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***In vivo* functional and myeloarchitectonic mapping of human primary auditory areas**

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Abstract

In contrast to vision, where retinotopic mapping alone can define areal borders, primary auditory areas such as A1 are best delineated by combining *in vivo* tonotopic mapping with post mortem cyto- or myelo-architectonics from the same individual. We combined high-resolution (800 μm) quantitative T₁ mapping with phase-encoded tonotopic methods to map primary auditory areas (A1 and R) within the ‘auditory core’ of human volunteers. We first quantitatively characterize the highly myelinated auditory core in terms of shape, area, cortical depth profile, and position, with our data showing considerable correspondence to post-mortem myeloarchitectonic studies, both in cross-participant averages and in individuals. The core region contains two ‘mirror-image’ tonotopic maps oriented along the same axis as observed in macaque and owl monkey. We suggest that these two maps within the core are the human analogues of primate auditory areas A1 and R. The core occupies a much smaller portion of tonotopically organized cortex on the superior temporal plane and gyrus than is generally supposed. The multi-modal approach to defining the auditory core will facilitate investigations of structure-function relationships, comparative neuroanatomical studies, and promises new biomarkers for diagnosis and clinical studies.

The parcellation of cortex into distinct areas is a long-standing program in neuroscience (Zilles and Amunts, 2010), but one more advanced in vision than in audition. While some early visual areas can be defined solely on the basis of architectonics (e.g., stria in V1) or non-invasive retinotopy (Engel et al., 1994), the borders of primary auditory cortex (A1) cannot easily be defined by anatomy or function alone. Areas A1 and R are two major fields of the ‘auditory core’ (Hackett, 2007), a narrow, keyhole-shaped region on the temporal plane. In *ex vivo* preparations, auditory core shows heavy, layer IIIb/IV-specific staining for cell bodies, myelin, acetylcholinesterase, cytochrome oxidase, and parvalbumin (Hackett, 2011). Although greater myelination in caudal core may correspond to A1 (Hackett et al., 2001), the boundary with R is difficult to define using myeloarchitectonic criteria (Morel et al., 1993).

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A1 and R are challenging to delineate using tonotopic mapping since tonotopy provides only one spatial axis, with no agreed-upon means to define borders perpendicular to isofrequency bands, which span multiple auditory areas (Hackett, 2011). In a small number of invasive experiments in animals, physiological measures (tonotopy, response properties) have been combined with *ex-vivo* histological mapping to define borders (Merzenich and Brugge, 1973; Imig et al., 1977; Pfingst and O'Connor, 1981; Morel and Kaas, 1992; Morel et al., 1993).

Studies of anesthetized macaques show best frequency (BF) progressions across and around auditory core. As seen in Figure 1 (most complete map from Morel et al., 1993, redrawn and contoured), A1 - in the highly myelinated caudomedial core - is characterized by a high-to-low BF progression, from a medial high BF corona descending to a rostrolateral low BF trough, which marks the A1/R border. In the narrower and more lightly myelinated area R in the lateral aspect of core, there is a gentler low-to-mid BF progression moving rostrolaterally.

There are multiple tonotopic maps across the temporal plane, as shown with fMRI in humans (Da Costa et al., 2011; Formisano et al., 2003; Talavage et al., 2004) and macaques (Petkov et al., 2006; Baumann et al., 2010). Whereas the A1/R boundary can be localized using tonotopic gradients (Formisano et al., 2003), the relation between frequency progressions and the position of core is disputed (Humphries et al., 2010). Current probabilistic post-mortem cytoarchitectonic maps of primary auditory areas (Morosan et al., 2001) are too blurry to localize core boundaries.

Anatomical MRI results (Walters et al., 2003; Bridge & Clare, 2006; Sigalovsky et al., 2006; Trampel et al., 2011; Glasser and Van Essen, 2011; Sereno et al., 2012) suggest that areal boundaries can be defined by measuring proxies for myelination. Here, we characterize the strongly myelinated auditory core using a high-resolution bias-free quantitative T_1 mapping protocol which provides estimates of myelination directly comparable across cortical regions or subjects. We combine these myelin maps with detailed tonotopic mapping in the same participants to characterize frequency progressions within and around auditory core.

Materials and Methods

Participants

6 adults (ages 22–55, 3 female) with normal hearing and vision participated in all parts of the study. 3 additional adults (ages 22–30, all male) participated only in the functional imaging part of the study. Experimental protocols were approved by local ethics committees, and all participants gave informed and signed written consent.

Structural imaging

Structural images were acquired on a whole-body Tim Trio system (2.89T, Siemens Healthcare, Erlangen, Germany), with radio-frequency (RF) body transmit and 32-channel receive head coil at the Wellcome Trust Centre for Neuroimaging. 3T scanners such as the Trio will not have as high signal-to-noise as higher field strength (e.g., 7T), but do benefit from greater uniformity of signal (less B_1 field inhomogeneities), smaller susceptibility artifacts (B_0 field inhomogeneities), lower acoustic noise, and less dropout in parts of the temporal lobe due primarily to B_1 field inhomogeneities. As part of the quantitative R_1 mapping protocol, proton density-weighted (PDw) and T_1 -weighted (T_1w) images were acquired using an in-house multi-echo 3D FLASH pulse sequence (Weiskopf et al., 2011): voxel size: $0.8 \times 0.8 \times 0.81 \text{ mm}^3$, FOV = $256 \times 216 \times 194 \text{ mm}$, matrix = $320 \times 270 \times 240$, TR = 23.7 ms, excitation flip angle: 6° (PDw) or 28° (T_1w). Acquisition was speeded up by $2 \times$ GRAPPA parallel imaging in the phase encoding and $6/8$ Partial Fourier in the partition

direction. To improve image quality (maximize SNR and minimize geometric distortion at the same time), 4 gradient echoes were acquired (TE=2.20, 4.75, 7.30, 9.85 ms) with high readout bandwidth after each excitation pulse. Each session consisted of four 10 min 31 sec acquisitions (two PDw and two T₁w) and two shorter scans to estimate B₀ and B₁ inhomogeneities (see below). Quantitative R₁ (=1/T₁) maps were estimated from the PDw and T₁w images according to the model developed by Helms et al., (2008a) including a correction for RF transmit field inhomogeneities (Lutti et al., 2010) and imperfect spoiling (Preibisch and Deichmann, 2009). Recent applications of this method have demonstrated its robustness and accuracy (Helms et al., 2008b; 2009; Weiskopf et al., 2011; Draganski et al., 2011). To correct for the effect of RF transmit inhomogeneities on R₁ maps, maps of the transmit field B₁⁺ were acquired using a 3D EPI spin-echo (SE)/stimulated echo (STE) method (Lutti et al., 2012; FOV = 256 × 192 × 192 mm³, matrix = 64 × 48 × 48, TE_{SE}/TE_{STE} = 39.38/72.62 ms, TR = 500 ms, α varying from 270° to 130° by steps of 10°, acquisition time 3 min 48 s), which was corrected for off-resonance effects using a standard B₀ fieldmap (double gradient echo FLASH, 3 mm isotropic resolution, whole-brain coverage (Lutti et al., 2010)).

Functional imaging

Functional images were acquired on a 1.5T whole-body Tim Avanto System (Siemens Healthcare), at the Birkbeck/UCL Centre for NeuroImaging, with RF body transmit and a 32-channel receive head coil. (Supplementary data for three participants was also acquired using a specialized single-loop coil positioned over the superior temporal gyrus). 1.5T has evolved dramatically over the last years with much more stable hardware and 32-channel head coils leading to 200–300% improvement in BOLD sensitivity in auditory regions (Wiggins et al., 2006); echo-planar imaging at this field strength also has less warping and distortion in auditory regions than higher-field magnets. Echo-planar images (EPI) were acquired with the following parameters: 24 slices, voxel resolution 3.2 × 3.2 × 3.2 mm (matrix size: 64 × 64), flip angle = 90°, bandwidth = 1474 Hz/pixel, TR = 2000ms, TE = 39ms, data acquired with prospective motion correction (Thesen et al., 2000). The voxel resolution is similar to that (~3×3×3mm) used in many recent retinotopy studies of small areas, such as V6, VIP, FEF, IPS1&2, V8, and LOC, e.g. Sereno & Huang (2006); Hagler et al., (2007); Larsson & Heeger (2006), Rajimehr & Tootell (2009); Konen et al., (2011). Individual scans had 262 volumes; to allow longitudinal relaxation to reach equilibrium, 6 initial volumes were discarded from each run (initial 2 not saved by scanner). For each imaging session, a short (3 min) T₁-weighted 3D MPRAGE (88 partitions, voxel resolution 1 × 1 × 2 mm³, flip angle = 7°, TE = 4 ms, TI = 1000 ms, TR = 8.2 ms, mSENSE acceleration = 2×, slab-selective excitation) was acquired with the same orientation and slice block center as the functional data for initial alignment with the high-resolution scans used to reconstruct the subject's cortical surface. For the 3 additional participants who did not take part in the quantitative imaging, we acquired a single high-resolution T₁-weighted MPRAGE for cortical surface reconstruction (1×1×1mm, 176 slices, TR=2730ms, TE=3.57ms, flip angle = 7°).

Auditory stimuli

Both retinotopic (Saygin and Sereno, 2008) and tonotopic (Woods et al., 2009) mapping studies have suggested that targeted attentional demands and increased stimulus complexity can significantly modulate fMRI activation in cortical maps. Thus, we used bandpass-swept complex and engaging non-linguistic vocalizations to map tonotopic regions. Base stimuli were adapted from the Montreal Affective Voices (Belin et al., 2008), a series of recordings of emotional non-verbal vocalizations elicited by actors producing sounds corresponding to a set of eight emotions (a ninth “neutral” emotion was not used in the current study, and only male voices were included). Tokens from this set were randomly selected such that no

two recordings were repeated until the entire set had been presented. These tokens were then spliced together with no intervening pause to form 8-minute, 32-second long passages. Next, these stimuli were amplitude-compressed in Adobe Audition 1.5 with additional manual editing; a bandpass filter was then cycled with a period of 64 seconds over the entire passage with center frequency logarithmically ascending from 150 Hz to 9600 Hz ($Q=2$, expanding to $Q=3$ at tails, where Q is the ratio of center frequency over bandwidth). Descending stimuli were created by reversing the entire waveform before filtering, then re-reversing at the final step. This initial filtering was cleaned using similarly ascending low- and high-pass filters positioned an octave above and below the center frequency. Finally, a dynamic amplitude envelope was imposed over the frequency sweeps to equilibrate perceived loudness in the scanner environment. For presentation with the Sensimetrics headphones, stimuli were passed through a final earbud-specific filter that compensated for the slight frequency peaks and phase offsets induced by the acoustic transfer functions of the earbuds. During scanning, subjects were asked to monitor the stimuli and press a button whenever they heard laughter (laughter stimuli were sparsely and non-periodically distributed through the stimulus train). For an example of a single stimulus sweep, see Audiofile 1.

Additional control stimuli were created to test the generality of the tonotopic maps across different types of sound. Similar bandpass-sweep filtering procedures were used on carefully edited musical excerpts (ABBA or Beatles songs), repetitive male speech samples, or amplitude modulated (16Hz) white noise (similar to Talavage et al., 2004).

Stimulus setups

Each subject was scanned using two different stimulus delivery setups in two to five different sessions. In the first setup, stimuli were delivered by an in-house manufactured electrodynamic, magnetless headphone (Josephs et al., 2009) using an isodynamically driven lightweight membrane either coupled to an ear insert that also acted as a passive attenuator of acoustical scanner noise, or presented directly at the ear. The headphone gives a high amplitude, smooth, and minimally distorted response over a wide bandwidth (approximately 14 kHz). Stimuli were delivered to only the left ear, as the size of the ear defender did not allow for subjects' heads to fit into the compact 32-channel head coil with two headphones. The number of 8 min 32 sec long runs collected per participant using this setup were 4, 8, 8, 8, 8, 12, 12, and 16. Three participants also took part in additional control stimulus conditions (see below), each with four 8'32" runs.

In the second setup, stimuli were delivered binaurally using in-house-safety-enhanced Sensimetrics (Malden, MA, USA) S14 earbuds, with the head centered within the 32-channel coil. NoMoCo (NoMoCo Inc, San Diego, CA, USA), cushions were placed around the head to provide additional passive scanner acoustical noise attenuation and to stabilize head position. All volunteers participated in four 8'32" long runs in this setup. Finally, a subset of 7 participants was scanned on an additional single run of a 8min 32sec block design 'auditory localizer' experiment where 16-second blocks of the tonotopy stimulus (with stimuli across blocks balanced for frequency range) were alternated with 16-second blocks of no stimulation.

Cortical surface reconstruction and sampling of R_1 values within cortical ribbon

Cortical surfaces were reconstructed with FreeSurfer (v5.0.0, Dale et al., 1999) from the aligned (AFNI 3dAllineate, Cox, 2011), hand-inspected average of the two high-resolution T_1 -weighted scans acquired for R_1 mapping. We initially attempted using quantitative R_1 scans for surface reconstruction but experienced localized segmentation failures because some boundaries between the pial surface, the CSF, and the skull have different contrast from FreeSurfer's priors in the segmentation algorithms. (Cortical surfaces for the three

participants who were not scanned using quantitative protocols were reconstructed using the single T_1 -weighted MPRAGE acquired at 1.5T). Each subject's reconstructed cortical surface was inflated to a sphere and registered to an average spherical surface atlas in FreeSurfer using a best-fit sulcal alignment (Fischl et al., 1999).

R_1 data sets were sampled along the normal to each gray/white matter surface vertex in steps of 10% of cortical thickness (thickness estimated in FreeSurfer, Fischl and Dale, 2000) and then smoothed tangentially at each depth with a 4 mm FWHM 2D kernel. The human cortex has deep sulci, but also a complex pattern of concavity and convexity, including many convex regions buried within major sulci. Postmortem studies in humans show that myeloarchitecture varies significantly with local cortical convexity (Annese et al., 2004), where more convex regions are thicker and more myelinated, especially in middle and upper layers. Recently we found that there is a moderately strong correlation between local curvature and R_1 at middle cortical depth fractions as well as a lesser but still significant correlation between cortical thickness and R_1 (Serenio et al., 2012). Thus, for each participant, we used the FreeSurfer estimate of local curvature (smoothed with a < 1 mm FWHM 2D kernel) and cortical thickness as hemisphere-wise linear predictors of R_1 values at each cortical depth to adjust for curvature-dependent changes in myelination as well as potential sampling artifacts due to local imperfections in reconstructed white or pial surfaces. Vertex-wise residuals from this regression were used as 'de-curved' and 'de-thickened' estimates of R_1 values whose units are directly comparable to raw demeaned R_1 values. Cross-subject cortical-surface-based averages of R_1 -derived values were calculated by projecting the values from each subject onto the unit sphere after 1 step (< 1 mm FWHM) of surface-based smoothing (Hagler et al., 2006), and averaging the values at each vertex. For visualization purposes, resulting average values were back-projected onto a single representative subject's entire inflated hemisphere or a resected temporal lobe showing the pial surface, with the latter graphically edited to remove the medial temporal lobe and aspects of the insula that were difficult to remove cleanly in FreeSurfer.

Creation of isofrequency contours from data of Figure 2a from Morel, Garraghty, & Kaas (1993)

These data were chosen for being (to our knowledge) the most complete and representative published tonotopic physiological recordings in a single macaque where auditory core was also characterized cyto- and myeloarchitectonically. Each best frequency (BF) datapoint along the physiological tracks in Figure 2A was entered into a 2D grid using the Draw Dataset plugin in AFNI (Cox, 2011) with a high-resolution jpeg of the original figure as an underlay. The sparse data were trilinearly interpolated onto a regular high-resolution grid using the 'TriScatteredInterp' function in Matlab R2010a (Mathworks, Natick, MA, USA); 'contourf' was used to calculate and draw logarithmically-spaced isofrequency contours over the entire recording area (see Recanzone et al., 1999, for a similar approach). Coronal slice profiles and areal borders were traced directly over the jpeg in Pages '09 (Apple, Cupertino, CA).

Analysis of R_1 regional differences and depth profiles using morphed probabilistic maps

In order to provide an initial assessment of R_1 -based myelination maps, and to examine R_1 cortical depth profiles using an independent set of regions-of-interest (ROIs), we created ROIs based on the 3D cytoarchitectonic probabilistic maps of subdivisions of Brodmann's area 41 (TE1.0, TE1.1, TE1.2) with TE1.0 corresponding to the auditory core (Morosan et al., 2001, pg 695; Hackett et al., 2001). The Morosan et al., (2001) raw probability maps provided in the current AFNI distribution (Cox, 2011) were projected to a FreeSurfer 'fsaverage' brain registered to the Talairach target brain, resampled onto the cortical surface, and thresholded at $p > 0.60$ to create ROI labels. The labels were ~ 2 mm FWHM (5 steps)

surface smoothed with manual removal of isolated marked vertices (due to ‘spillover’ from the 3D to 2D projection within the lateral fissure), then spherically morphed to each subject. The morphed (unthresholded) probability maps are illustrated in Figure 3e.

For each subject, average R_1 values for each ROI (TE1.0, TE1.1, TE1.2) were calculated in 10% steps of cortical thickness fraction (see above); these values were used to create R_1 depth profiles. Planned matched pairs t-tests comparing R_1 over the three ROIs were calculated from the data sampled at 50% of cortical thickness.

Functional analyses

Functional images were motion-corrected (AFNI 3dvolreg, heptic interpolation), and hand-registered (4×4 affine) with the high-resolution average T_1 -weighted volume used to create the cortical surface. Initial registration was carried out with the ‘align’ T_1 -weighted scan (same block center, slice plane direction as EPI scans); this initial registration was applied to the EPI volume, where final registration was finely adjusted by manual blink comparison (with contrast-reversed EPI images) to achieve a more exact overlay. Mapping data were analyzed using Fourier methods with individual and group analysis methods as previously described (Serenó et al., 1995; Sereno and Huang, 2006; Hagler et al., 2007), where voxels preferentially responding to a particular point in the stimulus cycle will show a higher amplitude at the frequency of stimulus cycling than at any other frequency. The phase of the signal, which corresponds to a particular point of the stimulus ramp, is then mapped to the color wheel, while the amplitude of the signal is mapped to the voxel’s color saturation. Runs with downward frequency sweeps were time-reversed and averaged with upward-swept scans in order to compensate for unspecified delays in the BOLD response.

Averaging of phase-encoded mapping data was performed using the methodology developed by Hagler and Sereno (2006) in which the real and imaginary components of the signal with respect to the stimulus ramp are averaged across subjects, preserving any phase information consistent between subjects. This was performed by projecting each participant’s phase-encoded map to the FreeSurfer spherical atlas, performing 1 step of surface-based smoothing (< 1 mm FWHM in 2D), averaging across subjects at each vertex, then painting back onto a single subject’s surface for viewing. Four cross-subject phase-encoded tonotopic averages were computed per hemisphere. Three of the four included only those subjects (N=6) with both functional and R_1 data, including separate averages for each auditory stimulus setup (binaural, left monoaural) as well as a grand average over these two setups. The fourth average drew from independent data collected using different coil, headphone, and stimulus arrangements, and served to test map replicability. The left hemisphere average (N=5) included data from three participants scanned with a 7 cm ring coil (Siemens Double Loop Array) over the left superior temporal gyrus, with monoaural right-ear stimulation using bandpass-swept musical and vocalization stimuli, and two participants scanned with the 32-channel head coil and binaural stimulation using bandpass-swept vocalization stimuli. All data for the right hemisphere average (N=4) were collected using binaural stimulation and the 32-channel coil and bandpass-swept musical, noise, or verbal stimuli. The three subjects without R_1 data contributed to both left and right hemisphere averages; two subjects included in the first three averages contributed different data to the left (S6) and left and right hemisphere (S1) averages.

To show uninterrupted phase data in the same cortical locations in all participants (da Costa et al., 2011), the cross-subject-average block design ‘auditory localizer’ data were used to create an independent functional mask for the tonotopic map data (N=7, 3.2mm 2D FWHM of surface smoothing for sampling each subject’s data to the icosahedral sphere, 4.5mm 2D FWHM when rendering to a representative subject). A single contiguous patch of suprathreshold vertices on the superior temporal plane was extracted for each hemisphere

(vertexwise $p < 0.01$, with 3054 (right) and 3431 (left) connected vertices), then projected via spherical registration to each subject. Within this mask, group average and individual subject tonotopic phase-encoded data are displayed using a pseudocolor scale representing the systematic phase offsets in hemodynamic response corresponding to the phase of the auditory sweeps (2.2mm 2D FWHM surface smoothing was applied to tonotopic phase maps to gently even out voxel edges).

Auditory frequency gradient directions were drawn by imposing isofrequency contours over the color phase map (20 steps per complete cycle), then overlaying arrows perpendicular to each contour line in and around core, with arrow length spanning $1/40^{\text{th}}$ of a complete phase cycle. The arrow direction thus represents the gradient direction while its length is proportional to the (logarithmic) tonotopic magnification factor (see below).

RESULTS

R_1 as a function of cortical depth in probabilistically defined core and adjacent regions

As noted above, primate auditory core can be differentiated from surrounding belt regions by the presence of heavy myelination in deep and middle cortical depths with particularly heavy deep and middle layer staining within medial core (Hackett et al., 2001). An increase in overall myelination as well as expansion of layers 5 and 6 along the crown of human Heschl's gyrus has also been noted (Wallace et al., 2002). Our *in vivo* human MRI samples much more coarsely than histology (800 μm versus $\sim 20 \mu\text{m}$), although we benefit from a relative super-resolution effect, in that at each vertex (whose position on the reconstructed cortical surface is at sub-voxel resolution), multiple voxels are sampled using linear interpolation. Nevertheless, we expected R_1 , which is proportional to the myelination level, to start high in deeper layers, drop to a moderate plateau at middle cortical depths, and then drop more steeply at superficial depth fractions. Given the continuously heavy myelination across middle layers of core ("astriate" profile, Hackett et al., 2001), the greatest R_1 difference between core and adjacent regions should be observed at middle cortical depths.

In order to test these predictions, we first created depth profiles of average R_1 in auditory ROIs likely to correspond to auditory core proper (TE1.0) and to adjacent regions lateral and medial to core (TE1.1 and TE1.2) in the published postmortem probabilistic maps of Brodmann's area 41 (Morosan et al., 2001). The cross-subject cross-hemisphere average R_1 depth profiles for TE1.0 in Figure 2a show the predicted steep drop in R_1 at the gray matter/white matter border, followed by a tilted plateau in R_1 at middle cortical depths and a second steep drop approaching the gray matter/pial boundary. The R_1 profiles for the medial (TE1.1) and lateral (TE1.2) regions are smoother and almost entirely overlapping, and clearly undershoot the TE1.0 'core' ROI profile at middle sampling depths as expected.

We verified these inter-ROI differences with hemisphere-wise paired t-tests comparing R_1 values sampled halfway through cortex (0.5). Here, R_1 in TE1.0 was greater than in corresponding TE1.1 (left hemisphere (lh), $t(5)=5.90$, $p<0.002$; right hemisphere (rh), $t(5)=5.60$, $p<0.0025$) and than in TE1.2 (lh, $t(5)=4.14$, $p<0.0090$, rh, $t(5)=-12.51$, $p<0.0001$), with no significant difference between TE1.1 and TE1.2 ($p > 0.4$).

Shape, position, and size of auditory core

Myeloarchitectonic studies (see Introduction) show that middle layer myelination of auditory core has a characteristic keyhole shape, with its long axis oriented diagonally on the superior temporal plane. In humans, it tends to lie along Heschl's gyrus (with darkest staining on the gyral crown - Wallace et al., 2002) but sometimes moves into the adjacent sulcus, with the broader rounder medial segment tapering to a thinner anterolateral finger (Hackett et al., 2001).

As an initial means of visualizing potential R_1 differences expected to relate the topography of increased myelin content, we inspected the spherical-surface-based cross-subject average maps of R_1 values sampled from voxels at middle cortical depths (depth fraction 0.5). When these average maps were thresholded at the mean R_1 value from the mid-probability TE1.0 ROI (0.66 sec^{-1}), a clear hyperintensity appeared in both hemispheres, spanning a thin strip of cortex ($\sim 1.9 \text{ cm} \times \sim 0.7 \text{ cm}$) running roughly along the posteromedial two thirds of Heschl's gyrus (see Figures 2b and 3a). The only other hyperintensities visible at this threshold were along the pre- and post-central gyri, corresponding to the strongly myelinated primary motor and somatosensory cortices (Glasser and Van Essen, 2011; Sereno et al., 2012). R_1 values within this presumptive core region (see Figure 4 & R_1 contours in Figure 6) were highest within a semicircular patch at its posteromedial aspect, with decreasing R_1 moving anterolaterally. A second smaller disconnected hyperintensity appeared further laterally along the superior temporal gyrus.

As noted above, local R_1 values, cortical myelin density, and laminar profile are known to vary with cortical thickness and local cortical curvature (Annese et al., 2004; Sereno et al., 2012), and thus may affect the R_1 -based localization of auditory core. As a validation measure, we compared the raw average R_1 maps to a cross-subject average of thickness-and R_1 values after regressing out the effects of curvature (see Methods and Figure 3d). This resulted in a picture of the keyhole-shaped maximum almost indistinguishable from that found with raw R_1 values. As an additional validation measure, comparison of cross-subject R_1 averages based on the two separate scan-rescan datasets also showed very close correspondence of auditory core localization (Figure 3b,c). A hyperintense R_1 strip over posteromedial Heschl's gyrus -- the presumptive auditory core -- could be easily identified in each participant, in 11 of 12 hemispheres (see Figure 4, and R_1 contours in Figure 6). As seen in Figure 4, there are also additional regions of moderately high R_1 values in individual participants more laterally and posteriorly to core; we speculate that these may correspond to area STA and potentially PA as identified in Wallace et al. (2002), but a conclusive homology will require directly comparable post mortem data.

Tonotopic maps and frequency gradients within and around the core region

Combined average human R_1 and tonotopic maps and gradients (Figure 5)—

The relationship between average tonotopic maps and average R_1 maps (sampled at 0.5 cortical depth fraction) from both hemispheres was similar to that seen in individual experiments in non-human primates (see inset, Figure 5). In both hemispheres, the posterior-medial end of presumptive core lies at the middle of an arch-shaped high frequency region (red). Moving anterolaterally along the long axis of the core, there is a steep high-to-low frequency progression (red->blue->green). Beyond the anterolateral end of the core, there is a large low frequency region (green), which progresses back to high frequencies (red). On the opposite anterior-medial edge of the core, selectivity progresses steeply back to high frequencies (red). Finally, at the anterior-most and posterior-most positions along the superior temporal gyrus, the characteristic frequencies descend once again (red->blue->green).

The arrangement of tonotopic maps can also be visualized by looking at the local frequency gradient direction (steepest uphill direction in frequency preference at a point; that is, ∇f with respect to cortical position). At each point, the gradient direction is perpendicular to the local isofrequency contour. The gradient direction has been drawn at a series of points along each isofrequency contour (iso-response-phase contour). The length of each vector is proportional to the scalar cortical magnification for logarithmic frequency (the reciprocal of the gradient vector amplitude). Thus, regions of slowest change in frequency sensitivity (smallest gradient amplitude) will have the longest arrows.

As noted above, the anterolateral part of the core lies within an elongated low-frequency trough in both hemispheres. Also in both hemispheres, the medial end of that low-frequency trough is located within region with the highest myelination. From this low-frequency trough, characteristic frequencies (CFs) rise in all directions. Along the anterolateral edge, there is a low-to-medium frequency progression, with higher CFs more lateral and anterior, as well as posterior. There is another progression from the low CF trough rising anteromedially, reaching a high CF maximum just anteromedial to the core. There is a third progression moving posteriorly from the middle of core that is interrupted by a possible discontinuity, where there is an extremely rapid progression through middle frequencies that ends in a high frequency maximum slightly posterior to core. Finally, there is some indication of a posteromedial progression within core in a direction parallel to the long edge that extends into higher frequency CFs, reaching a CF maximum posteromedially. It is interesting that CFs continue to rise for a short distance beyond the posteromedial end of the core (about 3 mm - see results for individual subjects below).

The pattern of tonotopic fields and progressions within and around core was consistent over the two different experimental setups (Figure 5 top and middle insets), although the stimulation (monaural vs. binaural) and head position in the head coil (shifted to right vs. centered) were different. In the right hemisphere, where auditory input was more closely matched, the average pattern of tonotopic maps over core is highly conserved over both scan types. In the left hemisphere (ipsilateral stimulation), we also see similar tonotopic mapping, but with a shallower transition to higher characteristic frequencies in and around the posteromedial cap of core. These tonotopic maps were also similar to cross-subject averages from data collected using different surface coil setups as well as individuals for whom R_1 data were not acquired (Figure 5, bottom insets). There was some divergence across maps (particularly medially), whereas the tonotopic progressions within and around the presumptive core region were very similar over all averages. The general layout of the tonotopic maps - where high CFs predominate along the medial superior temporal plane, with two to three high-CF 'fingers' extending laterally, and interdigitate with more lateral lower-CF regions extending medially - particularly paralleled the results of Formisano et al (2003, Figure 5), and Humphries et al (2010, Figures 5 & 6).

Individual R_1 and tonotopic maps—Since group average tonotopic map could potentially obscure individual differences, we also assessed individual maps relative to the position of auditory core (Fig. 6) and describe here four qualitative features of the average map (and of macaque monkey maps) that can be found in most individual subjects. A fifth feature involving correlations between tonotopy and R_1 is actually more obvious in individuals. For each feature, we list the number of occurrences by hemisphere (N.B.: left hemisphere results are less robust because of fewer runs and less optimal receive coil positioning; also, S4 was excluded on myeloarchitectonic grounds for showing no clear core). The five features are: (1) a corona of higher characteristic frequencies around the medial cap of core (rh N=5/6, lh N=4/5), (2) an elongated high-frequency patch adjoining the anteromedial edge of core, transitioning to lower frequencies moving anteriorly (rh N=6/6, lh N=5/5), (3) a mid-core trough of lowest characteristic frequencies (rh N=6/6, lh N=5/5), (4) a high-to-medium-to-low progression lateral to the posterior two thirds of the core (rh N=6/6, lh N=4/5), (5) a transition from ~2k-1k~0.5k characteristic frequency running from the anteromedial edge posterolaterally, with the ~0.5k characteristic frequency region in the highest R_1 region (highest 2–3 contour lines in each participant) (rh N=5/6, lh N=4/5).

It is also notable that, unlike the group average tonotopic map, the posteromedial aspect of core reaches into regions with the highest CF in individual maps; in addition, several participants showed a more extensive low-to-higher-frequency progressing anterolaterally to

the tip of core (as has been reported for some macaques, e.g., case 91–13 of Morel et al., 1993). In order to assess the stability of an individual's tonotopic maps, we compared repeated sessions (4 runs each) on one individual (see Figure 7), and found very similar patterns of results for the three sessions using bandpass-swept vocalizations and one session with bandpass-swept music; the session using the same experimental setup but with bandpass-swept amplitude-modulated white noise showed similar but much weaker maps. These results were in keeping with the robust maps shown in previous human tonotopy studies (e.g., Formisano et al., 2003; Talavage et al., 2004; Woods et al., 2009, 2011), in particular the consistency over individual runs and sessions (Humphries et al., 2010).

Summary of Results

Using quantitative anatomical R_1 ($=1/T_1$) maps as myelin markers, we found a small keyhole-shaped area running over the medial two thirds of Heschl's gyrus that corresponded to non-human primate and human post mortem descriptions of auditory core (Hopf, 1951; Jones et al., 1995; Hutsler and Gazzaniga, 1996; Kosaki et al., 1997; Hackett et al., 1998; Morosan et al., 2001; Hackett et al., 2001; Wallace et al., 2002; Fullerton and Pandya, 2007; Hackett and de la Mothe, 2009) in terms of its laminar profile, orientation, shape, and intensity differences. This was true both in the surface-based group average as well as in most individual participants. This pattern was consistent across scans and subjects, and was essentially unchanged when the influence of local thickness and curvature on myelination and R_1 measures was taken into account (Serenio et al., 2012).

On both a group and individual level, tonotopic mapping of this core region showed a pattern consistent with previous electrophysiological and architectonic studies in macaques and owl monkeys, with a progression from high to low frequency preference moving medially to laterally within the most densely myelinated part of core. We suggest that this region corresponds to area A1. Within the rostrolateral slightly less densely myelinated finger of the core, there is a low-to-medium frequency progression. Based on similarities with studies in non-human primates (Merzenich and Brugge, 1973; Imig et al., 1977; Morel et al., 1993) we suggest this region is the homologue of area R. It is also notable that the gradient field map position associated with A1 and R (Formisano et al., 2003, Figure 4) appears to be similar to the layout suggested by the R_1 -defined core and tonotopic maps. The small low-to-medium-to-high frequency gradient within R is more robust in individual participants than in the group average, probably a result of blurring due to anatomical inter-subject variation in this small area. Nonetheless, further high-resolution tonotopic imaging combined with quantitative R_1 mapping will be useful in clarifying the group average result, particularly as it is not entirely as expected given some current models of tonotopic mapping in macaques and humans (Baumann et al., 2010; Woods et al., 2009).

DISCUSSION

These results show that non-invasive *in vivo* MRI techniques in humans can be used to characterize basic auditory functional regionalization originally revealed in non-human primates via a labor-intensive combination of physiological recording followed by postmortem cortical flattening and histology. Since post-recording histology is often partly compromised (Sincich & Horton, 2005), the definition of auditory core boundaries has also relied on other physiological criteria such as tuning bandwidth or firing rate/latency (Recanzone et al., 2000). The current study demonstrates for the first time concordance between tonotopy and myelination on the surface of the human cortex comparable with earlier studies in macaque and owl monkeys (Merzenich and Brugge, 1973; Imig et al., 1977; Pfingst and O'Connor, 1981; Morel and Kaas, 1992; Morel et al., 1993), a result which should help to narrow down the location of the auditory core in humans (Humphries et al., 2010). The current results were consistent across subjects, sessions, auditory stimulus

setups and scanner field strength.. Given that the combined functional and structural protocol can be acquired using standard scanner hardware within approximately one hour, these techniques should be useful for defining primary auditory fields for further study (for a similar approach, see Talkington et al., 2012) as well as investigations of clinical populations with atypical auditory processes, such as congenital deafness (Karns et al., 2012) or in patients with schizophrenia or tinnitus.

One notable finding is the small size of core and its constituent areas A1 and R, compared to that of primary visual and somatosensory cortices (noted also in macaque by Merzenich and Brugge, 1973). Even when borders are liberally defined, the surface area of core itself rarely exceeds $\sim 2 \text{ cm}^2$ across individuals (see Figure 4), with A1 occupying somewhat more than half of that area. These dimensions are comparable to those reported for auditory core in human post-mortem studies ($\sim 2 \text{ cm} \times \sim 0.9 \text{ cm}$, as in Fig. 6a of Wallace et al., 2002), after corrections for shrinkage, distortion, and unfolding method.

One open question for identifying auditory core with either R_1 maps or post-mortem histology is how to set intensity thresholds, particularly at the medial and lateral extents. Borders based on post-mortem myelin staining methods tend to be defined by eye, with a sudden change in stain density corresponding to areal edges (after correcting for section-to-section intensity differences), which can then be correlated with borders derived from other measures such as laminar-specific patterns of Nissl concentrations. However, there are large differences in myelination *within* core (high medially, decreasing smoothly moving laterally), making a fixed threshold difficult to apply. For instance, as can be seen in Hackett et al., 1998 (their figures 2c&d), the grayscale level in the thin, lateral-most part of core is very similar to that in the belt region surrounding medial core.

In the present MRI-based *in vivo* study, an analogous case of equivalent R_1 values at the lateral tip of core and the region surrounding medial core can be observed in participant S5 (see Figure 4). We have addressed this issue by presenting both the graded and unmasked heatscale data (Figures 2–4), along with the R_1 contour plots (Figures 5&6), where low R_1 values well outside the probable contiguous boundaries of core are not shown. Gradient-based approaches potentially avoid fixed threshold problems, but the gradient is a local measure that is noisy with unsmoothed data yet highly sensitive to spatial smoothing parameters.

The current findings confirm several results and predictions of the pioneering study of Sigalovsky et al. (2006), who used a multi-scan whole-brain FLASH protocol (with different flip angles for each scan) to map R_1 variation across auditory cortex (averaged over cortical depth), finding consistent patterns of higher R_1 values in medial Heschl's gyrus in individual participants, along with hyperintensities along the superior temporal gyrus and planum temporale. As predicted by Sigalovsky et al., higher resolution (0.512 mm^3 voxel volume in the present study versus 1.69 mm^3 in Sigalovsky) and increased SNR (due to 32-channel coil, multi-echo readout, 3T field strength, more data points) made it possible to see laminar differences in R_1 between probabilistically defined core and adjacent regions in the present study (Figure 2a). The increased resolution and SNR also enabled us to resample and render R_1 values at specific cortical depths where core and non-core regions are most different. This sharpened the resulting maps and revealed a correlation between cortical curvature and myelination (Annese et al., 2004) at middle cortical depths (Serenio et al., 2012). Note that the average R_1 values of $\sim 0.75 \text{ sec}^{-1}$ measured by Sigalovsky et al. (2006) at 1.5T cannot be directly compared to the average value of 0.69 sec^{-1} within TE1.0 measured in this study at 3T, since R_1 decreases with field strength (Oros-Peusquens et al., 2008). However, the rather small R_1 measured at 1.5T suggests that it might have been affected by partial volume effects, particularly contamination with cerebral spinal fluid, whereas the R_1 measured in

this study is in line with typical R_1 values of gray matter at 3T (Oros-Peusquens et al., 2008).

The present study showed that the size and shape of the high- R_1 core varies subtly between hemispheres within one individual (e.g., S1 and S6 in Fig. 4) and more markedly among individuals. In addition, the absolute R_1 values in and around core varies. We think that most of these differences reflect real anatomical intra- and inter-individual differences. However, some may be due to technical limitations. For example, image artifacts due to head motion most likely caused misestimates of R_1 in participant S4, obscuring the hyperintensity in the auditory core of the left hemisphere and causing anomalies in the temporal lobe, and around highly myelinated areas such as primary motor and somatosensory areas. Quantitative mapping combining data from multiple image acquisitions and with higher-than-usual resolution requires extra vigilance to reduce head motion artifacts. Shortening of the acquisition time would help to reduce motion artifacts within and across the runs, but sufficient SNR must be preserved in order to preserve R_1 map precision.

Inaccuracies in the reconstruction of the white or pial surfaces originating in local misclassification of tissue types (and thus local misestimation of cortical thickness and depth) is another potential source of artifacts.. Since myelination (and R_1) vary as much with depth as they do tangentially, this can obscure areal boundaries. Surface imperfections can occur near large pial vessels, where the pial surface closely approaches itself, or where the gray/white matter surface closely approaches itself along thin strands of white matter under small gyri (although it should be noted that these comprise only a tiny fraction of temporal cortex).

Another unexpected finding was that the medial, high-frequency edge of A1 appeared to extend slightly medially (~3mm) beyond the boundary of highest R_1 values in the group average, and in some individuals. This possible over-estimated extent in the group average result may be due in part to blurring over somewhat variable functional anatomy, but may also be due to inherent difficulties in mapping of the BOLD response at the boundary between medial Heschl's gyrus and the deep, narrow circular sulcus. This region is highly vascularized, which may blur the BOLD response where primary cortex adjoins the caudal (CM) auditory field. This could be resolved by tonotopic mapping at higher field strengths and higher resolution (e.g., da Costa et al., 2011; Formisano et al., 2003, Humphries et al., 2010), or by using acquisition methods thought to suppress contributions from larger vessels (e.g., spin-echo EPI at 7T, Yacoub et al., 2005). However, this finding may also reflect true physiological and anatomical properties. CM, a very small auditory area which shares the medial high-frequency border with A1, shows much broader frequency tuning, and a more discontinuous tonotopic map (Kajikawa et al., 2005). CM is also highly myelinated, making it difficult to distinguish the A1/CM border based on myeloarchitectonics alone, even in post-mortem assays, where changes in cell type are a more unambiguous marker of the A1/CM border.

Our new quantitative, high-resolution, multi-modal definition of the auditory core will make it possible to investigate how musical expertise or auditory acuity are related to differences in myelination of A1 and R (see Duncan and Boynton, 2003; Schwarzkopf et al., 2011; Song et al., 2011; and Duncan and Boynton, 2007, for related studies in the visual, auditory, and somatosensory systems). Our methods can also be extended to non-human primates to improve our understanding of homologies as well as potential differences. Studies of auditory and language function in individuals whose core and primary auditory fields are precisely mapped will improve our understanding of the contribution of these early areas to perceptual and cognitive processes. Finally, the sensitive quantitative indices of cortical

microstructure developed here may help with early diagnosis of disease and serve as a biomarker to assess the effectiveness of new therapies.

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**Tonotopic contours across architecturally-defined auditory core in macaque.
Maps redrawn and rendered with data from Figure 2a of Morel, Garraghty, & Kaas (1993)**

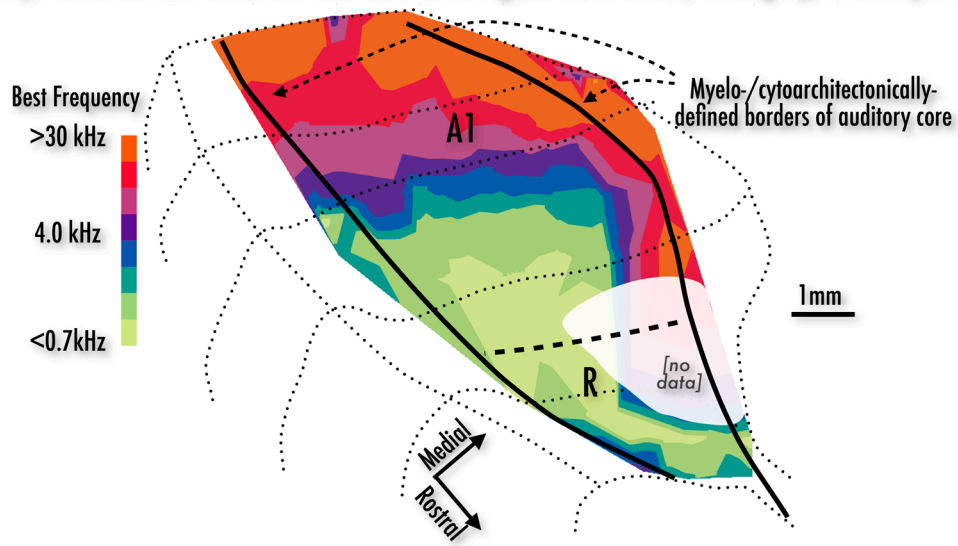


Figure 1.

Log-frequency isocontours within and around myelo- and cytoarchitecturally-defined auditory core (thick black lines), reconstructed from electrophysiological recording data reported in Figure 2A of Morel et al., 1993. (See Methods for details on contour rendering). Thin dotted lines show shape of underlying coronal sections of exposed temporal plane and superior temporal gyrus; thick dashed line is estimate of A1/R border. Figure 2A from Morel et al. was chosen for having the most extensive set of recording data over A1 and R, and for being representative of other datasets in Morel et al. and in other combined physiology/cytoarchitectonic experiments.

Tonotopic progressions in macaque can generally be described as follows: wrapping around the caudal cap of the core is a corona of high best-frequency (BF) neurons, with highest BF neurons near or slightly outside the core proper. This high-BF region tends to extend more than halfway along the medial edge of the core (moving rostrally), with BFs then dropping to mid- or low frequencies. Continuing along the medial edge toward the rostralmost tip of core, there is often a moderate increase from low to medium BFs. Progressing posteromedially to anterolaterally across core, there is a steep drop from higher to lower BFs ending in a low BF rostralateral trough. Along the lateral edge of core, moving posterior to anterior, there is a steeper descent in BF that joins the low-BF trough. Finally, in some cases there is an increase from low-to-medium/high BFs moving out from the anterolateral edge of the less densely myelinated aspect of core (for an almost complete low-to-mid BF gradient in R, see Figure 2B of Morel et al., 1993).

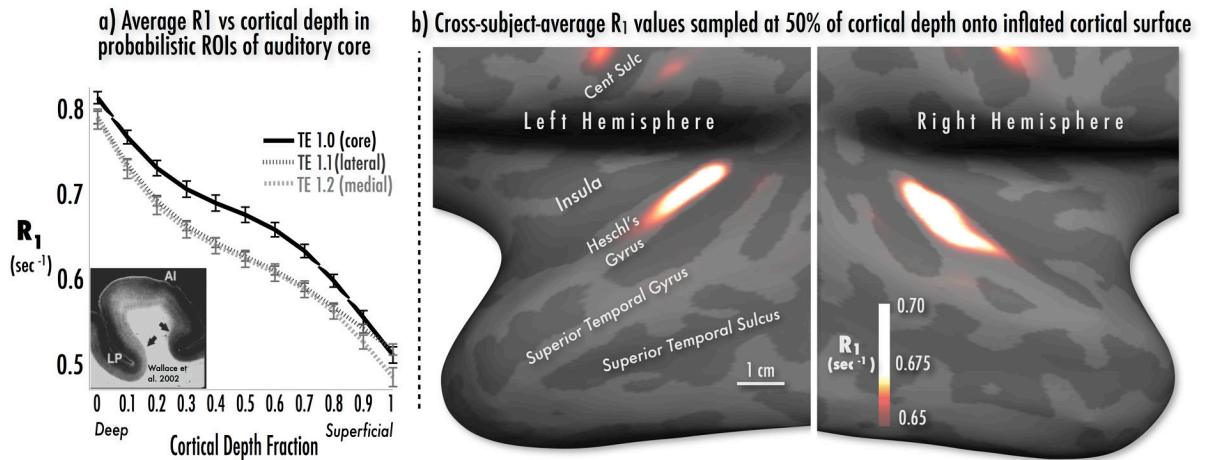


Figure 2.

(a, left panel) Relaxation rate ($R_1 \text{ sec}^{-1}$) as function of cortical depth, averaged within probabilistically defined subdivisions of Brodmann's area 41 (TE1.0, TE1.1, & TE1.2 according to Morosan et al., 2001). Average R_1 within TE1.0 (putative auditory core) decreases steeply from the gray/white boundary (depth fraction 0.0) to a tilted plateau at middle depths (0.3 to 0.6), then again drops steeply at superficial depths (0.7 to 1.0) (error bars: ± 1 SEM over subjects). R_1 within lateral (TE1.1) and medial (TE1.2) subdivisions shows a more gentle monotonic decrease from deep to superficial cortex. For comparison, the left inset is a myelin-stained section of human auditory core and belt cortex (from Wallace et al., 2002, contrast reversed) with a similar profile of myelination. **(b, right panel):** Group spherical average R_1 values sampled at 50% of cortical depth and projected onto a single subject's left and right inflated hemispheric surfaces. The auditory core is visible in both hemispheres as a keyhole-shaped hyperintensity maximum running posteromedially to anterolaterally over the medial half of Heschl's gyrus. Hyperintensity maxima can also be observed within the densely myelinated pre- and post-central gyri.

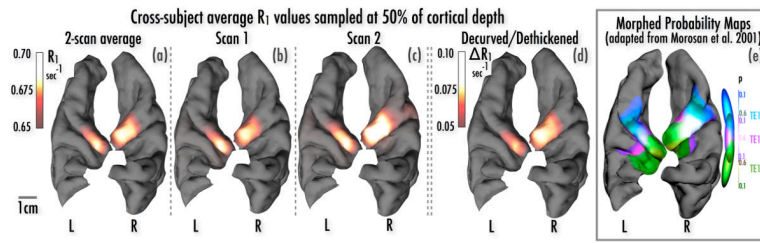


Figure 3.

Group average R_1 values from 50% of cortical depth, projected onto the pial surface of the digitally resected temporal lobes of a single subject; (a) local increases in R_1 values along medial Heschl's gyrus, averaged across both scans, same data as Figure 2b; (b, c) single-scan R_1 averages show excellent scan-rescan reproducibility; (d) maps of the change in R_1 after removing R_1 variance accounted for by local curvature and thickness show a very similar topology to raw R_1 maps; (e) probability maps of cytoarchitecturally defined TE1.0 ('core'), TE1.1, and TE1.2. derived from Morosan et al. (2001) Note that the overlap between probability distributions for TE1.0/TE1.1 and TE1.0/TE1.2 causes some probability maxima for TE1.1 (medial) and TE1.2 (lateral) to be darker colored, as shown in the overlapping probability ovoids at right.

All individual participants' R_1 values (S1-S6), sampled halfway through cortex, showing consistent myelination pattern corresponding to auditory core

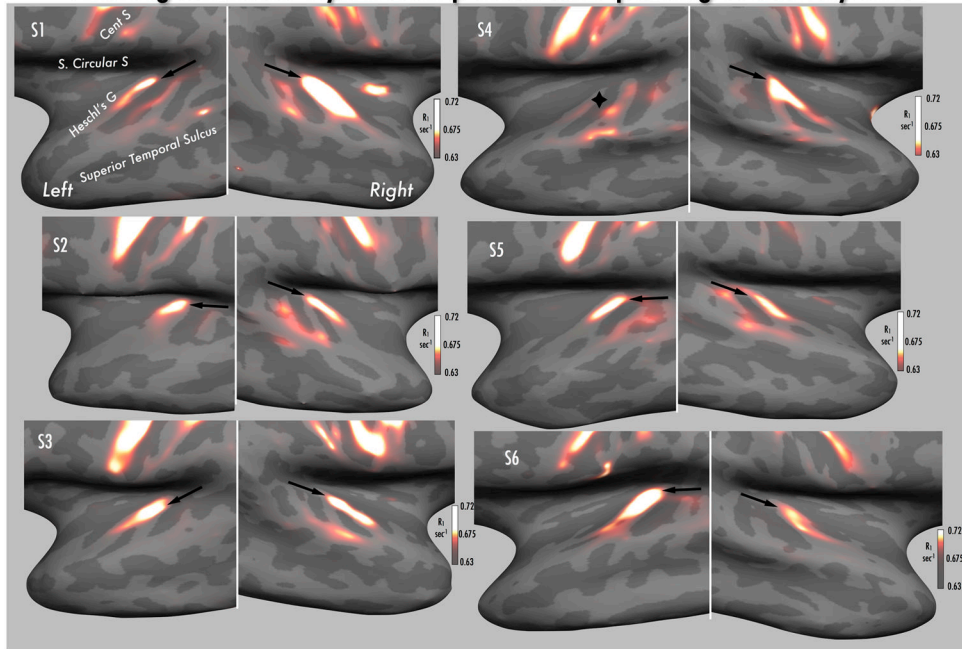


Figure 4.

Montage of individual participants' R_1 maps, with values sampled halfway through cortex and projected onto the inflated surface of temporal and frontal lobes. As shown in the colorbars, the range of projected R_1 values is constant over participants, with the colorscale slope adjusted slightly to show individual patterns of myelination. Keyhole-shaped regional increases in R_1 oriented posteromedially to anterolaterally across Heschl's gyrus were observed in all hemispheres (medialmost aspect indicated by arrow head) except the left hemisphere of participant S4 (marked with black diamond shape). R_1 maxima are also observed along the pre- and post-central gyri within strongly myelinated presumptive primary motor and somatosensory regions. Abbreviations: S. Circular S => superior circular sulcus of the insula; Cent S => central sulcus; Heschl's G => Heschl's gyrus.

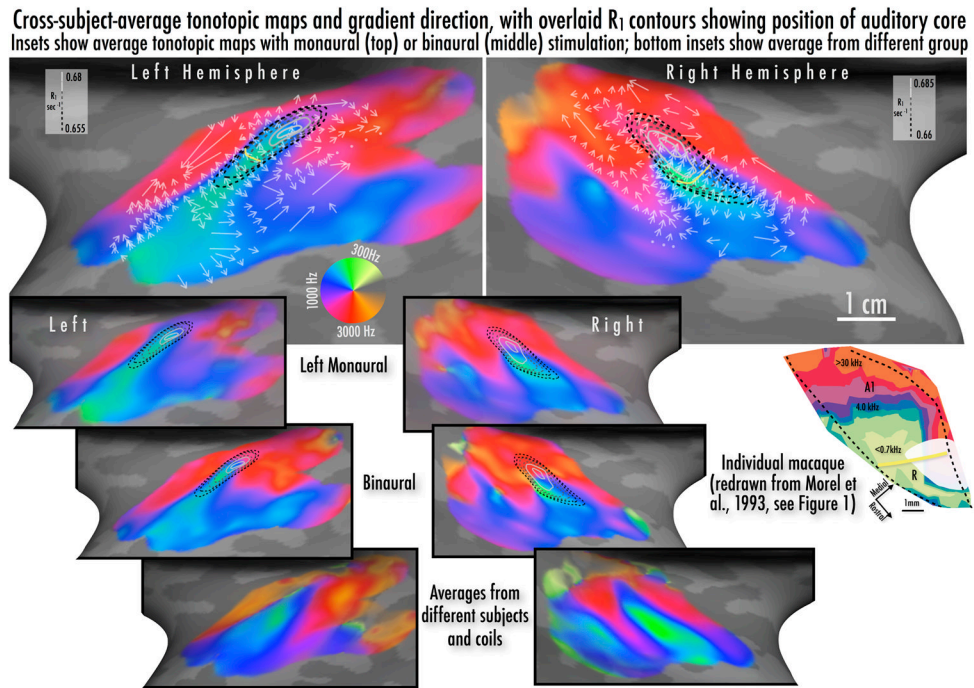


Figure 5.

Combined tonotopic maps, gradients and R_1 contours from group averages. Colormap shows fMRI results for characteristic frequency with logarithmic scaling in Hz around the color wheel. The angle of the overlaid white arrows shows the tonotopic gradient direction from low-to-high frequency preference at approximately 1/3 octave steps within tonotopic maps (perpendicular to isofrequency contour), with arrow length reflecting the approximate magnification factor (reciprocal of gradient vector). Dots indicate points where the gradient direction was unclear or interrupted. Dashed lines show R_1 values in grayscale-coded steps of 0.005 sec^{-1} (same data as Figure 4); curved yellow line is suggested A1/R border based on presence of low-frequency reversal and R_1 contour. Top and middle inset figures show tonotopic averages with left monaural and binaural stimulation from the same subjects as in the combined average. Bottom inset shows comparison tonotopic averages from different sets of participants and experimental setups; all tonotopic maps were identically masked using the independent auditory localizer (see Methods).

All individual participants' tonotopic maps (S1-S6) overlaid with 'dethickened' and 'decurved' R_1 contour

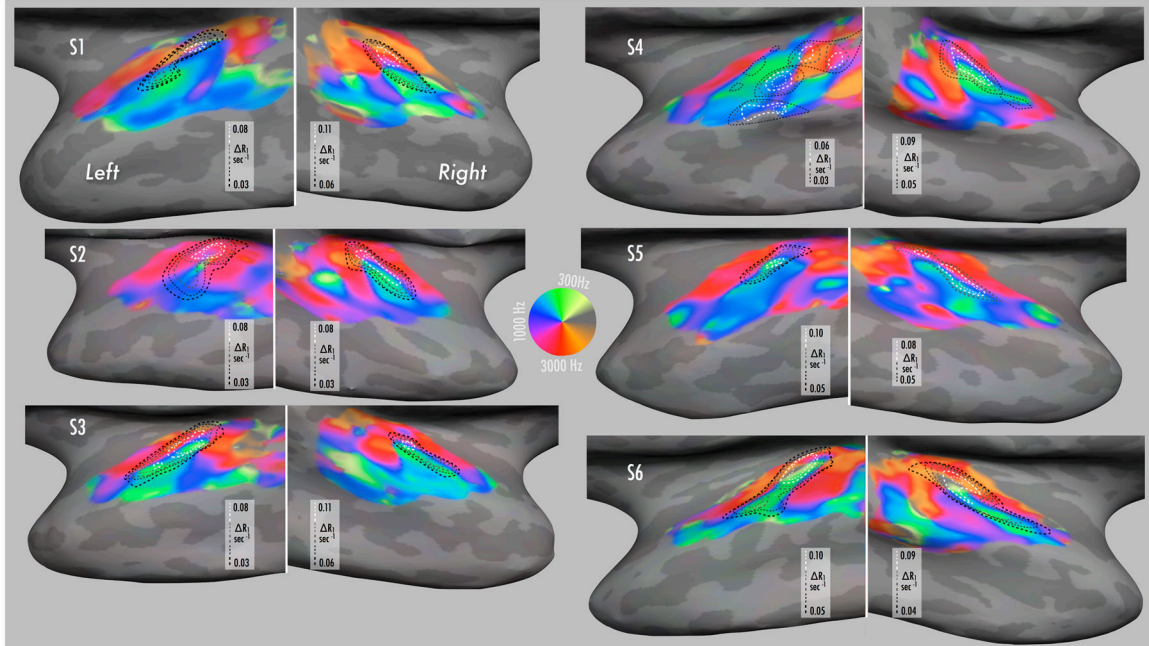


Figure 6.

Montage of individual participants' tonotopic maps and R_1 contours, sampled halfway through cortex and detrended for effects of local curvature and thickness (see Methods, and Figure 3d). Tonotopic colormaps are as in Figure 5. For visual clarity, contours indicate stepwise progression of R_1 values in probable auditory core only; see heatscale images in Figure 4 for raw R_1 data for each individual.

Repeated tonotopy sessions of a single participant, monaural presentation to left ear, 32-channel head coil, inflated right hemisphere shown. Top row, filtered vocalization stimulus, each panel from a different session. Bottom row, filtered music (left) and filtered white noise (right)

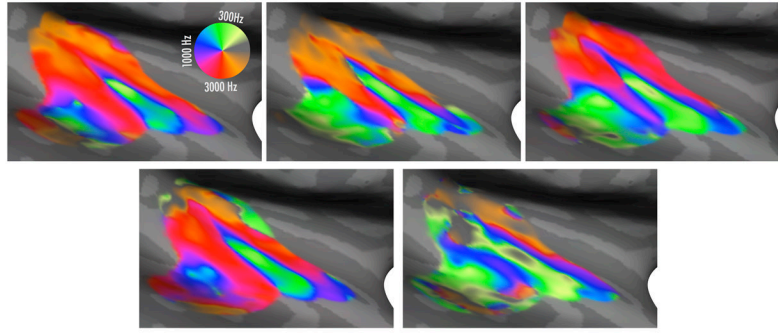


Figure 7. Functional tonotopic maps of an individual subject with focus on the right hemisphere of the inflated superior temporal lobe. Top row shows results from three different sessions of tonotopy using bandpass-swept vocalization stimuli (4 runs each); bottom left inset shows one session using bandpass-filtered music (4 runs), bottom right shows one session using bandpass-filtered, amplitude-modulated white noise (4 runs).