Chapter 2

Tools for Tinnitus Measurement: Development and Validity of Questionnaires to Assess Handicap and Treatment Effects

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Abstract

Tinnitus is a chronic condition that affects about 15% of the population and up to one in three older adults. For some it is a mild annoyance, for others it can be extremely distressing and can significantly deteriorate quality of life. Perceptual characteristics are a poor indicator of clinical need. Clinicians and researchers alike rely on self-report or questionnaires to quantify the severity of an individual’s tinnitus and to gauge the changes in tinnitus severity or tinnitus-related handicap over time or after clinical intervention. This book chapter evaluates the psychometric properties of five tinnitus questionnaires: Tinnitus Handicap Questionnaire, Tinnitus Reaction Questionnaire, Tinnitus Questionnaire, Tinnitus Handicap Inventory, and Tinnitus Functional
Index. We critically appraise the development process, validation, and responsiveness. We consider the true utility of each questionnaire to measure the short and long term consequences of tinnitus.

**Introduction**

Evidence-based assessment and treatment of tinnitus is important (e.g., Department of Health, 2009). However, tinnitus is notoriously difficult to measure objectively because it is an experiential phenomenon and can significantly differ between individuals. Objective measures include matching the pitch and loudness of tinnitus to an external sound. However, there is only a weak relationship between the psychoacoustic properties of sound (i.e., pitch or loudness) and its functional impact on the individual. Across those individuals with similar psychoacoustic attributes, there can be significant variability in self-reported handicap or the domains of impairment that are attributed to their tinnitus. It is not possible to determine or predict tinnitus distress based on the tinnitus pitch and loudness (Zeman et al., 2011; Andersson et al., 2005; Jakes et al., 1985). Perceptual characteristics therefore are not a good indicator of clinical need.

Tinnitus is multidimensional. Different symptom domains include emotional distress, problems with sleep, concentration or quality of life (QoL). Comorbidities in psychological well-being, such as stress, generalised anxiety and depression, and cognitive impairments profoundly impact on daily functioning (Stevens et al., 2007; Newman et al., 2011; Robinson et al., 2003). The degree to which tinnitus distress is perceived by patients can depend on the impact that tinnitus has on these factors or vice versa. For some it is a mild annoyance, for others it can be extremely distressing and significantly deteriorates quality of life.

Self-report measures, especially questionnaires, are the primary way to quantify the severity of an individual’s tinnitus and to assess the changes in tinnitus severity or tinnitus-related handicap over time or after clinical intervention (Meikle et al., 2007). Although, it is not compulsory to include a standard questionnaire in clinical practice (Hesser, 2000), most clinicians in NHS audiology departments in England do use questionnaires to assess tinnitus severity (67% out of 138 respondents; Hoare et al., 2012). The Department of Health (DH) guidelines recommend the use of validated tinnitus questionnaires; namely the Tinnitus Handicap Inventory (THI) (Newman et al., 1996) and Tinnitus Questionnaire (TQ) (Hallam, 1996; 2008)
for identifying tinnitus severity at intake assessment (Department of Health, 2009). Validated questionnaires can offer clinicians and researchers alike a systematic approach to quantifying tinnitus severity by distinguishing individuals who are bothered by their tinnitus from those who are not. This is a vital prerequisite for triaging patients effectively into the most appropriate interventions such as intensive management or education and advice. Questionnaires are also used for standardising selection criterion in research. For example, they identify people with similar degrees of tinnitus severity allowing the research to focus on a specific subtype.

Question items addressing different symptom domains can be categorised into subscales. Through analysing the responses in each subscale, clinicians can identify where the tinnitus distress lies and the specific domains of concern, so targeting the interventions. Tinnitus questionnaires can also be used as pre- and post-treatment outcome measures, therefore providing evidence of the changes that can occur in tinnitus severity. This is extremely important since there has been a move towards evidence-based commissioning within NHS services where healthcare professionals need to demonstrate to third-party payers the efficacy of management (NHS White Paper, 2010). Therefore there is a need to focus on evidence-based cost effective outcome measures that will improve patient experiences and the efficacy of interventions (NHS White Paper, 2010).

**Evaluating and Validating a Questionnaire**

Quality criterion for developing well validated questionnaires addresses four broad topics: reliability, validity, responsiveness and interpretation (Terwee et al., 2007). These are briefly summarised below.

**Reliability**

1. **Internal consistency.** The tinnitus questionnaire may assess different dimensions (subscales) and all the items in a subscale measure the same construct of tinnitus handicap.
2. **Reproducibility – reliability.** People with tinnitus can be distinguished from each other, despite measurement errors (relative measurement error)
3. **Reproducibility – agreement.** The tinnitus questionnaire score for individuals tested on multiple occasions over a short time period (1-2 weeks) are close to each other (absolute measurement error).
Validity
4. Content validity. The items in the tinnitus questionnaire are a correct and comprehensive reflection of the tinnitus handicap that the questionnaire is intended to measure.
5. Structural validity. The scores of the tinnitus questionnaire adequately reflect the construct of tinnitus handicap (i.e., the subscales explain at least 50% of the variance).
6. Construct validity. The scores demonstrate expected correlations between similar measures and expected differences between unrelated measures and these hypotheses should be defined a priori.

Responsiveness
7. Responsiveness. The tinnitus questionnaire is able to detect clinically important change over time.
8. Floor and ceiling effects. Few respondents achieve the lowest or highest possible score so that the tinnitus questionnaire does not compromise responsiveness, nor content validity and reliability.

Interpretation
9. Interpretability. It is possible to assign qualitative meanings to the quantitative scores, preferably with an indication of what change represents a minimal clinically important change for the patient group.

The remainder of this section considers and describes the techniques underlying each of the main quality criteria. It is recommended that the techniques are conducted from development to validation in the order presented below (Bland and Altman, 2002; Streiner & Norman, 2008). Table 1 provides a summary and definition of the different evaluation and validation techniques introduced in this review, and is intended to be used as a reference.

Step One: Ensuring the Questionnaire Content

As a useful general guide, developers should assess content validity to ensure that they (1) properly define what is being measured, (2) confirm that all aspects of the questionnaire (item choice, response options, and structure) are reviewed and judged by a panel of experts, (3) clearly report the development process, (4) ensure that any refinements to questionnaires
undergo further evaluations, and (5) empirically validate the factorial structure (Fitzpatrick, 1983; Haynes et al., 1995; Streiner and Norman, 2008).

**Table 1. Summary and definition of the evaluation and validation techniques described in this review**

<table>
<thead>
<tr>
<th>Validation technique</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Content validity</td>
<td>Content validity is used during the initial questionnaire development to assess the degree in which all facets, such as items, response formats, and instructions are relevant to and represent the target construct (concept, attribute or variable) of the questionnaire. It is based on the qualitative judgements of experts, preferably using a 5-7 point evaluation scales for agreement.</td>
</tr>
<tr>
<td>Frequency response distributions</td>
<td>Frequency response distribution provides information on the items ability to discriminate between individuals and to show sensitivity to change. It is the frequency count of a specific rating assigned to each item.</td>
</tr>
<tr>
<td>Cronbach’s alpha</td>
<td>Cronbach’s alpha provides a measure of the internal consistency. Inter-item correlations are an estimate of the proportion of variability due to the “true score” rather than measurement error. It is assumed that all of the items are tapping into the same construct and therefore positively correlate with each other. It is measured by essentially splitting the data in two in every possible way, and calculating the correlation coefficient for each split, therefore providing an average of these two values.</td>
</tr>
<tr>
<td>Inter-item correlations</td>
<td>Inter-item correlations provide a measure of internal consistency, assesses the association between pairs of items. This is an important component of item analysis as it highlights the inter-relatedness of the items and provides an indication of redundant items that are highly correlated. It is measured by correlating each item on the questionnaire with every other item.</td>
</tr>
<tr>
<td>Item-total correlations</td>
<td>Item-total correlations provide a measure of internal consistency, assessing the association between an item and the total score. It checks that the item is measuring the same construct as the test. It is measured by calculating the correlation coefficients of individual items with the questionnaire total, whilst omitting that item. It is computed using all the remaining items on the test.</td>
</tr>
<tr>
<td>PCA: Orthogonal varimax rotations</td>
<td>Orthogonal rotations improve factor extraction, maximising item loading. It is used to rotate the factors when the underlying factors are assumed to be independent of each other.</td>
</tr>
</tbody>
</table>
Table 1. (Continued)

<table>
<thead>
<tr>
<th>Validation technique</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCA: Oblique rotations</td>
<td>Oblique rotations improve factor extraction, maximising item loading. This is a linear rotation that assumes that underlying factors are related or correlated with each other.</td>
</tr>
<tr>
<td>Construct validity</td>
<td>Questionnaires designed to tap into health behaviour constructs should be based on underlying theories and therefore should measure aspects that are consistent with this theory and construct. Construct validity assesses the extent to which a questionnaire actually measures the target construct it purports to measure. This type of validity consists of two components; convergent and discriminant validity.</td>
</tr>
<tr>
<td>Convergent validity</td>
<td>A component of construct validity. Convergent validity measures the extent to which the construct of a new questionnaire corresponds with other questionnaire constructs that are theoretically similar. It is measured by calculating the correlations coefficients between the questionnaires, and assessing the strength of the association.</td>
</tr>
<tr>
<td>Discriminant validity</td>
<td>A measure of construct validity. Assesses the extent to which the underlying construct of the new questionnaire can differentiate between constructs that are theoretically independent. It is measured by calculating the correlations coefficients between the questionnaires, and assessing the strength of the association.</td>
</tr>
<tr>
<td>Test-retest reliability</td>
<td>Assesses the consistency and stability of the questionnaire over time. Test-retest reliability determines the extent to which the scores maintained standing from test to retest. It is measured by calculating the correlations coefficients of two set scores either for a single item, subscale or total scale from the two different administrations points in time.</td>
</tr>
<tr>
<td>Minimal clinically important change</td>
<td>The clinical significance of a treatment requires knowledge about the minimal change in total scores of the specific questionnaire used. The minimal clinically important change score refers to the smallest change in total scores between baseline and final assessment that can be considered clinically relevant and be attributed to treatment benefits. This score is considered important for the design of clinical trials, the interpretation of the scores for the clinical significance of a treatment and its ability to meet standards of efficacy.</td>
</tr>
<tr>
<td>Global question</td>
<td>Usually a single question that is used to measure one aspect of the health behaviour, such as symptom severity, using a 5-point or 7-point response scale. Global questions are anchor-based techniques that provide a way to assess the corresponding changes in questionnaire total scores.</td>
</tr>
</tbody>
</table>

The order of the information corresponds to each step in evaluating and validation section. PCA: Principal Component Analysis.
Step Two: Validating the Questionnaire Structure

Validation provides a theoretical basis for the questionnaire construct (i.e., the concept, attribute or variable that is the target of measurement). Principal Component Analysis (PCA) is a popular technique for identifying the underlying domains of a questionnaire (Field, 2009). Essentially PCA identifies sets of correlated variables (Factors) within the questionnaire items. The factorial structure is informed by the inter-item relationships, computed from the inter-item correlations (factor loadings) between each variable in the analysis (Floyd, 1995). Factor interpretations are improved by rotating the data. These rotations can either be oblique or orthogonal in form. They are designed to maximise item loading on the extracted factor whilst reducing the loading on the other factors. A ‘good’ tinnitus questionnaire would be expected to cover all the important domains of tinnitus handicap (e.g., Kennedy et al., 2004). In other words, it would have a multifactor structure.

The internal structure (“internal consistency”) is analysed using Cronbach’s alpha, inter-item and item-total correlations, to assess the consistency of the item content so ensuring that all the items in a questionnaire are measuring the same underlying construct (Clark and Watson, 1995; Cortina, 1993; Table 1). For instance, a Cronbach’s alpha score \( \alpha > 0.7 \) indicates acceptable internal consistency (Peterson, 1994; Table 2). In other words, the items all contribute to the overall construct.

Table 2. Category definitions of statistical tests: Cronbach’s alpha and correlation coefficient

<table>
<thead>
<tr>
<th>Type of test</th>
<th>Score</th>
<th>Category definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cronbach’s alpha (( \alpha ))</td>
<td>0.90 – 1.00</td>
<td>Extremely high</td>
</tr>
<tr>
<td></td>
<td>0.80 – 0.89</td>
<td>Good</td>
</tr>
<tr>
<td></td>
<td>0.70 - 0.79</td>
<td>Acceptable</td>
</tr>
<tr>
<td></td>
<td>0.60 – 0.69</td>
<td>Questionable</td>
</tr>
<tr>
<td></td>
<td>0.50 – 0.59</td>
<td>Poor</td>
</tr>
<tr>
<td></td>
<td>&lt; 0.50</td>
<td>Unacceptable</td>
</tr>
<tr>
<td>Correlation coefficient score (( r ))</td>
<td>0.80 – 1.00</td>
<td>Extremely strong</td>
</tr>
<tr>
<td></td>
<td>0.60 – 0.79</td>
<td>Strong</td>
</tr>
<tr>
<td></td>
<td>0.30 – 0.59</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td>0.00 – 0.29</td>
<td>Weak</td>
</tr>
</tbody>
</table>
Step Three: Validating the Questionnaire Construct

To ascertain relevant and useful interpretations that ‘truly’ reflect the individual’s perceived ability, the questionnaire should reliably measure the target construct health behaviour (Newman and Sandridge, 2004). A questionnaire should be assessed for the extent to which it actually measures the target construct it purports to measure (“construct validity”). Any new questionnaire should be compared, using correlation coefficients, to existing questionnaires that measuring constructs that are theoretically similar (“convergent validity”) or constructs that are independent (“discriminant validity”). For example, a new tinnitus questionnaire would be expected to show relatively high correlations (i.e., high convergence) with other tinnitus questionnaires, and relatively low correlations (i.e., high discriminance) with generalised depression, anxiety and hearing handicap questionnaires because these represent theoretically distinct constructs (see Table 2).

Step Four: Validating the Interpretations of the Scores

It is important to be able to reliably grade tinnitus severity if treatment efficacy or treatment needs are to be established. It is also beneficial to interpret questionnaire scores according to categories that provide a clinical meaning to the numerical score. This enables clinicians and researchers alike to identify and quantify tinnitus severity. There does not appear to be a universally accepted guideline on how best to define categories and develop a grading system. One technique is to conduct quartile analysis on normative datasets. The quartile analysis simply divides the patient population into four categories based on the distribution of the total score before treatment. However, quartile scores are limited. Although often based on large normative datasets, important information related to patient experience is omitted. This could be provided through using a tinnitus severity global question to compare the perceived tinnitus severity to the overall scores (i.e., a reference anchor).

Step Five: Validating the Questionnaires Sensitivity to Change (Responsiveness)

Validating sensitivity should initially begin during questionnaire development. Item selection should include items that are sensitive to changes
and the response scale should also be sensitive to small changes. Since the amount of response options are proportional to the items sensitivity to change, high resolution numerical scales (0-10) are recommended (Kirshner and Guyatt, 1995; Meikle et al., 2007). Changes in observed questionnaire scores should reliably and sensitively reflect change in the health behaviour being measured over time or after intervention.

To measure the consistency and stability of the questionnaire scores over time, test-retest reliability is used to assess the extent to which the scores are maintained from test to retest (without any interventions between the tests) (DeVon et al., 2007). Clinically meaningful treatment effects can impact upon the management of patients, and the ability to meet the requirements set by patients (Meikle et al., 2007). In order to measure the sensitivity of the questionnaire scores to changes in health behaviour, the variation that is clinically meaningful has to be separated from the measurement error that you naturally see between test and retest. To address this, the “minimal clinically important change” concept was developed (Jacobson et al., 1986; Jacobson et al., 1999; DeVet et al., 2006). This is defined as the value at which the change becomes clinically relevant to the patient. Two different approaches can be used to provide a comparison for the mean before-after treatment scores. (1) Distribution-based approaches are based on the statistical properties of the sample, i.e., the effect size (the difference in mean change and standard deviation change) (DeVet et al., 2006; Revicki et al., 2008). (2) Anchor-based approaches use an external indicator, such as global question response categories, to provide a reference of change (DeVet et al., 2006; Revicki et al., 2008).

**Tinnitus Questionnaires**

Over the past 30 years, a range of questionnaires have been developed to scale the severity and impact of tinnitus. These identify a number of dimensions associated with the complex construct of tinnitus severity, relating to sensory, behavioural and emotional reactions.

This section provides an in-depth evaluation of the psychometric properties of five tinnitus questionnaires, critically appraising their process of development, validation, and responsiveness. It focuses on the Tinnitus Handicap Questionnaire (THQ; Kuk et al., 1990), Tinnitus Questionnaire (TQ;
Hallam et al., 1988; Hiller and Goebel, 1992), Tinnitus Reaction Questionnaire (TRQ; Wilson et al., 1991), Tinnitus Handicap Inventory (THI; Newman et al., 1996) which appear in practice recommendations by the UK Department of Health and the international Tinnitus Research Initiative, and finally one of the most recent questionnaires to be developed; the Tinnitus Functional Index (TFI; Meikle et al., 2012).

In this section, the five questionnaires are presented in the order in which they were developed. The five questionnaires are evaluated following the validation steps above. Each questionnaire section starts with sections on initial questionnaire development (validation step one), whether the questionnaire covers all important tinnitus domains (validation step two), the two components of construct validity (validation step three), the ability of the questionnaire to grade tinnitus severity (validation step four) and the ability of the questionnaire to be responsive to treatment (validation step five).

**Tinnitus Questionnaire (TQ)**

From the best available information, it appears that the Tinnitus Questionnaire (TQ) was developed from the Tinnitus Effects Questionnaire (TEQ) (Hallam et al., 1988). However, the literature is unclear; there is no explicit description of the connection between these two questionnaires (see Henry and Wilson, 1998; Kennedy et al., 2004). Therefore, it is assumed that the TQ development started with Hallam et al. in 1988. Evidence from literature on the TEQ (i.e., Henry and Wilson, 1998) where the description of the questionnaire structure and scoring matches that of the TQ will be reviewed.

The TQ was developed by researchers at Royal National Throat Nose and Ear Hospital, London (Table 3; Hallam et al., 1988). Primarily designed to measure tinnitus severity, it is also used to evaluate change and to examine the relationship of different facets of complaint and other psychological variables to tinnitus (Hallam, 2008). For each item, individuals indicate the level of agreement using one of three response options; not true (0), partly true (1) and true (2). The total score is rescaled from the weighted sum of the items used in each subscale (41 items) so that the global score ranges from 0 –82, with higher scores indicating increased tinnitus distress (Table 4).
Table 3. Characteristics and psychometric properties of the five widely-used tinnitus questionnaires

<table>
<thead>
<tr>
<th>Questionnaire (Author, year)</th>
<th>Response options</th>
<th>No of items</th>
<th>Reported aims of measure</th>
<th>Psychometric properties</th>
<th>No. of participants</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tinnitus Questionnaire (TQ) (Hallam et al., 1988; Hallam, 1996, 2008)</td>
<td>3 levels: True, partly true, not true</td>
<td>52 items</td>
<td>5 subscales: Emotional and cognitive distress, Intrusiveness, Auditory Perceptual difficulties, Sleep disturbance, Somatic complaints</td>
<td>Measures psychological aspects of tinnitus complaints and distress</td>
<td>Cronbach’s Alpha $\alpha = 0.91$</td>
</tr>
<tr>
<td>Tinnitus Handicap Questionnaire (THQ) (Kuk et al., 1990)</td>
<td>100 levels: 100 = strongly agree, 0 = strongly disagree</td>
<td>27 items</td>
<td>3 subscales: Physical, emotional &amp; social effects of tinnitus, Hearing and communication ability, Individual’s perception of tinnitus</td>
<td>Measure of patients’ perceived degree of handicap due to tinnitus</td>
<td>Cronbach’s Alpha $\alpha = 0.94$</td>
</tr>
<tr>
<td>Tinnitus Reaction Questionnaire (TRQ) (Wilson et al., 1991)</td>
<td>5 levels: Not at all to almost all the time</td>
<td>26 items</td>
<td>No clear subscales</td>
<td>Measure psychological distress associated with tinnitus</td>
<td>Cronbach’s Alpha $\alpha = 0.96$</td>
</tr>
<tr>
<td>Tinnitus Handicap Inventory (THI) (Newman et al., 1996)</td>
<td>3 levels: Yes Sometimes No</td>
<td>25 items</td>
<td>3 subscales: Functional, Emotional, Catastrophic</td>
<td>Measures the level of perceived tinnitus severity</td>
<td>Cronbach’s Alpha $\alpha = 0.93$</td>
</tr>
<tr>
<td>Tinnitus Functional Index (TFI) (Meikle et al., 2012)</td>
<td>11 levels: 0 – 10 Descriptive anchors vary between items</td>
<td>25 items</td>
<td>8 subscales: Intrusiveness, Sense of control, Sleep, Cognition, Hearing, Relaxation, Emotional distress, Quality of life</td>
<td>Developed to measure both tinnitus severity and treatment-related changes</td>
<td>Cronbach’s Alpha $\alpha = 0.97^d$</td>
</tr>
</tbody>
</table>

$^a$ data from Newman et al., 1995.
$^b$ data from Hiller et al., 1994.
$^c$ data from Newman et al., 1998.
$^d$ 25-item prototype 2 data.
Table 4. The progression of the Tinnitus Questionnaire development

<table>
<thead>
<tr>
<th>Year Version (Author)</th>
<th>No. of Items (factors)</th>
<th>Subscales</th>
<th>No. of items in subscales</th>
<th>Cronbach’s alpha(α)</th>
<th>Scoring</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prototype (Hallam et al., 1988)</td>
<td>40 (3/4)</td>
<td>Emotional distress</td>
<td>20 items</td>
<td>N/A</td>
<td>N/A Remaining items:10</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Auditory perceptual difficulties</td>
<td>6 items</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sleep disturbance</td>
<td>4 items</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Final TQ (Hallam et al., 1988)</td>
<td>51 (3)</td>
<td>Sleep disturbance</td>
<td>13 items: 2, 4, 18, 20, 12, 31, 34, 35, 36, 39, 41, 48, 50</td>
<td>N/A</td>
<td>N/A Remaining items:17</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Emotional distress</td>
<td>15 items: 3, 5, 7, 10, 11, 13, 15, 19, 24, 27, 37, 39, 47, 43, 45</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Auditory perceptual difficulties</td>
<td>6 items: 9, 14, 18, 26, 33, 38</td>
<td></td>
<td></td>
</tr>
<tr>
<td>German version GHTQ (Hiller &amp; Goebel, 1992)</td>
<td>52 (5/6)</td>
<td>Emotional distress*</td>
<td>12 items: 1, 5, 8, 11, 16, 18, 19, 20, 28, 37, 39, 41</td>
<td>α = .85</td>
<td>Scores range: 0-84</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cognitive distress*</td>
<td>8 items: 3, 13, 17, 21, 27, 43, 44, 47</td>
<td>α = .85</td>
<td>Global score based on 42 items (α = .93)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Intrusiveness</td>
<td>8 items: 5, 7, 10, 15, 20, 34, 35, 48</td>
<td>α = .75</td>
<td>Remaining items:12</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Auditory perceptual difficulties</td>
<td>7 items: 2, 9, 14, 26, 33, 38, 50</td>
<td>α = .86</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sleep disturbance</td>
<td>4 items: 4, 12, 31, 36</td>
<td>α = .80</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Somatic complaints</td>
<td>3 items: 22, 25, 51</td>
<td>α = .74</td>
<td></td>
</tr>
<tr>
<td>Revised TQ (Hallam, 1996)</td>
<td>52 (5)</td>
<td>Emotional and cognitive distress</td>
<td>19 items: 3, 8, 13, 16, 17, 18, 19, 20, 21, 24, 27, 28, 30, 37, 39, 41, 43, 44, 47</td>
<td>α = .94</td>
<td>Scores range: 0-82</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Intrusiveness</td>
<td>7 items:</td>
<td>α = .87</td>
<td>Overall score</td>
</tr>
</tbody>
</table>
## Tools for Tinnitus Measurement

<table>
<thead>
<tr>
<th>Year Version (Author)</th>
<th>No. of Items (factors)</th>
<th>Subscales</th>
<th>No. of items in subscales</th>
<th>Cronbach's alpha(α)</th>
<th>Scoring</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>based on 41 items (α = .95)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Remaining items:11</td>
</tr>
<tr>
<td></td>
<td></td>
<td>auditory perceptual difficulties</td>
<td>5, 7, 10, 11, 15, 35, 45</td>
<td>α = .89</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>sleep disturbance</td>
<td>4 items: 2, 9, 14, 26, 33, 38, 50</td>
<td>α = .81</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>somatic complaints</td>
<td>4 items: 22, 25, 29, 34</td>
<td>α = .75</td>
<td></td>
</tr>
</tbody>
</table>

TQ: Tinnitus Questionnaire, GHTQ Goebel and Hiller Tinnitus Questionnaire.

*Emotional and Cognitive distress can also be combined. Numbers that appear in bold are featured twice in the questionnaire, in different subscales.

Unfortunately, the British version - the TQ (Hallam) – has little published information available. Apart from Hallam et al. (1988), the only evidence available is in the 2008 TQ manual (Hallam, 2008). The 1996 TQ manual (Hallam), where the revisions of TQ internal structure are reported, is out of print. Any evidence provided of discriminant and convergent validity in 1988 is now redundant because of subsequent modifications to the questionnaire. For example, the revisions to the TQ in 1996 mean that “The present scoring of subscales […] supersedes the earlier scoring system” (Hallam, 2008, p. 9). The revised TQ (Hallam 1996; 2008) presented an opportunity to conduct full psychometric validation. However, Hallam does not appear to have conducted any additional validation beyond that of validating the internal TQ structure. Evidence reported in the 2008 manual is based on unpublished data and redundant 1988 data (i.e., Hallam et al., 1988). In this review, we will refer to evaluation and validation evidence from the German translation (GHTQ; Hiller and Goebel, 1992; Goebel and Hiller, 1998). The global score ranges from 0 – 84 for the GHTQ, this is based on the 42 items in the subscales (out of a total 52 items) with two items are included twice (Q5 and Q20).

### Initial Questionnaire Development

From the information provided by Hallam et al. (1988) on the development of the TQ (prototype/final versions), it is difficult to ascertain the process and quality of development. There is only a reference to the choice of items being based on clinical observations of the most commonly reported
adverse effects and complaints associated with the multiple domains of tinnitus (see Hallam et al., 1988; Hallam, 2008).

The authors do refer back to the previous work by Jakes et al. (1985). They investigated the relationship between loudness and annoyance of tinnitus with a forced-choice questionnaire. Tinnitus handicap was found to have two dimensions (emotional distress and intrusiveness). It is not apparent from the literature how these two factors informed Hallam and colleagues’ decisions on item choice and yet they are regularly referred to as the basis of the TQ (Hiller and Goebel, 1992; Henry and Wilson, 1998; Gerhards et al., 2004). No further information was reported on the item choice.

No empirical validation was conducted until 1996 in the UK (Hallam, 1996). This work produced a revised TQ (Table 4). Previous items were unchanged, but the questionnaire now included an additional item that was “accidently missed” out of the 1988 version due to clerical error (Hallam, 2008). To compute the global and subscale scores for the TQ only 41 out of the possible 52 items are used, and for the GHTQ only 42 items are used (Table 4). The remaining items are considered as baseline information that may be clinically useful (Hallam, 2008, p.7). In fact they could be considered a hindrance as they increase the time required to administer the questionnaire and there is limited evidence for their clinical relevance (Hiller and Goebel, 2004; Newman and Sandridge, 2004).

**Questionnaire Structure**

The TQ purports to encompass the major domains of tinnitus (Hallam et al., 1988; Hallam, 2008). Over the years the factorial structure of the TQ has been revised and consequently the number of domains purported to be covered by TQ has changed (Table 4) (Hiller and Goebel, 2004).

**Factorial Structure**

After conducting PCA, the 1988 version (Hallam et al. 1988) reported three factors based on 34 of the 51 items. These were i) emotional distress, (ii) auditory difficulties and (iii) sleep disturbance. There was no further empirical validation. There is no evidence for the internal consistency of each subscale and the selected 34 items of the TQ.
In 1992, Hiller and Goebel investigated the psychometric properties of the GHTQ (German translation). PCA with orthogonal varimax rotations, followed by factorial stability analysis, revealed six main factors for interpretation: (i) emotional distress, (ii) auditory difficulties, (iii) sleep disturbance, (iv) cognitive distress, (v) intrusiveness, and (vi) somatic complaints (Hiller and Goebel, 1992). The first three factors had previously been identified by Hallam et al. (1988). The intrusiveness factor had previously been identified by Jakes et al. (1985) but not by Hallam et al. (1988).

In 1996, Hallam reinvestigated the factorial structure of the TQ. Using the same techniques as Hiller and Goebel (1992), a six factor solution was revealed, but only five factors were considered reliable. These were (i) emotional and cognitive distress, (ii) auditory difficulties, (iii) sleep disturbance, (iv) intrusiveness, and (v) somatic complaints. Here, the emotional and cognitive factors previously reported by Hiller and Goebel (1992) were combined into a single factor. The 2008 manual provides a brief explanation of the 1996 revalidation, but at times the information about how the questionnaire items load onto each factor is contradictory and vague. For instance, different items are listed under each factor throughout the manual (Seydel et al., 2012).

Nevertheless, the five factor structure of the TQ has also been replicated in its Cantonese (Kam et al., 2009), Dutch and French (Meeus et al., 2007) translations. The factorial structure of the 52-items would suggest that the TQ (Hallam, 1996; 2008) measures five separate domains of tinnitus distress, but some questions remain about the reliability of this conclusion.

It is important to highlight that the PCA analyses described GHTQ and TQ have been based on all 52 TQ items, not on the 41 items that contribute to calculate the global score. This aspect of the statistical methodology could have a major impact on the resulting factorial structure of the global TQ score (Gerhards et al., 2004).

Indeed, following PCA on the GHTQ’s 42 ‘overall score’ items, Gerhards reported only one or two factors instead of the previously reported five-six factors (Gerhards et al., 2004). Furthermore, just under half (47%) of the items focus on the emotional problems, although the remaining four tinnitus domains appear to be more evenly distributed (Figure 1; Kennedy et al., 2004).
Internal Consistency

The 2008 manual reported the internal consistency (Cronbach’s alpha) of the TQ. Alpha estimates were extremely high for the total score and sleep disturbance, good for emotional distress, auditory difficulties and intrusiveness, and acceptable for somatic complaints (Table 4). These high scores would suggest there is little variance between the individual items (Cronbach, 1951).
Questionnaire Construct

Construct Validity: Convergent Validity

The TQ measures constructs that are comparable to other tinnitus-related questionnaires. High convergent validity for TQ and TEQ has been shown by the strong correlations with the THQ, (r = 0.75), the TRQ (r = 0.74) (Henry and Wilson, 1998; Robinson et al., 2003), and the THI total (r = 0.89) and subscale scores (r = 0.79 to r = 0.86) (Baguley et al., 2000; Robinson et al., 2003). The TQ also showed strong correlations with the Tinnitus Cognitions Questionnaire (TCQ) negative subscale (r = 0.65) (Wilson and Henry, 1998).

Construct Validity: Discriminant Validity

The TQ demonstrates moderate to high discriminant validity indicating that it does measure constructs that are distinct from more generalised symptoms. The TQ shows moderately correlations with the Beck’s Depression Inventory (BDI; Beck et al., 1997) (r = 0.55) (Henry and Wilson, 1998; Robinson et al., 2003), the Hamilton Rating Scale for Depression (Hamilton, 1960) (r = 0.48), the Modified Somatic Perception Questionnaire scores (Main, 1983) (r = 0.46) and the Quality of Well-Being scale scores (Kaplan et al., 1996) (r = -0.37; Robinson et al., 2003). It has weak to moderate correlations with the Hopkins Symptom Checklist (SCL-90-R; Derogatis, 1977) (r = 0.26 to r = 0.39) (Hiller et al., 1994; Hiller and Goebel, 2004) and weak correlations with the Private Self-Consciousness Scale scores (Fenigstein et al., 1975) (r = 0.21; Robinson et al., 2003).

Interpretation of the Scores

Hallam (2008) suggests that quantifying tinnitus distress should be left to the clinicians’ discretion. Nevertheless, a grading system has been developed for the GHTQ (Goebel and Hiller, 1998) which provides clinical meaning to the scores. Gerhards et al. (2004) conducted quartile analysis on the data collected from 683 tinnitus patients in Germany and confirmed the same range of scores in the four categories. These data are summarised in Table 5.
Table 5. Grading systems providing qualitative meanings to the quantitative scores

<table>
<thead>
<tr>
<th>Score</th>
<th>Tinnitus handicap</th>
<th>Description of symptom severity</th>
</tr>
</thead>
<tbody>
<tr>
<td>THQ (Sullivan et al., 1993)</td>
<td>&gt;22</td>
<td>Bothersome tinnitus</td>
</tr>
<tr>
<td>GHTQ (Goebel &amp; Hiller, 1998)</td>
<td>0 – 30</td>
<td>Mild</td>
</tr>
<tr>
<td></td>
<td>31 – 46</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td>47 – 59</td>
<td>Severe</td>
</tr>
<tr>
<td></td>
<td>60 – 84</td>
<td>Very severe (heavy burden)</td>
</tr>
<tr>
<td>THI (McCombe et al., 2001)</td>
<td>0 – 16</td>
<td>Slight</td>
</tr>
<tr>
<td></td>
<td>18 – 36</td>
<td>Mild</td>
</tr>
<tr>
<td></td>
<td>38 – 56</td>
<td>Moderate</td>
</tr>
<tr>
<td></td>
<td>58 – 76</td>
<td>Severe</td>
</tr>
<tr>
<td></td>
<td>78 – 100</td>
<td>Catastrophic</td>
</tr>
<tr>
<td>TFI (Meikle et al., 2012)</td>
<td>&lt; 25</td>
<td>Mild</td>
</tr>
<tr>
<td></td>
<td>25 – 50</td>
<td>Significant problems with tinnitus</td>
</tr>
<tr>
<td></td>
<td>&gt; 50</td>
<td>Severe</td>
</tr>
</tbody>
</table>
Sensitivity to Change

Hallam (2008) claimed that the “Chief application of the TQ is in the evaluation and auditing of psychological interventions for tinnitus” (p.7). The evidence for this claim appears somewhat limited because the TQ items were not selected to be specifically responsive to treatment-related changes. It consequently lacks the required resolution to detect small changes in scores (Meikle et al., 2008). In all versions of the TQ (Hallam et al., 1988; Hallam, 1996; 2008), no data has been provided to determine a clinically significant change score following intervention (Newman and Sandridge, 2004). Therefore, clinicians and researchers alike would be advised not to rely on the TQ for auditing purposes or to determine minimal significant change.

Goebel et al. (2006) have categorised treatment effects using the TQ. They concluded 6-14 point change in TQ scores was indicative of improvement, i.e., the patient was “responding” to the treatment. They also classified a change of ≥15 points as a “winning response”, i.e., the patient was showing a large improvement. However, an alternative minimal clinically important change score for the GHTQ has been proposed by Adamchic et al. (2012). Using data from both the Randomized Evaluation of Sound Evoked Treatment of Tinnitus (RESET) trial study (N = 63) and Tinnitus Research Initiative (TRI) database (N = 694), Adamchic et al. (2012) analysed 757 patient responses on the GHTQ at baseline, clinical assessment and end of treatment. There was an average of 44 days between baseline and clinical assessment. Adamchic et al. (2012) used a combination of anchor-based and distribution-based techniques, such as Clinical Global Impression Improvement (CGI-I) rating, Receiver Operating Characteristic (ROC) curves and Standard Error Measurement (SEM) to produce the minimal clinically important change score for improvement. The CGI-I is an overall rating that uses up to seven response options to quantify treatment-related change in health behaviour, i.e., patients make a judgement on the total improvement of their tinnitus after treatment (Table 6a, 6b). For the analysis, the patient groups were formed according to the CGI scores. The seven CGI-I categories in the TRI database and the five CGI-I categories in the RESET database were combined into five groups; much better, minimally better, no change, minimally worse, much worse. The authors examined the predicted minimal clinically important change scores for each technique and choose the most representative score. The SEM scores estimated the lowest change score of -4.7 but the ROC curve predictions of a change score of -5 was considered most representative. This recommendation differs from that of Goebel et al. (2006). It’s derivation from a larger dataset
and using a combination of distribution-based and anchor-based methods, potentially offers a more precise estimate of meaningful change.

Table 6. Clinical Global Questions to determine patients judgement on perceived treatment-related change

<table>
<thead>
<tr>
<th>a) TRI Clinical Global Impression</th>
<th>b) RESET Clinical Global Impression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rate the total improvement of their tinnitus complaints compared to before the beginning of treatment.</td>
<td>Verbal rating of their tinnitus loudness and annoyance for each ear where the tinnitus was perceived as compared to baseline.</td>
</tr>
<tr>
<td>1. Very much better</td>
<td>1. Much better</td>
</tr>
<tr>
<td>2. Much better</td>
<td>2. Somewhat better</td>
</tr>
<tr>
<td>3. Minimally better</td>
<td>3. No change</td>
</tr>
<tr>
<td>4. No change</td>
<td>4. Somewhat worse</td>
</tr>
<tr>
<td>5. Minimally worse</td>
<td>5. Much worse</td>
</tr>
<tr>
<td>6. Much worse</td>
<td></td>
</tr>
<tr>
<td>7. Very much worse</td>
<td></td>
</tr>
</tbody>
</table>

c) TFI Global Question

All things considered, how is your overall tinnitus condition now, compared to your first visit to this clinic?

| 1. Much improved |
| 2. Moderately improved |
| 3. Slightly improved |
| 4. No change |
| 5. Slightly worse |
| 6. Moderately worse |
| 7. Much worse |

Summary

There is limited information available for the TQ, and the information that is available is somewhat poorly specified. The TQ appears to measure five separate domains of tinnitus distress. However, the validation of the factorial structure has generally not been conducted on the 41 items that contribute to the total TQ score and subscale scores. The TQ does have high construct
validity and would appear to measure tinnitus severity, although patient experience was potentially overlooked during the development of the grading system. Recently there have been attempts to overcome the lack of evidence for minimal clinically important change scores, with some success.

**Tinnitus Handicap Questionnaire (THQ)**

The 27-item THQ was developed to comprehensively measure a patient’s tinnitus handicap and to be sensitive to the changes in handicap over time (Kuk et al., 1990). It is claimed to measure three broad domains of tinnitus handicap: (i) the impact of tinnitus on social, emotional and physical aspects, (ii) hearing ability and unease, and (iii) the individuals’ perception of their tinnitus (Table 3). For each item on the THQ, the individual is asked to indicate how much he/she disagrees or agrees with the statement with a number between 0 (strong disagree) and 100 (strongly agree). The total score is rescaled from the weighted sum of all the items in each subscale so that the global score ranges from 0 – 100, with a higher score indicating greater handicap.

![Figure 2. Fifteen most common difficulties reported by tinnitus patients in open-ended questionnaire. Adapted from Tyler and Baker (1983).](image-url)
Initial Questionnaire Development

At the first step in the development, the authors obtained the 87 items directly from Tyler and Baker (1983; Figure 2) which were “arbitrarily” grouped into four domains that reflect tinnitus handicap (i.e., hearing, lifestyle, health, and emotion).

The authors briefly refer an additional domain on “others’ reaction to tinnitus”, but it is unclear how this additional domain was derived. Kuk et al. (1990) systematically reduced the initial 87 items to the final 27-item questionnaire, by sampling items that were sensitive to perceived tinnitus handicap. They did this by examining (i) response frequency distributions, (ii) inter-item correlations, and (iii) item-total correlations.

i. Items were eliminated based on their ability to differentiate between individual responses, i.e., the sensitivity of item. Items were assumed to be insensitive, if they were scored with either a 0 or 100 on the response options over 50% of the time. However, the authors do not clearly explain why they chose this criterion and it was applied inconsistently at this development stage. For example, the developers retained items with 50% “0” ratings because they “meet other criteria” (not specified).

ii. Highly correlated items were examined. The specific criterion value for high inter-item correlations was not reported, although an example of \( r > 0.60 \) was given as an example of redundant items. Multiple items that reflected the same situation, for example listening in quiet, were considered redundant. The most representative item of them was kept, whilst the rest of the redundant items were eliminated.

iii. Items were eliminated based on their “low” item-total correlations in order to maximise internal consistency. However, the specific cut-off value for a low item-total correlation was not reported. The authors considered the response frequency distributions (sensitivity) before eliminating the item based on the internal consistency scores. Two items were kept despite their low item-total scores.
Questionnaire Structure

Factorial Structure

Having conducted PCA and oblique rotations on the THQ, three factors were clearly identified (Kuk et al., 1990). Factor 1 represents social, emotional and physical functioning (15 items). Factor 2 represents hearing ability and unease (8 items), and factor 3 represents individual perception of tinnitus (4 items). Factors 1 and 2 were moderately correlated \( r = 0.49 \), but both were only very weakly correlated \( r = 0.10, r = 1.16 \) respectively) with factor 3. Items on factor 3 also resulted in extremely low item-total correlation scores \( r = 0.15 \) suggesting that it is potentially measuring separate variables to factors 1 and 2. Finally, the majority of items loaded onto factor 1 with some overlap of items onto factor 2 indicating that the THQ is particularly sensitive to the social, emotional and physical functioning aspects of tinnitus distress. This is consistent with Kennedy et al.’s (2004) critical review. Kennedy designated the items according to the six categories highlighted by Tyler and Baker’s open questionnaire study (1983) and found that the THQ places little emphasis on issues to do with sleep deprivation, cognitive disturbance or impact on health and well-being (Kennedy et al., 2004; Figure 1).

Internal Consistency

The THQ demonstrated extremely high Cronbach’s alpha for both the THQ total \( (\alpha = 0.94) \) and for factor 1 \( (\alpha = 0.94) \). These high alpha values could reflect the low uniqueness of the items content (Cortina, 1993). Some items could essentially be asking the same question in a different guise and therefore reflecting unnecessary duplication of content (Streiner, 2003). This provides additional evidence of a factorial structure that mainly focuses on a limited number of tinnitus domains. Factor 2 had a good Cronbach’s alpha \( (\alpha = 0.88) \) (Kuk et al., 1990). Factor 3 however, achieved unacceptable Cronbach’s alpha \( (\alpha = 0.47) \). Kuk et al. (1990) attributed this unacceptable score to the small number of items in the subscale and the low inter-item correlation. Alternatively, factor 3 might measure unrelated items that are not reflecting the overall concept of “Individual perception of tinnitus”. If this were true then it would not be contributing any additional value to the overall concept of tinnitus handicap (Cortina, 1993; Tavakol and Dennick, 2011).
Questionnaire Construct

Construct Validity: Convergent Validity

During the THQ development, the THQ displayed moderate correlations with average hearing thresholds (r = 0.52) and perceived tinnitus loudness (r = 0.57) (Kuk et al., 1990). Since then, the THQ has also been found to have strong correlations with the TQ (r = 0.75), the TRQ scores (r = 0.74) (Henry and Wilson, 1998) and the THI scores (r = 0.76; Robinson et al., 2003). Overall, the THQ demonstrates high convergent validity and therefore successfully measures aspects of tinnitus distress similar to other tinnitus questionnaires.

Construct Validity: Discriminant Validity

Low to moderate discriminant validity has been observed for the THQ. The THQ indicates moderate discriminant validity, displaying weak correlations with the Modified Somatic Perception Questionnaire scores (r = 0.37) the Private Self-Consciousness Scale (r = 0.11; Robinson et al., 2003), and moderate correlations with the life satisfaction scales (Lohmann, 1980) (r = 0.54), the physical health subscale (r = 0.54), the general health subscale (Duke-UNC Health profile; Parkerson et al., 1981) (r = 0.54; Kuk et al., 1990) and the Quality of Well-Being scale (r = 0.48; Robinson et al., 2003).

Nevertheless contradictory evidence indicates rather low discriminant validity. For example, the THQ has been shown to highly correlate with the Zung Self-Rating depression scale (Zung, 1965) (r = 0.63; Kuk et al., 1990) and the BDI (r = 0.62; Robinson et al., 2003). Robinson et al. (2003) also reported that the THQ moderately correlated with the Hamilton Rating Scale for Depression (r = 0.57).

Overall, the evidence suggests that the THQ has low to moderate discriminant validity and that it may be susceptible to generalised emotional distress, rather than distress that is specific to the condition of tinnitus.
Interpretation of the Scores

It is important to be able to reliably grade tinnitus severity if treatment efficacy or treatment needs are to be established. No grading system has been developed for the THQ measurement scale. Although a score of over 22/100 points (or 600/2700 points in raw scores) has been suggested as a lower boundary for bothersome tinnitus (Sullivan et al., 1993), there does not appear to be reliable empirical evidence to support this claim (see Table 5). Kuk et al. (1990) suggested that one possible method of determining the severity of symptoms for individuals is to compare a THQ score to their published normative data from 275 tinnitus patients. For example, a mean score of 60 would inform a clinician that the individual’s tinnitus was more severe than 80% of the tinnitus patients in this reference sample (Figure 3). Although this ranking is helpful in determining individual severity relative to others, it does not provide clinical interpretations of the scores (Newman and Sandridge, 2004; Kuk et al., 1990). For example, the mean score of 60 is not classified into a category, such as ‘moderate tinnitus’, that can inform the clinician of how severe the tinnitus is perceived and guide management.

Figure 3. Cumulative distribution of total scores on the Tinnitus Handicap Questionnaire (THQ). Data obtained from 275 patient responses (Kuk et al., 1990). Adapted from Kuk et al. (1990).
Sensitivity to Change

The THQ was primarily developed to effectively evaluate treatment-related change, with less emphasis placed on its ability to discriminate between diagnostic categories (Kuk et al., 1990). The THQ has high resolution response options (0-100), allowing for small changes in the scores to be detected. High test-retest reliability has been observed, with strong correlations between administrations for the total THQ scores ($r = 0.89$), factor 1 scores ($r = 0.89$) and factor 2 scores ($r = 0.90$) (Newman et al., 1995). Therefore, the global THQ score, and scores for factors 1 and 2 are stable across time. Changes can therefore probably be attributed to treatment-related effects. In contrast, factor 3 only showed moderate correlations ($r = 0.50$), suggesting that any changes shown over time could in fact be due to SEM rather than treatment-related effects.

Despite being specifically designed to maximise treatment responsiveness, Kuk et al. (1990) report no information on a minimal clinically important change score. The evidence for a minimal change score is dependent on the test-retest reliability data (Newman et al., 1995). From calculating the 95% confidence intervals (variance around the scores between administrations) with the SEM, Newman et al. (1995) conclude that a mean global THQ score would have to change by 21 points for the change to be considered clinically significant. However, this value maybe slightly compromised by lack of consistency and reproducibility shown by factor 3.

Summary

In conclusion, the development process of the THQ was informative and in general clearly reported, but there was a lack of clarity over the criterion scores for eliminating items. The THQ appears to cover some aspects of tinnitus distress but the emotional aspects seems to be the main contributor to global score. The usefulness and reliability of the three subscales is questionable. Factor 3 does not appear to add any further value to the questionnaire and is deemed to have a heterogeneous construct. Nevertheless, the THQ does show high convergent validity with other tinnitus questionnaires. There is no grading system to provide clinical meaning to the scores and a clinically meaningful change score has been suggested but it may be compromised by the limitations of factor 3.
Tinnitus Reaction Questionnaire (TRQ)

The 26-item Tinnitus Reaction Questionnaire was specifically designed to assess the psychological distress related to tinnitus, such as anger and depression (Wilson et al., 1991). Primarily developed to be an assessment of the effects of psychological interventions on tinnitus (treatment-related change), it is also claimed to distinguish between the levels of tinnitus-related distress. However, there is very little literature on the psychometric properties of the TRQ, with no apparent evidence for defining tinnitus distress values or a minimal clinically important change score. For each item, ratings are made on a 5-point scale with the anchors from: not at all (0) to almost all of the time (4). The total score ranges from 0 to 104, with a higher score denoting higher levels of distress (Table 3).

Initial Questionnaire Development

Critical evaluation of the development of the TRQ is difficult because of insufficient information provided in Wilson et al. (1991). For TRQ item choice, the authors briefly refer to the symptom categories reported by Tyler and Baker (1983; Figure 2) as the principal foundation, alongside the experience gained in previous work interviewing tinnitus patients on the efficacy of relaxation therapy (e.g., Ireland et al., 1985). However, it is not apparent exactly how this information was used to inform item development. It would appear that from the initial conception, the TRQ has only ever consisted of 26 items and so the number of items was never reduced like in the other cases. Although the authors did investigate response distributions, this was not used to inform the choice of items. No further information was provided on the item development and content.

Questionnaire Structure

Factorial Structure

The TRQ claims to measure aspects of psychological distress associated with tinnitus. It does not focus on the other domains of tinnitus distress discussed so far. The initial item-total correlations supported this, revealing
high correlations between the items (ranged from $r = 0.44$ to $0.81$) with the majority scoring $r > 0.70$, indicating that the TRQ measures a narrow range of tinnitus characteristics. PCA and orthogonal rotation revealed four factors related to psychological distress; (i) general distress (15 items), (ii) interference (9 items), (iii) severity (8 items), and (iv) avoidance (3 items) (Wilson et al., 1991). The analysis revealed that the majority of items loaded on more than one factor. For example, the majority of the items on the “interference” factor also load onto the “severity” factor. This indicates that the four factor solution is not representing distinctive subscales and that there is little value in looking at the factors individually.

Wilson et al. (1991) did also describe a two factor solution, where the majority of items loaded on factor 1, with six items loading on factor 2 (avoidance). The authors do not define factor 1. Again, there is evidence of cross loading, with the six items from factor 2 also loading on factor 1. However, Wilson et al. (1991) do not elaborate on the criterion value for factor loading and no loading scores were reported. Overall the evidence suggests a unidimensional structure to the TRQ. This interpretation is consistent with Kennedy et al.’s (2004) analysis of the individual items in the TRQ which found that >75% of items addressed the psychological and emotional effects (see Figure 1).

Wilson et al. (1991) recommend that the separate scores from the two factor solution, not the four factor solution, might be used to explain individual differences in tinnitus distress. Although this recommendation was made, it would appear not to have been followed by subsequent researchers. For example, the TRQ is often interpreted as having four valid subscales derived from the four factor solution (see Kennedy et al., 2004; Newman and Sandridge, 2004; Holgers et al., 2003).

**Internal Consistency**

The TRQ has also been shown to have excellent internal consistency for all 26 items in different populations ($\alpha = 0.96$) suggesting that all the items are measuring overlapping aspects of psychological distress (Wilson et al., 1991).
Questionnaire Construct

Construct Validity: Convergent Validity

Although there is limited evidence for convergent validity, we conclude that the TRQ appears to have high convergent validity. Robinson et al. (2003) reported that the TRQ highly correlates with the THQ (r = 0.78), the TQ (r = 0.82) and the THI (r = 0.88).

Construct Validity: Discriminant Validity

There is a large body of evidence which, taken overall, indicates that the TRQ has low discriminant validity and hence is sensitive to generalised emotional distress. Wilson et al. (1991) tested the TRQ in three different populations and found that in all three samples the TRQ strongly correlated with anxiety scales: Taylor Manifest anxiety scale (Bendig, 1956; r = 0.66) and Spielberger state-trait anxiety inventory (Spielberger et al., 1970; r = 0.58/60 to r = 0.71/74). Furthermore, that there were extremely strong correlations between the TRQ and the BDI (r = 0.63/87) (Wilson et al., 1991).

Strong correlations have also been replicated by Robinson et al. (2003) and Andersson et al. (2003). Robinson et al. (2003) reported that the TRQ strongly correlated with BDI (r = 0.66) and moderately correlated with the Hamilton Rating Scale for Depression (r = 0.52). Andersson et al. (2003) reported strong correlations with the Hospital Anxiety and Depression scales (r = 0.69 for depression and r = 0.72 for anxiety). Robinson et al. (2003) observed that the TRQ weakly correlates with the Private Self-Consciousness Scale (r = 0.20) and moderately correlates with the Quality of Well-Being scale (r = -0.39) and the Modified Somatic Perception Questionnaire (r = 0.53).

Interpretation of the Scores

The TRQ was developed to distinguish levels of tinnitus-related distress. Evidence for the TRQ as a measure of tinnitus severity is limited. Although, Wilson et al. (1991) claim that the TRQ is a useful screening questionnaire that can distinguish tinnitus sufferers, no grading system for categorising tinnitus severity has been explicitly developed.
Sensitivity to Change

Despite claims to effectively evaluate the effects of psychological interventions on tinnitus, Wilson et al. (1991) do not provide any evidence for this psychometric property. For example, no research has been conducted to find a minimal clinically important change score. Nevertheless the TRQ is consistently used as evidence for the efficacy of interventions.

Wilson et al. (1991) did provide evidence of a high test-retest reliability ($r = 0.88$) (over a retest period of 3 days to 3 weeks), suggesting that the TRQ is stable across time. However, without a minimal clinically important change score, the interpretation of any change in TRQ scores pre- and post-treatment is difficult. For example, there is no way of knowing whether a change in score of 2 points or 5 points is needed for the change to be considered meaningful.

Summary

In conclusion, the information available on the development and structure of the TRQ is unclear. The factorial structure and internal consistency seem to point towards a unidimensional structure focusing on the emotional impact of tinnitus, and the low discriminant validity indicates that this is generalised to the same psychological domains as depression and anxiety. Clinical interpretations of the TRQ scores are limited by the fact that no grading system was developed for quantifying tinnitus severity and no minimal clinically important change score has been recommended.

Tinnitus Handicap Inventory (THI)

The 25-item Tinnitus Handicap Inventory is reported to be a brief diagnostic and screening tool that measures the impact of tinnitus on everyday function (Newman et al., 1996). The THI consists of three response options; yes (4 points), sometimes (2 points) and no (0 points). The total score is rescaled from the weighted sum of all the items so that the global score ranges from 0 – 100, with a higher score denoting increased levels of tinnitus distress (Table 3). The THI was originally designed as a companion to the Hearing Handicap Inventory for Elderly (HHIE, Ventry and Weinstein, 1982) and the
Dizziness Handicap Inventory (DHI, Jacobson and Newman, 1990) to complete a set of tools to quantify perceived handicap in a variety of hearing related diseases, such as Ménière’s disease (Newman and Sandridge, 2004).

**Initial Questionnaire Development**

Again, limited information makes it difficult to ascertain the exact process behind the questionnaire development (Newman et al., 1996). Clinical expertise certainly played a role. For example, Newman stated: “The alpha-THI consisted of 45-items derived empirically from case histories of patients with tinnitus” (Newman et al., 1996, p. 144). Although, the authors infer that a combination of content and face validity were used to source and develop the items, the information provided is unclear. Some items were adapted from the HHIE and the DHI, and also from Tyler and Baker (1983; see Fig 2). The THI has adopted the identical response options and scoring as the DHI and HHIE. No further information was provided on the item development and content.

The 45 items were reduced to 25 items (final version) using response frequency distributions, item-total correlations, and content validity. Items were eliminated if high endorsement rates for one response option were found, or if item-total correlations were $r \leq 0.50$. Newman et al. (1996) believed these items were insensitive, discriminated little between subjects, and did not represent the scale concept. There is little clarity on who conducted the content validity, which items were removed and how the criteria were decided for the frequency distribution elimination score (i.e., 85% patients selecting same response) and for the item-total correlation scores ($r \leq 0.50$).

**Questionnaire Structure**

**Factorial Structure**

Following the examination of the item content, a three-domain model for the THI was proposed (Newman et al., 1996) to cover limitations in (i) functioning (12 items: mental, social and physical), (ii) emotional response to tinnitus (8 items), and (iii) the desperation associated with tinnitus (5 items: catastrophic). However, the domain content (subscales) were solely based on
content validity and the questionnaire structure was not subjected to empirical validation at the time.

In 2003, Baguley and Andersson investigated the theoretical factorial structure and three domains originally proposed by Newman et al. (1996). The THI scores were predefined into the three factor solution and subjected to PCA. Initial item-total correlations were rather high ($r = 0.60$) for a scale which assumes to be measuring a broad construct. This result suggests that the THI is tapping only into narrow characteristics (Clark and Watson, 1995). In contrast to the structure previously proposed, oblique rotation revealed that the majority of items loaded onto the one factor reflecting functioning (19 items). Factor 2 (emotional) and factor 3 (catastrophic) each consisted of 5 items, some of which also loaded onto factor 1. The high overlap between factors suggests that they are not distinctive from each other and are in fact measuring the same latent variables (Zachariae et al., 2000). This unidimensional structure is further evident in the THI Danish (Zachariae et al., 2000) and Italian translations (Monzani et al., 2008), and in Kennedy et al.'s (2004) analysis of tinnitus questionnaires where the THI was shown to favour psychological/emotional distress (68%; Figure 1). Therefore, despite Newman’s initial claim of a three-factor structure, the THI appears to be rather unidimensional.

Internal Consistency

During the initial development process, the only analysis conducted was Cronbach’s alpha (Newman et al., 1996). This measure alone does not provide a sufficient measure of the internal structure and its reliability. Validation of the factorial structure is required too (Streiner, 2003; Cortina, 1993). Excellent Cronbach’s alpha scores were reported for the THI 25-items ($\alpha = 0.93$) suggesting that all the items measure the same underlying construct. High scores such as this indicate redundancy of items and could be indicative of the unidimensional structure, although internal consistency is not sufficient evidence for homogeneity (Green et al., 1977; Schmitt, 1996). Both the functional and emotional subscales showed high internal consistency ($\alpha = 0.86$ and $\alpha = 0.87$ respectively). The internal consistency for the catastrophic subscale was questionable ($\alpha = 0.68$), potentially due to the small number of items in this scale. Coefficient alpha values are vulnerable to the number of test items (Schmitt, 1996). Recent evidence suggests that the questionable Cronbach’s alpha score for catastrophic subscale is more likely to indicate a
heterogeneous construct with poor inter-relatedness between the items (Tavakol and Dennick, 2011).

**Questionnaire Construct**

**Construct Validity: Convergent Validity**

Overall, the THI has high convergent validity demonstrating that it measures a construct that is comparable to other tinnitus questionnaires. The THI shows strong correlations with the THQ (r = 0.78), the symptom rating scales (r = 0.67 - 0.72) (Newman et al., 1996), the TQ (Spearman’s r = 0.89) (Baguley et al., 2000) and the TRQ (r = 0.89) (Robinson et al., 2003).

**Construct Validity: Discriminant Validity**

Overall, the THI measures a construct that is independent of generalised emotional distress. Moderate discriminant validity for the THI has been observed by the moderate correlations with the BDI (r = 0.32 - 0.58), the Hamilton Rating Scale for Depression (r = 0.49) and the Quality of Well-Being scale (r = -0.37) (Newman et al., 1996; Robinson et al., 2003). Weak to moderate correlation with the Modified Somatic Perception Questionnaire scores (r = 0.24 - 0.38) and the Private Self-Consciousness Scale (r = 0.23) (Robinson et al., 2003).

**Interpretation of the Scores**

The THI was developed to grade tinnitus severity at intake assessment. Developed by Newman et al. (1998) using the quartiles analysis on test-retest reliability data, the grading system originally defined four categories (Table 5). These categories were further developed by a UK working group (McCombe et al., 2001). The authors recommended adopting a five category grading system as a way to diagnose and screen tinnitus severity in clinical practice (Table 5). This recommendation is based on the expert opinions and knowledge. For example, “[…] take the known epidemiological and clinical data and with our own clinical, medico-legal and research experience to
produce a severity scale” (McCombe et al., 2001, p.26). No further evidence has been reported for the development of these categories other than a reference that the scores were based on Newman et al.’s 1998 analysis. In turn, no empirical evidence has been provided on the validity of the definitions. This prompts questions about the reliability of these categories to provide clinicians with valid meanings behind the scores.

**Sensitivity to Change**

The THI was not developed for use as an outcome measurement tool and does not explicitly maximise the responsiveness to treatment-related effects (Meikle et al., 2007). Nevertheless, it is used internationally as an outcome measure for testing the effectiveness of therapeutic interventions and for clinical research. Newman et al. (1998) conducted test-retest reliability to assess stability over short time intervals. The THI showed high test-retest reliability ($r = 0.92$) suggesting that a mean score would unlikely deviate more than 7.0 points (based on SEM) between tests (Newman et al., 1998).

The variance around these scores (test and retest) produced the 95% confidence intervals. From examining the 95% confidence interval and the SEM, Newman et al. (1998) concluded that a reduction of ≥20 points is required for the change to be classified as clinically meaningful (Newman et al., 1998). However, this reduction is dependent on the intake assessment score being >20 points, therefore potentially failing to observe the small changes that occur in patients with slight or mild tinnitus distress (Table 5).

In 2011, Zeman et al. investigated the minimal clinically important change required to show a noticeable improvement in the THI scores. They used an anchor-based technique; CGI-I rating with seven response options (Table 6a). Again, patient groups were formed according to the CGI-I scores (four groups: much better, minimally better, no change, worse). The effect size (Cohen $d$) separating the minimally better and no change groups ($d = 0.5$) was used alongside the estimated standard deviation of the before-after difference ($SD = 14$) to calculate the minimum significant change for improvement ($\Delta THI = 0.7$). Therefore, patients perceived a THI score reduction of 7 points as a meaningful improvement. This offers a more sensitive measure of responsiveness in the THI compared to the previous scores suggested by Newman et al. (1998), as it takes into account patient experience.
Despite these attempts to produce a criterion for a significant change score, the THI will always receive criticism for its inherent lack of sensitivity in the three response options.

**Summary**

In conclusion, the THI may not be as robust as first proposed. There is little clarity provided on the item development and it has a unidimensional structure which emphasises the emotional aspects of tinnitus distress. A grading system for quantifying tinnitus severity has been developed for the THI and does provide useful clinical interpretation. Finally, there have been attempts to provide a minimal clinically important change score.

**Tinnitus Functional Index (TFI)**

The 25-item TFI was a large international collaboration of 21 investigators (including 17 expert judges). It was specifically developed to be (i) discriminative to provide measures of tinnitus distress, (ii) evaluative to provide a responsive measure of treatment-related changes, and (iii) comprehensive to cover multiple domains of tinnitus severity. For each item, individuals are asked to rate their judgement on a scale of 0-10. Each item scale has descriptive anchors at either end that vary depending on the item content (Figure 4). The total score is rescaled from the weighted sum of the...
items in each subscale so that the global score ranges from 0 to 100, with a higher score denoting higher levels of distress (Table 3).

**Initial Questionnaire Development**

The report of the development is extremely detailed. Meikle et al. (2012) reports the development of the TFI through prototype 1 and 2 to the final 25-item questionnaire. Initially, 175 items were sourced from nine widely used tinnitus questionnaires excluding ambiguous items (i.e., referring to multiple subtopics) and overly negative items. The 175 items were assigned to 13 different domains of tinnitus handicap, on the consensus of all judges. Duplicate items were reduced to one. Judges completed a three-point responsiveness rating scale which also provided the basis for item selection. Prototypes 1 and 2 were administered on four occasions at five or four sites respectively. Both prototypes were completed by over 300 participants (the first occasion). However it should be noted that there was a high dropout rate after this first administration. The majority of the participants were recruited from Veterans’ Affairs (VA) hospitals. This is potentially problematic because VA patients are not representative of the wider clinical population. The majority of VA patients tend to be male and to experience severe tinnitus with co-morbidities.

Forty-three items were used to create prototype 1 with a 0 – 10 response scale. Thirteen items were removed after examination of response frequency distributions, effect size and exploratory factor analysis. Eight items were removed because the effect size for individuals whose tinnitus was unchanged or worse (based on global question) was unacceptably large (i.e., 0.89) and therefore considered unresponsive to perceived change. A variety of the factorial structure techniques were used on prototype 1 data to inform item choice and questionnaire structure. PCA followed by Principal Axis Factoring and oblique rotations revealed eight factors (8 of the proposed 13 domains were identified). A predefined ‘somatic’ domain was removed due to low factor loading for two items, one of which also showed floor effects (below the desired lower limit of 2.0). The remaining three items were removed due to low factor loadings or initial mean score. The criterion for low scores was not reported. Thirty items remained after this analysis to create prototype 2. Although the same examinations were conducted on prototype 2, none of these findings appear to contribute to the decision to remove the final five items.
This decision was based on “careful examination” of the items (content validity). The final version was a 25-item TFI with eight subscales (Table 3).

**Questionnaire Structure**

During the development of the TFI, reliability and validity testing (i.e., factor analysis, effect size and convergent/discriminant validity) were conducted on prototypes 1 and 2. The final 25-item TFI has not yet been subjected to a formal statistical validation. So far, evidence for its validation comes from a re-analysis of the relevant subset of data collected using the 30-item prototype 2. In this section, all the evidence provided is based on this re-analysis.

**Factorial Structure**

The TFI covers eight different domains of tinnitus distress. The 30-item prototype was subjected to PCA followed by Principal Axis Factoring, with orthogonal and oblique rotations informing both model’s factor extraction. The eight factor solution revealed in prototype 1 was also clearly revealed for the 30-item prototype 2, despite the reduction in items.

To validate the factorial structure, the 25-item prototype 2 data was again subjected to Principal Axis Factoring, with oblique rotations. The same eight factors were clearly identified; (i) cognition (3 items), (ii) auditory (3 items), (iii) intrusiveness (3 items), (iv) sleep (3 items), (v) relaxation (3 items), (vi) QoL (4 items), (vii) emotional (3 items), and (viii) sense of control (3 items). Only one item (TFI Q21) is shown to load on more than one factor (i.e., Cognitive and Quality of Life). The remaining 24 items load onto a single factor, demonstrated by the high loading values.

**Internal Consistency**

The 25-item prototype 2 was shown to have an excellent Cronbach’s alpha score for all 25 items ($\alpha = 0.97$) and for the eight subscales (range from $\alpha = 0.82 - 0.97$). This score suggests that all the items in the questionnaire are measuring a similar construct. The reliable factorial structure indicates that the
items are measuring different aspects of tinnitus distress, rather than the presence of redundant items.

**Questionnaire Construct**

**Construct Validity: Convergent Validity**

There is limited evidence for convergent validity for the TFI. Presently, there is only the evidence from the 25-item prototype 2 data. Meikle et al. (2012) reported extremely strong correlations between the 25-item prototype 2 and the THI ($r = 0.86$), i.e., high convergent validity. Further evidence is needed.

**Construct Validity: Discriminant Validity**

Again, there is limited evidence. The 25-item prototype 2 moderately correlates with the BDI-primary care ($r = 0.56$), demonstrating moderate discriminant validity with an independent measure of emotional distress.

**Interpretation of the Scores**

Meikle et al. (2012) did propose a grading system based on preliminary data analysis using the 25-item prototype 2 data (Table 5). Throughout development, the five categories (not a problem to a very big problem) from the global severity question (i.e., how much of a problem is your tinnitus?) have been compared to the mean scores from prototype 1 and 2. A strong association between these scores has been shown throughout ($p < 0.001$). Following these comparisons, the mean 25-item prototype 2 scores were categorised based on the individual responses to the global severity question. The response distributions of the total scores in these categories were examined to produce the grading system (Table 5).
Sensitivity to Change

The TFI was developed to effectively evaluate treatment-related change and this was explicitly considered at all steps of its design (Meikle et al., 2012). The items were judged on responsiveness by a panel of judges. The 0–10 response options allow for small changes in the scores to be detected.

The 25-item prototype 2 data demonstrated high test-retest reliability for the overall 25-item scores ($r = 0.76$) (retest period of 7 to 30 days). For the eight subscales, high test-retest reliability has been observed with the scores ranging from 0.66 to 0.90 (Meikle et al., 2012). Therefore, the 25-item prototype 2 and the subscales show stability over short time intervals.

Based on 25-item prototype 2 data, Meikle et al. (2012) did propose a minimal clinically important change score. The mean scores were categorised into five groups based on global perception of change scores at 3 and 6 months (Table 6c). Having examined the data in the plot for differences in scores between the much-to-moderately changed group and the unchanged at 3 and 6 months, Meikle et al. proposed a reduction of around 13 points to be meaningful to patients (Figure 5).

Figure 5. Overall mean TFI change scores at 3 and 6 month follow-ups after grouping patients based on Global Perception of Change score. A minimal clinically important change score of -13 points was determined based on the difference between the unchanged and much-to-moderately improved groups at 3 and 6 months. Adapted from Meikle et al. (2012).
Summary

The development process of the TFI is clearly described and the authors provide clear, in-depth information on all of the steps that were followed. The supplemental published information provides useful evidence from the different prototypes. So far the evidence would suggest that the TFI covers eight different domains of tinnitus distress, is comparable to at least one other tinnitus questionnaire and is measuring more than just generalised emotional distress. The authors have provided both a grading system and a minimal clinically important change score, therefore providing clinical interpretations to the TFI scores. However, this evidence is all based on preliminary data analysis with a rather limited sample of participants. Despite the rigorous validation conducted during the development of the TFI, the final TFI has not been subjected to formal validation.

Conclusion

Our review seeks to promote quality assurance in validation research concerning tinnitus questionnaires. Although beyond the scope of this book chapter, we recommend that the ‘gold standard’ would be to carry out a systematic review of the literature before selecting any given tinnitus questionnaire for a service audit or clinical trial. This would provide a comprehensive and critical overview of the existing questionnaires available. In addition to the measurement properties, selection might also give consideration to the suitability of the tinnitus questionnaire for the study population, the potential burden of completing the tinnitus questionnaire (e.g., length, question difficulty, emotional impact of certain questions), and the practical aspects (e.g., copyright costs, complexity of scoring method). We make the assumption that the questionnaire validation is conducted on a sample of people with tinnitus that is representative of the population in which the instrument is to be used.

Explicit quality criteria for studies on the development and evaluation of tinnitus questionnaires help to legitimise recommendations about what is the ‘best’ questionnaire for tinnitus handicap. A wide range of resources are available to help the researcher develop questionnaire instruments and evaluate existing tools. We draw the reader’s attention to one useful web resource that provides practical guidance on how to select, translate and
validate questionnaires (http://www.emgo.nl/kc/about.html), with an electronic quality assurance handbook available in English (see also Terwee et al., 2007). Following these guidelines, in Table 7, we summarise the evidence reviewed and reported in this chapter according to ten quality criteria for good measurement properties, arranged under the four key categories (see also Uijen et al., 2012). According to these criteria, the TFI comes out “best”, meeting eight of the ten criteria according to the current published development reports. However, one caveat to this recommendation is that the final 25-item version of the TFI has not been subjected to these formal validation criteria. This should be the subject of future research.

Table 7. Summary of the critical evaluation of the psychometric properties of the five tinnitus questionnaires

<table>
<thead>
<tr>
<th>Questionnaire</th>
<th>Content validity</th>
<th>Internal consistency</th>
<th>Structural validity</th>
<th>Construct validity</th>
<th>Reproducibility</th>
<th>Responsiveness</th>
<th>Floor or ceiling effects</th>
<th>Interpretablity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tinnitus Questionnaire</td>
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<td>0</td>
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<tr>
<td>Tinnitus Handicap Questionnaire</td>
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<tr>
<td>Tinnitus Reaction Questionnaire</td>
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<tr>
<td>Tinnitus Handicap Inventory</td>
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Rating: + = positive, 0 = no information available, − = poor, ? = indeterminate rating (doubtful design or method), where two symbols are given this indicates that several different validation criteria were used.
References


