Altered cochlear innervation in developing and mature naked and Damaraland mole rats

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Abstract
Compared to many other rodent species, naked mole rats (Heterocephalus glaber) have elevated auditory thresholds, poor frequency selectivity, and limited ability to localize sound. Because the cochlea is responsible for encoding and relaying auditory signals to the brain, we used immuno-fluorescence and quantitative image analysis to examine cochlear innervation in mature and developing naked mole rats compared to mice (Mus musculus), gerbils (Meriones unguiculatus), and Damaraland mole rats (Fukomys damarensis), another subterranean rodent. In comparison to mice and gerbils, we observed alterations in afferent and efferent innervation as well as their patterns of developmental refinement in naked and Damaraland mole rats. These alterations were, however, not always shared similarly between naked and Damaraland mole rats. Most conspicuously, in both naked and Damaraland mole rats, inner hair cell (IHC) afferent ribbon density was reduced, whereas outer hair cell afferent ribbon density was increased. Naked and Damaraland mole rats also showed reduced lateral and medial efferent terminal density. Developmentally, naked mole rats showed reduced and prolonged postnatal reorganization of afferent and efferent innervation. Damaraland mole rats showed no evidence of postnatal reorganization. Differences in cochlear innervation specifically between the two subterranean rodents and more broadly among rodents provides insight into the cochlear mechanisms that enhance frequency sensitivity and sound localization, maturation of the auditory system, and the evolutionary adaptations occurring in response to subterranean environments.

KEYWORDS
auditory, BK channels, cochlea, development, gerbil, inner and outer hair cells, inner ear, lateral and medial efferent olivocochlear innervation, mice, presynaptic afferent ribbons, RRID:AB_11150471, RRID:AB_143157, RRID:AB_2200400, RRID:AB_2313725, RRID:AB_2534069, RRID:AB_2535810, RRID:AB_399431, RRID:AB_90710, tonotopy

INTRODUCTION

Naked mole rats (Heterocephalus glaber) are long-lived rodents that live in large eusocial colonies in narrow, underground burrows. Their subterranean environment restricts oxygen levels, light exposure, and acoustic stimulation and has led to various physiological adaptations (Hetling et al., 2005; Park et al., 2017). Some of these adaptations might involve neotenous retention of immature characteristics by prolonged and/or arrested development (Larson & Park, 2009; Skulachev et al., 2017). The unique life history of naked mole rats has made them of enormous interest to comparative biologists and biomedical researchers. Nonetheless, their auditory system has been relatively uninvestigated. Further investigation of the structure and function of the auditory system in this species would provide comparative insight into the anatomical and physiological mechanisms that shape auditory responses throughout the lifespan of an organism.
Compared to many other rodent species, naked mole rats have relatively high auditory thresholds, poor high frequency hearing, and a limited ability to localize sound. For comparison, mice (Mus musculus) hear frequencies extending from 1 kHz to about 100 kHz, with the most sensitive frequency in the audiogram being approximately 16 kHz (Heffner & Heffner, 1993). Gerbils (Meriones unguiculatus) hear frequencies ranging from 125 Hz to about 60 kHz, with the most sensitive frequency being approximately 2 kHz (Müller, 1996; Ryan, 1976). Both mice and gerbils localize sound effectively, with mice using primarily interaural level differences and pinna cues (Allen & Ison, 2010; Heffner, Koay, & Heffner, 2001) and gerbils taking additional advantage of interaural time differences (Tolnai, Beutelmann, & Ison, 2010; Heffner, Koay, & Heffner, 2001) and gerbils taking additional advantage of interaural time differences (Tolnai, Beutelmann, & Ison, 2010; Heffner, Koay, & Heffner, 2001). Although hearing in Damaraland mole rats, another subterranean mole rat, has not been localized sound (Heffner & Heffner, 1993). Although hearing in Damaraland mole rats, another subterranean mole rat, has not been investigated, other Fukomys species show reduced cochlear tuning (Kössl, Frank, Burda, & Muller, 1996), elevated auditory thresholds and frequency ranges restricted to between 125 Hz and 4 kHz (five octaves), with best hearing within 0.8 and 1.4 kHz (Gerhardt, Henning, Begall, & Malkemper, 2017).

Previous work in naked mole rats has documented anatomical differences in their peripheral and central auditory structures, but how these differences contribute to comparative differences in their auditory function is unclear. Most conspicuously, naked and Damaraland mole rats lack pinnae, external structures of the outer ear that funnel sound and provide localization cues. Structural differences in the middle and inner ear of naked mole rats were also recently characterized by Mason, Cornwall, and Smith (2016) using microcomputed tomography and dissections. The authors of this study concluded that these structural differences were not by themselves sufficient to explain poor high frequency hearing in naked mole rats. Moreover, naked mole rats did not show specializations (most notably, hypertrophy of the malleus) that support low frequency hearing by inertial bone conduction (i.e., seismic hearing). Centrally, naked mole rats possess all major auditory brainstem nuclei (Gessele, Garcia-Pino, Omerbasic, Park, & Koch, 2016; Heffner & Heffner, 1993) except the superior paraolivary nucleus (Gessele et al., 2016). However, the structures involved in binaural processing of sound and sound localization are comparatively smaller (Gessele et al., 2016; Heffner & Heffner, 1993). The molecular organization of these structures has been investigated only more recently and only centrally. Perineuronal nets, specialized extracellular matrix structures responsible for synaptic stabilization in the adult brain, are observed in the auditory brainstem nuclei in naked mole rats and show no differences compared to other rodents (Beebe & Schofield, 2018). Nonetheless, molecular differences were reported in another study: specifically, the entire superior olivary complex lacked membrane-bound expression of the hyperpolarization-activated channel HCN1 (Gessele et al., 2016), an ion channel that greatly contributes to binaural temporal precision in the superior olivary complex of other rodents (Khurana et al., 2012; Koch, Braun, Kaper, & Grothe, 2004).

In this study, we were particularly interested in examining the anatomical and molecular organization of cochlear innervation in the auditory sensory epithelium (organ of Corti) of naked mole rats because this innervation is responsible for encoding and relaying auditory signals to the brain and, moreover, shapes the development of auditory brainstem structures (O’Neill, Connelly, Limb, & Ryugo, 2011). We hypothesized that differences in the innervation of the organ of Corti, perhaps resulting from reduced postnatal maturation, contribute to differences in auditory function in naked mole rats in comparison to other rodents. To begin to test this hypothesis, we used immunofluorescence to examine patterns of afferent and efferent innervation in excised organs of Corti from mature and developing naked mole rats compared to mice, gerbils, and also Damaraland mole rats (Fukomys damarensis). This selection of species allowed comparison of naked mole rats to rodents with sensitive high frequency hearing (mouse), sensitive low frequency hearing (gerbils) and another more closely related, subterranean, eusocial mole rate (Damaraland mole rats). We observed alterations in afferent and efferent innervation as well as their patterns of developmental refinement in naked mole rats compared to both mice and gerbils and also compared to Damaraland mole rats. Differences in cochlear innervation specifically between the two subterranean rodents and more broadly among rodents provides insight into the evolutionary adaptations to subterranean environments as well as the cochlear mechanisms that contribute to hearing sensitivity and sound localization.

2 | MATERIALS AND METHODS

2.1 | Animals

Animal protocols conformed to the respective legislation at the University of Illinois at Chicago, the University Medical Center Groningen (the Netherlands), the University of Groningen (the Netherlands), and the University of Oldenburg (Germany) outlined specifically in the Guide for the Care and Use of Laboratory Animals, the Policy on Humane Care and Use of Laboratory Animals, and Article 54 of Directive 2010/63/EU. Naked mole rats (H. glaber) and Damaraland mole rats (F. damarensis) were obtained from colonies maintained at the University of Illinois at Chicago. Mice (C57BL6 and FVB/NJ; M. musculus) were obtained from colonies maintained at the University Medical Center Groningen. Gerbils (M. unguiculatus) were obtained from colonies maintained at the University of Oldenburg. A total of 29 mice, 7 gerbils, 18 naked mole rats, and 4 Damaraland mole rats were used. Ages of animals are indicated in Table 1. For mice and gerbils exact ages were known and given in the text. For naked and Damaraland mole rats exact ages are given when known. Adult Damaraland mole rats were approximately 5 years of age and referred to as “mature” in the text.

For comparison, the onset of hearing in mice and gerbils occurs around 2 weeks of age but auditory maturation continues until weaning (Ehret, 1976; Finck, Schneek, & Hartman, 1972). Mice wean at 3–4 weeks of age and begin reproductive activity between 6 and 8 weeks of age (Prichett & Taft, 2007). Gerbils also wean between 3 and 5 weeks of age and reach reproductive activity at 4 months of
To obtain cochleae, animals were deeply anesthetized by either isoflurane inhalation (University of Groningen and University of Illinois at Chicago) or injection of a lethal dose of pentobarbital (University of Chicago) or injection of a lethal dose of pentobarbital (University of Oldenburg) and then rapidly decapitated. Procedures published previously were used to isolate intact organs of Corti from cochleae for immunostaining and confocal microscopy. To determine the number of afferent ribbons per HC at given ages in mice, gerbils, naked mole rats, and Damaraland mole rats.

| Species                  | Age          | Ribbons/IHC | Ribbons/OHC |
|--------------------------|--------------|-------------|-------------|
| Mouse (C57BL6)           | P6           | 41.5 ± 1.7  | 22.2 ± 0.9  |
|                          | P9           | 33.7 ± 0.7  | 18.7 ± 0.8  |
|                          | P12          | 19.5 ± 0.6  | 2.5 ± 0.03  |
|                          | P15          | 16.4 ± 1.1  | 2.3 ± 0.08  |
|                          | P21          | 17.7 ± 0.5  | 2.3 ± 0.08  |
|                          | P42 (6 weeks)| 17.0 ± 0.7  | 2.3 ± 0.04  |
| Gerbil                   | P9           | 41.0 ± 3.5  | 16.8 ± 0.9  |
|                          | P42 (6 weeks)| 25.2 ± 2.2  | 2.8 ± 0.08  |
| Naked mole rat           | P7           | 19.3 ± 3.7  | 24.4 ± 0.09 |
|                          | P11          | 19.2 ± 1.8  | 17.6 ± 0.2  |
|                          | P14          | 19.6 ± 1.8  | 13.2 ± 0.8  |
|                          | P28          | 15.5 ± 1.8  | 9.7 ± 0.8   |
|                          | P182 (6 months) | 12.9 ± 0.5 | 6.9 ± 0.2   |
|                          | P365 (1 year)| 13.6 ± 1.2  | 7.1 ± 0.7   |
| Damaraland mole rat      | P5           | 16.26 ± 1.1 | 12.0 ± 1.0  |
|                          | Mature       | 13.9 ± 1.0  | 14.5 ± 0.9  |

Abbreviations: HC = hair cell; IHC = inner hair cell; OHC = outer hair cell. *N = total number of individuals.

2.2 | Antibody characterization, immunofluorescence, and confocal microscopy

Immunofluorescent staining was performed using mouse monoclonal (IgG1) anti-CTBP2 (RRID:AB_399431; aa 361–445 from mouse CTBP2; 612044, BD Bioscience; 1:300), rabbit polyclonal antibody against the GluR2/3 (RRID:AB_90710; EGNYVYGIESVKI from rat GluR2; AB1506; Merck Millipore; 1:300), rabbit polyclonal anti-synapsin I (RRID:AB_2200400; Synapsin Ia/b from bovine brain; AB1543; Merck Millipore; 1:500), mouse monoclonal (IgG2a) anti-synapsin Ia/b (RRID:AB_11150471; aa 2–29 from human Synapsin Ia/b N-terminus; sc-376622; Santa Cruz Biotechnology; 1:300), and rabbit polyclonal anti-BK channel (RRID:AB_2313725; aa 1097–1196 from mouse KCa1.1; APC-021; Alomone Labs; 1:500). The specificity of the primary antibodies was verified by the vendors. Secondary antibodies, AlexaFluor 488 goat anti-mouse IgG1 (RRID:AB_2534069), AlexaFluor 568 goat anti-rabbit (RRID:AB_143157), and AlexaFluor 647 goat anti-mouse IgG2a (RRID:AB_2535810), were purchased from Invitrogen and used diluted 1:500.

Low magnification micrographs of entire organs of Corti were obtained with a Leica DM 4000B fluorescence microscope. High magnification confocal micrographs of identified tonotopic regions of the organ of Corti were acquired using a Leica SP8 confocal microscope with a 60x Olympus PlanApo oil-immersion lens (NA 1.42) under the control of the Leica Application Suite software. Image stacks were collected to encompass the entire region of interest from the specified frequencies. The step size (optical section thickness) was approximately 0.3 μm and determined by stepping at half the distance of the theoretical z-axis resolution (the Nyquist sampling frequency). Images were acquired in a 1,024 × 1,024 pixel raster at a sampling speed of 100 Hz. The laser power, PMT voltage, and gain and offset adjustments were adjusted to optimize the dynamic range of the detected intensity values. Images are presented as z-projections through the collected optical stacks.

Unless specifically stated, data are collected from the 16 kHz region in mice and from the 2 kHz region in gerbils. These regions represent the frequencies of best hearing and were identified using previously determined cochlear place-frequency maps (Müller, 1996; Müller, van Hunerbein, Hoidis, & Smolders, 2005). For both mice and gerbils, these regions are approximately 50% along the length of the basilar membrane. The total basilar membrane (cochlear canal) length is approximately 6.8 mm in mice and 12.1 mm in gerbils (Kirk & Gosselin-Ildari, 2009). Because similar cochlear place-frequency maps have not been determined for either naked or Damaraland mole rats, we compared to equivalent positions (50%) along the length of the basilar membrane. The total basilar membrane length is approximately 5.8 mm in naked mole rats (Mason et al., 2016) and 10.9 mm in Fukomys micklemi (Mason et al., 2016) and 11.1 mm in Fukomys anselli (Kirk & Gosselin-Ildari, 2009), two relatives of the Damaraland mole rat.

2.3 | Image analysis

Three-dimensional (3D) reconstructions were obtained using Imaris 6.4 software (Bitplane Inc.) and used to determine the number, volume, and location of synaptic elements. To determine the number of
synaptic elements per hair cell, the Imaris spots function was used to detect immunopuncta within a given field of view. This value was then divided by the total number of hair cells within that field of view. Hair cell counts were obtained from counts of immunofluorescently detected hair cell nuclei (for CTBP2 immunolabeling). For organs of Corti from naked and Damaraland mole rats, CTBP2 immunolabeling was not restricted to the nuclei of the hair cells but also apparent in the nuclei of other cell types. In 3D reconstructions, the hair cell nuclei were identified by their generally larger size and proximity to the CTBP2-immunolabeled afferent ribbons. In addition, differential interference contrast images obtained in parallel aided identification of the hair cells. Volumes ($\mu$m$^3$) of immunopuncta were determined using the Imaris surfaces function. All numerical values and their 3D locations were exported from Imaris for further statistical analyses.

Custom script developed in R was used to calculate Euclidean distances between immunopuncta (as described previously in Sadeghi, Pyott, Yu, & Glowatzki, 2014; Ye, Goutman, Pyott, & Glowatzki, 2017) and to classify inner hair cell (IHC) afferent ribbons (CTBP2 immunopuncta) as either pillar or modiolar. Briefly, for this later analysis, $x$, $y$, and $z$ coordinates were determined for the afferent ribbons and the nuclei using the spots function in Imaris. Using principal component analysis, (eigen) vectors were fit to the coordinates defining the row of afferent ribbons and also the coordinates defining the nuclei. The minimum and maximum $x$, $y$, $z$ coordinates for the vector fitting the afferent ribbons and the mean $x$, $y$, $z$ coordinate for the vector fitting the coordinates of the nuclei determined the plane that defined the pillar-modiolar axis. The pillar and modiolar sides of this plane were determined from other landmark structures (e.g., the spiral ganglion neurons and outer hair cells [OHCs]). The $x$, $y$, $z$ coordinates of the individual afferent ribbons were then used to classify ribbons as belonging to either the pillar or modiolar side. To allow comparison across z-stacks, ribbon volumes, determined using the Imaris surfaces function, from a given stack were normalized to the median volume from that stack.

2.4 | Statistics

Statistical analyses were performed using SPSS 24. Data for independent samples were analyzed using a general linear model ANOVA. In case of repeated measurements from the same individual subject (e.g., when obtaining data from multiple hair cells in a subject) a mixed model ANOVA was applied taking the subject into account. Post hoc pairwise tests used the Bonferroni correction, which was not applied in planned comparisons. Differences were considered significant when $P < 0.05$ (*) and highly significant when $P < 0.01$ (**). Group results are reported as the mean ± SEM. In most cases, $n$ represents the number of individuals.

3 | RESULTS

3.1 | Afferent innervation in the organs of Corti from mature mice, gerbils, naked, and Damaraland mole rats

In the mammalian organ of Corti, there are two classes of sensory hair cells (Goutman, Elgoyhen, & Gomez-Casati, 2015). The IHCs are responsible for relaying information to the brain for the perception of sound and make afferent connections to the type I spiral ganglion neurons. The OHCs are responsible for generating the cochlear amplifier and make afferent connections to the type II spiral ganglion neurons. We quantified the numbers of afferent synapses in these two types of sensory hair cells in the mature organs of Corti from mice (C57BL6, 6 weeks old) gerbils (6 weeks old), naked mole rats (1 year old), and Damaraland mole rats (mature) by immunolabeling with an antibody against CTBP2, a protein enriched in the presynaptic afferent ribbons. In all animals, both classes of hair cells show CTBP2-immunoreactivity indicative of presynaptic afferent ribbons (green, Figure 1a).

To compare afferent ribbon numbers per hair cell both within and across these four species, we quantified the number of afferent ribbons (CTBP2 immunopuncta) per IHC (gray bars, Figure 1b) and per OHC (white bars, Figure 1b). Quantification shows that, when comparing within species, the number of afferent ribbons per IHC compared to OHC was significantly greater in mice, gerbils, and naked mole rats but not Damaraland mole rats. In addition, there were differences in the numbers of afferent ribbons per IHC and OHC across species. Gerbils had significantly greater numbers of ribbons per IHC compared to all other species. Damaraland mole rats had significantly greater numbers of afferent ribbons per OHC compared to all other species. Naked mole rats also had significantly greater numbers of afferent ribbons per OHC compared to both mice and gerbils. Values are provided in Table 1. Although not quantified, there were no apparent systematic differences in the numbers of afferent ribbons per OHC row in any of the four species investigated.

We also investigated postsynaptic afferent glutamate receptors by immunolabeling with a rabbit polyclonal antibody against GluR2/3. In mice and gerbils, GluR2/3 immunoreactivity is punctate and juxtaposed to presynaptic afferent ribbons. In contrast to mice and gerbils, we observed no GluR2/3 immunoreactivity in naked or Damaraland mole rats (data not shown). Postsynaptic glutamate receptor labeling was not characterized further.

We were also interested in the distribution of the IHC afferent ribbons and ribbon sizes along the pillar-modiolar axis of the IHCs (Figure 1c). In cat, pillar-modiolar differences in IHC afferent ribbon sizes correlate with pillar-modiolar differences in afferent fiber thresholds (Merchan-Perez & Liberman, 1996). Pillar-modiolar differences in IHC afferent ribbon sizes have also been documented in mice (Liberman, Wang, & Liberman, 2011) and gerbils (Zhang, Engler, Koepcke, Steenken, & Koppl, 2018). To compare differences between the numbers and volumes of pillar and modiolar IHC afferent ribbons, we localized and quantified the volumes of ribbons (CTBP2 immunopuncta) as described in Section 2. In mice, there were fewer pillar ribbons (44%) than modiolar ribbons (56%). In contrast, in gerbils, naked mole rats, and Damaraland mole rats, the fractions of pillar and modiolar ribbons were equivalent. When comparing the mean normalized volumes of the pillar (gray circles, Figure 1c) versus modiolar (white circles, Figure 1c) IHC afferent ribbons, values within species were significantly different in mice, gerbils, and naked mole rats but not Damaraland mole rats. The steepness of this gradient (i.e., the ratio between the mean volumes of the pillar versus modiolar afferent ribbons) also appeared to vary across species, with the steepest gradients observed in mice. Values are provided in Table 2.
To summarize, Damaraland mole rats are particularly unique among the rodent species examined: unlike mice, gerbils, and naked mole rats, Damaraland mole rats show the same number of afferent ribbons per hair cell in both IHCs and OHCs and lack the pillar-modiolar gradient in IHC afferent ribbons sizes observed in other rodents.

### 3.2 Tonotopic variations in afferent ribbon density and BK channel expression in the IHCs from mature mice, gerbils, and naked mole rats

In both mice and gerbils, IHC afferent ribbon density peaks where the cochlea is most sensitive to sound (i.e., the most sensitive frequency in the audiogram; Meyer et al., 2009). Thus, reduced tonotopic variation in IHC afferent ribbon density could contribute to the poor sensitivity of hearing across frequencies observed in naked mole rats (Heffner & Heffner, 1993). Therefore, we analyzed IHC afferent ribbon density along the length of the cochlear spiral. In parallel, we investigated tonotopic changes in BK channel expression. BK channels are large conductance, voltage-, and calcium-activated potassium channels that are present in both inner and OHCs in the mammalian cochlea (reviewed in Pyott & Duncan, 2016). BK channel expression in both types of hair cells shows increasing expression from apical (low frequency) to basal (high frequency) regions following the onset of hearing in a variety of animals investigated (Pyott & Duncan, 2016). Therefore, the presence of BK channels in mammalian hair cells is associated with both frequency tuning of the cochlea and functional maturation of the cochlea. For these

### FIGURE 1
Comparison of inner and outer hair cell afferent innervation in mature mice (6 weeks old), gerbils (6 weeks old), naked mole rats (1 year old), and Damaraland mole rats (mature). (a) Projections through z-stacks of confocal sections spanning the regions of the organ of Corti containing the inner hair cell (IHC) and outer hair cell (OHC) afferent ribbons immunolabeled with an antibody against CTBP2 (green). (b) Comparison of afferent ribbon density CTBP2 immunopuncta per IHC (gray bars) and OHC (white bars). (c) Comparison of the mean normalized volume of pillar (gray circles) and modiolar (white circles) afferent ribbons. Asterisks indicate significantly different values (*$P < 0.05$, **$P < 0.01$). Values are provided in Tables 1 and 2.

### TABLE 2
Normalized volume of pillar and modiolar afferent ribbons in mature mice, gerbils, naked mole rats, and Damaraland mole rats

| Species                  | Age             | Pillar afferent ribbons | Modiolar afferent ribbons |
|--------------------------|-----------------|-------------------------|---------------------------|
| Mouse (C57BL6)           | P42 (6 weeks; N = 4 individuals) | 0.95 ± 0.05 (N = 232)  | 1.77 ± 0.11 (N = 293)    |
| Gerbil                   | P42 (6 weeks; N = 3 individuals) | 1.12 ± 0.08 (N = 270)  | 1.64 ± 0.09 (N = 262)    |
| Naked mole rat           | P182 (6 months; N = 3 individuals) | 1.18 ± 0.08 (N = 142)  | 1.48 ± 0.08 (N = 143)    |
| Damaraland mole rat      | Mature (N = 3 individuals) | 1.19 ± 0.05 (N = 172)  | 1.20 ± 0.06 (N = 176)    |

$N$ = total number of individuals and afferent ribbons examined.
reasons, we were interested in whether BK channels were also present in IHCs from naked mole rats, and, if so, if BK channel expression together with afferent ribbon density varied tonotopically.

To examine afferent ribbon density and BK channel expression tonotopically in the IHCs, we immunolabeled organs of Corti isolated from mice (6 weeks old), gerbils (6 weeks old), and naked mole rats (6 months) with a mouse monoclonal antibody against CTBP2 (green, Figure 2a) and a rabbit polyclonal antibody against the BK channel (red, Figure 2a). These experiments utilized wild-type littermates of BK channel knock-out mice, which are on the FVB/NJ background and have been previously used to examine BK channel expression in the IHCs (Pyott et al., 2007; Pyott, Glowatzki, Trimmer, & Aldrich, 2004; Rohmann, Wersinger, Braude, Pyott, & Fuchs, 2015; Wersinger, McLean, Fuchs, & Pyott, 2010). For all tonotopic regions examined in these animals, BK channel immunoreactivity was punctate and clustered around the neck of the hair cells and away from the synaptic pole where afferent ribbons were localized. (This separation is not always apparent in the 2D projections through the 3D z-stacks shown in Figure 2a.)

To examine expression quantitatively, we determined the number of afferent ribbons and BK channel immunopuncta per IHC at defined tonotopic locations in mice and gerbils and in apical, middle, and basal turns in naked mole rats (Figure 2b). These values are provided in Table 3. Although the sample size was not sufficient for statistical comparisons, a few trends were visible from the data. First, in mice and gerbils, the number of afferent ribbons and BK channel immunopuncta per IHC increased steadily with frequency and then dropped again at frequencies beyond the most sensitive frequency (16 kHz in mice and 2 kHz in gerbils). Second, in both mice and gerbils, the number of BK channel immunopuncta per IHC was greater than the number of afferent ribbons per IHC for frequencies at and below the most sensitive frequency before becoming smaller than the number of afferent ribbons per IHC for frequencies above the most sensitive frequency. Third, in naked mole rats, the number of afferent ribbons and BK channel immunopuncta per IHC showed little tonotopic variation from apical to middle and then middle to basal turns, and, furthermore, the number of BK channel immunopuncta per IHC was smaller than the number of afferent ribbons per IHC for all three turns. Thus, in contrast to mice and gerbils, naked mole rats show no evidence of tonotopic variation in either IHC afferent ribbon density or BK channel expression.

3.3 Lateral and medial efferent innervation in the organs of Corti from mature mice, gerbils, naked, and Damaraland mole rats

Efferent innervation of the hair cells begins during development and persists, with significant maturational modifications, in the adult cochlea. In the mature mammalian cochlea, there are two olivocochlear efferent

FIGURE 2 Tonotopic comparison of afferent ribbon density and BK channel expression in inner hair cells from mature mice, gerbils, and naked mole rats. (a) Projections through z-stacks of confocal sections spanning the regions of the organ of Corti containing the inner hair cell (IHC) afferent ribbons immunolabeled with an antibody against CTBP2 (green) and BK channels immunolabeled with an antibody against the C-terminus (red). Mice and gerbils are 6 weeks old, naked mole rats are 6 months old, and Damaraland mole rats are mature. Tonotopic regions are indicated. (b–d) Mean afferent ribbon density (CTBP2 immunopuncta/IHC, green) and BK channel density (BK immunopuncta/IHC, red) are plotted as a function of the tonotopic position in mice (b), gerbils (c), and naked mole rats (d). Values are provided in Table 3.
To examine the postnatal elimination of afferent synapses, we immunolabeled the afferent ribbons in organs of Corti isolated from variously aged mice, gerbils, naked mole rats, and Damaraland mole rats with a mouse monoclonal antibody against CTBP2 (green, Figure 4a). Both the IHCs and OHCs from mice and gerbils showed conspicuously more afferent ribbons at younger ages before the onset of hearing compared to older ages after the onset of hearing. In contrast, the difference in numbers of IHC and OHC afferent ribbons between younger ages, during the first postnatal week, and older ages, 6 months in naked mole rats and mature Damaraland mole rats, was much less conspicuous than in mice and gerbils.

To examine the time course of postnatal afferent ribbon pruning, we quantified the number of afferent ribbons (CTBP2 immunopuncta) per hair cell at various ages in both mice (Figure 4b) and naked mole rats (Figure 4c). In mice, between the first and third postnatal week (P6 and P21), afferent ribbons per IHC were reduced approximately two-fold and afferent ribbons per OHC were reduced more dramatically, approximately ten-fold. In mice, the numbers of afferent ribbons per IHC and afferent ribbons per OHC were significantly greater at P6 compared to P9 and the numbers at both P6 and P9 were significantly greater compared to P12, P21, and P42. These data indicate that the pruning of afferent ribbons per IHC and OHC is complete by the end of the second postnatal week (P12), and the numbers of afferent ribbons per hair cell remain stable thereafter. In naked mole rats, between the first postnatal week and the first year of life (P7 to P365), afferent ribbons per IHC and OHC showed less reduction compared to mice. No significant differences were observed comparing afferent ribbons per IHC across all ages but the small sample size may have reduced the statistical power. When comparing afferent ribbons per OHC, numbers were significantly reduced from P7 to P11, P11 to P14, and P14 to P28. After P28, no significant differences were
observed. These data indicate that the pruning of afferent ribbons per OHC is complete by the end of the fourth postnatal week (P28), and the numbers of afferent ribbons per OHC remain stable thereafter. Values are provided in Table 1.

For comparison across all four species, we additionally quantified the number of afferent ribbons (CTBP2 immunopuncta) per hair cell for immature and mature gerbils (P9 and P42/6 weeks) and Damaraland mole rats (P5 and mature) compared to immature and mature mice (P7 and P21) and immature and mature naked mole rats (P7 and 1 year). When comparing within species for these two ages (and, hence, without Bonferroni correction for multiple comparisons as in Figure 4c), mice, gerbils, and naked mole rats showed significant reductions in the number of afferent ribbons per IHC (Figure 4d). Mice, gerbils, and naked mole rats also showed significant reductions in the number of afferent ribbons per OHC (Figure 4e). In contrast, no evidence of developmental pruning of either IHC or OHC afferent ribbons was observed in Damaraland mole rats (Figure 4d,e). However, because only one P5 Damaraland mole rat sample was available, no statistical comparisons were made between immature and mature Damaraland mole rats. Values are provided in Table 5. The normalization of ribbon volume used throughout our study precludes any comparison across different ages. Therefore, absolute ribbon volumes were used only for this specific developmental question.

### Developmental changes in postnatal efferent innervation in the organs of Corti from mice, gerbils, naked, and Damaraland mole rats

Observations in mice and gerbils (as well as rats and hamsters) indicate that efferent innervation of the mammalian cochlea undergoes extensive modification shortly before the onset of hearing, generally around the second postnatal week (Simmons, 2002). As part of this maturation, efferent terminals retract from the IHCs as medial efferent contacts onto the IHC afferent terminals consolidate. To assess postnatal maturation of efferent innervation in the naked mole rat and Damaraland mole rat, we immunolabeled organs of Corti isolated from

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**FIGURE 3** Comparison of lateral and medial efferent innervation in mature mice, gerbils, naked mole rats, and Damaraland mole rats. (a) Projections through z-stacks of confocal sections spanning the regions of the organ of Corti containing the lateral and medial efferent terminals immunolabeled with an antibody against synapsin (red). Mice and gerbils are 6 weeks old, naked mole rats are 1 year old, and Damaraland mole rats are mature. (b) Comparison of lateral efferent density, synapsin immunopuncta per inner hair cell (IHC). (c) Comparison of medial efferent density, synapsin immunovolume per outer hair cell (OHC). Asterisks indicate significantly different values (*P* < 0.05, **P** < 0.01). Values are provided in Table 4.
P9 and P42 (6 week old) gerbils, P11 and 1 year old naked mole rats, and P5 and mature Damaraland mole rats with a rabbit polyclonal antibody against synapsin to label presynaptic efferent terminals (red, Figure 5a) and a mouse monoclonal antibody against CTBP2 to label IHC afferent ribbons and nuclei (green, Figure 5a). In both gerbils and naked mole rats, but less obviously in Damaraland mole rats, efferent terminals underwent extensive developmental reorganization. In young gerbils and naked mole rats, efferent terminals could be seen in the regions both above and below the IHC afferent ribbons and nuclei, consistent with efferent terminals contacting the IHCs. In contrast, in older gerbils and naked mole rats, efferent terminals were largely consolidated to the region below the IHC afferent ribbons and nuclei at the base of the IHCs. (This separation is not always apparent in the 2D projections through the 3D z-stacks shown in Figure 5a). In contrast, in Damaraland mole rats, efferent terminals were largely consolidated to the region below the IHC afferent ribbons and nuclei at the base of the IHC even at the younger age.

To quantify the degree of developmental retraction, the distance between each afferent ribbon (CTBP2 immunopunctum) and its nearest efferent terminal (synapsin immunopunctum) was quantified as described previously (Sadeghi et al., 2014; Ye et al., 2017). We predicted that this distance would increase with development. Cumulative distributions plotting the Euclidean distance between each IHC afferent ribbon and its nearest efferent terminal at the younger (gray line) and older (black line) ages showed a rightward shift with development for both gerbils (Figure 5b) and naked mole rats (Figure 5c). This rightward shift indicates an increase in the distance between nearest neighbor afferent ribbon and efferent terminal pairs, consistent with the loss of medial efferent contacts onto IHCs and consolidation of lateral efferent contacts onto the IHC afferent terminals below the IHCs. In contrast, this increase in the distance between nearest neighbor afferent ribbon and efferent terminal pairs was not observed in Damaraland mole rats (Figure 5d). When comparing within species, in gerbils and naked mole rats, the mean Euclidean distances were significantly different between the two ages examined (Figure 5e–f). In contrast, the mean Euclidean distances were not obviously different between the two ages in Damaraland mole rats (Figure 5g). However, because only one P5 Damaraland mole rat sample was available, no statistical comparisons were made between immature and mature Damaraland mole rats. Values are provided in Table 6.

**TABLE 4** Efferent innervation density in mature mice, gerbils, naked mole rats, and Damaraland mole rats

| Species          | Age (N =) | Lateral efferent terminals/IHC | Medial efferent terminal volumes/OHC |
|------------------|-----------|-------------------------------|-------------------------------------|
| Mouse (C57BL6)   | 6 weeks (3) | 31.8 ± 3.8                     | 57.4 ± 5.5                           |
| Gerbil           | 6 weeks (3) | 56.1 ± 8.4                     | 48.4 ± 8.7                           |
| Naked mole rat   | 1 year (3) | 21.7 ± 3.3                     | 17.9 ± 3.4                           |
| Damaraland mole  | Mature (3)  | 42.6 ± 4.1                     | 35.8 ± 4.3                           |

Abbreviations: IHC = inner hair cell; OHC = outer hair cell. *N = total number of individuals examined.

4 | DISCUSSION

4.1 | Overview

In this study, we used immunofluorescence and quantitative image analysis to examine cochlear innervation in mature and developing naked and Damaraland mole rats compared to mice and gerbils. This work was motivated by previous findings that indicate relatively poor hearing in naked mole rats compared to other rodents (Heffner & Heffner, 1993) despite the absence of overt differences in the anatomy of the middle and inner ear in naked mole rats (Mason et al., 2016). In comparison to mice and gerbils, both naked and Damaraland mole rats showed alterations in afferent and efferent innervation as well as their patterns of refinement during development. Differences in cochlear innervation in naked and Damaraland mole rats are likely a consequence of their unusual life histories and offer important comparative and biomedical insights into the mechanisms that underly normal auditory function as well as hearing loss.

4.2 | Afferent innervation in mature naked and Damaraland mole rats

Examination of afferent innervation in organs of Corti isolated from mature naked and Damaraland mole rats revealed both similarities and differences in comparison to mice and gerbils (Figure 1). Like mice and gerbils, both naked and Damaraland mole rats had segregated inner and OHCs and both types of hair cells contained presynaptic afferent ribbons. Our calculations of the average number of afferent ribbons per IHC in mice and gerbils closely match previously published values (Meyer et al., 2009). Moreover, this work is the first to quantify the average number of afferent ribbons per OHC across these four species. In comparison to mice and gerbils, the average number of afferent ribbons per IHC was reduced in naked and Damaraland mole rats whereas the average number of afferent ribbons per OHC was greater. The ratio of afferent ribbons per IHC compared to OHC is approximately 10:1 in mice and gerbils, 2:1 in naked mole rats, and 1:1 in Damaraland mole rats. Moreover, in contrast to both mice and gerbils, naked mole rats showed no tonotopic variation in the IHC afferent ribbon density (Figure 2).

These afferent ribbons are the presynaptic components of synapses formed with the auditory neurons. Based on extensive research, it is widely accepted that, in mammals, the type I auditory neurons are more abundant and contact single IHCs, whereas the much less abundant type II auditory neurons contact multiple OHCs (Spoendlin, 1972). Thus, the altered ratio of inner and OHC afferent innervation in naked and Damaraland mole rats could reflect a greater proportion of type II auditory neurons and perhaps also a greater number of contacts between the type II auditory neurons and OHCs compared to other mammals. Future work examining the relative abundance of types I and II auditory neurons and their contacts with the hair cells in naked and Damaraland mole rats would provide insight into the generalizability of the pattern of types I and II afferent innervation across mammals.

In addition to quantifying the number of afferent ribbons per hair cell, we also examined differences in the distributions and sizes (volumes) of afferent ribbons between the pillar and modiolar sides...
of the IHCs. When comparing the relative numbers of modiolar versus pillar afferent ribbons, we observed equivalent numbers of modiolar and pillar synapses in gerbils as well as naked and Damaraland mole rats. In contrast, we observed more modiolar than pillar synapses (56% compared to 44%) in mice, similar to previous reports (Liberman et al., 2011). However, these differences are minor and similar to the differences reported in different studies in the same species. For example, a previous study investigating gerbils reported 57% modiolar ribbons (Zhang et al., 2018) compared to our observation of 50%. Thus, these differences could easily result from subtle differences in the methods of defining the spatial coordinates of ribbons relative to the hair cells, and, therefore, further significance to these differences is not attributed.

When comparing the mean normalized volumes of modiolar versus pillar afferent ribbons, we observed statistically significant differences in mice, gerbils, and naked but not Damaraland mole rats. A difference
TABLE 5  Volume of inner hair cell ribbons at given developmental ages in mice, gerbils, naked mole rats, and Damaraland mole rats

| Species         | Age       | Volume (μm²) |
|-----------------|-----------|--------------|
| Mouse (C57BL6)  | P9        | 0.354 ± 0.006 (N = 2,270, 3 individuals) |
|                 | P21       | 0.551 ± 0.014 (N = 1,187, 3 individuals) |
| Gerbil          | P9        | 0.158 ± 0.005 (N = 1,920, 3 individuals) |
|                 | P42 (6 weeks) | 0.297 ± 0.009 (N = 1,301, 3 individuals) |
| Naked mole rat  | P7        | 0.303 ± 0.014 (N = 886, 3 individuals) |
|                 | P182 (6 months) | 0.601 ± 0.027 (N = 450, 3 individuals) |
| Damaraland mole rat | P5 | 0.190 ± 0.015 (N = 274, 1 individual) |
|                 | Mature    | 0.338 ± 0.011 (N = 363, 3 individuals) |

*N = total number of afferent ribbons and individuals examined.

In size in the same direction, relatively larger modiolar afferent ribbons, has been reported previously in cats (Merchan-Perez & Liberman, 1996), mice (Liberman et al., 2011), and gerbils (Zhang et al., 2018). Furthermore, in cats, the type I afferent fibers with high thresholds and low spontaneous rates tended to associate with relatively larger ribbons on the modiolar side of the IHC whereas afferent fibers with low thresholds and high spontaneous rates tended to associate with relatively smaller ribbons on the pillar side of the IHC (Merchan-Perez & Liberman, 1996). The presence of the same spatial gradient in ribbon volumes in naked mole rats suggests that this species may also display a similar distribution of type I afferent subtypes. Future work characterizing auditory responses, and especially the responses of single auditory nerve fibers, in naked and Damaraland mole rats would be necessary to definitively correlate the distribution of IHC ribbon volumes to afferent fiber physiology.

The significance of the increased afferent innervation of OHCs in naked and Damaraland mole rats compared to mice and gerbils is not clear because the function of afferent innervation of the OHCs is poorly understood. Recent in vitro experiments indicate that OHCs release vesicles infrequently and their postsynaptic effect is relatively small (Weisz, Glowatzki, & Fuchs, 2009), consistent with earlier in vivo recordings suggest that the type II auditory neurons respond only to the loudest sounds or not at all (Brown, 1994; Robertson, 1984; Robertson, Sellick, & Patuzzi, 1999). Increased afferent innervation of the OHCs in naked and Damaraland mole rats would suggest that the type II auditory neurons respond more robustly to sound compared to other rodents. Enhanced activation of type II auditory neurons might, therefore, contribute to the detection and transmission of sound stimuli centrally. The type II, like the type I, central axons project to the cochlear nucleus, where they bifurcate into ascending and descending branches (Brown, Berglund, Kiang, & Ryugo, 1988). Alternatively, or in addition, enhanced activation of type II auditory neurons might contribute peripherally to modulate the local neuronal circuitry of the OHCs, for example, via reciprocal synapses (Thiers, Nadol, & Liberman, 2008) that may play a role in regulating OHC electromotility. Previous observations of poor cochlear tuning in a related mole rat F. anselli (Kössl et al., 1996), suggest that the increased OHC afferent innervation (observed in this work) in naked and Damaraland mole rats would not contribute to enhanced cochlear tuning but might even serve to broaden it. Future work examining the structural and functional differences in OHC afferent innervation between naked and Damaraland mole rats and other mammals would help clarify the as yet undetermined role of OHC afferent innervation by type II auditory neurons.

4.3 | BK channel expression in naked mole rats

Because BK channels in IHCs show variations in their tonotopic distribution along the length of the cochlea and are also associated with the functional maturation of the cochlea, we investigated their expression and tonotopic distribution in naked mole rats (Figure 2). (Additional Damaraland mole rat samples were not available for tonotopic investigation.) Although BK channel immunoreactivity was present in IHCs from naked mole rats, the number and intensity of BK channel immunopuncta per IHC was reduced compared to both mice and gerbils. Furthermore, BK channel expression in IHCs varied tonotopically in both mice and gerbils, peaking in expression density at their respective most sensitive frequencies in the audiogram, 16 kHz in mice and 2 kHz in gerbils and then falling off sharply at higher frequencies. BK channel expression in IHCs was particularly enriched up to 2 kHz in gerbils. In contrast, BK channel expression in IHCs showed no tonotopic variation across apical, middle, and basal turns in naked mole rats. Correspondingly, audiograms of naked mole rats show no particular sensitivity across their range of audible frequencies (Heffner & Heffner, 1993). These findings suggest that BK channel density in the IHCs may contribute to differential sensitivity across frequencies in the cochlea in ways that require further investigation. Finally, BK channels are also expressed in the OHCs of mice (Engel et al., 2006; Maison, Pyott, Meredith, & Liberman, 2013; Rohmann et al., 2015; Wersinger et al., 2010). In this study, BK channels were not detected in the OHCs of naked mole rats. However, we have previously reported that BK channel immunoreactivity in OHCs is sensitive to the duration of fixation (Wersinger et al., 2010). Therefore, the absence of BK channel expression in OHCs in naked mole rats should be investigated more carefully in future experiments.

4.4 | Efferent innervation in mature naked and Damaraland mole rats

In addition to examining patterns of afferent innervation, we also examined lateral and medial efferent innervation in organs of Corti isolated from mature naked and Damaraland mole rats (Figure 3). All species showed lateral efferent innervation in the region below the IHCs (Figure 3 upper panels). Although the diffuse pattern of innervation made quantification of efferent density difficult, we did detect variations in lateral efferent density among the four species investigated and significantly reduced lateral efferent innervation in naked mole rats compared to both Damaraland mole rats and gerbils. The functional consequences of this reduced lateral efferent innervation are difficult to predict because the role of the lateral efferent system is not well understood. Lateral efferent innervation has been proposed to facilitate binaural hearing (Darrow, Maison, & Liberman, 2006; Irving, Moore, Liberman, & Sumner, 2011), modulate afferent excitability (Groff & Liberman, 2003), and protect the cochlea from noise-induced injury (Darrow, Maison, & Liberman, 2007). Finally, circuits within the lateral efferent system employ various neurotransmitters and show differences...
in their patterns of origination within the brain and termination within
the cochlea (reviewed in Reijntjes & Pyott, 2016). Therefore, future
work should additionally examine possible differences in the neurotransmitters and patterns of innervation employed by the lateral effer-
ent system in naked and Damaraland mole rats. Differences in these
patterns may permit a comparative approach to identify the functional
contributions of neurotransmitter-specific lateral efferent circuits.

In addition to lateral efferent innervation, all species examined
showed medial efferent innervation of the OHCs (Figure 3 lower
panels). Although we did observe medial efferent terminals in naked
and Damaraland mole rats, these terminals were smaller and less dis-
tinct than the medial efferent terminals observed in mice and gerbils.
The diffuse pattern of innervation made quantification of the efferent
density difficult; however, we did detect significantly reduced medial
efferent innervation density in naked mole rats compared to all other
species, including Damaraland mole rats. In comparison to lateral effer-
ent innervation, the role of medial efferent innervation is much better
understood. Specifically, medial efferent innervation provides inhibitory
input to the OHCs and, thereby, regulates the cochlear amplifier. In this
way, medial efferent innervation is believed to improve signal transduc-
tion, enhance signal detection, and protect the cochlea from noise-
induced injury (reviewed in both Guinan, 2018; Lopez-Poveda, 2018).
Thus, our findings of reduced medial efferent innervation density are
consistent with previous reports of reduced cochlear tuning in a related

FIGURE 5  Comparison of efferent development in gerbils, naked mole rats, and Damaraland mole rats. (a) Projections through z-stacks of
c Confocal sections spanning the regions of the organ of Corti containing the inner hair cell (IHC) and outer hair cell (OHC) afferent ribbons
immunolabeled with an antibody against CTBP2 (green) and efferent terminals immunolabeled with an antibody against synapsin (red).
Developmental ages are indicated. Note that medial efferent terminals on OHC are not included within the z-range shown. (b–d) Cumulative
histograms plotting the Euclidean distance between each CTBP2 immunopunctum and its nearest synapsin immunopunctum in immature (gray
lines) and mature (black lines) gerbils, naked mole rats, and Damaraland mole rats. (e–g) Mean Euclidean distance between each CTBP2
immunopunctum and its nearest synapsin immunopunctum in immature (gray bars) and mature (white bars) gerbils, naked mole rats, and
Damaraland mole rats. Asterisks indicate significantly different values (**P < 0.01). Values are provided in Table 6
mole rat species, *F. anselli* (Kössl et al., 1996). The observation of medial efferent innervation in the two mole rats examined in this study, however, contrasts previous ultrastructural observations in another mole rat species, *Spalax ehrenbergii*, in which no efferent innervation of the OHCs was observed (Raphael, Lenoir, Wroblewski, & Pujol, 1991). Thus, the patterns of medial efferent innervation appear to show considerable variation among subterranean rodents.

Finally, although the organization of the hair cells was not specifically investigated, the distribution of IHC afferent ribbons (Figure 1a upper panels) and lateral efferent terminals (Figure 3a upper panels) in naked and Damaraland mole rats indicate one row of IHC, like mice and gerbils. The clustered organization of OHC afferent ribbons (Figure 1a lower panels) in naked mole rats indicates that, at least in middle turns, there are three rows of OHCs, like mice and gerbils, although the medial efferent terminals (Figure 3a lower panels) are much less organized into distinct rows. In contrast, both the OHC afferent ribbons and efferent terminals in Damaraland mole rats were much less organized, at least in middle turns, and suggest three to four rows of OHCs. Precise organization of the OHCs and their hair bundles is necessary for cochlear amplification (LeMasurier & Gillespie, 2005). Therefore, future work in naked and Damaraland mole rats should specifically examine the organization of the OHCs and their bundles as part of physiological assessment of their function.

**4.5 Development of afferent and efferent innervation in the naked and Damaraland mole rats**

In mammals, afferent and efferent innervation of the cochlea undergo extensive modification during development. We examined changes in innervation at ages known or suspected to be before and after the onset of hearing. In this way, we sought to identify comparative differences in the patterns of maturation that might result in functional differences in mature naked and Damaraland mole rats and to obtain possible correlates for the onset of hearing in these animals to guide future experiments. We examined two specific indicators of postnatal maturation: pruning of IHC and OHC afferent ribbons (Figure 4) and loss of efferent innervation of the IHCs (Figure 5).

Pruning of afferent synapses in the developing organ of Corti is analogous to the pruning of supernumerary synapses that occurs elsewhere as a classic feature of nervous system development. This pruning is qualitatively and quantitatively observable in mice and gerbils (Figure 4). In contrast, naked and Damaraland mole rats undergo reduced postnatal pruning of IHC and OHC afferent ribbons. In fact, IHC afferent ribbons are not significantly reduced in Damaraland mole rats. Although we cannot exclude the possibility that pruning occurs prenatally, these findings indicate only subtle postnatal changes in IHC afferent innervation in naked and Damaraland mole rats. Nonetheless, the volume of individual IHC afferent ribbons were significantly increased postnatally in all four species examined. Developmental changes in ribbon numbers and volumes are consistent with previous observations in developing hair cells (Sobkowicz et al., 1982; Wong et al., 2014), which have led to the suggestion that smaller and more numerous ribbons in immature cells merge to form larger and less numerous ribbons in mature cells. Our findings in Damaraland mole rats, in which there is a developmental increase in ribbon size without a change in total ribbon number, suggest that the developmental increase in IHC afferent ribbon volumes occurs independently from the developmental pruning of IHC afferent ribbons.

We also examined postnatal pruning of OHC afferent ribbons. As in mice and gerbils, OHC afferent ribbons were significantly reduced in naked mole rats. In contrast, Damaraland mole rats showed similar numbers of OHC afferent ribbons at both the immature and mature ages. In our data, OHC afferent ribbon pruning is much more prolonged in naked mole rats but nevertheless certainly complete between P28 and 6 months of age. This prolonged period of maturation compared to mice and gerbils may reflect the prolonged retention of an immature pattern of afferent innervation into adulthood, paralleling observations in the naked mole rat brain (Penz et al., 2015).

As a second indicator of postnatal maturation, we examined retraction of efferent terminals from the IHCs (Figure 5). This retraction results from the loss of medial efferent terminals directly contacting the IHCs and consolidation of lateral efferent terminals beneath the IHCs. We quantified retraction as a developmental increase in the distance between the IHC afferent terminals (located at the base of the IHCs) and their nearest neighboring efferent terminals. As validation of this methodological approach, we observed significantly greater distances between the IHC afferent terminals and their nearest neighboring efferent terminals in mature compared to immature gerbil. The same was also observed in mature compared to immature naked mole rats. In contrast, there was no evidence of retraction in Damaraland mole rats. In fact, the mean distances suggest that, in Damaraland mole rats, efferent innervation is already consolidated early in postnatal development or that there is simply no postnatal axosomatic efferent innervation of the IHCs. Again, because only one immature (P5) Damaraland mole rat specimen was available, these conclusions are made cautiously. Transient efferent innervation of the IHCs likely serves to shape activity-dependent maturation of the auditory system (Clause et al., 2014). Future work would be necessary to determine how changes in the time course or pattern of this efferent innervation contribute to maturation of the auditory system, and specifically maturation of afferent innervation, in naked and Damaraland mole rats.

**5 Conclusion**

For comparative biologists, this work establishes naked and Damaraland mole rats as species worthy of future investigation to examine cochlear mechanisms that enhance frequency sensitivity.
and sound localization, maturation of the auditory system, and also evolutionary adaptations occurring in response to subterranean environments. For biomedical researchers, naked mole rats, which are extremely long-lived and highly resistant to various diseases of aging, may also prove particularly valuable to identify mechanisms that prevent or reduce age-related hearing loss, which involves loss of cochlear function and is one of the most prevalent chronic conditions of aging in humans (Gates & Mills, 2005).

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REFERENCES

Allen, P. D., & Ison, J. R. (2010). Sensitivity of the mouse to changes in azimuthal sound location: Angular separation, spectral composition, and sound level. Behavioral Neuroscience, 124(2), 265–277.

Beebe, N. L., & Schofield, B. R. (2018). Perineuronal nets in subcortical auditory nuclei of four rodent species with differing hearing ranges. The Journal of Comparative Neurology, 526(6), 972–989.

Braude, J. P., Vijayakumar, S., Baumgarter, K., Laurine, R., Jones, T. A., Jones, S. M., & Pyott, S. J. (2015). Deletion of Shank1 has minimal effects on the molecular composition and function of glutamatergic afferent postsynapses in the mouse inner ear. Hearing Research, 321, 52–64.

Brown, M. C. (1994). Antidromic responses of single units from the spiral ganglion. Journal of Neurophysiology, 71(5), 1835–1847.

Brown, M. C., Berglund, A. M., Kiang, N. Y., & Ryugo, D. K. (1988). Central back balances interaural sensitivity. The Journal of Comparative Neurology, 278(4), 581–590.

Clause, A., Kim, G., Sonntag, M., Weisz, C. J., Vetter, D. E., Rubsamen, R., & Kandler, K. (2014). The precise temporal pattern of prehearing spontaneous activity is necessary for tonotopic map refinement. Neuron, 82(4), 822–835.

Darrow, K. N., Maison, S. F., & Liberman, M. C. (2006). Cochlear efferent feedback balances interaural sensitivity. Nature Neuroscience, 9(12), 1474–1476.

Darrow, K. N., Maison, S. F., & Liberman, M. C. (2007). Selective removal of lateral olivocochlear efferents increases vulnerability to acute acoustic injury. Journal of Neurophysiology, 97(2), 1775–1785.

Edrey, Y. H., Hanes, M., Pinto, M., Mele, J., & Buffenstein, R. (2011). Successful aging and sustained good health in the naked mole rat: A long-lived mammalian model for biogerontology and biomedical research. JAR Journal, 52(3), 41–53.

Ehret, G. (1976). Development of absolute auditory thresholds in the house mouse (Mus musculus). Journal of the American Audiology Society, 1(5), 179–184.

Elwood, R. W. (1975). Paternal and maternal behaviour in the Mongolian gerbil. Animal Behaviour, 23(4), 766–772.

Engel, J., Braig, C., Ruttiger, L., Kuhn, S., Zimmermann, U., Blin, N., ... Knipper, M. (2006). Two classes of outer hair cells along the tonotopic axis of the cochlea. Neuroscience, 143(3), 837–849.

Finck, A., Schneck, C. D., & Hartman, A. F. (1972). Development of cochlear function in the neonate Mongolian gerbil (Meriones unguiculatus). Journal of Comparative and Physiological Psychology, 78(3), 375–380.

Gates, G. A., & Mills, J. H. (2005). Presbycusis. Lancet, 366(9491), 1111–1120.

Gerhardt, P., Henning, Y., Begall, S., & Malkemper, E. P. (2017). Audiograms of three subterranean rodent species (genus Fukomys) determined by auditory brainstem responses reveal extremely poor high-frequency cutoffs. The Journal of Experimental Biology, 220(Pt 23), 4377–4382.

Gessele, N., Garcia-Pino, E., Omerbasic, D., Park, T. J., & Koch, U. (2016). Structural changes and lack of HCN1 channels in the binaural auditory brainstem of the naked mole-rat (Heterocephalus glaber). PLoS One, 11(1), e0146428.

Goutman, J. D., Elgoeyhen, A. B., & Gomez-Casati, M. E. (2015). Cochlear hair cells: The sound-sensing machines. FEBs Letters, 589(22), 3354–3361.

Groff, J. A., & Liberman, M. C. (2003). Modulation of cochlear afferent response by the lateral olivocochlear system: Activation via electrical stimulation of the inferior colliculus. Journal of Neurophysiology, 90(5), 3178–3200.

Guinan, J. J., Jr. (2006). Olivocochlear efferents: Anatomy, physiology, function, and the measurement of efferent effects in humans. Ear and Hearing, 27(6), 589–607.

Guinan, J. J., Jr. (2018). Olivocochlear efferents: Their action, effects, measurement and uses, and the impact of the new conception of cochlear mechanical responses. Hearing Research, 362, 38–47.

Heffner, R. S., & Heffner, H. E. (1993). Degenerate hearing and sound localization in naked mole rats (Heterocephalus glaber), with an overview of central auditory structures. The Journal of Comparative Neurology, 331(3), 418–433.

Heffner, R. S., Koay, G., & Heffner, H. E. (2001). Audiograms of five species of rodents: Implications for the evolution of hearing and the perception of pitch. Hearing Research, 157(1–2), 138–152.

Hetling, J. R., Baig-Silva, M. S., Comer, C. M., Pardue, M. T., Samaan, D. Y., Qtaishat, N. M., ... Park, T. J. (2005). Features of visual function in the naked mole-rat Heterocephalus glaber. Journal of Comparative Physiology. A. Neuroethology, Sensory, Neural, and Behavioral Physiology, 191(4), 317–330.

Irving, S., Moore, D. R., Liberman, M. C., & Sumner, C. J. (2011). Olivocochlear efferent control in sound localization and experience-dependent learning. The Journal of Neuroscience, 31(7), 2493–2501.

Jarvis, J. U., O’Riain, M. J., Bennett, N. C., & Sherman, P. W. (1994). Mammalian eusociality: A family affair. Trends in Ecology & Evolution, 9(2), 47–51.

Khurana, S., Liu, Z., Lewis, A. S., Rosa, K., Chetkovich, D., & Golding, N. L. (2012). An essential role for modulation of hyperpolarization-activated current in the development of binaural temporal precision. The Journal of Neuroscience, 32(8), 2814–2823.

Kirk, E. C., & Gosselin-Ildari, A. D. (2009). Cochlear labyrinth volume and hearing abilities in primates. The Anatomical Record (Hoboken), 292(6), 765–776.

Koch, U., Braun, M., Kasper, C., & Grothe, B. (2004). Distribution of HCN1 and HCN2 in rat auditory brainstem nuclei. The European Journal of Neuroscience, 2001, 79–91.

Kössl, M., Frank, G., Burda, H., & Muller, M. (1996). Acoustic distortion products from the cochlea of the blind African mole rat, Cryptomys spec. Journal of Comparative Physiology A, 178(3), 427–434.

 Larson, J., & Park, T. J. (2009). Extreme hypoxia tolerance of naked mole-rat brain. Neoreport, 2018(3), 1634–1637.

LeMasurier, M., & Gillespie, P. G. (2005). Hair-cell mechanotransduction and cochlear amplification. Neuron, 48(3), 403–415.

Liberman, L. D., Wang, H., & Liberman, M. C. (2011). Opposing gradients of ribbon size and AMPA receptor expression underlie sensitivity differences among cochlear-nerve/hair-cell synapses. The Journal of Neuroscience, 31(3), 801–808.

Lopez-Poveda, E. A. (2018). Olivocochlear Efferents in animals and humans: From anatomy to clinical relevance. Frontiers in Neurology, 9, 197.

Maison, S. F., Pyott, S. J., Meredith, A. L., & Liberman, M. C. (2013). Olivocochlear suppression of outer hair cells in vivo: Evidence for combined action of BK and SK2 channels throughout the cochlea. Journal of Neurophysiology, 109(6), 1525–1534.

Mason, M. J., Cornwall, H. L., & Smith, E. S. (2016). Ear structures of the naked mole-rat, Heterocephalus glaber, and its relatives (Rodentia: Bathyergidae). PLoS One, 11(12), e0167079.

McLean, W. J., Smith, K. A., Glowatzki, E., & Pyott, S. J. (2009). Distribution of the NaK-ATPase alpha subunit in the rat spiral ganglion and organ of corti. Journal of the Association for Research in Otologyngology, 10(1), 37–49.

Merchant-Perez, A., & Liberman, M. C. (1996). Ultrastructural differences among afferent synapses on cochlear hair cells: Correlations with auditory brainstem responses reveal extremely poor high-frequency cut-offs. The Journal of Experimental Biology, 220(Pt 23), 4377–4382.
spontaneous discharge rate. The Journal of Comparative Neurology, 371(2), 208–221.
Meyer, A. C., Frank, T., Khinich, D., Hoch, G., Riedel, D., Chapochnikov, N. M., ... Moser, T. (2009). Tuning of synapse number, structure and function in the cochlea. Nature Neuroscience, 12(4), 444–453.
Müller, M. (1996). The cochlear place-frequency map of the adult and developing Mongolian gerbil. Hearing Research, 94(1–2), 148–156.
Müller, M., von Humerbein, K., Hoidis, S., & Smolders, J. W. (2005). A physiological place-frequency map of the cochlea in the CBA/J mouse. Hearing Research, 202(1–2), 63–73.
Okanoya, K., Yosida, S., Barone, C. M., Applegate, D. T., Brittan-Powell, E. F., Dooling, R. J., & Park, T. J. (2018). Auditory-vocal coupling in the naked mole-rat, a mammal with poor auditory thresholds. Journal of Comparative Physiological A, 204, 905–914.
O’Neil, J. N., Connelly, C. J., Limb, C. J., & Ryugo, D. K. (2011). Synaptic morphology and the influence of auditory experience. Hearing Research, 279(1–2), 118–130.
Park, T. J., Reznick, J., Peterson, B. L., Blass, G., Omerbasic, D., Bennett, N. C., ... Lewin, G. R. (2017). Fructose-driven glycolysis supports anoxia resistance in the naked mole-rat. Science, 356(6335), 307–311.
Penz, O. K., Fuzik, J., Kurek, A. B., Romanov, R., Larson, J., Park, T. J., ... Keimpema, E. (2015). Protracted brain development in a rodent model of extreme longevity. Scientific Reports, 5, 11592.
Prichett, K. R., & Taft, R. A. (2007). Reproductive biology of the mouse. In J. G. Fox, M. T. Davisson, F. W. Quimby, S. W. Barthold, C. E. Newcomer, & A. L. Smith (Eds.), The mouse in biomedical research: Normative biology, husbandry and models (Vol. 3, pp. 91–121). Amsterdam, the Netherlands: Academic Press.
Pyott, S. J., & Duncan, R. K. (2016). BK channels in the vertebrate inner ear. International Review of Neurobiology, 128, 369–399.
Pyott, S. J., Glowatzki, E., Trimmer, J. S., & Aldrich, R. W. (2004). Extrasynaptic localization of inactivating calcium-activated potassium channels in mouse inner hair cells. The Journal of Neuroscience, 24(43), 9469–9474.
Pyott, S. J., Meredith, A. L., Fodor, A. A., Vazquez, A. E., Yamoh, E. N., & Aldrich, R. W. (2007). Cochlear function in mice lacking the BK channel alpha, beta1, or beta4 subunits. The Journal of Biological Chemistry, 282(5), 3312–3324.
Raphael, Y., Lenoir, M., Wroblewski, R., & Pujol, R. (1991). The sensory epithelium and its innervation in the mole rat cocholea. The Journal of Comparative Neurology, 314(2), 367–382.
Reijntjes, D. O. J., & Pyott, S. J. (2016). The afferent signaling complex: Regulation of type I spiral ganglion neuron responses in the auditory periphery. Hearing Research, 336, 1–16.
Robertson, D. (1984). Horseradish peroxidase injection of physiologically characterized afferent and efferent neurones in the Guinea pig spiral ganglion. Hearing Research, 15(2), 113–121.
Robertson, D., Sellick, P. M., & Patuzzi, R. (1999). The continuing search for outer hair cell afferents in the Guinea pig spiral ganglion. Hearing Research, 136(1–2), 151–158.
Rohmann, K. N., Wersinger, E., Braude, J. P., Pyott, S. J., & Fuchs, P. A. (2015). Activation of BK and SK channels by efferent synapses on outer hair cells in high-frequency regions of the rodent cochlea. The Journal of Neuroscience, 35(5), 1821–1830.
Ryan, A. (1976). Hearing sensitivity of the mongolian gerbil. Meriones unguiculatus. The Journal of the Acoustical Society of America, 59(5), 1222–1226.
Sadeghi, S. G., Pyott, S. J., Yu, Z., & Glowatzki, E. (2014). Glutamatergic signaling at the vestibular hair cell calyx synapse. The Journal of Neuroscience, 34(44), 14536–14550.
Simmons, D. D. (2002). Development of the inner ear efferent system across vertebrate species. Journal of Neurobiology, 53(2), 228–250.
Skulachev, V. P., Holtze, S., Vyssokikh, M. Y., Bakeeva, L. E., Skulachev, M. V., Markov, A. V., ... Sadovnichii, V. A. (2017). Neoteny, prolongation of youth: From naked mole rats to “naked apes” (humans). Physiological Reviews, 97(2), 699–720.
Sobkowicz, H. M., Rose, J. E., Scott, G. E., & Slapnick, S. M. (1982). Ribbon synapses in the developing intact and cultured organ of Corti in the mouse. The Journal of Neuroscience, 2(7), 942–957.
Spoendlin, H. (1972). Inervation densities of the cochlea. Acta Oto-Laryngologica, 73(2), 235–248.
Thiers, F. A., Nadol, J. B., Jr., & Liberman, M. C. (2008). Reciprocal synapses between outer hair cells and their afferent terminals: Evidence for a local neural network in the mammalian cochlea. Journal of the Association for Research in Otolaryngology, 9(4), 477–489.
Tolnai, S., Beutelmann, R., & Klump, G. M. (2017). Exploring binaural hearing in gerbils (Meriones unguiculatus) using virtual headphones. PLoS One, 12(4), e0175142.
Weisz, C., Glowatzki, E., & Fuchs, P. (2009). The postsynaptic function of type II cochlear afferents. Nature, 461(7267), 1126–1129.
Wersinger, E., McLean, W. J., Fuchs, P. A., & Pyott, S. J. (2010). BK channels mediate cholinergic inhibition of high frequency cochlear hair cells. PLoS One, 5(11), e13836.
Wong, A. B., Rutherford, M. A., Gabrielsaitis, M., Pangnisc, T., Gottfert, F., Frank, T., ... Moser, T. (2014). Developmental refinement of hair cell synapses tightens the coupling of Ca2+ influx to exocytosis. The EMBO Journal, 33(3), 247–264.
Ye, Z., Goutman, J. D., Pyott, S. J., & Glowatzki, E. (2017). mGluR1 enhances efferent inhibition of inner hair cells in the developing rat cochlea. The Journal of Physiology, 595(11), 3483–3495.
Yu, W. M., & Goodrich, L. V. (2014). Morphological and physiological development of auditory synapses. Hearing Research, 311, 3–16.
Zhang, L., Engler, S., Koepcke, L., Steenken, F., & Koppl, C. (2018). Concurrent gradients of ribbon volume and AMPA-receptor patch volume in cochlear afferent synapses on gerbil inner hair cells. Hearing Research, 364, 81–89.

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