Reliability and validity of the patellofemoral disability index as a measure of functional performance and subjective pain in subjects with patellofemoral pain syndrome

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Objective: Patellofemoral pain syndrome (PFPS) is a condition that is characterized by patellar discomfort or pain that is aggravated during certain activities such as ascending/descending stairs. The Patellofemoral Disability Index (PDI) was developed to assess the effect of pain on functional activities in individuals with PFPS. The objectives of the current study were to determine the internal consistency, test-retest reliability, and validity of this index.

Design: Cross-sectional study.

Methods: Forty-one subjects who had PFPS with a mean age of 28.8±5.0 years and a mean body mass index of 25.6±4.7 kg/m\textsuperscript{2} participated in the study. All subjects were concurrently enrolled in a clinical trial for which they were instructed to complete hamstring-resistance exercises for 4 weeks. Over the course of the intervention, they completed both the PDI and the Oswestry Disability Index (ODI) at baseline after two weeks, and after four weeks. Pearson correlation coefficient was used to assess the criterion validity. Cronbach’s $\alpha$ was used to examine the internal consistency. Intraclass correlation coefficients with 95% confidence interval were computed to examine test-retest reliability.

Results: Subjects’ responses within both the PDI and the ODI yielded Pearson correlation coefficient values that were positive and highly significant (range, 0.73-0.97; \(p<0.001\)). There was a high level of internal consistency (Cronbach’s $\alpha \geq 0.8$), with the exception of stair climbing (Cronbach’s $\alpha = 0.65$). Intraclass correlation ranged from 0.87 to 0.92, indicating high levels of test-retest reliability.

Conclusions: The PDI is a valid, reliable, and feasible method of assessing pain and functional ability in patients with PFPS.

Key Words: Anterior knee pain syndrome, Oswestry Disability Index, Patellofemoral Disability Index

Introduction

Patellofemoral pain syndrome (PFPS) is a condition which is characterized by anterior or retropatellar discomfort or pain that is aggravated during certain activities such as squatting, prolonged sitting, or ascending/descending stairs [1]. PFPS may limit activities of daily living (ADL) due to a decrease in range of motion, knee stability, and the ability to carry out ordinary activities such as walking, running, sitting down, and squatting. The most recent statistics for PFPS in the United States show a steady increase in cases, with 1.5% to 7.3% of all medical care visits being attributed to this condition [2]. Among athletes, approximately 8% to 33% of knee-related injuries are due to
this condition [3,4].

Self-perceived pain in patients with PFPS is typically assessed using subjective tools such as the Numeric Pain Rating Scale (NPRS), the Anterior Knee Pain Scale, also known as the Kujala Score, and the Oswestry Disability Index (ODI) [5-7]. However, none of these tools are specific to PFPS patients, which shows a need for more appropriate evaluation tools that focus on lower extremity pain and function rather than a general functional assessment.

The ODI is designed to quantify patients’ low back pain intensity based on their level of disability [7]. Clinicians use this index in order to get an adequate assessment of patients’ disability arising from either lower back pain or knee pain, as the two have been shown to be highly correlated and often occur simultaneously [8,9]. The interrelatedness of lower back pain and knee pain has been attributed to several factors such as lumbar lordosis, a cause of lower back pain which may be worsened by degenerative changes in the knee that limit extension of the knee joint [10] and sacral inclination alters the knee flexion angle and is associated with patellofemoral pain [11]. In addition, individuals with lower back pain are simply more prone to experiencing different types of musculoskeletal pain, including knee pain [12].

The Kujala Score was developed in 1993 by Kujala et al. [13] and colleagues with the goal of evaluating the subjective symptoms and limitations in functionality that accompany patellofemoral disorders. There are 8 questions that assess functional limitations and 5 questions that assess impairment, and it has been shown in several studies to be a valid and reliable method of assessing anterior knee pain [14-16]. While the Kujala Score is perhaps the most targeted of these pain measurement tools, it has been reported that the language may be difficult for some patients to understand, especially with regards to the questions concerning knee flexion deficiency, muscle atrophy, and unusual patellar movements [17].

The Patellofemoral Disability Index (PDI) was developed in 1995 and was first published in 1998 by Lohman [18]. This assessment tool provides physical therapists with a method of assessing functional limitations caused by peripatellar pain specifically in patients with PFPS.

The objectives of the current study are to determine the validity of the PDI, measure the internal consistency of the questions regarding sitting, standing, and walking within the PDI, and assess the test-retest reliability of the PDI.

Methods

Subjects

This is a cross-sectional study of 41 subjects who reported patellofemoral pain. Subjects were screened by qualified physical therapists under the guidance of a certified orthopedic specialist. The screening exam for patellofemoral pain required subjects to complete the Patellar Apprehension Test [19], Waldron’s Test [1], and Clarke’s Test [1]. Measurements of patellar gliding/tilting, Q angle, and observations during the step-up and step-down test were also part of the screening exam. Meniscus and ligamentous pathologies were assessed using the McMurray Test [20] and the Anterior Drawer Test [21], respectively.

Recruitment was done through the use of emails, word of mouth, and flyers. In order to be included in the study, subjects must have exhibited patellofemoral pain for at least one month with a pain level of 3 or greater on the NPRS, experienced pain during at least two of the following activities: squatting, ascending/descending stairs, and running, and be between the ages of 18 and 45 years. Subjects who reported any of the following conditions were excluded from the study: traumatic injuries to the knee joint/lower limbs, meniscus lesion or ligamentous-related pathology, neurological disorders, past and current medical history of diabetes, osteoarthritis, osteoporosis or rheumatoid arthritis, and/or use of over-the-counter pain medication during the study period. Subjects were not screened for low back pain. All subjects were briefed on the purpose and methods of the study and were required to sign an informed consent form prior to randomization. The study was approved by the Institutional Review Board at Loma Linda University and was registered at http://clinicaltrials.gov (Registration No. NCT03042559).

Instruments

The Oswestry Low Back Pain Disability Questionnaire, also referred to as the ODI, is a tool employed by physical therapists in both clinical and research settings to assess a patient’s functional disability. It is currently considered one of the principal tools for assessing functional ability and quality of life in patients with low back pain [7]. The ODI consists of 10 items that cover a variety of activities which low back pain may limit, such as pain intensity, personal care, lifting, walking, sitting, standing, sexual activity, sleeping, traveling, and socializing. It also allows subjects to rate their pain level during each distinct activity. Under each
Table 1. Ranges of Patellofemoral Disability Index questionnaire scores and relationship to levels of disability

| Class | Score % | Disability rating          |
|-------|---------|---------------------------|
| 0     | 0       | No disability             |
| 1     | 1-20    | Minimal disability        |
| 2     | 21-40   | Minimal/moderate disability|
| 3     | 41-60   | Moderate disability       |
| 4     | 61-80   | Severe disability         |
| 5     | 81-100  | Bed-bound or symptom magnification |

The PDI questionnaire uses a similar format to the ODI. It is specifically designed to assess how a patient’s knee pain has affected his/her ability to manage in everyday life, while the ODI is designed to assess the same outcomes with respect to low back pain. Both indices effectively measure functionality and clinical progress of musculoskeletal pain in the low back and lower extremities, conditions which tend to manifest as coexisting morbidities [8,9].

The PDI includes 10 items about physical activities such as running and jumping, as well as other activities including sitting, kneeling, walking, limping, stair-climbing, squatting, standing, and instability. Like the ODI, there are six options within each category and the total score corresponds to a functional ability level ranging from no disability to complete disability. The maximal possible score is 50 and then multiplied by two in order to have a maximum score of 100 to match the disability score using the ODI scale. The results of this questionnaire also correspond to the disability rating scale in Table 1. This questionnaire differs from the ODI in that it targets activities that knee pain may limit, whereas the ODI covers activities that both knee and back pain may inhibit. It has been shown that knee-specific scales are more valid and reliable for subjects with knee pain when compared to more general health assessment instruments [22].

The present study was carried out in conjunction with a clinical trial in which subjects were prescribed hamstring-resistance exercises with either a knee brace or a sport cord. All subjects completed both the ODI and the PDI at baseline, after two weeks, and after four weeks of intervention.

Table 2. Characteristics of participants (N=41)

| Characteristic | Subject          |
|---------------|------------------|
| Sex           |                  |
| Female        | 20 (48.8)        |
| Male          | 21 (51.2)        |
| Affected leg  |                  |
| Right         | 23 (56.1)        |
| Left          | 18 (43.9)        |
| Age (y)       | 28.8 (5.0)       |
| Body mass index (kg/m²) | 25.6 (4.7) |
| Pain duration (d) | 545 (30-5,475)  |

Values are presented as n (%), mean (SD) or mean (min-max).

Data analyses

Data was analyzed using the statistical package IBM SPSS Statistics ver. 25.0 (IBM Co., Armonk, NY, USA). Pearson correlation coefficient was used to assess the criterion validity of the PDI at all times during sitting, walking, and standing. Cronbach’s $\alpha$ was used to examine the internal consistency of PDI items within each type of activity. Intraclass correlation coefficients (ICCs) with 95% confidence interval (CI) were computed to examine test-retest reliability. The level significance was set at $p \leq 0.05$.

Results

Subjects’ characteristics at baseline are summarized in Table 2.

Validity

At baseline, subjects’ responses to questions regarding sitting, walking, and standing within both the PDI and the ODI yielded Pearson correlation coefficient that were positive, and highly significant. Pearson correlation coefficient ranged from 0.91 to 0.97 at baseline, 0.78 to 0.90 after two weeks, and 0.73 to 0.90 after 4 weeks ($p<0.001$; Table 3)

Internal consistency

Measures of internal consistency were determined using all ten items within the PDI. Each type of activity was found to have an overall high level of consistency with the other items in the questionnaire (Table 4). Cronbach’s $\alpha$ values were $\geq 0.8$, indicating high internal consistency. The exception to this was stair climbing, which yielded an overall alpha value of 0.65.
Table 3. Pearson correlation coefficients (r) between Patellofemoral Disability Index and Oswestry Disability Index scores

| Item     | r    | p-value |
|----------|------|---------|
| Baseline |      |         |
| Sitting  | 0.91 | <0.001  |
| Walking  | 0.91 | <0.001  |
| Standing | 0.87 | <0.001  |
| Two weeks|      |         |
| Sitting  | 0.90 | <0.001  |
| Walking  | 0.78 | <0.001  |
| Standing | 0.89 | <0.001  |
| Four weeks|     |         |
| Sitting  | 0.77 | <0.001  |
| Walking  | 0.73 | <0.001  |
| Standing | 0.90 | <0.001  |

Test-Retest reliability

Intraclass correlation of the ten PDI items was assessed at baseline, two weeks, and four weeks. The overall values with 95% CI ranged from 0.87 (0.81-0.92) to 0.92 (0.89-0.95), indicating high levels of test-retest reliability (Table 5).

Discussion

Key results & interpretation

Based on our findings, the PDI is a valid tool for assessing patellofemoral pain in subjects with PFPS when compared to ODI (the gold standard). Internal consistency was generally satisfactory, in view of the fact that the Cronbach’s α scores for each item were generally high with the exception of stair climbing. Test-retest reliability was also satisfactory, though it should be noted that the ICC was slightly greater at two weeks than at four weeks, suggesting a possible decrease in reliability over time.

Generalizability

To the best of our knowledge, this is the only study that has been done to ascertain the validity and reliability of the PDI in patients with PFPS. Therefore, it is difficult to make a direct comparison of the present study with similar studies. However, the ODI has been widely adapted to suit a variety of languages and cultures, and these modified tools have been tested for validity and reliability. Like the PDI, these tools were derived from the ODI. Therefore, results from these studies may help to give some context to the findings.
Table 5. Intraclass correlation coefficient of the ten Patellofemoral Disability Index items within each time point

| Item            | Cronbach’s α (95% confidence interval) | Baseline | Two weeks | Four weeks |
|-----------------|----------------------------------------|----------|-----------|------------|
| All ten item    |                                        | 0.87     | 0.92      | 0.91       |
|                 |                                        | (0.81-0.92) | (0.89-0.95) | (0.86-0.94) |
| Running         |                                        | 0.87     | 0.91      | 0.92       |
| Jumping         |                                        | 0.87     | 0.92      | 0.89       |
| Kneeling        |                                        | 0.87     | 0.92      | 0.89       |
| Instability     |                                        | 0.86     | 0.91      | 0.90       |
| Limping         |                                        | 0.85     | 0.91      | 0.90       |
| Stair climbing  |                                        | 0.86     | 0.93      | 0.90       |
| Squatting       |                                        | 0.86     | 0.92      | 0.90       |
| Sitting         |                                        | 0.85     | 0.91      | 0.89       |
| Walking         |                                        | 0.86     | 0.92      | 0.89       |
| Standing        |                                        | 0.86     | 0.91      | 0.89       |

of the present study.

Vigatto et al. [23] developed and tested the reliability and validity of a Brazilian-Portuguese version for detecting clinical changes in subjects with low back pain. Internal consistency was high (Cronbach’s α=0.87), as was test-retest reliability (Cronbach’s α=0.99). Using the Roland-Morris Disability Questionnaire (RMDQ) as the standard for comparison, the test was also found to be valid (r=0.81, p<0.01). The Russian version of the ODI has also been shown to have high internal consistency (Cronbach’s α=0.82), though test-retest reliability was not as strong (ICC=0.70) [24]. The Hungarian ODI has been shown to have excellent internal consistency (Cronbach’s α=0.89), test-retest reliability (ICC=0.93), and validity, as compared to the physical subscale of the World Health Organization Quality of Life, which was also good (r=0.705, p<0.001) [25]. The ODI has also been translated and adapted for those who speak Tamil, and this version also showed high internal consistency (Cronbach’s α=0.92), test-retest reliability (ICC=0.92), and validity when compared to the RMDQ (r=0.82, p=0.01) [26]. A modified version of the ODI was tested in a Korean population, and tests of internal consistency and test-retest reliability, and convergent validity yielded much lower values than those of the present study (ICC=0.43-0.80, Cronbach’s α=0.69, r=−0.54, p<0.001 for highest value, respectively) [27]. Studies assessing the same properties in the Korean and Chinese versions have found similar positive results [28,29]. The values obtained from the present study are quite similar to those of the aforementioned studies, supporting the finding that the PDI is a valid and reliable tool.

It is important to note that while the PDI and the Kujala Score are similar in terms of what they are designed to measure, there are several key differences that make the PDI more apt for assessing functionality in the target population. The utility of the PDI lies principally in the targeted nature of the 10 questions within the index. Patients are asked to assess their condition during 10 ADL, among these running, jumping, sitting, kneeling, walking, limping, stair-climbing, squatting, standing, and instability. On the other hand, the Kujala Score consists of 8 questions that evaluate functional limitations and 5 questions that evaluate impairment. Patients are asked to assess their condition through items about limping, support, walking, stair-climbing, squatting, running, jumping, sitting, pain, swelling, subluxations, atrophy of thigh, and flexion deficiency. While there is some overlap, some of the questions within the Kujala score are not specific to patellofemoral pain. Furthermore, it has been reported that subjects may have difficulty understanding some of the jargon that is used in this questionnaire [17]. PDI items in each activity appear to be very specific and intelligible, and they target PFPS-related symptoms including pain level, functional tolerance, and performance of daily activities.

The results of this study must be considered in the context of its potential limitations. The time-frame of the study (4 weeks) was relatively short, and it is possible that changes in either the positive or negative direction may have been observed for test-retest reliability had the study period been extended. Furthermore, study participants were generally sedentary individuals, therefore the results may not be generalizable to athletes who, in particular, are prone to developing PFPS.

Based on our findings, the PDI is a sufficiently valid, reliable, and feasible method of assessing subjective pain and functional ability in patients with PFPS.

Conflict of Interest

The authors declared no potential conflicts of interest with respect to the authorship and/or publication of this article.

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