

1 **ABSTRACT**

2 **BACKGROUND CONTEXT:** Most spine patient-reported outcome (PRO) measures are divided into
3 neck and back subregions. This prevents their use in the assessment of the whole spine. By contrast,
4 whole-spine PRO measures assess the spine from cervical to lumbar as a single kinetic chain. However,
5 existing whole-spine PROs have been critiqued for clinimetric limitations including concerns with
6 practicality.

7 **PURPOSE:** To develop the Spine Functional Index (SFI) as a new whole-spine PRO measure that
8 addressed the limitations of existing whole-spine questionnaires; then to determine the SFI's
9 clinimetric and practical characteristics concurrently with a recognised criterion, the Functional Rating
10 Index (FRI).

11 **STUDY DESIGN/SETTING:** Observational cohort study within ten physical therapy outpatient
12 clinics.

13 **PATIENT SAMPLE:** Spine-injured patients were recruited from a convenience sample referred by a
14 medical practitioner to physical therapy. A pilot study (n=52, 57% female, age 47.6±17.5) followed by
15 the main study (n=203, 48% female, age 41.0±17.8) that had an average symptom duration of less than
16 five weeks.

17 **OUTCOME MEASURES:** SFI, FRI and Numerical Rating Scale (NRS).

18 **METHODS:** The SFI was developed through three stages: 1) item generation, 2) item reduction with
19 an expert panel and patient focus group, then 3) pilot field testing to provide provisional clinimetric
20 properties, sample size requirements and to determine suitability for a larger study. Participants
21 completed the SFI, FRI and NRS every two weeks for six weeks, then every four weeks until discharge
22 or study completion at six months. Responses were assessed to provide individual psychometric and
23 practical characteristics for both PROs, with the overall performance evaluated by the Measurement of
24 Outcome Measures and Bot clinimetric assessment scales.

1 **RESULTS:** The SFI demonstrated high criterion validity with the FRI, (Pearson's $r=0.87$, 95% CI),
2 equivalent internal consistency ($\alpha=0.91$) and a single-factor structure. The SFI and FRI demonstrated
3 suitable reliability ($ICC_{2,1}=0.97:0.95$), responsiveness (standardized response mean= $1.81:1.68$),
4 minimal detectable change with 90% CI ($6.4\%:9.7\%$), Flesch-scale reading ease ($64\%:47\%$) and user
5 errors ($1.5\%:5.3\%$). The clinimetric performance was higher for the SFI on the Measurement of
6 Outcome Measures ($96\%:64\%$) and on the Bot scale ($100\%:75\%$).

7 **CONCLUSIONS:** The SFI demonstrated sound clinimetric properties with lower response errors,
8 efficient completion and scoring, and improved responsiveness and overall clinimetric performance
9 compared to the FRI. These results indicated that the SFI was suitable for functional outcome
10 measurement of the whole-spine in both the research and clinical settings.

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1 clinimetric properties and performance, and subsequently compared to specific subregion spine PROs
2 for the back and neck [9-11]. The development and validation of a new whole-spine PRO requires two
3 phases: 1) initial development and evaluation of clinimetrics that includes concurrent validation with an
4 existing whole-spine PRO; then 2) subsequent concurrent validation with advocated criteria in separate
5 subregions and condition-specific back and neck populations. This study's purpose was phase 1.

6 There are at least 43 back-specific PROs with 13 that can be used to evaluate responsiveness to
7 change [2]. Among these the Oswestry Disability Index and Roland Morris Disability Questionnaire
8 are the most commonly advocated [2, 12]. For the neck, at least 13 PROs have been developed [13] but
9 there is limited agreement on which ones should be advocated [6, 8]. Five PROs purport validity for the
10 whole-spine: the Functional Rating Index (FRI) [10], the Bournemouth Questionnaire [14], the
11 Extended Aberdeen Spine Pain Scales [9], the Pain Disability Questionnaire [12] and the Core
12 Outcome Measures Index [3]. However, further testing is required of these whole-spine tools because
13 none have demonstrated an adequate factor structure through either Rasch analysis or factorial analysis
14 [2], and the capacity to measure the whole-spine as a single kinetic chain [15]. Of these five PROs, the
15 FRI is advocated most strongly due to its preferred administrative practicality and level of independent
16 research on comparative clinimetric properties for both low back pain [16] and neck pain [8].
17 Consequently, the FRI is the optimal choice as a criterion measure ahead of the other four available
18 whole-spine PROs when developing a new whole-spine PRO and in preference to generic PROs such
19 as the SF-36 or EuroQol.

20 The development of each of these five whole-spine tools has attempted to address the need for a
21 single whole-spine tool. The initial three were questioned due to poor methodology in development,
22 practicality, factor analysis and validation [2]. For example, the Pain Disability Questionnaire is not
23 spine specific, nor does it account for acute situations as it is for 'chronic disabling musculoskeletal
24 disorders' [12]. The eleven-item Core Outcome Measures Index has separate neck and back versions,
25 and is designed to measure patients after operative procedures within secondary and tertiary settings.

1 Completion involves several scoring techniques with computerized input [3] and independent
2 validation as a whole-spine measure is still required. Both the Aberdeen [9] and Pain Disability
3 Questionnaire [12] have dual-factor structures that limit their validity as a single summated score and
4 consequently are less than optimal measure [15]. The remaining three PROs have even less research in
5 this aspect as they have not had their factor structure determined by the recommended maximum
6 likelihood extraction method [17]. Consequently, a whole-spine PRO is needed that has been
7 appropriately developed [18], represents a single kinetic chain, has a single factor structure and
8 appropriate clinimetric properties for both the back and neck.

9 A PRO must be clinically practical, effective, efficient and validated with a recognized criterion
10 standard [19]. The Spine Functional Index (SFI) (Figure 1) was developed to comply and satisfy these
11 requirements. The aim of this study was to describe the development of the SFI, determine the
12 psychometric, practical and factor structure characteristics in a general spinal population and compare
13 the SFI to a whole-spine criterion measure, the FRI [10].
14
15

16 **MATERIALS AND METHODS**

17 A prospective observational study was completed in two phases (Figure 2):

- 18 1. SFI development in three-stages;
 - 19 2. SFI validation in a symptomatic spine cohort.
- 20

21 **Phase 1 - Development of the Spine Functional Index**

22 The established three-stage development process used 1) item generation, 2) item reduction and 3) field
23 testing [15, 18] (Figure 2).
24
25

1 ***Stage 1 Item Generation***

2 Electronic data bases, PubMed, Cinahl, Embase and Pedro, from 1980-2010 were searched by the
3 primary author (CPG) with key words ‘outcomes’, ‘self-report’, ‘function’, ‘disability’, ‘impairment’,
4 ‘spine’, ‘neck’, ‘back’, ‘thoracic’, ‘cervical’ and ‘lumbar’. An additional search included clinicians and
5 researchers for unpublished questionnaires. This produced 129 PROs. A four-person peer-panel was
6 formed, consisting of an occupational therapist, physical therapist with spine-specific post-graduate
7 qualifications, general practitioner physician and occupational medicine physician with spine-specific
8 consultancy work. The panel used consensus opinion that required a three vote minimum [20, 21] to
9 review and shorten the list to 29 PRO tools with 850 items that were directly cited in each of the PROs
10 and relevant to the spine injuries. The list was reduced to 409 items by the panel through binning and
11 winnowing methodology which removed duplicate and non-applicable items [22, 23].

12 ***Stage 2 Item Reduction***

13 The 409 items were reduced in five separate stages (2a-e) by the panel. Stage 2a reduced the list to 159
14 items by pooling items with a common construct (e.g. ‘sitting’, ‘sit in a chair’, ‘sit on a stool’ etc. were
15 collapsed to ‘sitting’). Stage 2b classified [18] items using the World Health Organisation-International
16 Classification of Functioning (WHO-ICF) [24] codes from the ICF Browser [25]: b=body functions,
17 s=body structures, d=activities and participation, e=environmental factors [26]. Stage 2c reduced the
18 159 items to 89 by combining the ICF codes to common descriptive construct titles (e.g. ‘stairs’ and
19 ‘ladders’ became ‘code d4551-climbing’). Stage 2d reduced the list to 74 by grouping and deletion (e.g.
20 ‘dressing’ and ‘putting on pants’ were retained but ‘fastening clothing’ was deleted). Stage 2e further
21 combined items via consensus of importance and relevance to achieve the final 25 items, 15 general
22 and ten spine-specific. The stems for each question were formulated: ‘Due to my spine: I have
23 difficulty/problems...’; or ‘I stay/change/avoid/get others...’.

1 To ensure current best practice epidemiological standards were met, each question's final
2 wording was achieved through peer panel consensus then given to two focus groups for feedback and
3 relevance for face and content validity [18]: a spine symptoms patient focus group (n=10, three
4 cervical, three thoracic and four lumbar); the four person author group that included a physical therapist
5 and an orthopaedic surgeon both with extensive experience in the spine, a biomechanist, and a physical
6 therapist with extensive clinimetric research experience. The ten person patient focus group and the
7 four person author panel supplemented the initial item reduction process performed by the 'expert
8 panel'. The focus groups were provided with the final 25 items list and the list of the 49 items excluded
9 in stages 2d. The mixed methods semi-structured interview process [27] was used to determine if the
10 25 items should be changed and if any of the 49 excluded items should be reinstated or included within
11 the final item list. The "Isikawa" qualitative methodological process [28] was used to supplement the
12 consensus agreement from both the patient and author focus groups and the expert panel. The format
13 and three-item response option, 'Yes', 'No' and 'Half' [15], [29] were selected.

14 ***Stage 3 Field Testing***

15 A pilot investigation enrolled 52 participants who provided a total of 85 responses (n^R). This ensured
16 $n=52$ baseline responses and an additional 33 responses: 13 for reliability ($n=13$; $n^R=26$); and 20 for
17 responsiveness, where two participants completed an additional third set of responses ($n=18$; $n^R=38$)
18 (Figure 2). This allowed for a preliminary assessment of floor and ceiling effects, sampling method
19 practicality and sample size calculations.

20 ***Sample Size***

21 From the pilot study, minimum samples were determined for an 80% chance of detecting actual
22 difference with 15% attrition ($p<0.05$) [30]. This compared favorably to previous FRI investigations
23 [10, 31] for concurrent validity ($n=106$), reliability ($n=56$), responsiveness ($n=84$) and predictive ability
24 through construct validity ($n=168$).

1 **Phase 2 - Validation of the SFI in a cohort population**

2 *Design*

3 A single stage, prospective observational study analyzed concurrent SFI and FRI responses. Each
4 participant was classified by subregion (cervical, thoracic or lumbar) where the percentage noted
5 ensured proportional reliability and responsiveness representation [15, 18].

6 *Setting and Participants*

7 Participants who complained of spinal pain or symptoms (n=203, responses=506) were consecutively
8 recruited from ten Australian physical therapy clinics. Inclusion criteria were referral by a medical
9 practitioner for a musculoskeletal spine condition or symptoms. Exclusion criteria were pregnancy, red
10 flag signs, <18 years and English language difficulty. Symptoms and classifications of spinal diagnoses
11 represent the entire spinal region, as described in Table 1.

12 Participants completed both the SFI and FRI PROs, however the number of FRI responses
13 (n=173; responses=386) was reduced due to a misunderstandings with one participating clinic that
14 returned only the SFI responses. Participants receiving ongoing treatment were re-measured every two
15 weeks for six weeks, then every four weeks until discharge. Status was classified as: acute at 0-6
16 weeks; subacute at >6-12 weeks; and chronic at >12 weeks. Pooled responses assessed criterion
17 validity, distribution and missing responses. Participants also completed an 11-point global numeric
18 rating scale of perceived present overall status [32, 33], whereby subjects rate their status on a scale
19 from 0-10 (0=worst possible, 10=normal). The global numeric rating scale was used as an external
20 criterion measure of clinical change, by calculating the difference in global perceived present status
21 over time.

22

23 **Questionnaires**

24 The FRI [10] is a single page PRO that contains 10 items, each rated on a five-point Likert scale
25 incorporating visual and descriptive response options. Five items on the FRI are common to the

1 Oswestry Disability Index and the Neck Disability Index with three additional Oswestry Disability
2 Index items, one Neck Disability Index item and a new 'pain' item [2]. The raw score of the FRI is
3 multiplied by 2.5 to generate a 0-100% score on the FRI (100%=no disability). One missing response is
4 permitted.

5 The SFI is a single page 25-item PRO, with a three-point Likert scale response option for each
6 item. The scores from the 25 items are tallied for the sum, the sum is multiplied by four and then
7 subtracted from 100 to generate a 0-100% score (100%= no disability). Two missing responses are
8 permitted.

9 An 11-points global numerical rating scale (0=worst possible, 10=normal or fully recovered) was
10 used to reflect individual perceived global functional status and act as an external criterion.

11

12 **Data Analysis - Psychometric Characteristics**

13 *Distribution and normality* were assessed from baseline histogram inspection and one-sample
14 Kolmogorov-Smirnov tests (significance >0.05) [30]. *Internal consistency* used baseline Cronbach's
15 Alpha ($\alpha=0-1.00$) calculations with an optimal value recommended as 0.90-0.95 [18, 30]. *Test-retest*
16 *reliability* was assessed through the Intraclass Correlation Coefficients (ICC) Type 2,1, and expressed
17 with 95% CI using scores on the PRO from acute/subacute participants at baseline and again on day
18 three during a non-treatment period. Participants rating on the global numerical rating scale of
19 perceived overall status at baseline and on day three provided the reference criterion to determine
20 change. Only those participants who had a change of 0, +/-1 were entered into analysis for test-retest
21 reliability (n=70) [15].

22 *Responsiveness* was assessed using the effect size (ES) and standardized response mean (SRM)
23 statistics [18]. Participants were classified by subregion with repeated measures analyzed (n=191 for
24 the SFI; n=144 for the FRI) for: acute at two weeks, subacute at four weeks and chronic at six weeks.
25 This accounted for variations in healing and therapists interventions [15]. There were participants that

1 received no follow-up or early discharge (SFI, n=12; FRI, n=7). The global numeric rating scale score
2 of a change ≥ 2.0 was the cut-off used to define patient-rated clinical change. *Error score* was
3 determined with the minimal detectable change with 90% confidence bounds (MDC_{90}) using the
4 standard error of the measurement formula and the ICC coefficients. *Minimal clinically important*
5 *difference (MCID)* was calculated using an anchor based method, with the anchor of patient-rated
6 change determined from the global numeric rating of change. Patients were classified as improved or
7 deteriorated if they had a minimum change of ≥ 2.0 points on the global numeric rating scale between
8 baseline and follow-up [18, 33, 34]. Consequently, the MDC appears as a statistically and clinically
9 appropriate MCID [35].

10 *Validity* was assessed for *face* and *content* through focus groups, panel feedback and readability
11 scores [36]; and *criterion* through Pearson's r coefficient (n=386). *Construct validity* used discriminant
12 validity with the external criterion global numeric rating scale of perceived self-rated change of health
13 status ≥ 2.0 points [34]. Additionally, an *a-priori* paired t-test statistical difference was required
14 between baseline and repeated test groups mean scores to categorize subjects as improved or
15 deteriorated when calculating the MCID. *Factor analysis* used baseline SFI and FRI data with loading
16 suppression at 0.30 and varimax rotation for maximum likelihood extraction [30] which required
17 assumptions of normality. Factor extraction had three *a-priori* requirements: scree-plot 'point of
18 inflection'; eigenvalue > 1.0 ; and variance $\geq 10\%$ [30].

19

20 **Data Analysis - Practical Characteristics, Readability and Summary Performance**

21 Practicality considered nine areas [15, 36] with being five self-evident: 1) self-administered; applicable
22 across a variety of 2) conditions; 3) severity levels; 4) relevance to defined populations; and 5) single-
23 page length. The remaining four areas were determined through focus groups for 6) interviews for ease
24 of understanding and completion; 7) questionnaire completion time; 8) therapist scoring-time from
25 three separate scores averaged from each clinic; and 9) missing responses as percentages of total

1 responses (SFI, n=506; FRI, n=386). *Readability* used the Flesch–Kincaid grade scales (range 0-12,
2 optimum<7) and reading-ease (optimum>60%) calculated from word-processing software. Summary
3 performance used the ‘Measurement of Outcome Measures’ scale that evaluated 25 essential properties
4 [5]; and the ‘Bot’ scale that evaluated twelve items [36]. The ‘Bot’ cut-off classifications were adjusted
5 [15, 29] for ‘time to administer’ at three minutes and ‘readability and comprehension’ determined by
6 the Flesch-Kincaid scale cut-offs [15]. Significance was set at $p<0.05$.

9 RESULTS

10 Participant demographics are reported in Table 1.

12 Psychometric Properties

13 Characteristics of internal consistency, reliability, responsiveness and error score are summarized in
14 Table 2, and construct validity in Table 3.

15 *Distribution and normality* were demonstrated through the Kolmogorov-Smirnov test
16 (SFI=1.163, significance=0.87; FRI=1.18, significance=0.87) with identical SFI and FRI baseline score
17 ranges (0% -98%). The SFI histogram shape was preferred particularly in the upper 90-100% interval
18 that contained 15 (7.5%) responses compared to the FRI with a single response (2%). The ‘Half Mark’
19 option was used by 57% of participants at baseline and in 43% of all responses. The baseline scores by
20 subregion were comparable between the SFI and FRI apart from the multi-area group (Table 4).

21 *Criterion validity* was high (Pearson’s $r=0.85$) between the SFI and FRI scores. *Construct*
22 *validity* through *discriminant validity* was demonstrated for the *a-priori* criterion (Table 3). The
23 subregion mean scores were different for both PROs and between both PROs, though the cervical,
24 thoracic and multi-area groups were of a similar value. However, none were statistically significant
25 apart from the multi-area group ($p<0.001$).

1 *Factor analysis* was suitable as the correlation matrix Kaiser-Meyer-Olkin value was 0.912 and
2 Barlett Test of Sphericity significant ($p<0.001$). A unidimensional structure was indicated for both
3 PROs as the three *a-priori* criteria were met with second point scree-plot inflection and one eigenvalue
4 >1.0 where variance was $>10\%$ (SFI=33.4%, FRI=55.6%). The SFI had six more factors with
5 eigenvalues >1.0 , but with variance $<10\%$ that accounted for 30.5%. Both PROs had four factors with
6 eigenvalues between 0.5 and 1.0 with the remaining factors all below a 0.5 eigenvalue.

7

8 **Practical Characteristics**

9 *Completion time* was SFI=122±37 seconds, and FRI=84±23 seconds; scoring time was SFI=16±4
10 seconds, FRI=27±13 seconds. The FRI required a computational aid, and with one missing response
11 the scoring time increased to 53±19 seconds. Combined completion and scoring was SFI=138±41
12 seconds, FRI=137±39 seconds. *Missing responses* were $<1.5\%$ for the SFI, and 5.3% for the FRI.
13 *Readability* for the SFI was grade=7, reading ease=64%; and for the FRI grade =7, reading ease
14 =47.2%. *Summary performance* on the Measurement of Outcome Measures was SFI=96%, FRI=64%;
15 and on the 'Bot' for the SFI=12/12 or 100%, FRI=9/12 or 75%.

16

17

18 **DISCUSSION**

19 The SFI was developed using a structured methodology. It demonstrated acceptable psychometric
20 properties, a single factor structure, and strong practical characteristics in patients with spinal pain and
21 symptoms of the cervical, thoracic and lumbar spine. Compared to the FRI, by visual comparison of
22 the results, the SFI had equal or preferable psychometric properties of reliability, validity,
23 responsiveness and error. The summary performance scores of practical characteristics on the
24 Measurement of Outcomes Measure and 'Bot' scales showed high scores for the SFI. The SFI was
25 demonstrated to be capable of assessing functional status at a single point in time and change over time

1 to determine the effectiveness of treatment interventions. The practical characteristics of short scoring
2 times, low missed responses and reading ease will reduce both the patient and administrative burden.

3 The SFI has a three-point response format, which was used by participants 57% of the time.
4 This response format provided a simple scoring format within a stable equally-spaced scale [37]. This
5 also enabled sound individual interpretation for the psychological perspective of an item's 'presence',
6 'absence' or an 'intermediate position' [38] as opposed to a dichotomous response option.

7 Normalised SFI distribution and subregion scores in this cohort of patients presenting to
8 physical therapy demonstrated no floor or ceiling tendency. The FRI had more missing responses at
9 the higher levels of functional loss, indicating reduced measurement capacity. This measurement
10 capacity of the SFI may improve the ability to discriminate change throughout the scale range. Internal
11 consistency, test-retest reliability and responsiveness values for the SFI were acceptable and
12 comparable to the FRI. The SFI demonstrated lower error values (SEM and MDC₉₀) which may allow
13 for improved sensitivity for detecting change over time in the assessment of intervention effectiveness
14 that may otherwise not show a valid effect [39]. Moreover, this may subsequently reduce the 'number
15 needed to treat' [40].

16 Responsiveness of the SFI in a cohort of patients undergoing physical therapy treatment was
17 acceptable and comparable to the FRI, despite the higher diversity in baseline impairment [41, 42]. As
18 an observational study in a cohort of patients undergoing physical therapy care, other influences on
19 responsiveness may have been present. These include variation in interventions provided, follow-up
20 duration (as responsiveness is less over a shorter follow-up period) and baseline severity (as acute and
21 chronic patients change at different rates) [34]. These variables were attempted to be minimize by using
22 the concurrent testing methodology. Factor analysis demonstrated a single-factor structure and
23 consistent variance levels for both the SFI and FRI. This study is the first to report the FRI factor
24 structure.

25

1 **Limitations and Strengths of the Study**

2 One limitation of this study was the recruitment of patients presenting for care at physical therapy
3 outpatient clinics only. Consequently, results cannot be generalized to inpatient or community settings.
4 Patients referred to physical therapy most likely represent the mid-range of spine conditions. The
5 study's strengths were the prospective, multi-center investigation that included patients from each
6 spinal region with varied degrees of severity and duration that represented both the general and work
7 injured populations with a large variation in diagnoses (Table 1). Furthermore, 191 subjects were
8 available for the responsiveness sample, measuring these subjects on repeated occasions over time.
9 This facilitated their measurement throughout the severity spectrum, as indicated by the suitable levels
10 of distribution within the histogram, including the least affected level at the point of discharge.

11

12 **Implications for Further Research**

13 The high SFI and FRI criterion validity implied generalizability to populations where the FRI has been
14 validated or compared to other spine related PROs. This includes the Oswestry Disability Index,
15 Roland Morris Disability Questionnaire and Neck Disability Index. However, independent
16 investigations are required where spine subregion PROs are concurrently compared through repeated
17 measures on diagnoses such as whiplash, acute and chronic low back and neck pain. The SFI had
18 several factors that accounted for substantial variance. This suggests that shortening to perhaps ten-
19 items may be possible. This may improve practicality and reduce both respondent and clinician burden.
20 A confirmatory factor analysis should be considered.

21

22

23 **CONCLUSIONS**

24 The SFI is a practical PRO for measurement of spine related patient status and change over time.
25 Compared to the FRI, an advocated whole-spine PRO, the SFI had comparable and sometimes

1 improved psychometric and practical characteristics and overall performance. The findings of this
2 study indicated the SFI is a viable PRO for measuring whole-spine functional status in both the clinical
3 and research settings.

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7

1 **FIGURE LEGENDS**

2

3 Figure 1: Spine Functional Index

4 Figure 2: Flow chart of SFI development and validation

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6

7 Table 1: Participant demographics for SFI: Stage 1, pilot and stage 2, validation

8 Table 2: Methodological characteristics of SFI and FRI criterion

9 Table 3: Construct validity: significant differences between baseline and repeated PRO scores

10 Table 4: Mean scores by subregion for SFI and FRI

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Table 1: Participant demographics for SFI: Stage 1, pilot and Stage 2, validation

	Stage 1	Stage 2
Demographic data	Development and pilot validation	Validation (main study)
Participants (n)	52	SFI=203, FRI=173
Responses (n)	85	SFI=506, FRI=386
Age (years)	47.6 ± 17.5	41.0 ± 17.8
Gender (% female)	57.1%	48.0%
Injury: duration (weeks)	71.5 ± 103.0	4.6 ± 8.4
time range (weeks)	1 - 300 (1 outlier = 1575 weeks)	1 - 45 (1 outlier = 520 weeks)
Subregion and diagnosis* :		
Cervical	n=24 (46%)	n=96 (47%)
Whiplash		32%
Joint		29%
Disc		7%
Soft tissue		10%
Nerve root and neural		3%
Other (nonspecific, pain, etc.)		<u>19%</u>
		<u>100%</u>
Thoracic	n=4 (8%)	n=48 (24%)
Joint		29%
Disc		6%
Soft Tissue		23%
Fracture		2%
Costovertebral		2%
Other (nonspecific, pain, etc.)		<u>38%</u>
		<u>100%</u>
Lumbar	n=24 (46%)	n=101 (50%)
Joint		30.5%
Disc		29.5%
Soft Tissue		5%
Nerve root or neural		1%
Postoperative		3%
Sacropelvic		3%
Osteoarthritis		4%
Other (nonspecific, pain, etc.)		<u>24%</u>
		<u>100%</u>
Multiarea	Nil	n=46 (23%)

* Subregion % values include multiarea individuals within each of their symptomatic regions making total >100%.

1 **Table 2: Methodological characteristics of SFI and FRI criterion**

	Reliability (ICC)	Internal consistency	Error score		Responsiveness			Missing responses
Stage 2	Rxx	Alpha	SEM	MDC₉₀	SD₁₀₀	ES	SRM	Percentage
SFI	0.972	0.911	2.76	6.44	24.80	1.25	1.81	1.5%
FRI	0.948	0.908	4.14	9.66	22.67	1.23	1.68	5.3%

2
3 Rxx: Test-retest reliability coefficient; Alpha: Cronbach's Alpha. SEM: Standard Error of the Measurement; MDC₉₀: Minimal
4 Detectable Change (90% CI); SD₁₀₀: Standard deviation at baseline (100% scale); ES: Effect Size; SRM: Standard Response
5 Mean.
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16 **Table 3: Construct validity comparing means for baseline, repeated scores and difference for the**
17 **SFI and FRI (n=113)**

Tool	Baseline mean	Repeat test mean *	Difference in means
SFI	43.6±24.8	17.3±19.1	31.0±17.1
FRI	43.0±22.7	17.1±18.6	28.0±16.7

18
19 * Repeated measures were made after a period of known natural healing: acute after two weeks; subacute after four weeks; and
20 chronic > 12 weeks, all with a p value <0.0001 for the *t*-statistic measures.
21
22

1 **Table 4: Mean baseline scores by subregion for SFI and FRI**

	Stage 2	
Subregion	SFI	FRI
Cervical	41.1±25.9 n=96 (47%), with Cx only n=78 (38%), Cx, Tx n=16, Cx, Lx n=2	39.2±21.6 n=67 (39%), with Cx only n=53 (31%), Cx, Tx n=13, Cx, Lx n=2
Thoracic	41.2±25.8 n=48 (24%), with Tx only n=8 (4%), Cx, Tx n=16, Tx, Lx n=24	38.2±23.4 n=38 (22%), with Tx only n=4 (2%), Cx, Tx n=13, Tx, Lx n=21
Lumbar	46.8±26.1 n=101 (50%) with Lx only n= 75 (37%), Cx, Lx n=2, Tx, Lx n=24	45.0±21.4 n=90 (52%) with Lx only n=67 (39%), Cx, Lx n=2, Tx, Lx n=21
Multi area	41.0±26.8 n=42 (21%) with Cx, Tx n=16, Cx, Lx n=2, Tx, Lx n=24	34.0±20.3 n=36 (21%) with Cx, Tx n=13, Cx, Lx n=2, Tx, Lx n=21
All – average for all data	44.8±26.1 n=203	41.4±17.2 n=173

2
3 Multi area included participants who reported symptoms in more than one area. Cervical = Cx, Thoracic =
4 Tx, Lumbar = Lx.

5

<u>SPINE FUNCTIONAL INDEX (SFI)</u>		DATE: _____
NAME: _____	INJURY: _____	<input type="checkbox"/> Neck <input type="checkbox"/> Mid Back <input type="checkbox"/> Lower Back

PLEASE COMPLETE: Your spine may make it difficult to do some things you normally do. This list contains sentences people use to describe themselves with such problems. Think of yourself over the last few days. **If an item describes you, mark the box. If not, leave the box blank. If an item partly describes you, Use a Half (½) Mark.**

DUE TO MY SPINE:

___ 1. I stay at home most of the time.
___ 2. I change position frequently for comfort.
___ 3. I avoid heavy jobs (e.g. cleaning, lifting more than 5kg or 10lbs, gardening, etc).
___ 4. I rest more often.
___ 5. I get others to do things for me.
___ 6. I have the pain / problem almost all the time.
___ 7. I have difficulty lifting and carrying (e.g. bags, shopping up to 5kg or 10lbs).
___ 8. My appetite is now different.

___ 9. My walking or normal recreation or sporting activity is affected.
___ 10. I have difficulty with normal home or family duties and chores.
___ 11. I sleep less well.
___ 12. I need assistance with personal care (e.g. washing and hygiene).
___ 13. My regular daily activities (work, social contacts) are affected.
___ 14. I am more irritable and / or bad tempered.
___ 15. I feel weaker and / or stiffer.
___ 16. My transport independence is affected (driving, public transport).

___ 17. I require assistance or am slower with dressing.
___ 18. I have difficulty moving in bed.
___ 19. I have difficulty concentrating and / or reading.
___ 20. My sitting is affected.
___ 21. I have difficulty getting in and out of chairs.
___ 22. I only stand for short periods of time.
___ 23. I have difficulty squatting and / or kneeling down.
___ 24. I have trouble reaching down (e.g. pick-up things, put on socks).
___ 25. I go up stairs slower or use a rail.

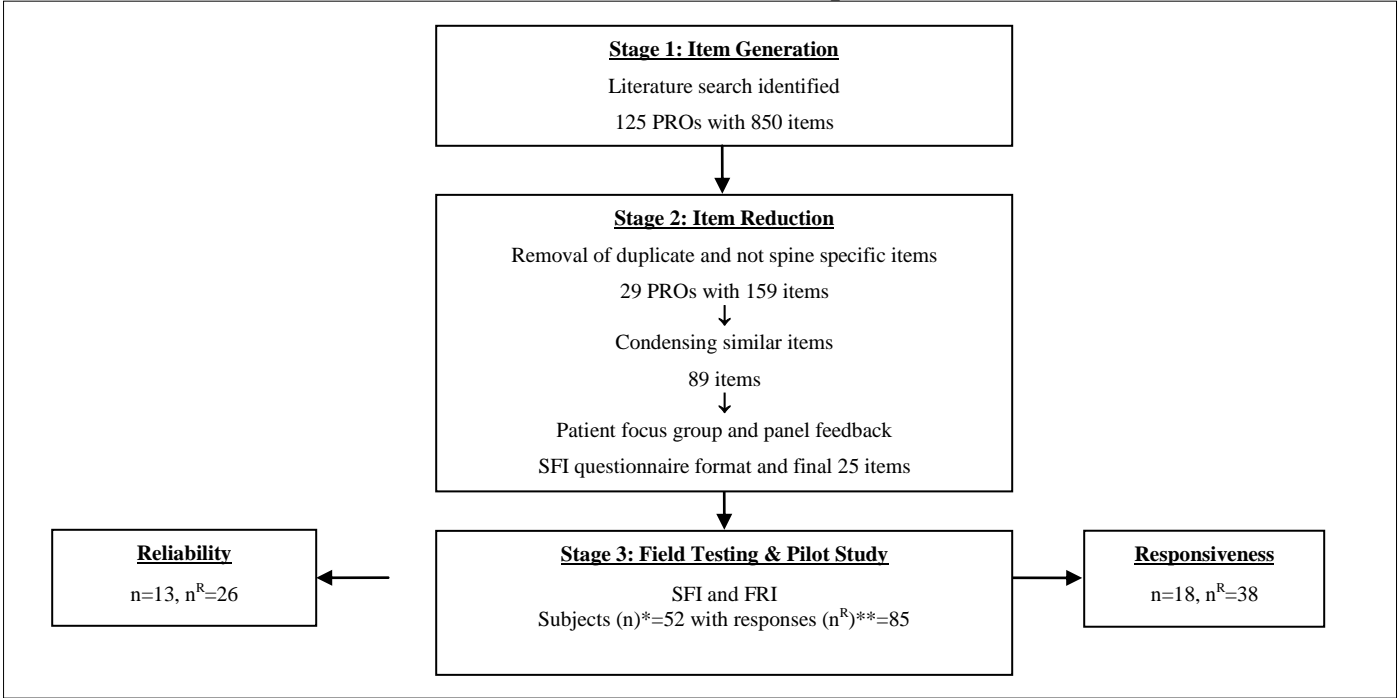
SFI SCORE: To score the upper part - add the marked boxes:

<input style="width: 80%;" type="text"/> TOTAL (SFI points)	100 Scale: 100 – (TOTAL x 4) =	<input style="width: 80%;" type="text"/> %
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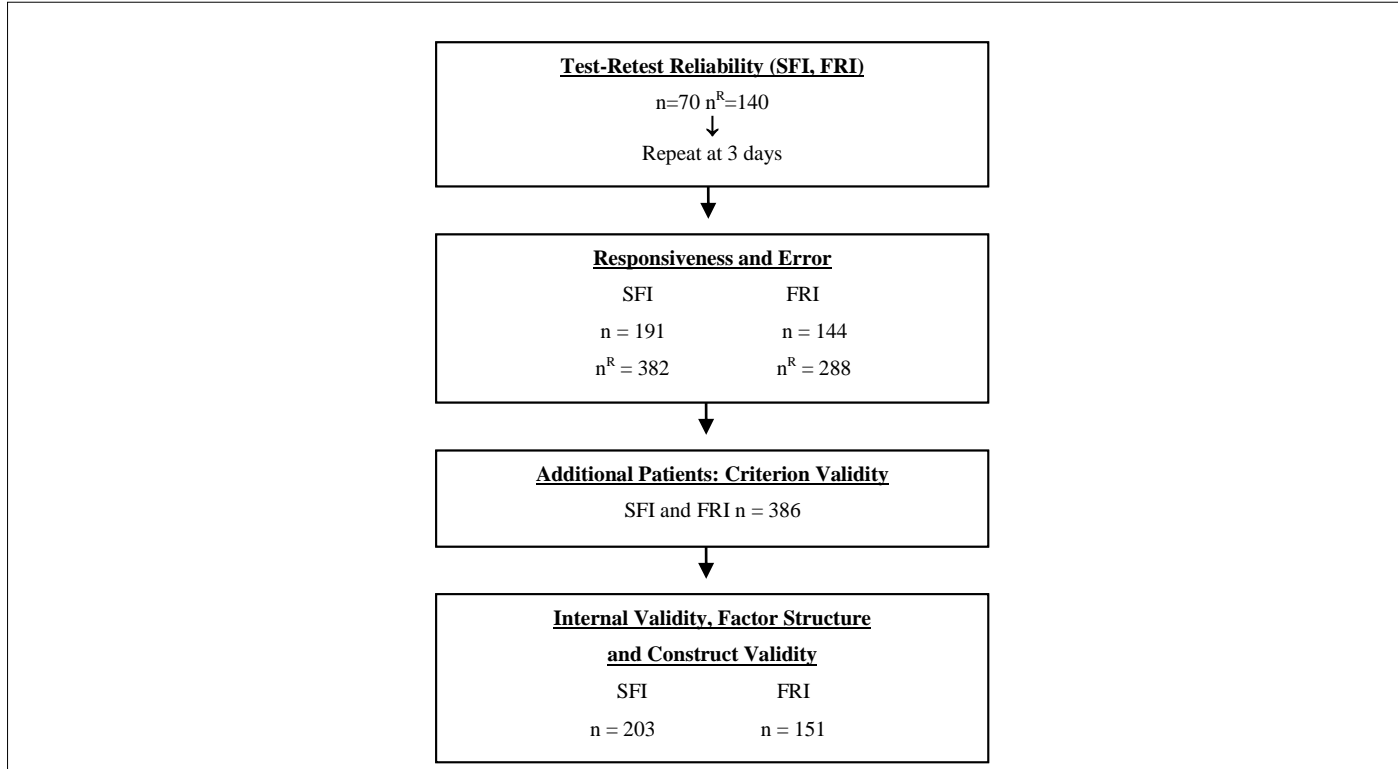
MDC (90% CI): Neck = 6.9% or 1.7 SFI points; Mid and Lower Back = 5.9% or 1.5 SFI points;

All spine = 6.5 % or 1.6 SFI points Change less than this may be due to error.

1 **Figure 2: Flow chart of SFI Development and Validation**
 2 **Phase 1: Questionnaire Development**



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5 **Phase 2: Validation Study (n=203 n^R=506)**



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8 * = n is number of subjects
9 ** = n^R is the number of responses
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