Index, and SF-36 = Short Form-36.*
patients as the first would have difficulty with everyday tasks such as getting into and out of a car, whereas the second would have little difficulty with these tasks. Is too much being measured to be reported in a single score? Should the Knee Society rating system scores be reported in a manner akin to the Harris hip score, which reports pain as its own value? Interestingly, at our institution, we found that the components of the Knee Society functional score had much higher postoperative intercorrelation, with moderate to marked correlation, than the Knee Society pain and clinical components, which had predominantly negligible intercorrelation.

To validate the construct of a given test, it is necessary for the scores of the test under investigation to positively correlate with the gold standard. For the knee with no such standard, we are left to measure the pain and functional components of the Knee Society rating system against those of the WOMAC and SF-36. There is a caveat for the WOMAC, as a better score on the WOMAC is the score closer to zero; hence, the Knee Society rating system must negatively correlate. Correlation coefficients for convergent construct validity are often in the range of 0.2 to 0.6, and infrequently are they >0.7. It should be noted that, even in doing this, the whole of the Knee Society rating system will remain not validated for revision patients as there is no clinical scale to measure against for instability, range of motion, extension lag, and flexion contracture.

The pain component of the Knee Society rating system preoperatively correlated more strongly with the SF-36 (0.45) than with the WOMAC (–0.37). This is interesting as the Knee Society rating system was intended for patients undergoing total knee arthroplasty, while the WOMAC was intended for patients with painful osteoarthritis of the knee or hip. However, the SF-36 possesses a more general assessment for bodily pain. The postoperative values were much more in line with what was expected as the Knee Society rating system correlated more highly with the WOMAC (–0.69) than with the SF-36 (0.52). Given these values and the guidelines established by McDowell and Newell, we can categorize the strength of the convergent construct validity as modest.

The functional component of the Knee Society rating system preoperatively correlated more strongly with the SF-36 (0.21) than with the WOMAC (–0.19), which may call into question the preoperative validity of the functional Knee Society rating system as per McDowell and Newell. Postoperatively, a similar theme arose as the Knee Society rating system again correlated more highly with the SF-36 (0.85) than with the WOMAC (–0.74). This is similar to the findings in the primary total knee arthroplasty validation process and, as discussed by Lingard et al., is likely due to the similarity of the questions regarding function for the SF-36 and Knee Society rating system, which predominantly concern stairs and walking ability. Given that the preoperative correlations of the Knee Society rating system subscales with the WOMAC and the SF-36 were lower than the postoperative associations, this indicates that the Knee Society rating system is not able to adequately measure the true preoperative status of the knee in a patient requiring revision total knee arthroplasty. Another possibility is that the Knee Society rating system is surgeon rated and as such the inherent bias of the rater may lead to lower scores. Furthermore, the SF-36 does not focus on the pain that is specific to an orthopaedic procedure but on bodily pain in general, while the WOMAC attributes pain to either the hip or knee joint.

Responsiveness was gauged with use of standardized response means. Congruent with the previous findings in primary total knee arthroplasty, we found that the Knee Society rating system for pain was shown to be the most responsive with a value of 1.6 compared with 1.3 for the WOMAC and 1.1 for the SF-36, despite the fact that the WOMAC was expected to be the most responsive with the greater depth of questions regarding pain. Conversely, the responsiveness of the Knee Society rating system with regard to function was shown to be the least at 0.7, while the WOMAC and the SF-36 tied at 0.9. This is congruent with the previously reported decreased responsiveness of the Knee Society rating system compared with the SF-36 and WOMAC.

Patient satisfaction following surgery was also analyzed to assess responsiveness of the above measures by correlation. Moderate correlation was shown to exist between satisfaction and the Knee Society clinical, pain, and functional assessments as well as the WOMAC pain and functional components. A lesser degree of association (low correlation) occurred between satisfaction and the SF-36 pain and functional scores. This is logical as the satisfaction index focuses on the knee, as do both the WOMAC and the Knee Society rating system, while the SF-36 is a more general assessment.

Although the construct of our study is sound and thorough, there are a few limitations worth noting. Our analysis of patients did not include a measure of patient expectations that has previously been noted to have a profound impact on patient expectations. Furthermore, our study was conducted at a single institution and as such may lack generalizability. However, a substantial portion of the patients having a revision came from other hospitals. Our study has similar findings to those from the assessment of primary total knee arthroplasty. Although we feel confident in our evaluation of the Knee Society rating system as both valid and responsive in its ability to gauge the pain of the revised knee, there are concerns that the Knee Society rating system does not adequately assess the preoperative functional state because of the low level of association with the WOMAC. We believe this to be a weakness of the Knee Society rating system as it is not able to adequately represent the possible breadth of the general state of patients following the failed total knee replacement. Currently, there continues to be no gold standard that optimally reflects the status of the knee as well as the patient prior to and following revision total knee arthroplasty. Ideally, numerous assessment scales should be administered to the patient in order to accurately reflect the patient characteristics for the purposes of academic study, but from a practical standpoint this may not be feasible. We encourage further research and development of a simple and concise standardized questionnaire for use before and after revision total knee arthroplasty.
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Ashish Joshi, MD, MPH

Javad Parvizi, MD, FRCS
Joint Reconstructive Research,
The Rothman Institute of Orthopaedics,
Thomas Jefferson University Hospital,
925 Chestnut Street, 2nd Floor, Philadelphia, PA 19107.
E-mail address for J. Parvizi: Parvij@aol.com

References