Development of the Italian Version of the Oswestry Disability Index (ODI-I)
A Cross-Cultural Adaptation, Reliability, and Validity Study

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Study Design. Evaluation of the psychometric properties of a translated, culturally adapted questionnaire.

Objective. Translating, culturally adapting, and validating the Italian version of the Oswestry Disability Index (ODI-I), allowing its use in Italian-speaking patients with low back pain inside and outside Italy.

Summary of Background Data. Growing attention is devoted to standardized outcome measures to improve interventions for low back pain. A translated form of the ODI in patients with low back pain has never been validated within the Italian population.

Methods. The ODI-I questionnaire was developed involving forward-backward translation, final review by an expert committee and test of the prefinal version to establish as better as possible proper correspondence with the original English latest version (2.1a). Psychometric testing included factor analysis, reliability by internal consistency (Cronbach α) and test-retest repeatability (Intra-class Coefficient Correlation), concurrent validity by comparing the ODI-I to Visual Analogue Scale, (Pearson correlation), and construct validity by comparing the ODI-I to Roland Morris Disability Questionnaire, RMDQ, and to Short Form Health Survey, Short Form Health Survey-36 (Pearson correlation).

Results. The authors required a 3-month period before achieving a shared version of the ODI-I. The questionnaire was administered to 126 subjects, showing satisfying acceptability. Factor analysis demonstrated a 1-factor structure (45% of explained variance). The questionnaire showed high internal consistency (α = 0.855) and good test-retest reliability (ICC = 0.961). Concurrent validity was confirmed by a high correlation with Visual Analogue Scale (r = 0.73, P < 0.001). Construct validity revealed high correlations with RMDQ (r = 0.819, P < 0.001), and with Short Form Health Survey-36 domains, highly significant with the exception of Mental Health (r = −0.139, P = 0.126).

Conclusion. The ODI outcome measure was successfully translated into Italian, showing good factorial structure and psychometric properties, replicating the results of existing language versions of the questionnaire. Its use is recommended in research practice.

Key words: Oswestry Disability Index, Low Back Pain, Italian validation, outcome measures, psycho-metric properties. Spine 2009;34:2090–2095

Low back pain (LBP) constitutes a major health burden among western countries as well as a major cause of medical expenses, work absenteeism and disability. Lifetime prevalence of LBP is reported as over 70% in industrialized populations; peak prevalence occurs between ages 35 and 55.1 Nearly 85% of the patients with a pain in their low back seen by primary care practitioners are affected by nonspecific LBP, defined as a pain not attributed to recognizable specific pathologies such as fractures, nerve root compressions, infections, tumors, inflammatory or systemic diseases.2,3 Nonspecific LBP is often influenced by heavy physical work, incorrect physical activities (e.g., frequent bending, twisting, lifting, pulling and pushing, and repetitive work), wrong postures, and psychosocial factors (e.g., distress, mood alterations, cognitive dysfunction, illness behavior, job dissatisfaction).2,4

With such a high epidemiological and clinical burden, it is of great importance to apply evidence-based, validated and comprehensive outcome measures to help clinicians to quantify and improve interventions for LBP.5 As supported by most researchers, a number of disease-specific measures are available for assessing functional outcomes related to LBP6: among back related function outcome measures Oswestry Disability Index (ODI) and Roland-Morris Disability Questionnaire (RMDQ) were recommended.7

Initially developed by John O’Brien in 1976, ODI 1.0 version was first published in 1980; the questionnaire was later adapted by the American Academy of Orthopedic Surgeons, omitting sections 1, 8, and 9 and changing the score of each item from 0 to 5 to 1 to 6.9 The 2.0 version (1989) was a modification of the original scale made by a Medical Research Council group in the United Kingdom.10 Minor revisions were introduced in ODI version 2.1 (2000) for section 4 (walking); on September 2006, ODI 2.1a version was introduced, modifying instruc-

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tions of the first paragraph to consistently use present tense, as outlined at the website of the developers.11

The ODI represents a ten 6-point questionnaire. The first section rates the intensity of pain and the remaining ones cover the disabling effect of pain on typical daily activities: personal care, lifting, walking, sitting, standing, sleeping, sex life, social life, and traveling. Each item ranges form 0 to 5 and the sum of the 10 scores is expressed as a percentage of the maximum scores, varying from 0 (no disability) to 100 (maximum disability). The questionnaire is completed in about 5 minutes and scored in less than 1 minute.7

The psychometric properties of the original version of the ODI were tested in a wide variety of clinical conditions demonstrating satisfying degree of internal consistency, reproducibility as well as face, content, concurrent, and construct validity.9

To authors' knowledge, the ODI was already validated in Greek,12 Norwegian,13 Japanese,14 Turkish,15 Korean,16 Arabic,17 German,18,19 Danish,20 Iranian,21 and Brazilian.22 These studies deserved great interest as they contributed to confirm reliability and validity of the translated forms of the questionnaire, allowing comparison of results, investigating pain, disability and functional status across different people and countries.

Though the Italian version of NASS/American Academy of Orthopedic Surgeons questionnaire was published,23 a validation trial of a translated form of the ODI was never conducted within an Italian population. As this lack represented a limit for clinicians and researchers of our country to share validated outcomes, the aim of this study was to describe translation, cultural adaptation, and validation (internal consistency, reproducibility, and validity) of the Italian version of the Oswestry Disability Index (ODI-I), in its latest version named 2.1a.

Materials and Methods

The Institutional Review Board of the Institute approved the trial, allowing the development of the ODI-I.

Subjects

Outpatients referring to the Physical and Rehabilitation Medicine Units of a Research Hospital and a University Hospital of Italy were included into the study from May to September 2008.

Inclusion criteria were: common low back pain in its subacute (pain lasting more than 4 weeks) and chronic phase (pain lasting more than 12 weeks), adult age (18 years or older), ability to read and speak Italian fluently. Exclusion criteria were: acute common low back pain (included recent thoracolumbar trauma), specific causes of low back pain (disc herniation, lumbar stenosis, spinal deformity, fracture, spondylolisthesis), central or peripheral neurologic signs, systemic illness (tumor and rheumatologic diseases), psychiatric and mental deficits. Patients with recent cerebro-vascular accidents and myocardial infarctions were also excluded.

All the patients included were investigated for demographic and clinical characteristics. A specific schedule was prepared to collect main comorbidities. All the patients eligible gave their written consent to be involved into the study.

Translation and Cross-Cultural Adaptation

This stage followed the Guidelines for the Process of Cross-Cultural Adaptation of Self-Report Measures.24,25

Step 1. Forward Translation. ODI 2.1a was initially forward translated from English into Italian. The purpose was to retain the concept of the original scale, using culturally and clinically fitting expressions. Two translations were performed independently by translators with Italian as their native tongue; one of them (naive translator) was not familiar with the measure. Keeping the language colloquial and compatible with a reading age level of 14 years, poorer wording choices were outlined and resolved in a discussion between the 2 translators. With regard to section 4 (walking), walking distances described in terms of miles or yards (e.g., “1 mile,” “½ mile,” “100 yards”) were judged unfamiliar to the Italians: the distances were changed to kilometers or meters (respectively, 1 kilometer, 500 m, and 100 m). Step 1 ended when a common adaptation was shared. None of the items was excluded.

Step 2. Backward Translation. Two bilingual native English-speaking translators backward translated the initial translation; the translators were selected because unaware of concepts explored and without medical background. Taking into account cultural diversities, conceptual equivalence, or vocabulary differences, the aim was to make sure that the Italian version reflected the same item contents of the original version.

Step 3. Expert Committee. The translated versions were submitted to a bilingual committee composed of clinicians, methodologists, and psychometricians; the 4 translators were included. To identify difficulties, inconsistencies or mistakes in translation, the committee explored semantic, idiomatic, experiential and conceptual equivalence of items and answers options. Moreover, the committee took into consideration the Italian version of ODI 2.1, formerly translated by one of the authors of this work (G.Z.),11 but neither back worded nor validated. Step 3 ended when a prefinal version was achieved.

Step 4. Test of the Prefinal Version. The scale was delivered to 30 low back pain sufferers. This field test had the aim to probe what was meant by each item and the chosen response; the distribution of responses was also checked for missing items. All the findings from this step were re-evaluated by the expert committee, although no adjustment was further required.

Step 5. Submission to the Developer. The final shared version of the questionnaire (see Appendix, Supplemental Digital Content 1, http://links.lww.com/BRS/A381) was sent to the developer.

Psychometric Scale Properties

Acceptability. Time needed to answer the questionnaire was registered. Once completed, patients were asked about troubles encountered; at the same time, examiners checked all data, included missing or multiple responses.

Factor Analysis. The factor structure of the ODI-I was analyzed by means of a factor analysis. Cattell Scree Test was used to determine the number of extracted factors (eigenvalues greater than 1).

Reliability. This psychometric property was performed by means of internal consistency and test-retest stability. The first characteristic was examined with Cronbach $\alpha$ estimated for the whole questionnaire. The second characteristic was analyzed asking the patients to complete the Questionnaire 7 days later.
the first fulfillment; Intraclass Correlation Coefficient (ICC) was applied for testing agreement between baseline and 7 day ODI-I scores as well as to evaluate item by item agreement.

**Validity.** Concurrent validity was achieved comparing (Pearson correlations) ODI-I to Visual Analogue Scale (VAS)\(^26\); construct validity was achieved comparing (Pearson correlations) ODI-I to Roland Morris Disability Questionnaire (RMDQ)\(^27,28\) and to Short Form Health Survey (SF-36).\(^{29,30}\)

**Statistical Methods**

All statistical analyses were performed using SPSS, 15.0 (Italian version, Appendix 1, Supplemental Digital Content 1, http://links.lww.com/BRS/A381).

**Results**

**Subjects**

The trial included a population of 126 subjects, 73 females (58%) and 53 males (42%), mean age 47 ± 14 years (range: 18–89). Mean low back pain duration was 20.9 ± 19.6 months (range: 1–108).

Additional socio-demographic characteristics of the enrolled population are described in Table 1.

**Translation and Cross-Cultural Adaptation**

A forward backward translation was used to translate the questionnaire into Italian, involving 4 translators. A 2-month period was necessary before a culturally adapted version was reached. Further revision conducted by selected experts in spinal and psychometric research as well as a prefinal version testing (1-month length) confirmed the work done.

**Psychometric Scale Properties**

**Acceptability.** All the questions were well accepted by the patients. The questionnaire was completed in 6.61 ± 2.84 minutes (range: 2–15). Item 8 (sex life) did not receive an answer in 11.9% of patients; no other missing responses were recorded and no multiple answers were found. There were no problems of comprehension with the questions.

**Factor Analysis.** Free factor analysis revealed a 1-factor structure on the basis of the numbers of eigenvalues superior to 1, as obtained by Cattel Scree Test (Figure 1). This factor explained 45% of variance. Item-factor loadings are reported in Table 2.

**Reliability.** Internal consistency: Cronbach α index for ODI-I was 0.855. Test-retest stability: correlations between ODI at day 1 and at day 7 demonstrated highly significant (ICC = 0.961; 95% CI: 0.943–0.972). Lowest ICC value was achieved for item 3. Repeatability results are fully reported in Table 3.

**Validity.** Table 4 presents correlations between ODI-I, VAS, RMDQ and SF-36 health survey. (1) Concurrent validity. High correlation was found when ODI-I and VAS were compared (\(r = 0.730, P < 0.001\)). (2) Construct validity. High correlation was found when ODI-I and RMDQ were compared (\(r = 0.819, P < 0.001\)); high

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**Table 1. Socio-Demographic Characteristics of the Population**

| Variable       | N   | %   |
|----------------|-----|-----|
| Married        | 87  | 69  |
| Employed       | 103 | 81.7|
| Smoking        | 27  | 21.4|
| Frequency of pain |     |     |
| Sometimes      | 33  | 26.2|
| Often          | 68  | 54.0|
| Persistent     | 25  | 19.8|
| Leg pain       | 67  | 53.1|
| Drugs utilization |     |     |
| Antidepressants| 3   | 2.4 |
| Analgesics     | 50  | 39.7|
| Muscle relaxants| 2  | 1.6 |
| NSAIDs         | 28  | 22.2|
| None           | 43  | 34.1|
| Education level|     |     |
| Elementary     | 5   | 4.0 |
| Mid School     | 17  | 13.5|
| Upper school   | 59  | 46.8|
| University     | 45  | 35.7|
| Comorbidities  |     |     |
| Hypertension   | 34  | 27.0|
| NIDDM          | 9   | 7.1 |
| Heart disease  | 14  | 11.1|
| Lung disease   | 10  | 8.0 |
| Enteric disease| 11  | 8.7 |
| Liver disease  | 6   | 4.7 |
| Renal failure  | 7   | 5.5 |
| Orthopaedic disease | 25 | 19.8|
| Others         | 10  | 8.0 |

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**Table 2. Factor Analysis Loadings**

| Component | 1   |
|-----------|-----|
| Item 9    | 0.770|
| Item 10   | 0.738|
| Item 4    | 0.684|
| Item 5    | 0.673|
| Item 6    | 0.664|
| Item 7    | 0.635|
| Item 2    | 0.617|
| Item 7    | 0.562|
Table 3. Repeatability by Intraclass Coefficient Correlation (ICC), 1–7 Day

| Repeatability | ICC  | 95% CI       |
|---------------|------|--------------|
| Item 1        | 0.815 | 0.736–0.870  |
| Item 2        | 0.925 | 0.895–0.946  |
| Item 3        | 0.804 | 0.725–0.861  |
| Item 4        | 0.944 | 0.920–0.960  |
| Item 5        | 0.918 | 0.885–0.941  |
| Item 6        | 0.901 | 0.862–0.929  |
| Item 7        | 0.831 | 0.768–0.878  |
| Item 8        | 0.959 | 0.942–0.972  |
| Item 9        | 0.984 | 0.951–0.994  |
| Item 10       | 0.930 | 0.902–0.950  |
| Total         | 0.961 | 0.943–0.972  |

Table 4. Concurrent and Construct Validity. Pearson Correlations Between ODI-I, VAS, RMDQ, and SF-36 Domains

| Comparison                        | r    | P      |
|-----------------------------------|------|--------|
| ODI score vs. VAS                 | r = 0.730 | P < 0.001 |
| ODI score vs. RMDQ                | r = 0.819 | P < 0.001 |
| ODI score vs. SF-36 domains       | r = 0.751 | P < 0.001 |
| SF-36 physical activity           | r = 0.607 | P < 0.001 |
| SF-36 physical role               | r = 0.691 | P < 0.001 |
| SF-36 health in general           | r = 0.300 | P < 0.001 |
| SF-36 vitality                    | r = 0.406 | P < 0.001 |
| SF-36 social activities           | r = 0.373 | P < 0.001 |
| SF-36 emotional role              | r = 0.406 | P < 0.001 |
| SF-36 mental health               | r = 0.139 | P = 0.121 |

Osthus et al\(^{19}\) defined their factors as “pain-related activity” and “pain intensity and pain-related participation.”

Internal consistency of ODI-I demonstrated strong correlations (\(\alpha = 0.855\)). Our finding was higher than Fisher and Johnson (0.76)\(^{31}\) and similar to Kopec et al (0.87)\(^{32}\) reports in which the original English 2.0 version was implemented. Our results were in agreement to most of the other cross-cultural and adapted versions, ranging from 0.75 (Iranian version)\(^{21}\) to 0.94 (Norwegian version)\(^{13}\).

Test-retest reliability of ODI-I assessed at day 1 and at day 7 demonstrated an highly significant correlation (ICC = 0.961); moreover, each item of ODI-I reported satisfying ICC correlations, ranging from 0.804 to 0.959. Our findings were higher than Gronblad et al (0.83)\(^{33}\) and Turkish version (0.938),\(^{15}\) retested after 1 week; higher results were also found when compared to Danish version (0.91),\(^{20}\) retested after 9.1 to 12 days, while similar results were obtained when compared to Swiss-German version (0.96),\(^{18}\) retested after 6 days. Due to the natural symptom fluctuation associated with the memory effects, weaker correlations should be considered when comparing our results to Fairbank (0.99, 1-day retest),\(^{8}\) Kopec et al (0.91, 4-day retest)\(^{32}\) as well as to other non-English ODI versions, such as Norwegian (0.88, 2-day retest),\(^{13}\) Japanese (0.93, 1-day retest),\(^{14}\) Korean (0.916, 2-day retest),\(^{16}\) Arabic (0.98, 3-day retest),\(^{17}\) Iranian (0.91, 1-day retest),\(^{21}\) and Brazilian (0.99, 1-day retest).\(^{22}\)

In the determination of concurrent validity, we demonstrated a significant correlation between ODI-I and VAS (r = 0.730, P < 0.001). Also Gronblad demonstrated a positive correlation when ODI-I was compared with a VAS (r = 0.62, chronic LBP).\(^{33}\) Among translated versions, moderate to good correlations were achieved in Greek (r = 0.865, acute/chronic LBP),\(^{12}\) Norwegian (r = 0.52, chronic LBP),\(^{13}\) Turkish (r = 0.367, chronic LBP),\(^{15}\) Korean (r = 0.425, chronic LBP),\(^{16}\) Swiss-German (r = 0.78, chronic LBP),\(^{18}\) Iranian (r = 0.54, chronic LBP),\(^{21}\) and Brazilian (r = 0.66, mainly chronic LBP)\(^{22}\) populations.

In the analysis for construct validity, we compared ODI-I to RMDQ and to SF-36, respectively. Several studies compared the Oswestry Disability Index with the Roland Morris Disability Questionnaire, showing that the measures could be related; Boscainos et al\(^{34}\) and Leclaire et al\(^{35}\) found a correlation of r = 0.73 and r = 0.72, respectively. Our results supported this evidence, as the ODI-I significantly related to the Italian version of the RMDQ (r = 0.819, P < 0.001). Remarkably, our value was higher than those presented in the other translated forms (Turkish: r = 0.815; Brazilian: \(r = 0.81\); Danish: r = 0.78; Swiss-German: r = 0.80; Japanese: r = 0.785; Greek: \(\rho = 0.729\); Iranian: r = 0.71)\(^{-1}\).

The ODI was also expected to reveal correlations with health-related quality of life scales. Satisfying results were achieved when ODI-I was compared with each of the 8 domains of SF-36 questionnaire, highly significant with the exception of Mental Health (r = −0.139, P = 0.121). These findings were in agreement with other ex-
isting studies in which good correlations between ODI and SF-36 were stated. Our findings were also supported by the existing non-English ODI versions, achieving highest correlations when reporting physical function domains and moderate to low correlations when psychosocial scales were analyzed. The Norwegian version showed better score in Physical Functioning \( (r = 0.77) \) and lowest scores in Mental Health \( (r = 0.37) \), Emotional Role \( (r = -0.33) \), and Vitality \( (r = 0.28) \). The Japanese version found higher values for Physical Role \( (r = 0.721) \) and lower scores for Mental Health \( (r = 0.603) \). The German version reported better scores for Physical Function \( (r = 0.78) \), lowering when Mental Health and Emotional Role were considered \( (r = 0.52 \text{ and } 0.48 \text{ respectively}) \). The Danish version stated a significant correlation when ODI was compared to Physical Function \( (r = 0.75) \) and to Bodily Pain \( (r = 0.65) \). The Iranian version \(^{21} \) demonstrated highest scores in Physical function \( (r = 0.68) \) and lowest scores in Emotional Role \( (r = 0.38) \), Mental Health \( (r = -0.36) \), and General Health \( (r = 0.23) \). The Brazilian version found highest coefficients when Physical Function \( (r = -0.83) \) and Bodily Pain \( (r = -0.58) \) were considered and lowest scores with Vitality \( (r = -0.19) \) and Mental Health \( (r = -0.22) \).

**Conclusion**

This study described the development of translation, cultural adaptation, reliability and validity of the Italian version of ODI 2.1a, which is expected to facilitate common low back pain examination and related disability. This form is recommended to be used for research purposes, with special care to subacute and chronic adult patients, both males and females. Although not examined in the present study, the authors do not find limitations for the implementation of ODI-I in acute subjects and in other causes of low back pain.

Further studies using ODI-I are suggested to focus on minimum detectable change, minimal clinical important difference, sensitivity with specific spinal conditions and discrimination among levels of severity, as already reported for the original version. \(^7,9\)

**Key Points**

- The validation of the Italian version of ODI showed good psychometric properties, in particular test-retest stability, internal consistency and validity.
- The factor analysis suggested a 1-factor structure.
- The ODI-I showed good correlation with theVAS as well as the Italian RMDQ and SF-36.
- The ODI-I is recommended to be introduced in the field of low back pain research in Italy.

Supplemental digital content is available for this article. Direct URL citations appear in the printed text, and links to the digital files are provided in the HTML text of this article on the journal’s Web site (www.spinejournal.com).

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