Spatial Sensitivity in Field PAF of Cat Auditory Cortex

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Abstract

We compared the spatial tuning properties of neurons in two fields (A1 and PAF) of cat auditory cortex. Broadband noise bursts of 80-ms duration were presented from loudspeakers throughout 360° in the horizontal plane (azimuth) or 260° in the vertical median plane (elevation). Sound levels varied from 20 to 40 dB above units' thresholds. We recorded neural spike activity simultaneously from 16 sites in field PAF and/or A1 of α-chloralose-anesthetized cats. We assessed spatial sensitivity by examining the dependence of spike count and response latency on stimulus location. In addition, we used an artificial neural network (ANN) to assess the information about stimulus location carried by spike patterns of single units and of ensembles of 2–32 units. The results indicate increased spatial sensitivity, more uniform distributions of preferred locations, and greater tolerance to changes in stimulus intensity among PAF units relative to A1 units. Compared to A1 units, PAF units responded at significantly longer latencies, and latencies varied more strongly with stimulus location. ANN analysis revealed significantly greater information transmission by spike patterns of PAF than A1 units, primarily reflecting the information transmitted by latency variation in PAF. Finally, information rates grew more rapidly with the number of units included in neural ensembles for PAF than A1. The latter finding suggests more accurate population coding of space in PAF, made possible by a more diverse population of neural response types.

INTRODUCTION

An intact auditory cortex is necessary for normal sound-localization behavior in humans (Greene, 1929; Klingon and Bontecou, 1966; Sanchez-Longo and Forster, 1958; Wortis and Pfeiffer, 1948; Zatorre and Penhune, 2001) and cats (Jenkins and Masterton, 1982; Jenkins and Merzenich, 1984). Generally, these studies have demonstrated contralesional deficits in sound localization following unilateral lesions of the auditory cortex and more global deficits following bilateral damage, although recent human data suggest a more complete representation of space in the right cerebral hemisphere (Zatorre and Penhune, 2001). In the midbrain, the superior colliculus presents an example in which space is represented through the topographic organization of spatial receptive fields (Middlebrooks and Knudsen, 1984; Palmer and King, 1982). These two sets of observations together raise the possibility that sound-source locations are represented topographically in the mammalian auditory cortex. The spike rates of many auditory cortical neurons are modulated by changes in sound-source location (Imig et al., 1990; Middlebrooks and Pettigrew, 1981; Rajan et al., 1990b) and adjacent neurons often show similar spatial sensitivities. Systematic traverses across the cortex, however, fail to reveal any systematic shifts in preferred location (Imig et al., 1990; Middlebrooks and Pettigrew, 1981; Rajan et al., 1990a). Moreover, the receptive fields of many cortical neurons encompass a hemifield or more of auditory space, and broaden further with increases of stimulus intensity (Middlebrooks and Pettigrew, 1981). Therefore, the notion that space is represented topographically in the auditory cortex currently is not supported.

Previous work in our laboratory has used statistical pattern-recognition algorithms to demonstrate that the temporal firing patterns of individual neurons carry significant amounts of information about the locations of sound sources throughout 360° of azimuth (Middlebrooks et al., 1998) and similarly broad ranges of elevation (Xu et al., 1998). The results suggest an alternative to the view that the representation of space in auditory cortex relies upon a topographic organization of sharply-tuned neurons. Specifically, the cortex appears to employ a distributed representation by panoramic neurons, whereby each neuron is involved in representing many different locations and each single location in space is represented by the coordinated responses of many different cortical neurons (Furukawa et al., 2000). Panoramic coding by individual neurons involves stimulus-related changes in both the firing rates and temporal patterns of spikes, with first-spike latencies carrying much of the relevant information (Furukawa and

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Middlebrooks, 2001; Middlebrooks et al., 1998).

Given the behavioral importance of sound localization, it is surprising that no cortical area has yet been distinguished as qualitatively specialized for localization. Certain auditory fields show connections with spatial structures—for example, field AES projects strongly to the superior colliculus (Meredith and Clemo, 1989)—but little physiological specificity for sound-source location. Rauschecker (1998, Rauschecker and Tian, 2000) has proposed a hypothesis, based on analogy with the primate visual system, that the primate auditory system might contain separate cortical processing “streams” specialized for auditory object identification (the “what” stream, involving fields rostral to A1) and spatial processing (the “where” stream, involving caudal fields). This view has received some quantitative support from physiological studies in macaques, demonstrating an increased prevalence of direction-selective neurons in the caudolateral (Tian et al., 2001), and caudomedial (Recanzone et al., 2000) belt areas of auditory cortex. Nevertheless, no evidence has yet been shown for obvious qualitative differences in spatial sensitivity indicative of a higher level of spatial processing in particular cortical fields.

A number of studies in our laboratory have examined the spatial sensitivity of auditory cortical neurons in the cat (Furukawa and Middlebrooks, 2001; Furukawa et al., 2000; Mickey and Middlebrooks, 2001; Middlebrooks et al., 1998; Xu et al., 1998), focusing on non-tonotopic field A2, the anterior ectosylvian sulcus (field AES), and A1. The results of these studies reveal that the general characteristics of spatial sensitivity are quite similar between cortical fields, although minor quantitative differences in tuning and panoramic location coding exist between fields. The lack of fundamental differences between cortical fields suggests that perhaps no area specialized for sound localization exists within the cortex. One possible explanation for this is that the processing of spatial information is essentially complete within the auditory brainstem (Middlebrooks et al., 2002). According to that view, the role of cortex in spatial processing is limited to the distribution of pre-processed information to appropriate perceptual, memory, and motor systems, rather than the actual computation of source location. That model accounts for the current lack of evidence for specialization for spatial processing. Another alternative, however, is that specialized location coding exists in one of several cortical areas in which spatial sensitivity has yet to be examined. The present study considers one of those areas, the posterior auditory field (PAF), located posterior to A1 along the caudal bank of the posterior ectosylvian sulcus of the cat cortex.

PAF is a reasonable candidate for spatial coding, as several aspects of PAF responses suggest increased sensitivity to stimulus features that covary with location. First, neurons in PAF show an increased prevalence of complex frequency-tuning characteristics (Heil and Irvine, 1998; Heil and Imig, 1987). Reciprocal corticocortical projections exist between PAF and ipsilateral fields A1, A2, Anterior Auditory Field (AAF), and Ventral-Posterior Auditory Field (VPAF), along with contralateral fields PAF and VPAF (Rouiller et al., 1991). PAF additionally projects to limbic structures including the cingulate and parahippocampal cortices and the claustrum (Rouiller et al., 1990).

Anatomically, PAF receives its principal thalamocortical projections from the ventral division of the medial geniculate body (MGB); it also receives input from the MGB’s medial and dorsal divisions (Huang and Winer, 2000; Morel and Imig, 1987). Reciprocal corticocortical projections exist between PAF and ipsilateral fields A1, A2, Anterior Auditory Field (AAF), and Ventral-Posterior Auditory Field (VPAF), along with contralateral fields PAF and VPAF (Rouiller et al., 1991). PAF additionally projects to limbic structures including the cingulate and parahippocampal cortices and the claustrum (Rouiller et al., 1990).

In the present study, we recorded responses of PAF and A1 units to sounds whose locations differed in both azimuth and elevation. We focussed on estimating the spatial sensitivity of neurons in the two cortical fields based on measures of response rate and latency. In addition, we analyzed the stimulus-related information conveyed by temporal firing patterns using a pattern-recognition algorithm based on artificial neural networks. The results of the study complement previous studies of azimuth and elevation tuning in areas A2 and AES (Furukawa and Middlebrooks, 2001; Furukawa et al., 2000; Middlebrooks et al., 1998; Xu et al., 1998), and reveal somewhat increased spatial sensitivity in PAF relative to other studied cortical fields, along with the appearance of an enhanced latency code for sound-source location.
MATERIALS AND METHODS

Animal preparation

Six purpose-bred male (4) and female (2) cats, weighing between 3.4 and 7.0 kg were used in this study. The two female cats were previously trained to detect acoustic stimuli in a behavioral study. Male cats participated only in this terminal experiment. All procedures complied with guidelines of the University of Michigan Committee on Use and Care of Animals, and were essentially identical to those described previously (Middlebrooks et al., 1998). Briefly, surgical anesthesia was induced and maintained with isoflurane (2-3%) in nitrous oxide (2 l/min) and oxygen (1 l/min). Following surgery, cats were transferred to intravenous α-chloralose (1.5 mg/ml) in Ringer’s solution for unit recording. Dosage was approximately 3 mg/kg/h, and adjusted to maintain an areflexic state. Atropine sulfate (0.1–0.2 ml, intramuscular) was administered at regular intervals throughout the experiment to suppress mucosal secretions. Following partial removal of the scalp and right temporalis muscle, a craniotomy of 1-cm diameter exposed the right middle ectosylvian gyrus and posterior ectosylvian sulcus (PES). The animal was positioned in the center of the sound chamber, with its head held by a bar attached to the skull fixture and its body suspended in a fabric sling. Thin wire supports maintained symmetric pinna placement throughout the experiment. A warm-water heating pad maintained body temperature at 37°C Celsius. An esophageal stethoscope fitted with a thermometer was used to continuously monitor the cat’s temperature, heart rate, and respiration. Experiments lasted from 2 to 4 days, after which the cats were euthanized. The right cortical hemisphere was then removed and immersed in buffered formalin for later visual confirmation of the region of cortex recorded.

Experimental apparatus and stimulus generation

The experimental apparatus and procedures for stimulus generation were essentially identical to those detailed previously (Middlebrooks et al., 1998; Xu et al., 1998). Recordings were made in a 2.6 x 2.6 x 2.5 m sound-attenuating chamber, the surfaces of which were lined with sound-absorbing foam to suppress reflections. Sounds were presented one at a time from calibrated loudspeakers located 1.2 m from the cat’s head. A circular hoop held 18 loudspeakers in the horizontal plane that spanned 360° of azimuth in 20° increments. The speaker location directly in front of the cat was labeled 0°, with positive azimuths to the cat’s right side (ipsilateral to the recording site) and negative azimuths to the left. A second circular hoop in the vertical median plane held 14 loudspeakers that spanned 260° of elevation in increments of 20°, from 60° below the frontal horizon (−60°) up and over the head to 20° below the rear horizon (+200°). Experiments were controlled by a personal computer, and acoustic stimuli were synthesized digitally using equipment from Tucker-Davis Technologies (TDT; Gainesville, FL). All stimuli were generated with 16-bit precision at a 100-kHz sampling rate. A computer-controlled multiplexer permitted any one loudspeaker to be activated at a time. Stimuli were either 80-ms Gaussian noise bursts with abrupt onsets and offsets or 80-ms pure tones with 5-ms raised-cosine onset/offset ramps.

Data acquisition and spike sorting

Extracellular unit activity was recorded using multichannel silicon-substrate microprobes. These probes, provided by the University of Michigan Center for Neural Communication Technology (Anderson et al., 1989), permit simultaneous recording from up to 16 cortical sites, and are fabricated in several formats. The data presented here were obtained using single-shanked probes with linear arrays of either 8 recording sites spaced every 200µm or 16 sites spaced every 100 or 150µm. Impedances were between 1 and 4 MΩ on 16-channel probes (site area: 177µm²) and 340–360 kΩ on 8-channel probes (site area: 1250µm²). In 8 instances, two such probes were placed simultaneously in different cortical areas, one in PAF and one in A1, and we recorded from 8 sites on each probe. Otherwise, a single probe was used and we recorded from all 8 or 16 sites at a time. Activity at each site was amplified and digitized with a TDT model RA16 biological amplifier/multichannel DSP system. Signals were sampled at 25 kHz, bandpass filtered (0.2–4 kHz), resampled at 12.5 kHz, and stored on a computer disk for offline analysis. Spikes were monitored online using custom software in order to estimate the thresholds and frequency tuning of units prior to data collection.

Off-line spike sorting was performed using custom software based on principal component analysis. This approach, based on that used by Furukawa et al. (2000), involved three steps. First, the signals—interpolated and resampled at 50 kHz—were “denoised” using a multi-channel array-processing technique (Bierer and Anderson, 1999). This procedure eliminates signal components that are correlated across multiple recording channels. Because many noise sources (e.g., instrumentation noise and background neural activity) are highly correlated across the recording array, denoising acts to improve the effective signal-to-noise ratio of the recorded waveforms. Second, candidate spikes were identified as waveform peaks that exceeded a criterion level proportional to the background RMS level recorded prior to stimulation. Third, candidate spike waveforms were sorted using principal component (PC) analysis with eight PCs. Clusters in PC-space were defined either by hand or by statistical cluster analysis to identify neural units. In ~2% of cases, two discrete clusters corresponding to reliably discriminable single units were identified from a single recording channel. Otherwise, at most one cluster was defined per channel. Poststimulus times of spikes accepted in the clustering procedure were stored with 20 μsec resolution.

In contrast to previous studies (Furukawa and Middlebrooks, 2001), we chose to record from as many sites as possible per penetration rather than to obtain recordings from clearly-isolated single neurons. The spike-sorting procedure described above was used to obtain the best possible isola-
tion of neural signals; however, a reasonable concern with multi-channel recordings is that estimates of neural responses may be distorted by the presence of signals from multiple neurons on individual recording channels. Using relatively strict criteria for waveshape separability and inter-spike timing (Furukawa and Middlebrooks, 2001), nineteen of the current recordings (5%) could be identified, with certainty, as isolated single neurons. The remainder were either clusters of 2 or more neighboring neurons having similar spike waveforms, or single units whose spikes varied in shape due to low signal-to-noise ratio. Consistent with our previous experience, we observed no systematic differences between the stimulus-tuning properties of well-isolated neurons and those with more limited isolation. Thus, we do not distinguish between them in this report; the term “unit” is used in reference to both. Correspondingly, the term “single unit” is used to describe one such unit, in contrast to “ensembles” of (2 or more) such units. Please note that this terminology differs from that of some studies using “single-unit” recordings, in which relative accurate isolation of single neurons is assumed. When necessary, we use the term “single neuron” to identify well-isolated single neurons; data for these are presented in several figures to allow comparison to the larger population of recorded units.

Units that responded with less than 1 spike per trial, on average, to their most effective stimulus were rejected from further analysis, as were units whose average response across all stimuli varied by more than a factor of two between the first and second halves of trials in a recording session. This screening procedure was carried out independently for responses to stimuli varying in azimuth and elevation (see Experimental Procedure) and resulted in a total of 117 units recorded from 13 penetrations in A1 and 267 units recorded from 31 penetrations in PAF. Of the 384 total units recorded, 349 were successfully screened for azimuth responses and 324 for elevation responses.

Experimental procedure

Recordings in this study focussed on cortical areas PAF and A1, which were identified initially by the cortical sulcal pattern and secondarily by their responsiveness to pure-tone stimulation, tonotopic organization, and response latencies. Penetrations in area PAF proceeded in the dorso-ventral or latero-medial direction along the caudal bank of the PES. In no case did we distinguish a clear reversal of tonotopy proceeding ventrally through PAF, indicative of entry into field VPAF (Reale and Imig, 1980). However, the most ventral penetrations tended to reveal broad regions of tuning to higher frequencies (> 12 kHz); suggesting that some penetrations may have been located in the transition region between ventral PAF and dorsal VPAF. Penetrations in A1 passed obliquely into the middle ectosylvian gyrus, generally proceeding in a rostro-caudal direction. Search stimuli, consisting of broadband noise bursts and 0.5–30 kHz pure tones, were presented from loudspeakers located at 0° or -40° (contralateral) azimuth or +80° elevation (10° from overhead). The penetration depth was adjusted to maximize the number of active recording sites, with typically 10–14 sites showing unit responses.

Study of the units in each penetration began by estimating their thresholds to noise bursts, tested in 5-dB increments of SPL. The stimuli were presented from a location at which units responded reliably, most often from loudspeakers at azimuths of 0° or -40° in the horizontal plane, or in the mid-saggittal plane at +80° elevation. Typically, unit thresholds varied by less than 10 dB across sites on a single penetration, and the modal threshold was adopted as the representative threshold for the penetration. Responses to pure-tone stimuli were tested using tone frequencies varying in 1/3- or 1/6-octave steps from 1 to 30 kHz; tone levels varied in 10 dB steps, typically from 0 to 50 dB SPL. Pure tones were always presented from 80° elevation; this overhead location was chosen because the spectrum of the cat’s DTF tended to be flatter there, minimizing the effects of filtering by the pinna on the units’ responses (Xu and Middlebrooks, 2000). Next, we measured the units’ spatial sensitivities using 80-ms noise bursts 20, 30, and 40 dB above threshold, presented from 18 locations in the horizontal plane (=180° to +160° in +20° steps) and 14 locations in the mid-saggittal plane (= 60° to +200°). Stimuli were presented in pseudorandom order, such that each combination of SPL and location was presented once before all combinations were repeated in a different random order; 40 repetitions were completed for each penetration. In some cases, locations in azimuth and elevation were tested in separate blocks; in others, all 30 locations were intermixed (0° and +180° appear in both sets). Neural activity was recorded from 20–50 ms before to 80–200 ms after the stimulus onset. Measurement of spatial sensitivity was often followed by presentations of additional stimuli related to other research questions, so that study at each penetration lasted from 2 to 10 hours. Experiments yielded data from 5–13 (median=9) penetrations.

Data analysis

Spatial sensitivity assessed by analysis of spike count and response latency.

Following spike-sorting, spike times were stored as latencies relative to the onset of sound at the loudspeaker. Arrival of sound at the cat’s head followed a delay of approximately 3.5 ms due to acoustical travel time. Spatial sensitivity was assessed by analyzing spike rates, response latencies, and the amount of stimulus-related information conveyed by spike patterns. We defined \( c_t \) and \( l_t \) as the spike count (number of spikes recorded) and the latency of the first spike, respectively, for a single trial \( t \). For each unit, we calculated the stimulus-specific spike count \( C \) as the arithmetic mean of \( c_t \) for trials matching a given combination of location and level. Similarly, the stimulus-specific response latency \( L \) was defined as the geometric mean of first spike latencies \( l_t \) for each stimulus. Trials which failed to elicit at least one spike were omitted from the calculation of \( L \) (following Furukawa and Middlebrooks, 2001). Examples of \( C \) and \( L \) calculated for single units...
are shown in Figure 3.

To facilitate the calculation of spatial tuning statistics, C was normalized to give the proportion of maximum response across location, ranging from 0 (at locations yielding no spikes) to 1 (at the location yielding the maximum number of spikes). Similarly, normalizing L by the minimum latency across location \( L_{\text{min}} \) and inverting the result gives us

\[
\Gamma = \frac{1}{L/L_{\text{min}}}.
\]

\( \Gamma \) ranges from a minimum of 0 (when \( L \approx \infty \)) to a maximum of 1 (when \( L = L_{\text{min}} \)). Values of \( C \) or \( \Gamma \) near 1 indicate effective stimulus locations, whereas values near 0 indicate stimuli that were ineffective at driving the unit. The common form of \( C \) and \( \Gamma \) facilitates the computation of several statistics of spatial sensitivity based on either the spike counts or response latencies of each unit. Spike-count modulation depth, tuning width, and spatial centroid (all defined below) were computed from \( C \) as in previous studies (Furukawa et al., 2000; Middlebrooks et al., 1998; Xu et al., 1998). Analogous measures based on latency were defined in a similar manner using \( \Gamma \) in place of \( C \). Note that the normalization of \( C \) and \( \Gamma \) across location was performed separately for azimuth and elevation at each stimulus level, as was the computation of spatial statistics described below. Where appropriate, subscripts are used to indicate the type of statistic (e.g., \( \Delta L_{\text{az}} \) or \( \Delta L_{\text{el}} \)).

**Depth of response modulation by location (\( \Delta \)).** The depth (or range) of response modulation is the degree to which response latencies or spike counts vary across space. It is designated by \( \Delta C \) for modulation of spike counts and \( \Delta L \) for modulation of response latency, and computed as the range of variation in \( C \) or \( L \):

\[
\Delta C = 1 - C_{\text{min}}
\]

\[
\Delta L = L_{\text{max}} - L_{\text{min}}
\]

where \( C_{\text{min}}, L_{\text{max}}, \) and \( L_{\text{min}} \) refer to the minimum and maximum values of \( C \) and \( L \) across location. Note that \( \Delta L \) is based on \( L \), not \( \Gamma \), and has units of milliseconds. \( \Delta C_{\text{az}} \) and \( \Delta L_{\text{az}} \) designate the modulation depth across azimuth; \( \Delta C_{\text{el}} \) and \( \Delta L_{\text{el}} \) correspond to variation in elevation.

**Spatial tuning width (\( W \)).** Spatial tuning width characterizes the range of locations that were effective in eliciting a strong or rapid response from a given unit. For each unit, values of \( C \) or \( \Gamma \) were linearly interpolated between locations at a resolution of 0.2°. Tuning width \( W_C \) or \( W_\Gamma \) was defined as the range of locations (not necessarily contiguous) associated with values of \( C > 0.5 \) (Middlebrooks et al., 1998) or \( \Gamma > 0.75 \). The stricter criterion was adopted for \( \Gamma \) because \( \Gamma \) tended to be modulated somewhat less than \( C \) overall. \( W \) has units of degrees, and is further identified by subscripts for count or latency and azimuth or elevation: \( W_{C,\text{az}}, W_{L,\text{el}}, \) etc.

**Spatial centroid (\( \xi \)).** Following Middlebrooks et al. (1998), we calculated a spatial centroid for each unit. The centroid is the spatial center of mass of a unit’s “peak response”. As for the calculation of \( W \), locations were interpolated to a resolution of 0.2°. The peak response was then defined as the group of contiguous locations with \( C \) or \( \Gamma \) greater than 0.75 and including the overall maximum \( C \) or \( \Gamma \). A further requirement for the calculation of spatial centroid was that the response fall to below 0.75 at some locations (i.e., some locations were not included in the peak). Units that were modulated by less than this amount were classified as having no centroid (“NC” in Figures 4 and 8). Otherwise, the spatial centroid, designated by \( \xi_C \) or \( \xi_\Gamma \), was computed by generating a set of vectors \( \vec{x} \), whose angles were the (interpolated) stimulus locations \( \theta_i \) included in the peak response and whose lengths were the values of \( C \) or \( \Gamma \) at those locations. The centroid \( \xi \) was defined as the angle of the resultant:

\[
\xi_C = \angle \sum \vec{x}_i, \text{where} \quad \frac{\sum \vec{x}_i}{|\vec{x}_i|} = C(\theta_i)
\]

\[
\xi_\Gamma = \angle \sum \vec{x}_i, \text{where} \quad \frac{\sum \vec{x}_i}{|\vec{x}_i|} = \Gamma(\theta_i)
\]

\( \xi \) is further denoted by subscripts for location type; e.g., \( \xi_{C,\text{az}} \) and \( \xi_{C,\text{el}} \) for azimuth and elevation, respectively.

**Spatial sensitivity assessed by ANN analysis of spike patterns.**

We estimated the spatial sensitivity afforded by temporal patterns of neural response using a pattern-recognition algorithm based on artificial neural networks (ANNs). The approach was similar to that described previously (Furukawa and Middlebrooks, 2001). Briefly, we sorted the neural response patterns into two sets obtained from even- and odd-numbered trials. One set (“training”) was used for setting ANN parameters; the other set (“test”) was used for testing the accuracy of the obtained ANN solution. The separation of training and test sets in this way provided for cross-validation of the pattern-recognition scheme. Times of spikes recorded on each trial were expressed with 100 μs precision. Next, average (“bootstrapped”) response patterns were formed from samples of spike patterns on eight trials, drawn randomly with replacement from the training or test set of 20 responses to each combination of stimulus location and sound level. Twenty such average response patterns were generated from the training set, and another 20 from the test set, for each unique stimulus combination. Average response patterns were then convolved with a Gaussian impulse (σ = 1 ms) and resampled at a resolution of 2 ms to produce spike density functions (SDFs). The bootstrapping and convolution operation served to low-pass filter the spike patterns below 137 Hz and smooth out the otherwise-sparse spike density functions used as input to the ANN. We refer to the resulting SDFs as “single-unit spike patterns;” they represent the smoothed responses
recorded at one recording site. To analyze the combined information in responses from multiple cortical sites, we concatenated single-unit spike patterns for each stimulus to form long vectors, referred to as "ensemble spike patterns."

In addition to these full spike patterns, we also generated test and training sets for two "reduced" input types. For these, the input to the network consisted of a single numerical value corresponding to the spike count or response latency associated with each set of bootstrapped trials. The latency input was computed as the geometric mean, across the 8 bootstrapped patterns, of first-spike latencies. Trials which elicited no spikes were excluded from the computation. Spike count was defined as the (arithmetic) mean number of spikes elicited during the 8 trials chosen for bootstrapping. Both input types were normalized for presentation to the network, such that a value of 1 corresponded to the maximum latency or spike count, across the entire stimulus ensemble, and 0 corresponded to the minimum. Aside from normalization, generation of network inputs and targets was identical to that used by Furukawa and Middlebrooks (2001).

Previous studies employed ANN architectures based on the feedforward multi-layer perceptron for localization (Furukawa et al., 2000; Middlebrooks et al., 1998; Xu et al., 1998) or learning vector quantization (LVQ) for classification (Furukawa and Middlebrooks, 2001). Here, a fixed architecture based on radial basis function (RBF) networks (Ghosh and Nag, 2001; Wasserman, 1993) is used for classification. The approach is similar to the LVQ networks used by Furukawa and Middlebrooks (2001), in that one classifier "unit" is assigned to each stimulus location, and SDFs are classified based on their similarity to these units' input weight vectors. In both cases, weight vectors were initially set to the mean of training-set SDFs at each location. Furukawa and Middlebrooks (2001) used the LVQ algorithm to further optimize the weight vectors, whereas in this case weight vectors remained fixed at the location-conditional mean SDFs. Each classifier unit outputs a scalar value inversely related to the distance (dot product) between its weight vector (the mean SDF for its location) and the input SDF. Normalized, these outputs can be interpreted as posterior probabilities, providing for a number of useful analyses. In the interest of comparing our results directly to earlier studies (Furukawa and Middlebrooks, 2001), however, we converted the output probabilities to a single classification by selecting the unit with maximum activation (i.e., the most likely stimulus location) as the network's response for each input.

Although there are precise correspondences between the classification approach used here and previous efforts, some may object to our use of the term "artificial neural network" in reference to this classifier because it lacks a number of traditional characteristics of ANNs, notably optimization of parameters and non-linear classification rules. Empirical verification of this method showed little benefit of introducing more complex architectures or network training with this dataset, so we adopted the simplest approach in this case; it is by no means guaranteed to generalize to other neural populations or stimulus conditions. While we recognize the limitations of the approach, we refer to the classifier as an ANN in the interest of relating to past and future work employing more general architectures.

Networks were designed and simulated using a customized version of the MATLAB Neural Network Toolbox (The Mathworks Inc, Natick MA). ANN estimates of stimulus locations were expressed as joint stimulus-response probability matrices (confusion matrices), from which we calculated the total stimulus-related (TSR) transmitted information, in bits, to assess the accuracy of ANN performance. The computation of transmitted information was identical to that described previously (Furukawa and Middlebrooks, 2001). Transmitted information (mutual information) reflects the amount of reduction in uncertainty about stimulus location given the set of network responses. One bit of transmitted information implies perfect discrimination of 2 regions of space (e.g., left vs. right) or more continuous discrimination with some error. Perfect identification of 18 locations corresponds to 4.17 bits. For the present study, we calculated the transmitted information from network classifications based on single-unit spike-patterns (TSR$_S$), ensemble spike patterns (TSR$_E$), and reduced spike-patterns consisting of only spike counts (TSR$_C$) or response latencies (TSR$_L$) obtained from single-unit responses.

Interpreting the TSR information rates estimated in this manner requires consideration of two potential sources of bias: overestimation due to finite samples and underestimation due to suboptimal ANN performance. First, because there are a limited number of SDFs to classify, the probabilities expressed in the confusion matrix are not exactly uniform; this results in a positively-biased information estimate. We studied the effects of this bias for each unit by randomly permuting the stimulus labels assigned to each SDF and recomputing TSR. The process excludes systematic stimulus-related sources of information from the TSR calculation. We computed the average of 100 such permutations to estimate bias for each unit, and found the biases to be sufficiently small that any effects on the current analyses were negligible (for azimuth, the mean and maximum bias across units were 0.025 and 0.057 bits, respectively; for elevation, they were 0.017 and 0.042). Second, it is rather unlikely that the ANN classifier used in this study could capitalize on all potential sources of information contained in the SDFs. The network architecture was designed for simplicity and computational efficiency rather than statistical optimality. Different (non-linear) architectures, approaches to parameter optimization, or methods of data representation would likely have resulted in somewhat different levels of classification performance. Hence, information estimates reported here represent lower bounds on the total stimulus-related information contained in the SDFs. Each of these factors (bias and sub-optimality) certainly had an effect on the absolute magnitude of TSR estimates, although neither was sufficient to justify modification of the method. More importantly, our focus in this report is on comparing information rates between cortical fields, rather than accurately estimating them in absolute terms. Because all information estimates are based on the same set of methods—and assuming that neither bias nor ANN performance differ between the neural populations being compared—these effects should have no effect on the interpretation of the current results.
Response patterns of auditory cortical neurons vary with stimulus level, potentially confounding location-related changes in neural responses. Because the ANN analysis depends upon recognition of stimulus-related response patterns, networks trained at one stimulus level can not perform accurately when tested at another level. Networks trained with responses to sounds that vary in level, however, learn to recognize level-invariant features of the response patterns, and thus to appropriately classify responses to sounds varying over a similar range of levels (Middlebrooks et al., 1998). In this case, the ANN’s fixed architecture forces it to recognize such features because each classifier unit is responsible for recognition of all SDFs corresponding to a single location, regardless of level. Without level-invariant features to distinguish stimuli of different locations, the network will fail to correctly classify some of the stimuli. Except where stated otherwise, ANN analyses in the present study were performed using varying levels, 20 to 40 dB above the unit’s threshold in 10-dB steps; estimates of transmitted information reflect the network’s dependence on features invariant over this range of levels.

Tests of statistical hypotheses.

Non-parametric permutation tests were used to compare distributions of various spatial statistics between cortical fields, stimulus levels, etc. Under the null hypothesis that there are no differences between the distributions, labels identifying the category membership (e.g., cortical field) can be reassigned freely without affecting any computable statistic of the distributions. We estimated the sampling distributions of statistics of interest (unless stated otherwise, the difference between two medians) under the null-hypothesis by randomly reassigning category labels in 5000 different permutations. The proportion of these exceeding (or falling below) the value computed with the original labelling gives the probability of type I error (“p-value”). Unless otherwise noted, p-values given in the text refer to this method. We adopted a fixed criterion of p < .05 for statements of statistical significance; in reporting the results of permutation tests, however, p-values refer to the computed proportion of type I errors, rounded to one significant digit. The permutation test has sensitivity limited by the number of permutations employed. Here, the maximum sensitivity is 0.0002 (1 type I error in 5000 permutations); “p < .0002” indicates that the actual value was more extreme than any obtained by random permutation. Other standard statistical tests (e.g., ANCOVA, linear regression) used the MATLAB statistics toolbox (The Mathworks, Inc., Natick MA).

RESULTS

General Observations

Consistent with previous studies, we found a higher proportion of units with nonmonotonic rate-level functions in PAF than A1. To quantify this proportion, we computed the monotonicity ratio (Sutter and Schreiner, 1995) for each unit; this is the ratio of mean spike count for stimuli presented at the highest tested level to the maximum mean spike count across stimulus level. Ratios of 1 are obtained for units with monotonic RLFs, while ratios less than 1 indicated some degree of nonmonotonicity. Based on responses to broadband noise, and adopting a criterion of 0.5, we found only 10/117 (9%) nonmonotonic units in A1, but 86/267 (32%) nonmonotonic units in PAF. This proportion is notably smaller than the 70–90% reported in previous studies of PAF using pure-tone stimulation and pentobarbital or ketamine anesthesia. (Heil and Irvine, 1998; Kitzes and Hollrigel, 1996; Phillips and Orman, 1984; Phillips et al., 1995).

PAF units generally responded better to tones than to noise stimuli, although PAF units in this study did not entirely fail to respond to noise as did those reported by Phillips et al. (1995). Some A1 units showed a similar preference for tones, but overall, A1 units were relatively more responsive to noise. To quantify this difference, we calculated the “Noise/Tone Ratio” for each unit. Briefly, we recorded unit responses to 80-ms noise bursts presented from a frontal or overhead location and varying over a range of at least 60 dB in level. Similarly, we recorded responses to 80-ms pure tones (rise/fall times were 5 ms) varying over 5 octaves in frequency (roughly 1–30 kHz) and 60 dB in level. The noise/tone ratio was defined as the ratio of mean spike counts elicited by the most effective noise and pure-tone stimuli. Distributions of this ratio for the two fields are shown in Fig. 1a. Distributions were computed by kernel density estimation (KDE), the result of convolving the data with a rectangular window of width 0.2; plotted values are equivalent to those of a histogram with continuously-varying bin centers. Overall, PAF units tended to favor optimal tones over optimal noises, whereas many A1 units responded to noise at least as well as they did to tones. However, the majority of PAF units responded to broadband noise with no less than half the spike count elicited by the best tonal stimulus. Figure 1b plots the monotonicity ratio against the noise/tone ratio for each unit, demonstrating a weak positive correlation between the two measures. Nonmonotonic units (falling below the dashed line) showed weaker responses to noise, whereas monotonic units (above the line) showed a broader range of noise response. That there are fewer nonmonotonic units in A1 (gray diamonds) than PAF (open circles) at least partially explains the difference in noise/tone ratio between the cortical fields. Each of these findings, that PAF shows a higher proportion of nonmonotonic units than A1 and that nonmonotonic units in PAF do not respond as well to broadband as narrowband stimuli, are consistent with the results of previous studies (Heil and Irvine, 1998; Phillips et al., 1995).

Also in agreement with previous studies (Heil and Irvine, 1998; Kitzes and Hollrigel, 1996; Loftus and Sutter, 2001; Phillips and Orman, 1984; Phillips et al., 1995), we found PAF units to respond with longer latency than A1 units. Distributions of overall response latency—median L across azimuth—are shown in Fig. 2 for PAF (solid lines), A1 (shaded region), and an additional population of 40 units previously recorded in A2 (unpublished observations from Furukawa and Middlebrooks, 2001) (dashed lines). Median overall latency in PAF
Stimulus-dependent variations in spike count

Azimuth and elevation centroids based on spike counts ($\xi_{ac}$ and $\xi_{el}$, see Methods) were calculated for each unit at 20 and 40 dB above threshold. Distributions of centroids in each cortical field appear in Figure 4. A proportion of units were so broadly tuned that no centroid could be determined; these are indicated as “NC” in the figure. In all cases, A1 showed higher proportions of such units than did PAF. The majority of units in both cortical fields had azimuth centroids located in the contralateral hemisphere (negative azimuths). Elevation centroids were distributed more uniformly, except at low sound levels, at which $\sim 40\%$ of A1 units had centroids between $+10^\circ$ and $+50^\circ$. We computed the $\chi^2$ “goodness of fit” statistic, using 20-degree bins, as an index of each distribution’s non-uniformity; we used standard permutation tests (see methods) to compare these. At 20 dB above threshold, the distribution of azimuth centroids was significantly more uniform in PAF than A1 ($p < .0006$, goodness of fit: $\chi^2_{(17df)} = 99.90$ in PAF, 135.29 in A1). Distributions of elevation centroids at the same level were not significantly different in uniformity ($p > .05$), although PAF and A1 distributions were significantly different from one another ($\chi^2$ contingency test: $\chi^2_{(3df)} = 37.95$, $p < .05$). Distributions of $\xi_{C}$ centroids in PAF were similar at 20 and 40 dB above threshold, whereas in A1 the distributions flattened somewhat at the higher level. As a result, centroid distributions 40 dB above threshold did not differ significantly between the two cortical fields ($\chi^2$ contingency test, azimuth: $\chi^2_{(17df)} = 25.25$, $p > .05$, elevation: $\chi^2_{(3df)} = 14.57$, $p > .05$).

Distributions of $\Delta C$, the depth of spike-count modulation across space, are plotted in Figure 5a. In both cortical fields, spike-rate modulation was shallower for stimuli presented at 40 dB than 20 dB above threshold ($p < .0002$), indicating a reduction of spatial selectivity at high stimulus levels. At the higher level, however, distributions of modulation...
FIG. 3 Examples of neural responses in PAF (left) and A1 (right) to stimuli varying in azimuth (a–f) and elevation (g–j). Each group of 3 panels summarizes the response of a single neural unit or multi-unit cluster in the following manner: Rasters of spikes recorded on trials at each stimulus location (over all stimulus levels) are plotted in the leftmost panel. Center and rightmost panels plot the stimulus-specific spike counts $C$ and response latencies $L$, respectively, computed for combinations of location and level. Solid, dashed, and dotted lines present data at levels 20, 30, and 40 dB above unit thresholds, respectively. Compared to PAF units, A1 units exhibit very little variation in response latency with location; the example units represent the 4th through 86th percentiles of latency variation in A1.
Stimulus-dependent variations in response latency

In addition to spike counts, we assessed spatial sensitivity by relating changes in response latency to stimulus locations. By analogy to the modulation of spike count (ΔC, Figure 5a), we characterized units by their range of variation in response latency across stimulus locations (ΔL). As shown in Fig. 2, overall response latencies were significantly (~20 ms) longer in PAF than in A1 or A2. Of particular relevance to this study, however, these increased latencies do not result simply from delayed overall response times in PAF, but are accompanied by increased stimulus-related variation. Figure 6 plots distributions of ΔL for PAF, A1, and A2 at 20 and 40 dB above threshold. In all cases, PAF units showed significantly more spatial variation of response latency than A1 (p < .0002 at all tested levels). Incidentally, identical results were obtained when the range of latency variation was expressed as a ratio relative to each unit’s median latency (p < .0002 at all tested levels). The majority of A1 and A2 units had response latencies that varied less than 10 ms across azimuth (median ΔLaz in A1: 3.5 ms, A2: 8.6 ms), while a majority of PAF units had latencies varying by twice these amounts (median ΔLaz = 19.3...
ms). Increasing stimulus level had the effect of reducing ΔL in both areas (p < .0002), but even at 40 dB above threshold, a significant number of PAF units demonstrated ranges of 10 ms or more.

Two issues arise with respect to stimulus-related variation in response latency. The first is whether response latencies vary independently of spike counts. To assess this, we computed, for each unit, the correlation between L and C across location. Negative values indicate that locations eliciting many spikes also elicit short-latency responses; this is generally to be expected, as high spike count and short response latency are both general indicators of effective stimulation. Distributions across azimuth of the correlation of L with C, 20 dB above threshold, are plotted for PAF and A1 in Figure 7a. Nearly all A1 units had strong negative correlations, most with values falling between −0.5 and −1. Most PAF units also showed negative correlations, but the distribution of values in PAF was centered nearer to zero (p < .002); a fair number of PAF units even showed positive correlations, indicating high spike-counts at locations eliciting long-latency responses. Consistent with this result, the two fields differed in the proportion of units whose azimuth centroids fell in opposite hemifields when computed by count or latency (Fig. 7b). Together, these findings suggest that in PAF—and not in A1—there exist a number of units (e.g., Figure 3c) whose response latencies and spike counts vary independently, and may therefore constitute independent spatial codes.

A second issue is whether latencies vary specifically with stimulus features related to space, or rather simply as a result of differences in the effective levels of stimuli located near or away from the pinna’s acoustic axis. Many auditory units show response latencies that vary systematically with stimulus level, and effective levels are subject to the directional acoustics of the cat’s pinnae (Middlebrooks and Pettigrew, 1981). We can predict the effects on response latency of these changes in level by computing the effective level for each stimulus position. To do so, we positioned an insert microphone in the ear canal contralateral to the recording site in one cat, and measured directional impulse responses for each loudspeaker position. From these, we calculated the RMS gain for each stimulus location (corrected for the loudspeaker response). We thus calculated the effective levels of individual stimuli presented to each cat, for stimuli varying in azimuth and elevation as well as a separate set of stimuli varying in level but not location. Generally, the latter were presented from either straight ahead (0°) or nearly overhead (+80° elevation). By interpolating the latencies of level-varying stimuli according to the effective levels of location-varying stimuli, we calculated ˆL, response latency predicted by effective level, for each stimulus. The corresponding range of latency variation ΔL was computed similarly to ΔL (Eq. 2). Overall, effective level predicted a smaller range of latency variation than was observed (permutation test on ΔL vs. $\Delta L$, p < .0002 for azimuth, elevation at all tested levels). This was true for both PAF and A1, and suggests that variation of response latency across space is not predicted by purely monaural effects.

Figure 8 plots the distributions of spatial centroids ($\xi_L$) based on latency. Across both azimuth and elevation, at both tested levels, the proportion of units for which no centroid could be calculated (NC or “untuned” units) was higher in A1 than in PAF, consistent with the reduced latency variation in A1 (Figure 6). Among tuned units, however, the distributions of centroids did not differ between the two fields ($\chi^2$ contingency test, azimuth: $\chi^2_{[17\text{df}]} = 18.4$ @ 20 dB, 11.8 @ 40 dB, $p > .05$; elevation: $\chi^2_{[13\text{df}]} = 18.0$ @ 20 dB, 10.5 @ 40 dB, $p > .05$). These generally followed the basic acoustics of the pinna, with shortest latencies observed in response to contralateral azimuths and around +60° in elevation. Distributions broadened somewhat and included more NC units 40 dB above threshold, consistent with the level effects apparent in figures 3 and 6. Additionally, distributions of azimuth centroids at the higher level appear shifted toward the midline, away from eccentric contralateral locations.

Distributions of tuning width $W_L$ are plotted in Figure 9. Tuning widths were narrower in PAF than A1 (p < .0002 at all
We used a pattern-recognition algorithm based on artificial neural networks (ANNs) to analyze the mutual or transmitted information between stimulus locations and network estimates of location based on neural spike patterns. The total stimulus-related ($TSR$) information provides an indication of the accuracy with which spike patterns serve to encode stimulus locations. The TSR information estimate served to quantify the units’ overall spatial sensitivity. Distributions of the single-unit transmitted information ($TSR_S$) provided by each unit’s full spike pattern are plotted in Figure 10. The results for networks trained on stimuli varying in azimuth compare favorably to those obtained in area A2 by Furukawa and Middlebrooks (2001), with the majority of units in PAF and A1 providing 0.5–1 bits of information. PAF units provided significantly more information than did A1 units (median $TSR_S$ in PAF: 0.70 bits, A1: 0.58 bits, $p < .0002$). A similar result was obtained when networks were trained to classify stimuli varying in elevation (median $TSR_S$ in PAF: 0.47 bits, A1: 0.41 bits, $p < .0002$).

When networks were trained using reduced inputs conveying only spike counts or response latencies, information rates decreased (Fig. 11). The spike counts of PAF and A1 units transmitted similar amounts of $TSR_C$ information regarding stimulus azimuth ($p > .05$). In contrast, the two fields differed significantly in $TSR_L$, estimated from response latencies (right panels, $p < .001$). The latter finding is consistent with the increased azimuth-dependent variation of response latency observed in PAF (Fig. 2), and strongly suggests that the difference in $TSR_S$ between PAF and A1 units (Fig. 10) is mediated by differences in latency coding.

We used stepwise regression to explore the relationship between information transmitted by full spike patterns ($TSR_S$) and information transmitted by count or latency ($TSR_C$ or $TSR_L$). Stepwise regression provides a method for computing the proportion of variance explained by one set of predictor variables, independent of that already explained by another set. The method involves computing differences in $R^2$ for models using different sets of predictor variables, and is often used to test the significance of each variable’s contribution to model fit. Here, our interest is in partitioning the total $TSR_S$ variance—across units in each area—according to the independent contributions of count and latency. Altogether, count and latency accounted for 84.9% of variance in $TSR_S$ across units in PAF and 80.9% in A1. This amount can be split into three independent classes: variance explained by count alone, variance explained by latency alone, and variance explained by both count and latency. Just over 51% of total variance in $TSR_S$ (in both areas) fell into the last category, suggesting that count and latency encode space more-or-less redundantly for a majority of units. In PAF, 26.7% of total variance was accounted for by latency ($TSR_L$) independent of count, whereas only 6.8% was explained by count ($TSR_C$) independent of latency. We conclude from this that the improved spatial coding in PAF is largely a result of information transmitted by latency. That is, PAF units that show a high degree of spatial sensitivity in PAF is largely a result of information transmitted by latency and count independently explained by latency and count, respectively. The remaining 15.1% (in PAF) or 19.1% (in A1) of total variance was not explained by either reduced measure. This proportion reflects both random and non-random differences between units; among these, most likely, are differences in information carried by other features of full spike-patterns, such as
FIG. 10 Distributions of the total stimulus-related information $TSR_{S}$, in bits, estimated by ANN analysis of full spike patterns recorded from single units. Stimuli varied in azimuth (left panels) or elevation (right panels), as well as in level (see Methods). ANNs classified each spike pattern to one of the possible stimulus locations, and confusion matrices of network classifications were used to compute the transmitted (mutual) information $TSR_{C}$ for each unit. Values on the ordinate are proportions of units falling per 0.1 bits on the abscissa. As in previous figures, results for PAF units are represented by the black line, and results for A1 by the shaded gray region. Median values for units in each cortical field are given by symbols above the distributions. Small circles below the abscissa indicate values for 17 well-isolated single neurons in PAF.

FIG. 11 Efficiency of azimuth coding by spike count and response latency. Distributions of transmitted information regarding stimulus azimuth are plotted as in Figure 10. Here, network inputs were normalized spike counts (left) or response latencies (right), as described in Methods.

the timing and magnitude of late responses (Loftus and Sutter, 2001), the overall dispersion of spike times (Furukawa and Middlebrooks, 2001), and the modulation of finer-scale temporal features of the main response (e.g., temporal banding in Fig. 3, e, f, and l).

Based on the increased diversity of spatial tuning properties observed in PAF (see Figure 4), we hypothesized that ensembles (in particular) of units recorded in that area might show superior coding of stimulus location (Furukawa et al., 2000). To test this, we trained ANNs to classify ensemble spike patterns (see Methods) according to stimulus location. The ensemble size $N$ varied from 2 to 32 units. Figure 12a plots $TSR_{C}$, the information transmitted by spike patterns of ensembles recorded in each cortical field. Data for ensembles composed of $N$ randomly-chosen units from a single field (“Random Ensembles”) are shown as symbols (PAF: circles, A1: diamonds). For each $N$, 10 different random ensembles were selected and analyzed. Bars represent data for ensembles composed of the $N$ units in each field having the highest single-unit $TSR_{S}$ (“Best Ensembles”). Analysis by ANCOVA using $\log_{2}(N)$ as a covariate confirmed that overall transmitted information rates were higher for random PAF ensembles than random A1 ensembles (azimuth: $F(1,96) = 33.23,$ $p < .05$; elevation: $F(1,96) = 18.78,$ $p < .05$). Additionally, information rates for azimuth increased more rapidly with ensemble size in PAF than in A1 ($F(1,96) = 4.24,$ $p < .05$; for elevation, $F(1,96) = 2.11,$ $p > .05$), suggesting less redundancy between units in PAF ensembles than A1 ensembles.

To assess the diversity of coding within ensembles more directly, we calculated—for each random ensemble in figure 12a—the difference between information transmitted by the complete ensemble and that transmitted by only the most informative unit contained therein. This diversity measure, which we designate $d_{TSR}$, reflects the degree to which units in the ensemble provide independent sources of information about location; its units are bits of information. Mean $d_{TSR}$ for random ensembles is plotted against $N$ in Figure 12b. For azimuth, PAF ensembles expressed greater diversity than A1 ensembles. In both cortical fields, $d_{TSR}$ increased with $N$ from 8–32, but was near zero for smaller ensembles. This suggests that the improvement in transmitted information from $N = 2$ to $N = 8$ (evident in Figure 12a) mainly reflects the increased likelihood of including an accurate unit in larger ensembles, rather than the combination of multiple independent sources of information. The increased diversity among larger ensembles is presumably due to the increased likelihood of sampling a second or third accurate unit that encodes spatial features differently than the first. In elevation, $d_{TSR}$ remained near zero for all ensemble sizes, regardless of cortical field. This suggests a dearth of effective coding strategies for elevation, whereby ensemble performance is limited mainly by performance of the best units.

DISCUSSION

Relationship to previous studies of field PAF

PAF units recorded in this study had significantly longer response latencies than did A1 units, and a larger proportion of PAF units exhibited non-monotonic rate-level functions than in A1. These basic response characteristics consistent with previous studies of PAF (Heil and Irvine, 1998; Kitzes and Hollrigel, 1996; Loftus and Sutter, 2001; Phillips and Orman, 1984; Phillips et al., 1995). It is interesting to note that these two aspects of PAF responsiveness appear similar for broadband (current study) and narrowband (previous studies) stimulation, especially considering that some previous reports have indicated a failure of PAF units to respond to stimulation by broadband noise (Phillips and Cynader, 1985; Phillips et al., 1995). The current results demonstrate that PAF units consistently respond to broadband stimuli presented in the freefield, albeit with fewer spikes than to pure-tone stimuli. It is possible that this difference reflects special sensitivity to spectral features imposed by directional filtering of sound by the head and pinna; these features are present for stimuli presented in the freefield (as in the current study), but absent when broad-
FIG. 12 Coding of spatial location by ensembles of units is more effective in PAF than A1. Transmitted stimulus-related information ($TSR_E$) was calculated based on ANN analysis of spike patterns recorded from ensembles of $N = 2–32$ randomly-selected units in each cortical field (“random ensembles”) or from the $N$ units in each field that exhibited the highest transmitted information when analyzed singly (“best ensembles”). 

**a:** $TSR_E$ is plotted as a function of $N$ for stimuli varying in azimuth (left) or elevation (right). Bars represent data for best ensembles and symbols represent data for random ensembles. 

**b:** diversity of spatial information within ensembles. Diversity ($d_{TSR}$, in bits) for a given ensemble is the difference between information transmitted by the complete ensemble and the maximum of information transmitted by the individual units comprising the ensemble. Mean $d_{TSR}$ for random ensembles of each $N$ and cortical field are represented by black-outlined (PAF) and solid gray (A1) bars.

band stimuli are presented using earphones. This notion is further supported by many of the other current results that indicate increased spatial sensitivity in PAF; other procedural differences (e.g., anesthesia), however, may also play a role in this difference.

With respect to binaural sensitivity, Orman and Phillips (1984) report that a majority of PAF neurons respond more strongly to diotic than to monotic stimulation, suggesting a facilitative role of binaural processing. Neurons displaying such “summative” responses also exhibited shorter response latencies for binaural stimuli than for monaural stimuli. Assuming little influence of differences in interaural timing, one might expect such neurons to respond most strongly to freefield stimuli presented on the interaural midline (i.e., 0° and/or 180° azimuth). The current results generally bear this out; although the majority of PAF units preferred contralateral azimuths, distributions of their azimuth centroids were somewhat skewed toward the midline (see Figs. 4 and 8).

**Superior spatial tuning in PAF**

A number of results of the current study reveal quantitatively sharper and more diverse tuning in PAF than A1. With respect to spike counts, the preferred locations ($\xi_C$) of PAF units sampled azimuth and elevation more evenly than did those of A1 units (Figure 4). This suggests that PAF as a whole may be better-suited to represent locations throughout space, rather than just within a single spatial region (e.g., the contralateral hemifield). In both fields, increasing sound level acted to decrease the depth of spike-count modulation by location and to increase the width of spatial tuning (Figure 5). However, this effect was significantly less severe in PAF than in A1. This finding could be related to the spatial tuning properties of units exhibiting nonmonotonic rate-level functions, which appeared in greater proportion in PAF than in A1. In the current study, as in previous studies of A1 (Barone et al., 1996; Imig et al., 1990), nonmonotonic units showed sharper spatial tuning that was more resistant to changes in SPL.

Latencies of PAF units showed substantially greater location sensitivity than did latencies of A1 units (Fig. 2). Previous work in A1 has demonstrated that first-spike latencies can reliably encode stimulus azimuths (Brugge et al., 1996; Eggermont, 1998), and Furukawa and Middlebrooks, 2001 showed that first-spike latency is responsible for the majority of spatial information transmitted by spike patterns of A2 neurons. Jenison (2001) has further proposed that neural ensembles might use relative latencies to encode space in a distributed fashion. The longer and more variable latencies of PAF units identify the area as qualitatively different from A1 in its ability to make use of such coding strategies; in a number of PAF units, response latencies vary independently of spike counts, providing an added coding dimension from which spatial tuning can be assessed. Tuning widths based on response latency $W_L$ were significantly narrower in PAF than A1 (Fig. 9) and, as for $W_C$, less variable with changes in stimulus level.
Significantly more PAF units had sufficient location-related latency variation for the calculation of spatial centroids ($\xi_L$, Fig. 8). Centroids based on latency were more broadly distributed in PAF than A1, although their distributions in both areas were highly concentrated along the acoustic axis of the cat pinna, which has its strongest response around $-60^\circ$ to $-40^\circ$ in azimuth and around $+60^\circ$ in elevation. This might seem to indicate that spatial variation in the latency of PAF responses does not represent acoustic space, but simply follows the effective level of stimulation. That idea is borne out somewhat by the reduction in spatial variation of latency ($\Delta L$, Fig. 2) at higher stimulus levels, but contradicted by the failure of effective level to account for the large $\Delta L$ values measured in PAF. Rather, it appears that PAF latencies are specifically modulated by binaural comparisons. Moreover, many PAF units have spike counts and response latencies that are not correlated across space, implying that the two features may constitute independent coding dimensions.

A number of factors probably contribute to spatial variation of PAF response latencies. Almost certainly, one is the modulation of acoustic intensity caused by the passive effects of acoustic transmission; for a number of reasons, however, such effects cannot explain the latency data on their own. The range of latency variation across location is too large to be accounted for by a slowing of response in the ascending pathway due to reduced effectiveness or increased “computation time.” There is wide variation in $\Delta L$ among PAF units responding at a common overall latency. Across identical ranges of location, units in A1 exhibit values of $\Delta L$ up to an order of magnitude smaller than PAF units. Finally, when stimuli were presented from a single loudspeaker, but with levels matching those of spatially-varying stimuli, the range of observed latencies was significantly reduced. It appears that some mechanism acts to impose significant delays—or at least to exaggerate them—upon PAF responses. These delays are sensitive to stimulus parameters including sound level and spatial location. Additionally, first-spike latencies in PAF appear to vary systematically with the frequency of pure-tone stimulation (Loftus and Sutter, 2001), and they could be sensitive to other parameters not yet examined. Regardless of the primacy of spatial variation in affecting response latency, PAF stands as an intriguing example of temporal coding of stimulus properties in the central nervous system. Further investigation into the neural mechanisms shaping response latencies in PAF—and their effects on responses to non-spatial stimuli featured—is indicated.

ANN analyses of neural firing patterns for stimuli varying in space provide another example of increased spatial sensitivity in PAF. As shown in Figure 10, responses of PAF units transmitted significantly more information about sound-source location than did those of A1 units. This advantage primarily reflects the added coding capability afforded by PAF units’ more-variable response latencies. Analysis of coding by spike count (Fig. 11) revealed no difference between the two fields, whereas coding by response latency showed an advantage for PAF units. Moreover, stepwise regression reveals that variation among PAF units in their coding of space by full spike patterns is more closely related to variation in coding by latency than spike count; the best units (in terms of spatial coding by spike patterns) in PAF are the ones whose latencies encode space accurately, and units with accurate rate-coding generally exhibit accurate latency-coding, as well.

**Response diversity and ensemble efficacy**

Analyses of spatial sensitivity of neural ensembles revealed significantly greater improvement with ensemble size in PAF than A1 in addition to greater spatial sensitivity of PAF ensembles overall. This result suggests that neurons in PAF are less redundant with respect to their spatial tuning than A1 neurons, a conclusion that is also supported by the greater diversity of spatial centroids (Fig. 4). Middlebrooks et al. (1998) have suggested a panoramic, distributed code for sound localization in the auditory cortex as an alternative to the currently unsupported concept of topographic spatial representation. One requirement for accurate coding in a distributed scheme is a diversity of spatial tuning. The current results demonstrate that area PAF of cat auditory cortex contains a population of units with such diversity in both azimuth and elevation. PAF units’ preferred locations sample space more uniformly than do those of A1 units, and their firing rates and response latencies are modulated more strongly by source location. In PAF, but not in A1, many units appear to use latency and spike count as independent coding dimensions. Coupled with sharper and more level-invariant spatial selectivity, these characteristics provide not only increased coding accuracy among single PAF units, but also improved cooperative coding by ensembles of neurons.

The differences reported in this paper strongly suggest that the stimulus-coding strategies employed by PAF units are particularly effective for the representation of spatial location. Before identifying neurons in PAF as having a specific role in spatial hearing, however, one must consider that many of these advantages must also apply to the coding of other aspects of sound. It is quite possible that the current results reflect more general properties of stimulus coding in PAF. For example, if variation of spectral features unrelated to sound localization produces modulation of response latency similar to that observed for location, then the population of neurons in PAF should be similarly effective at representing other types of complex sound (e.g., vocalizations). In that case, one could argue for a primary role of PAF in the general representation of complex sound, rather than in localization *per se*. Certain aspects of PAF responses seem particularly well-suited for guiding localization behavior; however, a definitive understanding of the functional significance of PAF will require further study.

**Speculation on the possible role of response latency in spatial integration**

Given the rather lengthy response latencies observed in PAF, one might speculate on the temporal relationship between responses in this field and those recorded in non-auditory and polysensory cortical fields, which respond at
longer latencies than A1—median latencies among macaque V1 neurons, for example, are on the order of 66 ms (Schmolesky et al., 1998). With the large difference between auditory and visual response latencies in mind, one might theorize that long latencies allow PAF to synchronize its input to polysensory cortices with that arriving from other sensory systems, and suggest a role for PAF in multisensory integration, attention, or learning. Further support for this idea comes from PAF’s pattern of projections (not shared by A1 or AAF) to high-order and association areas in the limbic system, including the claustrum, cingulate cortex, and parahippocampal cortex (Rouiller et al., 1990). The increased latency-variation in PAF could reflect the scaling of latencies so that slower cortical mechanisms can make use of temporal codes generated in the auditory system. This possibility fits well with the idea that the role of auditory cortex in sound localization is the distribution of preprocessed spatial information to other sensory, memory, and motor systems (Middlebrooks et al., 2001, 2002). Presumably, such a system would require proper scaling of response latencies to ensure appropriate temporal interactions, and one would expect to find units that respond to auditory stimulation with long—and possibly variable—latencies.

As to the mechanism responsible for generating and shaping delayed responses in PAF, one intriguing possibility is early non-specific inhibition of the excitatory response. In this conception, strong stimuli provide sufficient excitation to overcome inhibition and trigger a response with minimal delay. Weaker stimuli provide less effective or infrequent excitation, which takes longer to reach the inhibited response threshold. Such a mechanism could explain both the long response latencies in PAF and their variation with stimulus effectiveness. This idea is additionally supported by the results of Loftus and Sutter (2001) who showed that inhibition in PAF limited early responses to a narrow spectral region of high sensitivity. Stimuli outside this region produced spikes at longer latencies. A possible source of early inhibition in PAF is suggested by results of Huang and Winer (2000) that demonstrate significant projections from the medial division of the MGB to layer Ia of a number of cortical fields, including PAF. These projections take the form of large-caliber (about twice the diameter of other thalamocortical fibers) axons ascending through the cortex. Based on fiber diameter, Huang and Winer (2000) suggest that input arriving via these projections would generally arrive in layer I prior to excitatory input arriving in layers III–IV from the MGB’s ventral division. The pathway could play any of a large number of functional roles, but one possibility is the initiation of early inhibition of pyramidal cells in layers II, V and VI via inhibitory interneurons. Although medial division fibers project to all cortical fields studied by Huang and Winer (2000), PAF exhibited a higher proportion of such projections within layer I than did A1. In addition, projections from small regions of the medial division to PAF appeared more widespread—involving the majority of the dorso-ventral extent of the posterior ectosylvian sulcus—than projections to A1, which were more restricted (see Huang and Winer, 2000, figures 6, 9F).

Notes

1 As an example, the network classifies inputs based only on their similarity to the mean inputs associated with each location. For this reason, the network cannot recognize disjunctions in the input space (e.g., as would occur if a single stimulus elicited two different types of neural responses each dissimilar to their combined mean). Therefore, this network architecture would not be appropriate for analyzing neural responses that vary in a complex or context-dependent manner. Although visual inspection of measured SDFs revealed no obvious instances where this might be the case in general, outliers in the training data and disjunctions caused by level-variant SDFs probably acted to reduce network performance in this way.
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References


