Sensory and Motor Systems

Signature Patterns for Top-Down and Bottom-Up Information Processing via Cross-Frequency Coupling in Macaque Auditory Cortex

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Abstract

Predictive coding is a theoretical framework that provides a functional interpretation of top-down and bottom-up interactions in sensory processing. The theory suggests there are differences in message passing up versus down the cortical hierarchy. These differences result from the linear feedforward of prediction errors, and the nonlinear feedback of predictions. This implies that cross-frequency interactions should predominate top-down. But it remains unknown whether these differences are expressed in cross-frequency interactions in the brain. Here we examined bidirectional cross-frequency coupling across four sectors of the auditory hierarchy in the macaque. We computed two measures of cross-frequency coupling, phase–amplitude coupling (PAC) and amplitude–amplitude coupling (AAC). Our findings revealed distinct patterns for bottom-up and top-down information processing among cross-frequency interactions. Both top-down and bottom-up interactions made prominent use of low frequencies: low-to-low-frequency (theta, alpha, beta) and low-frequency-to-high-gamma couplings were predominant top-down, while low-frequency-to-low-gamma couplings were predominant bottom-up. These patterns were largely preserved across coupling types (PAC and AAC) and across stimulus types (natural and synthetic auditory stimuli), suggesting that they are a general feature of information processing in auditory cortex. Our findings suggest the modulatory effect of low frequencies on gamma-rhythms in distant regions is important for bidirectional information transfer. The finding of low-frequency-to-low-gamma interactions in the bottom-up direction suggest that nonlinearities may also play a role in feedforward message passing. Altogether, the patterns of cross-frequency interaction we observed across the auditory hierarchy are largely consistent with the predictive coding framework.

Key words: auditory cortex; cross-frequency coupling; information processing; neural coding; predictive coding; top down

Significance Statement

The brain consists of highly interconnected cortical areas, yet the patterns in directional cortical communication are not fully understood, in particular with regard to interactions between different signal components across frequencies. We used a novel, convenient, and computationally advantageous Granger-causal framework to examine bidirectional cross-frequency interactions across four sectors of the auditory cortical hierarchy in macaques. Our findings reveal that cross-frequency interactions are predominant in the top-down direction, with important implications for theories of information processing in the brain such as predictive coding. Our findings thereby extend the view of cross-frequency interactions in auditory cortex, suggesting that they also play a prominent role in top-down processing.
Introduction

One fundamental yet poorly understood component of cortical computation is the presence of widespread reciprocal cortical connections (Salin and Bullier, 1995; Kveraga et al., 2007; Romanski and Averbeck, 2009; Scott et al., 2017). Moreover, most previous work has focused on the visual pathways. It remains unknown whether there are fundamental differences between top-down (TD) and bottom-up (BU) information processing along the auditory pathway. Cross-regional top-down effects in particular have remained unexplored, as most work has focused on bottom-up (Giraud and Poeppel, 2012; Hyafil et al., 2015a) or intraregional effects (Lakatos et al., 2016). In the present study, we examined bidirectional information processing across the macaque auditory hierarchy through an analysis of cross-frequency coupling (CFC). We computed two types of CFC measures, amplitude–amplitude coupling (AAC) and phase–amplitude coupling (PAC), for both natural and synthetic stimuli.

Cross-frequency coupling refers to the correlation between the phase or amplitude component of one frequency and the amplitude component of other frequencies. This differs from linear analyses like coherence, which only examine coupling between signals at a single frequency. CFC, in contrast, considers interactions across the frequency spectrum. CFC may occur in several forms (Jirsa and Mueller, 2013) and has been observed during various tasks across several species including humans, macaques, and rodents. PAC, in particular, has received widespread attention. Coupling of low-frequency (e.g., theta, alpha) phase to high-frequency (e.g., gamma) amplitude has been associated with various behavioral mechanisms including learning, memory, and attention (Schack et al., 2002; Canolty et al., 2006; Jensen and Colgin, 2007; Cohen, 2008; Tort et al., 2008, 2009; Canolty and Knight, 2010; Kendrick et al., 2011; Doesburg et al., 2012; Colgin, 2015; Hyafil et al., 2015b). The reason for interest in PAC is that it may offer a mechanistic account of information coordination across neural populations and timescales. The phase of low-frequency oscillations could modulate the amplitude of high-frequency oscillations, putatively allowing control of spiking activity (Ray and Maunsell, 2011; Buzsáki et al., 2012). AAC has also been shown to correlate with behavior (Canolty and Knight, 2010), but has been less explored mechanistically. Here we obtained estimates of both PAC and AAC strength as measures of cross-regional information processing in the bottom-up and top-down directions, allowing the two measures to be contrasted and compared in terms of informational content.

One prevalent theory of information processing in the brain that separates bottom-up and top-down components is predictive coding. In this (Bayesian) view of the brain, expectations are formed about incoming sensory information. These top-down predictions (expectations) are compared with bottom-up sensory information (outcomes) and representations are then potentially updated based on a prediction error (surprise), which is the difference between sensory inputs and their expectations. Neural implementations of predictive coding suggest that predictions are formed in higher-order areas through a (linear) accumulation of prediction errors. The accumulation of prediction errors in higher-order areas results in the attenuation of high frequencies; thus, feedback connections can be cast as a low-pass or Bayesian filter. Conversely, prediction errors arising in lower-order areas are a nonlinear function of predictions. This means high frequencies can be created in lower-order areas due to the inherent nonlinearity. Hence, predictive coding implies that lower-order areas (where prediction errors arise) should display relatively higher frequencies (e.g., gamma range) than higher-order areas (where predictions are formed; Friston, 2008; Bastos et al., 2012). Several studies have supported this view (Bastos et al., 2012, 2015; Fontolan et al., 2014; Michalareas et al., 2016), though not without exceptions.

It remains unknown whether this prediction is expressed in cross-frequency interactions in the brain. Here we probe this prediction by examining differences in cross-frequency coupling strength in the bottom-up and top-down directions across the frequency spectrum. Based on the predictive coding framework (Friston, 2008; Bastos et al., 2012), we would expect there to be an asymmetry between bottom-up and top-down interactions. This asymmetry arises due to the linear effect of prediction errors on predictions in the bottom-up direction and the nonlinear effect of predictions on prediction errors in the top-down direction. Thus, we would expect nonlinear, cross-frequency interactions to predominate in the top-down direction.

Previous work has shown differences in how auditory cortical areas process natural and synthetic sounds (Fukushima et al., 2014). It is possible that these differences translate into systematic differences in CFC patterns. If the revealed CFC pattern is a more general hallmark of interareal communication, though, it should not be specific to stimulus type: natural and synthetic stimuli should show similar overall coupling patterns.

Our work revealed signature patterns of top-down and bottom-up processing in cross-regional PACs and AACs. These patterns were conserved across coupling types and across natural and synthetic stimuli. We found that low-to-low-frequency (theta, alpha, beta) and low-fre-
quency-to-high-gamma interactions were predominant top-down, while low-frequency-to-low-gamma couplings were predominant bottom-up. Our findings suggest that the modulation of gamma frequency power by low-frequency rhythms plays a role in bidirectional information transfer across the auditory hierarchy. The finding of pre-dominant cross-frequency interactions in the top-down direction across the auditory hierarchy largely agrees with the predictive coding framework.

Materials and Methods

Subjects

Three adult male rhesus monkeys (Macaca mulatta) weighing 5.5–10 kg were used for recordings. All procedures and animal care were conducted in accordance with the Institute of Laboratory Animal Resources Guide for the Care and Use of Laboratory Animals. All experimental procedures were approved by the National Institute of Mental Health Animal Care and Use Committee.

Stimuli

The stimuli used for the main experiment included 20 conspecific monkey vocalizations (VOCs) and two sets of 20 synthetic stimuli each [envelope-preserved sound (EPS); and spectrum-preserved sound (SPS)] derived from the original VOC stimuli. The VOC stimulus set consisted of 20 macaque vocalizations used in previous studies (Kikuchi et al., 2010; Fukushima et al., 2014).

To obtain EPS from VOC stimuli, the envelope of a particular vocalization was estimated based on the amplitude component of the Hilbert transform of the original stimulus. The amplitude envelope was then multiplied by broadband white noise to create the EPS stimulus. Thus, all 20 EPS stimuli exhibited flat spectral content; these stimuli could not be discriminated based on spectral features, while the temporal envelopes (and thus the durations) of the original vocalizations were preserved.

SPS stimuli were obtained by first generating broadband white noise with a duration of 500 ms and computing its Fourier transform. The amplitude of the SPS stimulus in the Fourier domain was then replaced by the average amplitude of the corresponding VOC stimulus before transforming back to the time domain by computing the inverse Fourier transform. This resulted in a sound wave form that preserved the average spectrum of the original vocalization, while exhibiting a flat temporal envelope, random phase, with a duration of 500 ms. A 2 ms cosine rise/fall was then imposed on the stimulus to avoid abrupt onset/offset effects. Hence, all 20 SPS stimuli exhibited nearly identical, flat temporal envelopes; these stimuli could not be discriminated using temporal features, while the average spectral power of the original vocalizations was preserved.

A total of 60 different stimuli were presented in pseudorandom order, with an interstimulus interval of 3 s. Each stimulus was presented 60 times. The sound pressure levels of the stimuli measured by a sound level meter ranged from 65 to 72 dB at a location close to the animal’s ear of the animal. Stimulus duration ranged from ~0.15 to 1 s (Fukushima et al., 2014).

Recordings

Custom-designed micro-electrocorticography (μECoG) arrays (NeuroNexus) were used to record field potentials from macaque auditory cortex. Arrays were machine fabricated on a very thin polyimide film (20 μm), with each array featuring 32 recording sites, 50 μm in diameter each, on a 4 × 8 grid with 1 mm spacing (i.e., 3 × 7 mm rectangular grid). Two animals, monkeys B and K, were both implanted with arrays in the left hemisphere, while one animal, monkey M, was implanted with arrays in the right hemisphere. Three of the arrays in each monkey were placed on top of supratemporal plane (STP) in a caudorostrally oriented row.

To implant the arrays, we removed a frontotemporal bone flap extending from the orbit ventrally toward the temporal pole and caudally behind the auditory meatus and then opened the dura to expose the lateral sulcus. The most caudal of the three ECoG arrays on the STP was placed first and aimed at area A1 by positioning it just caudal to an (imaginary) extension of the central sulcus and in close proximity to a small bump on the STP, both being markers of the approximate location of the A1. Each successively more rostral array was then placed immediately adjacent to the previous array to minimize interarray gaps. The arrays on the lateral surface of the STG were placed last. The probe connector attached to each array was temporarily attached with cyanoacrylate glue or Vetbond to the skull immediately above the cranial opening. Ceramic screws together with bone cement were used to fix the connectors to the skull. The skin was closed in anatomic layers. Postsurgical analgesics were provided as necessary, in consultation with the National Institute of Mental Health veterinarian.

Stimulus presentation and recording parameters

The monkey was placed in a sound-attenuating booth for the experiment (Biocooustics Instruments). The sound stimuli were presented while the monkey sat in a primate chair and listened passively with its head fixed. Auditory evoked potentials from the 128 channels of the ECoG array were bandpass filtered between 2 and 500 Hz, digitally sampled at 1500 Hz, and stored on hard-disk drives by a PZ2–128 preamplifier and the RZ2 base station (Tucker-Davis Technologies).

Data preprocessing

Data analysis was performed using MATLAB R2017a (MathWorks) software. Since there was little significant auditory evoked power >250 Hz, recordings were low-pass filtered and resampled at 500 Hz to enhance calculation speed and reduce memory requirements. The signal was bandfiltered at 60 Hz with a narrow-band filter to eliminate the possible presence of line noise; a sixth-order Butterworth filter was chosen to achieve a sharp drop-off in the stop-band, thus minimizing the effect of the filter on neighboring frequency bands.

The 96 sites on the STP were grouped based on the characteristic frequency maps obtained from the high-gamma power of the evoked response to a set of pure-tone stimuli. The change in frequency tuning along the STP reverses across areal boundaries, and these rever-
sals were therefore used to identify the areal boundaries. This resulted in a grouping of electrodes into four sectors (Fig. 1), which were estimated to correspond to the following subdivisions at a caudorostral level within macaque auditory cortex: sector (Sec) 1, A1/ML (primary auditory cortex/middle lateral belt); Sec 2, R (rostrotemporal core region of the auditory cortex)/AL (anterior lateral belt region of the auditory cortex); Sec 3, RTL (lateral rostrotemporal belt region of the auditory cortex); and Sec 4, RTp (rostrotemporal pole area). The recorded signal from each site partitioned into sectors 1–4 was rereferenced by subtracting the average of all
tex); and Sec 4, RTp (rostrotemporal pole area). The
(lateral rostrotemporal belt region of the auditory cor-
tral core region of the auditory cortex)/AL (anterior
ary auditory cortex/middle lateral belt); Sec 2, R (ros-
following subdivisions at a caudorostral level within

We defined the following terminology to refer to partic-
ular frequency spaces throughout this article: delta range,
0–2.5 Hz; theta range, 2.5–7.5 Hz; alpha range, 7.5–12.5
Hz; low beta range, 12.5–22.5 Hz; high beta range, 22.5–
42.5 Hz; low gamma range, 42.5–67.5 Hz; and high
gamma range, >67.5 Hz. The delta frequency band (0–
2.5 Hz) was not included in the analysis since the pream-
plifier used during the recording sessions does not allow
for recording frequency components <2 Hz and the
power in this band was expectedly low.

**Phase-amplitude (PAC) and amplitude-amplitude
(AAC) cross-frequency coupling calculation**

**Frequency transformation**

The field potential data was first transformed into fre-
quency space by computing the fast Fourier transform
(FFT) of the signal in every channel using a 200 ms
Hanning window, stepped by 200 ms. Thus, we obtained
a frequency representation of the data at a 5 Hz band-
width ranging from 2.5 to 132.5 Hz, which satisfies the
Nyquist criterion. We obtained the phase and amplitude
information in a given source and target region for every
channel and normalized the amplitude information in each
frequency band by the corresponding power in that fre-
quency band, averaged across the entire experiment, so
as to account for decreasing power with frequency (Ex-
tended Data Fig. 2–1).

**Obtaining CFCs using canonical correlation**

We computed two types of cross-frequency coupling,
PAC and AAC. Couplings were calculated for every chan-
el pair across the four sectors (S1–S4), resulting in six
possible cross-sector pairs (S1–S2, S1–S3, S1–S4, S2–
S3, S2–S4, and S3–S4). We obtained estimates of cou-
pling strength in two directions, bottom-up and top-down.
In the former case, the source component was derived
from a recording site in a sector lower in the auditory
hierarchy than the target component (referred to as
“bottom-up coupling”), while in the latter case the source
component came from a sector higher in the auditory
hierarchy than the target component (referred to as “top-
down coupling”; Fig. 1). Ultimately, we were interested
in examining the difference in coupling strength in the
top-down and bottom-up directions across the auditory
hierarchy. Results are presented collapsed across all
cross-sector pairs (Fig. 2), as well as separately for every
cross-sector pair (Fig. 3), averaged across all channel
pairs and across the three animals in both cases.

We used canonical correlation analysis (CCA) to com-
pute both types of coupling. The advantage of CCA is that
it computes coupling among multivariate observations in
the source and target regions. This framework, therefore,
allows CFC values to be computed across the entire
frequency range simultaneously. Furthermore, some ap-
proaches to calculating AAC and PAC first carry out
dimensionality reduction on the source and target region
signs separately, and then calculate correlations in the
low-dimensional space. If the variance preserving dimen-
sions in the source and target space are not the dimen-
sions that maximize correlations between the source and
target space, interactions may be missed. CCA on the
other hand finds the low-dimensional representations that
maximize coupling between the source (X) and target (Y)
space. In CCA, one is seeking projections (linear combi-
nations) \( U = X A \) and \( V = Y B \), such that the correlation
\( \text{corr}(U, V) \) among these derived directions is maximal
(Durstein, 2017).

We conducted our analyses using a Granger causal
framework. Therefore, we first conducted CCA between
amplitude in the target region and time-lagged versions of
the amplitude also within the target region, obtaining an
estimate of the coupling strength \( \hat{Y} \) that could be in-
ferred purely from coupling of the signal at the target
electrode onto itself. For computational reasons, we
used two time lags to estimate the target signal (we
found that using one time lag did not affect our results).
More explicitly, we transformed the current and lagged
signal into the frequency domain as described above
and computed the canonical correlation between the
lagged components and the current signal in the target.
We used the first 10 canonical values only to obtain \( \hat{Y} \)
(results did not differ from those obtained from includ-
ing more entries). We then obtained the residual \( \hat{Y} \) from
this analysis. Subsequent analysis was performed us-
ing the residuals \( \hat{Y} \) from this regression as the signal for
the target region.

We then computed PAC and AAC cross-frequency cou-
plings across the four sectors (Fig. 1) by regressing the
phase or amplitude component of the signal from the
source sector, respectively, on the amplitude component
of the signal in the target sector for every channel. More
specifically, we calculated instantaneous coupling in 200
ms windows between the source and the target region for
each electrode pair. PAC and AAC were calculated in the
CCA framework across all time windows and trials to-
gether, simultaneously across all frequency bands. In the
case of PAC, \( X_{n \times d} \) contained the sine and cosine trans-
formed phase component of the signal, while \( Y_{n \times d} \)
contained the squared and log-transformed amplitude
component of the signal, with \( n = 10,800 \) (1200 trials \( \times 9 
windows) observations for VOC and SPS stimuli and
\( n = 10,773 \) observations (1197 trials \( \times 9 
windows) for EPS stimuli, \( d2 = 25 \), and \( d1 = 25 \) or \( d1 = 25 \times 2 \) for PAC and
AAC, respectively (Eq. 1).

After obtaining the canonical coefficient matrices
\( B_{d2 \times d} \) and \( A_{d1 \times d} \) within the CCA framework (Eq. 1), one
Figure 1. Coupling analysis overview: recordings were made from four μECoG arrays spanning auditory cortex while monkeys listened to 1 of 20 natural VOCs, 20 synthetic EPSs, or 20 synthetic SPSs. Electrodes were partitioned into four sectors along the caudorostral axis (S1, A1/ML; S2, R/AL; S3, RTL; S4, RTp). We decomposed the signal into its time–frequency representation, and obtained amplitude and phase components for each sector. Plots of signal amplitude (in dB) are normalized to the baseline activity before stimulus onset. We investigated two types of coupling across sectors: amplitude–amplitude and phase–amplitude coupling. We computed both types of coupling in the bottom-up and top-down directions. Top-down coupling was defined as coupling in which...
can obtain the coupling matrix $P_{d_2 \times d_1}$ by multiplying the $B$ and $A$ matrices with the canonical values $S$ of the correlation matrix (Eq. 2). We used the first 10 directions ($d = 10$) that captured most of the interaction; results did not differ from those obtained from including all entries. For AAC, the final cross-frequency coupling matrix is $P$. For PAC, one combines the sine and cosine components of $P$ by computing their Euclidian norm, yielding the final coupling matrix $P_{d_2 \times d_1}$ with $d_2 = d_1$:

$$
U_{x \times d} = X_{x \times d_1} A_{d_1 \times d_2}, V_{x \times d} = \tilde{Y}_{x \times d_2} B_{d_2 \times d_1}, \arg \max_{A, B} \text{corr}(U, V).
$$

(1)

$$
P_{d_2 \times d_1} = B S^T A, P_{d_2 \times d_1} = \left( \sqrt{P_1 + P_{i_2}}, \ldots, \sqrt{P_m + P_{i_m}} \right).
$$

(2)

Obtaining the difference between TD and BU coupling

We obtained AAC and PAC matrices in this manner for both the bottom-up and the top-down direction. In bottom-up coupling, the source component was derived from a recording site in a sector lower in hierarchy than the target component (referred to as “bottom-up coupling”), while in top-down coupling, the source component came from a sector higher in hierarchy than the target component (Fig. 1). We then obtained the difference $\Delta P_{\text{same stim}}$ between the top-down and bottom-up coupling matrices $P_{\text{Top-Down}}$ and $P_{\text{Bottom-Up}}$ (Eq. 3). This was done separately for each of the cross-sector pairs. Results are presented collapsed across all cross-sector pairs (Fig. 2), as well as separately for every cross-sector pair (Fig. 3), as follows:

**Figure 2.** Top-down vs bottom-up phase–amplitude and amplitude–amplitude coupling in natural vocalizations (VOC). **A, B,** Phase-amplitude coupling (PAC) strength in the top-down (**A**) and bottom-up (**B**) directions. Depicted are canonical correlation-derived coupling coefficients (see Materials and Methods). **C,** Difference in top-down and bottom-up in PAC strength. Significant differences are enclosed by black rectangles ($p < 0.01$, cluster-corrected). **D, E,** Amplitude-amplitude coupling (AAC) strength in the top-down (**D**) and bottom-up (**E**) directions. Depicted are canonical correlation-derived coupling coefficients. **F,** Difference between top-down and bottom-up in AAC strength. Significant differences are marked by black outlines ($p < 0.01$, cluster corrected). Results are depicted averaged across all channels, cross-regional pairs, and animals. See Extended Data Figure 2-1, Extended Data Figure 2-2, and Extended Data Figure 2-3.
Obtaining the difference in coupling strength between original (VOC) and synthetic stimuli (EPS/SPS)

To examine the difference in coupling strength between natural and synthetic stimulus types (Figs. 4, 5), we computed the difference between VOC and EPS stimuli (Fig. 4) and between VOC and SPS stimuli (Fig. 5) separately in the TD (Eq. 4) and BU (Eq. 5) directions.

\[
\Delta P = P_{\text{Top-Down}} - P_{\text{Bottom-Up}}
\]

Equation 3

Statistical analysis

Paired \(t\) tests were performed to establish statistical significance of the difference between top-down and bottom-up coupling (\(\Delta P\); Figs. 2, 3, 4), as well as the differences between original and synthetic stimuli (\(\Delta P_{\text{VOC vs Synthetic}}^{\text{TD}}\), \(\Delta P_{\text{VOC vs Synthetic}}^{\text{BU}}\); Fig. 5). This was done separately for each frequency pair, by testing whether the mean of the difference between top-down and bottom-up coupling (or the difference between original and synthetic stimuli in each direction) was significantly different from zero (null hypothesis). Lilliefors test for normality was conducted for each frequency pair to ensure that the normality assumption was satisfied, and histogram plots were inspected for select frequency pairs (Extended Data Fig. 2-2, Extended Data Fig. 2-3). More specifically, to assess overall differences in coupling strength between the two directions (Figs. 2, 4) and to assess differences in coupling strength between natural and synthetic stimuli (Fig. 5), paired \(t\) tests were conducted across all channel pairs, cross-regional pairs, and animals.
together ($df = 9344$). To assess coupling differences for each regional pair separately (Fig. 3), paired $t$ tests were conducted separately for each regional pair, across all channel pairs and animals together ($df = 1056$).

Figure 4. Top-down vs bottom-up phase–amplitude and amplitude–amplitude coupling in synthetic envelope-preserved stimuli (EPSs) and synthetic spectrum-preserved stimuli (SPSs) A. Difference in top-down and bottom-up PAC strength in EPS stimuli. B. Difference in top-down and bottom-up AAC strength in EPS stimuli. C. Difference in top-down and bottom-up PAC strength in SPS stimuli. D. Difference in top-down and bottom-up AAC strength in SPS stimuli. Significant differences are marked by black outlines ($p \leq 0.01$, cluster corrected). Results are depicted averaged across all channels, cross-regional pairs, and animals. See Extended Data Figure 4-1, Extended Data Figure 4-2, Extended Data Figure 4-3, Extended Data Figure 4-4, Extended Data Figure 4-5, and Extended Data Figure 4-6.

Significant phase–amplitude and amplitude–amplitude coupling (highlighted by black outlines around the coupling values throughout the figures) was based on corrected $p$ values using nonparametric permutation tests.
that generated null distributions of the maximum cluster size (Nichols and Holmes, 2002; Maris and Oostenveld, 2007). This procedure implicitly adjusts for searching over multiple frequencies. The null distribution was obtained by randomly reassigning electrodes to source or target region and recalculating differences in coupling strength for 100 repeats. Clusters of the 99th percentile (corresponding to \( p < 0.01 \)) were considered significant and are reported throughout the manuscript (marked by black outlines).

**Results**

We analyzed surface potentials recorded simultaneously from multiple cortical areas in the auditory cortex of three macaques while the animals listened to auditory stimuli (Fig. 1). These auditory stimuli consisted of 20 natural conspecific vocalizations, 20 EPSs, and 20 SPSs, which were derived from the original vocalizations (Fukushima et al., 2014). A previous study established the approximate rostrocaudal location of implanted \( \mu \)ECoG arrays along the auditory hierarchy by the characteristic frequency of each electrode contact (Fukushima et al., 2012, 2014). This partitioned the recording sites into four sectors (S1–S4), putatively spanning the caudorostral levels from the core A1 (S1) to RTp (S4), including some of the surrounding belt areas (Fig. 1). We operationally defined feedforward to occur from earlier to later sectors (bottom-up direction), and feedback to occur from later to earlier sectors (top-down direction).

We wanted to examine cross-frequency interactions across the auditory hierarchy. To characterize phase–amplitude and amplitude–amplitude coupling, we first decomposed the signal into its spectral component using the short-time FFT and obtained phase and amplitude components (Fig. 1). We then used canonical correlation analysis (CCA) to parameterize the cross-frequency coupling (Materials and Methods). The fluctuations in amplitude in the target sector were predicted based on instantaneous amplitude fluctuations in the source sector within 200 ms windows. Furthermore, the target and source time series were adjusted to remove the effect of predicted fluctuations from the target region. In other words, by appealing to the notion of Granger causality, a significant mapping between the source and the target, which cannot be explained by the history of the target, is evidence of (cross-frequency) coupling.

We computed coupling strength across the frequency spectrum for each cross-regional pair, separately in the top-down and bottom-up direction (Fig. 1). Top-down...
coupling was defined as coupling in which the source electrode came from a sector of higher order than the target electrode, and vice versa for bottom-up coupling. For example, coupling between the phase (or amplitude) in S4 and the amplitude in S1 is defined as top-down, and coupling between the phase (or amplitude) in S1 and the amplitude in S4 is defined as bottom-up in the case of PAC (or AAC). This definition is in accordance with a previous approach (Fontolan et al., 2014).

**Distinct cross-frequency coupling signatures for bottom-up and top-down coupling, both involving low frequencies**

We first examined PAC and AAC across the four sectors, in auditory evoked potentials to 20 conspecific VOCs. Both CFC measures peaked in the theta frequency range and showed relatively strong coupling across all the target amplitude frequencies (Fig. 2A,B,D,E). The field potential signal showed the highest power in the same frequency band decaying with the typical 1/f pattern thereafter (Extended Data Fig. 2-1), thus satisfying a prerequisite for the presence of physiologically meaningful CFC (Aru et al., 2015).

Among PACs, both top-down and bottom-up coupling were dominant between theta phase and broadband amplitude (Fig. 2A,B). Examining differences in coupling strength in the two directions more closely (Fig. 2C), PACs showed dominant top-down coupling in the low-frequency space (theta, alpha, and beta phase to theta, alpha, and beta amplitude), as well as between low-frequency phase and high-gamma (>67.5 Hz) amplitude (p ≤ 0.01, cluster corrected). Bottom-up PAC was dominant between low-frequency phase (theta, alpha, beta) and low-gamma (47.5-62.5Hz) amplitude (p ≤ 0.01, cluster corrected).

AACs showed a pattern similar to that of PACs in both directions, but coupling strength and differences in coupling strength were overall by an order of magnitude lower than those for PACs (Fig. 2D,E). AACTs replicated the pattern observed in PACs, with dominant theta to broadband coupling in both directions. Additionally, and unlike in PACs, coupling strength among same-frequency couplings was pronounced in both directions with strongest coupling in the low-frequency range (Fig. 2D,E). Examining differences in coupling strength in the two directions more closely (Fig. 2F), AACTs displayed a similar pattern to PACs but showed smaller significant clusters than PACs across the entire frequency spectrum.

**Cross-regional coupling mainly driven by interactions between A1 and higher-order sectors**

To examine cross-regional patterns in coupling across the auditory hierarchy (Fig. 1) more closely, we analyzed differences between top-down and bottom-up coupling strength separately across the six cross-regional pairs (Fig. 3). This revealed that the overall coupling pattern (Fig. 2) was mostly driven by interactions between S1 (A1/ML) and S2-S4 (Fig. 3A–C), rather than by interactions among higher-order sectors (Fig. 3D–F).

More specifically, interactions between S1 (A1/ML) and S2 (R/AL)/ S3 (RTL)/ S4 (RTp) show dominant top-down coupling of low-frequency phase to low- and high-frequency amplitude (Fig. 3A–C), whereas these asymmetries are largely absent among interactions between higher order sectors (Fig. 3D–F). The interaction between S1 and S2 (Fig. 3B) shows overall less widespread top-down coupling strength compared with interactions between S1 and S3/S4. The full pattern (as manifested in Fig. 2) only gains full prominence among cross-regional pairs S1–S4 (Fig. 3C), which also includes predominant bottom-up coupling between low-frequency phase and low-gamma amplitude. Interactions between higher-order sectors (Fig. 3D–F) show fewer differences between top-down and bottom-up coupling, but they do show predominant bottom-up coupling between low-frequency phase and low-gamma amplitude (Fig. 3E,F). A similar trend was observed among AACs, although differences were overall smaller and less widespread compared with PACs (Extended Data Fig. 3-1).

**Top-down and bottom-up cross-frequency signatures generalize to synthetic stimuli**

To examine whether the observed coupling patterns are a general feature of interareal communication, we also analyzed coupling strength in two types of synthetic stimuli derived from the natural vocalizations. EPSs were obtained by leaving the original temporal envelope of the vocalization intact but flattening its temporal content, while SPSs were obtained by leaving the spectral information of the original vocalization intact but flattening its temporal envelope (Fukushima et al., 2014; see Stimuli in Materials and Methods).

EPS and SPS stimuli showed similar bottom-up and top-down coupling patterns to natural stimuli, with pronounced theta-to-broadband coupling strength in both directions (Extended Data Fig. 4-1, Extended Data Fig. 4-4). Examining differences between top-down and bottom-up coupling strength in both types of synthetic stimuli (Fig. 4, Extended Data Fig. 4-1, Extended Data Fig. 4-4), we found that they reflected the same coupling asymmetries across the frequency spectrum as did natural stimuli (Fig. 2). This was true for PAC (Fig. 4A,C) and AAC (Fig. 4B,D) alike. More specifically, both types of synthetic stimuli showed dominant top-down coupling among low-to-low-frequency and low-to-high-gamma interactions, and dominant bottom-up coupling among low-to-low-gamma interactions, as did natural stimuli. Both types of synthetic stimuli also showed similar coupling asymmetries to natural stimuli when examining cross-regional pairs separately (Extended Data Fig. 4-2, Extended Data 4-3, Extended Data 4-5, Extended Data 4-6), with most of the pattern driven by interactions between A1 and higher-order sectors rather than interactions among higher-order sectors, as in natural vocalizations.

**Synthetic stimuli show overall lower bidirectional coupling than natural stimuli**

To examine differences in coupling strength between natural and synthetic stimuli more closely, we compared coupling strength across stimuli separately in the top-down and bottom-up directions (Fig. 5).
EPS stimuli showed overall lower coupling strength than natural stimuli in both directions for both PACs and AACs (Fig. 5). Among PACs more specifically (Fig. 5A,B), natural stimuli showed higher coupling strength in the top-down direction for select frequency pairs in the low-frequency space as well as among low-frequency-to-high-gamma couplings (Fig. 5A). In the bottom-up direction, coupling strength for natural stimuli was enhanced among low-frequency-to-low- and high-gamma couplings (Fig. 5B). In the bottom-up direction, there were also select coupling values in the high-gamma frequency target amplitude space for which coupling was stronger in EPS stimuli.

Among AACs, coupling strength was increased for natural compared with EPS stimuli across the entire frequency spectrum in both directions (Fig. 5E,F). In particular, theta-gamma amplitude–amplitude interactions were significantly lower for EPS stimuli in both directions. Altogether, coupling strength is enhanced for natural stimuli over envelope-preserved but spectrally flat synthetic stimuli (i.e., EPS) in both directions and across the entire frequency spectrum.

SPS stimuli also showed overall lower coupling strength compared with natural stimuli. Among PACs, SPS stimuli showed lower coupling strength for select low-to-low- and low-to-high-frequency couplings in both directions (Fig. 5C,D). In the top-down direction, SPS stimuli showed higher coupling strength among gamma-gamma interactions (Fig. 5D). SPS and natural stimuli displayed similar differences among AACs (Fig. 5G,H). Altogether, coupling strength was mostly increased for natural stimuli compared with SPS stimuli across the entire frequency spectrum in both directions.

Comparing SPS and EPS stimuli directly (Extended Data Fig. 5-1), coupling strength was largely reduced for EPS stimuli across the frequency spectrum in both directions among PACs (Extended Data Fig. 5-1A,B). This was also true among AACs (Extended Data Fig. 5-1C,D), although there were fewer differences here than among PACs, especially in the high-frequency space.

Overall, top-down and bottom-up cross-frequency coupling signatures were preserved among synthetic stimuli, albeit at overall reduced coupling strength.

**Discussion**

We found distinct cross-frequency signatures for top-down and bottom-up information processing in auditory cortex. These patterns were similar across phase–amplitude and amplitude–amplitude couplings, and across stimulus types (albeit at lower coupling strength in synthetic spectrum-flat stimuli). More specifically, we observed dominant top-down coupling among low-to-low- and low-to-high-gamma frequency interactions, and dominant bottom-up coupling among low-to-low-gamma frequency interactions. These patterns were largely preserved across coupling types (PAC and AAC) and across stimulus types (natural and synthetic stimuli), and mostly driven by interactions between A1 and higher-order sectors. Differences in coupling strength across the auditory hierarchy are overall an order of magnitude smaller among AACs than among PACs. Although the observed effects are subtle, they are significant and highly preserved. Altogether, our results suggest that these cross-frequency signatures are a general hallmark of bottom-up and top-down information processing in auditory cortex.

The top-down coupling profile we observed is in accordance with previous results, but we also observed predominant low-frequency-to-low-gamma coupling in the bottom-up direction. Previous studies found that feedforward (or bottom-up) influences were mediated by gamma frequency range rhythms (Bosman et al., 2012; Fontolan et al., 2014; Bastos et al., 2015; Michalareas et al., 2016). However, across each of these studies, there were examples of a broader frequency range being used for bottom-up information transmission. Fontolan et al. (2014) reported higher bottom-up coupling of delta (1–3 Hz) frequency phase in left A1 to high-gamma (80–100 Hz) amplitude in left association auditory cortex using human depth recordings. Bastos et al. (2015) found that theta channels were being used in addition to gamma-frequency channels in feedforward information transmission among primate visual areas. Finally, Michalareas et al. (2016) did not observe stronger alpha–beta frequency range feedback (top-down) modulation in 8 of 21 primate visual cortex area pairs (with 2 pairs showing the opposite pattern), and no stronger gamma frequency range feedforward (bottom-up) modulation in 5 of 21 pairs. We observed predominant bottom-up coupling between low-frequency (theta, alpha, beta) phase (or amplitude) and low-gamma amplitude among phase–amplitude couplings (and amplitude–amplitude couplings, respectively). Differences in the exact frequency ranges reported may be due to regional (visual vs auditory cortex) variation (Canolty et al., 2006; Tort et al., 2009) and species-related variation. Overall, our findings agree with accounts of predominant low-to-high (cross-) frequency interactions top-down. But together with the exceptions above, our findings suggest that low-to-high (cross-) frequency interactions may also play a role in bottom-up processing, pointing to a picture of top-down and bottom-up processing that is more complex than the previously hypothesized “beta-down versus gamma-up.”

Bottom-up PAC can be relevant to stimulus encoding. For example, an oscillation-based theory of speech encoding (Giraud and Poeppel, 2012; Hyafil et al., 2015a) predicts that cortical theta oscillations track the syllabic rhythm of speech and, in turn, reset the spiking of neurons at a gamma frequency level through theta-gamma PAC. This ensures that neural excitability is optimally aligned with incoming speech. Our finding of dominant low-frequency phase to gamma amplitude coupling bottom-up fits within a framework that requires theta–gamma coupling for accurate speech encoding (Hyafil et al., 2015a). Cross-regional theta–gamma PAC as we observed here across A1 and higher-order sectors could be responsible for coordinating information processing along the auditory cortical hierarchy. Our observation that these patterns are consistent across stimulus classes implies that this may be a generalized mechanism of auditory information
processing that is co-opted during speech processing. However, our finding of enhanced low-to-high-frequency phase–amplitude coupling in the top-down direction shows that these couplings also mediate top-down influences on information processing.

Coupling in the top-down direction has been of particular interest in theories of the effect of attention on auditory information processing. Attention is thought to sample stimuli rhythmically, fluctuating at a 4–10 Hz rhythm (Landau and Fries, 2012), and was shown to enhance the processing of degraded speech in the anterior and posterior STS (Wild et al., 2012). Frontal top-down signals in the delta and theta frequency range have been shown to modulate low-frequency oscillations in human auditory cortex, increasing their coupling to continuous speech (Park et al., 2015). The same delta and theta frequency bands have been shown to be more strongly coupled to the gamma-range burst rate within monkey A1 during periods of task entrainment (Lakatos et al., 2016). In visual cortex, delta frequency oscillations have been shown to entrain with the rhythm of stimuli in the attended stream (Lakatos et al., 2008; Schroeder et al., 2010), and top-down beta-band modulation was found to be enhanced with attention (Bastos et al., 2015). Increased power in low-frequency range oscillations may also, in turn, enhance bottom-up rhythms through top-down (cross-frequency) coupling (Lee et al., 2013; Bressler and Richter, 2015); indeed, top-down beta-band activity was found to be maximally correlated with bottom-up gamma-band activity when top-down preceded bottom-up activity (Richter et al., 2017). In addition, top-down couplings may also reflect the modulatory activity of predictive processes on auditory/speech perception and processing (Poeppe et al., 2008; Wild et al., 2010; Davis et al., 2011; Arnal and Giraud, 2012; Gagnepain et al., 2012; Chao et al., 2018). Our findings of dominant low-to-low- and low-to-high-frequency coupling top-down are consistent with these accounts, and could well serve to enhance the engagement of the animal in the task (Park et al., 2015; Lakatos et al., 2016). In addition, the top-down coupling signal we observed may also be carrying predictive information, being consistent with frequency ranges that have been shown to carry prediction update signals (Chao et al., 2018). We cannot dissociate between attentive or predictive signals here, however.

We found PAC and AAC to offer similar accounts of information processing. This could be due to a high degree of shared informational content between the two measures; it remains unclear how these measures differ, and how AAC could function mechanistically in the brain. To the degree that PAC and AAC offer complementary, nonoverlapping information, it is possible that top-down influences are exerted more broadly through various CFC types, including both PAC and AAC, and others such as phase synchronization (Fries, 2005).

We observed the same pattern of results in synthetic stimuli. The overall lower coupling strength we observed in synthetic stimuli could be a consequence of the coding differences that were previously observed between synthetic and intact stimuli (Fukushima et al., 2014), or they could be driving these coding differences. Lower coupling strength in synthetic stimuli could also be a result of the modulatory effect of attention being lower for synthetic compared with natural stimuli, perhaps because animals pay less attention to unnatural, ecologically irrelevant stimuli. The observation of higher coupling strength in spectrally rich (SPS) over spectrally poor (EPS) stimuli, envelope-intact stimuli suggest enhanced encoding and processing of information-rich stimuli. This is also consistent with our previous results, which showed that information about spectrally rich stimuli was better maintained than information about spectrally flat stimuli, at least across the first two sectors (S1, S2) of the auditory hierarchy. In the highest-order auditory sector (S4), decoding performance showed differences across a broader frequency range among VOCs and EPSs than among VOCs and SPSs.

The fact that we find the same coupling patterns across natural and synthetic stimuli points to the existence of conserved cross-frequency signatures of bottom-up and top-down information processing in auditory cortex. These findings have important implications for theories of neural processing such as predictive coding. In theoretical implementations of predictive coding, the presence of nonlinearities in going from predictions to prediction errors may induce cross-frequency interactions in the top-down direction. These should be less prevalent if not absent in the bottom-up direction as prediction errors are thought to accumulate linearly. Our finding of predominant low-frequency to low-gamma cross-frequency coupling bottom-up suggests that nonlinearities may also play a role in bottom-up processing, allowing the modulation of gamma power in higher-order areas by the phase of low-frequency rhythms in lower-order areas, for example. Overall though, our finding of widespread predominant cross-frequency interactions in the top-down direction is in agreement with the predictive coding framework.

Many of the theoretical propositions for spectral asymmetries in forward and backwards message passing in predictive coding do not consider the role of attention, however. In predictive coding, attention is mediated by optimizing the precision of ascending prediction errors in a context-sensitive fashion. Physiologically, this corresponds to changing the gain of lower levels (usually considered to involve fast-spiking inhibitory interneurons and some form of coherence or synchronous gain). Crucially, the effect of attention adds an extra level of nonlinearity that could be mediated by backward connections, and, implicitly, cross-frequency coupling.

We used a method based on the canonical correlation framework that allowed us to estimate directed cross-frequency coupling strength in a computationally efficient framework. Many previous studies and methods to estimate cross-frequency coupling do not obtain directional estimates (Bruns and Eckhorn, 2004; Canolty et al., 2006; Cohen, 2008; Tort et al., 2010; Voytek et al., 2013; van Wijk et al., 2015), and thus do not allow for feedforward and feedback components to be teased apart. Studies that do take direction into account (Bastos et al., 2012, 2015; Fontolan et al., 2014; Michalareas et al., 2016) use
separate measures to look at coupling strength (e.g., through coherence) and directional information flow (e.g., through Granger causality), and do not take cross-frequency interactions into account. The dynamic causal modeling (DCM) framework (Friston et al., 2003, 2017; Chen et al., 2008) does allow for the computation of directed coupling across frequencies (Furl et al., 2014). Practically however, to enhance computational efficiency, current implementations of DCM reduce the dimensionality of the data by applying a singular value decomposition to the data from the source region X, and then projecting Y into this reduced space to compute coupling estimates (Chen et al., 2008). The variance preserving dimensions in the source and target space are not necessarily the dimensions that maximize correlations between source and target space, in which case interactions may be missed. In contrast, the canonical correlation framework used here has the advantage of finding low-dimensional representations that maximize coupling between the source (X) and target (Y) data. A similar approach has been used to obtain estimates of cross-frequency interactions in MEG data (Sato et al., 2010), however, without the directional Granger causal component included here.

In summary, we revealed distinct cross-frequency signatures of top-down and bottom-up information processing. These signatures are largely preserved across coupling types and stimulus types, suggesting that they are a general hallmark of information processing in auditory cortex. Our finding of prominent low-to-high-frequency coupling top-down extends the current view of low-to-high-frequency interactions in auditory cortex, which primarily emphasized their involvement in bottom-up processes. We used a method that allows for the computationally efficient estimation of directed cross-frequency interactions. Altogether, this extends our understanding of how information is propagated up and down the cortical hierarchy, with significant implications for theories of neural information processing such predictive coding.

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