Audiologist-Guided Internet-Based Cognitive Behavior Therapy for Adults With Tinnitus in the United Kingdom: A Randomized Controlled Trial

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INTRODUCTION

Most health care in the United Kingdom is provided by the publicly funded National Health Service (NHS) and is largely free at the point of use. General practitioners (GPs) provide primary health care and refer patients to specialist services as required. Recently the NHS has experienced challenges due to funding constraints together with an ever-growing demand for services (Smith et al. 2014). This has led to an increase in appointment waiting times, which has been associated with poorer outcomes for a variety of health issues (e.g., Pizer & Prentice, 2011; Smith et al. 2014). For patients experiencing significant levels of health-related distress, such as those with chronic tinnitus, minimizing waiting times should be prioritized (Gander et al. 2011).

Tinnitus is defined as the sensation of sound in the absence of a corresponding external acoustic stimulus (Baguley et al. 2013). It may be perceived on a spectrum from barely noticeable to debilitating (Brüggemann et al. 2016). Experiencing tinnitus is often associated with a wide range of associated symptoms such as sleep disturbance, concentration difficulties, irritation, frustration, anxiety, and depression (Langguth 2011). In England, there are an estimated ¾ of a million people per year who visit their GP with tinnitus as the primary complaint (El-Shunnar et al. 2011). Of these, only 37% are referred for specialist services (El-Shunnar et al. 2011). In addition, those referred often have a substantial wait of up to 18 weeks before an intervention pathway, such as obtaining tinnitus counseling, commences (Department of Health, 2009). A further constraint in tinnitus management in the United Kingdom is that the intervention with the most evidence of efficacy, namely cognitive behavioral therapy (CBT, see Hesser et al. 2011) is not readily available for those with tinnitus. This is largely due to a shortage of trained specialists (Baguley et al. 2013a). Moreover, specialist tinnitus services are not offered in all NHS hospitals across the United Kingdom, leaving many with distressing tinnitus without any specialized intervention options (Hoare et al. 2015).

The need for widely available, cost and clinically effective, tinnitus management is evident worldwide, and not isolated to the United Kingdom (Andersson, 2016). To increase access to
effective tinnitus intervention in Sweden, cognitive behavioral therapy is provided via the Internet (iCBT; Andersson, 2015). As iCBT has been found to be effective at reducing tinnitus and associated problems in clinical trials in Sweden and Germany (e.g., Andersson et al. 2002; Kaldo et al. 2008; Hesser et al. 2012; Nyenhuis et al. 2013; Jasper et al. 2014; Weise et al. 2016), it has been incorporated into regular clinical care in Sweden (Kaldo-Sandström et al. 2004; Kaldo et al. 2013). As iCBT could increase access to an evidence-based intervention in the United Kingdom, a comprehensive, user-friendly, intervention tailored for this population was designed by Beukes et al. (2016). Feasibility of iCBT in the United Kingdom was established in terms of recruitment, attrition, and compliance (Beukes et al. 2017a). The clinical efficacy of this redeveloped iCBT intervention in the United Kingdom has not yet been established. In this context, delivering iCBT guided by an audiologist would be optimal, but the efficacy of iCBT by a nonpsychological professional is unproven. This trial set out to explore the use of iCBT in the United Kingdom with the following objectives:

1. To evaluate the efficacy of audiology-guided iCBT in reducing tinnitus distress compared with weekly monitoring.
2. To ascertain the efficacy of iCBT for comorbidities associated with tinnitus.
3. To assess the stability of iCBT intervention effects 2-months postintervention.
4. To establish the on-going intervention effects during the course of iCBT.

The hypothesis was that iCBT for tinnitus would be more effective at reducing tinnitus distress and the associated comorbidities than weekly monitoring.

MATERIALS AND METHODS

Study Design

A delayed intervention efficacy trial with a 2-month follow-up was implemented to evaluate the efficacy of iCBT in the United Kingdom. This prospective, two-arm, randomized control trial was registered with Clinical Trials.gov: NCT02370810 on May 03, 2015. The Experimental Group received the iCBT intervention for 8 weeks, while the Control Group were monitored weekly. Once the experimental group completed the intervention, the control group underwent the same iCBT intervention. This study design, therefore, provided the opportunity to evaluate the intervention effects in two independent groups at two time points. Although the control group had a delay of 8 weeks before undertaking this intervention, this may be less than the 18-weeks wait they may have on standard treatment pathways on the NHS.

The Consolidated Standards of Reporting Trials eHealth guidelines (Eysenbach et al. 2011) were implemented to report the methods and results of this trial. For the full study protocol, see Beukes et al. (2015).

Ethical Considerations

Ethical approval was granted by the Faculty Research Ethics Panel of Anglia Ruskin University (FST/FREP/14/478). The trial was conducted in accordance with good clinical practice together with the ethical principles of the Declaration of Helsinki. A protocol was established to ensure the security of participants’ confidentiality when using the Web portal, complying with the following U.K. legislation: The Data Protection Act of 1998 and The Privacy and Electronic Communications (EC Directive) Regulations (Riach, 2003). There were no changes to the methods or assessment measures used after the trial commenced. No harms or unintended effects were reported.

Study Population

Recruitment • Recruitment was throughout the United Kingdom for a period of 2 months and targeted people from various demographical backgrounds with significant levels of tinnitus distress. Study information was available in various formats including online (e.g., the NHS Choices and clinicaltrials.gov Web sites), Twitter (British Tinnitus Association), Facebook forums (e.g., Action on Hearing loss, Thyroid UK), newspapers, and magazines (e.g., Mature Times, People’s Friend, Musicians Union bulletin, New Scientist, National Federation of Occupational Pensioners Magazine, Cambridge News), support groups (e.g., tinnitus, thyroid) and from health care professionals (GP surgeries, audiologists).

Participants • Those interested in the study registered interest on the study website (www.tacklingtinnitus.co.uk). They were informed of their right to withdraw at any stage without penalty. Eligibility for the study was determined in a two-stage process. Initially, participants completed the baseline assessment measurements online. Following completion, a telephonic screening was arranged, to ensure participants fulfilled the study requirements, which were as follows:

Inclusion Criteria

i) Aged 18 years and over living in the United Kingdom.
ii) Computer and Internet access and the ability to use these.
iii) The ability to read and type in English.
iv) Experiencing tinnitus for a minimum duration of 3 months.
v) A score of 25 or above on the Tinnitus Functional Index suggesting the need for tinnitus care (Meikle et al. 2012).

Exclusion Criteria

i) Reporting any major medical, psychiatric, or mental disorder, which may hamper commitment to the program.
ii) Reporting pulsatile, objective, or unilateral tinnitus, which have not been investigated medically.
iii) Tinnitus as a consequence of a medical disorder, still under investigation.
iv) Undergoing any tinnitus therapy concurrently with participating in this study.

Assessment Measures

A demographic questionnaire was used to obtain information related to gender, age, tinnitus duration, hearing aid use, medical examinations related to tinnitus, past or current tinnitus treatments, health and/or mental health conditions, and employment. Self-reported assessment measures were selected to establish tinnitus distress and identify associated difficulties, as these are generally used in clinical practice. The following assessment measures were completed at baseline (T₀), at post-intervention (T₁) and follow-up (T₂) in both groups.
Primary Assessment Measure • The Tinnitus Functional Index (TFI; Meikle et al. 2012) was selected as the primary assessment measure to quantify tinnitus distress. It was chosen above some other established tinnitus questionnaires, such as the Tinnitus Handicap Inventory (THI; Newman et al. 1996) because of its validation for assessing intervention responsiveness. It is a 25-item questionnaire, scored on a scale of 0 to 100. Scores less than 25 indicate mild tinnitus, with no need for intervention, whereas scores ranging from 25 to 50 signify significant tinnitus and the possible need for intervention. A score of 50 or greater demonstrates more severe tinnitus and indicates the need for more intensive intervention. A reduction in TFI scores shows improvement in tinnitus distress. Meikle et al. (2012) reported that meaningful changes occur when scores are reduced by 13 points or more. Due to regression to the mean artefacts, those with more severe scores are more likely to show significant changes on assessment measures than those reporting mild symptoms (Campbell & Kenny, 1999). The TFI has excellent psychometric properties with an internal consistency of 0.97 and test-retest reliability of 0.8 (Meikle et al. 2012).

Secondary Assessment Measures • The following secondary assessment measures were selected to identify difficulties that may be related to having tinnitus:

i) The Insomnia Severity Index (ISI; Bastien et al. 2001) was used to assess the presence of insomnia, as sleep difficulties are prevalent among those with tinnitus (Crönlein et al. 2016). The seven-item questionnaire is scored between 0 and 28 and has an acceptable internal consistency of 0.7 (Bastien et al. 2001).

ii) The Generalised Anxiety Disorder (GAD-7; Spitzer et al. 2006) was selected to quantify the level of anxiety, as the prevalence of anxiety is high in those with severe tinnitus (Pinto et al. 2014). This seven-item questionnaire is scored between 0 and 21 and has an internal validity of 0.9 (Löwe et al. 2006).

iii) The Patient Health Questionnaire (PHQ-9; Spitzer et al. 1999) was chosen to assess symptoms of depression, as depression among those with severe tinnitus is often reported (Pinto et al. 2014). Scoring is between 0 and 28 on this nine-item questionnaire with an internal validity of 0.8 (Spitzer et al. 1999).

iv) The Hearing Handicap Inventory for Adults Screening version (HHIA-S; Newman et al. 1991) was administered to assess difficulty hearing, which in this context may be related to the penetrating nature of tinnitus or the presence of hearing loss, commonly found in those with tinnitus (Langguth et al. 2017). This measure consists of 10 items, scored between 0 and 40 and has a good internal consistency of 0.9 (Newman et al. 1991).

v) The Hyperacusis Questionnaire (HQ; Khalfa et al. 2002) was administered to assess the presence of reduced tolerance of everyday sounds, otherwise known as hyperacusis, as there is a large overlap in the prevalence of tinnitus and hyperacusis (Schecklmann et al. 2014). This 14-item questionnaire is scored between 0 and 42. Fackrell et al. (2015) evaluated the psychometric properties of the HQ in a large population of participants with tinnitus in the United Kingdom and found a high internal consistency of 0.9 but were unable to confirm the original three factor solution proposed by Khalfa et al. (2002) and therefore suggested cautious use of the HQ until an alternative has been developed. To date, a questionnaire has yet to be developed so the HQ was used as a measure of sound sensitivity.

vi) The Cognitive Failures Questionnaire (CFQ; Broadbent et al. 1982) was administered to assess cognitive functions, as tinnitus may impact the control of attention leading to cognitive slips and errors in task completion (Tegg-Quinn et al. 2016). This 25-item questionnaire is scored between 0 and 100 and has a good internal consistency of 0.9 (Broadbent et al. 1982).

vii) The Satisfaction with Life Scales (SWLS; Diener et al. 1985) was administered as a quality of life measure assessing global life satisfaction as opposed to quality of life measures often related to self-care and mobility. Scoring is between 0 and 35 for five-items and has an internal consistency of 0.9 (Diener et al. 1985).

A low score signifies fewer problems than a high score and a reduction in score indicates improvement for all these measures except for the SWLS. For the SWLS, a higher score shows more life satisfaction than a lower score and an increase in score reveals improved life satisfaction.

Weekly Assessment Measure • The Tinnitus Handicap Inventory Screening version (THI-S; Newman et al. 2008) was selected to monitor tinnitus severity in both groups on a weekly basis during the 8-week period between T₀ and T₁. This measure was selected instead of the TFI or THI due to its concise nature as it consists of only 10 questions. The scores obtained are comparable (r = 0.9) with the full version of the THI (Newman et al. 2008), and good convergent validity (0.9) has been found between the TFI and THI (Meikle et al. 2012).

Data Collection

Data collection of the assessment measures was online throughout the trial for both groups. Results using an online format should be comparable with those using a paper presentation, as equivalent psychometric properties have been reported (Thorén et al. 2012). To minimize attrition postintervention, reminder e-mails were sent to encourage participants to complete the assessment measures. Assessment measures were used with permission of the copyright holders, and agreements were established for those that are not freely available to use, such as the TFI and ISI.

Study Intervention

The intervention was based on a self-help program originally developed by Andersson and Kaldo (2004). This content was redeveloped into an interactive e-learning version, to ensure it was visually stimulating and engaging (Beukes et al. 2016). The Web-based intervention platform used was designed in-house at Linköping University, Sweden and complied with a high level of data security and encrypted communications (Vlaescu et al. 2016). This 14-item questionnaire is scored between 0 and 42. The intervention ran over an 8-week period, during which 2–3 modules were released on a weekly basis. CBT principles such as goal setting, a clear structure, active participation, relapse prevention, and setting a time-frame for completing the intervention were incorporated (Andersson,
2002). There were 16 recommended modules and five optional modules. Recommended modules included CBT content such as applied relaxation, thought analysis, cognitive restructuring, imagery, and exposure techniques. Optional modules were available to add an element of tailoring, and participants could choose whether or not to do these modules. If initial baseline scores for the ISI indicated at least subthreshold insomnia (≥ 8) undertaking the optional sleep module was recommended. If the HHIA indicated a 50% probability of hearing disability (≥ 26), the hearing tactics module was suggested, and if scores were ≥ 30 on the CFQ, the module covering concentration guidelines was advised. The sound sensitivity module was recommended if scores were ≥ 28 on the HQ.

Guidance During the Intervention

Internet interventions are either independent of professional support (unguided) or offer some form of support (guided). A key element of this intervention was that it was guided, as better outcomes are reported for guided interventions (Baumeister et al. 2014). To maintain consistency with the standard approach of tinnitus interventions being delivered within the audiology community in the United Kingdom, an experienced audiological scientist guided the intervention. The audiologist was registered with the Health and Care Professions Council and appropriately trained to Masters Level in Audiology. The audiologist was experienced in managing tinnitus patients both in a clinical setting and online and had a suitable understanding of CBT principles but no formal CBT training. Supervision was provided by a clinical psychologist (specialized in providing tinnitus intervention) throughout the duration of the trial. Having audiological support for an iCBT intervention is unique to this study, as psychologists have guided participants in previous trials. The audiologist’s role was to conduct the telephone interviews, introduce weekly modules, provide feedback, answer queries, provide guidance, support, and encourage engagement. A secure encrypted messaging system was available to enable this two-way communication. Communication included feedback on progress, encouragement, and information about the content of new modules. A minimum of 10 minutes per week was spent on each participant and more time if required.

Data Analysis

Sample Size • Sample size estimation was calculated using G*Power version 3.1.6 (Faul et al. 2007) and based on achieving a clinically relevant change between baseline and postintervention using the primary assessment measure, the TFI. Calculations using the 13-point difference suggested during the development of the TFI indicated that 58 participants were required per group, with an allocation ratio of 1:1, to achieve a two-sided significance level of 0.05, with an effect size of 0.5 and 80% power. An additional 30 participants were recruited to ensure sufficient power during per-protocol analysis to account for possible dropouts. Therefore, 73 participants were recruited to each arm.

Enrolment and Randomization • Participants meeting the inclusion criteria were randomly assigned in the ratio of 1:1 and enrolled to either the experimental or control group. Allocation was based on a randomization sequence generated by computer algorithm (http://www.randomizer.org/) and done by an independent researcher. To prevent an unequal distribution among groups, participants were prestratified on the factors of age (≤ 60 or > 60 years) and tinnitus severity (TFI ≤ 50 or > 50). Block randomization, with blocks of four, were applied to ensure equal groups sizes within each stratum. Participants were informed when the intervention would commence by the principal investigator, but not which group they had been assigned to. The trial design resulted in the investigator not being masked to the assignment of interventions during the running of the trial. During the initial telephone screening, it was explained that the trial would start once registration was full and all participants were telephoned and randomized. Participants, therefore, expected a delay to the trial onset as no time-period was given. Participants may have realized their group assignment, but this was never explicitly stated.

Statistical Analysis • The Statistical Package for Social Sciences (SPSS) version 23.0 was used for statistical analysis, and the data analyst was masked to the groups to minimize bias. For all analyses, a two-tailed significance level of < 0.05 was considered statistically significant.

Missing Data Analysis • An intention-to-treat paradigm was used, as this analysis is less susceptible to bias than complete case analysis techniques. Missing value analysis was conducted to determine how to account for missing data. Little’s missing completely at random test (Little, 1988) indicated that data were likely to be missing completely at random (χ2 (55) = 42.4; p = 0.89). This suggested that missing values were probably randomly distributed across all observations and that there was no systematic pattern to the missing data. Missing data could thus be imputed through the multiple imputation procedure offered by SPSS using Markov Chain Monte Carlo method, which uses five imputation runs (Asendorpf et al. 2014). All preintervention assessment measure results were used as predictors. These results were compared with those obtained with a per-protocol analysis. As there was no difference, the intention-to-treat results are reported. Results obtained by averaging the five imputation runs (pooled results) were used where available. For some of the statistics, a pooling algorithm is not yet available. When this was the case, the first imputed set of results was reported.

Study Outcomes • The primary study outcome was a change in TFI score between the groups at postintervention (T1). Secondary study outcomes were changes in the scores of secondary assessment measures between groups at T1. A difference in scores between T1–T2 for the experimental group was used to assess the stability of intervention effects.

Group Differences and Stability of Intervention Effects • A mixed 2 × 3 analysis of variance for repeated measures with the within-subject variable of time (T0, T1, T2) and between-subject factor of group (experimental and control) was carried out to compare assessment measure results across the three-time points. Greenhouse-Geisser correction for nonsphericity was applied.

The main effects were followed up by paired samples t-tests to compare within-group differences at individual time points and independent sampled t-tests to compare results between the two groups at each time point.

Effect Sizes • Effect sizes and the 95% confidence intervals at postintervention were calculated by dividing the mean differences by the pooled SDs. Effect sizes below d = 0.5 represented small effect sizes; those of d = 0.5–0.79 medium effect sizes and those equal or greater than d = 0.8, large effect sizes (Cohen, 1992).
Monitoring Intervention Effects Between T₀ – T₁ • A mixed 2 × 8 analysis of variance for repeated measures was used to compare the results of the weekly THI-S scores with the within-subject variable of time (weeks 1–8) and between-subject factor of group (experimental and control). The main effects were followed up by paired samples t-tests to compare within-group differences at individual time points and independent sampled t-tests to compare results between the two groups at each time point.

Clinically Significant Change • A statistical significance of differences in group means is the standard analysis of clinical trials. Supplementing these results with an evaluation to determine whether the change in score is clinically meaningful is an indicator of the value of the intervention. The reliable change index (RCI; Jacobson & Truax, 1991) was used to determine clinical significance. It was calculated using the SD and means at T₀, the means at T₁, and the test-retest reliability coefficient or Chronbach’s alpha where this was not available.

Individual’s mean difference scores for those completing the intervention (i.e., both groups) between T₀ and T₁ were evaluated against the RCI criterion. Individual’s mean difference scores for those completing the intervention from the control group between T₀ and T₂ were also evaluated against the RCI criterion for the TFI.

RESULTS

Participant Characteristics
The baseline assessment measures were completed by 169 of the 244 adults on the trial waiting list. A total of 146 adults met the eligibility criteria and were randomly assigned to the experimental (n = 73) and control groups (n = 73) as shown in the Consolidated Standards of Reporting Trials diagram (Fig. 1). The mean age was 55.6 years (SD, 12.9), and there were more male participants overall (57%). The groups were well matched, as there were no clinically meaningful differences as seen in Table 1.

Attrition
There were four participants (5%) from the experimental group and three participants (4%) from the control group who withdrew while undertaking iCBT, generally due to time pressures or health problems. Significantly more participants [χ² (1, n = 146) = 5.8; p = 0.02] from the control group (99%) completed the assessment measures at T₁ compared with those from the experimental group (73%). There was no significant difference [χ² (1, n = 146) = 2.1; p = 0.16] in completion rates at T₂ with 73% from the experimental group and 82% from the control group completing these assessment measures. No significant baseline differences in terms of age, gender, employment status and level of education, tinnitus severity,
TABLE 1. Baseline Demographical and Clinical Characteristics of the Participants

<table>
<thead>
<tr>
<th>Category</th>
<th>Description</th>
<th>Experimental Group (n = 73) (%)</th>
<th>Control Group (n = 73) (%)</th>
<th>Overall (n = 146) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td>Male</td>
<td>43 (59)</td>
<td>40 (55)</td>
<td>83 (57)</td>
</tr>
<tr>
<td></td>
<td>Female</td>
<td>30 (41)</td>
<td>33 (45)</td>
<td>63 (43)</td>
</tr>
<tr>
<td>Age</td>
<td>Mean years (SD)</td>
<td>56.8 (12.2)</td>
<td>54.3 (13.5)</td>
<td>55.6 (12.9)</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>24–79 yr</td>
<td>22–83 yr</td>
<td>22–83 yr</td>
</tr>
<tr>
<td>Tinnitus duration</td>
<td>Mean years (SD)</td>
<td>11.1 (11.5)</td>
<td>12.4 (12.2)</td>
<td>12.4 (12.2)</td>
</tr>
<tr>
<td></td>
<td>Range</td>
<td>0.3–52 y</td>
<td>0.3–56 y</td>
<td>0.3–56 y</td>
</tr>
<tr>
<td>Using hearing aids</td>
<td>No</td>
<td>46 (63)</td>
<td>46 (63)</td>
<td>92 (63)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>27 (37)</td>
<td>27 (37)</td>
<td>54 (37)</td>
</tr>
<tr>
<td>Employment status</td>
<td>Retired/unemployed</td>
<td>30 (41)</td>
<td>32 (44)</td>
<td>62 (44)</td>
</tr>
<tr>
<td></td>
<td>Professional</td>
<td>18 (25)</td>
<td>23 (32)</td>
<td>41 (28)</td>
</tr>
<tr>
<td></td>
<td>Service occupation</td>
<td>9 (12)</td>
<td>6 (8)</td>
<td>15 (10)</td>
</tr>
<tr>
<td></td>
<td>Administrative/sales</td>
<td>8 (11)</td>
<td>9 (12)</td>
<td>17 (12)</td>
</tr>
<tr>
<td></td>
<td>Technical</td>
<td>8 (11)</td>
<td>3 (4)</td>
<td>11 (8)</td>
</tr>
<tr>
<td>TFI score</td>
<td>59.8 (18.0)</td>
<td>59.2 (19.0)</td>
<td>59.5 (18.4)</td>
<td></td>
</tr>
</tbody>
</table>

Efficacy of iCBT Versus Weekly Monitoring for Tinnitus Distress

Differences between the treatment arms were not constant over time (Table 2). Preintervention (T0) means were similar. At postintervention (T1), the mean TFI score was 21 points lower (SD, 14.9) compared with baseline among those in the experimental group. For the control group, the mean TFI score was 5 points lower (SD, 3.9) when compared with baseline. Although both groups exhibited reduced mean scores, the magnitude of the reduction in mean score in the experimental group was greater than in the control group, and this difference was statistically significant (Cohen’s d = 0.7) as seen in Table 2.

Figure 2 shows that the majority of the experimental group had a T0–T1 difference score reduction of 10–40 points, with a maximum reduction of 81 points. In comparison, the majority of the control group had smaller improvements with a T0–T1 difference score higher than baseline or up to 20 points reduced. The maximum improvement for the control group was 29 points. Both groups had similar means at follow-up (T2), indicating that the control group improved further after completing the intervention as summarized in Table 2.

Using the reliable change criterion of 23.3 in TFI score (i.e., 1.96 times the SE of 11.9), clinical significance was achieved by 51% of the experimental group and 5% of the control group at T2. A clinically significant change was found for 47% of the control group at T2 after they undertook the intervention. At T2, there was 41% of the experimental group and 1% of the control group with TFI scores below the level requiring intervention (< 25) who also had a reliable change of 23.3 after they completed the intervention. This was achieved by 38% of the control group at T2.

Efficacy of iCBT Versus Weekly Monitoring for Comorbidities Associated With Tinnitus

Differences between the secondary assessment measures were not constant over time for the treatment arms (Table 2). Preintervention (T0) means were similar. At postintervention (T1), the experimental group had a significantly greater reduction in insomnia, depression, hyperacusis, cognitive failures, and improvement in the quality of life in comparison with the control group. For anxiety and hearing disability, significant differences were not constant over time for the treatment arms (Table 2). Preintervention (T0) means were similar. At postintervention (T1), the experimental group had a significantly greater reduction in insomnia, depression, hyperacusis, cognitive failures, and improvement in the quality of life in comparison with the control group. For anxiety and hearing disability, significant differences were not constant over time for the treatment arms (Table 2).

TABLE 2. Within and Between Group Comparisons of the Assessment Measures Over Time

<table>
<thead>
<tr>
<th>Assessment Measure</th>
<th>Control vs. Experimental Group Mean Difference at Each Time Point (SD)</th>
<th>Group Comparison: F-Statistic*</th>
<th>Follow-Up Analysis: t-Statistic</th>
<th>Cohen's d (95% Confidence Interval)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T0</td>
<td>T1</td>
<td>T2</td>
<td>Time by Group Interaction</td>
</tr>
<tr>
<td>TFI</td>
<td>-0.6 (0.4)</td>
<td>15.1 (10.6)</td>
<td>3.5 (2.5)</td>
<td>15.8; p &lt; 0.001†</td>
</tr>
<tr>
<td>ISI</td>
<td>1.2 (0.8)</td>
<td>3.8 (2.7)</td>
<td>2.5 (1.7)</td>
<td>5.3; p = 0.006†</td>
</tr>
<tr>
<td>GAD-7</td>
<td>-0.4 (0.3)</td>
<td>1.4 (1.0)</td>
<td>0.4 (0.3)</td>
<td>3.1; p = 0.05</td>
</tr>
<tr>
<td>PHQ-9</td>
<td>0.2 (0.2)</td>
<td>1.9 (1.3)</td>
<td>0.4 (0.3)</td>
<td>3.7; p = 0.03†</td>
</tr>
<tr>
<td>HHIA-S</td>
<td>1.2 (0.8)</td>
<td>2.6 (1.8)</td>
<td>0.6 (0.4)</td>
<td>1.7; p = 0.18</td>
</tr>
<tr>
<td>HQ</td>
<td>0.2 (0.3)</td>
<td>3.2 (2.3)</td>
<td>1.4 (1.0)</td>
<td>3.1; p = 0.04†</td>
</tr>
<tr>
<td>CFQ</td>
<td>1.3 (0.9)</td>
<td>6.5 (4.6)</td>
<td>3.6 (2.5)</td>
<td>4.2; p = 0.01†</td>
</tr>
<tr>
<td>SWLS</td>
<td>-0.4 (0.3)</td>
<td>-2.2 (1.5)</td>
<td>-0.6 (0.4)</td>
<td>3.1; p = 0.04†</td>
</tr>
</tbody>
</table>

SD= Standard Deviation, T0= preintervention, T1= postintervention, T2= follow-up, TFI= Tinnitus Functional Index, ISI= Insomnia Severity Index, GAD= generalized anxiety disorder; PHQ= Patient Health Questionnaire. HHIA-S= Hearing Handicap Inventory for Adults-screening version, HQ= Hyperacusis Questionnaire, CFQ= Cognitive Failures Questionnaire, SWLS= Satisfaction with Life Scales.

†Significance at < 0.05.
within group differences were found postintervention, but no significant interaction between time and group was seen. For the assessment measures that were statistically significant, they were only clinically significant for a few participants at T1. Clinical significance (score change > 9.8) was reached by 22% of the experimental group and 4% of the control group for the ISI. For the PHQ-9, this was reached by 16% of the experimental group and 4% of the control group (score change of 6.4). For the HQ clinical significance (score change of 14.3) was reached by 11% of the experimental group and 4% of the control group. For the CFQ, it was 18% and 5% for the groups, respectively (score change of 14.1), whereas it reached 14% and 3% for the respective groups for the SWLS (score change of 6.3). The ISI had the highest percentage of participants having a clinically significant change among the secondary assessment measures.

Both groups had similar means at follow-up (T2), indicating that the control group had improved to the level of the experimental group after completing the intervention as summarized in Table 2.

Stability of Intervention Effects

There was no significant difference in the TFI scores between T1 and T2 for the experimental group, as seen in Figure 3. Likewise, improvements were maintained for all secondary assessment measures as no statistically significant differences were found. Intervention effects were, therefore, maintained 2-months postintervention for the experimental group.

Monitoring Intervention Effects Between T0 and T1

Differences between the intervention arms were not constant across the 8-time points between T0 and T1. The experimental group had a greater weekly reduction in tinnitus distress, as evidenced by the significant interaction \( F(7, 1008) = 19.5; p = 0.001^*; \) Cohen’s \( d = 0.9 \).
Follow-up analysis examining this main effect week-by-week indicated no group differences in weeks 1 to 3 of this period. From week 4 to 8, there were significant differences, as the experimental group’s tinnitus distress was rated significantly lower than that of the control group who were not undergoing the intervention, as seen in Figure 4.

DISCUSSION

This randomized trial found that iCBT guided by an audiologist was effective in reducing tinnitus distress. The symptoms of several tinnitus comorbidities, such as insomnia, depression, hyperacusis, and cognitive failures were also reduced, and an increase in life satisfaction was found. Results were stable 2-months postintervention. This discussion highlights the implications of the finding for each of the four research objectives.

Effects of iCBT for Tinnitus Distress

The main outcome measure for this trial was a change in tinnitus distress as measured by the TFI. Undertaking iCBT led to significantly greater improvements in tinnitus distress, compared with weekly monitoring. The small improvement found in the control group (5 points) at T1 may have been related to the positive effects of being included on an intervention pathway, despite not yet starting the intervention. The mean score reduction of 21 between T0 and T1 in the experimental group in the present study is comparable with the findings in the initial feasibility study with a mean difference of 19 points (Beukes et al. 2017). The TFI score improvements found in the experimental group were greater than those occurring in the control group.

To relate these findings to clinical significance, the RCI was calculated. Results indicated that a change of 23.3 on the TFI score was regarded as clinically significant. This was similar to the change of 23.9 found in the initial feasibility trial. At T1, clinical significance was reached by 51% of the experimental group and 5% of the control group. Earlier iCBT for tinnitus trials found that a clinically significant change was reached by 29–52% of participants (Andersson et al. 2002, Kaldo et al. 2008, Nyenhuis et al. 2013 and Jasper et al. 2014). A more recent study by Weise et al. (2016) reported that a higher proportion (73–81%) reached clinical significance following undertaking iCBT for tinnitus. Andersson et al. (2002) and Kaldo et al. (2008) reported finding 4–5% of a waiting-list control group achieved clinical significance, in line with the results of the present study. Discrepancies between different trials may be partly related to the differences in assessment measures used. Previous iCBT trials have used varying tinnitus assessment measures such as the THI, the Tinnitus Reactions Questionnaire (Wilson et al. 1991), or the Tinnitus Questionnaire (Hiller et al. 1994) with various study designs, thereby making direct comparisons difficult. Andersson (2015) reported that the pooled effect size of previous iCBT control studies (Andersson et al. 2002; Abbot et al. 2009; Hesser et al. 2012; Nyenhuis et al. 2013; Jasper et al. 2014) was Hedges’ g = 0.6, although a later study by Weise et al. (2016) was not included. Weise et al. (2016) found an effect size of Hedge’s g = 0.8 for tinnitus distress postintervention when using the THI. The medium effect size found of Cohen’s d = 0.7 (Hedge’s g = 0.7) for the present study is, therefore, between the values of previous iCBT tinnitus trials. This provides encouragement that the results of this study are consistent with those of previous iCBT trials.

In previous clinical trials, the intervention was guided by clinical psychologists trained in the provision of CBT. This trial is unique in providing this guidance using an audiologist, in line with tinnitus health care provision in the United Kingdom. Results indicate the efficacy of audiologist-guided iCBT to reduce tinnitus distress. Previous Internet-based trials for depression, anxiety, and social phobia have found comparable results, regardless of whether a clinician or an appropriately trained technical assistant guided the intervention (Titov et al. 2009; Robinson et al. 2010; Titov et al. 2010).

Effects of iCBT for Comorbidities Associated With Tinnitus

Significant improvements for insomnia, depression, hyperacusis, cognitive failures, and satisfaction with life were evident. Each group significantly improved in terms of anxiety and hearing disability following the completion of the iCBT intervention, but no main effect for the interaction between time and group was seen for these assessment measures. This may be related to the large variability in scores for these assessment measures.
between the groups over time. Low baseline scores were also evident for the anxiety assessment measure (7 points; SD, 0.3), which may have contributed to the nonsignificant interaction found. To relate these findings to clinical significance, the RCI was calculated for each secondary assessment measure at T1. For the ISI, 22% of the experimental group had a clinically significant change, compared with 4% of the control group. The range of clinical significance for the other secondary assessment measures were 11–18% of the experimental group and 3–5% of the control group. The proportions of those with clinically significant improvements with regard to the secondary assessment measures are, therefore, lower than those found for the TFI.

Previous trials of iCBT for tinnitus have used secondary outcome measures for insomnia (using the ISI), anxiety, and depression (using the Hospital Anxiety and Depression scale; Zigmund & Snaith 1983). Significant intervention effects have been reported for these tinnitus-associated comorbidities (Kaldo-Sandström et al. 2004; Kald et al. 2008; Jasper et al. 2014; Weise et al. 2016). These studies have not reported whether these results were clinically significant as they focused on statistical significance. Effect sizes in the present study for anxiety and depression (d = 0.3) were lower than those reported by Jasper et al. (2014) and Weise et al. (2016) of d = 0.5. This difference may partly be attributed to the difference in assessment measures used in these trials compared with that in the present trial. The result for insomnia (d = 0.6) for the present study was similar to that of Jasper et al. (2014) of d = 0.6 and lower than that reported by Weise et al. (2016) of g = 0.7.

Stability of Intervention Effects

Maintaining intervention effects is an important aspect of the efficacy of an intervention. It was found that the intervention effects were stable 2-months postintervention (T1) for both tinnitus severity and the secondary assessment measures in the experimental group. Stability of iCBT intervention effects have also been found in previous trials that monitored these effects over a longer period. Jasper et al. (2014) reported stability 6 months after completing iCBT for tinnitus severity, anxiety, depression, and insomnia. Kald et al. (2008), using a Swedish population, and Weise et al. (2016), using a German population, found results that were maintained 1-year after undertaking iCBT for tinnitus severity, anxiety, depression, and insomnia.

Intervention Effects During iCBT

A further objective of this trial was determining when intervention effects can be expected. Participant’s tinnitus severity was, therefore, monitored on a weekly basis by means of the THI-S. After the experimental group completed 4 weeks of the iCBT intervention, they had significantly lower tinnitus severity scores than those not undergoing the intervention. The likely delay in intervention effects are important to convey to future participants to adjust their expectations.

Study Limitations and Further Research

This study is not without limitations, which have implications for result interpretation. First, the participants were recruited from the general public due to interest in undertaking an Internet-intervention and not from a clinical setting. Therefore, these results may not be the same in a clinical sample. The demographical distribution of the participants in the present study showed more male participants, a slightly higher mean age distribution and longer tinnitus duration than those reported by previous iCBT trials on tinnitus (e.g., Andersson et al. 2002; Kald et al. 2007; Weise et al. 2016). This should be considered when assessing the generalizability of the results. Second, the likelihood of type I errors cannot be excluded due to multiplicity of testing. Third, not all participants completed the outcome measures at T1 and T2. Ways of encouraging more participants to complete these questionnaires and minimize attrition is required. A deeper understanding of factors affecting adherence is thus needed. The fourth limitation involves the assessment measures used. The HQ was used despite concerns raised regarding its psychometric properties (Fackrell et al. 2015) because of a lack of a better measure for hyperacusis. The TFI was selected as it was developed to evaluate intervention effects. There are, however, limitations in selecting this outcome measure as it has only been recently developed and further psychometric evaluations are required. Fackrell et al. (2016) raised concerns regarding substantial floor effects on many items and concluded that it may not be good at detecting treatment-related benefits in a research population. It may, therefore, have been a suboptimal assessment measure for a research volunteer population as used in the present trial. Lastly, data were not collected on treatment credibility, which is an important consideration regarding evaluating a new intervention.

Further research is required to gain more insights into iCBT for tinnitus. This includes determining the longer term results and participant’s experiences regarding this version of iCBT used and using an audiologist to guide the intervention. In addition, determining moderators and mediators of outcome (Hesser et al. 2014) and which specific aspects of iCBT result in positive outcomes, needs further exploration. Comparing intervention effects when guidance is provided by an audiologist versus a psychologist is required to determine the efficacy of using an audiologist for iCBT. Comparing these results with existing tinnitus clinical care is also required. A further RCT is therefore underway to compare iCBT with individualized face-to-face clinical care for tinnitus in the United Kingdom (Beukes et al. 2017).

CONCLUSIONS

Guided iCBT for tinnitus using audiological support resulted in statistically significant reductions in tinnitus distress and comorbidities (insomnia, depression, hyperacusis, cognitive failures) and improved quality of life. Including iCBT as an additional tinnitus intervention could be a cost-effective way of increasing access to CBT for tinnitus.

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All authors conceived and designed this study. GA developed the Swedish original iCBT intervention for tinnitus together with Viktor Kald, EB developed this version for use in the United Kingdom, carried out the study, and analyzed the data. The article was drafted by EB and critically revised and approved by all authors.
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