Pitch perception is adapted to species-specific cochlear filtering

Kerry MM Walker¹*, Ray Gonzalez², Joe Kang¹, Josh H McDermott²,³¶ & Andrew J King¹¶

*Corresponding author (kerry.walker@dpag.ox.ac.uk)

1 Department of Physiology, Anatomy & Genetics, University of Oxford
2 Department of Brain and Cognitive Sciences, Massachusetts Institute of Technology
3 Program in Speech and Hearing Biosciences and Technology, Harvard University

Keywords: pitch, ferret, human, psychophysics, auditory nerve, cochlea, resolved harmonics, hearing
Abstract

Pitch perception is critical for recognizing speech, music and animal vocalizations, but its neurobiological basis remains unsettled, in part because of divergent results from different species. We used a combination of behavioural measurements and cochlear modelling to investigate whether species-specific differences exist in the cues used to perceive pitch and whether these can be accounted for by differences in the auditory periphery. Ferrets performed a pitch discrimination task well whenever temporal envelope cues were robust, but not when resolved harmonics only were available. By contrast, human listeners exhibited the opposite pattern of results on an analogous task, consistent with previous studies. Simulated cochlear responses in the two species suggest that the relative salience of the two types of pitch cues can be attributed to differences in cochlear filter bandwidths. Cross-species variation in pitch perception may therefore reflect the constraints of estimating a sound’s fundamental frequency given species-specific cochlear tuning.
Introduction

Many of the sounds in our environment are periodic, and the rate at which such sounds repeat is known as their fundamental frequency, or F0. We perceive the F0 of a sound as its pitch, and this tonal quality is one of the most important features of our listening experience. The way that F0 changes encodes meaning in speech [1] and musical melody [2–4]. The F0 of a person’s voice provides a cue to their identity [5–7] and helps us attend to them in a noisy environment [8–10].

The vocal calls of non-human animals are also often periodic, and pitch is believed to help them to identify individuals and interpret communication calls [11,12]. Many mammalian species have been shown to discriminate the F0 of periodic sounds in experimental settings [13–17], and these animal models hold promise for understanding the neural mechanisms that underlie pitch perception. However, pitch acuity can differ markedly across species [16,18], raising the possibility that humans and other mammals may use different neural mechanisms to extract pitch.

The auditory cortex plays a key role in pitch processing, but it remains unclear how cortical neurons carry out the necessary computations to extract the F0 of a sound [19]. Neural correlates of F0 cues [20–22] and pitch judgments [23] have been observed across auditory cortical fields in some species, while a specialized pitch centre has been described in marmoset auditory cortex [24]. There is similar a lack of consensus regarding the neural code for pitch in the human brain [25]. A better understanding of the similarities and differences in pitch processing across species is essential for interpreting neurophysiological results in animals and relating them to human pitch perception.
Pitch discrimination in humans is driven by two acoustical cues that result from low-numbered ‘resolved’ harmonics and high-numbered ‘unresolved’ harmonics [26], and the relative importance of these cues offers a means to compare pitch mechanisms across species. In the frequency domain, F0 can be determined from the distribution of harmonics (Fig. 1A, upper panel) [27–29]. In the auditory nerve, the frequency spectrum is represented as a “place code” of activation across the tonotopic map as well as a “time code” of spikes that are phase-locked to the basilar membrane vibrations [30,31]. However, both these representations are limited by the cochlea’s frequency resolution [27]. Because cochlear filter bandwidths increase with frequency, only low-numbered harmonics produce discernible peaks of excitation and phase locked spikes at their centre frequency (Fig. 1A, middle panel). Such harmonics are said to be “resolved”. By contrast, high-numbered harmonics are not individually resolved, and instead produce beating in time at the F0, conveyed by phase-locking to their envelope [32] (Fig. 1A, bottom panel). For convenience and to be consistent with prior literature, we refer to these unresolved pitch cues as “temporal” cues, cognizant that the representation of resolved harmonics may also derive from a temporal neural code.
Figure 1: Simulated cochlear filters and their responses to a 500 Hz harmonic complex tone. A. Illustration of the role of unresolved and resolved harmonics in periodicity encoding. Upper plot: Amplitude spectrum for a tone complex that contains all harmonics of 500 Hz from 0 – 11.5 kHz. This sound will evoke a pitch corresponding to 500 Hz. Middle plot: Cartoon of the cochlear filters centred on every second harmonic of 500 Hz, based on data from Glasberg and Moore [41]. This illustrates that lower harmonics are resolved, while the cochlear filters corresponding to higher order harmonics respond to multiple harmonic components in the tone. Lower plot: The output of each of these cochlear filter is plotted throughout 5 ms of the tone complex. The resolved harmonics phase lock to the frequency of one harmonic, while unresolved harmonics beat at the sum of multiple harmonic components (i.e. 500 Hz), providing an explicit temporal representation of F0. B-E: A computational model of the cochlear filter bank was used to simulate representations of complex sounds in the ferret and human auditory nerve. Data are color-coded for the ferret (black) and human (blue). B. The frequency tuning of 15 example auditory nerve fibres is shown for the simulated human (left) and ferret (right) cochlea. C-E. Analyses of the responses of human and ferret cochlear filter banks to a 500 Hz tone complex. C. The response strengths of each of 500 auditory nerve fibres to a 500-Hz complex tone were averaged across the duration of the sound, and plotted across the full range of centre frequencies. Most harmonics produce clearly resolvable activation peaks across fibres in the human cochlea (upper plot), but only the lower harmonics are resolved in the ferret cochlea (lower plot). D. The temporal profile of the output of one simulated auditory nerve fibre with a centre frequency of 5000 Hz is shown for the human (upper plot) and ferret (lower plot) cochlea. E. The power at 500 Hz in the output of each frequency filter, averaged across the full duration of the tone complex, calculated for the human (blue) and ferret (black) auditory nerve. For each species, these values were normalized by the maximal F0 power across all channels. The plot shows the mean (+ standard error) normalized power at F0 across all auditory nerve fibres.
Although psychophysical experiments have demonstrated that humans can extract F0 using either resolved harmonics or unresolved harmonics alone [33–35], pitch perception is generally dominated by resolved harmonics [34,36]. Marmosets can also use resolved harmonics to detect F0 changes [37], whereas rodents (i.e. gerbils and chinchillas) rely upon temporal periodicity cues [38–40]. Why resolved harmonics are more important in humans is unknown, but this could relate to the availability of pitch cues following cochlear filtering. The growing evidence that cochlear bandwidths are broader in many other species [41–43] raises the possibility that they might process pitch cues in different ways from humans.

The behavioural studies carried out to date are difficult to compare across species. First, pitch in humans is defined as the percept through which sounds are ordered on a scale from low to high [44]. By contrast, animal studies often measure change detection in a go/no-go task, from which it is difficult to determine whether they experience a comparable ordered pitch percept or whether they are responding to a change in the perceived pitch as opposed to some aspect of timbre. A two-alternative forced choice (2AFC) task requiring “low” and “high” judgements analogous to those used in human psychophysical tasks would better enable cross-species comparisons [16], but has yet to be employed to examine the use of resolved and unresolved cues in animals. Second, the spectral range of stimuli was not fully controlled across F0 in previous studies (e.g. [16,37]), making it possible for animals to base their behavioural choices on the lower spectral edge of the sounds, rather than the sound’s overall F0. Finally, most animal studies [17,37,40] have not directly compared performance across human and non-human species on an equivalent task, so differences in task demands might therefore account for any apparent species
differences. For example, the pitch difference thresholds of ferrets can differ by orders of magnitude between a go/no-go and 2AFC task [45].

The present study overcomes these limitations by directly comparing the pitch cues used by humans and ferrets on a common 2AFC pitch classification task. We first use a computational model to simulate the representation of periodic sounds in the inner ear. The simulations generated predictions about the availability of periodicity cues in the auditory nerve of each species. We then tested these predictions by training ferrets and humans to classify the pitch of a harmonic complex tone. We find differences in their dependence on resolved and unresolved harmonics, which can be accounted by differences in cochlear tuning between ferrets and humans.

Results

Simulating the filtering of tones in the ferret and human cochlea

Humans are believed to have narrower cochlear filter bandwidths than ferrets and other non-human animals [17,41–43,46–49], and these physiological constraints may predispose them to rely on different acoustical cues to classify the pitch of complex tones. Specifically, individual auditory nerve fibres are believed to respond to a narrower range of frequency in humans than in ferrets, which should result in more resolvable harmonics across the human tonotopic map. On the other hand, if the bandwidth of an auditory nerve fibre is broader, its firing should phase lock more strongly to the beating that results from adjacent harmonics, potentially providing a stronger explicit representation of the temporal periodicity of F0 in ferrets than in humans.
To investigate this hypothesis, we modified a standard model of the cochlear filter bank [50] to simulate the representation of tones along the human and ferret basilar membrane. The output of each cochlear filter was half-wave rectified, compressed (by raising the rectified output to the 0.7 power), and lowpass filtered at 3kHz to simulate the transformation of basilar membrane motion into spiking output in the auditory nerve. The existing literature guided the design of this model [50–52] and parameters in the model were derived from either human psychophysics [41] or ferret auditory nerve recordings [48].

As shown in Figure 1B, the cochlear filters are wider for the ferret auditory nerve than the human. In Figure 1C-E, we compare the human and ferret simulated responses to a 500-Hz missing F0 tone complex that we used as a training sound in our ferret behavioural experiment (described below).

When the instantaneous power of the cochlear filters is summed across the duration of the sound and plotted as a function of centre frequency, the individual harmonics of the tone are more clearly resolved in the human cochlea than in the ferret (Fig. 1C). This takes the form of deeper troughs in the activation of nerve fibres whose centre frequencies lie between the harmonic components of the sound. To visualize the temporal representation of the same stimulus, we plotted the output of a single nerve fibre (here, a fibre with a centre frequency of 5 kHz) throughout time (Fig. 1D). In this case, the representation of the 500 Hz F0 is clearer in the ferret – the human cochlea produces weaker temporal modulation because fewer harmonics fall within the fibre’s bandwidth.

We also examined whether the temporal representation of F0 was enhanced in the ferret cochlea across the full range of frequency filters. A Fourier transform was performed on the output of each fibre throughout a 200ms steady-state portion of the
sound. The power of the response at F0 was then expressed as a proportion of the overall power for that fibre. The results of this metric averaged across all fibres in the model are shown in Fig. 1E. The average temporal representation of F0 was enhanced in the ferret compared to the human (Wilcoxon rank sum test; \( z = 8.286, p = 1.175 \times 10^{-16} \)). In fact, this F0 representation metric was higher in the ferret than the human cochlear model across every pair of individually simulated auditory nerve fibres.

These simulations suggest that the ferret cochlea provides an enhanced representation of the envelope periodicity of a complex tone, as conveyed by spikes that are phase-locked to the F0 in the auditory nerve. On the other hand, the human auditory nerve provides a better resolved representation of individual harmonics across the tonotopic array. It might thus be expected that these two types of cues would be utilized to different extents by the two species.

![Figure 2: Psychophysical task design. A. Schematic of ferret testing apparatus, viewed from above. B. Schematic of one trial in the 2-alternative forced choice pitch classification task. The target tone could be lower or higher in F0 than the reference tone (R). Dotted lines indicate time durations that are variable, depending on the animal’s behaviour.]

**Behavioural measures of pitch cue use in ferrets**

To test the role of different pitch cues in ferret pitch perception, we trained five animals on a two-alternative forced choice (2AFC) task that requires “low” and “high” pitch judgements analogous to those used in human psychophysical tasks (Fig. 2A,B). On each trial, a harmonic complex tone was presented at one of two possible fundamental frequencies. Ferrets were given water rewards for responding at the right
nose-poke port for a high F0, and at the left port for a low F0. Incorrect responses resulted in a time-out. We began by training four ferrets to classify harmonic complex tones with an F0 of 500 and 1000Hz, with a repeating pure tone presented at 707Hz (the midpoint on a logarithmic scale) for reference before each trial. Two of these animals, along with one naïve ferret, were then trained on the same task using target F0 values of 150 and 450Hz and a 260Hz pure tone reference. In both cases, the harmonics of the low and high stimuli to be discriminated were matched in spectral bandwidth, so that ferrets could not solve the task based on the frequency range of the sound (Fig. 3; left column). Rather, the animals had to discriminate sounds based on some cue to the F0. After completing several pre-training stages to habituate the animals to the apparatus and sound presentation (see Methods), the ferrets learned to perform the pitch classification task within 22 ± 3 (mean ± standard deviation) days of twice daily training.
Once the ferrets learned to perform this simple 2AFC task, we incorporated “probe trials” into the task in order to determine which acoustical cues they were using to categorize the trained target sounds. Probe trials made up 20% of trials in a given session, and were randomly interleaved with the “standard” trials described above. On probe trials, an untrained stimulus was presented, and the ferret received a water reward regardless of its behavioural choice. This task design discouraged ferrets from learning to use a different strategy to classify the probe sounds.

The inner ear is known to produce distortion in response to harmonic tones that can introduce energy at the fundamental frequency to the basilar membrane response, even for missing-fundamental sounds [53]. These distortion products could in principle counter our attempts to match the spectral bandwidths of the sounds,
since they could cause the lowest frequency present in the ear to differ as a function of F0. To determine if the ferrets relied on such cochlear distortion products to classify tones in our task, we added pink noise to the stimulus on 20% of randomly interleaved probe trials at an intensity that is known to be more than sufficient to mask cochlear distortion products in humans [54,55]. Ferrets performed more poorly on probe trials than on standard trials (paired t-test; \( t = 4.346, p = 0.005 \)), as expected for an auditory discrimination task performed in noise. However, they continued to perform the pitch classification at 71.85% ± 9.60% correct (mean ± standard deviation) with the noise masker, which is well above chance (1-sample t-test; \( t = 6.025, p = 0.001 \)). This suggests that ferrets did not rely on cochlear distortion products to solve our task.

We next moved to the main testing stage of our behavioural experiment, which aimed to determine if ferrets use resolved harmonics, temporal envelope periodicity, or both of these cues to identify the F0 of tones. All tone complexes, both the standard and probe stimuli, were superimposed on a pink noise masker. Our auditory nerve model (above) allowed us to estimate which harmonics in the tone complexes would be resolved in the ferret auditory nerve (Fig. 4A) [56]. This analysis suggests that our standard tones contained both resolved and unresolved harmonics for ferret listeners, as intended. We constructed four types of probe stimuli based on our resolvability estimates: (1) “Low Harmonic” tones containing only harmonics that we expected to be resolved; (2) “High Harmonic” tones containing harmonics presumed to be less well resolved; (3) “All Harmonics Random Phase” probes containing the full set of harmonics present in the standard tone, but whose phases were independently randomized in order to flatten the temporal envelope; and (4) “High Harmonics Random Phase” stimuli with the same randomization of harmonic
phases, but containing only presumptively unresolved harmonics. The spectral ranges of these stimuli are given in Figure 4B, and the spectra and audio waveforms (showing the temporal envelope periodicity) of the 500 and 1000 Hz stimuli are illustrated in Figure 3A. Ferrets were again given water rewards irrespective of their behavioural choice on probe trials, in order to avoid reinforcing different pitch classification strategies across probe stimuli.

The performance of ferrets on the standard and probe stimuli is shown in Figure 5A. A repeated-measures 3-way ANOVA indicated that performance varied with stimulus type (i.e., the standard and 4 probe stimuli) \((F = 10.540, p = 0.003)\), but not across subjects \((F = 1.060; p = 0.391)\) or the two reference conditions (i.e., 260 and 707 Hz) \((F = 0.438, p = 0.576)\). Scores did not significantly vary across individual ferrets in either the 260 Hz (2-way ANOVA; \(F = 0.366, p = 0.704\)) or 707

**Figure 4:** Harmonic content of stimuli. A. The number of resolved harmonics was estimated over a range of F0s, for ferret (black line) and human (blue line) cochlea. B. The frequency ranges and numbers of harmonic partials included in each stimulus. In this table, the F0 is deemed to be “0”, and further harmonics are counted from 1 onwards.
Hz condition (2-way ANOVA; \( F = 2.063, p = 0.158 \)), so data collected from the same
animals in these two conditions were treated as independent measurements.

![Figure 5: Pitch classification performance of ferrets and humans. A. Ferrets’ percent
correct scores on the pitch classification task are plotted for the standard tone trials
(left) and each of 4 probe stimuli (right). The results of testing with the 260 Hz
reference (150 and 450 Hz targets; red) and 707 Hz reference (500 and 1000 Hz
targets; black) are plotted separately. B. Humans’ pitch classification performance
is plotted, as in (A). C. Performance for each of 4 probe stimuli is expressed as the
ratio of the percentage correct score and that achieved with the standard training
tone stimulus. Data are shown for ferrets (black) and humans (blue). Values of 0
indicate that subjects performed at chance for the probe stimulus, while 1 indicates
that they classified the F0 of the probe as accurately as the training stimulus. Error
bars show mean ± SEM. Individual data for (A) and (B) are shown in Supp Figure
1.

To assess the acoustical cues used by animals to solve the pitch classification
task, we compared ferrets’ performance on the standard trials with that on each of the
four probe trial types (repeated measures 2-way ANOVA, Tukey’s HSD test). Ferrets
showed impaired performance on probes that contained only low harmonics (\( p =
0.001 \)), but performed as well as on standard trials when only high harmonics were
presented (\( p = 1.000 \)). Their performance was also impaired when we randomized the
phases of the high-harmonics (\( p = 0.002 \)). Phase randomization also impaired
performance when the full set of harmonics (both resolved and unresolved) were
present (\( p = 2.173 \times 10^{-5} \)). This pattern of results suggests that ferrets rely more
strongly on the temporal envelope periodicity (produced by unresolved harmonics)
than on resolved harmonics to classify the pitch of tones, unlike what would be expected for human listeners.

**Comparison of human and ferret pitch classification performance**

Humans were trained on a similar pitch classification task to the one described for ferrets in order to best compare the use of pitch cues between these two species. Participants were presented with harmonic complex tones and classified them as high or low. A training phase was used to teach participants the high and low F0s.

We tested human listeners using the same types of standard and probe stimuli as in the final stage of ferret testing described above. As the pitch discrimination thresholds of human listeners are known to be superior to those of ferrets [16], we adapted the target F0s (180 and 220 Hz) and harmonic cut-offs for human hearing (Fig. 4). The between-species comparison of interest here is therefore not the difference in absolute scores on the task, but the pattern of performance across probe conditions.

Human listeners also showed varied pitch classification performance across the standard and probe stimuli (repeated-measures 2-way ANOVA; \( F = 36.999, p = 1.443 \times 10^{-15} \)). However, a different pattern of performance across stimuli was observed for human subjects (Fig. 5B). Tukey’s HSD tests indicated that human listeners were significantly impaired when resolved harmonics were removed from the sounds, as demonstrated by impairments in the “High Harmonic” probes with (\( p = 9.922 \times 10^{-9} \)) and without (\( p = 1.029 \times 10^{-8} \)) randomized phases. Conversely, no impairment was observed when resolved harmonics were available, regardless of whether the phases of stimuli were randomized (“All Harmonics Random Phase” condition; \( p = 0.959 \)) or not (“Low Harmonics” condition; \( p = 0.101 \)). These results
are all consistent with the wealth of prior work on human pitch perception, but replicate previously reported effects in a task analogous to that used in ferrets.

The performance for each probe type relative to performance on the standard stimuli, is directly compared between the two species in Figure 5C. Here, a score of 1 indicates that the subject performed equally well for the standard tone and the probe condition, while a score of 0 indicates that the probe condition fully impaired their performance (reducing it to chance levels). This comparison illustrates the differences in acoustical cues underlying ferret and human pitch classifications. As our model simulations predicted, we found that while ferrets were impaired only when temporal envelope cues from unresolved harmonics were disrupted, humans continued to classify the target pitch well in the absence of temporal envelope cues, so long as resolved harmonics were present. This was confirmed statistically as a significant interaction between species and probe type on performance (repeated measures 3-way ANOVA; F = 14.802, p = 3.412 x 10^-9). The two species thus appear to predominantly rely on distinct cues to pitch.

Discussion

We used a combination of cochlear modelling and behavioural experiments to examine the use of pitch cues in ferrets and human listeners. Our model simulations illustrated how broader cochlear filter widths in ferrets result in fewer resolved harmonics and a more enhanced representation of temporal envelopes than the human cochlea. Based on this result, we predicted that the pitch judgments of ferrets would rely more strongly on temporal envelope cues than that of human listeners. Our behavioural experiments directly compared the use of pitch cues in the two species and found that this is indeed the case. Our results provide the first unambiguous
dissociation of pitch mechanisms across species, by utilizing the same task across species, and provide an illustration of the potential consequences of species differences in cochlear tuning.

Findings in other species

Human listeners have consistently been found to have better pitch discrimination thresholds when stimuli contain resolved harmonics [34–36,57,58]. Moreover, cortical responses to pitched sound in humans are stronger for resolved than unresolved harmonics, mirroring perceptual sensitivity [59,60]. The results of our human experiments are thus fully consistent with this large body of prior work, while enabling comparison with non-human animals. Because most natural sounds contain both low- and high-numbered harmonics, humans may learn to derive pitch primarily from resolved harmonics even when temporal envelope cues are also available, and are thus less equipped to derive pitch from unresolved harmonics alone. This would explain the drop in performance when resolved harmonic cues were removed on probe trials in our experiment.

Our cochlear simulations suggest that harmonic resolvability is worse for ferrets than human listeners, so they may conversely learn to rely more on temporal pitch cues when estimating pitch from natural sounds, leading to poorer performance for low harmonic tone complexes. Many non-human mammals are believed to have wider cochlear bandwidths than humans [42,43,61,62], and so we might expect temporal cues to dominate their pitch decisions as we have observed in ferrets. The few studies to directly address F0 cue use in pitch judgments by non-human animals have raised the possibility of species differences in pitch perception, but have relied on go/no-go tasks that differ from standard psychophysical tasks used in humans. For
instance, studies in gerbils suggest that they primarily use temporal cues to detect an
inharmonic component in a tone complex [38,39]. Chinchillas were similarly shown
to detect the onset of a periodic sound following a non-periodic sound using temporal,
rather than resolved harmonic, cues [40,63]. While these studies did not explicitly
compare the use of resolved and unresolved pitch cues, they are consistent with our
findings regarding the importance of temporal cues in non-human species.

Marmosets, on the other hand, appear to use the phase of harmonic components to
detect changes in the F0 of a repeating tone complex only when resolved harmonics
are omitted from the stimulus [17,37]. This suggests that temporal cues are only
salient for this species when they occur in unresolved harmonics. Similarly to
humans, marmosets were found to detect smaller changes in F0 when harmonics were
resolved than when only unresolved harmonics were available [37]. Comparable
studies have yet to be carried out in other non-human primates, so it remains unclear
whether primates are special in the animal kingdom in their dependence on resolved
harmonic cues. We note also that the behavioural task used in previous marmoset
experiments [17,37] required animals to detect a change in F0, whereas the task
employed in this study required ferrets to label the direction of F0 changes. Ferrets
show an order of magnitude difference in pitch acuity on these two tasks [45], raising
the possibility that primates might as well.

The use of probe trials without feedback in the present experiment allowed us to
determine which acoustical cues most strongly influenced listeners’ pitch judgements.
The ferrets relied predominantly on temporal cues under these conditions, but our
results do not preclude the possibility that they could also make pitch judgments
based on resolved harmonics if trained to do so. Indeed, although human listeners rely
on resolved harmonics under normal listening conditions, we can also extract pitch
from unresolved harmonics when they are isolated [34,36,57]. Our simulations show that up to 8 harmonics are resolved on the ferret cochlea, depending on the F0 (Fig. 4A). Consequently, if specifically trained to do so, one might expect ferrets to be able to derive F0 from these harmonics using the same template matching mechanism proposed for human listeners [27,29]. It is also important to note that the relationship between harmonic resolvability and auditory nerve tuning is not fully understood, and nonlinearities in response to multiple frequency components could cause resolvability to be worse than that inferred from isolated auditory nerve fibre measurements.

Overall, the available evidence fits with the idea that pitch judgments are adapted to the acoustical cues that are available and robust in a particular species, with differences in cochlear tuning thus producing cross-species diversity in pitch perception. A similar principle may be at work in human hearing, since listeners rely on harmonicity for some pitch tasks and spectral changes in others, potentially because of task-dependent differences in the utility of particular cues [7]. The application of normative models of pitch perception will likely provide further insight into the relative importance of these cues.

**Implications for neurophysiological work**

A better understanding of the similarities and differences in pitch processing across species is essential for relating the results of neurophysiological studies in animals to human pitch perception. The present experiments suggest that ferrets, a common animal model in studies of hearing (e.g. [23,64–67]), can estimate F0 from the temporal envelopes of harmonic complex tones. Our data indicate that ferrets generalize across sounds with different spectral properties (including wideband sounds, sounds in noise, and sounds containing only high harmonics) without relying
on explicit energy at the F0. In this respect, ferrets appear to have a pitch percept, even though the cues underlying it are apparently weighted differently than in human pitch perception.

The existing literature might be taken to suggest that primates are the most appropriate animal models for examining the role of resolved harmonics in human pitch perception, as they appear to be more like humans in their use of this cue [17,37]. On the other hand, our data suggest that ferrets are a powerful animal model for evaluating temporal models of pitch extraction (e.g. [50,68]). Like the ferret, cochlear implant users also have poor spectral resolution at the cochlea, and consequently these devices are severely limited in their ability to represent resolved harmonics. Using species such as the ferret to better understand the neural basis of temporal pitch processing could provide insight into why current implants produce impoverished pitch perception [69], and how they might be improved in the future.

Materials and Methods

EXPERIMENTAL SUBJECTS

Ferrets (Mustela putorius furo)

Five adult female pigmented ferrets (aged 6 – 24 months) were trained in this study. Power calculations estimated that 5 animals was the minimum appropriate sample size for 1-tailed paired comparisons with alpha = 5%, a medium (0.5) effect size, and beta = 20%. Ferrets were housed in groups of 2-3, with free access to food pellets. Training typically occurred in runs of 5 consecutive days, followed by two days rest. Ferrets could drink water freely from bottles in their home boxes on rest days. On training days, drinking water was received as positive reinforcement on the task, and was supplemented as wet food in the evening to ensure that each ferret received at
least 60 ml/kg of water daily. Regular otoscopic and typanometry examinations were carried out to ensure that the animals’ ears were clean and healthy, and veterinary checks upon arrival and yearly thereafter confirmed that animals were healthy. The animal procedures were approved by the University of Oxford Committee on Animal Care and Ethical Review and were carried out under license from the UK Home Office, in accordance with the Animals (Scientific Procedures) Act 1986.

Humans

The pitch classification performance of 16 adult humans (9 male, ages 18-53 years; mean age = 25.3 years) was also examined, which provided a 60% beta in the power calculations described for ferrets. All subjects reported having normal hearing. All experimental procedures on humans were approved by the Committee on the Use of Humans as Experimental Subjects at MIT.

METHOD DETAILS

Cochlear filter simulations

We used a cochlear filter bank previously developed by Patterson et al. [50] and implemented by Slaney [70] to simulate representations of sounds on the basilar membrane. The model simulates the response of the basilar membrane to complex sounds through two processing modules: (a) a set of parallel Gammatone filters, each with a different characteristic frequency and Equivalent Rectangular Bandwidth (ERB), produces a simulation of basilar membrane motion in response to the sound, and (b) a two-dimensional adaptation mechanism as observed in hair cell physiology. In order to compare the representation of harmonic tone complexes in the human and ferret cochlea, we modified this model to use filter constants derived from either psychophysical estimates of human cochlear filters [41], or ferret auditory nerve
recordings [48]. Based on these sources, the equivalent rectangular bandwidth of filter $i$ in the human cochlea was calculated as:

$$\text{ERB}_i = \frac{f_i}{(12.7 \times \frac{f_i}{1000})^{0.3}},$$

where $f_i$ is the centre frequency of the filter in Hz.

For the ferret cochlea, the equivalent rectangular bandwidth of each filter was estimated using the following linear fit to the data in Sumner and Palmer [48]:

$$\text{ERB}_i = \frac{f_i}{8.9047 + 209.6149}.$$

The output of each channel in the above Gammatone filter bank was half-wave rectified and then compressed (to the power of 0.7) to simulate transduction of sound by inner hair cells. Finally, the output was low-pass filtered at 3kHz to reflect the spike rate limit of auditory nerve fibres. This model architecture is similar to that used in previous studies (e.g. [51,52]).

**Training apparatus**

Ferrets were trained to discriminate sounds in custom-built testing chambers, constructed from a wire mesh cage (44 x 56 x 49 cm) with a solid plastic floor, placed inside a sound-insulated box lined with acoustic foam to attenuate echoes. Three plastic nose poke tubes containing an inner water spout were mounted along one wall of the cage: a central “start spout” and two “response spouts” to the left and right (Fig. 2A). Ferrets’ nose pokes were detected by breaking an infrared LED beam across the opening of the tube, and water was delivered from the spouts using solenoids. Sound stimuli, including acoustic feedback signals, were presented via a loudspeaker (FRS 8; Visaton, Crewe, UK) mounted above the central spout, which had a flat response ($\pm 2$ dB) from 0.2 – 20 kHz. The behavioural task, data acquisition, and stimulus generation were all automated using a laptop computer running custom Matlab (The
Mathworks, Natick, MA, USA) code, and a real-time processor (RP2; Tucker-Davis Technologies, Alachua, FL, USA).

Pre-training

Ferrets ran two training sessions daily, and typically completed 94 ± 24 trials per session (mean ± standard deviation). Several pre-training stages were carried out to shape animals’ behaviour for our classification task. In the first session, animals received a water reward whenever they nose poked at any of the spouts. Next, they received water rewards only when they alternated between the central and peripheral spouts. The water reward presented from the peripheral response spouts (0.3 - 0.5 ml per trial) was larger than that presented at the central start spout (0.1 - 0.2 ml per trial). The animal was required to remain in the central nose poke for 300 ms to receive a water reward from that spout.

Once animals performed this task efficiently, sound stimuli were introduced in the next session. At the start of each trial, a repeating pure tone “reference” (200 ms duration, 200 ms inter-tone interval, 60 dB SPL) was presented to indicate that the central spout could be activated. Nose poking at the central spout resulted in the presentation of a repeating complex tone “target” (200 ms duration, 200 ms inter-tone interval, 70 dB SPL) after a 100 ms delay. The animal was again required to remain at the centre for 300 ms, and early releases now resulted in the presentation of an “error” broadband noise burst (200 ms duration, and 60 dB SPL) and a 3 s timeout before a new trial began. The target tone could take one of two possible F0 values, which corresponded to rewards at one of the two peripheral spouts (right rewards for high F0 targets, and left for low F0s). For all training and testing stages, the target tones contained harmonics within the same frequency range, so that animals could not use spectral cut-offs to classify the sounds. The target tone continued to play until the
animal responded at the correct peripheral spout, resulting in a water reward. Once the animals could perform this final pretraining task with >70% accuracy across trials, they advanced to pitch classification testing.

Testing stages and stimuli

The complex tone target was presented only once per trial, and incorrect peripheral spout choices resulted in an error noise and a 10 s timeout (Fig. 2B). After such an error, the following trial was an error correction trial, in which the F0 presented was the same as that of the previous trial. These trials were included to discourage ferrets from always responding at the same peripheral spout. If the ferret failed to respond at either peripheral spout for 14 s after target presentation, the trial was restarted.

The reference pure tone’s frequency was set to halfway between the low and high target F0s on a log scale. We examined ferrets’ pitch classification performance using two pairs of complex tone targets in separate experimental blocks: the first with F0s of 500 and 1000 Hz (707 Hz reference), and the second with 150 and 450 Hz targets (260 Hz reference). Four ferrets were trained on the 707 Hz reference. Two of these animals, plus an additional naive animal, were trained on the 260 Hz reference. In each case, testing took place over 3 stages, in which the ferret’s task remained the same but a unique set of stimulus parameters was changed (Fig. 3 and 4), as outlined below. Ferrets were allocated to the 260 and 707 Hz reference conditions based on their availability at the time of testing.

Stage 1: Target sounds were tone complexes, containing all harmonics within a broad frequency range (specified in Fig. 4B). When an animal performed this task >75% correct on 3 consecutive sessions, (32.8 ± 7.1 sessions from the beginning of training; mean ± standard deviation; n = 4 ferrets), they moved to Stage 2.
Stage 2: On 80% of trials, the same standard target tones from Stage 1 were presented. The other 20% of trials were “probe trials”, in which the ferret was rewarded irrespective of the peripheral spout it chose, without a timeout or error correction trial. Probe trials were randomly interleaved with standard trials. The probe stimuli differed only by the addition of pink noise (0.1-10 kHz) to the target sounds, in order to mask possible cochlear distortion products at F0. The level of the noise masker was set so that the power at the output of a Gammatone filter centred at the F0 (with bandwidth matched to ferret auditory nerve measurements in that range [48]) was 5dB below the level of the pure tone components of the target. This is conservative because distortion products are expected to be at least 15 dB below the level of the stimulus components [54,55]. When an animal performed this task >75% correct on 3 consecutive sessions, they moved to stage 3.

Stage 3: The probe stimulus from Stage 2 served as the “Standard” sound on 80% of trials, and all stimuli (both the standard and probes) included the pink noise masker described above. Twenty percent of trials were probe trials, as in Stage 2, but this stage contained tones manipulated to vary the available pitch cues. We estimated the resolvability of individual harmonics using ERB measurements available in previously published auditory nerve recordings [48]. For a given F0, the number of resolved harmonics was approximated as the ratio of F0 and the bandwidth of auditory nerve fibres with a characteristic frequency at that F0, as described by Moore and Ohgushi [56], and applied by Osmanski et al. [17]. This measure yielded between 1 and 8 resolved harmonics for ferrets, depending on the F0 (Fig. 4A). Four types of probe stimuli were presented: (1) “Low Harmonics”, which contained only harmonics presumed to be resolved; (2) “High Harmonics”, comprised of harmonics presumed to be unresolved; (3) “All Harmonics Random Phase”, which contained the same set of
harmonics as the standard, but whose phases were independently randomized in order to reduce temporal envelope cues for pitch; and (4) “High Harmonics Random Phase”, which contained the harmonics present in “High Harmonics” stimuli, but with randomized phases. The bandpass cutoffs for the probe stimuli were chosen so that the “Low Harmonic”, but not “High Harmonic”, probes contained resolved harmonics for ferret listeners. Each probe stimulus was presented on at least 40 trials for each ferret, while the standard was tested on over 1000 trials per ferret.

**Human psychophysical task**

Human subjects were tested on a pitch classification task that was designed to be as similar as possible to Stage 3 of ferrets’ task (see above). Target F0s of 180 and 220 Hz were tested on 16 subjects.

In the psychophysical task, human listeners were presented with the same classes of stimuli described above for ferrets. The frequency ranges included in the probe stimuli are listed in Fig. 4B. Sounds were presented over headphones (Sennheiser HD280) in a sound attenuated booth (Industrial Acoustics, USA). A repeating reference pure tone (200 ms duration, 200 ms inter-tone interval, 60 dB SPL) was presented at the start of a trial, and the subject initiated the target harmonic tone complex (200 ms duration, 70 dB SPL) presentation with a keypress. Text on a computer monitor then asked the subject whether the sound heard was the low or high pitch, which the subjects answered via another keypress (1 = low, 0 = high). Feedback was given on the monitor after each trial to indicate whether or not the subject had responded correctly. Incorrect responses to the standard stimuli resulted in presentation of a broadband noise burst (200 ms duration, and 60 dB SPL) and a 3 s timeout before the start of the next trial. Error correction trials were not used for human subjects, as they did not have strong response biases. Standard harmonic
complex tones were presented on 80% of trials, and the 4 probes (“Low Harmonics”, “High Harmonics”, “All Harmonics Random Phase”, and “High Harmonics Random Phase”) were presented on 20% of randomly interleaved trials. Feedback for probe trials was always “correct”, irrespective of listeners’ responses. Humans were given 10 practice trials with the standard stimuli before testing, so that they could learn which stimuli were low and high, and how to respond with the keyboard. Each probe stimulus was tested on 40 trials for each subject, while the standard was tested on 680 trials per subject.

QUANTIFICATION AND STATISTICAL ANALYSIS

Psychophysical data analysis

Error correction trials were excluded from all data analysis, as were data from any testing session in which the subject scored less than 60% correct on standard trials. T-tests and ANOVAs with an alpha of 5% were used throughout to assess statistical significance, where the n indicates the number of subjects per group. Error bars in Figures 1 and 5 show mean ± standard errors. Further details of all statistical tests described here are provided as supplementary tables.

Because humans produced higher percent correct scores overall than ferrets on the behavioural task, we normalized probe scores against the standard scores when directly comparing performance between species. The score of each species in each probe condition was represented as:

\[ P_{\text{norm}} = \frac{P_a - 50}{S_a - 50} \]

where \( P_{\text{norm}} \) is the normalized probe score for species \( a \) on probe \( i \), \( P_a \) is the percent correct score for species \( a \) on probe \( i \), and \( S_a \) is the percent correct score of species \( a \) on the standard trials. If the performance of species \( a \) is unimpaired for a given probe
stimulus \( i \) relative to the standard stimulus, then \( P_{norm\,ai} \) will equal 1. If the listeners are completely unable to discriminate the F0 of the probe, then \( P_{norm\,ai} = 0 \).

The data and custom software developed in this manuscript are available on the Dryad archive.
Acknowledgments

This work was supported by a BBSRC New Investigator Award (BB/M010929/1) and a DPAG Early Career Fellowship (University of Oxford) to KMMW, a McDonnell Scholar Award to JHM, and a Wellcome Principal Research Fellowship to AJK (WT076508AIA, WT108369/Z/2015/Z), which included an Enhancement Award for JHM.

Declaration of Interests

The authors and funding bodies have no competing financial interests in the outcomes of this research.
References

1. Ohala, J.J. (1983). Cross-Language Use of Pitch: An Ethological View. Phonetica 40, 1–18.

2. Dowling, W.J., and Fujitani, D.S. (1971). Contour, interval, and pitch recognition in memory for melodies. J Acoust Soc Am 49, 524.

3. Krumhansl, C.L. (1990). Cognitive Foundations of Musical Pitch (New York: Oxford University Press).

4. Cousineau, M., Demany, L., and Pressnitzer, D. (2009). What makes a melody: The perceptual singularity of pitch sequences. J Acoust Soc Am 126, 3179–3187.

5. Gelfer, M.P., and Mikos, V.A. (2005). The relative contributions of speaking fundamental frequency and formant frequencies to gender identification based on isolated vowels. J Voice 19, 544–554.

6. Latinus, M., and Belin, P. (2011). Human voice perception. Curr Biol 21, R143–R145.

7. McPherson, M.J., and McDermott, J.H. (2018). Diversity in pitch perception revealed by task dependence. Nat Hum. Beh 2, 1–15.

8. Darwin, C.J. (2005). Pitch and Auditory Grouping. In Pitch: Neuial Coding and Perception, C. J. Plack, R. R. Fay, A. J. Oxenham, and A. N. Popper, eds. (New York: Springer-Verlag), pp. 278–305.

9. Miller, S.E., Schlauch, R.S., and Watson, P.J. (2010). The effects of fundamental frequency contour manipulations on speech intelligibility in background noise. J Acoust Soc Am 128, 435–443.

10. Popham, S., Boebinge, D., Ellis, D.P.W., Kawahara, H., and McDermott, J.H. (2018). Inharmonic speech reveals the role of harmonicity in the cocktail
11. Nelson, D.A. (1989). Song frequency as a cue for recognition of species and individuals in the field sparrow (Spizella pusilla). J Comp Psychol 103, 171–176.

12. Koda, H., and Masataka, N. (2002). A pattern of common acoustic modification by human mothers to gain attention of a child and by macaques of others in their group. Psychol Rep 91, 421–422.

13. Heffner, H., and Whitfield, I.C. (1976). Perception of the missing fundamental by cats. J Acoust Soc Am 59, 915–919.

14. Tomlinson, R.W., and Schwarz, D.W. (1988). Perception of the missing fundamental in nonhuman primates. J Acoust Soc Am 84, 560–565.

15. Shofner, W.P., Yost, W.A., and Whitmer, W.M. (2007). Pitch perception in chinchillas (Chinchilla laniger): Stimulus generalization using rippled noise. J Comp Psychol 121, 428–439.

16. Walker, K.M.M., Schnupp, J.W.H., Hart-Schnupp, S.M.B., King, A.J., and Bizley, J.K. (2009). Pitch discrimination by ferrets for simple and complex sounds. J Acoust Soc Am 126, 1321–1335.

17. Osmanski, M.S., Song, X., and Wang, X. (2013). The role of harmonic resolvability in pitch perception in a vocal nonhuman primate, the common marmoset (Callithrix jacchus). J Neurosci 33, 9161–9168.

18. Shofner, W.P. (2005). Comparative Aspects of Pitch Perception. In Pitch: Neural Coding and Perception Springer Handbook of Auditory Research,, C. J. Plack, A. J. Oxenham, R. R. Fay, and A. N. Popper, eds. (New York: Springer Science+Business Media, Inc.), pp. 56–98.

19. Wang, X., and Walker, K.M.M. (2012). Neural mechanisms for the party problem. Nat Comm 9, 2122.
abstraction and use of pitch information in auditory cortex. J Neurosci 32, 13339–13342.

20. Bizley, J.K.K., Walker, K.M.M., Silverman, B.W.W., King, A.J.J., and Schnupp, J.W.H. (2009). Interdependent encoding of pitch, timbre, and spatial location in auditory cortex. J Neurosci 29, 2064–2075.

21. Bendor, D., and Wang, X. (2010). Neural coding of periodicity in marmoset auditory cortex. J Neurophysiol 103, 1809–1822.

22. Fishman, Y.I., Micheyl, C., and Steinschneider, M. (2013). Neural representation of harmonic complex tones in primary auditory cortex of the awake monkey. J Neurosci 33, 10312–10323.

23. Bizley, J.K., Walker, K.M.M., Nodal, F.R., King, A.J., and Schnupp, J.W.H. (2013). Auditory cortex represents both pitch judgments and the corresponding acoustic cues. Curr Biol 23, 620–625.

24. Bendor, D., and Wang, X. (2005). The neuronal representation of pitch in primate auditory cortex. Nature 436, 1161–1165.

25. Griffiths, T.D., and Hall, D.A. (2012). Mapping pitch representation in neural ensembles with fMRI. J Neurosci 32, 13343–13347.

26. Moore, B.C.J., and Gockel, H.E. (2011). Resolvability of components in complex tones and implications for theories of pitch perception. Hear Res 276, 88–97.

27. Goldstein, J.L. (1974). An optimum processor theory for the central formation of the pitch of complex tones. J Acoust Soc Am 55, 1061–1069.

28. Terhardt, E. (1974). Pitch, consonance, and harmony. J Acoust Soc Am 55, 1061–1069.

29. Shamma, S., and Klein, D. (2000). The case of the missing pitch templates:
how harmonic templates emerge in the early auditory system. J Acoust Soc Am 107, 2631–2644.

30. Cariani, P.A., and Delgutte, B. (1996). Neural correlates of the pitch of complex tones. I. Pitch and pitch salience. J Neurophysiol 76, 1698–1716.

31. Schnupp, J.W.H., Nelken, I., and King, A.J. (2011). Auditory neuroscience: making sense of sound (MIT Press).

32. Schouten, J.F. (1970). The residue revisited. In Frequency Analysis and Periodicity Detection in Hearing, R. Plomp and G. F. Smoorenburg, eds. (Leiden: Sijthoff), pp. 41–58.

33. Houtsma, A.J.M., and Smurzynski, J. (1990). Pitch identification and discrimination for complex tones with many harmonics. J Acoust Soc Am 87, 304–310.

34. Shackleton, T.M., and Carlyon, R.P. (1994). The role of resolved and unresolved harmonics in pitch perception and frequency modulation discrimination. J Acoust Soc Am 95, 3529–3540.

35. Bernstein, J.G., and Oxenham, A.J. (2003). Pitch discrimination of diotic and dichotic tone complexes: harmonic resolvability or harmonic number? J Acoust Soc Am 113, 3323–33234.

36. Ritsma, R.J. (1967). Frequencies dominant in the perception of the pitch of complex sounds. J Acoust Soc Am 42, 191–198.

37. Song, X., Osmanski, M.S., Guo, Y., and Wang, X. (2016). Complex pitch perception mechanisms are shared by humans and a New World monkey. Proc Natl Acad Sci U S A 113, 781–786.

38. Klinge, A., and Klump, G.M. (2009). Frequency difference limens of pure tones and harmonics within complex stimuli in Mongolian gerbils and
humans. J Acoust Soc Am 125, 304–314.

Klinge, A., and Klump, G.M. (2010). Mistuning detection and onset asynchrony in harmonic complexes in Mongolian gerbils. J Acoust Soc Am 128, 280–290.

Shofner, W.P., and Chaney, M. (2013). Processing pitch in a nonhuman mammal (Chinchilla laniger). J Comp Psychol 127, 142–153.

Glasberg, B.R., and Moore, B.C.J. (1990). Derivation of auditory filter shapes from notched-noise data. Hear Res 47, 103–138.

Shera, C.A., Guinan, J.J., and Oxenham, A.J. (2002). Revised estimates of human cochlear tuning from otoacoustic and behavioral measurements. Proc Natl Acad Sci U S A 99, 3318–3323.

Joris, P.X., Bergevin, C., Kalluri, R., Mc Laughlin, M., Michelet, P., van der Heijden, M., and Shera, C.A. (2011). Frequency selectivity in Old-World monkeys corroborates sharp cochlear tuning in humans. Proc Natl Acad Sci U S A 108, 17516–17520.

Anon (1994). American National Standard Acoustical Terminology (American National Standards Inst.).

Walker, K.M.M., Bizley, J.K., King, A.J., and Schnupp, J.W.H. (2011). Cortical encoding of pitch: Recent results and open questions. Hear Res 271, 74–87.

Pickles, J.O., and Comis, S.D. (1976). Auditory-nerve-fiber bandwidths and critical bandwidths in the cat. J Acoust Soc Am 60, 1151–1156.

Kittel, M., Wagner, E., and Klump, G.M. (2002). An estimate of the auditory-filter bandwidth in the Mongolian gerbil. Hear Res 164, 69–76.

Sumner, C.J., and Palmer, A.R. (2012). Auditory nerve fibre responses in the ferret. Eur J Neurosci 36, 2428–2439.
49. Alves-pinto, A., Sollini, J., Wells, T., Sumnerqjf, C.J., Alves-pinto, A., Sollini, J., Wells, T., and Sumner, C.J. (2016). Behavioural estimates of auditory filter widths in ferrets using notched-noise maskers. J Acoust Soc Am 139, E19.

50. Patterson, R., Robinson, K., Holdworth, J., McKeown, D., Zhang, C., and Allerhand, M. (1992). Complex sounds and auditory images. In Auditory Physiology and Perception, Y. Cazals, K. Horner, and L. Demany, eds. (Elsevier Ltd), pp. 429–446.

51. Karjalainen, M. (1996). A binaural auditory model for sound quality measurements and spatial hearing studies. In IEEE International Conference on Acoustics, Speech, and Signal Processing (Atlanta, GA, USA: IEEE), pp. 985–988.

52. Roman, N., Wang, D., and Brown, G.J. (2003). Speech segregation based on sound localization. J Acoust Soc Am 114, 2236–2252.

53. Robles, L., Ruggero, M.A., and Rich, N.C. (1991). Two-tone distortion in the basilar membrane of the cochlea. Nature 349, 413–414.

54. Pressnitzer, D., and Patterson, R.D. (2001). Distortion products and the perceived pitch of harmonic complex tones. In Physiological and Psychophysical Bases of Auditory Function, D. Breebaart, A. Houtsma, A. Kohlrausch, V. Prijs, and R. Schoonhoven, eds. (Maastrict: Shaker BV), pp. 97–104.

55. Norman-Haignere, S., and McDermott, J.H. (2016). Distortion products in auditory fMRI research: Measurements and solutions. Neuroimage 129, 401–413.

56. Moore, B.C.J., and Ohgushi, K. (1993). Audibility of partials in inharmonic complex tones. J Acoust Soc Am 93, 452–461.
57. Moore, B.C.J., Peters, R.W., and Glasberg, B.R. (1985). Thresholds for the detection of inharmonicity in complex tones. J Acoust Soc Am 77, 1861–1867.

58. Kaernbach, C., and Bering, C. (2001). Exploring the temporal mechanism involved in the pitch of unresolved harmonics. J Acoust Soc Am 110, 1039–1048.

59. Penagos, H., Melcher, J.R., and Oxenham, A.J. (2004). A neural representation of pitch salience in nonprimary human auditory cortex revealed with functional magnetic resonance imaging. J Neurosci 24, 6810–6815.

60. Norman-Haignere, S., Kanwisher, N., and McDermott, J.H. (2013). Cortical pitch regions in humans respond primarily to resolved harmonics and are located in specific tonotopic regions of anterior auditory cortex. J Neurosci 33, 19451–19469.

61. Liberman, M.C. (1978). Auditory-nerve response from cats raised in a low-noise chamber. J Acoust Soc Am 63, 442–455.

62. Temchin, A.N., Rich, N.C., and Ruggero, M.A. (2008). Threshold tuning curves of chinchilla auditory-nerve fibers. I. Dependence on characteristic frequency and relation to the magnitudes of cochlear vibrations. J Neurophysiol 100, 2889–2898.

63. Shofner, W.P. (2002). Perception of the periodicity strength of complex sounds by the chinchilla. Hear Res 173, 69–81.

64. Fritz, J., Shamma, S., Elhilali, M., and Klein, D. (2003). Rapid task-related plasticity of spectrotemporal receptive fields in primary auditory cortex. Nat Neurosci 6, 1216–1223.
65. Walker, K.M.M., Bizley, J.K., King, A.J., and Schnupp, J.W.H. (2011). Multiplexed and robust representations of sound features in auditory cortex. J Neurosci 31, 14565–14576.

66. Atilgan, H., Town, S.M., Wood, K.C., Jones, G.P., Maddox, R.K., Lee, A.K.C., and Bizley, J.K. (2018). Integration of visual information in auditory cortex promotes auditory scene analysis through multisensory binding. Neuron 97, 640–655.

67. Schwartz, Z.P., and David, S. V (2018). Focal suppression of distractor sounds by selective attention in auditory cortex. Cereb Cortex 28, 323–339.

68. Meddis, R., and O’Mard, L (1997). A unitary model of pitch perception. J Acoust Soc Am 102, 1811–1820.

69. Gfeller, K., Turner, C., Oleson, J., Zhang, X., Gantz, B., Froman, R., and Olszewski, C. (2007). Accuracy of cochlear implant recipients on pitch perception, melody recognition, and speech reception in noise. Ear Hear 28, 412–423.

70. Slaney, M. (1993). An efficient implementation of the Patterson-Holdsworth auditory filter bank.
Supplemental Figure 1: Pitch classification performance of ferrets and humans, as shown in Figure 5A and B. Here, data are plotted for individual subjects. **A.** Ferrets’ percent correct scores on the pitch classification task are plotted for the standard tone trials (left) and each of 4 probe stimuli (right). The results of testing with the 260 Hz reference (150 and 450 Hz targets; red) and 707 Hz reference (500 and 1000 Hz targets; black) are plotted separately. Symbol shapes represent individual ferrets. **B.** Humans’ pitch classification performance, as plotted in (A) above. Data are randomly jittered on the x-axis to facilitate visualization of individual points. Each symbol and colour combination indicates an individual subject.
The figure illustrates the effects of varying harmonic content on signal processing, as indicated by Power (dB) across different frequencies. The power spectra show distinct patterns for different frequency ranges and harmonic configurations.

A table is included to summarize the resolved harmonic cues and temporal envelope cues for each condition:

| Condition                        | Resolved harmonic cues | Temporal envelope cues |
|----------------------------------|------------------------|------------------------|
| Training tone                    | Strong                 | Strong                 |
| Standard (all harmonics)         | Strong                 | Weak                   |
| Low harmonics                    | Weak                   | Strong                 |
| High harmonics                   | Weak                   | Weak                   |
| All harmonics, random phase      | Strong                 | Weak                   |
| High harmonics, random phase     | Weak                   | Weak                   |

The figure also includes time-domain representations for different frequencies, showing variations in amplitude over time (ms) for each condition.
### A

![Graph showing F0 vs. Resolved harmonics for Human and Ferret.](image)

- **Human** line
- **Ferret** line

### B

**Ferret**

| (reference = 260 Hz) | All harmonics  | Low harmonics | High harmonics |
|---------------------|----------------|---------------|----------------|
| Spectral range (Hz) | 450 - 6000     | 450 - 1500    | 1500 - 6000    |
| Low F0 harmonics    | 3 - 40         | 3 - 10        | 10 - 40        |
| High F0 harmonics   | 1 - 13         | 1 - 3         | 5 - 13         |

| (reference = 707 Hz) | All harmonics  | Low harmonics | High harmonics |
|---------------------|----------------|---------------|----------------|
| Spectral range (Hz) | 1000 - 10000   | 1000 - 4000   | 4000 - 10000   |
| Low F0 harmonics    | 2 - 20         | 2 - 8         | 8 - 20         |
| High F0 harmonics   | 1 - 10         | 1 - 4         | 4 - 10         |

**Human**

| (reference = 199 Hz) | All harmonics  | Low harmonics | High harmonics |
|---------------------|----------------|---------------|----------------|
| Spectral range (Hz) | 220 - 5940     | 220 - 1980    | 1980 - 5940    |
| Low F0 harmonics    | 2 - 33         | 2 - 11        | 11 - 33        |
| High F0 harmonics   | 1 - 27         | 1 - 9         | 9 - 27         |
