Prestin kinetics and corresponding frequency dependence augment during early development of the outer hair cell within the mouse organ of Corti

Jun-Ping Bai, Dhasakumar Navaratnam & Joseph Santos-Sacchi

Several studies have documented the early development of OHC electromechanical behavior. The mechanical response (electromotility, eM) and its electrical correlate (nonlinear capacitance, NLC), resulting from prestin’s voltage-sensor charge movement, increase over the course of several postnatal days in altricial animals. They increase until about p18, near the time of peripheral auditory maturity. The correspondence of auditory capabilities and prestin function indicates that mature activity of prestin occurs at this time. One of the major requirements of eM is its responsiveness across auditory frequencies. Here we evaluate the frequency response of prestin charge movement in mice over the course of development up to 8 months. We find that in apical turn OHCs prestin’s frequency response increases during postnatal development and stabilizes when mature hearing is established. The low frequency component of NLC, within in situ explants, agrees with previously reported results on isolated cells. If prestin activity is independent of cochlear place, as might be expected, then these observations suggest that prestin activity somehow influences cochlear amplification at high frequencies in spite of its low pass behavior.

The outer hair cell (OHC) enhances cochlear sensitivity up to 3 orders of magnitude, and the cellular basis of this feat resides within the OHCs that populate the organ of Corti. OHCs are electro-motile, responding to transmembrane current perturbations with changes in cell length, termed electromotility (eM)1,2. Though the pertinent electromechanical characteristics (e.g., voltage-dependence, and magnitude) of OHC eM had been studied since the mid-80s3,4, the identification of the membrane protein, SLC26a5 (prestin), driving this response required over a decade5. Various candidates were considered along the way, similar to the quest for the stereociliar MET channel6.

Prestin drives OHC electromotility (eM), known to be responsible for cochlear amplification (CA) in mammals7. The electrical signature of eM is a bell-shaped nonlinear capacitance (NLC), the first derivative of prestin sensor charge vs. membrane voltage (dQp/dVm), which peaks at a characteristic membrane voltage (Vh)8,9. We have previously studied the development of NLC in OHCs of the mouse10. Those studies demonstrated that increases in prestin charge (Qmax) continued after stabilization of linear capacitance (at p10), which corresponds to total membrane surface area (sum of membrane and embedded prestin surface area). Thus, though the number of prestin molecules appeared to stabilize, additional changes in NLC characteristics ensued, indicating some sort of maturational events. Here we study, in developing OHCs, the maturation of prestin kinetics, which has recently been shown to possess low pass characteristics in adult guinea pig OHCs11. We find that during development the frequency response of NLC increases, stabilizing near 6kHz in adults at the apical turn of the cochlea at room temperature. We also find that linear capacitance decreases with aging, and is indicative of reduced prestin activity.

1Department of Surgery (Otolaryngology), Yale University School of Medicine, 333 Cedar St, New Haven CT, USA.
2Department of Cellular and Molecular Physiology, Yale University School of Medicine, 333 Cedar St, New Haven CT, USA.
3Department of Neuroscience, Yale University School of Medicine, 333 Cedar St, New Haven CT, USA.
4Department of Neurology, Yale University School of Medicine, 333 Cedar St, New Haven CT, USA. *email: joseph.santos-sacchi@yale.edu
insertion in the membrane. Nevertheless, specific motor charge density (a metric for prestin density within the membrane) remains fairly constant.

Methods
C57/B6 pups aged postnatal days 6–18, and months 2 and 8 were used. All experimental protocols were approved by the Yale Animal Care and Use Committee, and were in accordance with relevant guidelines and regulations. Apical turns of the organ of Corti were dissected out and recorded with ionic current blocking solutions, thereby removing interference on measures of membrane capacitance. The extracellular solution contained (in mM): 100 NaCl, 20 tetraethylammonium (TEA)-Cl, 20 CsCl, 2 CoCl2, 1 MgCl2, 1 CaCl2, 10 HEPES, pH 7.2. The intracellular solution contained (in mM): 140 CsCl, 2 MgCl2, 10 HEPES, and 10 EGTA, pH 7.2. Pipettes were coated with M-coat to reduce stray capacitance, and had resistances of 3-5 MΩ. Gigohm seals were made and stray capacitance was balanced out with amplifier circuitry prior to establishing whole-cell conditions. A Nikon Eclipse E600-FN microscope with 40× water immersion lens was used to observe cells during voltage clamp. Whole cell voltage clamp recordings were performed from all rows of outer hair cells (OHCs) of whole mount organ of Corti. We limited the recording area within 1/4 turn. OHCs were recorded at room temperature using jClamp software and an Axopatch 200B amplifier. Data were low pass filtered at 10 kHz and digitized at 100 kHz with a Digidata 1320 A.

Chirp voltage stimuli were delivered across frequency and analyzed within 300–7000 Hz, where stray capacitance was removed. The voltage chirps were generated in jClamp using the Matlab logarithmic “chirp” function (10 mV pk; pts = 4096; F0 = 24.4141 Hz; F1 = 50 kHz; t1 = 0.04095 s). Chirp responses were not averaged at each voltage step (−160 to +120 in 40 mV steps). Cell currents were either averaged for analysis (6–8 cells/group) for surface plot presentations or analyzed individually for statistical measures. Detailed analysis of Cm was performed in Matlab. Capacitance was measured using dual-sine analysis at harmonic frequencies (Santos-Sacchi et al., 1998; Santos-Sacchi, 2004). Briefly, real and imaginary components of membrane current at harmonic frequencies were determined by FFT in jClamp, corrected for the roll-off of recording system admittance and residual stray capacitance. Rs, Rm and Cm were extracted using the dual-sine, 3-parameter solution of the standard patch clamp model, based on the original single sine solution. In order to extract Boltzmann parameters, capacitance-voltage data were fit to the first derivative of a two-state Boltzmann function.

\[
C_m = NLC + C_{\text{lin}} + C_{\text{lin}} = \frac{ze}{k_BT} \left( \frac{b}{1 + b^2} \right) + C_{\text{lin}} + C_{\text{lin}}
\]
\[ b = \exp \left( -ze \frac{V_m - V_h}{k_B T} \right), \quad C_{sa} = \frac{\Delta C_{sa}}{(1 + b^{-1})} \]

\( Q_{\text{max}} \) is the maximum nonlinear charge moved, \( V_h \) is voltage at peak capacitance or equivalently, at half-maximum charge transfer, \( V_m \) is \( R_s \)-corrected membrane potential, \( z \) is valence, \( C_{\text{lin}} \) is linear membrane capacitance, \( e \) is electron charge, \( k_B \) is Boltzmann's constant, and \( T \) is absolute temperature. \( C_{sa} \) is a component of capacitance that characterizes sigmoidal changes in specific membrane capacitance\(^{17} \). \( \Delta C_{sa} \) is the total sum of unitary changes per prestin motor protein. \( Q_{\text{sa}} \) denotes charge density, namely \( Q_{\text{max}}/C_{\text{lin}} \).

A Lorentzian function (or the sum of two Lorentzians) (Gale and Ashmore, 1997; Santos-Sacchi and Tan, 2018) was used to fit NLC across frequency.

\[ NLC(f) = NLC_0/\left[1 + \left(2\pi f\tau c\right)^2\right]^{1/2} \]

where \( NLC_0 \) is the zero frequency component, and \( \tau = 1/F_c \), the cut-off frequency. Additionally, a power fit of NLC across frequency was performed.

\[ NLC(f) = NLC_0 + a \cdot f^b \]

where \( NLC_0 \) is the zero frequency component, and \( a \) and \( b \) control the frequency response.

**Results**

Figure 1 illustrates our methodology to measure NLC across frequency. With our Boltzmann fit (Eq. 1) we obtain the usual Boltzmann parameters while extracting NLC, and additionally determine \( C_{sa} \), a component of capacitance that we believe arises from changes in membrane surface area or membrane thickness as prestin alters its conformational state across voltage\(^{17-20} \).

Figure 2 shows mean (+/− se) NLC at 3 selected frequencies within our measurement bandwidth, as a function of postnatal age. As determined previously, NLC reaches maximal values at p17/18 in rodents\(^{10,21} \). Here we find the same pattern across frequencies. \( C_{\text{lin}} \) increases until p12/13 as a result of surface area increase as the cells mature, lengthen and express increasing amounts of prestin within the membrane. However, \( C_{\text{lin}} \) decreases...
thereafter at all frequencies. By 8 months, C_{lin} has decreased by more than 25%. NLC also reduces following p17/18. The plots also highlight a substantial roll-off of peak NLC across frequency, which we investigate in more detail further below.

Because we limited our recordings to a quarter turn of the apical coil of the cochlea, we determined whether the variability in hair cell length within that region could impact our C_{lin} data. OHC lengths in different rows of the cochlear apex are indistinguishable. In the mouse strain we used, they determined that OHC lengths from middle (17.3 μm) and apical (21.3 μm) turn regions differed by 4 μm. Assuming that this progression will translate into our quarter turn (as their plots suggest) we would maximally expect about a 1 μm length variability in our hair cell recordings. Given the cylindrical geometry of the OHC and its diameter in mice (~7 μm), we calculate an expected maximal variability in linear capacitance (based on surface area arising from our measured membrane specific capacitance of 0.08 pF/μm^2) of 0.26 pF. This is well below our finding of ~1.6 pF (25% reduction from p12/13) during aging. Utilizing a more stringent estimate of 0.05 pF/μm^2 for specific membrane capacitance, giving 0.165 pF variability, our results are even less likely to be influenced by sampling within the quarter turn. We conclude that the C_{lin} changes during aging are real.

In Fig. 3A, the left panels show full NLC spectrum (0.39–7 kHz) surface plots as a function of postnatal age, illustrating the bell-shaped NLC and its roll-off with increasing frequency. Black dots indicate means and pink dots are +se, which sit atop the Boltzmann fits. The panels on the right show the same data top-down, and provide a clearer view of the frequency response changes during aging. (B) NLC_{Vh} for each age group is log-log plotted to more clearly show the low-pass behavior that changes during aging. The colored points were used for fitting in Fig. 5. For the 2 youngest groups grey lines show regions omitted from fits because of noise at high frequencies.

Figure 3. (A) Left panels show full NLC spectrum (0.39–7 kHz) surface plots as a function of postnatal age, illustrating the bell-shaped NLC and its roll-off with increasing frequency. Black dots indicate means and pink dots are +se, which sit atop the Boltzmann fits. The panels on the right show the same data top-down, and provide a clearer view of the frequency response changes during aging. (B) NLC_{Vh} for each age group is log-log plotted to more clearly show the low-pass behavior that changes during aging. The colored points were used for fitting in Fig. 5. For the 2 youngest groups grey lines show regions omitted from fits because of noise at high frequencies.
Quantification of $Q_{\text{max}}$ and $C_{\text{lin}}$ confirms their initial rise between p6/7 to p12/13 as the cells increase in surface area, and mirror measures made in isolated cells. The development of $Q_{\text{sp}}$ in the low frequency region also corresponds to measures from isolated cells. After p17/18, $Q_{\text{max}}$ decreases, while $Q_{\text{sp}}$ appears to stabilize, due to simultaneous decrease in $C_{\text{lin}}$. The additional frequency information provided by chirp analysis shows that both $Q_{\text{max}}$ and $Q_{\text{sp}}$ following p17/18 depend on frequency, with higher frequencies showing reduced values. Interestingly, $V_h$ within the explant differs from isolated cells during early development, and may result from forces that normally exist in the organ of Corti, e.g., membrane tension. It is well known that many factors can influence $V_h$, including membrane tension and intracellular $\text{Cl}^-$,

$z$, which characterizes the voltage sensitivity of prestin or the distance sensor charge is moved within the membrane field, is variable in p6/7 animals, but clearly increases after this time period over age. Some frequency dependence of $z$ in the adult is evident, and reduction in $z$ at high frequencies in the adult guinea pig was originally noted by Gale and Ashmore. Finally, in Fig. 5 we explore the changes in frequency cut-off (Fc) of NLC at $V_h$. Since the reduction of $NLC_{Vh}$ across frequency is clearly not a simple process, three methods of fitting were evaluated to estimate frequency roll-off (Fig. 5A). We provide fits to average data. Examples of the 3 fit types for p17/18 OHCs are shown, including single Lorentzian, dual Lorentzian (sum of 2 Lorentzians) and power fits. The blue line depicts the fit and red lines show 95% confidence predictions of the fits (fitting performed in Sigmaplot). The poorest fit is with a single Lorentzian, followed by dual Lorentzian and power fits. In regard to the power fit, we have previously found evidence of multi/stretched exponential behavior in NLC. The stretched exponential (known as the Kohlrausch-Williams-Watts [KWW] relaxation function), first devised by Kohlrausch (1854) to describe the behavior of capacitor discharge in a Leyden jar, is also applicable to the viscoelastic behavior of glasses, and other systems where time-dependent behavior results from overcoming a multitude of energy barriers. Interestingly, cytoskeletal dynamics is one such system. The KWW function has a related counterpart in the frequency domain, termed the Havriliak-Negami [HN] relaxation function. Fitting our data with that counterpart, we obtain similar quality fits to the power functions (for example, at p17,18 the fit $R^2$ is 0.9975 and 0.9974 for power and HN). Since the power function fit has fewer parameters (one less) than HN, we have chosen to

Figure 4. Boltzmann characteristics (mean +/- se) for selected bandwidths. The dashed line denotes NLC measured at low frequencies from isolated OHCs (Abe et al., 2010). Note abrupt decline of $C_{\text{lin}}$, while $C_{\text{sa}}$ remains steady. $V_h$ is initially at a more positive potential than isolated cells. $Q_{\text{max}}$ (pC) and $Q_{\text{sp}}$ (pC/pF) increase rapidly during development. $Q_{\text{max}}$ decreases after p17/18, with a clear difference between low and high frequencies, but $Q_{\text{sp}}$ stabilizes because of the $C_{\text{lin}}$ drop. Following some variability at p6/7, $z$ increases with age, and also tends to be smaller at higher frequencies. P6,7 n = 7; p8,9 n = 7; p12,13 n = 6; p17,18 n = 8; 2 month n = 8; 8 month n = 9.
illustrate power fits. The usage of the term *low-pass* for the power function should not be construed in a strict engineering sense, but here denotes domination of low pass components whose slope changes over maturation. Figure 5B shows that regardless of the utilized fitting function, each provides evidence for increases in frequency responsiveness during development.
since they are above the sampled frequency. In Fig. 5C, we provide another metric of frequency response roll-off by finding the –3dB magnitude of $NLC_{\text{ref}}$ relative to 350 Hz. During the life span, $F_c$ increases and appears to stabilize near 6 kHz in our older animals. Interestingly, the cut-offs roughly correspond to those of the single Lorentzian fits. We are cognizant that these cut-off estimates cannot be considered true cut-off frequencies, but only indicate that the $NLC_{\text{response}}$ has lost much of its steady state magnitude even at low frequencies. That is, the $F_c$s simply reflect the relative roll-off among aging groups, and their absolute values will differ depending on the reference frequency. Thus, all methods to quantify roll-off indicate that frequency responsiveness increases with age. Possible reasons for this are discussed below.

Discussion

Studying the development of OHC function can reveal important determinants of adult auditory function. A variety of OHC characteristics mature during the course of post-natal development in altricial animals, including stereociliar transduction and eM. By using such animals, these characteristics can be conveniently studied in isolated cells or explants following birth. The development of eM and its corresponding electrical correlate, nonlinear membrane capacitance (NLC), have been found to reach maturity near p18 in the rodent10,21,37, close to the onset of hearing. These studies did not include an assessment of prestin frequency dependence during development, a likely important factor for OHC performance in the adult animal’s acoustic environment. Here we evaluated the development of NLC frequency response in the in situ mouse cochlea, and find that its frequency cut-off ($F_c$) increases from p6 to p18, thereafter stabilizing out to 8 months at about 6 kHz at room temperature. Assuming that the frequency response is solely limited by the kinetics of prestin (however, see below) and given a $Q_{\text{m}}$ of 23, the $F_c$ would approach 15 kHz. The data indicate that the $F_c$ is lower pass in nature than the full auditory capabilities of the mouse, since prestin’s intrinsic kinetics is likely place independent. That is, prestin’s molecular structure, molecular interactions, and especially molecular behavior have yet to show significant differences among OHCs from various turns of the cochlea24,40.

Importantly, most developmental prestin characteristics measurable within the in situ organ of Corti correspond to those measures obtained in isolated cells10,21, i.e., at low measurement frequencies. However, we find that $V_h$ in young animals differs from isolated cell studies, and this may be due, for example, to changes in turgor pressure or membrane tension upon isolation. It is notable that for a simple two-state voltage-dependent protein, $V_h$ is determined by the ratio of forward and backward transition rates of conformational change. Thus, $V_h$ alterations may inform on changes in prestin kinetics. Considering the differences of $V_h$ in situ vs. in isolated cells, it may be desirable to analyze the effects of prestin mutations, e.g. the “499” knock-in mutation in the explant, rather than isolated cells.

We find that $C_{\text{m}}$ decreases following the attainment of mature NLC. The decrease corresponds to a decrease in $Q_{\text{m, max}}$, indicating that the absolute number of prestin motors in OHCs decreases with age. This result implies that the surface area of the cells decrease because prestin occupies less surface area, and this is expected since in the prestin KO OHC surface area/length is markedly reduced41. The reduction of surface area and prestin in OHCs may have influential effects on the mechanical properties of the cochlear partition. Indeed, in the prestin KO mouse, Mellado Lagarde et al. have found that frequency tuning of the partition changes42. Thus, we might expect altered hearing in the aging mouse based on the reduction of prestin content that we find.

Why does the frequency response of prestin change during aging? Several possibilities exist. Prestin function itself could change during maturation due to changes in 1) phosphorylation or glycosylation35,36; 2) multimerization35–37; or 3) molecular crowding within the lateral membrane. Extrinsic influences may also change; for example, during the development of the spaces of Nuel supporting cell rearrangements may occur, and less cell-cell contact could reduce interactive mechanical constraints. Certainly, such types of interactions can affect auditory performance48. Finally, cytoskeletal influences on prestin39,40 may change during development. We recently determined that MAP1S, a small microtubule binding protein interacts with both actin and prestin, and co-expression in heterologous cells promotes targeting and alteration in prestin characteristics51. Cytoskeletal forces acting on prestin may work to influence conformational switching by the well-known piezolectric nature of the protein30–32,34–36, which has been extensively modelled32–34. Thus, there may well be mechanical filtering of prestin performance depending on whole-cell mechanics that simultaneously impacts the $F_c$ of both NLC and eM, as the two are well coupled35,36. In any case, whatever mechanisms that enhance prestin’s frequency response during the developmental likely limits its frequency response in adults. Interestingly, quite low pass estimates ($F_c$ ~1.5 kHz) of adult OHC eM frequency response have been found in the high frequency region of the living gerbil using optical coherence microscopy (OCT) vibrometry37. Thus, at this point, the mechanism whereby OHC eM influences very high frequency hearing is under question.

References

Received: 30 May 2019; Accepted: 25 October 2019;
Published online: 11 November 2019

1. Kachar, B., Brownell, W. E., Altschuler, R. & Fex, J. Electrokinetic shape changes of cochlear outer hair cells. Nature 322, 365–368 (1986).
2. Brownell, W. E., Bader, C. R., Bertrand, D. & de Ribaupierre, Y. Evoked mechanical responses of isolated cochlear outer hair cells. Science 227, 194–196 (1985).
3. Ashmore, J. F. A fast motile response in guinea-pig outer hair cells: the cellular basis of the cochlear amplifier. J. Physiol. 388, 323–347 (1987).
4. Santos-Sacchi, J. & Dilger, J. P. Whole cell currents and mechanical responses of isolated outer hair cells. Hear. Res. 35, 143–150 (1988).
5. Zheng, J. et al. Prestin is the motor protein of cochlear outer hair cells. Nature 405, 149–155 (2000).
6. Corey, D. P. What is the hair cell transduction channel? J. Physiol. 576, 23–28 (2006).
www.nature.com/scientificreports/ 7. Dallos, P. et al. Prestin-based outer hair cell motility is necessary for mammalian cochlear amplification. *Neuron* 58, 333–339 (2008).
8. Ashmore, J. F. Forward transduction in the mammalian cochlea. *Neurosci. Res. Suppl.* 12, S39–S50 (1990).
9. Santos-Sacchi, J. Reversible inhibition of voltage-dependent outer hair cell motility and capacitance. *J. Neurosci.* 11, 3096–3110 (1991).

Abe, T. et al. Developmental expression of the outer hair cell motor protein in the mouse. *J. Membr. Biol.* 215, 49–56 (2007).
11. Santos-Sacchi, J. & Song, L. Chloride anions regulate kinetics but not voltage-sensor Qmax of the solute carrier SLC26a5. *Biophys. J.* 110, 1–11 (2016).
12. Gillis, K. D. In Single Channel Recording (eds Sakmann, B. & Neher, E.) Ch. 7, 155–198 (Plenum Press, 1995).
13. Santos-Sacchi, J. High frequency measures of OHC nonlinear capacitance (NLC) and their significance: Why measures stray away from predictions. *AIP Conference Proceedings* 2015, 060004-060001–060004-060005 (2018).
14. Santos-Sacchi, J., Kekeha, S. & Takahashi, S. Effects of membrane potential on the voltage dependence of motility-related charge in outer hair cells of the guinea-pig. *J. Physiol.* 510(Pt 1), 225–233 (1998).
15. Santos-Sacchi, J. Determination of cell capacitance using the exact empirical solution of dV/dCm and its phase angle. *Biophys. J.* 87, 714–727 (2004).
16. Busch, M. & Neher, E. Rates of diffusional exchange between small cells and a measuring patch pipette. *Pflugers Arch.* 411, 204–211 (1988).
17. Santos-Sacchi, J. & Navarrete, E. Voltage-dependent changes in specific membrane capacitance caused by prestin, the outer hair cell lateral membrane motor. *Pflugers Arch.* 411, 99–106 (2002).
18. Chessum, L. et al. Helios is a key transcriptional regulator of outer hair cell maturation. *Nature* 563, 696–700 (2018).
19. Duret, G., Pereira, F. A. & Raphael, R. M. Diffusional inhibits prestin by chloride-dependent mechanism. *PloS One* 12, e0183046 (2017).
20. Harasztosi, C. & Gummer, A. W. The chloride-channel blocker 9-anthracenecarboxylic acid reduces the nonlinear capacitance of prestin-associated charge movement. *Eur. J. Neurosci.* 43, 1062–1074 (2016).
21. Oliver, D. & Falker, B. Expression density and functional characteristics of the outer hair cell motor protein are regulated during postnatal development in rat. [In Process Citation]. *J. Physiol. (Lond)* 519(Pt 3), 791–800 (1999).
22. Keilen, S. & Richter, C. P. Cochlear dimensions obtained in hemicochlea of four different strains of mice: CBA/Ca, 129/CD1, 129/SvEv and C57BL/6J. *Hear. Res.* 162, 91–104 (2001).
23. Solsona, C., Innocenti, B. & Fernandez, J. M. Regulation of exocytotic fusion by cell inflation. *Biophys. J.* 74, 1061–1073 (1998).
24. Santos-Sacchi, J., Navarattam, D., Raphael, R. & Oliver, D. *The Cochlea Chapter 5: Prestin - molecular mechanisms underlying outer hair cell electromotility. Springer Handbook Of Auditory Research*. Vol. The Cochlea (Springer, 2017).
25. Kybalchenko, V. & Santos-Sacchi, J. C3- flux through a non-selective, stretch-sensitive conductance influences the outer hair cell motor of the guinea-pig. *J. Physiol.* 547, 873–891 (2003).
26. Oliver, D. et al. Intracellular anions as the voltage sensor of prestin, the outer hair cell motor protein. *Science* 292, 2340–2343 (2001).
27. Iwasa, K. H. Effect of stress on the membrane capacitance of the auditory outer hair cell. *Biophys. J.* 65, 492–498 (1993).
28. Gale, J. E. & Ashmore, J. F. Charge placement induced by rapid stretch in the basolateral membrane of the guinea-pig outer hair cell. *Proc. R. Soc. Lond. B. Biol. Sci.* 255, 243–249 (1994).
29. Kybalchenko, V. & Santos-Sacchi, J. Anion control of voltage sensing by the motor protein prestin in outer hair cells. *Biophys. J.* 95, 4439–4447 (2008).
30. Kekeha, S. & Santos-Sacchi, J. Membrane tension directly shifts voltage dependence of outer hair cell motility and associated gating charge. *Biophys. J.* 68, 2190–2197 (1995).
31. Gale, J. E. & Ashmore, J. F. An intrinsic frequency limit to the cochlear amplifier. *Nature* 289, 63–66 (1997).
32. Santos-Sacchi, J., Navarrete, E. & Song, L. Fast electromechanical amplification in the lateral membrane of the outer hair cell. *Biophys. J.* 96, 739–747 (2009).
33. Santos-Sacchi, J. & Song, L. Chloride-driven Electromechanical Phase Lags at Acoustic Frequencies Are Generated by SLC26a5, the Outer Hair Cell Motor Protein. *Biophys. J.* 107, 126–133 (2014).
34. Langer, J. S. Shear-transformation-zone theory of viscosity, diffusion, and stretched exponential relaxation in amorphous solids. *Phys. Rev. E Stat. Nonlin. Soft Matter Phys.* 78, 040102 (2008).
35. Lieleg, O., Kayser, J., Brambilla, G., Cipelletti, L. & Rausch, A. R. Slow dynamics and internal stress relaxation in bundled cytoskeletal networks. *Nat Mater* 10, 236–242 (2011).
36. Alverez, F., Alegria, A. & Colmenero, J. Relationship between the time-domain Kohlrausch-Williams-Watts and frequency-domain Havrilak-Negami relaxation functions. *Physical Review B* 44, 7306–7312 (1991).
37. He, D. Z., Evans, B. N. & Dallos, P. First appearance and development of electromotility in neonatal gerbil outer hair cells. *Hear. Res.* 78, 77–90 (1994).
38. Okumade, O. & Santos-Sacchi, J. IR laser-induced perturbations of the voltage-dependent solute carrier protein SLC26a5. *Biophys. J.* 105, 1822–1828 (2013).
39. Santos-Sacchi, J., Kekeha, S., Kikuchi, T., Katori, Y. & Takasaka, T. Density of motility-related charge in the outer hair cell of the guinea pig is inversely related to best frequency. *Neurosci. Lett.* 256, 155–158 (1998).
40. Mahendrasingam, S., Beurg, M., Fettplace, R. & Hackney, C. M. The ultrastructural distribution of prestin in outer hair cells: a post-embryonic immunogold embedding investigation of low-frequency and high-frequency regions of the rat cochlea. *Eur. J. Neurosci.* 31, 1595–1605 (2010).
41. Liberman, M. C. et al. Prestin is required for electromotility of the outer hair cell and for the cochlear amplifier. *Nature* 419, 300–304 (2002).
42. Mellado Lagarde, M. M., Drexl, M., Lukashkin, A. N., Zuo, J. & Russell, I. J. Prestin’s role in cochlear frequency tuning and transmission of mechanical responses to neural excitation. *Eur. Biophys. J.* 18, 200–202 (2008).
43. Rajagopalan, L. et al. Glycosylation regulates prestin cellular activity. *J.ASC.Res.Otolaryngol.* 11, 39–51 (2010).
44. Drak, L. et al. Effects of cyclic nucleotides on the function of prestin. *J. Physiol.* 563, 483–496 (2005).
45. Hallworth, R., Stark, K., Zholudeva, L., Currall, B. B. & Nichols, M. G. The conserved tetrameric subunit stoichiometry of SLC26 proteins. *Microsc. Microanal.* 19, 799–807 (2013).
46. Navarattam, D., Bai, J. P., Samarayake, H. & Santos-Sacchi, J. N-terminal-mediated homomultimerization of prestin, the outer hair cell motor protein. *Biophys. J.* 89, 3345–3352 (2005).
47. Detro-Dassen, S. et al. Conserved dimeric subunit stoichiometry of SLC26 multifunctional anion exchangers. *J.Biol.Chem.* 283, 4177–4188 (2008).
48. Lukashkina, V. A., Yamashita, T., Zuo, J., Lukashkin, A. N. & Russell, I. J. Amplification mode differs along the length of the mouse cochlea as revealed by connexin 26 deletion from specific gap junctions. *Sci. Rep.* 7, 5185 (2017).
49. Keller, J. P. et al. Functional regulation of the SLC26-family protein prestin by calcium/calmodulin. *J. Neurosci.* 34, 1325–1332 (2014).
50. Zhang, M., Kalinec, G. M., Urrutia, R., Billadeau, D. D. & Kalinec, F. ROCK-dependent and ROCK-independent control of cochlear outer hair cell electromotility. *J.Biol.Chem.* 278, 35444–35465 (2003).
51. Bai, J. P. et al. Prestin surface expression and activity are augmented by interaction with MAP1S, a microtubule-associated protein. *J.Biol.Chem.* 285, 20834–20843 (2010).
52. Spector, A. A., Brownell, W. E. & Popel, A. S. Effect of outer hair cell piezoelectricity on high-frequency receptor potentials. J. Acoust. Soc. Am. 113, 453–461 (2003).
53. Rabbitt, R. D., Clifford, S., Breneman, K. D., Farrell, B. & Brownell, W. E. Power efficiency of outer hair cell somatic electromotility. PLoS Comput. Biol. 5, e1000444 (2009).
54. Iwasa, K. H. A two-state piezoelectric model for outer hair cell motility. Biophys. J. 81, 2495–2506 (2001).
55. Santos-Sacchi, J. & Tan, W. The Frequency Response of Outer Hair Cell Voltage-Dependent Motility Is Limited by Kinetics of Prestin. J. Neurosci. 38, 5493–5506 (2018).
56. Dong, X. X. & Iwasa, K. H. Tension sensitivity of prestin: comparison with the membrane motor in outer hair cells. Biophys. J. 86, 1201–1208 (2004).
57. Vavakou, A., Cooper, N. P. & van der Heijden, M. The frequency limit of outer hair cell motility measured in vivo. Elife 8, e47667 (2019).

Acknowledgements
This work was supported by National Institutes of Health–National Institute on Deafness and Other Communication Disorders Grants R01 DC000273 (JSS), R01 DC016318 (JSS), and R01 DC008130 (JSS and DN).

Author contributions
All authors contributed to the design and analysis of experiments. J.S.S. wrote the manuscript; J.P.B. and D.N. edited the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Correspondence and requests for materials should be addressed to J.S.-S.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019