Estimation of Prognosis in Non Specific Low Back Pain from Biopsychosocial Perspectives

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1. Introduction

The significance of low back pain and associated disability across many western nations continues to be an important issue. Pain and disability associated with musculoskeletal conditions represents a significant health and economic burden in Australia, with total direct and indirect costs of musculoskeletal disorders including arthritis being in excess of 15 billion dollars per annum (AAMPG. 2003).

In the USA the data from the 1998 Medical Expenditure Panel Survey, reported that health care expenditure incurred by individuals with back pain reached in excess of 90 billion dollars. Individuals with back pain also incurred 60% more health care costs than those without (Luo, Pietrobon et al. 2003).

There is also a growing literature that supports the contention that low back pain is a complex, multidimensional health issue and should be reviewed within the context of the individual (Young, Waisiak, et al 2011).

Evidence-based clinical practice guidelines, such as those produced in Australia in 2003 emphasise the importance of the biopsychosocial model in the understanding of pain in general and musculoskeletal pain in particular (AAMPG. 2003). The biopsychosocial model focuses on illness, communicating or behaving in a manner that suggests the individual is not well, rather than on disease, the person’s experience of illness being influenced by physical, psychological and social factors (Engel. 1980). Engel’s model highlighted that in order to respond adequately to an individuals suffering, and give them a sense of being understood, the clinician needed to be able to respond simultaneously to the biological, psychological and social dimensions of illness (Borrell-Carrio, Suchman et al 2004)

The model incorporates the biomedical understanding of nociception as part of the wider holistic biopsychosocial view, rather than treating psychosocial factors as an ‘overlay’ to the biomedical model.

Philosophically the model is a way of understanding how suffering, disease and illness may be affected by multiple levels of organisation, from the molecular to societal. Practically, the model provides a way of understanding the patient’s subjective experience as an essential contributor to accurate clinical diagnosis and assessment (Borrell-Carrio et al 2004).
There has been increasing acceptance that psychosocial factors play a significant role in the transition from an acute episode, or episodes, of low back pain to a chronic disorder (Pincus, Burton, et al. 2002). There is also some evidence that these factors may play an aetiological role (Pincus et al. 2002) (Linton, 2000) (Trouchon, and Fillion, L.2000).

The acceptance of the wider biopsychosocial model has been further extended by the World Health Organisation’s (WHO) International Classification of Functioning, Disability and Health (ICF) (WHO, 2000).

Many individual psychological factors have been reported over the years as potential obstacles to recovery, with much of the early work centred around fear. Initially the work focussed around the fear avoidance model of exaggerated pain perception in chronic low back developed by Lethem and co-workers (Lethem, Slade, Troup, Bentley. 1983). The Fear-Avoidance Belief Questionnaire developed by Waddell and co-workers published in 1993 measured beliefs about physical activity and work (Waddell, Somerville, Henderson, Newton, Main 1993). Workers with low back pain were shown to believe that physical activity including work was feared to increase both spinal pain and damage. These fear avoidant beliefs helped to explain self reported disability in normal daily activities including work (Waddell et al 1993).

Fear of pain together with fear of hurt or harm were further postulated as a fundamental mechanism of disability associated with low back pain were explored by Vlaeyen and Linton. The natural human reaction to pain is an automatic one to try and avoid what is believed to be the cause of the pain, ongoing fear then potentially drives further avoidant behavior(Vlaeyen and Linton 2000).

Contrary to the weight of evidence however, surveys of primary contact practitioners continue to indicate that general practitioners for instance, may only be partially managing low back pain from this evidence-based perspective. This may be due in part to the difficulties reported in changing physician behaviour (Buchbinder, Staples et al. 2009) and then further highlighted by the difficulty reported in integrating the biopsychosocial model into clinical disciplines, such as physiotherapy (Harland and Lavallee, 2003). The potential difficulties in fully incorporating a biopsychosocial approach to patients, appear to transcend professional training and boundaries, as Harding and co-workers reported in 2010. Whilst pain clinic practitioners in the UK embrace behavioural based management as part of a biopsychosocial pain management model, little consideration was given to social factors (Harding, Campbell et al 2010).

The estimation of prognosis (identifying those at risk of a poor outcome such as ongoing pain or disability, or failing to return to work) of an episode of low back pain is particularly important to clinicians, patients, employers and third party payers alike. However despite the weight of evidence there remains a tendency to rely on instruments that quantify prognostic factors from a biomedical perspective alone, rather than the contemporary biopsychosocial model, whilst focusing on return to work as a single outcome. A predictive instrument that may help clinicians fully incorporate the biopsychosocial model into clinical practice clearly has some utility, and whilst it has been suggested that such an instrument needs to capture all the potential biopsychosocial risk factors that may adversely affect functional outcomes (Hilficker, Bachmann, et al. 2007), an initial attempt to produce such a draft questionnaire resulted in a long and unwieldy instrument.
The study reported in this chapter tested the hypothesis that it would be feasible to develop a valid and reliable instrument for estimating the prognosis of non-specific low back pain from within the biopsychosocial domain, suitable for routine clinical use.

Ethical approval for each step of the instrument’s development was obtained from the ethics committee, University of Queensland, Australia.

The components of the resultant instrument (termed the Biopsychosocial Index of Prognosis (BPIP)) were developed from the literature review and perspective of a logical course of clinical enquiry into the current and past episodes of low back pain. The nature and character of the pain together with the severity, spread and duration of the symptoms were also included in the draft questionnaire. Past and present medical and psychological histories were considered together with demographic details including occupation. The items generated were combined into a content map derived from the International Classification of Function (ICF). Initial item generation included items from previously validated instruments identified from the literature where possible. The draft index initially contained 145 items. The large pool of items were initially included in the prototype questionnaire, which was progressively and systematically subjected to item reduction. The necessity for this process of item reduction was established a priori, and driven by the lack of feasibility of carrying a large number of potentially redundant items through to the validation stage of the instruments development (Bellamy, Campbell et al. 2002). The development of the BPIP potentially filled a gap in measurement, estimating prognosis from a variety of biopsychosocial domains and driven by clinical, rather than theoretical perspectives.

This chapter describes the development of the BPIP including the steps taken in item reduction and reports on testing the validity of the final instrument, and its reliability against the ‘gold standard’ measure identified as the Roland and Morris disability questionnaire (RDQ) (Roland and Morris, 1983). The RDQ is one of the most frequently reported instruments used to assess the disability associated with low back pain. The RDQ was also the criterion measure selected for the previous work of Burton and co-workers on psychosocial predictors (Burton, Tillotson, Main and Hollis 1995). A variety of other instruments were considered from the literature including the Orebro Musculoskeletal Pain Questionnaire (Linton and Hallden, 1998) as the potential gold standard. However the focus was on the wider community with low back pain, not just workers with acute low back pain, so the RDQ remained the instrument of choice.

2. Methods

2.1 Development of the BPIP

The initial assessments of the prototype index were undertaken in three distinct phases. In phase one, piloting was undertaken with a small group without low back pain, consisting of five health care professionals from a variety of disciplines and from different English speaking countries including England, Australia and New Zealand and chosen for their academic and general life experience. The primary focus of the first pilot trial was to obtain feedback on the pool of items selected for evidence of any ambiguity or repetition, ease of administration and the overall comprehensiveness of the questionnaire.
In phase two, a further pilot study was undertaken with a group of low back pain patients recruited from a variety of primary health care practices on the Sunshine Coast in Australia. Patients were provided with a pack containing the prototype questionnaire, and instruction sheet and two copies of the informed consent form together with two stamped addressed envelopes. Each stage of data collection was subjected to ethical clearance through The University of Queensland’s ethics committee.

Phase three was a 24 hour test-retest of the reliability of the amended instrument was completed. BPIP packs containing 2 identical questionnaires were distributed with the instruction to complete the first questionnaire and then the second one, ideally at the same time of day, 24 hours later without reference to the answers provided the day before. The intention was to establish reliability in a sample of 25% of the total of respondents required for the trial of the main instrument.

No specific exclusion criteria were stated other than children or adolescents, pregnancy or three months post partum. The instrument was designed for use in adults over the age of 18 years with English as a first language.

The overall development process of the BPIP is summarised in Figure 1

BPIP item generation
↓
Draft Index
↓
(Prototype BPIP)
BPIP item reduction
↓
Reliability (test re-test)
↓
BPIP item reduction
↓
Reliability & validity elucidation (Australian) sample
↓
Reliability & validity confirmation (New Zealand) sample

Fig. 1.

A total of one hundred and forty five items were generated from the literature review and clinical perspectives for the draft questionnaire. Redundant and non response items were eliminated following consultation with a panel of five independent healthcare practitioners as the first stage of item reduction. The questionnaire was subsequently reduced to a ninety nine item prototype.
A 24 hour test-retest of the stability of the ninety-nine item questionnaire was undertaken by twenty five respondents who were asked to complete the first questionnaire (A) and then the second (B) at the same time twenty four hours later, without referring back to their previous answers.

2.2 Data collection

Questionnaire packs containing the prototype BPIP, RDQ, an informed consent form and instruction sheet were assembled. The data captured at baseline and twelve weeks, outcomes at twelve weeks having been previously reported as being predictive of outcome at one year and beyond (Burton, McClune, et al. 2004).

Recruitment of respondents originally intended to capture a homogeneous cohort drawn from general practice proved impossible despite repeated requests via the faculty of Health Sciences, University of Queensland. Recruitment difficulties therefore resulted in two distinct cohorts, a larger Australian cohort (n = 91) recruited from an osteopathic practice and a diagnostic imaging centre. The smaller New Zealand cohort (n = 27) was recruited from an osteopathic practice and an osteopathic teaching clinic.

The larger Australian group was utilised as an elucidation (or clarification) cohort, and the smaller New Zealand cohort was treated as a small prospective validation of the final version of the BPIP.

2.3 Statistical analysis

The continued process of item reduction, including correlation analysis of baseline BPIP data with the 12 week score on the RDQ resulted in an ordinal scale of initially 24 items, which was further reduced to 12 items for correlation with the RDQ. The BPIP was tested for internal consistency in both the Australian and New Zealand cohorts.

The baseline data from the Australian cohort were assessed for concurrent, predictive and construct validity against the RDQ.

Reliability of the index was evaluated by calculation of Cronbach’s Alpha of the baseline BPIP score with the twelve week score on the RDQ in the both cohorts. The mean scores fell within the range of possible scores indicating that the scores were entered into the data set correctly.

The BPIP baseline data from the New Zealand cohort was treated as a small prospective validation of the BPIP scale, with the baseline data from the BPIP correlated with the change score from the baseline to twelve weeks of the RDQ.

The questions retained their original numbers from the draft questionnaire until item reduction was completed, and were then renumbered 1-12 in the final BPIP scale.

Analysis was undertaken using the Statistical Package for Social Sciences (SPSS).

3. Results

The results are reported in the same order as they contributed to the item reduction process, and final assessment of the 12 item BPIP.
3.1 Test-re-test of the ninety nine item prototype BPIP

A positive correlation between A and B questionnaires was demonstrated in a one tailed test, df 11-2 = 9. The Pearson product moment correlation coefficient was calculated in Excel: Pearson $r = 0.98$, $P < 0.0005$.

The calculation of $R^2 \ 0.98 \times 0.98 = 0.96$, demonstrated that 96% of the shared variance between the two test samples was accounted for by the result.

One hundred and eighteen respondents completed the twelve week data collection period. This was comprised of 91 in the Australian cohort and 27 in New Zealand.

The demographics including gender and age, and the percentage make up of the two cohorts are contained in Tables 1 and 2. Table 3 reports on the duration of low back pain measured at baseline in both cohorts.

<table>
<thead>
<tr>
<th>Gender</th>
<th>Australian</th>
<th>New Zealand</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>Aus: 26 (29%)</td>
<td>NZ: 6 (22%)</td>
</tr>
<tr>
<td>Female</td>
<td>Aus: 65 (71%)</td>
<td>NZ: 21 (78%)</td>
</tr>
</tbody>
</table>

Table 1.

<table>
<thead>
<tr>
<th>Age Range</th>
<th>Australian</th>
<th>New Zealand</th>
</tr>
</thead>
<tbody>
<tr>
<td>20-29</td>
<td>6 (7%)</td>
<td>3 (11%)</td>
</tr>
<tr>
<td>30-39</td>
<td>13 (14%)</td>
<td>6 (22%)</td>
</tr>
<tr>
<td>40-49</td>
<td>16 (18%)</td>
<td>5 (19%)</td>
</tr>
<tr>
<td>50-59</td>
<td>30 (33%)</td>
<td>7 (26%)</td>
</tr>
<tr>
<td>60 +</td>
<td>26 (29%)</td>
<td>6 (22%)</td>
</tr>
</tbody>
</table>

Table 2.

<table>
<thead>
<tr>
<th>Australian Cohort Duration of LBP at Baseline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weeks duration</td>
</tr>
<tr>
<td>Totals</td>
</tr>
<tr>
<td>Percentage</td>
</tr>
</tbody>
</table>

Table 3.

<table>
<thead>
<tr>
<th>New Zealand Cohort Duration of LBP at Baseline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weeks duration</td>
</tr>
<tr>
<td>Totals</td>
</tr>
<tr>
<td>Percentage</td>
</tr>
</tbody>
</table>

Table 4.
### 3.2 Item reduction and reliability

Correlation analysis (used to describe the strength and direction of the linear relationship between the two variables) of the baseline BPIP Australian data with the twelve week RDQ Australian data resulted in reducing the prototype scale to questions which correlated at or above 0.3. Items which correlated below 0.3 were considered too weak for inclusion, and a total of twenty four questions were retained for further analysis.

Reliability coefficients for internal consistency of the twenty four item BPIP scale were as follows:

- The Australian cohort, Cronbach’s Alpha = 0.8736.
- The New Zealand cohort, Cronbach’s Alpha = 0.8628.

A further review of the correlation analysis of the baseline BPIP Australian data with twelve week RDQ Australian data for items that correlated at or above 0.4 resulted in further item reduction of the BPIP to twelve questions. Following this further item reduction, reliability coefficients for internal consistency were calculated and are included in Tables 5 and 6.

#### Table 5. Inter-item Statistics Baseline BPIP/12 week RDQ; Australian cohort

<table>
<thead>
<tr>
<th>Item</th>
<th>Scale Mean if Item Deleted</th>
<th>Scale Variance if Item Deleted</th>
<th>Corrected Item-Total Correlation</th>
<th>Cronbach’s Alpha if Item Deleted</th>
</tr>
</thead>
<tbody>
<tr>
<td>severity</td>
<td>29.97</td>
<td>89.921</td>
<td>.557</td>
<td>.867</td>
</tr>
<tr>
<td>Workintrude</td>
<td>30.66</td>
<td>85.438</td>
<td>.741</td>
<td>.857</td>
</tr>
<tr>
<td>Socintrude</td>
<td>30.58</td>
<td>81.670</td>
<td>.728</td>
<td>.854</td>
</tr>
<tr>
<td>q19</td>
<td>30.50</td>
<td>82.527</td>
<td>.634</td>
<td>.860</td>
</tr>
<tr>
<td>q24</td>
<td>29.95</td>
<td>83.289</td>
<td>.606</td>
<td>.862</td>
</tr>
<tr>
<td>q25</td>
<td>30.36</td>
<td>80.859</td>
<td>.687</td>
<td>.857</td>
</tr>
<tr>
<td>q26</td>
<td>29.86</td>
<td>91.889</td>
<td>.282</td>
<td>.882</td>
</tr>
<tr>
<td>Worklimit</td>
<td>30.25</td>
<td>87.524</td>
<td>.553</td>
<td>.866</td>
</tr>
<tr>
<td>soclimit</td>
<td>30.21</td>
<td>88.238</td>
<td>.526</td>
<td>.867</td>
</tr>
<tr>
<td>q69</td>
<td>30.61</td>
<td>88.213</td>
<td>.517</td>
<td>.868</td>
</tr>
<tr>
<td>q73</td>
<td>30.32</td>
<td>84.827</td>
<td>.546</td>
<td>.866</td>
</tr>
<tr>
<td>q74</td>
<td>30.84</td>
<td>85.945</td>
<td>.489</td>
<td>.870</td>
</tr>
</tbody>
</table>

The Australian cohort, Cronbach’s Alpha = 0.875.

The New Zealand cohort, Cronbach’s Alpha = 0.776.

Once it was established that the reduced scale reliably measured the underlying construct, regression analysis was undertaken to establish levels of statistical significance.

Regression analysis of the Australian cohort based on the twelve item scale demonstrated that 61.7% of the variance in the RDQ score at twelve weeks was accounted for by the BPIP scale, with \( p = 0.0005 \).
Regression analysis of the change score of the RDQ with the twelve item BPIP in the New Zealand cohort demonstrated that 78.2% of the variance in RDQ scores was accounted for by the BPIP scale, with \( p = 0.006 \).

### 3.3 Multiple regression analysis (Australian cohort)

Standard multiple regression analysis was undertaken as it allows for a more sophisticated examination of the relationship between the dependent and independent variables, in this case how well the BPIP was able to predict outcome of the RDQ.

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>Std. Deviation</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDQ change 13</td>
<td>6.4074</td>
<td>7.05069</td>
<td>27</td>
</tr>
<tr>
<td>severity</td>
<td>3.59</td>
<td>.970</td>
<td>27</td>
</tr>
<tr>
<td>WorkIntrude</td>
<td>3.15</td>
<td>1.199</td>
<td>27</td>
</tr>
<tr>
<td>SocIntrude</td>
<td>2.70</td>
<td>1.235</td>
<td>27</td>
</tr>
<tr>
<td>q19</td>
<td>2.52</td>
<td>1.553</td>
<td>27</td>
</tr>
<tr>
<td>q24</td>
<td>3.30</td>
<td>1.235</td>
<td>27</td>
</tr>
<tr>
<td>q25</td>
<td>2.52</td>
<td>1.503</td>
<td>27</td>
</tr>
<tr>
<td>q26</td>
<td>3.44</td>
<td>1.502</td>
<td>27</td>
</tr>
<tr>
<td>WorkLimit</td>
<td>2.81</td>
<td>.879</td>
<td>27</td>
</tr>
<tr>
<td>SOCLIMIT</td>
<td>2.78</td>
<td>.801</td>
<td>27</td>
</tr>
<tr>
<td>Q69</td>
<td>3.11</td>
<td>1.311</td>
<td>27</td>
</tr>
<tr>
<td>Q73</td>
<td>1.96</td>
<td>1.192</td>
<td>27</td>
</tr>
<tr>
<td>Q74</td>
<td>1.70</td>
<td>1.137</td>
<td>27</td>
</tr>
</tbody>
</table>

Table 7. Multiple Regression; Australian cohort: Baseline 12 Item BPIP/12 wk RDQ.
Multiple regression analysis was then performed on the Australian cohort to establish the predictive value of the shortened BPIP scale score at baseline for the 12 week RDQ.

The results of the analysis of the Australian cohort are presented in Table 7, Table 8 contains the ANOVA result which demonstrates levels of significance.

<table>
<thead>
<tr>
<th>Model</th>
<th>Sum of Squares</th>
<th>df</th>
<th>Mean Square</th>
<th>F</th>
<th>Sig.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Regression</td>
<td>1505.555</td>
<td>12</td>
<td>125.463</td>
<td>10.186</td>
<td>.000 a</td>
</tr>
<tr>
<td>Residual</td>
<td>936.063</td>
<td>76</td>
<td>12.317</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>2441.618</td>
<td>88</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a. Predictors: (Constant), q74, q26, q69, severity, soclimit, q73, q24, q19, SocIntrude, q25, WorkIntrude, WorkLimit

b. Dependent Variable: RDQ12wk aus

Table 8.

The variance in the RDQ score (61.7%) at 12 weeks was accounted for by the 12 item BPIP. The null hypothesis that multiple R in this population equals zero was also confirmed by significance in the ANOVA, \( p = 0.0005 \). Multiple R is a measure of how strongly or weakly the criterion variables (BPIP variables) are related to the dependant variable (RDQ)(Stalin. 2003).

3.4 Regression analysis, (New Zealand cohort)

Regression analysis of the 12 item BPIP with the RDQ change scores baseline to 12 weeks was then performed on the New Zealand cohort, as part of the prospective validation of the

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>Std. Deviation</th>
<th>N</th>
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<tbody>
<tr>
<td>RDQchange 13</td>
<td>6.4074</td>
<td>7.05069</td>
<td>27</td>
</tr>
<tr>
<td>severity</td>
<td>3.59</td>
<td>.970</td>
<td>27</td>
</tr>
<tr>
<td>WorkIntrude</td>
<td>3.15</td>
<td>1.199</td>
<td>27</td>
</tr>
<tr>
<td>SocIntrude</td>
<td>2.70</td>
<td>1.235</td>
<td>27</td>
</tr>
<tr>
<td>q19</td>
<td>2.52</td>
<td>1.553</td>
<td>27</td>
</tr>
<tr>
<td>q24</td>
<td>3.30</td>
<td>1.235</td>
<td>27</td>
</tr>
<tr>
<td>q25</td>
<td>2.52</td>
<td>1.503</td>
<td>27</td>
</tr>
<tr>
<td>q26</td>
<td>3.44</td>
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<td>27</td>
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<tr>
<td>WorkLimit</td>
<td>2.81</td>
<td>.879</td>
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<tr>
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<td>2.78</td>
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<td>1.311</td>
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<td>Q74</td>
<td>1.70</td>
<td>1.137</td>
<td>27</td>
</tr>
</tbody>
</table>

Table 9. Multiple Regression of 12 Item BPIP scores with the change score Roland Morris Disability Index, baseline to 12 weeks; New Zealand Cohort
BPIP following the statistical analysis of the Australian data. The change score from baseline to 12 weeks was utilised as the raw baseline to 12 week score in the New Zealand cohort and did not reach statistical significance. The results are summarised in the same order as the Australian results in Tables 9 and 10.

The variance in the change score from baseline to 12 weeks of the Roland Morris Disability Index (78.2%) was accounted for by the 12 item BPIP.

The null hypothesis that multiple R in this New Zealand population equals zero was also confirmed by significance in the ANOVA $p = 0.006$

<table>
<thead>
<tr>
<th>Model</th>
<th>Sum of Squares</th>
<th>df</th>
<th>Mean Square</th>
<th>F</th>
<th>Sig.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Regression</td>
<td>1011.152</td>
<td>12</td>
<td>84.263</td>
<td>4.193</td>
</tr>
<tr>
<td></td>
<td>Residual</td>
<td>281.367</td>
<td>14</td>
<td>20.098</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>1292.519</td>
<td>26</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a. Predictors: (Constant), Q74, Q69, WorkLimit, q24, SocIntrude, q26, severity, SOCLIMIT, Q73, q19, WorkIntrude, q25
b. Dependent Variable: RDQchange 13

Table 10.

4. Discussion

The development of the BPIP from the original hypothesis, that it was feasible to develop a biopsychosocial prognostic instrument for eventual use in clinical practice, produced a promising result. The utility of the scale may also enable clinicians to consider the possibility that a slow or poor prognosis may be the result of a number of psychosocial obstacles to recovery being present, rather than assuming that underlying pathophysiological constraints to recovery are the culprits.

The study built on the seminal study of Burton and co-workers (Burton, Tillotson et al. 1993) which utilised a small number of items from previously validated measures to correlate with the RDQ, to positively demonstrate that psychosocial factors were indeed predictive of outcome. Other biopsychosocial instruments tend to focus on single psychosocial predictors (Hurley, Dusoir. et al 2000) or are designed as screening instruments for identifying those specifically at risk of developing long term incapacity from work such as the Orebro questionnaire (Linton and Hallden. 1997). These instruments also differ from the BPIP in that they primarily focus on screening workers with acute low back pain, rather than the wider patient population often consulted in primary care.

The BPIP was developed from clinical perspectives rather than isolating a single construct such as fear avoidance as a predictive factor of disability. The BPIP’s focus is consistent with the current idea of low back pain as a fluctuating and disabling condition (Young, Wasiak,
Phillips and Gross. 2011) that needs to be considered within the context of the individuals experience. Recent studies suggest that perceptions of personal control, pain self efficacy, the acute/chronic timeline and illness identity are distinctive psychological obstacles to recovery in primary care patients with low back pain, with depression, catastrophising and fear avoidance being reported as less significant (Foster, Thomas, Bishop, Dunn and Main. 2011). These findings being broadly consistent with the content of the BPIP.

The approach to data collection was to keep exclusion criteria to a minimum and not discriminate against respondents from the arbitrary time dependent classification of 12 weeks between acute and chronic low back pain patients.

4.1 Summary of key findings

A logical sequence of item reduction resulted in the 24 item BPIP scale, which was assessed both for its validity and reliability in separate cohorts, with a similar and strong result in both cases, with alpha value above 0.85.

The Pearson product moment correlation of the total score of the 24 item BPIP scale at baseline with the 12 week RDQ score demonstrated that the New Zealand scores did not reach statistical levels of significance. The small numbers in the cohort (n =27) or the more acute nature of the respondents, (21.9% reported pain of more than twelve weeks duration) may have potentially impacted on the result. However as the two cohorts tested differently, the decision to deal with the two groups as separate cohorts appears to have been justified.

The BPIP was shortened to 12 items following further correlation analysis of the Australian cohort’s BPIP scores and questions demonstrating the stronger correlation of 0.4 were retained, reducing the scale to a single sheet of A4 paper, potentially improving the scales clinical utility.

To check that the reduced scale could still be considered reliable, reliability analysis using Cronbach’s alpha was again undertaken in the Australian cohort. There was no appreciable difference in the reliability of the 24-item scale over the much shorter 12-item scale.

There was a small decrease in Cronbach’s alpha score in the 12 item BPIP in the New Zealand cohort of 0.09 over the 24 item scale although the scale can still be considered as reliable. The correlation analysis of the items was combined with all the other steps of item reduction, to produce the definitive BPIP scale, which proved to be statistically reliable.

The results demonstrate that a high proportion of the variance in the RDQ score at 12 weeks was accounted for by the 12 item BPIP, and statistically significant. The retained questions appear at the end the chapter. The BPIP question which made the strongest unique contribution to the dependent variable was; ‘The pain makes me feel that I can’t go on with my normal activities’. The individual’s perception of their low back pain being the greatest single contributor to their level of functioning, as measured by the RDQ, and interference with work and between episode limitations at work were the next two strongest single contributors.

Interference with work was the strongest single contributor to predicting the dependent variable, which may be expected when the more acute nature of the New Zealand cohort’s low back pain is taken into account. Interestingly the next strongest contributor to the prediction of the dependent variable in the New Zealand cohort was the statement that; ‘My
low back pain is dominating my life’, with the statement: ‘I have no hope of ever getting back to normal activities’ the next strongest predictor. Despite the New Zealand group being a more acute group by duration of low back pain reported at baseline, these perceptions, and mal-adaptive cognitions about the low back pain appear to be evident. This suggests that early on in the duration of an episode of low back pain, perception and mal-adaptive cognitions may well play a part in influencing prognosis, and can be readily quantified by the BPIP.

4.2 Comparison with other studies

The variance in RDQ scores was accounted for by the 12 item BPIP in the New Zealand cohort, (78.2%) which was considerably higher than the result of Burton and co-workers (Burton et al. 1995). This is potentially a reflection on the wider biopsychosocial domains of the BPIP capturing a greater range of influences on function when correlated against the RDQ. The result was also statistically significant.

The variance in RDQ score accounted for by the BPIP is similarly higher than that reported by Foster and co-workers in 2010. The prospective cohort study of Foster and co-workers of 3019 adults in the UK assessed distinctiveness of psychosocial obstacles to recovery and their multivariate model accounted for 47.7% in variance of RDQ score at 6 months.

A poor perception of outcome, high disability levels, and high self rated pain intensity were also identified as indicators of a poor prognosis in a large inception cohort study reported by Costa and co-workers (Costa, Maher et al. 2009). The large cohort demonstrated more than 50% of respondents had not recovered fully from pain or disability after 12 months and highlighted the desirability of an early biopsychosocial estimation of prognosis to inform patient management.

The size of the cohort compared favourably with other predictive instrument studies reported in the literature such as Hurley and co-workers (n=118) (Hurley, Dusoir et al 2001) and Stratford and co-workers (n=88) (Stratford, Binkley et al. 1996).

4.3 Limitations

The biopsychosocial model itself is used in a variety of ways within the literature, with any number of interchangeable variables being included from time to time. This makes a definitive statement about the model and which variables to include for statistical analysis somewhat difficult.

There is also some debate about the relative weight that should be attributed to the biomedical or psychosocial dimensions and whether the literature has become too focused on individual aspects of the model, rather than the overall inclusive biopsychosocial concept. However the literature continues to reflect the need for clinicians to be able to consider their patients from a biopsychosocial perspective in order to inform both prognosis and target interventions.

Self-report and perception of the patient’s condition has also been identified as being vulnerable to the individual’s social experience (Sen.2002).

The sample size also potentially limits the generalisability of the results.
The difficulties in recruitment of respondents may also be considered a limiting factor to the research.

The resulting scale will also require alteration to an entirely ordinal scale to be user friendly for routine clinical use. Scales need to be quick and easy to both administer and score to be a useful clinical resource. These issues will need to be addressed and the results confirmed in other populations in future research.

5. Conclusion

Predicting the outcome from an episode of low back pain has previously been reported and described in terms of the uni-dimensional biomedical perspective, rather than the contemporary biopsychosocial model, although it is evident that psychosocial factors or contextual factors can be a major obstacle to recovery. Clinicians are often encouraged to seek further information such as diagnostic imaging when a patient does not meet their expectation for recovery, rather than considering psychosocial obstacles to recovery. Routine clinical measurement as part of prognostication potentially encourages consideration of all the potential variables that may impact on recovery, and contributes to evidence-based ‘best practice’.

When psychosocial obstacles are not identified or duly considered in clinical practice, there is clearly a risk that some patients will go on to develop chronic pain and disability. Early identification of poor prognostic factors may potentially help target specific intervention and improve case management, and an objective measure may potentially contribute to this process. The BPIP potentially fills a gap in measurement, as a valid and reliable prognostic instrument, developed from a clinical perspective and intended for routine clinical use, following further validation in differing groups of low back pain patients.

6. 12-Item BPIP

Q1. Please score the severity of your low back pain as an average over the last 7 days where 0 equals no pain, and 10 equals worst pain.

0 1 2 3 4 5 6 7 8 9 10

Q2. How much has your low back pain interfered with your normal work (including housework) over the last 7 days?

Please circle one answer

Not at all: A little: Quite a bit: A lot: Totally

Q3. How much has your low back pain interfered with your normal social (inc sport) life over the last 7 days?

Please circle one answer

Not at all: A little: Quite a bit: A lot: Totally

Q4. I can only walk short distances because of my low back pain

1 2 3 4 5
Q5. Everything I do I consider how it will affect my low back pain  
1 2 3 4 5
Q6. My low back pain is dominating my life  
1 2 3 4 5
Q7. My low back pain disturbs my sleep  
1 2 3 4 5
Q8. The pain makes me feel that I can’t go on with my normal activities  
1 2 3 4 5
Q9. I believe I will get back to my normal level of activities  
1 2 3 4 5
Q10. I have no hope of ever getting back to normal activities  
1 2 3 4 5
Q11. In between episodes does your back limit what you can do at work?  
Please circle one response
Q12. In between episodes does your back limit what you can do socially (inc sport)?  
Please circle one response
Not at all: Occasionally: Often: All the time

7. References

Harding, G., Campbell, J., Parsons, S., Rachman, A., Underwood, M. British pain clinic practitioners' recognition and use of the biopsychosocial pain management model


Young, A.E., Wasiak, R., Phillips, L., and Gross D.P. Workers perspectives on low back pain recurrence: „It comes and it goes and comes and goes, but it’s always there’ 2011. Pain:152:204-211
This book includes two sections. Section one is about basic science, epidemiology, risk factors and evaluation, section two is about clinical science especially different approach in exercise therapy. I envisage that this book will provide helpful information and guidance for all those practitioners involved with managing people with back pain-physiotherapists, osteopaths, chiropractors and doctors of orthopedics, rheumatology, rehabilitation and manual medicine. Likewise for students of movement and those who are involved in re-educating movement-exercise physiologists, Pilates and yoga teachers etc.

How to reference
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