Cross-Modal Plasticity in Higher-Order Auditory Cortex of Congenitally Deaf Cats Does Not Limit Auditory Responsiveness to Cochlear Implants

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Congenital sensory deprivation can lead to reorganization of the deprived cortical regions by another sensory system. Such cross-modal reorganization may either compete with or complement the “original” inputs to the deprived area after sensory restoration and can thus be either adverse or beneficial for sensory restoration. In congenital deafness, a previous inactivation study documented that supranormal visual behavior was mediated by higher-order auditory fields in congenitally deaf cats (CDCs). However, both the auditory responsiveness of “deaf” higher-order fields and interactions between the reorganized and the original sensory input remain unknown. Here, we studied a higher-order auditory field responsible for the supranormal visual function in CDCs, the auditory dorsal zone (DZ). Hearing cats and visual cortical areas served as a control. Using mapping with microelectrode arrays, we demonstrate spatially scattered visual (cross-modal) responsiveness in the DZ, but show that this did not interfere substantially with robust auditory responsiveness elicited through cochlear implants. Visually responsive and auditory-responsive neurons in the deaf auditory cortex formed two distinct populations that did not show bimodal interactions. Therefore, cross-modal plasticity in the deaf higher-order auditory cortex had limited effects on auditory inputs. The moderate number of scattered cross-modally responsive neurons could be the consequence of exuberant connections formed during development that were not pruned postnatally in deaf cats. Although juvenile brain circuits are modified extensively by experience, the main driving input to the cross-modally reorganized higher-order auditory cortex remained auditory in congenital deafness.

Key words: cochlear implant; cross-modal plasticity; deaf white cat; deafness; deprivation; development

Significance Statement

In a common view, the “unused” auditory cortex of deaf individuals is reorganized to a compensatory sensory function during development. According to this view, cross-modal plasticity takes over the unused cortex and reassigns it to the remaining senses. Therefore, cross-modal plasticity might conflict with restoration of auditory function with cochlear implants. It is unclear whether the cross-modally reorganized auditory areas lose auditory responsiveness. We show that the presence of cross-modal plasticity in a higher-order auditory area does not reduce auditory responsiveness of that area. Visual reorganization was moderate, spatially scattered and there were no interactions between cross-modally reorganized visual and auditory inputs. These results indicate that cross-modal reorganization is less detrimental for neurosensory restoration than previously thought.

Introduction

Congenital sensory loss triggers cross-modal plasticity in the deprived sensory cortex (Rauschecker, 1995; Bavelier et al., 2006; Stein and Stanford, 2008; Merabet and Pascual-Leone, 2010; Stein et al., 2014). Cross-modal reorganization is likely related to enhanced abilities in the remaining senses as best documented in deaf individuals (Neville and Lawson, 1987; Levänen et al., 1998;
A reorganization toward a new modality might interfere with restoration of the original sensory input and thus could close sensitive periods for the therapy, for example, with cochlear implants (CIs) (Kral, 2007). This is related to a fundamental question of developmental neuroscience: can experience cause a sensory area to change the major driving modality and thus restructure the brain’s connectome extensively? A differential role of auditory areas in cross-modal reorganization has been demonstrated, with visual reorganization specific to higher-order auditory fields in congenital deafness (Kral et al., 2005; Lomber et al., 2010; Leonard et al., 2012). Although the primary auditory cortex has been the focus of many studies (for review, see Kral and Sharma, 2012), so far, no information is available on auditory responsiveness of higher-order auditory fields in congenital deafness. Furthermore, it remains unclear whether cross-modal responses are even distributed within a reorganized field or if reorganized neurons cluster spatially. To answer these questions, the present study determined the auditory responsiveness and the quantitative relation between visual responsiveness and original auditory responsiveness in a secondary sensory area known to take over a cross-modal behavioral function in deafness (Lomber et al., 2010), the auditory dorsal zone (DZ). The investigation was performed at the level of individual neurons using large-scale microelectrode mapping.

The functional impact of cross-modal reorganization on the re-stored auditory responsiveness using CIs on neuronal responses in area DZ was compared between adult hearing cats (HCs) and congenitally deaf cats (CDCs). To allow large-scale mapping of the whole field with multielectrode arrays, instead of searching the optimal stimulus (possible only in few neurons), we used stimuli likely to evoke responses in the majority of neurons related to the given modality and mapped the whole fields. We used an acute preparation, allowing us to separate reorganizations of bottom-up and lateral interactions from other influences. We recorded responses to both modalities at the same positions.

The present study demonstrates a modest visual cross-modal reorganization and a dominance of auditory responsiveness in the cross-modally reorganized DZ. The data do not indicate interference between the reorganized and original modality and show a substantial preservation of the original sensory inputs, even after cross-modal reorganization, with an absence of bi-modal interactions.

Materials and Methods

Animals. Experiments were performed in five adult congenitally deaf white cats (Heid et al., 1998) and four adult hearing controls (>12 months of age, 7 female and 2 male). HCs had normal hearing with click-evoked auditory brainstem response (ABR) thresholds <40 dB SPLpeak. CDCs had been identified from the colony of deaf white cats using ABR procedures at 1.3–1.5 volume percentage isoflurane concentration in a 1:2 mixture of O2/N2O. Adequacy of anesthesia depth and the animals’ physiological state was monitored by means of ECG, heart rate, end-tidal CO2, muscle tone, and EEG signals. End-tidal CO2 was maintained at <4.5%. Core temperature was kept >37.5°C using a homeothermic blanket. Physiological state was additionally monitored by analyzing capillary blood every 12 h for blood gas concentration, pH, bicarbonate concentration, base excess, glycemia, and oxygen saturation. A modified Ringer’s solution containing bicarbonate (according to the base excess) was infused intravenously. Each animal’s head was fixed in a stereotactic holder (Horsley–Clarke).

We then retested the hearing status in all animals by measuring ABR. For this purpose, a small trephination was drilled at the vertex and ABR responses were recorded with an epidural silver-ball electrode (diameter 1 mm) referenced to a silver-wire neck electrode. HCs had click response thresholds <40 dB SPL. The absence of ABR responses in deaf animals confirmed deafness as diagnosed during early hearing screening soon after birth (Fig. 1A).

Stimulation was performed using feline CIs (MED-EL, custom-made, five channels, distance between contacts 1 mm) inserted bilaterally through the round window into scalae tympani of both ears. This involved exposing both bullae and ear canals. To prevent electrophyony in HCs, the hair cells were destroyed pharmacologically before cochlear implantation. This was achieved by intracochlear instillation of 300 μl of 2.5% neomycin sulfate solution over a 5 min period and subsequent rinsing using Ringer’s solution. The absence of hearing was subsequently confirmed by the absence of ABRs (Fig. 1). To test the functionality of the CIs and to determine the stimulation threshold, we then determined electrical ABRs (eABRs; Fig. 1). eABR thresholds were measured between the epidural silver electrode to a reference in the neck (amplification 100,000×, sixth-order band-pass filter 10–10,000 Hz). Electrical brainstem responses were recorded for a biphasic pulse (200 μs/phase) at different current levels with bipolar stimulation between all possible bipolar electrode contact combinations.

As in a previous study in a larger group of animals (Tillein et al., 2012), the eABR thresholds were not different between HCs and CDCs [Fig. 1D; HCs: left ear 154 ± 43 μA; right ear: 139 ± 28 μA; CDCs: left ear 161 ± 79 μA; right ear 152 ± 38 μA; Wilcoxon–Mann–Whitney (WMW) test, α = 5%].

Recording of electrophysiological activity. For electrophysiological recording, a trephination above the lateral (suprasylvian) sulcus was performed, exposing the dorsal auditory cortex (DZ) on the lateral bank and visual medial suprasylvian sulcus areas (anterior and posterior medial suprasylvian sulcus areas, AMLS/PMLS, subsequently referred to jointly as MLS; Fig. 2A). The dura was opened and the cortex surface was covered with silicone oil. A modified Davies chamber was positioned around the trephination site to stop a layer of agarose and a closure was created melted bone wax after the electrode arrays were set in place. Cortical activity was recorded with two linear 16 site multielectrode arrays for which the interelectrode distance was 150 μm, surface area 177 μm², impedance 1–2 MΩ (NeuroNexus). The multielectrode arrays were positioned and inserted using micromanipulators, which were attached to the stereotactic frame (TSE Systems). The penetration angle was kept constant throughout the experiment. At least one penetration in each investigator area was stained using DiI (1,1'-dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine perchlorate; Invitrogen) that was applied to the noncontact side of the multielectrode array using a micropipette (Eppendorf). An epidural vertex silver-ball electrode served as an electric reference for both multi electrode arrays. The recorded signals were amplified 5000–10,000 times with a Neurolynx amplifier, band-pass filtered (1 Hz–9 kHz), digitized (at a sampling rate of 30 kHz), and stored on a computer.

Mapping procedure and stimulation design. We mapped the dorsal auditory cortex and visual areas along the medial part of the suprasylvian sulcus in deaf and HCs (Fig. 1B). Multielectrode arrays were inserted on both sides of the sulcus at a distance of ~500 μm from the midline of the sulcus, thus penetrating the dorsal auditory cortex and the two visual area in MLS. With an interelectrode distance of 150 μm and the uppermost site inserted just into the cortex, the tip of the electrode shank was inserted

Bavelier and Neville, 2002; Bavelier et al., 2006; Auer et al., 2007; Bottari et al., 2010, 2014.)
with a micromanipulator to a depth of $\sim 2400 \mu m$ ($\pm 100 \mu m$) from the cortical surface. The depth or position of the multielectrode array was not changed after insertion to search for activity. At each penetration position and after the closure of the modified Davies chamber with agarose and bone wax, we allowed the multielectrode array to settle and stabilize the recordings for 10–20 min. Each block of sensory stimulation was initiated by 15 min of recording of spontaneous activity and was concluded by 15 min recording of spontaneous activity to exclude drifts of the general state of the animals. This approach allowed us to ensure a constant light anesthetic state. We paid attention to avoid deep anesthesia with burst suppression to prevent possible abnormal heteromodal responses in the cortex (Land et al., 2012). During the neuronal recordings, isoflurane concentration was reduced to 1.0–1.2 volume percentage, and adequacy of anesthesia depth was monitored to ensure comparable anesthesia levels and to avoid periods of burst suppression.

The number of spontaneously active sites was similar for HCs and CDCs, with more spontaneous activity in the visual cortex than in the auditory dorsal cortex in both groups (CDCs: 63% in DZ vs 75% in MLS; WMW test, $p = 0.03$; HCs: 53% in DZ vs 75% in MLS; WMW test, $p = 0.029$).

We analyzed and included all electrode sites in the statistics. In the text, "position" refers to the penetration location in the cortex (Fig. 2B) and "site" refers to electrode sites deep in the cortex, of which there were 16 for each electrode array. In total, we analyzed 1440 recording sites (720 in the auditory and 720 in the visual cortex) in HCs and 1632 sites (816 in the auditory and 720 in the visual cortex) in CDCs.

**Sensory stimulation.** eABR thresholds were determined at the beginning of the experiment (Fig. 1). The eABR threshold of the respective ears then was used as a reference current level. Electrical stimulation was wide bipolar between the apical-most and the basal-most contact of the implant, covering cochlear positions with characteristic frequencies $> 10$ kHz (Kral et al., 2009).

**Auditory stimulation.** For intracortical recordings, pulses were presented binaurally, from 2 dB below to 6 dB above the eABR threshold in 1 dB steps for each ear. The electrical stimulus was a triplet of biphasic pulses (200 $\mu s$/phase at 500 pulses/s, giving a total stimulation time of 4.4 ms) applied in bipolar configuration between the basal-most electrode and the apical-most electrode of the CI. Pulse levels were randomized and the interstimulus intervals were 1000 ms. Each electrical stimulus was repeated 30 times.

**Visual stimulation.** Visual stimuli were generated in MATLAB (The MathWorks) with the Psychophysics Toolbox (Brainard, 1997). Stimuli were presented on a TFT display (Model 2009wdt; Dell) at a 28 cm distance in front of the contralateral eye. In analogy to the electrical pulse, we used a visual flash stimulus to study general visual responses. This stimulus is simple and broadly activates neurons in the visual system, both in the magnocellular and parvocellular subsystems. We presented 100 ms full-field flashes with positive contrast (white flash) or negative contrast (black flash). Each type of flash was repeated 50 times with an interstimulus interval of 1000 ms consisting of a gray background. Furthermore, to include apparent movement into the stimulus, square-wave phase reversal gratings of different orientations ($0^\circ$, $45^\circ$, $90^\circ$, $135^\circ$) and spatial frequencies (0.1–2.0 cycles/degree) were used for visual stimulation.

**Bimodal stimulation.** To investigate interactions between the visual and auditory responses, bimodal stimulation was used. Visual stimulus (full-field flash, 16.7 ms duration, one frame, 60 Hz refresh rate) was combined with auditory stimulation (triplet of biphasic pulses, 200 $\mu s$/phase, 500 Hz) at 6 dB above threshold. The onset of the stimuli varied across a range from $\sim 30$ to 30 ms.

**Histology.** After the experiment, the animals were transcardially perfused. After thoracotomy, 0.5 mL of heparin (Liquemin; Hoffman-La Roche) was injected into the left ventricle. Then, 2 L of phosphate buffer (0.1 M, pH 7.4) and 2 L of fixative (2.5% glutaraldehyde and 2.0% formaldehyde) were infused transcardially with pressure $> 100$ mmHg. After 24 h of postfixation in 4% formaldehyde, the brain was excised from the skull, photographed, and a block containing the investigated cortical areas was cryoprotected in 30% sucrose, frozen, and cut in frontal plane

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**Figure 1.** Cochlear implants in HCs and CDCs. A, In HCs, acoustic stimulation generated evoked brainstem responses that disappeared after destruction of hair cells by intracochlear application of neomycin. This treatment was required to eliminate electrophonic effects (see text for details). B, In CDCs, the conclusion of the juvenile hearing screening procedure was confirmed at the beginning of the acute experiments by an absence of brainstem responses for stimulation at high sound pressure levels. C, Cochlear implants used for the present experiments were donated by MED-EL. Stimulation was with apical-most and basal-most contact (5 mm distance) in wide bipolar configuration with charge-balanced biphasic pulses at 200 $\mu s$/phase. D, eABRs in HCs after implantation. E, eABRs in CDCs after implantation. F, eABR thresholds (connected by the line) show no systematic difference in threshold between deaf and hearing animals or between the ears.
The firing rate was defined as the mean number of spikes in the interval 30 ms after stimulus onset. We subsequently excluded sites if neuronal activity was modulated by electrical stimulation via the CI satisfying a fixed statistical criterion for all sites (DZ and PMLS). Mean firing rates. We additionally analyzed “continuous multiunit activity” (cMUA), including the all spike amplitudes (also the so-called “hash”) in 50 μm sections using a cryotome (Leica). The sections were first photographed in fluorescence mode to reveal the DiI (Keyence, BZ-9000). Subsequently, the sections were alternatively stained with Nissl and antibodies against SMI 32 (Mellott et al., 2010), allowing us to identify the borders of field A1, DZ, and lateral sulcus regions (LLS and MLS). All stained sections were then digitized and the penetrations were reconstructed (Keyence, BZ-9000). The DiI-stained penetrations were combined with photographs of SMI-32-stained (same) sections.

**Data analysis.** Multitunits (MUs) were derived by band-pass filtering the raw signal between 700 Hz and 9 kHz. First, we determined all spike activities with amplitudes that exceeded a fixed threshold of 50 μV (amplifier noise level <15 μV), separating large spikes. A fixed spike threshold was used to ensure comparability between groups and multitunit firing rates. We additionally analyzed “continuous multitunit activity” (cMUA), including all the spike amplitudes (also the so-called “hash”) using the 700 Hz high-pass filtered, rectified, and squared signal without thresholding. This signal was denoted as cMUA.

**Analysis of ongoing activity.** To derive ongoing multiunit rate from 15 min intervals before and after stimulation, for each site, we randomly selected 100 intervals with a 1 s window length and calculated the mean rate of these 100 intervals for all sites. We subsequently excluded sites if firing rate was <0.1 Hz during the entire period (nonactive sites).

**Analysis of responses.** Sites in CDCs and HCs were defined as responsive if neuronal activity was modulated by electrical stimulation via the CI satisfying a fixed statistical criterion for all sites (DZ and PMLS). Mean auditory responses were calculated for each of the 9 stimulation levels (−2 dB to 6 dB above the eABR threshold). Response strength was defined as the mean number of spikes in the interval 30 ms after stimulus onset; that is, the time window when auditory responses occurred. Auditory responses were ordered by stimulus level and the correlation coefficient between response strength and stimulation level was determined. If the coefficient was significant (p < 0.05) or if the unit significantly responded above baseline in five of the nine stimulation levels (two-sided t test against baseline activity, p < 0.05), then the site was considered responsive to auditory input. For all responsive sites, the response latency was defined as the peak of the response at 6 dB above threshold. Visual responsiveness was considered as present in those neurons that showed a significant increase in firing rate within the 60 ms after the stimulus (α = 5%). Visually evoked activity was tested against baseline multitunit activity before stimulus onset. Both rates were compared with a two-sided t test (p < 0.05) and, if found significant, were collected as a response for further analysis. Response latency was defined as peak latency.

**Presence of bimodal enhancement was tested at those stimulus delays where peak responses overlapped. Quantification was performed using the enhancement index (EI) (Meredith and Stein, 1983)

\[
E.I. = \frac{VA - \max(V, A)}{\max(V, A)}
\]

where VA is the firing rate with bimodal stimulation, V and A are the firing rates of visual alone and auditory alone stimulation, respectively, and max denotes the maximum function. To determine the additive or superadditive character of bimodal responses, the additive index (AI) was also used (King and Palmer, 1985)

\[
A.I. = \frac{VA - (V + A)}{V + A}
\]

If not stated otherwise, all data are presented in the form of mean ± SD. Data from animals were not pooled, statistical comparisons were performed using the enhancement index (EI) (Meredith and Stein, 1983)
formed at the animal level (5 CDCs vs 4 HCs). We used a nonparametric two-tailed WMW test with 5% significance level to compare data between cortical areas and between HCs and CDCs.

**Results**

The present study investigates the sensory function of a higher-order auditory area (DZ) in adult CDCs. Cooling inactivation of this area previously demonstrated a causal relation to supranormal visual behavior in CDCs ([Lomber et al., 2010](#)). Here, neuronal responses were collected simultaneously with two DiI-stained single-shank multielectrode arrays (16-channel) inserted to cover all cortical layers in DZ and the visual fields in the middle suprasylvian region (MLS; Palmer et al., 1978; Fig. 2). In total, we analyzed 1632 sites in CDCs \( (n_{\text{visual}} = 816 \text{ and } n_{\text{auditory}} = 816) \) and 1440 recording sites in HCs \( (n_{\text{visual}} = 720 \text{ and } n_{\text{auditory}} = 720) \).

**Visual reorganization of the dorsal auditory cortex of CDCs**

Visual stimuli allowing standardized large-scale mapping of cortical areas with multielectrode arrays (flashes and phase-reversal gratings) were presented using a screen covering the frontal visual field.

Statistical significance of visual responsiveness was determined by comparing prestimulus and poststimulus firing rate. Visual responsiveness did not differ between flashes and gratings. Because flashes represented a stimulus better corresponding to the auditory stimulus, in the following, we focused on the responses to flash stimuli. From all responsive sites, mean time histograms were computed (Fig. 3A, B). Response to the visual flash showed prominent ON and OFF response typical for visual areas. A distinct difference between CDCs and HCs in visual responsiveness was only found in DZ (Fig. 3A), where CDCs
exhibited visual responses, whereas HCs did not. In contrast, the visual areas showed similar visual responses in CDCs and HCs (Fig. 3B). This demonstrates a dissociation of the effect of congenital deafness on MLS neurons and DZ neurons.

The latencies of visual responses in the “deaf” DZ were higher than response latencies in the MLS, the control visual area (Fig. 3C; CDCDZ = 41; median 35 ms with median absolute deviation, MAD, of 7.6 ms; CDCMLSL = 165; median 30 ms with MAD 7.3 ms; HCMMLSL = 184; median 30 ms with MAD 11.2 ms; WMW test, p = 0.005). The visual evoked firing rates in the deaf DZ were lower than the rates in the visual areas, but differed significantly from activity in the HCs, where no detectable evoked responses were present (Fig. 3D; CDCDZ = 41; 1.1 ± 1.6 spikes/stimulus; HCDZ = 0.2 ± 0.2 spikes/stimulus; WMW test, p = 0.007). In contrast, in the visual cortex, latency and firing rate of responses did not differ significantly (Fig. 3C,D; CDCMLSL = 165; 2.4 ± 2.9 spikes/stimulus; HCMMLSL = 184; 2.4 ± 3.0 spikes/stimulus; WMW test, p = 0.37) and the latencies corresponded to those observed in MLS previously (Dinse and Krüger, 1994; Nowak and Bullier, 1997).

We found a moderate number of visually responsive recording sites in DZ of CDCs (Fig. 3E; CDC:5.1 ± 3.0%; HC:1.3 ± 0.9%; WMW test, p = 0.03; for higher responsiveness with units with smaller amplitude, see below). Whereas visual responsiveness in DZ was thus emphasized in CDCs, visual responsiveness in the visual cortex was similar to that in HCs (Fig. 3E; CDC:24 ± 13%; HC:26 ± 10%; WMW test, p = 0.85). The overall number of DZ units responsive to visual stimulation in CDCs constituted ~20% of visually responsive units in the visual areas. Together, these results demonstrate cross-modal reorganization in field DZ of CDCs, a possible correlate of previous behavioral observations (Lomber et al., 2010).

The position of visually responsive sites across the “deaf” auditory cortex did not show a systematic gradient (Fig. 3F). Visual responses in the deaf animals were sparsely distributed in the dorsal auditory cortex, whereas, in the visual cortex, visual responses were abundant both in CDCs and HCs (Fig. 3F).

In total, the absence of a difference in visual responsiveness in the control visual areas (MLS) between the investigated groups demonstrate that the general state of the animals was comparable between the two groups and cannot account for the difference in DZ responsiveness. Therefore, the difference in visual responsiveness in field DZ between HCs and CDCs was the result of the different hearing experience.

Preserved auditory responsiveness in DZ

To activate the auditory system and test auditory responsiveness, we used electrical stimulation with CIs in both groups of animals (Fig. 1). Implantation was performed acutely during the final experiments: none of the animals had previous experience with the electrical stimulation. However, HCs had normal hearing and their auditory system developed under the influence of hearing. Electrical “hearing thresholds” were quantified by eABRs (Fig. 1). For comparability, we also stimulated HCs with CIs (Fig. 1). Their hair cells were destroyed pharmacologically before implantation to prevent electrophonic responses.

The responsiveness to the auditory stimulus was again determined statistically and mean poststimulus time histograms were computed from responsive sites (Fig. 4A,B). CI-evoked responses were found in DZ of both CDCs and HCs (Fig. 4A,B). CDCs had slightly shorter auditory response latencies than HCs (Fig. 4C, CDCDZ = 149; median 11 ms with MAD 3.7 ms and HCMLSL = 113; median 12 ms with MAD 3.6 ms; WMW test, p = 0.03), indicating a change in the response properties in the deaf DZ compared with HCs. CI-evoked firing rates in DZ of CDCs and HCs did not differ significantly (Fig. 4D; CDCDZ = 149; 0.69 ± 0.7 spikes/stimulus; HCDZ = 113; 0.64 ± 0.65 spikes/stimulus; WMW test, p = 0.74). We found no significant differences in the number of responsive sites in DZ between groups (Fig. 4E; CDCDZ = 149; 20 ± 9%; HCDZ = 113; 16 ± 7%; WMW test, p = 0.38). Auditory responsive sites were thus similarly present throughout the entire field DZ in both groups of animals (Fig. 4F); however, there was a tendency toward cluttering in the rostral part.

Both in HCs and CDCs, our selection criterion identified sites in the visual cortex as being auditory responsive. However, these were chance-level selections as a consequence of the automatic statistical criterion because poststimulus time histograms in the responsive sites did not reveal a response (Fig. 4B,F).

The results demonstrate that field DZ preserves its auditory responsiveness after congenital deafness despite cross-modal reorganization. The data show a dissociation of the effects of deafness on visual and auditory responsiveness in field DZ, with an increase in visual responsiveness in deaf DZ and no effect in MLS and similar auditory responsiveness in both investigated fields and both groups of animals, but moderate differences in the properties of the responses between deaf and hearing DZ (response latency).

Auditory responsiveness in DZ was similar in CDCs and HCs, with CDCs differing from HCs in the additional presence of visual responses. To exclude an influence of our spike selection criterion (large units), we additionally included an analysis of all units by analyzing the 700 Hz high-pass-filtered squared signal without thresholding (cMUA), including also small units. With cMUA, the incidence of visual responses in DZ increased substantially, but the difference between the groups remained (Fig. 5A, CDC: 17 ± 4%; HC: 5 ± 1%; Mann–Whitney test, p = 0.03). Also for cMUA, auditory responsive sites in the visually reorganized deaf DZ exceeded visually responsive neurons (Fig. 5B, cMUA: auditory: 48 ± 23%, visual: 17 ± 9%; MU: auditory: 20 ± 9%, visual: 5.1 ± 4%). Therefore, whereas the sensitivity of both measures differed, the auditory-responsive sites remained more frequent than visually responsive sites, so this observation was not due to the previous spike selection criterion (Fig. 5C).

Absence of bimodal interactions

Notably, only 1.4% (n = 12) of all recorded sites in DZ of CDCs responded to both auditory and visual stimuli (i.e., were classified as bimodal units). We computed the bimodal EI for those stimulus timings where peak unimodal responses would overlap. The EI was between −20% and 20% in 10 of 12 units with a mean of −10 ± 18 (n = 12). The bimodal EI was in most cases negative with a mean of −20 ± 15 (n = 12), indicating a subtractive character of the interactions. However, it should be kept in mind that these bimodal units represented only a negligible number of the total sites recorded and had rather weak responses.

In addition, we tested possible interactions of bimodal stimuli on firing rates within the visually reorganized deaf DZ in a more general approach. Because visual and auditory responses had different latencies and bimodal interactions depend on the timing between stimuli, visual and auditory stimuli were presented at different time delays relative to each other. In doing so, we measured bimodal interactions and responsiveness as a function of different time delays of bimodal stimuli. Provided a true modulation of one response by the other modality was present, a distinct peak in the responsiveness for certain time delays should
show up in this analysis. We inspected each site and then pooled the units to compare the mean results statistically.

First, we tested auditory neurons in the deaf DZ that did not respond to visual stimulation alone. These might respond to a combination of visual and auditory input. For the auditory-responsive units, the auditory stimulus was kept constant at 6 dB above threshold and the visual flash (16.7 ms duration) was shifted in time relative to the auditory stimulus. The visual stimulus did not influence auditory responsiveness in the set of purely auditory-responsive units in the deaf DZ ($n_{1005}/H11022/0.3$ spikes/stimulus). We computed the average for each cat and then tested the modulation by time delay ($\text{Fig. 5D}$, comparing modulation of average responses for each cat, ANOVA, $F_{(15,64)} = 0.04, p = 0.98$). Therefore, the set of auditory-

Figure 4. CI eABRs in CDCs and HCs. A, Comparison of responses to CI stimulation in DZ of CDCs and HCs. Poststimulus time histograms show mean responses with SEM error bars for responsive sites at 6 dB above eABR thresholds in the deaf animals. B, Visual areas in MLS did not respond to CI stimulation. C, Comparison of latencies of responsive sites in CDCs and HCs. Tukey boxplots show the median with 1.5 interquartile range whiskers. D, Response rates were similar in CDCs and HCs in all investigated fields. Plots denote mean with SD error bars. E, Number of auditory-responsive sites was significantly higher in the auditory dorsal cortex than in the visual MLS region. Plots denote mean with SD error bars. F, Responsiveness to flashes arranged along the straightened lateral sulcus (dashed line) in CDCs and HCs (left vs right) relative to the dorsal end of the posterior ectosylvian sulcus (distance = 0 mm). Number of responsive sites is depicted by the size of the rectangles and nonresponsive sites by dots; animals are coded by color. Size-count examples are shown on the left side of the plot.
responsive sites and the set of visually responsive sites in the dorsal auditory cortex seem to be essentially distinct without interactions between sensory inputs when tested with electrical stimuli in the auditory domain.

Discussion

The present data demonstrate cross-modal reorganization in the secondary auditory field of CDCs, a possible correlate of previous behavioral observations. The cross-modal visual reorganization, however, did not reduce auditory responsiveness substantially even after long-term congenital deafness (\textsuperscript{[1]}H11022 1 year). Cross-modally responsive units were sparsely distributed in DZ and were approximately three times less frequent than auditory responsive units. The few units with audiovisual responses did not show bimodal enhancement.

The comprehensive mapping of auditory and visual fields demonstrated a dissociation of deprivation effects between the areas and sensory inputs: the higher-order auditory cortex developed visual responsiveness in CDCs. In contrast, there was no effect of lifelong auditory deprivation on the adjacent visual cortex. The data thus demonstrate a dissociation of the effect of deafness on responsiveness of auditory and visual areas.

Although cross-modal reorganization is well documented (Rauschecker, 1995; Bavelier et al., 2006; Auer et al., 2007; Merabet and Pascual-Leone, 2010; Ding et al., 2015) and related to enhanced abilities in the remaining senses in deaf individuals (Neville and Lawson, 1987; Levänen et al., 1998; Bavelier et al., 2006; Bottari et al., 2014; Stein et al., 2014), the original function of the reorganized neural tissue has not been explored before at the level of neurons. The absence of adverse effects of cross-modal reorganization on auditory responsiveness has implications for visual communication before cochlear implantation in children born deaf.

Methodological considerations

The cross-modally reorganized DZ is known to be involved in visual motion detection (Lomber et al., 2010). Here, we used flashes and phase-reversal gratings known to evoke strong responses in motion-sensitive and pattern-sensitive neurons (Kral et al., 2003). Strong responsiveness with such stimuli in neighboring motion-sensitive visual areas confirmed their effectiveness in driving the neurons (cf. Dinse and Krüger, 1994; Nowak and Bullier, 1997). The responsiveness to gratings and flashes was similar. Although we cannot exclude that a more specific visual stimulus could have caused stronger visual responsiveness in field DZ, this argument can never be disproven. The present standardized approach is optimal for large-scale mapping purposes. Furthermore, the flash stimulus is a good visual correlate of the wide bipolar electrical stimulus used for auditory stimulation, allowing direct comparison of the inter-modal balance in responsiveness.

We did not investigate the detailed functional properties of CI responses. The present study, as a first step, focused on responsiveness; more detailed future studies might reveal further differences in auditory properties of deaf DZ beyond the latency differences observed here. Here, the deaf brains were untrained, completely naive to auditory stimuli, and still stronger auditory than visual responsiveness was observed in their field DZ. Hearing experience with CI would likely further increase the auditory

Figure 5. Absence of bimodal responses in the visually reorganized deaf DZ. A, cMUA responses (including all spikes) also reveal differences in visually responsive sites in the dorsal auditory cortex between CDCs and HCs. Plots denote mean with SD error bars. B, Direct comparison of auditory and visual responses in the deaf auditory cortex. Responsiveness of the visually reorganized dorsal auditory cortex of CDCs showed more auditory-responsive sites than visually responsive sites. Plots denote mean with SD error bars. C, Comparison of ratio between auditory responsive and visually responsive sites for MUA and cMUA. D, Responsiveness to auditory stimulation was not affected by visual stimulation at different onset asynchronies. Plots denote mean with SD error bars. E, Responsiveness of visual response was not affected by auditory stimulation at different onset asynchronies. Plots denote mean with SD error bars. F, Nonresponsive sites were not affected by different stimulus asynchronies. Plots denote mean with SD error bars.
Cross-modal reorganization in congenital deafness

Cross-modal reorganization in DZ corresponds to data from humans (Finney et al., 2001, 2003; Bavelier et al., 2006). Ectopic subcortical and lateral projections into DZ (from the adjacent MLS, area 7, and areas 19/20/21) are present in CDCs (Barone et al., 2013). Cross-modal visual responses in the deaf DZ had latencies slightly longer than responses in the adjacent MLS (Palmer et al., 1978), consistent with a direct connection from MLS to the deaf DZ in CDCs. Visually responsive sites in the deaf DZ were moderate in number and accounted for ∼30% of the auditory-responsive sites. The present findings are thus consistent with the perceptual-narrowing hypothesis (Ghazanfar and Schroeder, 2006; Lewkowicz, 2014; Maurer and Werker, 2014) because such sparse responsiveness may result from patchy excitatory connections that appear developmentally and are normally pruned during postnatal life (Innocenti and Price, 2005; Innocenti, 2011).

Here, we have provided data for visual flash stimulation because this stimulus is independent of the receptive field and other properties of the individual neuron. With regard to the moderate 5% or 17% of visually responsive sites in the cross-modally reorganized cortex (compared with 25% or 60% of auditory-responsive sites), it needs to be emphasized that it is not clear what minimal amount of cross-modal reorganization is necessary to enhance performance. Distributed attentional and top-down modulations are likely to complement (boost) the cross-modal responsiveness described here. Conflicting findings in the direction of change of sensory capabilities in blind or deaf individuals, which can range from an increase to a decrease (Pavani and Bottari, 2012), indicate that cortical cross-modal reorganization may be variable depending on a given individual’s (top-down) compensation strategy.

Auditory activation of the secondary auditory cortex in deafness

The deaf DZ remained responsive to the auditory input. The auditory responsiveness was not different from HC, demonstrating that the secondary auditory cortex develops its basic functional responsiveness even in absence of any hearing. This corresponds to persistence of basic functionality within the deprived sense of congenitally sensory-deprived humans (Collignon et al., 2009; Leonard et al., 2012; Striemen-Amit et al., 2012a, 2012b; Cardin et al., 2013, 2016; Gandhi et al., 2015). The present study provides the physiological substrate of this phenomenon. Supporting this, sighted and congenitally blind subjects show specificity for motion or object form in the occipital cortex regardless of input modality, even in absence of visual experience (Voss and Zatorre, 2012; Peelen et al., 2014).

Some studies have reported negative interference from visual inputs for auditory processing in deaf subjects (Doucet et al., 2006; Buckley and Tobey, 2011; Sandmann et al., 2012; Chen et al., 2016). Conversely, beneficial cross-modal effects for speech understanding have been reported, mainly in the postlingually deaf (Giraud et al., 2000; Rouger et al., 2007; Stroopahl et al., 2015). These latter studies, however, investigated postlingually deafened groups, whereas the present study investigated the congenitally deaf.

The increase in visual responsiveness in the present study was not related quantitatively to a similar decrease in auditory responsiveness. Therefore, units that were visually responsive were not strongly coupled to the auditory system. We did not find bimodal interactions in the deaf DZ. In cats, visual deprivation interferes with development of multimodal integration (Wallace et al., 2004, 2006; Carriere et al., 2007, 2008). The absence of vision during development interferes with the development of multimodal interactions in the superior colliculus, but less with visual responsiveness (Yu et al., 2013; Stein et al., 2014). Even rearing under omnidirectional noise has a similar effect (Xu et al., 2014). Furthermore, early visual deprivation has detrimental effects on multimodal integration in association area 7 (adjoining the MLS areas; Carlson et al., 1987). A loss of bimodal interactions and a dominance of visual inputs have been documented in prelingually deaf children after cochlear implantation (Schorr et al., 2005). Moreover, congenital cataracts prevented audiovisual integration in humans (Guerreiro et al., 2015). The absence of bimodal interactions in the present study is explicable by the absence of early auditory experience that is essential for development of audiovisual interactions.

Visual recruitment of higher-order auditory areas might close the sensitive periods for therapy of deafness (Kral, 2007; Lomber et al., 2010). Even though some neurons took over visual functions and were auditory nonresponsive in the present study, the general auditory responsiveness was not affected significantly. Therefore, the major limiting factors after sensitive periods appear to be the developmentally reduced plasticity and central processing deficits within the auditory modality itself (Kral et al., 2002, 2009, 2013; Fallon et al., 2009). It needs to be emphasized that absence of a difference in auditory DZ responsiveness in the present study does not demonstrate the absence of functional auditory deficits. Such deficits have been described numerous times in the auditory system (for review, see Kral, 2013) and are likely also present in the dorsal auditory cortex. The present study, however, did focus only on responsiveness and its balance between visual and auditory stimulation.

Bimodal training has improved auditory spatial localization and changed auditory responses in field A1 of deaf ferrets equipped with CIs (Isaiah et al., 2014). This demonstrates that after focused training bimodal processing can become beneficial for some (but maybe not all, see Schorr et al., 2005; Guerreiro et al., 2015) functions after restoration of hearing, probably via top-down influences from the frontal cortex (Isaiah et al., 2014). The present data are consistent with these findings.

The present study has an important translational aspect. Communication before cochlear implantation in deaf children is an important predictive factor of implantation outcomes (Tait et al., 2000). Some argue that deaf children should communicate via cued speech or sign language before cochlear implantation (Lyness et al., 2013; Campbell et al., 2014; Mellon et al., 2015). However, many clinicians are concerned with potential negative interactions between modalities (Geers et al., 2011; Mellon et al., 2015). Indeed, the cross-modally reorganized units in the present study did not respond to auditory stimulation. However, they represented only a minor portion of neurons in DZ. Consequently, the present data disprove that cross-modal reorganization eliminates functional auditory inputs to the higher-order dorsal auditory cortex.
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