Age-related degradation of tectorial membrane dynamics with loss of CEACAM16

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ABSTRACT Studies of genetic disorders of sensorineural hearing loss have been instrumental in delineating mechanisms that underlie the remarkable sensitivity and selectivity that are hallmarks of mammalian hearing. For example, genetic modifications of TECTA and TECTB, which are principal proteins that comprise the tectorial membrane (TM), have been shown to alter auditory thresholds and frequency tuning in ways that can be understood in terms of changes in the mechanical properties of the TM. Here, we investigate effects of genetic modification targeting CEACAM16, a third important TM protein. Loss of CEACAM16 has been recently shown to lead to progressive reductions in sensitivity. Whereas age-related hearing losses have previously been linked to changes in sensory receptor cells, the role of the TM in progressive hearing loss is largely unknown. Here, we show that TM stiffness and viscosity are significantly reduced in adult mice that lack functional CEACAM16 relative to age-matched wild-type controls. By contrast, these same mechanical properties of TMs from juvenile mice that lack functional CEACAM16 are more similar to those of wild-type mice. Thus, changes in hearing phenotype align with changes in TM material properties and can be understood in terms of the same TM wave properties that were previously used to characterize modifications of TECTA and TECTB. These results demonstrate that CEACAM16 is essential for maintaining TM mechanical and wave properties, which in turn are necessary for sustaining the remarkable sensitivity and selectivity of mammalian hearing with increasing age.

INTRODUCTION

The mammalian cochlea is a remarkable sensor that can reliably detect vibrations on the order of picometers (1), and perform high-quality frequency analysis such that the frequency of sound is mapped to place along the cochlear partition (2). Mechanical measurements have established that high sensitivity, sharp tuning, and nonlinearity are already manifest in the peripheral stages of auditory processing (3). It is now widely accepted that these remarkable properties ultimately derive from active mechanical amplification residing in the cochlea. Although there is ongoing debate about the nature of the cochlear amplifier, many observations point to somatic motility of outer hair cells (4) generated by prestin (5,6) as the principal component of the amplifier. Genetic manipulations that eliminate prestin-based somatic electromotility cause significant hearing loss (7). Hair-bundle-based adaptation mechanisms, which are important for amplifying the response to sound in non-mammalian cochleae, are also present in mammalian hair cells (8–10), suggesting that bundle-based amplification also plays a role in cochlear mechanics. In addition, genetic manipulations of proteins in the tectorial membrane (TM)
have been shown to cause hearing loss (11–13). Recent studies suggest complex phenomena associated with the TM, such as resonance (14,15), pulsatile radial fluid flow (16), TM traveling waves (17–23), and poroelastic effects (22,24) are all linked with cochlear sensitivity and tuning. The development of mouse models of genetic hearing disorders that exclusively target the TM have linked changes in sensitivity and tuning in mutant mice with changes to the TM (11–13,25–28). The wide range of functional deficits associated with these mouse models illustrates the importance of this accessory structure to the integrity of the amplifying feedback loop in which outer hair cells’ somatic and bundle motility reside. Although there has been progress in elucidating the importance of TM traveling waves in the mechanical excitation of cochlear hair cells, the mechanisms by which the dynamic mechanical properties of the TM contribute to system-level amplification and how alterations might influence age-related hearing loss remain unclear. Studies examining protein turnover in the cochlea have revealed that the TM is a relatively stable structure (29). However, recent studies have shown that loss of CEACAM16, an abundant protein that comprises the TM’s striated sheet structure along with TECTA and TECTB (12,13,30–33), leads to anatomical changes and progressive decline in hearing function. Apart from the increased incidence of spontaneous otoacoustic emissions (SOAEs), cochlear function in Ceacam16<sup>gal/gal</sup> mutant mice is near normal in juveniles at~1 month of age. However, distortion product otoacoustic emissions (DPOAE) decrease and auditory brainstem response (ABR) thresholds increase at older ages when compared with Ceacam16<sup>+/+</sup> mice (34). By one year of age, mice lacking CEACAM16 have significantly elevated ABR thresholds at all frequencies and no emissions of any kind. These changes have been attributed to a progressive loss of matrix from the core of the TM and to accelerated age-related degeneration of the TM in Ceacam16<sup>gal/gal</sup> mice (28,34).

CEACAM16 has been posited to interact with TECTA and TECTB (28,34), thereby forming the TM’s striated sheet matrix (33). Although transcription of Tecta and Tectb is critical for the development of TM structure (35), mRNA for these two genes is not measurable after weaning in mice (36). In contrast, Ceacam16 is transcribed and CEACAM16 is secreted in adult mice by a variety of nonsensory and supporting cells, including epithelial cells of the spiral limbus and inner sulcus, border cells, inner and outer pillar cells, and Deiters’ cells (28). Continued expression of CEACAM16 by a variety of cell types and its interaction with other TM striated sheet proteins suggests that it plays an essential role in maintaining the structure of the TM with increasing age.

To better understand how the loss of functional CEACAM16 affects cochlear mechanisms, we explore dynamic wave properties of TMs from Ceacam16<sup>gal/gal</sup> and Ceacam16<sup>+/+</sup> mice at various ages. We show that TM trav-eling wave decay constants and wave speeds, measured in isolated TM segments excised from the middle cochlear turn, are reduced in adult Ceacam16<sup>gal/gal</sup> mice (~12–14 weeks of age) compared with juvenile Ceacam16<sup>gal/gal</sup> (~4–6 weeks of age) and adult Ceacam16<sup>+/+</sup> mice. Loss of sensitivity in mice lacking CEACAM16 could therefore relate to changes in the mechanical interactions between motion patterns along the basilar membrane and TM. We analyze traveling wave properties of the TM and determine the corresponding material properties, including shear storage modulus (G<sup>s</sup>) and shear viscosity (η). In addition to determining wave properties, these material properties may also have a direct effect on the stimulation of hair bundles of sensory hair cells and on the magnitudes of DPOAEs produced in these mutants.

**MATERIALS AND METHODS**

**Isolated TM preparation**

TM segments were isolated from mice ranging from 4 to 14 weeks of age using previously published techniques (37). One TM segment was isolated from each of six Ceacam16<sup>+/+</sup> mice (12–14 weeks of age), from each of two juvenile Ceacam16<sup>gal/gal</sup> mice (4–6 weeks of age) and from each of five adult Ceacam16<sup>gal/gal</sup> (12–14 weeks of age) mice. All of the TM segments were from the mid apical region of the cochlea. All of these experimental animals originated from C57Bl/6J background strains. Cochleae were surgically excised and immersed in artificial endolymph (AE) containing 174 mM KCl, 5 mM HEPES, 3 mM dextrose, 2 mM NaCl, and 0.02 mM CaCl<sub>2</sub>. The AE bath was equilibrated at room temperature to pH 7.15. The bone encasing the cochlea was removed with a #11 scalpel blade to expose the Organ of Corti. Bright- and dark-field illumination using a dissection microscope (Zeiss, Oberkochen, Germany) allowed for visualization of the TM along the cochlear spiral. A sterile eyelash was then used to remove the membrane from its limbal attachment to the Organ of Corti. TM segments from the middle cochlear turn were then removed using a micropipette and placed in fresh AE in preparation for wave chamber experiments. The care and use of animals in this study were approved by the Massachusetts Institute of Technology Committee on Animal Care.

**TM wave chamber**

Isolated TM segments were suspended between vibrating and stationary supports in a wave chamber containing AE (Fig. 1 A). The vibrating support was attached to the underlying glass slide through a piezoelectric actuator (Thorlabs, Newton, NJ) that delivered sinusoidal motions in the radial direction at audio frequencies (10–20 kHz). The stationary support was attached directly to the underlying glass slide. Using a sterile eyelash, a TM segment was carefully attached to the top surfaces of the supports, which had previously been coated with 3 μL of tissue adhesive (Cell-Tak; Collaborative Research, Bedford, MA).

**Optical imaging and analysis**

Stop-action images of sinusoidally excited TM segments were obtained using stroboscopic illumination from a light-emitting diode that was focused on TM samples with the transmitted-light condenser (Zeiss Axiosplan; Carl Zeiss, Oberkochen, Germany). The resulting images from a 20× water-immersion objective (0.5 N.A.) were captured with a five-megapixel charge-coupled device camera (Stingray; Allied Vision Technologies, Singapore, Singapore). Images were obtained...
A segment of an isolated TM is suspended in artificial endolymph (AE) between two glass cover slips so that vibrations of the left cover slip excite radial motions of the TM that propagate as a wave traveling in the longitudinal direction. The amplitude and phase of motion as a function of pixel location is determined from stroboscopic images obtained with a video microscope at eight phases of the sinusoidal stimulus. (B and C) Representative results. Images of the TM are shaded (cyan) to indicate the region where the amplitude of motion is attenuated by less than a factor of 2 relative to the amplitude of at the edge of the vibrating support. The width of this region is given by the decay constant \( \sigma \) and is represented in this figure by the length of the cyan line. The smaller decay constant in (C) (135 \( \mu \text{m} \)) relative to (B) (221 \( \mu \text{m} \)) indicates less spread of excitation in the Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) preparation than in the wild-type preparation. The colored lines represent lines of constant phase separated by \( 2 \pi /16 \) radians, which is equal to the wavelength \( \lambda \) divided by 16. The smaller separation between these lines in (C) (95 \( \mu \text{m} \)) relative to those in (B) (116 \( \mu \text{m} \)) indicates that the wave in the Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) preparation travels more slowly (3.80 m/s) than that in the wild-type preparation (4.64 m/s). The stimulus frequency was 10 kHz. To see this figure in color, go online.

**RESULTS AND DISCUSSION**

**TM wave parameters for Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) and wild-type TMs**

We measured TM wave motions in both wild-type mice (Ceacam16\(^{+/+} \)) and mice without functional CEACAM16 (Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \)). Representative results for TMs from adult mice are shown in Fig. 1, B and C illustrate two important trends; TM waves propagate more slowly and dissipate over shorter longitudinal distances in TMs from adult Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) mice than in TMs from adult wild-type mice. As the TM wave propagates, the amplitude of motion tends to decrease with distance. The shaded regions of Fig. 1, B and C highlight the portions of the TM for which the amplitude of the radial motion is attenuated by less than a factor of 2 relative to that of the vibrating support. The width of this region corresponds to one decay constant \( \sigma \), which is illustrated by the horizontal cyan bars in Fig. 1, B and C. The average decay constant is larger for this wild-type TM (\( \sigma = 221 \mu \text{m} \)) than it is for this Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) TM (\( \sigma = 135 \mu \text{m} \)). These results suggest that mechanical spread of excitation through the TM would be smaller in Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) TMs than in wild-type TMs. As the TM wave propagates, the phase of motion also tends to decrease. The colored lines in Fig. 1, B and C illustrate lines of constant phase separated by \( 2 \pi /16 \) radians. Wave speed \( v \) can be calculated from the distance between adjacent lines because wave speed \( v = f \lambda \). Because the wavelength \( \lambda \) is smaller for the Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) TM (\( \lambda = 380 \mu \text{m} \)) than for the wild-type TM (\( \lambda = 380 \mu \text{m} \)), it follows that the speed is also smaller (\( v = 3.80 \text{ m/s} \) for the Ceacam16\(^{\beta \text{gal}/\beta \text{gal}} \) TM vs. 4.64 m/s for the wild-type).

Similar results for multiple preparations and for frequencies from 10 to 15 kHz are presented in Fig. 2. Across this range of frequencies, wave speeds (Fig. 2 A) for TMs...
from adult Ceacam16\textsuperscript{gal/gal} mice (green circles) tend to be smaller than those from juvenile Ceacam16\textsuperscript{gal/gal} mice (orange Y signs) and smaller than those from wild-type mice (blue plus signs). These trends are summarized in bar plots (Fig. 2 C), where the heights of the colored bars represent the median speeds, and the black lines represent interquartile ranges (iqr’s).

Wave speeds were generally slower for adult Ceacam16\textsuperscript{gal/gal} TMs (median: 4.55 m/s; iqr: 3.68–4.76 m/s; n = 49 measurements from five preparations) than for wild-type TMs (median: 5.88 m/s; iqr: 5.12–6.62 m/s; n = 50 measurements from six preparations), and the difference was highly significant (p < 10\textsuperscript{-6} in Welch’s t-test, with t = 6.55 and 68.1 (dof), computed using the Satterthwaite approximation (40)). Median wave speeds for young Ceacam16\textsuperscript{gal/gal} TMs (median: 5.17 m/s; iqr: 4.50–6.89 m/s; n = 12 measurements from two preparations) were between those for adult Ceacam16\textsuperscript{gal/gal} TMs and those for wild-type TMs. Differences between young Ceacam16\textsuperscript{gal/gal} and adult Ceacam16\textsuperscript{gal/gal} TMs (p < 0.0061, with t = 2.92 and 12.8 dof) were highly significant (i.e., p < 0.01). Differences between young Ceacam16\textsuperscript{gal/gal} TMs and wild-type TMs were not statistically significant (p = 0.11, with t = 1.28 and 20.8 dof).

Decay constants for TMs from adult Ceacam16\textsuperscript{gal/gal} mice (Fig. 2 B, green circles) also tend to be smaller than those for juvenile Ceacam16\textsuperscript{gal/gal} mice (orange Y signs) or those for wild-type mice (blue plus signs), and decay constants for TMs from wild-type mice tend to be larger than those for juvenile or adult Ceacam16\textsuperscript{gal/gal} mice. These trends are summarized in the bar plots shown in Fig. 2 D. Decay constants were generally smaller for adult Ceacam16\textsuperscript{gal/gal} TMs (median: 94 μm; iqr: 77–121 μm; n = 49 measurements from five preparations) than for wild-type TMs (median: 196 μm; iqr: 164–239 μm; n = 50 measurements from six preparations), and the difference was highly significant (p < 10\textsuperscript{-6}, with t = 8.19 and 96.3 dof). Median decay constants for young Ceacam16\textsuperscript{gal/gal} TMs (median: 123 μm; iqr: 91–207 μm; n = 12 measurements from two preparations) were between those for adult Ceacam16\textsuperscript{gal/gal} TMs and those for wild-type TMs. Decay constants for TMs from juvenile Ceacam16\textsuperscript{gal/gal} mice were not significantly different from those for wild-type mice (p = 0.21, with t = 0.83 and 12.0 dof) or adult Ceacam16\textsuperscript{gal/gal} mice (p = 0.079, with t = 1.50 and 11.9 dof).

In summary, we measured wave parameters of three mouse populations: adult Ceacam16\textsuperscript{gal/gal} mice, juvenile Ceacam16\textsuperscript{gal/gal} mice, and wild-type mice. Both the wave speeds and decay constants were smaller in adult Ceacam16\textsuperscript{gal/gal} TMs than in wild-type TMs, and those differences were highly significant, demonstrating important mechanical differences between Ceacam16\textsuperscript{gal/gal} and wild-type TMs. Also, the wave speeds in adult Ceacam16\textsuperscript{gal/gal} TMs are smaller than those in juvenile Ceacam16\textsuperscript{gal/gal} TMs, and these differences are highly significant, demonstrating age-related changes in the mechanical properties of Ceacam16\textsuperscript{gal/gal} TMs.

**TM material parameters for Ceacam16\textsuperscript{gal/gal} and wild-type TMs**

Wave properties of viscoelastic materials derive from their material properties according to the following relationship (41,42):

\[
\left( \frac{2\pi}{\lambda} - j \frac{1}{\sigma} \right)^2 = k^2 = \frac{\rho \omega^2}{G' + j\omega\eta},
\]

where ρ is density, G’ is shear modulus, η is shear viscosity, and ω is angular frequency in radians/s. We can use this relationship to compute the material properties (G’ and η) from the wave properties (λ and σ) presented in the previous section. Notice, however, that these material properties also depend on both density ρ and angular frequency ω. We can account for the dependence on ρ and ω by defining normalized material properties G’/ρω\textsuperscript{2} and η/ρω:

\[
\left( \frac{2\pi}{\lambda} - j \frac{1}{\sigma} \right)^2 = k^2 = \frac{1}{\rho \omega^2} + j \frac{\eta}{\rho \omega},
\]

which depend on only λ and σ. In Fig. 3 we illustrate the dependence of normalized shear modulus (Fig. 3 B) and normalized shear viscosity (Fig. 3 C) on wave parameters. We use these maps to convert the range of observed wave parameters (Fig. 3 A) to corresponding ranges of normalized material properties (Fig. 3 D) for wild-type and Ceacam16\textsuperscript{gal/gal} TMs.

The two-dimensional maps in Fig. 3 A provide a concise representation of differences between the wave properties of adult Ceacam16\textsuperscript{gal/gal} and wild-type TMs, which have nonoverlapping interquartile ranges in both wavelength and decay constant dimensions. By contrast, the interquartile ranges of these wave properties for juvenile Ceacam16\textsuperscript{gal/gal} TMs overlap with both adult Ceacam16\textsuperscript{gal/gal} and wild-type TMs. There is considerable overlap of the corresponding material properties in Fig. 3 D. The median values of normalized shear viscosity for the TMs of juvenile Ceacam16\textsuperscript{gal/gal} and wild-type mice are similar (median: 3.52 (μm\textsuperscript{2}); iqr: 2.33–3.86 (μm\textsuperscript{2}); n = 12 measurements from two preparations for the former; median: 3.45 (μm\textsuperscript{2}); iqr: 2.29–5.20 (μm\textsuperscript{2}); n = 50 measurements for the latter). However, the median value of normalized shear viscosity of adult Ceacam16\textsuperscript{gal/gal} TMs (median: 1.90 (μm\textsuperscript{2}); iqr: 1.46–2.36 (μm\textsuperscript{2}); n = 49 measurements from five preparations) is smaller than that of juvenile Ceacam16\textsuperscript{gal/gal} TMs by a factor of 1.85. The median value of normalized shear modulus for juvenile Ceacam16\textsuperscript{gal/gal} TMs (median: 1.89 (μm\textsuperscript{2}); iqr: 0.69–3.73 (μm\textsuperscript{2}); n = 12 measurements...
from two preparations) is nearly a factor of two smaller than that for wild-type TMs (median: 3.74 (μm)²; iqr: 3.07–4.40 (μm)²; n = 50 measurements from six preparations). The median value of normalized shear modulus for adult Ceacam16gal/gal TMs (median: 1.11 (μm)²; iqr: 0.66–1.63 (μm)²; n = 49 measurements from five preparations) is more than three times smaller than that for wild-type TMs.

In summary, the normalized shear modulus of adult Ceacam16gal/gal TMs tend to be smaller than those of juvenile Ceacam16gal/gal TMs, and those of juvenile Ceacam16gal/gal TMs then tend to be smaller than those of wild-type TMs. These trends are consistent with the increasing prominence of holes in the TMs of mice lacking Ceacam16 (34). However, the normalized shear viscosity of juvenile Ceacam16gal/gal TMs is comparable with that of wild-type TMs, suggesting that different mechanisms may contribute shear viscosity and shear stiffness.

Comparisons of properties of Ceacam16gal/gal, TectaY1870C+ and Tectb–/– TMs

We have previously measured wave and material properties in mice with mutations that target α-tectorin (TectaY1870C+) and β-tectorin (Tectb–/–). Both of these mutations reduce TM shear modulus relative to wild-type TMs (18,43). However, the two mutations are associated with different hearing phenotypes; Tectb–/– mice have sharpened basilar membrane tuning by a factor of two to three at mid and high frequencies (13), whereas TectaY1870C+ mice have normal basilar membrane tuning and even broader neural tuning (11). Because the stiffnesses of TectaY1870C+ and Tectb–/– TMs are similar, stiffness alone cannot account for observed differences in hearing phenotypes. However, there are also differences in viscous loss. The viscous component of TectaY1870C+ TMs is smaller than that of wild-types (43). In contrast, the shear viscosity of Tectb–/– TMs is similar to that of wild-types (18). Paradoxically, the larger viscosity in Tectb–/– TMs is associated with sharper tuning, which is the opposite of predictions from conventional models of viscous loss.

Wave and material properties of adult Ceacam16gal/gal and wild-type TMs are compared with previously published results for TectaY1870C+ and Tectb–/– TMs (22) in Fig. 4. Decay constants (Fig. 4 A) for TectaY1870C+ TMs (median: 247 μm; iqr: 181–314 μm; n = 12 measurements from seven preparations) are generally greater than those for Tectb–/– TMs (median: 162 μm; iqr: 121–204 μm; n = 8 measurements from four preparations), and those for Tectb–/– TMs are generally greater than those for Ceacam16gal/gal TMs (median: 94 μm; iqr: 77–121 μm; n = 49 measurements from five preparations). Both of these relations are statistically significant (p = 0.014, with τ = 2.38 and 18.0 dof for the former, and p = 0.029, with t = 2.18 and 8.9 df for the latter).

Wavelengths for Tectb–/– TMs (median: 422 μm; iqr: 370–474 μm; n = 12 measurements from four preparations) are generally greater than those for both Ceacam16gal/gal TMs (median: 339 μm; iqr: 300–381 μm; n = 49 measurements from five preparations) and TectaY1870C+ TMs (median: 310 μm; iqr: 259–360 μm; n = 12 measurements from seven preparations). Both of these relations are highly significant (p < 0.01). To see this figure in color, go online.
significant ($p = 0.0021$, with $t = 3.41$ and 14.4 dof for the former; $p = 0.00078$, with $t = 3.61$ and 22.0 dof for the latter). The small differences in wavelengths for Ceacam16$^{gal/gal}$ TMs and Tecta$^{Y1870C/+}$ TMs are not statistically significant ($p = 0.10$, with $t = 1.33$ and 14.6 dof).

Fig. 4, B and C illustrate how the preceding wave parameters map to normalized material properties, with the results shown in Fig. 4 D. The normalized shear moduli for Tecta$^{Y1870C/+}$ (median: 2.16 ($\mu$m)$^2$; iqr: 1.57–2.80 ($\mu$m)$^2$; $n = 12$ measurements from seven preparations) and Tectb$^{-/}$ TMs (median: 2.72 ($\mu$m)$^2$; iqr: 1.81–3.27 ($\mu$m)$^2$; $n = 8$ measurements from four preparations) are not significantly different ($p = 0.13$, with $t = 1.18$ and 13.4 dof). However, both are smaller than those of wild-type TMs (median: 3.74 ($\mu$m)$^2$; iqr: 3.07–4.40 ($\mu$m)$^2$; $n = 50$ measurements from six preparations) and larger than those of adult Ceacam16$^{gal/gal}$ TMs (median: 1.11 ($\mu$m)$^2$; iqr: 0.66–1.63 ($\mu$m)$^2$; $n = 49$ measurements from five preparations), and these differences are statistically significant ($p = 0.019$, with $t = 2.44$ and 9.0 dof for the comparison of wild-type and Tectb$^{-/}$ TMs, and $p = 0.0014$, with $t = 3.60$ and 14.5 dof for the comparison of Tecta$^{Y1870C/+}$ and Ceacam16$^{gal/gal}$ TMs).

The normalized shear viscosities of Tectb$^{-/}$ TMs (median: 2.72 ($\mu$m)$^2$; iqr: 1.96–3.58 ($\mu$m)$^2$; $n = 8$ measurements from four preparations) and wild-type TMs (median: 3.45 ($\mu$m)$^2$; iqr: 2.29–5.20 ($\mu$m)$^2$; $n = 50$ measurements from six preparations) are not significantly different ($p = 0.097$, with $t = 1.35$ and 15.5 dof). But the normalized shear viscosities of Ceacam16$^{gal/gal}$ TMs (median: 1.90 ($\mu$m)$^2$; iqr: 1.46–2.36 ($\mu$m)$^2$; $n = 49$ measurements from five preparations) tend to be smaller than those of Tectb$^{-/}$ TMs ($p = 0.051$, with $t = 1.85$ and 7.7 dof), and those of Tecta$^{Y1870C/+}$ TMs (median: 0.90 ($\mu$m)$^2$; iqr: 0.54–1.37 ($\mu$m)$^2$; $n = 12$ measurements from seven preparations) are smaller than those of Ceacam16$^{gal/gal}$ TMs ($p < 10^{-4}$, with $t = 4.79$ and 17.8 dof).

Interestingly, the progression from largest to smallest values of normalized shear modulus is different from that for normalized shear viscosity. In particular, the normalized shear modulus of the Tecta$^{Y1870C/+}$ TMs was greater than that of the Ceacam16$^{gal/gal}$ TMs, whereas the normalized shear viscosity of the Tecta$^{Y1870C/+}$ TMs was smaller than that of the Ceacam16$^{gal/gal}$ TMs. These results make it clear that the mechanisms that underlie viscosity and stiffness differ.

Implications of differences in material properties

The median value of normalized shear modulus for Ceacam16$^{gal/gal}$ TMs is smaller than that of Tectb$^{-/}$ TMs, and both of these are smaller than that of wild-type TMs. The similarity of these trends with those for normalized shear viscosity suggests that both trends may result from a decrease in striated sheet matrix that contributes to both of these material properties. Interestingly, a similar trend does not hold for Tecta$^{Y1870C/+}$ TMs. Whereas the median shear modulus for Ceacam16$^{gal/gal}$ TMs is approximately half that for Tecta$^{Y1870C/+}$ TMs, the median shear viscosity for Ceacam16$^{gal/gal}$ TMs is more than a factor of two greater than that for Tecta$^{Y1870C/+}$ TMs. This prominent difference suggests that other important structural changes (such as protein cross-linking) are likely to be important.
for understanding differences in the shear viscosities of these mutant TMs.

**Implications for cochlear mechanisms**

The hearing phenotype associated with mice without functional CEACAM16 differs from that associated with wild-type mice in (at least) two important ways: SOAEs are much more prevalent in juvenile Ceacam16<sup>gal/gal</sup> mice than in age-matched wild-types, and adult Ceacam16<sup>gal/gal</sup> mice have progressive elevation of hearing thresholds relative to age-matched wild-types (28,34). The decreases in shear storage modulus and shear viscosity shown in Fig. 3 could play important roles in both of these characteristics of the Ceacam16<sup>gal/gal</sup> phenotype. Recent models (44) suggest that reducing viscous and elastic coupling through the TM increases the prevalence of unstable modes (and presumably the prevalence of SOAEs) and decreases cochlear sensitivity to low-level stimuli.

The median values of normalized shear modulus and normalized shear viscosity progress from 3.74 (μm)<sup>2</sup> and 3.45 (μm)<sup>2</sup> for wild-type TMs to 1.89 (μm)<sup>2</sup> and 3.52 (μm)<sup>2</sup> for juvenile Ceacam16<sup>gal/gal</sup> TMs to 1.11 (μm)<sup>2</sup> and 1.90 (μm)<sup>2</sup> for adult Ceacam16<sup>gal/gal</sup> TMs, corresponding to an average of 0.5, 2, and 30 unstable modes, respectively (44). Whereas this modeling predicts an age-related increase in unstable modes, the reverse is seen in experiments in which SOEs were observed in 70% of juvenile Ceacam16<sup>gal/gal</sup> mice but in only 10% of Ceacam16<sup>gal/gal</sup> mice at 6–7 months of age. Although young mutants retain wild-type-like sensitivity (as assessed with ABR thresholds), they have reduced DPOAEs at 6–7 months of age, as do Tecta<sup>Y1870C/+</sup> mice. In fact, both mutants show a partial loss of gain (as assessed ABR thresholds), yet the Tecta<sup>Y1870C/+</sup> mice are prolific emitters (45), whereas the older mice lacking Ceacam16 are not.

TM properties other than shear storage modulus (G') and shear viscosity (η) could further complicate these comparisons. Our experimental chamber was designed to observe the longitudinal spread of radial excitation of the TM. However, other modes of motion could also be important given that the TM is morphologically (46) and functionally anisotropic (47–49). For example, it has been suggested that length changes of outer hair cells can induce transverse motions of the subtectorial fluid and the overlying TM, thereby enhancing inner hair cell excitations (16). Longitudinal motions have also been reported (50) and may play a role in creating vibration hotspots (see discussion in (50)). All of these factors point to the importance of considering the three-dimensional nature of mechanical interactions within the cochlear partition.

The anisotropic structure of the TM mirrors its anisotropic architecture, with collagenous proteins contributing to its network of radial fibers coursing through a striated sheet matrix composed primarily of two noncollagenous proteins: α-tectorin (TECTA) and β-tectorin (TECTB) (51). Whereas Tecta and Tectb are expressed at high levels during development, their expression is not detectable after postnatal day 22 (36). In contrast, Ceacam16 is expressed from postnatal day 12 into adulthood (32), suggesting that CEACAM16 may stabilize TECTA in the TMs of adults because the two proteins are known to interact (31).

Other mechanical properties may also contribute to changes in hearing associated with Ceacam16<sup>gal/gal</sup>

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**FIGURE 4** Comparison of wave and material properties of Ceacam16<sup>gal/gal</sup>, Tecta<sup>Y1870C/+</sup>, and Tectb<sup>−/−</sup> TMs. (A) Wave parameters for wild-type (WT), Tecta<sup>Y1870C/+</sup> (A), Tectb<sup>−/−</sup> (B), and adult Ceacam16<sup>gal/gal</sup> (C) TMs. (B and C) Dependence of normalized shear modulus and normalized shear viscosity on wave parameters. (D) Material properties of Ceacam16<sup>gal/gal</sup>, Tecta<sup>Y1870C/+</sup>, and Tectb<sup>−/−</sup> TMs. Other aspects of this figure are described in the caption of Fig. 3. To see this figure in color, go online.
mice. For example, differences in coupling between neighboring outer hair cells, loss of Hensen’s stripe, changes in the subtectorial space, and/or the emergence of holes in the TMs of mutant mice could influence the degree to which SOAEs are generated (28, 44). Furthermore, the recent demonstration (24) that nanomechanical properties of the TM differ substantially from the micromechanical properties measured in this study may be especially relevant because they predict that mechanical interactions between the TM and individual hair bundles may differ significantly from those that govern longitudinal coupling within the core of the TM. It has also been suggested that the TM may act as a calcium reservoir (52). Given the several calcium-dependent processes that influence the tip-link and transducer complex, the implications of changes to the structure, the material properties, and the wave characteristics of the TM are not yet fully understood.

CONCLUSIONS

CEACAM16 is a noncollagenous glycoprotein that is essential to normal hearing and to the structure of the striated sheet matrix that comprises the core of the TM. Mice that lack Ceacam16 exhibit an increased incidence of SOAEs as juveniles and progressive hearing loss as adults. To better understand the cochlear mechanisms that underlie these behavioral changes, we have measured wave and material properties of TMs isolated from Ceacam16^gal/gal and wild-type mice and determined that adult but not juvenile mutants have statistically different wave speeds and decay constants relative to controls. Additionally, we compared those results to previous measurements in Tecta^Y1870C^+ and Tectb^–/– mutants. Results show a clear separation of wave properties. Interestingly, there is a monotonic progression, with both median wave speed and median decay constants being larger in wild-type TMs than in Tectb^–/– TMs and larger in Tectb^–/– TMs than in adult Ceacam16^gal/gal TMs. However, Tecta^Y1870C^+ TMs do not follow this same monotonic progression; instead, they have significantly slower speeds and larger decay constants than would be expected from the trends for the other groups.

Correlations between these results and previously measured threshold shifts suggest that the slower speeds observed in adult Ceacam16^gal/gal as well as in Tecta^Y1870C^+ and Tectb^–/– TMs may contribute to the increase in hearing thresholds, as suggested in some cochlear models (53). Furthermore, whereas relatively small differences in material properties were observed in juvenile Ceacam16^gal/gal TMs relative to wild-type TMs, recent models (44) show that these differences are sufficient to increase the number of SOAEs in juvenile Ceacam16^gal/gal TMs. By contrast, the relatively large differences in material properties in adult Ceacam16^gal/gal TMs would decrease sensitivity (as seen in behavioral tests) and thereby also inhibit SOAEs that would otherwise be even more numerous than in juveniles.

In conclusion, comparisons of Ceacam16^gal/gal, Tecta^Y1870C^+, and Tectb^–/– TMs suggest that the behavior of the TM is a result of a combination of properties that interact in complicated ways to assure proper hair cell activation and to stabilize the active process. It follows that properties of the hearing phenotype can depend in complicated ways on the many properties of the TM.

AUTHOR CONTRIBUTIONS

J.B.S., A.M., R.G., M.A.C., and D.M.F. designed the research. A.M. and J.B.S. performed the research. A.M., D.F., and J.B.S. analyzed the data. J.B.S., A.M., R.G., M.A.C., and D.M.F. wrote the study.

ACKNOWLEDGMENTS

This work was supported by National Institutes of Health grant R01-DC00238 and R01-DC00089.

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