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Task-modulated “what” and “where” pathways in human auditory cortex

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Human neuroimaging studies suggest that localization and identification of relevant auditory objects are accomplished via parallel parietal-to-lateral-prefrontal “where” and anterior-temporal-to-inferior-frontal “what” pathways, respectively. Using combined hemodynamic (functional MRI) and electromagnetic (magnetoencephalography) measurements, we investigated whether such dual pathways exist already in the human nonprimary auditory cortex, as suggested by animal models, and whether selective attention facilitates sound localization and identification by modulating these pathways in a feature-specific fashion. We found a double dissociation in response adaptation to sound pairs with phonetic vs. spatial sound changes, demonstrating that the human nonprimary auditory cortex indeed processes speech-sound identity and location in parallel anterior “what” (in anterolateral Heschl’s gyrus, anterior superior temporal gyrus, and posterior planum polare) and posterior “where” (in planum temporale and posterior superior temporal gyrus) pathways as early as ≈ 70 – 150 ms from stimulus onset. Our data further show that the “where” pathway is activated ≈ 30 ms earlier than the “what” pathway, possibly enabling the brain to use top-down spatial information in auditory object perception. Notably, selectively attending to phonetic content modulated response adaptation in the “what” pathway, whereas attending to sound location produced analogous effects in the “where” pathway. This finding suggests that selective-attention effects are feature-specific in the human nonprimary auditory cortex and that they arise from enhanced tuning of receptive fields of task-relevant neuronal populations.

functional MRI | magnetoencephalography | selective attention | spatiotemporal brain imaging

One’s ability to perceive auditory objects in everyday acoustic environments depends on localization and identification of relevant sounds. Primate models (1, 2) suggest that this task is accomplished via parallel anterolateral “what” and caudolateral “where” nonprimary auditory cortex streams, resembling the functional subdivisions of the visual system (3, 4). Human neuropsychological (5–8) and neuroimaging (9–20) studies have consistently shown anterior-temporal-to-inferior frontal “what” and parietal-to-lateral-prefrontal “where” auditory pathways, but whether such dual pathways exist also in the human nonprimary auditory cortex has remained a more controversial issue. Although there is accumulating evidence that nonprimary auditory cortex regions posterior to the Heschl’s gyrus (HG) are involved in spatial processing (21–26) and that areas anterior to HG process sound-identity cues such as speech (27, 28) and pitch (29), the posterior nonprimary auditory cortex areas have been reported to respond strongly to phonetic stimuli as well (30, 31). This observation has raised hypotheses alternative to the dual pathway model, suggesting that the posterior nonprimary auditory cortex processes rapid spectrotemporal changes (30, 32), common to both speech sounds and sound motion/location cues (32), and that the anterior pathway concentrates on invariant sound features (32). Evidence of a double dissociation between

processing of phonetic vs. spatial features is thus needed to determine whether the dual pathway model is valid for anterior vs. posterior human nonprimary auditory cortex areas.

Selective attention is known to support both sound localization and recognition, but it is unclear how representations of auditory space and identity are top-down modulated in the human auditory cortex. Overall enhancement of human auditory cortex activity by selective attention has been verified by functional MRI (fMRI) (14, 33–37), positron emission tomography (38–40), electroencephalography (41), and magnetoencephalography (MEG) (42) studies, and recent fMRI results further implied that these effects mainly occur in the nonprimary auditory areas (37). Dichotic listening studies of spatial attention suggest signal enhancements in auditory areas contralateral to the attended ear (38, 42, 43). However, although distinct prefrontal and parietal activations to attentional processing of “what” vs. “where” auditory information have been consistently reported (9, 13–16), previous positron emission tomography and fMRI studies have failed to find evidence for feature-specific attentional effects for sound identity and location in the auditory cortex (37, 39). Whereas detailed neuronal mechanisms, including amplification of relevant object representations (44) and enhanced selectiveness for attended stimuli (45), have been characterized in the human visual cortices (see also refs. 4 and 46), it remains unclear how selective attention affects neuronal representations of sounds to facilitate auditory perception. Recent animal models suggested task-dependent modulation of spectrotemporal receptive fields in the auditory cortex (47), but such effects have so far not been shown in humans.

Functional neuroimaging of human auditory cortex is challenging because of its relatively small size. However, recent studies suggest that rapid attenuation of neuronal activity after two or more successive auditory stimuli, termed “neuronal adaptation” (48, 49), can be used to probe the selectivity of auditory cortex neurons for a particular type of information (50, 51), helping circumvent limited resolution of noninvasive neuroimaging methods. To measure stimulus-feature tuning properties of the human auditory cortex, one can vary the physical similarity between a pair of sounds (Adaptor and Probe) and measure the adaptation of a response as a function of the difference between the sounds (51). Specifically, a Probe sound

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Abbreviations: ECD, equivalent-current dipole; fMRI, functional MRI; HG, Heschl’s gyrus; MEG, magnetoencephalography; ROI, region of interest; STG, superior temporal gyrus; PT, planum temporale; PP, planum polare.

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produces a strongly attenuated, i.e., adapted response after a physically identical Adaptor. Furthermore, if Adaptor and Probe differ in one sound attribute, adaptation is more prominent in a neuronal population broadly tuned (i.e., nonselective) than in a population sharply tuned (i.e., selective) to that attribute (50). Interestingly, visual selective attention appears to modulate adaptation of fMRI signals (45), suggesting that the phenomenon of adaptation can also be used to probe the neural basis of selective attention.

Neuronal adaptation in the human auditory cortex can be measured with the N1 response, peaking at ≈ 100 ms after stimulus onset in trial-averaged MEG. The N1 “adaptation cycle” is closely coupled with the cellular-level “very long” adaptation time constant of 1–10 s, purportedly necessary for representing temporally distributed auditory objects (49). Importantly, the N1 response has separate anterior and posterior auditory cortex sources (50, 52, 53). These sources adapt differently to sound-feature changes, the anterior source showing sharp and the posterior source showing broad frequency tuning (50). Given the necessity of fine frequency analysis for sound-object processing and our dependency on broadband spectral cues in auditory localization (54), these two N1 sources could reflect the “what” and “where” pathways of the human nonprimary auditory cortex.

We hypothesized that the human nonprimary auditory cortex includes parallel anterior “what” and posterior “where” pathways (1, 2), and that selective attention to sound identity vs. location modulates these pathways in a task-dependent fashion. We further hypothesized that these effects would be revealed by differential adaptation in the putative “what” and “where” auditory streams ≈ 100 ms after stimulus (50), as measured by a spatiotemporal brain imaging approach that combines the temporal resolution (milliseconds) of MEG with the spatial resolution (millimeters) of fMRI.

Results

Fig. 1 shows the stimulus/task paradigm used in the fMRI and MEG measurements. The reaction times (mean $\pm$ SEM) were not significantly different between the Attend Location (740 ± 75 ms) and Attend Phoneme (706 ± 70 ms) conditions, but the hit rate was higher [$F(1,8) = 28.8$, $P < 0.01$] in the Attend Phoneme ($92 \pm 3\%$) than Attend Location ($83 \pm 3\%$) condition. The false alarm rate to “sham targets” (i.e., a phonetic target during Attend Location condition and vice versa; $P = 0.12$) was slightly higher [$F(1,8) = 9.7$, $P < 0.05$] in Attend Location ($5 \pm 1\%$) than Attend Phoneme ($1 \pm 1\%$) condition. During the Ignore condition, the rate of false responses ($0.6 \pm 0.3\%$) was not significantly different from 0%.

Differential Adaptation to Phonetic vs. Spatial Information in Auditory Cortex. To test our first hypothesis of differential adaptation to “what” and “where” information in the anterior vs. posterior auditory cortex, we compared brain responses to Probes preceded by identical, phonetically different, or spatially different Adaptors (see Fig. 1c for a schematic illustration of response adaptation). In support of our hypothesis, the fMRI-weighted MEG source estimates (Figs. 2 and 3) showed that, although the auditory cortex activity to Adaptors was similar in all conditions, for Probe responses the main effect of stimulus change [$F(2,16) = 11.5$, $P < 0.01$] and the interaction of stimulus change and source location [$F(2,16) = 15.9$, $P < 0.001$] were significant. (The ANOVA effects of hemisphere laterality were nonsignificant.) Specifically, regions posterior to HG, including the planum temporale (PT) and posterior aspects of the superior temporal gyrus (STG), responded more strongly to Probes preceded by spatially different (but phonetically similar) Adaptors, in comparison to those preceded by phonetically different or identical Adaptors [right hemisphere: $F(1,8) = 16.3$, $P < 0.01$;

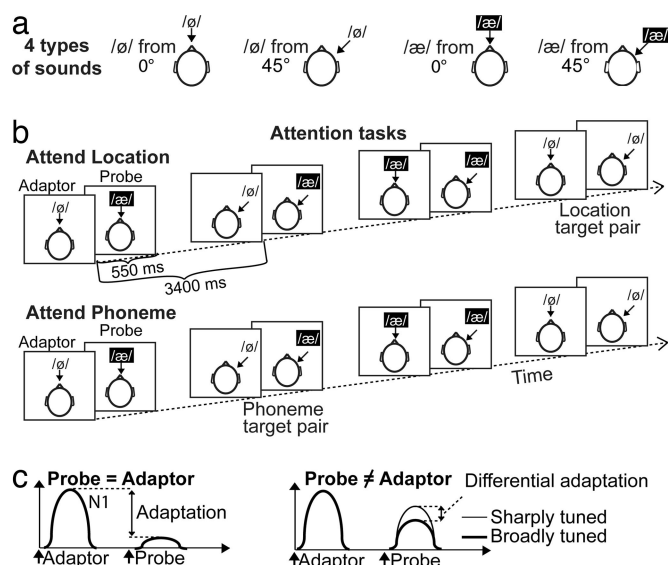


Fig. 1. Schematic illustration of the experimental paradigm and phenomenon of neuronal adaptation. (a) Sound stimuli. Pairs of Finnish vowels /æ/ and /o/ were presented from straight ahead or 45° to the right. (b) Sound sequence and tasks. Vowel pairs (i.e., Adaptor followed by Probe) were spatially discordant, phonetically discordant, or identical. During consecutive blocks, subjects were instructed to attend to either spatial (Attend Location) or phonetic (Attend Phoneme) similarities between successive sound pairs or to ignore the presented stimuli (Ignore condition not shown here). In the Attend Location condition, the subject responded to sound pairs that matched the spatial pattern of the preceding sound pair (same directions in same order), irrespective of the phonetic content. In the Attend Phoneme condition, the targets were, in turn, sound pairs being phonetically similar to the preceding sound pair (same phonemes in same order), irrespective of the spatial content. (c) Schematic illustration of response adaptation. A Probe sound preceded by an identical Adaptor produces a strongly adapted response. Adaptation is weakest when Probe differs from Adaptor in a feature to which the neuronal population is sharply tuned.

left hemisphere: $F(1,8) = 20.2$, $P < 0.01$]. That is, activity in the posterior nonprimary auditory cortex was adapted less after location than phoneme changes (Figs. 2 and 3), suggesting that this region is more sharply tuned to spatial than phonetic information. In contrast, the anterior nonprimary auditory cortex regions, encompassing the anterolateral HG and parts of the anterior STG and the planum polare (PP), exhibited stronger activity when Adaptor and Probe differed phonetically than spatially, suggesting sharper phoneme tuning within these areas [right hemisphere: $F(1,8) = 11.2$, $P < 0.05$; left hemisphere: $F(1,8) = 20.4$, $P < 0.01$] (Figs. 2 and 3).

Selective Attention and Phoneme vs. Location Processing in Auditory Cortex. Our second hypothesis was that selective attention to phonetic vs. spatial features differentially modulates adaptation in the anterior “what” vs. posterior “where” auditory pathways, respectively. To quantify this, we modeled the anterior and posterior N1 sources (50, 53) as equivalent-current dipoles (ECD), which are less sensitive to crosstalk across different sources (e.g., anterior and posterior N1) than distributed estimates (55). Consistent with previous observations (50, 52, 53), the Adaptor N1 responses were explained by an earlier (left hemisphere, 92 ± 4 ms; right hemisphere, 89 ± 3 ms) posterior and a later (left hemisphere, 118 ± 4 ms; right hemisphere, 120 ± 5 ms) anterior ECD (see *Supporting Results* in *Supporting Text*, which is published as supporting information on the PNAS web site). Fig. 4 shows that these ECD loci, with significantly different origins along the anterior–posterior y axis in both hemispheres [$F(1,8) = 164.3$, $P < 0.001$], were in agreement with

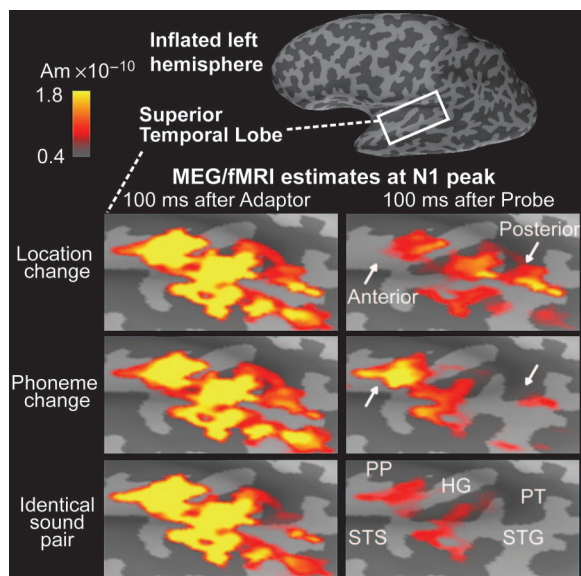


Fig. 2. Differential adaptation to phonemes vs. sound locations in nonprimary auditory cortex. Cortical fMRI-weighted MEG source estimates are shown in a representative subject at the N1 peak latency. The auditory cortex areas activated by the Adaptor (the first stimulus of the pair) are identical across the conditions, but specific adaptation-induced differences in activity patterns elicited by Probes (the second stimulus of the pair) are observed: The posterior activity is strongest (i.e., least adapted) when Adaptor and Probe differ spatially, and the anterior activity is strongest when Adaptor and Probe differ phonetically. The results were similar in the right hemisphere of this subject (not shown here). STS, superior temporal sulcus.

the fMRI-weighted MEG source estimates of N1 activity (Figs. 2 and 3). These ECD sources were used to model attentional modulation of feature-specific adaptation in the anterior and posterior nonprimary auditory cortex.

The ECD analysis corroborated the fMRI-weighted MEG results, suggesting differential adaptation to “what” vs. “where” information in the anterior and posterior regions of nonprimary auditory cortex (see *Supporting Results*). Furthermore, supporting our attentional hypothesis, there was a significant interaction [$F(2,16) = 4.4$, $P < 0.05$] among the type of attention (Attend Location vs. Attend Phoneme vs. Ignore), source location (anterior vs. posterior nonprimary auditory cortex), and stimulus

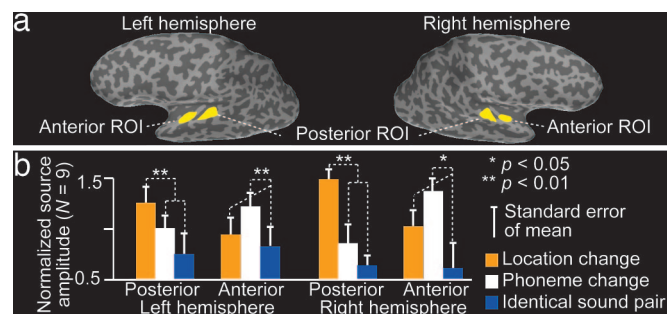


Fig. 3. ROI analysis of fMRI-weighted MEG source estimates to Probe sounds, showing differential adaptation after location vs. phoneme changes in the posterior and anterior auditory cortex, respectively. (a) The ROI locations in a representative subject are represented on the inflated cortex. (b) The ROI group average results suggest sharper spatial tuning in the posterior and sharper phoneme tuning in the anterior auditory cortex regions. The statistical significances refer to *a priori* Helmert contrast between the condition of interest (Location Change in the posterior and Phoneme Change in the anterior ROI) vs. other conditions.

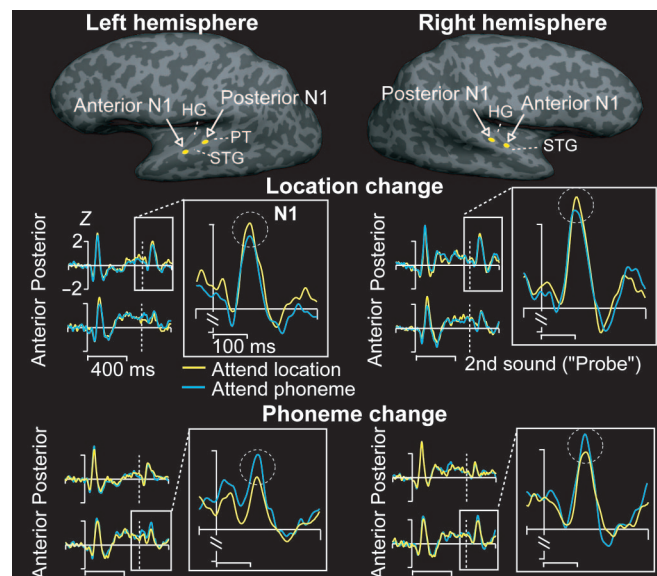


Fig. 4. Group-average ECD results of the two N1 subcomponents (50, 52, 53) showing the attentional modulation of the anterior and posterior auditory cortex activity to Probes following spatially or phonetically different Adaptors. The ECD locations (Top), group-averaged in a spherical standard space (60), are displayed on the inflated brain hemispheres of one subject. (Middle and Bottom) The source waveforms were amplitude-normalized within each subject before calculating the group averages (shown as Z-score values). This procedure retains the within-subject amplitude proportions, and each subject contributes equally to the group mean. Insets demonstrate the responses at -50 – 400 ms around Probes from sources showing peak attention effects. The N1 response amplitudes to Probes (encircled) are modulated task-dependently. The posterior N1 activity to Probes following spatially different Adaptors is enhanced by spatial attention. The anterior N1 activity to Probes following phonetically different Adaptors is enhanced by phonetic attention.

order (Adaptor vs. Probe) (Figs. 4 and 5). Thus, selective attention modulated feature adaptation of anterior and posterior N1 responses to Probes in a task-specific fashion, without affecting the responses to Adaptors themselves. Based on *a priori* comparisons of means, the right-hemispheric posterior N1 response to Probes, when preceded by spatially different Adaptors, was significantly stronger in Attend Location vs. other conditions [$F(1,8) = 30.7$, $P < 0.01$]. That is, spatial attention reduced response adaptation to sound pairs with location changes in the putative “where” pathway in the right hemisphere. When phonetic attributes were attended, the anterior N1 activation to

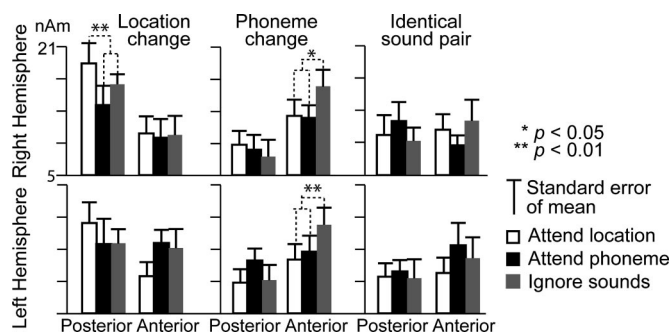


Fig. 5. Group-average selective attention effects in the right (Upper) and left (Lower) auditory cortices in ECD estimates. The response amplitudes to Probes are modulated task-dependently: The posterior N1 activity is enhanced by spatial attention, and the anterior N1 activity is enhanced by phonetic attention. The figure also shows the differential adaptation in the anterior and posterior N1 sources for phonetic vs. spatial information, respectively.

Although the ECD locations approximate the center of gravity of underlying neural activity (55), this approach is less sensitive to crosstalk (59) across different sources than distributed estimates (55), thus offering a robust model for contrasting attention-dependent modulations of the anterior and posterior auditory cortex N1 sources.

The N1 signals recorded from a subset of gradiometer channels (on average 40 per hemisphere) covering the left and right temporal lobes were used in the ECD modeling (55) (see *Supporting Methods*). The posterior N1 was fitted at the ascending phase (≈ 90 ms of the sound onset) and the anterior N1 was fitted at the descending phase (≈ 120 ms) of the N1 response to Adaptors (the goodness-of-fit was $>80\%$ for each ECD fitted). The resulting posterior and anterior ECDs (see *Supporting Results*) were then entered into a time-varying multidipole model to explain the recorded MEG responses. One multi-ECD estimate per hemisphere (based on the same channel selection across all conditions) was used to model each attentional condition in each subject. The attentional effects were tested by using an ANOVA with *a priori* contrasts (with Greenhouse–Geisser correction), including the following factors: serial position of stimuli (Adaptor vs. Probe), hemisphere, type of attention (Attend Location vs. Attend Phoneme vs. Ignore), dipole location (anterior vs. posterior), and type of stimulus change within a pair.

fMRI Analysis. After preprocessing, the fMRI time series were analyzed by using a general linear model (see *Supporting Methods*). Each subject's functional volumes were aligned with their anatomical images. The corresponding cortical surface representations were coregistered to a spherical standard space (60) for a surface-based random-effects model of group activations, calculated in addition to the individual activation maps used in the MEG source analysis. Six ROIs per hemisphere (anterior auditory cortex, posterior auditory cortex, posterior parietal, inferior frontal, dorsolateral prefrontal, and premotor) were selected based on the significant group activations (see Fig. 7, which is published as supporting information on the PNAS web site). The ROI activations, constrained by voxels showing significant individual activations ($P < 0.01$), were entered into a random-effects model to compare activations in different task conditions.

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