Mapping the human subcortical auditory system using histology, post mortem MRI and in vivo MRI at 7T

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Abstract Studying the human subcortical auditory system non-invasively is challenging due to its small, densely packed structures deep within the brain. Additionally, the elaborate

- three-dimensional (3-D) structure of the system can be difficult to understand based on currently
- available 2-D schematics and animal models. We addressed these issues using a combination of
- histological data, post mortem magnetic resonance imaging (MRI), and in vivo MRI at 7 Tesla. We
- 18 created anatomical atlases based on state-of-the-art human histology (BigBrain) and post mortem
- $_{19}$ MRI (50 μ m). We measured functional MRI (fMRI) responses to natural sounds and demonstrate
- that the functional localization of subcortical structures is reliable within individual participants
- ²¹ who were scanned in two different experiments. Further, a group functional atlas derived from the
- ²² functional data locates these structures with a median distance below 2mm. Using diffusion MRI
- ²³ tractography, we revealed structural connectivity maps of the human subcortical auditory pathway
- both in vivo (1050 μ m isotropic resolution) and post mortem (200 μ m isotropic resolution). This
- ²⁵ work captures current MRI capabilities for investigating the human subcortical auditory system,
- describes challenges that remain, and contributes novel, openly available data, atlases, and tools
- ²⁷ for researching the human auditory system.
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29 Introduction

³⁰ Understanding the structure of the human subcortical auditory pathway is a necessary step to

- ³¹ research its role in hearing, speech communication, and music. However, due to methodological
- issues in human research, most of our understanding of the subcortical (thalamic, midbrain, and
- ³³ brainstem) auditory pathway arises from research conducted in animal models. This might be
- ³⁴ problematic because, while the organization of the auditory pathway is largely conserved across
- ³⁵ mammalian species (*Malmierca and Hackett, 2010; Schofield, 2010*), the form and function of each
- ³⁶ structure may not be analogous (*Moore, 1987*). In this paper we show that three human imaging
- ³⁷ modalities -histology, post mortem magnetic resonance imaging (MRI), and in vivo MRI at ultra
- ³⁸ high-field (7 Tesla)- can identify the structures of the subcortical auditory pathway at high spatial
- resolution (between 50 and 1100 μ m).
- ⁴⁰ Although MRI has become increasingly powerful at imaging deep brain structures, anatomical

investigation of the human subcortical auditory pathway has been primarily conducted in post
 mortem tissue dissection and staining. *Moore* (1987) stained both myelin and the cell bodies of
 subcortical auditory structures in four post mortem human brainstem samples and compared them

subcortical auditory structures in four post mortem human brainstem samples and compared them
 to the analogous structures in cats (a common model for auditory investigations at the time). Later

investigations from the same group (*Moore et al., 1995*) used myelin and Nissl cell body staining to

⁴⁵ investigations from the same group (*Moore et al., 1995*) used myelin and Nissl cell body staining to ⁴⁶ investigate the timeline of myelination in human auditory brainstem development. More recently,

Kulesza (2007) stained six human brainstems for Nissl substance, focusing on the superior olivary

complex, finding evidence of a substructure (the medial nucleus of the trapezoid body) whose

existence in the human auditory system has been debated for decades.

Advances in post mortem human MRI allow for investigating three-dimensional (3-D) brain anatomy with increasingly high resolution (100 µm and below). This points to "magnetic resonance histology" (*Johnson et al., 1993*) as a promising avenue for identifying the small, deep subcortical auditory structures. However, to the best of our knowledge, post mortem MRI has not been utilized within the subcortical auditory system, although it has provided useful information about laminar structure in the auditory cortex (*Wallace et al., 2016*). To study the subcortical auditory system in living humans. MRI is the best available tool due to

56 its high spatial resolution. Anatomical in vivo MRI investigations of the human subcortical auditory 57 pathway so far have focused on thalamic nuclei (Devlin et al., 2006: Moerel et al., 2015), and the 58 identification of the acoustic radiations between the auditory cortex and medial geniculate nucleus 59 of the thalamus with diffusion-weighted MRI tractography (Devlin et al., 2006: Behrens et al., 2007: 60 *layad et al.*, 2014: Maffei et al., 2018). The inferior colliculus of the midbrain can also be identified 61 using anatomical MRI—for instance, Tourdias et al. (2014) and Moerel et al. (2015) show the inferior 62 colliculus using short inversion time T1-weighted anatomical MRI at 7 Tesla, although neither 63 investigation focused on anatomical segmentation of the inferior colliculus. Due to their small 64 size and deep locations, identification of more caudal subcortical structures-the superior olivary 65 complex and cochlear nucleus-remain challenging with in vivo anatomical MRI. 66

Although lower spatial resolution than anatomical MRI, functional MRI (fMRI) has been used to 67 investigate the relevance of subcortical processing of auditory information in humans, but it has 68 been limited by the small size of the structures involved and the relatively low resolution attainable 69 at conventional field strengths (3 Tesla and below) (Guimaraes et al., 1998; Harms and Melcher, 70 2002: Griffiths et al., 2001: Hawley et al., 2005). These acquisitions required trade-offs, such as low 71 through-plane resolution (7 mm) in exchange for moderate in-plane resolution (1.6 mm), and in 72 some cases researchers synchronized image collection to the cardiac cycle in order to overcame 73 the physiological noise associated with blood pulsation in the brainstem (Guimaraes et al., 1998: 74 Sigalovsky and Melcher, 2006). 75 More recent advances in MRI, especially the increased signal-to-noise ratio (SNR) available at

More recent advances in MRI, especially the increased signal-to-noise ratio (SNR) available at ultra-high magnetic fields (7 Tesla and above), have enabled higher resolution functional imaging of subcortical structures and more advanced localization of human auditory subcortical structures as well as their functional characterization. Using MRI at 7 Tesla (7T), *De Martino et al.* (2013) and *Moerel et al.* (2015) collected relatively high resolution (1.1-1.5 mm isotropic) fMRI with an auditory paradigm to identify tonotopic gradients in the inferior colliculus and medial geniculate nucleus. In these studies, high isotropic resolution and SNR provided an opportunity to investigate auditory responses throughout the subcortical auditory system.

⁸⁴ Despite the methodological advances in investigating the human brain, a systematic comparison ⁸⁵ of their capabilities for imaging the subcortical auditory system has not yet been undertaken. Here ⁸⁶ we use publicly available histological data (*Amunts et al., 2013*) to segment the main nuclei along ⁸⁷ the subcortical auditory pathway. Using state-of-the-art acquisition and analysis techniques, we ⁸⁸ evaluate the ability to identify the same structures through post mortem anatomical MRI, through ⁸⁹ functional MRI using natural sounds, and through estimating the connectivity between subcortical ⁹⁰ auditory structures with post mortem and in vivo diffusion MRI tractography. To compare the ⁹¹ histological, post mortem, and in vivo data, we project all images to MNI common reference space 92 (Fonov et al., 2009, 2011). Finally, to facilitate dissemination of our results, we have made the post

mortem anatomical data, in vivo functional and diffusion data, and the resulting atlases publicly
 available.

⁹⁵Where histology provides ground truth information about neural anatomy, we show that post ⁹⁶mortem MRI can provide similarly useful 3-D anatomical information with less risk of tissue damage ⁹⁷and warping. We also show that in vivo functional MRI can reliably identify the subcortical auditory ⁹⁸structures within individuals, even across experiments. Overall, we found that each methodology

⁹⁹ successfully localized each of the small structures of the subcortical auditory system, and while

¹⁰⁰ known issues in image registration hindered direct comparisons between methodologies, each

¹⁰¹ method provides complementary information about the human auditory pathway.

102 **Results**

¹⁰³ Definition of a subcortical auditory atlas from histology

To obtain a spatially accurate reference for all the subcortical auditory structures, we manually
 segmented publicly available histological data (100 µm version of the BigBrain 3-D Volume Data
 Release 2015 in MNI space from https://bigbrain.loris.ca (*Amunts et al., 2013*)).

¹⁰⁷ Upon inspecting this dataset, we noticed that the area around the inferior colliculus was in-¹⁰⁸ correctly transformed into MNI space. This was causing the colliculi to be larger and more caudal ¹⁰⁹ than in the MNI reference brain (*Figure 7*, second and third panels). Thus, our first step was to ¹¹⁰ correctly register the area around the colliculi (*Figure 7*, fourth panel; see Methods for details on ¹¹¹ the correction procedure).

The results of our BigBrain subcortical auditory segmentation in corrected MNI space are reported in *Figure 1* together with schematics redrawn from *Moore* (1987) (for the cochlear nucleus, superior olivary complex, and inferior colliculus) and the Allen Human Brain Atlas (*Hawrylycz et al.*, 2012; *Ding et al.*, 2016) (for the medial geniculate body). These schematics were used as reference during the segmentation. The 3-D rendering of the segmented structures highlighting the complex shape of the cochlear nucleus and superior olivary complex is also presented in *Figure 1*. The rendering is presented from a posterior lateral view in order to compare it with the Grav's Anatomy.

¹¹⁹ Plate 719 (*Gray and Lewis, 1918*).

120 Post mortem MRI

121 Post mortem MRI atlas of the human subcortical auditory system

Magnetic resonance histology—i.e., the study of tissue at microscopic resolution using MRI—provides 122 several unique advantages over conventional histology; 1) it is non-destructive; 2) it suffers minimal 123 distortion from physical sectioning and dehydration: 3) it yields unique contrast based on water 124 in the tissue and how it is bound (e.g., diffusion); and 4) it produces 3-D data. These advantages 125 make it an ideal medium for visualizing the 3-D organization of the deep brain structures (Johnson 126 et al., 1993). To delineate the subcortical auditory structures with MR histology, we acquired 50 µm 127 isotropic voxel size 3-D gradient echo (GRE) MRI on a human post mortem brainstem and thalamus 128 (described previously in (*Calabrese et al.*, 2015): see Methods for additional details). These data are 129 presented in Figure 2 (second column) after transformation to MNI space and resampling to 100 130 um isotropic resolution (see Methods section for details). The post mortem MRI data are presented 131 together with the histological data for comparison (first column). 132

Based on our segmentations of the subcortical auditory structures in the post mortem MRI data, the resulting 3-D model is presented in *Figure 2*. A volumetric quantification of the identified structures (in the BigBrain and post mortem MRI) is reported in *Table 1* and the overlap between the segmentations computed after projection in MNI space are reported in Table 2 (as inset in *Figure 2*)

137 Figure 2).



Figure 1. Literature diagrams (left columns) redrawn from *Moore* (1987) for the cochlear nucleus (CN), superior olivary complex (SOC), inferior colliculus (IC) and from the Allen Human Brain Atlas (*Hawrylycz et al., 2012*) for the medial geniculate body (MGB) compared to similar cuts from histology (BigBrain) in MNI (central column) and 3-D reconstructions of the segmented structures from the histology (bottom right column). The auditory structures are highlighted in gray in the left column, by a dotted line in the central column and in red on the modified Gray's anatomy Plate 719 (*Gray and Lewis, 1918*) and rendered as solid red surface meshes within the surface point cloud render of BigBrain MNI brainstem (right column). See *Figure 9* for 3-D animated videos of these auditory structures.









Table 2. Segmentation similarity comparison between BigBrain, postmortem and in-vivo auditory nuclei

		DICE	DICE Coeff.		dorff Dist.
		Left	Right	Left	Right
brain across menters	MGN	0.72	0.75	1.43	1.34
	IC	0.83	0.77	0.48	0.85
	SOC	0.61	0.51	4.30	5.80
seg	CN	0.80	0.74	0.77	2.63
	MGN	0.5	0.5	3.3	5.1
rtem	IC	0.3	0.4	6.4	5.8
t-mo	SOC	0.2	0.01	5.8	7.8
sod	CN	0.2	0.2	7.1	6.6
	MGN	0.4	0.5	3.9	4.7
bigbrain vs in-vivo	IC	0.4	0.3	6.6	7.3
	SOC	0.1	0.03	8.9	11.5
	CN	0.04	0.1	14.6	11.6

Figure 2. BigBrain–7T post mortem MRI image comparisons. Histological data (BigBrain) (left column) and T2*-weighted post mortem MRI data (100 µm - central column) in MNI space. Panels from bottom to top are chosen to highlight subcortical auditory structures (CN [bottom] to MGB [top]). Arrows (white with red outline) indicate the location of the subcortical auditory nuclei. The 3-D structures resulting from the segmentation of the post mortem data is presented on the top right panel. Table 2 quantifies (using DICE coefficient and average Hausdorff distance) the agreement (in MNI space) for all subcortical structures between: 1) segmentations performed on the BigBrain dataset by the two raters (KS and OFG) [top]; 2) segmentations obtained from the BigBrain dataset and from the post mortem MRI data [middle]; 3) segmentations obtained from the BigBrain dataset and from in vivo functional MRI data [bottom]). See *Figure 9* for 3-D animated videos of these auditory structures.

Table 1. Comparisons between the volume (mm³) of auditory subcortical structures reported in the literature (*Glendenning and Masterton, 1998*) and the volume obtained in our BigBrain segmentation (in MNI space), post mortem MRI data segmentation and in vivo functional clusters (defined based on voxels that are significant in at least three, four, or five participants out of the ten included in Experiment 1).

	Literature	BigBrain	Post mortem	In vivo (thr=3)	ln vivo (thr=4)	In vivo (thr=5)
CN	46	32	11	54	24	11
SOC	7	6	4	124	63	29
IC	65	63	73	263	189	146
MGN	58	75	134	304	207	152

¹³⁸ 3-D connectivity map of the human subcortical auditory system from post mortem ¹³⁹ diffusion MRI

Identifying the connectivity between subcortical auditory nuclei is crucial for understanding the 140 structure of the pathway. However, methods for tracing neuronal pathways that are available in 141 other animal models are generally not available in human studies, even post mortem. Diffusion-142 weighted MRI (dMRI) can be used to measure the orientation and magnitude of molecular motion 143 and infer patterns of white matter in brain tissue (both post mortem and in vivo). Using 200 um 144 diffusion-weighted MRI data acquired on the same post mortem sample (see Methods for details). 145 we modeled diffusion orientations and estimated likely connectivity pathways (or streamlines) using 146 tractography. Constraining the streamlines to only those that pass through auditory structures (as 147 identified from the anatomical MRI data and dilated 500 um to include adjacent white matter) we 148 visualized the connectivity map of the subcortical auditory pathway in *Figure 3*, left panel. 149 Connectivity closely resembles the expected pattern of the human subcortical auditory wiring 150 In particular, streamlines predominantly pass through the lateral lemniscus, the primary subcortical 151 auditory tract. Additional streamlines run through the brachium of the inferior colliculus, connecting 152 the inferior colliculus with the medial geniculate of the thalamus. Many streamlines then course 153

rostrally toward the auditory cortex (not present in this specimen).

At the caudal extent of the lateral lemniscus, streamlines pass through the superior olivary complex. Streamlines also run through the root of CNVIII. In total, each expected step along the subcortical auditory pathway is represented in this connectivity map.

Figure 3 (top right panel) shows the percentage of total streamlines connecting each of the subcortical auditory structures as estimated from this post mortem diffusion MRI sample. Overall, connections tend to be between ipsilateral structures, with weak connectivity to contralateral structures other than commissural connections to the contralateral homolog (except for between the cochlear nuclei). Still, the majority of streamlines pass through just one region (shown along the diagonal).

To investigate the relationship between streamline connectivity and ROI definition strictness. 164 we conducted two additional analyses. In *Figure 3*, we dilated the anatomical ROIs by 500 µm (2.5 165 voxels at 200 µm resolution), thereby including nearby white matter tracts (as well as adjacent 166 subcortical structures). In contrast, Figure 3-Figure Supplement 1 shows streamlines based on the 167 anatomical ROIs without dilation to account for white matter. As regions were defined as the core 168 nuclei in the anatomical MRL they largely exclude white matter tracts (such as the lateral lemniscus 169 and brachium of the inferior colliculus), leading to much sparser connectivity between subcortical 170 auditory nuclei. 171

Next, we resampled the diffusion MRI images to an in vivo-like resolution (1.05 mm isotropic). We
 again estimated fiber ODFs using CSD and estimated white matter connections with deterministic
 tractography. Using the (undilated but downsampled) anatomically defined ROIs as tractogra phy waypoints, we can visualize streamline estimates connecting subcortical auditory structures
 (*Figure 3-Figure Supplement 2*). Similar to the dilated ROI connectivity estimates, we see greater
 ipsilateral connectivity estimates between structures, particularly between left structures.



Figure 3. Post mortem diffusion MRI tractography. Left: streamlines passing through subcortical auditory structures, defined from 50 µm post mortem MRI in the same specimen, warped to 200µm isotropic diffusion image space and dilated 2.5 voxels (500 µm) to include neighboring white matter. Colors represent the local orientation at each specific point along the streamline: blue is inferior-superior, red green is anterior-posterior, and red is left-right. Ten percent of streamlines are represented in this image. A rotating animation is available in the online resources. Top right: Connectivity heatmap of subcortical auditory structures. Bottom right: Diffusion orientation distribution functions (ODFs) for each voxel; axial slice at the level of the rostral inferior colliculus (IC), including the commissure of the IC (bottom center arrow) and brachium of the IC (top left arrow). A video of the streamlines is available online: https://osf.io/kmbp8/

Figure 3-Figure supplement 1. Post mortem tractography with undilated ROIs.

Figure 3-Figure supplement 2. Post mortem tractography using data downsampled to in vivo resolution (1.05 mm).

Figure 3-video 1. 360° rotation video of post mortem streamlines.

178 Vasculature representations from post mortem MRI

179 Because T2*-weighted GRE imaging is sensitive to blood vessels, we processed our anatomical

¹⁸⁰ MR image to highlight brainstem vasculature (*Figure 5*, right column, base image). These 3-D

- vasculature images bear striking resemblance to post mortem data acquired with a stereoscopic
- microscope after full clearing method (see *Duvernoy* (2013) for detailed diagrams of human brain-
- stem vasculature). These vasculature images in the MNI space can be helpful to understand the
- nature of the in vivo functional signals (see next section).

185 In vivo MRI

We next sought to identify the structures and connections of the human subcortical auditory 186 system in living participants. By leveraging the increase signal and contrast to noise available at 187 ultra-high magnetic fields (7 Tesla) (Vaughan et al., 2001: Ugurbil et al., 2003: Ugurbil, 2016), we 188 collected high resolution anatomical (0.7 mm isotropic), diffusion-weighted (1.05 mm isotropic; 189 198 diffusion gradient directions across 3 gradient strengths) and functional (1.1 mm isotropic) 190 MRI in ten participants (see Methods for details). Leveraging the increased SNR available at high 191 fields, we aimed to collect data that would allow a functional definition of the auditory pathway 192 in individual participants. For this reason, we collected a large quantity of functional data in all 193 individuals: two sessions with 12 runs each in Experiment 1 and two sessions with eight runs each 194 in Experiment 2 (totalling 8 hours of functional data for each participant who completed both 195 experiments). All statistical analyses were performed at the single subject level. Group analyses 196 were used to evaluate the correspondence across subjects of individually defined regions (i.e., the 197 definition of a probabilistic atlas across participants) as well as the ability to generalize to new 198 participants by means of a leave-one-out analysis. 199

200 Anatomical MRI

Visual inspection and comparison to the MNI dataset (*Figure 5–Figure Supplement 2*) showed that the MGB and IC could be identified on the basis of the anatomical contrast, especially in the short inversion time T1-weighted data (*Tourdias et al., 2014; Moerel et al., 2015*). However, while the superior olivary complex (SOC) could be identified in the MNI dataset (*Figure 5–Figure Supplement 2*), it could not be identified in average anatomical image from our 7T data. This is possibly due to the limited number of subjects leading to the lower signal to noise in the average image. We have also explored the combination of image contrasts within each individual using a

²⁰⁸ compositional method proposed in (*Gulban et al., 2018b*), but the results were inconclusive.

209 Functional MRI

The difficulty in delineating the CN and SOC from anatomical in vivo MRI data (see Figure 5-Figure 210 *Supplement 1* for the average anatomical images obtained from our in vivo data) oriented our 211 investigation towards the possibility to identify the subcortical auditory pathway—in vivo and in 212 single individuals—on the basis of the functional responses to sounds. Functional responses to 213 168 natural sounds (Experiment 1) were collected at 7T using a sparse acquisition scheme and a 214 fast event-related design. We additionally report the reproducibility of the individual functional 215 delineations in six out of the ten participants who participated in a follow up experiment in which 216 responses to 96 natural sounds (Experiment 2) were collected at 7T using a sparse acquisition 217 scheme and a fast event-related design. 218 Statistical analysis of the functional responses allowed us to define voxels with significant 219

Statistical analysis of the functional responses allowed us to define voxels with significant activation in response to sounds in each individual. Additionally, we created a probabilistic functional atlas based on the overlap of statistically significant maps across individuals (after anatomical registration to a reference subject). To evaluate the generalization to new data we also computed leave-one-out probabilistic functional atlases each time leaving one one of our participants (see Methods for details).



Figure 4. Single subject functional activation maps obtained from Experiment 1 thresholded for significance (FDR-q = 0.05 and p<0.001; see Methods for details) and leave-one-out probabilistic functional maps highlighting voxels that are significant in at least three of the other nine subjects. For each participant, CN/SOC and IC are shown in transversal cuts, MGB is shown in a coronal cut. See single subject videos for 3-D view of these maps in *Figure 10* supplements. Unthresholded maps can be found in our online resources (see Data Availability section).

Figure 4–Figure supplement 1. Correspondence between single subject activation maps and leave-one-out probabilistic maps. Figure 4–Figure supplement 2. Effect of threshold on leave-one-out probabilistic maps on correspondence with single subject activations Figure 4–Figure supplement 3. Reproducibility across experiments of the functional activation maps in six participants (also see *Figure 11*). Figure 4–Figure supplement 4. Correspondence between single subject activation maps across experiments. Figure 4–Figure supplement 5. Effect of spatial smoothing in the analysis of the data collected from two of the participants.

Figure 4 shows, for each individual participant, the statistically thresholded (see Methods) 225 activation maps together with leave-one-out probabilistic functional maps obtained considering 226 all other individuals. The unthresholded maps are reported in supplement videos to *Figure 4* 227 and available for inspection in the online repository of the data. In all our participants, we could 228 identify clusters of significant activation in response to sounds in the MGB. IC. SOC, and CN. In each 229 individual and for each auditory nucleus, these activation clusters correspond to locations that are 230 significantly active in at least three out of the other nine participants to the experiment Figure 4-231 Figure Supplement 1 reports the overlap and distance between functional centroids of the single 232 subject activation maps and the leave-one-out probabilistic maps. In addition, Figure 4-Figure 233 **Supplement 3** shows the reproducibility of the functional responses across experiments in six of 234 the participants. The analysis of the overlap and distance between the centroids of activation across 235 experiments within each of these six participants is reported in *Figure 4-Figure Supplement 4*. The 236 higher signal-to-noise ratio attainable in regions corresponding to the IC and MGB results in highly 237 reproducible functional responses both within and across participants in these regions. Activation 238 clusters identified at the level of CN and SOC in single individuals also reproduce (albeit to a smaller 239 degree with respect to IC and MGB), both within subjects (i.e., across experiments) and across 240 subjects. 241

The left column of *Figure 5* shows the probabilistic functional map obtained from all participants in Experiment 1 (i.e., representing the number of subjects in which each voxel was identified as significantly responding to sounds-the map is thresholded to display voxels that are significantly activated in at least three out of the ten participants) overlaid on the in vivo average anatomical MRI image (short inversion time T1-weighted image (*Tourdias et al., 2014*); see Methods for details).

Projecting these data to the reference MNI space allowed evaluating the correspondence between in vivo functionally defined regions and histological data (Big Brain - *Figure 5*, center column).

At the level of the CN, the clusters of voxels active in at least three out of the ten participants correspond mostly to the ventral part of CN. The dorsal subdivision of the CN is not recovered in these probabilistic maps (at least not in at least three volunteers consistently) possibly due to partial voluming with the nearby cerebrospinal fluid in combination with thinness (thickness around 0.5 mm) of the dorsal CN as it wraps around inferior cerebellar peduncle (see *Figure 1*). Nearby, the location of the activation clusters identifying the SOC overlaps with the SOC as identified in the BigBrain data.

As the next step, we qualitatively investigated if the orientation of the vasculature at the level of 257 the SOC may have an effect on size (and location) of the functionally defined regions. As a visual aid 258 in this evaluation, we overlaid the functionally defined regions with the vasculature image obtained 250 from the post mortem data (Figure 5, right column). In all subcortical regions the vasculature 260 appears to have a specific orientation, and, at the level of the SOC. vessels drain blood from the 261 center in a ventral direction (i.e., the direction of draining is towards the surface of the brainstem in 262 the top of the image reported in the transverse view, bottom in *Figure 5*). This specific vasculature 263 architecture may result in the displacement or enlargement of the functionally defined clusters 264 towards the ventral surface of the brainstem (as highlighted in the correspondence with histological 265 data in Figure 5). 266

The probability of the same voxel to be significantly modulated by sound presentation across 267 subjects increased at the level of the IC and MGB, where the histologically defined regions cor-268 responded (for the large part) to all subjects exhibiting significant responses to sounds. At the 269 threshold of three subjects in the probabilistic maps, the IC seems to extend towards the superior 270 direction, bordering and sometimes including parts of superior colliculus. On the other hand, 27 similarly to what may happen in the SOC, the general directions of the vasculature penetrating 272 the IC and draining blood towards the dorsal surface of brainstem angled in a superior direction 273 (Figure 5 right panel) may also impact the functional definition of the IC. 274

The functional responses in the MGB cover an area that is in agreement with histological data.

Interestingly, compared to the IC or SOC, there is no major direction of extension of functional
 responses as well as no clear direction (in comparison to SOC and IC) of vascular draining.

A quantification of the volume of functionally defined structures is reported in *Table 1* for different thresholds of the probabilistic group map (from a threshold that defines the regions based on voxels that are significant in at least three out of the ten participants to a threshold that define the regions based on voxels that are significant in at least five out of the ten participants). The overlap between functional regions and the BigBrain segmentations after projection in MNI space is reported in Table 2 (as bottom right inset in *Figure 2* - computed using a threshold for the probabilistic maps that defines the regions based on voxels that are significant in at least three of

285 the ten participants).

286 Diffusion MRI

With the successful identification of the subcortical auditory structures with functional MRI, we next
 sought to estimate the likely connections between these structures in vivo. We analyzed the high
 spatial and angular resolution diffusion data to estimate streamlines of white matter connectivity
 following a similar process as the post mortem MRI (see Methods for further details).

Figure 6 shows diffusion tractography streamlines that pass through at least one subcortical
 auditory structure (as defined by group-level probabilistic functional activation [significant response
 in at least three out of ten subjects]; see section above). The high spatial and angular resolution of
 these data allow for vastly improved estimation of white matter connections between these deep,
 small structures.

While not a measure of actual physical brain connections—and therefore requiring caution in interpretation—connectivity patterns resemble what we would expect to see based on animal model tracer investigations. Overall, the connectivity network appears to be dominated by laterality, in that left hemisphere structures are generally more connected with other left hemisphere structures.

However, there are a few notable exceptions to this pattern: the cochlear nuclei and superior olivary complexes are strongly connected bilaterally, which fits with animal research suggesting one-half to two-thirds of ascending auditory connections cross the midline at these early stages. Additionally, there are a small number of connections between left and right inferior colliculi, likely along the anatomical commissure of the inferior colliculus.

305 **Discussion**

The auditory pathway includes a number of subcortical structures and connections, but identifying 306 these components in humans has been challenging with existing in vivo imaging methods. We 307 showed that functional localization of the subcortical auditory system is achievable within each 308 participant, and that localization is consistent across experimental sessions. To further facilitate 309 research on the anatomy and function of the human subcortical auditory system, we created 310 3-D atlases of the human auditory pathway based on gold standard histology, 50 µm isotropic 31 resolution post mortem anatomical MRI, and in vivo functional MRI at 7T. In addition, we created 312 3-D connectivity maps of the human subcortical auditory pathway using diffusion MRI tractography 313 in a post mortem MRI sample and in living participants. 314 These atlases and connectivity maps are the first fully 3-D representations of the human 315 subcortical auditory pathway and are publicly available to make the localization of subcortical

³¹⁶ subcortical auditory pathway and are publicly available to make the localization of subcortical
 ³¹⁷ auditory nuclei easier. In particular, the atlases are available in a common reference space (MNI152)
 ³¹⁸ to make registration to other MRI data as straightforward as possible. As part of this registration
 ³¹⁹ process, we have improved the registration of the brainstem of BigBrain histological data to the
 ³²⁰ MNI space, where the original MNI version presented a significant misregistration of the colliculi
 ³²¹ (as noticeable in *Figure 7*). The result of our new registration allows to more correctly localize the

³²² colliculi of BigBrain data in MNI without compromising the registration of other brainstem and

323 thalamic nuclei.



Figure 5. In vivo functional MRI responses to auditory stimuli, combined across ten participants. Left column: Conjunction of participants plotted on top of one participant's short inversion T1-weighted anatomical MRI. Center column: Conjunction of participants' fMRI responses warped to MNI space and plotted on top of BigBrain MNI (corrected) image. Right column: Conjunction of fMRI responses plotted on top of post mortem MRI vasculature images (1.1 mm minimum intensity projection).

Figure 5-Figure supplement 1. In vivo anatomical group average images in MNI space.

Figure 5-Figure supplement 2. Anatomical images from MNI ICBM 152 compared to BigBrain in MNI space



Figure 6. In vivo tractography of the subcortical auditory system from 7T diffusion-weighted MRI. Left: 3-D images from one participant. Fiber orientation distribution functions were estimated from diffusion-weighted MRI images of the brainstem and were used for deterministic tractography. Streamlines that passed through functionally defined auditory ROIs (dark grey) are shown here (excluding streamlines through the medulla). Colors represent the local orientation at each specific point along the streamline: blue is inferior-superior, red green is anterior-posterior, and red is left-right. A rotating animation is available in the online resources. Top right: connectivity between subcortical auditory ROIs as a percentage of total brainstem streamlines, averaged over 10 participants. Bottom right: schematic of auditory brainstem connectivity from Gray's Anatomy of the Human Body. A video of the streamlines is available online: https://osf.io/ykd24/

Figure 6-Figure supplement 1. Bar plot of streamline counts through each ROI. **Figure 6-video 1.** 360° rotation video of in vivo streamlines.

In creating the atlas with three distinct modalities, we were able to assess the reliability of each 324 of the methods in identifying the human subcortical auditory pathway. Each modality provided 325 useful information to the segmentation of the auditory nuclei. All regions could be identified in 326 the BigBrain histological data, that also allowed us to identify small auditory sub-nuclei such as 327 the medial superior olive and lateral superior olive. High-resolution post mortem MRI also clearly 328 delineated the medial geniculate and inferior colliculus (with less contrast for the superior olive 329 and cochlear nucleus) while the overall image contrast facilitated registration with in vivo MRI 330 High-resolution in vivo functional MRI exhibited greater sensitivity to auditory structures than in 331 vivo anatomical MRI that was even higher resolution. We showed that functional MRI is useful to 332 localize structures throughout the auditory pathway despite their small size. In each participant we 333 identified voxels significantly responding to sound presentation in regions corresponding to the CN. 334 SOC. IC and MGB. We validated these definition by evaluating both the within-subject reproducibility 335 (i.e., by comparing functional maps across two experiments in six individuals) and the ability of a 336 probabilistic atlas defined on nine out of our ten participants to generalize to the left out volunteer. 337 In total, we found that each of the methods described here provides information to the delin-338 eation of the human subcortical auditory pathway. Our post mortem and in vivo data suggest that 339 MRI is a capable tool for investigating this system across spatial scales providing a bridge to the 340 gold standard, histology. 341

While not representing specific cells, MRI holds a number of advantages over the gold standard method, histology (*Johnson et al., 1993*). First, MRI allows for visualization and analysis of an entire 3-D structure at once, with minimal geometric warping from (virtual) slice to slice (which can occur in slice-based histology if individual slices contract on a slide or are damaged during the physical slicing). Second, MRI can be used in vivo in human participants, opening up the possibility to address research questions on the functional and anatomical properties of human subcortical structures, their correspondence, and their involvement in human behavior.

Probing the connectivity of the human subcortical auditory pathway has been extremely limited, since gold standard (but invasive) tracer studies are largely unavailable for human specimens. In this study, we show that diffusion MRI tractography is sensitive to connections within the human subcortical auditory system, both post mortem and in vivo. In addition to streamlines corresponding to the lateral lemniscus-the major ascending auditory white matter tract-we can see streamlines crossing the midline at the level of the superior olivary complex and the inferior colliculus.

Interestingly, with the highest resolution data (200 µm post mortem diffusion-weighted MRI). 355 we were able to estimate streamlines visually resembling the expected auditory pathway, but 356 missing putative key connections between subcortical auditory structures themselves when using 357 the strictly defined ROIs as tractography seeds. In contrast, the relatively lower resolution in vivo 358 diffusion-weighted MRI produced estimates of connectivity more like what we expected from the 359 literature. We had two hypotheses as to why these results appeared. First, the higher resolution 360 anatomical definition of the nuclei not including the immediately surrounding white matter could 361 miss streamlines that terminate at the immediate proximity of the structures' borders (similar 362 to issues in cortex (Reveley et al., 2015)). Second, partial volume effects in the lower resolution 363 data—combining white matter and grev matter in the same voxels—could actually *increase* stream-36/ lines terminating within the anatomical ROIs. Dilating the post mortem ROIs and downsampling the 365 data to the in vivo resolution both resulted in greater streamline connectivity between subcortical 366 auditory structures, suggesting that our hypotheses were likely. Thus, while high spatial resolution 367 diffusion-weighted MRI allowed for much finer, higher quality streamline estimates, it also places 368 constraints on tractography analyses that must be accounted for and investigated further. 360

More generally, the density of brainstem and midbrain nuclei and frequent crossings between perpendicular white matter bundles pose a challenge to diffusion tractography estimations of white matter connectivity, so it was not clear beforehand if this methodology would be sufficient for visualizing these connections. Additionally, because a gold-standard connectivity method is not available in humans, we could not directly validate our tractography findings (as can be done in the macaque, though with limited success; see *Thomas et al.* (2014)). However, our results suggest that,
 with continually improving diffusion-weighted MRI acquisition and analysis techniques, focused
 investigations on the human subcortical auditory pathway can-and should-become more prominent
 in the near future.

In addition to high resolution anatomical post mortem MRI and diffusion MRI tractography 379 we were also able to identify the subcortical auditory system in vivo with functional MRI. Previous 380 studies have identified these structures with functional MRL but they typically required constrained 381 acquisition parameters—for instance, they used single slices with low through-plane resolution 382 in order to support high in-plane resolution (Guimaraes et al., 1998; Harms and Melcher, 2002; 383 Griffiths et al., 2001: Hawley et al., 2005: Sigalovsky and Melcher, 2006). In the present study, by 38/ taking advantage of the increased signal of high-field (7-Tesla) MRI, we were able to image the 385 brainstem using isotropic voxels at high resolution across a wider field-of-view that covers the 386 human auditory pathway in coronal oblique slices. The use of slice acceleration (*Moeller et al.* 387 2010: Setsompop et al., 2012) allowed us to acquire enough slices to cover the whole brainstem. 388 thalamus and cortical regions around Heschl's gyrus with the exclusion of anterior portions of the 389 superior temporal gyrus and sulcus. Using isotropic voxels allowed us to better evaluate the 3-D 390 volume of significantly activated regions, limiting partial volume effects that are inevitable when 391 using thick anisotropic slices. 392

Similar to previous research at lower magnetic fields (Hawley et al., 2005; Sigalovsky and 393 Melcher, 2006), the 7T MR images did not allow for an anatomical definition of the CN and SOC 394 (although IC and MGB were clearly visible). A possible reason for this is the reduced signal- and 395 contrast-to-noise ratio in these regions. Only very recently has 7T MRI enabled anatomical localiza-396 tion at the level of the SOC in individual subjects (Garcia-Gomar et al., 2019). It should be noted 397 that we could identify the SOC in the MNI ICBM 152 dataset that results from the average of a much 398 larger cohort. Therefore, future investigations should be tailored to optimize anatomical image 399 contrasts to auditory brainstem regions in single subjects. The (post mortem) atlases we provide 400 here will prove a useful tool for these investigations by providing a reference for the expected 401 location (and size) of these regions 402

In contrast to in vivo anatomical localization, our data—in agreement with previous reports 403 (Hawley et al., 2005: Sigalovsky and Melcher, 2006)—show that functional mapping of the subcor-404 tical auditory pathway is an effective method for localizing these structures. While histologically 405 defined CN and SOC regions have been previously used to sample functional responses from in vivo 406 fMRI data (Hawley et al., 2005: Sigalovsky and Melcher, 2006), the overlap between functionally 407 and histologically defined subcortical auditory structures has not been reported before. Here 408 we investigated the ability of BOLD fMRI (as an indirect measure of neuronal activity) to localize 400 subcortical auditory regions. We show that functional definitions are possible as distinct clusters 410 of activation were detected in all subjects across the subcortical auditory pathway. These regions 411 were reproducible both within subjects (across experiments) and across subjects (comparing single 412 participants functional maps to the leave-one-out atlas obtained with all other participants). We 413 could identify the subcortical auditory nuclei despite not using cardiac gating, a method that previ-414 ous studies showed to increase the signal-to-noise ratio in subcortical regions (Guimaraes et al., 415 1998: Harms and Melcher, 2002: Griffiths et al., 2001: Hawley et al., 2005: Sigalovsky and Melcher, 416 2006). We instead increased statistical power by presenting a large number of natural sounds 417 with multiple repetitions. Using smaller voxels also reduced partial volume effects between cere-418 brospinal fluid (which is heavily affected by physiological noise) and the brain tissue (Triantafyllou 419 et al., 2016). In addition, the correspondence of functionally defined regions across ten participants 420 after anatomical alignment allowed us to build a functional probabilistic atlas. 421

Despite these positive outcomes, functionally defined regions exhibited overall larger volumes compared to the histological ones (see Table 1 in *Table 1*). Although we acquired data at relatively high resolution (1.1 mm isotropic), our functional voxel size and the mild spatial smoothing (1.5mm) might be the source of this observation. Another factor that may have impacted the increased

volume of the in vivo probabilistic regions can be the residual anatomical misalignment across 426 subjects that also contributes especially to the lower degree of overlap at CN and SOC. In this case, 427 the individual anatomical images not showing enough contrast might be the cause. Partial volume 428 also most likely impacted small regions such as the CN and SOC, and draining effects due to the 429 vascular architecture could also have an impact on the size and localization of the in vivo defined 430 regions. Further, because we used only the overall response to sounds as functional definition, the 431 regions we defined may include sub-regions not specific to the system under investigation (e.g., the 432 inclusion of multisensory deep layers of the superior colliculus at the border with the IC) (Sparks 433 and Hartwich-Young, 1989: liang et al., 1997). This effect could be reduced by using different 434 stimuli and statistical contrasts. For instance, one could contrast uni-sensory and multi-sensory 435 stimuli to identify—within the current functional definition—the IC voxels that respond to visual 436 stimulation and thus may represent multi-sensory superior colliculus. For the IC and MGB, where 437 signal-to-noise ratio in the functional data is larger, a higher threshold in the probabilistic maps 438 results in a more accurate volumetric definition as well as more correct anatomical localization (see 439 e.g., *Figure 5*). It should also be noted that direct comparison of post-mortem and in vivo results 440 suffers from the additional problem of aligning data with very diverse contrasts and resolutions. 441 For the IC and MGB our procedure could be verified on the basis of the anatomical contrast in the 447 in vivo data, for the CN and SOC the lack of anatomical contrast (to be leveraged by the alignment 443 procedure) in the in vivo data may be the source of some of the misalignment between the data. 444

We also investigated the possibility of defining anatomical connections between subcortical 445 auditory nuclei using diffusion-weighted MRI. While affected by similar confounds as functional 446 MRI (e.g., partial voluming effects, physiological noise, and relative signal weighting), this technique 447 faces additional complications introduced by the number of orientations required, the gradient 448 strength (b-value) selected, the modeling of diffusion or fiber orientations within each voxel, and 449 the estimation of streamlines across brain regions, especially within the subcortical auditory sys-450 tem (Zanin et al., 2019). The post mortem and in vivo diffusion MRI datasets in this study each 451 implemented state-of-the-art acquisition techniques to optimize the MRI signal-to-noise ratio and 452 minimize MRI modeling errors. For example, as the fixation process likely changes the diffusion 453 characteristics of the tissue (Pfefferbaum et al., 2004; Miller et al., 2011), we compensated for this 454 effect by increasing the diffusion gradient strength (b-value). The constrained spherical deconvolu-455 tion modeling method takes advantage of the high angular resolution of each dataset to provide 45F fine-grained estimations of fiber orientation distributions. Additionally, the Euler Delta Crossings 457 (FuDX) deterministic tractography method is effective at generating streamlines through voxels with 458 multiple fiber orientation peaks, such as where white matter bundles cross. However, as diffusion 459 MRI and tractography are not measuring true neuronal connections, there is still room for error in 460 diffusion orientation and streamline estimation (*Schilling et al., 2019a*,b). 461

Our BigBrain histological segmentations are very similar in volume to those reported previously in the literature (*Moore, 1987; Glendenning and Masterton, 1998*), with slightly smaller cochlear nuclei and slightly larger medial geniculate bodies, but similar SOC and IC volumes. It has to be noted that the physical slicing process potentially introduces deformations in the tissue, and while the publicly available BigBrain dataset is of extremely high quality (with good registration from slice to slice), subtle deformations may have affected the shape or volume of the structures we identified.

Post mortem MRI segmentations differed more greatly, with smaller CN and SOC definitions but 469 larger MGB definitions compared to both the literature and BigBrain histological segmentations. 470 These differences could possibly be caused by the reduced contrast-to-noise ratio in the post 471 mortem MRI data compared to the histological data (despite their high spatial resolution). This 472 reduced contrast-to-noise ratio may be caused by both reduced differences in magnetic properties 473 between the regions and their surrounding tissues as well as from residual partial volume effects 474 (especially for the very small sections of the dorsal CN, for example) that may have blurred the 47 borders of the auditory nuclei in the post mortem MRI data. Contrast-to-noise ratio may be 476

ameliorated by different acquisition/reconstruction techniques (Wang et al., 2018), and optimizing 477 parameters may improve the definition of auditory nuclei on the basis of post mortem MRI data. 478 Finally, slight misregistration between specimens (e.g. the histological data and the post mortem 479 MRI data) likely still affect our comparisons, as registration between images (particularly from 480 different modalities) remains a challenge. For instance *Figure 2* shows slightly different shapes 48 and locations for the inferior colliculus between the two datasets, despite non-linear registration to 482 the same template. Although non-linear methods significantly improve gross registration between 483 specimens, large misregistrations are still possible (as shown for the colliculi in the original BigBrain 181 MNI registration). These issues can be addressed manually using additional image registration 485 techniques, as we did here with the BigBrain MNI registration (see our "corrected" version above). 486 but such hands-on, time-intensive edits are not always possible. Further, yastly different image 487 contrasts (like histology and MRI) result in different regions or subregions being emphasized in the 488 signal, creating an additional challenges in the registration procedure. 489 More generally, post mortem imaging—whether MRI or histology—is prone to modest defor-490

More generally, post mortem imaging—whether MRI or histology—is prone to modest defor mation of the specimen. Additionally, both post mortem specimens in this paper (BigBrain and
 post mortem MRI) were from 65-year-old male donors, and age may have additionally affected the
 volume of the brain structures we investigated.

Despite these limitations, the inter-rater and inter-experiment reliability in this study suggest 494 that each method is effective for localizing the subcortical auditory pathway. The reliable functional 495 localization of subcortical auditory structures opens the door to future investigations of more 496 complex human auditory processing. The atlases derived from each localization method is publicly 497 available (see "Data and code availability" in Methods) to facilitate further investigations into the 498 structure, function, and connectivity of the human subcortical auditory system in vivo. Lastly, the 499 3-D representations found in this paper and in the available data should be beneficial to others 500 in understanding the immensely complex, but identifiable, structure of the human subcortical 50 auditory pathway. 502

503 Methods

See Supplementary *Figure 8* for a summary of data sources, data processing steps, and software
 used in these analyses.

506 MRI acquisition parameters

507 In vivo MRI

The experimental procedures were approved by the ethics committee of the Faculty for Psychology and Neuroscience at Maastricht University (reference number: ERCPN-167_09_05_2016), and were performed in accordance with the approved guidelines and the Declaration of Helsinki. Written informed consent was obtained for every participant before conducting the experiments. All participants reported to have normal hearing, had no history of hearing disorder/impairments or neurological disease.

Images were acquired on a 7T Siemens MAGNETOM scanner (Siemens Medical Solutions, Erlangen, Germany), with 70 mT/m gradients and a head RF coil (Nova Medical, Wilmington, MA,

⁵¹⁶ USA; single transmit, 32 receive channels) at Maastricht University, Maastricht, Netherlands.

⁵¹⁷ We conducted two separate experiments. In Experiment 1, data were collected for n=10 partici-⁵¹⁸ pants (age range 25 to 30, 6 females), in three separate sessions. In the first session, we acquired ⁵¹⁹ the in vivo anatomical data set consisting of: 1) a T1-weighted (T1w) image acquired using a 3-D ⁵²⁰ MPRAGE sequence (repetition time [TR] = 3100 ms; time to inversion [TI] = 1500 ms [adiabatic ⁵²¹ non-selective inversion pulse]; echo time [TE] = 2.42 ms; flip angle = 5°; generalized auto-calibrating ⁵²² partially parallel acquisitions [GRAPPA] = 3 (*Griswold et al., 2002*); field of view [FOV] = 224 × 224 ⁵²³ mm²; matrix size = 320 × 320; 256 slices; 0.7 mm isotropic voxels; pixel bandwidth = 182 Hz/pixel;

524 first phase encode direction anterior to posterior; second phase encode direction superior to

inferior): 2) a Proton Density weighted (PDw) image (0.7 mm iso.) with the same 3-D MPRAGE 525 as for the T1w image but without the inversion pulse (TR = 1380 ms; TE = 2.42 ms; flip angle = 526 5°: GRAPPA = 3: FOV = 224 × 224 mm²: matrix size = 320 × 320: 256 slices: 0.7 mm iso. voxels: 527 pixel bandwidth = 182 Hz/pixel: first phase encode direction anterior to posterior; second phase 528 encode direction superior to inferior): 3) a T2*-weighted (T2w) anatomical image acquired using 529 a modified 3-D MPRAGE sequence (*De Martino et al., 2015*) that allows freely setting the TE (TR = 530 4910 ms⁻ TF = 16 ms⁻ flip angle = 5° GRAPPA = 3⁻ FOV = 224 × 224 mm² matrix size = 320 × 320⁻ 531 256 slices: 0.7 mm iso, voxels: pixel bandwidth = 473 Hz/pixel: first phase encode direction anterior 532 to posterior; second phase encode superior to inferior) and 4) a T1-weighted images acquired with 533 a short inversion time (SI-T1w) using a 3-D MPRAGE (*Tourdias et al.*, 2014) (TR = 4500 ms; TI = 670 534 ms [adiabatic non-selective inversion pulse]; TE = 3.37 ms; flip angle = 4°; GRAPPA = 3; FOV = 224 535 \times 224 mm²; matrix size = 320 \times 320; 256 slices; 0.7 mm isotropic voxels; pixel bandwidth = 178 536 Hz/pixel: first phase encode direction anterior to posterior; second phase encode direction superior 537 to inferior). To improve transmit efficiency in temporal areas when acquiring these anatomical 538 images we used dielectric pads (*Teeuwisse et al.*, 2012). 539

In the same session we acquired, for each participant, a diffusion-weighted MRI data set using a 540 multi-band diffusion-weighted spin-echo FPI protocol originating from the 7T Human Connectome 541 Project (1.05 mm isotropic acquisition and b-values = 1000 and 2000 s/mm²) (Vu et al., 2015). 542 extended in order to collect one additional shell at b-value at b = 3000 s/mm² (Gulban et al., 543 **2018***a*). Other relevant imaging parameters were (FOV = 200×200 mm² with partial Fourier 6/8. 544 132 slices, nominal voxel size = 1.05 mm isotropic, TR/TE = 7080/75.6 ms, MB = 2, phase encoding 545 acceleration (GRAPPA) = 3, 66 directions and 11 additional b = 0 volumes for every b-value). A 546 total of 462 volumes were obtained (231 in each phase encoding direction anterior-posterior and 547 posterior-anterior) for a total acquisition time of 60 minutes. 548

The other two sessions were used to collect functional data in order to identify sound responsive 549 regions in the human thalamus and brainstem. Participants listened to 168 natural sounds (1 550 second long) coming from seven categories (speech, voice, nature, tools, music, animals and 551 monkey calls) presented in silent gaps in between the acquisition of functional volumes and were 552 asked to press a button every time the same sound was repeated. The experimental paradigm 553 followed a rapid-event-related design in which sounds were presented with a mean inter stimulus 554 interval of four volumes (minimum three maximum five volumes). The two sessions were identical 555 and each session consisted of twelve functional runs and across the twelve runs each sound was 556 presented three times (i.e., each sounds was presented six times across the two sessions). The 168 557 sounds were divided in four sets of 42 sounds, each set was presented in three (non consecutive) 558 runs. As a result, the twelve functional runs of each session formed four cross validation sets each 550 one consisting of nine training runs and three testing runs (i.e., 126 training and 42 testing sounds). 560 Note that the testing runs were non overlapping across the cross validations. Catch trials (i.e., sound 561 repetitions) were added to each run, and were excluded from all analyses. 562

Functional MRI data were acquired with a 2-D Multi-Band Echo Planar Imaging (2D-MBEPI) 563 sequence (Moeller et al., 2010; Setsompop et al., 2012) with slices prescribed in a coronal oblique 564 orientation in order to cover the entire brainstem and thalamus and covering primary and secondary 565 cortical regions (TR = 2600 ms; Gap = 1400 ms; TE = 20 ms; flip angle = 80°; GRAPPA = 3; Multi-Band 566 factor = 2; FOV = 206 × 206 mm²; matrix size = 188 × 188; 46 slices; 1.1 mm isotropic voxels; phase 567 encode direction inferior to superior). Reverses phase encode polarity acquisitions were used for 568 distortion correction. Respiration and cardiac information were collected during acquisition using a 569 respiration belt and pulse oximeter respectively. 570

In experiment 2, six of the volunteers that participated in experiment 1 were recalled and functional data were acquired with the same slice prescription and functional MRI parameters as in experiment 1 (2D-MBEPI; TR = 2600 ms; Gap = 1400 ms; TE = 20 ms; flip angle = 80°; GRAPPA = 3; Multi-Band factor = 2; FOV = 206 × 206 mm²; matrix size = 188 × 188; 46 slices; 1.1 mm isotropic voxels; phase encode direction inferior to superior). Experiment 2 consisted of two sessions

in which participants listened to 96 natural sounds (1 second long) coming from six categories (speech, voice, nature, tools, music, animals) together with ripples (bandwidth = 1 octave; center 577 frequency = [300 Hz, 4 kHz]; AM rate = [3 Hz, 10 Hz]). Some ripple sounds contain a short noise 578 burst ('target') and participants were asked to detect such target in either low frequency ripples 579 or high frequency ripples in the two sessions respectively (the target occurrence varied (70 vs. 30 percent) for ripples whose center frequency did or did not match the current attention condition) 581 All sounds were presented in silent gaps in between the acquisition of functional volumes. The 582 experimental paradigm followed a rapid-event-related design in which sounds were presented 583 with a mean inter stimulus interval of four volumes (minimum three maximum five volumes). The 584 two sessions consisted of eight functional runs and across the eight runs each natural sound was 585 presented three times (i.e., each sounds was presented six times across the two sessions) while the 586 ripples were presented seven times per run. The 96 natural sounds were divided in four sets of 587 24 sounds, each set was presented in two (non consecutive) runs. As a result, the eight functional 588 runs of each session formed four cross validation sets each one consisting of six training runs 589 and two testing runs (i.e., 72 training natural sounds and 24 testing natural sounds). Note that 590 the testing runs were non overlapping across the cross validations. In each session of experiment 591 two we also collected a lower resolution (1 mm isotropic) anatomical reference images (T1 and PD 592 weighted) using the 3D MPRAGE sequence for alignment purposes and included reverses phase 593 encode polarity acquisitions for distortion correction. Respiration and cardiac information were 594 collected during acquisition using a respiration belt and pulse oximeter respectively. 595 Both in-vivo datasets acquired for experiment 1 and experiment 2 have never been published 596

⁵⁹⁷ before. This is the first work that uses this dataset.

598 Post mortem MRI

⁵⁹⁹ A human brainstem and thalamus specimen were dissected at autopsy from a 65-year-old anony-

mous male. The specimen was flushed with saline and immersed for two weeks in 10% solution of

neutral buffered formalin. Following this, the specimen was re-hydrated for one week in 0.1 M solu-

tion of phosphate buffered saline doped with 1% (5 mM) gadoteridol. Before the MRI acquisition,

the specimen was placed in custom MRI-compatible tube immersed in liquid fluorocarbon.

Magnetic resonance imaging was conducted in a 210 mm small-bore Magnex/Agilent MRI at the Duke University Center for In Vivo Microscopy. 3-D gradient echo images were collected at 50 μ m³ spatial resolution over a period of fourteen hours, with FOV = 80 × 55 × 45 mm, repetition time (TR) = 50 ms, echo time (TE) = 10 ms, flip angle = 60°, and bandwidth = 78 Hz/pixel.

⁶⁰⁸ Diffusion-weighted spin echo images were collected at 200 μ m³ spatial resolution with 120 ⁶⁰⁹ diffusion gradient directions at strength b=4000 s/m² and 11 b=0 s/m² volumes over 208 hours. ⁶¹⁰ The FOV was 90 × 55 × 45 mm with TR = 100 ms, TE = 33.6 ms, and bandwidth = 278 Hz/pixel.

611 Anatomical image registration

SI-T1w, T1w, T2*w and PDw images (700 µm iso.) were transformed to Talairach space (500 µm iso.) using BrainvoyagerQX version 2.8.4 (*Goebel, 2012*). Intensity inhomogeneity correction as
 implemented in SPM12 unified segmentation (*Ashburner and Friston, 2005*) was used for all images.
 A smaller volume containing brainstem and thalamus in each image was extracted (in the Talairach space) using FSL version 5.0.9 (*Jenkinson et al., 2012*) and histogram matched using percentile
 clipping (1% and 99%).

Individual masks for each 10 brainstems were created semi-automatically using ITK-SNAP version 3.6.0 active contour segmentation mode followed by manual edits. These masks included regions starting from 2 cm below the inferior part of pons to 0.5 cm above the medial geniculate nucleus (MGN), with a lateral extend reaching until the lateral geniculate nucleus (LGN) and 3 cm anterior from MGN, not including cerebellum or large arteries that lie on the surface of brainstem. These brainstem masks were then used with FSL-FNIRT (*Andersson et al., 2007*) to warp nine of the ten brainstems to the reference brainstem (subject 1) using SI-T1w images. We used the SI- T1w images to drive the non linear registration due to the enhanced anatomical contrast across

structures within the thalamus and brainstem present in these images (*Tourdias et al., 2014*; *Moerel et al., 2015*). The FNIRT parameters were subsamp = 2, 2, 1, 1, miter = 100, 100, 50, 50,

infwhm = 2, 2, 1, 1, reffwhm = 2, 2, 0, 0, lambda = 100, 50, 20, 5, estint = 0, 0, 0, 0, warpres = 2, 2, 2 with spline interpolation (parameters not mentioned here were the defaults as set in FSL 5.0.9).

To compare in vivo with post mortem MRI and histology data, we projected the averaged SI-T1w, T1w, T2*w and PDw images to the MNI reference space (ICBM 152 2009b non-linear symmetric, 500 µm iso.) (*Fonov et al., 2009, 2011*)¹. The ICBM 152 reference includes T1w, T2w and PDw data and projecting in vivo and post mortem MRI as well as histology data to this space allowed us also to evaluate the contrast that these commonly used template images have in subcortical auditory areas. To register our in vivo MRI data set to MNI, we used FSL-FNIRT but this time driven by the T1w images (available both in our data set and in the MNI ICBM 152 2009b data).

The post mortem diffusion b0 image was transformed to the post mortem anatomical image space with an affine transformation in ANTs. Anatomical-space images (including the manually segmented atlas) could then be transformed into diffusion space using the 'antsApplyTransforms' command, with the affine transform matrix, a super-sampled diffusion image (from 200 µm to 50 µm to match the anatomical image resolution) as the reference image, and denoting the warp as an inverse transform.

In vivo and post mortem images were registered non-linearly using ANTs. The in vivo SI-T1w
 image was warped to the post mortem diffusion b0 image following a rigid, then affine, then
 non-linear SyN algorithm. This produced an in vivo brainstem image in post mortem diffusion
 space.

The ANTs non-linear registration also created warp and inverse warp transforms that could then be used to transform atlases from one space to another. To preserve the higher resolution of the post mortem MRI when inverse warping post mortem images to in vivo space, we supersampled the in vivo SI-T1w image to 200 μm (matching the post mortem diffusion image) or 50 μm (matching the post mortem anatomical image).

Finally, to transform the post mortem anatomical image (50 μm) to MNI space, we applied the
 inverse transform from post mortem anatomical to diffusion space (resampled to 50 μm), then the
 inverse transform from diffusion space to in vivo space (similarly upsampled to 50 μm), and finally
 from in vivo space to MNI space using the FSL-FNIRT inverse transform (described above).

656 BigBrain histology segmentation

In what follows we describe the main anatomical observations related to the auditory structures as segmented in the 100 μm histological data. Images were segmented independently by two raters (KRS, OFG). Overlap between the two raters was high (see Table 2 [top row - Big Brain across segmenters] in *Figure 2*); in the figures we show the regions that were consistently segmented by both raters.

662 Vestibulocochlear nerve

625

⁶⁶³ The vestibulocochlear nerve (the eighth cranial nerve, or CNVIII) enters the brainstem where

the medulla and the pons meet (the pontomedullary junction). The cochlear component of the

vestibulocochlear nerve is composed of spiral ganglion neurons, whose cell bodies are within the

⁶⁶⁶ cochlea and which carry frequency-specific information to the brainstem.

In the BigBrain histology, CNVIII extends primarily laterally (but also anteriorly and inferiorly)

⁶⁶⁸ from the pontomedullary junction, bound posteriorly by the cerebellum. Parts of the nerve root are

⁶⁶⁹ still visible in the images although being cut. It is therefore not labeled in our histological atlas (but

see the post mortem MRI atlas below).

¹http://www.bic.mni.mcgill.ca/ServicesAtlases/ICBM152NLin2009

Cochlear nucleus 671

Once reaching the brainstem, the auditory nerves split into two main routes-one to the anterior 672

ventral cochlear nucleus (AVCN), and one to the posterior ventral cochlear nucleus (PVCN) and then 673 on to the dorsal cochlear nucleus (DCN) (Webster, 1992). Within each subnucleus, the neurons

674 maintain the tonotopic frequency representation they receive from the cochlea via the cochlear 675

nerve (De No. 1933b.a: Rose et al., 1960: Sando, 1965: Evans, 1975: Rvugo and May, 1993: Rvugo 676

and Parks, 2003) (see bottom panels of the two left most columns in Figure 2). 677

In the BigBrain data, the AVCN is situated anterior and medial to the root of CNVIII, while the 678 PVCN continues from the root of CNVIII and extends posteriorly towards the DCN. The DCN is clearly 679 visible as a dark band wrapping around the cerebellar peduncle posteriorly, becoming exposed on 680

the dorsal surface of the pons. 68

Superior olivary complex 682

The next structure along the auditory pathway is the superior olivary complex (SOC), which in 683 humans is located in the inferior pons. The SOC receives the majority of its ascending inputs 684 from the contralateral cochlear nucleus, although it also receives ipsilateral inputs as well. The 685 contralateral dominance is maintained throughout the remaining ascending pathway. The SOC is 686 comprised of the lateral superior olive (LSO) medial superior olive (MSO) and the medial nucleus of the trapezoid body (MNTB). The size of each of these nuclei varies between species, and it 688 is debated whether the trapezoid body exists in the human SOC (Moore, 1987: Strominger and 689 Hurwitz, 1976) (but see Kulesza and Grothe (2015) review of recent findings affirming the existence 690

of the human MNTB). 691

Although the individual substructures within the SOC have unique anatomy that can be identified 692 from histology (Moore, 1987; Kulesza, 2007), here we outline the structure of the SOC as a whole 693 in order to include all identifiable substructures (namely the MSO and LSO - see second panel 694 from the bottom of the two left most columns in *Figure 1*). The MSO is the largest SOC nucleus in 695 humans. unlike in other animals. The MSO receives inputs from both the left and right AVCN and 696 sends outputs to the ipsilateral lateral lemniscus. The LSO receives inputs from the ipsilateral AVCN 697 and from the ipsilateral MNTB. Outputs are sent to both ipsilateral and contralateral lateral lemnisci. 698 The MNTB receives inputs from the contralateral AVCN, and its axons terminate in the ipsilateral 690 LSO. 700

The MSO and LSO are visible in the BigBrain images, despite their small size. The MSO is a 701 thin pencil-like collection of nuclei whose caudalmost point begins around the same axial plane 702 as the rostralmost extent of the AVCN, about 4 mm medial (and slightly anterior) to the AVCN. It 703 then extends about 1 cm rostrally (angled slightly laterally), where it eventually meets the lateral 704 lemniscal tract. The LSO neighbors the MSO near its caudalmost portion, forming a "V" shape 705 when viewed axially. In our histological atlas, these two structures are combined into a single SOC 706 segmentation. Cells of the MNTB are not clear to us in this sample, so we do not segment it in our 707 atlas. 708

Inferior colliculus 709

The inferior colliculus (IC) is a large, spherical structure in the dorsal midbrain and receives ascending 710 inputs from the auditory brainstem via the lateral lemniscus (see second panel from the top of the 711 two left most columns in *Figure 1*). The central nucleus of the inferior colliculus receives most of 712 these connections, with external nuclei primarily receiving descending connections (*Webster, 1992*). 713

The inferior colliculus sends axons to the medial geniculate body of the thalamus via the brachium 714 of the inferior colliculus. 715

In the BigBrain data, the inferior colliculus is clearly identifiable as the lower two of the four 716 bumps along the dorsal portion of the midbrain (or tectum). The darkest staining within these 717 structures corresponds to the central nucleus of the inferior colliculus. An intensity gradient 718 outside of the central nucleus likely corresponds to the external and dorsal nuclei, which were 710

- included in our segmentation of the IC. Bounding the IC superiorly is the superior colliculus;
- medially, the commissure of the IC connecting the two inferior colliculi, as well as the aqueduct and
- periaqueductal grey; and anteriorly, other midbrain nuclei such as the cuneiform nucleus (lateral
- ⁷²³ and inferior to the IC are the borders of the midbrain).
- 724 Medial geniculate of the thalamus
- The medial geniculate body (MGB) of the thalamus is the final subcortical auditory structure that
- ⁷²⁶ sends auditory signals to the auditory cortex via the acoustic radiations (*Winer, 1984*) (see top panel
- ⁷²⁷ of the two left most columns in *Figure 1*). The MGB contains two or three major subdivisions: the
- ventral MGB receives the majority of IC inputs, while the dorsal and medial subdivisions (at times
- ⁷²⁹ grouped together, at times separately) receive more varied inputs from auditory and non-auditory
 ⁷³⁰ subcortical structures.
- In the BigBrain sample, the MGB is visible as a dark patch medial to the lateral geniculate nucleus (which can be easily identified by its striations) in a coronal view. Axially, the MGB takes an ovoid shape with a clear dorsolateral boundary next to the brachium of the superior colliculus, which appears light due to lack of cell nuclei being stained. Ventromedially, the MGB is bordered by a light
- $_{735}$ band corresponding to the medial lemniscus. Rostrally, we marked the edge of the MGB where cell
- staining decreases, at the border with the pulvinar nucleus and ventral posterolateral nucleus of
 the thalamus.

738 Post mortem MRI segmentation

- ⁷³⁹ In what follows we describe the anatomical contrast that can be leveraged from these post mortem
- 740 MRI data in order to identify structures in the auditory brainstem. We then used these segmenta-
- tions to create an MRI-based atlas of the subcortical auditory system, separate from the BigBrain
- ⁷⁴² histology-based atlas.
- 743 Vestibulocochlear nerve
- The CNVIII is visible in the post mortem MRI near the pontomedullary junction, extending laterally
- ⁷⁴⁵ and anteriorly from the brainstem (see the lower panels in *Figure 2*).
- 746 Cochlear nucleus
- The cochlear nuclei are challenging to identify in the post mortem MRI data, although the presence
- of the CNVIII root provides a landmark for localizing the other structures. Due to low signal contrast
- $_{749}$ around the ventral cochlear nucleus area in the T2*-weighted GRE MRI, we segmented the VCN
- according to the literature: bound by the cochlear nerve root and wall of the pons laterally, and by cerebellar white matter tracks medially. We were able to segment the dorsal cochlear nucleus
- ⁷⁵¹ by cerebellar white matter tracks medially. We were able to segment the dorsal cochlear nucleus ⁷⁵² based on the T2*-weighted image, where it appears brighter and can be identified as running
- posteriorly from the VCN and dorsally along the surface of the pons. distal to the inferior cerebellar
- 754 peduncle.

755 Superior olivary complex

- As with the cochlear nuclei, the SOC are more difficult to identify in the post mortem MRI than in the
- ⁷⁵⁷ histology, likely since the individual subnuclei like the MSO and LSO approach the size of a voxel in
- ⁷⁵⁸ at least one direction and are therefore prone to partial voluming effects. However, the pencil-like
- ⁷⁵⁹ MSO can still be identified in the coronal plane as a dark, elongated structure in the T2*-weighted
- ⁷⁶⁰ image, starting around the level of the ventral cochlear nucleus. In the axial plane, the SOC (but not
- ⁷⁶¹ its individual subnuclei) can be seen as a dark spot in the T2*-weighted image between the facial
- nucleus and the trapezoid body (see the second row from the bottom in *Figure 2*).

763 Inferior colliculus

- As in the BigBrain data, the inferior colliculus is relatively easy to identify based on its gross anatomical structure on the dorsal aspect of the midbrain. Additionally, the MR contrast provides

- relatively clear boundaries between the colliculi and surrounding structures. Indeed, it may even
- ⁷⁶⁷ be possible to segment the inferior colliculus into its subnuclei-the central, external, and dorsal
- nuclei-based on T2*-weighted MR signal intensities (see the second row from the top in *Figure 2*).
- ⁷⁶⁹ The external nucleus of the IC appears dark in the T2*-weighted image, on the lateral aspect
- of the IC. Medial to the external nucleus is the central nucleus, which has higher T2*-weighted
- intensity (appears brighter) in our MR images, and has clear boundaries on its ventral, medial, and
- dorsolateral sides. The dorsal nucleus is along the dorsal aspect of the IC and is the brightest
- ⁷⁷³ subcomponent within the IC in terms of T2*-weighted MR signal.

774 Medial geniculate

Although the borders of the MGB are less clear in the post mortem MRI than in the BigBrain images. 779 the structure itself is again relatively easy to identify by its gross anatomical location as well as 776 MR signal intensity. In the coronal plane, the medial geniculate is medial to the lateral geniculate 777 at the junction of the midbrain and thalamus. Axially, the medial geniculate has circular or ovoid 778 shape, again medial to the lateral geniculate. In the axial plane, the medial geniculate is largely 770 bordered dorsolaterally by the brachium of the superior colliculus which appears as a thick dark 780 band of fibers in the T2*-weighted image. Medially, the medial geniculate is bound by the brachium 781 of the inferior colliculus (also appearing as a dark fiber band), at least through the caudal half 782 of the structure. We have included the portions of this fiber bundle in the segmentation of the 783 medial geniculate, as the auditory fibers connecting the IC and the MGB are quite relevant to MRI 784 connectivity investigations (including our own; post mortem tractography results below). 785

As with the inferior colliculus, it may be possible to identify separate divisions within the medial 786 geniculate. Within the overall structure, there are two identifiable substructures based on T2*-787 weighted MR image intensity. Dorsomedially (and somewhat caudally), about half of the medial 788 geniculate has high T2*-weighted contrast and appears bright; the ventrolateral (and slightly rostral) 789 half appears darker in the T2*-weighted image. These segmentations largely (but not perfectly) 790 align with the ventral and dorsal/medial nuclei of the medial geniculate in the Allen Human Brain 791 Atlas (Hawrylycz et al., 2012), as well as with those of Paxinos et al. (2019). However, they vary 792 somewhat from the the axial slice segmentation from *Merker* (1983) shown in *Amunts et al.* (2012). 793

⁷⁹⁴ which show a largely horizontal delineation between the substructures.

795 Functional MRI analysis

In both functional experiments, data were preprocessed using BrainvoyagerOX version 2.8.4 796 (Goebel, 2012). Slice-scan-time correction, motion correction, temporal high-pass filtering (GLM-797 Fourier, 6 sines/cosines) and temporal smoothing (Gaussian, width of kernel 5.2 s). The defaults 798 in BrainvoyagerOX v2.8.4 were used for these steps aside from the explicitly stated values. The 799 functional images were then distortion corrected using the opposite phase encoding direction 800 images using FSL-TOPUP (Andersson et al., 2003). Conversion between Brainvoyager file types 801 to NIfTI which was required to perform distortion correction was done using Neuroelf version 802 1.1 (release candidate 2)² in Matlab version 2016a. For alignment across experiments (i.e., to 803 co-register the data of experiment 2 to the ones collected in experiment 1) we used FSL-FLIRT. In 804 this procedure the alignment between the functional data of the two experiments was tailored to a 805 mask that included the brainstem, thalamus and auditory cortex. 806

After pre-processing, functional images were then transformed to Talairach space using Brainvoyager at a resolution of 0.5 mm isotropic. We have previously used this procedure in order to reveal tonotopic maps in both the inferior colliculus and medial geniculate nucleus (*De Martino et al., 2013; Moerel et al., 2015*) and have shown that the upsampling has no consequence on the spatial distribution of the responses. Upsampling can also reduce effects of interpolation that is common during resampling in many image processing steps. After upsampling, mild spatial smoothing (Gaussian, FWHM 1.5mm) was also applied. *Figure 4–Figure Supplement 5* shows the

²http://neuroelf.net/

effect that spatial smoothing has on the activation maps obtained from two participants data in 814 experiment 1. 815

GLM-denoise (Kay et al., 2013) was used to estimate noise regressors. In brief, for each cross 816 validation a noise pool of non responsive voxels (i.e., voxels with a response to sound representation 817 determined by an F-statistic below a given threshold) was determined on the training data set (16 81 runs across the two sessions of experiment 1 and 12 runs across the two sessions of experiment 2) 819 and used to obtain noise regressors defined as the principal components of the noise pool time 820 course matrix that added to a GLM analysis (*Friston et al.*, 1994) of the training data would result 821 in an increased activation. The number of noise regressors was optimized using cross validation 822 within the training set. The selected noise regressor spatial maps were projected on the test data to 823 obtain the regressors for the test data. 824

Similarly, the hemodynamic response function (HRF) best characterizing the response of each 825 voxel in the brainstem was obtained using a deconvolution GLM (with 9 stick predictors) on the 826 training data. Note that this procedure, while possibly overfitting information in the training data, 827 produces noise regressors and an HRF for each test run (e.g. the noise regressors for runs 4, 6 and 828 9 of session one in experiment 1 comes from an analysis performed on all other runs in the same 829 session) that are not overfitted. 830

The resulting HRF and noise regressors were used in a GLM analysis of the test runs. We 831 combined all test runs (for each individual voxel) using a fixed effect analysis. 832

Statistical maps of responses to sounds vs silence were corrected for multiple comparisons 833 at the individual level using False Discovery Rate (FDR: g-FDR = 0.05). An additional threshold on 834 the uncorrected p-value of each voxel (i.e., p<0.001) was applied to further reduce the number of 835 false positive activation that can be expected when applying FDR. Unless otherwise stated, single 836 subject statistical maps are displayed by color coding yoxels that surpass these statistical thresholds. 837 Unthresholded statistical maps are visualized in *Figure 10* and are available at the online repository 838 of the data (https://osf.jo/hxekn/?view_only=be9ec398304344e8bb694a0658d77ed6) for inspection 839 The functional activation maps of the six participants that took part in both experiments have 840 been analyzed to demonstrate within participant reproducibility of effects. Since the stimuli were dif-841 ferent and the number of runs were different, this second experiment shows a generalization of the 842 first experiment, thereby additionally validating the detection of these structures. Figure 4-Figure 843 **Supplement 3** shows the statistically thresholded activation maps for each of this six participants 844 for the two experiments in three anatomical cuts (two transversal for CN/SOC and IC and one 845 coronal for the MGB). The percentage of statistically significant voxels in experiment 1 that are 846 statistically significant in experiment 2 is reported together with the distance between the centroids 847 of activations between the two experiments in Figure 4-Figure Supplement 4 (for each individual 848 and in average across individuals). The unthresholded maps of both experiments (for each of 8/19 the six participants) are also visualized in *Figure 11* and are available at the online repository 850 851

(https://osf.io/hxekn/?view_only=be9ec398304344e8bb694a0658d77ed6) for inspection.

To produce group level results, the single subject statistical maps were warped to the reference 852 brainstem (subject 1) by applying the warping field obtained on the anatomical data. After projection 853 to the common space, single subject statistical maps were binarized and converted to a probabilistic 854 map by: 1) applying of a cluster size threshold of 3.37 mm³ (27 voxels in the 0.5 mm isotropic 855 anatomical space 2.5 voxels in the original functional resolution) and 2) summing maps across 856 subjects at each single voxel (i.e., a value of 10 indicates that all 10 subjects exhibited a statistically 857 significant response to sound presentation corrected for multiple comparisons and belonging to 858 a cluster of at least 27 yoxels in the anatomical space). The additional clustering allowed us to 850 further control for possible false positives by imposing a neuroanatomically plausible hypothesis 860 (i.e., none of our region of interest is smaller than 3.37 mm³ in volume). The same procedure was 861 also repeated by leaving one subject out (i.e., we generated probabilistic maps from 9 out of the 862 ten subjects each time leave one subject out). The leave-one-out probabilistic maps were then 863 back-projected to the anatomical space of the left out subject (i.e., the probabilistic map obtained from subjects 1 to 9 was back-projected to the anatomical space of subject 10). Unless otherwise stated, probabilistic maps are displayed with minimum threshold of at least three out of ten (or nine

⁸⁶⁷ for the leave one out maps) subjects exhibiting significant responses at each voxel. Unthresholded

⁸⁶⁸ probabilistic maps are available for inspection at the online repository.

We evaluated how well cluster localized on the basis of our probabilistic maps generalize 869 to new data. Figure 4 displays the statistically thresholded activation maps for each of the ten 870 participants in experiment 1 in three anatomical cuts (two transversal for CN/SOC and IC and one 871 coronal for the MGB) together with the probabilistic map obtained from the other nine participants 872 (thresholded by displaying voxels that are functionally significant in at least three out of nince 873 participants). In *Figure 4–Figure Supplement 1* we report the percentage of voxels in the leave 874 one out probabilistic maps that are statistically significant in the left out subject. The overlap 875 is reported together with the distance between the centroids of activations in the leave one 876 out probabilistic maps and the left out subject. The effect of the threshold on the probabilistic 877 maps is analyzed in Figure 4-Figure Supplement 2. The unthresholded maps (leave one subject 878 out and single subject) are also visualized in *Figure 10* and available at the online repository 879 (https://osf.io/hxekn/?view_only=be9ec398304344e8bb694a0658d77ed6) for inspection. 880 To compare the functional activation maps with histology data and post mortem MRI data, the

To compare the functional activation maps with histology data and post mortem MRI data, the probabilistic maps were projected to the MNI space using the warping field obtained from the anatomical dataset.

884 BigBrain data

Histology data were obtained by downloading the 100 µm version of the BigBrain (*Amunts et al.*,
2013) 3-D Volume Data Release 2015 (from https://bigbrain.loris.ca). We downloaded both the
original images and the dataset already aligned to MNI ICBM 152. The nuclei along the auditory
pathway (cochlear nucleus, superior olive, inferior colliculus and medial geniculate nucleus) were
manually segmented in the histology space image using ITK-SNAP (*Yushkevich et al.*, 2006) largely
following the definitions in *Moore* (1987) when possible.

⁸⁹¹ Correction of the alignment of the inferior colliculi to MNI

Upon visual inspection of the BigBrain image in the MNI ICM 152 space, we detected a major regis-892 tration error around the inferior colliculi (see *Figure 7* - second panel from the left). The registration 893 guality to MNI ICMBM 152 space in the rest of the brainstem was deemed satisfactory, but the the 894 region of the inferior colliculus required correction in order to perform a valid comparison with the 895 MRI data (in vivo and post mortem). Interestingly, the region of the colliculi of the BigBrain in the 896 original histology space appeared to be closer in location to the position of the inferior colliculus in 897 the MNI dataset (compare panel 1 and 3 in *Figure 7*) indicating that the highlighted misalignment 898 in the original BigBrain MNI dataset originated during the registration procedure. 890

To perform a new registration to MNI of the brainstem and thalamus of the BigBrain data that 900 observed the already correctly registered boundaries (e.g. the Pons) but corrected the region 901 around the inferior colliculus bilaterally, we followed N steps. First, we defined a region of interest 902 around the inferior colliculus using common anatomical landmarks that were visible in the BigBrain 903 MNI and MNI (2009b) T1. PD. T2 images and where aligned satisfactorily. Second, this region was cut 90/ out from the BigBrain MNI and replaced by the same region (i.e., defined by the same anatomical 905 landmarks) in the BigBrain histology space data (before projection to MNI). The convex hulls of the 906 region of interest in the BigBrain histology and in the MNI space were matched using 3-D optimal 907 transport as implemented in Geogram version 1.6.7 (Lévy. 2015: Lévy and Schwindt, 2018). Third. 908 the convex hull matched region of the the BigBrain histology space was used to replace the incorrect 909 region which was cut out at step 2. As a result of these three steps we obtained a version of the 910 BigBrain in MNI (BigBrain MNI - implanted) that had the inferior colliculus in the right position but 911 where the transitions between outside to inside of the region of interest that was corrected were 912 visible and not respecting of the topology. To correct for these residual errors, we performed a 913



Figure 7. The registration error around the inferior colliculus is visible bilaterally when comparing Panel 2 and Panel 3. The dashed lines indicate the correct shape (and location) of the colliculi in MNI space. The arrows point to the inferior colliculus (IC). The last panel shows the corrected BigBrain MNI dataset.

new FSL-FNIRT alignment between the original BigBrain in histology space and the BigBrain MNI 914 - implanted image. The resulting image (BigBrain MNI - corrected) preserved the actual topology 915 inside the brainstem and at the same time resulted in a correct alignment of the regions around 916

the inferior colliculus bilaterally (see *Figure 7* - right panel).

917

Post mortem MRI vasculature analysis 918

Gradient echo (GRE) MRI is sensitive to vasculature within the imaged tissue. To highlight vasculature 919

in the post mortem brainstem specimen, we computed the minimum intensity projection in coronal 920

sagittal and axial direction from the 50 µm isotropic voxel GRE MRI data over slabs of 1.1 mm in 921

thickness using Nibabel (Brett et al., 2017) and Numpy (Van Der Walt et al., 2011)). This image can 922

be seen in Figure 5 right column. 923

Diffusion MRI analysis 924

Post mortem diffusion 925

Before analysis, post mortem diffusion volumes were each registered to the first b0 volume using 926 an affine transformation in ANTs version 2.1.0 (Avants et al., 2011). To estimate white matter fiber 927

orientations, we used the constrained spherical deconvolution (CSD) model as implemented in DIPY 928 0.14 (Gorgolewski et al., 2011: Garvfallidis et al., 2014: Tournier et al., 2007) as a Nipype pipeline 929 (Gorgolewski et al., 2011). CSD posits that the observed diffusion signal is a convolution of the 930 true fiber orientation distribution (FOD) with a response function. DIPY's 'auto-response' function 931 estimates the fiber response function from a sphere of 10 voxels in the center of the sample above 932

a given fractional anisotropy (FA) threshold (0.5 in our study). We then estimated FOD peaks in 933 each voxel using DIPY's 'peaks-from-model' method with a 10° minimum separation angle and a 934

maximum of 5 peaks per voxel. 935

White matter fiber streamlines were estimated deterministically with DIPY's EudX method (Mori 936 et al., 1999: Garvfallidis, 2013) with 1,000,000 seeds per voxel, a 75° streamline angle threshold. 937 and an FA termination threshold of 0.001 (since data outside the specimen sample were already 938 masked to 0). 939

To define regions of interest (ROIs) for the fiber display, the auditory structures manually 940 delineated in the post mortem T2*-weighted MR images were transformed to diffusion space 941 using ANTs, and global streamlines were filtered by considering only the voxels in each one of the 942 ROIs as a seed and further constrained by using all auditory ROIs as tractography waypoints. This 943 resulted in a high-resolution, high-quality auditory-specific subcortical tractogram, which were then 944 visualized in TrackVis 0.6.1 (Wang et al., 2007). 945

946 In vivo diffusion

7T in vivo dMRI data was corrected for distortions with the HCP pipeline *Glasser et al.* (2016);
 Sotiropoulos et al. (2013). Specifically, geometric and eddy-current distortions, as well as head
 motion, were corrected by modeling and combining data acquired with opposite phase encoding
 directions Andersson et al. (2003); Andersson and Sotiropoulos (2015, 2016). The data were then

masked to include just the brainstem and thalamus, matching the post mortem specimen.

⁹⁵² Similar to the post mortem analysis, we estimated diffusion FODs with a CSD model imple-⁹⁵³ mented in DIPY with response function FA threshold of 0.5. Peaks were extracted with a minimum

⁹⁵³ mented in DIPY with response function FA threshold of 0.5. Peaks were extracted with a minimum ⁹⁵⁴ separation angle of 25°. White matter connectivity was estimated with deterministic tractography

⁹⁵⁴ separation angle of 25°. White matter connectivity was estimated with deterministic tractography ⁹⁵⁵ throughout the brainstem and thalamus, again using DIPY's EudX algorithm (*Mori et al., 1999*:

Garyfallidis, 2013) with 1,000,000 seeds per voxel, a 45° streamline angle threshold, and an FA

957 termination threshold of 0.023.

For the tractography in the in vivo data we used subcortical auditory ROIs as defined by the analysis of the functional data (i.e., regions that exhibited significant [corrected for multiple comparisons] response to sound presentation in at least three out of ten subjects). The functional ROIs were transformed to individual diffusion space and used as tractography seeds, with all other

⁹⁶² auditory ROIs as waypoints, producing a subcortical auditory tractogram for each in vivo subject.

963 Data and code availability

⁹⁶⁴ Unprocessed in vivo data are available at (https://openneuro.org/datasets/ds001942). Atlas seg ⁹⁶⁵ mentations and tractography streamlines are available through the Open Science Framework
 ⁹⁶⁶ (https://osf.io/hxekn/). Processing and analysis resources, including links to all data and software
 ⁹⁶⁷ used in this paper, are available at https://github.com/sitek/subcortical-auditory-atlas (*Sitek and*

⁹⁶⁸ Gulban, 2019). See Figure 8 for an overview of currently available data and code (full resolution

⁹⁶⁹ version available at our code repository).

970 Animated 3D volume renderings

Video animations in *Figure 9, Figure 10* and *Figure 11* were created using pyqtgraph (v0.10.0, http://www.pyqtgraph.org/) volume rendering. The t-value maps were clipped to 0-20 range and scaled to 0-255 range. These t-values are 3D volume rendered by assigning the corresponding gray value to each voxel as well as the alpha channel (transparency). Which means that lower values are

closer to black and translucent. Animation frames were generated by rotating camera one degree at a time for 360 degrees. Additive rendering was used for 2D projections to provide depth vision

at a time for 360 degrees. Additive rendering was used for 2D projections to provide depti
 (i.e., for preventing voxels closest to the camera from seeing values inside the clusters.).

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Figure 8. Summary of data processing steps, including availability of data and code.



Figure 9. One frame of volume rendered animations for comparing histology (BigBrain), post-mortem MRI, in-vivo MRI unthresholded positive t-values group average and in-vivo MRI clusters of significant activity overlapping in at least 4 subjects in each voxel.

Figure 9-video 1. 3D volume rendered comparisons in MNI space.



Figure 10. One frame of volume rendered animations for single subject statistical maps. (Left)positive t-values (middle) after thresholding (right) leave-one-out probabilistic map (\geq 4)). Viewing angle here is similar to Figure *Figure 1*.

Figure 10-video 1. Subject 01 Figure 10-video 2. Subject 02 Figure 10-video 3. Subject 03 Figure 10-video 4. Subject 05 Figure 10-video 5. Subject 06 Figure 10-video 6. Subject 07 Figure 10-video 7. Subject 08 Figure 10-video 8. Subject 09 Figure 10-video 9. Subject 10 Figure 10-video 10. Subject 11



Figure 11. One frame of volume rendered animations for Subject 01 statistical maps (experiment 1 positive t-values & thresholded (col 1-2) and experiment 2 positive t-values & thresholded (col 3-4)). Viewing angle here is similar to Figure *1*.

Figure 11-video 1. Subject 01 experiment 1 vs experiment 2.

Figure 11-video 2. Subject 02 experiment 1 vs experiment 2.

- Figure 11-video 3. Subject 05 experiment 1 vs experiment 2.
- Figure 11-video 4. Subject 09 experiment 1 vs experiment 2.

Figure 11-video 5. Subject 10 experiment 1 vs experiment 2.

Figure 11-video 6. Subject 11 experiment 1 vs experiment 2.

Figure 11-video 7. Group average (N=6) unthresholded positive t-values for experiment 1 vs experiment 2.

988 Glossary

989 Anatomical abbreviations

- **AVCN** Anteroventral cochlear nucleus.
- **CN** Cochlear nucleus.
- **CNVIII** 8th nerve, vestibulocochlear nerve.
- **DCN** Dorsal cochclear nucleus.
- IC Inferior colliculus.
- LGN Lateral geniculate nucleus.
- ⁹⁹⁰ **LSO** Lateral superior olive.
 - **MGB/MGN** Medial geniculate body/nucleus.
 - **MNTB** Medial nucleus of the trapezoid body.
 - MSO Medial superior olive.
 - **PVCN** Posteroventral cochlear nucleus.
 - **SOC** Superior olivary complex.

991 MRI acquisition abbreviations

- **7T** 7 Tesla.
- dMRI diffusion magnetic resonance imaging.
- FOV Field of view.
- **fMRI** functional magnetic resonance imaging.
- **GRAPPA** Generalized auto-calibrating partially parallel acquisitions.
- MB Multi-band.
- **MPRAGE** Magnetization prepared rapid acquisition gradient echo.
- ⁹⁹² MRI Magnetic resonance imaging.
 - **PDw** Proton density weighted.
 - **SI-T1w** Short inversion time T1-weighted.
 - **T1w** T1-weighted.
 - T2*w T2*-weighted.
 - TE Echo time.
 - **TR** Repetition time.

993 Data analysis abbreviations

- **CSD** Constrained spherical deconvolution.
- **FA** Fractional anisotropy.
- **FDR** False discovery rate.
- **FOD** Fiber orientation distribution.
- **GLM** General linear model.
- **HCP** Human connectome project.
- **HRF** Hemodynamic response function.
- ⁹⁹⁴ **ICBM** Internation Consortium for Brain Mapping.
 - M0 T2 signal with no diffusion weighting.
 - MD Mean diffusivity.
 - MNI Montreal Neurological Institude.
 - MSMT Multi-shell multi-tissue
 - **ODFs** Orientation distribution functions.
 - **ROI** Region of interest.

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Figure 3–Figure supplement 1. Post mortem human diffusion-weighted MRI tractography (from 200 µm isotropic voxels) with anatomically defined subcortical auditory seeds, downsampled to 200 µm but undilated. Streamlines that passed through manual segmentations of the medulla and optic tracts were excluded. 10 percent of streamlines are visualized for clarity. Top right: connectivity heatmap of subcortical auditory structures. Bottom right: Streamlines that pass through the right inferior colliculus.

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Figure 3-Figure supplement 2. Post mortem human diffusion-weighted MRI tractography with anatomically defined subcortical auditory seeds. MRI data were downsampled from 200 μ m to 1050 μ m to match in vivo data acquisition and then processed in the same manner as other diffusion tractography analyses. Streamlines that passed through manual segmentations of the medulla and optic tracts were excluded. 10 percent of streamlines are visualized for clarity. Top right: Connectivity heatmap of subcortical auditory structures.



Figure 4-Figure supplement 1. Correspondence between single subject activation maps and leaveone-out functional probabilistic maps. Leave-one-out probabilistic functional maps are thresholded to identify voxels that are significantly responding to sounds in at least three of nine participants. The overlap represents (per region of interest) the percentage of the voxels on the leave-one-out probabilistic maps that is significantly responding to sounds in the left out subject. For each region of interest we also report the distance in mm between the centroids of the leave-one-out probabilistic maps and the centroids of the regions significantly responding to sounds in the left out subject. The last column represents the average overlap and distance across participants per region and error bars represent the standard error across the participants.



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Figure 4-Figure supplement 4. Correspondence between single subject activation maps Experiment 1 and Experiment 2. All maps are thresholded for significance (FDR-q=0.05 and p<0.001; see methods for details). The overlap represents (per region of interest) the percentage of the voxels significantly active in Experiment 1 that is significantly responding to sounds in Experiment 2. For each region of interest we also report the distance in mm between the centroids of the regions significantly responding to sounds in both experiments. Videos are provided in the appendix that visualize thresholded and unthresholded maps for each of the individual participants. The last column represents the average overlap and distance across participants per region and error bars represent the standard error across the participants.



Figure 4–Figure supplement 5. Effect of spatial smoothing on functional activation maps. Functional activation maps obtained from Experiment 1 in two participants with and without applying spatial smoothing (1.5mm FWHM Gaussian smoothing) prior to the statistical analysis. Maps are thresholded for statistical significance (FDR-q = 0.05 & p<0.001; see Methods for details)). For each participant, CN/SOC and IC are shown in transversal cuts, MGB is shown in a coronal cut.





Figure 5-Figure supplement 1. In vivo anatomical group average images in MNI space.



Figure 5–Figure supplement 2. Anatomical images from MNI ICBM 152 2009b dataset compared to BigBrain histology in MNIspace (left column).



