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The Development of a Single Frequency Place in the Mammalian Cochlea: The Cochlear Resonance in the Mustached Bat *Pteronotus parnellii*

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Cochlear microphonic potentials (CMs) were recorded from the sharply tuned, strongly resonant auditory foveae of 1- to 5-week-old mustached bats that were anesthetized with Rompun and Ketavet. The fovea processes Doppler-shifted echo responses of the constant-frequency component of echolocation calls. During development, the frequency and tuning sharpness of the cochlear resonance increases, and CM ringing persists for longer after the tone. CM is relatively insensitive at tone onset and grows linearly with increased stimulus level. During the tone, the CM is more sensitive and grows compressively with increased stimulus level and phase leads onset CM by $90^\circ$ for frequencies below the resonance. CM during the ringing is also sensitive and compressive and phase leads onset CM by $180^\circ$ below the resonance and lags it by $180^\circ$ above the resonance. Throughout postnatal development, CMs measured during the tone and in the ringing increase both in sensitivity and compression. The cochlear resonance appears to be attributable to interaction between two oscillators. The more broadly tuned oscillator dominates the onset response, and the narrowly tuned oscillator dominates the ringing. Early in development, mechanical coupling between the oscillators results in a relatively broadly tuned system with several frequency modes in the CM at tone onset and in the CM ringing. Beating occurs between the resonance and the stimulus response during the tone and between two components of the narrowly tuned oscillator at tone offset. At maturity, the CM has three modes for frequencies within 10 kHz of the resonance at tone onset and a single, sharply tuned mode in the ringing.

**Key words:** mustached bat; cochlea; microphonic potential; frequency tuning; resonance; synchronization; basilar membrane; tectorial membrane; cochlear amplifier; auditory fovea

Introduction

The mustached bat’s auditory fovea is perhaps the most sharply tuned of all biological mechanical structures. This region occupies 40% of the cochlea and is used to detect the Doppler-shifted echo response of the dominant harmonic constant-frequency component (CF2) of the echolocation call at $\sim 61$ kHz. In new world bats, the acquisition of an acoustic fovea is unique to mustached bats, and there is evidence that this event happened within the last 10 million years (Kössl et al., 1999). Mustached bats exploit the auditory fovea to measure target velocity and insect wing beats in the acoustic clutter of dense vegetation (Neuweiler, 1990; Schnitzler and Kalkow, 1998). The massive overexpression of the 61 kHz region provides an opportunity to study the development of frequency tuning and amplification in a single segment of the mammalian cochlea.

The auditory fovea comprises a basal sparsely innervated (SI) region and a more apical densely innervated CF2 region (Fig. 1). The SI region where signals between 61 and 72 kHz are processed is morphologically specialized with a thickened basilar membrane (BM) and club-shaped tectorial membrane (TM) (Henson and Henson, 1991), which interact together to produce a sharply tuned resonance (Suga et al., 1975; Russell et al., 2003). This resonance is detected by hair cells in the CF2 region (Kössl and Vater, 1995).

We describe the developmental changes in the auditory fovea based on measurements of the cochlear microphonic (CM). CM responses are dominated by outer hair cell (OHC) receptor potentials (Dallos et al., 1972; Russell and Sellick, 1983; Patuzzi et al., 1989). OHC cell bodies are active components of the organ of Corti (Brownell et al., 1985; Dallos, 1992), and the OHC hair bundles are inserted into the TM (Kimura, 1966; Engström and Engström, 1978). Thus, OHCs form an active mechanical link between the BM, which supports the organ of Corti, and the TM, against which the OHC bundles react. Measurement of the CM should reflect how OHCs actively respond to acoustic stimulation and contribute to tuning of the mustached bat’s cochlea. Through appropriately timed electromechanical feedback, OHCs are believed to amplify and sharpen frequency tuning in
the mammalian cochlea (for review, see Robles and Ruggero, 2001), and they do this by reacting with the TM (Davis, 1965; Gummer et al., 1996; Legan et al., 2000; Lukashkin and Russell, 2003). We have recorded the CM from the cochlear aqueduct of the mustached bat (Fig. 1), which provides a direct way of recording from the auditory fovea (Henson and Pollak, 1972; Henson et al., 1985). The strong resonance of the mustached bat’s auditory fovea builds slowly and provides an opportunity to compare the magnitude and phase of OHC responses at tone onset, when cochlear amplification is weak, with those during the full manifestation of the resonance, when amplification is strong (Russell et al., 2003). Temporal analysis of the CM also provides an opportunity to examine the mechanical properties of the different cochlear components that contribute to the resonance.

Mustached bats hear in the first postnatal week. Throughout the next 5 weeks of development, the cochlear resonance shifts upward in frequency by 0.45–0.5 octaves in parallel with the CF2 component of the echolocation call. CM becomes more sensitive, and the resonance becomes sharply tuned. Maturation of the cochlea is correlated with developmental changes in the harmonic composition and lengthening and stabilization of the CF2 component of the echolocation call (Kössl et al., 2003; Vater et al., 2003). Here, we describe the development of the underlying mechanisms that give rise to the remarkable frequency tuning of the auditory fovea.

Materials and Methods

Young mustached bats were collected in July and August in hot caves in Cuba during a period of 4–5 weeks of postnatal development. As an estimate of age, we measured forearm length (FAL), which provides a reliable age-related index during the prevalent period of several bat species (Kunz and Anthony, 1982; Rubäsmen, 1987). Newborn mustached bats have an FAL of 20–22 mm, and, at an approximate growth rate of 1 mm/d, the mature forearm length of 51–52 mm is attained after 4–5 weeks (Vater et al., 2003). In the caves, the bats have a very hot and humid environment (close to 38°C and 100% humidity), which in this respect is comparable with the conditions in their mothers’ wombs. The bats are hairless for at least 4 weeks, and they start to fly around in the cave at an age that corresponds to 46–48 mm forearm length. As described by Silva Taboada (1979), immature bats start to hunt close to the cave entrance region, but their insect diet is at this time still complemented by mother’s milk.

Nineteen young bats were brought to a laboratory in Havana and kept 1–14 d before measurement of cochlear potentials: first postnatal week, n = 2; second postnatal week, n = 3; third postnatal week, n = 4; fourth postnatal week, n = 4; and fifth postnatal week, n = 6. During this period of captivity, the bats’ environment was kept at a high temperature of >35°C and at high humidity. To measure CM and the compound action potential of the auditory nerve (CAP), young bats were anesthetized with Rompun (0.01–0.06 ml of 0.2% solution/10 gm of body weight) alone or with a Rompun and Ketavet mixture (ketamine-HCl, 100 mg/ml, 2% Rompun, and water in the proportion 9:1:190; dose, 0.005–0.02 ml/10 gm of body weight). The skull of the bats was fixed by dental acrylic and the resonance becomes sharply tuned. Maturation of the cochlea is correlated with developmental changes in the harmonic composition and lengthening and stabilization of the CF2 component of the echolocation call (Kössl et al., 2003; Vater et al., 2003). Here, we describe the development of the underlying mechanisms that give rise to the remarkable frequency tuning of the auditory fovea.

Results

Sound-evoked electrical potentials from the cochlear duct

The electrical signal recorded from the cochlear duct in Pteronotus parnellii in response to a tone burst consists of the CM, the CAP of the auditory nerve, and a brainstem field potential (FP; Fig. 1B). The symmetry of the CM waveform and, hence, the generation of a DC component (summing potential; Fig 2B,D), is frequency- and level-dependent and depends on the developmental stage of the bat.

The waveform of the CM recorded from adult bats for frequencies at and close to the CF echo response is symmetrical and without a measurable DC component for low- to moderate-level tones (Fig. 1B), although these traces and all others reported here are slightly skewed by the 50 Hz high-pass filter of the recording system. A positive DC component is evident only for tone levels above ~90 dB SPL (data not shown). Typically, the CM contains a strong resonance with a frequency close to the Doppler-shifted CF echo response frequency (Suga et al., 1975). The CM of immature bats also contains a resonant component that outlasts the stimulus tone but usually for only a few milliseconds, depending on the tone frequency and on the developmental stage of the bat. The CM recorded from the cochlea of a 3-week-old bat resonates for longer at the offset of a resonance frequency tone (Fig. 2B,E) than from the cochleas of bats at earlier developmental stages. The waveform of the CM depends on both the frequency and developmental stage of the bat. For immature bats, the resonance frequency represents a transition point for the symmetry of the response. In the example shown in Figure 2A, the waveform is symmetrical at all levels <85 dB SPL in response to frequencies below the resonance. Hence, the DC component is negligible. For tones at the resonance frequency (e.g., 58.5 kHz) (Fig. 2B) that exceed a critical level, which depends on the developmental stage...
of the bat, the CM waveform becomes asymmetrical, and a positive DC component is generated. This is 65 dB SPL for the CM recording shown in Figure 2 B. For frequencies of >1 kHz above the resonance, the CM is asymmetrical at all levels above the detection threshold. Therefore, for frequencies above the resonance frequency, the CM always has a DC component (Fig. 2 D). CM responses recorded from the cochlear duct of cmbat21 to resonance frequency tones. Both the time and magnitude scales have been magnified to reveal that level-dependent differences in the onset of the DC component (smooth trace) are negligible. AC (B, D), AC component of the CM; DC (B, D), DC component of the CM. E, CM recorded from an immature bat (cmbat31; 47 mm) in response to 2 msec, 75 dB SPL tones to show the frequency dependence of the cochlear resonance (res; 57 kHz). Electrical recordings in A–D are high-pass-filtered at 50 Hz; those in E are bandpass-filtered between 20 and 70 kHz.

**Time dependence of CM frequency selectivity and amplitude sensitivity**

Mechanical tuning in the SI region of the cochlea is so sharp (Kössl and Russell, 1995) that the CM takes several milliseconds to build up in response to stimulus tones at and close to the resonance frequency, and then the CM continues to resonate when the tone has stopped (see Figs. 1, 2). This slowing down of the responses of the cochlea provides an opportunity to investigate temporal changes in the sensory processing of acoustic signals within the first millisecond of a tone, during the steady state of the tone, and 0.5 msec after the offset of the tone, when contributions from the stimulus tone have died away, leaving only the ringing. Measurement periods are indicated by the vertical gray bars in Figure 3 A. CM measurements during the first millisecond of the tone are made during the 1 msec rise time of the tone envelope that was used in these experiments to suppress
stimulus onset transients. These dynamic stimulus conditions have been taken into account when calculating the magnitude of the CM during tone onset.

Amplification and compression of resonance frequency CM magnitude level functions increase with time from tone onset and during development

CM magnitude level functions were measured within 1 msec of tone onset (Fig. 3A, squares), 4–9 msec from tone onset (steady state) (Fig. 3A, circles), and 0.5 msec after tone offset, during the resonance (Fig. 3A, triangles) for a bat of ~1 week in age (Fig. 3B) and another ~5 weeks old (Fig. 3C). Level functions recorded from the 1-week-old bat in response to resonance frequency tones (45.5 kHz) (Fig. 3B) are relatively insensitive, appearing above the noise floor at 70 dB SPL, and do not show compression for tone levels of ~95 dB SPL, which was the maximum stimulus level. Measurements made during tone onset were 10–15 dB less sensitive than those measured during the steady state. CMs measured from the older bat (59 kHz) (Fig. 3C) during the ongoing tone and after the offset tones close to the resonance were compressive at high SPLs and nearly 40 dB more sensitive than the CM magnitude level function measured at tone onset. Thus CM measured at tone onset has the characteristics of responses recorded from a passive cochlea without amplification. CM measured during the tone and after tone offset has the characteristics of responses from a cochlea with amplification. In more mature bats, the CM level functions display the compression that has been observed in the BM displacement and CM level functions of adult bats (Russell and Kössl, 1999; Russell et al., 2003). The gain of cochlear amplification, as measured by the relative displacements along the y-axis of the onset and steady-state level functions (Fig. 3B, C), increases with increasing age and an upward shift in resonance frequency (Kössl et al., 2003), as shown in Figure 3D. The relationship between the gain of cochlear amplification and the frequency of the cochlear resonance is not constant. From birth to 2 weeks of age, when the forearm length is 32–34 mm and the resonance is ~53 kHz (Kössl et al., 2003), the gain of cochlear amplification increases by 1 dB for each 1 kHz increase in cochlear resonance. From 2 weeks of age until adulthood, the cochlear gain increases by 5 dB for each 1 kHz increase in the cochlear resonance.

The resonance peak of the magnitude–frequency function is amplified, sharpened, and shifted upward in frequency with time from tone onset

Magnitude–frequency curves measured at constant sound pressure levels and at different times from tone onset are shown in Figure 4 for six bats at developmental stages that advance from newborn (Fig. 4A) to mature (Fig. 4F). The data are based on fast Fourier transforms (FFTs) applied to 1 msec time windows with delays that are incremented in 0.2 msec steps from 0.4 to 2 msec and then in 1 msec intervals to 9 msec from tone onset. CM magnitude–frequency curves were measured during a 1 msec window in the ringing that follows the tone commencing 0.5 msec from tone offset (Fig. 4, thick traces). Magnitude–frequency functions measured from the youngest bat (Fig. 4A) change only in magnitude and not in shape with increasing time from tone onset. The frequency–magnitude curves are also very noisy at sound levels that do not elicit noisy responses from the cochlea of more mature bats (compare Fig. 4B–F). Frequency–magnitude curves measured from bats at later developmental stages also increase in magnitude with time from tone onset. Within 1 msec of tone onset, the curves resemble low-pass filter functions. In a slightly older specimen (Fig. 4B), the curves continue to resemble low-pass filter functions throughout the ongoing tone, whereas the ringing that follows the tone offset behaves as a resonance with a peak at 51 kHz (dashed line). The low-pass characteristics of the frequency–magnitude curves of older bats (Fig. 4C–F) at tone onset evolve into narrowly tuned bandpass functions with increasing time from tone onset with resonance peaks (dashed lines) that match the resonance of the CM ringing (thick traces). In bats close to maturity (Fig. 4E), the CM magnitude–frequency functions have two peaks (Kössl et al., 2003) (dashed and dotted lines). The lower-frequency peak appears within 0.8 msec of tone onset and becomes narrower with increasing measurement delay from tone onset. The higher-frequency peak, which is tuned to 61 kHz (dotted line), appears 1.8–2 msec from tone onset and is lower in frequency than the second peak in the ringing, which is tuned to 61.2 kHz.
Beating

Beating can be recorded from the cochleas of immature bats during the ongoing tone at frequencies close to the resonance (Fig. 5A). By measuring the period of this beating, which decreases as the stimulus frequency departs from the resonance (Fig. 5A, B), it is apparent that the periodicity is attributable to beating between the stimulus and resonance. Beating disappears, and hence the beating period tends to infinity, at the resonance frequency, and this is indicated by the vertical dotted line and arrow in Figure 5B. The resonance can also be deduced by adding (subtracting) the difference between the stimulus frequency and the resonance frequency as the stimulus frequency passes through the resonance (Fig. 5C). The periodicity in the offset resonance is independent. For frequencies above resonance, CM phase becomes almost constant and frequency-independent. For frequencies above resonance, CM phase returns to a phase lag of 100°–120°/kHz with increasing frequency.

CM Phase measurements

Measurements of CM phase from the developing auditory fovea can provide information about the nature and development of cochlear amplification and the properties and development of the cochlear resonance. According to models and direct measurements from the cochlea (Geisler and Sang, 1995; Markin and Hudspeth, 1995; Gummer et al., 1996; Nilsen and Russell, 1999), cochlear amplification is optimal when energy from the OHCs is fed back at maximum basilar membrane velocity. Thus, one might expect that CM from the acoustic fovea with feedback would phase lead by ~90° that from the fovea without feedback. Phase measurements also provide an indication of the relative movements of the structures that form the basis of the cochlear resonance and of the frequency and sharpness of the resonance, as indicated by the appearance and slopes of the phase transitions that should accompany the resonance.

Representative examples of phase as a function of frequency measured from the CM recorded from the cochleas of bats at different developmental stages, as indicated by their FALs, are shown in Figure 6 [Fig. 6A, FAL, 21.5 mm (week 1), 80 dB SPL; B, FAL, 29.5 mm (week 2), 60 dB SPL; C, FAL, 40 mm (week 3–4), 55 dB SPL; D, FAL, 46 mm (week 4–5), 60 dB SPL]. The corresponding magnitude functions are plotted in the bottom parts. Phase and magnitude were measured within 1.5 msec of tone onset (squares, CM_{ON}), during the steady state of the tone (4–9 msec from tone onset; filled circles, CM_{STeady}), and 0.5–1.5 msec after tone offset (triangles, CM_{OFF}). Phase differences (CM_{STeady} – CM_{ON}, thin line; CM_{OFF} – CM_{ON}, thick line) are shown in the top-most parts.

Phase–frequency relations of CM_{ON}

For frequencies within 5 kHz below the cochlear resonance, the slope of the CM_{ON} phase–frequency relationship is 100–120°/kHz. For frequencies within 0.2–0.4 kHz of the resonance frequency, CM_{ON} phase becomes almost constant and frequency-independent. For frequencies above resonance, CM_{ON} returns to a phase lag of 100–120°/kHz with increasing frequency.

CM_{STeady} phase leads CM_{OFF} below the resonance

The thin solid lines in the top parts of Figure 6 represent the phase difference between the CM_{ON} and CM_{STeady}. CM_{STeady} leads CM_{ON} by ~90° for frequencies below the resonance in all bats of different developmental stages. The slope of CM_{STeady} is similar to that of CM_{ON} (100–120°/kHz); CM_{ON} continues to lag CM_{STeady} for frequencies above the resonance in recordings made from the two most immature bats (Fig. 6A, B). CM_{OFF} and CM_{STeady} are in phase for frequencies within 1 kHz above the resonance in the more mature bats (Fig. 6C, D).

CM_{OFF} phase leads and then lags CM_{ON} below and above the resonance

The thick lines in the top parts of Figures 6 represent the phase difference between CM_{OFF} and CM_{ON}. The phase of CM_{OFF} leads
CMOFF by \(\sim 180^\circ\) for frequencies within 1 kHz below the resonance and lags CMON by a similar amount within 1 kHz above the resonance (Fig. 6). The slope of the CMOFF phase–stimulus frequency relationship is very steep through the resonance, and it intersects the phase–stimulus frequency relationships of CMON and CMsteady, an indication that a very sharp resonator produces CMOFF.

At all developmental stages and for frequencies within 500 Hz of the resonance, CMON phase lags CMsteady for frequencies below the resonance by 97.7 \(\pm\) 13.4° (mean \(\pm\) SD; \(n = 11\)). This is an indication that at tone onset in the cochlear fovea, amplification is minimal and occurs at maximum BM velocity during CMsteady. In all but the two bats from which we made measurements (data from one of which is shown in Fig. 6A) CMsteady and CMON are almost in phase for frequencies immediately above the resonance; i.e., CMON phase lags CMsteady by 6.6 \(\pm\) 9.3° (mean \(\pm\) SD; \(n = 9\)). In the two youngest bats, CMON phase lags CMsteady for frequencies above the resonance by 81.6° (Fig. 6A) and 92.3° (data not shown), respectively. In all bats, including the youngest, for which this analysis was performed, CMOFF phase leads CMON by \(\sim 180^\circ\) for frequencies below the resonance (176.7 \(\pm\) 23.5°, mean \(\pm\) SD; \(n = 11\)) and lags CMON by a similar amount for frequencies above the resonance (180.8 \(\pm\) 17.3°, mean \(\pm\) SD; \(n = 11\)), thereby resulting in a sharp transition in phase of CMOFF at the resonance frequency.

Temporal changes in CM frequency functions

Temporal changes in CM responses were measured to tones at frequencies near the resonance. We measured the frequency of the CM at 0.1 msec intervals during the first 2 msec of the tone and then at intervals of 1 msec during the remainder of the 10 msec tone and for up to 5 msec after the cessation of the tone. These measurements were made for a series of tones stepped either in 0.1 or 0.2 kHz increments from 2 to 5 kHz above and below the resonance frequency. The frequency was measured from 8192-point FFTs that were applied to 1 msec sample periods of the CM. Figure 7A gives an example of CM frequency as a function of time recorded from a 4-week-old immature, bat (FAL, 48 mm) in response to 10 msec tone bursts that are incremented in 0.2 kHz steps from 53 to 63 kHz. The vertical gray bars indicate the sample periods at tone onset, at steady state, and after the cessation of the tone during which measurements were made to provide the data shown in Figure 7B. In Figure 7B is shown the frequency (top part, thin line) and amplitude (bottom part, stars) of the CM measured at tone onset (CMON) and the frequency (top part, thick line) and amplitude (bottom part, triangles) of the CM in the ringing that follows the tone (CMOFF) as functions of stimulus frequency. At tone onset, CMON responds not to the stimulus frequency (diagonal dotted line) but is several hundred hertz less than this. The frequency of CMOFF changes with increasing stimulus frequency at an average rate of 0.83 \(\pm\) 0.2 Hz/Hz from 53 to 63 kHz (Fig. 7B). The change is irregular and occurs in discrete steps (arrows). Between frequency steps, CMOFF frequency remains constant for several hundred hertz, and in the stimulus frequency range of 55.5–59.5 kHz, the frequency of CMOFF remains constant at the frequency of the principal resonance, which for this bat is 58.2 kHz.

In Figure 8, CM frequency as a function of stimulus frequency is shown for four bats at increasing stages of development (Fig. 8A, FAL, 21.5 mm, 80 dB SPL; B, 29.5 mm, 65 dB SPL; C, 48 mm, 65 dB SPL; D, 50 mm, 65 dB SPL). The frequency of CMON (top parts, thin lines) recorded from the youngest bat (Fig. 8A) increases with increasing stimulus frequency in a series of discrete steps, which are indicated by arrows in Figure 8A. The transitions in the steps are associated with minima in the magnitude frequency functions of CMON (stars), as indicated by arrows in the bottom traces of Figure 8A. The frequency of CMOFF (which, for clarity, has been displaced by 2 kHz to higher frequencies) also increases as a series of steps. These steps are not as defined as those in CMON, and the interstep frequency is 1.5 times that of CMON. There is a small region that extends \(- 1\) kHz above the resonance (bottom part, filled triangles), indicated by the horizontal dashed line in the top part, where the frequency of CMOFF is independent of stimulus frequency. With advancing development, the dependence of CMON on stimulus frequency remains stepwise, although the steps may not always be so clearly apparent as that seen in the youngest bat. Thus, the frequency of CMON hunts at approximately the stimulus frequency either with transient excursions (see Figs. 7, 8A,B, 10A) as in the youngest bats or with smooth excursions (Fig. 8C,D). These differences are attrib-
utable to subtle differences between cochleae in the development of CM from tone onset because CM \textsubscript{ON} progresses from stepwise to smooth excursion at approximately the stimulus frequency with increasing time from tone onset (see Fig. 10C). In some bats, the mean of CM \textsubscript{ON} tends to be proportional to stimulus frequency; e.g., the regression of CM \textsubscript{ON} recorded fromcmbat21 (Fig. 7B) is 0.83 \pm 0.02 Hz/Hz, and that recorded from cmbat26 (Fig. 8B) is 0.94 \pm 0.02 Hz/Hz. With advancing development, the frequency of CM \textsubscript{OFF} becomes more stable and constant in response to stimulus frequencies at approximately the resonance frequencies (Figs. 8B–D, 9A). Even stimulus frequencies \( \geq 2 \) kHz above the resonance cause ringing at the primary resonance frequency (59.54 kHz) at the tone offset. A second resonance peak in the CM \textsubscript{OFF} frequency–stimulus frequency relationship shown in Figure 8C in response to a stimulus frequency of 63.5 kHz causes a ringing at the primary resonance frequency of 59.54 kHz.

In all specimens irrespective of age, CM \textsubscript{ON} is always a few hundred hertz below that of CM \textsubscript{OFF} when driven by a resonance frequency tone. This difference in frequency response is made more apparent in Figure 9A. The frequency difference between CM \textsubscript{ON} and CM \textsubscript{OFF} in response to stimulus frequency tones tends to decrease during development, as shown in Figure 9B, where CM \textsubscript{OFF} – CM \textsubscript{ON} is plotted as a function of cochlear resonance frequency, which increases during development (Kössl et al., 2003). The regression slope is \(-0.0273 \pm 0.006\), and the relationship is given as CM \textsubscript{OFF} = CM \textsubscript{ON} \(-0.0273(CM \textsubscript{OFF}) + 1.834\), which provides an intercept of \(-65 \) kHz, which is \( \sim 3 \) kHz above the resonance for a mature bat.

The frequency dependence of CM \textsubscript{ON} and CM \textsubscript{OFF} during development is very similar to that measured from the cochleas of adult bats (Russell et al., 2003). This similarity is exemplified by measurements made from a bat at an advanced developmental stage (FAL, 47 mm; Fig. 10). The frequency of CM \textsubscript{OFF} increases as a series of discrete steps with increasing stimulus frequency (Fig. 10A, horizontal dashed lines), an indication that the oscillator that is responsible for the sustained ringing that follows tone offset can switch from one frequency mode to another depending on the frequency of the driving tone. CM \textsubscript{ON} behaves as if it is generated by an oscillator that is attracted to the stimulus frequency and has several frequency modes that are related to those of the CM \textsubscript{OFF} oscillator but with frequencies \( \sim 1 \) kHz below those of the CM \textsubscript{OFF} oscillator. Coupling between the CM \textsubscript{ON} oscillator and the stimulus frequency increases with the time from the measurement of the frequency of CM \textsubscript{ON} from tone onset. This is shown in Figure 10B, where CM frequency is plotted as a function of stimulus frequency for CM frequency measured with delays of 0.3–0.9 msec from tone onset. With increasing delay, the CM frequency approaches more closely the stimulus frequency (diagonal dotted line). Within the 52–62 kHz frequency range, CM \textsubscript{ON} frequency intersects and crosses the stimulus frequency at 52.3, 56.8, and 60.8 kHz (vertical dotted lines, transition frequencies of the principal modes of the onset oscillator). The transition frequencies indicate the bandwidth of the principal mode of the cochlear resonance region (Russell et al., 2003).

The relation between CM frequency and stimulus frequency is more apparent in Figure 10C, where the frequency difference between the CM and the stimulus is plotted against stimulus frequency. When measured only 0.3 msec from tone onset, CM \textsubscript{ON} is most strongly attracted away from the stimulus at the transition frequency close to 55 kHz and progresses from stepwise to smooth excursions at approximately the stimulus frequency with increasing time from tone onset (Fig. 10C). Thus, the differences seen in frequency dependence of CM \textsubscript{ON} among different specimens of young bats (Figs. 7, 8D, 10A) may be attributable to subtle differences between cochleae in the development of CM from tone onset. In Figure 10D, the phase, frequency, and magnitude of CM \textsubscript{ON} measured 0.6 msec from tone onset are plotted as functions of stimulus frequency. The frequency transitions (vertical dashed lines) are associated with magnitude minima and phase transitions. Within the stimulus frequency range of 52–62 kHz, there are three frequency modes

![Figure 6. A–F, CM phase (top part) and CM magnitude (bottom part) as functions of frequency for four immature bats (A, FAL, 21.5 mm, 80 dB SPL; B, FAL, 29.5 mm, 60 dB SPL; C, FAL, 40 mm, 55 dB SPL; D, FAL, 46 mm, 60 dB SPL.) Open squares, measurements within 1 msec of tone onset of a 10 msec tone (onset); filled circles, measurements beginning 6 msec of tone onset (steady state); open triangles, measurements beginning 0.5 msec after the tone offset (offset). Top part, Thick line, Offset phase – onset phase; thin line, steady state – onset phase; vertical dotted line, resonance frequency. All measurements were based on 8196-point FFTs applied to 1 msec sample periods of the CM responses to 10 msec tones.](image-url)
of CMON at 52.3, 56.8, and 60.8 kHz, which are shown by the vertical dotted lines in Figure 10D. When measured with a delay of 0.6 msec from tone onset, the relative coupling between the frequency modes of the onset oscillator and the stimulus frequency causes the frequency of CMON to change at a rate of 0.39 Hz/Hz with stimulus frequency between the transition points.

Discussion
Amplification and compression increase with cochlear maturation
The echo-processing 61 kHz region of the mustached bat’s cochlea (auditory fovea) provides a unique opportunity to investigate the development of a single frequency place on the cochlea by using the relatively noninvasive technique of measuring CM from the cochlear aqueduct, which is dominated by OHC responses from the fovea.

In response to resonance frequency tones, CMsteady, recorded from the mature bat cochlea grows compressively with increasing level (Russell et al., 2003). CMsteady is 50 – 60 dB more sensitive at low levels and is more sharply tuned than the linear growth of CMON, which CMsteady phrase leads by 90°. The increased sensitivity and compression of CMsteady compared with CMON has been taken to indicate that at tone onset, the CM is from a passive cochlea. Over a > 1 msec, OHCs amplify the responses of the cochlea by boosting the vibration of the BM during maximum velocity, and they compress the responses of the cochlea at high levels (for review, see Robles and Ruggero, 2001; Russell et al., 2003). This process begins in the first week of a mustached bat’s life, when CMsteady is 10 – 15 dB more sensitive than CMON. By the fifth week, CMsteady is 35 – 40 dB more sensitive than CMON. Amplification develops slowly for the first 2 weeks of life and then accelerates in the second to fifth weeks (Fig. 3D). By contrast, tuning of the resonance develops rapidly over the first 3 weeks of development and then slows down (Kössl et al., 2003). Thus, the development of cochlear amplification occurs separately and continues after the development of frequency tuning has begun to asymptote.

Transducer operating point shifts during development
From birth, CMON phase lead CMsteady by 90° below the resonance, an indication that the timing of cochlear feedback appears to be close to optimal at maximum BM velocity. However, neither the gain nor the frequency tuning of the immature bat cochlea matches that of the mature cochlea. One possible reason for this discrepancy is that OHCs in the immature bat cochlea do not always operate at approximately the most sensitive point of the hair cell transducer function (Russell et al., 1986). The CM recorded from mature bats, is symmetrical for frequencies above and below the resonance for levels of < 80 dB SPL (Russell et al., 2003). This finding agrees with basal turn OHC responses from noncholocating mammals (Russell et al., 1986), in which OHC hair bundles react against the TM (Legan et al., 2000) so that ~50% of the transducer channels are open at rest, OHCs operate at the steepest point on the transducer function, and the DC component is negligible (Russell and Kössl, 1992). OHCs appear not to function at approximately their most sensitive operating point in immature bats, as indicated by a DC component that is present at all levels for frequencies at and above the resonance; with the consequence that cochlear feedback is not optimal. It is suggested that the axial OHC motility (Brownell et al., 1985) is responsible for maintaining the OHC operating point (Russell and Kössl, 1992). Perhaps in the cochleas of immature bats (< 4 – 5 weeks old), at and above the resonance frequency, the mechanical impedance of the cochlear partition is too great for OHCs to exert sufficient force against the TM to minimize the receptor potential DC component and thereby to maintain the operating point of the transducer function at its most sensitive position.

The oscillator frequencies move together and upward during development
CM and micromechanical measurements (Kössl and Russell, 1995; Russell and Kössl, 1999; Russell et al., 2003) from the SI region of the mustached bat’s cochlea have led to the proposal that the cochlear resonance is attributable to OHC-mediated feedback between two oscillators that have been attributed to the TM and the BM.

During development, the frequency of the cochlear resonance shifts upward from ~ 45 kHz in postnatal week 1 to ~ 60 kHz in postnatal week 5, and the frequency separation between CMON and CMOFF decreases from 0.6 to 0.2 kHz. The large frequency gap between the two oscillators in the first week of life may partially account for the finding that CMsteady continues to lead CMON for frequencies above the resonance in the two youngest bats in this study. In older bats, the oscillators are closer in frequency and become mass-limited and, hence, in phase at approximately the same frequency. However, the individual characteristics of the two oscillators tend to remain constant throughout development, at least from week 2 onward, in that the offset-oscillator (TM) is more sharply tuned and takes longer to build up its responses to the resonance frequency than does the broadly
tuned onset oscillator. The offset oscillator phase leads the onset oscillator for frequencies below the resonance and lags it at frequencies above the resonance. As a consequence, there is an abrupt 360° phase transition at the resonance frequency, which is also apparent in distortion product otoacoustic emission measured at the external auditory meatus (Russell and Kössl, 1999).

Development of the cochlear place
It is now well established, from measurements made in the cochleas of other CF bats (Rübsamen, 1987; Vater and Rübsamen, 1989), nonecholocating mammals (Harris and Dallos, 1984; Rübsamen and Lippe, 1998), and birds (Rubel and Ryals, 1983), that the characteristic frequency of a place on the BM shifts upward in frequency during development. If the natural frequency of the resonance is governed by the mass and stiffness of the cochlear partition, then \( \omega = \frac{k}{m}^{1/2} \), where \( k \) is stiffness, and \( m \) is mass. The stiffness of the cochlear partition would have to almost double for an increase in resonance of 0.5 octaves. There are no large-scale postnatal morphological changes in the cochlea of the mustached bat. However, differences in the physiological responses of immature and mature bat cochleas could be explained by differences in cellular skeletal and extracellular matrix morphology (Vater et al., 2003). An upward shift in frequency and enhanced resonance are associated with a stiffening and decreased damping of the cochlea. These changes would be facilitated by mechanical stiffening of cellular elements of the organ of Corti through the incorporation of cytoskeletal proteins, increased radial tension of the cochlear partition through the activity of the tensioning fibroblasts of the outer spiral ligament (Henson and Henson, 1988), and a reduction in the thickness of the tympanal layer of the BM (Vater et al., 2003).

Beating in the cochlea
Frequency beating to near-resonance ongoing tones is seen in CM responses from both mature and juvenile bats and is attributable to interaction between the sharply tuned cochlear resonance and the stimulus tone (Suga et al., 1975). The discovery of beating in the CM recorded from the cochleas of bats that are close to maturity, but not in the resonance of sensitive mature bats, is a new finding. Similar beating has been observed in the CM ringing recorded from mature bats that have undergone gaseous anesthesia (Russell et al., 2003).
al., 2003) and after activation of the cochlear efferents (Suga et al., 2000). The mechanical changes recorded in the cochleas of mature bats have been suggested to reflect mechanical uncoupling between two structural components of the cochlear partition attributable to dephosphorylation of OHC structural proteins (Russell et al., 2003). The more broadly tuned onset oscillator, whose responses build up swiftly to resonance frequency tones, dominates the onset response. The narrowly tuned offset oscillator, which takes time to build up responses to resonance frequency tones, dominates the ringing. CM frequency measured at tone onset from the foveal region of the cochlea is not that of the tone but one of several frequency modes of the acoustic laser. The ringing has only a single mode. Thus, over a few milliseconds from tone onset, OHC-mediated interaction between the two oscillators (presumably the BM and TM) results in acoustic lasing. The lasing modes change according to the stimulus frequency, the relative contributions of the two oscillators, and gain of OHC amplification to the acoustic laser.

In the first week of postnatal development, the mustached bat’s cochlea behaves like an acoustic laser. The onset oscillator has many modes, which depend on stimulus frequency, but the offset oscillator is barely developed and is broadly tuned. The mode-hopping behavior at tone onset is very clear, with sharply defined frequency transitions between the frequency “plateaus.” By the second week, the number of frequency modes that can be elicited by stimulus tones that differ qualitatively throughout the remaining period of postnatal development, although the resonance shifts to higher frequencies with advancing development. The increase in the ringing of CM_{OFF} appears to be correlated with an increase in the rate of development of cochlear amplification in the second postnatal week (see Fig. 3D).

The behavior of the onset oscillator remains qualitatively the same throughout the entire period of postnatal development and resembles that of a synchronized oscillator (Strogatz, 1994; Russell et al., 2003). CM frequency in response to tones at and within 10 kHz (at least) of the resonance is a compromise frequency that depends on the stimulus frequency, the frequency mode of the oscillator, and tone duration. CM_{ON} attempts to “hunt” the stimulus frequency, and close to tone offset, the frequency of CM_{ON} sharply undershoots and overshoots the stimulus frequency. These sudden frequency changes are associated with phase reversals and magnitude minima (Fig. 10D). A few milliseconds later, the system become more synchronized, and the CM_{ON} frequency overshoots and undershoots become smoother (Fig. 10C). This is possibly attributable to the temporally related increase in OHC gain, which mediates the interaction between the onset and offset oscillators. In some bats (see Fig. 7B, 8B), CM_{ON} hunts not the stimulus frequency but a frequency proportional to it (indicated by the dashed lines). This might be because the frequencies of the onset and offset oscillators in these bats are far apart, and CM_{ON}
is a compromise frequency that is driven by both oscillators (Russell et al., 2003).

Thus, the foveal region of the mustached bat cochlea functions as an acoustic laser from birth. The foveal region of the cochlea is present, and the specialized basoapical gradients in morphology of the basilar and tectorial membranes that are essential components of the acoustic laser (Russell and Kössl, 1999) are established (Vater et al., 2003). Changes in the structure of the extracellular matrices and cytoskeletons of the cellular components of the cochlea, rather than in gross morphology, appear to be responsible for the maturation and sharp tuning of this highly resonant biological structure.

References


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