PROCEEDINGS B

rspb.royalsocietypublishing.org

Research



Cite this article: Womack MC, Christensen-Dalsgaard J, Coloma LA, Chaparro JC, Hoke KL. 2017 Earless toads sense low frequencies but miss the high notes. *Proc. R. Soc. B* **284**: 20171670. http://dx.doi.org/10.1098/rspb.2017.1670

Received: 24 July 2017 Accepted: 4 September 2017

Subject Category:

Evolution

Subject Areas: evolution, physiology, ecology

Keywords:

tympanic middle ear, sensory loss, hearing, vibration, auditory brainstem recordings, Bufonidae

Author for correspondence:

Molly C. Womack e-mail: mollywo@berkeley.edu

[†]Present address: Environmental Science, Policy, and Management, University of California, Berkeley, CA, USA.

Electronic supplementary material is available online at https://dx.doi.org/10.6084/m9. figshare.c.3881782.



Earless toads sense low frequencies but miss the high notes

Molly C. Womack^{1,†}, Jakob Christensen-Dalsgaard², Luis A. Coloma^{3,4}, Juan C. Chaparro^{5,6} and Kim L. Hoke¹

¹Department of Biology, Colorado State University, Fort Collins, CO, USA

²Department of Biology, University of Southern Denmark, Campusvej 55, 5230 Odense M, Denmark ³Centro Jambatu de Investigación y Conservación de Anfibios, Fundación Otonga, Giovanni Farina 566 y Baltra, San Rafael, Quito, Ecuador

⁴Universidad Regional Amazónica Ikiam, Muyuna, Tena, Ecuador

⁵Museo de Biodiversidad del Peru, Cusco, Peru

⁶Museo de Historia Natural de la Universidad Nacional de San Antonio Abad del Cusco, Cusco, Peru

(D) MCW, 0000-0002-3346-021X

Sensory losses or reductions are frequently attributed to relaxed selection. However, anuran species have lost tympanic middle ears many times, despite anurans' use of acoustic communication and the benefit of middle ears for hearing airborne sound. Here we determine whether pre-existing alternative sensory pathways enable anurans lacking tympanic middle ears (termed earless anurans) to hear airborne sound as well as eared species or to better sense vibrations in the environment. We used auditory brainstem recordings to compare hearing and vibrational sensitivity among 10 species (six eared, four earless) within the Neotropical true toad family (Bufonidae). We found that species lacking middle ears are less sensitive to highfrequency sounds, however, low-frequency hearing and vibrational sensitivity are equivalent between eared and earless species. Furthermore, extratympanic hearing sensitivity varies among earless species, highlighting potential species differences in extratympanic hearing mechanisms. We argue that ancestral bufonids may have sufficient extratympanic hearing and vibrational sensitivity such that earless lineages tolerated the loss of high frequency hearing sensitivity by adopting species-specific behavioural strategies to detect conspecifics, predators and prey.

1. Introduction

Evolutionary biologists typically attribute sensory loss or reduction to the colonization of an environment that relaxes selection on that sensory system [1,2] and/or to selection for another trait that inhibits development of sensory structures [3-8]. However, many species of frogs and toads have lost tympanic middle ears, a condition termed earlessness, without a consistent adaptive explanation. The tympanic middle ear functions in hearing airborne sound by transferring sound energy to fluid vibrations sensed by hair cells in the inner ear (reviewed in [8]). That similar tympanic hearing systems evolved independently in the major tetrapod lineages [8-11] signifies the importance of tympanic middle ears for hearing on land. Despite these benefits of tympanic hearing and fossil evidence that the middle ear evolved in the latest common ancestor of all extant anurans [12-16], at least 38 lineages within the order Anura secondarily lost middle ears [17]. These independently evolved (convergent) losses of middle ears are puzzling given that anurans use acoustic communication for mating, defense and territoriality (reviewed in [18]). Furthermore, earless species occupy diverse habitats, making it unlikely that earless species share common environmental selection pressures that relax selection on their tympanic middle ears [19-22]. Here we examine the hypothesis that ancestral alternative sensory pathways maintain hearing sensitivity in earless species, relaxing selection on the tympanic middle ear.





Figure 1. Species of eared and earless toads covered a range of body sizes and four ear loss events. The *x*-axis shows snout – vent length (SVL) of each individual used in the hearing study, represented by an oval. The species are arranged on the *y*-axis based on hypothesized phylogenetic relationships from molecular data (MC Womack 2017, unpublished data). Eared species are shown in orange and earless species in blue. Genus names are abbreviated: *R, Rhinella; Rha, Rhaebo; O, Osornophryne*.

Extratympanic hearing pathways do not require a tympanic middle ear and provide some degree of hearing sensitivity in all anurans [23-30]. The extent to which extratympanic mechanisms preserve airborne hearing sensitivity in anurans lacking tympanic ears is debated. Experimentally removing the tympanum of two eared frog species, Hyliola regilla (=Hyla regilla) and Dryophytes versicolor (=Hyla versicolor), decreased hearing sensitivity by 25 dB above 1 kHz [23]. Yet, closely related eared and earless species from the genus Atelopus differ in sensitivity by only 8-13 dB between 2000 and 2500 Hz [26], and similarly sized, but distantly related, H. regilla (eared) and A. chiriquiensis (earless) differ by only 5 dB in sensitivity [31]. The limited number of species tested makes it hard to generalize the extent to which ancestral extratympanic pathways transmit low and high frequency vibrations in earless anurans. Moreover, earless anurans may not all have equivalent extratympanic sensitivity. Lineage-specific hearing mechanisms may allow some species better mid-to-high frequency hearing sensitivity than others, however, few studies compare hearing among earless species [26].

In addition to preserved airborne hearing, earless anurans may have increased sensitivity to substrate vibrations through extratympanic pathways. Anurans are thought to sense substrate vibrations via the operculum, a cartilaginous disc in the oval window of the inner ear in amphibians [32-34]. Low-frequency auditory fibres are sensitive to both sound and substrate vibrations [30,35]. The middle ear and the operculum are in physical contact within the oval window of the inner ear and their movement is coupled [36]. Therefore, loss of the middle ear could allow freer movement of the operculum, increasing sensitivity to substrate vibrations in earless species. Earless species that call when very close or in direct contact with conspecifics [37,38] could communicate via substrate vibrations that travel through the ground or plants. However, no one has compared vibrational sensitivity between eared and earless species to determine whether earless species have enhanced vibrational sensitivity.

Here we determine whether alternative sensory pathways could enable conspecific communication and predator and prey detection in earless anurans by (1) uniformly preserving airborne hearing sensitivity; (2) providing lineage-specific hearing sensitivity; or (3) enhancing sensitivity to substrate vibrations. To address these hypotheses, we compared hearing and vibrational sensitivity among six eared and four earless species within the true toad family (Bufonidae). Bufonidae has the highest prevalence of ear transitions within a single family and may also have examples of ear regain, and moreover, enable comparisons of closely related species that differ in ear structures [17]. Our earless species completely lacked all middle ear structures, as verified histologically (electronic supplementary material, S1). The 10 species of bufonids used in this study encompass four independent ear loss or regain events [17] as well as a range of body sizes (figure 1). First, we compared the hearing sensitivities of eared and earless species to determine whether extratympanic hearing mechanisms buffer earless bufonids from hearing loss and hence relax selection on tympanic middle ears. Second, we considered hearing sensitivity differences among earless species to verify the existence of species-specific extratympanic hearing mechanisms. Third, we tested whether earless species have enhanced sensitivity to substrate vibrations compared with eared bufonids. Commonalities across independent transitions in ear structures inform us about sensory capabilities associated with middle ear lability in bufonids.

2. Material and methods

(a) Animal collection

Ten species of bufonids (*Rhaebo haematiticus* n = 14, *Rhinella alata* n = 10, *Rhinella arborescandens*, n = 9, *Rhinella festae* n = 4, *Rhinella leptoscelis* n = 10, *Rhinella horribilis* n = 10, *Rhinella spinu- losa* n = 11, *Rhinella tacana* n = 5, *Rhinella yunga* n = 13 and *Osornophryne guacamayo* n = 9) were collected from field sites in Andes and Amazonian forests from Ecuador and Peru (electronic supplementary material, S2). We measured the snout-vent length (SVL) of each animal using a dial caliper (31-415-3, Swiss Precision Instruments Inc., Garden Grove, CA, USA).

(b) Auditory brainstem recordings to test hearing and vibrational sensitivity

We tested the hearing and vibrational sensitivities of each species to measure sensitivity of animals to tones across a range of frequencies. All auditory brainstem recordings (ABRs) were performed at laboratories in Ecuador and Peru. We measured

model no.	model (y \sim fixed effects + (1 \mid random effects)	model purpose
1	hearing thresholds \sim frequency $+$ ear (Y/N) $+$	eared versus earless airborne
	frequency $ imes$ ear $+$ (1 species) $+$ (1 species:frequency) $+$ (1 individual)	hearing sensitivity
2	hearing thresholds \sim frequency $+$ ear (Y/N) $+$	model $1 + \text{control}$ for click
	frequency $ imes$ ear $+$ click $+$ (1 species) $+$ (1 species:frequency) $+$ (1 individual)	
3	hearing thresholds \sim frequency $+$ ear (Y/N) $+$	model $1 + \text{control}$ for SVL
	frequency $ imes$ ear $+$ SVL $+$ (1 species) $+$ (1 species:frequency) $+$ (1 individual)	
4	vibration thresholds \sim frequency $+$ ear (Y/N) $+$	eared versus earless vibration
	frequency $ imes$ ear $+$ (1 species) $+$ (1 species:frequency) $+$ (1 individual)	sensitivity
5	earless hearing thresholds \sim frequency $ imes$ species $+$ (1 $ $ individual)	interspecific differences in earless
		airborne hearing sensitivity
6	earless hearing thresholds \sim frequency $ imes$ species $+$ SVL $+$ (1 $ $ individual)	model 5 $+$ control for SVL

sensitivity to airborne sound in four bufonid species as in Womack et al. [39] and used hearing sensitivity data from Womack et al. [39] for an additional six species (adult animals only, R. haematiticus, Rhinella alata, Rhinella leptoscelis, Rhinella horribilis, Rhinella spinulosa, Rhinella tacana). Briefly, we paralysed bufonids with 0.05% succinylcholine chloride (Sigma-Aldrich, St Louis, MO, USA) at a dosage of 7.5 μ l g⁻¹. We placed three 28-gauge stainless steel electrodes (Model F-E2, GRASS Technologies, Warwick, RI, USA) subdermally and amplified (RA4PA, Tucker-Davis Technologies, Alachua, FL, USA) and measured the electrical signals generated by the auditory nerve (the VIIIth nerve). We broadcast acoustic stimuli to the bufonids using a three-inch speaker (FF85 K, Fostex, Tokyo, Japan) and a digital AC/DC amplifier (DTA-1 Class T, Dayton Audio, Springboro, OH, USA). We calibrated the amplifier using a 0.5 inch free-field microphone (46AE, G.R.A.S. Sound and Vibration A/S, Skovlytoften, Denmark) and a pistonphone (Type 42AA, G.R.A.S. Sound and Vibration A/S). We used customized software that controlled calibrated stimulus presentation and data acquisition using a mobile processor (RM2, Tucker-Davis Technologies). We presented 25-ms tone bursts at a rate of 25 Hz. Tone bursts ranged in frequency from 200 to 4000 Hz and were played in order from low to high frequency. We increased sound level in 5 dB increments and averaged response signals over 400 tone bursts. We measured background noise level within the testing chamber to verify that it was below the hearing thresholds of the animals and not interfering with the testing.

To test sensitivity to vibrational stimuli of all 10 bufonid species, we placed the bufonids on a small plastic platform connected to a mini-shaker (Type 4810, Brüel & Kjær Sound & Vibration Measurement A/S, Nærum, Denmark) within the audio chamber. We calibrated the mini-shaker with a charge accelerometer (Type 4527-C, Brüel & Kjær Sound & Vibration Measurement A/S) using customized software that controlled stimulus presentation and data acquisition using the RM2 processor. We stimulated the platform with 25-ms tone bursts, ranging in frequency from 200 to 900 Hz. Responses to each stimulus were averaged over 400 tone bursts presented at a 25 Hz rate. We measured responses to increasing amplitude tone bursts at each frequency in 5 dB increments and tested frequencies in order from low to high frequency.

During both the auditory and vibrational ABRs, we measured the response to a transient signal between every two frequencies tested. For the auditory tests, the transient was generated from a half cycle 4 kHz sinusoid at 105 dB re SPL, and for vibrational tests we used a half cycle 2 kHz sinusoid at -30 dB re 1 m s⁻². We verified that the individual's responsiveness to this signal remained stable throughout the testing session. If transient

responses dropped below 25% of the original signal, we omitted all subsequent measurements from analyses (reducing datasets for nine of the 69 total individuals in the hearing study and six of the total 76 individuals in the vibrational study). We visually determined thresholds for each frequency by finding the minimum stimulus decibel level that evoked a response signal amplitude of 0.002 mV (two times the average noise level) or greater from the auditory nerve.

(c) Testing effects of middle ear presence on hearing and vibrational sensitivity

To effectively visualize hearing differences among numerous species with overlapping hearing thresholds, we graphed smoothed splines, which fit a smoothed curve to the estimated hearing thresholds, using the package sme [40] in R [41]. To quantify the overall effect of ear presence/absence on hearing thresholds, we used linear mixed-effects models in the package lme4 [42]. We ran a model with hearing threshold as the dependent variable, ear (presence/absence), frequency (as a factor), and their interaction as fixed effects, and included species, species by frequency interaction and individual as nested random effects (table 1: Model 1). We then used Type III tests of fixed effects to determine the effect of ear presence/absence on overall hearing sensitivity and on hearing sensitivity across frequency. Finally, to examine differences in hearing sensitivity between eared and earless bufonids at each frequency, we calculated least-squares means of the hearing thresholds for eared and earless species at each frequency and estimated Tukey's honestly significant differences using the package lsmeans [43]. Because our goal was to determine overall species differences in hearing sensitivity, we did not examine patterns of sex differences among species despite the possibility that males and females may differ in hearing (as in [44-46]).

Because overall ABR signal amplitudes might influence the signal : noise ratio and hence hearing thresholds, we tested whether click response (the response to the transient signal played between every two frequencies tested) accounted for hearing sensitivity differences between eared and earless species. Therefore, we added click response level as an additional fixed effect to our hearing threshold model (table 1: Model 2). Type III fixed effects showed that the main effect of ear (presence/absence) was still significant ($F_{(1,12)} = 9.6$, p = 0.009) as was the interaction between ear (presence/absence) and frequency ($F_{(15,130.1)} = 3.5$, p < 0.001), consistent with our results from Model 1. We do not discuss this model in the main text or interpret least-squares means differences using this

4

model because click response may be confounded with overall hearing sensitivity.

Body size can also affect anuran hearing sensitivity [47,48], and the diversity of body sizes among eared and earless species in this study allowed us to verify that hearing sensitivity differences between eared and earless species were not being driven by differences in body size. We ran a model to control for the effects of body size on hearing sensitivity differences between eared and earless species by adding SVL as an additional fixed effect to our hearing threshold model (table 1: Model 3). Type III fixed effects showed that, when controlling for body size $(F_{1,41.4} = 8.5, p = 0.006)$, the main effect of ear (presence/ absence) was still significant ($F_{1,12.5} = 8.0$, p = 0.015) as was the interaction between ear (presence/absence) and frequency $(F_{15,128,4} = 3.4, p < 0.001)$, consistent with our results from Model 1. We do not interpret overall differences in hearing sensitivity between eared and earless species using this model because the species differ substantially in their ranges of body size (figure 1).

We used similar mixed-effects models to compare vibrational sensitivity between eared and earless species. We ran a model with vibrational thresholds as the dependent variable, ear (presence/absence), frequency (as a factor) and their interaction as fixed effects, including species, species by frequency interaction and individual as nested random effects (table 1: Model 4). We then calculated the effect of ear presence/absence on vibrational sensitivity thresholds using Type III tests of fixed effects. Because we found no differences between eared and earless species, we did not assess vibration sensitivity using models that controlled for body size or click response.

(d) Comparison of hearing sensitivity among earless species

We next examined species differences in airborne hearing sensitivities among earless species to test for lineage-specific adaptations in extratympanic mechanisms. We ran mixed-effects models (table 1: Models 5) in which hearing thresholds for earless species were the dependent variable, species (as a factor), frequency (as a factor) and their interaction were fixed effects, and individual was a random variable (table 1: Model 5). We checked the Type III fixed effects for species to determine whether airborne hearing sensitivity differed among earless species. We verified that hearing sensitivity differences among earless species were not being driven by differences in body size by running an additional mixed-effects model that controlled for SVL (table 1: Model 6). Type III fixed effects showed that, when controlling for body size ($F_{1,36} = 6.8$, p = 0.013), the main effect of ear (presence/absence) was still significant $(F_{3,36} = 13.6, p < 0.001)$ as was the interaction between ear (presence/absence) and frequency ($F_{44,430} = 4.4$, p < 0.001), consistent with our results from Model 5. We do not interpret overall differences in hearing sensitivity between eared and earless species using this model because the species differ substantially in their ranges of body size (figure 1). Lastly, to examine differences in hearing sensitivity among earless bufonids at each frequency, we calculated least-squares means of the hearing thresholds for each earless species at each frequency and estimated Tukey's honestly significant differences.

3. Results

(a) Hearing sensitivity differences between eared

and earless species

To determine whether earless species maintain sensitivity to airborne sound through extratympanic hearing pathways, **Table 2.** Least-squares means differences between hearing sensitivity thresholds of eared and earless species. Least-squares means differences and standard error (s.e.) rounded to the nearest dB are given for each frequency with Tukey's honest significant differences between hearing stages italic. A negative difference indicates that the eared species had a lower threshold (were more sensitive) at that frequency while a positive difference indicates that eared species had a higher hearing threshold (were less sensitive) at that frequency.

	frequency (Hz)	airborne sound sensitivity differences between eared and earless species, dB (\pm s.e.)
	200	-2 (<u>+</u> 6)
	300	0 (<u>+</u> 6)
	400	-7 (<u>+</u> 6)
	500	-6 (±6)
	700	-8 (±6)
	900	-12 (±6)
	1100	-16 (±6)*
	1300	-19 (±6)**
	1500	-20 (±6)**
	1750	-19 (±6)**
	2000	-23 (±6)***
	2250	-25 (±6)***
	2500	-24 (±6)***
	3000	-16 (±6)*
	3500	-18 (<u>+</u> 7)*
	4000	-10 (<u>+</u> 7)
Ì,	· · · · · · · · · · · · · · · · · · ·	

p* < 0.05, *p* < 0.01, ****p* < 0.001.

we compared hearing sensitivity thresholds between eared and earless species. Overall, we found that earless species were less sensitive to airborne sound compared with eared species and that hearing differences between eared and earless species varied by frequency ($F_{15,859.6} = 12.3$, p < 0.001; table 1: Model 1). At and below 900 Hz and at 4000 Hz, eared and earless species had equivalent hearing, but between 1100 and 3500 Hz, earless species were 16–25 dB less sensitive than eared species (table 2; figure 2*a*).

(b) Hearing sensitivity differences among earless species

To determine whether earless species have species-specific extratympanic hearing sensitivity, we compared hearing among earless lineages. We found differences in hearing sensitivity among earless species that varied by frequency ($F_{45,439} = 4.9$, p < 0.001, table 1: Model 5). Most differences in hearing among earless species occurred at higher frequencies (above 1100 Hz; table 3). Figure 2*b* shows the variation in hearing sensitivity among earless species across the frequency range tested.

(c) Vibrational sensitivity differences between eared and earless species

To determine whether earless species had enhanced vibrational sensitivity, we compared sensitivity to substrate



Figure 2. Airborne hearing sensitivity differs between eared and earless species and among earless species. (*a*) Average eared (orange) and earless (blue) species' airborne hearing thresholds are plotted as a function of frequency. The least-squares means thresholds are plotted with error bars showing 95% confidence intervals. (*b*) Smoothed spline estimates of the hearing thresholds of individual species are plotted as a function of frequency. Each eared species' average hearing thresholds (bold lines) and standard errors (shaded area) are represented by a unique shade of orange/red, and earless species by unique shades of blue/green. Genus names are abbreviated: *R*, *Rhinella*; *Rha*, *Rhaebo*, and *O*, *Osornophryne*. Background noise level in the testing chamber (black line) is shown in both panels.

vibrations between eared and earless species. We found no consistent differences in vibrational sensitivity ($F_{1,108.5} = 2.4$, p = 0.121; table 1: Model 4; figure 3). Furthermore, responsiveness to substrate vibrations did not vary between eared and earless species depending on frequency ($F_{5,52.5} = 1.4$, p = 0.239; table 1: Model 4; table 2).

4. Discussion

Overall, we found support that alternative sensory pathways provide sensitivity to low frequency sound in all earless species and facilitate species-specific sensitivity at high frequencies. Although earless species were generally less sensitive to sound at high frequencies (above 900 Hz), species varied in hearing sensitivity in ways that could affect conspecific communication. Additionally, earless species did not exhibit higher sensitivity to substrate vibrations than eared species. Below we discuss the implications of these results for proposed extratympanic hearing pathways and social communication in earless anurans.

(a) Extratympanic hearing may reduce stabilizing

selection on middle ear structures in anurans

Earless bufonid species have a 16-25 dB hearing sensitivity reduction at high frequencies. Earless species show no reduction in hearing sensitivity at low frequencies ($\leq 900 \text{ Hz}$), consistent with the idea that tympanic hearing is mainly important for anurans above 1 kHz and that extratympanic pathways play a larger role in hearing below 1 kHz [23].

6

Table 3. Least squares means differences among hearing sensitivity thresholds of earless species. All pairs of earless species with differences in least-squares means hearing thresholds are reported for each frequency. The species with less sensitive hearing is always to the left of the less-than symbol. Least-squares means differences and standard error (s.e.) rounded to the nearest dB are given for each pair.

frequency (Hz)	airborne sound sensitivity differences among earless species, dB (\pm s.e.)			
200	Rhinella arborescandens < 0 . guacamayo,			
300	(± 4)			
400	<i>R. arborescandens</i> $< R$. yunga, 10 (+4)			
500	NA			
700	NA			
900	R. fesate $<$ R. yunga, 21(\pm 5)			
1100	NA			
1300	0. guacamayo $<$ R. arborescandens, 15 (\pm 5)	R. festae $<$ R. yunga, 21 (\pm 5)		
1500	0. guacamayo $<$ R. arborescandens	0. guacamayo $<$ R. festae,	R. festae $<$ R. yunga,	
	20 (<u>+</u> 5)	27 (<u>+</u> 6)	15 (<u>+</u> 5)	
1750	NA			
2000	R. arborescandens $<$ R. yunga, 17 (\pm 4)			
2250	0. guacamayo $<$ R. festae, 21 (\pm 7)	R. arborescandens $<$ R. yunga 14 (\pm 4)		
2500	R. arborescandens $<$ R. yunga, 13 (\pm 4)	0. guacamayo $<$ R. yunga, 21 (\pm 5)		
3000	R. arborescandens $<$ R. yunga, 10 (\pm 4)	0. guacamayo $<$ R. yunga, 13 (\pm 5)		
3500	NA			
4000	NA			

The decreases in hearing sensitivity we describe are comparable to previous studies in anurans (discussed below) and not as dramatic as might be expected from the reduction in sensitivity in gekkonid lizards lacking middle ears (up to 62 dB) [49]. Equivalent measurements in other tetrapod groups have not been published to more broadly quantify the hearing sensitivity benefits of tympanic ear structures. As at least some vocalizations of earless bufonids have energy primarily above 900 Hz (see below), the lack of a tympanic middle ear would restrict the distance over which conspecifics acoustically communicate. Reduced hearing sensitivity might similarly limit predator or prey detection in earless lineages, although the auditory system's contribution to these functions has not been examined in bufonids. Moreover, the moderate reduction in airborne hearing sensitivity we describe may not capture all of the sensory consequences associated with earlessness, as current models of sound localization in eared anurans require the middle ear cavity and Eustachian tubes [50]. We argue that extratympanic hearing does not completely compensate for hearing reduction associated with tympanic middle ear loss in anurans, but the moderate reduction in sensitivity compared with those in lizards prompt future study to test whether anurans have more effective extratympanic hearing pathways than other terrestrial vertebrates.

Although the magnitude of hearing loss associated with earlessness in our study (16–25 dB between 900 and 4000 Hz) is less dramatic than that measured in lizards, our results are generally consistent with previous estimates of extratympanic

hearing sensitivity in other anurans. Lombard & Straughan [23] experimentally removed the tympanum in two hylid species and measured up to 25-dB reductions in sensitivity above 1 kHz [23]. Comparisons among species concluded that earless bufonids from the genus Atelopus were only 5 dB less sensitive than the eared *Hyliola regilla* (=*Hyla regilla*) [26] and 8-13 dB less sensitive than Atelopus species that have a columella but lack a tympanic membrane [31]. Although these previous estimates of hearing loss associated with earlessness within bufonids are smaller than our estimates, they are difficult to compare directly given differences in methodology and sampling design. That hylids [23] and bufonids (this study) both gain approximately 25 dB sensitivity from tympanic ears suggests that extratympanic hearing provides similar sensitivity to airborne sound in both groups. Our results thus provide no evidence that middle ears are especially labile in bufonids compared with other anurans because extratympanic hearing in this family is exceptional.

(b) Enhanced vibrational sensitivity unlikely to affect ear lability

Owing to the proximity of the operculum and columellar footplate in the oval window, loss of the middle ear could impact the free-motion of the operculum and increase sensitivity to low frequency sound sensed via vibration, either induced in the skull or in the substrate and transduced by opercular motion. Bufonids, like other anurans [30,51–53] and salamanders [54–56], are at least as sensitive to substrate



Figure 3. Eared and earless species have equivalent vibrational sensitivity. Average eared (orange) and earless (blue) species' vibrational thresholds are plotted as a function of frequency. The least-squares means thresholds are plotted with error bars showing 95% confidence intervals.

vibrations as most other vertebrates [57–63]. Substrate vibrations may thus be an important sensory modality in all batrachians. Studies in a few species have characterized the use of substrate vibrations by adult anurans for communication and feeding (e.g. [64–66]), but more extensive analyses are necessary to draw conclusions about possible fitness consequences of vibrational sensitivity. We found that eared and earless bufonid species are equally sensitive to vibration, providing no support for the hypothesis that enhanced vibrational sensitivity compensated for the reduced airborne sensitivity in animals lacking tympanic middle ears.

(c) Diverse communication strategies may enable species to tolerate reduced hearing sensitivity

Earless bufonids may adopt different communication strategies to compensate for their reduced hearing sensitivity. Broadly speaking, species could either vocalize at frequencies to which they are most sensitive, or reserve acoustic communication for short-range interactions. Earless anurans might vocalize at lower frequencies at which their extratympanic hearing is effective, a strategy adopted by R. yunga (JL Stynoski, FA Trama, FL Rizo Patrón, E Tapia, KL Hoke, 2017, unpublished data). By contrast, species might communicate at short range, either employing substrate vibrations or airborne hearing. For example, R. arborescandens has been observed calling when close to conspecifics perched on ferns and other plants that would be excellent vibrational conduits (JC Chaparro and JL Stynoski 2016, personal communication). More natural history studies that characterize how airborne hearing and substrate vibrations function in conspecific communication, predator detection and feeding across species will enable us to infer likely fitness consequences of ear loss in early eared ancestors.

5. Concluding remarks

Our results show that earless bufonids have reduced highfrequency hearing sensitivity but may rely on their sensitivity to low frequencies and/or species-specific extratympanic hearing mechanisms for conspecific communication and for predator and prey detection. These and other behavioural strategies could relax selection on tympanic middle ears within some lineages if ancestral, or compensate for ear loss if derived. To infer possible fitness consequences of reduced highfrequency hearing sensitivity in early earless ancestors, more natural history studies are needed to characterize broadly how extratympanic pathways for both airborne and substrate borne vibrations function in conspecific communication, predator detection, and feeding across species. Alternative sensory pathways, acoustic environments and signalling behaviour may jointly create unique selection regimes within different clades that enabled the widespread convergent loss of what is a vital sensory structure to most terrestrial tetrapods.

Ethics. The Institutional Animal Care and Use Committee at Colorado State University approved all experiments (IACUC Protocol #12-3484A) and the Ministerio del Ambiente in Ecuador and the Servicio Forestal y de Fauna Silvestre in Peru (SERFOR) approved collection, research and export permits (electronic supplementary material, S2). Data accessibility. The hearing and vibration datasets used in this study can be found in electronic supplementary material, S3.

Authors' contributions. M.C.W. and K.L.H. contributed to the conception of the study, data collection, data analysis and writing. J.C.-D. contributed to the ABR methodological design and data collection. L.A.C. and J.C.C. contributed to the sampling design and animal collection. All authors contributed to manuscript editing and gave final approval for publication.

Competing interests. We have no competing interests.

Funding. This study was supported by the U.S. National Science Foundation (NSF #IOS-13503461350346 and OISE-1157779) as well as the grants-in-aid-of-research fellowship from Sigma Xi (G20111015158047) and the Danish National Research Council (grant no. 1323-00132B to J.C.D.).

Acknowledgements. We thank Elicio E. Tapia (CJ), Peter Condori (MUBI) and Amanda Delgado (MHNC) who helped to obtain the wild-caught animals for this study. Special thanks to María Dolores Guarderas (CJ), Amanda Delgado and Florencia Trama (Center for Conservation and Sustainable Development, CDS, Oxapampa, Peru) for facilitating collection and transportation permits. We also thank Jennifer Stynoski for sharing her observations on conspecific communication used by earless species in this study.

- Jeffery WR. 2005 Adaptive evolution of eye degeneration in the Mexican blind cavefish. J. Hered. 96, 185–196. (doi:10.1093/jhered/esi028)
- Lehmann GU, Strauß J, Lakes-Harlan R. 2007 Listening when there is no sexual signalling? maintenance of hearing in the asexual bushcricket *Poecilimon intermedius. J. Comp. Physiol. A.* 193, 537–545. (doi:10.1007/s00359-007-0209-y)
- 3. Norris KS, Lowe CH. 1951 *A study of the osteology and musculature of* Phrynosoma mcallii *pertinent to its systematic status*. Chicago, IL: Chicago Academy of Sciences.
- Wright S. 1964 Pleiotropy in the evolution of structural reduction and of dominance. *Am. Nat.* 98, 65–69. (doi:10.1086/282301)
- Berman DS, Regal PJ. 1967 The loss of the ophidian middle ear. *Evolution* 21, 641–643. (doi:10.1111/j. 1558-5646.1967.tb03418.x)
- Nijhout HF, Emlen DJ. 1998 Competition among body parts in the development and evolution of insect morphology. *Proc. Natl Acad. Sci. USA* 95, 3685–3689. (doi:10.1073/pnas.95.7.3685)
- Yamamoto Y, Byerly MS, Jackman WR, Jeffery WR. 2009 Pleiotropic functions of embryonic sonic hedgehog expression link jaw and taste bud amplification with eye loss during cavefish evolution. *Dev. Biol.* 330, 200–211. (doi:10.1016/j. ydbio.2009.03.003)
- Manley GA. 2010 An evolutionary perspective on middle ears. *Hear. Res.* 263, 3-8. (doi:10.1016/j. heares.2010.08.012)
- Clack JA. 1997 The evolution of tetrapod ears and the fossil record. *Brain Behav. Evol.* 50, 198–212. (doi:10.1159/000113334)
- Christensen-Dalsgaard J, Carr CE. 2008 Evolution of a sensory novelty: tympanic ears and the associated neural processing. *Brain Res. Bull.* **75**, 365–370. (doi:10.1016/j.brainresbull.2007.10.044)
- Manley GA, Sienknecht UJ. 2013 The evolution and development of middle ears in land vertebrates. In *The middle ear* (eds S Puria, RR Fay, AN Popper), pp. 7–30. New York, NY: Springer.
- Rage JC, Rocek Z. 1989 Redescription of *Triadobatrachus massinoti* (Piveteau, 1936) an anuran amphibian from the early Triassic. *Palaeontographica A* 206, 1–16.
- Shubin NH, Jenkins Jr FA. 1995 An early Jurassic jumping frog. *Nature* **377**, 49. (doi:10.1038/ 377049a0)
- Gao KQ, Wang Y. 2001 Mesozoic anurans from Liaoning Province, China, and phylogenetic relationships of archaeobatrachian anuran clades. *J. Vert. Paleontol.* **21**, 460–476. (doi:10.1671/0272-4634(2001)021[0460:MAFLPC]2.0.C0;2)
- Gao KQ, Chen S. 2004 A new frog (Amphibia: Anura) from the Lower Cretaceous of western Liaoning, China. *Cretaceous Res.* 25, 761–769. (doi:10.1016/j.cretres.2004.06.011)
- 16. Báez AM, Nicoli L. 2008 A new species of *Notobatrachus* (Amphibia, Salientia) from the

Middle Jurassic of northwestern Patagonia. J. Paleontol. **82**, 372–376. (doi:10.1666/06-117.1)

- Pereyra MO *et al.* 2016 The complex evolutionary history of the tympanic middle ear in frogs and toads (Anura). *Sci. Rep.* 6, 34130. (doi:10.1038/ srep34130)
- Wells KD, Schwartz JJ. 2007 The behavioral ecology of anuran communication. In *Hearing and sound communication in amphibians* (eds P Narins, AS Feng, RR Fay), pp. 44–86. New York, NY: Springer.
- Vorobyeva E, Smirnov S. 1987 Characteristic features in the formation of anuran sound conducting systems. *J. Morphol.* **192**, 1–11. (doi:10.1002/jmor. 1051920102)
- Jaslow AP, Hetherington TE, Lombard RE. 1988 Structure and function of the amphibian middle ear. In *The evolution of the amphibian auditory system* (ed. B Fritzsch), pp. 69–91. New York, NY: Wiley.
- Smirnov SV. 1991 The anuran middle ear: developmental heterochronies and adult morphology diversification. *Belg. J. Zool.* 121, 99–110.
- Hetherington TE. 1992 The effects of body size on the evolution of the amphibian middle ear. In *The evolutionary biology of hearing* (eds DB Webster, AN Popper, RR Fay), pp. 421–437. New York, NY: Springer.
- Lombard RE, Straughan IR. 1974 Functional aspects of anuran middle ear structures. J. Exp. Biol. 61, 71-93. (doi:10.1159/000114111)
- Hetherington TE, Lombard RE. 1983 Electromyography of the opercularis muscle of *Rana catesbeiana*: an amphibian tonic muscle. *J. Morphol.* **175**, 17–26. (doi:10.1002/jmor.1051750103)
- Ehret GÜ, Keilwerth EL, Kamada TS. 1994 The lung – eardrum pathway in three treefrog and four dendrobatid frog species: some properties of sound transmission. J. Exp. Biol. 195, 329–343.
- 26. Lindquist ED, Hetherington TE, Volman, SF. 1998 Biomechanical and neurophysiological studies on audition in eared and earless harlequin frogs (*Atelopus*). *J. Compar. Physiol. A* **183**, 265–271. (doi:10.1007/s003590050254)
- Hetherington TE, Lindquist ED. 1999 Lung-based hearing in an 'earless' anuran amphibian. *J. Compar. Physiol. A* 184, 395–401. (doi:10.1007/ s003590050338)
- Wilczynski W, Resler C, Capranica RR. 1987 Tympanic and extratympanic sound transmission in the leopard frog. *J. Compar. Physiol. A* 161, 659–669. (doi:10.1007/bf00605007)
- Narins PM, Ehret G, Tautz J. 1988 Accessory pathway for sound transfer in a neotropical frog. *Proc. Natl Acad. Sci. USA* 85, 1508–1512. (doi:10. 1073/pnas.85.5.1508)
- Christensen-Dalsgaard J, Narins PM. 1993 Sound and vibration sensitivity of VIIIth nerve fibers in the frogs *Leptodactylus albilabris* and *Rana pipiens pipiens. J. Compar. Physiol. A* **172**, 653–662. (doi:10.1007/bf00195391)

- Jaslow AP, Lombard RE. 1996 Hearing in the neotropical frog, *Atelopus chiriquiensis. Copeia* 1996, 428–432. (doi:10.2307/1446859)
- Kingsbury BF, Reed HD. 1909 The columella auris in Amphibia. Second contribution. *J. Morphol.* 20, 549–628. (doi:10.1002/jmor.1050200403)
- Hetherington TE. 1985 Role of the opercularis muscle in seismic sensitivity in the bullfrog *Rana catesbeiana. J. Exp. Zool.* 235, 27–34. (doi:10.1002/ jez.1402350105)
- Hetherington TE. 1988 Biomechanics of vibration reception in the bullfrog, *Rana catesbeiana*. *J. Compar. Physiol. A* 163, 43-52. (doi:10.1007/ bf00611995)
- Christensen-Dalsgaard J, Jørgensen MB. 1996 Sound and vibration sensitivity of VIIIth nerve fibers in the grassfrog, *Rana temporaria. J. Compar. Physiol. A* 179, 437–445. (doi:10.1007/bf00192311)
- Mason, MJ, Narins, PM. 2002 Vibrometric studies of the middle ear of the bullfrog *Rana catesbeiana* II. The operculum. *J. Exp. Biol.* 205, 3167–3176.
- Cocroft RB, McDiarmid RW, Jaslow AP, Ruiz-Carranza PM. 1990 Vocalizations of eight species of *Atelopus* (Anura: Bufonidae) with comments on communication in the genus. *Copeia* **1990**, 631–643. (doi:10.2307/1446428)
- Gluesenkamp AG, Acosta N. 2001 Sexual dimorphism in *Osornophryne guacamayo* with notes on natural history and reproduction in the species. *J. Herp.* 35, 148–151. (doi:10.2307/1566040)
- Womack MC, Christensen-Dalsgaard J, Hoke KL.
 2016 Better late than never: effective air-borne hearing of toads delayed by late maturation of the tympanic middle ear structures. J. Exp. Biol. 219, 3246-3252. (doi:10.1242/jeb.143446)
- Berk M. sme: Smoothing-splines mixed-effects models. R package version 0.8. h. See https:// CRAN.R-project.org/package=sme.
- R Core Team. 2015 *R: a language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing. See http:// www.R-project.org/.
- Bates D, Mächler M, Bolker B, Walker S. 2014 Fitting linear mixed-effects models using lme4. *J. Statistic. Softw.* 67, 1–48. (doi:10.18637/jss.v067.i01)
- Lenth RV. 2016 Least-squares means: the R package Ismeans. J. Stat. Softw. 69, 1–33. (doi:10.18637/ jss.v069.i01)
- Boatright-Horowitz SS, Simmons AM. 1995 Postmetamorphic changes in auditory sensitivity of the bullfrog midbrain. *J. Comp. Physiol. A* **177**, 577–590. (doi:10.1007/bf0020718)
- Miranda JA, Wilczynski W. 2009 Sex differences and androgen influences on midbrain auditory thresholds in the green treefrog, *Hyla cinerea*. *Hear. Res.* 252, 79–88. (doi:10.1016/j.heares.2009.04.004)
- Shen JX, Xu ZM, Yu ZL, Wang S, Zheng DZ, Fan SC. 2011 Ultrasonic frogs show extraordinary sex differences in auditory frequency sensitivity. *Nat. Commun.* 2, 342. (doi:10.1038/ncomms1339)

rspb.royalsocietypublishing.org Proc. R. Soc. B 284: 20171670

9

- Hetherington, TE. 1992 The effects of body size on functional properties of middle ear systems of anuran amphibians. *Brain Behav. Evol.* **39**, 133–142. (doi:10.1159/000114111)
- Fox, JH. 1995 Morphological correlates of auditory sensitivity in anuran amphibians. *Brain Behav. Evol.* 45, 327–338. (doi:10.1159/000113560)
- Wever EG, Vernon JA, Peterson EA, Crowley DE. 1963 Auditory responses in the Tokay gecko. *Proc. Natl Acad. Sci. USA* 50, 806-811. (doi:10.1073/ pnas.50.5.806)
- Christensen-Dalsgaard, J. 2005 Directional hearing in nonmammalian tetrapods. In *Sound source localization* (eds RR Fay, AN Popper), pp. 67–123. New York, NY: Springer.
- Koyama H, Lewis ER, Leverenz EL, Baird RA. 1982 Acute seismic sensitivity in the bullfrog ear. *Brain Res.* 250, 168–172. (doi:10.1016/0006-8993(82)90964-7)
- Narins PM, Lewis ER. 1984 The vertebrate ear as an exquisite seismic sensor. *J. Acoust. Soc. Am.* **76**, 1384–1387. (doi:10.1121/1.391455)
- Christensen-Dalsgaard J, Jørgensen MB. 1988 The response characteristics of vibration-sensitive saccular fibers in the grassfrog, *Rana temporaria*. *J. Compar. Physiol. A* **162**, 633–638. (doi:10.1007/ bf01342638)

- Ross RJ, Smith JJB. 1978 Detection of substrate vibrations by salamanders: inner ear sense organ activity. *Can. J. Zool.* 56, 1156–1162. (doi:10.1139/ z78-159)
- Ross RJ, Smith JJB. 1980 Detection of substrate vibrations by salamanders: frequency sensitivity of the ear. *Comp. Biochem. Physiol. Part A Mol. Integr. Physiol.* 65, 167 – 172. (doi:10.1016/0300-9629(80)90218-2)
- Ross RJ, Smith JJB. 1982 Responses of the salamander inner ear to vibrations of the middle ear. *Can. J. Zool.* 60, 220–226. (doi:10.1139/z82-030)
- 57. Wever EG, Vernon JA. 1960 The problem of hearing in snakes. *J. Aud. Res* **1**, 77–83.
- Hartline PH. 1971 Physiological basis for detection of sound and vibration in snakes. J. Exp. Biol. 54, 349-371.
- Lewis ER, Narins PM. 1999 The acoustic periphery of amphibians: anatomy and physiology. In *Comparative hearing: fish and amphibians* (eds RR Fay, AN Popper), pp. 101–154. New York, NY: Springer.
- Heffner RS, Koay G, Heffner HE. 2001 Audiograms of five species of rodents: implications for the evolution of hearing and the perception of pitch. *Hear. Res.* 157, 138–152. (doi:10.1016/s0378-5955(01)00298-2)

- Brittan-Powell EF, Christensen-Dalsgaard J, Tang Y, Carr C, Dooling RJ. 2010 The auditory brainstem response in two lizard species. *J. Acoust. Soc. Am.* 128, 787–794. (doi:10.1121/1.3458813)
- Christensen CB, Christensen-Dalsgaard J, Brandt C, Madsen PT. 2012 Hearing with an atympanic ear: good vibration and poor sound-pressure detection in the royal python, *Python regius. J. Exp. Biol.* 215, 331–342. (doi:10.1242/jeb.062539)
- Capshaw G, Soares D. 2016 Hearing in plethodontid salamanders: a review. *Copeia* **104**, 157–164. (doi:10.1643/ot-15-310)
- Caldwell MS, Johnston GR, McDaniel JG, Warkentin KM. 2010 Vibrational signaling in the agonistic interactions of red-eyed treefrogs. *Curr. Biol.* 20, 1012–1017. (doi:10.1016/j.cub.2010. 03.069)
- Halfwerk W, Jones PL, Taylor RC, Ryan MJ, Page RA. 2014 Risky ripples allow bats and frogs to eavesdrop on a multisensory sexual display. *Science* 343, 413–416. (doi:10.1126/science.1244812)
- Rueda Solano LA, Warkentin KM. 2016 Foraging behavior with possible use of substrate-borne vibrational cues for prey localization in *Atelopus laetissimus* (Ruiz-Carranza, Ardila-Robayo, and Hernández-Camacho, 1994). *Herpetol. Notes* 9, 191–195.