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Noise-sensitive but more precise subcortical representations co-exist with robust cortical encoding of natural vocalizations

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Abstract

Humans and animals maintain accurate sound discrimination in the presence of loud sources of background noise. It is commonly assumed that this ability relies on the robustness of auditory cortex responses. However, only a few attempts have been made to characterize neural discrimination of communication sounds masked by noise at each stage of the auditory system and to quantify the noise effects on the neuronal discrimination in terms of alterations in amplitude modulations. Here, we measured neural discrimination between communication sounds masked by a vocalization-shaped stationary noise from multiunit responses recorded in the cochlear nucleus, inferior colliculus, auditory thalamus, primary and secondary auditory cortex at several signal-to-noise ratios (SNR) in anesthetized male or female guinea pigs. Masking noise decreased sound discrimination of neuronal populations in each auditory structure, but collicular and thalamic populations showed better performance than cortical populations at each SNR. In contrast, in each auditory structure, discrimination by neuronal populations was slightly decreased when tone-vocoded vocalizations were tested. These results shed new light on the specific contributions of subcortical structures to robust sound encoding, and suggest that the distortion of slow amplitude modulation cues conveyed by communication sounds is one of the factors constraining the neuronal discrimination in subcortical and cortical levels.

Significance statement

Dissecting how auditory neurons discriminate communication sounds in noise is a major goal in auditory neuroscience. Robust sound coding in noise is often viewed as a specific property of cortical networks although this remains to be demonstrated. Here, we tested the discrimination performance of neuronal populations at five levels of the auditory system in response to conspecific vocalizations masked by noise. In each acoustic condition, subcortical neurons better discriminated target vocalizations than cortical ones and in each structure, the reduction in discrimination performance was related to the reduction in slow amplitude modulation cues.

Introduction

Understanding the neural mechanisms used by the auditory system to extract and represent relevant information for discriminating communication sounds in a variety of acoustic environments is a major goal in auditory neurosciences.

Several studies have prompted the view that the perceptual robustness mainly relies on the capacity of cortical neurons to extract invariant acoustic features (Narayan et al., 2007; Schneider and Woolley, 2013; Carruthers et al., 2015; Ni et al., 2017; Town et al., 2018), and it was proposed that this capacity is due to a larger adaptation of cortical cells to the noise statistics compared with subcortical cells (Rabinowitz et al., 2013). Indeed, in the cortical field L - the analogous of primary auditory cortex (A1) in bird - the percentage of correct neuronal discrimination between zebra-finch songs embedded in different types of acoustic maskers decreases proportionally to the target-to-masker ratio and parallels behavioral performance (Narayan et al., 2007). Also, consistent to behavioral data (for review see Verhey et al., 2003), the co-modulation of different frequency bands in background noise improved tone detection in noise of auditory cortical, thalamic and collicular neurons (Nelken et al., 1999; Las et al., 2005). Moreover, between-vowels discrimination performance of neuronal populations located in A1 resists to a large range of acoustic alterations (including changes in fundamental frequency, spatial location or level) and is similar to behavioral performance (Town et al., 2018).

The goal of the present study was to challenge this view by identifying the auditory structures responsible for this robust neural discrimination. Background noise has three disruptive effects on communication sounds (Noordhoek and Drullman, 1997; Dubbelboer and Houtgast, 2007): it attenuates the power of their amplitude modulation components (AM, also called “temporal-envelope”; Houtgast and Steeneken, 1985; Ewert and Dau, 2000; Biberger and Ewert, 2017), corrupts their frequency modulation components (FM, also called “temporal fine structure”; Shamma and Lorenzi, 2013; Varnet et al., 2017) and introduces stochastic fluctuations in AM power which generate temporal irregularities (from bin to bin) in the signal temporal envelopes (Ewert and Dau, 2000). Here, electrophysiological recordings were collected from the cochlear nucleus up to a secondary auditory cortical area in anesthetized guinea pigs and the discrimination performance of neuronal populations was assessed for four utterances of the same vocalization category (the whistle, e.g. the guinea pig alarm call) presented against a vocalization-shaped stationary noise at three signal-to-noise ratios (SNRs: +10, 0, -10 dB). An increased discrimination performance may result from the specialization

109 of cortical responses for detecting crucial vocalization features (Wang et al., 1995; Wang and
110 Kadia, 2001; Schneider and Woolley, 2013), whereas a decreased discrimination performance
111 may result from the loss of spectro-temporal details promoting the categorization of sounds
112 into auditory objects (Nelken and Bar-Yosef, 2008; Chechik and Nelken, 2012). Mutual
113 information was used to determine if the temporal patterns of neuronal responses to the four
114 vocalizations sufficiently differed to assign each response to a particular vocalization. The
115 results obtained in noise were compared to the effects of a deterministic signal-processing
116 scheme, namely a tone vocoder, which markedly altered the FM cues and progressively
117 attenuated the AM cues (within 38 to 10 frequency bands). The AM spectra were computed at
118 the output of simulated guinea pig auditory filters for each acoustic alteration. Our results
119 suggest that, the attenuation of slow AM cues is one of the factors explaining the decrease in
120 discrimination performance in cortical and subcortical structures. In addition, this study
121 revealed that, for each acoustic distortion tested, the highest level of discrimination was found
122 in subcortical structures, either at the collicular level (in masking-noise conditions) or at the
123 thalamic level (in vocoder conditions).

124

125

126

Materials and Methods

Subjects

These experiments were performed under the national license A-91-557 (project 2014-25, authorization 05202.02) and using the procedures N° 32-2011 and 34-2012 validated by the Ethic committee N°59 (CEEA Paris Centre et Sud). All surgical procedures were performed in accordance with the guidelines established by the European Communities Council Directive (2010/63/EU Council Directive Decree).

Extracellular recordings were obtained from 47 adult pigmented guinea pigs (aged 3 to 16 months, 36 males, 11 females) at five different levels of the auditory system: the cochlear nucleus (CN), the inferior colliculus (IC), the medial geniculate body (MGB), the primary (AI) and secondary (area VRB) auditory cortex. Animals, weighting from 515 to 1100 g (mean 856 g), came from our own colony housed in a humidity (50-55%) and temperature (22-24°C)-controlled facility on a 12 h/12 h light/dark cycle (light on at 7:30 A.M.) with free access to food and water.

Two to three days before each experiment, the animal's pure-tone audiogram was determined by testing auditory brainstem responses (ABR) under isoflurane anaesthesia (2.5 %) as described in Gourévitch et colleagues (2009). The ABR was obtained by differential recordings between two subdermal electrodes (SC25-NeuroService) placed at the vertex and behind the mastoid bone. A software (RTLab, Echodia, Clermont-Ferrand, France) allowed averaging 500 responses during the presentation of nine pure-tone frequencies (between 0.5 and 32 kHz) delivered by a speaker (Knowles Electronics) placed in the animal right ear. The auditory threshold of each ABR was the lowest intensity where a small ABR wave can still be detected (usually wave III). For each frequency, the threshold was determined by gradually decreasing the sound intensity (from 80 dB down to -10 dB SPL). All animals used in this study had normal pure-tone audiograms (Gourévitch et al., 2009; Gourévitch and Edeline, 2011; Aushana et al., 2018).

Surgical procedures

All animals were anesthetized by an initial injection of urethane (1.2 g/kg, i.p.) supplemented by additional doses of urethane (0.5 g/kg, i.p.) when reflex movements were observed after pinching the hind paw (usually 2-4 times during the recording session). A single dose of

157 atropine sulphate (0.06mg/kg, s.c.) was given to reduce bronchial secretions and a small dose
 158 of buprenorphine was administrated (0.05mg/kg, s.c.) as urethane has no antalgic properties.
 159 After placing the animal in a stereotaxic frame, a craniotomy was performed and a local
 160 anesthetic (Xylocain 2%) was liberally injected in the wound.
 161 For auditory cortex recordings (area A1 and VRB), a craniotomy was performed above the
 162 left temporal cortex. The opening was 8 mm wide starting at the intersection point between
 163 parietal and temporal bones and 8-10 mm height. The dura above the auditory cortex was
 164 removed under binocular control and the cerebrospinal fluid was drained through the cisterna
 165 to prevent the occurrence of oedema.
 166 For the recordings in MGB, a craniotomy was performed above the most posterior part of the
 167 MGB (8 mm posterior to Bregma) to reach the left auditory thalamus at a location where the
 168 MGB is mainly composed of its ventral, tonotopic, division (Redies et al., 1989; Edeline et
 169 al.; 1999, 2000; Anderson et al., 2007; Wallace et al., 2007).
 170 For IC recordings, a craniotomy was performed above the IC and portions of the cortex were
 171 aspirated to expose the surface of the left IC. For CN recordings, after opening the skull above
 172 the right cerebellum, portions of the cerebellum were aspirated to expose the surface of the
 173 right CN (Paraouty et al., 2018).
 174 After all surgeries, a pedestal in dental acrylic cement was built to allow an atraumatic
 175 fixation of the animal's head during the recording session. The stereotaxic frame supporting
 176 the animal was placed in a sound-attenuating chamber (IAC, model AC1). At the end of the
 177 recording session, a lethal dose of Exagon (pentobarbital >200 mg/kg, i.p.) was administered
 178 to the animal.

179 **Recording procedures**

180 Data from multi-unit recordings were collected in 5 auditory structures, the non-primary
 181 cortical area VRB, the primary cortical area A1, the medial geniculate body (MGB), the
 182 inferior colliculus (IC) and the cochlear nucleus (CN). In a given animal, neuronal recordings
 183 were only collected in one auditory structure. Cortical extracellular recordings were obtained
 184 from arrays of 16 tungsten electrodes ($\varnothing$: 33 μm , <1 M Ω) composed of two rows of 8
 185 electrodes separated by 1000 μm (350 μm between electrodes of the same row). A silver wire,
 186 used as ground, was inserted between the temporal bone and the dura matter on the
 187 contralateral side. The location of the primary auditory cortex was estimated based on the
 188 pattern of vasculature observed in previous studies (Edeline and Weinberger, 1993; Manunta

189 and Edeline, 1999; Edeline et al., 2001; Wallace et al., 2000). The non-primary cortical area
 190 VRB was located ventral to A1 and distinguished because of its long latencies to pure tones
 191 (Grimsley et al., 2012; Rutkowski et al., 2002). For each experiment, the position of the
 192 electrode array was set in such a way that the two rows of eight electrodes sample neurons
 193 responding from low to high frequency when progressing in the rostro-caudal direction [see
 194 examples of tonotopic gradients recorded with such arrays in figure 1 of Gaucher and
 195 colleagues (2012) and in figure 6A of Occelli and colleagues (2016)].

196 All the remaining extracellular recordings (in MGB, IC and CN) were obtained using 16
 197 channel multi-electrode arrays (NeuroNexus) composed of one shank (10 mm) of 16
 198 electrodes spaced by 110 μm and with conductive site areas of $177\mu\text{m}^2$. The electrodes were
 199 advanced vertically (for MGB and IC) or with a 40° angle (for CN) until evoked responses to
 200 pure tones could be detected on at least 10 electrodes.

201 All thalamic recordings were from the ventral part of MGB (see above surgical procedures)
 202 and all displayed latencies $< 9\text{ms}$. At the collicular level, we distinguished the lemniscal and
 203 non-lemniscal divisions of IC based on depth and on the latencies of pure tone responses. We
 204 excluded the most superficial recordings (until a depth of $1500\mu\text{m}$) and those exhibiting
 205 latency $\geq 20\text{ms}$ in an attempt to select recordings from the central nucleus of IC (CNIC). At
 206 the level of the cochlear nucleus, the recordings were collected from both the dorsal and
 207 ventral divisions.

208 The raw signal was amplified 10,000 times (TDT Medusa). It was then processed by an RX5
 209 multichannel data acquisition system (TDT). The signal collected from each electrode was
 210 filtered (610-10000 Hz) to extract multi-unit activity (MUA). The trigger level was set for
 211 each electrode to select the largest action potentials from the signal. On-line and off-line
 212 examination of the waveforms suggests that the MUA collected here was made of action
 213 potentials generated by a few neurons at the vicinity of the electrode. However, as we did not
 214 used tetrodes, the result of several clustering algorithms (Pouzat et al., 2002; Quiroga et al.,
 215 2004; Franke et al., 2015) based on spike waveform analyses were not reliable enough to
 216 isolate single units with good confidence. Although these are not direct proofs, the fact that
 217 the electrodes were of similar impedance (0.5-1M Ω) and that the spike amplitudes had
 218 similar values (100-300 μV) for the cortical and the subcortical recordings, were two
 219 indications suggesting that the cluster recordings obtained in each structure included a similar
 220 number of neurons.

221

222 Acoustic stimuli

223 Acoustic stimuli were generated using MATLAB (The Mathworks, Natick, MA), transferred
 224 to a RP2.1-based sound delivery system (TDT) and sent to a Fostex speaker (FE87E). The
 225 speaker was placed at 2 cm from the guinea pig's right ear, a distance at which the speaker
 226 produced a flat spectrum (± 3 dB) between 140 Hz and 36 kHz. The stimulation was not
 227 purely monaural, but the animal's head and body largely attenuated binaural cues. Calibration
 228 of the speaker was made using noise and pure tones recorded by a Bruel & Kjaer microphone
 229 4133 coupled to a preamplifier B&K 2169 and a digital recorder Marantz PMD671.

230 The Time-Frequency Response Profiles (TFRP) were determined using 129 pure-tones
 231 frequencies covering eight octaves (0.14-36 kHz) and presented at 75 dB SPL. The tones had
 232 a gamma envelop given by $\gamma(t) = (\frac{t}{4})^2 e^{-\frac{t}{4}}$, where t is time in ms. At a given level, each
 233 frequency was repeated eight times at a rate of 2.35 Hz in pseudorandom order. The duration
 234 of these tones over half-peak amplitude was 15 ms and the total duration of the tone was 50
 235 ms, so there was no overlap between tones.

236 A set of four conspecific vocalizations was used to assess the neuronal responses to
 237 communication sounds. These vocalizations were recorded from animals of our colony. Pairs
 238 of animals were placed in the acoustic chamber and their vocalizations were recorded by a
 239 Bruel & Kjaer microphone 4133 coupled to a preamplifier B&K 2169 and a digital recorder
 240 Marantz PMD671. A large set of whistle calls was loaded in the Audition software (Adobe
 241 Audition 3) and four representative examples of whistle were selected. As shown in figure
 242 1A, despite the fact the maximal energy of the four selected whistles was in the same
 243 frequency range (typically between 4 and 26 kHz), these calls displayed slight differences in
 244 their spectrogram and spectrum (Fig. 1A-B). In addition, their global temporal envelopes
 245 clearly differed (Fig. 1C). The four selected whistles were processed by three tone vocoders
 246 (Gnansia et al., 2009, 2010). In the following figures, the unprocessed whistles will be
 247 referred to as the original versions, and the vocoded versions as Voc38 (Voc20, Voc10
 248 respectively) for the 38-band (20-band, 10-band, respectively) vocoded whistles. In contrast
 249 to previous studies that used noise-excited vocoders (Nagarajan et al., 2002; Ranasinghe et
 250 al., 2012; Ter-Mikaelian et al., 2013), a tone vocoder was used here, because noise vocoders
 251 introduce random (i.e., non-informative) intrinsic temporal-envelope fluctuations distorting
 252 the crucial spectro-temporal modulation features of communication sounds (Shamma and
 253 Lorenzi, 2013; Kates, 2011; Stone et al., 2011).

Figure 1D displays the spectrograms of the 38-band vocoded (first column), the 20-band vocoded (second column) and the 10-band vocoded (third column) of the four whistles. The three vocoders differed only in terms of the number of frequency bands (i.e., analysis filters) used to decompose the whistles (38, 20 or 10 bands). The 38-band vocoding process is briefly described below, but the same principles apply to the 20-band or the 10-band vocoders. Each digitized signal was passed through a bank of 38 fourth-order Gammatone filters (Patterson, 1987) with center frequencies uniformly spaced along a guinea-pig adapted ERB (Equivalent Rectangular Bandwidth) scale ranging from 20 to 35505 Hz (Sayles and Winter, 2010). In each frequency band, the temporal envelope was extracted using full-wave rectification and lowpass filtering at 64 Hz with a zero-phase, sixth-order Butterworth filter. The resulting envelopes were used to amplitude modulate sine-wave carriers with frequencies at the center frequency of the Gammatone filters, and with random starting phase. Impulse responses were peak-aligned for the envelope (using a group delay of 16 ms) and the temporal fine structure across frequency channels (Hohmann, 2002). The modulated signals were finally weighted and summed over the 38 frequency bands. The weighting compensated for imperfect superposition of the bands' impulse responses at the desired group delay. The weights were optimized numerically to achieve a flat frequency response. Figure 1E shows the long-term power spectrum of the 38-band, 20-band and 10-band vocoded whistles, and figure 1F shows their global temporal envelopes (which were relatively well preserved by the vocoding process).

The four whistles were also presented in a frozen noise ranging from 10 to 24,000 Hz. To generate this noise, recordings were performed in the colony room where a large group of guinea pigs were housed (30-40; 2-4 animals/cage). Several 4-seconds of audio recordings were added up to generate a "chorus noise", which power spectrum was computed using the Fourier transform. This spectrum was then used to shape the spectrum of a white Gaussian noise. The resulting vocalization-shaped stationary noise therefore matched the "chorus-noise" audio spectrum, which explains why some frequency bands were over-represented in the vocalization-shaped stationary noise. Figure 1G displays the spectrograms of the four whistles in the vocalization-shaped stationary noise with a SNR of +10 dB SPL, 0 dB SPL, -10 dB SPL. Figure 1H shows the long-term power spectrum of the four whistles at the +10 dB, 0 dB and -10 dB SNR, and figure 1I shows their global temporal envelopes (which were severely altered at the 0 dB and -10 dB SNR).

Amplitude-modulation (AM) spectra were computed for the original, vocoded and noisy versions of each vocalization by decomposing each sound with the same bank of 50

288 gammatone filters than for the vocoding (spanning the range 0.1-50 kHz). The AM
 289 component (envelope) thus corresponds to the magnitude of the analytic signal, whereas the
 290 TFS corresponds to its unwrapped instantaneous phase.

291 For the AM spectrum, we analyzed the temporal envelope in each frequency band through a
 292 bank of AM filters using a method adapted from the human study by Varnet and colleagues
 293 (2017) for the guinea pigs' hearing range (1/3-octave wide first-order Butterworth bandpass
 294 filters overlapping at -3 dB, with center frequencies between 0.1 Hz and 410 Hz). The root-
 295 mean-square amplitude of the filtered output was multiplied by a factor of $\sqrt{2}$. For each AM
 296 filter, a modulation index was calculated by dividing the output by the mean amplitude of the
 297 AM component for the vocalization sample in the corresponding gammatone filter.

298 **Experimental protocol**

299 As inserting an array of 16 electrodes in a brain structure almost systematically induces a
 300 deformation of this structure, a 30-minutes recovering time lapse was allowed for the
 301 structure to return to its initial shape, then the array was slowly lowered. Tests based on
 302 measures of time-frequency response profiles (TFRPs) were used to assess the quality of our
 303 recordings and to adjust the electrodes' depth. For auditory cortex recordings (AI and VRB),
 304 the recording depth was 500-1000 μm , which corresponds to layer III and the upper part of
 305 layer IV according to Wallace and Palmer (2008). For thalamic recordings, the NeuroNexus
 306 probe was lowered about 7mm below pia before the first responses to pure tones were
 307 detected.

308 When a clear frequency tuning was obtained for at least 10 of the 16 electrodes, the stability
 309 of the tuning was assessed: we required that the recorded neurons displayed at least three
 310 successive similar TFRPs (each lasting 6 minutes) before starting the protocol. When the
 311 stability was satisfactory, the protocol was started by presenting the acoustic stimuli in the
 312 following order: We first presented the 4 original whistles in their natural versions, followed
 313 by the vocoded versions with 38, 20 and 10 bands at 75 dB SPL. The same set of original
 314 whistles was then presented in the vocalization-shaped stationary noise presented at 65, 75
 315 and 85 dB SPL. Thus, the level of the original vocalizations was kept constant (75 dB SPL),
 316 and the noise level was increased (65, 75 and 85 dB SPL). In all cases, each vocalization was
 317 repeated 20 times. Presentation of this entire stimulus set lasted 45 minutes. The protocol was
 318 re-started either after moving the electrode arrays on the cortical map or after lowering the
 319 electrode at least by 300 μm for subcortical structures.

320 **Data analysis**

321 *Quantification of responses to pure tones*

322 The TFRP were obtained by constructing post-stimulus time histograms (PSTH) for each
323 frequency with 1 ms time bins. The firing rate evoked by each frequency was quantified by
324 summing all the action potentials from the tone onset up to 100 ms after this onset. Thus,
325 TFRP are matrices of 100 bins in abscissa (time) multiplied by 129 bins in ordinate
326 (frequency). All TFRPs were smoothed with a uniform 5x5 bin window.

327 For each TFRP, the Best Frequency (BF) was defined as the frequency at which the highest
328 firing rate was recorded. Peaks of significant excitatory response were automatically
329 identified using the following procedure: An excitatory peak in the TFRP was defined as a
330 contour of firing rate above spontaneous activity plus six times the standard deviation of the
331 spontaneous activity. Recordings without significant excitatory peak of responses or with only
332 inhibitory responses were excluded from the data analyses. The bandwidth (BW) was defined
333 as the sum of all peak widths in octaves. The response duration was the time difference
334 between the first and last spikes of the significant peaks. The response
335 strength was the total number of spikes falling in the significant peaks.

336 *Quantification of responses evoked by vocalizations*

337 The responses to vocalizations were quantified using two parameters: (i) The firing rate of the
338 evoked response, which corresponds to the total number of action potentials occurring during
339 the presentation of the stimulus minus spontaneous activity; (ii) the trial-to-trial temporal
340 reliability coefficient (named CorrCoef as in our previous studies: Gaucher et al., 2013a;
341 Huetz et al., 2014; Gaucher and Edeline, 2015; Aushana et al., 2018) which quantifies the
342 trial-to-trial reliability of the responses over the 20 repetitions of the same stimulus. This
343 index was computed for each vocalization: it corresponds to the normalized covariance
344 between each pair of spike trains recorded at presentation of this vocalization and was
345 calculated as follows:

$$346 \text{CorrCoef} = \frac{1}{N(N-1)} \sum_{i=1}^{N-1} \sum_{j=i+1}^N \frac{\sigma_{x_i x_j}}{\sigma_{x_i} \sigma_{x_j}}$$

347 where N is the number of trials and $\sigma_{x_i x_j}$ is the normalized covariance at zero lag between
348 spike trains x_i and x_j where i and j are the trial numbers. Spike trains x_i and x_j were previously
349 convolved with a 10-msec width Gaussian window. Based upon computer simulations, we

350 have previously shown that this CorrCoef index is not a function of the neurons' firing rate
351 (Gaucher et al., 2013a).

352 *Quantification of mutual information from the responses to vocalizations*

353 The method developed by Schnupp and colleagues (2006) was used to quantify the amount of
354 information (Shannon, 1948) contained in the responses to vocalizations obtained with natural
355 vocoded and noise stimuli. This method allows quantifying how well the vocalization's
356 identity can be inferred from neuronal responses. Here, "neuronal responses" refer either to (i)
357 the spike trains obtained from a small group of neurons below one electrode (for the
358 computation of the individual Mutual Information, $MI_{\text{Individual}}$), or to (ii) a concatenation of
359 spike trains simultaneously recorded under several electrodes (for the computation of the
360 population, $MI_{\text{Population}}$). In both cases, the following computation steps were the same.
361 Neuronal responses were represented using different time scales ranging from the duration of
362 the whole response (firing rate) to a 1-ms precision (precise temporal patterns), which allows
363 analyzing how much the spike timing contributes to the information. As this method is
364 exhaustively described in Schnupp and colleagues (2006) and in Gaucher and colleagues
365 (2013a), we only present below the main principles.

366 The method relies on a pattern-recognition algorithm that is designed to "guess which
367 stimulus evoked a particular response pattern" (Schnupp et al., 2006) by going through the
368 following steps: From all the responses of a cortical site to the different stimuli, a single
369 response (test pattern) is extracted and represented as a PSTH with a given bin size (different
370 sizes were considered as indicated in the Results section). Then, a mean response pattern is
371 computed from the remaining responses (training set) for each stimulus class. The test pattern
372 is then assigned to the stimulus class of the closest mean response pattern. This operation is
373 repeated for all the responses, generating a confusion matrix where each response is assigned
374 to a given stimulus class. From this confusion matrix, the Mutual Information (MI) is given
375 by Shannon's formula:

$$376 \quad MI = \sum_{x,y} p(x,y) \times \log_2 \left(\frac{p(x,y)}{p(x) \times p(y)} \right)$$

377 where x and y are the rows and columns of the confusion matrix, or in other words, the values
378 taken by the random variables "presented stimulus class" and "assigned stimulus class".

379 In our case, we used responses to the 4 whistles and selected the first 280 ms of these
380 responses to work on spike trains of exactly the same duration (the shortest whistle being 280

ms long). In a scenario where the responses do not carry information, the assignments of each response to a mean response pattern is equivalent to chance level (here 0.25 because we used 4 different stimuli and each stimulus was presented the same number of times) and the MI would be close to zero. In the opposite case, when responses are very different between stimulus classes and very similar within a stimulus class, the confusion matrix would be diagonal and the mutual information would tend to $\log_2(4) = 2$ bits.

This algorithm was applied with different bin sizes ranging from 1 to 280 ms (see figure 2B for the evolution of MI with temporal precisions ranging from 1 to 40 ms).

The MI estimates are subject to non-negligible positive sampling biases. Therefore, as in Schnupp and colleagues (2006), we estimated the expected size of this bias by calculating MI values for “shuffled” data, in which the response patterns were randomly reassigned to stimulus classes. The shuffling was repeated 100 times, resulting in 100 MI estimates of the bias (MI_{bias}). These MI_{bias} estimates are then used as estimators for the computation of the statistical significance of the MI estimate for the real (unshuffled) datasets: the real estimate is considered as significant if its value is statistically different from the distribution of MI_{bias} shuffled estimates. Significant MI estimates were computed for MI calculated from neuronal responses under one electrode. The range of MI_{bias} values was very similar between auditory structures: depending on the conditions (original, vocoded, noisy vocalizations), it was from 0.102 to 0.107 in the CN, from 0.107 to 0.110 in the IC, from 0.105 to 0.114 in the MGB, 0.107 to 0.111 in A1 and from 0.106 to 0.116 in VRB. There was no significant difference between the mean MI_{bias} values in the different structures (unpaired t-test, all $p > 0.25$).

The information carried by a group of recordings was estimated by the population MI ($MI_{Population}$), using the same method described above: responses of several simultaneous multiunit recordings were concatenated and considered as a single pattern. To assess the influence of the group size of simultaneous multiunit recordings on the information carried by that group ($MI_{Population}$), the number of sites used for computing $MI_{Population}$ varied from 2 to the maximal possible size (which is equal to 16 minus the non-responsive sites). As the number of possible combinations could be extremely large (C_n^k , where k is the group size and n the number of responsive sites in a recording session), a threshold was fixed to save computation time: when the number of possible combinations exceeded one hundred, 100 combinations were randomly chosen, and the mean of all combinations was taken as the $MI_{Population}$ for this group size.

For the $MI_{Population}$, the values of bias were also computed: on average and for all sets of 9 simultaneous recordings, it was 0.104 in the CN, 0.111 in the IC, 0.114 in the MGB, 0.107 in

415 AI and 0.106 in VRB. There was no significant difference between the mean $MI_{\text{Population}}$ bias
416 values in the different structures (unpaired t-test, all $p > 0.20$).

417

418

419 *Statistics*

420 To assess the significance of the multiple comparisons (vocoding process: four levels;
421 masking noise conditions: three levels; auditory structure: five levels), we used an analysis of
422 variance (ANOVA) for multiple factors to analyze the whole data set. Post-hoc pair-wise tests
423 were performed between the original condition and the vocoding or noisy conditions. They
424 were corrected for multiple comparisons using Bonferroni corrections and were considered as
425 significant if their p value was below 0.05. All data are presented as mean values $\pm$ standard
426 error (s.e.m.).

427

428

Results

From a database of 2334 recordings collected in the five auditory structures, two criteria were used to include neuronal recordings in our analyses. A recording had to show significant responses to pure tones (see Methods) and an evoked firing rate significantly above spontaneous firing rate (200 ms before each original vocalization) for at least one of the four original vocalizations. Applying these two criteria led to the inclusion of 499 recordings in CN, 386 recordings in CNIC, 262 recordings in MGv, 354 recordings in A1 and 95 recordings in VRB. Table 1 summarizes the range of best frequencies, mean bandwidth, response duration and response strength obtained when testing pure tone responses in each auditory structure. In the following sections, the neuronal responses to the original vocalizations presented in quiet are compared across brain structures and the discrimination performance are described at the individual and population levels. The neuronal discrimination tested with tone-vocoded vocalizations and vocalizations presented against different levels of masking noise are described and compared next.

Determination of optimal parameters for temporal analyses of spike trains in the five auditory structures

Before quantifying the neuronal discrimination performance in the five investigated structures, we first looked for the optimal parameters for analyzing the temporal patterns of spike trains in the five structures.

First, the CorrCoef index which quantifies the trial-to-trial temporal reliability, was computed with a Gaussian window ranging from 1 to 50 ms. As a general rule, the largest the Gaussian window, the largest the CorrCoef mean value whatever the structure was. We questioned if selecting a particular value for the Gaussian window influenced the between-structure differences in CorrCoef mean values. Based upon the responses to the original vocalizations, figure 2A shows that the relative ranking between auditory structures remained unchanged whatever the width of the Gaussian window was. Therefore, we kept the value of 10 ms for the Gaussian window (dashed line in Fig. 2A) as it was used in previous cortical studies (Huetz et al., 2009; Gaucher et al., 2013a; Gaucher and Edeline, 2015; Aushana et al., 2018).

Second, at the cortical level, it was previously showed that the maximal value of mutual information (MI) based on temporal patterns was obtained, on average, with a bin size of 8ms (Schnupp et al., 2006; Gaucher et al., 2013a). However, it has never been demonstrated that the same bin size was optimal at all levels of the auditory system. Figure 2B shows the

evolution of MI as a function of temporal precision ranging from 1 to 40 ms based on the responses to the original vocalizations. In our experimental conditions, and with our set of acoustic stimuli, the 8-ms temporal precision was found to be optimal for all auditory structures, in the original (dashed line in Fig. 2B), vocoded and noisy conditions (data not shown). Therefore, the MI value obtained for a temporal precision of 8 ms was subsequently used in our analyses.

Subcortical structures better discriminate the original vocalizations

Figure 3A displays neuronal responses of two simultaneous multiunit recordings obtained at five levels of the auditory pathway (CN, CNIC, MGv, AI and VRB). The neuronal responses were strong and sustained in the CN and CNIC, more transient in MGv, phasic in AI and more diffuse in VRB. For most of the recordings, temporal patterns of response were clearly reproducible from trial-to-trial, but they differed from one vocalization to another both at the cortical and subcortical level. The PSTHs displayed in figure 3B show that at each level of the auditory system, the four whistles triggered distinct temporal patterns of responses.

Quantifications of evoked responses to original vocalizations over all the recordings are presented on figures 3C-F for each auditory structure. These analyses clearly pointed out large differences between the mean values of evoked firing rate, CorrCoef and MI quantified at the cortical vs. at the subcortical level. First, the evoked firing rate was significantly higher in the subcortical structures than in the cortex (unpaired t-test, lowest p value $p < 0.001$). It was also higher in CN compared to the other subcortical structures (Fig. 3C). Second, the CorrCoef values were significantly higher in CNIC and MGv compared to AI and VRB (Fig. 3D), indicating that the trial-to-trial reliability of evoked responses was stronger in these structures than in CN, AI and VRB. Third, the $MI_{\text{Individual}}$ values obtained at the subcortical level were significantly higher than at the cortical level (unpaired t-test, highest $p < 0.001$ between the cortex and the other structures; Fig. 3E). At the subcortical level, the $MI_{\text{Individual}}$ values were significantly higher in MGv than in CNIC and in CN (unpaired t-test, $p < 0.01$) with the CN exhibiting the lowest MI values at the subcortical level. The $MI_{\text{Individual}}$ values were also significantly lower in VRB than in AI ($p = 0.037$). Recordings in MGv displayed the highest $MI_{\text{Individual}}$ mean values, suggesting that, on average, thalamic neurons discriminate better the four original whistles than the other auditory structures. As shown in figure 3G, in each auditory structure, high $MI_{\text{Individual}}$ values were strongly correlated with high values of trial-to-trial temporal reliability (indexed by the CorrCoef value; $0.77 < r < 0.88$; $p < 0.001$). Finally,

MI was also computed based on the temporal patterns obtained from two to sixteen simultaneous multiunit recordings to determine whether the discrimination performance of neural networks confirm the results obtained at the individual (i.e., single recording) level. $MI_{Population}$ quantifies how well the four whistles can be discriminated based on temporal patterns expressed by neuronal populations distributed on the tonotopic map. The $MI_{Population}$ computed from 9 simultaneous multiunit recordings shows that neural populations in subcortical structures discriminate the four original whistles better than the cortical populations (unpaired t-test, highest p value $p < 0.002$ between CN and VRB) without any statistical difference between the three subcortical structures (Fig. 3F).

We next investigated the diversity of the $MI_{Individual}$ and $MI_{Population}$ values obtained in the different structures. The distributions of $MI_{Individual}$ values were plotted as a function of temporal precision for each structure (Fig. 4 A1-A5). It showed that whatever the temporal precision, there were more curves with high $MI_{Individual}$ values in the subcortical structures than in the cortical areas (see red curves on Fig. 4 A1-A5). The examination of the evolution of the $MI_{Population}$ as a function of the number of simultaneous multiunit recordings in the different structures revealed that the growth functions rapidly reached high values in all subcortical structures, whereas there were only a few of such curves in AI and VRB whatever the number of recordings considered (Fig. 4 B1-B5).

With a temporal resolution of 8 ms, we presented the cumulative percentages of neurons for the $MI_{Individual}$ (Fig. 5A) and the $MI_{Population}$ values (Fig. 5B) in each structure. Above a value of 1.5 bits (indicating that at least 3 stimuli can be discriminated), there were 39% of MGv neurons, 18% and 14% of the neurons in CNIC and CN respectively; but only 3.5% and 2% of the neurons in A1 and VRB respectively. This proportion was significantly higher in MGv than in CN and CNIC ($p = 0.017$ and $p = 0.04$) and was also significantly higher in subcortical structures compared with the cortical ones (all p values < 0.01). The same conclusions were reached for the $MI_{Population}$ values: More than 90% of the MGv neuronal populations were above 1.5 bits, 83 % and 75% of the populations in CNIC and CN respectively, whereas these populations represented less than 40% at the cortical level (36 % and 34% in A1 and VRB respectively).

Thus, both at the level of individual recordings, and at the population of simultaneous multiunit recordings, subcortical neurons are more accurate in discriminating the four original whistles than cortical ones.

Modest effects of tone vocoding

Figure 6A displays rasters of recordings obtained in the five structures in response to the original and tone-vocoded vocalizations using 38 (Voc38), 20 (Voc20) and 10 (Voc10) frequency bands. As illustrated by the rasters and the PSTHs presented in figure 6B, in all structures, neurons still vigorously responded to the vocoded stimuli even for 10-band vocoded stimuli.

Figure 6C-F summarizes the vocoding effects on the four parameters quantifying neuronal responses. Compared to the responses to the original vocalizations, the evoked firing rate obtained in all structures in response to vocoded stimuli only showed modest variations (Fig. 6C): apart from an increase in firing rate in the CN with the 38-band vocoded stimuli, a significant decrease in evoked firing rate in response to the 10-band vocoded vocalizations was only found at the subcortical level (for all subcortical structures, one-way ANOVA: $F_{CN(3,1995)}=22.6$; $F_{CNIC(3,1543)}=8.85$; $F_{MGv(3,1047)}=6.55$, $p<0.001$; with post-hoc paired t tests, $p<0.05$), whereas there was no decrease in either AI or VRB. Vocoding also decreased the CorrCoef mean values in every structure except in VRB (Fig. 6D). This decrease was significant with the 10-band vocoded vocalizations in CN, MGv and in AI (one-way ANOVA: $F_{CN(3,1930)}=26.48$; $F_{MGv(3,889)}=7.7$; $F_{AI(3,1125)}=3.42$, highest p value, $p<0.02$; with post-hoc paired t tests, $p<0.05$). The decrease in CorrCoef value was already significant with 20-band vocoded vocalizations in the CNIC (one-way ANOVA: $F_{(3,1391)}=26.19$, $p<0.001$; with post-hoc paired t tests, $p<0.05$).

Similarly, vocoding decreased the $MI_{Individual}$ values in each structure except in VRB (Fig. 6E). Here too, the decrease was significant with the 10-band vocoded vocalizations in CN, MGv and AI (one-way ANOVA: $F_{CN(3,1445)}=12.23$, $F_{MGv(3,810)}=3.75$, $F_{AI(3,720)}=3.59$, highest p value, $p<0.02$; with post-hoc paired t tests, $p<0.05$) and it was already significant with 20-band vocoded vocalizations in the CNIC (one-way ANOVA: $F_{CNIC(3,1231)}=13.17$, $p<0.001$; with post-hoc paired t tests, $p<0.05$). At the population level ($MI_{Population}$), compared to the values obtained in response to the original vocalizations, the $MI_{Population}$ values computed with the 10-band vocoded vocalizations were significantly lower in the subcortical structures (one-way ANOVA: $F_{CN(3,127)}=6.46$, $F_{CNIC(3,115)}=6.28$, $F_{MGv(3,67)}=4.62$, highest p value, $p<0.005$; with post-hoc paired t tests, $p<0.05$) but not at the cortical level (Fig. 6F). The evolution of $MI_{Population}$ as a function of the number of simultaneous multiunit recordings (Fig. 7 A-E) showed that in each subcortical structure, the curves rapidly reached high $MI_{Population}$ values

(close to the maximal value of 2 bits) in each vocoding conditions, whereas in AI and VRB the curves slowly reached the maximum $MI_{\text{Population}}$ values.

In conclusion, for the five auditory structures, the neuronal responses to 10-band vocoded vocalizations were slightly weaker, temporally less accurate and less discriminative than the responses to the original vocalizations. Nonetheless, on average, subcortical neurons still maintained the highest discrimination performance between tone-vocoded vocalizations, both at the level of individual recordings and at the population level.

Pronounced effects of masking noise on neuronal discrimination

The rasters presented in figure 8A illustrate the effects induced by presenting the original vocalizations against a vocalization-shaped stationary noise at three SNRs (+10, 0 and -10 dB). As illustrated by the rasters and the PSTHs presented in figure 8B, masking noise attenuated neuronal responses at each level of the auditory system. However, the auditory structures were differentially affected by noise. The responses in the CNIC did not change up to a 0 dB SNR, decreasing only at a -10 dB SNR. This was not the case in the other auditory structures where the responses decreased either at a +10 dB SNR (MGv and CN) or at a 0 dB SNR (AI and VRB).

Figure 8C-F summarizes the effects of masking noise on the different parameters quantifying neuronal responses. Masking noise significantly reduced the evoked firing rate in each auditory structure already at the +10 dB SNR (Fig. 8C, one-way ANOVA: $F_{\text{CN}(3,1995)}=309.33$, $F_{\text{CNIC}(3,1543)}=220.64$, $F_{\text{MGv}(3,1047)}=155.07$, $F_{\text{AI}(3,1415)}=96.27$, $p<0.001$; with post-hoc paired t tests, $p<0.05$), except in VRB.

At the subcortical level, masking noise strongly reduced the CorrCoef values in CN and MGv at the highest SNR (+10 dB) tested here (Fig. 8D; one-way ANOVA: $F_{\text{CN}(3,1884)}=382.22$, $F_{\text{MGv}(3,791)}=155.82$, $p<0.001$; with post-hoc paired t tests, $p<0.05$) whereas in the CNIC, this reduction was significant only at the 0 dB SNR (one-way ANOVA: $F_{\text{CNIC}(3,1357)}=154.12$, $p<0.001$; with post-hoc paired t tests, $p<0.05$). At the cortical level, the CorrCoef values were significantly reduced in AI at the +10 dB SNR and in VRB at the 0 dB SNR (one-way ANOVA: $F_{\text{AI}(3,1093)}=60.83$, $F_{\text{VRB}(3,335)}=29.56$, $p<0.001$; with post-hoc paired t tests, $p<0.05$).

At the subcortical level, noise reduced the $MI_{\text{Individual}}$ values but again, there was a marked difference between the CNIC and the other subcortical structures: the $MI_{\text{Individual}}$ mean value in CN and MGv was significantly reduced at the +10 dB SNR (Fig. 8E; one-way ANOVA: $F_{\text{CN}(3,819)}=56.75$, $F_{\text{MGv}(3,621)}=63.61$, $p<0.001$; with post-hoc paired t tests, $p<0.05$), whereas the

MI_{Individual} value in the CNIC was only significantly reduced at the 0 dB SNR (one-way ANOVA: $F_{(3,1078)}=32.08$, $p<0.001$; with post-hoc paired t tests, $p<0.05$). At the cortical level, noise significantly reduced the average MI_{Individual} in AI only at the -10 dB SNR (one-way ANOVA: $F_{(3,649)}=9.49$, $p<0.001$; with post-hoc paired t tests, $p<0.05$) whereas the average MI_{Individual} in VRB remained unchanged (Fig 8E).

The effects of masking noise on the network discrimination performance were quantified with the MI_{Population} (Fig. 8F). At the cortical level, there was a significant reduction of MI_{Population} values only at the -10 dB SNR (one-way ANOVA: $F_{A1(3,111)}=16.63$, $F_{VRB(3,23)}=11.41$, $p<0.001$; with post-hoc paired t tests, $p<0.05$) whereas there was a significant decrease in CN already at the +10 dB SNR (one-way ANOVA: $F_{CN(3,127)}=51.49$, $p<0.001$; with post-hoc paired t tests, $p<0.05$). In MGv and CNIC, neuronal populations still displayed the highest discrimination performance although the decrease in MI_{Population} value was significant at the 0 dB SNR (one-way ANOVA: $F_{MGv(3,67)}=41.59$, $F_{CNIC(3,115)}=22.59$, $p<0.001$; with post-hoc paired t tests, $p<0.05$).

Note that, in VRB, the CorrCoef and MI_{Population} were much more decreased in the noise conditions than in the vocoding conditions, suggesting that the lack of significant decreases in vocoding conditions was not a “floor effect” due to the low initial values.

The evolution of the MI_{Population} as a function of the number of simultaneous multiunit recordings in the different structures (Fig. 9A-E) revealed that regardless of the number of neurons considered, noise effects were similar up to the 0 dB SNR: the population curves in CNIC and MGv grew up relatively rapidly and reached higher values than the curves obtained in CN and in the two cortical areas. At the -10 dB SNR, the MI_{Population} from the CNIC remained higher (regardless of the number of neurons considered) than in the other structures; whereas there was no increase of the MI_{Population} with the number of neurons in VRB.

One puzzling result came from the fact that on average, the values of MI_{Individual} and MI_{Population} decreased more for CN recordings than for the two subsequent subcortical relays. However, at least 20% of the CN recordings at the +10 dB SNR maintained MI_{Individual} values above 1 bit (Fig. 10A, red curves) and MI_{Population} values above 1.5 bits (Fig. 10C, red curves), suggesting that a sub-population of CN neurons were still able to send information about the vocalization identity at higher brainstem centers. This also suggests that the discrimination performed by a group of a fixed number of neurons deteriorates with noise faster in the CN and consequently, more CN neurons are necessary to obtain an equivalent amount of information observed in CNIC.

632 The distributions of the TFRP parameters (best frequency, bandwidth, response duration,
 633 response strength) from this specific sub-population of CN neurons did not differ from the
 634 neurons exhibiting $MI_{\text{Individual}}$ values below 1 bit at the +10 dB SNR in terms of best
 635 frequency and bandwidth but significantly differ in terms of response duration and response
 636 strength (chi-square tests; $p < 0.05$, Fig. 10B). More precisely, the CN recordings exhibiting
 637 higher $MI_{\text{Individual}}$ values at +10 dB SNR had longer duration responses and stronger evoked
 638 firing rate to pure tones.

639 A more general question is to evaluate whether the TFRP characteristics in the different
 640 auditory structures (see examples in Fig. 11A) influenced the noise effects quantified by the
 641 $MI_{\text{Individual}}$ values (Fig. 11B-C). As indicated in figure 11, there was no relationship between
 642 the best frequency values and the changes in $MI_{\text{Individual}}$ values (Fig. 11B) and no relationship
 643 between the frequency bandwidth and the changes in $MI_{\text{Individual}}$ values (Fig. 11C). Thus, in all
 644 structures, the noise-induced alterations in $MI_{\text{Individual}}$ values seem to be independent from the
 645 characteristics of pure tone responses.

646 To summarize, masking noise differently impacted the neurons' discrimination performance
 647 at the subcortical and cortical levels. Although cortical neurons were more resistant to
 648 changes in noise level, the thalamic and collicular neurons maintained higher MI values, with
 649 the CNIC neurons displaying the highest discrimination performance both at the individual
 650 and population level in the most challenging condition (i.e., at the -10 dB SNR).

651

652 **Alteration of slow amplitude modulations as one of the factors explaining the changes in** 653 **neuronal discrimination**

654

655 Masking noise produced spectro-temporal degradations: it reduced the AM cues in the
 656 different audio-frequency bands, introduced irrelevant envelope fluctuations and altered the
 657 temporal fine structure (TFS) of the sound. Tone vocoding removed almost all the TFS but
 658 also progressively filtered out the fast AM. As a vast literature demonstrated that slow AM
 659 cues are crucial for speech understanding in normal and degraded conditions (Houtgast and
 660 Steeneken, 1985; Drullman et al., 1994, 1995; Shannon et al., 1995; Dubbelboer and
 661 Houtgast, 2007; Jorgensen and Dau, 2011), we quantified the alterations of AM cues (due to
 662 masking noise and to vocoding) and looked for potential relationships with the alterations in
 663 neural discrimination ($MI_{\text{Population}}$) in the five structures.

664 The AM spectra obtained in vocoding and noise conditions showed that the AM cues were
 665 attenuated compared to the original condition (Fig. 12A). The +10 dB SNR condition

666 produced a flattening of the AM modulation spectrum, which was further accentuated in the 0
 667 dB and -10 dB SNR conditions. In these two most degraded conditions, noise also introduced
 668 non-relevant fluctuations at high rates. In contrast, vocoding preserved the general shape of
 669 the AM spectra while progressively filtering out the AM fluctuations.

670 We investigated the relationships between these degradations of AM cues and neural
 671 discrimination ($MI_{\text{Population}}$) in the five structures for each experimental condition (Fig. 12B).
 672 More precisely, for all conditions, figure 12B shows the changes in $MI_{\text{Population}}$ for each
 673 auditory structure as a function of the attenuation of AM cues (computed as the mean
 674 modulation index between 1 and 20 Hz). Figure 12B reveals that in all structures other than
 675 the CN, $MI_{\text{Population}}$ is barely affected as long as the reduction of the AM index ($\Delta\text{modulation}$
 676 index) remains lower than 25%; beyond this limit, the $MI_{\text{Population}}$ is reduced (i.e., at the 0 dB
 677 and -10 dB SNR). The straightforward conclusion is that the reduction of slow AM cues is
 678 one of the factors controlling the decrease in $MI_{\text{Population}}$ at the cortical and subcortical levels.
 679 In the cochlear nucleus, the decrease on the $MI_{\text{Population}}$ is much larger than in the other
 680 structures, suggesting that the alteration of AM cues has more impact on the $MI_{\text{Population}}$ at the
 681 most peripheral level. Alternatively, one should keep in mind that the neuronal discrimination
 682 in noise can be based upon other acoustic cues such as the FM cues (in particular pitch cues),
 683 spectral regularity and harmonicity cues, and the simultaneous rising slope of energy across
 684 channels. Thus, in the cochlear nucleus, but also in the other structures, the strong decrease in
 685 $MI_{\text{Population}}$ can potentially result from alterations of one, or several, of these parameters.

686 Dissecting the contributions of each of these parameters to neuronal discrimination and its
 687 decrease in degraded conditions will require manipulations of controlled stimuli in
 688 independent conditions. Confirming that the slow AM cues are the main factor for
 689 discrimination in degraded conditions could theoretically be achieved by keeping the exact
 690 same AM cues and modifying only one of the acoustic parameter listed above. Using a
 691 computational model of the peripheral auditory system will help to estimate the respective
 692 representations of the envelope and temporal fine structure after acoustic degradations (Moon
 693 et al., 2014; Wirtzfeld et al., 2017). For example, the search for "equivalent" experimental
 694 conditions in terms of amounts of neural degradation of AM and FM cues could be performed
 695 by using the FAME vocoder (Zeng et al., 2005) to alter systematically AM and FM
 696 parameters (i.e., cutoff frequency, modulation strength, modulation phase) of the
 697 vocalizations used as stimuli. The results of this type of experiments should also be
 698 generalized with other categories of guinea pig calls, other types of communication sounds
 699 from other species and should included in other types of masking noises.

Discussion

Here, we demonstrated that for each acoustic distortion, subcortical neurons displayed the highest level of discrimination performance of natural vocalizations, either at the collicular level (in masking noise conditions) or at the thalamic level (in vocoder conditions). More precisely, background noise markedly reduces neural discrimination performance in all auditory structures with larger effects in the cochlear nucleus, whereas the vocoder induced little effect in each auditory structure. Interestingly, the discrimination performance of cortical neurons was less impacted making these neurons more robust to all acoustic alterations. Moreover, comparison of neural data collected in response to noisy versus vocoded vocalizations suggests that the transmission of slow (< 20 Hz) amplitude modulation information is one of the factors contributing to the neural discrimination decrease in noise at the cortical and subcortical levels.

Subcortical structures represent natural vocalizations more precisely than primary and non-primary cortical areas

In contrast with previous cortical studies, which have quantified the discrimination between calls that belong to different categories making the discrimination easy for cortical neurons (Narayan et al., 2006, 2007; Ter-Mikaelian et al., 2013; Ni et al., 2017), we used four vocalizations that belong to the same category making the discrimination more difficult for cortical neurons. We showed that on average subcortical populations discriminated the original vocalizations better than cortical populations. Moreover, smaller populations of subcortical neurons compared to cortical ones were sufficient to discriminate between the stimuli used in this study. These results corroborate the finding by Chechik and colleagues (2006) that the MGB and AI responses contain 2-to-4 fold less information than the responses of IC neurons. Here, the discrimination performance in MGv was closer than the ones displayed by the other subcortical structures. A potential explanation is that Chechik and colleagues (2006) recorded from all MGB divisions, including the medial and dorsal divisions, whereas our thalamic recordings were limited to MGv and exhibited tonic responses to vocalizations similar to those observed in the CNIC and the CN (Fig. 3A and 5A). The stimulus sets also differ, as we used four utterances of the same category (the Whistle), whereas Chechik and colleagues (2006) used three birds' chirps and variants of these stimuli leading potentially to an easier classification between groups of stimuli

734 compared to our protocol. An interesting result was that the optimal bin size for computing
 735 MI was similar for all structures (8 ms bin, Fig. 2B). Importantly, with a smaller or a larger
 736 bin, the mutual information would have been underestimated, but this would not have
 737 changed the differences reported here: whatever the bin size, subcortical neurons will still
 738 discriminate better the original vocalizations than the cortical areas (Fig. 2B). Potentially, the
 739 optimal bin size depends more upon the stimuli durations than upon the auditory structure.
 740 When computing mutual information from IC, MGB and A1 neuronal responses, Chechik and
 741 colleagues (2006) usually found an optimal bin size of 4 ms, different from ours, probably
 742 because their stimulus durations are shorter than our stimuli (67-111 ms vs. to 280-363 ms
 743 here). Recently, we also found shorter optimal bin size when computing MI with shorter (12-
 744 65 ms) communication sounds (Royer et al., 2019).

745 Our original stimuli differed in terms of temporal envelope and, as a consequence, the most
 746 efficient way to discriminate them is probably to follow the time course of AM cues. It is well
 747 known that when progressing from the lower to the upper stages of the auditory system, the
 748 neurons' ability to follow AM cues considerably changes (Joris et al, 2004; Escabi and Read,
 749 2005). Brainstem neurons phase-lock on AM modulations up to hundreds of Hertz (Frisina et
 750 al., 1990; Rhode and Greenberg, 1994), whereas thalamic neurons do so for a few tens of
 751 Hertz (Creutzfeldt et al., 1990; Preuss and Müller-Preuss, 1990) and cortical neurons for even
 752 lower rates (Gaese and Ostwald, 1995; Schreiner and Urbas, 1998). As a consequence,
 753 subcortical neurons, (but not cortical ones) can follow the largest and fastest AM cues (7-15
 754 Hz) contained in the original vocalizations (see the peak of the black curve in AM spectra,
 755 Fig. 12A). This likely explains why subcortical neurons better discriminate the original
 756 stimuli both at the individual and population levels. Cortical neurons only follow the weakest
 757 and slowest AM cues (1-5 Hz) of the original vocalizations, which potentially explains why
 758 cortical neurons weakly discriminate the original stimuli and tend to encode them as a single
 759 category (Mesgarani et al., 2014b).

760

761 **Alterations of slow amplitude modulation cues is one of the factors explaining the**
 762 **changes in cortical and subcortical discrimination**

763

764 Previous studies using vocoded vocalizations reported that cortical responses were not
 765 drastically reduced even with two frequency bands (Nagarajan et al., 2002; Ranasinghe et al.,
 766 2012; Ter-Mikaelian et al., 2013; Aushana et al., 2018). At the level of AI, studies have

767 pointed out the relationships between the noise impact on the cortical and behavioral
 768 discrimination performance. In bird field L (homologous to AI), neuronal responses to song
 769 motifs were strongly reduced by three types of masking noises, and the neural discrimination
 770 performance was progressively reduced when the SNR decreased, in parallel with the
 771 behavioral performance (Narayan et al., 2007). Our VRB results are reminiscent of those
 772 obtained in area NCM (homologous to a secondary area) where feed-forward inhibition
 773 allowed the emergence of invariant neural representations of target songs in noise conditions
 774 (Schneider and Woolley, 2013). Similarly to the results by Ranasinghe and colleagues (2012),
 775 our IC neuronal responses were found to be resistant to drastic spectral degradations.
 776 Only one study directly compared the impact of vocoding and masking noise on cortical
 777 responses to vocalizations (Nagarajan et al., 2002). In this study, auditory cortex responses
 778 were robust to spectral degradations even in response to 2-band vocoded vocalizations. Also,
 779 broadband white noise reduced neuronal responses at 0 dB SNR. Last, temporal-envelope
 780 degradations strongly reduced the evoked firing rate and the neural synchronization to the
 781 vocalization envelope. Importantly, band-pass filtering the vocalizations between 2-30 Hz did
 782 not reduce firing rate and neural synchronization to the vocalization envelope. This is in
 783 agreement with the results in our conditions: when the Δ modulation index (computed between
 784 1 and 20 Hz) revealed modest AM alterations, there was little effect on the neuronal
 785 discrimination, but when the AM alterations reached about 20-30% or more, the neuronal
 786 discriminations were reduced (Fig. 12B). Thus, our results are consistent with the hypothesis
 787 that one of the factors constraining auditory discrimination at the cortical and subcortical level
 788 is the fidelity of transmission and processing of slow AM cues.

789 When quantifying how different noise levels alter neuronal coding in the auditory system, it
 790 was found that the neural representation of natural sounds becomes progressively independent
 791 of the level of background noise from the auditory nerve to the IC and AI (Rabinowitz et al.,
 792 2013). It was proposed that this tolerance to background noise results from an adaptation to
 793 the noise statistics, which is more pronounced at the cortical than at the subcortical level. In
 794 agreement with this study, we found that populations of cortical neurons (AI and VRB) were
 795 more resistant to noise than subcortical ones. However, we did not observe a monotonic
 796 evolution of resistance to noise in the auditory system: at the subcortical level, the
 797 discrimination performance of CN neuronal populations drastically dropped as early as +10
 798 dB SNR, the populations of CNIC neurons maintained the highest discrimination performance
 799 even at the -10 dB SNR, those of thalamic ones largely decreased at 0 dB SNR, whereas

cortical neurons showed the lowest discrimination performance at all SNRs but were more robust to noise. In the IC, previous work showed that background noise changes the shape of the temporal modulation transfer function of individual neurons from bandpass to lowpass (Lesica and Grothe, 2008). The CNIC is a massive hub receiving probably the highest diversity of inhibitory and excitatory inputs (Malmierca, 2004; Ayala et al., 2016) and potentially the large diversity of these inputs allows this structure to extract crucial temporal information about the stimulus temporal envelope, even at relatively low SNR.

Limitations of the study

We previously did not find evidence for higher cortical discrimination in awake animals compared with anesthetized animals (Huetz et al., 2009): with normal and reversed whistle stimuli, the percentage of cortical cells with significant MI values was higher in anesthetized (71%) than in awake animals (44%, Table 1 in Huetz et al., 2009). In addition, the Hmax value (equivalent of MI) was higher in anesthetized than in awake animals (0.38 vs. 0.24, Table 2 in Huetz et al., 2009). Last, the trial-to-trial temporal reliability of cortical cells to whistle calls was not different in anesthetized and awake guinea pigs (anesthetized 0.48 vs. awake 0.42; Fig. 8 in Huetz et al., 2009). A recent study (Town et al., 2018) revealed that the cortical discrimination performance between vowels observed in awake animals using acoustic degradations were similar in anesthetized animals (Bizley et al., 2009). Therefore, based on these two studies, the cortical discrimination performance can only be slightly lower or similar in awake compared to anesthetized animals. At the subcortical level, it seems that there is not a large difference between the phase-locking properties of neurons in anesthetized and awake animals (Joris et al., 2004). Temporal properties of IC neurons are only mildly affected by anesthesia (Ter-Mikaelian et al., 2007), indicating that collicular neurons will still be far better than cortical ones to follow the 10-20 Hz temporal cues contained in the four vocalizations. Together, these studies suggest that the hierarchy between cortical and subcortical structures in discriminating communication sounds should be more pronounced or should remained the same in awake animals.

Another limitation of the present study lies in the use of a limited set of stimuli that is restricted to the four same whistles. However, the four whistles used here were clearly representative of our whole database of whistles in terms of frequency range, duration, range of frequency and amplitude modulations. Changing the four whistles from one recording to another can help generalizing the results, but the main advantage of using exactly the same four whistles is that from one recording to the next, and from one structure to another, we

were sure that the same acoustic cues were available for the neural discrimination. However, the whistles are a subset of the guinea pig repertoire, and therefore the present results may not generalize to other communication sounds, and larger sets of stimuli should be used to confirm that the slow AM cues control the neural discrimination. Even if amplitude modulations seem the main cues for speech understanding (Drullman et al., 1994; Shannon et al., 1995), other factors (the pitch, the frequency modulation, the harmonicity cues) can also be involved.

As our results are based on multiunit recordings, we do not know whether the same number of neurons were present in the cluster recordings from the different structures, and whether the individual discrimination performance of the cell types found in each structure are equivalent. On the other hand, the MI evaluated here is the reflection of a local computation performed by a small population of individual neurons, which gives us a good estimation of the whole discrimination performance of a given structure.

Functional implications

In humans, speech sounds (such as phonemes) showing similar acoustic properties trigger similar responses and are represented as a single category in the superior temporal gyrus (Mesgarani et al., 2014b). As already proposed by Chechick and Nelken (2012), auditory cortex neurons extract abstract auditory entities rather than detailed spectro-temporal features. Obviously, this urges to define the acoustic features that form a category of auditory objects. It is relatively easy to delimit broad categories such as environmental sounds, animal vocalizations, music and speech (Gygi and Shafiro, 2013; Gygi et al., 2004, 2007; Woolley et al., 2005; Singh and Theunissen, 2003) in terms of modulation cues, but within these categories, defining invariant features is a difficult task. Here, the use of vocalizations belonging to the same category of the guinea pig repertoire, i.e. “whistles”, may explain both the relatively poor discrimination abilities of cortical neurons compared to subcortical ones and the robustness of cortical responses to vocoding and background noise.

From the present study, it appears that the subcortical structures engage significantly more neurons (20-40%) with high discrimination performance than the cortical areas (2-3% see Fig. 5A), confirming that the neural code is rather sparse at the cortical level (Hromádka et al., 2008), which might not be the case at the subcortical level. However, it is also possible that top-down projections coming from auditory cortex and reaching the thalamus, inferior colliculus and cochlear nucleus (Jacomme et al., 2003; Malmierca and Ryugo, 2011)

868 influence the neural discrimination at the subcortical level, especially in awake, behaving,
869 animals. Thus, we can envision that in behaving animals, learning-induced cortical plasticity
870 also contributes to enhancing the subcortical neural discrimination via the corticofugal
871 projections. Further studies are required to determine to what extent these subcortical
872 representations influence auditory abilities in animals and humans.
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	CN	Lemniscal pathway			Non-lemniscal pathway
		CNIC	MGv	A1	VRB
Number of animals	10	11	10	11	5
Number of recordings tested	672	478	448	544	192
TFRP only	560	421	285	455	126
TFRP and significant response to at least one vocalization	499	386	262	354	95
TFRP quantifications					
BF Range (kHz): min-max	0.18 - 18	0.34 - 36	0.33 - 33	0.14 - 36	0.67 - 36
Mean bandwidth (octave)	3.91	2.88	4.16	2.07	1.79
Mean response duration (ms)	26.83	35.37	17.31	43.73	44.83
Response strength (AP/sec)	77.23	82.25	41.61	37.69	19.97

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Table 1. Summary of the number of animals, number of selected recordings and TFRP quantifications in each structure.

Figure Legends

Figure 1. Spectrograms, spectra and temporal envelopes of the acoustic stimuli. **A-C.** Spectrograms (A), spectra (B) and temporal envelopes (C) of the four original whistles used in this study. **D-F.** From left to right, spectrograms (D), spectra (E) and temporal envelopes (F) of the four vocoded whistles using 38, 20 and 10 frequency bands. **G-I.** From left to right, spectrograms (G), spectra (H) and temporal envelopes (I) of the four original whistles embedded in a vocalization-shaped stationary noise at three SNRs (+10, 0 and -10 dB).

Figure 2. Evolution of the CorrCoef and MI mean values as a function of temporal precision in each structure. **A.** The trial-to-trial temporal reliability, quantified by the CorrCoef, was calculated from responses to original vocalizations with a width Gaussian window varying from 1 to 50 ms in CN (in black), CNIC (in green), MGv (in orange), A1 (in blue) and VRB (in purple). In our study, a 10-ms width Gaussian window (dashed black line) was selected for the data analysis in each structure. **B.** Mutual information (in bits) was calculated from neuronal responses to original vocalizations with a bin size varying from 1 to 40 ms in CN (in black), CNIC (in green), MGv (in orange), A1 (in blue) and VRB (in purple). In this study, the value of 8 ms was selected for the data analysis because in each structure, the MI value was maximal (dashed black line). This hold true also in the different conditions of acoustic alterations, both in noise and vocoded conditions (data not shown).

Figure 3. Subcortical neurons discriminate better the original vocalizations than cortical neurons. **A.** From bottom to top, neuronal responses were recorded in CN, CNIC, MGv, A1 and VRB simultaneously under 16 electrodes but only two are represented here, with alternated black and red colors. Each dot represents the emission of an action potential and each line corresponds to each presentation of one of four original whistles. The grey part of rasters corresponds to evoked activity. For each example, the values of the best frequency (BF in kHz) and of the bandwidth (BW in octave) obtained when testing the responses to pure tones are indicated in the left side. The waveforms of the four original whistles are displayed under the rasters. **B.** Peristimulus time histograms (PSTHs) of each neuronal response presented in A. For each neuronal recording, the four PSTHs of the four original whistles have been overlaid.

C-F. The panels show the mean values of (C) the evoked firing rate (spikes/sec), (D) the trial-to-trial temporal reliability quantified by the CorrCoef value, (E) the neuronal discrimination assessed by the mutual information (MI) computed at the level of the individual recording ($MI_{\text{Individual}}$, bits) and (F) the neuronal discrimination at the population level ($MI_{\text{Population}}$, bits) with populations of 9 simultaneous multiunit recordings obtained with the four original vocalizations in CN (in black), CNIC (in green), MGv (in orange), A1 (in blue) and VRB (in purple). The evoked firing rate corresponds to the total number of action potentials occurring during the presentation of the stimulus minus spontaneous activity (200 ms before each acoustic stimulus). In each structure, error bars represent the SEM of the mean values and black lines represent significant differences between the mean values (unpaired t test, $p < 0.05$). The evoked firing rate decreases from the CN to VRB but both the trial-to-trial temporal reliability (CorrCoef) and the discrimination performance (MI) reach a maximal value in MGv. Note also that at the population level, all the subcortical structures discriminate better the original vocalizations than cortical areas. **G.** Scatter plots showing in each structure, the strong correlations ($0.77 < r < 0.88$) between the CorrCoef and the $MI_{\text{Individual}}$ (bits) values obtained in original conditions.

Figure 4. Diversity of neuronal discrimination performance in quiet for each structure at the individual and population level.

A. Neural discrimination performance in response to original vocalizations in each auditory structure. Waterfall plots show the mutual information (MI, *bits*) as a function of temporal resolution (1 to 256 ms) for the selected recordings in CN (A1), CNIC (A2), MGv (A3), A1 (A4) and VRB (A5). In each structure, the units are ranked by the mean MI value obtained for all bin sizes. Note that there was a larger proportion of neurons with high values of MI (close from the maximal value of 2 bits) in MGv, CNIC and CN (*red curves*) compared to a much lower proportion in the cortical areas AI and VRB.

B. Population information quickly reaches high values with simultaneous multiunit recordings at the subcortical but not cortical level. For each auditory structure, each thin line represents a particular case of simultaneous recording with a maximum number of electrodes (maximum 16 simultaneous multiunit recordings) and each thick line represents the mean value of $MI_{\text{Population}}$ in CN (B1, in black), CNIC (B2, in green), MGv (B3, in orange), A1 (B4, in blue) and VRB (B5, in purple). Note that the mean $MI_{\text{Population}}$ value quickly reaches high values close from the maximum value of 2 bits in the subcortical structures (CN, CNIC and MGv) compared to the two cortical areas (A1 and VRB).

Figure 5. High discrimination performance neurons are more numerous in subcortical structures than in auditory cortex in original conditions. **A.** Cumulative percentage of the neuronal discrimination performance obtained in original vocalizations assessed by the mutual information (MI) computed at the level of the individual recordings ($MI_{\text{Individual}}$, bits) and (**B**) at the population level ($MI_{\text{Population}}$, bits) with populations of 9 simultaneous multiunit recordings in CN (in black), CNIC (in green), MGv (in orange), A1 (in blue) and VRB (in purple).

Figure 6. Vocoding slightly alters neuronal responses at each stage of the auditory system. **A.** From left to right, examples of raster plots representing the responses to the four original whistles (*Original*) and their vocoded versions (*Voc38*, *Voc20* and *Voc10*). The grey part of rasters corresponds to evoked activity. From bottom to top, neuronal responses were recorded in CN, CNIC, MGv, A1 and VRB. For each example, the values of the best frequency (BF in kHz) and of the bandwidth (BW in octave) obtained when testing the responses to pure tones are indicated in the left side. For each example, the mean evoked firing rate (spikes/sec) obtained in each condition is indicated below the rasters. **B.** Peristimulus time histograms (PSTHs) of each neuronal response presented in A. For each neuronal recording, the four PSTHs of the original and vocoded conditions have been overlayed. The grey part of the PSTHs corresponds to evoked activity. **C-F.** The mean values ($\pm$ SEM) represent the vocoding effects on (**C**) the evoked firing rate (spikes/sec), (**D**) the temporal reliability represented by the CorrCoef value, (**E**) the neuronal discrimination assessed by the mutual information (MI) computed at the level of the individual recordings ($MI_{\text{Individual}}$, bits) and (**F**) the neuronal discrimination at the population level ($MI_{\text{Population}}$, bits) with populations of 9 simultaneous multiunit recordings in CN (in black), CNIC (in green), MGv (in orange), A1 (in blue) and VRB (in purple) (one-way ANOVA, $P < 0.05$; with post-hoc paired t tests, $*P < 0.05$). The evoked firing rate corresponds to the total number of action potentials occurring during the presentation of the stimulus minus spontaneous activity (200 ms before each acoustic stimulus). At the population level, the discrimination performance significantly decreased only for 10 frequency bands in subcortical structures and did not decrease in cortical areas.

Figure 7. Vocoding effects on the $MI_{\text{Population}}$ growth functions in each auditory structure. The curves display the average growth functions of the $MI_{\text{Population}}$ for each structure in each vocoding condition (indicated by a gradient colors) in CN (**A**, in black), CNIC (**B**, in green), MGv (**C**, in orange), A1 (**D**, in blue) and VRB (**E**, in purple). In each structure, the vocoding slightly reduced the $MI_{\text{Population}}$ values. At the cortical level, the reduction induced by vocoding was similar at 38 and 20 bands, then a stronger reduction was observed at 10 bands. At the thalamic level, there was almost no change in the growth function of the $MI_{\text{Population}}$ with 38 and 20 bands vocalizations, but there was a large decrease in $MI_{\text{Population}}$ with the 10-band vocoded stimuli. In the CNIC, the vocoding only induced a reduction of the $MI_{\text{Population}}$ for 20 and 10 bands; a similar scenario was observed at the CN level.

Figure 8. Noise strongly reduces neuronal responses in all structures but to a lesser extent in the central nucleus of the inferior colliculus. **A.** From left to right, raster plots of responses of four original whistles (*Original*) and their noisy versions embedded in the vocalization-shaped stationary noise at three SNRs: +10, 0 and -10 dB. The grey part of rasters corresponds to evoked activity. From bottom to top, neuronal responses were recorded in CN, CNIC, MGv, A1 and VRB. For each example, the values of the best frequency (BF in kHz) and of the bandwidth (BW in octave) obtained when testing the responses to pure tones are indicated in the left side. For each example, the mean evoked firing rate (spikes/sec) obtained in each condition is indicated below the rasters. The green dashed box indicates a typical example of CNIC neuronal responses that are resistant to the noise addition. **B.** Peristimulus time histograms (PSTHs) of each neuronal response presented in A. For each neuronal recording, the four PSTHs of the original and noisy conditions have been overlaid. The grey part of the PSTHs corresponds to evoked activity. **C-F.** The mean values ($\pm$ SEM) represent the noise effects on (**C**) the evoked firing rate (spikes/sec), (**D**) the temporal reliability represented by the CorrCoef value, (**E**) the neuronal discrimination assessed by the mutual information (MI) computed at the level of the individual recordings ($MI_{\text{Individual}}$, bits) and (**F**) the neuronal discrimination at the population level ($MI_{\text{Population}}$, bits) with populations of 9 simultaneous multiunit recordings in CN (in black), CNIC (in green), MGv (in orange), A1 (in blue) and VRB (in purple) (one-way ANOVA, $P < 0.05$; with post-hoc paired t tests, $*P < 0.05$). The evoked firing rate corresponds to the total number of action potentials occurring during the presentation of the stimulus minus spontaneous activity (200 ms before each acoustic stimulus). At the population level, the discrimination performance significantly decreased in all structures when the SNR decreased, with on average the CNIC populations still able to discriminate 2 out of 4 stimuli ($MI_{\text{Population value}} > 1\text{bit}$).

Figure 9. Noise effects on the $MI_{\text{Population}}$ growth functions in each auditory structure. The curves display the noise effects on the $MI_{\text{Population}}$ growth functions for each structure and at each SNR (indicated by a gradient colors) in CN (**A**, in black), CNIC (**B**, in green), MGv (**C**, in orange), A1 (**D**, in blue) and VRB (**E**, in purple). In general, background noise largely altered the growth functions of the $MI_{\text{Population}}$ in each structure (but to a lesser extent in the CNIC). In the CN, noise induced a stronger reduction of the $MI_{\text{Population}}$, which was clearly a function of SNR. In the CNIC, noise induced SNR-dependent reduction in the $MI_{\text{Population}}$ values, the reduction being modest at a +10 and 0 dB SNR but more important at a -10 dB SNR. In the MGv, noise progressively lowered the curves of the $MI_{\text{Population}}$. In the cortex, the $MI_{\text{Population}}$ growth functions were not strongly impacted except at the -10 dB SNR.

Figure 10. A subpopulation of CN neurons maintains good neuronal discrimination performance at a +10 dB SNR.

A. Waterfall plot shows the mutual information ($MI_{\text{Individual}}$, bits) as a function of temporal resolution (1 to 256 ms) for the CN recordings at +10 dB SNR. The recordings are ranked by the mean MI value obtained for all bin sizes. Note that at this particular SNR, 20% of the CN recordings still maintained $MI_{\text{Individual}}$ values above 1 bit, indicating that some CN neurons still send information about the vocalization identity at higher brainstem centers such as the CNIC. **B.** Distributions of the Time-Frequency Response Profile (TFRP) parameters (best frequency, bandwidth, response duration and response strength) for the two neuronal populations in CN depending of the MI value (*in grey*, $MI \geq 1\text{bit}$ and *in black*, $MI < 1\text{bit}$). Note that, there were significant differences in terms of response duration and the response strength. **C.** The curves display the individual and average growth functions of the $MI_{\text{Population}}$ for the simultaneous CN recordings at the +10 dB SNR. Note that despite the fact that the mean $MI_{\text{Population}}$ value was much lower than in the original condition (see figure 4B1), about 20% of the simultaneously recorded populations reached a value of 1.5 bits with 9 neurons or less (*red curve lines*).

Figure 11. No relationship between the mutual information and the parameters of TFRPs (the best frequency, BF and the bandwidth, BW) at each stage of the auditory system.

A. Typical examples of Time-Frequency Response Profile (TFRP) recorded in VRB, AI, MGv, CNIC and CN. These TFRPs are examples of responses to pure tones and the first column also corresponds to the same neurons as those presented in figures 3, 5 and 7. *From left to right*, the maximal firing rate (in spikes/sec) was 100 and 220 in VRB, 195 and 200 in AI, 460 and 420 in MGv, 315 and 250 in CNIC and 340 and 330 in CN. From these TFRPs, we extracted parameters such as the best frequency (in kHz), the bandwidth (in octave), the response duration (in ms) and the response strength (in spikes/sec). **B.** Noise effect on neuronal discrimination ($MI_{\text{Individual}}$, bits) according to the best frequency (BF). Scattergrams of the $MI_{\text{Individual}}$ values obtained at the +10 dB SNR as a function of the $MI_{\text{Individual}}$ values obtained with the original vocalizations based on neuronal responses recorded in CN, CNIC, MGv, A1 and VRB. We separated the recordings in three groups according to the best frequency: $BF < 5\text{kHz}$ (in red), $5 \leq BF \leq 15\text{kHz}$ (in blue) and $BF > 15\text{kHz}$ (in green). $MI_{\text{Individual}}$ mean values are represented with a black cross. **C.** Noise effect on neuronal discrimination ($MI_{\text{Individual}}$, bits) according to the bandwidth (BW). Scattergrams of the $MI_{\text{Individual}}$ values obtained at the +10 dB SNR as a function of the $MI_{\text{Individual}}$ values obtained with the original vocalizations based on neuronal responses recorded in CN, CNIC, MGv, A1 and VRB. We separated the recordings in three groups according to the bandwidth: $BW \leq 2$ octaves (in red), $2 \leq BW \leq 4$ octaves (in blue) and $BW \geq 4$ octaves (in green). $MI_{\text{Individual}}$ mean values are represented with a black cross. Note that, in all structures, the decrease in $MI_{\text{Individual}}$ value from the original conditions to the +10 dB SNR occurred whatever the BF and the BW values.

Figure 12. Reduction of slow AM cues as one of the factors explaining the neuronal discrimination performance at the subcortical and cortical levels. **A.** Vocoding and noise effects on the amplitude-modulation (AM) spectra. The plot represents the averaged modulation spectra of the four original vocalizations (*in black*), vocoded vocalizations (Voc38, Voc20 and Voc10: *red, green and blue respectively, solid lines*) and vocalizations in noise at three SNRs (+10, 0 and -10 dB: *red, green and blue respectively, dashed lines*).

1296 Vertical black dashed line corresponds to the maximum frequency (20 Hz) selected for the
1297 data analysis.
1298 **B.** Percentage of $\Delta MI_{\text{Population}}$ as a function of $\Delta \text{modulation index}$ computed for each structure
1299 from mean $MI_{\text{Population}}$ or mean modulation-index values obtained in all adverse conditions and
1300 mean values in the original condition. Each dot represents neuronal data ($\Delta MI_{\text{Population}}$) in CN
1301 (*in black*), CNIC (*in green*), MGv (*in orange*), A1 (*in blue*) and VRB (*in purple*). Polynomial
1302 curves fitting all acoustic conditions have been generated (*color lines*). In all conditions
1303 (vocoding or noise), there is a limit of AM reduction from which the $\Delta MI_{\text{Population}}$ decreases in
1304 cortical and subcortical structures.
1305
1306

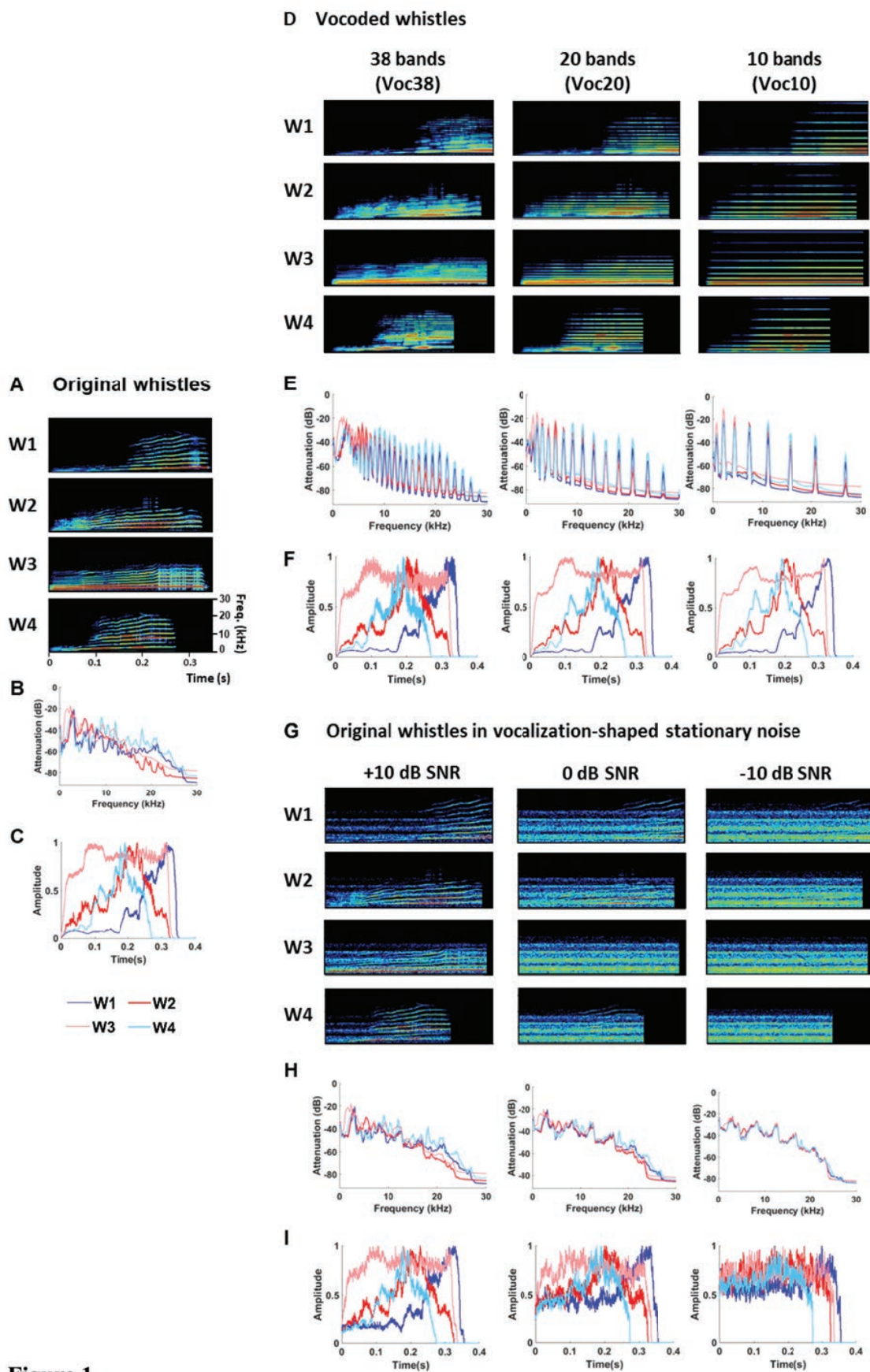


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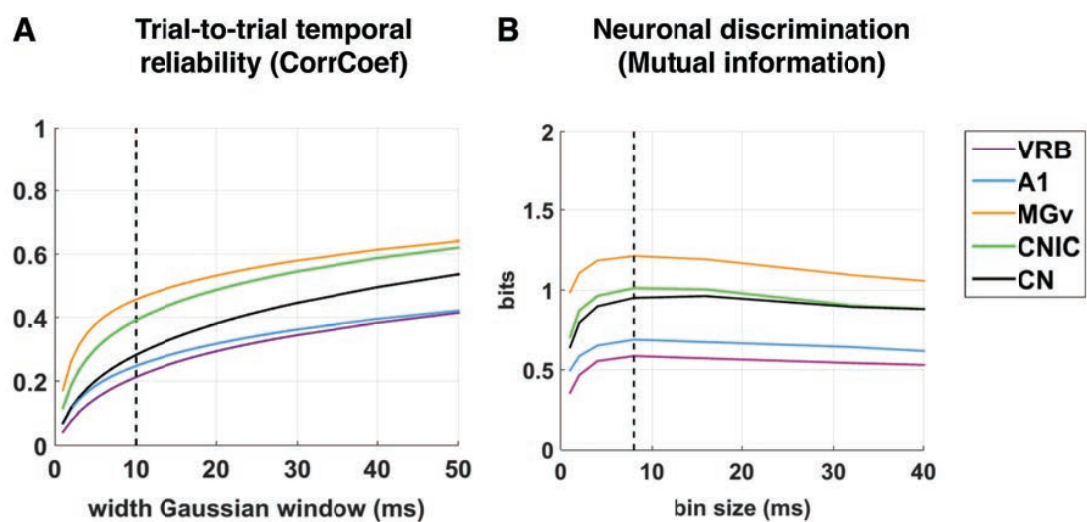


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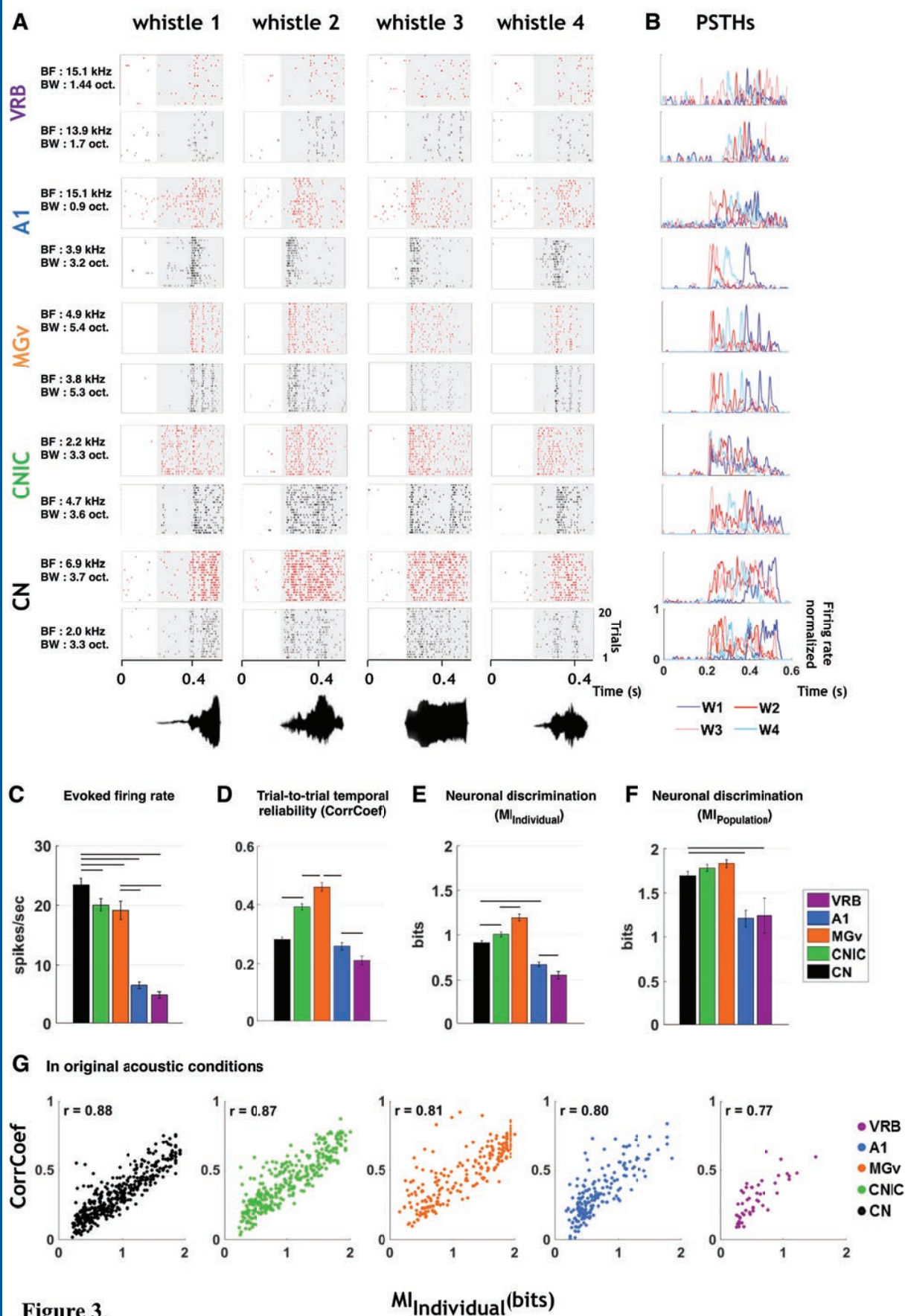


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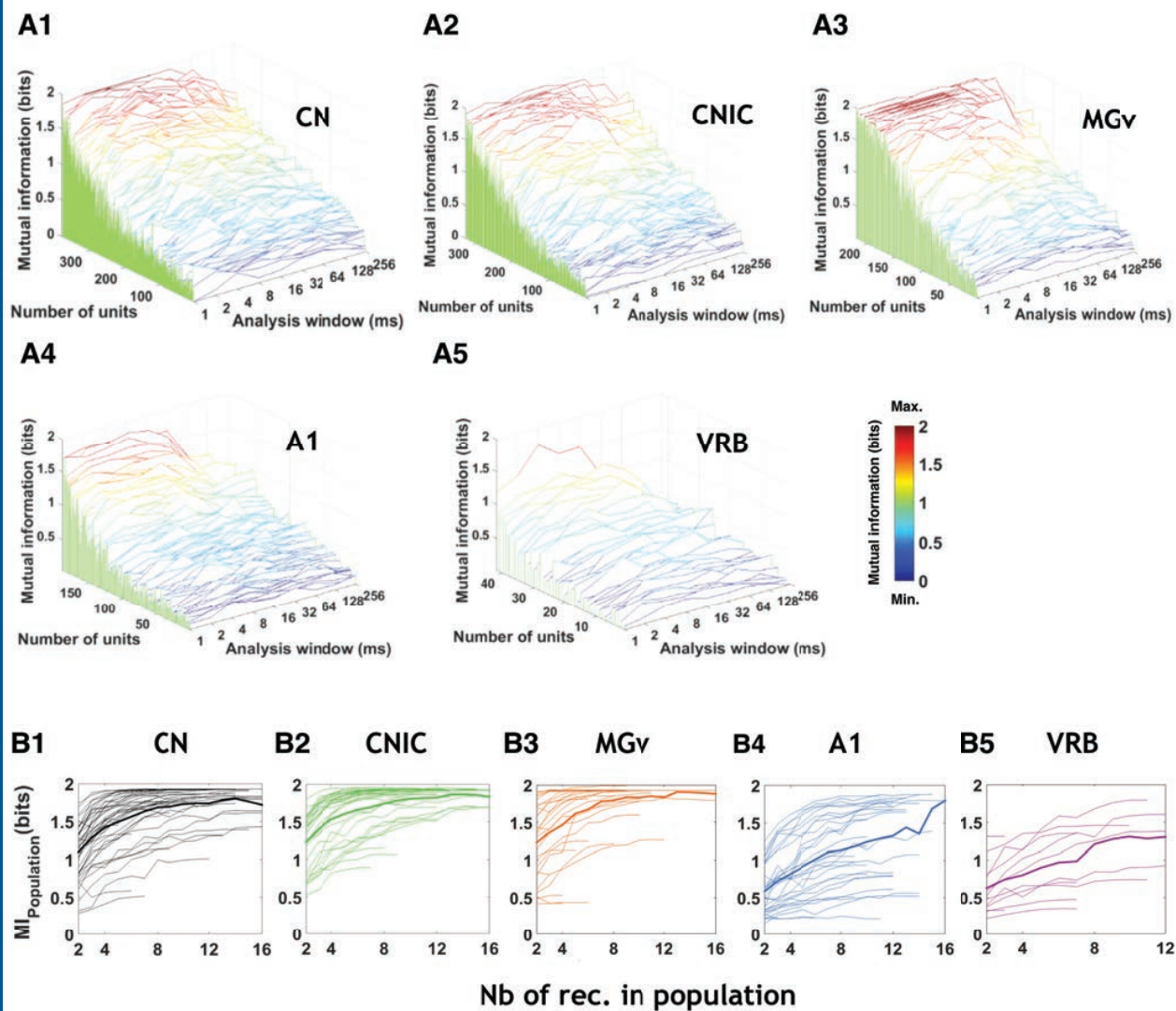


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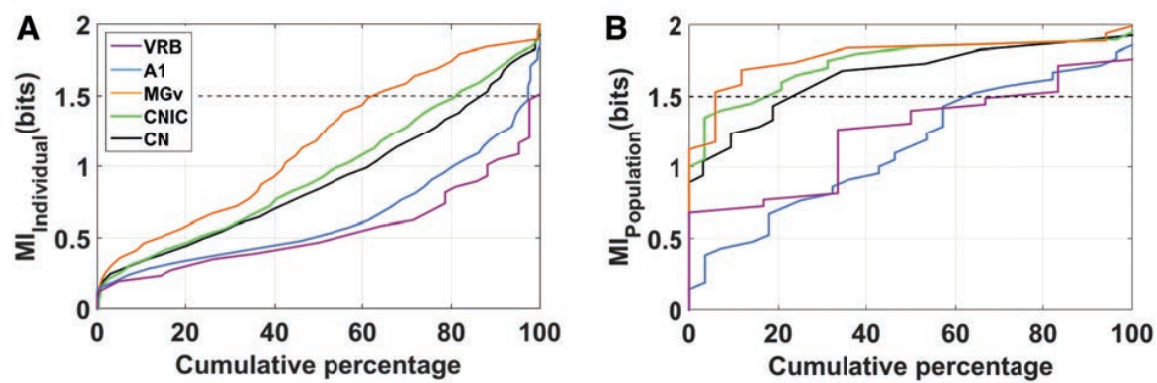


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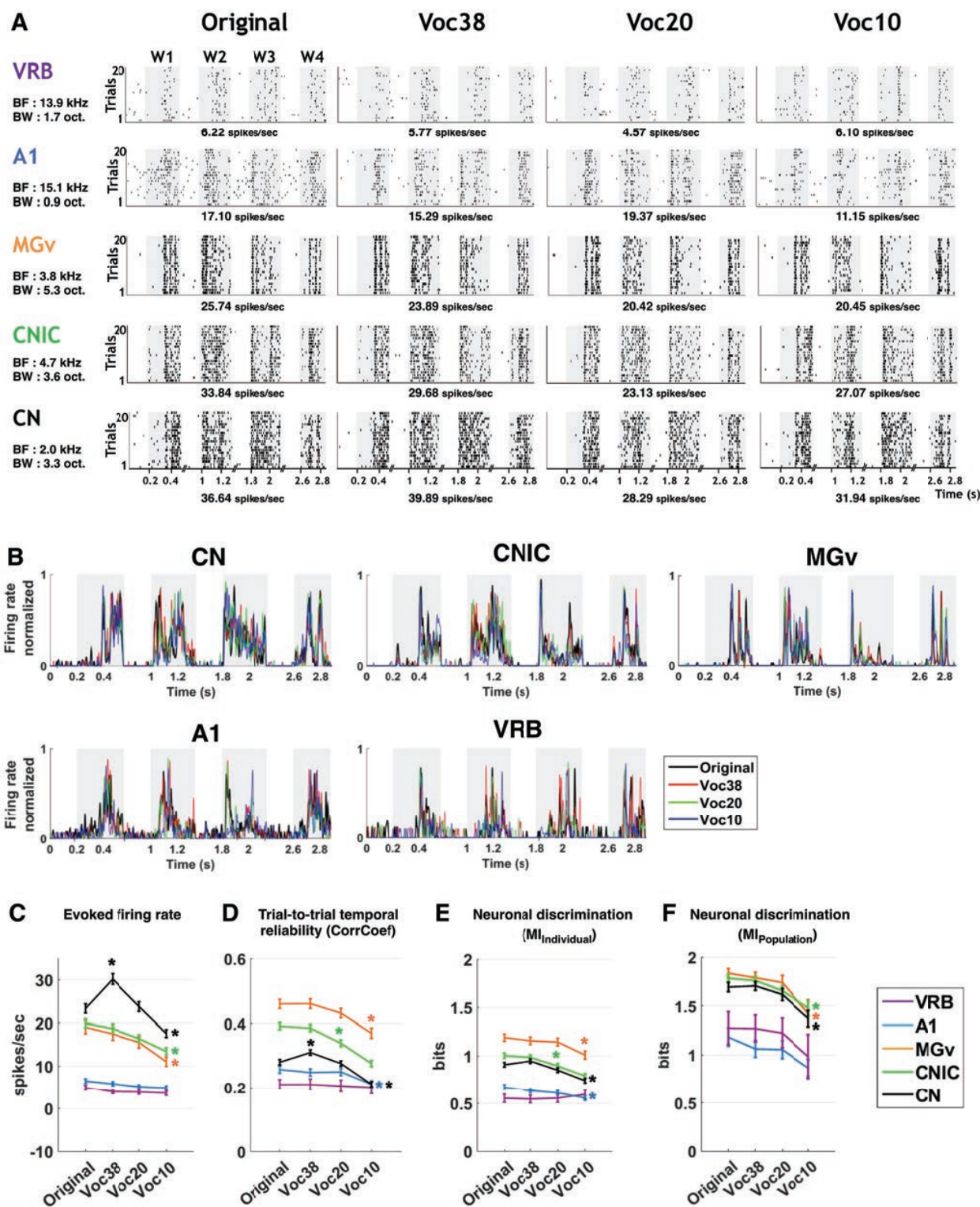


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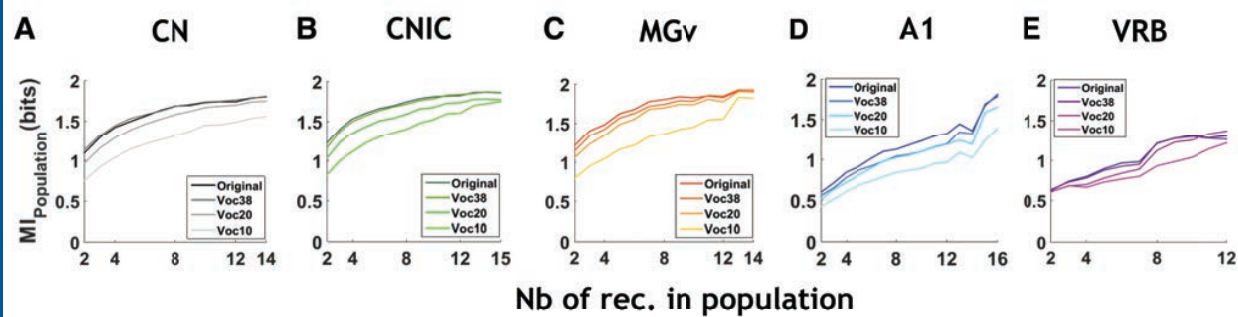


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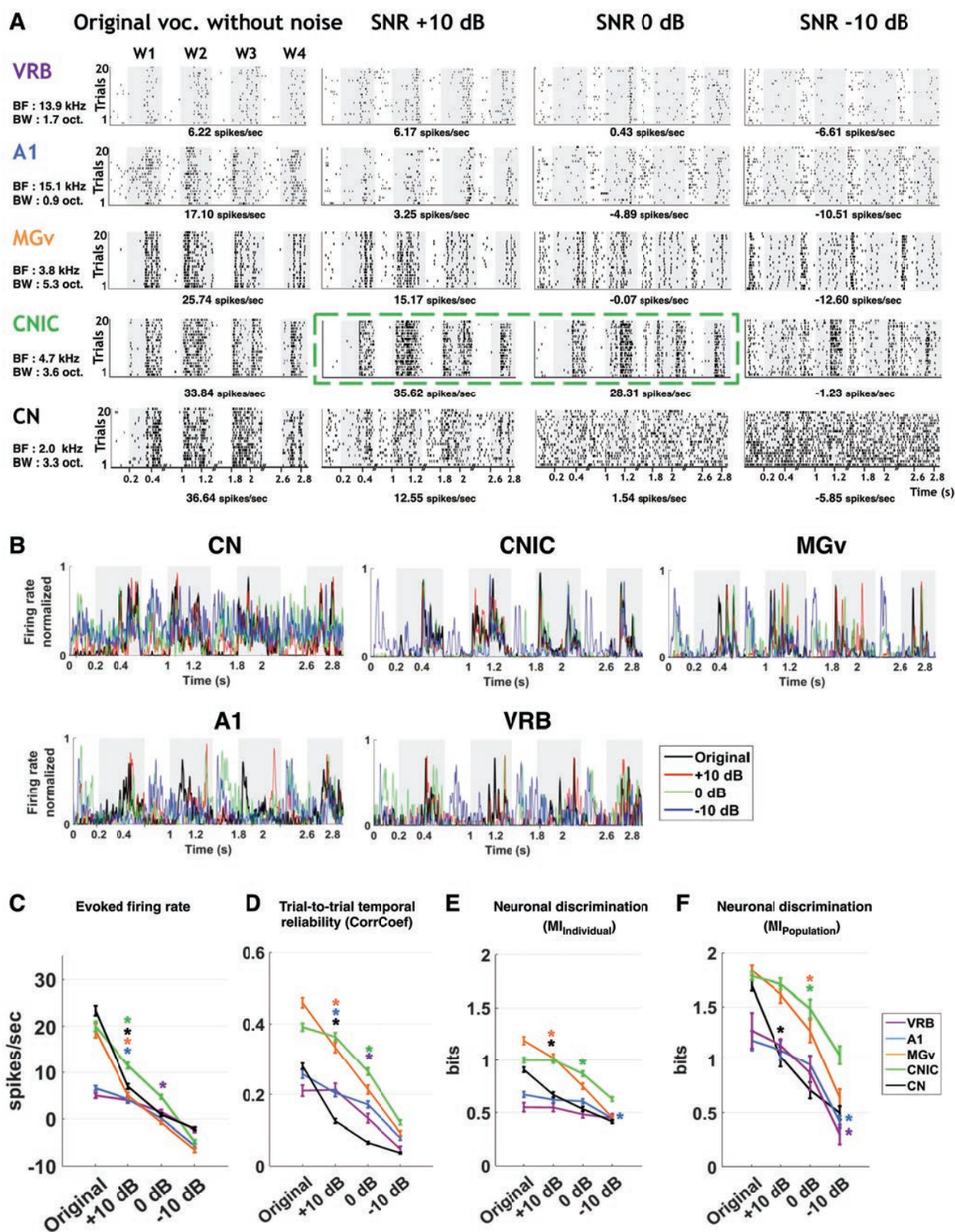


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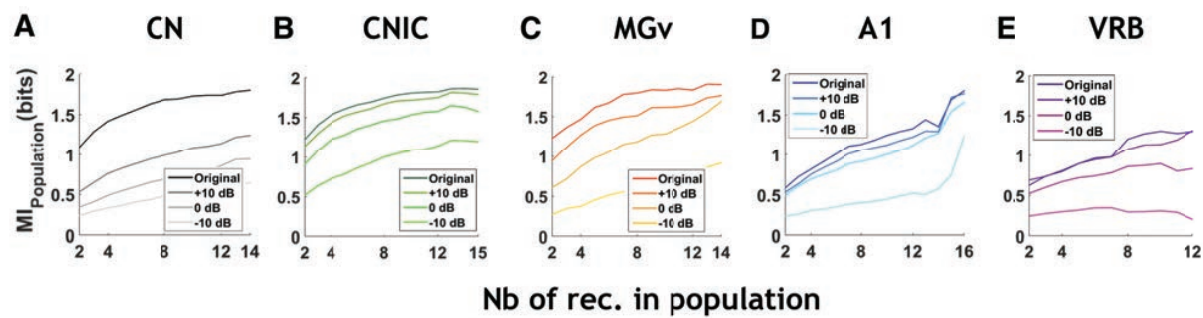


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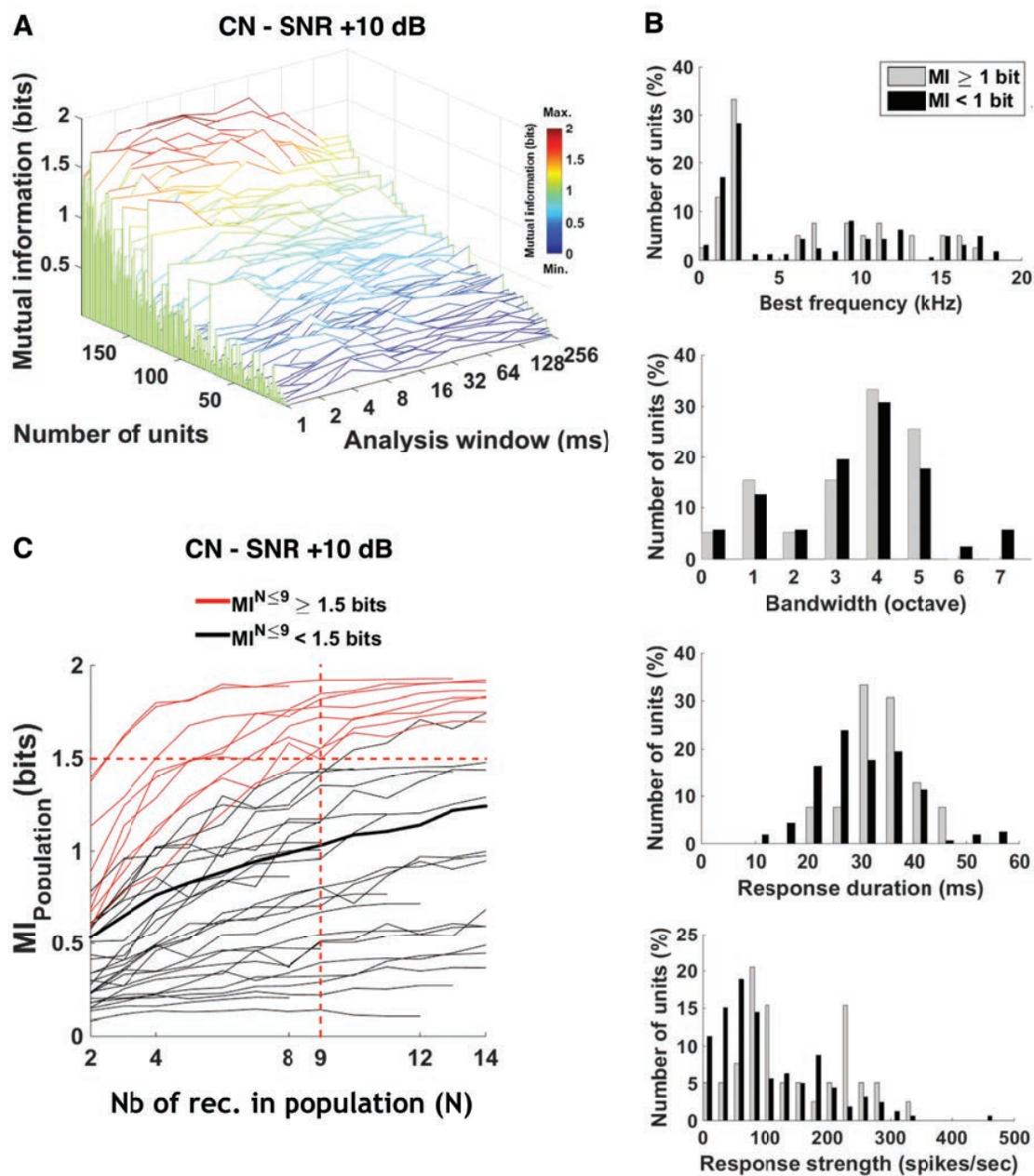


Figure 10.

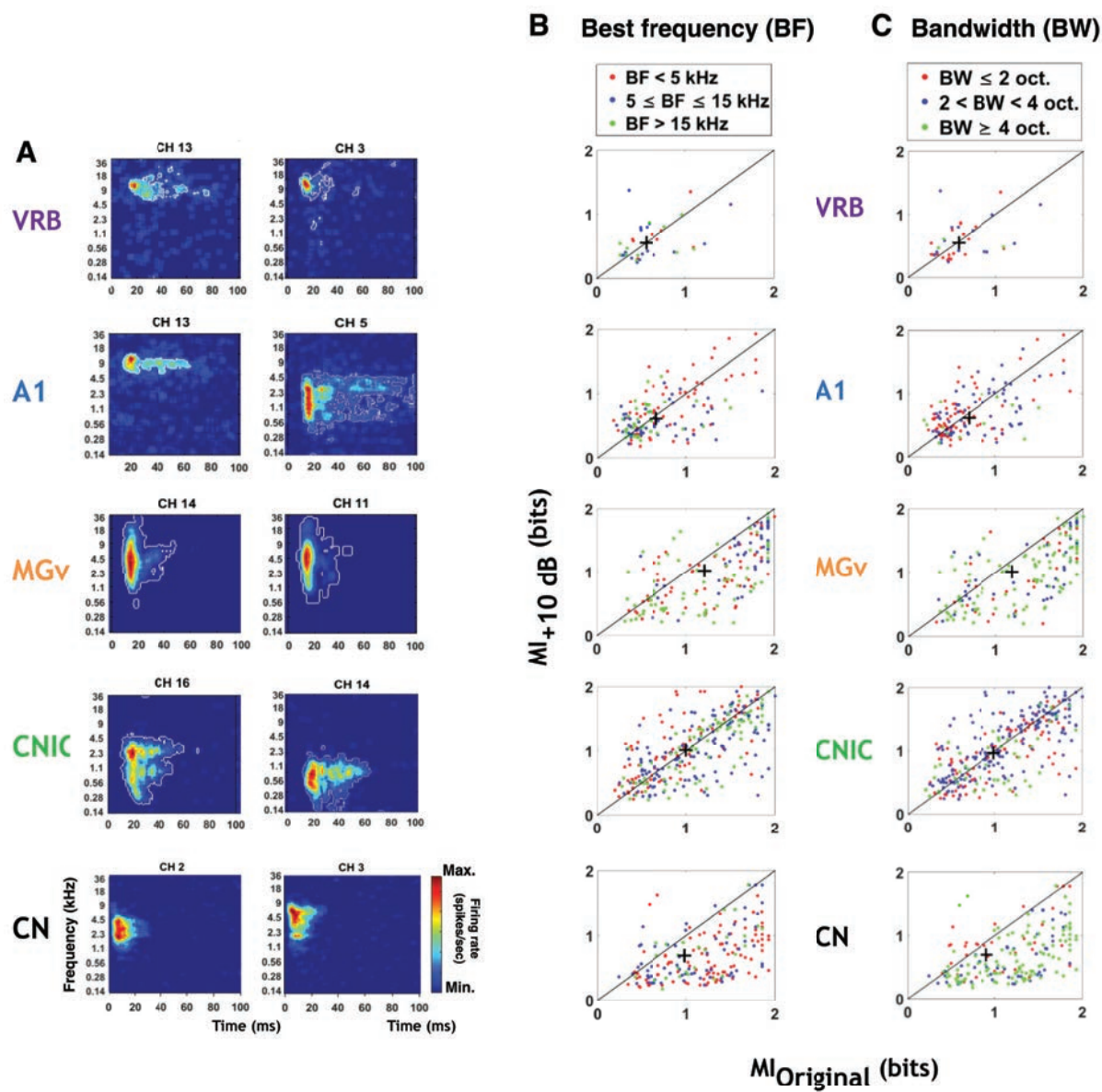


Figure 11.

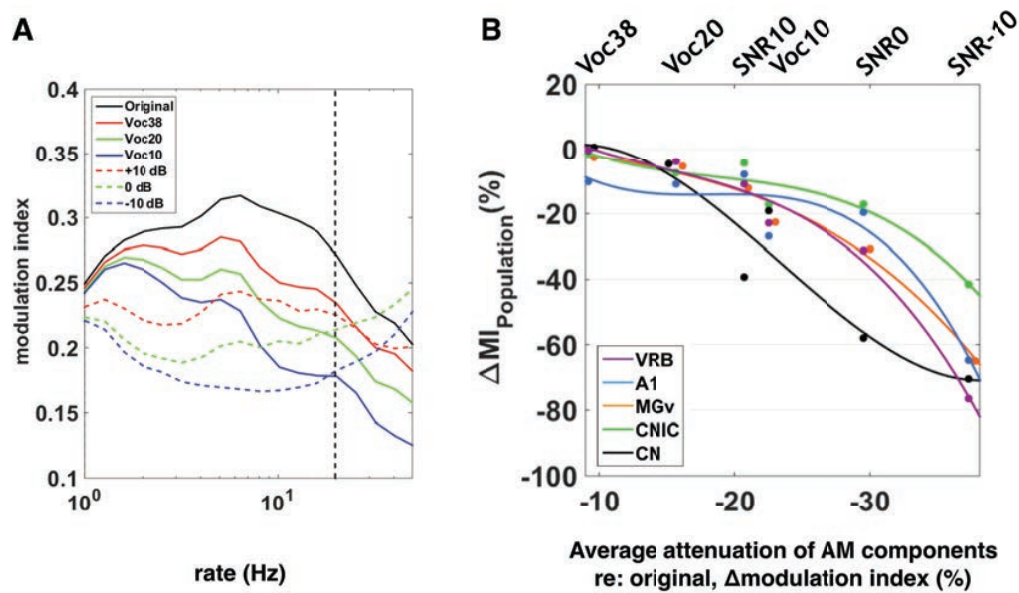


Figure 12.