

TINNITUS

Bimodal neuromodulation combining sound and tongue stimulation reduces tinnitus symptoms in a large randomized clinical study

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Tinnitus is a phantom auditory perception coded in the brain that can be bothersome or debilitating, affecting 10 to 15% of the population. Currently, there is no clinically recommended drug or device treatment for this major health condition. Animal research has revealed that sound paired with electrical somatosensory stimulation can drive extensive plasticity within the brain for tinnitus treatment. To investigate this bimodal neuromodulation approach in humans, we evaluated a noninvasive device that delivers sound to the ears and electrical stimulation to the tongue in a randomized, double-blinded, exploratory study that enrolled 326 adults with chronic subjective tinnitus. Participants were randomized into three parallel arms with different stimulation settings. Clinical outcomes were evaluated over a 12-week treatment period and a 12-month posttreatment phase. For the primary endpoints, participants achieved a statistically significant reduction in tinnitus symptom severity at the end of treatment based on two commonly used outcome measures, Tinnitus Handicap Inventory (Cohen's *d* effect size: -0.87 to -0.92 across arms; $P < 0.001$) and Tinnitus Functional Index (-0.77 to -0.87 ; $P < 0.001$). Therapeutic improvements continued for 12 months after treatment for specific bimodal stimulation settings, which had not previously been demonstrated in a large cohort for a tinnitus intervention. The treatment also achieved high compliance and satisfaction rates with no treatment-related serious adverse events. These positive therapeutic and long-term results motivate further clinical trials toward establishing bimodal neuromodulation as a clinically recommended device treatment for tinnitus.

INTRODUCTION

The brain is a complex and robust network of cells that mediates our everyday functions from sensations and movements to consciousness and learning. In certain circumstances, the brain enters deviant states that can lead to debilitating or even life-threatening conditions (1–5). Neuromodulation has emerged as a promising approach to treat many different brain disorders, in which electrical or magnetic stimulation of the nervous system can be used to modulate aberrant neurons to improve a patient's symptoms (6–10). One widely used approach applies surface electrical stimulation on the scalp or body, including activation of the trigeminal or vagus nerve, to treat a wide range of health conditions, such as tinnitus, seizures, depression, anxiety, memory loss, inflammation, and pain (6–9, 11–16). There is a growing consensus that this type of stimulation activates or modulates a broad area of the brain; however, greater or more consistent efficacy may be achieved through electrical stimulation paired

with a targeted input to modulate specific brain regions for improving symptoms (7, 8, 17–23). In particular, electrical stimulation applied to the scalp leads to current that spreads across the skull and activates a broad volume of the brain, or noninvasive electrical stimulation of a peripheral nerve can excite widespread projections throughout the brain that broadly activate multiple interconnected neural regions. A targeted input (e.g., a pure tone or movement of a finger) can be designed to activate a local population of neurons that overlaps with the broad activation caused by surface electrical stimulation; thus, this coordinated activation by both modalities can lead to enhanced modulation or paired plasticity within just the overlapping localized region (7, 8, 17–23).

Paired stimulation has recently shown promise for treating a brain condition known as tinnitus, which is a phantom sound perception that occurs without any external acoustic input. Tinnitus can be bothersome or debilitating for 10 to 15% of the population and is a major health issue in society (24–27). There is still no clinically recommended device or drug treatment for this condition (28, 29). Animal research and pilot human studies have shown that bimodal neuromodulation combining sound and electrical somatosensory stimulation (30), including tongue stimulation (17, 31), can drive extensive plasticity in the auditory system and improve tinnitus symptoms. Electrical somatosensory stimulation can broadly activate or modulate neurons throughout the auditory pathway, from the brainstem up to the auditory cortex, and in emotional and attentional centers (13, 17, 30, 32–40). Pairing specific acoustic and somatosensory stimuli can drive enhanced modulation within brain regions linked to tinnitus (17, 30, 31), in which repeating stimulation of two inputs can lead to long-term therapeutic plasticity (7, 8, 17, 20, 22, 23).

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There is a globally recognized clinical need for new treatments for tinnitus because of the lack of effective options for this debilitating condition. On the basis of the positive efficacy and safety outcomes in a previous human pilot study presenting sound and tongue stimulation to treat tinnitus (31), along with supporting animal data (17), we pursued a double-blind randomized clinical study (ClinicalTrials.gov, NCT02669069) (41) to further evaluate different parameter settings of bimodal neuromodulation in a broad tinnitus population. Bimodal stimulation was delivered using a medical device (referred to as MBT in the study; marketed as Lenire) that pairs sound with electrical stimulation of the tongue (Fig. 1A). The main objective of this exploratory study was to build further supporting data for the safety and efficacy of bimodal neuromodulation. More specifically, the study aimed to evaluate the effects of three different parameter settings of bimodal stimulation on tinnitus symptoms, both in the general tinnitus population and in subtypes of patients that may be differentially responsive to the tested parameters. A large number of tinnitus participants were screened, randomized, and enrolled into one of three study arms (Fig. 1B).

One challenge with these types of studies is the large range of parameters that can be evaluated for a treatment paradigm such as our bimodal neuromodulation, including types of acoustic stimuli, electrical pulse patterns and their location on the tongue, and durations and timings of stimulation. On the basis of previous animal and human research (30, 42), effective therapeutic benefit with bimodal neuromodulation may be achieved by implementing a specific delay between sound and electrical somatosensory stimulation, as well as matching the frequency spectrum of the acoustic stimulus to the tinnitus percept. Other studies in animals and patients with tinnitus (17, 31, 43) suggest that a broad spectrum of acoustic frequencies and/or a range of delays between sound and electrical somatosensory stimulation can be effective in driving extensive plasticity within auditory regions to alleviate tinnitus symptoms. Considering the parameters that could be effective for bimodal neuromodulation, attempting to investigate the contribution of each parameter with a sufficient number of subjects would require thousands of participants. Our strategy was to reduce the parameter space into three groups, in which several stimulation parameters differed between groups. The advantage of this approach is that

multiple parameters can be explored with fewer treatment arms and a smaller sample size. The disadvantage of this approach is that if a significant difference is observed between the treatment groups, then distinguishing the main determinant of the therapeutic effect would require a follow-up study. Because this is the first in a series of planned clinical studies exploring the parameter space of bimodal neuromodulation for tinnitus treatment, we pursued this three-arm study design to initially narrow in on a smaller and more manageable parameter space that would then be systematically evaluated in future clinical trials.

The study was conducted at two clinical sites: the Wellcome Trust—Health Research Board Clinical Research Facility, St. James's Hospital, Dublin, Ireland and the Tinnituszentrum of the University of Regensburg, Germany. Tinnitus participants were provided and instructed to use a bimodal neuromodulation device for 60 min daily for 12 weeks. They were clinically evaluated at 6 and at 12 weeks during the treatment period (Fig. 1B). When treatment was completed, participants were assessed at three follow-up visits up to 12 months after returning their device. The primary endpoints of the study composed of within-arm and between-arm comparisons of tinnitus symptom severity during treatment. The outcome measures were two widely used and validated instruments to assess changes in tinnitus symptom severity: Tinnitus Handicap Inventory (THI) and Tinnitus Functional Index (TFI) (44–47). Long-term changes in tinnitus symptom severity were also assessed during the 12-month posttreatment phase. Exploratory analyses described in the published protocol paper (41), including the effects of different stimulation parameters on subtypes of tinnitus participants and comparisons of alternative methods for measuring the impact of tinnitus, will be completed and presented in a subsequent publication.

RESULTS

Characteristics and summary of study participants

As previously described in the published protocol paper (41), study participants were recruited to the clinical trial using media advertising and directed to an online eligibility assessment. Eligibility was initially assessed through a set of general prescreening questions to manage the large number of candidates expected to respond to the advertising (48). As anticipated, a substantial number of candidates (7627) registered their interest in the trial. Individuals (5826) completed the online assessment, and 698 were invited for screening at a clinical site. On the basis of the inclusion criteria, 333 individuals were randomized into the three study arms (1:1:1), with 326 successfully completing enrollment and device fitting. Figure 2 presents the participant flow diagram for the study.

Enrolled participants were randomized into the three treatment arms according to four binary categories of stratification (41) using the method of minimization (49), as shown in Table 1. The four stratification categories included atonal tinnitus, hyperacusis, normal hearing, and noise-induced hearing

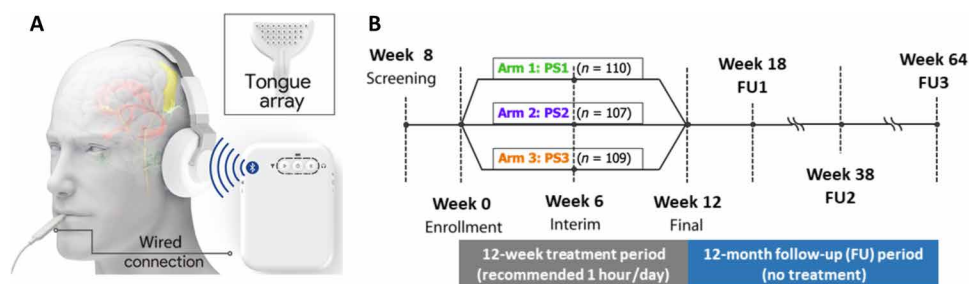


Fig. 1. Treatment approach and study design. (A) Schematic of CE-marked bimodal neuromodulation device (marketed as Lenire) developed by Neuromod Devices (Dublin, Ireland) consisting of wireless (Bluetooth) headphones that deliver acoustic stimuli, a wire-connected 32-electrode array for presenting electrical stimulation patterns to the anterior dorsal surface of the tongue, and a battery-powered controller that coordinates both stimulus modalities. Bimodal stimulation activates auditory and somatosensory pathways and converging centers in auditory, limbic, and attentional brain regions implicated in tinnitus. (B) Schematic of the clinical study design in which 698 participants were screened at the clinic to enroll 326 eligible participants into one of the three treatment arms. Participants were evaluated for changes in tinnitus symptom severity during the 12-week treatment period and at three follow-up (FU) visits up to 12 months after treatment.

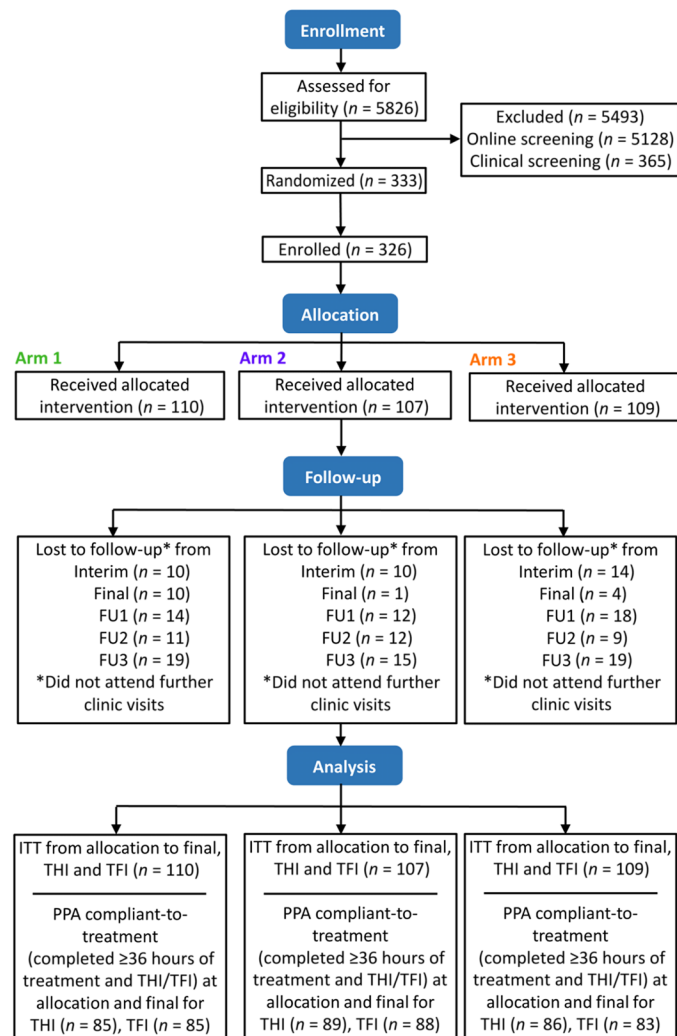


Fig. 2. Participant flow diagram. For the primary endpoints, within-arm comparisons were performed with a per-protocol analysis (PPA) for those who were compliant to treatment (device usage of at least 36 hours over the 12-week treatment period), whereas between-arm comparisons were performed with an intention-to-treat (ITT) analysis with imputation for missing values described in Materials and Methods. Follow-up visits are depicted in Fig. 1.

loss. The participants were distributed across treatment arms with no significant differences ($P > 0.05$) in relation to the four stratification categories. There were also no significant differences ($P > 0.05$) across arms for the other characteristics and groups shown in Table 1. In addition to the stratification categories, the “other groups” listed in Table 1 were intentionally included in the study design to perform exploratory analyses and identify different subtypes of patients with tinnitus who could be differentially responsive to certain bimodal stimulation parameters. These subtyping results are outside the scope of this paper and will be presented in a subsequent publication.

As listed in Table 1, 84.0% of enrolled participants returned for their final visit at the end of their 12-week treatment. The high retention rate is mirrored in the high treatment compliance rate of 83.7% (fig. S1). Minimum treatment compliance was defined as device usage of at least 36 hours over the intended 12-week treatment period, which was specified in the design of the clinical trial (41)

and derived from a previous bimodal neuromodulation study (31). The device logged the time and date of daily usage by each participant. There were no significant differences in the compliance rate or number of participants at the final visit between treatment arms ($P > 0.05$; fig. S1). Participants also generally attended their visits at the intended time points specified in the study design (fig. S2).

Overall, there were no major differences in characteristics across study arms that would bias the primary endpoint analyses. There were also a high number of treatment-compliant participants that could be included into the primary endpoint analyses.

Within-arm and between-arm therapeutic effects during treatment

Each participant was fitted with a device delivering one of three bimodal stimulation settings (PS1, PS2, and PS3; Fig. 1B). In a previous pilot study (31), paired sound and electrical tongue stimulation provided significant improvements in tinnitus symptoms. To explore which stimulation parameters contribute to the therapeutic effects, three different stimulation settings were designed for this study. All stimuli were presented at suprathreshold intensities, as in the previous pilot study, which enabled blinding of participants and researchers to the treatment arms. Details and schematics of the different stimulation settings are provided in Materials and Methods and table S1.

PS1 consisted of a wide range of pure tones (500 to 8000 Hz) synchronized in time with a burst of electrical pulses delivered to the tongue surface. Because a wideband noise component was included in the paired stimuli used in the previous pilot study, the acoustic stimulus in PS1 also incorporated a broadband noise background. Previous animal research suggests that there may be a specific delay or range of delays between sound and somatosensory stimulation for driving greater improvements in tinnitus (17, 30, 42). However, it is not yet known how the effective delays identified in animals translate to humans. Presenting two stimuli synchronously is consistent with the concept of paired plasticity (7, 8, 18–21), where a person can perceptually integrate and associate meaning between two inputs that are physically presented at the same time (50, 51). In keeping consistent with the paired plasticity approach, we presented acoustic and tongue stimuli at the same time for PS1. PS2 was similar to PS1, except that there were short interstimulus delays between sound and tongue stimulation that were randomly distributed between 30 and 50 ms. PS1 also had a fixed relationship between pure-tone frequencies and stimulation locations on the tongue array, whereas PS2 did not have a fixed tone-to-tongue mapping. The rationale for PS2 was to assess whether therapeutic effects required a specific acoustic feature to be systematically and synchronously mapped to a specific tongue location for a given session, as occurred in the previously published pilot study (31). The 30- to 50-ms range was selected so that the onset of the electrical pulses fell midway between the onset of the preceding and proceeding acoustic stimuli (i.e., for a repetition period of ~80 ms). Considering that an offset of 30 to 50 ms may not be sufficient to disrupt paired plasticity on a higher cortical or cognitive level that could function on longer time scales, greater interstimulus delays (550 to 950 ms) were introduced in PS3. These longer delays also required a longer repetition period, which was set to ~2 s for PS3 and resulted in a stimulation rate that was about 25 times less than the rate used for PS1 and PS2. To further assess whether the frequency range of acoustic stimuli influenced therapeutic effects, only low-frequency pure tones

Table 1. Characteristics of enrolled participants and key numbers for each visit. dB HL, decibel hearing level; dB SL, decibel sensation level (equals dB HL minus hearing threshold level of acoustic stimulus); FU, follow-up; LDL, loudness discomfort level; MML, minimum masking level. Somatic tinnitus is defined as a tinnitus that can be directly modulated by one or more head or neck maneuvers. MML was determined by presenting a wideband noise at increasing intensities until the sound masked the participant's tinnitus. A description of the stratification categories is provided in Materials and Methods. *P* value is based on the Fisher's exact test for count variables or a linear regression for continuous variables.

Characteristics	Units	Full cohort	Arm 1	Arm 2	Arm 3	<i>P</i> value
Total participants enrolled (device fitted)	# Participants	326	110	107	109	0.990
Ireland: participants enrolled (device fitted)	# Participants	261	86	85	90	0.975
Germany: participants enrolled (device fitted)	# Participants	65	24	22	19	0.883
Gender: Male	# Participants (% of enrolled)	212 (65.0%)	65 (59.1%)	79 (73.8%)	68 (62.4%)	0.694
Gender: Female	# Participants (% of enrolled)	114 (35.0%)	45 (40.9%)	28 (26.2%)	41 (37.6%)	0.351
Age at baseline	Years [mean (SD)]	48.1 (11.6)	46.3 (12.0)	48.7 (12.6)	49.5 (10.0)	0.102
Tinnitus duration	Years [mean (SD)]	2.6 (1.6)	2.5 (1.7)	2.7 (1.6)	2.6 (1.6)	0.777
THI at baseline	Points [mean (SD)]	43.5 (13.2)	44.5 (14.2)	44.0 (12.7)	42.0 (12.6)	0.325
TFI at baseline	Points [mean (SD)]	47.9 (14.7)	47.7 (14.5)	48.1 (14.9)	47.7 (14.9)	0.969
Mean hearing loss (250 Hz to 8 kHz) at screening	dB HL [mean (SD)]	17.8 (10.3)	17.2 (9.9)	18.1 (9.7)	18.2 (11.2)	0.699
Attended interim visit	# Participants (% of enrolled)	277 (85.0%)	97 (88.2%)	92 (86.0%)	88 (80.7%)	0.898
Attended final visit	# Participants (% of enrolled)	274 (84.0%)	89 (80.9%)	94 (87.9%)	91 (83.5%)	0.964
Attended FU1 visit	# Participants (% of enrolled)	185 (56.8%)	59 (53.6%)	70 (66.4%)	56 (51.4%)	0.620
Attended FU2 visit	# Participants (% of enrolled)	183 (56.1%)	57 (51.8%)	69 (64.5%)	57 (52.3%)	0.716
Attended FU3 visit	# Participants (% of enrolled)	156 (47.9%)	46 (41.8%)	57 (53.3%)	53 (48.6%)	0.760
Stratification						
Atonal tinnitus	# Participants (% of enrolled)	50 (15.3%)	18 (16.4%)	16 (15.0%)	16 (14.7%)	1.000
Hyperacusis (LDL <60-dB SL at 500 Hz)	# Participants (% of enrolled)	66 (20.2%)	19 (17.3%)	21 (19.6%)	26 (23.9%)	0.743
Normal hearing (≤20-dB HL, 250 Hz to 8 kHz)	# Participants (% of enrolled)	78 (23.9%)	26 (23.6%)	26 (24.3%)	26 (23.9%)	1.000
Noise-induced hearing loss	# Participants (% of enrolled)	107 (32.8%)	35 (31.8%)	37 (34.6%)	35 (32.1%)	1.000
Other groups						
Somatic tinnitus (≥1 maneuver)	# Participants (% of enrolled)	201 (61.7%)	67 (60.9%)	63 (58.9%)	71 (65.1%)	0.906
Substantial MML (≥15-dB SL)	# Participants (% of enrolled)	240 (73.6%)	80 (72.7%)	84 (78.5%)	76 (69.7%)	0.902
Tinnitus duration (<3 years)	# Participants (% of enrolled)	170 (52.1%)	58 (52.7%)	54 (50.5%)	58 (53.2%)	0.953

(100 to 500 Hz) were presented in PS3. Similar to PS2, PS3 did not have a fixed tone-to-tongue mapping.

During device fitting, sound volume was adjusted to each participant's hearing thresholds, and electrical stimulation intensity was adjusted to each participant's tongue sensitivity. Hearing thresholds across all enrolled participants for each arm are shown in fig. S3. Treatment was self-administered by participants at home using the device for a recommended 60 min/day with a minimum compliance criterion of 36 hours over the full treatment period. The device was returned at the 12-week visit, and participants were assessed at three follow-up visits over 12 months.

As described in the previously published protocol paper (41), the primary endpoints composed of: (i) within-arm changes in the THI and TFI outcome measures from baseline (average of scores from screening and enrollment visits) to the end of treatment for participants who were treatment-compliant (i.e., a per-protocol analysis in which participants used the device for at least 36 hours during the intended 12-week period) and (ii) between-arm differences in the changes in THI and TFI outcome measures from baseline to the end of treatment for the full cohort of participants (i.e., an intention-to-treat analysis where missing data points were estimated using a multiple imputation method). Both THI and TFI are questionnaires with global scores from 0 to 100, with a higher score indicating greater tinnitus symptom severity.

To initially visualize the therapeutic effects experienced by each participant, Fig. 3 displays a scatterplot for each treatment arm. The final THI or TFI score for each treatment-compliant participant is

plotted against the baseline score. Most of the participants exhibited an improvement in THI or TFI during treatment, ranging between 74.7 and 88.8% across arms. There was also a trend toward greater improvement in THI or TFI for those who had worse tinnitus symptoms at baseline, with many participants achieving large improvements (e.g., a decrease in 20 or 30 points; fig. S4). These positive therapeutic effects were observed for all three stimulation settings.

For the primary endpoints of the study, Fig. 4 presents the averaged results across participants for the within-arm and between-arm comparisons in a forest plot format. All within-arm comparisons, for both THI and TFI, were highly significant on the basis of paired two-tailed *t* tests ($P < 0.001$). Within-arm primary endpoints were successfully achieved and statistically significant when accounting for multiple comparisons. Arms 1, 2, and 3 exhibited a decrease of 14.6, 14.5, and 13.5 points in THI and 13.9, 13.8, and 13.2 points in TFI, respectively. In terms of Cohen's *d* effect sizes and 95% confidence intervals (CIs), arms 1, 2, and 3 exhibited values of -0.92 [$-1.14, -0.71$], -0.87 [$-1.08, -0.67$], and -0.92 [$-1.16, -0.69$] for THI and -0.87 [$-1.10, -0.65$], -0.77 [$-0.98, -0.56$], and -0.78 [$-1.05, -0.51$] for TFI, respectively. We did not observe a significant difference between the arms during the treatment phase [$P > 0.05$ for between-arm comparisons; analysis of covariance (ANCOVA): $F_{3, 270} = 58.45$, $P > 0.05$; description and justification of statistical analyses are provided in Materials and Methods].

Long-term treatment outcomes

The study was designed to track changes in tinnitus symptom severity at multiple time points during treatment and up to 12 months after the end of treatment. Figure 5 plots the mean change in THI and TFI relative to baseline value for each assessment visit for the three study arms. There were three interesting observations from the data presented in Fig. 5. First, all three stimulation settings resulted in rapid improvement within the first 6 weeks of treatment. Second, there were minimal improvements in the second 6 weeks of treatment, which may be attributed to habituation processes in the brain. Third, improvements were significantly sustained out to 12 months after treatment ($P < 0.05$; 95% CI does not cross the zero line), except for arm 3. There were also a greater proportion of participants retaining an improvement 12 months after treatment for arms 1 and 2 compared to arm 3 (although not significant based on Fisher's exact test, $P > 0.05$; figs. S5 and S6), which corresponded to greater sustained effects for arms 1 and 2 compared to arm 3 (significant differences between arms are listed in Fig. 5). These long-term findings suggest that bimodal neuromodulation using higher-frequency tones with synchronized or shorter delayed tongue stimulation could be more effective in delivering a sustained therapeutic benefit in a higher percentage of patients with tinnitus compared to bimodal neuromodulation with low-frequency tones and long delayed tongue stimulation. Because there were no apparent differences in outcomes between arms 1 and 2, the fixed tone-to-tongue mapping is not necessary to drive therapeutic effects. Arms 1 and 2 delivered a greater number of stimulus bursts per second than in arm 3, which could have also contributed to the differences in long-term outcomes.

Previous studies have shown that many people with tinnitus have high-frequency hearing loss and a dominant tinnitus pitch that falls within the compromised spectral region (25, 52, 53). Presenting sounds with high-frequency components and/or matched

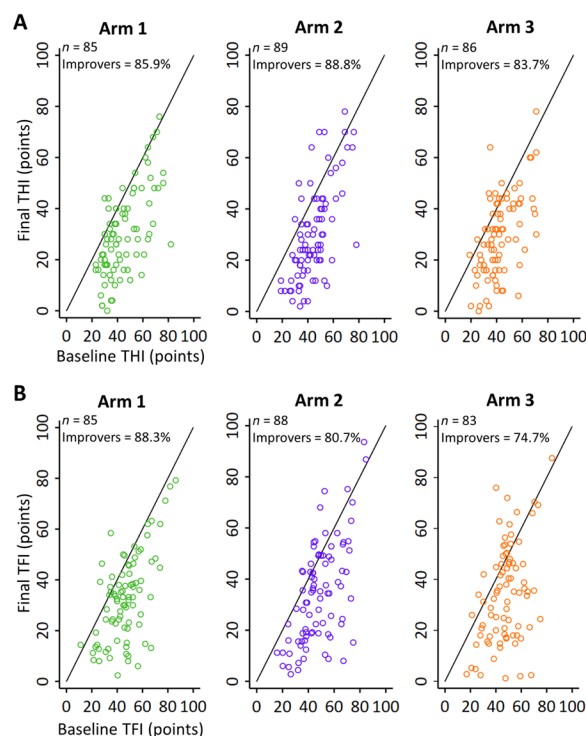


Fig. 3. Changes in tinnitus symptom severity for each participant. (A) Scatterplots represent THI scores for each treatment-compliant individual at baseline (average of score at screening and enrollment) versus the end of treatment (12-week final visit) for each arm. (B) Scatterplots represent TFI scores. Points below diagonal black line indicate an improvement (i.e., a decrease in score) in outcome measure of THI or TFI. Data points were jittered for visibility.

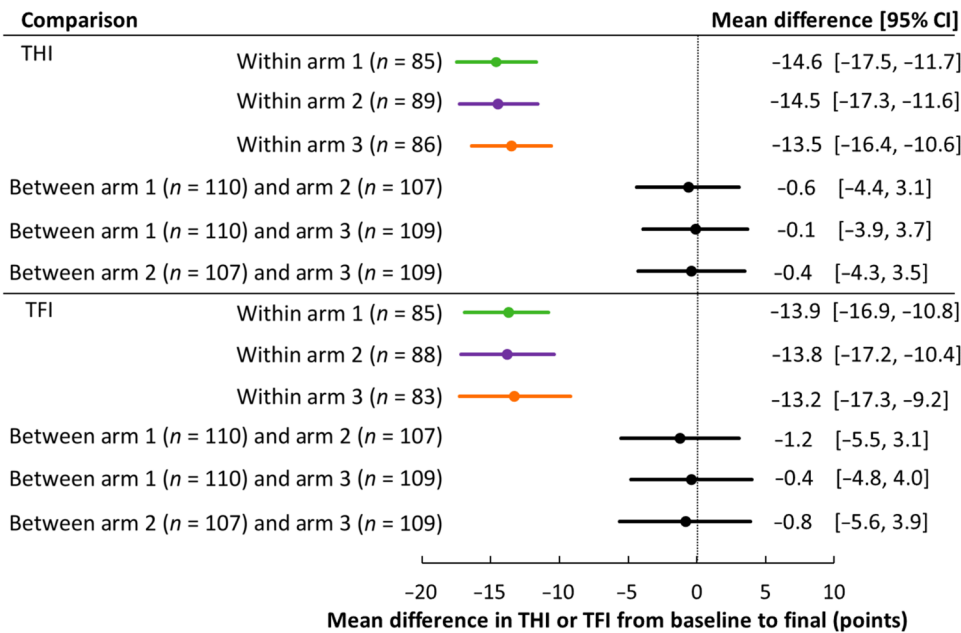


Fig. 4. Within-arm and between-arm analyses for primary endpoints at 12-week assessment. The mean difference in THI or TFI score across participants for each arm from baseline to the final assessment (end of treatment) for the within-arm cases or the mean difference between two arms for the change in THI or TFI score from baseline to the final assessment for the between-arm cases is plotted with the $\pm 95\%$ CI. For the primary endpoints, the within-arm statistical analyses were based on a per-protocol estimand, which consists of all treatment-compliant participants with baseline and final scores. All within-arm comparisons for THI and TFI were highly significant on the basis of paired two-tailed *t* tests ($P < 0.001$; 95% CI are nonoverlapping with the vertical line at zero). The between-arm statistical analyses were based on an intention-to-treat estimand and tested with multiple regression using baseline scores as a covariate. Missing data were handled by using a Markov chain Monte Carlo multiple imputation method (further details are provided in Materials and Methods), which leads to *n* values that match the enrolled numbers for each arm. There were no significant differences between any arms ($P > 0.05$; 95% CI crosses zero line).

to the dominant tinnitus pitch has been proposed as an effective method for treating tinnitus, both for sound-only approaches and for bimodal neuromodulation (30, 54–56). Alleviating the tinnitus percept has also been achieved through cochlear implant stimulation by reactivating compromised hearing pathways (57, 58). To further assess the possible contribution of high-frequency tones in the long-term therapeutic effects shown in Fig. 5, we plotted the change in THI or TFI as a function of high-frequency hearing loss (averaged across 4 to 8 kHz; figs. S7 and S8). If high-frequency tones are critical, especially because many of our participants had high-frequency hearing loss across arms, we would expect less therapeutic benefit for those with greater high-frequency hearing loss, particularly for arm 3 where PS3 consisted of only low-frequency sounds (~100 to 500 Hz). We did not observe any relationship between therapeutic effects and high-frequency hearing loss, either at the end of treatment or at the 12-month posttreatment assessment. When plotting the change in THI or TFI as a function of hearing loss only for 8 kHz (i.e., the band with the greatest high-frequency hearing loss across participants; figs. S9 and S10), a trend emerged for arm 3 at the end of treatment, but it was not significant ($P > 0.05$) at the 12-month posttreatment assessment. These trends were weak with high variability; thus, the high-frequency component of PS1 and PS2 could be contributing to therapeutic outcomes for those with high-frequency hearing loss, but it was not the main driver.

In terms of overall stimulation rate, there was no clear indication that the reduced number of stimulus bursts per second in PS3 compared to PS1 and PS2 affected overall efficacy (fig. S11). This finding is consistent with the observation that all arms exhibited rapid and equivalent reductions in THI and TFI within 6 weeks of treatment with minimal additional reductions during the second 6-week period, suggesting that participants using PS3 were receiving more than the minimum number of stimulus bursts for the intervention to be effective. However, from these data, the contribution of stimulation rate on long-term outcomes cannot be completely ruled out and should be further investigated in a follow-up study.

To summarize the overall short-term and long-term therapeutic benefits of bimodal neuromodulation, Fig. 6 collapses the three study arms and pools data for all treatment-compliant participants. Pooled THI and TFI scores are plotted for baseline versus the final treatment assessment (Fig. 6, A and B) and baseline versus the 12-month follow-up assessment (Fig. 6, C and D). These data provide evidence that, irrespective of stimulation parameters, the majority of treatment-compliant participants experienced an improvement in tinnitus symptoms after 12 weeks of treatment

(>81%) and at 12 months after treatment (>77%) based on a large sample size of $n \geq 151$. The remaining ~20% of participants experienced no change or an increase in their tinnitus symptoms (Fig. 6 shows the extent of increase in THI or TFI for those individuals). On the basis of the data from Fig. 6, there was a decrease of 14.2 points (SD: 13.4; Cohen's *d*: -0.91 [-1.04, -0.79]) at the end of treatment that was sustained to 12.7 points (SD: 16.2; Cohen's *d*: -0.81 [-1.00, -0.62]) at 12 months after treatment for THI. There was also a decrease of 13.6 points (SD: 16.2; Cohen's *d*: -0.81 [-0.94, -0.67]) at the end of treatment that was sustained to 14.5 points (SD: 18.6; Cohen's *d*: -0.80 [-0.99, -0.61]) at 12 months after treatment for TFI. These results represent one of the largest and longest posttreatment datasets for a medical device study in the tinnitus field.

Safety and acceptability of device

A Medical Research Organization (NAMSA, Minneapolis, MN, USA) guided and assisted the close-out process of our clinical trial, in addition to reviewing and categorizing safety data together with the Medical Review Board of the study. An adverse event (AE) was classified in line with in line with MEDDEV 2.7/3 (Medical Devices Guidance Documents) and defined as any untoward medical occurrence; unintended disease or injury; or any untoward clinical signs in a subject, user, or other person whether or not related to the investigational medical device. A serious AE (SAE) was defined as an AE that led to death, injury, or permanent impairment to a body structure or a body

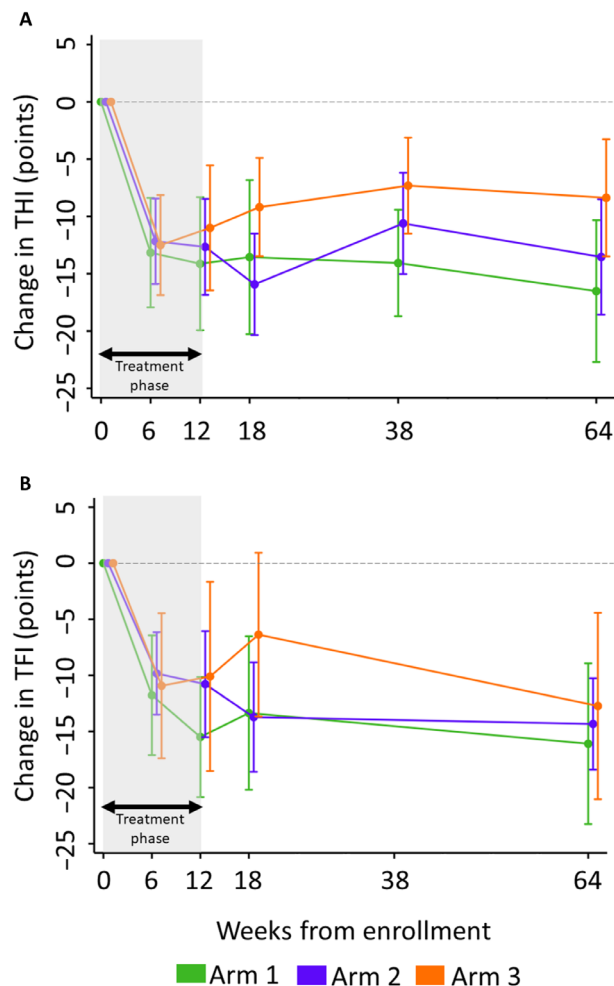


Fig. 5. Long-term clinical efficacy across treatment arms. (A) Change in THI relative to baseline for all treatment-compliant participants who attended every assessment up to the 12-month posttreatment visit (week 64). The change in THI equals the global score (0 to 100) at a given time point minus the global score at baseline. Mean change values and 95% CIs are plotted for each arm (arm 1, $n = 31$; arm 2, $n = 41$; and arm 3: $n = 32$; same individuals at every visit for each arm). (B) Change in TFI relative to baseline for all treatment-compliant participants who attended every assessment up to the 12-month posttreatment visit. The change in TFI equals the global score (0 to 100) at a given time point minus the global score at baseline. Mean change values and 95% CIs are plotted for each arm (arm 1, $n = 31$; arm 2, $n = 40$; and arm 3: $n = 31$; same individuals at every visit for each arm). TFI assessment was not included in the protocol for the 38-week visit. During and after treatment, all arms exhibit a significant reduction in THI and TFI that is sustained out to 12 months after treatment, except for arm 3 at 6 weeks after treatment for TFI ($P > 0.05$; 95% CI crosses the zero line). On the basis of post hoc analyses, arm 1 was significantly different from arm 3 at 38 weeks (6 months after treatment; $P = 0.031$) and at 64 weeks (12 months after treatment; $P = 0.043$) for THI. Arm 2 was significantly different from arm 3 at 18 weeks (6 weeks after treatment; $P = 0.030$) for THI. Statistical significance was determined using an unequal variance, two-tailed t test appropriate for these comparisons based on the method presented in (76). For clearer visualization, the data points and error bars were jittered in time.

function or led to a serious deterioration in health of the subject that either resulted in: (i) a life-threatening illness or injury, (ii) a permanent impairment of a body structure or a body function, (iii) an in-patient hospitalization or prolongation of existing hospitalization, or (iv) a medical or surgical intervention to prevent life-threatening illness.

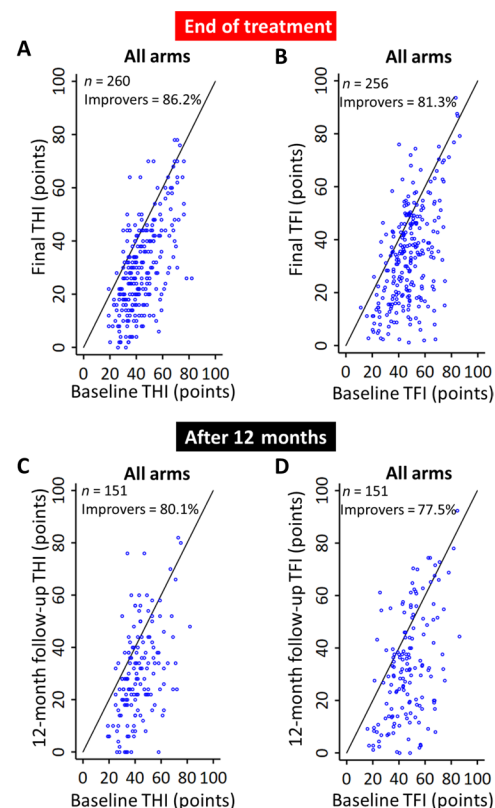


Fig. 6. Long-term clinical efficacy of bimodal neuromodulation. (A and B) Scatterplots represent THI or TFI scores for each treatment-compliant individual at baseline (average of score at screening and enrollment) versus the end of treatment (12-week final visit) for all arms pooled together to represent bimodal neuromodulation in general. (C and D) Scatterplots represent THI or TFI scores for each treatment-compliant individual at baseline versus 12-month posttreatment visit for all arms pooled together. All individuals who completed at least the assessments displayed in each figure were included to maximize the total number of subjects plotted in each figure, leading to a large sample size for each plot. Data points are jittered for visibility.

There were no treatment-related SAEs in this study. Table 2 lists 155 AEs that were potentially related to the treatment. These AEs were categorized into three causality groups as (i) probably device related, (ii) possibly device related, and (iii) probably not device related and further classified as moderate or negligible in terms of severity. One hundred six of the 155 AEs were expected AEs and were described in the clinical study protocol, the device labeling (user manual), and informed consent documents. The most commonly reported expected AE was an increase in tinnitus symptoms and was further subdivided into two categories: a dramatic increase in tinnitus symptoms or a subjective increase in tinnitus symptoms. A dramatic increase corresponded to a substantial increase in tinnitus that was noticeably bothersome to the participant. A subjective increase corresponded to a participant experiencing a change in their tinnitus, which was not considered dramatic but could be occasionally or moderately bothersome or non-bothersome. Other expected AEs included discomfort or pain in the head, ear, or mouth area, as well as discomfort or an ulceration within the oral cavity or on the tongue. There were also 17 unanticipated AEs (Table 2 and 3). Table 2 includes a separate category listed as an “event caused by patient conditions,” which were AEs attributed to

Table 2. Safety data recorded throughout study.

Potential treatment AEs	Total	Negligible	Moderate
(Numbers based on 326 enrolled participants)	155	96	59
Probably device related	35	19	16
Subjective increase in tinnitus symptoms	9	3	6
Dramatic increase in tinnitus symptoms	2	0	2
Fluctuating tinnitus	1	0	1
Headache or pain in the head area	1	0	1
Pain in the ear or mouth area	4	4	0
Temporary swelling of gums	1	1	0
Sensitivity of tongue or oral mucosa	2	2	0
Transient discomfort associated with tongue part	2	2	0
Ulceration of oral cavity	2	1	1
Ulceration of the tongue	4	1	3
Unanticipated AE (see Table 3)	2	1	1
Misuse of device*	5	4	1
Possibly device related	85	46	39
Subjective increase in tinnitus symptoms	45	26	19
Dramatic increase in tinnitus symptoms	3	0	3
Fluctuating tinnitus	9	5	4
Headache or pain in the head area	3	2	1
Pain in the ear or mouth area	6	4	2
Transient discomfort associated with tongue part	2	2	0
Ulceration of oral cavity	2	0	2
Ulceration of the tongue	1	0	1
Unanticipated AE (see Table 3)	11	5	6
Event caused by patient conditions (see table S2)	3	2	1
Probably not device related	35	31	4
Subjective increase in tinnitus symptoms	3	3	0
Dramatic increase in tinnitus symptoms	2	0	2
Pain in the ear or mouth area	1	1	0
Increase in hearing threshold#	1	1	0
Unanticipated AE (see Table 3)	4	4	0
Event caused by patient conditions (see table S2)	24	22	2

*Moderate case caused by an individual overly increasing the sound volume to attempt to mask the tinnitus that led to a bothersome increase in tinnitus, but the individual was instructed to reduce loudness to a comfortable intensity that improved the situation. There were four negligible AEs in which participants increased the tongue stimulation intensity too high that caused discomfort in the oral cavity and tongue area, in which they were instructed to reduce the intensity to improve the situation. #One individual had a 40-dB increase in hearing threshold at 1 kHz in the right ear between screening and final assessments, but remeasuring thresholds at the 6-week follow-up showed the same thresholds as in screening visit. Because there were no obvious cases of compromised hearing across participants attributed to the treatment, this case was likely attributed to a data entry error.

Table 3. Unanticipated AEs listed in Table 2		
Probably device related	Possibly device related	Probably not device related
Moderate:	Moderate:	Moderate:
-Small red fissure on right buccal mucosa	-Cold sores (4 cases)	-None
	-Angular cheilitis	
	-Increased nervousness and shakiness	
Negligible:	Negligible:	Negligible:
-Metallic taste from tongue component	-Light headedness for a few seconds	-Ulceration of lip but not reported during treatment
	-Temporary dizziness	-Pooling of saliva and difficulty swallowing but participant does not believe due to device
	-Fullness in ears for a few seconds	-Difficulty sleeping but reported a cold during the same period
	-Tingling in ears periodically	-Difficulty sleeping but reported cessation of alcohol intake
	-Temporary tiredness and nausea	

other health or personal conditions. Most of these were negligible AEs (24 of 27) with three moderate cases (see table S2).

Comparing differences in each AE subcategory across treatment arms was not possible because of the low number of instances in each arm. However, there was a sufficient number of cases for subjective increase in tinnitus by pooling across causality groups (57 total cases), in which there was no significant difference among the treatment arms (arm 1, 21; arm 2, 17; and arm 3, 19; $P = 0.920$, Fisher’s exact test).

In summing all cases for dramatic increase or subjective increase in tinnitus symptoms, there is a total of 64 AEs listed in Table 2, which correspond to 52 unique individuals (i.e., a single participant could have more than one AE for increased tinnitus symptoms that occurred throughout the study). Forty-seven of these 52 unique AEs happened during treatment, which is consistent with the number of participants who did not exhibit an improvement in THI or TFI at the end of treatment (36 for THI and 48 for TFI based on Fig. 6).

Overall, the treatment proved to be safe with no SAEs and a high satisfaction rate across a large cohort of participants. There were various AEs listed in Table 2 that may have caused discomfort to the participants, but the majority of the participants (273 of 326) still used the device for at least the minimum compliance of 36 hours over the 12-week treatment period. At the end of treatment, participants were asked, “Overall, would you say you have benefitted from using this device?” Out of 272 responses, 66.5% indicated “yes.” They were also asked, “If you knew someone with tinnitus would you recommend they try this treatment?” Out of 270 responses, 77.8% indicated “yes”. Those who responded “yes” to these two questions generally exhibited a greater reduction in tinnitus symptoms compared to those who responded “no” (see figs. S12 and S13). These high compliance and satisfaction rates, when compared to the reported AEs, support a strong benefit-to-risk profile for this medical device treatment for tinnitus.

DISCUSSION

The main objective of this study was to build further safety and efficacy data based on a previous pilot study in humans (31). This study was designed to explore the effects of different stimulation param-

eters on tinnitus symptoms in a broad tinnitus population and in specific subtypes of patients to determine whether they are differentially responsive to certain stimulation settings. Overall, the study represents one of the largest medical device clinical trials in the tinnitus field, supporting the safety, patient tolerability, and efficacy of bimodal neuromodulation for the treatment of tinnitus.

Of the 326 enrolled participants, 84% used the device at or above the minimum compliance criterion of 36 hours over the 12-week treatment period. Of the treatment-compliant participants, 74.7 to 88.8% demonstrated improvements after 12 weeks of treatment with an average decrease of 13.2 to 14.6 points for different arms in THI and TFI, corresponding to a large Cohen’s d effect size ranging from -0.77 to -0.92 . The extent of improvement observed in this study is comparable to the clinically meaningful effects reported in other well-designed clinical trials (28, 45, 47, 59). Currently, the only clinically validated and recommended treatment for tinnitus is cognitive behavioral therapy (CBT) (28, 59). A seminal randomized controlled trial reported improvements in THI of about 10 points after 8 months of a specialized CBT treatment (60). A recently published Cochrane review for CBT also highlighted an average improvement in THI of 10.9 points by the end of treatment (61). In our study, bimodal neuromodulation achieved 14.2 points mean improvement in THI within 6 to 12 weeks of self-administered treatment that could be sustained for 12 months after treatment at 12.7 points. To the best of our knowledge, no other tinnitus treatment has demonstrated such long-term posttreatment effects in a large-scale clinical trial.

In the follow-up phase of the study, we observed different trends that emerged between the three stimulation settings. The more synchronized settings (PS1 and PS2) exhibited sustained effects that were more consistent over time compared to that of PS3, suggesting that shorter interstimulus delays or synchronization between sound and tongue stimulation could be a key feature for driving long-term plasticity. PS1 and PS2 consisted of higher-frequency pure tones and a greater number of stimulus bursts per second (i.e., about 25 times greater) compared to the stimulus features in PS3, which may have also contributed to the long-term benefit observed.

Further support for the efficacy of bimodal neuromodulation was provided in a previous pilot study by Marks *et al.* (30) in 2018

that evaluated a device presenting sound combined with electrical somatosensory stimulation via the cheek or neck region. Twenty tinnitus participants were randomized (1:1) to a bimodal treatment group or an auditory-only control group and underwent 30 min of daily treatment for 4 weeks. The bimodal stimulation group achieved significant improvements in tinnitus symptom severity (e.g., an average reduction of 6.3 points in TFI), whereas the auditory-only group did not achieve a noticeable improvement in symptoms. Therapeutic effects remained for about 3 weeks after treatment. In our study, TFI decreased by 13.6 points with residual therapeutic effects that were sustained out to 12 months after treatment at 14.5 points. There are several possible reasons for the differences in long-term effects. Marks *et al.*'s (30) study used bimodal stimulation consisting of electrical stimulation of the cheek or neck, in contrast to stimulation of the tongue in our study. They also used different acoustic stimuli (i.e., a matched tinnitus spectrum for each participant) and a specific interstimulus timing of 5 ms. We used a diverse range of acoustic stimuli (i.e., different frequency ranges) and delays across treatment arms yet still observed long-term benefits for all arms, suggesting that these stimulus features are not the main reason for the discrepancy. Treatment in the study by Marks *et al.* (30) was only presented for 30 min/day for 4 weeks (a total of 14 hours of stimulation), whereas we used a treatment paradigm with a minimum of 36 hours of stimulation for 12 weeks. On the basis of the concept of paired plasticity (7, 8, 17, 20, 23), repeated bimodal stimulation over a longer period of time may have driven greater and/or longer-lasting changes in the brain in our study. Together with our results, these observations suggest that a diverse range of acoustic and somatosensory stimulation patterns are effective in reducing tinnitus symptoms but that a minimum duration or repetition of paired stimulation is required to achieve long-lasting therapeutic effects. Furthermore, long-term improvements in tinnitus remain robust to interstimulus delays between sound and somatosensory stimulation that vary on the order of tens of milliseconds but appear to diminish with delays exceeding hundreds of milliseconds.

The study by Marks *et al.* (30) reported that repeated paired stimulation could drive continued improvements in tinnitus symptoms with each week of bimodal neuromodulation for their 4-week regimen. However, our study indicates that there may be a limit as to the degree of improvement achieved when using the same stimulation setting. During the second 6 weeks of stimulation, we did not observe as large of an improvement in tinnitus symptoms as that occurred during the first 6 weeks of treatment. There may be habituation effects where the brain becomes less sensitive in response to repetitive stimuli over time (22, 62–65). One way to overcome habituation could be to change the stimulation setting over the course of a treatment regimen (e.g., after 6 weeks of treatment). This stimulus adjustment may potentially lead to greater improvements in tinnitus symptoms that could be sustained over a longer posttreatment period. We are currently engaged in another large-scale clinical trial evaluating the effects of varying bimodal stimulation parameters over time for tinnitus treatment (66).

One limitation in the study is that all three arms consisted of bimodal neuromodulation with suprathreshold intensities of sound and tongue stimulation that were required to maintain blinding, and they showed similar improvements in THI and TFI scores at the end of treatment. On the basis of the concept of paired plasticity, a significant difference would have been expected between the synchronized arm 1 and less synchronized arm 3. Although

this between-arm endpoint was not achieved at the end of treatment, differences in THI and TFI scores emerged between arms during the posttreatment phase, suggesting that different stimulation settings can lead to different long-term treatment outcomes. Prolonged residual effects lasting 12 months would also not be expected for placebo effects. The bimodal stimulation paradigm used by Marks *et al.* (30) led to a significant improvement in tinnitus symptoms that returned back to baseline within 3 weeks. This intervention was believed to be a legitimate treatment by the tinnitus participants, demonstrating that simply informing participants that they are receiving an effective bimodal treatment does not lead to a 12-month residual effect. As described above, there appears to be a minimum duration of bimodal neuromodulation (>14 to 36 hours) required to achieve long-term effects, which can be further sustained by using shorter delays between sound and tongue stimulation (<50 to 550 ms). Therefore, future clinical studies can be designed to reveal differences in long-term outcomes between treatment arms that diverge toward opposite extremes of those two parameter dimensions.

MATERIALS AND METHODS

Study design

We conducted a parallel-arm, double-blind, randomized exploratory study investigating the safety and efficacy of three different stimulation settings. The study was performed at two different clinical sites. One site was the Wellcome Trust—Health Research Board Clinical Research Facility at St. James's Hospital in Dublin, Ireland, and the second site was the Tinnituszentrum of the University of Regensburg at the Bezirksklinikum Regensburg in Germany. The study protocol was independently reviewed and approved by the Research Ethics Committees of the Tallaght Hospital–St. James's Hospital [protocol number: 2016-03-List-11(3)] and by the University Hospital Regensburg (protocol number: 16-101-0186). A detailed description of the study design, protocol, and planned analyses were previously published in a protocol paper (41), and the clinical trial is registered at ClinicalTrials.gov [NCT02669069; (67)]. The CONSORT checklist is provided in data file S1. Informed consent was obtained from each of the 698 participants at the screening visit. One candidate decided not to sign the consent form during the consent process and was not enrolled in the study. The candidate had completed three questionnaires [THI, State-Trait Anxiety Inventory (STAI), and patient history]. This protocol deviation was documented and reported to the appropriate Research Ethics Committee.

A total of 698 candidates were clinically screened at the two clinic sites. Of those candidates, 326 tinnitus participants were enrolled in the study. About 80% of the participants were enrolled at the Ireland clinical site, and the remaining 20% were enrolled at the Germany clinical site (Table 1). The screening visit included various tinnitus, hearing, oral cavity, and health assessments conducted by designated audiologists, clinicians, and researchers to identify participants that satisfied the inclusion and exclusion criteria described in the next section. Each participant was assigned a unique identifier code (UIC) so that participant files and data were managed in a deidentified manner throughout the study. Only designated investigators had access to the keys to decrypt UIC codes. Eligible participants were randomized to one of the three different treatment arms. All participants and participant-facing investigators were blinded to the treatment assignment for each participant. When participants returned for their enrollment visit, we performed several tinnitus

assessments and health evaluations relevant for monitoring AEs. Each participant was then fitted with a bimodal neuromodulation device and provided a training session on how to use the device. We recommended two daily 30-min treatment sessions over a 12-week period. The two sessions could be performed consecutively or at different times during the day. If the participants used the device for at least 36 hours over the full treatment period, then they were considered compliant to treatment, as described in the published protocol paper (41). During the treatment period, the participants attended the clinic for interim assessments at a 6-week visit and, at the end of treatment, at a 12-week visit. Tinnitus assessments and health evaluations were repeated across all visits. Participants returned their devices at the 12-week visit and were asked to return for three follow-up visits at week 18 (6-week follow-up), week 38 (6-month follow-up), and week 64 (12-month follow-up) to assess the long-term safety and efficacy of the treatment. If there were cases that required additional medical assistance during the study, referrals were made to an appropriate clinician (three cases are described in table S3). The study timeline is depicted in Fig. 1B, with the number of participants attending and completing assessments listed in Fig. 2 and Table 1.

For randomization of participants to the three different treatment arms (1:1:1), stratification was performed on the basis of four binary categories using the method of minimization (49), as described in the published protocol paper (41). These categories included atonal tinnitus, hyperacusis, normal hearing, and noise-induced hearing loss. Atonal tinnitus refers to participants who did not have a dominant tonal tinnitus. Hyperacusis was defined as a sound level tolerance [or loudness discomfort level (LDL)] that was less than 60-dB sensation level (SL) for a pure tone presented at 500 Hz in either ear [dB SL equals dB hearing level (HL) minus hearing threshold level of acoustic stimulus]. Because we did not incorporate additional clinical criteria and questionnaire data relevant for classifying hyperacusis into the stratification process, this category should be interpreted more strictly as a sound sensitivity condition specific to 500 Hz rather than what is typically considered as hyperacusis. The word “hyperacusis” is still used throughout this paper to remain consistent with the previously published stratification terminology for the clinical study (41). This hyperacusis criterion of LDL <60-dB SL is also summarized in units of decibel HL to show that consistent stratification occurred for both types of units in our study (table S4). Normal hearing was defined as having pure-tone thresholds of ≤ 20 -dB HL in all audiometric test frequencies between 250 Hz and 8 kHz in both ears. Noise-induced hearing loss was defined as having a dip in pure-tone hearing thresholds of 10 dB or more in any of the measurement frequency bands of 3, 4, or 6 kHz in either ear. The classification was performed manually by a trained audiometrist wherein patients with either symmetrical dips or asymmetrical dips (e.g., 10 dB on the low-frequency side and 5 dB on the high-frequency side) were included. In some cases, participants could have substantial presbycusis that altered the audiograms used for classification, in which case these participants were included in the noise-induced hearing loss category if they self-reported that their tinnitus was caused by noise exposure.

The tinnitus treatment device is a proprietary CE-marked class IIa medical device developed by Neuromod Devices (Dublin, Ireland). The device comprises bimodal auditory and tongue stimulation (Fig. 1A). The auditory stimulus includes sequences of tone bursts with a continuous structured wideband noise in the background that are presented to both ears. The tongue is stimulated electrically

via a 32-electrode transmucosal array placed on the anterior dorsal surface of the tongue. Electrical stimulation is delivered in the form of biphasic, anodic-leading pulses between about 5- and 210- μ s duration and with a fixed amplitude. Bimodal stimulation with suprathreshold intensities for both sound and tongue stimulation was used for all three study arms, which enabled participant and investigator blinding to their assigned treatment. Suprathreshold settings of bimodal neuromodulation also achieved significant improvements in tinnitus symptom severity in a previous pilot study (31). Schematics and details of the stimulation parameters for each arm are presented in table S1. Briefly, arm 1 (PS1 setting) consisted of synchronized sound and tongue stimulation. A given pure tone (~ 15 -ms duration, 500 to 8000 Hz, repetition period of ~ 80 ms) was presented at the same time that a train of pulses was presented to a specific location on the tongue (~ 12 to 15 ms). For arm 1, the stimulation location on the tongue array was fixed for a given pure-tone frequency; this resulted in a fixed tone-to-tongue mapping. Note that the 32-electrode tongue array shown in Fig. 1A was divided into two halves (left and right) such that symmetrical locations on each side of the tongue were electrically stimulated as the corresponding pure tone was presented binaurally (table S1). Arm 2 (PS2 setting) used the same auditory stimulus as arm 1 but had an interstimulus delay that varied in the range of 30 to 50 ms across each stimulus presentation. The tone-to-tongue mapping was also randomized across stimulus presentations. Arm 3 (PS3 setting) used lower-frequency tones (100 to 500 Hz), longer varying interstimulus delays (550 to 950 ms), and a randomized tone-to-tongue mapping as in arm 2. These longer delays also required a longer repetition period, which was set to ~ 2 s for PS3. Arm 3 with PS3 was designed to consist of a much less synchronized and less coordinated setting compared to PS1 used in arm 1. On the basis of the concept of paired stimulation described in Introduction (17, 20, 23, 30, 68–70), the greatest treatment differences would be expected between arms 1 and 3. Greater synchrony or more coordinated activation of bimodal neuromodulation should drive stronger therapeutic plasticity within brain regions related to tinnitus.

Sound volume range and electrical tongue stimulation intensities were customized to each participant's sensation thresholds. The participant's pure-tone audiometric thresholds (250 Hz to 8 kHz) were measured at the screening visit (fig. S3) and subsequently used to configure the intensity of the auditory stimuli to ~ 10 dB in SL (dB SL) above their hearing thresholds at each tone frequency. The participant could adjust the default auditory stimulus intensity between -12 and $+12$ dB during treatment using volume buttons on the controller. For safety reasons, the upper stimulus intensity was limited for participants with profound hearing loss (up to ~ 90 dBA). Electrical stimulation intensity (i.e., via changes in pulse duration for a fixed amplitude) was configured for each participant by adjusting the intensity from subthreshold to suprathreshold sensations to a comfortable intensity across different electrodes, which was when the participant could feel sensations on the tongue but below an intensity that was bothersome or painful. This comfortable intensity was used as the calibrated setting, and the participant could adjust the electrical stimulation intensity down to 60% and up to 160% of this calibrated setting using the buttons on the controller. The treatment device reverted to the default intensities at the start of each new treatment session.

Participants

The study recruited participants with chronic subjective tinnitus, which is a phantom auditory percept that is attributable to abnormal firing

patterns in the brain (25). Subjective tinnitus differs from objective tinnitus, which is typically associated with sounds generated from vasculature or pulsation anomalies, abnormal muscle contractions, or head/jaw movements (25). Participants in the age range of 18 and 70 with a tinnitus duration between 3 months and 5 years were included in the study. We recruited participants with THI scores of 28 to 76 points (44). We considered a score less than 28 as being sufficiently mild as to be less amenable to therapeutic intervention. A score greater than 76 is classified as catastrophic (44) and typically associated with other severe comorbidities including depression, anxiety, and suicidal tendencies; these participants were not included because of the challenges in managing these complex comorbidities in a large-scale clinical trial. Only participants with a minimum masking level (MML) between 20 and 80 dB HL were included in the study. MML was determined by presenting a wideband noise at increasing intensities until the sound masked the participant's tinnitus. Additional inclusion criteria required that the participant was able to read and understand English at the Ireland site or read and understand German at the Germany site, was willing and able to provide informed consent, and was willing to commit to the full duration of the study.

Candidates were excluded if they had objective tinnitus or somatic tinnitus caused by a head or neck injury or if their tinnitus was comorbid with a neurological condition as assessed by an audiologist or clinician. Conductive hearing loss demonstrated by abnormal otoscopy or abnormal tympanometry was an exclusion criterion. Substantial sensorineural hearing loss, either unilaterally or bilaterally, was also an exclusion criterion because the headphones had an upper volume limit. Specifically, those candidates having >40-dB HL in at least one measurement frequency in the range of 250 Hz to 1 kHz or >80-dB HL in at least one measurement frequency in the range of 2 to 8 kHz were excluded. These hearing loss criteria corresponded to 10.2% or 5.3% of the screened individuals (of the 698) who were excluded from the study, respectively. Further exclusions included candidates with the following: a hearing aid used within 90 days before eligibility assessment, any type of electro-active implantable device (e.g., vagal nerve stimulator, cochlear implant, or a cardiac pacemaker), a LDL of <30-dB SL at 500 Hz for either ear, a temporomandibular joint disorder, anxiety determined by a score >120 of 160 on the STAI (71, 72), or moderate to severe dementia as indicated by a score <20 on the Mini-Mental State Examination (MMSE) (73). A final set of exclusion criteria based on a medical history taken at the screening assessment were Meniere's disease, oral piercings, pregnancy, involvement in medicolegal cases, history of auditory hallucinations, current prescription of a drug for a central nervous system pathology (e.g., epilepsy), and previous use of a Neuromod Devices' product.

Primary endpoints

The THI (44, 45) and TFI (47) are clinical outcome measures commonly used to assess tinnitus symptom severity (46). These outcome measures have been used across multiple studies that have supported clinical guidelines for tinnitus interventions (28, 61). The only clinically recommended treatment for tinnitus, CBT, has also leveraged these multi-item questionnaires for evidence that it is an effective tinnitus intervention (61). The THI predominantly assesses the emotional and functional impact of tinnitus, in which 25 items are scored 4/2/0 on a categorical scale corresponding to yes/sometimes/no. The global score of the THI (i.e., sum of points across all 25 items)

has a value from 0 to 100, with a higher score indicating a greater negative impact of tinnitus. The TFI assesses a range of tinnitus-related functional complaints experienced over the week before assessment. Each of the 25 items is assessed on an 11-point Likert scale, and the sum of the scores is normalized to give a global score from 0 to 100, with a higher score indicating a greater negative impact of tinnitus.

Changes and deviations to the trial design

The target sample size assumed 91 participants in each treatment arm at the 12-week endpoint, with the design accounting for an expected dropout rate of 20% for a total of 342 participants. The achieved retention rate was better than predicted, and so, recruitment was stopped at 326 participants based on the accumulating data.

Of the 326 enrolled participants, 34 of them did not meet the eligibility criteria and are considered deviations to the protocol. Ten had a tinnitus duration greater than 5 years, 3 had a THI score less than 28 points, 16 scored below 20 points on the MMSE, 2 had an MML of greater than 80-dB HL, 2 had previously been involved in a medical legal case, and 1 indicated they had a neurological disorder. In addition to these 34 participants, there were 30 who had at least one missing entry for their eligibility assessment such that it was not possible to confirm if there was a deviation to the inclusion and exclusion criteria. Analyses were conducted with and without these participants. The sensitivity analysis demonstrated no significant effect on the primary endpoint findings ($P > 0.05$); thus, all of these participants were included in the analyses presented in this paper.

During the database validation process, there were 10 participants whose audiogram data were identified as being incorrectly inputted into the database at the screening visit, in which their hearing threshold values were supposed to be higher than what was inputted into the database (i.e., by 5 to 25 dB across one to five frequencies in the range of 500 to 8000 Hz). In these cases, the sound volume was not set to be loud enough for some frequencies. Analyses were conducted with and without these participants. The sensitivity analysis demonstrated no significant effect on the primary endpoint findings ($P > 0.05$); thus, all of these participants were also included in the analyses presented in this paper.

Statistical analyses

The primary endpoints of the study were designed together with statistical experts from NAMS Medical Research Organization (Minneapolis, MN, USA) and are described in a previously published protocol paper (41). These endpoints included within-arm changes from baseline to the end of treatment at 12 weeks (final) for THI and TFI and between-arm differences in changes from baseline to final for THI and TFI. Baseline was calculated as the average of the THI or TFI scores at screening and enrollment. The within-arm analyses were based on a per-protocol estimand and tested with paired two-tailed t tests, where normality assumption was confirmed with the Shapiro-Wilk test. Within-arm effect sizes reported in the paper were based on Cohen's d and are calculated as the mean score at the end of treatment minus the mean score at baseline, divided by the pooled SD for the two scores. The between-arm analyses were based on an intention-to-treat estimand with multiple imputation and tested with multiple regression using baseline scores as a covariate, where normality and equal variance assumptions were confirmed with the Shapiro-Wilk test and Bartlett's test, respectively.

An ANCOVA was also performed across arms. Missing data were handled by using a Markov chain Monte Carlo multiple imputation method (74, 75) that filled in the missing predictor variables for 10 multiple imputed datasets. This procedure was followed by a second imputation process that estimated the final outcome variable within each imputed dataset. The imputation was based on a regression model with predictor variables of age; gender; tinnitus tonality; degree of somatic modulation; duration of tinnitus; treatment arm; treatment compliance; high-frequency hearing loss; degree of sound intolerance; and baseline and posttreatment THI, TFI, and MML scores. Inferences for the between-arm endpoints were evaluated on each imputed dataset and the results combined to yield the estimates, CIs, and associated significance values. The use of per-protocol estimand for within-arm analyses ensured that the changes in outcome measures within each treatment arm were reflective of daily use scenarios where the participants use the treatment as directed and thus included only the participants who were compliant to treatment. The threshold for inclusion in the per-protocol analysis was set at a minimum compliance threshold of 36 hours across the intended 12-week treatment period, as described in the protocol paper (41). We were able to track daily usage and stimulus adjustments that were logged automatically by the device, such as the time and date when the device was in use, the duration of electrode contact with the tongue, and the intensities of both the auditory and tongue stimuli for each treatment session.

The study was conducted at two clinical sites with the intention of pooling data from both sites for the primary endpoint inferences. A test for site interaction on the primary between-arm endpoints (comparing mean change in scores from baseline to final assessments) was conducted and found to be not statistically significant ($P > 0.05$ for both THI and TFI, using linear regression with independent variable of treatment arm and interaction term of clinical site, and with baseline score as a covariate). Therefore, the data from each clinical site were combined as intended in the study design to perform the analyses presented in this paper. Figures S14 and S15 further show scatterplots for the baseline and final THI and TFI scores, respectively, for each treatment arm and clinical site.

SUPPLEMENTARY MATERIALS

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Fig. S1. Compliance and satisfaction rates in using treatment device.

Fig. S2. Histogram of time (in weeks) when participants attended each visit relative to enrollment.

Fig. S3. Hearing thresholds for enrolled participants.

Fig. S4. Scatterplots for change in THI or TFI score versus baseline score for each arm.

Fig. S5. Scatterplots for THI scores at baseline versus 12-month posttreatment assessment for each arm.

Fig. S6. Scatterplots for TFI scores at baseline versus 12-month posttreatment assessment for each arm.

Fig. S7. Scatterplots for change in THI or TFI score (from baseline to the end of 12-week treatment) versus high-frequency hearing loss (4 to 8 kHz) for each arm.

Fig. S8. Scatterplots for change in THI or TFI score (from baseline to 12-month posttreatment assessment) versus high-frequency hearing loss (4 to 8 kHz) for each arm.

Fig. S9. Scatterplots for change in THI or TFI score (from baseline to the end of 12-week treatment) versus 8-kHz hearing loss for each arm.

Fig. S10. Scatterplots for change in THI or TFI score (from baseline to 12-month posttreatment assessment) versus 8-kHz hearing loss for each arm.

Fig. S11. Scatterplots for change in THI or TFI score versus duration of device usage.

Fig. S12. Changes in tinnitus symptoms based on responses to first satisfaction question.

Fig. S13. Changes in tinnitus symptoms based on responses to second satisfaction question.

Fig. S14. Scatterplots for THI scores for each treatment-compliant individual at baseline versus the end of treatment (12-week “final” visit) for each arm.

Fig. S15. Scatterplots for TFI scores for each treatment-compliant individual at baseline versus the end of treatment (12-week final visit) for each arm.

Table S1. Schematics and description of stimulation setting used in each arm.

Table S2. AEs attributed to patient conditions from Table 2.

Table S3. Patient referrals for AEs.

Table S4. Stratification category of hyperacusis in Table 1 calculated in units of dB HL.

Data file S1. CONSORT checklist.

[View/request a protocol for this paper from Bio-protocol.](#)

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allocation sequence and assigned participants to each arm. C.H., C.O.C., M.S., and B.L. coordinated and/or performed the study assessments and documentation tasks. S.H. programmed the devices and managed technical support. B.C., B.L., C.H., T.S., S.H., E.M., and H.H.L. evaluated and helped to categorize the AEs. H.H.L., E.M., and S.H. were involved with overseeing and closing out the study. H.H.L., E.M., S.H., B.C., C.H., B.L., D.A.H., S.V., M.S., S.L.L., and T.S. analyzed and/or interpreted the data. H.H.L. drafted the manuscript with contributions from the other authors. **Competing interests:** B.C., C.H., S.H., E.M., C.O.C., S.L.L., S.D., and HHL are or were employees, consultants, and/or shareholders of Neuromod Devices. B.L., S.V., and D.A.H. serve on the clinical advisory board of Neuromod Devices and receive monetary compensation for their contribution. **Data and materials availability:** All data associated with this study are present in the paper or the Supplementary Materials. Participant-level data can be accessed with reasonable request and appropriate ethics approval by contacting H.H.L. (hubert.lim@neuromoddevices.com).

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Bimodal neuromodulation combining sound and tongue stimulation reduces tinnitus symptoms in a large randomized clinical study

Brendan Conlon, Berthold Langguth, Caroline Hamilton, Stephen Hughes, Emma Meade, Ciara O Connor, Martin Scheckmann, Deborah A. Hall, Sven Vanneste, Sook Ling Leong, Thavakumar Subramaniam, Shona D'Arcy and Hubert H. Lim

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Relief from ringing

There is no universally effective treatment for tinnitus, phantom perceived sound, or ringing in the ears. Conlon *et al.* studied the effects of bimodal neuromodulation using a device that delivers electrical stimulation to the tongue and sound to the ears. Participants reported reduced symptom severity at the end of 12 weeks of treatment, which persisted for 12 months. Results support the safety and potential utility of bimodal neuromodulation for tinnitus.

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