Tonotopic variations of calcium signalling in turtle auditory hair cells

A. J. Ricci, M. Gray-Keller and R. Fettiplace

*J. Physiol.* 2000;524;423-436

This information is current as of February 21, 2008

This is the final published version of this article; it is available at:

http://jp.physoc.org/cgi/content/full/524/2/423

This version of the article may not be posted on a public website for 12 months after publication unless article is open access.
Tonotopic variations of calcium signalling in turtle auditory hair cells

A. J. Ricci, M. Gray-Keller and R. Fettiplace

Department of Physiology, University of Wisconsin Medical School, Madison, WI 53706, USA

(Received 12 October 1999; accepted after revision 13 January 2000)

1. Turtle cochlear hair cells are electrically tuned by a voltage-dependent Ca\textsuperscript{2+} current and a Ca\textsuperscript{2+}-dependent K\textsuperscript{+} current (I\textsubscript{BKCa}). The effects of intracellular calcium buffering on electrical tuning were studied in hair cells at apical and basal cochlear locations tuned to 100 and 300 Hz, respectively.

2. Increasing the intracellular BAPTA concentration changed the hair cell’s resonant frequency little, but optimized tuning at more depolarized membrane potentials due to a positive shift in the half-activation voltage (V\textsubscript{A}) of the I\textsubscript{BKCa}.

3. The shift in V\textsubscript{A} depended similarly on BAPTA concentration in basal and apical hair cells despite a 2-4-fold difference in the size of the Ca\textsuperscript{2+} current at the two positions. The Ca\textsuperscript{2+} current amplitude increased exponentially with distance along the cochlea.

4. Comparison of V\textsubscript{A} values and tuning properties using different BAPTA concentrations with values measured in perforated-patch recordings gave the endogenous calcium buffer as equivalent to 0.21 mm BAPTA in low-frequency cells, and 0.46 mm BAPTA in high-frequency cells.

5. High conductance Ca\textsuperscript{2+}-activated K\textsuperscript{+} (BK\textsubscript{Ca}) channels recorded in inside-out membrane patches were 2-fold less Ca\textsuperscript{2+} sensitive in high-frequency than in low-frequency cells.

6. Confocal Ca\textsuperscript{2+} imaging using the fluorescent indicator Calcium Green-1 revealed about twice as many hotspots of Ca\textsuperscript{2+} entry during depolarization in high-frequency compared to low-frequency hair cells.

7. We suggest that each BK\textsubscript{Ca} channel is gated by Ca\textsuperscript{2+} entry through a few nearby Ca\textsuperscript{2+} channels, and that Ca\textsuperscript{2+} and BK\textsubscript{Ca} channels occupy, at constant channel density, a greater fraction of the membrane area in high-frequency cells than in low-frequency cells.

Hair cells of the turtle cochlea are frequency tuned through the gating of large-conductance Ca\textsuperscript{2+}-dependent K\textsuperscript{+} (BK\textsubscript{Ca}) channels that are activated by Ca\textsuperscript{2+} influx through voltage-dependent L-type Ca\textsuperscript{2+} channels (Art & Fettiplace, 1987). The resonant frequency of a hair cell is correlated with its complement of Ca\textsuperscript{2+} and BK\textsubscript{Ca} channels, cells tuned to higher frequencies possessing more channels of both types (Wu et al. 1995). To achieve local Ca\textsuperscript{2+} concentrations of sufficient magnitude to activate the BK\textsubscript{Ca}, the Ca\textsuperscript{2+} and BK\textsubscript{Ca} channels are thought to be co-localized in clusters of high channel density proposed to coincide with the synaptic release sites (Roberts et al. 1990). The multiple hotspots of Ca\textsuperscript{2+} entry observed with confocal microscopy (Issa & Hudspeth, 1994; Tucker & Fettiplace, 1995; Hall et al. 1997) are consistent with the clustered arrangement of Ca\textsuperscript{2+} channels on the basolateral aspect of hair cells. With such an arrangement, how are the numbers of Ca\textsuperscript{2+} channels regulated with changes in the cell’s resonant frequency? For example, an increase in resonant frequency might be associated with either a higher Ca\textsuperscript{2+} channel density in a cluster or an enlargement of the membrane area devoted to the clusters. In support of the latter hypothesis, there is evidence in both turtle (Sneary, 1988) and chicken (Martinez-Dunat et al. 1997) that the number of hair cell transmitter release sites and, by implication, Ca\textsuperscript{2+}-channel clusters, increases along the tonotopic axis of the cochlea.

Both the amplitude and timing of Ca\textsuperscript{2+} excursions at the BK\textsubscript{Ca} channel will depend on the microstructure of the clusters, especially the density of the two channel types. Local Ca\textsuperscript{2+} gradients will also be accentuated by the concentration and kinetics of the mobile intracellular calcium buffers (Stern, 1992; Roberts, 1994; Naraghi & Neher, 1997). Depending on the overall channel density and the buffer concentration, each BK\textsubscript{Ca} channel might be influenced by all Ca\textsuperscript{2+} channels in the cluster or by only a few nearest neighbours. To approach this problem we have examined the effects of exogenous calcium buffers on the activation of the I\textsubscript{BKCa} and the hair cell’s tuning properties. Comparison
with results derived from perforated-patch recordings have allowed us to estimate the effective concentration of endogenous calcium buffer (Zhou & Neher, 1993; Roberts, 1993; Tucker & Pettitplace, 1996; Ricci et al. 1998). We have collected data from hair cells at two cochlear locations, enabling us to study cells with different magnitudes of $\text{Ca}^{2+}$ current. Our results suggest that each BKCa channel is gated by $\text{Ca}^{2+}$ entry through a few adjacent $\text{Ca}^{2+}$ channels, and that an increase in channel numbers is accomplished by augmentation of clusters at constant channel density.

**METHODS**

**Preparation**

The preparation and method of hair cell recording in the intact basilar papilla have been described previously (Ricci & Pettitplace, 1997). Turtle (Trachemys scripta elegans, carapace length 75–100 mm) were decapitated and the cochlea dissected out using procedures approved by the Animal Care Committee at the University of Wisconsin (protocol number A3368-01). The tectorial membrane was removed following 20 min incubation in turtle saline, composition (mM): NaCl, 125; KCl, 4; CaCl2, 2.8; MgCl2, 2.2; sodium pyruvate, 2; glucose, 8; NaHepes, 10 (pH 7.6) containing up to 50 μM of protease (Sigma type XXIV). The preparation was transferred to a recording chamber on the stage of a Zeiss AxioSkop FS microscope and viewed through a ×63 water immersion objective (NA 0.9) and a Hamamatsu C2400 CCD camera. The chamber was continuously perfused with turtle saline. For voltage-clamp measurements at the low-frequency location, 1 mM 4-aminopyridine and 0.1 μM ampin (Calbiochem, San Diego, CA, USA) were added to the saline to block K+ channels other than the large-conductance BKCa channel. Other K+ channels are known to occur in low frequency hair cells (Goodman & Art, 1996), and their presence complicates the analysis of the BKCa current and $I_{\text{BKCa}}$.

Whole-cell currents were measured with a List EPC-7 amplifier attached to a borosilicate patch electrode that was advanced from the abneural edge of the basilar papilla to seal on to the basolateral membrane of a hair cell (Ricci & Pettitplace, 1997). Recordings usually came from cells in the middle of the papilla two to three cells in from the abneural edge. Hair cell location and total length of the basilar papilla were documented at the end of an experiment. Most recordings came from cells in two regions, at about 0.3 (the apical location) or 0.6 (the basal location) of the distance along the cochlea from the low-frequency end. Measurements were given as means ±1 standard error of the mean (s.e.m.). Membrane currents were stored on a Sony PCM108 recorder at a bandwidth of 0–20 kHz. Experiments were performed at 21–23 °C.

**Electrical recordings**

Whole-cell electrodes were normally filled with a solution containing (mM): KCl, 125; Na$_2$ATP, 3; MgCl$_2$, 2; K$_2$Hepes, 10, adjusted to pH 7.2 with KOH, with the addition of various concentrations (0.1–30 μM) of the calcium buffers BAPTA, niro-BAPTA, (Molecular Probes, Eugene, OR, USA) or EOTA (Fluka, Ronkonkoma, NY, USA). With 10 or 30 μM calcium buffer, the KCl concentration was reduced to maintain constant osmolarity. To measure voltage-dependent $\text{Ca}^{2+}$ currents, $\text{Ca}^{2+}$ was substituted for $K^+$ as the major monovalent ion (see below). After applying up to 50% series resistance compensation, electrode access resistances were 3–10 MΩ, giving recording time constants of 30–150 μs. Membrane potentials were corrected for the uncompensated series resistance and junction potential.

The method of perforated-patch recording was identical to that used previously (Horn & Marty, 1988; Ricci et al. 1998). The electrode solution contained (mM): potassium aspartate, 110; KCl, 15; MgCl$_2$, 2; K$_2$BAPTA, 0.1 or 1; Hepes, 10, neutralized to pH 7.2 with KOH. For each experiment, 2.4 mg nystatin (Calbiochem, San Diego, CA, USA) was dissolved in 10 μl dry dimethyl sulfoxide and diluted 1:1000 in the stock intracellular solution. The patch pipette was tip filled with antibiotic-free stock solution, and back-filled with the nystatin solution to prevent the antibiotic leaking into the bath during penetration of the papilla and sealing to the membrane. In general, it took at least 15 min from sealing to achieve a stable low-resistance patch perfusion, after which time the recordings were stable for at least a further 30 min. During this period there was no evidence of either run down in the currents or $\text{Ca}^{2+}$ leakage. Although the perforated-patch recordings did not allow wash in of ATP, the $\text{Ca}^{2+}$-ATPase was clearly functional, since there was never any indication during repeated depolarizations of the slow component of the $I_{\text{BKCa}}$ tail current that appeared when the $\text{Ca}^{2+}$-ATPase was blocked with vanadate (see Results). Series resistance for the perforated-patch recordings were 7–20 MΩ after applying up to 40% series resistance compensation. Potentials were adjusted for a 10 mV junction potential between the potassium aspartate solution and the external saline.

The amplitude of the $\text{Ca}^{2+}$ current in a given hair cell showed up to a 2-fold increase during the first 5 min after attaining the whole-cell configuration. A time-dependent growth of the $I_{\text{BKCa}}$ was also observed in both whole-cell and perforated-patch measurements, which may reflect the augmentation of the $\text{Ca}^{2+}$ current. However, the variations in $I_{\text{BKCa}}$ size were not accompanied by any significant shift in the channel’s activation curve. For example, 6 of the 11 cells recorded in perforated-patch mode showed a 35% mean increase in $I_{\text{BKCa}}$ during the course of the recording, but below 1 mV mean shift in the half-activating voltage ($V_1/2$, see eqn (1)). It was difficult to study the phenomenon systematically due to the time required for equilibration of the patch electrode solution with the cytoplasm in whole-cell recording, and the time to achieve low-resistance access in perforated-patch mode. However, measurements were not normally taken until the current amplitude had stabilized, and the activation curve parameters quoted in the Results are those obtained when a steady state had been reached.

**Macropatch recordings**

$I_{\text{BKCa}}$ were measured in inside-out membrane macropatches excised from hair cells at known locations (Art et al., 1995). Electrodes (resistance 0.5–1 MΩ), connected to an Axopatch 200 amplifier, were filled with a solution (mM): KCl, 130; K$_2$EOTA, 5; K$_2$Hepes, 10; pH 7.4. The intracellular face of the patch was exposed to solutions (mM): KCl, 130; K$_2$dibromobisBAPTA, 2 (Molecular Probes, Eugene, OR, USA); dithiothreitol, 1; K$_2$Hepes, 10, pH 7.4, with different amounts of CaCl$_2$ added to yield free $\text{Ca}^{2+}$ concentrations from 0.3–20 μM. Solutions with free $\text{Ca}^{2+}$ greater than 20 μM contained no dibromobisBAPTA. $\text{Ca}^{2+}$ activities in all samples were measured with a MI-600 Calcium electrode (Microelectrodes Inc., Londonderry, NH, USA) calibrated in a series of standard $\text{Ca}^{2+}$ buffer solutions (WPI, Sarasota, FL, USA). Leak currents, measured in a nominally zero-$\text{Ca}^{2+}$ solution containing 10 mM EOTA, were subtracted from all traces. Current responses were filtered with an eight-pole Bessel filter at 5 kHz prior to digitization and analysis.

**Confocal imaging**

The composition of the patch-electrode solution, both for imaging and for measuring voltage-dependent $\text{Ca}^{2+}$ currents, was (mM): CaCl$_2$, 110; Na$_2$ATP, 3; MgCl$_2$, 2; Na$_2$GTP, 0.3; creatine
phosphate, 5; ascorbic acid, 1; EGTA (or BAPTA), 1; Hepes, 10; adjusted to pH 7.2 with CsOH. Spatial distributions of intracellular Ca\(^{2+}\) transients were measured with 0.1 mM K\(_4\) Calcium Green-1 as the 3000 Da–dextran conjugate (Molecular Probes, Eugene, OR, USA) in the patch-electrode solution. Cells were illuminated with the 488 nm line of an argon laser and fluorescence images, passed through a 515 nm long pass filter, were collected with an Odyssey real-time laser scanning confocal microscope (Norrin Instruments, Middleton, WI, USA) attached to the TV port of the Axioskop FS. Images were stored on an S-VHS videocassette recorder (Sony SVO 9500 MD) or an optical disc recorder (Panasonic TQ 3031F). Images were later analysed with Metamorph software (Universal Imaging).

### Figure 1: Tonotopic organization of the turtle basilar papilla

Left, surface view of the basilar papilla during an experiment showing the epithelial strip of hair cells on the right-hand side, apical (lagena) end at the top and basal (saccular) end at the bottom. Scale bar, 100 \(\mu\)m. Fractional distance \(d\) from lagena end shown on left. Most of the measurements of the buffer effects were taken at values of \(d\) of approximately 0.3 or 0.6. Right, examples of electrical resonance recorded in hair cells at the locations indicated. Each record is the response to 25 presentations of a small depolarizing current step evoking oscillations in membrane potential at the start and end of the step. Recordings were obtained with electrodes containing 0.1 or 1 mM BAPTA as the calcium buffer. For some of the cells, the resting potential was about −70 mV, and thus the current step was superimposed on a standing current to depolarize the cell into the range where tuning was optimal. Resonant frequencies are given next to traces. Membrane potentials prior to current step: −51 mV (63 Hz), −47 mV (97 Hz), −47 mV (124 Hz), −39 mV (254 Hz), −44 mV (345 Hz) and −44 mV (400 Hz).
RESULTS

Map of the voltage-dependent Ca$^{2+}$ current along the tonotopic axis

Recording in the isolated basilar papilla allows the ionic properties of a hair cell to be measured in cells of known location. The turtle basilar papilla is tonotopically organized with a hair cells resonant frequency increasing along the long axis of the papilla. Figure 1 shows examples of the tuning properties of hair cells, the resonant frequency being deduced from the period of oscillations at the onset of an injected current step. The inferred tonotopic map is similar to that described in the turtle half-head (Crawford & Pettitplace, 1980). Figure 2 shows that the magnitude of the voltage-dependent Ca$^{2+}$ current increased in parallel with the cell's resonant frequency. To construct the plot, results from cells within each 100 μm length of the papilla were pooled, giving mean values that show an exponential increase in current size with fractional distance along the papilla from the low-frequency end (Fig. 2C). The value of 0.35 for the space constant $\lambda$, is similar to the average value of $\lambda$ for the frequency map (0.37; Wu & Pettitplace, 1996), indicating that the hair cell Ca$^{2+}$ current increases in proportion to resonant frequency. For subsequent analysis of $I_{\text{BK,ca}}$, we compared measurements at two locations with mean fractional distances along the papilla of 0.29 ± 0.05 (low frequency) and 0.6 ± 0.04 (high frequency). The peak Ca$^{2+}$ current was 0.30 nA at the low-frequency location and 0.72 nA at the high-frequency location. The Ca$^{2+}$ current activation curve was similar for 1 and 10 mM BAPTA intracellular calcium buffer (Fig. 2B).

Effects of calcium buffers on activation of the BK$_{\text{ca}}$ channel

Ca$^{2+}$ influx via the voltage-dependent Ca$^{2+}$ channel serves the dual function in hair cells of gating the BK$_{\text{ca}}$ channel involved in frequency tuning (Art & Pettitplace, 1987; Roberts et al. 1990), and controlling exocytosis of neuro-
transmitter (Parsons et al. 1994). The effects of intracellular calcium buffering on the BK$_{Ca}$ channel was assessed from shifts in the channel's voltage-activation curve obtained from tail current measurements. The procedure is illustrated in Fig. 3A for perforated-patch recordings where the cells retain their native mobile buffer. The results demonstrate differences in BK$_{Ca}$ channel performance at the two papillary locations. $I_{BKCa}$ were approximately 2-fold larger and faster at the high-frequency location (Fig. 3A). At a holding potential of $-60$ mV, the maximum tail currents and time constants of deactivation for perforated-patch were $0.46 \pm 0.04$ nA and $4.45 \pm 0.06$ ms ($n = 6$) and $0.96 \pm 0.08$ nA and $0.65 \pm 0.03$ ms ($n = 5$) at the low-frequency and high-frequency locations, respectively. Thus cells tuned to higher frequencies possesses larger and faster $I_{BK(Ca)}$, which agrees with previous observations on isolated hair cells (Art & Fettiplace, 1987; Art et al. 1993). Figure 3B shows plots of the tail current $I$, against membrane potential $V$, which have been fitted with the Boltzmann equation:

$$I = I_{max}/(1 + \exp((V - V_0)/V_S)), \quad (1)$$

where $I_{max}$ is the maximum tail current, $V_0$ is the half-activation voltage and $V_S$ is the slope factor. $V_0$ and $V_S$ had mean values of $-46 \pm 0.8$ mV and $1.8 \pm 0.2$ mV ($n = 6$; low-frequency location) and $-42 \pm 1.0$ mV and $2.3 \pm 0.5$ mV ($n = 5$; high-frequency location).

$I_{KCa}$ activation was similarly characterized with exogenous calcium buffers in whole-cell recording (Fig. 3C). At both

![Figure 3. BK$_{Ca}$ channel currents under different calcium buffering conditions](https://example.com/figure3.png)

Figure 3. BK$_{Ca}$ channel currents under different calcium buffering conditions

A, families of tail currents obtained with perforated-patch recordings for 25 ms depolarizing voltage steps of increasing amplitude from a holding potential of $-60$ mV. The top family was obtained for a high-frequency location ($d = 0.63$) and the bottom family from a low-frequency location ($d = 0.31$). Note that the tail currents, representing deactivation of the BK$_{Ca}$ channel, are larger and faster in the high-frequency hair cell. Each trace is the average of 5–25 presentations. B, activation curves of the $I_{BKCa}$ current obtained from plots of the tail current ($I$), against membrane potential ($V$), during the voltage step for the two cells shown in A. The smooth curves are fits to eqn (1) with values of maximum current ($I_{max}$), half-activation voltage ($V_0$) and slope factor ($V_S$) of high frequency, 1.22 nA, $-41$ mV, 1.4 mV (■); low frequency, 0.58 nA, $-46$ mV, 2.1 mV (○). C, average $I_{BKCa}$ in three different cells recorded with 0.1, 10 and 30 mM intracellular BAPTA. The membrane potential during the voltage step is given above each trace, holding potential $-60$ mV. Note that more depolarization is needed to activate the outward current in higher concentrations of BAPTA. In 30 mM BAPTA, the smallest depolarizations elicit only inward Ca$^{2+}$ current. Each trace is the average of 5–25 presentations.
papillary locations, raising the BAPTA concentration from 1 to 10 mM shifted the current activation to more depolarized potentials (Fig. 4). Values of $V_{1/2}$ from whole cell measurements were similar to those obtained in perforated-patch recordings (Fig. 4D). The $V_{1/2}$ values were independent of the size of the maximum current, but there was an ~4 mV difference between the $V_{1/2}$ values at the low- and high-frequency locations for all BAPTA concentrations. A possible reason for this disparity is examined later. The effective concentration of endogenous mobile buffer can be estimated by comparison with the $V_{1/2}$ values obtained in the perforated-patch recordings. Interpolation from the plots in Fig. 4C gave the endogenous buffer as equivalent to 0–20 mM BAPTA at the low-frequency location and 0–47 mM BAPTA at the high-frequency location. The results suggest an increase in endogenous buffer concentration towards the high-frequency end of the cochlea.

The effect of buffer concentration most probably stems from the mobile buffer restricting the spread of Ca$^{2+}$ away from its source, the Ca$^{2+}$ channel. Thus greater depolarization is needed to achieve the same concentration of Ca$^{2+}$ at its binding site on the BK$_{Ca}$ channel. With up to 10 mM BAPTA,

![Figure 4](https://example.com/fig4.png)

**Figure 4. Effects of intracellular calcium buffer on $I_{BK(Ca)}$ activation**
A, example of activation curves obtained from tail current measurements as in Fig. 3A, for cells at a high-frequency location ($d = 0.62$). Intracellular calcium buffer: 0.1 mM BAPTA (○); 1 mM BAPTA (△); 10 mM BAPTA (×) and perforated-patch measurements (PP) for endogenous buffer (■). $B$, activation curves obtained from tail current measurements for cells at a low-frequency location ($d = 0.32$). Symbols as in A. For each set of points in A and B, the tail current has been normalized to its maximum value and fitted with a Boltzmann equation (eqn (1)) to obtain half-activation voltage $V_{1/2}$, and slope factor $V_{S}$, for each condition. $C$, half-activation voltage $V_{1/2}$, plotted against BAPTA concentration. Each point is the mean ± 1 s.e.m. for the low-frequency location (○) and the high-frequency location (■), the number of measurements averaged being given beside each point. The construction lines are interpolations to derive the endogenous buffer from $V_{1/2}$ values of −42 mV (high frequency) and −46 mV (low frequency) obtained with perforated-patch recording (see text). $D$, slope factor $V_{S}$, plotted against BAPTA concentration for the low-frequency location (○) and the high-frequency location (■). Perforated-patch values for $V_{S}$ were 1.8 ± 0.2 mV (low frequency) and 2.3 ± 0.5 mV (high frequency).
it was possible to activate fully the $I_{\text{BK(Ca)}}$ by suitable depolarization. However, when the BAPTA concentration was raised to 30 mM, the $I_{\text{BK(Ca)}}$ could not be completely activated with depolarization, even to the peak of the $\text{Ca}^{2+}$ current at $-20$ mV, and the smallest depolarizations evoked only inward $\text{Ca}^{2+}$ current (Fig. 3C). $V_{\text{th}}$ values were measured for two other calcium buffers, the slow buffer EGTA and the low-affinity buffer nitroBAPTA. At the high-frequency location, the mean $V_{\text{th}}$ value was $-41 \pm 1.0$ mV ($n = 11$) for 10 mM EGTA, and $-40 \pm 0.7$ mV ($n = 7$) for 1 mM nitroBAPTA. These results indicate that nitroBAPTA has a comparable efficacy to BAPTA but EGTA is about 15-fold less effective. NitroBAPTA, although having a low $\text{Ca}^{2+}$ affinity ($K_d 40 \mu M$), has the same forward rate constant as BAPTA, whereas EGTA has a similar affinity to BAPTA ($K_d 0.2 \mu M$) but binds $\text{Ca}^{2+}$ at least 100-fold slower (Naraghi, 1997). Our results agree with those of Roberts (1993), arguing for the importance of the forward rate constant rather than the buffer affinity in influencing $\text{Ca}^{2+}$ activation of the hair cell BK$_{\text{Ca}}$ channel. The results also imply, based on the analysis of Naraghi & Neher (1997), that the two channels must be close neighbours.

**Ca$^{2+}$ accumulation near the BK$_{\text{Ca}}$ channel**

$\text{Ca}^{2+}$ entering through the voltage-dependent channels will be rapidly bound by cytoplasmic buffers and then extruded on a slower time scale by a $\text{Ca}^{2+}$-ATPase (Tucker & Fettiplace 1995). During a prolonged depolarization, $\text{Ca}^{2+}$ might be expected to accumulate beneath the membrane, its concentration growing due to diffusion from neighbouring channels. The BK$_{\text{Ca}}$ channel-activation curves in Fig. 4 were constructed from 25 ms depolarizing voltage steps. Figure 5 shows results from a cell where the duration of the voltage step was varied from 10 to 100 ms. For all amplitudes of depolarization, the outward current attained a maximum level within 2 ms, little longer than it takes the $\text{Ca}^{2+}$ current to fully activate (Art & Fettiplace, 1987), and then remained constant for the duration of the step. Activation

![Figure 5](https://example.com/figure5.png)

**Figure 5. BK$_{\text{Ca}}$ channel activation is independent of the voltage step duration**

A, average currents in response to depolarizing voltage steps of duration 10 ms (left) and 100 ms (right), holding potential of $-60$ mV. B, $I_{\text{BK(Ca)}}$ activation curves for cell in A, derived from the tail currents at the end of voltage steps of duration $10 \text{ ms} (\square), 25 \text{ ms} (\triangle)$ and $100 \text{ ms} (\Delta)$. The tail currents ($I$) have been normalized to their maximum value ($I_{\text{max}}$), which was $4.8 \text{nA}$, independent of step duration. Smooth curve is a fit to eqn (1) with half-activation voltage $V_{\text{th}} = -44$ mV and slope factor $V_S = 3.5$ mV. C, average currents in response to sub-saturating depolarizing voltage steps, with durations from 25 to 1500 ms. Note that the maximum current is independent of the step duration. D, same experiment as in C in another cell, with electrode solution containing 1 mM vanadate to block $\text{Ca}^{2+}$-ATPase pumps. Note the increase in steady current with step duration and the slow component of the tail current, which may reflect clearance of $\text{Ca}^{2+}$ accumulated during the step. In both C and D, the voltage was stepped from a holding potential of $-60$ to $-44$ mV corresponding to the $V_{\text{th}}$ of the channel. Each trace is the average of 5–25 responses. In all panels, the intracellular calcium buffer was 0.1 mM BAPTA.
curves deduced from tail currents were identical for different stimulus durations (Fig. 5B). In other experiments (not illustrated), including a perforated-patch recording, the amplitude of the $I_{K(Ca)}$ was found to be invariant with duration of the voltage step between 2 ms and more than 1000 ms. These results suggest that the time course of free Ca$^{2+}$ at the BK$_{Ca}$ channel resembles that of the Ca$^{2+}$ current itself, and implies that each Ca$^{2+}$ channel influences only nearby BK$_{Ca}$ channels.

Rapid equilibration of Ca$^{2+}$ in the vicinity of its target the BK$_{Ca}$ channel, must partly depend on fast diffusion away from the source without local saturation of the calcium buffer. A constant $I_{K(Ca)}$ was found for sub-maximal stimulation even with the lowest buffer concentration, 0.1 mM BAPTA. However, when the Ca$^{2+}$ extrusion mechanism was blocked by adding 1 mM vanadate to the internal solution, the current acquired a secondary growth phase and a slow component of the tail current on

![Figure 6. Effects of intracellular calcium buffer on electrical tuning](http://example.com/figure6.png)

**Figure 6. Effects of intracellular calcium buffer on electrical tuning**

A, hair cell resonant frequencies plotted against concentration of BAPTA in electrode solution. The resonant frequency was obtained from the period of the oscillations at the onset of a current step that produced the sharpest tuning (the largest quality factor). The membrane potential at which tuning was sharpest is referred to as the 'best resonant voltage'. Each point is the mean ± 1 s.e.m of measurements at the low-frequency location (●) and at the high-frequency location (○). The number of values averaged is given beside each point, the same numbers apply also to the measurements in B and C. B, maximal quality factors of electrical tuning plotted against concentration of BAPTA in electrode solution, conventions as in A. Definition of quality factor is given in text. The small changes in $K_r$ and $Q$ with increasing BAPTA concentration were not statistically significant using a one-way ANOVA test at the 0.01 confidence level. C, the membrane potential at which the quality factor was maximal (best resonant voltage) plotted against internal concentration of BAPTA for cells at the low-frequency location (●) and at the high-frequency location (○). Construction lines are interpolations to derive the endogenous buffer from the 'best resonant voltage' of −40.2 and −46 mV measured with perforated-patch recording at the high- and low-frequency locations, respectively. D, the half-activation voltage of the BK$_{Ca}$ channel ($V_{1/2}$) is plotted against 'best resonant voltage' for all buffering conditions. The straight line is the least squares fit, slope 0.7 and regression coefficient, $r = 0.85$. 

*Figure 6. Effects of intracellular calcium buffer on electrical tuning*

A, hair cell resonant frequencies plotted against concentration of BAPTA in electrode solution. The resonant frequency was obtained from the period of the oscillations at the onset of a current step that produced the sharpest tuning (the largest quality factor). The membrane potential at which tuning was sharpest is referred to as the 'best resonant voltage'. Each point is the mean ± 1 s.e.m of measurements at the low-frequency location (●) and at the high-frequency location (○). The number of values averaged is given beside each point, the same numbers apply also to the measurements in B and C. B, maximal quality factors of electrical tuning plotted against concentration of BAPTA in electrode solution, conventions as in A. Definition of quality factor is given in text. The small changes in $K_r$ and $Q$ with increasing BAPTA concentration were not statistically significant using a one-way ANOVA test at the 0.01 confidence level. C, the membrane potential at which the quality factor was maximal (best resonant voltage) plotted against internal concentration of BAPTA for cells at the low-frequency location (●) and at the high-frequency location (○). Construction lines are interpolations to derive the endogenous buffer from the 'best resonant voltage' of −40.2 and −46 mV measured with perforated-patch recording at the high- and low-frequency locations, respectively. D, the half-activation voltage of the BK$_{Ca}$ channel ($V_{1/2}$) is plotted against 'best resonant voltage' for all buffering conditions. The straight line is the least squares fit, slope 0.7 and regression coefficient, $r = 0.85$. 

*Figure 6. Effects of intracellular calcium buffer on electrical tuning*
repolarization (Fig. 5D). Both features are symptomatic of lack of clearance of Ca$^{2+}$ in the absence of an extrusion process, implying that the Ca$^{2+}$-ATPase is vital for maintaining intracellular gradients away from the channel cluster to prevent local buffer saturation and accumulation of Ca$^{2+}$. Slow components of the tail current were never seen in control recordings, either whole cell or perforated patch, suggesting that in those recordings, Ca$^{2+}$ extrusion via the Ca$^{2+}$-ATPase was fully operational.

**Effects of the calcium buffer on hair cell tuning**

Calium buffer effects on electrical tuning were studied in current-clamp conditions, where under-clamped voltage resonance could be evoked by injection of small current pulses (Fig. 1). Both the frequency of the voltage oscillations ($f_0$) and their time constant of decay ($\tau$) depend on membrane potential, (Crawford & Fettiplace, 1981). Therefore a hair cells resonant frequency was defined as that frequency at which the quality factor ($Q$) was maximal. $Q$ is given by $(\pi f_0 \tau)^2 + 0.25)^{1/2}$, where $\tau$ is the decay time constant of the oscillations (Crawford & Fettiplace, 1981). The membrane potential for maximal $Q$ referred to as the best resonant voltage, became systematically more depolarized with higher BAPTA concentrations. The best resonant voltage correlated with the $V_h$ for the BK$_{Ca}$ channel (Fig. 6D), which reflects the balance between the inward Ca$^{2+}$ current and the outward K$^+$ current needed to achieve optimal tuning. As with the $V_h$ values, there was an ~5 mV difference between the best resonant voltages in the low- and high-frequency cells.

With perforated-patch recordings, the best resonant voltages for the low-frequency and high-frequency locations were $-45.8 \pm 1.0$ mV ($f_0 = 97 \pm 27$ Hz; $n = 5$) and $-40.2 \pm 2.3$ mV ($f_0 = 265 \pm 14$ Hz; $n = 6$). Using values for the best resonant voltage at the two locations, it was also possible to obtain a second estimate of the endogenous calcium buffer (Fig. 6C). Expressed as an equivalent BAPTA concentration, the mobile buffer was 0.22 mÒ in the low-frequency cells and 0.45 mÒ in the high-frequency cells. The concentrations are similar to those deduced from half-activation voltages for the $I_{BKCa}$, supporting the notion of a cochlear gradient of endogenous calcium buffer. Owing to the correlation between the best resonant voltage and the $V_h$ for the BK$_{Ca}$ channel (Fig. 6D), the two methods for estimating endogenous buffer concentration are not independent.

Tuning was assessed during a period of several minutes in current clamp when the cell was depolarized to membrane potentials between −50 and −40 mV. Usually, in the absence of mechanotransduction, it was necessary to impose a holding current to bias the cell into the range where it was optimally tuned (Art & Fettiplace, 1987). On return to voltage clamp we observed a consistent increase in membrane capacitance that may reflect exocytosis of synaptic vesicles (Parsons et al. 1994). The cells initially had an average capacitance of approximately 12 pF, and responded to the period in current clamp with an increase of 2–3 pF. For those cells showing a capacitance increase, the magnitude of the change varied with cochlear location and was 1.9 ± 0.2 pF in 12 low-frequency cells and 2.9 ± 0.3 in 10 high-frequency cells. The likelihood of observing the capacitance increasing also depended on the nature of the exogenous calcium buffer. The fraction of cells showing an increase was 0.75 (0·1 BAPTA; $n_T = 8$), 0·56 (1·0 mM BAPTA; $n_T = 18$), 0·17 (10 mM BAPTA; $n_T = 12$) and 0·70 (10 mM EGTA; $n_T = 13$) where $n_T$ is the total number of cells in each group. Thus the capacitance change displayed a similar dependence on calcium buffer concentration to BK$_{Ca}$ channel activation. BAPTA at 10 mM was needed to attenuate it significantly, but 10 mM EGTA had little effect. In 11 perforated-patch recordings, no increase in capacitance was observed ($0.2 \pm 0.09$ pF), which may be due to concurrent re-uptake of exocytosed membrane, a property lost in whole cell recordings from hair cells (Parsons et al. 1994). Our results lack the temporal resolution to distinguish between different pools of exocytosed vesicles which may possess different dependencies on calcium buffer concentration. Nevertheless, they suggest a component of the transmitter-release apparatus experiences a Ca$^{2+}$ signal comparable to that activating the BK$_{Ca}$ channel.

**Ca$^{2+}$ sensitivity of the BK$_{Ca}$ channel**

Figure 4 demonstrated that the $I_{BKCa}$ activates at more depolarized potentials in high-frequency hair cells compared to low-frequency cells. One factor that might contribute to this disparity is a difference in the Ca$^{2+}$ sensitivity of the BK$_{Ca}$ channel. To test for this possibility, $I_{BKCa}$ were recorded in inside-out membrane macro patches from hair cells at two locations with fractional distances along the papilla of 0.34 ± 0.03 (low frequency) and 0.60 ± 0.01 (high frequency). Data were obtained on seven hair cells at each location with maximal patch currents of 0·12–0·68 nA (low frequency) and 0·2–1·4 nA (high frequency) at −50 mV. These currents correspond approximately to 10–100 BK$_{Ca}$ channel per patch. Each patch was exteriorized from the hair cell epithelium so that its intracellular face could be exposed to a range of Ca$^{2+}$ concentrations between 1 and 1000 mÒ, and $I_{BKCa}$ were evoked with depolarizing voltage steps from −50 to +50 mV. Figure 7 illustrates the major differences observed in BK$_{Ca}$ channels at the two positions. These were most conspicuous at −50 mV, where the high-frequency channels were less Ca$^{2+}$ sensitive and deactivated more rapidly. For each patch, the current at −50 mV, normalized to its maximum value, was plotted against Ca$^{2+}$ concentration (Fig. 7C). Fits to the Hill equation gave a half-saturating Ca$^{2+}$ concentration (Ca$_{50}$) of 5.8 ± 0.6 mÒ (low-frequency position) and 10·3 ± 0.6 mÒ (high-frequency position) with mean Hill coefficients of 2·9 and 1·9 at the low- and high-frequency positions, respectively. The deactivation of the currents at −50 mV in 2·5 mÒ Ca$^{2+}$ could be fitted with a single exponential decay with a time constant of 2.4 ± 0.4 ms in low-frequency and 0.74 ± 0.05 ms in high-frequency cells.
Further evidence supporting location-dependent variations in Ca$^{2+}$ sensitivity of the BK$_{Ca}$ channel was obtained from voltage-activation curves in a fixed Ca$^{2+}$ concentration. For one patch at each position, it was possible to obtain a complete activation curve in 5 $\mu$M Ca$^{2+}$ (Fig. 7D), from which a half-activation voltage of $-68$ mV was inferred for the low-frequency patch and $-26$ mV for the high-frequency patch. Owing to the combined Ca$^{2+}$ and voltage dependence of the channels, the need for greater depolarization to activate the high-frequency channels is consistent with them being less Ca$^{2+}$ sensitive. The reported Ca$^{2+}$ sensitivities and deactivation time constants are both within the range of values previously reported for single BK$_{Ca}$ channels in turtle isolated hair cells (Art et al. 1995). However, in the earlier measurements it was not possible to demonstrate a correlation between the two parameters. Such a correlation would fit with the notion that hair cells at different locations express distinct variants of the BK$_{Ca}$ channel (Jones et al. 1999). Indeed, the alternatively spliced variants cloned from turtle hair cells possess the property that those with faster kinetics are less Ca$^{2+}$ sensitive, which accords with the present results on the native channels.

**Hotspots of Ca$^{2+}$ influx**

In isolated turtle hair cells, Ca$^{2+}$ influx via voltage-gated channels occurs over small regions or 'hotspots' confined to the basal half of the cell (Tucker & Fetipplace, 1995). The ability to record from hair cells at specific papillary locations allowed us to examine whether the structure of the hotspots varied with location to reflect the difference in maximum Ca$^{2+}$ current. Regions of Ca$^{2+}$ elevation were defined using the fluorescent dye Calcium Green-1. Following attachment of the whole-cell electrode, the cell was drawn onto its side in order to optimize spatial resolution in the confocal images. Figure 8A–C shows single images of a high-frequency hair cell captured before and during a 200 ms depolarization to

---

**Figure 7. BK$_{Ca}$ channel properties in inside out patches**

A, average currents recorded in an inside-out membrane patch detached from a low-frequency hair cell ($d = 0.3$) for voltage steps from $-50$ to $+50$ mV in the presence of Ca$^{2+}$ concentrations from 2.2 to 470 $\mu$M. Dashed line denotes zero current level. B, currents recorded in an inside-out patch from a high-frequency hair cell ($d = 0.6$) for voltage steps from $-50$ to $+50$ mV in Ca$^{2+}$ concentrations from 2.2 to 1000 $\mu$M. Each trace in A and B is the average of between 100 and 250 responses. C, $I_{BK,Ca}(I)$ scaled to its maximum value ($I_{\text{max}}$) is plotted against Ca$^{2+}$ concentration in low-frequency ($\bullet$) and high-frequency (O) hair cells. Each point is the mean ± 1 s.e.m. of measurement on seven inside-out patches; for most of the points the s.e.m. is less than the symbol size. Smooth curves are fits to the Hill equation, $I/\text{max} = 1/(1 + (\text{Ca}_{\text{eq}}/\text{Ca})^n)$, with Ca$_{\text{eq}}$ and Hill coefficient ($n$), respectively, of 5.8 $\mu$M and 2.9 ($\bullet$) and 10.3 $\mu$M and 1.9 (O). D, BK$_{Ca}$ channel activation curves for channels in a low-frequency hair cell ($d = 0.33$; $\bullet$) and a high-frequency hair cell ($d = 0.61$; O), derived from tail-current measurement ($I$) from a holding potential of $-80$ mV. Smooth curves are fits to eqn (1) with values of $V_b$ and $V_a$, respectively, of $-68$ and 27 mV (low frequency) and $-26$ and 24 mV (high frequency).
−20 mV designed to maximally activate the Ca\(^{2+}\) current. The increase in intracellular Ca\(^{2+}\) was distributed over a ring around the nucleus, but there was evidence of punctate regions or ‘hotspots’ of fluorescence.

The lack of temporal resolution imposed by the frame rate hinders an accurate determination of the number of hotspots. To approach this problem, the time course of the fluorescence changes was characterized by constructing around the hotspots contours to correspond to a fixed Ca\(^{2+}\) level (Tucker & Fettiplace, 1995, 1996). Contours were initially drawn starting with the areas of bright fluorescence evident in the first image, and the fluorescence change mapped over several seconds. For example, at least seven regions are discernable in Fig. 8B, and their associated time courses are given in Fig. 8D. Two pieces of evidence were used to support the enumeration of hotspots. Firstly, all hotspots should demonstrate a similar time course. If the contour was not appropriately centred, or if it covered two spots, the fluorescence change would rise more slowly or with a delay. Secondly, the size of the maximal fluorescence change should be comparable for all spots. In some cells, where the contours were initially incorrectly contrived, the magnitude of the peak fluorescence exhibited ‘quantization’, so that some spots had two or three times larger peaks than the average, suggesting that they encompassed multiple sites of Ca\(^{2+}\) entry. Applying these procedures showed that each high frequency hair cell possessed between five and eight hotspots. As previously reported, the maximum number of spots was visible in a central section through the cell, and few extra spots appeared de novo on focusing

![Figure 8](image.png)

**Figure 8. Hotspots of Ca\(^{2+}\) influx in a high-frequency hair cell**

Confocal images of a hair cell filled with Calcium Green-1 before (A) and during (B and C) a 200 ms depolarizing current step to −20 mV. Times at which the images were captured are shown as arrows above the Ca\(^{2+}\) current (inset). Pseudocolor scale on left corresponds to pixel intensities from 0 to 255. D, time course of the fluorescence changes in seven regions in response to the depolarization. Different traces correspond to the regions of the same color shown in the schematic hair cell on the left. Orange region may contain multiple hotspots, but was judged to be a single spot on the criteria given in the text.
through the cell, even though such focusing sometimes improved the sharpness of a given spot.

Measurements on cells at the two papilla locations demonstrated that the mean number of Ca\(^{2+}\) hotspots in high-frequency hair cells was 1.8 times that in low-frequency cells (Fig. 9B). As an alternative method of comparing cells, the total area occupied by the hotspots was calculated from the sum of the areas encompassed by the contours. These areas were normalized to the total cross-sectional area of the cell, which was similar for the two positions (154 ± 16 \(\mu m^2\) in four low-frequency cells and 151 ± 14 \(\mu m^2\) in six high-frequency cells). This method of analysis avoided the difficulty in hotspot counting of distinguishing between closely spaced spots, but confirmed that twice the cell area was occupied by hotspots in high-frequency compared to low-frequency cells (Fig. 9C). The fluorescence hotspots require Ca\(^{2+}\) entry through voltage-dependent channels (Tucker & Fettiplace, 1995). The imaging results therefore suggest that Ca\(^{2+}\) channels are distributed over twice the membrane area in high-frequency cells compared to low-frequency cells. Since cells at the high-frequency location possess 2.4 times the number of Ca\(^{2+}\) channels, these results are consistent with Ca\(^{2+}\) channels being present at a constant density irrespective of location.

**DISCUSSION**

The endogenous buffer

By recording in the intact papilla, we have been able to compare the properties of hair cells at two specific locations tuned to approximately 100 and 300 Hz. We have provided evidence about Ca\(^{2+}\) entry, buffering and action at one target, the large-conductance Ca\(^{2+}\)-activated K\(^+\) channel BK\(_{Ca}\), in the soma of auditory hair cells. All aspects of this pathway were found to vary with the cochlear location of the hair cell, and hence the frequency to which it was tuned. The concentration of mobile endogenous calcium buffer in the hair cell soma was estimated as 0.21 mM BAPTA at the low-frequency location, and 0.46 mM BAPTA at the high-frequency location (means of the values in Fig. 4 and 6). These values are in good agreement with estimates of buffer in the hair bundle of 0.1 mM BAPTA at the same low-frequency position and 0.4 mM BAPTA at the high-frequency location (Ricci et al. 1998). Taken together the results suggest that the calcium buffer has a uniform concentration throughout the cell, and that this concentration increases along the cochlea's tonotopic axis.

The variation in buffer concentration is consistent with the gradient in the expression of calbindin-28k, a likely candidate for the endogenous buffer (Navaratnam et al. 1998).
found in the chick cochlea. A larger value of 0.9 mM BAPTA for the endogenous buffer was previously measured in isolated turtle hair cells using the small conductance Ca\(^{2+}\)-activated K\(^{+}\) (SK) channel as the Ca\(^{2+}\) sensor (Tucker & Fettiplace, 1999). The cochlear origin of those cells was unknown, but their properties, including size of Ca\(^{2+}\) current, suggest they were tuned to high frequencies. Roberts (1993) estimated the native calcium buffer in isolated frog scala cea hair cells to be in excess of 1 mM BAPTA, though he also did not distinguish the epithelial location or frequency specificity of the cells.

### Ca\(^{2+}\) and BK\(_{Ca}\) channel interactions

Both the insensitivity of the BK\(_{Ca}\) channel activation to BAPTA (Fig. 4) and the lack of accumulation of Ca\(^{2+}\) near the BK\(_{Ca}\) channel (Fig. 5) argue for the BK\(_{Ca}\) channels being in close proximity to the Ca\(^{2+}\) channels. At least 10 mM BAPTA was required to alter significantly the activation range of the BK\(_{Ca}\) channel. Since the voltage-activation curve for the channel was independent of pulse duration, the BAPTA effect must approximate the steady-state condition. Naraghi & Neher (1997) have estimated that, for a steady-state Ca\(^{2+}\) gradient, 2 mM BAPTA would have a space constant of 28 nm, indicative of its buffering range near a Ca\(^{2+}\) source. For 10 mM BAPTA, the space constant would be roughly halved. Thus it is likely that each Ca\(^{2+}\) channel influences only its immediate neighbouring BK\(_{Ca}\) channels. Since the Ca\(^{2+}\) and BK\(_{Ca}\) channels maintain a constant stoichiometry in cells tuned across the spectrum, (Art et al. 1993), it is conceivable that the two channel types are assembled into a single complex in the membrane. The proximity of the Ca\(^{2+}\) channel to its target ensures that the rate of BK\(_{Ca}\) channel activation is limited solely by intrinsic channel kinetics and not by Ca\(^{2+}\) diffusion.

There was a difference of 4 mV between the half-activation voltage (\(V_{1/2}\)) for the BK\(_{Ca}\) channels at the low- and high-frequency locations and at least part of this difference may be due to the lower Ca\(^{2+}\) sensitivity of BK\(_{Ca}\) channels at the high-frequency location (Fig. 7). Assuming that the Ca\(^{2+}\) current activation increases e-fold in 6-7 mV (Art & Fettiplace, 1987), the fraction of the total Ca\(^{2+}\) current required to half-activate the BK\(_{Ca}\) channels in high-frequency cells is 1.8 (\(= \exp(4/6.7)\)) that in low-frequency cells. This is close to the ratio of the Ca\(^{2+}\) sensitivities of the BK\(_{Ca}\) channels in detached patches, which was 1.78 (\(= 10^{-3}/5-8\)). After correction for the 4 mV difference, the \(V_{1/2}\) values at the two locations (Fig. 4C) possess an identical dependence on BAPTA concentration. Such a result would be expected if each BK\(_{Ca}\) channel is influenced by the same number of Ca\(^{2+}\) channels in cells at both cochlear locations.

### Structure of the Ca\(^{2+}\) microdomains

Hair cells tuned to the higher frequency had approximately twice the number of Ca\(^{2+}\) channels and also twice the number of Ca\(^{2+}\) entry zones or ‘hotspots’. The number of hotspots is less than the number of sites of transmitter release in turtle hair cells (at least 17, Sneary, 1988). It is possible that the number of ‘hotspots’ was underestimated due to the limited temporal resolution of the imaging experiments causing neighbouring Ca\(^{2+}\) microdomains to fuse. However, the area of the hotspots also differed by the same 2-fold ratio between the two locations. This suggests that the Ca\(^{2+}\) channels that cluster to form the hotspots are present at a constant density in both high-frequency and low-frequency cells. An increase in the number of channels per cell is then accomplished by addition of clusters at constant channel density.

If the Ca\(^{2+}\) and BK\(_{Ca}\) channels are aggregated at synaptic release sites (Roberts et al. 1990; Isa & Hudspeth, 1994), changes in intracellular Ca\(^{2+}\) by influx through Ca\(^{2+}\) channels will regulate exocytosis as well as BK\(_{Ca}\) channel activation. Similar elevations in Ca\(^{2+}\) may be required to control the two processes, both of which must be fast and continuously graded from the resting potential near −50 mV, to −20 mV. The need for multiple clusters of Ca\(^{2+}\) channels is most likely linked to their role in exocytosis. The maximum size of each cluster may then be constrained by the area of membrane adjacent to the synaptic body, allowing the release site to be rapidly replenished with vesicles. The synaptic body in frog hair cells has a mean diameter of about 0.5 μm (Lunzi et al. 1999). An increase in the hair cell’s complement of Ca\(^{2+}\) channels may therefore serve a dual role in signalling. In conjunction with changes in the BK\(_{Ca}\) channels, it will augment the electrical resonant frequency. The increased number of release sites may also enhance the temporal fidelity of synaptic transmission by allowing the release sites to be used asynchronously, one site being refilled while another discharges.

---


Acknowledgements
The work was supported by NIH grants RO1 DC01362 to R.F. and RO1-DC03896 to A.J.R. and a Deafness Research Foundation grant to A.J.R.

Corresponding author
R. Fettiplace: 185 Medical Sciences Building, 1300 University Avenue, Madison, WI 53706, USA.

Email: fettiplace@physiology.wisc.edu

Author’s present address
A. J. Ricci: Neuroscience Center for Excellence, Louisiana State University Medical Center, 2020 Gravier Street, New Orleans, LA 70112, USA.
Tonotopic variations of calcium signalling in turtle auditory hair cells
A. J. Ricci, M. Gray-Keller and R. Fettiplace

J. Physiol. 2000;524;423-436

This information is current as of February 21, 2008