1	The Derived-Band Envelope Following Response and its Sens		
2	tivity to Sensorineural Hearing Deficits		
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28 Abstract

The envelope following response (EFR) has been proposed as a non-invasive 29 marker of synaptopathy in animal models. However, its amplitude is affected 30 by the spread of basilar-membrane excitation and other coexisting sensorineu-31 ral hearing deficits. This study aims to (i) improve frequency specificity of the 32 EFR by introducing a derived-band EFR (DBEFR) technique and (ii) investigate 33 the effect of lifetime noise exposure, age and outer-hair-cell (OHC) damage on 34 DBEFR magnitudes. Additionally, we adopt a modelling approach to validate the 35 frequency-specificity of the DBEFR and test how different aspects of sensorineural 36 hearing loss affect peripheral generators. The combined analysis of simulations 37 and experimental data proposes that the DBEFRs extracted from the [2-6]-kHz 38 frequency band is a sensitive and frequency-specific measure of synaptopathy in hu-39 mans. Individual variability in DBEFR magnitudes among listeners with normal 40 audiograms was explained by their self-reported amount of experienced lifetime 41 noise-exposure and corresponded to amplitude variability predicted by synaptopa-42 thy. Older listeners consistently had reduced DBEFR magnitudes in comparison 43 to young normal-hearing listeners, in correspondence to how age-induced synap-44 topathy affects EFRs and compromises temporal envelope encoding. Lastly, OHC 45 damage was also seen to affect the DBEFR magnitude, hence this marker should be 46 combined with a sensitive marker of OHC-damage to offer a differential diagnosis 47 of synaptopathy in listeners with impaired audiograms. 48

49 Keywords

derived-band envelope following response; cochlear synaptopathy; sensorienu ral hearing-loss; supra-threshold hearing deficits

52 1. Introduction

Struggling to understand speech in noisy environments is a prevalent complaint 53 of the ageing population, even when they have normal audiometric thresholds. 54 Although hearing thresholds are informative about the sensory function of the 55 cochlea, they are insensitive to auditory-nerve (AN) fiber loss, which is the first 56 sign of permanent hearing damage (Kujawa and Liberman, 2009; Liberman and 57 Kujawa, 2017) and related to supra-threshold hearing (Bharadwaj et al., 2014). 58 Recent animal studies have shown that ageing, ototoxicity and overexposure to 59 noise can lead to an irreversible loss of AN synapses, i.e. cochlear synaptopathy 60 (CS), and delayed degeneration of cochlear neurons, while leaving the cochlear 61 sensory hair cells intact (Kujawa and Liberman, 2009; Lin et al., 2011; Liu et al., 62 2012; Furman et al., 2013; Lobarinas et al., 2017; Valero et al., 2017). Even 63 when the noise exposure dose only causes a temporary threshold shift (Kujawa 64 and Liberman, 2009), noise-induced AN fibers degeneration can progress through 65 the lifespan and yield an increased sensitivity of the ear to age-induced hearing 66 dysfunction (Fernandez et al., 2015). Additionally, reduced numbers of spiral 67 ganglion cells in post-mortem histology of human temporal bones with preserved 68 sensory cells, confirmed the existence of age-related CS in humans (Makary et al., 69 2011; Viana et al., 2015; Wu et al., 2019). Thus, noise exposure and ageing are 70 important causes of CS, a deficit which compromises the temporal coding fidelity 71 of supra-threshold sound as a result of a reduced number of afferent AN synapses 72 innervating the inner hair cell (Bharadwaj et al., 2014, 2015). 73

Since the discovery of CS, several attempts have been made to associate changes in indirect and non-invasive measures of auditory function such as scalp-recorded auditory evoked potentials (AEPs) to the histologically quantified degree of AN fibers loss in animals. For example, auditory brainstem responses (ABRs), evoked by transient stimuli and reflecting the synchronized onset responses of AN fibers

(Don and Eggermont, 1978) showed a decreased supra-threshold wave-I amplitude 79 after synaptopathy due to noise-exposure (Kujawa and Liberman, 2009; Lobarinas 80 et al., 2017; Lin et al., 2011), despite recovered normal distortion product otoa-81 coustic emission (DPOAE) and ABR thresholds. The number of AN fibers can 82 also be quantified using envelope following responses (EFRs), which capture how 83 well AN fibers can phase-lock to the stimulus envelope (Joris and Yin, 1992). The 84 EFR can be extracted from scalp-electrodes in response to a sinusoidally ampli-85 tude modulated (SAM) pure-tone stimulus (Bharadwaj et al., 2014), and has been 86 proposed as an AEP-based measure of CS (Shaheen et al., 2015; Parthasarathy 87 and Kujawa, 2018). 88

Despite the strong relation between AEP markers and CS in animal studies, the 89 indirect nature of AEP recordings hinders a clear and direct interpretation of re-90 sponse strength in terms of CS. First of all, a mixture of sources contribute to scalp 91 potentials, some of which are electrical activity induced by subject-specific factors 92 and unrelated to the sound-driven response (e.g. head size, age, sex, geometry 93 of the generators and physiological noise level; Trune et al., 1988; Mitchell et al., 94 1989; Bharadwaj et al., 2014; Plack et al., 2016). Other sources relate to the sound-95 driven response but depend on outer-hair-cell (OHC) health (Gorga et al., 1985) 96 or cochlear tonotopy (Don and Eggermont, 1978). Lastly, the scalp-recorded AEP 97 is strongly influenced by stimulus characteristics and the corresponding spread 98 of basilar-membrane (BM) excitation, which can confound a frequency-specific 99 diagnosis of CS (Bharadwaj et al., 2014, 2015; Verhulst et al., 2018a; Encina-100 Llamas et al., 2019). To address these issues, several studies have proposed dif-101 ferential/relative AEP-based metrics: the EFR amplitude slope as a function of 102 modulation depth (Bharadwaj et al., 2014, 2015; Guest et al., 2018), ABR wave-V 103 latency changes in different levels of background noise (Mehraei et al., 2016), or the 104 combined use of noise-floor corrected EFRs with ABRs to segregate mixed hear-105

ing pathologies and normalize inter-individual variabilities (Vasilkov and Verhulst, 106 2019, preprint). Secondly, a number of techniques have been proposed to confine 107 ABR generation to specific frequency bands: the use of simultaneous off-frequency 108 masking paradigms, i.e. the derived-band ABR (Eggermont, 1976; Don and Eg-100 germont, 1978), tone-burst ABRs (Rasetshwane et al., 2013) and notched noise 110 paradigms (Abdala and Folsom, 1995). Lastly, asynchrony of low-spontaneous 111 rate (LSR) AN fibers to the transient stimulus (Bourien et al., 2014) may limit 112 the use of the ABR wave-I amplitude to capture all aspects of CS, as noise-induced 113 CS might preferentially affect LSR AN fibers (Furman et al., 2013). 114

This study proposes the use of a relative derived-band EFR method (DBEFR), 115 to confine the EFR to a specific frequency band. To construct DBEFRs, we 116 changed the bandwidth of the stimulus on the low-frequency side rather than 117 using off-frequency masking methods. Thus, a consecutive subtraction of re-118 sponses to stimuli with various bandwidths will yield a relative measure of supra-119 threshold sound coding. We further hypothesize that the relative metric design 120 of the DBEFR reduces the impact of subject-specific factors and increases its 121 sensitivity to individual sensorineural hearing deficits. DBEFR magnitudes were 122 extracted from individuals in four groups to study their applicability to diagnose 123 sensorineural hearing deficits: (1) a young normal-hearing control group, (2) a 124 group with self-reported hearing difficulties in noisy environments, (3) a group 125 of older listeners with normal audiograms and (4) an age-matched group with 126 sloping high-frequency audiograms. We assumed that the second group might be 127 affected by CS due to noise overexposure or ageing and that the third group might 128 be affected by age-induced CS, without co-occuring OHC damage. Aside from 129 collecting DBEFRs, we assessed individual OHC function using audiometric and 130 DPOAE thresholds. In line with animal studies of age-related and noise-induced 131 synaptopathy, we expect that the DBEFR will be reduced in all but the control 132

133 group.

Because, a direct assessment of the individual degree of OHC and AN damage 134 is presently experimentally impossible, we complemented our experimental work 135 with a modelling approach to better understand the relationship between sen-136 sorineural pathologies and their effect on the peripheral generators of the DBEFR. 137 Models can study how AN fiber and sensory hair cell damage impacts the EFR 138 generators to understand their respective roles for DBEFR generation (Verhulst 139 et al., 2016, 2018a,b). We adopt a biophysically inspired model of the human au-140 ditory periphery calibrated for ABR and EFR simulation (Verhulst et al., 2018a) 141 and considered the simulations together with the data to interpret the implications 142 of our findings for DBEFR-based hearing diagnostics. 143

¹⁴⁴ 2. Materials and Methods

Two experiments were conducted at two recording locations. In the first exper-145 iment (Ghent University), normal-hearing (NH) and listeners with self-reported 146 hearing difficulties (NHSR) participated. In the second experiment (Oldenburg 147 University), a total of 43 participants were recruited in three groups: a young NH 148 control group (vNH), an older NH group (oNH) and an older group with sloping 140 high-frequency audiogram (oHI). Ethical approvals were obtained from Ghent and 150 Oldenburg Universities and all participants were informed about the experimental 151 procedures and signed an informed consent before the experiment. 152

153 2.1. Participants

16 NH listeners with ages between 18 and 30 (NH: 24.21±4.10 years, five females) and 9 NH subjects with self-reported hearing difficulties (NHSR) with ages between 23 to 49 (NHSR: 33.78±8.57 years, three females) participated in the first experiment. The NHSR participants were recruited using a flyer asking

whether they had speech understanding difficulties in the presence of background 158 noise, while not presently being treated for hearing disorders. Measurements were 159 conducted in two sessions per subject, with a maximum sound exposure time of 90 160 minutes per session. The participants filled out a questionnaire, in which they were 161 asked how often (yearly, monthly, weekly or daily) they had been playing a musical 162 instrument in a band, attended festivals, concerts or discotheques and used noisy 163 tools during their lifetime. Moreover, the total number of noise-exposed sessions, 164 their duration and estimated noise loudness (a score between 1 to 5) were also 165 assessed (Degeest et al., 2014). Audiograms were measured with an Interacoustics 166 Clinical Computer Audiometer (AC5) at ten standard frequencies between 0.25167 and 8 kHz. 168

The second experiment was conducted with three participant groups composed 169 of: 15 young normal-hearing (yNH: 24.53 ± 2.26 years, eight female), 16 old normal-170 hearing (oNH: 64.25 ± 1.88 years, eight female) and 12 old hearing-impaired (oHI: 171 65.33 ± 1.87 years, seven female) participants. All yNH participants had pure-172 tone thresholds below 20 dB-HL at all measured frequencies between 0.125 and 173 10 kHz (Auritec AT900, Hamburg, Germany audiometer). In both experiments, 174 the audiometrically better ear was chosen for the experiment and stimuli were 175 presented monaurally while participants were seated in a comfortable chair in an 176 acoustically and electrically shielded sound booth, watching silent movies with 177 subtitles to stay awake. Figure 1 shows audiograms of the subjects in all groups. 178 From here on, \triangle stands for the NH group in the first experiment, \Box for NHSR 179 group, \Diamond for yNH in the second experiment, \bigcirc for oNH and \triangleleft for oHI group. 180

181 2.2. Distortion Product Otoacoustic Emissions (DPOAEs)

In the first experiment, DPOAEs were recorded to ten primary-level pairs, (L_1, L_2) , at nine primary-frequency pairs: $f_2 = [546, 780, 1002, 1476, 1998, 3012, 3996, 1002, 1476, 1998, 3012, 3996]$

6006, 8003] and $f_1 = f_2/1.2$. L₂ ranged from 20 to 65 dB-SPL in 5 dB steps and L₁ 184 $= 0.4L_2 + 39$ dB, according to the scissors paradigm (Kummer et al., 1998). The 185 nine primary frequency pairs were chosen to have complete stimulus periods of the 186 primaries in each pair. For each frequency and level pair, 45 repetitions were gener-187 ated in MATLAB 2016b and an ER-10X extended-bandwidth Etymotic Research 188 probe system was used to deliver the two pure tones via a loudspeaker/microphone 189 probe inserted in the ear-canal using a silicone eartip. The response was recorded 190 and digitized using a Fireface UCX external sound card (RME). The pure tones 191 were calibrated separately using a B&K artificial ear and B&K sound level meter 192 at each primary frequency, separately. The time-domain ear-canal recordings were 193 converted to pressure using the microphone sensitivity (50 $\frac{mV}{Pa}$) and pre-amplifier 194 gain (40 dB). Then, I/O functions were calculated for the measured primary-195 frequency pairs by defining the L_{DP} as the averaged spectrum magnitude at the 196 $2f_1$ - f_2 cubic distortion frequency, multiplied by $\frac{2}{N\sqrt{2}}$, where N is the number of 197 samples at each f_2 response. Finally, a linear function, i.e. $L_{DP} = aL_2 + b$, was fit 198 to the bootstrapped data-points and the crossing point with $L_{DP}=0$ Pa was defined 199 as the DPOAE threshold at the measured f_2 frequency. DPOAEs in the second ex-200 periment were acquired using a custom-made software (Mauermann, 2013) which 201 implements a primary frequency sweep method at a fixed f_2/f_1 of 1.2 (Long et al., 202 2008). The primary frequencies were swept across an 1/3 octave range around the 203 $f_2 = 4$ kHz geometric mean with a duration of 2s/octave. Primary levels were cho-204 sen according to the scissors paradigm (Kummer et al., 1998). DPOAE threshold 205 at each frequency was calculated by fitting a linear function to the bootstrapped 206 data-points and was extrapolated to cross $L_{DP}=0$ Pa. Additional details on the 207 experimental procedure can be found in Verhulst et al. (2016). 208

209 2.3. Envelope Following Responses (EFRs)

The EFR stimuli in the first experiment were five filtered white noise carriers, 210 which were 100% modulated with a 120-Hz sinusoid. To generate them, the white 211 noise was filtered between the following frequency regions: [0.25-22], [0.5-22], [1-212 22, [2-22] and [4-22] kHz, using a 1024th order FIR band-pass filter designed by 213 the Blackman-window method. In each frequency band, a stimulus with a duration 214 of 1.25 s was generated in MATLAB 2016b, windowed with a 1.25% cosine-tapered 215 window and delivered monaurally over ER-2 earphones, connected to a Fireface 216 UCX external sound card (RME) and a TDT-HB7 headphone driver. A uniformly-217 distributed random silence jitter was applied between consecutive epochs (200 218 $ms\pm 20 ms$) of the 370 stimulus presentations. Stimuli with various bandwidths 219 were calibrated to have the same spectral magnitude, i.e. the widest bandwidth 220 stimulus was presented at 70-dB-SPL, while narrower bandwidth stimuli had lower 221 sound pressure levels to preserve an equal spectral level in all conditions. The 222 calibration was performed using a B&K sound-level-meter type 2606. Figure 2a 223 illustrates the designed stimuli in the frequency domain. Scalp-recorded potentials 224 were obtained with a 64-Channel Biosemi EEG recording system and a custom-225 built trigger box using a sampling frequency of 16384 Hz. The electrodes were 226 placed according to the 10-20 standard, using highly conductive gel (Signa gel). 227 The Common Mode Sense (CMD) and Driven Right Leg (DRL) electrodes were 228 placed on top of the head. Six external channels were used as well, i.e. two 229 earlobe electrodes as reference and the remaining electrodes were placed on the 230 forehead and cheeks to record electrical activity induced by horizontal and vertical 231 eye movements. All channels were re-referenced to the average of the two earlobe 232 electrodes. 233

In the second experiment, four EFR stimuli with white noise carriers were band-pass filtered using the same filter as in the first experiment in [0.3-16], [0.7-

16], [2.8-16] and [5.6-16] kHz frequency regions. The precise lower cut-off fre-236 quencies employed in the band-pass filtering were $\frac{0.5}{\sqrt{2}}$, $0.5\sqrt{2}$, $\frac{4}{\sqrt{2}}$ and $4\sqrt{2}$ kHz, 237 respectively. Stimuli were 95% modulated with a 120-Hz pure tone and presented 238 at 70 dB SPL using the same configuration as the first experiment. The stimuli 230 had a duration of 400 ms, were 2.5% ramped with a tapered-cosine window and 240 presented 1000 times using a uniformly distributed random inter-stimulus silence 241 jitter of 100 ms ± 10 ms. The calibration was performed in the same way as for 242 the first experiment, but using B&K sound level meter type 2610. A 64-channel 243 Biosemi EEG system was adopted to record the responses using EEG caps with 244 equidistant electrode spacing. The CMS and DRL electrodes were located on the 245 fronto-central midline and on the tip of the nose of the participants, respectively. 246

247 **3.** EFR Analysis

Acquired EFRs were first filtered using an 800th order Blackman window-based 248 FIR filter between 60 and 600 Hz, using the filt filt function of MATLAB to avoid 240 time delays and phase shifts. Signals were broken into 1-s long epochs relative to 250 the trigger onset, from 0.25 to 1.25 s in the first and into 0.3-s long epochs, from 251 0.1 to 0.4 s in the second experiment. Baseline correction was applied before the 252 epochs were averaged across trials. 30 and 100 epochs were rejected on the basis of 253 the highest peak-to-trough values in the first and second experiment, respectively. 254 Since the firing patterns of neurons are influenced by factors such as instantaneous 255 external inputs, previous firing patterns and the general state of the system, the 256 interpretation of the raw EFR spectrum resulting from the Fast Fourier Transform 257 (FFT) of the averaged epochs is challenging. Synaptic delays and axon conduc-258 tion limitations cause a $\frac{1}{f}$ behaviour in EEG (Buzsaki, 2006, Chapter 10) and it is 259 crucial to suppress this noise-floor to analyse the stimulus-driven spectrum. The 260 bootstrapping approach proposed in Zhu et al. (2013) was employed to estimate 261

the $\frac{1}{f}$ noise-floor component. First, 340 epochs were drawn randomly with re-262 placement, among the 340 epochs (900 epochs in the second experiment). Then, 263 the FFT of these epochs were averaged. This procedure was repeated $N_1=200$ 264 times ($N_2=400$ for the second experiment), resulting in a nearly Gaussian dis-265 tribution of raw, averaged spectra. The average value of this distribution yielded 266 the frequency domain representation of the EFRs. Afterwards, the same procedure 267 with $M_1 = 1000$ repetitions ($M_2 = 1200$ for the second experiment) and phase-flipped 268 (180°) odd epochs was followed to estimate the spectral noise-floor as a function 269 of frequency. The idea behind this approach is that the time-locked response is 270 suppressed if the averaging is repeated sufficiently across phase-inverted epochs. 271 Finally, the averaged absolute values of the estimated noise floors were subtracted 272 from the averaged absolute values of the EFR spectra amplitudes to obtain the 273 stimulus-driven EFR spectrum: 274

$$EFR_{raw}(f) = \frac{2}{n_p} \left| \frac{\sum_{i=1}^{N} FFT(X_i)}{N_p} \right|$$
(1)

Noisefloor(f) =
$$\frac{2}{n_p} \left| \frac{\sum_{j=1}^{M} FFT([-1]^j X_j)}{M_p} \right|$$
 (2)

$$EFR_{Spec}(f) = EFR_{raw}(f) - Noisefloor(f)$$
 (3)

275

X represents the epochs vector, N the number of bootstrap repetitions, M the number of repetitions to estimate the noise-floor, p the experiment number (i.e. one or two) and n equals the number of FFT points ($n_1=16384$ and $n_2=8192$). Figure 3 represents EFR_{raw}, Noisefloor and EFR_{Spec} spectra of subject No. 8 from NH group in the first experiment. All EFR_{Spec} peak values which were four standard deviations above the noise-floor (EFR_{SpecSD}) for frequencies corresponding to the modulation frequency (120 Hz) and its following two harmonics (240 and 360 Hz) were added to yield EFR magnitude of the corresponding condition.

$$EFR_{PtN} = \sum_{k=0}^{2} EFR_{SpecSD}(f_k), \qquad f_k = 120 \times (k+1)$$
(4)

To construct DBEFRs, the calculated EFR_{PtN} for each narrower-band condition was subtracted from the following wider-band condition using:

$$DBEFR_{PtN} = \begin{cases} (EFR_{PtN})_{wide} - (EFR_{PtN})_{narrow}, & (EFR_{PtN})_{wide} > (EFR_{PtN})_{narrow} \\ 0, & (EFR_{PtN})_{wide} \le (EFR_{PtN})_{narrow} \end{cases}$$
(5)

Derived frequency bands from EFRs to the first experimental stimuli are shown schematically in Fig. 2b.

²⁸⁸ 4. Questionnaire analysis

The completed questionnaires from the participants in the first experiment 289 were used to estimate the individual life-time noise exposure dose. To this end, the 290 collected individual data related to the frequency and duration of experienced noise 291 exposure were converted to a number of sessions per year multiplied by the duration 292 and the personal estimated noise loudness scores, i.e. a number between 1 and 5. 293 We followed the procedures as described in Degeest et al. (2014). The scores were 294 separately calculated for questionnaire categories: (i) playing musical instrument 295 in a band, (ii) attending festivals, concerts and discotheques and (iii) using noisy 296 tools. Outcomes were normalized across NH and NHSR groups participants by 297 the highest reported dose, i.e. 30600, 18480 and 26000 hours in each category, 298

²⁹⁹ respectively.

300 5. Model Simulations

A biophysical model of the human auditory periphery (Verhulst et al., 2018a), 301 schematically shown in Fig. 4, was adopted to simulate the experimental con-302 ditions and to investigate the effect of different aspects of sensorineural hearing 303 deficits on the EFR_{PtN} and DBEFR_{PtN} magnitudes. The original implementation 304 of the model is described in Verhulst et al. (2018a) and can be downloaded from 305 "https://github.com/HearingTechnology/Verhulstetal2018Model". The parameters 306 which determine the weights between the population AN, cochlear nucleus (CN) 307 and inferior colliculus (IC) responses were adjusted along with the AN innervation 308 patterns across CF for the purpose of this study. 309

310 5.1. Auditory nerve-fiber distribution

The original model implementation introduced the same number of synapses 311 between inner-hair-cells (IHCs) and AN fibers for all simulated characteristic fre-312 quencies (CF), whereas human and rhesus monkey innervation patterns show a 313 bell-shaped pattern across CF. To make the model more realistic, the averaged 314 synaptic counts of four control rhesus monkeys (seven ears) and nine frequencies 315 (Valero et al., 2017) were mapped to corresponding fractional distances of the 316 human cochlea using the monkey place-frequency map (Greenwood, 1990). Frac-317 tional distances from the base of cochlea, d_i , were calculated according to the 318 measured frequency points (f_{RM_i}) : 319

$$f_{\rm RM_i}[\text{in Hz}] = 360(10^{2.1(1-d_i)} - 0.85), \qquad i = 1, 2, ..., 9$$
 (6)

320

321 The obtained d_is were substituted into the analogous Greenwood map equation

for humans, yielding the corresponding frequency points (f_{H_i}) :

$$f_{\rm H_i}[{\rm in \ Hz}] = 165.4(10^{2.1(1-d_i)} - 0.88), \qquad i = 1, 2, ..., 9$$
 (7)

To calibrate the model with the applied AN pattern, a 70 dB-nHL click-train containing both stimulus polarities was presented at a rate of 11 Hz. To perform this calibration, simulated ABR wave amplitudes were matched to the experimental data on the basis of 55 averages. Specifically, the $M_1 = 4.6729 \times 10^{-14}$, $M_3 = 5.6885 \times 10^{-14}$ and $M_5 = 14.641 \times 10^{-14}$ parameters were adjusted on the basis of average NH ABR wave-I, III and V reference data from Picton (2010), i.e. $w_I = 0.15 \mu V_p$, $w_{III} = 0.17 \mu V_p$ and $w_V = 0.61 \mu V_{pp}$.

Using the synapse counts from rhesus monkey and the mapped frequency points 330 for the human cochlea (f_{H_i}) , a "smoothing spline" curve was fit to estimate the 331 number of synapses across all frequency channels in the model. Finally, to simulate 332 different AN fiber types, i.e. high spontaneous-rate (HSR), medium spontaneous-333 rate (MSR) and LSR fibers, and their properties, the obtained population dis-334 tribution was multiplied by the corresponding AN type proportion factor C, i.e. 335 $\mathrm{C}_{\mathrm{HSR}}=0.60,\ \mathrm{C}_{\mathrm{MSR}}=0.25$ and $\mathrm{C}_{\mathrm{LSR}}=0.15$ (Liberman, 1978, cat data), before 336 responses were summed at each simulated CF and fed to the CN model. The sim-337 ulated frequency-specific AN fibers distribution is shown on the top-right column 338 of Fig. 4. 339

340 5.2. Stimuli

The model stimuli were matched to the experimental conditions and had a duration of 600 and 400 ms for the first and second experiment, respectively. Twenty stimulus repetitions with different white noise iterations were applied to the model and simulations were averaged before the EFR_{PtN} was calculated using the same procedure as in Eq. 4. The amplitudes of the model stimuli were set based on the broadest condition, i.e. 0.25 to 22 kHz for the first experiment and 0.3 to
16 kHz for the second experiment to yield an input of 70 dB SPL. The narrower
band stimuli were calibrated relative to the broadest condition, such that they had
the same spectral level as the broadband condition but with a different SPL.

³⁵⁰ 5.3. Simulating sensorineural hearing loss

The simulated CS profiles and their corresponding AN fiber types are shown 351 in Fig. 4. Different degrees of CS were modelled by manipulating the number 352 and types of AN fibers. The table in Fig. 4 shows the simulated synaptopathy 353 profiles. OHC damage was simulated by changing the CF-dependent mechanical 354 gain of the cochlea by moving poles of the BM admittance function to yield a filter 355 gain reduction corresponding to a desired dB-HL-loss, which also yielded wider 356 cochlear filters. The inset in Fig. 4 shows the simulated cochlear gain loss profiles. 357 Procedures are further detailed in Verhulst et al. (2016, 2018a). 358

359

360 6. Results

³⁶¹ 6.1. EFR and dependence on stimulus frequency

Figure 5 shows individual and group-mean EFR_{PtN} magnitudes to different 362 frequency bandwidths in the first (panel a) and second (panel b) experiments. 363 Despite within-group individual variability, experimental group-means revealed 364 approximately constant EFR_{PtN} magnitudes to stimuli with frequencies below 365 2 kHz and reduced magnitudes to frequencies above 2 kHz and 2.8 kHz in the 366 first and second experiment, respectively. A paired-sample t-test with Bonferroni 367 correction was applied to compare EFR_{PtN} magnitudes to stimuli with different 368 frequency bandwidths in each group. In the first experiment, a single significant 369 difference was observed between the $EFR_{[2-22]}$ and $EFR_{[4-22]}$ conditions in NH 370

group (t(11)=7.02, p<0.0000; specificed by # in Fig. 5a), which disappeared for 371 the NHSR group (t(8)=3.13, p=0.014). In the second experiment, a paired-sample 372 t-test with Bonferroni correction gave a significant difference between $EFR_{[2.8-16]}$ 373 and $EFR_{[5.6-16]}$ in yNH (t(12)=7.86, p<0.0000; specificed by + in Fig. 5b) and 374 oNH groups (t(12)=6.21, p<0.0000; specificed by ++ in Fig. 5b), but not in the 375 oHI group (t(9)=2.03, p=0.072). Simulated NH-EFRs are shown in hexagons in 376 Fig. 5 and corroborate experimental findings by showing a minor contribution of 377 stimulus frequencies below 2 kHz on the EFR generation. 378

379 6.2. Derived-Band Envelope Following Responses (DBEFRs)

DBEFR_{PtN} magnitudes calculated using Eq. 5 are shown in Fig. 6 for the first 380 (panel a) and second (panel b) experiment. A paired-sample t-test with Bonferroni 381 correction comparing the $DBEFR_{PtN}$ magnitudes in each group revealed only a 382 significant difference between the [1-2] and [2-4] kHz condition in the NH group 383 (t(11)=-3.99, p=0.002; specificed by # in Fig. 6a). In the second experiment, 384 paired-sample t-test showed significant difference between [0.3-0.7] and [2.8-5.6]-385 kHz conditions only in yNH group (t(12)=-7.00, p<0.000; specificed by + in Fig.386 6b). In support of our experimental findings, simulated NH-DBEFR magnitudes 387 in both experiments (shown by hexagons in Fig. 6a and b) were equal for derived-388 bands below 2-kHz and increased for $\text{DBEFR}_{[2-4]}$ (in the first experiment) and 389 $DBEFR_{[2.8-5.6]}$ (in the second experiment). In line with EFR_{PtN} findings in Sec-390 tion 6.1, experimental and simulated $DBEFR_{PtN}$ magnitudes in both experiments 391 showed an increased contribution of the [2-6] kHz derived frequency band to the 392 EFR generation. 393

³⁹⁴ 6.3. Possible origins of individual EFR differences

Previous studies have shown a dependency of the scalp-recorded AEP magnitude to head size, sex and age (Trune et al., 1988; Mitchell et al., 1989; Vasilkov and Verhulst, 2019, preprint). Hence, the spread of data-points within different recorded test-groups and spectral bandwidths could be explained by subjectspecific factors unrelated to hearing or hearing-related factors associated with the main factors for grouping: (i) self-reported hearing difficulties in noisy environments in the first experiment, (ii) age and (iii) elevated hearing thresholds in the second experiment.

Pooling together the NH and NHSR EFR_{PtN} magnitudes, a regression analysis 403 was conducted to investigate the effect of age, 4 kHz threshold, head size and 404 $DPTH_{3000}$ on the $EFR_{[2-22]}$ (Fig. 7, left column) and $DBEFR_{[2-4]}$ magnitude 405 (Fig. 8, left column). None of the regressions showed a relation between tested 406 variables, suggesting that other factors than those reported were responsible for 407 the individual variability among listeners. The regression analysis on EFR_{PtN} and 408 DBEFR_{PtN} magnitudes combined from all experimental groups in the second ex-409 periment (Fig. 7 and 8, right column) showed a meaningful correlation of age, 410 threshold, head size and $DPTH_{4000}$ with the $EFR_{[2.8-16]}$ magnitude. However, ex-411 tracting the $\text{DBEFR}_{[2.8-5.6]}$, reduced the correlation with age and 4-kHz threshold 412 and suppressed any meaningful correlation with head-size and $DPTH_{4000}$. More-413 over, excluding the oHI group from the correlation analysis, led to a reduced and 414 insignificant correlation coefficient (R=-0.382, p=0.083) between 4-kHz threshold 415 and $DBEFR_{[2.8-5.6]}$. These results suggest that the proposed DBEFR metric is not 416 affected by head size. Moreover, individual variabilities between the yNH and oNH 417 groups in the second experiment might be related to degraded temporal envelope 418 coding as a consequence of CS (Bharadwaj et al., 2015), given the insignificant 419 correlations of DBEFRs with the 4-kHz threshold, DPTH₄₀₀₀ and head size. 420

421 6.4. EFR_{PtN} and $DBEFR_{PtN}$ magnitude variability across tested groups

To investigate the separability of the recruited groups by means of their DBEFR 422 magnitudes, we analysed the group-mean differences in each experiment. In the 423 first experiment, an independent two-sample t-test comparison between the means 424 of stimulated frequency bandwidths in the NH and NHSR group (Fig. 5a), showed a 425 significant difference only between the [2-22] and [4-22]-kHz conditions (EFR_[2-22]: 426 t(19)=3.36, p=0.003 and EFR_[4-22]: t(19)=2.76, p=0.012). However, significant 427 mean-differences disappeared between similar conditions in the NH and NHSR 428 groups after extracting DBEFR magnitudes in Fig. 6a (DBEFR_[2-4]: t(19)=0.90, 429 p=0.338). The insignificant difference across groups and insignificant correlation 430 coefficients of $DBEFR_{[2-4]}$ with subject-specific factors observed in Fig. 8, might 431 partly be explained by the different amounts of experienced lifetime noise exposure 432 reported in the questionnaires and might point to various degrees of noise-induced 433 CS. Calculated noise scores in Fig. 9 revealed an insignificant correlation with 434 $DBEFR_{[2-4]}$ magnitudes (R=0.13, p=0.089). However, certain cases appeared 435 to be inconsistent with our noise-induced synaptopathy hypothesis, i.e., (i) high 436 noise scores in the NH group, e.g. subject No. 12 and (ii) low noise scores in the 437 NHSR group, e.g. subject No. 1. We suggest that the insignificant group-mean 438 differences can be explained by (i) subject-dependent unreliable discriminating fac-439 tor between NH and NHSR group (Coughlin, 1990), (ii) variability in answering 440 lifetime noise-exposure dose in questionnares (Prendergast et al., 2017; Bramhall 441 et al., 2017), (iii) an insufficient number of samples and (iv) a limited sensitivity 442 of the DBEFR_{PtN} metric to noise-induced CS. 443

In the second experiment, an independent two-sample t-test was applied to investigate the effect of age between the yNH and oNH groups, and elevated highfrequency thresholds between the oNH and oHI groups. This comparison showed a significant effect of age on all frequency bandwidths and a significant effect of hearing threshold on all frequency bands except for the [5.6-16] kHz band (t(21) = -1.81, p = 0.084). The same comparison for the DBEFR magnitudes revealed a significant effect of age and hearing threshold only in the [2.8-5.6]-kHz derived band condition (t(24) = 3.13, p=0.004 and t(21) = -4.60, p = 0.002, respectively), consistent with the correlation presented in Fig. 8. Detailed t and p values of independent two-sample t-tests, evaluating the effect of age and hearing thresholds on EFR and DBEFR magnitudes, are listed in Table. 1.

Our group-mean results combined with the correlation analysis in Section 6.3 455 suggests that the DBEFR metric removes inter-subject variability unrelated to 456 hearing between yNH and oNH groups, but leaves individual magnitude differences 457 within a group meaningful, given the often non-overlapping standard deviations. 458 Consequently, the significant group-mean difference between vNH and oNH might 459 reflect individual degrees of sensorineural hearing loss. To investigate the diagnos-460 tic sensitivity, it is of course necessary to understand the respective role of OHC 461 deficits and CS on DBEFR magnitudes. Given that oHI listeners may suffer from 462 both OHC deficits and CS, it is important to study the impact of OHC-damage 463 and CS, both independently and concomitantly. 464

⁴⁶⁵ 6.5. The EFR relationship to different aspects of sensory hearing-loss

Since OHC-damage and CS might both affect the EFR magnitude (Garrett 466 and Verhulst, 2019: Vasilkov and Verhulst, 2019, preprint), we employed a compu-467 tational model of the auditory periphery to simulate how different degrees of CS 468 affected the EFR_{PtN} magnitude, both in presence and absence of high-frequency 469 sloping OHC-loss above 1 kHz (simulated high-frequency sloping audiograms in 470 Fig. 4). The most sensitive regions of the cochlea responding to a 120-Hz mod-471 ulated broadband noise were identified to lie between the CFs of 2 and 6 kHz 472 (Keshishzadeh et al., 2019). As a result, we only considered two EFR condi-473

tions of each experiment, namely $EFR_{[2-22]}$ and $EFR_{[4-22]}$ in the first experiment 474 (Fig. 10a) and $\text{EFR}_{[2.8-16]}$ and $\text{EFR}_{[5.6-16]}$ in the second experiment (Fig. 10b). 475 Model simulations showed that CS, when no other hearing deficits co-occur, re-476 duces the EFR and DBEFR magnitudes. Applying sloping high-frequency OHC-477 damage increased the DBEFR magnitudes in both experiments (Fig. 10c and d). 478 According to the simulations, the NH DBEFR magnitude reduced by 46% as a 479 consequence of removing 47% of the AN fibers (i.e., the 10-0-0 CS profile defined 480 in Fig. 4), while the Slope20 OHC-damage (defined in Fig. 4) increased the NH 481 DBEFR magnitude by 27%. Hence, the effect of OHC-damage on the DBEFR 482 magnitude is smaller than that of CS alone, however it is not negligible. There-483 fore, the experimental range of individual EFR and DBEFR magnitudes can be 484 explained by different degrees of variation simulated by CS and OHC-damage. 485

Our simulations predicted the experimental observed absolute range of DBEFR 486 magnitudes and explained the experimental differences between vNH and oNH 487 groups on the basis of age-induced CS, not OHC-damage induced differences. Fur-488 thermore, the simulations suggest that oNH and oHI listeners might both suffer 489 from CS. Results are less clear for the NHSR group where there is a strong overlap 490 with the NH group. However, the noise scores from the questionnaires in Fig. 9, 491 could ascribe some of the spread in DBEFR magnitudes within the NH and NHSR 492 groups to noise-induced CS, and to a lesser degree to OHC-damage given all had 493 normal hearing thresholds. 494

It is worthwhile to note that EFR magnitudes in both experiments (Fig. 10a and b), decreased as a result of CS alone and increased by applying high-frequency OHC-damage with a severity of less than 20 dB-HL at 8 kHz. However, higher degrees of OHC-damage reduced the EFR magnitudes. We explain this nonmonotonic behaviour on the basis of the AN fiber discharge rate-level curve, where increased simulated EFR_{PtN} magnitudes (Fig. 10 c and d) and amplitude-

modulated (AM) responses (Fig. 11b) to supra-threshold stimuli (70 dB-SPL) 501 caused by OHC-damage, might stem from the extended dynamic range of the AN 502 fibers for less effective AN-driving levels (Bharadwaj et al., 2014, their Fig. 3c). 503 Given that experimental and simulated stimuli were calibrated to have equal spec-504 tral magnitudes for all stimulus bandwidths, the narrowest bandwidth stimulus 505 was presented at a lower overall sound level than the 70 dB-SPL broadband stim-506 ulus. Thus, applying more severe OHC-loss, lowered the AN discharge rate and 507 envelope synchrony strength (Verhulst et al., 2018a, Fig. 5) and decreased the 508 EFR magnitudes (Verhulst et al., 2018a, their Fig. 7). However, DBEFR magni-509 tudes increased monotonically for all simulated degrees of OHC damage (Fig. 10c 510 and d). 511

512 7. Discussion

513 7.1. Tonotopic sensitivity of the EFR generators

Despite the individual variability within groups, experimental group-mean 514 EFR_{PtN} magnitudes to broadband stimuli with different bandwidths (Fig. 5a), 515 were equal at frequencies below 4 kHz and reduced in response to [4-22] kHz 516 condition. In the second experiment (Fig. 5b), the EFRs remained equal at fre-517 quencies below 5.6 kHz and degraded when the [5.6-16] kHz band was added. 518 Consequently, equal DBEFR_{PtN} magnitudes were obtained for frequencies below 519 2 kHz. Individual variability was best observed for the DBEFR_{PtN} extracted from 520 the [2-4] kHz (first experiment, Fig. 6a) and [2.8-5.6] kHz (second experiment, 521 Fig. 6b) frequency bands. Simulated EFRs to the experimental stimuli shown 522 with hexagons in Fig. 5 and 6, confirmed observed experimental EFR_{PtN} and 523 $DBEFR_{PtN}$ frequency-dependent behaviour. In addition, the model can be used 524 to study which CF regions along the cochlea contributed strongly to the population 525

⁵²⁶ EFR response. To this end, we calculated the AM (Fig. 11a) and derived-band ⁵²⁷ AM (DBAM) responses at each CF (Fig. 11b) as follows:

$$AM_{AN}(N_{CF}) = \frac{1}{n} \sum_{i=0}^{2} \left[2 \left| FFT(AN_{N_{CF}}) \right| \right]_{f_i},$$

$$N_{CF} = 1, 2, ..., 401, f_i = 120 \times (i+1)$$
(8)

$$DBAM_{AN} = |AM_{AN}(wide) - AM_{AN}(narrow)|$$
(9)

AN_{NCF} is the AN-response at N_{CF} channel and $n = n_1$ as was defined in Eq. 1. These simulations corroborate the experimentally-observed minor contribution of low-frequency CF channels to the EFR generation.

In a previous modelling study (Keshishzadeh et al., 2019), we investigated 531 the tonotopic sensitivity of EFR_{PtN} to broadband stimuli and ascribed the poor 532 low-frequency AM coding to a combination of the chosen modulation frequency 533 (120 Hz) and the narrower bandwidth of apical cochlear filters compared to the 534 higher CF filters (Moore and Glasberg, 1983). Model simulations in response to 535 the spectrally broadest condition, i.e. [0.25-22] kHz, modulated with a range of 536 lower modulation frequencies than 120 Hz, showed that the saturation proper-537 ties of AN fibers limited the modulation response at all modulation frequencies 538 at higher CFs despite an enhanced modulated response at the BM. This resulted 539 in a degraded response at CFs above 4 kHz and shifted the frequency sensitivity 540 of AM coding to the lower CFs at low modulation frequencies. Since the brain 541 response to modulation frequencies below 70 Hz may contain cortical as well as 542 brainstem contribution (Purcell et al., 2004; Picton, 2010, Chapter 10), employing 543 low modulation frequencies might render EFR-based CS diagnosis insensitive, even 544 though an improved frequency-sensitivity can be obtained from the apical regions 545 using these lower modulation frequencies. Therefore, the employed experimen-546

tal modulation frequency, i.e. 120-Hz in combination with a broadband carrier,
might be able to establish a frequency-specific CS diagnosis at frequencies above
2 kHz. In this context, the proposed DBEFR method showed a notable contribution of the [2-4] kHz CF region to the EFR generation by showing a significantly
stronger DBEFR_{PtN} magnitude compared to lower derived-band conditions in the
NH group.

553 7.2. Diagnostic Applications

The measured DBEFR magnitudes are individually separable and above the 554 noise-floor even for HI listeners, whose group-mean was significantly above the 555 noise-floor. In addition, the DBEFR offers a frequency-specific metric to assess 556 supra-threshold temporal coding of the population of AN fibers and brainstem 557 neurons in the [2-6] kHz region. Despite these promising results, the diagnostic 558 sensitivity of DBEFRs also has limitations. The proposed DBEFR magnitude is 559 sensitive to CS alone, when no other coexisting hearing deficits occur and is hence 560 applicable for use in ageing listeners with normal audiograms and those with self-561 reported hearing difficulties or prone to noise exposure. However, DBEFRs are 562 also affected by OHC damage (Fig. 10). The metric hence needs to be comple-563 mented with another supra-threshold metric sensitive to OHC damage within the 564 tonotopic range of interest to allow a separation of the CS and OHC aspect of 565 sensorine learning damage from the recorded DBEFRs from listeners with im-566 paired audiograms. 567

Lastly, the employed high modulation frequency, i.e. 120 Hz, suppresses cortical contributions to the EFR_{PtN} magnitudes, but also degrades AM-coding from lower CFs and thereby limits the tonotopic sensitivity of the EFR_{PtN} to frequencies above 2 kHz. Consequently, apical-end supra-threshold hearing deficits would not be reflected in the proposed DBEFR_{PtN} metric even for stimuli which contain frequencies below 2 kHz. These results are consistent with the source generators of
derived-band ABRs (DBABR), which reduce in amplitude for bands below 2 kHz
(Don and Eggermont, 1978). This predominant basal origin of the ABR also confines the potential of ABR/DBABR-based CS diagnosis to basal cochlear regions
(e.g. wave-I amplitude).

578 8. Conclusion

We proposed the use of a relative $DBEFR_{PtN}$ metric to render the EFR_{PtN} 579 frequency-specific and rule out subject-specific factors unrelated to hearing to ap-580 ply it in the study of identifying the origins of sensorineural hearing deficits and 581 clarifying their role in supra-threshold temporal envelope encoding. DBEFR_{PtN} 582 magnitudes from two experiments were analysed and compared to model sim-583 ulations to conclude that the frequency-sensitivity of DBEFR_{PtN} magnitudes to 584 broadband stimuli is limited to the [2-6] kHz bandwidth. Secondly, we showed that 585 the DBEFR metric eliminates inter-subject variability caused by hearing-unrelated 586 sources. Model simulations (Fig. 10) explained the significant difference between 587 vNH and oNH listeners on the basis of CS, which could result from age-induced 588 CS as identified from human post-mortem studies (Makary et al., 2011: Viana 589 et al., 2015; Wu et al., 2019). Supported by model predictions (Fig. 10d), the 590 significant difference between age-matched oNH and oHI groups was explained by 591 OHC-damage and coexisting CS as a consequence of ageing. Accordingly, profound 592 OHC damage may confound DBEFR-based clinical applications of CS diagnosis. 593 Despite this limitation in the differential diagnosis of CS and OHC deficits on 594 the basis of the DBEFR magnitude, the proposed metric can be used to diagnose 595 CS in a frequency-specific manner in listeners with thresholds below 20 dB-HL. 596 Moreover, it provides an objective marker of supra-threshold temporal envelope 597 coding, which can be used to study its role in sound perception studies. Lastly, 598

⁵⁹⁹ our results clearly demonstrate that older listeners with or without impaired audiograms suffer from degraded temporal envelope coding at frequencies above 2 ⁶⁰⁰ kHz.

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759 Tables

Table 1: The results of a two-tailed t test show the effect of age and hearing threshold on EFR and DBEFR magnitudes in the second experiment.

Metric	Frequency Bandwidth	Age Effect	Threshold Effect
	[kHz]	yNH vs. oNH	oNH vs. oHI
	[0.3-16]	t(24) = 5.812	t(21) = -3.020
_		p=0.000	p=0.006
	[0.7-16]	t(24) = 6.632	t(21) = -2.175
EFR		p=0.000	p=0.041
	[2.8-16]	t(24) = 5.836	t(21) = -4.498
		p=0.000	p=0.000
	[5.6-16]	t(24) = 4.734	t(21) = -1.811
		p=0.000	p=0.084
	[0.3-0.7]	t(24) = -2.09	t(21)=-0.86
DBEFR		p=0.050	p=0.40
		t(24)=3.13	t(21) = -4.60
	[2.8-5.6]	p=0.004	p=0.002

760 Figure Captions

Figure 1. Measured audiograms in the first (left) and second (right) experiment. Markers indicate the audiometric threshold at 4 kHz. The dashed line is the averaged audiometric threshold at each group and the yellow shading the standard deviation.

Figure 2. Spectra of the 120-Hz modulated stimuli and derived bands. (a) Designed stimulus spectra in different frequency bands and specified cut-off frequencies of the bandpass filter. (b) Derived bands from the EFRs recorded to the stimuli shown in (a) obtained by spectral subtraction.

Figure 3. Magnitude spectrum of the $\text{EFR}_{\text{raw}}(f)$ (in blue), Noisefloor(f) (in red) and $\text{EFR}_{\text{Spec}}(f)$ (in black) calculated for subject No. 8 from the first experiment. EFR spectra were evoked by the stimulus with the broadest bandwidth, i.e. [0.25-22] kHz. Peaks at the stimulus modulation frequency, and two harmonics (i.e. $f_0 = 120$ Hz, $f_1 = 240$ Hz and $f_2 = 360$ Hz) are clearly visible above the noise-floor.

Figure 4. Modeling approach. The block-diagram shows different levels of the auditory pathway modelled in the employed biophysical model of the hearing periphery (Verhulst et al., 2018a). The top-right graph indicates the simulated distribution of different types of AN fibers across CF. The table shows simulated CS profiles and the graph on the bottom right depicts simulated different degrees of cochlear gain loss. The corresponding simulated thresholds at 8 kHz are indicated by the legend.

Figure 5. EFR_{PtN} magnitudes to 120-Hz modulated stimuli with different white noise carrier bandwidths in the (a) first and (b) second experiment. Individual data-points are depicted with open symbols and standard deviations were obtained using a bootstrapping procedure (Zhu et al., 2013). Filled symbols reflect the group-means and their corresponding standard deviations. Simulated EFRs from a NH model were added in filled hexagons. Significant effects of considered frequency-band on EFR_{PtN} magnitudes are specified by: (#) in the NH-group (first experiment), (+) in the yNH-group and (++) in the oNH-group (second experiment). To enhance the visualization of differences, panel (a) was plotted on narrower y-axis range, therefore the real values of lowered EFR_{PtN} magnitudes were specified next to the corresponding data-points.

Figure 6. DBEFR_{PtN} magnitudes derived using Eq. 5 for 120 Hz modulated stimuli with different white-noise-carrier bandwidths in the (a) first and (b) second experiment. DBEFR_{PtN} for each frequency band was obtained from a wider and narrower width stimulus. Standard deviations were calculated using a bootstrapping procedure and stemmed from averaged responses from 20 stimulus iterations in the model simulations. Group means and standard deviations are depicted using filled symbols. Significant effects of considered frequency-band on NH-group in the first experiment and yNH-group in the second experiment are specified by (#) and (+), respectively. To enhance the visualization of differences, figures were plotted on narrower y-axis range, therefore the real values of lowered $DBEFR_{PtN}$ magnitudes were specified next to the corresponding data-points.

Figure 7. Correlation analysis of $\text{EFR}_{[2-22]}$ ($\text{EFR}_{[2.8-16]}$) with age, audiometric threshold at 4 kHz, head-size and DPTH_{3000} (DPTH_{4000}) in the first (left) and second (right) experiments. Correlation between EFR magnitudes and all factors but age were reported using the Pearson's correlation coefficient. The Spearman's correlation coefficient was calculated to study the effect of age in the second experiment.

Figure 8. Correlation analysis of $\text{DBEFR}_{[2-4]}$ ($\text{DBEFR}_{[2.8-5.6]}$) with age, audiometric threshold at 4 kHz, head-size and DPTH_{3000} (DPTH_{4000}) in the first (left) and second (right) experiments. Correlation between DBEFR magnitudes and all factors but age were reported using the Pearson's correlation coefficient. The Spearman's correlation coefficient was calculated to study the effect of age in the second experiment.

Figure 9. Bar-plots of noise scores acquired from questionnaires of NH and NHSR groups, classified in three categories, i.e. experience noise as a consequence of (i) playing a musical instrument in a band, (ii) attending festivals or concerts and (iii) using noisy tools. Results are shown normalised, where the score of 1 corresponds to 30600, 18480 and 26000 hours of accumulated noise dose on the considered categories, respectively.

Figure 10. Experimental EFR_{PtN} and $DBEFR_{PtN}$ magnitudes (colored open symbols): (a) EFR_{PtN} to [2-22] and [4-22] kHz, (b) EFR_{PtN} to [2.8-16] and

[5.6-16] kHz and (c) DBEFR_{PtN} at [2-4] kHz and (d) DBEFR_{PtN} at [2.8-5.6] kHz. Simulated EFR_{PtN} (a,b) and DBEFR_{PtN} (c,d) magnitudes are shown in each panel using filled hexagons and degrees of CS as indicated on the X axis and CF-dependent patterns of OHC damage as given by the legend.

Figure 11. Modulated responses calculated at each CF using Eq. 8 and 9 to different experimental conditions for normal listeners and different sensorineural hearing losses at the AN processing level of the model, (a) broadband and (b) derived-band. In both panels, dotted lines show AM-responses to sloping 10 dB-HL OHC-loss at 8 kHz and lighter colors indicate AM responses to certain degree of CS.

761 Figures

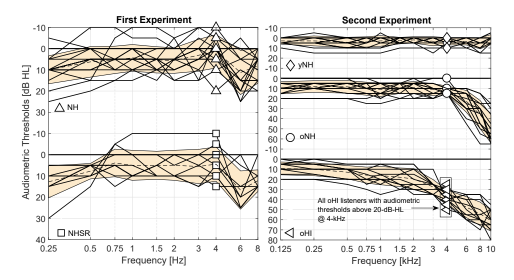


Figure 1

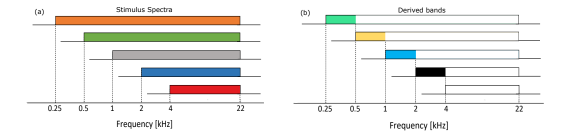


Figure 2

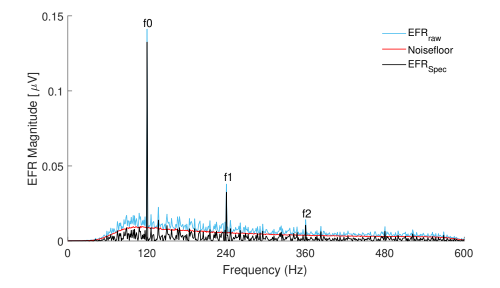


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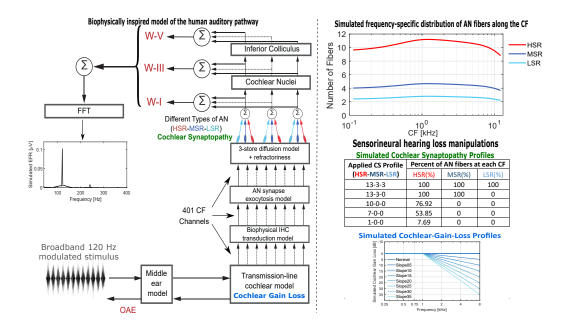


Figure 4

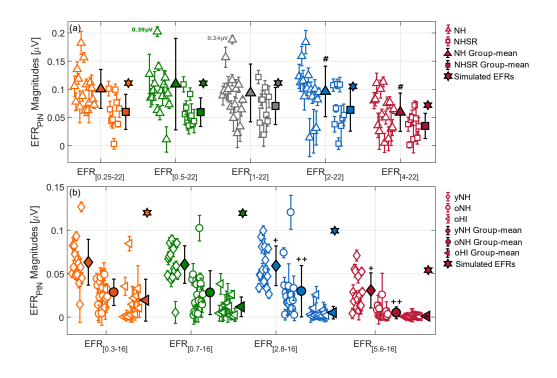


Figure 5

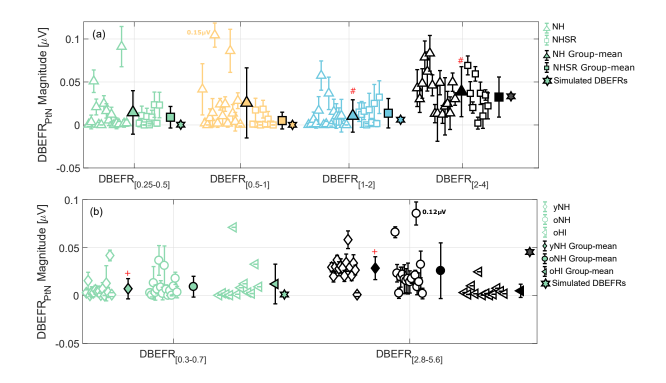


Figure 6

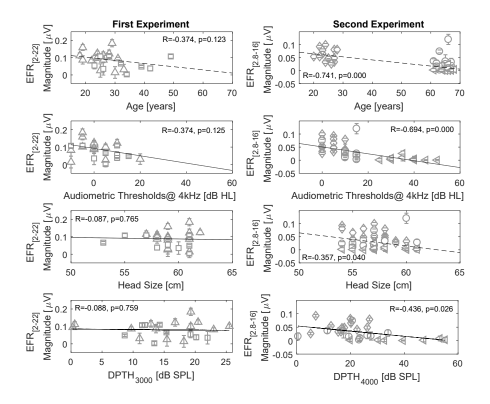


Figure 7

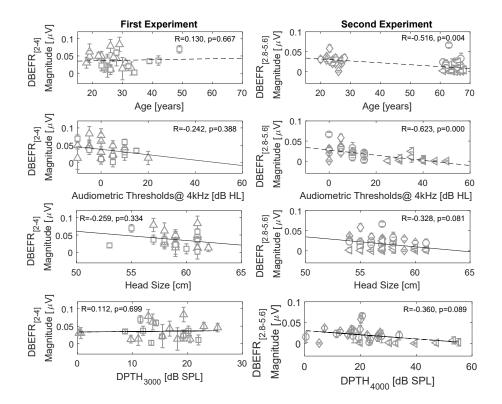


Figure 8

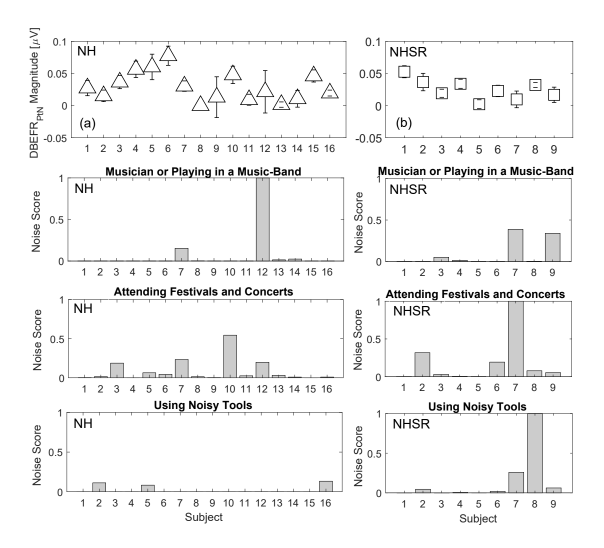


Figure 9

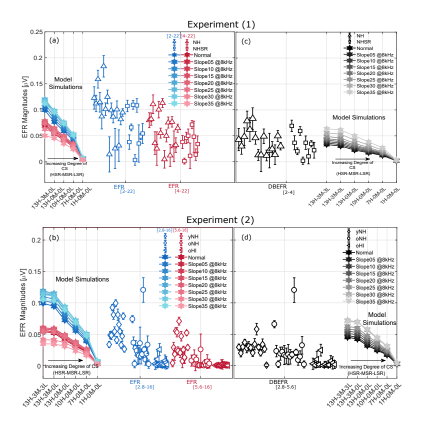


Figure 10

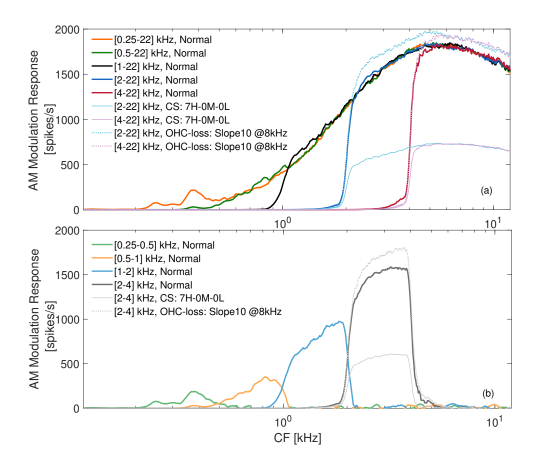


Figure 11