- 1 Clinical effectiveness and cost minimisation model of Alpha-Stim cranial electrotherapy
- 2 stimulation in treatment seeking patients with moderate to severe generalised anxiety disorder.
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Abstract

- 19 Background. Cranial electrotherapy stimulation (CES) is a well-tolerated neuromodulation treatment
- 20 with demonstrated trial efficacy in anxiety disorders. The aim of the current study was to
- 21 demonstrate its clinical and cost effectiveness during and after CES in people with generalised
- anxiety disorder (GAD) who had not responded to low intensity psychological treatment in a routine
 health service.
- 24 Methods. Consecutive sample of eligible patients with GAD waiting for individual cognitive
- 25 behaviour therapy (CBT) selected from two publicly funded services in England. They received 60
- 26 minutes per day Alpha-Stim CES for 6-12 weeks. Primary outcome was remission on the GAD-7 scale
- at 12 and 24 weeks. Cost effectiveness was examined using a cost minimisation model of directhealth costs.
- 29 **Results.** Of 161 patients recruited, 72 (44.7%) and 77 (47.8%) achieved remission on the GAD-7 at 12
- 30 and 24 weeks respectively with 122 (75.8%) receiving at least 6 weeks CES. Mean (sd) GAD-7 score at
- 31 baseline significantly improved from 15.77 (3.21) to 8.92 (5.42) and 8.99 (6.18) at 12 and 24 weeks
- 32 respectively (p<0.001). 80 (49.7%) participants required further individual CBT. CES provided a
- 33 saving of £540.88 per patient (95% CI -£327.12, £648.69).
- 34 Limitations. Participants were not randomised and there was no control group. Only 48 (29.9%)
- 35 participants completed every assessment.
- 36 **Conclusion.** In patients with generalised anxiety disorder not responding to low intensity
- 37 psychological treatment, 6-12 weeks daily Alpha Stim CES may be effective after treatment and 3
- 38 months later, thereby reducing the need for individual CBT and saving health costs.
- 39 247 words

40 Keywords

41 Cranial electrotherapy; neuromodulation; generalised anxiety disorder; cost effectiveness.

42 Abbreviations

- 43 AIS Athens Insomnia Scale; Alpha-Stim AID cranial electrotherapy stimulator for control of anxiety,
- 44 insomnia and depression; CBT cognitive behaviour therapy; CE Conformité Européene, European
- 45 Union regulatory marking; CES cranial electrotherapy stimulation; CI confidence interval; CSRI -
- 46 Client Service Receipt Questionnaire; DSM-IV- Diagnostic and Statistical Manual of Mental Disorders
- 47 4th Edition; EEG electroencephalography; EQ5D-5L Eurogol; FIML full information maximum
- 48 likelihood estimation; fMRI functional magnetic resonance imaging; GAD generalised anxiety
- 49 disorder; GAD-7 self-rated measure of generalised anxiety disorder symptoms; GLM general
- 50 linear model; HE health economics; iCBT individual cognitive behaviour therapy; IAPT Improving
- 51 Access to Psychological Treatment service; IRAS Integrated Research Application Service; ITT -
- 52 intention to treat; MCAR missing completely at random; NHS National Health Service; NICE -
- 53 National Institute for Clinical Excellence; NRES National Research Ethics Service; PHQ-9 Personal
- 54 Health Questionnaire 9 item; PSA Probabilistic sensitivity analysis; PSSRU Personal Social Services
- 55 Research Unit; RCT randomised controlled trial; RM ANOVA repeat measures analysis of variance;
- 56 WASA Work and Social Adjustment Scale

59 Introduction

60 Generalised anxiety disorder (GAD) is a common and persistent mental disorder with a point or 61 annual prevalence of 2.1 to 4.4% (Hunt et al, 2002; Grant et al, 2005; Remes et al, 2017; Ruscio et al, 62 2017). GAD is often present with other mental disorders such as depression, other anxiety disorders, 63 insomnia and physical illness (Chapman et al 2010; Ruscio et al, 2017), all of which can lead to 64 considerable health expenditure (Sandelin et al, 2013). According to the National Institute for 65 Clinical Excellence (NICE) Guideline for Generalised Anxiety Disorder for England and Wales (NICE, 66 2011), the first step in the management of GAD is education about the condition and monitoring 67 delivered in primary care. The second step is low intensity psychological intervention of the person's 68 choice, which is provided by the Improving Access to Psychological Treatment service (IAPT) in all 69 parts of the National Health Service in England (NICE, 2011), usually in the form of facilitated 70 computerised cognitive behaviour therapy or bibliotherapy (Gyani et al, 2013). While these 71 approaches are relatively cheap and effective, many people with GAD do not improve and require 72 additional treatment (Andrews et al, 2018). The third step NICE recommended intervention is either 73 a high intensity psychological intervention such as individual cognitive behaviour therapy (iCBT), also 74 delivered by IAPT services and relatively expensive, or drug treatment, initially with selective 75 serotonin reuptake inhibitor antidepressants but if these are ineffective then more expensive drugs 76 such as pregabalin are used. There can be a substantial delay before iCBT can be offered (Sandelin et 77 al, 2013).

78

79 Cranial electrotherapy stimulation (CES) was first utilised to induce sleep and relaxation using bursts 80 of small electric currents applied to the head in the 1900s (Guleyupoglu et al, 2013). Improvements 81 have taken place in electrode placement, use of battery driven devices and understanding of dose, 82 frequency of treatment and waveform that is required to improve anxiety symptoms. Single courses 83 of CES are associated with changes in electroencephalography (EEG) from delta (0-3.5Hz) and beta 84 (12.5-30Hz) frequencies to more relaxing and alerting alpha frequencies (8-12 Hz) (Kennerly, 2004). 85 Cortical and subcortical brain activation on fMRI have been demonstrated in people with high levels 86 of anxiety (Feusner et al, 2012) and increases in plasma beta endorphins, adrenocorticotrophic 87 hormone and cortisol (Liss and Liss, 1996; Shealy et al, 1998) after a single 20 minute CES treatment. 88 89 A recently published systematic review funded by the United States Department of Veteran Affairs 90 identified five randomised controlled trials (RCTs) with 198 participants for anxiety disorders 91 comparing active CES to sham CES (Shekelle et al, 2018). It concluded that there was low quality 92 evidence of the effectiveness of CES for anxiety and depression symptoms in people with anxiety 93 disorders at the end of treatment as well as evidence that CES does not cause serious side effects. A 94 randomised controlled trial in 115 volunteers with a primary anxiety disorder showed the 95 effectiveness of 5 weeks of active CES versus sham CES on anxiety and depression symptoms at the 96 end of treatment (Barclay and Barclay, 2014). However, there have been no studies of the 97 maintenance of clinical improvement or cost effectiveness of CES in treatment seeking patients with 98 GAD who had not responded to second-line treatment as recommended by NICE (2011). Therefore 99 we examined the clinical and cost effectiveness of 6-12 weeks CES treatment for treatment seeking 100 patients with GAD who had not responded to facilitated computerised cognitive behaviour therapy 101 or bibliotherapy over 24 weeks. These patients were all waiting for iCBT for GAD. 102 103 There are four aims to the current study to determine:

- The proportion of patients treated with CES in IAPT services who reach the clinical threshold for remission (GAD-7 score of 7 or less; Spitzer et al, 2016), reliable improvement and recovery after treatment at 12 weeks.
- The proportion of patients treated with CES in IAPT services who maintain the clinical
 threshold for remission (GAD-7 score of 7 or less), reliable improvement and recovery at 24
 weeks.
- If there are significant changes over 24 weeks in generalised anxiety, depression, insomnia,
 social adjustment and quality of life.
- If the cost of CES offsets the cost of psychological treatment and other treatment over 24 weeks.

115 Method

- **Design.** This is a study in routine care carried out after efficacy has been established against sham
- treatment in a meta-analysis of RCTs (Shekelle et al, 2018) to establish the effectiveness and costs in
- 118 routine care settings as outlined by the United Kingdom Medical Research Council Complex
- 119 Intervention Framework (2000) and the National Institute for Health and Care Excellence (2018). An
- open consecutive patient cohort design with 24 week follow up in National Health Service (NHS)
- 121 mental health treatment settings in England was employed where all participants were offered
- 122 Alpha-Stim cranial electrotherapy stimulation (CES) for 6-12 weeks if they had not reached remission
- 123 with therapist or full guided self-help and were waiting to receive individual cognitive behaviour
- 124 therapy (iCBT).
- 125 Setting. Two NHS Improving Access to Psychological Treatment (IAPT) services in the same county in
- 126 England covering a more affluent urban and rural area and a less affluent inner city area. The
- 127 services were run by two different NHS organisations. All data and treatment were delivered by staff
- 128 who were independent of the company who makes Alpha Stim CES. Ethical approval for the study
- 129 was granted by the Nottingham 2 NRES committee (IRAS206555).

130 Inclusion/exclusion criteria:

- 131 1. A score of 8 or more on GAD-7 scale, a 7-item self-rated measure of symptoms of
- 132 generalised anxiety disorder (Spitzer et al, 2016), because nationally IAPT services determined that
- 133 further treatment should be offered after full or guided computerised self-management or
- bibliotherapy if a person scores above the threshold for remission i.e. a total score of 8 or more.
- A clinical diagnosis of generalised anxiety disorder alone or in combination with a comorbid depression or other anxiety disorder e.g. obsessive compulsive disorder or physical health morbidity.
 Excluded was a diagnosis of any other mental disorder e.g. substance use disorder, eating disorder, bipolar disorder, non-affective psychosis. In keeping with an implementation study the diagnostic information used for the inclusion and exclusion criteria were made on clinical grounds without using any standardised psychiatric interviews by clinically qualified mental health professionals independently of the research team.
- 142 3. On waiting list for individual CBT (high intensity psychological intervention).
- 143 4. Does not require urgent clinical care.
- 144 5. If female not known to be pregnant.
- 145 6. Implantation with a pace maker or an implantable cardioverter device (ICD) are exclusions.
- 146 6. Gives informed written and oral consent to the study.
- 147 7. Agrees to return Alpha-Stim equipment at the end of the study.

148 Being on medication did not lead to exclusion.

149 **Outcome measures:**

- 150 These are standard clinical outcome measure employed routinely by the NHS IAPT services with the
- addition of measures of insomnia, quality of life and an economic interview to assess health costs.
- 152 They were collected face to face at baseline. Clinical outcome and quality of life measure were
- 153 collected at four, six, eight, 12 and 24 weeks by e-mail, telephone or post according to participant
- 154 preference. A second economic interview was conducted by telephone or Skype at six months
- according to participant preference. All participants who completed the economic interview were
- given a £10 gift voucher in recognition of the time given to completing the research outcome
- 157 assessments.

158

159 **Primary outcome measure:**

- 160 The primary outcome is the proportion of participants who reach remission (7 points or less) at 12
- and 24 weeks on the GAD-7 since IAPT services are paid according to the proportion of patients who
- 162 reach this threshold after treatment in their service (Richards and Borglin, 2011). Other key
- 163 outcomes are the proportion of cases who meet a clinically important ("reliable improvement") 5
- point improvement on the GAD-7 at 12 and 24 weeks (Richards and Borglin, 2011), the proportion
- 165 who meet criteria for recovery (GAD-7 score of 7 or less and also exhibiting a 5 point drop in GAD-7
- score) at 12 and 24 weeks (<u>Richards and Borglin, 2011</u>), and the effect size of the change in GAD-7
- score over 12-24 weeks. A clinically important deterioration is an increase in GAD-7 score of 5 points
- 168 at 12 and 24 weeks (Richards and Borglin, 2011).

169 Secondary outcome measures:

170 1. Personal Health Questionnaire, 9-item (PHQ-9; Kroenke et al, 2001), a 9-item self-rated 171 measure of the severity of depression symptoms. Remission is a total score of 9 or less at 12 or 24 172 weeks in those who had scored 10 or more at baseline, reliable improvement is a drop of 6 points or 173 more, and recovery is a score of 9 or less and a 6 point drop at 12 and 24 weeks <u>(Richards and 174 Borglin, 2011)</u>. We also examined the effect size of the change in PHQ-9 score symptoms from 175 baseline to 12 and 24 weeks.

1762.Athens Insomnia Scale (AIS; Soldatos et al, 2000). This scale has 8 items with a maximum177score of 24. A score of 6 indicates a possible sleep problem and 4 indicates recovery (Soldatos et al,1782003). Therefore Rremission is defined as the proportion of people who score a total of 4 or less at17912 and 24 weeks. No data exists on reliable improvement so a drop of 50% in baseline score by 12180and 24 weeks was used. Recovery is the proportion of people who showed a drop of 50% in baseline181score and scored 4 or less at 12 and 24 weeks. We also examined the effect size of the change in182insomnia symptoms from baseline to 12 and 24 weeks.

3. Work and Social Adjustment Scale (WASA; Mundt et al, 2002), an 8-item self-rated measure
of work and social function. A total score of 20 or more indicates considerable impairment in
function (Mundt et al, 2002). A return to normal function requires a total score of 10 or less and
functional recovery requires a total score of 11 or more at baseline with a drop to 10 points or below
by 12 and 24 weeks (Mundt et al, 2002). We also examined the effect size of the change in WASA
score from baseline to 12 and 24 weeks.

4. EQ5D-5L (EuroQol, van Hout et al, 2012), a 6- item self-rated measure of health utility and
quality of life. We examined the effect size of the change in EQ5D-5L from baseline to 12 and 24
weeks.

193 Economic interview:

We used the Client Service Receipt Interview (CSRI; Beecham and Knapp, 1992) adapted for use in
studies of anxiety disorders in primary care and community settings. It was completed at baseline
and 24 weeks.

197

198 Procedure.

199 Consecutive treatment seeking patients who received low intensity IAPT interventions (therapist 200 guided self-management on a computerised CBT programme or bibliotherapy for GAD) but had not 201 reached a total score of 8 or more, were unlikely to meet any exclusion criteria for the study, and 202 were willing to be placed on a waiting list for iCBT, were identified from IAPT service records. IAPT 203 staff contacted a potential participant to seek permission for their contact details to be passed to 204 the study team who checked their eligibility over the phone. A face to face meeting was arranged 205 with a member of the study team who checked the inclusion/exclusion criteria and sought written 206 informed consent. If the participant consented study staff showed the participants how to use the 207 Alpha-Stim CES device, outlined how to obtain support while using it, and negotiated the return of 208 the CES device at the end of 6-12 weeks treatment. Women of child-bearing potential completed a

- 209 urine pregnancy dipstick human chorionic gonadotropin test.
- 210 Alpha-Stim AID is a CE marked medical device which is marketed for the alleviation of psychological
- 211 conditions including anxiety, insomnia and depression, through using cranial electrotherapy
- stimulations (CES) which are tiny electric currents applied through ear clips worn for 60 minutes per
- 213 day. The treatment provided by the device is therefore non-invasive, non-pharmacological, and can
- be used as adjunctive treatment to drug or psychological treatment or a treatment on its own. All
- participants were offered 60 minutes per day of alpha-stim CES treatment at a current of one
- 216 <u>hundred micro amps</u>-per day 7 days per week for 6 consecutive weeks. <u>The 60 minutes session</u>
- 217 <u>starts when the ear clips are attached and stops automatically when the hour is finished. The device</u>
 218 <u>was not locked -because it would not be in usual clinical practice. The device did not automatically</u>
- 219 <u>record adherence to treatment.</u> Participants could choose to continue with the same CES treatment
- for a further 6 weeks, thereby completing 12 weeks CES treatment in total. At the end of 12 weeks
- the participants could not receive any further CES treatment. Since this was a naturalistic study,
- 222 decisions concerning if and when iCBT might be received by the participant were made by IAPT staff
- with the participants; the study team did not influence this decision. If participants started iCBT
- during the 6-12 weeks of CES, they could continue with CES while receiving iCBT at the same time.
- 225 Similarly general practitioners could independently decide to place the patient on medication for
- GAD at the same time as participants continued to receive CES. A summary of the procedures of the
- study is shown in Table 1; as well as outcome measures, adherence to CES and side-effects were
- recorded at each study visit.

229 Sample size.

A meta-analysis of 5 CES RCTs estimates an effect size of at least 0.60 (Shekelle et al, 2018). On this basis remission might be expected in 26.5% patients with GAD receiving alpha stim CES in IAPT settings. The aim was to recruit a sample with at least 25 participants achieving remission after alpha stim CES at 12 weeks and followed up at 24 weeks; a sample of 160 would be required assuming 40% loss to follow up by 24 weeks.

235 Statistical analyses.

- 236 Prior to statistical analyses, data screening was conducted to evaluate the tenability of assumptions
- 237 specific to the general linear model (GLM). These assumptions included normally distributed outcome

238 variables, independence of observations for different subjects, and homogeneity of covariance 239 matrices within subjects across repeated measurements. The assumptions of the GLM were tenable 240 except for homogeneity of covariance within subjects on their measurements over time. The 241 Greenhouse-Geiser adjustment was applied to F-statistics and degrees of freedom when violations 242 appeared. After data screening, analyses proceeded using a within-subjects repeated measures 243 analysis of variance (RM ANOVA) for the primary outcome and secondary outcome variables. Additionally, regarding aims 1 and 2, descriptive analyses were conducted to determine remission, 244 245 reliably improvement and recovery.

To answer our research aim 3, we used a within-subjects univariate repeated measures analysis of variance (RM ANOVA). Separate univariate RM ANOVAs were conducted for each outcome variable in two distinct phases. The first set of analyses proceeded using data from the empirical sample. The second set of RM ANOVA analyses included an intention-to-treat (ITT) analysis strategy using a full complement of scores on each outcome variable. The following section includes information specific to the ITT analytic approach.

Intention-to-treat (ITT) analysis avoids overoptimistic estimates of the efficiency of an intervention 252 253 resulting from the removal of non-compliers by accepting that noncompliance and protocol deviations 254 are likely to occur in clinical practice. Intention-to-treat analyses was applied including all patients as 255 they were assigned at baseline, regardless of their adherence to treatment, the treatment they 256 received or any subsequent withdrawal from the study (Fisher, 1990). To evaluate the type or pattern 257 of missing scores for each outcome measure, the missing completely at random (MCAR) test was 258 employed (Little and Rubin, 2002; Enders, 2010). Once the data was determined to adhere to MCAR 259 (i.e. p >.05), replacement of scores proceeded using model-based full information maximum likelihood 260 (.FIML) estimation.

261

262 Health economics

263 In order to determine the cost impact of introducing CES into the pathway as a second-line 264 treatment instead of or prior to individual CBT (iCBT), a cost minimisation analysis was undertaken 265 using a health economic (HE) model decision tree (see Figure 1). In both branches of the HE model 266 the patient population was non-responders to low-intensity guided or full computerised self-help or bibliotherapy given as the first-line treatment. The decision tree was populated with the 267 268 probabilities of response to second line CES treatment from the study versus second-line iCBT with 269 the remission rate of 54.2% from Gyani et al (2013) which is the average remission rate between 270 guided and full self-help groups in that study. In addition, the same probability of outcome from 271 subsequent iCBT sessions given to non-responders in both arms was modelled as in the current 272 pathway (treatment as usual) such that for non-responders to second-line iCBT a further course of 273 the same number of iCBT sessions would follow. For non-responders to second-line CES up to two 274 further courses of iCBT were included in the decision tree. In all cases successful response was 275 measured by the achievement of the GAD-7 threshold of remission as used in the IAPT programme 276 (Richards and Borglin, 2011). Neither a cost-utility analysis nor a cost-consequences analysis was 277 employed because the study did not have a comparator for outcomes although EQ-5D results are 278 reported here separately for Alpha-stim CES treatment.

The hypothesis tested in the HE model was that adding CES as a second-line treatment in the pathway will eliminate, for the proportion of patients who respond to CES, the need for the more expensive iCBT leading to cost savings. Although not included in the model, it would also potentially reduce waiting times for those patients who would still progress to iCBT since early response to

available CES therapy promises to free up therapist resource for iCBT as well as potentially the

- number of iCBT sessions each participant would need after receiving CES. The HE model used a 6-
- 285 month time horizon, reflecting the expected duration of GAD response (NICE, 2011) and including
- the time period for consecutive treatments of CES and/or iCBT. Given this short time horizon, costs
- 287 were not discounted.

The modelling was undertaken from the United Kingdom NHS payer perspective with prices uplifted using the most recent national annually published resource, the PSSRU Unit Costs of Health and Social Care 2017 (Curtis and Burns, 2017) which gave compounded ratios for an uplift up to 2016.

291 Costs were derived for CBT from Radhakrishnan et al (2013) for 60 or 90 minutes of iCBT (£98.59 or £

- 292 176.97 per session) uplifted from 2010 to 2016 prices using the appropriate ratio of 1.09 yielding £
- 293 £110.96 and £199.17 respectively. Overall treatment costs were computed for 8 sessions of 60
- 294 minutes iCBT, as in the 'standard of care' model, yielding a total cost of £887.68. For comparison, the 295 model was also constructed with alternative choices of two additional more expensive iCBT regimes:
- the 'Clark and Wells model' with 14 sessions of 90 minutes sessions of iCBT, costing £2788.43 in total
- and the 'Heimberg model' with one session of 90 minute iCBT followed by 15 sessions of 60 minutes
- 298 iCBT, costing £1863.57 in total (NICE, 2013).
- Alpha-stim CES cost per treatment was a manufacturer estimate from the unit cost of the device of
- 300 £450.00 (excluding valued added tax) with a utilisation of 15 patients over an average product
- 301 lifetime of 3 years (based on a 10 week sole use per patient). It allowed for losses with respect to the

302 quoted 5 year warranty that was estimated to reduce average product lifetime by 2 years. A

- Additional therapist time, postage and consumables was estimated at £40, yielding £70 per
- 304 treatment.

305 A probabilistic sensitivity analysis (PSA) was undertaken on cost of treatment, probability of 306 response and utilisation of response with parameters as shown in Table 2 (York Health Economics 307 Consortium, 2016). In addition a one-way deterministic threshold analysis was performed on cost to 308 find the price at which the intervention would no longer be cost saving. Probabilistic sensitivity 309 analysis (PSA) is a technique used in economic modelling that allows the quantification of the level of 310 confidence in the output parameters of the analysis, in relation to the uncertainty in the model 311 inputs. In the probabilistic analysis, the parameters' value from clinical trials, observational studies 312 or in some cases expert opinion are represented as distributions around their deterministic value. A 313 set of input parameter values is drawn by random sampling from each distribution, and the model generates outputs (cost and health outcome), which are stored. This is repeated in many iterations 314 315 of the model (typically 1,000 to 10,000), resulting in a distribution of outputs that can be graphed on 316 the cost-effectiveness plane, and analysed.

317

318 Results

Figure 2 shows the flow of participants through the study. Only 22% of potentially eligible patients
agreed to take in the study. All 161 participants started CES treatment and 112 (69.6%) completed at

321 least 6 weeks treatment. Of the 49 (30.4%) participants who withdrew from treatment by 12 weeks,

- nine (5.6%) could not find the time to complete the treatment, four (2.5%) withdrew because of no
- improvement, four (2.5%) withdrew because of side effects (two with headaches and insomnia, one
- with nausea and one with a strange feeling after use), two (1.2%) withdrew because they felt better,
- and 30 (18.6%) gave no reason. Of the 161 participants, 80 (49.7%) had iCBT. Eighty-one (50.3%)
- 326 completed follow ups to 12 weeks and 72 (44.7%) to 24 weeks.

- 327 Table 2 shows that participants were drawn from a broad range of ages and nearly three quarters
- 328 were female. The overwhelming majority were white British, most had at least high school
- 329 education, married and were in employment. However, the mean baseline scores were in the severe
- range for GAD (Spitzer et al, 2001), moderately severe range for depression (Kroenke et al, 1999),
- showed significant sleep difficulties (Soldatos et al, 2004), substantial functional impairment (Mundt
- et al, 2002), and low health utility comparable to scores for out-patients with a broad range of
- 333 physical and mental disorders (van Hout et al, 2012).
- 334 Table 3 shows the primary outcome. By 12 weeks, 72 (44.7%) participants achieved remission and 335 recovery on the GAD-7 at 12 weeks and 76 (47.2%) at 24 weeks. The proportions of participants 336 achieving reliable improvement on the GAD-7 were 102 (63.4%) and 105 (65.2%) at 12 and 24 weeks 337 respectively. No patient showed reliable deterioration at 12 or 24 weeks. There was a drop in GAD-7 338 score from mean (sd) 15.77 (3.21) to 8.92 (5.42) by 12 weeks and this is maintained to 8.99 (6.18) at 339 24 weeks, a mild degree of GAD-7 symptoms by 12 and 24 weeks. The within-subjects effects is 340 statistically significant (F=72.02, df1=3.7/df2=563.74, p<0.001) and the effect size is medium (partial 341 eta square=0.31). The vast majority of the drop in GAD-7 is experienced in the first 6 weeks and 342 there is no statistically significant difference between week 6 and any subsequent time point up to week 24. The same pattern is seen in 48 participants with assessments at every time point except 343 344 the effect size of the within subjects treatment effect was large rather than medium (Appendix Table 345 1). Of the 81 participants who only received CES, 49 (60.3%) achieved remission on the GAD-7 at 12 weeks and 53 (65.4%) achieved remission on the GAD-7 at 24 weeks. Of the 25 participants who 346
- received both CES and iCBT, 17 (68%) achieved remission and recovery on the GAD-7 and 23 (92%)
- 348 achieved reliable improvement at 12 and 24 weeks.
- Table 3 shows that the effects on the PHQ-9 were similar in relation to the GAD-7 although a lower
- proportion achieved a reliable improvement at 12 and 24 weeks. The within subjects effect was
- significant (F=42.89, df1=3.9/df=559.01, p<0.001) with the mean PHQ-9 score dropping from the
- 352 moderately severe range to the mild range but the effect size was small (partial Eta square=0.21).
- 353 There was some worsening of depression symptoms by week 24 and the fall in PHQ-9 score was only
- 354 significant between baseline and 12 weeks but not 24 weeks. Only around a quarter of participants
- achieved remission on the Athens Insomnia Scale at 12 and 24 weeks. There was a statistically
- significant within-subjects drop in insomnia over the 24 period (F=42.69, df1=5.0/df=542.9, p<0.001)
- and the effect size was medium (partial Eta square=0.21).
- 358 Table 3 also demonstrates that just over a quarter of participants made a functional recovery on the
- 359 WASA at 12 and 24 weeks with CES. Figure 2 and Table 3 show that there is a significant within-
- 360 subjects effect of Alpha-Stim CES over the 24 weeks (F=17.35, df1=3.5/df=557.45, p<0.001) but the
- 361 effect size is small (partial Eta square=0.10). The effects of Alpha-Stim CES on the EQ-5D-5L were
- very similar to the WASA with a significant within subjects effect over 24 weeks (F=13.94,
- df1=4.1/df2=651.3, p<0.0001) but the effect size is also small (partial Eta square=0.08).
- 364 The results of the health economics decision tree model populated with the costs and probabilities 365 for the 8 session standard care model of CBT yielded the results as shown in Table 4. The costs and 366 responses are presented for a cohort of 1000 patients. CES provided a saving of -£540,878 (95% CI [-£648,692, -£327,117]) and the number of responses to treatment were increased by 187.56 per 367 368 1000 (95% CI [141.03, 227.82]). Using the "Clark and Wells model" of iCBT as comparator, CES 369 provided a saving of -£1,637,410 (95% Cls -£1,914,463, -£1,175,437]) and the number of responses 370 to treatment were increased by 187.56 per 1000 (95% CIs [141.58, 226.12]). With the Heimberg 371 Model as a comparator, CES provided a saving of -£1,212,463 (95% CIs -£1,429,369, -£843,394]) and

- the number of responses to treatment were increased by 187.56 per 1000 (95% CIs [140.79.,
- 227.71]). Cost-outcome scatterplots for each model are shown in the Appendix.
- 374

376 Discussion

377 This study shows that in moderate to severe treatment seeking patients with GAD, nearly 45 per 378 cent of patients achieved remission and 63 per cent reliable improvement in their self-rated anxiety 379 symptoms with Alpha-Stim CES treatment. These improvements were maintained for a further 12 380 weeks after CES was completed whether or not patients received iCBT in addition. Most of the 381 improvement with CES was seen in the first 4 weeks. It had a moderate effect size. Remission rates 382 are lower than reported for iCBT in routine IAPT services in the UK (Radhakrishnan et al, 2013); 383 however our sample had substantially higher scores than routinely reported for IAPT services 384 (Radhakrishnan et al, 2013; NHS Digital, 2018). . Approximately 50 per cent of patients on the 385 waiting list for iCBT received iCBT, thereby enabling the NHS IAPT services to treat other patients on 386 the waiting list for iCBT. The mean severity of GAD-7 symptoms decreased from severe to mild and 387 below case threshold over 12 weeks and remained at that level for 24 weeks. There were similar 388 drops in depression symptoms and insomnia symptoms as well as improvements in function and 389 quality of life although all of these effects were smaller with some slippage between 12 and 24 390 weeks. Although there was a significant drop in depression symptoms between baseline and 12 391 weeks, it was not significant at 24 weeks indicating that the effects of CES on depression symptoms 392 had started to wane by 24 weeks. Overall a quarter of patients receiving CES regained a functional 393 recovery. Alpha-Stim CES was well tolerated with only six (4%) patients stopping it because of side-394 effects and four (3%) because they were not making any progress. Compared to a standard course of 395 iCBT (eight sessions or longer), Alpha-stim CES reduced costs of care by £540 or more per patient 396 and it was also cost effective.

- 397 The strengths of the study were that clinical and cost effectiveness was examined in a consecutive 398 large sample of treatment seeking patients in universally available publicly funded services provided 399 by the state irrespective of the ability to pay or health insurance. Inclusion criteria were set to reflect 400 the criteria used by IAPT services to offer individual CBT. This criteria was set at 8 or more on the 401 GAD-7 reflecting the upper end of mild severity compared to the usual clinical thresholds for mild, 402 moderate and severe anxiety of 5, 10 and 15 on the GAD-7 (Spitzer et al, 2006). However 95 per cent 403 of the sample had moderate or severe symptoms of GAD at baseline, well above the minimum 404 threshold for entry to the study and the national NHS IAPT criteria for remission. They had already 405 failed to improve with facilitated bibliography or computerised psychological treatment for GAD_L-so 406 spontaneous improvement was unlikely. Placebo responses are less frequent frequent in research 407 participants with less severe anxiety or depression and in those who have not responded to previous 408 active treatment for their condition (Stein et al, 2006; Weimer et al, 2015). Therefore the study 409 shows the effectiveness of CES in a clinical treatment seeking sample of patients with moderate to 410 severe treatment resistant generalised anxiety disorder(.
- There are important limitations of the study. There was no control group and the study was not a
 randomised controlled trial. However meta-analysis of previous RCTs of active CES versus sham CES
 already provides evidence that CES is effective in treating anxiety and depression symptoms
 (Shekelle et al, 2018). The United Kingdom Medical Research Council (2000) and National Institute
 for Health and Care Excellence (2018) recommend that implementation studies are completed in
- 416 routine treatment settings to check that the efficacy seen in RCTs is translated into routine clinical
- 417 practice settings. This study was therefore designed to meet this requirement, to examine if

effectiveness is maintained after CES treatment completion, and if there were any cost savings from
CES treatment. Such studies do not necessarily utilise control groups; they must enrol treatment
seeking patients studied under routine care delivery. Alpha-Stim CES was more effective at achieving
remission than we expected from the effect size in a meta-analysis of RCTs (Shekelle et al, 2018)
with 44.7% patients achieving remission, comparable to iCBT in routine treatment settings, rather

423 than 26.5% patients as we had planned.

The sample recruited only 22 per cent of those eligible to take part in the study. However, the offer to take part in this research and to receive this treatment came through cold calling by the clinical <u>team through letter, e-mail or telephone call</u>. If participants were prepared for the possibility of receiving CES by the IAPT services then uptake of CES might be higher. <u>A strength of cold calling and</u> <u>lack of research team contact is that placebo responses to CES may have been low because of</u> <u>infrequent contact of the research team so that the effectiveness of CES in the study was not</u> inflated compared to clinical practice.

431

432 Another limitation of the study was that the sample lacked ethnic diversity. The sample was drawn

433 from all ages although there were greater proportions of younger and middle aged participants in

the study, reflecting the composition of age groups in routine IAPT NHS services. As expected the

435 vast majority of patients with GAD were female. There was a broad representation of education,

436 marital status and employment status reflecting the age composition of the sample.

There was a high degree of attrition of the study to follow up with the loss of 55.2% by 24 weeks

- despite financial incentive to provide data as opposed to 40% that we had anticipated. The study
- 439 was adequately powered because CES was more effective than we had expected. The results are
- similar between the ITT sample with imputed results and those completing all follow up assessmentssuggesting that the conclusions drawn from the whole sample using imputation are probably safe to
- suggesting that the conclusions drawn from the whole sample using imputation are probably safe to
 make. We also only have a limited amount of information on the reasons that participants withdrew
- from CES or follow up. The most common reason given for withdrawal from CES is not being able to
- find the time to use CES for 60 minutes per day. <u>The CES device was also not locked so some</u>
- 445 participants may have used a higher current than we instructed them to and got adverse effects that
- 446 <u>they chose not to report. We have no evidence that anyone did this.</u> Almost as many dropped out of
- 447 CES because it had worked as those who stopped because it did not. A limitation of the health
- economics analysis is that we did not consider the possibility that CES might have reduced the delay
- 449 in receiving iCBT by freeing up capacity in other CBT therapists or that those patients who received
- 450 both CES and iCBT might have had fewer iCBT sessions. Therefore cost savings from CES may be
- 451 underestimated in treatment settings offering iCBT for GAD.

452 We did not personalise CES to each individual. It is possible that different waveforms of current,

453 stimulus intensity and stimulation location might have been more efficacious for some participants

454 (Guleyupoglu et al, 2013). Some participants may have tolerated 5 days of treatment with CES per

455 week better than 7 days per week with higher completion rates of 6-12 weeks CES treatment.

456 As well as improvements in anxiety, there were improvements in depression and insomnia, two

457 other potential indications for CES. Although the results are encouraging, further research is needed

in patients with primary depression and primary insomnia disorders. There were also high remission,

- 459 recovery and reliable improvement rates in GAD-7 score when participants received both iCBT and
- 460 CES in the first 12 weeks. Research might explore if higher and more sustained rates of remission are
- in generalised anxiety disorder in trials of iCBT plus active CES versus iCBT plus sham CES.

- 462 In conclusion, we provide evidence that CES may be clinically effective and cost reducing during
- administration and for three months afterwards in routine treatment settings offering psychological
- treatments for moderate to severe GAD. CES improves the efficiency of these services, a critical issue
- because of the shortage and high turnover of psychological treatment staff, allowing them to reach
- their targets for remission with fewer highly skilled staff. As a result, it is also cost saving to such
- services even when a range of different assumptions are made about the delivery of psychological
- 468 treatment.
- 469 5,268 words

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474 **Conflict of interest**

- 475 The chief investigator (RM) and MC report no financial or other conflicts of interest for their
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- 479

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610 Table 1. Procedure and assessments in the study (n=161)

ASSESSMENT	VISIT 1 BASELINE	VISIT 2 WEEK 4	VISIT 3 WEEK 6	VISIT 4 WEEK 8	VISIT 5 WEEK 12	VISIT 6 WEEK 24
CONSENT	Х					
TRAINING TO USE CES	Х					
PREGNANCY TEST	X (*)					
GAD-7	Х	Х	Х	Х	Х	Х
EQ-5D-5L	X	Х	Х	Х	Х	Х
CSRI	Х				Х	Х
WASA	X	Х	Х	Х	Х	Х
PHQ-9	Х	Х	Х	Х	Х	Х
AIS	X	Х	Х	Х	Х	Х
ALPHA-STIM CES	Ongoing	Ongoing	Ongoing (**)	Ongoing (**)	Ongoing (**)	
ADHERENCE		Х	Х	X (**)	X (**)	
ADVERSE EVENTS		Х	Х	X (**)	X (**)	

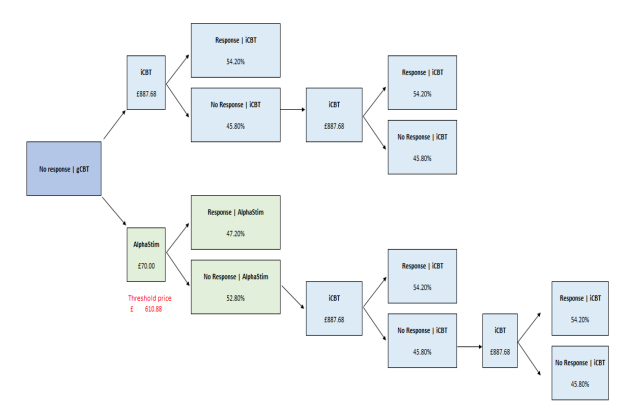
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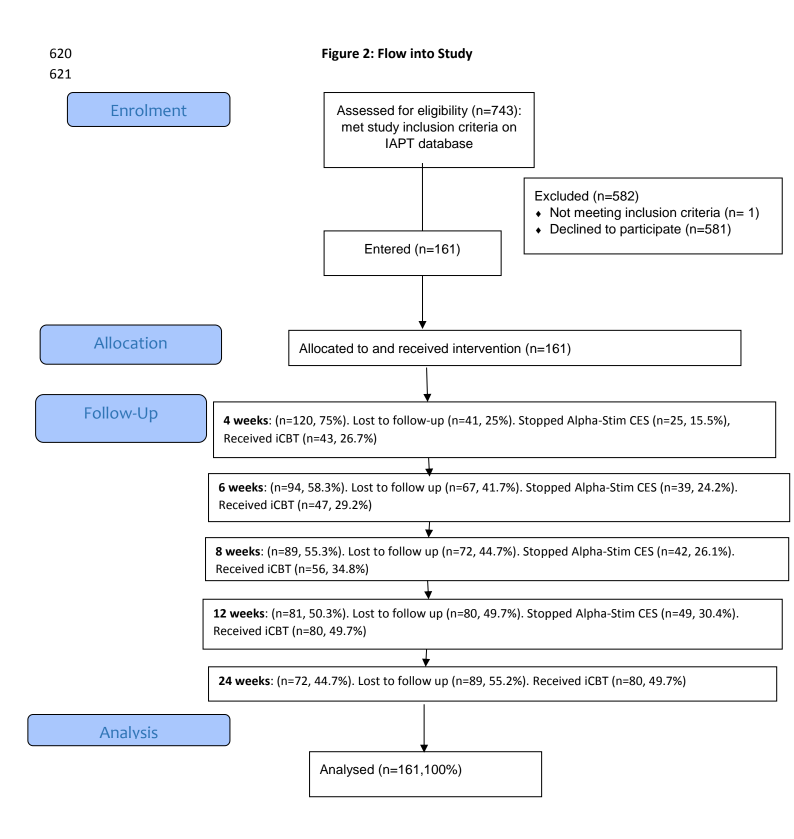
612 (*) If a female of child-bearing potential

613 (**) If continuing with Alpha-Stim AID CES treatment between week 6 – week 12.

615 Figure 1. Decision Tree model for comparison of Alpha-stim CES pathway with individual cognitive

behaviour therapy (iCBT) treatment as usual.





622 Table 2. Baseline characteristics of participants (n=161).

Variable	Mean (sd) or n(%)
Age, years	38.00 (14.2) (min=18, max=76)
Gender, female	118 (73.3%)
Ethnicity, white British	153 (95.0%)
Marital status: Married or cohabiting	95 (59.0%)
Single	50 (31.1%)
Divorced	12 (7.5%)
Widowed	4 (2.5%)
Education: No qualifications	5 (3.1%)
GCSE (left school at 16 years)	39 (24.2%)
A level or other non-degree higher qualification	67 (41.6%)
Degree	50 (31.1%)
Employment: Employed	106 (65.8%)
Unemployed	33 (20.5%)
Retired	11 (6.8%)
Student	7 (4.3%)
Homemaker	4 (2.5%)
GAD-7	15.77 (3.21)
PHQ-9	16.07 (4.94)
Athens Insomnia Scale	12.91 (4.82)
WASA	20.81 (7.74)
EQ-5D-5L	51.61 (19.0)

Table 3: Intention to treat analysis of remission, reliable improvement, recovery and mean (sd)

626 continuous outcomes with Alpha-stim CES at 12 and 24 weeks (n=161).

Outcome		Remissio 12 week n (%)		Reliable improve 12 weeks n (%)	Recovery 12 weeks n (%)	Remission 24 weeks n (%)	Reliable improve 24 weeks n (%)	Recovery 24 weeks n (%)
GAD-7								
Overall, n=161	L	72 (44.7)	102 (63.4)	72 (44.7)	77 (47.8)	105 (65.2)	77 (47.8)
No CBT, n=81		49 (60.5)	67 (82.7)	49 (60.5)	53 (65.4)	70 (86.4)	53 (65.4)
PHQ-9								
Overall, n=161	.61 73 (45.3))	76 (47.2)	61 (37.9)	82 (50.9)	80 (49.7)	67 (41.6)
GAD-7 and PH	IQ-9							
Overall, n=161	L	62 (38.5)		75 (46.6)	54 (37.5)	69 (42.9)	75 (46.6)	59 (36.5)
AIS								
Overall, n=161	L	39 (24.2)	53 (32.9)	37 (23.0)	45 (28.0)	60 (37.3)	43 (26.7)
		Normal		Functional		Normal	Functional	
		Function	า	recovery		Function	recovery	
		12 weeks n (%)		12 weeks n (%)		24 weeks n (%)	24 weeks n (%)	
WASA								
Overall, n=161		28 (17.4)	43 (26.7)		29 (18.0)	48 (29.8	
Outcome	Base	line	4 v	veeks	6 weeks	8 weeks	12 weeks	24 weeks
GAD-7 ¹	15.77	7 (3.21) 10		.14 (4.86)	9.73 (4.89)	9.34 (4.58)	8.92 (5.42)	8.99 (6.18)
PHQ-9 ²	16.07	7 (4.94) 11		.22 (6.09)	10.38 (5.91)	10.04 (6.46)	8.91 (5.78)	10.42 (6.97)
AIS ³	12.91	1 (4.82) 10		.27 (5.27)	10.18 (5.20)	9.72 (5.16)	8.81 (4.86)	7.94 (4.62)
WSAS ⁴	20.81	81 (7.74) 18		.27 (8.89)	16.95 (9.56)	15.94 (9.22)	14.89 (9.99)	15.98 (9.18)
EQ-5D-5L⁵	51.61 (19.00)		57.90 (20.15)		61.00 (20.47)	62.99 (21.08)	64.80 (21.72)	62.50 (22.97)

627 ¹ Effect of treatment over time significant F =88.12, df1=5.0/df2=156.0, p < .001, partial Eta square = 0.74 (large); within

628 subjects effect over time significant F=72.02, df1=3.7/df2=563.74, p < .001, partial Eta square = 0.31 (medium)

 2 Effect of treatment over time significant F=28.38, df1=5.0/df2=156.0, p < .001, partial Eta square = 0.48 (medium); within

630 subjects effect over time significant F=42.89, df1=3.9/df=559.01, p < .001, partial Eta square = 0.21 (small)

631 ³ Effect of treatment over time significant F=40.85, df1=5.0/df2=156.0, p < .001, partial Eta square = 0.57 (large); within

632 subjects effect over time significant F=42.69, df1=3.8/df=542.9, p < .001, partial Eta square = 0.21 (medium)

 4 Effect of treatment over time significant F=17.18, df1=5.0/df2=156.0, p < .001, partial Eta square = 0.36 (medium); within

634 subjects effect over time significant F=17.35, df1=3.5/df=557.45, p < .001, partial Eta square = 0.10 (small)

⁵ Effect of treatment over time not significant F=16.11, df1=5.0/df2=156.0, p < .001, partial Eta square = 0.34 (medium);

636 within subjects effect over time significant F=13.94, df1=4.1/df2=651.3, p < .001, partial Eta square = 0.08 (small)

637

638

640 Table 4: Costs and responses of Alpha-Stim CES in relation to the eight session standard care

641 model of CBT

	Deterministic	Probabilistic	Distribution	Alpha	Beta	N int	N control
Cost of							
Individual CBT				£			
	£887.68	£923.64		887.68			
			Gamma		1		
Probability of			Gamma		1		
Response to							
Individual CBT							
	54.2%	56%	Beta	199.46	168.54	368	679
Patients per	0	20/0			200.01		
Alpha-Stim CES							
lifetime							
	5.00	5.41	Gamma	5	1		
Per patient cost							
of Alpha-Stim							
CES	£70.00	£64.75	Calculated				
Probability of							
Response to							
Alpha-Stim CES							
	47%	39%	Beta	45	55		
	Expected	Lower	Upper	Expec	ted	Lower	Upper
	Cost	95% CI	95% CI	Responses		95% CI	95% CI
iCBT only	£1,294,233	£1,198,677	£1,392,923	701.68		650.29	751.85
AlphaStim	£753,355	£651,653	£981,087	889.24		860.29	907.14
Net	-£540,878	-£648,692	-£327,117	187.56		141.03	227.82