# Fronto-temporal coupling dynamics during spontaneous activity and auditory processing

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- 13 coupling, coherence, sensory coding, auditory processing
- 14 Abstract
- 15 Most mammals rely on the extraction of acoustic information from the environment in order to
- 16 survive. However, the mechanisms that support sound representation in auditory neural networks
- 17 involving sensory and association brain areas remain underexplored. In this study, we address the
- 18 functional connectivity between an auditory region in frontal cortex (the frontal auditory field, FAF)
- 19 and the auditory cortex (AC) in the bat *Carollia perspicillata*. The AC is a classic sensory area
- 20 central for the processing of acoustic information. On the other hand, the FAF belongs to the frontal
- 21 lobe, a brain region involved in the integration of sensory inputs, modulation of cognitive states, and
- in the coordination of behavioural outputs. The FAF-AC network was examined in terms of
- 23 oscillatory coherence (local-field potentials, LFPs), and within an information theoretical framework
- 24 linking FAF and AC spiking activity. We show that in the absence of acoustic stimulation,
- 25 simultaneously recorded LFPs from FAF and AC are coherent in low frequencies (1-12 Hz). This
- 26 "default" coupling was strongest in deep AC layers and was unaltered by acoustic stimulation.
- 27 However, presenting auditory stimuli did trigger the emergence of coherent auditory-evoked gamma-
- 28 band activity (>25 Hz) between the FAF and AC. In terms of spiking, our results suggest that FAF
- 29 and AC engage in distinct coding strategies for representing artificial and natural sounds. Taken
- 30 together, our findings shed light onto the neuronal coding strategies and functional coupling
- 31 mechanisms that enable sound representation at the network level in the mammalian brain.

#### 32 1 Introduction

33 Many animals rely on the processing of acoustic information for survival. Nevertheless, the 34 mechanisms by which sounds are represented in neural networks involving distant areas in the brain 35 remain obscure. In the mammalian cortex, sensory and integration areas have been described as part 36 of putative neural networks tasked with sound processing. The auditory cortex (AC), for example, 37 plays an important role in sound analysis and even in coordinating acoustically guided behaviours 38 (Song et al., 2010; Li et al., 2017). Neuronal activity within the AC represents a large range of 39 acoustic properties including spectrotemporal structure (Gaese and Ostwald, 1995; Lu et al., 2001; 40 Yin et al., 2011; Gaucher et al., 2013; Kanold et al., 2014; Gao and Wehr, 2015; Lu et al., 2016; 41 Martin et al., 2017; Sheikh et al., 2019), sound source location encompassing azimuth/elevation coding (Recanzone, 2000; Mrsic-Flogel et al., 2005; Salminen et al., 2015; Trapeau and 42 43 Schonwiesner, 2018) and target distance processing (Suga and O'Neill, 1979; Hechavarria et al., 44 2013; Bartenstein et al., 2014; Beetz et al., 2016), as well as abstract properties such as sound 45 "emotional valence" (Concina et al., 2019) and future behavioural outcomes based on auditory

46 stimuli (Francis et al., 2018).

47 Regions within the frontal lobe of the mammalian brain also participate in auditory processing and 48 could in principle synchronize their activity with that of canonical auditory areas, such as the AC (see 49 (Winkowski et al., 2018)). Frontal and AC regions are strongly connected through feedforward and 50 feedback anatomical pathways (Kobler et al., 1987; Medalla and Barbas, 2014; Plakke and 51 Romanski, 2014; Winkowski et al., 2018). Neurons within the prefrontal cortex (PFC, a region in the frontal lobe) respond to sounds when the latter possess rich spectrotemporal dynamics (Eiermann and 52 53 Esser, 2000; Kanwal et al., 2000; Romanski and Goldman-Rakic, 2002). Additionally, PFC is 54 thought to engage in cognitive processes ranging from attention, learning, and memory 55 formation/retrieval, to decision making (Miller, 2000; Floresco and Ghods-Sharifi, 2007; St Onge et 56 al., 2011; Gourley et al., 2013; Gilmartin et al., 2014; Pezze et al., 2014; Helfrich and Knight, 2016; 57 Kim et al., 2016; Werchan et al., 2016; Helfrich et al., 2017). This area could thus be a fundamental 58 node for sound evaluation in auditory networks, and even for the implementation of acoustically 59 guided behaviours.

Though there is increasing evidence supporting the idea of frontal-AC functional networks for
auditory processing, specifics regarding activity coupling within this network remain unknown. For
example, it remains unclear if different types of neural signals (i.e. spikes and local field potentials,

LFPs) measured simultaneously in frontal and AC areas synchronize during spontaneous activity and
during listening. Moreover, it is unknown whether frontal activity displays preferential coupling
patterns with certain layers of the AC. Assessing the latter can only be achieved by conducting
simultaneous measurements from frontal and AC regions using layer-specific intracranial recordings
to study spikes and LFPs.

68 In the current study, we address the functional connectivity in a fronto-auditory cortical circuit in bats 69 (species *Carollia perspicillata*). The bat AC has been studied extensively, and it has been shown that 70 oscillatory and spiking activity patterns in the bat cortex are in accordance with those observed 71 during the processing of artificial and naturalistic sounds in other animal models, including speech in 72 humans. This comprises phenomena such as multiscale temporal processing of acoustic streams 73 (Giraud and Poeppel, 2012; Hyafil et al., 2015; Hechavarria et al., 2016b; Teng et al., 2017; Garcia-74 Rosales et al., 2018a), interactions between spikes and LFPs for audition (Lakatos et al., 2005; 75 Kayser et al., 2009; Arnal and Giraud, 2012; Kayser et al., 2012; Gilmartin et al., 2014; Garcia-76 Rosales et al., 2018b; Garcia-Rosales et al., 2019), and gamma-band activity for communication call 77 processing (Medvedev and Kanwal, 2008). In bats, there exists a region within the frontal lobe that is 78 responsive to sounds: the frontal auditory field (FAF; (Kobler et al., 1987; Eiermann and Esser, 2000; 79 Kanwal et al., 2000)). This region is anatomically connected with the AC (Kobler et al., 1987), but it 80 also receives auditory afferents via a non-lemniscal pathway through the suprageniculate nucleus of 81 the thalamus, bypassing main auditory centres in the midbrain (Kobler et al., 1987). In addition to 82 pure tones, neurons in the FAF encode spectrotemporally complex sounds with variable latencies and 83 response properties (Eiermann and Esser, 2000; Kanwal et al., 2000; Lopez-Jury et al., 2019). Our 84 goal was to examine specifics of fronto-AC activity in awake bats during the processing of acoustic 85 streams. We tackled this question by quantifying neural synchronization in the FAF-AC network in 86 terms of oscillatory coherence (a mechanism underlying interareal communication; (Fries, 2015)), 87 during both spontaneous activity and the processing of natural and artificial acoustic sequences.

We found that the FAF-AC network is synchronized by default (i.e. without sensory stimulation) in low-frequencies (up to 12 Hz), and that coherence with the FAF is strongest in deep laminae of the AC. In addition, low-frequency coherence between the two structures remains unchanged during acoustic processing, and auditory-evoked gamma-band synchronization emerges at stimulus onset without clear layer specificity. Finally, based on an information theoretical framework, our data suggest that the neuronal coding of acoustic streams in FAF and AC may occur with non-overlapping

94 neural codes. Taken together, the data presented in this manuscript offer insights into the strategies
95 for sound representation in fronto-AC neural networks.

#### 96 2 Results

# 97 2.1 Stimulus-related LFPs in AC lag relative to those in FAF

98 We recorded electrophysiological data from the primary AC, paired with penetrations from the FAF, 99 in 5 awake *Carollia perspicillata* bats (all males; n = 50 penetrations). Recordings in the AC were 100 performed with laminar electrodes inserted perpendicularly into the brain, spanning depths of 0-750 101 µm as measured from the cortical surface. Each penetration in AC was paired with a simultaneous 102 recording from the FAF using a single carbon electrode at an average depth of  $313 \pm 56 \,\mu\text{m}$  (mean  $\pm$ 103 std). Auditory stimuli consisted of artificially constructed syllabic trains with repetition rates of 5.28 104 and 97 Hz in order to test for slow and fast periodicities in acoustic streams, plus another syllabic 105 train that had no clear rhythmicity as syllables were presented in a Poisson-like sequence with 70 Hz average rate (see Methods). The trains consisted of a repeated short duration, broadband distress 106 107 syllable from C. perspicillata, recorded in previous work (Hechavarria et al., 2016a), whose 108 spectrotemporal design is typical of distress syllables emitted by this species. In addition, we 109 presented a natural distress vocalization ("nat" throughout the text) that has been used in previous 110 research (Hechavarria et al., 2016b; Garcia-Rosales et al., 2018a; Garcia-Rosales et al., 2019), and 111 which comprises temporal modulations in low (ca. 4 Hz) and high-frequency ranges (> 50 Hz), 112 corresponding to the bout and syllabic periodicities, respectively, typical of this animal's distress 113 vocalizations (Hechavarria et al., 2016a).

114 Grand average traces of simultaneously recorded LFPs from FAF and AC (the latter at a depth of 450 115 μm, corresponding to input layers in AC) during acoustic stimulation are shown in Fig. 1A. LFP 116 responses from frontal and auditory cortical regions showed clear modulation by the acoustic streams 117 that were well-correlated across structures for each stimulus tested, particularly at depths  $> 200 \,\mu m$ 118 (Fig. S1). Remarkably, population averaged LFPs appeared "faster" in the FAF than in the AC (see 119 Fig 1A, right column; blue: FAF; orange: AC at 450 µm) relative to stimulus onset. In fact, cross-120 correlation analyses of the traces depicted in Fig. 1A showed that LFPs recorded in frontal regions 121 preceded those recorded from the AC by at least 4 ms (Fig. 1B), effectively indicating that primary 122 auditory cortical stimulus-related LFPs in input layers "lag" relative to those in the FAF. We 123 confirmed this trend by systematically determining the temporal lag between LFPs in the AC at

124 various depths, and LFPs from the FAF. Figure 1C summarizes the results by illustrating, for each 125 stimulus, the distribution of lags between FAF and AC across depths (note that negative lags indicate 126 FAF "leading"). Indeed, we observed a significant effect of LFPs in the FAF being "faster" than 127 those in the AC, robust across depths in the latter structure, and also across stimuli (Fig 1C; FDR-128 corrected tailed Wilcoxon signed rank tests, testing that the medians of the lag distributions were 129 significantly less than 0 across electrodes;  $p_{corr} < 0.05$ ; precise  $p_{corr}$  values are indicated next to each 130 heatmap). These results were corroborated by testing FAF-AC lags in LFPs but considering only LFP 131 pairs that were well correlated across structures (correlation coefficients higher than 0.5, shown in

132 **Fig. S1**).

133 That LFPs in the frontal auditory field "lead" relative to those in the auditory cortex suggests the 134 presence of fast inputs into frontal auditory areas, agreeing with a non-lemniscal auditory pathway 135 converging into the FAF and consisting of as few as four synapses in bats (Kobler et al., 1987). We 136 sought for evidence of fast neuronal responses in FAF that would support these observations by 137 measuring response latencies of the neuronal spiking recorded simultaneously in both structures (see 138 Methods). We observed that spiking responses from the FAF could in fact be as fast as spiking 139 responses from the AC, although this effect was found only in a subpopulation of neurons (on 140 average 5.65%  $\pm$  3.19% of the units considered, across stimuli and channels; see Fig. 2A, B, which 141 depicts example responses from one FAF and one AC unit at 450 µm recorded simultaneously). 142 Measured neuronal response latencies from both structures yielded that AC spiking was on average 143 faster than the FAF spiking (Fig. 2C; for illustrative purposes AC responses are those recorded at 144 450 µm). Still, some FAF units exhibited response latencies below 10 ms, indicative of fast acoustic 145 inputs into this structure. By subtracting latencies from simultaneously recorded units in the FAF and 146 the AC (across depths; latencies were pooled from all tested stimuli, but paring was only done within 147 a particular stimulus), it became evident that auditory cortical spiking was typically faster than its 148 FAF counterpart, although some latencies from frontal regions were shorter than those in the AC. 149 Figure 2D shows the distribution of latency differences (across cortical depths in the AC), depicting 150 the abovementioned observations. Latency differences were significantly higher than 0 for all 151 recording depths in the AC, except in the case of the most superficial channel (FDR-corrected 152 Wilcoxon signed rank tests, significance when  $p_{corr} < 0.05$ ; corrected p values across channels are 153 given to the right of the latency distribution heatmap in **Fig. 2D**).

154 Altogether, these results provide evidence supporting that the FAF receives fast auditory inputs.

155 Notwithstanding, such inputs do not necessarily elicit equally fast spiking, suggesting that the

156 neuronal dynamics in frontal areas are "sluggish" in comparison to primary AC. Sluggish dynamics

157 can arise from multiple factors, and may be essential for sensory integration in the frontal cortex (see

158 Discussion).

#### 159 2.2 FAF and AC synchronize in low frequency LFP bands during spontaneous activity

160 Local-field potentials recorded in the FAF and the AC typically showed visible phase

synchronization, even in the absence of acoustic stimulation (see Fig. 3A, where LFP traces from

162 both structures during a single 3 s epoch of spontaneous activity are depicted). We quantified phase

163 coherence between the two structures by means of the imaginary part of the coherency ("iCoh" in

164 this manuscript; see Methods and (Nolte et al., 2004)), for data recorded both during spontaneous and

165 sound-driven activities. The iCoh metric allows to minimize spurious phase-synchrony attributable,

166 for example, to common referencing and passive spreading of field potentials, by effectively

167 removing non-lagged phase correlations (Bastos and Schoffelen, 2015).

168 Coherence analyses revealed that, as shown in **Fig. 3A**, FAF and AC were synchronized in low

169 frequencies. **Figure 3B** depicts population averaged z-normalized (to a surrogate distribution where

170 phase relationships were abolished; see Methods) iCoh values (z-iCoh) across electrode depths in the

171 AC. Elevated low-frequency coherence is evident in deep layers of the AC, suggesting as well that

172 FAF-AC synchrony was depth-dependent. A time-resolved analysis of iCoh (**Fig. 3C**) over the same

173 LFP traces used to calculate values in **Fig. 3B** also showed that low-frequency phase synchrony was

174 strongest in deeper channels, and furthermore limited to the low-frequency region of the coherence

175 spectrum. The data depicted in **Fig. 3C** are shown here for illustrative purposes and serve as a

176 comparison with the time-resolved coherence estimations performed on LFPs recorded during

177 acoustic processing (see below).

178 To statistically corroborate our observations, we divided the coherence spectrum into canonical

179 frequency bands encompassing delta (1-4 Hz), theta (4-8 Hz), alpha (8-12 Hz), beta (12-25 Hz) and

180 low-gamma (25-45 Hz). We then tested if z-normalized iCoh values in each band were significantly

181 different than 0 across the population. Because of the nature of the surrogate analyses (see Methods),

182 non-consistent phase synchronization in the data would yield a distribution of z-iCoh statistically

183 indistinguishable from 0. In order words, z-iCoh values significantly higher than 0 suggest consistent,

184 population-wise phase-locking between LFPs in FAF and AC during spontaneous activity. Figure

- 185 **3D** (top) depicts the z-iCoh calculated between FAF and AC oscillations for each frequency band at
- 186 various AC depths. For frequency bands between 1-12 Hz (i.e. delta to alpha), we observed
- 187 significantly higher than 0 z-iCoh estimates (FDR-corrected Wilcoxon signed rank tests,  $p_{corr} < 0.05$
- 188 for significance; log-converted p values are shown in **Fig. 3D**, bottom), which became gradually
- 189 lower towards higher frequencies (note also the decay in Fig. 3B). Beta or gamma z-iCoh
- 190 distributions were not significantly different than 0 at any cortical depth.
- 191 It was also apparent that coherence values were depth dependent in the AC, particularly in the delta
- band (where z-iCoh was strongest, see Fig. 3B, D). We tested the depth dependence of coherence by
- 193 comparing the distributions of z-iCoh for all pairs of channels, across all penetrations and frequency
- bands. Results are summarized in significance matrices depicted in **Fig. 3E**. Each cell (*i*, *j*) in a
- 195 matrix represents the log-converted, corrected p value (FDR Wilcoxon signed rank tests), obtained
- 196 after statistically comparing FAF-AC coherence using an AC channel at depth *i*, and another at depth
- 197 *j*. In the matrices, cells within red contour lines correspond to statistically significant p<sub>corr</sub> values (p<sub>corr</sub>
- 198 < 0.05). As shown in **Fig. 3E**, deep electrodes in the AC were significantly better synchronized with
- 199 the FAF, an effect only visible in the delta band.

# 200 2.3 FAF-AC synchronization during acoustic processing

201 To quantify synchronization during acoustic processing in the FAF-AC circuit, we calculated time-202 resolved iCoh values in response to four acoustic stimuli (a natural call, syllabic trains of 5.28 and 97 203 Hz, and a syllabic train with Poisson temporal structure; see above). Population average coherograms 204 (i.e. time-frequency representations of iCoh values) are shown in Fig. 4A-D for all stimuli tested, and 205 at representative depths in the AC (50, 450 and 700 µm). The most conspicuous pattern across 206 stimuli was the appearance of low-gamma coherence (typically in the range of 25-45 Hz), which was 207 associated with stimulus onset (at time 0) and apparently independent of auditory cortical depth. 208 Gamma synchrony was auditory-evoked, a notion strengthened when considering responses to the 209 5.28 Hz syllabic train (Fig. 4B), where the evoked coherence tracked individual syllable 210 presentations. Remarkably, we observed little evidence for an increase of low-frequency synchrony 211 when compared to spontaneous activity (compare heatmaps in **Fig. 4** with **Fig. 3C**). That is, even 212 though low-frequency coherence was present before sound presentation (in line with our results using 213 spontaneous LFPs), it did not change visibly after stimulus onset. To illustrate the occurrence of low 214 and high frequency coherence, Figure 4E-H depict single-trial LFPs from a representative pair of

215 simultaneous penetrations in the AC (450 µm depth) and FAF. Raw LFPs are shown in their 216 broadband form (0.1-300 Hz), and filtered in frequency ranges of 4-12 Hz and 25-45 Hz (i.e. low and 217 high frequencies oscillations). The range 4-12 Hz was chosen for low-frequency activity because 218 spectral parameters at lower frequencies could not be reliably estimated with the window size chosen 219 for time-resolved coherence analyses (200 ms; see Methods). Note that the occurrence of 220 synchronized waves in the AC and FAF is clear after stimulus onset (0-ms mark). 221 The defined low- and high-frequency ranges were then used to quantify changes from spontaneous to 222 stimulus-driven coherence in the FAF-AC network. A systematic, time-resolved analysis of low-223 frequency synchrony revealed that, across stimuli and auditory cortical depths, there was little change 224 (calculated as percentage increase from spontaneous to sound-driven activity: [iCoh<sub>stim</sub>-iCoh<sub>spont</sub>]/ 225 iCoh<sub>spont</sub>\* 100) in coherence between both structures (Fig. 5A, D, G, J, top heatmaps). The data 226 showed that low frequency synchrony preceded stimulus presentation in deep layers, seldom reached 227 50% increase from spontaneous across stimuli (black contour lines in Fig. 5A, D, G, J), and when it 228 did, it typically happened at middle AC depths. We did not observe statistical evidence showing 229 significant increase of low frequency coherence at stimulus onset (i.e. first 100 ms after sound 230 presentation; Fig. 5B, E, H, K, black traces; FDR-corrected Wilcoxon signed rank tests, significance

231 when  $p_{corr} < 0.05$ ).

232 Figure 5C, F, I, L (top heatmaps) illustrate effect size calculations (r; see Methods) for the low-233 frequency coherence increase in a time resolved manner across channels. In the heatmaps, only time 234 points where the increase was significantly different from 0 (uncorrected Wilcoxon singed-rank test, 235 p < 0.05) are shown. From this analysis the following was evident: (i) the pattern of significance was 236 inconsistent across stimuli for low frequency coherence; and (ii) effect sizes were typically small, 237 with areas of medium effect size (those within grey contour lines) appearing also with an inconsistent 238 pattern across sounds. Large effect sizes (within red contour lines) were overall only observed in 239 small clusters, lacking consistency throughout the stimulus set. The boundaries between small, 240 medium and large effect sizes were defined as follows: r < 0.3, small;  $0.3 \le r < 0.5$ , medium; r > 0.5, 241 large (Fritz et al., 2012). Altogether, these results corroborate a lack of reliable increase in low-242 frequency coherence between FAF and AC during passive listening, compared to spontaneous 243 activity.

High frequency FAF-AC coherence was considerably more sensitive to acoustic stimulation. As
expected from the data depicted in Fig. 4, we observed a strong increase of low-gamma interareal

246 synchronization associated to the stimulus onset (Fig. 5A, D, G, J, bottom heatmaps). Stimulus-247 evoked gamma synchrony was typically higher than 50% (reaching values as high as 80%) and 248 tracked the syllable presentations of the 5.28 Hz syllabic train (Fig. 5D, bottom). Indeed, the increase 249 of low-gamma synchrony at stimulus onset was significantly above 0 for all stimuli (Fig. 5B, E, H, 250 **K**, grey traces; same statistical analysis as for low-frequencies;  $p_{corr} < 0.05$ ), and more reliably so for 251 channels located in input layers of the AC (in this context, electrodes at depths of 250-350 µm; layers 252 III-IV in C. perspicillata's AC span depths of 200-450 µm, see (Garcia-Rosales et al., 2019)). For the 253 5.28 Hz and the Poisson syllabic trains, the onset-related increase in low-gamma coherence occurred 254 essentially along all AC depths studied. In terms of effect size (Fig. 5C, F, I, L; bottom heatmaps), 255 we observed large effects of increased low-gamma coherence at stimulation onset across stimuli (less 256 clearly in the case of the natural vocalization), with sustained, seemingly periodic increases along the 257 time-course of the 5.28 Hz syllabic sequence. Taken together, coherence analysis results indicate that 258 acoustic stimulation elicits auditory-evoked low-gamma synchronization between the frontal auditory 259 field and the auditory cortex.

#### 260 2.4 Gamma-band activity in FAF and AC

261 Previous studies showed the occurrence of gamma-band activity in the AC of primates, bats, and rats 262 (Brosch et al., 2002; Medvedev and Kanwal, 2008; Vianney-Rodrigues et al., 2011). These studies 263 reported auditory cortical gamma which was not time-locked to the onset of a stimulus, and that 264 appeared even hundreds of milliseconds after sound presentation. Given the nature of the coherence 265 analyses performed here, it is possible that the presence of non-locked gamma oscillations could have 266 been overlooked in the FAF and the AC of C. perspicillata. To explore the occurrence of these 267 rhythms in our dataset, we focused on the onset period of the 5.28 Hz syllable train as it was the 268 stimulus that permitted the analysis of an onset window without the influence of subsequent sounds 269 (after the first syllable presentation), for a sufficiently long time-lapse of at least 180 ms. The time 270 period around the first syllable presentation in the 5.28 Hz train was subdivided into three segments: 271 (1) a window of 90 ms spanning times before stimulus onset (pre); (2) a window of 90 ms starting at 272 stimulus onset (onset); (3) a window of 90 ms starting 90 ms past stimulus onset (late); and (4) a 273 window of 180 ms starting at stimulus onset (*full*). The span of these segments is illustrated in **Fig.** 274 6A together with representative LFP traces from a penetration pair. The segments were chosen in 275 order to contrast LFP power at different frequency bands (low frequencies, 0-15 Hz; low-gamma, 25-

45 Hz; high-gamma 45-80 Hz; and broad gamma 25-80 Hz) with spontaneous LFP power before
stimulus presentation (*pre* window).

278 There was a consistent increase of onset-related gamma activity in FAF and AC, particularly during 279 the *onset* period, which was potentially linked to an evoked activation in cortex as it was associated 280 with a broadband increase in LFP power (see Fig. 6B, green traces, and Fig. S2A). To uncover the 281 presence of gamma within later time periods in our data, the power of different frequency bands in 282 the *late* period was statistically compared (Wilcoxon signed rank test, significance threshold at p =283 0.01; see Methods) with the power in the *pre* period on a trial-by-trial basis, per penetration (Fig. 284 6C). The percentage of penetrations in FAF where there was a power increase in low frequencies was 285 of 8%, reaching between 14-24% in the AC. We also observed a relatively small number of 286 penetrations (<10%) either in AC or FAF in which there was a significant power increase in gamma 287 for the *late* period as compared to the *pre* segment. We further calculated the time course of the 288 percentage of penetrations showing significant power increase at times surrounding the stimulus 289 presentation for both FAF and AC, at various LFP frequencies (Fig. 6D; see Methods). As expected 290 from the data shown in **Fig. 6B-C** and **Fig. S2A**, up to about 50 ms after stimulus onset there was a 291 high percentage of penetrations (ca. 80%; cf. with Fig. S2A) where the power in gamma increased 292 significantly in either structure. This number was relatively low (< 20%) for times beyond 50-60 ms 293 after stimulus onset.

294 The FAF-AC circuit exhibited increased auditory-evoked gamma band coherence, related to the onset 295 of acoustic stimulation (Figs. 4, 5). We tested to what extent we could disentangle gamma activity in 296 our data from a non-specific broadband response, by means of previously used approach which relies 297 on comparing the relative power distributions of gamma and low frequency LFPs (Medvedev and 298 Kanwal, 2008). In this case, evidence for the gamma-band activity being a different component from 299 the broadband evoked-related potentials relies on the statistical independence between low- and high-300 frequency power in the LFPs. For this analysis the *full* window was used (see **Fig. 6A**). The power 301 distributions of gamma (either 25-45 or 45-60 Hz) and low frequencies typically did not differ in our 302 dataset (see Fig. 7A, E for a representative penetration). A systematic population analysis was 303 performed to quantify the percentage of penetrations in the data for which there was evidence of 304 statistical independence between the power of gamma and that of low-frequency potentials. As 305 depicted in Fig. 7B, F, power distributions of gamma (in the 25-45 and 45-60 Hz ranges) and low 306 frequencies (0-15 Hz) were significantly different from each other (2-sample Kolmogorov Smirnov

tests, significance when p < 0.01) only in a small proportion (< 15%) of the total amount of

308 penetrations, either in FAF or AC. This could suggest that gamma-band activity in these frequency

309 ranges cannot be readily disentangled from a broadband power increase related to an onset response,

- 310 assuming that if they were separable processes (i.e. gamma and low frequency activities) their power
- 311 distributions would differ significantly (Medvedev and Kanwal, 2008).

312 We reasoned, however, that a lack of significant differences between the distributions does not

- 313 necessarily imply that the relative powers of gamma and low-frequencies are well correlated when
- 314 considering trial specific information. Strong correlations would occur if low and high frequency
- 315 relative powers were tightly determined by the strength of the broadband activation. Therefore, given
- a weak correlation, it could be argued that the dynamics of low- and high-frequency (gamma) LFPs
- 317 might be more complex than an unspecific power increase related to the evoked response. We
- 318 observed poor correlations, across penetrations in FAF and AC, between low-frequency and gamma-
- 319 band relative power on a trial-by-trial basis (see, for example, Fig. 7C, G). The distribution of
- 320 correlation coefficients for the population data is depicted in Fig. 7D, H. Overall, correlation
- 321 coefficients were low, having a median in the FAF of 0.22 (25<sup>th</sup> and 75<sup>th</sup> percentiles: 0.12 and 0.35)
- for the 25-45 Hz gamma range, and of  $0.07 (25^{\text{th}} \text{ and } 75^{\text{th}} \text{ percentiles: } -0.05 \text{ and } 0.23)$  for the 45-60
- Hz band. In the AC, the median across channels was of 0.17 (25<sup>th</sup> and 75<sup>th</sup> percentiles: 0.03 and 0.31)
- for the 25-45 Hz gamma, and of 0.08 (25<sup>th</sup> and 75<sup>th</sup> percentiles: -0.04 and 0.1) for the band of 45-60
- Hz. Typically, no more than 20-25% of the penetrations in AC and FAF showed a significant
- 326 correlation (significance when p < 0.01) between relative power at low-frequencies and gamma (25-
- 327 45 Hz, median 20% of sites; 45-60 Hz, median 8% of sites; see **Fig. S2B**).
- 328 We also quantified how the overall energy of early activation correlates with the gamma coherence
- 329 increase in the FAF-AC circuit. Gamma-band coherence increase (compared to spontaneous activity,
- 330 see **Fig. 5**) was poorly correlated with evoked potential energy for all AC channels (median across
- channels: 0.15; 25<sup>th</sup> and 75<sup>th</sup> percentiles: 0.08 and 0.22), as illustrated in **Fig. 7I** for representative
- depths in AC, and in **Supplementary Fig. S3** for all depths. Although gamma-band activity is not
- 333 straightforwardly separable from a broadband activation pattern, the data shown in **Fig. S3** and the
- poor trial-by-trial correlation between gamma and low-frequency powers (see above) suggest the
- 335 possibility of interesting gamma-band dynamics in *C. perspicillata*'s FAF and AC.

# 336 2.5 Mutual information in FAF and AC spiking

337 We investigated spike-spike interactions in the FAF-AC network within an information theoretical 338 framework (Shannon, 2001). Mutual information (MI or "information" throughout the text) between 339 the stimuli and neuronal responses allows to quantify the theoretical ability of a neuron (or a set 340 thereof) to represent the acoustic input on a single trial basis (see Methods). MI captures all non-341 linear dependencies of any statistical order in the data, and its quantification depends on the neural 342 code being considered (Kayser et al., 2009). Here, we aimed to determine the coding abilities in AC, 343 FAF, and a joint response from both structures based on a spike rate code (I<sub>rate</sub> for single units, I<sub>ioint</sub> 344 considering responses from AC and FAF together; (Kayser et al., 2009; Garcia-Rosales et al., 345 2018a)). A rate code was considered so that our results could be comparable with previous data 346 obtained from C. perspicillata's AC (Garcia-Rosales et al., 2018a).

347 Overall, we relied on an information theoretic approach because we observed that representations in

AC and FAF were quite different and on occasions, at least in appearance, complementary. For

349 example, Fig. 6A depicts spiking from two simultaneously recorded FAF and AC units in response to

350 the natural stimulus. Note how the firing rate of the FAF unit increases as the stimulus progresses,

351 whereas the AC unit is time-locked to slow temporal modulations in the stimulus and does not

352 respond like its FAF counterpart. Information theory would allow to measure possible interactions

353 between these responses in a quantitative manner.

**Figure 6B** shows a schematic of the rate code used to quantify MI for single units and joint

355 responses. For subsequent analyses and in order to guarantee that all units considered were auditory-

responsive (both in FAF and AC), we used only responses that provided at least 0.1 bit/s of

information (with a window size of 4 ms this corresponds to  $4 \times 10^{-3}$  bits; see Methods). The number

358 of units used to quantify I<sub>rate</sub>, per channel (including the FAF electrode, and across the AC linear

probe) is depicted in **Fig. 6C**, left. Note that the number of penetrations is equivalent to the number

of units per channel in this case:  $n \ge 13$  in FAF, and  $n \ge 29$  in AC. For paired responses (FAF-AC)

361 the number of units considered was less because the inclusion criterion ( $I_{rate} \ge 0.1$  bit/s) had to be

fulfilled by units in AC and FAF simultaneously ( $n \ge 8$  pairs; Fig. 6C, right).

363 Population values of I<sub>rate</sub> for the FAF and the AC at different depths are depicted in **Fig. 6D**, for each

364 stimulus. Two main conclusions can be drafted from this figure. (i) In the AC, the highest

365 information about the stimuli was found at depths between 200-650 µm. (ii) Neurons in the FAF

366 were on average less informative than AC ones, but were well above the limit set in the inclusion

367 criterion across stimuli (*nat*:  $0.56 \pm 0.09$  bits/s, 5.28 Hz train:  $0.45 \pm 0.06$  bits/s, 97 Hz train:  $0.48 \pm$ 

368 0.04 bits/s, Poisson train:  $0.42 \pm 0.05$  bits/s; given as mean  $\pm$  s.e.m). We also quantified the

- 369 information provided by paired neuronal responses from both regions (I<sub>joint</sub>), which is illustrated as
- black traces in **Fig. 6E**. This figure also shows a direct comparison between I<sub>joint</sub> and the I<sub>rate</sub> of FAF
- 371 and AC neurons that conform each pair (I<sub>rate\_pair\_FAF</sub> and I<sub>rate\_pair\_AC</sub>, respectively). The contribution of
- 372 I<sub>rate</sub> from the FAF was significantly smaller than the contribution of I<sub>rate</sub> from the AC (FDR-corrected
- 373 Wilcoxon signed rank tests,  $p_{corr} < 0.05$ ; corrected p values of all comparisons are given in **Fig. S4**).
- 374 Although  $I_{rate}$  in the FAF was always > 0.1 bit/s, we did not observe  $I_{joint}$  to be significantly higher
- than the I<sub>rate</sub> from the AC (I<sub>rate\_AC</sub>; orange traces in the **Fig. 6E**), across electrodes and stimuli tested
- 376 (Fig. S7). We did observe  $I_{joint}$  to be well correlated with the sum of  $I_{rate}$  calculated using the
- information of FAF and AC spiking ( $I_{sum} = I_{rate_pair_FAF} + I_{rate_pair_AC}$ ; Figs. S5-S9).

When considering the interactions of spiking across structures in terms of the codes defined in this 378 379 study, the linear relationship between I<sub>sum</sub> and I<sub>joint</sub>, evident in Figs. S5-S9, would support the notion 380 of "independence" in the information provided by the spiking in each structure. Independence arises 381 theoretically when  $I_{joint} = I_{sum}$ , and implies that each structure represents different aspects of the 382 sensory stimulus. However, the results illustrated in **Fig. 6E** suggest that independence cannot be 383 inferred from the data with certainty. Mathematically, independence would require I<sub>joint</sub> to be higher 384 than both Irate in AC and FAF simultaneously, which was not fulfilled at a population level: Ijoint was 385 not significantly different than I<sub>rate</sub> obtained from AC units (see Fig. S4). Calculating information 386 estimates using time windows of up to 12 ms yielded comparable results, indicating that the temporal 387 resolution of the codes had little effect in the described outcomes (data not shown). Overall, our 388 quantification of stimulus-related spiking information, occurring simultaneously in the AC and FAF, 389 suggests different sound coding strategies in these two structures. These observations are further 390 addressed in the Discussion section.

#### **391 3 Discussion**

In this study we investigated the functional connectivity between frontal and auditory cortical regions of the bat *Carollia perspicillata*. Specifically, we examined the coupling dynamics of the FAF, a frontal area which receives auditory afferents from cortical and subcortical structures, and the primary AC. Functional connectivity was assessed during spontaneous activity and during the processing of natural and artificially generated acoustic sequences. Our main results are: i) LFPs recorded simultaneously in both regions suggest that the FAF receives faster auditory inputs relative to the AC, yet these inputs do not necessarily elicit faster spiking in the FAF; ii) during spontaneous

399 activity, the FAF-AC network is coupled in low frequencies (up to 12 Hz), with stronger coherence 400 values at deep layers of the AC; iii) while acoustic stimulation does not considerably alter the default 401 low-frequency coupling, auditory-evoked gamma-band synchronization in the FAF-AC circuit 402 emerges upon sound presentation; iv) considering a spiking rate code, the FAF is less informative 403 than the AC about the acoustic stimuli, while a joint code using simultaneous spiking from both 404 regions suggests that FAF and AC engage in distinct coding dynamics. Altogether, our data shed 405 light onto how distant brain areas in the mammalian brain engage in sound representation. The results 406 of this paper are summarized in Fig. 9.

#### 407 **3.1** Auditory afferents into the FAF

408 Stimulus-related LFPs recorded simultaneously from FAF and primary AC, at various depths in the 409 latter structure, indicate the presence of fast synaptic inputs into the frontal region that precede those 410 arriving into the AC even at input layers. The work of Kobler, Casseday, and colleagues in the late 411 1980s (Kobler et al., 1987; Casseday et al., 1989), showed that the FAF receives auditory afferents 412 via a non-canonical pathway that bypasses major auditory centres in the midbrain, including the 413 inferior colliculus (IC). In this pathway, acoustic information from neurons in the cochlear nucleus is 414 sequentially relayed to the anterolateral olivary complex, the suprageniculate nucleus of the thalamus 415 (SGN), and from there into the FAF. Thus, although auditory inputs reach the frontal region also 416 through the AC, acoustic information may reach the frontal field, directly from the cochlea, in as few 417 as four synapses (Kobler et al., 1987). Synaptic currents related to inputs from the SGN into the FAF 418 could lead to changes in LFPs (Buzsaki et al., 2012) faster than their counterparts at input layers of the AC, which are predominantly driven by thalamocortical synapses originating in the ventral region 419 420 of the medial geniculate body. Therefore, a rapid, non-canonical pathway into FAF accounts for our 421 observations regarding the temporal relation between LFPs in both regions studied.

422 The results described in this manuscript indicate that, although sound-related LFPs from the FAF

423 lead those from the AC, the neuronal spiking latencies are shorter in AC. In other words, our data

424 indicate that faster inputs in FAF are not sufficient to elicit faster spiking in most cases.

425 Electrophysiological studies in frontal auditory areas have shown that neuronal responses are usually

426 of relatively large latencies (although short latencies are also to be found), sparse and of high

427 variability (Newman and Lindsley, 1976; Eiermann and Esser, 2000; Kanwal et al., 2000; Plakke and

428 Romanski, 2014). Recent data from *C. perspicillata*'s FAF highlighted the possibility that the

429 sparseness in the response properties of frontal neurons could be explained by slow, low-threshold,

and long-lasting synaptic dynamics, at least considering projections from the AC (Lopez-Jury et al.,
2019). These slow, long-lasting synaptic dynamics could support sensory integration, by conditioning
FAF neurons to spike after accumulating synaptic inputs over time, and/or to integrate multiple
synaptic inputs originating from different sensory modalities. Certainly, cross-modal sensory

434 integration occurs in the frontal cortex (Fuster et al., 2000; Romanski, 2007; Hwang and Romanski,

435 2015), whereas integration over relatively long timescales appears to be a feature of higher-order

436 cortical areas in general (Runyan et al., 2017).

437 Fast acoustic afferents, even without eliciting reliable stimulus-evoked spiking, imply nonetheless 438 that auditory information is already present in the FAF before it receives inputs from the AC. Within 439 the predictive coding framework (Arnal and Giraud, 2012; Friston, 2018), it is then possible to 440 hypothesize that the frontal field may be a region where prediction errors could also be generated. 441 That is, the rapid non-lemniscal pathway could relay faithful information into FAF about the auditory 442 stimuli, which can in turn be compared with information received from primary AC and the 443 thalamus. Alternatively, a further proposition would be that prediction errors, relayed "upwards" 444 along the auditory hierarchy (Carbajal and Malmierca, 2018), and generated in the SGN and the AC, 445 are integrated in the FAF. The result of such integration could in turn be used to update the 446 "expectations" of the system. In rodents, prediction error signals appear all along the auditory 447 pathway, but occur more strongly in higher-order structures (Parras et al., 2017), while prediction 448 error signals related to sounds have also been described in the frontal cortex of rats (Imada et al., 449 2012) and humans (Durschmid et al., 2016; Durschmid et al., 2018). The involvement of the FAF in

450 predictive coding could be thoroughly tested in future experimental work.

451 A number of FAF neurons project directly to the superior colliculus (SC), a structure related to motor 452 control of head and pinna movement in bats (Kobler et al., 1987). The presence of early acoustic 453 information in FAF, the fact that frontal neurons in *C. perspicillata* show preference for naturalistic 454 echolocation acoustic stimuli (high-frequency pulses used to navigate by bats; (Eiermann and Esser, 455 2000)), and the existence of projections from FAF into motor-related structures such as the SC, 456 suggest that the FAF plays an important role in coordinating auditory-guided behaviour. This would 457 be in line with proposed roles of prefrontal cortex in motor control (Risterucci et al., 2003), including 458 volitional motor (vocal) production after acoustic stimulation (Hage and Nieder, 2015). However, 459 whether neuronal activity in the FAF mediates motor outputs is still to be tested in depth.

#### 460 **3.2** FAF-AC synchronization during spontaneous activity

461 We examined the default (i.e. in the absence of external stimulation) functional connectivity in the 462 FAF-AC network in terms of oscillatory coherence (Fig. 3). The data presented in this manuscript 463 show that simultaneously recorded LFPs from both structures are phase-synchronized in low-464 frequencies of the spectrum (1-12 Hz), although more strongly so in the delta band (1-4 Hz). 465 Empirical evidence points towards a role of oscillatory coherence in the coupling of distant brain 466 regions, a view that is summarized in proposed theoretical mechanisms such as the communication-467 through-coherence framework (Fries, 2015). Several studies have pinpointed an involvement of low-468 frequency synchronization in the functional coupling between the frontal cortex and a variety of brain 469 structures. Coherent oscillations between distant cortical areas (including the frontal cortex) in the 470 low-frequency range correlate with working memory (Daume et al., 2017), fear memory 471 consolidation (Popa et al., 2010), attentional selection (Womelsdorf and Everling, 2015), and long-472 term fear recall (Cambiaghi et al., 2016; Karalis et al., 2016). In the auditory domain, top-down 473 control exerted from frontal cortical areas, through low-frequency oscillatory activity, increases 474 coupling to speech signals in the human AC (Park et al., 2015). Our results indicate that, in the bat 475 brain, frontal areas that participate in audition are functionally interconnected by means of low-476 frequency LFPs with primary auditory cortex. Critically, such coupling does not require external 477 input, which hints towards the presence of a default synchrony in the fronto-auditory cortical 478 circuitry. The latter could constitute a functional basis for high-order, interareal auditory processing 479 in the mammalian brain.

480 Because recordings in the AC were performed with a laminar probe, we were able to study the 481 laminar dependence of FAF-AC synchrony in AC. During spontaneous activity, low-frequency 482 coherence was strongest in deep layers (depth > 700  $\mu$ m), but the strength of coherence was only 483 affected by depth in the delta band. Interestingly, a recent study revealed that, also during 484 spontaneous activity, spike-LFP synchronization was strongest in deep layers of the AC (Garcia-485 Rosales et al., 2019). Such spike-LFP coupling was associated to the presence of discrete 486 spontaneous states of increased spiking rate (UP-states) in laminae V and VI. The origins of deep 487 layer UP-states are unclear, but it has been hypothesized that they could be driven by higher-order 488 structures (Sakata and Harris, 2009). In the current study, we observed a putative higher-order 489 auditory structure (the FAF in the frontal lobe) synchronized via a low-frequency oscillatory channel 490 with deep layers of a primary sensory area (the AC). From the phase correlation of delta-band LFPs 491 in the FAF-AC network, it is possible to speculate that delta oscillations in FAF could modulate UP-

492 states in AC. However, we note that causality cannot be inferred from our current dataset and needs493 to be addressed thoroughly with further experimental approaches.

In all, the "default" coupling in the FAF-AC circuit is supported by the presence of anatomical
connections between frontal and auditory cortices (Kobler et al., 1987)). Although it has to be
properly addressed, we propose that homologous frontal regions tasked with audition in other species
may be functionally interconnected with AC in a similar manner. This possibility is still unexplored,
yet addressing this question might be crucial for unravelling the mechanisms of high-order auditory
processing, cognition, and behaviour based on audition.

# 500 **3.3** Functional coupling in the FAF-AC network during acoustic processing

501 Functional coupling between FAF and AC was also addressed in the context of acoustic processing 502 (Figs. 4, 5). Animals were exposed to four distinct acoustic streams, including a conspecific distress 503 vocalization, and three "trains" constructed by repeating a distress syllable at distinct rates. 504 Independently of the stimulus used, we observed little change in the low-frequency phase synchrony 505 of the FAF-AC network, as compared to spontaneous activity. This is a puzzling result which, 506 assuming that low-frequency oscillatory coupling in the network is useful for auditory perception, 507 could in principle be explained largely by our experimental design. Although the animals were awake 508 during the experiments, they listened to the sequences passively: i.e. they were not expected to 509 behave in response to the stimulus, and other variables (e.g. attentional processes) were not 510 modulated according to a controlled experimental approach. Thus, statistically negligible and 511 unreliable changes in the low-frequency dynamics of FAF-AC connectivity could be explained by 512 the fact that the passive listening of acoustic streams is not sufficient to alter the default functional 513 coupling in the network. Whether attention (a top-down process) or behavioural planning could 514 modify the neuronal connectivity in the circuit by either enhancing it or decreasing it, as compared to 515 spontaneous activity, needs to be tested in further research.

Experimental evidence suggests that oscillatory activity in the gamma range is crucial for neuronal
computations, including sensory processing and cognitive mechanisms (Fries, 2009). Previous
studies have shown the presence of gamma-band activity in primary auditory cortex of rats (VianneyRodrigues et al., 2011), monkeys (Brosch et al., 2002) and bats (Medvedev and Kanwal, 2008)
during passive listening. These studies reported the presence of gamma oscillations at even later time

520 during passive listening. These studies reported the presence of gamma oscillations at even later time

521 periods (150-300 ms) in what could be considered "induced" (as opposed to "evoked") activity, not-

522 locked to the auditory stimulus. Our results show that, past 90 ms and up to 180 ms after stimulus 523 presentation, such late, non-locked gamma oscillations are relatively scarce (~5% of out of 50 524 penetrations). These results should not be taken as evidence for the lack of late gamma oscillations in 525 the AC or FAF of C. perspicillata, because later periods (190-300 ms after sound presentation) in 526 which oscillatory activity may have occurred were not analysed due to the nature of the stimulus 527 (note that the second syllable presentation of the 5.28 Hz occurs already at ~189.4 ms). Further 528 research can be aimed at detecting gamma activity at later time points in the cortex of C. 529 perspicillata.

530 In humans, sources of auditory-evoked gamma-band activity (aeGBA) can be found both in primary

auditory and frontal (anterior cingulate cortex, ACC) cortices (Mulert et al., 2007; Polomac et al.,

532 2015). Frontal aeGBA is modulated by attention and correlates with performance in auditory

detection tasks (Debener et al., 2003; Gurtubay et al., 2004). Moreover, aeGBA in the frontal lobe

and gamma-band synchronization between frontal and auditory cortical regions correlate too with

task difficulty (Mulert et al., 2007; Polomac et al., 2015), suggesting a role of gamma-band

536 coherence in a fronto-auditory cortical circuit for cognitive control in audition. In fact, disorders of

537 the central nervous system such as schizophrenia are marked by a dysregulation of aeGBA in frontal

regions (Cho et al., 2006; Leicht et al., 2010; Curic et al., 2019), further supporting the importance of

539 gamma-band activity for cognition.

540 Our data indicate that low-gamma (25-45 Hz) coherence in the FAF-AC circuit significantly

541 increases with sound presentation, independently of the stimulus considered. This supports a role of

542 gamma synchronization between frontal and auditory cortices for auditory processing, although the

543 functional significance of aeGBA and its coherence across cortical areas should be considered with

544 care. Coherent activity does not imply directly that there exists effective communication between two

545 given regions. Moreover, the nature of the gamma activity is also to be examined cautiously. Here we

546 attempted to disentangle gamma oscillations from a frequency unspecific power surge related to

547 auditory evoked responses (Fig. 7), which would in principle explain gamma coherence between

548 FAF and AC. Based on a method proposed by Medvedev and Kanwal (2008), we did not observe

549 statistical evidence supporting that the distributions of low-frequency and gamma-band LFP power

550 were significantly different from one another across penetrations in FAF and AC (**Fig. 7B, F**).

551 However, we did observe, across multiple penetrations, a lack of trial-by-trial correlation between the

552 LFP power in the abovementioned frequency bands (Fig. 7C, D, G, H, and Fig. S2B). In addition,

553 our data also showed a very weak correlation between the energy of the evoked response in the LFP

and the gamma-band coherence increase (Fig. 7I and Fig. S3). While these results do not

555 conclusively demonstrate that gamma-band activity can be separated from a frequency unspecific

onset response, they hint towards the possibility of evoked gamma activity being an important

557 component for audition, as suggested by previous work (Brosch et al., 2002; Medvedev and Kanwal,

558 2008; Vianney-Rodrigues et al., 2011). The functional roles of onset-related gamma activity in the

559 FAF-AC circuit of *C. perspicillata* (and in the mammalian auditory system in general) should be

thoroughly addressed with dedicated experimental approaches in the future.

561 We would like to note that it remains speculative what might be potentially signalled by the FAF-AC 562 gamma synchronization reported in this study. Short onset-related coherence increase might convey 563 information about the presence of an acoustic stimulus, but not necessarily allow to elucidate the 564 stimulus' spectrotemporal features. In the AC of the bat *Pteronotus parnelii*, Medvedev and Kanwal 565 (2008) reported that the spectral properties of the gamma component of the response could be used to 566 differentiate among a battery of conspecific communication calls. In primary visual cortex, for 567 example, distinct characteristics of stimulus-induced gamma rhythms (e.g. peak frequency or 568 amplitude) encode for distinct properties of presented visual stimuli (e.g. contrast or orientation; 569 (Hermes et al., 2015; Murty et al., 2018)), although it has been argued that the variability of gamma 570 based on stimulus properties may constrain the utility of the rhythm for complex integrative 571 computations, at least in early visual cortex (Henrie and Shapley, 2005; Ray and Maunsell, 2010; 572 Bartoli et al., 2019). Our stimulus set is not ideal to determine in an unbiased manner whether the 573 nature of FAF-AC gamma synchronization changes given the spectrotemporal characteristics of the 574 stimuli, in particular because LFPs synchronize to a stimulus' temporal structure also in the gamma 575 range (see (Hechavarria et al., 2016b; Garcia-Rosales et al., 2018a). The latter could alter the spectral 576 patterns of coherence without necessarily meaning that the nature of the underlying coherence is 577 changing, which is an artefact that needs to be controlled for. The careful and systematic variation of 578 acoustic properties of sounds could be used in further research to explore in full the patterns of FAF-579 AC gamma-band coherence and its role for audition.

# 580 **3.4** Mutual information in the FAF-AC circuit

581 Quantifying the amount of information provided by FAF and AC about the acoustic stimuli revealed 582 that the former structure was, in comparison, significantly less informative than the latter. In addition, 583 when considering responses from frontal or auditory cortices in a joint rate code, it was not possible

584 to determine a clear population trend towards independence, redundancy or synergy between the 585 spiking activities of both structures, from an information theoretic perspective. We propose two 586 candidate explanations for our results, which need not be mutually exclusive. First, it is possible that 587 the way in which FAF encodes incoming auditory stimuli is not sufficiently well-captured by means 588 of a code based on spiking rate, and therefore the true encoding capabilities could be underestimated 589 by assuming such scheme (see (Masquelier, 2013; Insanally et al., 2019)). Second, we note that the 590 fact that the FAF conveys, in comparison, significantly less information than the AC, is an expected 591 result as the AC is a structure specialized for auditory computations. As part of the frontal cortex, it is 592 plausible that the FAF encodes for other variables that go beyond acoustic features (e.g. ethological 593 relevance of the sound, multimodal sensory information, etc.). In that case, FAF units would yield 594 low  $I_{rate}$  values when attempting to quantify their abilities to encode a sound based on relatively 595 simple methodological approaches, which rely solely on acoustic processing. The former allows to 596 hypothesize that FAF and AC might engage in distinct, non-overlapping coding strategies.

#### 597 4 Methods

#### 598 4.1 Animal preparation and surgical procedures

All experimental procedures were performed in compliance with current German and European
regulations on animal experimentation. Experiments were approved by the Regierungspräsidium
Darmstadt (experimental permit #FU-1126). The study was performed on 5 adult bats of the species *Carollia perspicillata* (all males). Animals were obtained from a colony in the Institute for Cell
Biology and Neuroscience, Goethe University, Frankfurt am Main. Bats used for experiments were
kept separately from the main colony.

605 Before undergoing surgical procedures, bats were anaesthetized with a mixture of ketamine (10 mg\*kg<sup>-1</sup>, Ketavet, Pfizer) and xylazine (38 mg\*kg<sup>-1</sup>, Rompun, Bayer). For surgery and any 606 607 subsequent handling of the wounds, local anaesthesia (ropivacaine hydrochloride, 2 mg/ml, Fresenius 608 Kabi, Germany) was applied subcutaneously in the scalp area. A rostro-caudal midline incision was 609 made in the scalp, after which skin and muscle tissues were removed carefully in order to expose the 610 skull. A sufficiently large area of the bone was also exposed to make possible the attachment of a 611 custom-made metal rode (1 cm length, 0.1 cm diameter), used during recordings to fixate the 612 animal's head. The rod was attached with dental cement (Paladur, Heraeus Kulzer GmbH, Germany). 613 Animals were given at least one full day of recovery after surgery, and before experiments were

614 performed upon them. The AC and the FAF were located based on well-establish landmarks such as

blood vessel patterns, and the sulcus anterior (see (Esser and Eiermann, 1999; Eiermann and Esser,

616 2000)). On the first day of recordings each cortical region was exposed by cutting a small hole (~ 1

 $mm^2$ ) in the skull with a scalpel blade.

618 Recordings, which lasted no more than 4 hours a day, were performed chronically on awake bats.

619 Water was given to the animals at a period of approximately 1-1.5 hours. Between recording

620 sessions, a bat was allowed to recover for at least a full day. Experiments for the day were halted if

621 the bat showed any sign of discomfort.

### 622 4.2 Electrophysiological recordings

Recordings were made inside an electrically isolated and sound-proofed chamber. Inside the chamber, bats were placed upon a custom-made holder which was kept at a constant temperature of 30 °C with a heating blanket (Harvard, Homeothermic blanket control unit). A speaker (NeoCD 1.0 Ribbon Tweeter; Fountek Electronics, China), located inside of the chamber 12 cm away from the animal's right ear (contralateral to the hemisphere were recordings were performed), was used for free-field stimulation. The speaker was calibrated using a <sup>1</sup>/<sub>4</sub>-inch microphone (Brüel & Kjær, model 4135, Denmark), which was connected to a custom-made amplifier.

630 Data were acquired from the bat's left AC as described in a previous study (Garcia-Rosales et al.,

631 2019). Neurophysiological data were recorded from the AC using 16-channel laminar electrodes

632 (Model A1x16, NeuroNexus, MI; impedance:  $0.5-3 \text{ M}\Omega$ ), with a channel separation of 50 µm. The

633 probe was carefully inserted into the brain perpendicular to the cortical surface using a piezo

634 manipulator (PM-101, Science 455 products GmbH, Hofheim, Germany) until the top-channel was

barely visible on the surface of the tissue. Thus, we were able to record from depths ranging 0-750

636 μm, reaching all layers in AC. Histological confirmation of the extent of the electrodes inside the

637 cortex are detailed elsewhere (Garcia-Rosales et al., 2019). Recordings were made in primary AC,

638 although we cannot discard the presence of columns from high frequency fields (Esser and Eiermann,

639 1999). The laminar probes were connected to a micro-amplifier (MPA 16, Multichannel Systems

- 640 MCS GmbH, Reutlingen, Germany), and acquisition was done via a portable multichannel system
- 641 with integrated analogue-to-digital converter (Multi Channel Systems MCS GmbH, model ME32
- 642 System, Germany) with a sampling frequency 20 kHz and a precision of 16 bits. Data acquisition was

on-line monitored and stored in a computer using MC\_Rack\_Software (Multi Channel Systems MCS
GmbH, Reutlingen, Germany; version 4.6.2).

645 For recordings in the FAF, a single carbon electrode (Carbostar-1, Kation scientific; Impedance at 1 646 kHz:  $0.4-1.2 \text{ M}\Omega$ ) was inserted into the frontal region of the left hemisphere and lowered to depths 647 of ~300-450 µm with the aid of a second piezo manipulator (same characteristics as the previous 648 one). The electrode was connected to a micro-amplifier which was also connected to the integrated 649 multichannel recording system as described above. It was possible to use the same hardware because 650 the integrated system accommodates up to 32 simultaneous channel recordings. Ground and 651 reference electrodes (silver wires) were inserted as to only touch the dura mater of non-auditory 652 regions of the bat's brain, preferentially located in occipital areas of the contralateral hemisphere.

#### 653 4.3 Acoustic stimulation

654 Acoustic stimulation was controlled from the recording computer using a custom-written Matlab 655 (version 7.9.0.529 (R2009b), MathWorks, Natick, MA) software. As acoustic stimuli we used a 656 natural distress call from C. perspicillata and three synthetic trains constructed from a single distress 657 syllable, repeated at different rates. Procedures for recording the natural sequence are described in a 658 previous study (Hechavarria et al., 2016a). The call is representative of C. perspicillata's vocal 659 repertoire, and has been used by us in previous studies addressing auditory processing at the level of 660 the AC (Hechavarria et al., 2016b; Garcia-Rosales et al., 2018a; Garcia-Rosales et al., 2019). Distress 661 calls of C. perspicillata exhibit two prominent, coexistent temporal modulations: the syllabic- and bout rhythmicities. Syllabic rates in C. perspicillata's distress utterances are in the range of > 30 Hz 662 663 (median, 71.4 Hz; iqr: 57.1 Hz), whereas bouts (groups of syllables emitted in close sequence) are 664 repeated with rates typically < 12 Hz (Hechavarria et al., 2016a). In the natural call used here, 665 syllables are repeated on average with a rate of 63.7 Hz (see (Garcia-Rosales et al., 2018a)), whereas 666 bouts are uttered with a rate of 4 Hz (i.e. 8 bouts in 1.96 s).

667 To emulate the temporal dynamics of the communication sequences, a stereotypical distress syllable

668 was used to construct artificial acoustic sequences. A first syllabic train had a repetition rate of 5.28

669 Hz, matching the slow temporal dynamics of *C. perspicillata*'s distress utterances. A second one,

670 with a repetition rate of 97 Hz, was used to simulate fast temporal dynamics in communication

671 streams. Finally, we constructed a syllabic train where syllables were repeated in a Poisson-like

672 manner, with an average rate of 70 Hz. This simulated fast-repetition rates without any periodicity

and without a slow temporal structure. The 5.28 and 97 Hz trains had a duration of 2 s, while the

674 Poisson train was 4 s long. The syllables had an intensity of 70 dB SPL (root-mean square), close to

675 the intensity of the natural call (see (Garcia-Rosales et al., 2018a)).

676 Sounds were digital-to-analogue converted by means of a sound card (M2Tech Hi-face DAC, 384

677 kHz, 32 bit) and amplified (Rotel power amplifier, model RB-1050) in order to be presented through

the speaker inside of the chamber. Prior to presentation the call and syllabic trains were down-

679 sampled to 192 kHz, and low-pass filtered (80 kHz cut-off). All sounds were pseudorandomly

presented 50 times each, with and inter-stimulus interval of 1 s. A period of 300 ms, and another of

500 ms, was appended at the beginning and the end of each sequence, respectively.

682 Prior to any acoustic stimulation, per penetration, electrophysiological data were acquired for a

683 period of 180 s. These data were used for coherence analyses during spontaneous activity.

#### 684 **4.4** Separation of spiking activity and local-field potentials

All analyses were performed offline with custom-written Matlab (version 8.6.0.267246 (R2015b))

686 scripts. Initially, the raw electrophysiological signal from each channel (all electrodes in AC and

687 FAF) was bandpass filtered (fourth-order Butterworth filter) in order to extract traces pertaining

688 spiking activity (300-3000 Hz cut-off frequencies) and LFPs (0.1-300 Hz cut-off frequencies). For

689 computational reasons, LFP data were down-sampled to 1 kHz and stored for subsequent analyses.

690 Spike detection and sorting from FAF and AC electrodes were performed using the SpyKING

691 CIRCUS toolbox (Yger et al., 2018). Spike detection threshold was set at 5 median absolute

deviations from the noise baseline, and spike sorting was done automatically by the SpyKING

693 CIRCUS algorithm based on the probe's geometry to avoid detecting the same templates in two

694 adjacent electrodes. Each template was assigned to the electrode where its amplitude was the

strongest. Per electrode (either in AC or FAF), we chose as representative spiking the template with
the highest spike count. Spiking responses relative to a single electrode are referred to as a "unit" in
the manuscript.

# 698 **4.5 Spike latency estimation**

Spike latency was defined as the time point in which a unit's spiking rate was statistically different
from the expected rate during spontaneous activity, based on a previous study (Chase and Young,
2007). In brief, the algorithm proposed by Chase and Young compares a unit's response to a stimulus

702 across several time windows, with the expected spiking rate under the assumption that the unit fires 703 spontaneously with Poisson statistics, given a certain rate. A unit's firing rate in the 250 ms silence 704 period before stimulus onset, across the 50 repetitions from all stimuli tested (a total 200 trials), was 705 considered its spontaneous spiking rate for the abovementioned assumption. The response of a unit to 706 a certain stimulus (i.e. spiking after stimulus onset) was pooled across trials. Taking this pooled 707 response, the probability of observing at least *n* spikes in a given window  $t_n$  (after stimulus onset), 708 assuming Poisson firing in the absence of acoustic inputs, can be defined as follows (Chase and 709 Young, 2007):

710 
$$P_{t_n}(\geq n) = 1 - \sum_{m=0}^{n-1} \frac{(N\lambda t_n)^m e^{-N\lambda t_n}}{m!}, \qquad [1]$$

711 where N is the number of repetitions of the given stimulus, and  $\lambda$  is the spontaneous firing rate. 712 Starting from stimulation onset, the probability that each elicited spike indicates a stronger than 713 chance deviation in rate from the firing rate estimated in the absence of stimulation (the 250 ms 714 window), is taken as the probability that the spontaneous firing rate would have produced that 715 particular spike as the last of *n* spikes in a window  $t_n$ . In this context,  $t_n$  is the width of the window 716 containing the *n* spikes observed so far. Hence, the time of the first spike for which the aforementioned probability is sufficiently low (here,  $P_{t_n} (\geq n) < 10^{-5}$ ) is considered as the unit's 717 718 latency. This method circumvents caveats regarding classical peak latency estimations using peri-719 stimulus time histograms or spike-density functions over time (Levakova et al., 2015).

# 720 **4.6 Interareal coherence analyses**

All coherence analyses were done using the Chronux toolbox (Bokil et al., 2010). As a metric of

722 interareal phase synchronization we used the imaginary part of the coherency ("iCoh" in the

manuscript; (Nolte et al., 2004)), both during spontaneous activity and acoustic processing.

Coherency is complex value that measures phase consistency between two time series, across several trials. The coherency between two signals *x* and *y*, at a certain frequency  $\omega$ , can be defined as follows

726 (Bastos and Schoffelen, 2015):

727 
$$coh_{xy}(\omega) = \frac{\frac{1}{n}\sum_{k=1}^{n}A_{x}(\omega,k)A_{y}(\omega,k)e^{i(\phi_{x}(\omega,k)-\phi_{y}(\omega,k))}}{\sqrt{(\frac{1}{n}\sum_{k=1}^{n}A_{x}^{2}(\omega,k))(\frac{1}{n}\sum_{k=1}^{n}A_{y}^{2}(\omega,k))}}, \quad [2]$$

728 where  $A_x(\omega, k)$  and  $A_y(\omega, k)$  are the amplitudes of signals x and y at frequency  $\omega$  and trial k, while 729  $coh_{xy}$  represents the coherency between both signals, and n is the number of trials per stimulus (n = 730 50). Coherency is a complex quantity, but its absolute value ranges from 0 to 1, indicating the 731 relative, normalized strength of phase synchronization between time series. Taking the imaginary 732 part of  $coh_{xy}$  is a straightforward manner to remove non phase-lagged interactions that could be 733 attributable to, for example, passive field spread or common referencing (Bastos and Schoffelen, 734 2015). In order to minimize further common influences related to the temporal structure of the 735 stimuli, we subtracted from each trial the mean (across all trials of a given stimulus) LFP of each 736 channel (Kikuchi et al., 2017), and calculated coherence using the de-meaned traces. Note that this 737 could affect low frequencies more than high frequencies, because the latter are more sensitive to 738 temporal jitter. Additionally, while the approach alleviates obtaining simply stimulus-evoked 739 coherence, it could also mask phase-locking that is time-locked to the stimulus but not entirely 740 attributable to acoustic temporal features.

741 From the 180 s trace of spontaneous activity recorded simultaneously from FAF and AC

742 penetrations, 50 chunks of 3 s length each were taken. The precise time at which chunks started was 743 chosen randomly in a way that the resulting sub-segments would still be non-overlapping. Each of 744 these paired chunks from FAF and AC were treated as a trial, and iCoh was estimated from all 50 of 745 them for the corresponding penetration (data shown in Fig. 3B). A surrogate calculation was 746 performed whereby the precise phase relationship between FAF and AC "trials" in spontaneous 747 activity was abolished. This was accomplished by pairing FAF chunks with AC chunks randomly. 748 The former affects the timing of the phase-relationships but maintains the overall power across 749 selected chunks. The pairing was performed randomly a sufficiently large number of times (250), and 750 iCoh was calculated at each repetition of the surrogate analysis. Thus, it was possible to obtain a 751 distribution of iCoh values that represented coherence estimates in the absence of consistent phase 752 relationships between AC and FAF during spontaneous activity. The iCoh calculated from the 753 original data was then related to the surrogate iCoh values by means of z-normalization (z-iCoh). At 754 a population level, lack of consistent phase coherence between FAF and AC at a certain frequency 755 would yield z-iCoh values close to 0.

756 Time-frequency resolved iCoh values were obtained by means of coherogram calculations (*cohgram* 757 function in Chronux; data depicted in Figs. 3C, 4 and 5). A time-resolved approach allowed us to 758 examine changes of coherence over time while animals listened to acoustic streams. Each

759 coherogram was constructed by calculating coherence in a sliding window of 200 ms length, which 760 was advanced in steps of 2 ms. Because of the spectral resolution due to window length, the coherence spectrum for frequencies below 4 Hz could not be estimated with precision. As with any 761 762 time-frequency resolved approach, there is a compromise between temporal and spectral resolutions, 763 which we empirically found to be best balanced with a 200 ms window. All power spectra in the 764 time-resolved analysis were obtained with the multitaper method (Percival and Walden, 1993), 765 available in the Chronux toolbox, using 3 tapers and a time-bandwidth (TW) product 2. 766 The frequency range of 4-12 Hz was used as a representative of low-frequencies in the spectrum, and 767 the 25-45 Hz band was considered as low-gamma. For comparing iCoh values during sound 768 presentation vs. iCoh values during spontaneous activity, we calculated time-resolved iCoh during 769 spontaneous activity using the same segments with which non-time resolved coherency was

calculated (shown in **Fig. 3C**). Because the length of the spontaneous segments (3 s) was not

precisely equal to the length of the stimuli (the Poisson process, for example, was 4 s long), we

collapsed the time-resolved spontaneous iCoh values in the temporal dimension (median across

- timepoints per frequency). Thus, it was possible estimate the percentage increase during sound
- processing in a time-resolved manner as follows:

775 
$$iCoh_{increase}(\omega, t) = \frac{iCoh_{stim}(\omega, t) - iCoh_{spont}(\omega)}{iCoh_{spont}(\omega)} * 100, \quad [3]$$

where  $iCoh_{stim}(\omega, t)$  is the iCoh value during stimulus presentation at frequency  $\omega$  ant time *t*, while *iCoh\_{spont}(\omega)* is the collapsed time-resolved iCoh during spontaneous activity at the same frequency. The percentage increases were narrowed to the frequency bands of interest (i.e. 4-12 and 25-45 Hz) by calculating the median iCoh increase in the corresponding frequency range over time (data depicted in **Fig. 5A, D, G, J**). For evaluating iCoh increase at stimulus onset, the median was calculated not only for the frequency range, but also across time in the first 100 ms after the sequence onset.

783 **4.7 LFP onset power analyses** 

To test to what extent coherent gamma oscillations in the FAF-AC network could be attributable to a broadband evoked response in the LFP, we explored the statistical dependence of gamma power on a trial by trial basis. Spectral properties were obtained using three different temporal windows (**Fig. 6**): pre (-110 ms – -20 ms relative to stimulus onset), *onset* (0 – 90 ms relative to stimulus onset), *late* 

788 (90 - 180 ms relative to stimulus onset), and *full* (0 - 180 ms relative to stimulus onset). All analyses 789 were performed using the 5.28 Hz syllabic train, thereby guaranteeing that responses to only one 790 syllable (i.e. the first syllable presented) were considered in the time windows of choice. All spectra 791 were calculated using the Chronux toolbox, with 2 tapers and a TW product of 2, on a trial-by-trial 792 basis. Power spectra were compared for every penetration, per trial, statistically probing changes in 793 the power of low-frequency (0-15 Hz) and gamma bands (25-45, 45-80, and 25-80 Hz) in the pre vs. 794 *onset* windows (**Fig. S2A**; Wilcoxon signed rank tests, significance when p < 0.01), as well as during the pre vs. late periods (Fig. 6C). A percentage of significant difference (ratio across 50 penetrations) 795 796 is depicted in the abovementioned figures. The time-frequency analyses shown in Fig. 6D was 797 obtained by evaluating significance differences, per penetration, at given time windows (90 ms 798 length) which were slid (10 ms steps) over times surrounding stimulus onset. Each time window 799 spectra were compared, on a trial-by-trial basis given a penetration, with a window located before 800 stimulus onset (center at ca. -90 ms relative to onset; Wilcoxon signed rank tests, significance when p 801 < 0.01).

802 As per Medvedev and Kanwal (2008), the power spectra from each penetration were z-normalized 803 across trials, and the power in a given band was calculated by integrating (*trapz* function, Matlab) 804 over the z-normalized spectrum (per trial). Care was taken that the number of frequency samples 805 were comparable when integrating at different bands; the gamma band was therefore divided into 25-806 45 and 45-60 Hz sub-bands. We then determined whether the distribution of power in gamma and 807 low-frequencies were different, per penetration, by means of a 2-sample Kolmogorov Smirnov test 808 (alpha at 0.01). Because differences or lack of differences in the power distributions do not 809 necessarily imply the existence (or lack) of trial-by-trial correlation between the power of low and 810 gamma frequency bands, we tested whether these powers were correlated for every penetration, on a 811 trial by trial basis. In this context, correlations were significant given a p < 0.01.

#### 812 **4.8 Information theoretic analyses**

Information in the neuronal response regarding the acoustic stimuli was quantified by means of
Shannon's mutual information (MI; (Shannon, 2001)). The MI between a stimulus set *S* and a
response set *R* is mathematically expressed as follows:

816 
$$I(R;S) = H(R) - H(R|S),$$
 [4]

817 where H(R) is the response entropy (i.e. the overall variability of the response set), which is

818 expressed as:

819 
$$H(R) = -\sum_{r \in R} P(r) \log_2[P(r)], \quad [5]$$

820 while

821 
$$H(R|S) = -\sum_{s \in S} P(s) \sum_{r \in R} P(r|s) \log_2[P(r|s)], [6]$$

822 is referred to as the "noise entropy", representing the irreproducibility of the response given a 823 stimulus. The probabilities P(r), P(s) and P(r/s) indicate the probability of observing response r taken 824 from the set R, the probability of observing stimulus s from the set S, and the probability of observing 825 response r given stimulus s, respectively. If the logarithm in Eqs. 5 and 6 is of base 2, the MI has 826 units of bits. Each bit of information means that an external observer is able to reduce, by observing 827 the response, the uncertainty about the stimulus by a factor of 2 on a single trial basis. These 828 quantities were estimated by means of the Information Breakdown Toolbox (ibTB; (Magri et al., 829 2009)).

#### 830 4.8.1 Stimuli for MI computations

831 With aims of quantifying the amount of information provided by FAF and AC spiking regarding a 832 specific acoustic stream, we calculated each unit's ability to discriminate consecutive chunks of the 833 stimulus from each other (de Ruyter van Steveninck et al., 1997; Kayser et al., 2009; Kayser et al., 834 2010; Garcia-Rosales et al., 2018a). A particular sequence S (be it, for example, the natural the 835 distress call) was subdivided into non-overlapping, consecutive segments  $s_k$  (k = 1, 2, 3, ..., M), all of 836 length T = 4 ms. We chose this segment length so that results from this paper would be comparable 837 with previous data from the AC of C. perspicillata (Garcia-Rosales et al., 2018a). Using lengths in 838 the range of 2-12 ms did not alter the results qualitatively. Each segment s was treated as an 839 independent substimulus from the set S. Note that, in this framework, all substimuli are equiprobable.

840 **4.8.2 Rate and joint neuronal codes** 

The manner in which P(r) is quantified depends directly on the assumptions made to characterize the neuronal response (i.e. the neural code considered). Here we used a rate code (I<sub>rate</sub>), which determines how well a unit discriminates between each substimulus *s*, based on its spiking rate. The response set represented whether a spike occurred or not, and can be characterized as follows:  $R = \{0, 1\}$ , where 1

and 0 represent the occurrence or absence of a spike, respectively. P(r) was then the probability that a

- unit fired or not a spike across all trials, whereas P(r/s) was the probability of firing to a certain
- substimulus. The time window was sufficiently short to assume that, in general, a single spike would
- 848 occur within each time segment, and therefore binarized responses were used for MI calculations.
- 849 The information provided by joint responses from the FAF and the AC (I<sub>joint</sub>) was calculated by
- taking into account which unit elicited a spike in a merged response (see Fig. 6B; also referred to as
- 851 "line code" in the literature (Panzeri et al., 2007; Kayser et al., 2009)). That is, the response set was
- defined as  $R = \{ (0, 0); (0, 1); (1, 0); (1, 1) \}$ , where each member of the set represents whether and
- 853 which unit fired a spike (e.g. (0, 1) could indicate that the FAF unit did not fire, whereas the AC one
- did; (1, 0) would represent the opposite).

#### **4.8.3 Quantifying information from limited samples**

856 The probabilities in **Eqs. 5** and 6 are estimated empirically from the data, based on the representation 857 of neuronal responses described above (i.e. the neural codes). These empirically estimated 858 probabilities (such as P(r), or P(r/s)) are biased because it is impossible in practice to sample all 859 possible values of R a sufficiently large number of times (ideally, infinite). A number of methods 860 have been developed to deal with the sampling bias (Panzeri et al., 2007). In this study, we used the 861 Quadratic Extrapolation (QE) procedure (Strong et al., 1998), implemented in the ibTB. In addition 862 to the QE, we subtracted possible remaining biases by means of a bootstrap procedure (Montemurro 863 et al., 2008; Garcia-Rosales et al., 2018a), using 250 repetitions. For paired responses, we also used 864 the shuffling procedure (Panzeri et al., 2007) implemented in the iBTB together with the bootstrap 865 method. To corroborate that the information estimates presented in the results were not affected by 866 the limited sampling bias, we conducted numerical simulations in order to measure the dependence 867 of the bias on the number of trials. The results of these simulations are shown in **Fig. S8**, and indicate 868 that the number of trials used in this study (50) was sufficient to robustly estimate the information 869 quantities presented.

#### 870 4.9 Statistical analyses

All statistical analyses were conducted in Matlab (version 8.6.0.267246 (R2015b)), with custom-

- 872 written scripts using the Statistics and Machine Learning Toolbox. Tests for comparisons between
- the distributions of the quantities described above were always indicated in the main text. When
- 874 multiple comparisons were done, we performed False-Discovery Rate (FDR) corrections (e.g.,
- 875 comparing across multiple channel pairs in Fig. 3E) with the Benjamini and Hochberg procedure

876 (Benjamini and Hochberg, 1995). The significance threshold was set at an alpha of 0.05. If the p

values reported were uncorrected, it is stated so in the text. Effect sizes were calculated with the r

878 metric, which is defined as follows (Fritz et al., 2012):

$$r = \frac{W}{\sqrt{N}}, \qquad [7]$$

880 where *r* is the effect size, *W* is the test statistic of the Wilcoxon signed rank test used in this context,

and N is the sample size of the quantities being compared (N = 50). Values of  $r \le 0.3$  were

considered small effects, while  $0.3 \le r \le 0.5$  were considered as medium effects, and large effects

883 were considered when r > 0.5 (Fritz et al., 2012).

#### 884 **5** Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financialrelationships that could be construed as a potential conflict of interest.

# 887 6 Author Contributions

F.G.R and J.C.H designed the study. F.G.R collected the data. F.G.R analyzed the data and wrote
the manuscript. F.G.R, L.L.J, E.G.P, Y.C.C, and J.C.H discussed the results and reviewed the
manuscript.

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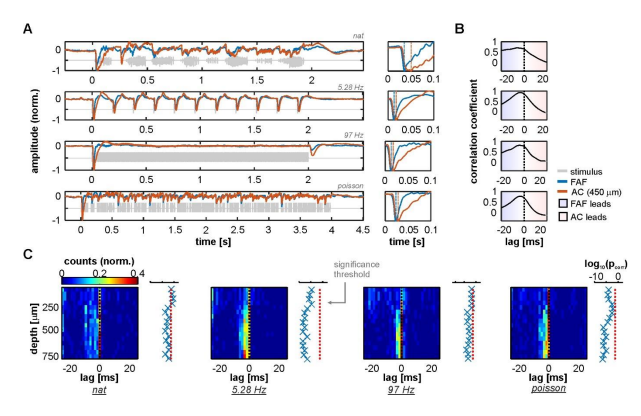
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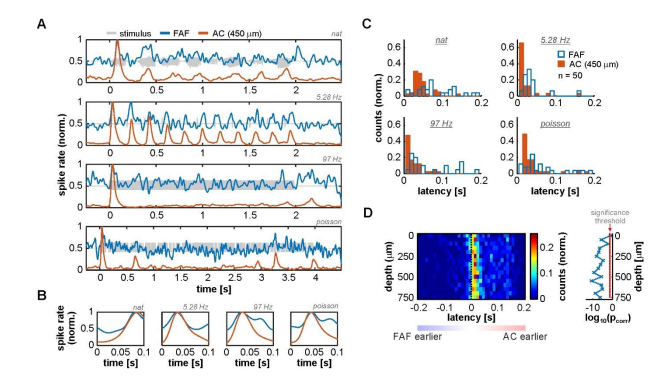
## 1152 **1 Data Availability Statement**

- 1153 The data that support the findings of this study are available from the corresponding authors upon
- 1154 reasonable request.



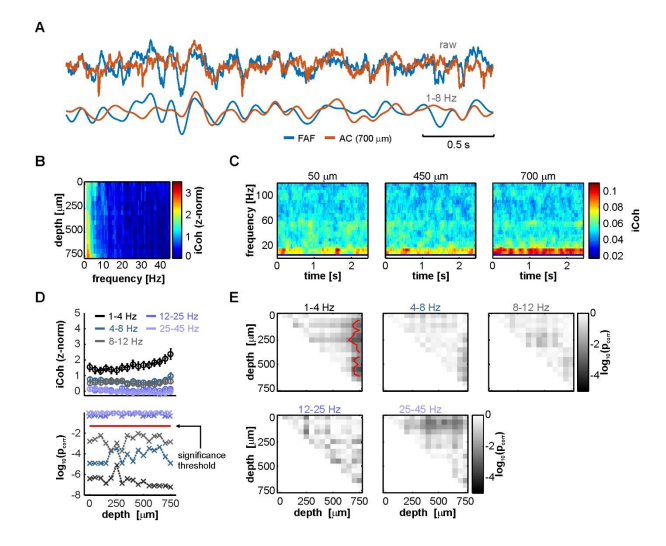
1157 Fig. 1. LFP stimulus-related activity in the frontal auditory field (FAF) precedes that of the auditory cortex (AC). (A) Left column: Grand average across all 50 penetrations of LFPs 1158 recorded from the FAF (blue) and the AC at a depth of 450 µm (orange) in response to the four 1159 1160 stimuli tested (ordered from top to bottom: natural sequence (nat), 5.28 Hz train, 97 Hz train, and 1161 the Poisson syllabic sequence (poisson); grey traces). Right column: zoom into the first 100 ms 1162 after stimulus onset. Negative peaks in the evoked potential are marked with vertical dashed 1163 lines. Note that peaks in the FAF occur earlier than in the AC. (B) Cross-correlation between 1164 traces in A (*left column*). Peaks in negative lags indicate that FAF field-potentials lead those in 1165 the AC. (C) Peak lags between the cross-correlation of FAF LFPs and AC LFPs, for all 1166 penetrations (n = 50) and across recording depths. Next to each heatmap, log-scaled corrected p values testing that the peak distribution is significantly below 0 (FDR corrected tailed Wilcoxon 1167 1168 signed rank tests;  $p_{corr} < 0.05$  for significance).

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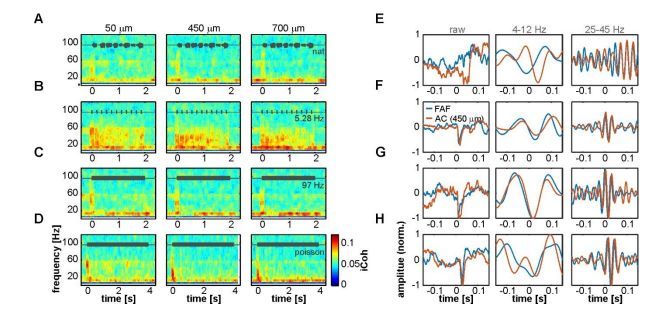
1171 Fig. 2. Spiking activity suggests the presence of fast inputs into the FAF. (A) Spiking responses from two simultaneously recorded units in the FAF (blue) and the AC (at 450 µm; blue), in 1172 1173 response to all stimuli tested (top to bottom). (B) Zoom-in into the first 100 ms after stimulus 1174 onset of the examples shown in A. Note that, for this pair, the peak response for the FAF unit 1175 was at least as fast as the for auditory cortical one. (C) Latency distribution of FAF (blue) units 1176 and AC units at 450  $\mu$ m, for all stimuli (n = 50 penetrations). The FAF was, overall, sluggish in 1177 comparison to the AC. (**D**) Response latency difference between simultaneously recorded 1178 spiking for FAF and AC at different depths (positive difference, FAF slower than AC; negative 1179 difference indicates the opposite). In some cases, FAF spiking responses occurred earlier than 1180 AC responses, although the AC was in general significantly faster than the FAF across channels, 1181 except in the case of the most superficial contact (FDR-corrected Wilcoxon signed rank tests, 1182 significance when  $p_{corr} < 0.05$ ). Log-converted p values and significance threshold are shown to 1183 the right of the latency distributions. The threshold is indicated as a red dashed line.

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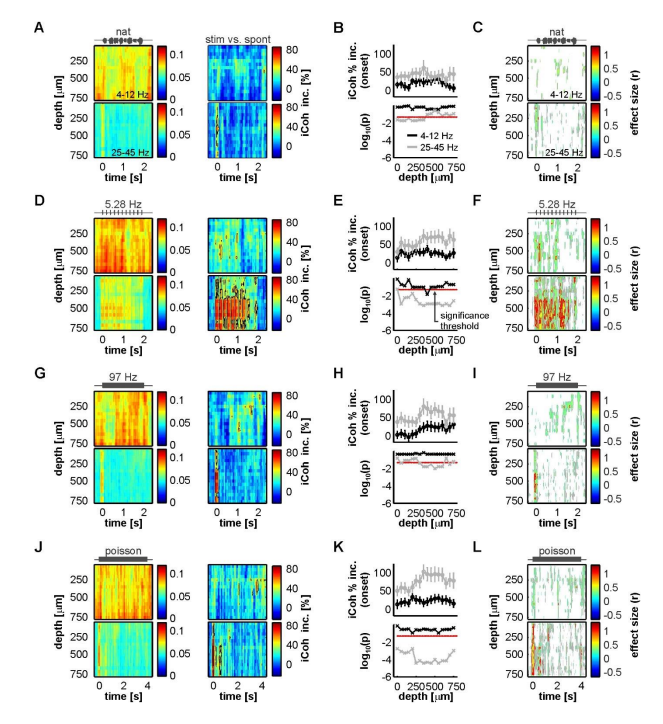
1186 Fig. 3. LFPs from FAF and AC are low-frequency coherent during spontaneous activity in a 1187 depth-dependent manner. (A) Simultaneously recorded LFP trace from the FAF (blue) and the 1188 AC at a depth of 700  $\mu$ m (orange). Raw (top pair) and low-frequency filtered (bottom pair) traces are shown. (B) Frequency-dependent average imaginary coherence (iCoh), z-normalized 1189 1190 to a surrogate distribution, across recording depths in the AC. Deep channels showed, on 1191 average, the strongest coherence values at low frequencies. (C) Time-resolved iCoh using the 1192 same segments as in **B**, with a sliding window of 200 ms (see Methods), the same used for 1193 analyzing stimulus-related synchronization. (**D**) Top: depth-dependent population z-normalized 1194 iCoh from **B**, for distinct frequency bands (delta, theta, alpha, beta and low gamma, see 1195 Methods; frequency ranges indicated in the plot), across all penetrations (shown as mean  $\pm$ 1196 SEM). *Bottom*: log-scaled corrected p values after testing, per frequency band, whether z-1197 normalized iCoh values were significantly higher than 0 across penetrations, per AC depth 1198 (FDR-corrected Wilcoxon signed rank tests, p<sub>corr</sub> < 0.05 for significance; threshold indicated as a 1199 horizontal red dashed line). (E) Significance matrices comparing, per frequency band, z-

- 1200 normalized iCoh values across different depths. Each cell (*i*, *j*) in a matrix depicts the log-scaled
- 1201  $p_{corr}$  obtained from statistically comparing coherence at channels with depths *i* and *j* in the AC.
- 1202 Red contour lines delimit regions of statistical significance (FDR-corrected Wilcoxon signed
- 1203 rank tests, significance when  $p_{corr} < 0.05$ ).



1206 Fig. 4. Interareal phase synchrony during acoustic sequence processing. (A-D) Mean time-1207 resolved coherence between LFPs from the FAF and the AC at three representative depths (50, 1208 450 and 700  $\mu$ m), in response to the natural sequence (A), a syllabic train of 5.28 Hz (B), a 1209 syllabic train of 97 Hz (C), and the syllabic train with a Poisson structure (D). Note that low 1210 frequency synchrony is high even without acoustic stimulation, and the appearance of gamma-1211 band evoked synchronization at the stimulus onset (time 0), albeit more weakly in response to 1212 the natural call in A. (E-H) LFP recordings from the AC (orange) and FAF (blue) around the 1213 time of stimulus onset (at 0 s; order in E-H corresponds to order in A-D), from single trials in a 1214 representative penetration. Left column depicts the raw LFP, whereas middle and right columns 1215 depict field-potentials filtered in 4-12 and 25-45 Hz low-frequency and gamma-bands, 1216 respectively.

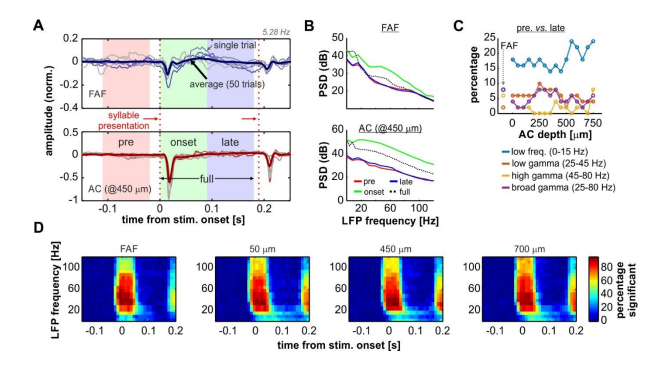
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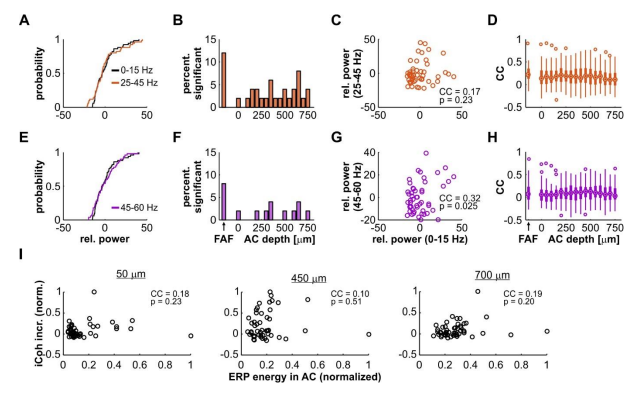
Fig. 5. Acoustic stimulation alters FAF-AC coherence mostly in the gamma-band. (A) *Top*:
average time course of iCoh while animals listened to the natural sequence (left) across
recording depths for low frequencies (4-12 Hz), and percentage increase of coherence in that
range relative to the spontaneous activity (right). *Bottom*: same as *Top*, but for iCoh values in the
gamma range (25-45 Hz). Black contour lines delimit regions with average increase of coherence
> 50%. (B) Population onset-related iCoh increase (median in the period of 0-150 ms after
stimulus onset) across depths in the AC (black traces, low-frequency band iCoh; grey traces,

1226	gamma-band iCoh; shown as mean +- SEM). On the bottom subpanel, log-scaled corrected p
1227	values obtained after testing that such increase was significantly different from 0% (black, low-
1228	frequency band; grey, gamma-band; FDR-corrected Wilcoxon signed rank tests, significance
1229	when $p_{corr} < 0.05$ , indicated as a red dashed line). (C) Time-resolved effect size of population
1230	iCoh percentage increase (r; see Methods) for the low-frequency band (top) and the gamma-
1231	range (bottom). Grey contour lines delimit regions of $r > 0.3$ , whereas red contours mark regions
1232	of $r > 0.5$ (medium and large effect sizes, respectively). r values are only shown for time points,
1233	across channels, where the coherence increase was significantly higher than 0% (Wilcoxon
1234	signed rank test, $p < 0.05$ ). ( <b>D-F</b> ) Same as <b>A-C</b> but considering a syllabic train at 5.28 Hz as
1235	stimulus. (G-I) Same as A-C, the stimulus being a syllabic train at 97 Hz. (J-L) Same as A-C,
1236	except the stimulus was the Poisson syllabic train.



1239 Fig. 6. Onset related power increase in AC and FAF. (A) Representative LFP recordings of one 1240 penetration pair (FAF, top; AC at 450 µm, bottom), depicting 6 single trials for illustrative 1241 purposes (thin lines) and the average across all 50 trials. The time segments of *pre*, *onset*, *late*, 1242 and *full*, used for analyses (see main text), are indicated in the graphs. First and second syllable 1243 presentations of the 5.28 Hz train are indicated with vertical, red dashed lines. (B) Power spectral 1244 density (PSD) of the pre (red), onset (green), late (blue) and full (black, dashed) periods from the 1245 data depicted in A (average over the 50 trials), in the FAF (top) and AC (bottom). (C) Percentage 1246 of penetrations (after a total of 50) for which the power (at several frequency bands, indicated in 1247 the figure) was significantly different during the *late* period than during the *pre* period. (**D**) 1248 Time-frequency analysis illustrating the percentage of penetrations in FAF and AC (at three 1249 representative depths: 50, 450 and 700  $\mu$ m) in which the power at a given time window was 1250 significantly higher than the power at a window preceding the stimulus onset.

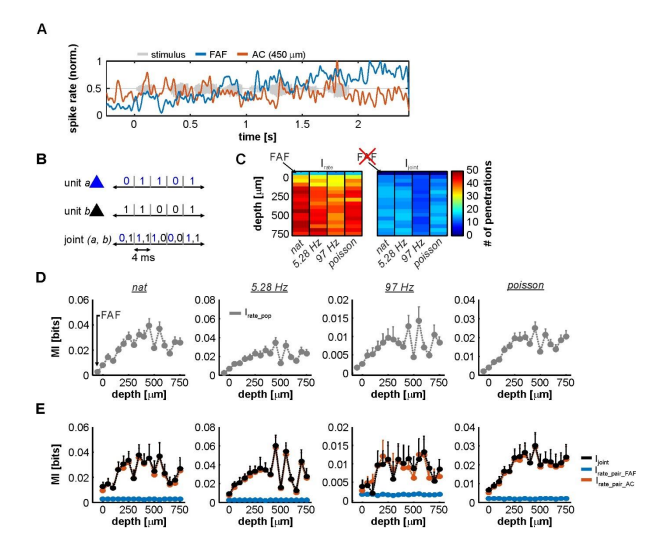
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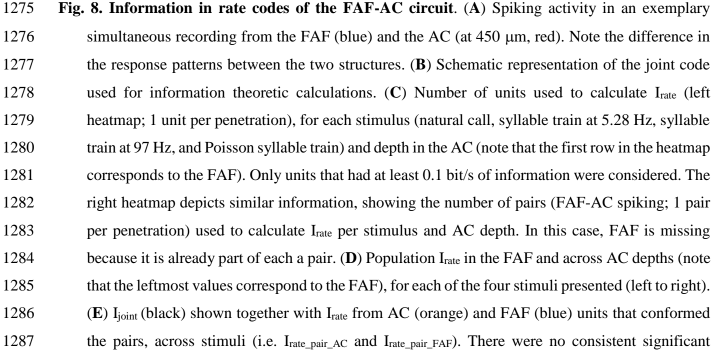
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1253 Fig. 7. Power distributions of gamma-band and low-frequency LFPs in AC and FAF. (A) 1254 Distributions of the relative power of low-frequency (0-15 Hz; black) and gamma-band activity 1255 (25-45 Hz; orange) across trials, recorded from a single representative penetration in the FAF. 1256 These distributions were not significantly different from each other (2-sample Kolmogorov-1257 Smirnov test, p = 0.84). (B) Percentage from the total number of penetrations (n = 50) for which 1258 the distributions of low-frequency and gamma (25-45 Hz) power were significantly different 1259 from each other at an alpha of 0.01, in FAF and at different depths of the AC. (C) Scatter plot 1260 and correlation coefficient (CC) of the trial-by-trial relationship between low-frequency and 1261 gamma band (25-45 Hz) power (n = 50 trials), for the same representative penetration shown in 1262 **A**. The CC was of 0.17, and it was not significant: p = 0.23. (**D**) Distribution, in FAF and at all 1263 AC depths, of CCs between gamma-band (25-45 Hz) and low-frequency power. The median in the AC across depths was of 0.17, whereas the median in the FAF was of 0.22. (E-H) Similar to 1264 1265 **A-D**, but the gamma range considered was of 45-60 Hz (signaled in purple). In panel **E**, both distributions were also not significant from each other (2-sample Kolmogorov-Smirnov test, p = 1266 1267 0.84). The CC in G was of 0.32, and it was not significant at an alpha of 0.01 (p = 0.025). In 1268 panel H, the median across depths in the AC was of 0.08, whereas the median CC in the FAF 1269 was of 0.07. (I) Correlations between evoked-potential (ERP) energy in AC and gamma-band 1270 coherence increase (same as in Fig. 5) for three representative depths in the AC (at 50, 450 and

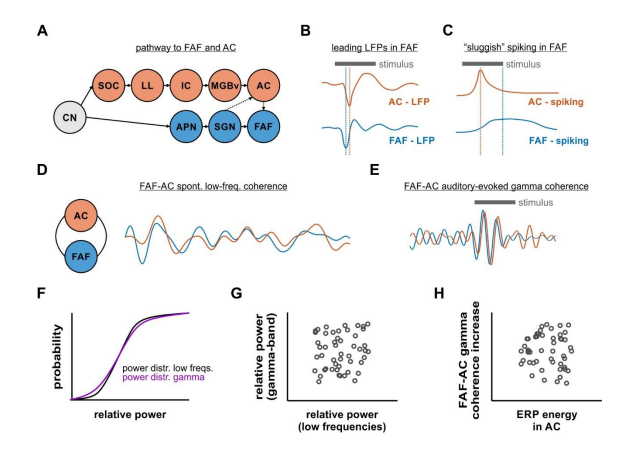
- 1271 700 μm; all depths are shown in **Fig. S3**). Values are normalized for clarity. There were no
- 1272 significant correlations at any of the depths shown ( $p \ge 0.2$ ).



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- 1288 differences between Ijoint and Irate, when the latter was calculated for AC units (Wilcoxon signed
- 1289 rank tests, p > 0.05).



1292 Fig. 9. Functional coupling dynamics in the FAF-AC circuit of C. perspicillata. (A) Representation of the auditory pathway to the FAF and the AC (CN, cochlear nucleus; SOC, superior olivary 1293 1294 complex: LL, lateral lemniscus: IC, inferior colliculus: MGBv, ventral division of the medial 1295 geniculate body of the thalamus; AC, auditory cortex; APN, anterolateral periolivary nucleus; 1296 SGN, suprageniculate nucleus of the thalamus; FAF, frontal auditory field). (B) Schematic 1297 representation illustrating that stimulus-related LFPs in FAF lead relative to those in the AC (see 1298 Fig. 1). (C) Fast LFP responses in FAF do not necessarily elicit fast spiking responses (schematic). Neurons in the frontal region typically respond more "sluggishly" than their auditory cortical 1299 1300 counterparts. (**D**) FAF and AC were coherent in low-frequencies during spontaneous activity (i.e. 1301 in the absence of sound stimulation). (E) During acoustic processing in passive listening animals, 1302 low-frequency coherence was unaltered in the FAF-AC circuit, although there was an emergence 1303 of auditory-evoked gamma band coherence in the network. Traces in panels D and E are based on 1304 data shown in Figs. 3 and 4. Note that, for illustrative purposes, the temporal scales and amplitudes 1305 in **D** and **E** are not comparable. (F) Distributions of power in low and gamma-band frequencies 1306 were typically not significantly different from each other. However, there was very weak trial-by-1307 trial correlation between low-frequency and gamma power ( $\mathbf{G}$ ), as well as very low correlations

- 1308 between event-related potential (ERP) energy and gamma coherence increase across penetrations
- 1309 (**H**).
- 1310
- 1311