Complementary adaptive processes contribute to the developmental plasticity of spatial hearing

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Spatial hearing evolved independently in mammals and birds and is thought to adapt to altered developmental input in different ways. We found, however, that ferrets possess multiple forms of plasticity that are expressed according to which spatial cues are available, suggesting that the basis for adaptation may be similar across species. Our results also provide insight into the way sound source location is represented by populations of cortical neurons.

Sound localization mechanisms are remarkably plastic early in life, as demonstrated by the effects of monaural deprivation on the development of spatial hearing. Different forms of adaptation, however, have been demonstrated in different species. Barn owls show a compensatory adjustment in neuronal sensitivity to the binaural cues that are altered by hearing loss. In contrast, ferrets become more dependent on the unchanged spectral cues provided by the intact ear for horizontal localization. These results imply that the basis for developmental plasticity may vary across species, which is consistent with evidence that sound localization evolved independently and differs in several respects between mammals and birds. However, if the auditory system can express multiple forms of plasticity, spatial processing in different species may be more similar than previously thought. Here we show that monaurally deprived ferrets can also adapt to abnormal binaural cues, a process comparable to that used by barn owls. The nature of this adaptation has important implications for the encoding of spatial information by neuronal populations in primary auditory cortex (A1).

We reared ferrets, each with an earplug in one ear. This alters the binaural cues that normally dominate how animals perceive the azimuth of a sound but preserves the monaural localization cues at the intact ear. Because monaural cues require a comparison of input at different sound frequencies to provide reliable spatial information, they cannot be used to localize narrowband sounds. If ferrets can adapt to unilateral hearing loss only by becoming more reliant on the intact monaural spectral cues, they should therefore be unable to localize narrowband stimuli.

We measured the ability of ferrets to judge the azimuth of high-frequency narrowband sounds. Like humans and other mammals, ferrets rely primarily on interaural level differences (ILDs) to localize these sounds. We observed clear differences in performance across groups, even for individual animals (Fig. 1a–c). Control ferrets performed this task accurately under normal hearing conditions (Fig. 1a,d), but made large errors when the left ear was occluded with an earplug (Fig. 1b,d). However, much smaller errors were made by monaurally deprived ferrets whose left ears had been plugged from hearing onset (juvenile-plugged ferrets) (Fig. 1c,d), suggesting that an adaptive change in ILD processing had occurred in these animals.

We tested this by performing bilateral extracellular recordings in A1 of juvenile-plugged and control ferrets (Fig. 2a). We measured ILD sensitivity by calculating the slope of each neuron's firing rate–ILD function (Fig. 2b and Online Methods). A positive ILD indicates that the sound level was higher in the right ear (right-ear greater); thus, a positive ILD slope indicates that the neuron preferred right-ear-greater stimuli.

Neurons recorded in left and right A1s of seven control ferrets under normal hearing conditions had, on average, positive and negative ILD slopes, respectively, demonstrating that they preferred stimuli whose levels were higher in the opposite ear (Fig. 2c). When we used digital filtering to introduce a virtual earplug to the left ear, ILD sensitivity in both hemispheres declined (Fig. 2c), indicated by ILD slopes that approached 0. In contrast, neurons recorded from seven juvenile-plugged ferrets in the presence of a virtual earplug had steeper ILD slopes, showing that they retained sensitivity to ILD. In these animals, however, neurons recorded from both hemispheres preferred right-ear-greater stimuli (slope > 0) (Fig. 2c), suggesting that juvenile plugging shifted ILD sensitivity toward the nonplugged ear.

We next measured the way inputs from the two ears interact in the absence of the virtual earplug by varying the sound level in the ipsilateral ear while keeping the contralateral level fixed. In controls, this revealed the presence of binaural suppression (Online Methods) at large positive and negative ILDs, resulting in mean binaural interaction functions that peaked (best ILDs) when the sound level was greater in the ear opposite the recording site (Fig. 2d). We hypothesized that compensation for the attenuating effects of the earplug in the juvenile-plugged ferrets would manifest as shifts in the best ILD (Supplementary Fig. 1). Although the changes observed were insufficient to fully compensate for the ~45–dB attenuation produced by the earplug, we found that the best ILD shifted in the predicted direction in both left (10–dB shift in median) and right (20–dB shift in median) A1 (Fig. 2e,f), confirming that adaptive changes in binaural processing had taken place.

A popular model of sound localization in mammals is based on the relative activity of neuronal populations in each hemisphere, each with a preference for contralateral sound sources. We therefore built a population decoder (Supplementary Fig. 2) that classifies ILDs on single trials using the difference in mean activity between the two hemispheres (‘hemispheric decoder’; Online Methods), and found that this decoder worked well for controls under normal hearing.

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BRIEF COMMUNICATIONS

Figure 1 Behavioral judgments of sound azimuth using narrowband high-frequency stimuli. (a–c) Joint distributions of stimulus and response, expressed as degrees (deg) azimuth, for a control ferret with normal hearing (a) and a control (b) and juvenile-plugged (JP) ferret (c) wearing an earplug in the left ear. Grayscale represents the number of trials (n) corresponding to each stimulus-response combination. (d) Mean unsigned error for control and earplugged ferrets, normalized so that 0 and 1 correspond to perfect and chance performance, respectively. Error bars show bootstrapped 95% confidence intervals. Controls wearing an earplug (n = 6 ferrets) made larger errors than normal hearing controls (n = 4; P < 0.001, bootstrap test). While wearing an earplug, juvenile-plugged ferrets (n = 2) made smaller errors than acutely plugged controls (P < 0.001, bootstrap test).

conditions (Fig. 3 and Supplementary Fig. 3). However, in the presence of a virtual earplug, neurons in each A1 of juvenile-plugged animals tended to prefer ILDs that favor the nonplugged right ear (Figs. 2c and 3a). Hemispheric differences in activity therefore remained relatively constant as a function of ILD (Fig. 3a), causing the hemispheric decoder to perform very poorly for juvenile-plugged animals (Fig. 3b and Supplementary Fig. 4).

Within each hemisphere of the juvenile-plugged ferrets, however, we observed some neurons (left A1, 71/375, 18.9%; right A1, 43/209, 20.6%) that preferred left-ear-greater stimuli. We therefore created a second population decoder (‘opponent-process decoder’) that classified ILDs using the difference in mean activity between neurons that preferred right-ear–greater and left-ear–greater stimuli, irrespective of the hemisphere in which neurons were recorded. The opponent-process decoder performed much better than the hemispheric decoder (P < 0.001; bootstrap test) (Fig. 3c,d). This representation of ILD was robust to changes in both population size and stimulus-independent interneuronal correlation, and remained when the analysis was restricted to neurons recorded from a single hemisphere (Supplementary Fig. 5) or when the decoder was required to first learn the ILD preferences of different neurons (Supplementary Figs. 6 and 7).

Previous studies have indicated that intra-hemispheric opponent processing and population codes that exploit tuning-curve heterogeneity might have an important role in spatial hearing. Our results show that this may be even more important after unilateral hearing loss, as a decoder that compares activity between neurons with opposing ILD preferences provided a close match to the animals’ behavioral performance (Fig. 3d). Despite differences in the cortical representation of ILD, the opponent-process decoder also provided a better match to the behavioral performance of controls, particularly in the presence of the virtual

Figure 2 Adaptive processing of ILDs in A1. (a) Bilateral extracellular recordings were made under normal hearing conditions or with a virtual earplug in the left ear. (b) Spike rate versus ILD, for one unit (i.e., activity recorded on a single electrode), for different average binaural levels (thin lines) plus mean rate-ILD function (thick). Linear regression (dotted line) was used to quantify the slope of the mean rate-ILD function, which was then normalized (Online Methods). (c) ILD slopes (±95% confidence intervals) obtained for neurons recorded ipsilaterally (left A1; black symbols) or contralaterally (right A1; gray symbols) to the virtual earplug. In normal hearing controls, neurons in both hemispheres preferred stimuli whose levels were higher in the opposite ear (total n = 292 units; P < 0.001; bootstrap test). ILD sensitivity in controls declined when a virtual earplug was introduced (left hemisphere, n = 126 units, P < 0.001; bootstrap test; right hemisphere, n = 166 units, P < 0.001; bootstrap test). In contrast, ILD slopes in juvenile-plugged animals were steeper under plugged conditions (left hemisphere, n = 375 units, P < 0.001; right hemisphere, n = 209 units, P < 0.001; bootstrap test). (d) Mean binaural interaction (±s.e.m.) as a function of ILD across all units recorded from controls under normal hearing conditions. Data are plotted separately for left (n = 142 units, black) and right (n = 177 units, gray) A1. Best ILDs for each hemisphere are indicated by arrows. (e) Binaural interaction functions (mean ± s.e.m.) in juvenile-plugged ferrets under normal hearing conditions. (f) Unit best ILDs in each A1 of control and juvenile-plugged (JP) animals. Horizontal lines indicate medians (± interquartile range). Range (whiskers) and approximate 95% confidence intervals (tapers) are also shown. Arrows indicate the adaptive direction in which binaural interaction functions should shift, relative to controls, for JP animals to compensate for the hearing loss experienced during development. In juvenile-plugged ferrets, best ILDs shifted in the predicted direction in both left (n = 379 units; P < 0.001; bootstrap test) and right (n = 226 units; P < 0.001; bootstrap test) A1.
Thus, some form of opponent processing may be a crucial aspect of representing auditory space, particularly in A1, irrespective of the animals’ experience.

Our results show that ferrets can localize high-frequency sounds accurately using abnormal binaural cues and that a concomitant remodeling of ILD representations in auditory cortex may underlie this process. Because ferrets also adapt to unilateral hearing loss, both during development and in adulthood, by becoming more dependent on the unchanged spectral cues provided by the intact ear, these data show that different processes can contribute to the developmental plasticity of spatial hearing within the same species. If spectral cues provide sufficiently precise information about object location, abnormal binaural cues may be ignored or given very little weight. Conversely, if, as in this study, spectral cues are compromised or unavailable, the auditory system may rely more on the abnormal binaural cues to localize sounds accurately. Our finding that the brain can utilize different strategies to adapt to altered inputs therefore highlights the remarkable flexibility of auditory spatial processing.

**METHODS**

Methods and any associated references are available in the online version of the paper.

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**AUTHOR CONTRIBUTIONS**

This study was conceived and designed by P.K. and A.J.K. P.K. did the behavioral experiments, modeling and data analysis. P.K. and J.C.D. made the electrophysiological recordings. P.K., J.C.D. and A.J.K. wrote the manuscript.

**COMPETING FINANCIAL INTERESTS**

The authors declare no competing financial interests.

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ONLINE METHODS

General. Twenty-three ferrets, both male and female, were used in this study, of which seven were raised with an earplug and the rest raised normally. All sample sizes were chosen on the basis of previous behavioral and neurophysiological studies in our lab. Animals were assigned to different experimental groups by technicians who were blind to the purposes of the study. Portions of the data were collected by people who were unaware of either the project aims or the experimental group. However, group allocation was by necessity known to the lead investigator throughout the study. All behavioral and neurophysiological testing was performed in animals that were at least 5 months old, some of which had been tested previously on a broadband sound-localization task. All procedures were performed under licenses granted by the UK Home Office and met with ethical standards approved by the University of Oxford. General details concerning animal welfare and behavioral testing have been described previously.

Briefly, ferrets were housed either individually or in small groups (<8) within environmentally enriched laboratory cages. Each period of behavioral testing lasted for up to 14 d, during which drinking water was provided through correct performance of the behavioral task and via additional supplements provided at the end of each day. Dry food was made available at all times, and free access to water was given between testing periods.

The left ear of the juvenile-plugged ferrets was first fitted with an earmold (EAR Classic) between postnatal day 25 (P25) and P29, shortly after the age of hearing onset in this species, and was thereafter monitored routinely and replaced as necessary. All details relating to earplugging have been described previously. Briefly, earplugs were replaced either with or without sedation (Domitor, 0.1 mg per kilogram of body weight (mg/kg); Pfizer). To secure the earplug in place, the concha of the external ear was additionally filled with an ear mold impression compound (Otoform-K2, Dreve Otoplastik). The status of both ears was checked routinely using otoscopy and tympanometry, with any accumulation of cerumen removed under sedation. Earplugs were periodically removed so that normal hearing was experienced ~20% of the time, amounting to a total of 3 d in any 15-d period, with periods of nonocclusion evenly spread through-out the diurnal light cycle, which exhibited seasonal variation that sought to replicate that found naturally. This intermittent hearing-loss paradigm is the same as that used in our previous study in which adaptive reweighting of spatial cues was demonstrated and was followed because (i) previous studies have reported maladaptive changes in neuronal ILD sensitivity in mammals raised with a continuous unilateral hearing loss and (ii) we wanted to more closely replicate the type of fluctuating hearing loss that is prevalent among children with otitis media with effusion. This procedure was repeated continuously until the conclusion of the experiments. Earplugs were kept in place for at least 4 d before terminal electrophysiological experiments, and were only removed immediately before surgery.

Behavioral testing. Behavioral testing was performed during the hours of daylight in a soundproof chamber, within which was situated a custom-built circular mesh enclosure. Details of this setup have been described elsewhere. Animals initiated each trial by standing on a platform located in the middle of the testing chamber and licking a spout positioned at the front of this platform for a variable delay, after which a stimulus was presented from one of 12 loudspeakers (FRS-8; Visaton) equally spaced around the periphery of the enclosure. Animals registered their response by approaching the location of one of the loudspeakers. Behavioral responses and trial initiation were monitored using infrared detectors, with experimental contingencies controlled by Matlab (Mathworks). Correct responses were rewarded with water delivered via spouts situated beneath each loudspeaker. Animals were trained from ~P150 onward to approach the location of sounds to receive a water reward.

All stimuli were 200 ms in duration, generated with a sampling rate of 97.6 kHz, and had 10 ms cosine ramps applied to the onset and offset. Stimuli were either broadband (0.5–30 kHz) or narrowband (1/6 of an octave wide with a center frequency of 15 kHz) bursts of noise, with different stimulus types randomly interleaved across trials. On the majority (~75%) of trials, broadband noise stimuli were presented, the level of which varied between 56 and 84 dB SPL in increments of 7 dB. For these stimuli, and these stimuli alone, incorrect responses were followed by correction trials on which the same stimulus was presented. Persistent failure to respond accurately to correction trials was followed by ‘easy trials’ on which the stimulus was repeated continuously until the animal made a response. The addition of broadband stimuli, which are easier to localize because all available cues are present, was necessary because some animals found it very difficult to localize narrowband stimuli while wearing an earplug, performing very close to chance. The provision of broadband stimuli, and corresponding easy trials, therefore helped keep animals motivated to perform the sound-localization task accurately.

In most test sessions, narrowband stimuli were presented at 84 dB SPL. In some sessions, however, narrowband stimuli were also presented at 56 dB SPL. This was done primarily to determine whether plugged animals were relying on monaural level cues available to the nonplugged ear to guide behavioral responses. If this was the case, changes in sound level would be expected to produce shifts in response bias, which can be measured by the mean signed error. Although controls wearing an earplug showed shifts in the mean signed error when sound level was changed (P < 0.05; post hoc test; interaction between group and sound level, P = 0.039; ANOVA), implying some reliance on monaural level cues available to the intact ear, this was not the case in juvenile-plugged ferrets wearing an earplug or normally reared animals tested under normal hearing conditions (P > 0.05; post hoc test).

Behavioral analyses. For analysis purposes, stimulus and response locations in the front and rear hemifields were collapsed. This provided a measure of performance that is unaffected by front-back errors, which are unavoidable when localizing short-duration narrowband stimuli, and which are thought to primarily reflect a failure in spectral, rather than binaural, processing. We then calculated the discrepancy between the lateral position of the stimulus and that of the response to generate estimates of behavioral error for each trial. The mean unsigned error, which measures error magnitude, was then calculated to determine the ability of animals to judge the lateral position of narrowband sounds. When testing controls wearing an earplug, we wanted to measure performance in animals that were relatively naïve to the earplug. For this reason, each of these animals was tested in only a few sessions, which meant that very few narrowband trials were obtained for each individual. It was therefore necessary to pool the data across animals before applying bootstrap procedures to the data obtained on individual trials.

Neurophysiological recordings. Ferrets were anesthetized for recordings to provide the stability needed for presenting large stimulus sets via earphones a sufficient number of times to obtain reliable estimates of ILD sensitivity. This also meant that we could eliminate changes in arousal level or attentional modulation as factors contributing to any differences observed between the control and juvenile-plugged ferrets. For analysis purposes, stimulus and response locations in the front and rear hemifields were collapsed. This provided a measure of the response to generate estimates of behavioral error for each trial. The mean unsigned error, which measures error magnitude, was then calculated to determine the ability of animals to judge the lateral position of narrowband sounds. When testing controls wearing an earplug, we wanted to measure performance in animals that were relatively naïve to the earplug. For this reason, each of these animals was tested in only a few sessions, which meant that very few narrowband trials were obtained for each individual. It was therefore necessary to pool the data across animals before applying bootstrap procedures to the data obtained on individual trials.

Neural signals were band-pass filtered (500 Hz to 3 kHz), amplified, and digitized (25 kHz) using TDT System 3 processors (Tucker-Davis Technologies). Stimuli were generated in Matlab, amplified (TDT headphone amplifier) and presented via earphones (Panasonic, RP-HV298) that were situated at the entrance to each ear canal. Action potentials were extracted in Brainwave (Tucker-Davis...
Technologies) from a mixture of single units and small multi-unit clusters, which were pooled for subsequent analyses to increase statistical power. All further data analyses were carried out using Matlab.

Stimuli consisted of 200-ms bursts of spectrally-flat broadband (0.5–30 kHz) noise, generated at a sampling rate of 97.6 kHz, and with 10 ms cosine ramps applied to the onset and offset. Average binaural level (ABL) was varied between 62 and 78 dB SPL in increments of 4 dB and interaural level differences (ILDs) were set to vary between ±16 dB (before any effects of a virtual earplug), varying in increments of 8 dB. Specific ILDs were created by adjusting the sound level in opposite directions in the two ears, which meant that the sound level presented to a given ear varied between 54 and 86 dB SPL. ILD stimuli were presented either under normal hearing conditions or with a virtual earplug in the left ear. Virtual earplugs were created by applying an additional filter to the signal presented to the left ear, which delayed the input by 110 μs and attenuated it by 15–45 dB (varying as a function of frequency, as shown in ref. 6), similar to the effects of a real earplug. Each unique stimulus was repeated 30 times, and different acoustical conditions were presented in blocks.

**Neurophysiological analyses.** Spike rates were calculated for each trial within a 75-ms window spanning the onset response of each acoustically driven unit. These were then used to assess the ILD sensitivity of individual units as well as neuronal populations. ILD slope was estimated, for each unit, by calculating the mean spike rate as a function of ILD. The slope of this mean rate-ILD function was then quantified using linear regression, with the resulting slope value expressed, as in previous studies of ILD coding27,28, as a percentage of the maximal response for each unit (r_{max}). The equation defining this procedure was therefore as follows:

\[
\text{ILDslope} = \left( \frac{r_{max} - r_{min}}{\text{ILD}_{max} - \text{ILD}_{min}} \right) \times 100 \frac{r_{max}}{r_{max}}
\]

where r_{max} and r_{min} represent the maximum and minimum values of the linear fit and ILD_{max} and ILD_{min} denote the maximum and minimum ILD values presented. Positive ILDs were more intense in the right ear (right-greater) and negative ILDs were therefore assigned to units that responded more strongly to right-greater ILDs, with negative slope values assigned to units that preferred left-greater ILDs.

Binaural interaction (β) was calculated, for each unit and ILD, as follows:

\[
\beta_{\text{ILD}} = \frac{2 \times (\text{ILD}_{\text{PSI}} - \text{r}_{\text{Contra}})}{r_{\text{max}}}
\]

where r_{PSI} and r_{Contra} indicate the spike rates elicited by equivalent monaural stimulation of the ears ipsilaterally and contralaterally to the recording site (i.e., presenting the same left- and right-ear inputs separately), respectively, and r_{max} denotes the maximum spike rate exhibited by a particular unit. Negative binaural interaction values therefore reflected sublinear summation of motor units, indicating more suppressive binaural interaction, with positive values indicating supralinear summation and, therefore, binaural facilitation. Under normal hearing conditions, similar mean firing rates were observed for both hemispheres in juvenile-plugged ferrets (n = 379, n = 226; P = 0.38; bootstrap test) and controls (n = 177, n = 142; P = 0.12; bootstrap test).

Population decoders classified responses by comparing the mean activity between two distinct subpopulations of units in a manner similar to that described previously16. In all cases, population decoders were implemented using custom-written Matlab code. Subpopulations of units were defined either on the basis of the hemisphere in which they were recorded (hemispheric decoder) or whether they preferred right-ear-greater ILDs or left-ear-greater ILDs (opponent-process decoder) (Supplementary Fig. 2). For the opponent-process decoder, analyses were applied to the data obtained from either one or both hemispheres. In all cases, spike rates were normalized for each unit by dividing by the maximum single-trial response exhibited by that unit across trials. This normalization step effectively regularized the weight given to each unit. Without it, units with low firing rates would have contributed very little to the decoder’s decision, even if those units were very sensitive to ILDs. Although the precise details of this normalization procedure had very little impact on decoder performance, we opted for the procedure reported here because it produced marginally smaller errors (across all groups of animals) than alternative approaches. In the training phase, a population-averaged rate-ILD function was computed by calculating, for each ILD, the mean normalized spike rate across all units belonging to each sub-population. In doing so, data were averaged across different trials and ABLs.

The difference between these population-averaged rate-ILD functions was then used, during the test phase, to infer stimulus ILD from single-trial population responses as both ILD and ABL were varied. This was done by calculating, for a single trial, the difference in mean activity between the two neuronal subpopulations, comparing this value with the difference between population-averaged rate-ILD functions, and identifying the ILD that was associated with the most similar population-averaged response. In all cases, data were cross-validated using a leave-one-out procedure. Decoder performance was quantified by calculating the discrepancy between the stimulus ILD and the ILD-classification given by the decoder on each trial. The mean unsigned error across trials was then calculated and normalized so that values of 0 and 1 respectively indicated perfect and chance performance.

Because different numbers of units were recorded for different subpopulation types and groups of animals, which would be expected to affect decoder performance, we ran the population analysis using a fixed number of units chosen at random from the total number recorded for each subpopulation type in a particular group. We then repeated this sampling procedure 1,000 times and averaged performance across repetitions to generate a mean estimate of decoder performance for each group. To assess the relationship between decoder performance and the number of units used, this analysis was repeated for different population sizes (Supplementary Fig. 5). For formal comparisons between different decoder types (Fig. 3), we elected to use a population size of 200 units. Although this choice was to a certain extent arbitrary, it was guided by the fact that the performance associated with this population size lies on the asymptotic portion of the exponential fits. In other words, performance does not improve dramatically as population size is increased further. On the other hand, smaller population sizes enable better estimates of the variability produced by sampling different units.

Averaging the activity of many neurons can improve decoder performance because the activity of individual neurons is noisy. When this noise is independently distributed for each neuron, it can be cancelled out by averaging. However, noise that is shared by many neurons recorded simultaneously does not cancel when averaged, which can limit decoder performance29. This type of correlated noise can be produced in neuronal populations by common fluctuations in background activity that is independent of the stimulus. We investigated the effects of these noise correlations by measuring decoder performance while the order of stimulus presentation was randomized independently (‘shuffled’) for each simultaneously recorded unit, a procedure which removes stimulus-independent correlations between units30. Performance was then compared with that of decoders applied to the unshuffled data (Supplementary Fig. 5).

It is important to note that, whereas our unshuffled data contain noise correlations in the activity of simultaneously recorded data, noise correlations are necessarily absent for units recorded at different times. Our unshuffled data can therefore be understood as a set of neuronal ensembles recorded separately, with noise correlations existing within but not between these different ensembles. This means that our data most likely underestimate the impact of noise correlations, which means that real neural performance may be slightly worse than that obtained by applying our decoders to unshuffled data. Previous work, however, has shown that noise correlations diminish as the spatial separation of neurons is increased31. It is therefore likely that noise correlations in the brain exist within, but to a much lesser extent between, local neuronal ensembles.

In real neural systems, the readout of neural activity is also likely to be corrupted by noise intrinsic to the readout process. To simulate this, we therefore added a small amount of normally-distributed random noise to the decision-making process described above in order to estimate the robustness of population representations. In the presence of small amounts of noise, however, performance changed very little from that observed in its absence.

**Statistical analyses.** Confidence intervals at the 95% level were estimated empirically for different measures using 10,000 bootstrapped samples, each of which was obtained by resampling with replacement from the original data. These samples

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were then used to construct bootstrapped distributions of the desired measure, from which confidence intervals were derived32. A bootstrap procedure was also used to assess the significance of group differences. First, the difference between two groups was measured using an appropriate statistic (for example, difference in means, t-statistic, or rank-sum statistic). The data from different groups were then pooled and resampled with replacement to produce two new samples, and the difference between these samples was measured using the same statistic as before. This procedure was subsequently repeated 10,000 times, which provided an empirical estimate of the distribution that would be expected for the statistic of interest under the null hypothesis. This bootstrapped distribution was then used to derive a P value for the difference observed in the original sample. In all cases, two-sided tests of significance were used. Although bootstrap tests were used because they make fewer distributional assumptions about the data, conventional parametric and non-parametric statistical tests were also performed and produced very similar results to those obtained by bootstrapping.

**Code availability.** Matlab code was used to control behavioral testing, create stimuli for neurophysiological testing, and perform all analyses. In all cases, Matlab version 7.0.4 or 2010a was used (Mathworks). Portions of the code used, for which Mathworks does not own the copyright, can be made available on request.

A Supplementary Methods Checklist is available.

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