Customised enriched acoustic environment for sound therapy of tinnitus

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Abstract – Tinnitus is an auditory disorder very difficult to treat. Whereas up until now there is not a “cure” for tinnitus, the most extended treatment combines counselling with sound therapy. When this sound is a broadband noise in the audio frequency band, this protocol is named tinnitus retraining therapy. Even though broadband noise was proposed at the beginning as the stimulus for sound therapy, many other sounds have been subsequently proposed and used, including tones, noise bands, music, and nature sounds. Although any sound, low enough to avoid annoyance, discomfort or hearing damage, is better than silence for tinnitus treatment, it is not still clear the relationship of the success of the therapy with the properties of the sound stimuli. The aim of this article is to propose an optimal sound treatment that provides a precise and selective stimulation of the whole auditory system. The proposed sound stimulus, Enriched Acoustic Environment, consists of sequential tones or broadband noise matched to the HL curves of the patients. The acoustical characteristics of these stimuli are analyzed and their positive effects in the treatment of subjects with tinnitus are reported.

Keywords: Tinnitus, Hearing loss, Sound therapy, Enriched Acoustic Environment

1 Introduction

Tinnitus is the medical term for the auditory perception of sounds in the absence of sound sources internal or external to the body [1, 2]. This phantom auditory perception presents mainly as a continuous and fluctuant tone, ringing, or hissing of high or low pitch, with loud or soft sensation level. Tinnitus is an uncomfortable auditory disorder affecting severely the quality of life of subjects [3], causing many audiological, cognitive, and neurological concerns including altered sleep patterns, attention deficits, annoyance, irritability, panic, stress, anxiety, and depression [4–6]. Epidemiological studies afford that approximately 10% of the adult population in Western industrialized countries have been eventually affected by tinnitus [7]. For many of these subjects (approximately 1–2%), tinnitus is a handicap producing distress and deteriorating significantly their quality of life. Approximately 5 million of European people have suffered of moderate-to-severe tinnitus [8]. In Spain, 17% of consultations in specialised audiological clinics are due to tinnitus [9], whilst 1% of population are affected of severe tinnitus [10].

It has been suggested that hearing loss (HL), the rise of hearing thresholds of a subject expressed in decibel (dB), is a significant risk factor for developing tinnitus. The occurrence of hearing problems in the developed countries has increased over the past last years, due to aging of the population (age related hearing loss, ARHL) and overexposition to noise (noise related hearing loss, NRHL) [11]. The incidence of tinnitus, as ARHL does, increases with age. Loud noises, such as those from industry, traffic in urban environment, and portable music players in young population, are common sources of NRHL and increase the risk of suffering tinnitus. Exposure to noise during short time, such as attending a disco session, can produce tinnitus that usually goes away, but living with long-term harmful noise can produce permanent damage to auditory system and hence raises the risk of developing tinnitus [12]. For developed countries, this renders a higher incidence of tinnitus, which in turn has a significant economic impact on the public health system [13]. Although HL can be a risk factor for tinnitus, it may apparently occur in people with normal hearing thresholds. Recent studies have disclosed that high frequency HL (above 8 kHz), which is not currently measured in hearing screening, and hidden hearing loss (HHL), a type of permanent cochlear damage without an elevation of hearing thresholds, can afford tinnitus [14, 15]. Therefore, tinnitus is a quite prevalent disorder challenging to deal with. There is evidence that tinnitus is associated to adjustments of the normal operation of the neural part of the auditory system which attempts to compensate for some kind of deficit at the auditory periphery [16]. When there is some deafferentiation from the peripheral downstream to the neural auditory system (a loss of hearing,
for instance), the neural part reorganises to compensate for this deficit. This capacity of reorganization is a characteristic of brain plasticity. There exists physiological evidence of functional changes in the neural part as a reaction to damages in the peripheral part of the auditory system, such as those produced by an acoustic trauma [17], head trauma, ear infection, ototoxicity, vestibular troubles, and others, giving rise to tinnitus.

The deafferentation of a normal input to central auditory system represents a potential trigger of plastic changes, such as increasing of the spontaneous activity (hyperactivity), hypersynchrony or tonotopic map reorganization [16]. These three mechanisms could underpin the origin of tinnitus at sub-cortical and/or cortical levels of the auditory system through the following sequence [18]: (1) a problem in the auditory periphery (for instance, hearing loss due to noise overexposition or aging) is produced; (2) deafferentation in the damaged frequencies could afford an overrepresentation of the frequencies at the edge of the lesioned band, which causes hyperactivity and/or possible hypersynchrony in cortical pathways, giving rise to the primary tinnitus signal; (3) under normal functioning, the tinnitus signal is inhibited at the thalamic gate by a feedback loop (inhibitory gating mechanism). When this gating mechanism is compromised, inhibition fails, and the neural signal is transmitted to higher order structures, where it causes permanent reorganization and chronic tinnitus.

Symptomatic tinnitus can be temporarily masked by sound stimulation aimed to produce residual inhibition (RI) [19]. The tinnitus signal usually returns back some time after (from minutes to hours) the acoustic signal ceases. An acoustic signal which yields a long lasting RI after the stimulus is ceased should become an effective sound therapy. The dependence of the inhibition grade on the time after the stimulus ceases is still under study [19]. Many sound therapies for tinnitus treatment have been suggested [20–22] including tinnitus re-training (TRT) [23, 24], auditory discrimination (ADT) [25–29], phase-shift [30, 31], neuromonics [32], notched-music [33–35], neuromodulation [2, 36–38], fractal tones [38–40], and many others.

Sound therapies assume that exposing the auditory system to an enriched sound can rearrange it to normal functioning due to plasticity. Although any sound is better than silence for tinnitus treatment, provided that it does no annoy, create discomfort, or damage hearing [24], it is still not clear the relationship of the success of the therapy with the properties of the sound stimuli. The aim of this article is to propose an optimal sound treatment that provides a precise and selective stimulation of the whole auditory system. Since this sound therapy is applied combined with counselling, in the context of a variation of the TRT, the bases of this tinnitus treatment are reviewed in Section 2. The fundamentals and acoustic properties of the proposed therapy, named Enriched Acoustic Environment (EAE), are exposed in Section 3. Some results are presented in Section 4. And finally, the main conclusions of this article are outlined in Section 5.

2 Tinnitus retraining treatment (TRT)

Tinnitus is a hearing disorder that arises by aberrant plastic compensation mechanisms in the neural auditory system to some peripheral deafferentation. Thus, tinnitus is in fact a phantom auditory perception generated at subcortical or cortical levels of the auditory system. In spite of many efforts carried out to find a “cure” for tinnitus, including pharmacological drugs, sound, magnetic/electrical stimulation, physiotherapy, relaxation, and other complex interventions (defined as a combination or two or more of the preceding modalities) [41], such a cure there not exists nowadays. Rather than looking for removing it, the practical guideline for treating tinnitus is trying to reduce its distress.

The most extended treatment of tinnitus is TRT, a comprehensive, non-invasive protocol applied in most clinics that has demonstrated to achieve about 80% of distress reduction in tinnitus patients, based in the neurophysiological model of Jastreboff [23]. This model is based on that, whilst the auditory system is involved in the trigger and propagation of the tinnitus signal, other brain systems, mainly the limbic and the autonomic nervous systems, are involved in its deleterious reactions, beginning with annoyance, panic, and difficulties with sleep and concentration, and, in the worst cases, evolving into anxiety, stress, and depression.

Sound, an acoustic signal at the peripheral auditory system, is transduced into a sequence of spikes at the hair cells-auditory nerve interface. The sound evoked spikes propagate downstream the afferent auditory way from the auditory nerve (AN) to the auditory cortex (AC) through the cochlear nucleus (CN), the olivary complex (OC), the lateral lemniscus (LL), the inferior colliculus (IC), and the medial geniculate body (MGB) [11]. Sound processing also activates other brain systems, such as the amygdala and the hippocampus, two major limbic structures. The amygdala is particularly sensitive to voice and music and plays a significant role in conditioning to auditory fear and regulation of the response to sound startle [42]. A stressful acoustic stimulus, may release amygdala-mediated stress hormones via the HPA-axis, which may have negative effects on health. A short-term exposure to stress hormones may produce hearing protection [42]. Noise exposure can affect long term plasticity of hearing [42]. The hippocampus can add temporal dimension to the auditory processing.

Sound patterns originate subcortically and enter both auditory and limbic systems via the MGN. Under normal functioning, the limbic system may recognize a transient tinnitus as perceptually unsuitable and inhibit it at the MGN [43], this in turn reducing the propagation of the unwanted tinnitus signal in both circuits. When inhibition of this tinnitus signal is prevented, continued thalamocortical activity results and the constant perception of the tinnitus signal is produced (chronic tinnitus). Tinnitus, thus induced by hearing disorder, is associated to changes in the hippocampus. The dysregulation of limbic and auditory networks is in the basis of chronic tinnitus, indicating that an advanced comprehension of auditory-limbic interactions
may be crucial for an ultimate cure of tinnitus in the future [42, 43].

The aim of TRT is to habituate to the tinnitus signal by reducing its negative response. When the tinnitus signal propagates to other brain systems, and in particular triggers the limbic and autonomic nervous systems, brings a number of aversive reactions and tinnitus becomes annoying. Consequently, the aim of treatment should be taking advantage of brain plasticity to achieve the habituation of negative tinnitus reactions, by blocking the functional connections that spread the tinnitus signal from the auditory to the other brain systems [23].

In practice, TRT combines counselling with sound therapy, both based on the above described neurophysiological model of tinnitus [23]. The main goals of sound therapy and counselling are to lessen the strength of neuronal activity related to tinnitus and to demystify it by unravelling the mechanisms underlying tinnitus, respectively [44]. Successful counselling will diminish the negative associations with tinnitus, which is necessary to facilitate its habituation. The sound stimulus is intended to maintain an enriched environment of non-annoying sounds that improve the treatment result by modifying auditory processing at sub-conscious levels and facilitating the habituation to the tinnitus perception [23].

An important aspect of TRT is that the aetiology of tinnitus is irrelevant, since it does not attempt to suppress the tinnitus signal itself but to attenuate its negative reactions. Any type of tinnitus can be successfully treated by TRT. As the tinnitus signal is prevented from reaching brain systems, the negative reactions significantly attenuate, being not necessary to remove them. This feature differentiates TRT from other methods more oriented to the suppression of the tinnitus source [23].

### 3 Enriched Acoustic Environment (EAE)

#### 3.1 Antecedents

Noreña and Eggermont [17] demonstrated that the reorganization of the tonotopic map of cats produced by the exposition to an injuring noise could be inverted by exposing them to an appropriate EAE. Fourteen laboratory cats were exposed first to a loud noise environment. Seven of these cats (group 1) were not subjected to additional acoustic stimulation, whereas a second group of other seven cats (group 2) were immersed into an EAE inside a room. The EAE consisted of a high-pass filtered tone-pip sequence (see Sect. 3.2) of random frequency between 625 Hz and 20 kHz and sound pressure level of 80 dB. This level was above the expected HL in the high frequency range, but low enough to avoid further impairment to their hearing system. The cats of group 2 were stimulated for 24 h/day during 35 days to obtain the best effect on the auditory system after the traumatizing hearing loss. Other control group of 5 non-exposed cats was used. They mapped the auditory AI cortex of the three group cats. The tonotopic map of the traumatized non-EAE exposed cats (group 1) showed a profound reorganization compatible with high frequency hearing loss. However, the cortical map of the traumatized EAE exposed cats (group 2) showed a frequency organization similar to that of the cats in the control group. Therefore, they concluded that the cortical tonotopic map reorganization of cats produced by the traumatizing noise could be compensated by stimulating with a tone-pip EAE in the damaged frequency region.

Schaette and Kempter [45, 46] used a computational model of auditory neurons to demonstrate that tinnitus-induced hyperactivity produced by homeostatic plasticity could be reversed by appropriate acoustic stimulation. Using their model, they were able to calculate the firing rate and gain of downstream auditory neurons. The effects of homeostatic plasticity in these neurons were taken into account by introducing changes in the gain factor triggered by deviations of the main activity from a certain target activity (hyperactivity), so that the gain adjustments mimicked the changes in effective response by homeostatic scaling. As a result of this model, when the excitatory input to one of these neurons was reduced (deafferentation), homeostatic plasticity increased the effective gain resulting in tinnitus. They were also able to evaluate the effect of different sound stimuli (sound therapy) to these tinnitus-damaged neurons. When a white noise was used, none effect was observed in the aberrant gain produced by tinnitus. If, however, a matched-noise stimulation (a stimulus with spectrum matched to the HL curves) was applied to the damaged neuron, the aberrant gain was eventually removed.

Noreña and Chery-Croze [47] studied if the aberrant central gain distinctive of hyperacusis patients could be reduced by subjecting them to an EAE, consisting of a sequence of tone-burst. The auditory hypersensitivity of these patients could be caused by hearing losses, which produced a deficit of sensory inputs to the central auditory system, and hence plasticity-induced increase in central gain. The aim of the study was to test whether this aberrant central gain could be compensated by the use of an EAE and the auditory hypersensitivity consequently reduced. The EAE was composed of a sequence of tone-bursts of random frequency within the hearing range, and amplitude weighted as a function of the hearing loss. The participants were asked to hear the EAE for 1–3 h/day at a just audible level. After 15 weeks, all participants reported a statistically significant sensitivity decrease. Herráiz et al. [48] demonstrated that this therapeutic tone-burst EAE was able to produce clinically relevant relief in 21 tinnitus patients.

#### 3.2 Gammatone sequence

Therefore, sequences of EAE have been used until now with either tone-pips or tone-bursts. Both consist of random frequency tones with amplitude weighted by a factor proportional to the HL and shaped by a window that is symmetrical, for the case of tone-burst, and asymmetrical for the case of tone-pips. The equation for such an EAE sequence is then,
\[
    eae(t) = \sum_n A_n(f_n) W(t - \tau_n) \cos[2\pi f_n(t - \tau_n)],
\]
where,
\[
    A_n(f_n) = 10^{\text{HL}(f_n)/20},
\]
is the amplitude of the tone,
\[
    W(t) = \begin{cases} 
        W_T(t) & \text{for tone - burst} \\
        t^{-\alpha} e^{-\gamma t} & \text{for tone - pips}
    \end{cases}
\]
is the envelope function, \( W_T \) is a time window, \((\alpha, \gamma)\) are parameters, \( f_n \) is a frequency within the hearing range, \( \text{HL}(f_n) \) is the hearing loss at this frequency, and \( \tau_n \) is the interlatency between tones (the inverse of the rate, or number of pulses per second).

The envelope of tone-bursts corresponds to a time window (Hanning, Hamming, Blackman–Harris, etc.). The rising and decaying parts of the tone-pip window are controlled by two parameters \((\alpha, \gamma)\), which can be chosen so that its magnitude spectrum matches the tuning curve of different parts of the auditory system. Figure 1a shows two tones, a tone-burst and a tone-pip, of the same frequency (500 Hz) and with similar duration (0.1 s). Figure 1b shows the corresponding log-spectra. The main difference between both spectra is the presence of strong side-lobes in the case of the tone-burst. The spectrum of the tone-pip is more "clean," and asymmetrical, similarly to the frequency response curves of different sites of the auditory system. Therefore, it seems that the tone-pips are more refined sounds for stimulating the human auditory system. Korn studied a similar problem in the context of the theory of audio information [51]. According to the uncertainty principle, \( \Delta f \Delta t = \text{constant} \), any measurement of the frequency of a signal is affected by a zone of uncertainty, which is inversely proportional to the time (duration) of observation. The way our auditory system processes a signal is closely related with the uncertainty principle. A click is perceived as one of two consecutive sounds depending on its length. The mechanism allowing our auditory system to change from the time to the frequency domain in the processing of sounds is related with the concept of masking, which consists of the increment of the hearing threshold of a tone in the presence of another tone. The masking curve at each frequency is considered as the representative curve of this frequency. In other words, masking can be considered as a discretiser of the auditory system allowing to measure the discrete value of such frequency for a finite length stimulus. Therefore, the stimulus corresponding to the inverse Fourier transform of a masking curve can be considered as the elemental message at discrete frequency (EMDIF) [51]. When the stimulus is shorter than an EMDIF, its spectrum is wider than the corresponding masking curve and it is processed in the frequency domain. If, on the other hand, the stimulus is larger than an EMDIF, its spectrum is shorter than the masking curve, and it will be processed in the time domain.
domain. Korn [51] showed that masking curves can be modelled by the filter,

$$S(x) = \frac{x^2 + (\omega - \omega_0^2)}{\omega^{x-1}},$$  

(4)

which inverse Fourier transform is,

$$e(t) = t^{-1}e^{-\pi t}\cos \omega_0 t,$$  

(5)

which corresponds exactly to a tone-pip.

An appropriate therapy for tinnitus subjects should be that with the two parameters ($\alpha$, $\gamma$) tuned to the curves at various locations of the human auditory system. These are, in fact, the gamma filters [52–55]. The inverse Fourier transform of the gamma filters are the gammatones. Let,

$$\text{ERB} = 0.108 f_0 + 24.7,$$  

(6)

be the equivalent rectangular bandwidth of an auditory filter, also known as critical band, where $f_0$ is the central frequency of the band. The corresponding gammatone is then given by Equation (5) with the pair of parameters ($\alpha$, $\gamma$),

$$\alpha = 2\pi \text{ERB} = 2\pi (0.108 f_0 + 24.7)$$

$$\gamma = 4$$

(7)

Therefore, just a parameter, the central frequency of the band $f_0$ determines the gamma filter at each frequency.

A bank of six normalised gamma filters at octave band frequencies between 250 Hz and 8 kHz, similar to the filters currently used for simulating the basilar membrane movement in the cochlea, is shown in Figure 3. Thus, a sequence of gammatones should be given by,

$$\text{eax}(t) = \sum_{m} A_m \frac{(2\pi \text{ERB})^4}{6} (t - \tau_m)^3 e^{-2\pi (0.108 f_m + 24.7)(t - \tau_m)}$$

$$\times \cos[2\pi f_m(t - \tau_m)],$$  

(8)

where $A_m$ are the amplitudes, proportional to the HL values (Eq. (2)), and $(2\pi \text{ERB})^4/6$ is a normalization factor [56].

Let us illustrate the design process of an EAE by gammatones for a subject with the pure tone audiogram shown in Figure 4. The subject exhibits mild losses (30 dB maximum) in the right ear and mild to moderate losses (minimum 20 dB at 125 Hz and maximum 50 dB at 8 kHz) in the left ear. Figure 5 shows the 40 gammatones of the first 10 s of each ear (10 s, 4 pulses/s). The maximum amplitude of the gammatones is normalised to 0.95 to avoid clipping effect. The amplitudes of the left ear are larger, as the HLs for this ear are higher. For each ear, the gammatones have amplitude proportional to the HL at each frequency and random frequency. The varying amplitude nature of the gammatones is best appreciated in Figure 6, where the spectra of the first 240 gammatones (60 s, 4 pulses/s) are shown superimposed to the left and right HL curves. Notice that there are gammatones at frequencies other than these measured by the audiogram. In fact, $\text{HL}(f_m)$ is a function measured at some discrete frequencies (for instance [125, 250, 500, 750, 1000, 1500, 2000, 3000 4000, 6000, 8000] Hz). Thus, this discrete function must be interpolated on a constant frequency bin axis. The contour maps of the short-time Fourier transforms of the first 8 gammatones (2 s, 4 pulses/s) of each ear (Fig. 7) illustrate their random frequency characteristic. Notice that, whereas the gammatones have random nature along time, the left and right ears are stimulated at the same frequencies for each instant (diotic listening).

Hence, the sequence with gammatones provides an EAE customised for each patient, as it matches their HL. Furthermore, it affords the most selective and advanced...
stimulus for tinnitus therapy, as it drives selectively the damaged auditory system with the corresponding auditory filter, in the way suggested by the previous works reviewed in Section 3.1.

### 3.3 Matched broadband noise

With gammatone sequences the hearing frequency band is stimulated with tones at frequencies randomly chosen and amplitudes proportional to the hearing losses at these frequencies. A broadband noise filtered by the hearing loss curve could alternatively be used. As proposed by Schaette and Kempter [45, 46] such a filtered (or coloured) broadband signal should also stimulate optimally the auditory system of the subject with amplitude proportional to the hearing loss.

A filtered broadband noise, $f_{bb}(t)$, is then defined as,

$$f_{bb}(t) = \mathcal{F}^{-1}\{\text{RAND}(f) \cdot \text{HL}(f) \cdot A(f)\},$$

being $\text{RAND}(f)$ a random signal in the hearing frequency band, $\text{HL}(f)$ is an interpolated hearing loss curve in the same frequency band, $\mathcal{F}^{-1}$ denotes inverse Fourier transform, and $A(f)$ is the constant frequency bin version of the amplitude weighting function, Equation (2). The log-spectra corresponding to the right and left ears of the subject, superimposed to the hearing loss curves, are shown in Figure 8.

Although HL is considered a high risk factor to suffer of tinnitus, a percentage varying between 10 and 30% of tinnitus can occur without simultaneous hearing loss [41, 57]. Nevertheless, hearing losses above 8 kHz [14] or cochlear synaptopathy [58], can co-occur with normal audiograms up to 8 kHz. Notice that, for these patients with flat HL, the proposed EAE with filtered broadband noise becomes a conventional TRT.

### 4 Results

The effectiveness of EAE as an optimal sound stimulus for tinnitus therapy is illustrated here with three cases extracted from our currently in process clinical study [59, 60]. This study, approved by the CSIC Ethical Subcommittee, runs since January 2018, and is aimed to demonstrate the efficacy of short treatment (4 months, 1 h/day) with customised EAE in a selected cohort of Spanish tinnitus patients (150 patients until today). All the participants gave their signed informed consent to use their data for publication maintaining their anonymity.

The treatment consisted of a combination of counselling and sound therapy. Counselling was provided at the beginning of treatment, during a session of roughly 90 min, and consisted of a presentation of the functioning of the auditory system, the mechanisms, epidemiology, and possible treatments of tinnitus with realistic expectations. Participants fulfilling the inclusion criteria were informed about the fundamentals of this therapy and requested to choose, with the help of a custom-designed graphical user interface, between either sequential (gammatones) or broadband sound, both matched to their HL curves. The principal criteria for the choice were the listening comfortability and the need to adjust the playing volume just below their tinnitus level sensation (mixing point) [23]. Most participants chose matched broadband noise, since is easier to adjust the volume to the mixing point with continuous sound.
The sound stimuli are designed in our laboratory, taking into account the HL curves of the patients, and are given to them in an easily playable audio format, with the prescription of listening for 1 h per day using high quality headphones.

Tinnitus severity of participants was assessed using a validated Spanish version of the Tinnitus Handicap Inventory (THI) [61, 62]. The handicap produced by tinnitus is graded as slight for THI < 16, mild for 18 ≤ THI < 36, moderate for 38 ≤ THI < 56, severe for 58 ≤ THI < 76, and catastrophic for THI ≥ 78 [63]. It is accepted that a reduction of THI of 20 points is clinically relevant [23].

Participant #1 is a 39-year-old man who was included in our clinical study in December 2019. He suffered from fluctuant, severe tinnitus (THI = 58) at the right ear since 30 months ago, likely due to stress. His audiogram (Fig. 9) showed normal hearing except for a slight HL (25 dB HL) in the right ear at 8 kHz. His tinnitus sound was matched to a hissing of 3 kHz with bandwidth of 15%. He was treated with a matched broadband noise along four months. The time progress of his THI is illustrated in Figure 10. As it can be seen, his tinnitus handicap was reduced in two grades (from severe to mild) amounting a total THI decrease of ΔTHI = −34.

Participant #2 is a 66-year-old man who initiated our tinnitus treatment in June 2020. He was ached of fluctuant, severe tinnitus (THI = 70) for 6 months, likely triggered by a diagnosed ankylosing spondylitis which had caused a moderately severe conductive loss in his left ear (Fig. 11) below 1 kHz. He also displayed mild losses in both ears at middle frequencies (between 2 and 4 kHz). He perceived in his left ear a hissing of 300 Hz with bandwidth of 25%. He was treated with a personalised gammatone sequence during four months. His tinnitus handicap, Figure 12, was reduced in three grades (from severe to slight) amounting a total THI decrease of ΔTHI = −56.

Participant #3 illustrates a mixed treatment case. She is a 45-year-old woman who enters our study in January 2018. She suffered of fluctuant, bilateral, moderate tinnitus (THI = 56) since 14 months ago. She presented an increasing with frequency HL which varied from slight at low frequencies to moderately severe at high frequencies, Figure 13. Her tinnitus sounded as a mix of hissing at 100 and 2000 Hz (with bandwidth of 50% in both cases) and a kind of cymbals. She was subjected to an EAE consisting of a gammatone sequence during the first two months. After that, she changed to a matched broadband noise along the last two months. After four months of mixing.
EAE treatment, her THI decreased two grades (from moderate to slight), Figure 14, with a total THI reduction of $\Delta$THI = 44.

Therefore, independent of patient age, gender, tinnitus aetiology, tinnitus sound, or baseline handicap degree, EAE with either matched broadband noise or gammatone sequence was able to provide clinically significant tinnitus relief on these patients.

5 Conclusions

A sound stimulus consisting of either a sequence of gammatones or a broadband noise, both matched to the hearing loss curves of the subject, is proposed as a personalised and refined sound therapy for tinnitus. The sequential EAE already demonstrated its capability to restore the reorganized tonotopic map of cats produced by traumatizing noise, or to rescale the aberrant gain sensitivity of hyperacusis patients. The HL-matched broadband noise showed to be the optimal sound stimulation for reverse the tinnitus induced hypersensitivity in a computational model of the functioning of neurons in the auditory system. This EAE sound therapy have been applied to patients of different aetiology, age, gender or tinnitus sound, and lateralization, providing a clinically relevant handicap relief in just four months.

Conflict of interest

The authors declare that they do not have any conflict of interest.

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